

Package ‘lrrstat’

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Type Package

Title Power and Sample Size Calculation for Non-Proportional Hazards and Beyond

Version 0.3.4

Description Performs power and sample size calculation for non-proportional hazards model using the Fleming-Harrington family of weighted log-rank tests. The sequentially calculated log-rank test score statistics are assumed to have independent increments as characterized in Anastasios A. Tsiatis (1982) <doi:10.1080/01621459.1982.10477898>. The mean and variance of log-rank test score statistics are calculated based on Kaifeng Lu (2021) <doi:10.1002/pst.2069>. The boundary crossing probabilities are calculated using the recursive integration algorithm described in Christopher Jennison and Bruce W. Turnbull (2000, ISBN:0849303168). The package can also be used for continuous, binary, and count data. For continuous data, it can handle missing data through mixed-model for repeated measures (MMRM). In crossover designs, it can estimate direct treatment effects while accounting for carryover effects. For binary data, it can design Simon's 2-stage, modified toxicity probability-2 (mTPI-2), and Bayesian optimal interval (BOIN) trials. For count data, it can design group sequential trials for negative binomial endpoints with censoring. Additionally, it facilitates group sequential equivalence trials for all supported data types. Moreover, it can design adaptive group sequential trials for changes in sample size, error spending function, number and spacing of future looks. Finally, it offers various options for adjusted p-values, including graphical and gatekeeping procedures.

License GPL (>= 2)

Imports Rcpp (>= 1.0.9), RcppParallel (>= 5.1.5), parallel (>= 3.5), rlang (>= 1.0.6), ggplot2 (>= 3.3.6), plotly (>= 4.10.1), shiny (>= 1.7.1), stats (>= 3.5.0)

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<https://github.com/kaifenglu/lrstat>

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lrstat-package	<i>Power and Sample Size Calculation for Non-Proportional Hazards and Beyond</i>
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Description

A comprehensive clinical trial design and analysis package with a focus on non-proportional hazards and weighted log-rank methods, plus broad support for group sequential design, adaptive design, multiplicity, dose-finding, and endpoint-specific power and sample size calculations.

Details

For proportional hazards, power is determined by the total number of events and the constant hazard ratio together with information rates and spending functions. For non-proportional hazards, the hazard ratio varies over time and calendar time determines the mean and variance of the weighted log-rank score statistic. The package uses the analytic approach in Lu (2021) and approximates the variance-covariance matrix of sequential statistics under the alternative by that under the null to leverage the independent increments structure in Tsiatis (1982) for the Fleming-Harrington family.

The package capabilities can be grouped as follows:

- **Time-to-event design under proportional and non-proportional hazards:** weighted log-rank statistics and design operating characteristics (for example `lrstat`, `lrpower`, `lrsamplesize`, `lrsim`, `lrschoenfeld`), including equivalence settings, event prediction, accrual modeling, and calendar time determination (for example `accrual`, `caltime`, `nevent`, `pevent`, `patrisk`, `natrisk`, `getDurationFromNevents`, `getNeventsFromHazardRatio`).
- **Group sequential and adaptive designs:** standard, adaptive, multi-arm multi-stage (MAMS), and seamless frameworks, including boundaries, conditional power, confidence intervals, and exit probabilities (for example `getDesign`, `adaptDesign`, `getBound`, `getCP`, `getCI`, `getRCI`, `getADCI`, `getADRCI`, `exitprob`, and their `_multiarm` and `_seamless` variants).
- **Fixed and sequential sample size/power across endpoint types:** binary, continuous, ordinal/multinomial, paired, crossover/carryover, MMRM, repeated measures ANOVA, two-way ANOVA, Wilcoxon, Fisher's exact, McNemar, one-sample exact, agreement, logistic regression, and negative binomial endpoints (for example `getDesign*`, `power*Exact`, `samplesize*Exact`, `kmpower`, `rmpower`, `nbpower`, and associated `*samplesize` and `*equiv` functions).
- **Multiplicity and graphical procedures:** weighted Bonferroni and graph-based updates, Bonferroni mixtures of weighted Simes and Dunnett, and gatekeeping procedures for multiple hypotheses (for example `fadjpbbon`, `fadjpsim`, `fadjpdun`, `fseqbon`, `fstp2seq`, `fstdmix`, `fmodmix`, `fwgtmat`, `updateGraph`).
- **Early-phase and dose-finding tools:** Simon's two-stage, Bayesian Simon designs, mTPI-2, and BOIN utilities (for example `simon2stage`, `simonBayesAnalysis`, `simonBayesSim`, `mTPI2Table`, `BOINTable`).
- **Estimation, confidence intervals, and supporting utilities:** Clopper-Pearson, exact risk ratio/risk difference, Brookmeyer-Crowley survival quantiles, Miettinen-Nurminen and REML intervals/statistics for stratified measures, Hedges' g, piecewise-exponential distribution tools, multivariate normal integration, and model diagnostics (for example `ClopperPearsonCI`, `riskRatioExactCI`,

riskDiffExactCI, survQuantile, mnRiskDiffCI, remlRiskDiff, hedgesg, dtpwexp, ptpwexp, qtpwexp, rtpwexp, pbvnorm, pmvnormr, qmvnormr, phregr, liferegr, residuals_phregr, survfit_phregr, zph_phregr).

- **Interactive use:** a bundled Shiny interface for exploratory design workflows (runShinyApp_lrstat).

The development of lrstat was strongly influenced by rpact, with argument naming aligned where possible for ease of adoption. Key differences include direct approximation (rather than Schoenfeld-based approximation) for weighted log-rank design calculations, explicit use of accrualDuration to define the accrual end, and treatment of final stage trial outcome when early futility stopping does not occur.

Author(s)

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References

Anastasios A. Tsiatis. Repeated significance testing for a general class of statistics used in censored survival analysis. J Am Stat Assoc. 1982;77:855-861.

Christopher Jennison, Bruce W. Turnbull. Group Sequential Methods with Applications to Clinical Trials. Chapman & Hall/CRC: Boca Raton, 2000, ISBN:0849303168

Kaifeng Lu. Sample size calculation for logrank test and prediction of number of events over time. Pharm Stat. 2021;20:229-244.

See Also

rpact, gsDesign

Examples

```
lrpower(kMax = 2, informationRates = c(0.8, 1),
       criticalValues = c(2.250, 2.025), accrualIntensity = 20,
       piecewiseSurvivalTime = c(0, 6),
       lambda1 = c(0.0533, 0.0309), lambda2 = c(0.0533, 0.0533),
       gamma1 = 0.00427, gamma2 = 0.00427,
       accrualDuration = 22, followupTime = 18)
```

accrual	<i>Number of Enrolled Subjects</i>
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Description

Obtains the number of subjects enrolled by given calendar times.

Usage

```
accrual(time, accrualTime, accrualIntensity, accrualDuration)
```

Arguments

time	A vector of calendar times at which to calculate the number of enrolled subjects.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
accrualDuration	Duration of the enrollment period.

Value

A vector of total number of subjects enrolled by the specified calendar times.

Author(s)

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Examples

```
# Example 1: Uniform enrollment with 20 patients per month for 12 months.
```

```
accrual(time = 3, accrualTime = 0, accrualIntensity = 20,
        accrualDuration = 12)
```

```
# Example 2: Piecewise accrual, 10 patients per month for the first
# 3 months, and 20 patients per month thereafter. Patient recruitment
# ends at 12 months for the study.
```

```
accrual(time = c(2, 9), accrualTime = c(0, 3),
        accrualIntensity = c(10, 20), accrualDuration = 12)
```

adaptDesign

Adaptive Design at an Interim Look

Description

Calculates the conditional power for specified incremental information, given the interim results, parameter value, data-dependent changes in the error spending function, and the number and spacing of interim looks. Conversely, calculates the incremental information required to attain a specified conditional power, given the interim results, parameter value, data-dependent changes in the error spending function, and the number and spacing of interim looks.

Usage

```

adaptDesign(
  betaNew = NA_real_,
  INew = NA_real_,
  L = NA_integer_,
  zL = NA_real_,
  theta = NA_real_,
  IMax = NA_real_,
  kMax = NA_integer_,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,
  futilityCP = NULL,
  futilityTheta = NULL,
  spendingTime = NA_real_,
  MullerSchafer = FALSE,
  kNew = NA_integer_,
  informationRatesNew = NA_real_,
  efficacyStoppingNew = NA_integer_,
  futilityStoppingNew = NA_integer_,
  typeAlphaSpendingNew = "sfOF",
  parameterAlphaSpendingNew = NA_real_,
  futilityBoundsInt = NULL,
  futilityCPInt = NULL,
  futilityThetaInt = NULL,
  typeBetaSpendingNew = "none",
  parameterBetaSpendingNew = NA_real_,
  userBetaSpendingNew = NA_real_,
  spendingTimeNew = NA_real_,
  varianceRatio = 1
)

```

Arguments

betaNew	The type II error for the secondary trial.
INew	The maximum information of the secondary trial. Either betaNew or INew should be provided, while the other must be missing.
L	The interim adaptation look of the primary trial.
zL	The z-test statistic at the interim adaptation look of the primary trial.
theta	The assumed parameter value.
IMax	The maximum information of the primary trial. Must be provided.

kMax	The maximum number of stages of the primary trial.
informationRates	The information rates of the primary trial.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage of the primary trial. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage of the primary trial. Defaults to TRUE if left unspecified.
criticalValues	The upper boundaries on the z-test statistic scale for efficacy stopping for the primary trial. If missing, boundaries will be computed based on the specified alpha spending function.
alpha	The significance level of the primary trial. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending for the primary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsiatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value of alpha spending for the primary trial. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user-defined alpha spending for the primary trial. Represents the cumulative alpha spent up to each stage.
futilityBounds	The lower boundaries on the z-test statistic scale for futility stopping for the primary trial. Defaults to $\text{rep}(-8, \text{kMax}-1)$ if left unspecified.
futilityCP	The conditional power-based futility bounds for the primary trial.
futilityTheta	The parameter value-based futility bounds for the primary trial.
spendingTime	The error spending time of the primary trial. Defaults to missing, in which case it is assumed to be the same as informationRates.
MullerSchafer	Whether to use the Muller and Schafer (2001) method for trial adaptation.
kNew	The number of looks of the secondary trial.
informationRatesNew	The spacing of looks of the secondary trial.
efficacyStoppingNew	The indicators of whether efficacy stopping is allowed at each look of the secondary trial. Defaults to TRUE if left unspecified.
futilityStoppingNew	The indicators of whether futility stopping is allowed at each look of the secondary trial. Defaults to TRUE if left unspecified.

typeAlphaSpendingNew	The type of alpha spending for the secondary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpendingNew	The parameter value of alpha spending for the secondary trial. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
futilityBoundsInt	The futility boundaries on the z statistic scale for new stages of the integrated trial.
futilityCPInt	The conditional power-based futility bounds for new stages of the integrated trial.
futilityThetaInt	The parameter value-based futility bounds for the new stages of the integrated trial.
typeBetaSpendingNew	The type of beta spending for the secondary trial. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpendingNew	The parameter value of beta spending for the secondary trial. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpendingNew	The user-defined cumulative beta spending. Represents the cumulative beta spent up to each stage of the secondary trial.
spendingTimeNew	The error spending time of the secondary trial. Defaults to missing, in which case it is assumed to be the same as informationRatesNew.
varianceRatio	The ratio of the variance under H0 to the variance under H1.

Value

An adaptDesign object with three list components:

- primaryTrial: A list of selected information for the primary trial, including L, zL, theta, maxInformation, kMax, informationRates, efficacyBounds, futilityBounds, information, alpha, conditionalAlpha, conditionalPower, predictivePower, and and MullerSchafer.
- secondaryTrial: A list of selected information for the secondary trial, including overallReject, alpha, kMax, maxInformation, informationRates, efficacyBounds, futilityBounds, cumulativeRejection, cumulativeFutility, cumulativeAlphaSpent, information, typeAlphaSpending, parameterAlphaSpending, typeBetaSpending, parameterBetaSpending, userBetaSpending, and spendingTime.

- `integratedTrial`: A list of selected information for the integrated trial, including `L`, `zL`, `theta`, `maxInformation`, `kMax`, `informationRates`, `efficacyBounds`, `futilityBounds`, and `information`.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Lu Chi, H. M. James Hung, and Sue-Jane Wang. Modification of sample size in group sequential clinical trials. *Biometrics* 1999;55:853-857.

Hans-Helge Muller and Helmut Schafer. Adaptive group sequential designs for clinical trials: Combining the advantages of adaptive and of classical group sequential approaches. *Biometrics* 2001;57:886-891.

See Also

[getDesign](#)

Examples

```
# two-arm randomized clinical trial with a normally distributed endpoint
# 90% power to detect mean difference of 15 with a standard deviation of 50
# Design the Stage I Trial with 3 looks and Lan-DeMets O'Brien-Fleming type
# spending function
delta <- 15
sigma <- 50

(des1 <- getDesignMeanDiff(
  beta = 0.1, meanDiff = delta, stDev = sigma,
  kMax = 3, alpha = 0.025, typeAlphaSpending = "sfOF"
))

s1 <- des1$byStageResults$informationRates
b1 <- des1$byStageResults$efficacyBounds
n <- des1$overallResults$numberOfSubjects

# Monitoring the Stage I Trial
L <- 1
nL <- des1$byStageResults$numberOfSubjects[L]
deltahat <- 8
sigmahat <- 55
sedeltahat <- sigmahat * sqrt( 4 / nL)
zL <- deltahat / sedeltahat

# Making an Adaptive Change: Stage I to Stage II
# revised clinically meaningful difference downward to 10
# retain the standard deviation at the design stage
# Muller & Schafer (2001) method to design the secondary trial
# with 2 looks and Lan-DeMets Pocock type spending function
# re-estimate sample size to reach 90% conditional power
```

```

deltaNew <- 10

(des2 <- adaptDesign(
  betaNew = 0.1, L = L, zL = zL, theta = deltaNew,
  IMax = n / (4 * sigma^2), kMax = 3, informationRates = s1,
  alpha = 0.025, typeAlphaSpending = "sfOF",
  MullerSchafer = TRUE, kNew = 2, typeAlphaSpendingNew = "sfP"
))

INew <- des2$maxInformation
(nNew <- ceiling(INew * 4 * sigma^2))
(nTotal <- nL + nNew)

```

adaptDesign_multiarm *Adaptive Multi-Arm Multi-Stage Design*

Description

Calculates the conditional power for specified incremental information, given the interim results, parameter value, data-dependent changes in treatment selection, the error spending function, and the number and spacing of interim looks. Conversely, calculates the incremental information required to attain a specified conditional power, given the interim results, parameter value, data-dependent changes in treatment selection, the error spending function, and the number and spacing of interim looks.

Usage

```

adaptDesign_multiarm(
  betaNew = NA_real_,
  INew = NA_real_,
  M = NA_integer_,
  r = 1,
  corr_known = TRUE,
  L = NA_integer_,
  zL = NA_real_,
  theta = NA_real_,
  IMax = NA_real_,
  kMax = NA_integer_,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,

```

```

    futilityCP = NULL,
    futilityTheta = NULL,
    spendingTime = NA_real_,
    MullerSchafer = FALSE,
    MNew = NA_integer_,
    selected = NA_integer_,
    rNew = 1,
    kNew = NA_integer_,
    informationRatesNew = NA_real_,
    efficacyStoppingNew = NA_integer_,
    futilityStoppingNew = NA_integer_,
    typeAlphaSpendingNew = "sfOF",
    parameterAlphaSpendingNew = NA_real_,
    futilityBoundsInt = NULL,
    futilityCPInt = NULL,
    futilityThetaInt = NULL,
    typeBetaSpendingNew = "none",
    parameterBetaSpendingNew = NA_real_,
    userBetaSpendingNew = NA_real_,
    spendingTimeNew = NA_real_,
    nthreads = 0
)

```

Arguments

betaNew	The type II error for the secondary trial.
INew	The maximum information for any active arm versus the common control in the secondary trial. Either betaNew or INew should be provided, while the other must be missing.
M	Number of active treatment arms in the primary trial.
r	Randomization ratio of each active arm to the common control in the primary trial.
corr_known	Logical. If TRUE, the correlation between Wald statistics is derived from the randomization ratio r as $r/(r+1)$. If FALSE, a conservative correlation of 0 is assumed.
L	The interim adaptation look of the primary trial.
zL	The z-test statistics at the interim adaptation look of the primary trial.
theta	A vector of length M representing the assumed treatment effects for each active arm versus the common control. The global null is $\theta_i = 0$ for all i , and alternatives are one-sided: $\theta_i > 0$ for at least one $i = 1, \dots, M$.
IMax	Maximum information for any active arm versus the common control for the primary trial. Must be provided.
kMax	The maximum number of stages of the primary trial.
informationRates	The information rates of the primary trial.

efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage of the primary trial. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage of the primary trial.
criticalValues	The matrix of by-level upper boundaries on the max z-test statistic scale for efficacy stopping for the primary trial. The first column is for level M, the second column is for level M - 1, and so on, with the last column for level 1. If left unspecified, the critical values will be computed based on the specified alpha spending function.
alpha	The significance level of the primary trial. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending for the primary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsiatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value of alpha spending for the primary trial. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user-defined alpha spending for the primary trial. Represents the cumulative alpha spent up to each stage.
futilityBounds	The futility boundaries on the max-z statistic scale for the primary trial. Defaults to rep(-8, kMax-1) if left unspecified.
futilityCP	The conditional power-based futility bounds for the primary trial.
futilityTheta	The parameter value-based futility bounds for the primary trial.
spendingTime	The error spending time of the primary trial. Defaults to missing, in which case it is assumed to be the same as informationRates.
MullerSchafer	Whether to use the Muller and Schafer (2001) method for trial adaptation.
MNew	Number of active treatment arms in the secondary trial.
selected	The indices of the selected active treatment arms for the secondary trial.
rNew	Randomization ratio of each active arm to the common control in the secondary trial.
kNew	The number of looks of the secondary trial.
informationRatesNew	The spacing of looks of the secondary trial.
efficacyStoppingNew	The indicators of whether efficacy stopping is allowed at each look of the secondary trial. Defaults to TRUE if left unspecified.
futilityStoppingNew	The indicators of whether futility stopping is allowed at each look of the secondary trial. Defaults to TRUE if left unspecified.

typeAlphaSpendingNew	The type of alpha spending for the secondary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpendingNew	The parameter value of alpha spending for the secondary trial. Corresponds to ρ for "sfKD", and γ for "sfHSD".
futilityBoundsInt	The futility boundaries on the max-z statistic scale for new stages of the integrated trial.
futilityCPInt	The conditional power-based futility bounds for new stages of the integrated trial.
futilityThetaInt	The parameter value-based futility bounds for the new stages of the integrated trial.
typeBetaSpendingNew	The type of beta spending for the secondary trial. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpendingNew	The parameter value of beta spending for the secondary trial. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpendingNew	The user-defined cumulative beta spending. Represents the cumulative beta spent up to each stage of the secondary trial.
spendingTimeNew	The error spending time of the secondary trial. Defaults to missing, in which case it is assumed to be the same as informationRatesNew.
nthreads	The number of threads to use (0 leaves the RcppParallel setting unchanged).

Value

An adaptDesign_multiarm object with three list components:

- **primaryTrial**: A list of selected information for the primary trial, including M, r, corr_known, L, zL, theta, maxInformation, kMax, informationRates, efficacyBounds, futilityBounds, information, alpha, conditionalAlpha, conditionalPower, MullerSchafer, and byLevelBounds, where byLevelBounds is a data frame with columns level, stage, and efficacyBounds, representing the efficacy bounds for each combination of the number of active arms and the stage of analysis in the primary trial.
- **secondaryTrial**: A list of selected information for the secondary trial, including overallReject, alpha, M, r, selected, corr_known, kMax, maxInformation, informationRates, cumulativeRejection, cumulativeAlphaSpent, information, typeAlphaSpending, parameterAlphaSpending,

typeBetaSpending, parameterBetaSpending, userBetaSpending, spendingTime, and byHypothesisBounds, where byHypothesisBounds is a data frame with columns hypothesis, stage, efficacyBounds, and futilityBounds, representing the efficacy and futility bounds for each hypothesis and each stage of analysis in the secondary trial.

- integratedTrial: A list of selected information for the integrated trial, including M, r, corr_known, MNew, rNew, selected, L, zL, theta, maxInformation, kMax, informationRates, efficacyBounds, futilityBounds, information, and byIntersectionBounds, where byIntersectionBounds is a data frame with columns intersectionHypothesis, stage, and efficacyBounds, representing the efficacy bounds for each intersection hypothesis and each stage of analysis in the integrated trial.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

References

Ping Gao, Yingqiu Li. Adaptive multiple comparison sequential design (AMCSD) for clinical trials. Journal of Biopharmaceutical Statistics, 2024, 34(3), 424-440.

See Also

[getDesign_multiarm](#)

Examples

```
# Two active treatment arms are compared with a common control in a
# two-look time-to-event design using O'Brien-Fleming-type alpha spending.
# Suppose each active arm has a true hazard ratio of 0.75 versus control,
# and the total number of events across all three arms at the final analysis
# is 486. This corresponds to approximately 324 events for each active arm
# versus the common control. Under these assumptions, the trial has about
# 80% power to detect the treatment effect in at least one active arm.
```

```
(des1 <- getDesign_multiarm(
  IMax = 324 / 4, theta = c(-log(0.75), -log(0.75)),
  M = 2, r = 1, kMax = 2, informationRates = c(1/2, 1),
  alpha = 0.025, typeAlphaSpending = "OF", nthreads = 1))
```

```
# Now assume that, at the interim analysis, the observed hazard ratios for
# the two active arms versus control are 0.91 and 0.78, respectively. Using
# the rule "drop any arm with an observed hazard ratio > 0.9", arm 1 is
# dropped. We then aim to achieve 80% conditional power to detect a hazard
# ratio of 0.78 for the remaining arm at the final look. The analysis below
# indicates that the required total number of events for arm 2 versus control
# at the final analysis should be increased from 324 to 535.
```

```
(des2 <- adaptDesign_multiarm(
  betaNew = 0.2, M = 2, r = 1, corr_known = FALSE,
  L = 1, zL = c(-log(0.91), -log(0.78)) * sqrt(324 / 4 / 2),
  theta = c(-log(0.91), -log(0.78)),
```

```
IMax = 324 / 4, kMax = 2, informationRates = c(1/2, 1),
alpha = 0.025, typeAlphaSpending = "OF",
MNew = 1, selected = 2, rNew = 1, nthreads = 1))
```

adaptDesign_seamless *Adaptive Phase 2/3 Seamless Design*

Description

Calculates the conditional power for specified incremental information, given the interim results, parameter value, data-dependent changes in the error spending function, and the number and spacing of interim looks. Conversely, calculates the incremental information required to attain a specified conditional power, given the interim results, parameter value, data-dependent changes in the error spending function, and the number and spacing of interim looks.

Usage

```
adaptDesign_seamless(
  betaNew = NA_real_,
  INew = NA_real_,
  M = NA_integer_,
  r = 1,
  corr_known = TRUE,
  L = NA_integer_,
  zL = NA_real_,
  theta = NA_real_,
  IMax = NA_real_,
  K = NA_integer_,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,
  futilityCP = NULL,
  futilityTheta = NULL,
  spendingTime = NA_real_,
  MullerSchafer = FALSE,
  kNew = NA_integer_,
  informationRatesNew = NA_real_,
  efficacyStoppingNew = NA_integer_,
  futilityStoppingNew = NA_integer_,
  typeAlphaSpendingNew = "sfOF",
```

```

parameterAlphaSpendingNew = NA_real_,
futilityBoundsInt = NULL,
futilityCPInt = NULL,
futilityThetaInt = NULL,
typeBetaSpendingNew = "none",
parameterBetaSpendingNew = NA_real_,
userBetaSpendingNew = NA_real_,
spendingTimeNew = NA_real_,
nthreads = 0
)

```

Arguments

betaNew	The type II error for the secondary trial.
INew	The maximum information for the active arm versus the common control in the secondary trial. Either betaNew or INew should be provided, while the other must be missing.
M	Number of active treatment arms in Phase 2.
r	Randomization ratio of each active arm to the common control in Phase 2.
corr_known	Logical. If TRUE, the correlation between Wald statistics in Phase 2 is derived from the randomization ratio r as $r/(r+1)$. If FALSE, a conservative correlation of 0 is used. This option is only used for critical value calculations.
L	The interim adaptation look in Phase 3.
zL	The z-test statistic at the interim adaptation look of Phase 3.
theta	The assumed treatment effect for the selected arm versus the common control.
IMax	Maximum information for the active arm versus the common control for the original trial. Must be provided.
K	Number of sequential looks in Phase 3.
informationRates	A numeric vector of information rates fixed before the trial. If unspecified, defaults to $(1 : (K + 1))/(K + 1)$.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage of the primary trial. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage of the primary trial. Defaults to TRUE if left unspecified.
criticalValues	The upper boundaries on the z-test statistic scale for the most promising arm in Phase 2 and the z-test statistics for the selected arm in Phase 3 for the primary trial. If missing, boundaries will be computed based on the specified alpha spending function.
alpha	The significance level of the primary trial. Defaults to 0.025.

typeAlphaSpending	The type of alpha spending for the primary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value of alpha spending for the primary trial. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user-defined alpha spending for the primary trial. Represents the cumulative alpha spent up to each stage.
futilityBounds	The lower boundaries on the z-test statistic scale at the end of phase 2 for the most promising arm and on the z-test statistic scale in phase 3 for futility stopping for the primary trial. Defaults to $\text{rep}(-8, k_{\text{Max}}-1)$ if left unspecified.
futilityCP	The conditional power-based futility bounds for the primary trial.
futilityTheta	The parameter value-based futility bounds for the primary trial.
spendingTime	The error spending time of the primary trial. Defaults to missing, in which case it is assumed to be the same as informationRates.
MullerSchafer	Whether to use the Muller and Schafer (2001) method for trial adaptation.
kNew	The number of looks of the secondary trial.
informationRatesNew	The spacing of looks of the secondary trial.
efficacyStoppingNew	The indicators of whether efficacy stopping is allowed at each look of the secondary trial. Defaults to TRUE if left unspecified.
futilityStoppingNew	The indicators of whether futility stopping is allowed at each look of the secondary trial. Defaults to TRUE if left unspecified.
typeAlphaSpendingNew	The type of alpha spending for the secondary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpendingNew	The parameter value of alpha spending for the secondary trial. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
futilityBoundsInt	The futility boundaries on the z statistic scale for new stages of the integrated trial.
futilityCPInt	The conditional power-based futility bounds for new stages of the integrated trial.

futilityThetaInt	The parameter value-based futility bounds for the new stages of the integrated trial.
typeBetaSpendingNew	The type of beta spending for the secondary trial. One of the following: "sf0F" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpendingNew	The parameter value of beta spending for the secondary trial. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpendingNew	The user-defined cumulative beta spending. Represents the cumulative beta spent up to each stage of the secondary trial.
spendingTimeNew	The error spending time of the secondary trial. Defaults to missing, in which case it is assumed to be the same as informationRatesNew.
nthreads	The number of threads to use (0 leaves the RcppParallel setting unchanged).

Value

An adaptDesign_seamless object with three list components:

- primaryTrial: A list of selected information for the primary trial, including M, r, corr_known, K, L, zL, theta, maxInformation, kMax, informationRates, efficacyBounds, futilityBounds, information, alpha, conditionalAlpha, conditionalPower, and MullerSchafer.
- secondaryTrial: A list of selected information for the secondary trial, including overallReject, alpha, kMax, maxInformation, informationRates, efficacyBounds, futilityBounds, cumulativeRejection, cumulativeFutility, cumulativeAlphaSpent, information, typeAlphaSpending, parameterAlphaSpending, typeBetaSpending, parameterBetaSpending, userBetaSpending, and spendingTime.
- integratedTrial: A list of selected information for the integrated trial, including M, r, corr_known, K, L, zL, theta, maxInformation, kMax, informationRates, efficacyBounds, futilityBounds, and information.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Ping Gao, Yingqiu Li. Adaptive two-stage seamless sequential design for clinical trials. Journal of Biopharmaceutical Statistics, 2025, 35(4), 565-587.

See Also

[getDesign_seamless](#)

Examples

```
(des1 <- adaptDesign_seamless(
  betaNew = 0.1, M = 2, r = 1, corr_known = FALSE,
  L = 1, zL = -log(0.67) * sqrt(80 / 4), theta = -log(0.691),
  IMax = 120 / 4, K = 2, informationRates = c(1/3, 2/3, 1),
  alpha = 0.025, typeAlphaSpending = "OF", kNew = 1, nthreads = 1))
```

aml

*Acute myelogenous leukemia survival data from the survival package***Description**

Survival in patients with acute myelogenous leukemia.

time Survival or censoring time

status censoring status

x maintenance chemotherapy given or not

Usage

```
aml
```

Format

An object of class `data.frame` with 23 rows and 3 columns.

assess_phregr

*Assess Proportional Hazards Assumption Based on Supremum Test***Description**

Obtains the standardized score processes and the simulated distribution under the null hypothesis as well as the p-values for the supremum tests.

Usage

```
assess_phregr(object, resample = 1000, seed = 12345)
```

Arguments

object The output from the phregr call.

resample The number of simulation samples for the supremem test.

seed The random seed for the simulations.

Details

The supremum test corresponds to the ASSESS statement with ph option of SAS PROC PHREG.

Value

A list with the following components:

- time the unique event times.
- score_t the observed standardized score process.
- score_t_list a list of simulated standardized score processes under the null hypothesis.
- max_abs_value the supremum of the absolute value of the observed standardized score process for each covariate and the supremum of the sum of absolute values of the observed standardized score processes across all covariates.
- p_value the p-values for the supremum tests for each covariate and the global test.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

References

D. Y. Lin, L. J. Wei, and Z. Ying. Checking the Cox model with cumulative sums of martingale-based residuals. *Biometrika* 1993; 80:557-572.

Examples

```
fit <- phregr(data = liver, time = "Time", event = "Status",
             covariates = c("log(Bilirubin)", "log(Protime)",
                           "log(Albumin)", "Age", "Edema"),
             ties = "breslow")

aph <- assess_phregr(fit, resample = 1000, seed = 314159)

aph

plot(aph, nsim = 20)
```

Description

Performs simulation for two-endpoint two-arm group sequential trials.

- Endpoint 1: Binary endpoint, analyzed using the Mantel-Haenszel test for risk difference.
- Endpoint 2: Time-to-event endpoint, analyzed using the log-rank test for treatment effect.

The analysis times for the binary endpoint are based on calendar times, while the time-to-event analyses are triggered by reaching the pre-specified number of events. The binary endpoint is assessed at the first post-treatment follow-up visit (PTFU1).

Usage

```
binary_tte_sim(
  kMax1 = 1,
  kMax2 = 1,
  riskDiffH0 = 0,
  hazardRatioH0 = 1,
  allocation1 = 1,
  allocation2 = 1,
  accrualTime = 0,
  accrualIntensity = NA,
  piecewiseSurvivalTime = 0,
  stratumFraction = 1,
  globalOddsRatio = NA,
  pi1 = NA,
  pi2 = NA,
  lambda1 = NA,
  lambda2 = NA,
  gamma1 = NA,
  gamma2 = NA,
  delta1 = NA,
  delta2 = NA,
  upper1 = NA,
  upper2 = NA,
  n = NA,
  plannedTime = NA,
  plannedEvents = NA,
  maxNumberOfIterations = 1000,
  maxNumberOfRawDatasetsPerStage = 0,
  seed = 0,
  nthreads = 0
)
```

Arguments

kMax1	Number of stages for the binary endpoint.
kMax2	Number of stages for the time-to-event endpoint.
riskDiffH0	Risk difference under the null hypothesis for the binary endpoint.

hazardRatioH0	Hazard ratio under the null hypothesis for the time-to-event endpoint.
allocation1	Number of subjects in the treatment group in a randomization block. Defaults to 1 for equal randomization.
allocation2	Number of subjects in the control group in a randomization block. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
globalOddsRatio	Global odds ratio of the Plackett copula linking the two endpoints.
pi1	Response probabilities by stratum for the treatment group for the binary endpoint.
pi2	Response probabilities by stratum for the control group for the binary endpoint.
lambda1	A vector of hazard rates for the event in each analysis time interval by stratum for the treatment group for the time-to-event endpoint.
lambda2	A vector of hazard rates for the event in each analysis time interval by stratum for the control group for the time-to-event endpoint.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
delta1	The hazard rate for exponential treatment discontinuation, a vector of hazard rates for piecewise exponential treatment discontinuation applicable for all strata, or a vector of hazard rates for treatment discontinuation in each analysis time interval by stratum for the treatment group for the binary endpoint.
delta2	The hazard rate for exponential treatment discontinuation, a vector of hazard rates for piecewise exponential treatment discontinuation applicable for all strata, or a vector of hazard rates for treatment discontinuation in each analysis time interval by stratum for the control group for the binary endpoint.
upper1	Maximum protocol-specified treatment duration for the treatment group.
upper2	Maximum protocol-specified treatment duration for the control group.
n	Sample size.
plannedTime	Calendar times for the analyses of the binary endpoint.

plannedEvents	Target cumulative number of events for the time-to-event analyses.
maxNumberOfIterations	Number of simulation iterations to perform.
maxNumberOfRawDatasetsPerStage	Number of subject-level datasets to retain per stage. Set to 0 to skip raw data saving.
seed	The seed to reproduce the simulation results.
nthreads	The number of threads to use in simulations (0 means the default RcppParallel behavior).

Details

We consider dual primary endpoints with endpoint 1 being a binary endpoint and endpoint 2 being a time-to-event endpoint. The analyses of endpoint 1 will be based on calendar times, while the analyses of endpoint 2 will be based on the number of events. Therefore, the analyses of the two endpoints are not at the same time points. The correlation between the two endpoints is characterized by the global odds ratio of the Plackett copula. In addition, the time-to-event endpoint will render the binary endpoint as a non-responder, and so does the dropout. In addition, the treatment discontinuation will impact the number of available subjects for analysis. The administrative censoring will exclude subjects from the analysis of the binary endpoint.

Value

A list with 4 components:

- sumdataBIN: A data frame of summary data by iteration and stage for the binary endpoint:
 - iterationNumber: The iteration number.
 - stageNumber: The stage number, covering all stages even if the trial stops at an interim look.
 - analysisTime: The time for the stage since trial start.
 - accruals1: The number of subjects enrolled at the stage for the treatment group.
 - accruals2: The number of subjects enrolled at the stage for the control group.
 - totalAccruals: The total number of subjects enrolled at the stage.
 - source1: The total number of subjects with response status determined by the underlying latent response variable.
 - source2: The total number of subjects with response status (non-responder) determined by experiencing the event for the time-to-event endpoint.
 - source3: The total number of subjects with response status (non-responder) determined by dropping out prior to the PTFU1 visit.
 - n1: The number of subjects included in the analysis of the binary endpoint for the treatment group.
 - n2: The number of subjects included in the analysis of the binary endpoint for the control group.
 - n: The total number of subjects included in the analysis of the binary endpoint at the stage.
 - y1: The number of responders for the binary endpoint in the treatment group.

- y2: The number of responders for the binary endpoint in the control group.
- y: The total number of responders for the binary endpoint at the stage.
- riskDiff: The estimated risk difference for the binary endpoint.
- seRiskDiff: The standard error for risk difference based on the Sato approximation.
- mhStatistic: The Mantel-Haenszel test Z-statistic for the binary endpoint.
- sumdataTTE: A data frame of summary data by iteration and stage for the time-to-event endpoint:
 - iterationNumber: The iteration number.
 - eventsNotAchieved: Whether the target number of events is not achieved for the iteration.
 - stageNumber: The stage number, covering all stages even if the trial stops at an interim look.
 - analysisTime: The time for the stage since trial start.
 - accruals1: The number of subjects enrolled at the stage for the treatment group.
 - accruals2: The number of subjects enrolled at the stage for the control group.
 - totalAccruals: The total number of subjects enrolled at the stage.
 - events1: The number of events at the stage for the treatment group.
 - events2: The number of events at the stage for the control group.
 - totalEvents: The total number of events at the stage.
 - dropouts1: The number of dropouts at the stage for the treatment group.
 - dropouts2: The number of dropouts at the stage for the control group.
 - totalDropouts: The total number of dropouts at the stage.
 - uscore: The numerator of the log-rank test statistic for the time-to-event endpoint.
 - vscore: The variance of the log-rank test statistic for the time-to-event endpoint.
 - logRankStatistic: The log-rank test Z-statistic for the time-to-event endpoint.
- rawdataBIN (exists if maxNumberOfRawDatasetsPerStage is a positive integer): A data frame for subject-level data for the binary endpoint for selected replications, containing the following variables:
 - iterationNumber: The iteration number.
 - stageNumber: The stage under consideration.
 - analysisTime: The time for the stage since trial start.
 - subjectId: The subject ID.
 - arrivalTime: The enrollment time for the subject.
 - stratum: The stratum for the subject.
 - treatmentGroup: The treatment group (1 or 2) for the subject.
 - survivalTime: The underlying survival time for the time-to-event endpoint for the subject.
 - dropoutTime: The underlying dropout time for the time-to-event endpoint for the subject.
 - trtDiscTime: The underlying treatment discontinuation time for the binary endpoint for the subject.
 - trtDurUpperLimit: The maximum protocol-specified treatment duration for the subject based on the treatment group assignment.
 - ptfu1Time: The underlying assessment time for the binary endpoint for the subject.

- timeUnderObservation: The time under observation since randomization for the binary endpoint for the subject.
- latentResponse: The underlying latent response variable for the binary endpoint for the subject, which determines the response status for the binary endpoint at PTFU1 visit.
- responder: Whether the subject is a responder for the binary endpoint.
- source: The source of the determination of responder status for the binary endpoint: = 1 based on the underlying latent response variable, = 2 based on the occurrence of the time-to-event endpoint before the assessment time of the binary endpoint (imputed as a non-responder), = 3 based on the dropout before the assessment time of the binary endpoint (imputed as a non-responder), = 4 excluded from analysis due to administrative censoring.
- rawDataTTE (exists if maxNumberOfRawDatasetsPerStage is a positive integer): A data frame for subject-level data for the time-to-event endpoint for selected replications, containing the following variables:
 - iterationNumber: The iteration number.
 - stageNumber: The stage under consideration.
 - analysisTime: The time for the stage since trial start.
 - subjectId: The subject ID.
 - arrivalTime: The enrollment time for the subject.
 - stratum: The stratum for the subject.
 - treatmentGroup: The treatment group (1 or 2) for the subject.
 - survivalTime: The underlying survival time for the time-to-event endpoint for the subject.
 - dropoutTime: The underlying dropout time for the time-to-event endpoint for the subject.
 - timeUnderObservation: The time under observation since randomization for the time-to-event endpoint for the subject.
 - event: Whether the subject experienced the event for the time-to-event endpoint.
 - dropoutEvent: Whether the subject dropped out for the time-to-event endpoint.

Author(s)

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Examples

```
tcut <- c(0, 12, 36, 48)
surv <- c(1, 0.95, 0.82, 0.74)
lambda2 <- (log(surv[1:3]) - log(surv[2:4]))/(tcut[2:4] - tcut[1:3])

sim1 <- binary_tte_sim(
  kMax1 = 1,
  kMax2 = 2,
  accrualTime = seq(0, 8),
  accrualIntensity = 40/9 * seq(1, 9),
  piecewiseSurvivalTime = c(0, 12, 36),
  globalOddsRatio = 1,
  pi1 = 0.80,
```

```

pi2 = 0.65,
lambda1 = 0.65*lambda2,
lambda2 = lambda2,
gamma1 = -log(1-0.04)/12,
gamma2 = -log(1-0.04)/12,
delta1 = -log(1-0.02)/12,
delta2 = -log(1-0.02)/12,
upper1 = 15*28/30.4,
upper2 = 12*28/30.4,
n = 640,
plannedTime = 20 + 15*28/30.4,
plannedEvents = c(130, 173),
maxNumberOfIterations = 1000,
maxNumberOfRawDatasetsPerStage = 1,
seed = 314159,
nthreads = 1)

```

BOINTable

*BOIN Decision Table for Dose-Finding Trials***Description**

Generates the decision table for the Bayesian Optimal Interval (BOIN) design, a widely used approach for dose-escalation trials that guides dose-finding decisions based on observed toxicity rates.

Usage

```

BOINTable(
  nMax = NA_integer_,
  pT = 0.3,
  phi1 = 0.6 * pT,
  phi2 = 1.4 * pT,
  a = 1,
  b = 1,
  pExcessTox = 0.95
)

```

Arguments

nMax	The maximum number of subjects allowed in a dose cohort.
pT	The target toxicity probability. Defaults to 0.3.
phi1	The lower equivalence limit for the target toxicity probability.
phi2	The upper equivalence limit for the target toxicity probability.
a	The prior toxicity shape parameter for the Beta prior.
b	The prior non-toxicity shape parameter for the Beta prior.

pExcessTox The threshold for excessive toxicity. If the posterior probability that the true toxicity rate exceeds p_T is greater than **pExcessTox**, the current and all higher doses will be excluded from further use to protect future participants. Defaults to 0.95.

Value

An S3 class **BOINTable** object with the following components:

- **settings**: The input settings data frame with the following variables:
 - **nMax**: The maximum number of subjects in a dose cohort.
 - **pT**: The target toxicity probability.
 - **phi1**: The lower equivalence limit for target toxicity probability.
 - **phi2**: The upper equivalence limit for target toxicity probability.
 - **lambda1**: The lower decision boundary for observed toxicity probability.
 - **lambda2**: The upper decision boundary for observed toxicity probability.
 - **a**: The prior toxicity parameter for the beta prior.
 - **b**: The prior non-toxicity parameter for the beta prior.
 - **pExcessTox**: The threshold for excessive toxicity.
- **decisionDataFrame**: A data frame listing dose-finding decisions for each combination of sample size (n) and number of observed toxicities (y):
 - **n**: Cohort size.
 - **y**: Number of observed toxicities.
 - **decision**: Recommended action: escalate, de-escalate, or stay at the current dose.
- **decisionMatrix**: A matrix version of the decision table showing the recommended action based on the number of toxicities for each possible cohort size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Liu, S., & Yuan, Y. (2015). Bayesian optimal interval designs for phase I clinical trials. *Journal of the Royal Statistical Society: Series C (Applied Statistics)*, 64(3), 507-523.

Examples

```
BOINTable(nMax = 18, pT = 0.3, phi = 0.6*0.3, phi2 = 1.4*0.3)
```

caltime

*Calendar Times for Target Number of Events***Description**

Obtains the calendar times needed to reach the target number of subjects experiencing an event.

Usage

```
caltime(
  nevents = NA_real_,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = FALSE
)
```

Arguments

nevents	A vector of target number of events.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
lambda1	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.
lambda2	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.

gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.

Value

A vector of calendar times expected to yield the target number of events.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
# Piecewise accrual, piecewise exponential survivals, and 5% dropout by
# the end of 1 year.

caltime(nevents = c(24, 80), allocationRatioPlanned = 1,
        accrualTime = seq(0, 8),
        accrualIntensity = 26/9*seq(1, 9),
        piecewiseSurvivalTime = c(0, 6),
        lambda1 = c(0.0533, 0.0309),
        lambda2 = c(0.0533, 0.0533),
        gamma1 = -log(1-0.05)/12,
        gamma2 = -log(1-0.05)/12,
        accrualDuration = 22,
        followupTime = 18, fixedFollowup = FALSE)
```

ClopperPearsonCI	<i>Clopper-Pearson Confidence Interval for One-Sample Proportion</i>
------------------	--

Description

Obtains the Clopper-Pearson exact confidence interval for a one-sample proportion.

Usage

```
ClopperPearsonCI(n, y, cilevel = 0.95)
```

Arguments

n	The sample size.
y	The number of responses.
cilevel	The confidence interval level.

Value

A data frame with the following variables:

- n: The sample size.
- y: The number of responses.
- phat: The observed proportion of responses.
- lower: The lower limit of the confidence interval.
- upper: The upper limit of the confidence interval.
- cilevel: The confidence interval level.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Clopper, C. J., & Pearson, E. S. (1934). The use of confidence or fiducial limits illustrated in the case of the binomial. *Biometrika*, 26(4), 404-413.

Examples

```
ClopperPearsonCI(20, 3)
```

corr_pfs_os	<i>Correlation Between PFS and OS Given Correlation Between PD and OS</i>
-------------	---

Description

Computes the correlation between PFS and OS given the correlation between PD and OS.

Usage

```
corr_pfs_os(
  piecewiseSurvivalTime = 0L,
  hazard_pfs = NA_real_,
  hazard_os = NA_real_,
  rho_pd_os = NA_real_
)
```

Arguments

<code>piecewiseSurvivalTime</code>	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., <code>c(0, 6)</code> breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
<code>hazard_pfs</code>	A scalar or numeric vector specifying the hazard(s) for PFS based on a piecewise exponential distribution.
<code>hazard_os</code>	A scalar or numeric vector specifying the hazard(s) for overall survival (OS) based on a piecewise exponential distribution.
<code>rho_pd_os</code>	A numeric value specifying the correlation between PD and OS times.

Details

This function first determines the piecewise exponential distribution for PD such that the implied survival function for PFS time, $T_{\text{pfs}} = \min(T_{\text{pd}}, T_{\text{os}})$, closely matches the specified piecewise exponential distribution for PFS with hazard vector λ_{pfs} . Then, it calculates the correlation between PFS and OS times based on the derived piecewise exponential distribution for PD and the given piecewise exponential distribution for OS.

Value

The estimated correlation between PFS and OS.

Author(s)

Kaifeng Lu (<kaifenglu@gmail.com>)

Examples

```
u <- c(0, 1, 3, 4)
lambda1 <- c(0.0151, 0.0403, 0.0501, 0.0558)
lambda2 <- 0.0145
rho_pd_os <- 0.5
corr_pfs_os(u, lambda1, lambda2, rho_pd_os)
```

Description

Obtains the covariance between restricted mean survival times at two different time points.

Usage

```

covrmst(
  t2 = NA_real_,
  tau1 = NA_real_,
  tau2 = NA_real_,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  accrualDuration = NA_real_,
  maxFollowupTime = NA_real_
)

```

Arguments

t2	The calendar time for analysis 2.
tau1	The milestone time for analysis 1.
tau2	The milestone time for analysis 2.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
lambda1	A vector of hazard rates for the event for the active treatment group. One for each analysis time interval.
lambda2	A vector of hazard rates for the event for the control group. One for each analysis time interval.
gamma1	The hazard rate for exponential dropout, or a vector of hazard rates for piecewise exponential dropout for the active treatment group.
gamma2	The hazard rate for exponential dropout, or a vector of hazard rates for piecewise exponential dropout for the control group.
accrualDuration	Duration of the enrollment period.

maxFollowupTime

Follow-up time for the first enrolled subject. For fixed follow-up, maxFollowupTime = minFollowupTime. For variable follow-up, maxFollowupTime = accrualDuration + minFollowupTime.

Value

The covariance between the restricted mean survival times for each treatment group.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
covrmst(t2 = 25, tau1 = 16, tau2 = 18, allocationRatioPlanned = 1,
  accrualTime = c(0, 3), accrualIntensity = c(10, 20),
  piecewiseSurvivalTime = c(0, 6),
  lambda1 = c(0.0533, 0.0309), lambda2 = c(0.0533, 0.0533),
  gamma1 = -log(1-0.05)/12, gamma2 = -log(1-0.05)/12,
  accrualDuration = 12, maxFollowupTime = 30)
```

dtpwexp

Density Function of Truncated Piecewise Exponential Distribution

Description

Obtains the density of a truncated piecewise exponential distribution.

Usage

```
dtpwexp(
  q,
  piecewiseSurvivalTime = 0,
  lambda = 0.0578,
  lowerBound = 0,
  log.d = FALSE
)
```

Arguments

q The vector of quantiles.

piecewiseSurvivalTime A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., `c(0, 6)` breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.

lambda A vector of hazard rates for the event. One for each analysis time interval.

lowerBound	The left truncation time point for the survival time. Defaults to 0 for no truncation.
log.d	Logical; if TRUE, densities d are given as log(d).

Value

The density d such that $d = \lambda(q) * P(X > q | X > \text{lowerBound})$.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
dtpwexp(q = c(8, 18), piecewiseSurvivalTime = c(0, 6, 9, 15),
        lambda = c(0.025, 0.04, 0.015, 0.007))
```

errorSpent	<i>Error Spending</i>
------------	-----------------------

Description

Obtains the error spent at given spending times for the specified error spending function.

Usage

```
errorSpent(t, error = 0.025, sf = "sf0F", sfpar = NA_real_)
```

Arguments

t	A vector of spending times, typically equal to information fractions.
error	The total error to spend.
sf	The spending function. One of the following: "sf0F" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, and "sfHSD" for Hwang, Shi & DeCani spending function. Defaults to "sf0F".
sfpar	The parameter for the spending function. Corresponds to ρ for "sfKD" and γ for "sfHSD".

Details

This function implements a variety of error spending functions commonly used in group sequential designs, assuming one-sided hypothesis testing.

O'Brien-Fleming-Type Spending Function

This spending function allocates very little alpha early on and more alpha later in the trial. It is defined as:

$$\alpha(t) = 2 - 2\Phi\left(\frac{z_{\alpha/2}}{\sqrt{t}}\right),$$

where Φ is the standard normal cumulative distribution function, $z_{\alpha/2}$ is the critical value from the standard normal distribution, and $t \in [0, 1]$ denotes the information fraction.

Pocock-Type Spending Function

This function spends alpha more evenly throughout the study:

$$\alpha(t) = \alpha \log(1 + (e - 1)t),$$

where e is Euler's number (approximately 2.718).

Kim and DeMets Power-Type Spending Function

This family of spending functions is defined as:

$$\alpha(t) = \alpha t^\rho, \quad \rho > 0.$$

- When $\rho = 1$, the function mimics Pocock-type boundaries.
- When $\rho = 3$, it approximates O'Brien-Fleming-type boundaries.

Hwang, Shih, and DeCani Spending Function

This flexible family of functions is given by:

$$\alpha(t) = \begin{cases} \alpha \frac{1 - e^{-\gamma t}}{1 - e^{-\gamma}}, & \text{if } \gamma \neq 0 \\ \alpha t, & \text{if } \gamma = 0. \end{cases}$$

- When $\gamma = -4$, the spending function resembles O'Brien-Fleming boundaries.
- When $\gamma = 1$, it resembles Pocock boundaries.

Value

A vector of errors spent up to the interim look.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
errorSpent(t = 0.5, error = 0.025, sf = "sfOF")
```

```
errorSpent(t = c(0.5, 0.75, 1), error = 0.025, sf = "sfHSD", sfpar = -4)
```

exitprob	<i>Stagewise Exit Probabilities</i>
----------	-------------------------------------

Description

Obtains the stagewise exit probabilities for both efficacy and futility stopping.

Usage

```
exitprob(b, a = NA_real_, theta = 0L, I = NA_real_)
```

Arguments

b	Upper boundaries on the z-test statistic scale.
a	Lower boundaries on the z-test statistic scale. Defaults to <code>c(rep(-8.0, kMax-1), b[kMax])</code> if left unspecified, where <code>kMax = length(b)</code> .
theta	Stagewise parameter of interest, e.g., $-U/V$ for weighted log-rank test, where U is the mean and V is the variance of the weighted log-rank test score statistic at each stage. For proportional hazards and conventional log-rank test, use the scalar input, $\theta = -\log(HR)$. Defaults to 0 corresponding to the null hypothesis.
I	Stagewise cumulative information, e.g., V , the variance of the weighted log-rank test score statistic at each stage. For conventional log-rank test, information can be approximated by $\phi \cdot (1-\phi) \cdot D$, where ϕ is the probability of being allocated to the active arm, and D is the total number of events at each stage. Defaults to <code>seq(1, kMax)</code> if left unspecified.

Value

A list of stagewise exit probabilities:

- `exitProbUpper`: The vector of efficacy stopping probabilities
- `exitProbLower`: The vector of futility stopping probabilities.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
exitprob(b = c(3.471, 2.454, 2.004), theta = -log(0.6),
         I = c(50, 100, 150)/4)

exitprob(b = c(2.963, 2.359, 2.014),
         a = c(-0.264, 0.599, 2.014),
         theta = c(0.141, 0.204, 0.289),
         I = c(81, 121, 160))
```

Description

Computes the exit (rejection) probabilities for a multi-arm multi-stage design.

Usage

```
exitprob_multiarm(
  M = NA_integer_,
  r = 1,
  theta = NA_real_,
  corr_known = TRUE,
  kMax = NA_integer_,
  b = NULL,
  a = NULL,
  I = NULL,
  nthreads = 0
)
```

Arguments

M	Number of active treatment arms.
r	Randomization ratio of each active arm to the common control.
theta	A vector of length M representing the true treatment effects for each active arm versus the common control.
corr_known	Logical. If TRUE, the correlation between Wald statistics is derived from the randomization ratio r as $r/(r + 1)$. If FALSE, a conservative correlation of 0 is used.
kMax	Number of sequential looks.
b	A vector of efficacy boundaries for the max-Z statistics.
a	A vector of futility boundaries for the max-Z statistics.
I	A vector of information levels for any active arm versus the common control.
nthreads	The number of threads to use (0 leaves the RcppParallel setting unchanged).

Details

The function assumes a multivariate normal distribution for the Wald statistics and all active arms share the same information level.

Value

A vector `exitProb` of length `kMax` containing the probability of rejection at each look.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Ping Gao, Yingqiu Li. Adaptive multiple comparison sequential design (AMCSD) for clinical trials. Journal of Biopharmaceutical Statistics, 2024, 34(3), 424-440.

Examples

```
# Setup: 2 active arms vs control and 3 sequential looks.

# Information levels: equal spacing over 3 looks based on a maximum of
# 95 patients per arm, SD = 1.0
I <- 95 / (2 * 1.0^2) * seq(1, 3)/3

# O'Brien-Fleming critical values
b <- c(3.886562, 2.748214, 2.243907)

# Type I error under the global null hypothesis
p0 <- exitprob_multiarm(M = 2, theta = c(0, 0), kMax = 3,
                        b = b, I = I, nthreads = 1)
cumsum(p0$exitProbUpper)

# Power under alternative: Treatment effects of 0.3 and 0.5
p1 <- exitprob_multiarm(M = 2, theta = c(0.3, 0.5), kMax = 3,
                        b = b, I = I, nthreads = 1)
cumsum(p1$exitProbUpper)
```

exitprob_seamless	<i>Exit Probabilities for Phase 2/3 Seamless Design</i>
-------------------	---

Description

Computes the upper and lower exit probabilities for a phase 2/3 seamless design. In Phase 2, multiple active arms are compared against a common control arm. If the largest test statistic at the end of Phase 2 crosses the efficacy boundary, the trial stops early for efficacy; if it falls below the futility boundary, the trial stops early for futility. Otherwise, the arm is selected to proceed to Phase 3, where it is tested against the control over multiple looks with upper and optional lower stopping boundaries.

Usage

```
exitprob_seamless(
  M = NA_integer_,
  r = 1,
  theta = NA_real_,
```

```

    corr_known = TRUE,
    K = NA_integer_,
    b = NULL,
    a = NULL,
    I = NULL,
    nthreads = 0
)

```

Arguments

M	Number of active treatment arms in Phase 2.
r	Randomization ratio of each active arm to the common control in Phase 2.
theta	A vector of length M representing the true treatment effects for each active arm versus the common control.
corr_known	Logical. If TRUE, the correlation between Wald statistics in Phase 2 is derived from the randomization ratio r as $r/(r+1)$. If FALSE, a conservative correlation of 0 is used.
K	Number of sequential looks in Phase 3.
b	A vector of efficacy boundaries (length $K+1$). The first element is the efficacy boundary for the Phase-2 test statistic; the remaining K elements are efficacy boundaries for the selected arm in Phase 3.
a	An optional vector of futility boundaries (length $K+1$). The first element is the futility boundary for the Phase-2 test statistic; the remaining K elements are futility boundaries for the selected arm in Phase 3. If omitted, no futility stopping is applied.
I	A vector of information levels (length $K+1$) for any active arm versus the common control. The first element is for Phase 2; the remaining K elements are for the looks in Phase 3.
nthreads	The number of threads to use (0 leaves the RcppParallel setting unchanged).

Details

The function assumes a multivariate normal distribution for the Wald statistics. Among designs that continue beyond the Phase-2 analysis, the arm with the largest Z-statistic at the end of Phase 2 is carried forward.

Decision Rules:

- **Phase 2 efficacy stop:** reject if the largest Phase-2 test statistic satisfies $Z_{(M)}(I_0) \geq b_0$.
- **Phase 2 futility stop:** stop for futility if the largest Phase-2 test statistic satisfies $Z_{(M)}(I_0) \leq a_0$.
- **Continue to Phase 3:** if $a_0 < Z_{(M)}(I_0) < b_0$, continue with the arm having the largest Phase-2 test statistic.
- **Phase 3 efficacy stop:** at look k , reject if the selected arm's Z-statistic exceeds the efficacy boundary and no earlier stop has occurred.
- **Phase 3 futility stop:** at look k , stop for futility if the selected arm's Z-statistic is below the futility boundary and no earlier stop has occurred.

Design Assumptions:

- All active arms share the same information level in Phase 2.
- Exactly one active arm is selected at the end of Phase 2 based on the largest observed Z-statistic when the trial continues to Phase 3.

Value

A list containing the following components:

- `exitProbUpper`: A vector of length $K + 1$. The first element is the probability of stopping for efficacy in Phase 2; the remaining elements are the probabilities of stopping for efficacy at each look in Phase 3.
- `exitProbLower`: A vector of length $K + 1$. The first element is the probability of stopping for futility in Phase 2; the remaining elements are the probabilities of stopping for futility at each look in Phase 3.
- `exitProbByArmUpper`: A $(K + 1) \times M$ matrix. The (k, m) -th entry gives the probability of stopping for efficacy at look k given that arm m is selected as the most promising.
- `exitProbByArmLower`: A $(K + 1) \times M$ matrix. The (k, m) -th entry gives the probability of stopping for futility at look k given that arm m is selected as the most promising.
- `selectionProb`: A vector of length M containing the probability that each active arm is selected as the most promising.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

References

Ping Gao, Yingqiu Li. Adaptive two-stage seamless sequential design for clinical trials. *Journal of Biopharmaceutical Statistics*, 2025, 35(4), 565-587.

Examples

```
# Setup: 2 active arms vs control in Phase 2; 1 selected arm vs control
# in Phase 3. Phase 3 has 2 sequential looks.

# Information levels: equal spacing over 3 looks based on a maximum of
# 110 patients per arm, SD = 1.0
I <- c(110 / (2 * 1.0^2) * seq(1, 3)/3)

# O'Brien-Fleming efficacy boundaries
b <- c(3.776605, 2.670463, 2.180424)

# No futility stopping
p0 <- exitprob_seamless(M = 2, theta = c(0, 0), K = 2,
                       b = b, I = I, nthreads = 1)
cumsum(p0$exitProbUpper)

# Add futility stopping
```

```

a <- c(0, 0.5, b[3])
p1 <- exitprob_seamless(M = 2, theta = c(0.3, 0.5), K = 2,
                        b = b, a = a, I = I, nthreads = 1)
cbind(
  cumulativeEfficacy = cumsum(p1$exitProbUpper),
  cumulativeFutility = cumsum(p1$exitProbLower)
)

```

fadjpbbon

*Adjusted p-Values for Bonferroni-Based Graphical Approaches***Description**

Obtains the adjusted p-values for graphical approaches using weighted Bonferroni tests.

Usage

```
fadjpbbon(p, wgtmat = NULL)
```

Arguments

p	The raw p-values for elementary hypotheses.
wgtmat	A list containing the weight matrix and the indicator matrix for intersection hypotheses. If NULL, equal weights are assigned within each intersection hypothesis.

Value

A list with the following components:

- inthyp: The indicator matrix for the intersection hypotheses.
- pinter: The local p-values for the intersection hypotheses.
- padj: The adjusted p-values for the elementary hypotheses.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Frank Bretz, Willi Maurer, Werner Brannath and Martin Posch. A graphical approach to sequentially rejective multiple test procedures. *Statistics in Medicine*. 2009; 28:586-604.

Examples

```
pvalues <- matrix(c(0.01,0.005,0.015,0.022, 0.02,0.015,0.010,0.023),
                  nrow=2, ncol=4, byrow=TRUE)
w <- c(0.5,0.5,0,0)
G <- matrix(c(0,0,1,0,0,0,0,1,0,1,0,0,1,0,0,0),
            nrow=4, ncol=4, byrow=TRUE)
wgtmat <- fwgtmat(w,G)
fadjpbon(pvalues, wgtmat)
```

fadjpdun

*Adjusted p-Values for Dunnett-Based Graphical Approaches***Description**

Obtains the adjusted p-values for graphical approaches using weighted Dunnett tests.

Usage

```
fadjpdun(p, wgtmat = NULL, family = NULL, corr = NULL, nthreads = 0)
```

Arguments

p	The raw p-values for elementary hypotheses.
wgtmat	A list containing the weight matrix and the indicator matrix for intersection hypotheses. If NULL, equal weights are assigned within each intersection hypothesis.
family	The matrix of family indicators for elementary hypotheses. Defaults to one family containing all elementary hypotheses.
corr	The correlation matrix that should be used for the parametric test. Can contain NAs for unknown correlations between families. By default, within-family correlations are 0.5 and between-family correlations are missing.
nthreads	The number of threads to use in simulations (0 means the default RcppParallel behavior).

Value

A list with the following components:

- inthyp: The indicator matrix for the intersection hypotheses.
- pinter: The local p-values for the intersection hypotheses.
- padj: The adjusted p-values for the elementary hypotheses.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Frank Bretz, Martin Posch, Ekkehard Glimm, Florian Klinglmueller, Willi Maurer, and Kornelius Rohmeyer. Graphical approach for multiple comparison procedures using weighted Bonferroni, Simes, or parameter tests. *Biometrical Journal*. 2011; 53:894-913.

Examples

```
pvalues <- matrix(c(0.01,0.005,0.015,0.022, 0.02,0.015,0.010,0.023),
                  nrow=2, ncol=4, byrow=TRUE)
w <- c(0.5,0.5,0,0)
G <- matrix(c(0,0,1,0,0,0,0,1,0,1,0,0,1,0,0,0),
            nrow=4, ncol=4, byrow=TRUE)
wgtmat <- fwgtmat(w,G)

family <- matrix(c(1,1,0,0,0,0,1,1), nrow=2, ncol=4, byrow=TRUE)
corr <- matrix(c(1,0.5,NA,NA, 0.5,1,NA,NA,
                NA,NA,1,0.5, NA,NA,0.5,1),
              nrow = 4, byrow = TRUE)
fadjpdun(pvalues, wgtmat, family, corr, nthreads = 1)
```

fadjpsim

Adjusted p-Values for Simes-Based Graphical Approaches

Description

Obtains the adjusted p-values for graphical approaches using weighted Simes tests.

Usage

```
fadjpsim(p, wgtmat = NULL, family = NULL)
```

Arguments

p	The raw p-values for elementary hypotheses.
wgtmat	A list containing the weight matrix and the indicator matrix for intersection hypotheses. If NULL, equal weights are assigned within each intersection hypothesis.
family	The matrix of family indicators for elementary hypotheses. Defaults to one family containing all elementary hypotheses.

Value

A list with the following components:

- inthyp: The indicator matrix for the intersection hypotheses.
- pinter: The local p-values for the intersection hypotheses.
- padj: The adjusted p-values for the elementary hypotheses.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Frank Bretz, Martin Posch, Ekkehard Glimm, Florian Klinglmueller, Willi Maurer, and Kornelius Rohmeyer. Graphical approach for multiple comparison procedures using weighted Bonferroni, Simes, or parameter tests. *Biometrical Journal*. 2011; 53:894-913.

Kaifeng Lu. Graphical approaches using a Bonferroni mixture of weighted Simes tests. *Statistics in Medicine*. 2016; 35:4041-4055.

Examples

```
pvalues <- matrix(c(0.01,0.005,0.015,0.022, 0.02,0.015,0.010,0.023),
                  nrow=2, ncol=4, byrow=TRUE)
w <- c(0.5,0.5,0,0)
G <- matrix(c(0,0,1,0,0,0,0,1,0,1,0,0,1,0,0,0),
            nrow=4, ncol=4, byrow=TRUE)
wgtmat <- fwgtmat(w,G)

family <- matrix(c(1,1,0,0,0,0,1,1), nrow=2, ncol=4, byrow=TRUE)
fadjpsim(pvalues, wgtmat, family)
```

fCERCer

Compute conditional error rates

Description

Compute conditional error rates for the intersection hypotheses after the first stage of a two-stage multiple testing procedure.

Usage

```
fCERCer(
  stg1_p,
  wgtmat,
  family = NULL,
  corr = NULL,
  info_frac,
  stg1_bnd,
  stg2_bnd,
  nthreads = 0
)
```

Arguments

stg1_p	P-values for the elementary hypotheses in stage 1.
wgtmat	Weight matrix for the intersection hypotheses.
family	Family matrix indicating which hypotheses belong to which families. If NULL, all hypotheses are assumed to belong to the same family.
corr	Correlation matrix for the test statistics.
info_frac	Information fraction for the first stage.
stg1_bnd	Stage 1 bounds for the elementary hypotheses in each intersection hypothesis.
stg2_bnd	Stage 2 bounds for the elementary hypotheses in each intersection hypothesis.
nthreads	The number of threads to use in simulations (0 means the default RcppParallel behavior).

Value

A list containing:

- stg1_elemhyp_r_idx: The 1-based indices of elementary hypotheses rejected in stage 1.
- stg1_inthyp_nr_idx: The 1-based indices of intersection hypotheses not rejected in stage 1.
- inthyp: The indicator matrix of those non-rejected intersection hypotheses.
- CER: The conditional error rate for each non-rejected intersection hypothesis.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Cyrus Mehta, Ajoy Mukhopadhyay, and Martin Posch. Graph Based, Adaptive, Multiarm, Multiple Endpoint, Two-Stage Designs. *Statistics in Medicine*. 2025.

Examples

```
initial_weights <- c(0.5, 0.5, 0, 0)
transition_matrix <- matrix(c(0, 0.5, 0.5, 0,
                             0.5, 0, 0, 0.5,
                             0, 1, 0, 0,
                             1, 0, 0, 0),
                           nrow = 4, byrow = TRUE)
wgtmat <- fwgtmat(initial_weights, transition_matrix)
family <- matrix(c(1, 1, 0, 0,
                  0, 0, 1, 1),
                 nrow = 2, byrow = TRUE)
corr <- matrix(c(1, 0.5, NA, NA,
                0.5, 1, NA, NA,
                NA, NA, 1, 0.5,
                NA, NA, 0.5, 1),
               nrow = 4, byrow = TRUE)
```

```

bounds <- fCERStageBound(
  wgtmat, family, corr, alpha = 0.025,
  alpha1 = errorSpent(0.5, 0.025, "sf0F"),
  info_frac = 0.5, nthreads = 1)
fCERCer(stg1_p = c(0.00045, 0.0952, 0.0225, 0.1104),
  wgtmat, family, corr, info_frac = 0.5,
  bounds$stg1_bnd, bounds$stg2_bnd, nthreads = 1)

```

fCERNewBound

Compute new stage-wise bounds after adaptation

Description

Compute new stage 2 bounds after an adaptation of the hypotheses tested or their weights.

Usage

```

fCERNewBound(
  stg1_p,
  wgtmat,
  family = NULL,
  corr = NULL,
  stg1_inthyp_nr_idx,
  CER,
  stg2_elemhyp_idx,
  stg2_wgtmat,
  info_frac_new,
  nthreads = 0
)

```

Arguments

stg1_p	P-values for the elementary hypotheses in stage 1.
wgtmat	Weight matrix for the original intersection hypotheses.
family	Family matrix indicating which hypotheses belong to which families. If NULL, all hypotheses are assumed to belong to the same family.
corr	Correlation matrix for the test statistics.
stg1_inthyp_nr_idx	Indices of the intersection hypotheses not rejected in stage 1.
CER	Conditional error rates for the intersection hypotheses.
stg2_elemhyp_idx	Indices of the elementary hypotheses in stage 2.
stg2_wgtmat	Weight matrix for the intersection hypotheses in stage 2.
info_frac_new	New information fraction for stage 1 after adaptation.
nthreads	The number of threads to use in simulations (0 means the default RcppParallel behavior).

Value

A list containing:

- `inhyp`: The indicator matrix of the adapted intersection hypotheses.
- `stg2_coef_new`: The new stage 2 coefficient for each adapted intersection hypothesis.
- `stg2_bnd_new`: The new stage 2 bounds for the elementary hypotheses in each adapted intersection hypothesis.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

References

Cyrus Mehta, Ajoy Mukhopadhyay, and Martin Posch. Graph Based, Adaptive, Multiarm, Multiple Endpoint, Two-Stage Designs. *Statistics in Medicine*. 2025.

Examples

```
initial_weights <- c(0.5, 0.5, 0, 0)
transition_matrix <- matrix(c(0, 0.5, 0.5, 0,
                             0.5, 0, 0, 0.5,
                             0, 1, 0, 0,
                             1, 0, 0, 0),
                           nrow = 4, byrow = TRUE)
wgtmat <- fwgtmat(initial_weights, transition_matrix)
family <- matrix(c(1, 1, 0, 0,
                  0, 0, 1, 1),
                nrow = 2, byrow = TRUE)
corr <- matrix(c(1, 0.5, NA, NA,
                0.5, 1, NA, NA,
                NA, NA, 1, 0.5,
                NA, NA, 0.5, 1),
              nrow = 4, byrow = TRUE)
stage1_pvalues <- c(0.00045, 0.0952, 0.0225, 0.1104)
bounds <- fCERStageBound(
  wgtmat, family, corr, alpha = 0.025,
  alpha1 = errorSpent(0.5, 0.025, "sfOF"),
  info_frac = 0.5, nthreads = 1)
conditional_error_rates <- fCERCer(
  stage1_pvalues, wgtmat, family, corr, info_frac = 0.5,
  bounds$stg1_bnd, bounds$stg2_bnd, nthreads = 1)
stage2_weight_matrix <- fwgtmat(
  w = c(0.5, 0.5),
  G = matrix(c(0, 1, 1, 0), 2, 2, byrow = TRUE))
fCERNewBound(stage1_pvalues, wgtmat, family, corr,
  conditional_error_rates$stg1_inhyp_nr_idx,
  conditional_error_rates$CER,
  stg2_elemhyp_idx = c(2, 4),
  stage2_weight_matrix, info_frac_new = 0.4,
  nthreads = 1)
```

fCERRej

*Determine stage 2 rejections from conditional error rates***Description**

Determine the elementary hypotheses rejected after stage 2 using adapted stage 2 bounds and cumulative p-values.

Usage

```
fCERRej(cum_p, stg1_elemhyp_r_idx, stg2_elemhyp_idx, stg2_inthyp, stg2_bnd_new)
```

Arguments

cum_p	Cumulative p-values for the elementary hypotheses in stage 2.
stg1_elemhyp_r_idx	Indices of the elementary hypotheses rejected in stage 1.
stg2_elemhyp_idx	Indices of the elementary hypotheses in stage 2.
stg2_inthyp	Indicator matrix of the intersection hypotheses in stage 2.
stg2_bnd_new	New stage 2 bounds for the elementary hypotheses in each intersection hypothesis.

Value

A logical vector containing:

- One value for each elementary hypothesis, indicating whether it is rejected after stage 2.
- Values are ordered by the elementary-hypothesis indices of the original procedure.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Cyrus Mehta, Ajoy Mukhopadhyay, and Martin Posch. Graph Based, Adaptive, Multiarm, Multiple Endpoint, Two-Stage Designs. *Statistics in Medicine*. 2025.

Examples

```

initial_weights <- c(0.5, 0.5, 0, 0)
transition_matrix <- matrix(c(0, 0.5, 0.5, 0,
                             0.5, 0, 0, 0.5,
                             0, 1, 0, 0,
                             1, 0, 0, 0),
                             nrow = 4, byrow = TRUE)
wgtmat <- fwgtmat(initial_weights, transition_matrix)
family <- matrix(c(1, 1, 0, 0,
                  0, 0, 1, 1), nrow = 2, byrow = TRUE)
corr <- matrix(c(1, 0.5, NA, NA,
                0.5, 1, NA, NA,
                NA, NA, 1, 0.5,
                NA, NA, 0.5, 1),
                nrow = 4, byrow = TRUE)
stage1_pvalues <- c(0.00045, 0.0952, 0.0225, 0.1104)
bounds <- fCERStageBound(
  wgtmat, family, corr, alpha = 0.025,
  alpha1 = errorSpent(0.5, 0.025, "sfOF"),
  info_frac = 0.5, nthreads = 1)
conditional_error_rates <- fCERCer(
  stage1_pvalues, wgtmat, family, corr, info_frac = 0.5,
  bounds$stg1_bnd, bounds$stg2_bnd, nthreads = 1)
stage2_weight_matrix <- fwgtmat(
  w = c(0.5, 0.5),
  G = matrix(c(0, 1, 1, 0), 2, 2, byrow = TRUE))
adapted_bounds <- fCERNewBound(
  stage1_pvalues, wgtmat, family, corr,
  conditional_error_rates$stg1_inhyp_nr_idx,
  conditional_error_rates$CER,
  stg2_elemhyp_idx = c(2, 4), stage2_weight_matrix,
  info_frac_new = 0.4, nthreads = 1)
stage2_cumulative_pvalues <-
  1 - pnorm(sqrt(0.4) * qnorm(1 - c(0.0952, 0.1104))) +
  sqrt(1 - 0.4) * qnorm(1 - c(0.0299, 0.0586)))
fCERRej(stage2_cumulative_pvalues,
  conditional_error_rates$stg1_elemhyp_r_idx,
  stg2_elemhyp_idx = c(2, 4),
  adapted_bounds$inhyp, adapted_bounds$stg2_bnd_new)

```

fCERStageBound

*Compute stage-wise bounds for multiple testing***Description**

Compute stage-wise bounds for a two-stage multiple testing procedure.

Usage

```
fCERStageBound(
  wgtmat,
  family = NULL,
  corr = NULL,
  alpha,
  alpha1,
  info_frac,
  nthreads = 0
)
```

Arguments

<code>wgtmat</code>	Weight matrix for the intersection hypotheses
<code>family</code>	Family matrix indicating which hypotheses belong to which families. The correlation is known only for hypotheses belonging to the same family. If <code>NULL</code> , all hypotheses are assumed to belong to the same family.
<code>corr</code>	Correlation matrix for the test statistics
<code>alpha</code>	Overall significance level
<code>alpha1</code>	Significance level for the first stage
<code>info_frac</code>	Information fraction for the first stage
<code>nthreads</code>	The number of threads to use in simulations (0 means the default <code>RcppParallel</code> behavior).

Value

A list containing:

- `inthy`: The indicator matrix of the intersection hypotheses.
- `stg1_coef`: The stage 1 coefficient for each intersection hypothesis.
- `stg2_coef`: The stage 2 coefficient for each intersection hypothesis.
- `stg1_bnd`: The stage 1 bounds for the elementary hypotheses in each intersection hypothesis.
- `stg2_bnd`: The stage 2 bounds for the elementary hypotheses in each intersection hypothesis.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Cyrus Mehta, Ajoy Mukhopadhyay, and Martin Posch. Graph Based, Adaptive, Multiarm, Multiple Endpoint, Two-Stage Designs. *Statistics in Medicine*. 2025.

Examples

```

initial_weights <- c(0.5, 0.5, 0, 0)
transition_matrix <- matrix(c(0, 0.5, 0.5, 0,
                             0.5, 0, 0, 0.5,
                             0, 1, 0, 0,
                             1, 0, 0, 0),
                           nrow = 4, byrow = TRUE)
wgtmat <- fwgtmat(initial_weights, transition_matrix)
family <- matrix(c(1, 1, 0, 0,
                  0, 0, 1, 1),
                nrow = 2, byrow = TRUE)
corr <- matrix(c(1, 0.5, NA, NA,
                0.5, 1, NA, NA,
                NA, NA, 1, 0.5,
                NA, NA, 0.5, 1),
              nrow = 4, byrow = TRUE)
alpha <- 0.025
alpha1 <- errorSpent(0.5, alpha, "sfOF")
fCERStageBound(wgtmat, family, corr, alpha,
              alpha1, info_frac = 0.5, nthreads = 1)

```

fDefaultWgtmat

*Default Weight Matrix for All Intersection Hypotheses***Description**

Obtains the default weight matrix for all intersection hypotheses, assigning equal weights to the elementary hypotheses within each intersection hypothesis.

Usage

```
fDefaultWgtmat(m)
```

Arguments

m The number of elementary hypotheses.

Value

A list with the following components:

- **inthy**: The indicator matrix for the intersection hypotheses.
- **wgtmat**: The default weight matrix for the elementary hypotheses.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
fDefaultWgtmat(3)
```

finthyp

Indicator Matrix for All Intersection Hypotheses

Description

Obtains the indicator matrix for all intersection hypotheses formed from a given number of elementary hypotheses, matching the row order used by fwgtmat and fDefaultWgtmat.

Usage

```
finthyp(m)
```

Arguments

`m` The number of elementary hypotheses.

Value

The $(2^m - 1) \times m$ indicator matrix for the intersection hypotheses, with row 1 the global intersection (all hypotheses active) and the last row the singleton $\{m\}$.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
finthyp(3)
```

float_to_fraction

Converting a decimal to a fraction

Description

Converts a decimal to a fraction based on the algorithm from <http://stackoverflow.com/a/5128558/221955>.

Usage

```
float_to_fraction(x, tol = 1e-06)
```

Arguments

x	The fraction in decimal form.
tol	The tolerance level for the conversion error.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
float_to_fraction(5/3)
```

fmodmix

Adjusted p-Values for Modified Mixture Gatekeeping Procedures

Description

Obtains the adjusted p-values for the modified gatekeeping procedures for multiplicity problems involving serial and parallel logical restrictions.

Usage

```
fmodmix(
  p,
  family = NULL,
  serial,
  parallel = NULL,
  gamma,
  test = "hommel",
  exhaust = TRUE
)
```

Arguments

p	The raw p-values for elementary hypotheses.
family	The matrix of family indicators for the hypotheses.
serial	The matrix of serial rejection set for the hypotheses.
parallel	The matrix of parallel rejection set for the hypotheses.
gamma	The truncation parameters for each family. The truncation parameter for the last family is automatically set to 1.
test	The component multiple testing procedure. Options include "holm", "hochberg", or "hommel". Defaults to "hommel".
exhaust	Whether to use alpha-exhausting component testing procedure for the last family with active hypotheses. It defaults to TRUE.

Value

A list with the following components:

- `inthyp`: The indicator matrix for the intersection hypotheses.
- `pinter`: The local p-values for the intersection hypotheses.
- `padj`: The adjusted p-values for the elementary hypotheses.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Alex Dmitrienko, George Kordzakhia, and Thomas Brechenmacher. Mixture-based gatekeeping procedures for multiplicity problems with multiple sequences of hypotheses. *Journal of Biopharmaceutical Statistics*. 2016; 26(4):758–780.

George Kordzakhia, Thomas Brechenmacher, Eiji Ishida, Alex Dmitrienko, Winston Wenxiang Zheng, and David Fuyuan Li. An enhanced mixture method for constructing gatekeeping procedures in clinical trials. *Journal of Biopharmaceutical Statistics*. 2018; 28(1):113–128.

Examples

```
p <- c(0.0194, 0.0068, 0.0271, 0.0088, 0.0370, 0.0018, 0.0814, 0.0066)
family <- matrix(c(1, 1, 0, 0, 0, 0, 0, 0,
                  0, 0, 1, 1, 0, 0, 0, 0,
                  0, 0, 0, 0, 1, 1, 0, 0,
                  0, 0, 0, 0, 0, 0, 1, 1),
                nrow=4, byrow=TRUE)

serial <- matrix(c(0, 0, 0, 0, 0, 0, 0, 0,
                 0, 0, 0, 0, 0, 0, 0, 0,
                 1, 0, 0, 0, 0, 0, 0, 0,
                 0, 1, 0, 0, 0, 0, 0, 0,
                 0, 0, 1, 0, 0, 0, 0, 0,
                 0, 0, 0, 1, 0, 0, 0, 0,
                 0, 0, 0, 0, 1, 0, 0, 0,
                 0, 0, 0, 0, 0, 1, 0, 0),
                nrow=8, byrow=TRUE)

parallel <- matrix(0, 8, 8)
gamma <- c(0.6, 0.6, 0.6, 1)
fmodmix(p, family, serial, parallel, gamma, test = "hommel", exhaust = TRUE)
```

fPCRej

*Determine stage 2 rejections using p-value combination***Description**

Combine stage 1 and stage 2 local p-values and determine the elementary hypotheses rejected after stage 2.

Usage

```
fPCRej(
  stg1_loc_p,
  stg2_loc_p,
  stg1_elemhyp_r_idx,
  stg2_elemhyp_idx,
  alpha,
  info_frac
)
```

Arguments

stg1_loc_p	A list returned by fPCStagewise for stage 1.
stg2_loc_p	A list returned by fPCStagewise for stage 2.
stg1_elemhyp_r_idx	Indices of the elementary hypotheses rejected at stage 1.
stg2_elemhyp_idx	Indices of the elementary hypotheses tested at stage 2.
alpha	Overall significance level.
info_frac	Information fraction for stage 1.

Value

A list containing:

- stg1_inthyp_nr_idx: The 1-based indices of intersection hypotheses retained from stage 1.
- stg2_elemhyp_idx: The 1-based indices of elementary hypotheses tested in stage 2.
- inthyp: The indicator matrix of the stage 2 intersection hypotheses.
- stg1_pinter: The stage 1 local p-values.
- stg2_pinter: The stage 2 local p-values.
- comb_pinter: The combined local p-values.
- rej_elem: A logical vector indicating the elementary hypotheses rejected after stage 2.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

References

Cyrus Mehta, Ajoy Mukhopadhyay, and Martin Posch. Graph Based, Adaptive, Multiarm, Multiple Endpoint, Two-Stage Designs. *Statistics in Medicine*. 2025.

Examples

```
initial_weights <- c(0.5, 0.5, 0, 0)
transition_matrix <- matrix(c(0, 0.5, 0.5, 0,
                             0.5, 0, 0, 0.5,
                             0, 1, 0, 0,
                             1, 0, 0, 0),
                             nrow = 4, byrow = TRUE)
wgtmat <- fwgtmat(initial_weights, transition_matrix)
family <- matrix(c(1, 1, 0, 0,
                  0, 0, 1, 1),
                  nrow = 2, byrow = TRUE)
corr <- matrix(c(1, 0.5, NA, NA,
                0.5, 1, NA, NA,
                NA, NA, 1, 0.5,
                NA, NA, 0.5, 1),
                nrow = 4, byrow = TRUE)
alpha <- 0.025
alpha1 <- errorSpent(0.5, alpha, "sfOF")
stage1_pvalues <- c(0.00045, 0.0952, 0.0225, 0.1104)
stage1_loc_p <- fPCStagewise(
  stg2_p = stage1_pvalues, wgtmat = wgtmat,
  family = family, corr = corr,
  stg1_inhyp_nr_idx = 1:15, stg2_elemhyp_idx = 1:4,
  stg2_wgtmat = wgtmat, test = "dunnett",
  nthreads = 1)
stage1_rejections <- fPCStage1(stage1_loc_p, alpha1)

adapted_graph <- updateGraph(initial_weights,
                             transition_matrix,
                             I = 1:4, j = 1)
stage2_weight_matrix <- fwgtmat(
  w = adapted_graph$w[adapted_graph$I],
  G = adapted_graph$G[adapted_graph$I, adapted_graph$I])
stage2_pvalues <- c(0.1121, 0.0112, 0.1153)
stage2_loc_p <- fPCStagewise(
  stg2_p = stage2_pvalues, wgtmat = wgtmat,
  family = family, corr = corr,
  stg1_inhyp_nr_idx = stage1_rejections$stg1_inhyp_nr_idx,
  stg2_elemhyp_idx = adapted_graph$I,
  stg2_wgtmat = stage2_weight_matrix,
  test = "dunnett", nthreads = 1)
fPCRej(stg1_loc_p = stage1_loc_p, stg2_loc_p = stage2_loc_p,
       stg1_elemhyp_r_idx = stage1_rejections$stg1_elemhyp_r_idx,
       stg2_elemhyp_idx = adapted_graph$I, alpha = alpha,
       info_frac = 0.5)

# Change weights for the elementary hypotheses.
```

```

stage2_weight_matrix_reweighted <- fwgtmat(
  w = c(0.5, 0.25, 0.25),
  G = adapted_graph$G[adapted_graph$I, adapted_graph$I])
stage2_loc_p_reweighted <- fPCStagewise(
  stg2_p = stage2_pvalues, wgtmat = wgtmat,
  family = family, corr = corr,
  stg1_inthyp_nr_idx = stage1_rejections$stg1_inthyp_nr_idx,
  stg2_elemhyp_idx = adapted_graph$I,
  stg2_wgtmat = stage2_weight_matrix_reweighted,
  test = "dunnett", nthreads = 1)
fPCRej(stg1_loc_p = stage1_loc_p,
  stg2_loc_p = stage2_loc_p_reweighted,
  stg1_elemhyp_r_idx = stage1_rejections$stg1_elemhyp_r_idx,
  stg2_elemhyp_idx = adapted_graph$I,
  alpha = alpha, info_frac = 0.5)

```

fPCStage1

*Determine stage 1 rejections for adaptive multiple testing***Description**

Determine the elementary hypotheses rejected at stage 1 and the intersection hypotheses retained for stage 2.

Usage

```
fPCStage1(stg1_loc_p, alpha1)
```

Arguments

stg1_loc_p	A list returned by fPCStagewise, containing the stage 1 intersection hypotheses and local p-values.
alpha1	Stage 1 significance level.

Value

A list containing:

- stg1_elemhyp_r_idx: The 1-based indices of elementary hypotheses rejected in stage 1.
- stg1_inthyp_nr_idx: The 1-based indices of intersection hypotheses not rejected in stage 1.
- inthyp: The indicator matrix of those non-rejected intersection hypotheses.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

References

Cyrus Mehta, Ajoy Mukhopadhyay, and Martin Posch. Graph Based, Adaptive, Multiarm, Multiple Endpoint, Two-Stage Designs. *Statistics in Medicine*. 2025.

Examples

```
wgtmat <- fwgtmat(c(0.5, 0.5, 0, 0),
                  matrix(c(0, 0.5, 0.5, 0,
                          0.5, 0, 0, 0.5,
                          0, 1, 0, 0,
                          1, 0, 0, 0),
                        nrow = 4, byrow = TRUE))
family <- matrix(c(1, 1, 0, 0,
                  0, 0, 1, 1),
                nrow = 2, byrow = TRUE)
corr <- matrix(c(1, 0.5, NA, NA,
                0.5, 1, NA, NA,
                NA, NA, 1, 0.5,
                NA, NA, 0.5, 1),
              nrow = 4, byrow = TRUE)
stage1_local_pvalues <- fPCStagewise(
  stg2_p = c(0.00045, 0.0952, 0.0225, 0.1104),
  wgtmat = wgtmat, family = family, corr = corr,
  stg1_inthyp_nr_idx = 1:15, stg2_elemhyp_idx = 1:4,
  stg2_wgtmat = wgtmat, test = "dunnett",
  nthreads = 1)
fPCStage1(stg1_loc_p = stage1_local_pvalues,
          alpha1 = errorSpent(0.5, 0.025, "sfOF"))
```

fPCStagewise

Compute stage-wise local p-values

Description

Compute local p-values for the intersection hypotheses in a two-stage adaptive multiple testing procedure using a p-value combination method.

Usage

```
fPCStagewise(
  stg2_p,
  wgtmat = NULL,
  family = NULL,
  corr = NULL,
  stg1_inthyp_nr_idx,
  stg2_elemhyp_idx,
  stg2_wgtmat = NULL,
  test = "dunnett",
```

```

    nthreads = 0
  )

```

Arguments

stg2_p	Stage 2 p-values for the elementary hypotheses.
wgtmat	Weight matrix for the stage 1 intersection hypotheses. If NULL, equal weights are assigned within each intersection hypothesis.
family	Family matrix indicating which hypotheses belong to which families. The correlation is known only for hypotheses belonging to the same family. Defaults to one family containing all elementary hypotheses.
corr	Correlation matrix for the test statistics. If NULL, within-family correlations are 0.5 and between-family correlations are missing.
stg1_inthyp_nr_idx	Indices of the stage 1 intersection hypotheses that were not rejected.
stg2_elemhyp_idx	Indices of the elementary hypotheses tested at stage 2.
stg2_wgtmat	Weight matrix for the stage 2 intersection hypotheses. If NULL, equal weights are assigned within each intersection hypothesis.
test	P-value combination method. It can start with "bon", "sim", or "dun"; the default is "dunnett".
nthreads	The number of threads to use in simulations (0 means the default RcppParallel behavior).

Details

Despite the stage-oriented parameter names, this function can also be used to generate stage 1 local p-values. In that case, provide the complete set of intersection hypotheses in `stg1_inthyp_nr_idx`, all elementary hypotheses in `stg2_elemhyp_idx`, and the corresponding stage 1 p-values and weight matrix. The example illustrates this use.

Value

A list containing:

- `inthyp_idx`: The 1-based indices of the stage 1 intersection hypotheses represented in the output.
- `inthyp`: Their intersection-hypothesis indicator matrix.
- `pinter`: Their local p-values.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Cyrus Mehta, Ajoy Mukhopadhyay, and Martin Posch. Graph Based, Adaptive, Multiarm, Multiple Endpoint, Two-Stage Designs. *Statistics in Medicine*. 2025.

Examples

```

initial_weights <- c(0.5, 0.5, 0, 0)
transition_matrix <- matrix(c(0, 0.5, 0.5, 0,
                             0.5, 0, 0, 0.5,
                             0, 1, 0, 0,
                             1, 0, 0, 0),
                           nrow = 4, byrow = TRUE)
wgtmat <- fwgtmat(initial_weights, transition_matrix)
family <- matrix(c(1, 1, 0, 0,
                  0, 0, 1, 1),
                nrow = 2, byrow = TRUE)
corr <- matrix(c(1, 0.5, NA, NA,
                0.5, 1, NA, NA,
                NA, NA, 1, 0.5,
                NA, NA, 0.5, 1),
              nrow = 4, byrow = TRUE)
fPCStagewise(stg2_p = c(0.00045, 0.0952, 0.0225, 0.1104),
             wgtmat = wgtmat, family = family, corr = corr,
             stg1_inhyp_nr_idx = 1:15, stg2_elehyp_idx = 1:4,
             stg2_wgtmat = wgtmat, test = "dunnett",
             nthreads = 1)

```

fquantile

*The Quantiles of a Survival Distribution***Description**

Obtains the quantiles of a survival distribution.

Usage

```
fquantile(S, probs, ...)
```

Arguments

S	The survival function of a univariate survival time.
probs	The numeric vector of probabilities.
...	Additional arguments to be passed to S.

Value

A vector of length(probs) for the quantiles.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
fquantile(pweibull, probs = c(0.25, 0.5, 0.75),
          shape = 1.37, scale = 1/0.818, lower.tail = FALSE)
```

fseqbon	<i>Group Sequential Trials Using Bonferroni-Based Graphical Approaches</i>
---------	--

Description

Obtains the test results for group sequential trials using graphical approaches based on weighted Bonferroni tests.

Usage

```
fseqbon(
  w,
  G,
  alpha = 0.025,
  kMax,
  typeAlphaSpending = NULL,
  parameterAlphaSpending = NULL,
  maxInformation = NULL,
  incidenceMatrix = NULL,
  k1,
  p,
  information,
  spendingTime = NULL,
  lookback = TRUE,
  nthreads = 0
)
```

Arguments

w	The vector of initial weights for elementary hypotheses.
G	The initial transition matrix.
alpha	The significance level. Defaults to 0.025.
kMax	The maximum number of stages.
typeAlphaSpending	The vector of alpha spending functions for the hypotheses. Each element is one of the following: "sf0F" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function. Defaults to "sf0F" if not provided.

parameterAlphaSpending	The vector of parameter values for the alpha spending functions for the hypotheses. Each element corresponds to the value of ρ for "sfKD" or γ for "sfHSD". Defaults to missing if not provided.
maxInformation	The vector of target maximum information for each hypothesis. Defaults to a vector of 1s if not provided.
incidenceMatrix	The $k_{\text{Max}} \times m$ incidence matrix indicating whether the specific hypothesis will be tested at the given look. Here m is the number of hypotheses. If not provided, defaults to testing each hypothesis at all study looks.
k1	The number of study looks at the interim analysis.
p	The $Bk_1 \times m$ matrix of raw p-values for each hypothesis by study look. Here B is the number of replications for simulations. In other words, the number of rows must be a multiple of k_1 .
information	The $Bk_1 \times m$ matrix of observed information for each hypothesis by study look.
spendingTime	The $Bk_1 \times m$ matrix of spending time for alpha spending by study look. Each element must be between 0 and 1, and the spending time must be increasing by study look for each hypothesis. If not provided, it is the same as <code>informationRates</code> calculated from <code>information</code> and <code>maxInformation</code> .
lookback	Whether to allow retesting at earlier looks. It defaults to TRUE.
nthreads	The number of threads to use in simulations (0 means the default RcppParallel behavior).

Details

When `lookback = TRUE`, the procedure allows the user to retest an unrejected hypothesis at earlier looks if its weight increased after rejection of other hypotheses. The procedure will return the first look at which the specific hypothesis is rejected. If the hypothesis is not rejected at any look, it will return 0.

Value

A vector to indicate the first look the specific hypothesis is rejected (0 if the hypothesis is not rejected).

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Willi Maurer and Frank Bretz. Multiple testing in group sequential trials using graphical approaches. *Statistics in Biopharmaceutical Research*. 2013; 5:311-320.

Examples

```
# Case study from Maurer & Bretz (2013)

fseqbon(
  w = c(0.5, 0.5, 0, 0),
  G = matrix(c(0, 0.5, 0.5, 0, 0.5, 0, 0, 0.5,
               0, 1, 0, 0, 1, 0, 0, 0),
             nrow=4, ncol=4, byrow=TRUE),
  alpha = 0.025,
  kMax = 3,
  typeAlphaSpending = rep("sf0F", 4),
  maxInformation = rep(1, 4),
  k1 = 2,
  p = matrix(c(0.0062, 0.017, 0.009, 0.13,
               0.0002, 0.0035, 0.002, 0.06),
             nrow=2, ncol=4, byrow=TRUE),
  information = matrix(c(rep(1/3, 4), rep(2/3, 4)),
                      nrow=2, ncol=4, byrow=TRUE),
  nthreads = 1)
```

fstdmix

Adjusted p-Values for Standard Mixture Gatekeeping Procedures

Description

Obtains the adjusted p-values for the standard gatekeeping procedures for multiplicity problems involving serial and parallel logical restrictions.

Usage

```
fstdmix(
  p,
  family = NULL,
  serial,
  parallel = NULL,
  gamma,
  test = "hommel",
  exhaust = TRUE
)
```

Arguments

p	The raw p-values for elementary hypotheses.
family	The matrix of family indicators for the hypotheses.
serial	The matrix of serial rejection set for the hypotheses.

<code>parallel</code>	The matrix of parallel rejection set for the hypotheses.
<code>gamma</code>	The truncation parameters for each family. The truncation parameter for the last family is automatically set to 1.
<code>test</code>	The component multiple testing procedure. Options include "holm", "hochberg", or "hommel". Defaults to "hommel".
<code>exhaust</code>	Whether to use alpha-exhausting component testing procedure for the last family with active hypotheses. It defaults to TRUE.

Value

A list with the following components:

- `inthy`: The indicator matrix for the intersection hypotheses.
- `pinter`: The local p-values for the intersection hypotheses.
- `padj`: The adjusted p-values for the elementary hypotheses.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Alex Dmitrienko and Ajit C Tamhane. Mixtures of multiple testing procedures for gatekeeping applications in clinical trials. *Statistics in Medicine*. 2011; 30(13):1473–1488.

Examples

```
p <- c(0.0194, 0.0068, 0.0271, 0.0088, 0.0370, 0.0018, 0.0814, 0.0066)
family <- matrix(c(1, 1, 0, 0, 0, 0, 0, 0,
                  0, 0, 1, 1, 0, 0, 0, 0,
                  0, 0, 0, 0, 1, 1, 0, 0,
                  0, 0, 0, 0, 0, 0, 1, 1),
                nrow=4, byrow=TRUE)

serial <- matrix(c(0, 0, 0, 0, 0, 0, 0, 0,
                  0, 0, 0, 0, 0, 0, 0, 0,
                  1, 0, 0, 0, 0, 0, 0, 0,
                  0, 1, 0, 0, 0, 0, 0, 0,
                  0, 0, 1, 0, 0, 0, 0, 0,
                  0, 0, 0, 1, 0, 0, 0, 0,
                  0, 0, 0, 0, 1, 0, 0, 0,
                  0, 0, 0, 0, 0, 1, 0, 0),
                nrow=8, byrow=TRUE)

parallel <- matrix(0, 8, 8)
gamma <- c(0.6, 0.6, 0.6, 1)
fstdmix(p, family, serial, parallel, gamma, test = "hommel",
        exhaust = FALSE)
```

fstp2seq	<i>Adjusted p-Values for Stepwise Testing Procedures for Two Sequences</i>
----------	--

Description

Obtains the adjusted p-values for the stepwise gatekeeping procedures for multiplicity problems involving two sequences of hypotheses.

Usage

```
fstp2seq(p, gamma, test = "hochberg", retest = TRUE)
```

Arguments

p	The raw p-values for elementary hypotheses.
gamma	The truncation parameters for each family. The truncation parameter for the last family is automatically set to 1.
test	The component multiple testing procedure. It is either "Holm" or "Hochberg", and it defaults to "Hochberg".
retest	Whether to allow retesting. It defaults to TRUE.

Value

A matrix of adjusted p-values.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
p <- c(0.0194, 0.0068, 0.0271, 0.0088, 0.0370, 0.0018, 0.0814, 0.0066)
gamma <- c(0.6, 0.6, 0.6, 1)
fstp2seq(p, gamma, test="hochberg", retest=1)
```

ftrunc

*Adjusted p-Values for Holm, Hochberg, and Hommel Procedures***Description**

Obtains the adjusted p-values for possibly truncated Holm, Hochberg, and Hommel procedures.

Usage

```
ftrunc(p, test = "hommel", gamma = 1)
```

Arguments

p	The raw p-values for elementary hypotheses.
test	The test to use, e.g., "holm", "hochberg", or "hommel" (default).
gamma	The value of the truncation parameter. Defaults to 1 for the regular Holm, Hochberg, or Hommel procedure.

Value

A list with the following components:

- inthyp: The indicator matrix for the intersection hypotheses.
- pinter: The local p-values for the intersection hypotheses.
- padj: The adjusted p-values for the elementary hypotheses.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Alex Dmitrienko, Ajit C. Tamhane, and Brian L. Wiens. General multistage gatekeeping procedures. *Biometrical Journal*. 2008; 5:667-677.

Examples

```
pvalues <- matrix(c(0.01,0.005,0.015,0.022, 0.02,0.015,0.010,0.023),
                  nrow=2, ncol=4, byrow=TRUE)
ftrunc(pvalues, "hochberg")
```

fwgtmat

*Weight Matrix for All Intersection Hypotheses***Description**

Obtains the weight matrix for all intersection hypotheses.

Usage

```
fwgtmat(w, G)
```

Arguments

w The vector of weights for elementary hypotheses.
G The transition matrix.

Value

The weight matrix starting with the global null hypothesis.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
w <- c(0.5, 0.5, 0, 0)
G <- matrix(c(0, 0, 1, 0, 0, 0, 0, 1, 0, 1, 0, 0, 1, 0, 0, 0),
            nrow=4, ncol=4, byrow=TRUE)
(wgtmat <- fwgtmat(w, G))
```

getAccrualDurationFromN

*Accrual Duration to Enroll Target Number of Subjects***Description**

Obtains the accrual duration to enroll the target number of subjects.

Usage

```
getAccrualDurationFromN(nsubjects, accrualTime, accrualIntensity)
```

Arguments

`nsubjects` The vector of target number of subjects.

`accrualTime` A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., `c(0, 3)` breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.

`accrualIntensity` A vector of accrual intensities. One for each accrual time interval.

Value

A vector of accrual durations.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
getAccrualDurationFromN(nsubjects = c(20, 150), accrualTime = c(0, 3),
                        accrualIntensity = c(10, 20))
```

getADCI

Confidence Interval After Adaptation

Description

Obtains the p-value, median unbiased point estimate, and confidence interval after the end of an adaptive trial.

Usage

```
getADCI(
  L = NA_integer_,
  zL = NA_real_,
  IMax = NA_real_,
  kMax = NA_integer_,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  criticalValues = NA_real_,
  alpha = 0.25,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  spendingTime = NA_real_,
  MullerSchafer = FALSE,
  Lc = NA_integer_,
  zLc = NA_real_,
```

```

    INew = NA_real_,
    informationRatesNew = NA_real_,
    efficacyStoppingNew = NA_integer_,
    typeAlphaSpendingNew = "sfOF",
    parameterAlphaSpendingNew = NA_real_,
    spendingTimeNew = NA_real_
)

```

Arguments

L	The interim adaptation look of the primary trial.
zL	The z-test statistic at the interim adaptation look of the primary trial.
IMax	The maximum information of the primary trial.
kMax	The maximum number of stages of the primary trial.
informationRates	The information rates of the primary trial.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage of the primary trial. Defaults to true if left unspecified.
criticalValues	The upper boundaries on the z-test statistic scale for efficacy stopping for the primary trial.
alpha	The significance level of the primary trial. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending for the primary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value of alpha spending for the primary trial. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
spendingTime	The error spending time of the primary trial. Defaults to missing, in which case, it is the same as informationRates.
MullerSchafer	Whether to use the Muller and Schafer (2001) method for trial adaptation.
Lc	The termination look of the integrated trial.
zLc	The z-test statistic at the termination look of the integrated trial.
INew	The maximum information of the secondary trial.
informationRatesNew	The spacing of looks of the secondary trial up to look L2.
efficacyStoppingNew	The indicators of whether efficacy stopping is allowed at each look of the secondary trial up to look L2. Defaults to true if left unspecified.

typeAlphaSpendingNew

The type of alpha spending for the secondary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early efficacy stopping. Defaults to "sfOF".

parameterAlphaSpendingNew

The parameter value of alpha spending for the secondary trial. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".

spendingTimeNew

The error spending time of the secondary trial up to look L2. Defaults to missing, in which case, it is the same as informationRatesNew.

Details

If typeAlphaSpendingNew is "OF", "P", "WT", or "none", then informationRatesNew, efficacyStoppingNew, and spendingTimeNew must be of full length kNew, and informationRatesNew and spendingTimeNew must end with 1.

Value

A data frame with the following variables:

- pvalue: p-value for rejecting the null hypothesis.
- thetihat: Median unbiased point estimate of the parameter.
- cilevel: Confidence interval level.
- lower: Lower bound of confidence interval.
- upper: Upper bound of confidence interval.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Ping Gao, Lingyun Liu and Cyrus Mehta. Exact inference for adaptive group sequential designs. Stat Med. 2013;32(23):3991-4005.

See Also

[adaptDesign](#)

Examples

```
# two-arm randomized clinical trial with a normally distributed endpoint
# 90% power to detect mean difference of 15 with a standard deviation of 50
# Design the Stage I Trial with 3 looks and Lan-DeMets O'Brien-Fleming type
# spending function
```

```

delta <- 15
sigma <- 50

(des1 <- getDesignMeanDiff(
  beta = 0.1, meanDiff = delta, stDev = sigma,
  kMax = 3, alpha = 0.025, typeAlphaSpending = "sfOF"
))

s1 <- des1$byStageResults$informationRates
b1 <- des1$byStageResults$efficacyBounds
n <- des1$overallResults$numberOfSubjects

# Monitoring the Stage I Trial
L <- 1
nL <- des1$byStageResults$numberOfSubjects[L]
deltahat <- 8
sigmahat <- 55
sedeltahat <- sigmahat * sqrt( 4 / nL)
zL <- deltahat / sedeltahat

# Making an Adaptive Change: Stage I to Stage II
# revised clinically meaningful difference downward to 10 power the study
# retain the standard deviation at the design stage
# Muller & Schafer (2001) method to design the secondary trial
# with 2 looks and Lan-DeMets Pocock type spending function
# re-estimate sample size to reach 90% conditional power
deltaNew <- 10

(des2 <- adaptDesign(
  betaNew = 0.1, L = L, zL = zL, theta = deltaNew,
  IMax = n / (4 * sigma^2), kMax = 3, informationRates = s1,
  alpha = 0.025, typeAlphaSpending = "sfOF",
  MullerSchafer = TRUE, kNew = 2, typeAlphaSpendingNew = "sfP"
))

INew <- des2$secondaryTrial$maxInformation
(nNew <- ceiling(INew * 4 * sigma^2))
(nTotal <- nL + nNew)

# Monitoring the Integrated Trial
s2 <- des2$secondaryTrial$informationRates

Lc <- 2
deltahatc <- 9.5
sigmahatc <- 52.759
L2 <- Lc - L
nL2 <- nNew * s2[L2]
nc <- nL + nL2
sedeltahatc <- sigmahatc * sqrt(4 / nc)
zLc <- deltahatc / sedeltahatc
zL2 <- (zLc * sqrt(nc) - zL * sqrt(nL)) / sqrt(nL2)

getADCI(

```

```

L = L, zL = zL, IMax = n / (4 * sigmahatc^2), kMax = 3,
informationRates = s1, alpha = 0.025, typeAlphaSpending = "sfOF",
MullerSchafer = TRUE, Lc = Lc, zLc = zLc,
INew = nNew / (4 * sigmahatc^2), informationRatesNew = s2,
typeAlphaSpendingNew = "sfP")

```

getADCI_multiarm	<i>Confidence Interval After Adaptation for Multi-Arm Multi-Stage Design</i>
------------------	--

Description

Obtains the p-value, conservative point estimate, and confidence interval after the end of an adaptive multi-arm multi-stage trial.

Usage

```

getADCI_multiarm(
  M = NA_integer_,
  r = 1,
  corr_known = TRUE,
  L = NA_integer_,
  zL = NA_real_,
  IMax = NA_real_,
  kMax = NA_integer_,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.25,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  spendingTime = NA_real_,
  MullerSchafer = FALSE,
  MNew = NA_integer_,
  selected = NA_integer_,
  rNew = 1,
  Lc = NA_integer_,
  zLc = NA_real_,
  INew = NA_real_,
  informationRatesNew = NA_real_,
  efficacyStoppingNew = NA_integer_,
  typeAlphaSpendingNew = "sfOF",
  parameterAlphaSpendingNew = NA_real_,
  spendingTimeNew = NA_real_,
  nthreads = 0
)

```

Arguments

M	Number of active treatment arms in the primary trial.
r	Randomization ratio of each active arm to the common control in the primary trial.
corr_known	Logical. If TRUE, the correlation between Wald statistics is derived from the randomization ratio r as $r/(r + 1)$. If FALSE, a conservative correlation of 0 is assumed.
L	The interim adaptation look of the primary trial.
zL	The z-test statistics at the interim adaptation look of the primary trial.
IMax	Maximum information for any active arm versus the common control for the primary trial. Must be provided.
kMax	The maximum number of stages of the primary trial.
informationRates	The information rates of the primary trial.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage of the primary trial. Defaults to TRUE if left unspecified.
criticalValues	The matrix of by-level upper boundaries on the max z-test statistic scale for efficacy stopping up to look L for the primary trial. The first column is for level M, the second column is for level M - 1, and so on, with the last column for level 1. If left unspecified, the critical values will be computed based on the specified alpha spending function.
alpha	The significance level of the primary trial. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending for the primary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value of alpha spending for the primary trial. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
spendingTime	The error spending time of the primary trial. Defaults to missing, in which case, it is the same as informationRates.
MullerSchafer	Whether to use the Muller and Schafer (2001) method for trial adaptation.
MNew	The number of active treatment arms in the secondary trial.
selected	The indices of the selected treatment arms for the secondary trial among the M active arms in the primary trial.
rNew	The randomization ratio of each active arm to the common control in the secondary trial.
Lc	The termination look of the integrated trial.
zLc	The z-test statistics at the termination look of the integrated trial.

I _{New}	The maximum information for any active arm versus the common control in the secondary trial.
informationRates _{New}	The spacing of looks of the secondary trial.
efficacyStopping _{New}	The indicators of whether efficacy stopping is allowed at each look of the secondary trial. Defaults to TRUE if left unspecified.
typeAlphaSpending _{New}	The type of alpha spending for the secondary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending _{New}	The parameter value of alpha spending for the secondary trial. Corresponds to ρ for "sfKD", and γ for "sfHSD".
spendingTime _{New}	The error spending time of the secondary trial. Defaults to missing, in which case, it is the same as informationRates _{New} .
nthreads	The number of threads to use (0 leaves the RcppParallel setting unchanged).

Details

If typeAlphaSpending_{New} is "OF" or "none", then informationRates_{New}, efficacyStopping_{New}, and spendingTime_{New} must be of full length k_{New}, and informationRates_{New} and spendingTime_{New} must end with 1.

Value

A data frame with the following variables:

- level: Number of elementary hypotheses considered for multiplicity.
- index: The treatment arm with max Z among the active arms.
- pvalue: p-value for rejecting the null hypothesis.
- thetahat: Point estimate of the parameter.
- cilevel: Confidence interval level.
- lower: Lower bound of confidence interval.
- upper: Upper bound of confidence interval.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Ping Gao, Yingqiu Li. Adaptive multiple comparison sequential design (AMCSD) for clinical trials. Journal of Biopharmaceutical Statistics, 2024, 34(3), 424-440.

Examples

```
getADCI_multiarm(
  M = 2, r = 1, corr_known = FALSE, L = 1, zL = c(2.075, 2.264),
  IMax = 300 / 4, kMax = 2, informationRates = c(0.5, 1),
  alpha = 0.025, typeAlphaSpending = "sfOF",
  MNew = 1, selected = 2, rNew = 1,
  Lc = 2, zLc = 1.667, INew = 374 / 4, nthreads = 1)
```

getADCI_seamless	<i>Confidence Interval After Adaptation for Phase 2/3 Seamless Design</i>
------------------	---

Description

Obtains the p-value, conservative point estimate, and confidence interval after the end of an adaptive phase 2/3 seamless design.

Usage

```
getADCI_seamless(
  M = NA_integer_,
  r = 1,
  corr_known = TRUE,
  L = NA_integer_,
  zL = NA_real_,
  IMax = NA_real_,
  K = NA_integer_,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  criticalValues = NA_real_,
  alpha = 0.25,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  spendingTime = NA_real_,
  MullerSchafer = FALSE,
  Lc = NA_integer_,
  zLc = NA_real_,
  INew = NA_real_,
  informationRatesNew = NA_real_,
  efficacyStoppingNew = NA_integer_,
  typeAlphaSpendingNew = "sfOF",
  parameterAlphaSpendingNew = NA_real_,
  spendingTimeNew = NA_real_,
  nthreads = 0
)
```

Arguments

M	Number of active treatment arms in Phase 2.
r	Randomization ratio of each active arm to the common control in Phase 2.
corr_known	Logical. If TRUE, the correlation between Wald statistics in Phase 2 is derived from the randomization ratio r as $r/(r+1)$. If FALSE, a conservative correlation of 0 is used. This option is only used for critical value calculations.
L	The interim adaptation look in Phase 3.
zL	The z-test statistic at the interim adaptation look of Phase 3.
IMax	Maximum information for the active arm versus the common control for the original trial. Must be provided.
K	Number of sequential looks in Phase 3.
informationRates	A numeric vector of information rates fixed before the trial. If unspecified, defaults to $(1 : (K + 1))/(K + 1)$.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage of the primary trial. Defaults to TRUE if left unspecified.
criticalValues	The upper boundaries on the z-test statistic scale for the most promising arm in Phase 2 and the z-test statistics for the selected arm in Phase 3 for the primary trial. If missing, boundaries will be computed based on the specified alpha spending function.
alpha	The significance level of the primary trial. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending for the primary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsiatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value of alpha spending for the primary trial. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
spendingTime	The error spending time of the primary trial. Defaults to missing, in which case, it is the same as informationRates.
MullerSchafer	Whether to use the Muller and Schafer (2001) method for trial adaptation.
Lc	The termination look of the integrated trial.
zLc	The z-test statistic at the termination look of the integrated trial.
INew	The maximum information for the active arm versus the common control in the secondary trial.
informationRatesNew	The spacing of looks of the secondary trial.
efficacyStoppingNew	The indicators of whether efficacy stopping is allowed at each look of the secondary trial. Defaults to TRUE if left unspecified.

typeAlphaSpendingNew	The type of alpha spending for the secondary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpendingNew	The parameter value of alpha spending for the secondary trial. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
spendingTimeNew	The error spending time of the secondary trial. Defaults to missing, in which case, it is the same as informationRatesNew.
nthreads	The number of threads to use (0 leaves the RcppParallel setting unchanged).

Details

If typeAlphaSpendingNew is "OF", "P", "WT", or "none", then informationRatesNew, efficacyStoppingNew, and spendingTimeNew must be of full length kNew, and informationRatesNew and spendingTimeNew must end with 1.

Value

A data frame with the following variables:

- pvalue: p-value for rejecting the null hypothesis.
- thetahat: Point estimate of the parameter.
- cilevel: Confidence interval level.
- lower: Lower bound of confidence interval.
- upper: Upper bound of confidence interval.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

References

Ping Gao, Yingqiu Li. Adaptive multiple comparison sequential design (AMCSD) for clinical trials. Journal of Biopharmaceutical Statistics, 2024, 34(3), 424-440.

Examples

```
getADCI_seamless(
  M = 2, r = 1, corr_known = FALSE,
  L = 1, zL = -log(0.67) * sqrt(80 / 4),
  IMax = 120 / 4, K = 2, informationRates = c(1/3, 2/3, 1),
  alpha = 0.025, typeAlphaSpending = "OF",
  Lc = 2, zLc = -log(0.677) * sqrt(236 / 4), INew = 236 / 4,
  nthreads = 1)
```

getADRCI

*Repeated Confidence Interval After Adaptation***Description**

Obtains the repeated p-value, conservative point estimate, and repeated confidence interval for an adaptive group sequential trial.

Usage

```
getADRCI(
  L = NA_integer_,
  zL = NA_real_,
  IMax = NA_real_,
  kMax = NA_integer_,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  criticalValues = NA_real_,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  spendingTime = NA_real_,
  MullerSchafer = FALSE,
  Lc = NA_integer_,
  zLc = NA_real_,
  INew = NA_real_,
  informationRatesNew = NA_real_,
  efficacyStoppingNew = NA_integer_,
  typeAlphaSpendingNew = "sfOF",
  parameterAlphaSpendingNew = NA_real_,
  spendingTimeNew = NA_real_
)
```

Arguments

L	The interim adaptation look of the primary trial.
zL	The z-test statistic at the interim adaptation look of the primary trial.
IMax	The maximum information of the primary trial.
kMax	The maximum number of stages of the primary trial.
informationRates	The information rates of the primary trial.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage of the primary trial. Defaults to true if left unspecified.
criticalValues	The upper boundaries on the z-test statistic scale for efficacy stopping for the primary trial.

alpha	The significance level of the primary trial. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending for the primary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value of alpha spending for the primary trial. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
spendingTime	The error spending time of the primary trial. Defaults to missing, in which case, it is the same as informationRates.
MullerSchafer	Whether to use the Muller and Schafer (2001) method for trial adaptation.
Lc	The look of interest in the integrated trial.
zLc	The z-test statistic at the look of the integrated trial.
INew	The maximum information of the secondary trial.
informationRatesNew	The spacing of looks of the secondary trial.
efficacyStoppingNew	The indicators of whether efficacy stopping is allowed at each look of the secondary trial up to look L2. Defaults to true if left unspecified.
typeAlphaSpendingNew	The type of alpha spending for the secondary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpendingNew	The parameter value of alpha spending for the secondary trial. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
spendingTimeNew	The error spending time of the secondary trial. up to look L2. Defaults to missing, in which case, it is the same as informationRatesNew.

Details

If typeAlphaSpendingNew is "OF", "P", "WT", or "none", then informationRatesNew, efficacyStoppingNew, and spendingTimeNew must be of full length kNew, and informationRatesNew and spendingTimeNew must end with 1.

Value

A data frame with the following variables:

- pvalue: Repeated p-value for rejecting the null hypothesis.

- `thetahat`: Point estimate of the parameter.
- `cilevel`: Confidence interval level.
- `lower`: Lower bound of repeated confidence interval.
- `upper`: Upper bound of repeated confidence interval.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Cyrus R. Mehta, Peter Bauer, Martin Posch and Werner Brannath. Repeated confidence intervals for adaptive group sequential trials. *Stat Med.* 2007;26:5422–5433.

See Also

[adaptDesign](#)

Examples

```
# two-arm randomized clinical trial with a normally distributed endpoint
# 90% power to detect mean difference of 15 with a standard deviation of 50
# Design the Stage I Trial with 3 looks and Lan-DeMets O'Brien-Fleming type
# spending function
delta <- 15
sigma <- 50

(des1 <- getDesignMeanDiff(
  beta = 0.1, meanDiff = delta, stDev = sigma,
  kMax = 3, alpha = 0.025, typeAlphaSpending = "sfOF"
))

s1 <- des1$byStageResults$informationRates
b1 <- des1$byStageResults$efficacyBounds
n <- des1$overallResults$numberOfSubjects

# Monitoring the Stage I Trial
L <- 1
nL <- des1$byStageResults$numberOfSubjects[L]
deltahat <- 8
sigmahat <- 55
sedeltahat <- sigmahat * sqrt( 4 / nL)
zL <- deltahat / sedeltahat

# Making an Adaptive Change: Stage I to Stage II
# revised clinically meaningful difference downward to 10 power the study
# retain the standard deviation at the design stage
# Muller & Schafer (2001) method to design the secondary trial
# with 2 looks and Lan-DeMets Pocock type spending function
# re-estimate sample size to reach 90% conditional power
deltaNew <- 10
```

```

(des2 <- adaptDesign(
  betaNew = 0.1, L = L, zL = zL, theta = deltaNew,
  IMax = n / (4 * sigma^2), kMax = 3, informationRates = s1,
  alpha = 0.025, typeAlphaSpending = "sf0F",
  MullerSchafer = TRUE, kNew = 2, typeAlphaSpendingNew = "sfP"
))

INew <- des2$secondaryTrial$maxInformation
(nNew <- ceiling(INew * 4 * sigma^2))
(nTotal <- nL + nNew)

# Monitoring the Integrated Trial
s2 <- des2$secondaryTrial$informationRates

Lc <- 2
deltahatc <- 9.5
sigmahatc <- 52.759
L2 <- Lc - L
nL2 <- nNew * s2[L2]
nc <- nL + nL2
sedeltahatc <- sigmahatc * sqrt(4 / nc)
zLc <- deltahatc / sedeltahatc
zL2 <- (zLc * sqrt(nc) - zL * sqrt(nL)) / sqrt(nL2)

getADRCI(
  L = L, zL = zL, IMax = n / (4 * sigmahatc^2), kMax = 3,
  informationRates = s1, alpha = 0.025, typeAlphaSpending = "sf0F",
  MullerSchafer = TRUE, Lc = Lc, zLc = zLc,
  INew = nNew / (4 * sigmahatc^2), informationRatesNew = s2,
  typeAlphaSpendingNew = "sfP")

```

getBound

Efficacy Boundaries for Group Sequential Design

Description

Obtains the efficacy stopping boundaries for a group sequential design.

Usage

```

getBound(
  k = NA_integer_,
  informationRates = NA_real_,
  alpha = 0.025,
  typeAlphaSpending = "sf0F",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  spendingTime = NA_real_,

```

```
    efficacyStopping = NA_integer_
  )
```

Arguments

k Look number for the current analysis.

informationRates Information rates up to the current look. Must be increasing and less than or equal to 1.

alpha The significance level. Defaults to 0.025.

typeAlphaSpending The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsiatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".

parameterAlphaSpending The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".

userAlphaSpending The user defined alpha spending. Cumulative alpha spent up to each stage.

spendingTime A vector of length k for the error spending time at each analysis. Must be increasing and less than or equal to 1. Defaults to missing, in which case, it is the same as informationRates.

efficacyStopping Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.

Details

If typeAlphaSpending is "OF", "P", "WT", or "none", then informationRates, efficacyStopping, and spendingTime must be of full length kMax, and informationRates and spendingTime must end with 1.

Value

A numeric vector of critical values up to the current look.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
getBound(k = 2, informationRates = c(0.5,1),
        alpha = 0.025, typeAlphaSpending = "sfOF")
```

Description

Calculates the efficacy stopping boundaries for a multi-arm multi-stage design.

Usage

```
getBound_multiarm(
  M = NA_integer_,
  r = 1,
  corr_known = TRUE,
  k = NA_integer_,
  informationRates = NA_real_,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  spendingTime = NA_real_,
  efficacyStopping = NA_integer_,
  nthreads = 0
)
```

Arguments

M	Number of active treatment arms.
r	Randomization ratio of each active arm to the common control.
corr_known	Logical. If TRUE, the correlation between Wald statistics is derived from the randomization ratio r as $r/(r + 1)$. If FALSE, a conservative correlation of 0 is assumed.
k	The index of the current look.
informationRates	A numeric vector of information rates up to the current look. Values must be strictly increasing and ≤ 1 .
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".

<code>userAlphaSpending</code>	The user defined alpha spending. Cumulative alpha spent up to each stage.
<code>spendingTime</code>	A numeric vector of length k specifying the error spending time at each analysis. Values must be strictly increasing and ≤ 1 . If omitted, defaults to <code>informationRates</code> .
<code>efficacyStopping</code>	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
<code>nthreads</code>	The number of threads to use (0 leaves the <code>RcppParallel</code> setting unchanged).

Details

The function determines critical values by solving for the boundary that satisfies the alpha-spending requirement.

If `typeAlphaSpending` is "OF", "P", "WT", or "none", then `informationRates`, `efficacyStopping`, and `spendingTime` must be of full length `kMax`, and `informationRates` and `spendingTime` must end with 1.

Value

A numeric vector of length k containing the critical values (on the standard normal Z-scale) for each analysis up to the current look.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Ping Gao, Yingqiu Li. Adaptive multiple comparison sequential design (AMCSD) for clinical trials. *Journal of Biopharmaceutical Statistics*, 2024, 34(3), 424-440.

Examples

```
# Determine O'Brien-Fleming boundaries for a TSSSD with
# 2 active arms and 3 looks.
getBound_multiarm(M = 2, k = 3, informationRates = seq(1, 3)/3,
                  alpha = 0.025, typeAlphaSpending = "OF",
                  nthreads = 1)
```

getBound_seamless	<i>Efficacy Boundaries for Phase 2/3 Seamless Design</i>
-------------------	--

Description

Calculates the efficacy stopping boundaries for a phase 2/3 seamless design, accounting for selection of the most promising treatment at the end of Phase 2 and sequential testing in Phase 3.

Usage

```
getBound_seamless(
  M = NA_integer_,
  r = 1,
  corr_known = TRUE,
  k = NA_integer_,
  informationRates = NA_real_,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  spendingTime = NA_real_,
  efficacyStopping = NA_integer_,
  nthreads = 0
)
```

Arguments

M	Number of active treatment arms in Phase 2.
r	Randomization ratio of each active arm to the common control in Phase 2.
corr_known	Logical. If TRUE, the correlation between Wald statistics in Phase 2 is derived from the randomization ratio r as $r/(r+1)$. If FALSE, a conservative correlation of 0 is used.
k	The index of the current look in Phase 3.
informationRates	A numeric vector of information rates up to the current look. Values must be strictly increasing and ≤ 1 .
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".

<code>parameterAlphaSpending</code>	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
<code>userAlphaSpending</code>	The user defined alpha spending. Cumulative alpha spent up to each stage.
<code>spendingTime</code>	A numeric vector of length $k + 1$ specifying the error spending time at each analysis. Values must be strictly increasing and ≤ 1 . If omitted, defaults to <code>informationRates</code> .
<code>efficacyStopping</code>	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
<code>nthreads</code>	The number of threads to use (0 leaves the <code>RcppParallel</code> setting unchanged).

Details

The function determines critical values by solving for the boundary that satisfies the alpha-spending requirement, given selection of the arm with the largest Z-statistic at the end of Phase 2.

If `typeAlphaSpending` is "OF", "P", "WT", or "none", then `informationRates`, `efficacyStopping`, and `spendingTime` must be of full length `kMax`, and `informationRates` and `spendingTime` must end with 1.

Value

A numeric vector of length $k + 1$ containing the critical values (on the standard normal Z-scale) for each analysis up to the current look.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Ping Gao, Yingqiu Li. Adaptive two-stage seamless sequential design for clinical trials. *Journal of Biopharmaceutical Statistics*, 2025, 35(4), 565-587.

Examples

```
# Determine O'Brien-Fleming boundaries for a seamless design with
# 2 active arms in Phase 2 and 2 looks in Phase 3 (3 looks total).
getBound_seamless(M = 2, k = 2, informationRates = seq(1, 3)/3,
                  alpha = 0.025, typeAlphaSpending = "OF",
                  nthreads = 1)
```

getCI

*Confidence Interval After Trial Termination***Description**

Obtains the p-value, median unbiased point estimate, and confidence interval after the end of a group sequential trial.

Usage

```
getCI(
  L = NA_integer_,
  zL = NA_real_,
  IMax = NA_real_,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  criticalValues = NA_real_,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  spendingTime = NA_real_
)
```

Arguments

L	The termination look.
zL	The z-test statistic at the termination look.
IMax	The maximum information of the trial.
informationRates	The information rates up to look L.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage up to look L. Defaults to true if left unspecified.
criticalValues	The upper boundaries on the z-test statistic scale for efficacy stopping up to look L.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending for the trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCan spending function, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".

spendingTime The error spending time up to look L. Defaults to missing, in which case, it is the same as informationRates.

Details

If typeAlphaSpending is "OF", "P", "WT", or "none", then informationRates, efficacyStopping, and spendingTime must be of full length kMax, and informationRates and spendingTime must end with 1.

Value

A data frame with the following components:

- pvalue: p-value for rejecting the null hypothesis.
- thetahat: Median unbiased point estimate of the parameter.
- cilevel: Confidence interval level.
- lower: Lower bound of confidence interval.
- upper: Upper bound of confidence interval.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

References

Anastasios A. Tsiatis, Gary L. Rosner and Cyrus R. Mehta. Exact confidence intervals following a group sequential test. Biometrics 1984;40:797-803.

Examples

```
# group sequential design with 90% power to detect delta = 6
delta <- 6
sigma <- 17
n <- 282
(des1 <- getDesign(IMax = n/(4*sigma^2), theta = delta, kMax = 3,
  alpha = 0.05, typeAlphaSpending = "sfHSD",
  parameterAlphaSpending = -4))

# crossed the boundary at the second look
L <- 2
n1 <- n*2/3
delta1 <- 7
sigma1 <- 20
zL <- delta1/sqrt(4/n1*sigma1^2)

# confidence interval
getCI(L = L, zL = zL, IMax = n/(4*sigma1^2),
  informationRates = c(1/3, 2/3), alpha = 0.05,
  typeAlphaSpending = "sfHSD", parameterAlphaSpending = -4)
```

getCI_multiarm	<i>Confidence Interval After Trial Termination for Multi-Arm Multi-Stage Design</i>
----------------	---

Description

Obtains the p-value, conservative point estimate, and confidence interval after the end of a multi-arm multi-stage trial.

Usage

```
getCI_multiarm(
  M = NA_integer_,
  r = 1,
  corr_known = TRUE,
  L = NA_integer_,
  zL = NA_real_,
  IMax = NA_real_,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  spendingTime = NA_real_,
  nthreads = 0
)
```

Arguments

M	Number of active treatment arms.
r	Randomization ratio of each active arm to the common control.
corr_known	Logical. If TRUE, the correlation between Wald statistics is derived from the randomization ratio r as $r/(r + 1)$. If FALSE, a conservative correlation of 0 is used.
L	The termination look.
zL	The vector of z-test statistics at the termination look.
IMax	Maximum information for any active arm versus the common control.
informationRates	The information rates up to look L.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage up to look L. Defaults to TRUE if left unspecified.

criticalValues	The matrix of by-level upper boundaries on the max z-test statistic scale for efficacy stopping up to look L. The first column is for level M, the second column is for level M – 1, and so on, with the last column for level 1. If left unspecified, the critical values will be computed based on the specified alpha spending function.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending for the trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
spendingTime	The error spending time up to look L. Defaults to missing, in which case, it is the same as informationRates.
nthreads	The number of threads to use (0 leaves the RcppParallel setting unchanged).

Details

If typeAlphaSpending is "OF", "P", "WT", or "none", then informationRates, efficacyStopping, and spendingTime must be of full length kMax, and informationRates and spendingTime must end with 1.

Value

A data frame with the following components:

- level: Number of elementary hypotheses considered for multiplicity.
- index: The treatment arm with max Z among the active arms.
- pvalue: p-value for rejecting the null hypothesis.
- thetahat: Point estimate of the parameter.
- cilevel: Confidence interval level.
- lower: Lower bound of confidence interval.
- upper: Upper bound of confidence interval.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Ping Gao, Yingqiu Li. Adaptive multiple comparison sequential design (AMCSD) for clinical trials. Journal of Biopharmaceutical Statistics, 2024, 34(3), 424-440.

Examples

```
getCI_multiarm(
  L = 2, zL = c(2.075, 2.264),
  M = 2, r = 1, corr_known = FALSE,
  IMax = 300 / 4, informationRates = c(1/2, 1),
  alpha = 0.025, typeAlphaSpending = "sf0F",
  nthreads = 1)
```

getCI_seamless	<i>Confidence Interval After Trial Termination for Phase 2/3 Seamless Design</i>
----------------	--

Description

Obtains the p-value, point estimate, and confidence interval after the end of a phase 2/3 seamless trial.

Usage

```
getCI_seamless(
  M = NA_integer_,
  r = 1,
  corr_known = TRUE,
  L = NA_integer_,
  zL = NA_real_,
  IMax = NA_real_,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  criticalValues = NA_real_,
  alpha = 0.025,
  typeAlphaSpending = "sf0F",
  parameterAlphaSpending = NA_real_,
  spendingTime = NA_real_,
  nthreads = 0
)
```

Arguments

M	Number of active treatment arms in Phase 2.
r	Randomization ratio of each active arm to the common control in Phase 2.
corr_known	Logical. If TRUE, the correlation between Wald statistics in Phase 2 is derived from the randomization ratio r as $r/(r+1)$. If FALSE, a conservative correlation of 0 is used. This option is only used for critical value calculations.
L	The termination look in Phase 3.
zL	The z-test statistic at the termination look.

IMax	Maximum information for any active arm versus the common control.
informationRates	The information rates up to look L.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage up to look L. Defaults to TRUE if left unspecified.
criticalValues	The upper boundaries on the z-test statistic scale for the most promising arm in Phase 2 and the z-test statistics for the selected arm in Phase 3 up to look L. If missing, boundaries will be computed based on the specified alpha spending function.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending for the trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
spendingTime	The error spending time up to look L. Defaults to missing, in which case, it is the same as informationRates.
nthreads	The number of threads to use (0 leaves the RcppParallel setting unchanged).

Details

If typeAlphaSpending is "OF", "P", "WT", or "none", then informationRates, efficacyStopping, and spendingTime must be of full length $K + 1$, and informationRates and spendingTime must end with 1.

Value

A data frame with the following components:

- pvalue: p-value for rejecting the null hypothesis.
- thetahat: Point estimate of the parameter.
- cilevel: Confidence interval level.
- lower: Lower bound of confidence interval.
- upper: Upper bound of confidence interval.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

References

Ping Gao, Yingqiu Li. Adaptive two-stage seamless sequential design for clinical trials. Journal of Biopharmaceutical Statistics, 2025, 35(4), 565-587.

Examples

```
getCI_seamless(
  L = 2, zL = 2.075, M = 2, r = 1, corr_known = FALSE,
  IMax = 300 / 4, informationRates = c(1/3, 2/3, 1),
  alpha = 0.025, typeAlphaSpending = "sfOF", nthreads = 1)
```

getCP

Conditional Power for Generic Group Sequential Design

Description

Obtains the conditional power for specified incremental information given the interim results, parameter values, and data-dependent changes in the error spending function, as well as the number and spacing of interim looks.

Usage

```
getCP(
  INew = NA_real_,
  L = NA_integer_,
  zL = NA_real_,
  theta = NA_real_,
  IMax = NA_real_,
  kMax = NA_integer_,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,
  futilityCP = NULL,
  futilityTheta = NULL,
  spendingTime = NA_real_,
  MullerSchafer = FALSE,
  kNew = NA_integer_,
  informationRatesNew = NA_real_,
  efficacyStoppingNew = NA_integer_,
  futilityStoppingNew = NA_integer_,
```

```

typeAlphaSpendingNew = "sfOF",
parameterAlphaSpendingNew = NA_real_,
futilityBoundsInt = NULL,
futilityCPInt = NULL,
futilityThetaInt = NULL,
typeBetaSpendingNew = "none",
parameterBetaSpendingNew = NA_real_,
spendingTimeNew = NA_real_,
varianceRatio = 1
)

```

Arguments

<code>INew</code>	The maximum information of the secondary trial.
<code>L</code>	The interim adaptation look of the primary trial.
<code>zL</code>	The z-test statistic at the interim adaptation look of the primary trial.
<code>theta</code>	A scalar or a vector of parameter values of length $k_{\text{Max}} + k_{\text{Max}} - L$ if <code>MullerSchafer</code> = FALSE or length $k_{\text{Max}} + k_{\text{New}}$ if <code>MullerSchafer</code> = TRUE.
<code>IMax</code>	The maximum information of the primary trial.
<code>kMax</code>	The maximum number of stages of the primary trial.
<code>informationRates</code>	The information rates of the primary trial.
<code>efficacyStopping</code>	Indicators of whether efficacy stopping is allowed at each stage of the primary trial. Defaults to true if left unspecified.
<code>futilityStopping</code>	Indicators of whether futility stopping is allowed at each stage of the primary trial. Defaults to true if left unspecified.
<code>criticalValues</code>	The upper boundaries on the z-test statistic scale for efficacy stopping for the primary trial.
<code>alpha</code>	The significance level of the primary trial. Defaults to 0.025.
<code>typeAlphaSpending</code>	The type of alpha spending for the primary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsiatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
<code>parameterAlphaSpending</code>	The parameter value of alpha spending for the primary trial. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
<code>userAlphaSpending</code>	The user defined alpha spending for the primary trial. Cumulative alpha spent up to each stage.
<code>futilityBounds</code>	The lower boundaries on the z-test statistic scale for futility stopping for the primary trial. Defaults to <code>rep(-8, kMax-1)</code> if left unspecified.

futilityCP	The conditional power-based futility bounds for the primary trial.
futilityTheta	The parameter value-based futility bounds for the primary trial.
spendingTime	The error spending time of the primary trial. Defaults to missing, in which case, it is the same as informationRates.
MullerSchafer	Whether to use the Muller and Schafer (2001) method for trial adaptation.
kNew	The number of looks of the secondary trial.
informationRatesNew	The spacing of looks of the secondary trial.
efficacyStoppingNew	The indicators of whether efficacy stopping is allowed at each look of the secondary trial. Defaults to true if left unspecified.
futilityStoppingNew	The indicators of whether futility stopping is allowed at each look of the secondary trial. Defaults to true if left unspecified.
typeAlphaSpendingNew	The type of alpha spending for the secondary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpendingNew	The parameter value of alpha spending for the secondary trial. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
futilityBoundsInt	The futility boundaries on the z statistic scale for new stages of the integrated trial.
futilityCPInt	The conditional power-based futility bounds for new stages of the integrated trial.
futilityThetaInt	The parameter value-based futility bounds for the new stages of the integrated trial.
typeBetaSpendingNew	The type of beta spending for the secondary trial. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpendingNew	The parameter value of beta spending for the secondary trial. Corresponds to ρ for "sfKD", and γ for "sfHSD".
spendingTimeNew	The error spending time of the secondary trial. Defaults to missing, in which case, it is the same as informationRatesNew.
varianceRatio	The ratio of the variance under H0 to the variance under H1.

Value

A vector of two conditional powers given the interim results and parameter values, one without design change and the other with data-dependent design changes.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

References

Cyrus R. Mehta and Stuart J. Pocock. Adaptive increase in sample size when interim results are promising: A practical guide with examples. *Stat Med.* 2011;30:3267–3284.

See Also

[adaptDesign](#)

Examples

```
# Conditional power calculation with delayed treatment effect

# Two interim analyses have occurred with 179 and 266 events,
# respectively. The observed hazard ratio at the second interim
# look is 0.81.

trialsdt <- as.Date("2020-03-04")           # trial start date
iadt <- c(as.Date("2022-02-01"), as.Date("2022-11-01")) # interim dates
mo1 <- as.numeric(iadt - trialsdt + 1)/30.4375 # interim months

# Assume a piecewise Poisson enrollment process with a 8-month ramp-up
# and 521 patients were enrolled after 17.94 months
N <- 521                                # total number of patients
Ta <- 17.94                             # enrollment duration
Ta1 <- 8                                # assumed end of enrollment ramp-up
enrate <- N / (Ta - Ta1/2) # enrollment rate after ramp-up

# Assume a median survival of 16.7 months for the control group, a
# 5-month delay in treatment effect, and a hazard ratio of 0.7 after
# the delay
lam1 <- log(2)/16.7 # control group hazard of exponential distribution
t1 <- 5             # months of delay in treatment effect
hr <- 0.7           # hazard ratio after delay
lam2 <- hr*lam1     # treatment group hazard after delay

# Assume an annual dropout rate of 5%
gam <- -log(1-0.05)/12 # hazard for dropout

# The original target number of events was 298 and the new target is 335
mo2 <- caltime(
  nevents = c(298, 335),
  allocationRatioPlanned = 1,
  accrualTime = seq(0, Ta1),
```

```

    accrualIntensity = enrate*seq(1, Ta1+1)/(Ta1+1),
    piecewiseSurvivalTime = c(0, t1),
    lambda1 = c(lam1, lam2),
    lambda2 = c(lam1, lam1),
    gamma1 = gam,
    gamma2 = gam,
    accrualDuration = Ta,
    followupTime = 1000)

# expected number of events and average hazard ratios
(lr1 <- lrstat(
  time = c(mo1, mo2),
  accrualTime = seq(0, Ta1),
  accrualIntensity = enrate*seq(1, Ta1+1)/(Ta1+1),
  piecewiseSurvivalTime = c(0, t1),
  lambda1 = c(lam1, lam2),
  lambda2 = c(lam1, lam1),
  gamma1 = gam,
  gamma2 = gam,
  accrualDuration = Ta,
  followupTime = 1000,
  predictTarget = 3))

hr2 <- 0.81 # observed hazard ratio at interim 2
z2 <- (-log(hr2))*sqrt(266/4) # corresponding z-test statistic value

# expected mean of -log(HR) at the original looks and the new final look
theta <- -log(lr1$HR[c(1,2,3,4)])

# conditional power with sample size increase
getCP(INew = (335 - 266)/4,
      L = 2, zL = z2, theta = theta,
      IMax = 298/4, kMax = 3,
      informationRates = c(179, 266, 298)/298,
      alpha = 0.025, typeAlphaSpending = "sfOF")

```

getCP_multiarm

Conditional Power for Multi-Arm Multi-Stage Design

Description

Obtains the conditional power for specified incremental information given the interim results, parameter values, and data-dependent changes in the selected treatment(s), the error spending function, as well as the number and spacing of interim looks.

Usage

```
getCP_multiarm(
```

```

INew = NA_real_,
M = NA_integer_,
r = 1,
corr_known = TRUE,
L = NA_integer_,
zL = NA_real_,
theta = NA_real_,
IMax = NA_real_,
kMax = NA_integer_,
informationRates = NA_real_,
efficacyStopping = NA_integer_,
futilityStopping = NA_integer_,
criticalValues = NULL,
alpha = 0.025,
typeAlphaSpending = "sfOF",
parameterAlphaSpending = NA_real_,
userAlphaSpending = NA_real_,
futilityBounds = NULL,
futilityCP = NULL,
futilityTheta = NULL,
spendingTime = NA_real_,
MullerSchafer = FALSE,
MNew = NA_integer_,
selected = NA_integer_,
rNew = 1,
kNew = NA_integer_,
informationRatesNew = NA_real_,
efficacyStoppingNew = NA_integer_,
futilityStoppingNew = NA_integer_,
typeAlphaSpendingNew = "sfOF",
parameterAlphaSpendingNew = NA_real_,
futilityBoundsInt = NULL,
futilityCPInt = NULL,
futilityThetaInt = NULL,
typeBetaSpendingNew = "none",
parameterBetaSpendingNew = NA_real_,
spendingTimeNew = NA_real_,
nthreads = 0
)

```

Arguments

INew	The maximum information for any active arm versus the common control in the secondary trial.
M	Number of active treatment arms in the primary trial.
r	Randomization ratio of each active arm to the common control in the primary trial.
corr_known	Logical. If TRUE, the correlation between Wald statistics is derived from the

	randomization ratio r as $r/(r+1)$. If FALSE, a conservative correlation of 0 is assumed.
L	The interim adaptation look of the primary trial.
zL	The z-test statistics at the interim adaptation look of the primary trial.
theta	A vector of length M representing the assumed treatment effects for each active arm versus the common control. The global null is $\theta_i = 0$ for all i , and alternatives are one-sided: $\theta_i > 0$ for at least one $i = 1, \dots, M$.
IMax	Maximum information for any active arm versus the common control for the primary trial. Must be provided.
kMax	The maximum number of stages of the primary trial.
informationRates	The information rates of the primary trial.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage of the primary trial. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage of the primary trial.
criticalValues	The upper boundaries on the max z-test statistic scale for efficacy stopping for the primary trial. If missing, boundaries will be computed based on the specified alpha spending function.
alpha	The significance level of the primary trial. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending for the primary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsiatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value of alpha spending for the primary trial. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user-defined alpha spending for the primary trial. Represents the cumulative alpha spent up to each stage.
futilityBounds	The futility boundaries on the max-z statistic scale for the primary trial. Defaults to $\text{rep}(-8, \text{kMax}-1)$ if left unspecified.
futilityCP	The conditional power-based futility bounds for the primary trial.
futilityTheta	The parameter value-based futility bounds for the primary trial.
spendingTime	The error spending time of the primary trial. Defaults to missing, in which case it is assumed to be the same as informationRates.
MullerSchafer	Whether to use the Muller and Schafer (2001) method for trial adaptation.
MNew	Number of active treatment arms in the secondary trial.

selected	The indices of the selected active treatment arms for the secondary trial.
rNew	Randomization ratio of each active arm to the common control in the secondary trial.
kNew	The number of looks of the secondary trial.
informationRatesNew	The spacing of looks of the secondary trial.
efficacyStoppingNew	The indicators of whether efficacy stopping is allowed at each look of the secondary trial. Defaults to TRUE if left unspecified.
futilityStoppingNew	The indicators of whether futility stopping is allowed at each look of the secondary trial. Defaults to TRUE if left unspecified.
typeAlphaSpendingNew	The type of alpha spending for the secondary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpendingNew	The parameter value of alpha spending for the secondary trial. Corresponds to ρ for "sfKD", and γ for "sfHSD".
futilityBoundsInt	The futility boundaries on the max-z statistic scale for new stages of the integrated trial.
futilityCPInt	The conditional power-based futility bounds for new stages of the integrated trial.
futilityThetaInt	The parameter value-based futility bounds for the new stages of the integrated trial.
typeBetaSpendingNew	The type of beta spending for the secondary trial. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "none" for no early futility stopping. Defaults to "none".
parameterBetaSpendingNew	The parameter value of beta spending for the secondary trial. Corresponds to ρ for "sfKD", and γ for "sfHSD".
spendingTimeNew	The error spending time of the secondary trial. Defaults to missing, in which case it is assumed to be the same as informationRatesNew.
nthreads	The number of threads to use (0 leaves the RcppParallel setting unchanged).

Value

A vector of two conditional powers given the interim results and parameter values, one without design change and the other with data-dependent design changes.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Ping Gao, Yingqiu Li. Adaptive multiple comparison sequential design (AMCSD) for clinical trials. Journal of Biopharmaceutical Statistics, 2024, 34(3), 424-440.

See Also

[adaptDesign_multiarm](#)

Examples

```
getCP_multiarm(  
  INew = 373 / 4, M = 2, r = 1, corr_known = FALSE,  
  L = 1, zL = c(-log(0.91), -log(0.78)) * sqrt(324 / 4 / 2),  
  theta = c(-log(0.91), -log(0.78)),  
  IMax = 324 / 4, kMax = 2, informationRates = c(1/2, 1),  
  alpha = 0.025, typeAlphaSpending = "OF",  
  MNew = 1, selected = 2, rNew = 1, nthreads = 1)
```

getCP_seamless	<i>Conditional Power for Phase 2/3 Seamless Design</i>
----------------	--

Description

Obtains the conditional power for specified incremental information given the interim results, parameter values, and data-dependent changes in the error spending function, as well as the number and spacing of interim looks.

Usage

```
getCP_seamless(  
  INew = NA_real_,  
  M = NA_integer_,  
  r = 1,  
  corr_known = TRUE,  
  L = NA_integer_,  
  zL = NA_real_,  
  theta = NA_real_,  
  IMax = NA_real_,  
  K = NA_integer_,  
  informationRates = NA_real_,  
  efficacyStopping = NA_integer_,  
  futilityStopping = NA_integer_,  
  criticalValues = NULL,
```

```

alpha = 0.025,
typeAlphaSpending = "sfOF",
parameterAlphaSpending = NA_real_,
userAlphaSpending = NA_real_,
futilityBounds = NULL,
futilityCP = NULL,
futilityTheta = NULL,
spendingTime = NA_real_,
MullerSchafer = FALSE,
kNew = NA_integer_,
informationRatesNew = NA_real_,
efficacyStoppingNew = NA_integer_,
futilityStoppingNew = NA_integer_,
typeAlphaSpendingNew = "sfOF",
parameterAlphaSpendingNew = NA_real_,
futilityBoundsInt = NULL,
futilityCPInt = NULL,
futilityThetaInt = NULL,
typeBetaSpendingNew = "none",
parameterBetaSpendingNew = NA_real_,
spendingTimeNew = NA_real_,
nthreads = 0
)

```

Arguments

I _{New}	The maximum information for the active arm versus the common control in the secondary trial.
M	Number of active treatment arms in Phase 2.
r	Randomization ratio of each active arm to the common control in Phase 2.
corr_known	Logical. If TRUE, the correlation between Wald statistics in Phase 2 is derived from the randomization ratio r as $r/(r+1)$. If FALSE, a conservative correlation of 0 is used. This option is only used for critical value calculations.
L	The interim adaptation look in Phase 3.
z _L	The z-test statistic at the interim adaptation look of Phase 3.
theta	The assumed treatment effect for the selected arm versus the common control.
I _{Max}	Maximum information for the active arm versus the common control for the original trial. Must be provided.
K	Number of sequential looks in Phase 3.
informationRates	A numeric vector of information rates fixed before the trial. If unspecified, defaults to $(1 : (K+1))/(K+1)$.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage of the primary trial. Defaults to TRUE if left unspecified.

futilityStopping	Indicators of whether futility stopping is allowed at each stage of the primary trial. Defaults to true if left unspecified.
criticalValues	The upper boundaries on the z-test statistic scale for the most promising arm in Phase 2 and the z-test statistics for the selected arm in Phase 3 for the primary trial. If missing, boundaries will be computed based on the specified alpha spending function.
alpha	The significance level of the primary trial. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending for the primary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value of alpha spending for the primary trial. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user-defined alpha spending for the primary trial. Represents the cumulative alpha spent up to each stage.
futilityBounds	The lower boundaries on the z-test statistic scale for the most promising arm in Phase 2 and the z-test statistics for the selected arm in Phase 3 for the primary trial.
futilityCP	The conditional power-based futility bounds for the primary trial.
futilityTheta	The parameter value-based futility bounds for the primary trial.
spendingTime	The error spending time of the primary trial. Defaults to missing, in which case it is assumed to be the same as informationRates.
MullerSchafer	Whether to use the Muller and Schafer (2001) method for trial adaptation.
kNew	The number of looks of the secondary trial.
informationRatesNew	The spacing of looks of the secondary trial.
efficacyStoppingNew	The indicators of whether efficacy stopping is allowed at each look of the secondary trial. Defaults to TRUE if left unspecified.
futilityStoppingNew	The indicators of whether futility stopping is allowed at each look of the secondary trial. Defaults to true if left unspecified.
typeAlphaSpendingNew	The type of alpha spending for the secondary trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early efficacy stopping. Defaults to "sfOF".

parameterAlphaSpendingNew	The parameter value of alpha spending for the secondary trial. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
futilityBoundsInt	The futility boundaries on the z statistic scale for new stages of the integrated trial.
futilityCPInt	The conditional power-based futility bounds for new stages of the integrated trial.
futilityThetaInt	The parameter value-based futility bounds for the new stages of the integrated trial.
typeBetaSpendingNew	The type of beta spending for the secondary trial. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpendingNew	The parameter value of beta spending for the secondary trial. Corresponds to ρ for "sfKD", and γ for "sfHSD".
spendingTimeNew	The error spending time of the secondary trial. Defaults to missing, in which case it is assumed to be the same as informationRatesNew.
nthreads	The number of threads to use (0 leaves the RcppParallel setting unchanged).

Value

A vector of two conditional powers given the interim results and parameter values, one without design change and the other with data-dependent design changes.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Ping Gao, Yingqiu Li. Adaptive two-stage seamless sequential design for clinical trials. Journal of Biopharmaceutical Statistics, 2025, 35(4), 565-587.

See Also

[adaptDesign_seamless](#)

Examples

```
getCP_seamless(
  INew = 198 / 4, M = 2, r = 1, corr_known = FALSE,
  L = 1, zL = -log(0.67) * sqrt(80 / 4), theta = -log(0.691),
  IMax = 120 / 4, K = 2, informationRates = c(1/3, 2/3, 1),
```

```
alpha = 0.025, typeAlphaSpending = "OF", kNew = 1,
nthreads = 1)
```

getDesign

Power and Sample Size for Generic Group Sequential Design

Description

Obtains the maximum information and stopping boundaries for a generic group sequential design assuming a constant treatment effect, or obtains the power given the maximum information and stopping boundaries.

Usage

```
getDesign(
  beta = NA_real_,
  IMax = NA_real_,
  theta = NA_real_,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,
  futilityCP = NULL,
  futilityTheta = NULL,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
  spendingTime = NA_real_,
  varianceRatio = 1
)
```

Arguments

beta	The type II error.
IMax	The maximum information. Either beta or IMax should be provided while the other one should be missing.
theta	The parameter value. Null hypothesis is at $\theta = 0$, and the alternative hypothesis is one-sided for $\theta > 0$.
kMax	The maximum number of stages.

informationRates	The information rates. Fixed prior to the trial. Defaults to $(1:kMax) / kMax$ if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., $kMax-1$. Defaults to $\text{rep}(-8, kMax-1)$ if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	The futility bounds on the conditional power scale.
futilityTheta	The futility bounds on the parameter scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length $kMax$ for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.
varianceRatio	The ratio of the variance under H_0 to the variance under H_1 .

Details

The function determines efficacy and futility bounds based on the inputs provided, following a clear priority order.

Efficacy bounds: If `criticalValues` are supplied, they take precedence and all alpha-spending parameters are ignored. Otherwise, efficacy bounds are derived from the specified alpha-spending function.

Futility bounds: Futility inputs are evaluated in the following order of priority:

1. If `futilityBounds` are provided, they override all other futility-related inputs (`futilityCP`, `futilityTheta`, and beta-spending parameters).
2. If `futilityBounds` are not provided but `futilityCP` is specified, then `futilityTheta` and beta-spending parameters are ignored.
3. If only `futilityTheta` is provided, beta-spending parameters are ignored.
4. If none of `futilityBounds`, `futilityCP`, or `futilityTheta` are specified, futility bounds are computed using the beta-spending approach.

Value

An S3 class design object with three components:

- `overallResults`: A data frame containing the following variables:
 - `overallReject`: The overall rejection probability.
 - `alpha`: The overall significance level.
 - `attainedAlpha`: The attained significance level, which is different from the overall significance level in the presence of futility stopping.
 - `kMax`: The number of stages.
 - `theta`: The parameter value.
 - `information`: The maximum information.
 - `expectedInformationH1`: The expected information under H1.
 - `expectedInformationH0`: The expected information under H0.
 - `drift`: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
 - `inflationFactor`: The inflation factor (relative to the fixed design).
- `byStageResults`: A data frame containing the following variables:
 - `informationRates`: The information rates.
 - `efficacyBounds`: The efficacy boundaries on the Z-scale.
 - `futilityBounds`: The futility boundaries on the Z-scale.
 - `rejectPerStage`: The probability for efficacy stopping.
 - `futilityPerStage`: The probability for futility stopping.
 - `cumulativeRejection`: The cumulative probability for efficacy stopping.
 - `cumulativeFutility`: The cumulative probability for futility stopping.
 - `cumulativeAlphaSpent`: The cumulative alpha spent.
 - `efficacyTheta`: The efficacy boundaries on the parameter scale.
 - `futilityTheta`: The futility boundaries on the parameter scale.

- efficacyP: The efficacy boundaries on the p-value scale.
- futilityP: The futility boundaries on the p-value scale.
- information: The cumulative information.
- efficacyStopping: Whether to allow efficacy stopping.
- futilityStopping: Whether to allow futility stopping.
- rejectPerStageH0: The probability for efficacy stopping under H0.
- futilityPerStageH0: The probability for futility stopping under H0.
- cumulativeRejectionH0: The cumulative probability for efficacy stopping under H0.
- cumulativeFutilityH0: The cumulative probability for futility stopping under H0.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - typeBetaSpending: The type of beta spending.
 - parameterBetaSpending: The parameter value for beta spending.
 - userBetaSpending: The user defined beta spending.
 - spendingTime: The error spending time at each analysis.
 - varianceRatio: The ratio of the variance under H0 to the variance under H1.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Christopher Jennison, Bruce W. Turnbull. Group Sequential Methods with Applications to Clinical Trials. Chapman & Hall/CRC: Boca Raton, 2000, ISBN:0849303168

Examples

```
# Example 1: obtain the maximum information given power
(design1 <- getDesign(
  beta = 0.2, theta = -log(0.7),
  kMax = 2, informationRates = c(0.5,1),
  alpha = 0.025, typeAlphaSpending = "sf0F",
  typeBetaSpending = "sfP"))

# Example 2: obtain power given the maximum information
(design2 <- getDesign(
  IMax = 72.5, theta = -log(0.7),
  kMax = 3, informationRates = c(0.5, 0.75, 1),
  alpha = 0.025, typeAlphaSpending = "sf0F",
  typeBetaSpending = "sfP"))
```

getDesign2	<i>Power and Sample Size for Group Sequential Design With Futility Stopping Under Null Hypothesis</i>
------------	---

Description

Obtains the maximum information and stopping boundaries for a generic group sequential design with futility stopping under the null hypothesis assuming a constant treatment effect, or obtains the power given the maximum information and stopping boundaries.

Usage

```
getDesign2(
  beta = NA_real_,
  IMax = NA_real_,
  theta = NA_real_,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sf0F",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  symmetricBounds = TRUE,
  astar = 0.025,
  futilityBounds = NULL,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
  spendingTime = NA_real_,
  varianceRatio = 1
)
```

Arguments

beta	The type II error.
IMax	The maximum information. Either beta or IMax should be provided while the other one should be missing.
theta	The parameter value. Null hypothesis is at $\theta = 0$, and the alternative hypothesis is one-sided for $\theta > 0$.
kMax	The maximum number of stages.
informationRates	The information rates. Fixed prior to the trial. Defaults to $(1:kMax) / kMax$ if left unspecified.

efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
symmetricBounds	If TRUE, futility bounds are set to the negative of efficacy bounds at each analysis (subject to futilityStopping). If FALSE, futility bounds are determined by futilityBounds or beta spending.
astar	The overall futility stopping probability under the null hypothesis.
futilityBounds	A vector of length kMax for the futility stopping boundaries on the Z-scale under the null hypothesis. Defaults to rep(-8, kMax) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
typeBetaSpending	The type of beta spending function for determining futility bounds under the null hypothesis when futilityBounds is not provided. The same types as typeAlphaSpending are allowed, except that "none" corresponds to no futility stopping under the null hypothesis.
parameterBetaSpending	The parameter for the beta spending function. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	A vector of length kMax for the user defined beta spending function when typeBetaSpending == "user". The last element must be equal to astar and the vector must be increasing.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.
varianceRatio	The ratio of the variance under H0 to the variance under H1.

Details

The futility stopping boundaries under the null hypothesis are non-binding. The function determines efficacy and futility bounds based on the inputs provided, following a clear priority order.

Efficacy bounds: If `criticalValues` are supplied, they take precedence and all alpha-spending parameters are ignored. Otherwise, efficacy bounds are derived from the specified alpha-spending function.

Futility bounds: Futility inputs are evaluated in the following order of priority:

1. If `futilityBounds` are provided, they override beta-spending parameters.
2. If `futilityBounds` are not specified, futility bounds are computed using the beta-spending approach with `astar` being the maximum futility stopping probability under the null hypothesis. If `typeBetaSpending == "none"`, then there is no futility stopping under the null hypothesis.

If `symmetricBounds = TRUE`, the futility bounds are set to `-efficacyBounds` and beta-spending inputs are ignored.

Value

An S3 class design object with three components:

- `overallResults`: A data frame containing the following variables:
 - `overallReject`: The overall rejection probability.
 - `alpha`: The overall significance level.
 - `attainedAlpha`: The attained significance level, which is different from the overall significance level in the presence of futility stopping.
 - `astar`: The overall futility stopping probability under the null hypothesis.
 - `kMax`: The number of stages.
 - `theta`: The parameter value.
 - `information`: The maximum information.
 - `expectedInformationH1`: The expected information under H1.
 - `expectedInformationH0`: The expected information under H0.
 - `drift`: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
 - `inflationFactor`: The inflation factor (relative to the fixed design).
- `byStageResults`: A data frame containing the following variables:
 - `informationRates`: The information rates.
 - `efficacyBounds`: The efficacy boundaries on the Z-scale.
 - `futilityBounds`: The futility boundaries on the Z-scale.
 - `rejectPerStage`: The probability for efficacy stopping.
 - `futilityPerStage`: The probability for futility stopping.
 - `cumulativeRejection`: The cumulative probability for efficacy stopping.
 - `cumulativeFutility`: The cumulative probability for futility stopping.
 - `cumulativeAlphaSpent`: The cumulative alpha spent.
 - `efficacyTheta`: The efficacy boundaries on the parameter scale.
 - `futilityTheta`: The futility boundaries on the parameter scale.

- efficacyP: The efficacy boundaries on the p-value scale.
- futilityP: The futility boundaries on the p-value scale.
- information: The cumulative information.
- efficacyStopping: Whether to allow efficacy stopping.
- futilityStopping: Whether to allow futility stopping.
- rejectPerStageH0: The probability for efficacy stopping under H0.
- futilityPerStageH0: The probability for futility stopping under H0.
- cumulativeRejectionH0: The cumulative probability for efficacy stopping under H0.
- cumulativeFutilityH0: The cumulative probability for futility stopping under H0.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - typeBetaSpending: The type of beta spending.
 - parameterBetaSpending: The parameter value for beta spending.
 - userBetaSpending: The user defined beta spending.
 - spendingTime: The error spending time at each analysis.
 - varianceRatio: The ratio of the variance under H0 to the variance under H1.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Christopher Jennison, Bruce W. Turnbull. Group Sequential Methods with Applications to Clinical Trials. Chapman & Hall/CRC: Boca Raton, 2000, ISBN:0849303168

Examples

```
(design1 <- getDesign2(
  beta = 0.149, theta = -log(0.65),
  kMax = 2, informationRates = c(0.87, 1),
  alpha = 0.004, typeAlphaSpending = "sfOF",
  astar = 0.1, typeBetaSpending = "sfHSD",
  parameterBetaSpending = -8))
```

getDesignAgreement	<i>Power and Sample Size for Cohen's kappa</i>
--------------------	--

Description

Obtains the power given sample size or obtains the sample size given power for Cohen's kappa.

Usage

```
getDesignAgreement(
  beta = NA_real_,
  n = NA_real_,
  ncats = NA_integer_,
  kappaH0 = NA_real_,
  kappa = NA_real_,
  p1 = NA_real_,
  p2 = NA_real_,
  rounding = TRUE,
  alpha = 0.025
)
```

Arguments

beta	The type II error.
n	The total sample size.
ncats	The number of categories.
kappaH0	The kappa coefficient under the null hypothesis.
kappa	The kappa coefficient under the alternative hypothesis.
p1	The marginal probabilities for the first rater.
p2	The marginal probabilities for the second rater. Defaults to be equal to the marginal probabilities for the first rater if not provided.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
alpha	The one-sided significance level. Defaults to 0.025.

Details

The kappa coefficient is defined as

$$\kappa = \frac{\pi_o - \pi_e}{1 - \pi_e},$$

where $\pi_o = \sum_i \pi_{ii}$ is the observed agreement, and $\pi_e = \sum_i \pi_{i.} \pi_{.i}$ is the expected agreement by chance.

By Fleiss et al. (1969), the variance of $\hat{\kappa}$ is given by

$$Var(\hat{\kappa}) = \frac{v_1}{n},$$

where

$$v_1 = \frac{Q_1 + Q_2 - Q_3 - Q_4}{(1 - \pi_e)^4},$$

$$Q_1 = \pi_o(1 - \pi_e)^2,$$

$$Q_2 = (1 - \pi_o)^2 \sum_i \sum_j \pi_{ij}(\pi_{i.} + \pi_{.j})^2,$$

$$Q_3 = 2(1 - \pi_o)(1 - \pi_e) \sum_i \pi_{ii}(\pi_{i.} + \pi_{.i}),$$

$$Q_4 = (\pi_o \pi_e - 2\pi_e + \pi_o)^2.$$

Given κ and marginals $\{(\pi_{i.}, \pi_{.i}) : i = 1, \dots, k\}$, we obtain π_o . The only unknowns are the double summation in Q_2 and the single summation in Q_3 .

We find the optimal configuration of cell probabilities that yield the maximum variance of $\hat{\kappa}$ by treating the problem as a linear programming problem with constraints to match the given marginal probabilities and the observed agreement and ensure that the cell probabilities are nonnegative. This is an extension of Flack et al. (1988) by allowing unequal marginal probabilities of the two raters.

We perform the optimization under both the null and alternative hypotheses to obtain $\max Var(\hat{\kappa}|\kappa = \kappa_0)$ and $\max Var(\hat{\kappa}|\kappa = \kappa_1)$ for a single subject, and then calculate the sample size or power according to the following equation:

$$\sqrt{n}|\kappa - \kappa_0| = z_{1-\alpha} \sqrt{\max Var(\hat{\kappa}|\kappa = \kappa_0)} + z_{1-\beta} \sqrt{\max Var(\hat{\kappa}|\kappa = \kappa_1)}.$$

Value

An S3 class designAgreement object with the following components:

- power: The power to reject the null hypothesis.
- alpha: The one-sided significance level.
- n: The total sample size.
- ncats: The number of categories.
- kappaH0: The kappa coefficient under the null hypothesis.
- kappa: The kappa coefficient under the alternative hypothesis.
- p1: The marginal probabilities for the first rater.
- p2: The marginal probabilities for the second rater.
- piH0: The cell probabilities that maximize the variance of estimated kappa under H0.
- pi: The cell probabilities that maximize the variance of estimated kappa under H1.
- rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

V. F. Flack, A. A. Afifi, and P. A. Lachenbruch. Sample size determinations for the two rater kappa statistic. *Psychometrika* 1988; 53:321-325.

Examples

```
(design1 <- getDesignAgreement(
  beta = 0.2, n = NA, ncats = 4, kappaH0 = 0.4, kappa = 0.6,
  p1 = c(0.1, 0.2, 0.3, 0.4), p2 = c(0.15, 0.2, 0.24, 0.41),
  rounding = TRUE, alpha = 0.05))
```

Description

Obtains the power and sample size for one-way analysis of variance.

Usage

```
getDesignANOVA(
  beta = NA_real_,
  n = NA_real_,
  ngroups = 2,
  means = NA_real_,
  stDev = 1,
  allocationRatioPlanned = NA_real_,
  rounding = TRUE,
  alpha = 0.05
)
```

Arguments

beta	The type II error.
n	The total sample size.
ngroups	The number of treatment groups.
means	The treatment group means.
stDev	The common standard deviation.
allocationRatioPlanned	Allocation ratio for the treatment groups. It has length ngroups - 1 or ngroups. If it is of length ngroups - 1, then the last treatment group will assume value 1 for allocation ratio.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
alpha	The two-sided significance level. Defaults to 0.05.

Details

Let $\{\mu_i : i = 1, \dots, k\}$ denote the group means, and $\{r_i : i = 1, \dots, k\}$ denote the randomization probabilities to the k treatment groups. Let σ denote the common standard deviation, and n denote the total sample size. Then the F -statistic

$$F = \frac{SSR/(k-1)}{SSE/(n-k)} \sim F_{k-1, n-k, \lambda}$$

where

$$\lambda = n \sum_{i=1}^k r_i (\mu_i - \bar{\mu})^2 / \sigma^2$$

is the noncentrality parameter, and $\bar{\mu} = \sum_{i=1}^k r_i \mu_i$.

Value

An S3 class designANOVA object with the following components:

- power: The power to reject the null hypothesis that there is no difference among the treatment groups.
- alpha: The two-sided significance level.
- n: The number of subjects.
- ngroups: The number of treatment groups.
- means: The treatment group means.
- stDev: The common standard deviation.
- effectsize: The effect size.
- allocationRatioPlanned: Allocation ratio for the treatment groups.
- rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignANOVA(
  beta = 0.1, ngroups = 4, means = c(1.5, 2.5, 2, 0),
  stDev = 3.5, allocationRatioPlanned = c(2, 2, 2, 1),
  alpha = 0.05))
```

getDesignANOVAContrast

Power and Sample Size for One-Way ANOVA Contrast

Description

Obtains the power and sample size for a single contrast in one-way analysis of variance.

Usage

```
getDesignANOVAContrast(
  beta = NA_real_,
  n = NA_real_,
  ngroups = 2,
  means = NA_real_,
  stDev = 1,
  contrast = NA_real_,
  meanContrastH0 = 0,
  allocationRatioPlanned = NA_real_,
  rounding = TRUE,
  alpha = 0.025
)
```

Arguments

beta	The type II error.
n	The total sample size.
ngroups	The number of treatment groups.
means	The treatment group means.
stDev	The common standard deviation.
contrast	The coefficients for the single contrast.
meanContrastH0	The mean of the contrast under the null hypothesis.
allocationRatioPlanned	Allocation ratio for the treatment groups. It has length ngroups - 1 or ngroups. If it is of length ngroups - 1, then the last treatment group will assume value 1 for allocation ratio.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
alpha	The one-sided significance level. Defaults to 0.025.

Value

An S3 class designANOVAContrast object with the following components:

- power: The power to reject the null hypothesis for the treatment contrast.
- alpha: The one-sided significance level.
- n: The number of subjects.
- ngroups: The number of treatment groups.
- means: The treatment group means.
- stDev: The common standard deviation.
- contrast: The coefficients for the single contrast.
- meanContrastH0: The mean of the contrast under the null hypothesis.
- meanContrast: The mean of the contrast under the alternative hypothesis.
- effectsize: The effect size.
- allocationRatioPlanned: Allocation ratio for the treatment groups.
- rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
(design1 <- getDesignANOVAContrast(
  beta = 0.1, ngroups = 4, means = c(1.5, 2.5, 2, 0),
  stDev = 3.5, contrast = c(1, 1, 1, -3),
  allocationRatioPlanned = c(2, 2, 2, 1),
  alpha = 0.025))
```

getDesignEquiv	<i>Power and Sample Size for Generic Group Sequential Equivalence Design</i>
----------------	--

Description

Obtains the maximum information and stopping boundaries for a generic group sequential equivalence design assuming a constant treatment effect, or obtains the power given the maximum information and stopping boundaries.

Usage

```
getDesignEquiv(
  beta = NA_real_,
  IMax = NA_real_,
  thetaLower = NA_real_,
  thetaUpper = NA_real_,
  theta = 0,
  kMax = 1L,
  informationRates = NA_real_,
  criticalValues = NULL,
  alpha = 0.05,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  spendingTime = NA_real_
)
```

Arguments

beta	The type II error.
IMax	The maximum information. Either beta or IMax should be provided while the other one should be missing.
thetaLower	The parameter value at the lower equivalence limit.
thetaUpper	The parameter value at the upper equivalence limit.
theta	The parameter value under the alternative hypothesis.
kMax	The maximum number of stages.
informationRates	The information rates. Fixed prior to the trial. Defaults to $(1:kMax) / kMax$ if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level for each of the two one-sided tests, e.g., 0.05.

typeAlphaSpending

The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".

parameterAlphaSpending

The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".

userAlphaSpending

The user defined alpha spending. Cumulative alpha spent up to each stage.

spendingTime

A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Details

Consider the equivalence design with two one-sided hypotheses:

$$H_{10} : \theta \leq \theta_{10},$$

$$H_{20} : \theta \geq \theta_{20}.$$

We reject H_{10} at or before look k if

$$Z_{1j} = (\hat{\theta}_j - \theta_{10})\sqrt{I_j} \geq b_j$$

for some $j = 1, \dots, k$, where $\{b_j : j = 1, \dots, K\}$ are the critical values associated with the specified alpha-spending function, and I_j is the information for θ (inverse variance of $\hat{\theta}$) at the j th look. For example, for estimating the risk difference $\theta = \pi_1 - \pi_2$,

$$I_j = \left\{ \frac{\pi_1(1 - \pi_1)}{n_{1j}} + \frac{\pi_2(1 - \pi_2)}{n_{2j}} \right\}^{-1}.$$

It follows that

$$(Z_{1j} \geq b_j) = (Z_j \geq b_j + \theta_{10}\sqrt{I_j}),$$

where $Z_j = \hat{\theta}_j\sqrt{I_j}$.

Similarly, we reject H_{20} at or before look k if

$$Z_{2j} = (\hat{\theta}_j - \theta_{20})\sqrt{I_j} \leq -b_j$$

for some $j = 1, \dots, k$. We have

$$(Z_{2j} \leq -b_j) = (Z_j \leq -b_j + \theta_{20}\sqrt{I_j}).$$

Let $l_j = b_j + \theta_{10}\sqrt{I_j}$, and $u_j = -b_j + \theta_{20}\sqrt{I_j}$. The cumulative probability to reject $H_0 = H_{10} \cup H_{20}$ at or before look k under the alternative hypothesis H_1 is given by

$$P_\theta \left(\bigcup_{j=1}^k (Z_{1j} \geq b_j) \cap \bigcup_{j=1}^k (Z_{2j} \leq -b_j) \right) = p_1 + p_2 - p_{12},$$

where

$$p_1 = P_{\theta} \left(\bigcup_{j=1}^k (Z_{1j} \geq b_j) \right) = P_{\theta} \left(\bigcup_{j=1}^k (Z_j \geq l_j) \right),$$

$$p_2 = P_{\theta} \left(\bigcup_{j=1}^k (Z_{2j} \leq -b_j) \right) = P_{\theta} \left(\bigcup_{j=1}^k (Z_j \leq u_j) \right),$$

and

$$p_{12} = P_{\theta} \left(\bigcup_{j=1}^k (Z_j \geq l_j) \cup (Z_j \leq u_j) \right).$$

Of note, both p_1 and p_2 can be evaluated using one-sided exit probabilities for group sequential designs. If there exists $j \leq k$ such that $l_j \leq u_j$, then $p_{12} = 1$. Otherwise, p_{12} can be evaluated using two-sided exit probabilities for group sequential designs.

Since the equivalent hypothesis is tested using two one-sided tests, the type I error is controlled. To evaluate the attained type I error of the equivalence trial under H_{10} (or H_{20}), we simply fix the control group parameters, update the active treatment group parameters according to the null hypothesis, and use the parameters in the power calculation outlined above.

Value

An S3 class `designEquiv` object with three components:

- **overallResults**: A data frame containing the following variables:
 - **overallReject**: The overall rejection probability.
 - **alpha**: The overall significance level.
 - **attainedAlphaH10**: The attained significance level under H10.
 - **attainedAlphaH20**: The attained significance level under H20.
 - **kMax**: The number of stages.
 - **thetaLower**: The parameter value at the lower equivalence limit.
 - **thetaUpper**: The parameter value at the upper equivalence limit.
 - **theta**: The parameter value under the alternative hypothesis.
 - **information**: The maximum information.
 - **expectedInformationH1**: The expected information under H1.
 - **expectedInformationH10**: The expected information under H10.
 - **expectedInformationH20**: The expected information under H20.
- **byStageResults**: A data frame containing the following variables:
 - **informationRates**: The information rates.
 - **efficacyBounds**: The efficacy boundaries on the Z-scale for each of the two one-sided tests.
 - **rejectPerStage**: The probability for efficacy stopping.
 - **cumulativeRejection**: The cumulative probability for efficacy stopping.
 - **cumulativeAlphaSpent**: The cumulative alpha for each of the two one-sided tests.
 - **cumulativeAttainedAlphaH10**: The cumulative probability for efficacy stopping under H10.
 - **cumulativeAttainedAlphaH20**: The cumulative probability for efficacy stopping under H20.
 - **efficacyThetaLower**: The efficacy boundaries on the parameter scale for the one-sided null hypothesis at the lower equivalence limit.

- efficacyThetaUpper: The efficacy boundaries on the parameter scale for the one-sided null hypothesis at the upper equivalence limit.
- efficacyP: The efficacy bounds on the p-value scale for each of the two one-sided tests.
- information: The cumulative information.
- settings: A list containing the following components:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - spendingTime: The error spending time at each analysis.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Example 1: obtain the maximum information given power
(design1 <- getDesignEquiv(
  beta = 0.2, thetaLower = log(0.8), thetaUpper = log(1.25),
  kMax = 2, informationRates = c(0.5, 1),
  alpha = 0.05, typeAlphaSpending = "sfOF"))
```

```
# Example 2: obtain power given the maximum information
(design2 <- getDesignEquiv(
  IMax = 72.5, thetaLower = log(0.7), thetaUpper = -log(0.7),
  kMax = 3, informationRates = c(0.5, 0.75, 1),
  alpha = 0.05, typeAlphaSpending = "sfOF"))
```

getDesignFisherExact *Power and Sample Size for Fisher's Exact Test for Two Proportions*

Description

Obtains the power given sample size or obtains the sample size given power for Fisher's exact test for two proportions.

Usage

```
getDesignFisherExact(
  beta = NA_real_,
  n = NA_real_,
  pi1 = NA_real_,
  pi2 = NA_real_,
  allocationRatioPlanned = 1,
  alpha = 0.05
)
```

Arguments

beta	The type II error.
n	The total sample size.
pi1	The assumed probability for the active treatment group.
pi2	The assumed probability for the control group.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
alpha	The two-sided significance level. Defaults to 0.05.

Value

A data frame with the following variables:

- alpha: The two-sided significance level.
- power: The power.
- n: The sample size.
- pi1: The assumed probability for the active treatment group.
- pi2: The assumed probability for the control group.
- allocationRatioPlanned: Allocation ratio for the active treatment versus control.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignFisherExact(
  beta = 0.2, pi1 = 0.5, pi2 = 0.2, alpha = 0.05))
```

getDesignLogistic

Power and Sample Size for Logistic Regression

Description

Obtains the power given sample size or obtains the sample size given power for logistic regression of a binary response given the covariate of interest and other covariates.

Usage

```

getDesignLogistic(
  beta = NA_real_,
  n = NA_real_,
  ncovariates = NA_integer_,
  nconfigs = NA_integer_,
  x = NA_real_,
  pconfigs = NA_real_,
  corr = 0,
  oddsratios = NA_real_,
  responseprob = NA_real_,
  rounding = TRUE,
  alpha = 0.05
)

```

Arguments

beta	The type II error.
n	The total sample size.
ncovariates	The number of covariates.
nconfigs	The number of configurations of discretized covariate values.
x	The matrix of covariate values.
pconfigs	The vector of probabilities for the configurations.
corr	The multiple correlation between the predictor and other covariates. Defaults to 0.
oddsratios	The odds ratios for one unit increase in the covariates.
responseprob	The response probability in the full model when all predictor variables are equal to their means.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
alpha	The two-sided significance level. Defaults to 0.05.

Details

We consider the logistic regression of a binary response variable Y on a set of predictor variables $x = (x_1, \dots, x_K)^T$ with x_1 being the covariate of interest: $\log \frac{P(Y_i=1)}{1-P(Y_i=1)} = \psi_0 + x_i^T \psi$, where $\psi = (\psi_1, \dots, \psi_K)^T$. Similar to Self et al (1992), we assume that all covariates are either inherently discrete or discretized from continuous distributions (e.g. using the quantiles). Let m denote the total number of configurations of the covariate values. Let

$$\pi_i = P(x = x_i), i = 1, \dots, m$$

denote the probabilities for the configurations of the covariates under independence. The likelihood ratio test statistic for testing $H_0 : \psi_1 = 0$ can be approximated by a noncentral chi-square distribution with one degree of freedom and noncentrality parameter

$$\Delta = 2 \sum_{i=1}^m \pi_i [b'(\theta_i)(\theta_i - \theta_i^*) - \{b(\theta_i) - b(\theta_i^*)\}],$$

where

$$\theta_i = \psi_0 + \sum_{j=1}^k \psi_j x_{ij},$$

$$\theta_i^* = \psi_0^* + \sum_{j=2}^k \psi_j^* x_{ij},$$

for $\psi_0^* = \psi_0 + \psi_1 \mu_1$, and $\psi_j^* = \psi_j$ for $j = 2, \dots, K$. Here μ_1 is the mean of x_1 , e.g., $\mu_1 = \sum_i \pi_i x_{i1}$. In addition, by formulating the logistic regression in the framework of generalized linear models,

$$b(\theta) = \log(1 + \exp(\theta)),$$

and

$$b'(\theta) = \frac{\exp(\theta)}{1 + \exp(\theta)}.$$

The regression coefficients ψ can be obtained by taking the log of the odds ratios for the covariates. The intercept ψ_0 can be derived as

$$\psi_0 = \log(\bar{\mu}/(1 - \bar{\mu})) - \sum_{j=1}^K \psi_j \mu_j,$$

where $\bar{\mu}$ denotes the response probability when all predictor variables are equal to their means.

Finally, let ρ denote the multiple correlation between the predictor and other covariates. The non-centrality parameter of the chi-square test is adjusted downward by multiplying by $1 - \rho^2$.

Value

An S3 class `designLogistic` object with the following components:

- `power`: The power to reject the null hypothesis.
- `alpha`: The two-sided significance level.
- `n`: The total sample size.
- `ncovariates`: The number of covariates.
- `nconfigs`: The number of configurations of discretized covariate values.
- `x`: The matrix of covariate values.
- `pconfigs`: The vector of probabilities for the configurations.
- `corr`: The multiple correlation between the predictor and other covariates.
- `oddsratios`: The odds ratios for one unit increase in the covariates.
- `responseprob`: The response probability in the full model when all predictor variables are equal to their means.
- `effectsize`: The effect size for the chi-square test.
- `rounding`: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Steven G. Self, Robert H. Mauritsen and Jill Ohara. Power calculations for likelihood ratio tests in generalized linear models. *Biometrics* 1992; 48:31-39.

Examples

```
# two ordinal covariates
x1 = c(5, 10, 15, 20)
px1 = c(0.2, 0.3, 0.3, 0.2)

x2 = c(2, 4, 6)
px2 = c(0.4, 0.4, 0.2)

# discretizing a normal distribution with mean 4 and standard deviation 2
nbins = 10
x3 = qnorm(((1:nbins) - 0.5)/nbins)*2 + 4
px3 = rep(1/nbins, nbins)

# combination of covariate values
nconfigs = length(x1)*length(x2)*length(x3)
x = expand.grid(x3 = x3, x2 = x2, x1 = x1)
x = as.matrix(x[, ncol(x):1])

# probabilities for the covariate configurations under independence
pconfigs = as.numeric(px1 %x% px2 %x% px3)

# convert the odds ratio for the predictor variable in 5-unit change
# to the odds ratio in 1-unit change
(design1 <- getDesignLogistic(
  beta = 0.1, ncovariates = 3,
  nconfigs = nconfigs,
  x = x,
  pconfigs = pconfigs,
  oddsratios = c(1.2^(1/5), 1.4, 1.3),
  responseprob = 0.25,
  alpha = 0.1))
```

getDesignMeanDiff

Group Sequential Design for Two-Sample Mean Difference

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for two-sample mean difference.

Usage

```

getDesignMeanDiff(
  beta = NA_real_,
  n = NA_real_,
  meanDiffH0 = 0,
  meanDiff = 0.5,
  stDev = 1,
  allocationRatioPlanned = 1,
  normalApproximation = TRUE,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NA_real_,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NA_real_,
  futilityCP = NA_real_,
  futilityMeanDiff = NA_real_,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
  spendingTime = NA_real_
)

```

Arguments

<code>beta</code>	The type II error.
<code>n</code>	The total sample size.
<code>meanDiffH0</code>	The mean difference under the null hypothesis. Defaults to 0.
<code>meanDiff</code>	The mean difference under the alternative hypothesis.
<code>stDev</code>	The standard deviation.
<code>allocationRatioPlanned</code>	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
<code>normalApproximation</code>	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
<code>rounding</code>	Whether to round up sample size. Defaults to 1 for sample size rounding.
<code>kMax</code>	The maximum number of stages.
<code>informationRates</code>	The information rates. Fixed prior to the trial. Defaults to (1:kMax) / kMax if left unspecified.

efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsiatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	The futility boundary on the conditional power scale.
futilityMeanDiff	The futility boundary on the mean difference scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Value

An S3 class `designMeanDiff` object with three components:

- `overallResults`: A data frame containing the following variables:
 - `overallReject`: The overall rejection probability.

- alpha: The overall significance level.
- attainedAlpha: The attained significance level, which is different from the overall significance level in the presence of futility stopping.
- kMax: The number of stages.
- theta: The parameter value.
- information: The maximum information.
- expectedInformationH1: The expected information under H1.
- expectedInformationH0: The expected information under H0.
- drift: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
- inflationFactor: The inflation factor (relative to the fixed design).
- numberOfSubjects: The maximum number of subjects.
- expectedNumberOfSubjectsH1: The expected number of subjects under H1.
- expectedNumberOfSubjectsH0: The expected number of subjects under H0.
- meanDiffH0: The mean difference under the null hypothesis.
- meanDiff: The mean difference under the alternative hypothesis.
- stDev: The standard deviation.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale.
 - futilityBounds: The futility boundaries on the Z-scale.
 - rejectPerStage: The probability for efficacy stopping.
 - futilityPerStage: The probability for futility stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeFutility: The cumulative probability for futility stopping.
 - cumulativeAlphaSpent: The cumulative alpha spent.
 - efficacyP: The efficacy boundaries on the p-value scale.
 - futilityP: The futility boundaries on the p-value scale.
 - information: The cumulative information.
 - efficacyStopping: Whether to allow efficacy stopping.
 - futilityStopping: Whether to allow futility stopping.
 - rejectPerStageH0: The probability for efficacy stopping under H0.
 - futilityPerStageH0: The probability for futility stopping under H0.
 - cumulativeRejectionH0: The cumulative probability for efficacy stopping under H0.
 - cumulativeFutilityH0: The cumulative probability for futility stopping under H0.
 - efficacyMeanDiff: The efficacy boundaries on the mean difference scale.
 - futilityMeanDiff: The futility boundaries on the mean difference scale.
 - numberOfSubjects: The number of subjects.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - typeBetaSpending: The type of beta spending.

- parameterBetaSpending: The parameter value for beta spending.
- userBetaSpending: The user defined beta spending.
- spendingTime: The error spending time at each analysis.
- allocationRatioPlanned: Allocation ratio for the active treatment versus control.
- normalApproximation: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution.
- rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Example 1: group sequential trial power calculation
(design1 <- getDesignMeanDiff(
  beta = NA, n = 456, meanDiff = 9, stDev = 32,
  kMax = 5, alpha = 0.025, typeAlphaSpending = "sfOF",
  typeBetaSpending = "sfP"))

# Example 2: sample size calculation for two-sample t-test
(design2 <- getDesignMeanDiff(
  beta = 0.1, n = NA, meanDiff = 0.3, stDev = 1,
  normalApproximation = FALSE, alpha = 0.025))
```

getDesignMeanDiffCarryover

Power and Sample Size for Direct Treatment Effects in Crossover Trials

Description

Obtains the power and sample size for direct treatment effects in crossover trials accounting or without accounting for carryover effects.

Usage

```
getDesignMeanDiffCarryover(
  beta = NA_real_,
  n = NA_real_,
  trtpair = NA_real_,
  carryover = TRUE,
  meanDiffH0 = 0,
  meanDiff = 0.5,
  stDev = 1,
  corr = 0.5,
  design = NA_real_,
```

```

    cumdrop = NA_real_,
    allocationRatioPlanned = NA_real_,
    normalApproximation = FALSE,
    rounding = TRUE,
    alpha = 0.025
  )

```

Arguments

beta	The type II error.
n	The total sample size.
trtpair	The treatment pair of interest to power the study. If not given, it defaults to comparing the first treatment to the last treatment.
carryover	Whether to account for carryover effects in the power calculation. Defaults to TRUE.
meanDiffH0	The mean difference for the treatment pair of interest under the null hypothesis. Defaults to 0.
meanDiff	The mean difference for the treatment pair of interest under the alternative hypothesis.
stDev	The standard deviation for within-subject random error.
corr	The intra-subject correlation due to subject random effect.
design	The crossover design represented by a matrix with rows indexing the sequences, columns indexing the periods, and matrix entries indicating the treatments.
cumdrop	The cumulative dropout rate over periods.
allocationRatioPlanned	Allocation ratio for the sequences. Defaults to equal randomization if not provided.
normalApproximation	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution.
rounding	Whether to round up the sample size. Defaults to TRUE for sample size rounding.
alpha	The one-sided significance level. Defaults to 0.025.

Details

The linear mixed-effects model to assess the direct treatment effects in the presence of carryover treatment effects is given by

$$y_{ijk} = \mu + \alpha_i + b_{ij} + \gamma_k + \tau_{d(i,k)} + \lambda_{c(i,k-1)} + e_{ijk}$$

$$i = 1, \dots, n; j = 1, \dots, r_i; k = 1, \dots, p; d, c = 1, \dots, t$$

where μ is the general mean, α_i is the effect of the i th treatment sequence, b_{ij} is the random effect with variance σ_b^2 for the j th subject of the i th treatment sequence, γ_k is the period effect, and e_{ijk} is the random error with variance σ^2 for the subject in period k . The direct effect of the

treatment administered in period k of sequence i is $\tau_{d(i,k)}$, and $\lambda_{c(i,k-1)}$ is the carryover effect of the treatment administered in period $k-1$ of sequence i . The value of the carryover effect for the observed response in the first period is $\lambda_{c(i,0)} = 0$ since there is no carryover effect in the first period. The intra-subject correlation due to the subject random effect is

$$\rho = \frac{\sigma_b^2}{\sigma_b^2 + \sigma^2}.$$

Therefore, $\text{stDev} = \sigma^2$ and $\text{corr} = \rho$. By constructing the design matrix X for the linear model with a compound symmetry covariance matrix for the response vector of a subject, we can obtain

$$\text{Var}(\hat{\beta}) = (X'V^{-1}X)^{-1}.$$

The covariance matrix for the direct treatment effects and carryover treatment effects can be extracted from the appropriate sub-matrices. The covariance matrix for the direct treatment effects without accounting for the carryover treatment effects can be obtained by omitting the carryover effect terms from the model.

The power is for the direct treatment effect for the treatment pair of interest with or without accounting for carryover effects as determined by the input parameter `carryover`. The relative efficiency is for the direct treatment effect for the treatment pair of interest accounting for carryover effects relative to that without accounting for carryover effects.

The degrees of freedom for the t-test accounting for carryover effects can be calculated as the total number of observations minus the number of subjects minus $p-1$ minus $2(t-1)$ to account for the subject effect, period effect, and direct and carryover treatment effects. The degrees of freedom for the t-test without accounting for carryover effects is equal to the total number of observations minus the number of subjects minus $p-1$ minus $t-1$ to account for the subject effect, period effect, and direct treatment effects.

Value

An S3 class `designMeanDiffCarryover` object with the following components:

- `power`: The power to reject the null hypothesis.
- `alpha`: The one-sided significance level.
- `numberOfSubjects`: The maximum number of subjects.
- `trtpair`: The treatment pair of interest to power the study.
- `carryover`: Whether to account for carryover effects in the power calculation.
- `meanDiffH0`: The mean difference for the treatment pair of interest under the null hypothesis.
- `meanDiff`: The mean difference for the treatment pair of interest under the alternative hypothesis.
- `stDev`: The standard deviation for within-subject random error.
- `corr`: The intra-subject correlation due to subject random effect.
- `design`: The crossover design represented by a matrix with rows indexing the sequences, columns indexing the periods, and matrix entries indicating the treatments.

- **designMatrix**: The design matrix accounting for intercept, sequence, period, direct treatment effects and carryover treatment effects when `carryover = TRUE`, or the design matrix accounting for intercept, sequence, period, and direct treatment effects when `carryover = FALSE`.
- **nseq**: The number of sequences.
- **nprd**: The number of periods.
- **ntrt**: The number of treatments.
- **cumdrop**: The cumulative dropout rate over periods.
- **V_direct_only**: The covariance matrix for direct treatment effects without accounting for carryover effects. The treatment comparisons for the covariance matrix are for the first $t - 1$ treatments relative to the last treatment.
- **V_direct_carry**: The covariance matrix for direct and carryover treatment effects.
- **v_direct_only**: The variance of the direct treatment effect for the treatment pair of interest without accounting for carryover effects.
- **v_direct**: The variance of the direct treatment effect for the treatment pair of interest accounting for carryover effects.
- **v_carry**: The variance of the carryover treatment effect for the treatment pair of interest.
- **releff_direct**: The relative efficiency of the design for estimating the direct treatment effect for the treatment pair of interest after accounting for carryover effects with respect to that without accounting for carryover effects. This is equal to v_direct_only/v_direct .
- **releff_carry**: The relative efficiency of the design for estimating the carryover effect for the treatment pair of interest. This is equal to v_direct_only/v_carry .
- **half_width**: The half-width of the confidence interval for the direct treatment effect for the treatment pair of interest.
- **nu**: Degrees of freedom for the t-test.
- **allocationRatioPlanned**: Allocation ratio for the sequences.
- **normalApproximation**: The type of computation of the p-values. If `TRUE`, the variance is assumed to be known, otherwise the calculations are performed with the t distribution.
- **rounding**: Whether to round up the sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Robert O. Kuehl. Design of Experiments: Statistical Principles of Research Design and Analysis. Brooks/Cole: Pacific Grove, CA. 2000.

Examples

```
# Williams design for 4 treatments

(design1 <- getDesignMeanDiffCarryover(
```

```

beta = 0.2, n = NA,
meanDiff = 0.5, stDev = 1,
design = matrix(c(1, 4, 2, 3,
                 2, 1, 3, 4,
                 3, 2, 4, 1,
                 4, 3, 1, 2),
               4, 4, byrow = TRUE),
alpha = 0.025))

```

getDesignMeanDiffCarryoverEquiv

Power and Sample Size for Equivalence in Direct Treatment Effects in Crossover Trials

Description

Obtains the power and sample size for equivalence in direct treatment effects in crossover trials accounting or without accounting for carryover effects.

Usage

```

getDesignMeanDiffCarryoverEquiv(
  beta = NA_real_,
  n = NA_real_,
  trtpair = NA_real_,
  carryover = TRUE,
  meanDiffLower = NA_real_,
  meanDiffUpper = NA_real_,
  meanDiff = 0,
  stDev = 1,
  corr = 0.5,
  design = NA_real_,
  cumdrop = NA_real_,
  allocationRatioPlanned = NA_real_,
  normalApproximation = FALSE,
  rounding = TRUE,
  alpha = 0.025
)

```

Arguments

beta	The type II error.
n	The total sample size.
trtpair	The treatment pair of interest to power the study. If not given, it defaults to comparing the first treatment to the last treatment.

carryover	Whether to account for carryover effects in the power calculation. Defaults to TRUE.
meanDiffLower	The lower equivalence limit of mean difference for the treatment pair of interest.
meanDiffUpper	The upper equivalence limit of mean difference for the treatment pair of interest.
meanDiff	The mean difference under the alternative hypothesis,
stDev	The standard deviation for within-subject random error.
corr	The intra-subject correlation due to subject random effect.
design	The crossover design represented by a matrix with rows indexing the sequences, columns indexing the periods, and matrix entries indicating the treatments.
cumdrop	The cumulative dropout rate over periods.
allocationRatioPlanned	Allocation ratio for the sequences. Defaults to equal randomization if not provided.
normalApproximation	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution.
rounding	Whether to round up the sample size. Defaults to TRUE for sample size rounding.
alpha	The one-sided significance level. Defaults to 0.025.

Details

The linear mixed-effects model to assess the direct treatment effects in the presence of carryover treatment effects is given by

$$y_{ijk} = \mu + \alpha_i + b_{ij} + \gamma_k + \tau_{d(i,k)} + \lambda_{c(i,k-1)} + e_{ijk}$$

$$i = 1, \dots, n; j = 1, \dots, r_i; k = 1, \dots, p; d, c = 1, \dots, t$$

where μ is the general mean, α_i is the effect of the i th treatment sequence, b_{ij} is the random effect with variance σ_b^2 for the j th subject of the i th treatment sequence, γ_k is the period effect, and e_{ijk} is the random error with variance σ^2 for the subject in period k . The direct effect of the treatment administered in period k of sequence i is $\tau_{d(i,k)}$, and $\lambda_{c(i,k-1)}$ is the carryover effect of the treatment administered in period $k-1$ of sequence i . The value of the carryover effect for the observed response in the first period is $\lambda_{c(i,0)} = 0$ since there is no carryover effect in the first period. The intra-subject correlation due to the subject random effect is

$$\rho = \frac{\sigma_b^2}{\sigma_b^2 + \sigma^2}.$$

Therefore, $\text{stDev} = \sigma$ and $\text{corr} = \rho$. By constructing the design matrix X for the linear model with a compound symmetry covariance matrix for the response vector of a subject, we can obtain

$$\text{Var}(\hat{\beta}) = (X'V^{-1}X)^{-1}.$$

The covariance matrix for the direct treatment effects and carryover treatment effects can be extracted from the appropriate sub-matrices. The covariance matrix for the direct treatment effects

without accounting for the carryover treatment effects can be obtained by omitting the carryover effect terms from the model.

The power is for the direct treatment effect for the treatment pair of interest with or without accounting for carryover effects as determined by the input parameter `carryover`. The relative efficiency is for the direct treatment effect for the treatment pair of interest accounting for carryover effects relative to that without accounting for carryover effects.

The degrees of freedom for the t-test accounting for carryover effects can be calculated as the total number of observations minus the number of subjects minus $p - 1$ minus $2(t - 1)$ to account for the subject effect, period effect, and direct and carryover treatment effects. The degrees of freedom for the t-test without accounting for carryover effects is equal to the total number of observations minus the number of subjects minus $p - 1$ minus $t - 1$ to account for the subject effect, period effect, and direct treatment effects.

Value

An S3 class `designMeanDiffCarryover` object with the following components:

- `power`: The power to reject the null hypothesis.
- `alpha`: The one-sided significance level.
- `numberOfSubjects`: The maximum number of subjects.
- `trtpair`: The treatment pair of interest to power the study.
- `carryover`: Whether to account for carryover effects in the power calculation.
- `meanDiffLower`: The lower equivalence limit of mean difference for the treatment pair of interest.
- `meanDiffUpper`: The upper equivalence limit of mean difference for the treatment pair of interest.
- `meanDiff`: The mean difference for the treatment pair of interest under the alternative hypothesis.
- `stDev`: The standard deviation for within-subject random error.
- `corr`: The intra-subject correlation due to subject random effect.
- `design`: The crossover design represented by a matrix with rows indexing the sequences, columns indexing the periods, and matrix entries indicating the treatments.
- `designMatrix`: The design matrix accounting for intercept, sequence, period, direct treatment effects and carryover treatment effects when `carryover = TRUE`, or the design matrix accounting for intercept, sequence, period, and direct treatment effects when `carryover = FALSE`.
- `nseq`: The number of sequences.
- `nprd`: The number of periods.
- `ntrt`: The number of treatments.
- `cumdrop`: The cumulative dropout rate over periods.
- `V_direct_only`: The covariance matrix for direct treatment effects without accounting for carryover effects. The treatment comparisons for the covariance matrix are for the first $t - 1$ treatments relative to the last treatment.

- `V_direct_carry`: The covariance matrix for direct and carryover treatment effects.
- `v_direct_only`: The variance of the direct treatment effect for the treatment pair of interest without accounting for carryover effects.
- `v_direct`: The variance of the direct treatment effect for the treatment pair of interest accounting for carryover effects.
- `v_carry`: The variance of the carryover treatment effect for the treatment pair of interest.
- `releff_direct`: The relative efficiency of the design for estimating the direct treatment effect for the treatment pair of interest after accounting for carryover effects with respect to that without accounting for carryover effects. This is equal to v_direct_only/v_direct .
- `releff_carry`: The relative efficiency of the design for estimating the carryover effect for the treatment pair of interest. This is equal to v_direct_only/v_carry .
- `half_width`: The half-width of the confidence interval for the direct treatment effect for the treatment pair of interest.
- `nu`: Degrees of freedom for the t-test.
- `allocationRatioPlanned`: Allocation ratio for the sequences.
- `normalApproximation`: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution.
- `rounding`: Whether to round up the sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Robert O. Kuehl. Design of Experiments: Statistical Principles of Research Design and Analysis. Brooks/Cole: Pacific Grove, CA. 2000.

Examples

```
# Williams design for 4 treatments

(design1 <- getDesignMeanDiffCarryoverEquiv(
  beta = 0.2, n = NA,
  meanDiffLower = -1.3, meanDiffUpper = 1.3,
  meanDiff = 0, stDev = 2.2,
  design = matrix(c(1, 4, 2, 3,
                    2, 1, 3, 4,
                    3, 2, 4, 1,
                    4, 3, 1, 2),
                  4, 4, byrow = TRUE),
  alpha = 0.025))
```

getDesignMeanDiffEquiv

Group Sequential Design for Equivalence in Two-Sample Mean Difference

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for equivalence in two-sample mean difference.

Usage

```
getDesignMeanDiffEquiv(
  beta = NA_real_,
  n = NA_real_,
  meanDiffLower = NA_real_,
  meanDiffUpper = NA_real_,
  meanDiff = 0,
  stDev = 1,
  allocationRatioPlanned = 1,
  normalApproximation = TRUE,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  alpha = 0.05,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  spendingTime = NA_real_
)
```

Arguments

beta	The type II error.
n	The total sample size.
meanDiffLower	The lower equivalence limit of mean difference.
meanDiffUpper	The upper equivalence limit of mean difference.
meanDiff	The mean difference under the alternative hypothesis.
stDev	The standard deviation.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
normalApproximation	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.

<code>rounding</code>	Whether to round up sample size. Defaults to 1 for sample size rounding.
<code>kMax</code>	The maximum number of stages.
<code>informationRates</code>	The information rates. Fixed prior to the trial. Defaults to $(1:kMax) / kMax$ if left unspecified.
<code>alpha</code>	The significance level for each of the two one-sided tests. Defaults to 0.05.
<code>typeAlphaSpending</code>	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
<code>parameterAlphaSpending</code>	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
<code>userAlphaSpending</code>	The user defined alpha spending. Cumulative alpha spent up to each stage.
<code>spendingTime</code>	A vector of length <code>kMax</code> for the error spending time at each analysis. Defaults to missing, in which case, it is the same as <code>informationRates</code> .

Value

An S3 class `designMeanDiffEquiv` object with three components:

- `overallResults`: A data frame containing the following variables:
 - `overallReject`: The overall rejection probability.
 - `alpha`: The significance level for each of the two one-sided tests. Defaults to 0.05.
 - `attainedAlpha`: The attained significance level.
 - `kMax`: The number of stages.
 - `information`: The maximum information.
 - `expectedInformationH1`: The expected information under H1.
 - `expectedInformationH0`: The expected information under H0.
 - `numberOfSubjects`: The maximum number of subjects.
 - `expectedNumberOfSubjectsH1`: The expected number of subjects under H1.
 - `expectedNumberOfSubjectsH0`: The expected number of subjects under H0.
 - `meanDiffLower`: The lower equivalence limit of mean difference.
 - `meanDiffUpper`: The upper equivalence limit of mean difference.
 - `meanDiff`: The mean difference under the alternative hypothesis.
 - `stDev`: The standard deviation.
- `byStageResults`: A data frame containing the following variables:
 - `informationRates`: The information rates.
 - `efficacyBounds`: The efficacy boundaries on the Z-scale for each of the two one-sided tests.

- rejectPerStage: The probability for efficacy stopping.
- cumulativeRejection: The cumulative probability for efficacy stopping.
- cumulativeAlphaSpent: The cumulative alpha for each of the two one-sided tests.
- cumulativeAttainedAlpha: The cumulative probability for efficacy stopping under H0.
- efficacyMeanDiffLower: The efficacy boundaries on the mean difference scale for the one-sided null hypothesis on the lower equivalence limit.
- efficacyMeanDiffUpper: The efficacy boundaries on the mean difference scale for the one-sided null hypothesis on the upper equivalence limit.
- efficacyP: The efficacy bounds on the p-value scale for each of the two one-sided tests.
- information: The cumulative information.
- numberOfSubjects: The number of subjects.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - spendingTime: The error spending time at each analysis.
 - allocationRatioPlanned: Allocation ratio for the active treatment versus control.
 - normalApproximation: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
 - rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Example 1: group sequential trial power calculation
(design1 <- getDesignMeanDiffEquiv(
  beta = 0.1, n = NA, meanDiffLower = -1.3, meanDiffUpper = 1.3,
  meanDiff = 0, stDev = 2.2,
  kMax = 4, alpha = 0.05, typeAlphaSpending = "sf0F"))

# Example 2: sample size calculation for t-test
(design2 <- getDesignMeanDiffEquiv(
  beta = 0.1, n = NA, meanDiffLower = -1.3, meanDiffUpper = 1.3,
  meanDiff = 0, stDev = 2.2,
  normalApproximation = FALSE, alpha = 0.05))
```

getDesignMeanDiffMMRM *Group Sequential Design for Two-Sample Mean Difference From the MMRM Model*

Description

Obtains the power and sample size for two-sample mean difference at the last time point from the mixed-model for repeated measures (MMRM) model.

Usage

```
getDesignMeanDiffMMRM(
  beta = NA_real_,
  meanDiffH0 = 0,
  meanDiff = 0.5,
  k = 1,
  t = NA_real_,
  covar1 = diag(k),
  covar2 = NA_real_,
  accrualTime = 0,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0,
  gamma1 = 0,
  gamma2 = 0,
  accrualDuration = NA_real_,
  allocationRatioPlanned = 1,
  normalApproximation = TRUE,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NA_real_,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NA_real_,
  futilityCP = NA_real_,
  futilityMeanDiff = NA_real_,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
  spendingTime = NA_real_
)
```

Arguments

beta	The type II error.
meanDiffH0	The mean difference at the last time point under the null hypothesis. Defaults to 0.
meanDiff	The mean difference at the last time point under the alternative hypothesis.
k	The number of postbaseline time points.
t	The postbaseline time points.
covar1	The covariance matrix for the repeated measures given baseline for the active treatment group.
covar2	The covariance matrix for the repeated measures given baseline for the control group. If missing, it will be set equal to the covariance matrix for the active treatment group.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
gamma1	The hazard rate for exponential dropout, or a vector of hazard rates for piecewise exponential dropout for the active treatment group.
gamma2	The hazard rate for exponential dropout, or a vector of hazard rates for piecewise exponential dropout for the control group.
accrualDuration	Duration of the enrollment period.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
normalApproximation	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The degrees of freedom for the t-distribution is the total effective sample size minus 2. The exact calculation using the t distribution is only implemented for the fixed design.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
kMax	The maximum number of stages.
informationRates	The information rates. Defaults to $(1:kMax) / kMax$ if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.

futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	The futility bounds on the conditional power scale.
futilityMeanDiff	The futility bounds on the mean difference scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Details

Consider a longitudinal study with two treatment groups. The outcome is measured at baseline and at k postbaseline time points. For each treatment group, the outcomes are assumed to follow a multivariate normal distribution. Conditional on baseline, the covariance matrix of the post-baseline outcomes is denoted by Σ_1 for the active treatment group and Σ_2 for the control group. Let μ_1 and μ_2 denote the mean vectors of post-baseline outcomes for the active and control groups, respectively. We are interested in testing the null hypothesis $H_0 : \mu_{1,k} - \mu_{2,k} = \delta_0$ against the alternative $H_1 : \mu_{1,k} - \mu_{2,k} = \delta$.

The study design is based on the information for treatment difference at the last postbaseline time point. This information is given by

$$I = 1/\text{Var}(\hat{\mu}_{1,k} - \hat{\mu}_{2,k})$$

In the presence of monotone missing data, let $p_{g,1}, \dots, p_{g,k}$ denote the proportions of subjects in observed data patterns 1 through k for treatment group $g = 1$ (active) or 2 (control). A subject in pattern j has complete data up to time t_j , i.e., the observed outcomes are $y_{i,1}, \dots, y_{i,j}$, with missing values for $y_{i,j+1}, \dots, y_{i,k}$.

According to Lu et al. (2008), the information matrix for the post-baseline mean vector in group g is

$$I_g = n\pi_g J_g$$

where π_g is the proportion of subjects in group g , and

$$J_g = \sum_{j=1}^k p_{g,j} \begin{pmatrix} \Sigma_{g,j}^{-1} & 0 \\ 0 & 0 \end{pmatrix}$$

Here, $\Sigma_{g,j}$ is the leading $j \times j$ principal submatrix of Σ_g . It follows that

$$\text{Var}(\hat{\mu}_{1,k} - \hat{\mu}_{2,k}) = \frac{1}{n} \left(\frac{1}{\pi_1} J_1^{-1}[k, k] + \frac{1}{\pi_2} J_2^{-1}[k, k] \right)$$

The observed data pattern probabilities depend on the accrual and dropout distributions. Let $H(u)$ denote the cumulative distribution function of enrollment time u , $G_g(t)$ denote the survival function of dropout time t for treatment group g , and τ denote the calendar time at interim or final analysis. Then, for $j = 1, \dots, k-1$, the probability that a subject in group g falls into observed data pattern j is

$$p_{g,j} = H(\tau - t_j)G_g(t_j) - H(\tau - t_{j+1})G_g(t_{j+1})$$

For the last pattern ($j = k$, i.e., completers),

$$p_{g,k} = H(\tau - t_k)G_g(t_k)$$

For the final analysis, τ is the study duration, so $H(\tau - t_j) = 1$ for all j . Therefore, the pattern probabilities depend only on the dropout distribution:

$$p_{g,j} = G_g(t_j) - G_g(t_{j+1}), \quad j = 1, \dots, k-1$$

and

$$p_{g,k} = G_g(t_k)$$

Cumulative dropout probabilities at post-baseline time points can be used to define a piecewise exponential dropout distribution. Let $F_g(t_j)$ denote the cumulative probability of dropout by time t_j for treatment group g . The left endpoints of the piecewise survival time intervals are given by $t_0 = 0, t_1, \dots, t_{k-1}$. The hazard rate in the interval $(t_{j-1}, t_j]$ is given by

$$\gamma_{g,j} = -\log \left(\frac{1 - F_g(t_j)}{1 - F_g(t_{j-1})} \right) / (t_j - t_{j-1}), \quad j = 1, \dots, k$$

Value

An S3 class `designMeanDiffMMRM` object with three components:

- `overallResults`: A data frame containing the following variables:
 - `overallReject`: The overall rejection probability.
 - `alpha`: The overall significance level.
 - `attainedAlpha`: The attained significance level, which is different from the overall significance level in the presence of futility stopping.
 - `kMax`: The number of stages.
 - `theta`: The parameter value.
 - `information`: The maximum information.
 - `expectedInformationH1`: The expected information under H1.
 - `expectedInformationH0`: The expected information under H0.
 - `drift`: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
 - `inflationFactor`: The inflation factor (relative to the fixed design).
 - `numberOfSubjects`: The maximum number of subjects.
 - `studyDuration`: The maximum study duration.
 - `expectedNumberOfSubjectsH1`: The expected number of subjects under H1.
 - `expectedNumberOfSubjectsH0`: The expected number of subjects under H0.
 - `expectedStudyDurationH1`: The expected study duration under H1.
 - `expectedStudyDurationH0`: The expected study duration under H0.
 - `accrualDuration`: The accrual duration.
 - `followupTime`: The follow-up time.
 - `fixedFollowup`: Whether a fixed follow-up design is used.
 - `meanDiffH0`: The mean difference under H0.
 - `meanDiff`: The mean difference under H1.
- `byStageResults`: A data frame containing the following variables:
 - `informationRates`: The information rates.
 - `efficacyBounds`: The efficacy boundaries on the Z-scale.
 - `futilityBounds`: The futility boundaries on the Z-scale.
 - `rejectPerStage`: The probability for efficacy stopping.
 - `futilityPerStage`: The probability for futility stopping.
 - `cumulativeRejection`: The cumulative probability for efficacy stopping.
 - `cumulativeFutility`: The cumulative probability for futility stopping.
 - `cumulativeAlphaSpent`: The cumulative alpha spent.
 - `efficacyP`: The efficacy boundaries on the p-value scale.
 - `futilityP`: The futility boundaries on the p-value scale.
 - `information`: The cumulative information.
 - `efficacyStopping`: Whether to allow efficacy stopping.
 - `futilityStopping`: Whether to allow futility stopping.
 - `rejectPerStageH0`: The probability for efficacy stopping under H0.
 - `futilityPerStageH0`: The probability for futility stopping under H0.

- cumulativeRejectionH0: The cumulative probability for efficacy stopping under H0.
- cumulativeFutilityH0: The cumulative probability for futility stopping under H0.
- efficacyMeanDiff: The efficacy boundaries on the mean difference scale.
- futilityMeanDiff: The futility boundaries on the mean difference scale.
- numberOfSubjects: The number of subjects.
- numberOfCompleters: The number of completers.
- analysisTime: The average time since trial start.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - typeBetaSpending: The type of beta spending.
 - parameterBetaSpending: The parameter value for beta spending.
 - userBetaSpending: The user defined beta spending.
 - spendingTime: The error spending time at each analysis.
 - allocationRatioPlanned: The allocation ratio for the active treatment versus control.
 - accrualTime: A vector that specifies the starting time of piecewise Poisson enrollment time intervals.
 - accrualIntensity: A vector of accrual intensities. One for each accrual time interval.
 - piecewiseSurvivalTime: A vector that specifies the starting time of piecewise exponential survival time intervals.
 - gamma1: The hazard rate for exponential dropout or a vector of hazard rates for piecewise exponential dropout for the active treatment group.
 - gamma2: The hazard rate for exponential dropout or a vector of hazard rates for piecewise exponential dropout for the control group.
 - k: The number of postbaseline time points.
 - t: The postbaseline time points.
 - covar1: The covariance matrix for the repeated measures given baseline for the active treatment group.
 - covar2: The covariance matrix for the repeated measures given baseline for the control group.
 - normalApproximation: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution.
 - rounding: Whether to round up sample size.

Author(s)

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References

Kaifeng Lu, Xiaohui Luo, and Pei-Yun Chen. Sample size estimation for repeated measures analysis in randomized clinical trials with missing data. *The International Journal of Biostatistics* 2008; 14(1), Article 9.

Examples

```
# function to generate the AR(1) correlation matrix
ar1_cor <- function(n, corr) {
  exponent <- abs(matrix((1:n) - 1, n, n, byrow = TRUE) - ((1:n) - 1))
  corr^exponent
}

(design1 = getDesignMeanDiffMMRM(
  beta = 0.2,
  meanDiffH0 = 0,
  meanDiff = 0.5,
  k = 4,
  t = c(1,2,3,4),
  covar1 = ar1_cor(4, 0.7),
  accrualIntensity = 10,
  gamma1 = 0.02634013,
  gamma2 = 0.02634013,
  accrualDuration = NA,
  allocationRatioPlanned = 1,
  kMax = 3,
  alpha = 0.025,
  typeAlphaSpending = "sf0F"))
```

getDesignMeanDiffXO	<i>Group Sequential Design for Mean Difference in 2x2 Crossover</i>
---------------------	---

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for two-sample mean difference in 2x2 crossover.

Usage

```
getDesignMeanDiffXO(
  beta = NA_real_,
  n = NA_real_,
  meanDiffH0 = 0,
  meanDiff = 0.5,
  stDev = 1,
  allocationRatioPlanned = 1,
  normalApproximation = TRUE,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NA_real_,
```

```

alpha = 0.025,
typeAlphaSpending = "sfOF",
parameterAlphaSpending = NA_real_,
userAlphaSpending = NA_real_,
futilityBounds = NA_real_,
futilityCP = NA_real_,
futilityMeanDiff = NA_real_,
typeBetaSpending = "none",
parameterBetaSpending = NA_real_,
userBetaSpending = NA_real_,
spendingTime = NA_real_
)

```

Arguments

beta	The type II error.
n	The total sample size.
meanDiffH0	The mean difference under the null hypothesis. Defaults to 0.
meanDiff	The mean difference under the alternative hypothesis.
stDev	The standard deviation for within-subject random error.
allocationRatioPlanned	Allocation ratio for sequence A/B versus sequence B/A. Defaults to 1 for equal randomization.
normalApproximation	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
kMax	The maximum number of stages.
informationRates	The information rates. Fixed prior to the trial. Defaults to (1:kMax) / kMax if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".

parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	The futility boundary on the conditional power scale.
futilityMeanDiff	The futility boundary on the mean difference scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Value

An S3 class designMeanDiffXO object with three components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The overall significance level.
 - attainedAlpha: The attained significance level, which is different from the overall significance level in the presence of futility stopping.
 - kMax: The number of stages.
 - theta: The parameter value.
 - information: The maximum information.
 - expectedInformationH1: The expected information under H1.
 - expectedInformationH0: The expected information under H0.
 - drift: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
 - inflationFactor: The inflation factor (relative to the fixed design).
 - numberOfSubjects: The maximum number of subjects.
 - expectedNumberOfSubjectsH1: The expected number of subjects under H1.
 - expectedNumberOfSubjectsH0: The expected number of subjects under H0.
 - meanDiffH0: The mean difference under the null hypothesis.

- meanDiff: The mean difference under the alternative hypothesis.
- stDev: The standard deviation for within-subject random error.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale.
 - futilityBounds: The futility boundaries on the Z-scale.
 - rejectPerStage: The probability for efficacy stopping.
 - futilityPerStage: The probability for futility stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeFutility: The cumulative probability for futility stopping.
 - cumulativeAlphaSpent: The cumulative alpha spent.
 - efficacyP: The efficacy boundaries on the p-value scale.
 - futilityP: The futility boundaries on the p-value scale.
 - information: The cumulative information.
 - efficacyStopping: Whether to allow efficacy stopping.
 - futilityStopping: Whether to allow futility stopping.
 - rejectPerStageH0: The probability for efficacy stopping under H0.
 - futilityPerStageH0: The probability for futility stopping under H0.
 - cumulativeRejectionH0: The cumulative probability for efficacy stopping under H0.
 - cumulativeFutilityH0: The cumulative probability for futility stopping under H0.
 - efficacyMeanDiff: The efficacy boundaries on the mean difference scale.
 - futilityMeanDiff: The futility boundaries on the mean difference scale.
 - numberOfSubjects: The number of subjects.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - typeBetaSpending: The type of beta spending.
 - parameterBetaSpending: The parameter value for beta spending.
 - userBetaSpending: The user defined beta spending.
 - spendingTime: The error spending time at each analysis.
 - allocationRatioPlanned: Allocation ratio for sequence A/B versus sequence B/A.
 - normalApproximation: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution.
 - rounding: Whether to round up sample size.

Author(s)

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Examples

```
(design1 <- getDesignMeanDiffXO(
  beta = 0.2, n = NA, meanDiff = 75, stDev = 150,
  normalApproximation = FALSE, alpha = 0.05))
```

getDesignMeanDiffXOEquiv

Group Sequential Design for Equivalence in Mean Difference in 2x2 Crossover

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for equivalence in mean difference in 2x2 crossover.

Usage

```
getDesignMeanDiffXOEquiv(
  beta = NA_real_,
  n = NA_real_,
  meanDiffLower = NA_real_,
  meanDiffUpper = NA_real_,
  meanDiff = 0,
  stDev = 1,
  allocationRatioPlanned = 1,
  normalApproximation = TRUE,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  alpha = 0.05,
  typeAlphaSpending = "sf0F",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  spendingTime = NA_real_
)
```

Arguments

beta	The type II error.
n	The total sample size.
meanDiffLower	The lower equivalence limit of mean difference.
meanDiffUpper	The upper equivalence limit of mean difference.
meanDiff	The mean difference under the alternative hypothesis.
stDev	The standard deviation for within-subject random error.
allocationRatioPlanned	Allocation ratio for sequence A/B versus sequence B/A. Defaults to 1 for equal randomization.
normalApproximation	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.

rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
kMax	The maximum number of stages.
informationRates	The information rates. Fixed prior to the trial. Defaults to $(1:kMax) / kMax$ if left unspecified.
alpha	The significance level for each of the two one-sided tests. Defaults to 0.05.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Value

An S3 class designMeanDiffXOEquiv object with three components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The overall significance level.
 - attainedAlpha: The attained significance level.
 - kMax: The number of stages.
 - information: The maximum information.
 - expectedInformationH1: The expected information under H1.
 - expectedInformationH0: The expected information under H0.
 - numberOfSubjects: The maximum number of subjects.
 - expectedNumberOfSubjectsH1: The expected number of subjects under H1.
 - expectedNumberOfSubjectsH0: The expected number of subjects under H0.
 - meanDiffLower: The lower equivalence limit of mean difference.
 - meanDiffUpper: The upper equivalence limit of mean difference.
 - meanDiff: The mean difference under the alternative hypothesis.
 - stDev: The standard deviation for within-subject random error.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale for each of the two one-sided tests.

- rejectPerStage: The probability for efficacy stopping.
- cumulativeRejection: The cumulative probability for efficacy stopping.
- cumulativeAlphaSpent: The cumulative alpha for each of the two one-sided tests.
- cumulativeAttainedAlpha: The cumulative probability for efficacy stopping under H0.
- efficacyMeanDiffLower: The efficacy boundaries on the mean difference scale for the one-sided null hypothesis on the lower equivalence limit.
- efficacyMeanDiffUpper: The efficacy boundaries on the mean difference scale for the one-sided null hypothesis on the upper equivalence limit.
- efficacyP: The efficacy bounds on the p-value scale for each of the two one-sided tests.
- information: The cumulative information.
- numberOfSubjects: The number of subjects.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - spendingTime: The error spending time at each analysis.
 - allocationRatioPlanned: Allocation ratio for sequence A/B versus sequence B/A.
 - normalApproximation: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
 - rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Example 1: group sequential trial power calculation
(design1 <- getDesignMeanDiffXOEquiv(
  beta = 0.1, n = NA, meanDiffLower = -1.3, meanDiffUpper = 1.3,
  meanDiff = 0, stDev = 2.2,
  kMax = 4, alpha = 0.05, typeAlphaSpending = "sf0F"))

# Example 2: sample size calculation for t-test
(design2 <- getDesignMeanDiffXOEquiv(
  beta = 0.1, n = NA, meanDiffLower = -1.3, meanDiffUpper = 1.3,
  meanDiff = 0, stDev = 2.2,
  normalApproximation = FALSE, alpha = 0.05))
```

getDesignMeanRatio	<i>Group Sequential Design for Two-Sample Mean Ratio</i>
--------------------	--

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for two-sample mean ratio.

Usage

```
getDesignMeanRatio(
  beta = NA_real_,
  n = NA_real_,
  meanRatioH0 = 1,
  meanRatio = 1.25,
  CV = 1,
  allocationRatioPlanned = 1,
  normalApproximation = TRUE,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NA_real_,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NA_real_,
  futilityCP = NA_real_,
  futilityMeanRatio = NA_real_,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
  spendingTime = NA_real_
)
```

Arguments

beta	The type II error.
n	The total sample size.
meanRatioH0	The mean ratio under the null hypothesis. Defaults to 1.
meanRatio	The mean ratio under the alternative hypothesis.
CV	The coefficient of variation. The standard deviation on the log scale is equal to $\sqrt{\log(1 + CV^2)}$.

<code>allocationRatioPlanned</code>	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
<code>normalApproximation</code>	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
<code>rounding</code>	Whether to round up sample size. Defaults to 1 for sample size rounding.
<code>kMax</code>	The maximum number of stages.
<code>informationRates</code>	The information rates. Fixed prior to the trial. Defaults to $(1:kMax) / kMax$ if left unspecified.
<code>efficacyStopping</code>	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
<code>futilityStopping</code>	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
<code>criticalValues</code>	Upper boundaries on the z-test statistic scale for stopping for efficacy.
<code>alpha</code>	The significance level. Defaults to 0.025.
<code>typeAlphaSpending</code>	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
<code>parameterAlphaSpending</code>	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
<code>userAlphaSpending</code>	The user defined alpha spending. Cumulative alpha spent up to each stage.
<code>futilityBounds</code>	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., $kMax-1$. Defaults to $rep(-8, kMax-1)$ if left unspecified. The futility bounds are non-binding for the calculation of critical values.
<code>futilityCP</code>	The futility boundary on the conditional power scale.
<code>futilityMeanRatio</code>	The futility boundary on the mean ratio scale.
<code>typeBetaSpending</code>	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".

parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Value

An S3 class designMeanRatio object with three components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The overall significance level.
 - attainedAlpha: The attained significance level, which is different from the overall significance level in the presence of futility stopping.
 - kMax: The number of stages.
 - theta: The parameter value.
 - information: The maximum information.
 - expectedInformationH1: The expected information under H1.
 - expectedInformationH0: The expected information under H0.
 - drift: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
 - inflationFactor: The inflation factor (relative to the fixed design).
 - numberOfSubjects: The maximum number of subjects.
 - expectedNumberOfSubjectsH1: The expected number of subjects under H1.
 - expectedNumberOfSubjectsH0: The expected number of subjects under H0.
 - meanRatioH0: The mean ratio under the null hypothesis.
 - meanRatio: The mean ratio under the alternative hypothesis.
 - CV: The coefficient of variation.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale.
 - futilityBounds: The futility boundaries on the Z-scale.
 - rejectPerStage: The probability for efficacy stopping.
 - futilityPerStage: The probability for futility stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeFutility: The cumulative probability for futility stopping.
 - cumulativeAlphaSpent: The cumulative alpha spent.
 - efficacyP: The efficacy boundaries on the p-value scale.
 - futilityP: The futility boundaries on the p-value scale.
 - information: The cumulative information.
 - efficacyStopping: Whether to allow efficacy stopping.
 - futilityStopping: Whether to allow futility stopping.

- rejectPerStageH0: The probability for efficacy stopping under H0.
- futilityPerStageH0: The probability for futility stopping under H0.
- cumulativeRejectionH0: The cumulative probability for efficacy stopping under H0.
- cumulativeFutilityH0: The cumulative probability for futility stopping under H0.
- numberOfSubjects: The number of subjects.
- efficacyMeanRatio: The efficacy boundaries on the mean ratio scale.
- futilityMeanRatio: The futility boundaries on the mean ratio scale.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - typeBetaSpending: The type of beta spending.
 - parameterBetaSpending: The parameter value for beta spending.
 - userBetaSpending: The user defined beta spending.
 - spendingTime: The error spending time at each analysis.
 - allocationRatioPlanned: Allocation ratio for the active treatment versus control.
 - normalApproximation: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution.
 - rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignMeanRatio(
  beta = 0.1, n = NA, meanRatio = 1.25, CV = 0.25,
  alpha = 0.05, normalApproximation = FALSE))
```

getDesignMeanRatioEquiv

Group Sequential Design for Equivalence in Two-Sample Mean Ratio

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for equivalence in two-sample mean ratio.

Usage

```

getDesignMeanRatioEquiv(
  beta = NA_real_,
  n = NA_real_,
  meanRatioLower = NA_real_,
  meanRatioUpper = NA_real_,
  meanRatio = 1,
  CV = 1,
  allocationRatioPlanned = 1,
  normalApproximation = TRUE,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  alpha = 0.05,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  spendingTime = NA_real_
)

```

Arguments

beta	The type II error.
n	The total sample size.
meanRatioLower	The lower equivalence limit of mean ratio.
meanRatioUpper	The upper equivalence limit of mean ratio.
meanRatio	The mean ratio under the alternative hypothesis.
CV	The coefficient of variation.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
normalApproximation	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
kMax	The maximum number of stages.
informationRates	The information rates. Fixed prior to the trial. Defaults to (1:kMax) / kMax if left unspecified.
alpha	The significance level for each of the two one-sided tests. Defaults to 0.05.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type

	spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Value

An S3 class designMeanRatioEquiv object with three components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The significance level for each of the two one-sided tests. Defaults to 0.05.
 - attainedAlpha: The attained significance level.
 - kMax: The number of stages.
 - information: The maximum information.
 - expectedInformationH1: The expected information under H1.
 - expectedInformationH0: The expected information under H0.
 - numberOfSubjects: The maximum number of subjects.
 - expectedNumberOfSubjectsH1: The expected number of subjects under H1.
 - expectedNumberOfSubjectsH0: The expected number of subjects under H0.
 - meanRatioLower: The lower equivalence limit of mean ratio.
 - meanRatioUpper: The upper equivalence limit of mean ratio.
 - meanRatio: The mean ratio under the alternative hypothesis.
 - CV: The coefficient of variation.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale for each of the two one-sided tests.
 - rejectPerStage: The probability for efficacy stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeAlphaSpent: The cumulative alpha for each of the two one-sided tests.
 - cumulativeAttainedAlpha: The cumulative probability for efficacy stopping under H0.
 - efficacyP: The efficacy bounds on the p-value scale for each of the two one-sided tests.
 - information: The cumulative information.
 - numberOfSubjects: The number of subjects.
 - efficacyMeanRatioLower: The efficacy boundaries on the mean ratio scale for the one-sided null hypothesis on the lower equivalence limit.

- efficacyMeanRatioUpper: The efficacy boundaries on the mean ratio scale for the one-sided null hypothesis on the upper equivalence limit.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - spendingTime: The error spending time at each analysis.
 - allocationRatioPlanned: Allocation ratio for the active treatment versus control.
 - normalApproximation: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
 - rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Example 1: group sequential trial power calculation
(design1 <- getDesignMeanRatioEquiv(
  beta = 0.1, n = NA, meanRatioLower = 0.8, meanRatioUpper = 1.25,
  meanRatio = 1, CV = 0.35,
  kMax = 4, alpha = 0.05, typeAlphaSpending = "sf0F"))

# Example 2: sample size calculation for t-test
(design2 <- getDesignMeanRatioEquiv(
  beta = 0.1, n = NA, meanRatioLower = 0.8, meanRatioUpper = 1.25,
  meanRatio = 1, CV = 0.35,
  normalApproximation = FALSE, alpha = 0.05))
```

getDesignMeanRatioXO *Group Sequential Design for Mean Ratio in 2x2 Crossover*

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for two-sample mean ratio in 2x2 crossover.

Usage

```
getDesignMeanRatioXO(
  beta = NA_real_,
  n = NA_real_,
  meanRatioH0 = 1,
  meanRatio = 1.25,
```

```

CV = 1,
allocationRatioPlanned = 1,
normalApproximation = TRUE,
rounding = TRUE,
kMax = 1L,
informationRates = NA_real_,
efficacyStopping = NA_integer_,
futilityStopping = NA_integer_,
criticalValues = NA_real_,
alpha = 0.025,
typeAlphaSpending = "sf0F",
parameterAlphaSpending = NA_real_,
userAlphaSpending = NA_real_,
futilityBounds = NA_real_,
futilityCP = NA_real_,
futilityMeanRatio = NA_real_,
typeBetaSpending = "none",
parameterBetaSpending = NA_real_,
userBetaSpending = NA_real_,
spendingTime = NA_real_
)

```

Arguments

beta	The type II error.
n	The total sample size.
meanRatioH0	The mean ratio under the null hypothesis. Defaults to 1.
meanRatio	The mean ratio under the alternative hypothesis.
CV	The coefficient of variation. The standard deviation on the log scale is equal to $\sqrt{\log(1 + CV^2)}$.
allocationRatioPlanned	Allocation ratio for sequence A/B versus sequence B/A. Defaults to 1 for equal randomization.
normalApproximation	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
kMax	The maximum number of stages.
informationRates	The information rates. Fixed prior to the trial. Defaults to $(1:kMax) / kMax$ if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.

futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	The futility boundary on the conditional power scale.
futilityMeanRatio	The futility boundary on the mean ratio scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Value

An S3 class designMeanRatioXO object with three components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The overall significance level.
 - attainedAlpha: The attained significance level, which is different from the overall significance level in the presence of futility stopping.

- kMax: The number of stages.
- theta: The parameter value.
- information: The maximum information.
- expectedInformationH1: The expected information under H1.
- expectedInformationH0: The expected information under H0.
- drift: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
- inflationFactor: The inflation factor (relative to the fixed design).
- numberOfSubjects: The maximum number of subjects.
- expectedNumberOfSubjectsH1: The expected number of subjects under H1.
- expectedNumberOfSubjectsH0: The expected number of subjects under H0.
- meanRatioH0: The mean ratio under the null hypothesis.
- meanRatio: The mean ratio under the alternative hypothesis.
- CV: The coefficient of variation.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale.
 - futilityBounds: The futility boundaries on the Z-scale.
 - rejectPerStage: The probability for efficacy stopping.
 - futilityPerStage: The probability for futility stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeFutility: The cumulative probability for futility stopping.
 - cumulativeAlphaSpent: The cumulative alpha spent.
 - efficacyMeanRatio: The efficacy boundaries on the mean ratio scale.
 - futilityMeanRatio: The futility boundaries on the mean ratio scale.
 - efficacyP: The efficacy boundaries on the p-value scale.
 - futilityP: The futility boundaries on the p-value scale.
 - information: The cumulative information.
 - efficacyStopping: Whether to allow efficacy stopping.
 - futilityStopping: Whether to allow futility stopping.
 - rejectPerStageH0: The probability for efficacy stopping under H0.
 - futilityPerStageH0: The probability for futility stopping under H0.
 - cumulativeRejectionH0: The cumulative probability for efficacy stopping under H0.
 - cumulativeFutilityH0: The cumulative probability for futility stopping under H0.
 - numberOfSubjects: The number of subjects.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - typeBetaSpending: The type of beta spending.
 - parameterBetaSpending: The parameter value for beta spending.
 - userBetaSpending: The user defined beta spending.

- spendingTime: The error spending time at each analysis.
- allocationRatioPlanned: Allocation ratio for sequence A/B versus sequence B/A.
- normalApproximation: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution.
- rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignMeanRatioXO(
  beta = 0.1, n = NA, meanRatio = 1.25, CV = 0.25,
  alpha = 0.05, normalApproximation = FALSE))
```

getDesignMeanRatioXOEquiv

Group Sequential Design for Equivalence in Mean Ratio in 2x2 Crossover

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for equivalence mean ratio in 2x2 crossover.

Usage

```
getDesignMeanRatioXOEquiv(
  beta = NA_real_,
  n = NA_real_,
  meanRatioLower = NA_real_,
  meanRatioUpper = NA_real_,
  meanRatio = 1,
  CV = 1,
  allocationRatioPlanned = 1,
  normalApproximation = TRUE,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  alpha = 0.05,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  spendingTime = NA_real_
)
```

Arguments

<code>beta</code>	The type II error.
<code>n</code>	The total sample size.
<code>meanRatioLower</code>	The lower equivalence limit of mean ratio.
<code>meanRatioUpper</code>	The upper equivalence limit of mean ratio.
<code>meanRatio</code>	The mean ratio under the alternative hypothesis.
<code>CV</code>	The coefficient of variation.
<code>allocationRatioPlanned</code>	Allocation ratio for sequence A/B versus sequence B/A. Defaults to 1 for equal randomization.
<code>normalApproximation</code>	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
<code>rounding</code>	Whether to round up sample size. Defaults to 1 for sample size rounding.
<code>kMax</code>	The maximum number of stages.
<code>informationRates</code>	The information rates. Fixed prior to the trial. Defaults to $(1:kMax) / kMax$ if left unspecified.
<code>alpha</code>	The significance level for each of the two one-sided tests. Defaults to 0.05.
<code>typeAlphaSpending</code>	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
<code>parameterAlphaSpending</code>	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
<code>userAlphaSpending</code>	The user defined alpha spending. Cumulative alpha spent up to each stage.
<code>spendingTime</code>	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as <code>informationRates</code> .

Value

An S3 class `designMeanRatioEquiv` object with three components:

- `overallResults`: A data frame containing the following variables:
 - `overallReject`: The overall rejection probability.
 - `alpha`: The overall significance level.
 - `attainedAlpha`: The attained significance level.
 - `kMax`: The number of stages.

- information: The maximum information.
 - expectedInformationH1: The expected information under H1.
 - expectedInformationH0: The expected information under H0.
 - numberOfSubjects: The maximum number of subjects.
 - expectedNumberOfSubjectsH1: The expected number of subjects under H1.
 - expectedNumberOfSubjectsH0: The expected number of subjects under H0.
 - meanRatioLower: The lower equivalence limit of mean ratio.
 - meanRatioUpper: The upper equivalence limit of mean ratio.
 - meanRatio: The mean ratio under the alternative hypothesis.
 - CV: The coefficient of variation.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale for each of the two one-sided tests.
 - rejectPerStage: The probability for efficacy stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeAlphaSpent: The cumulative alpha for each of the two one-sided tests.
 - cumulativeAttainedAlpha: The cumulative probability for efficacy stopping under H0.
 - efficacyMeanRatioLower: The efficacy boundaries on the mean ratio scale for the one-sided null hypothesis on the lower equivalence limit.
 - efficacyMeanRatioUpper: The efficacy boundaries on the mean ratio scale for the one-sided null hypothesis on the upper equivalence limit.
 - efficacyP: The efficacy bounds on the p-value scale for each of the two one-sided tests.
 - information: The cumulative information.
 - numberOfSubjects: The number of subjects.
 - settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - spendingTime: The error spending time at each analysis.
 - allocationRatioPlanned: Allocation ratio for sequence A/B versus sequence B/A.
 - normalApproximation: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
 - rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
# Example 1: group sequential trial power calculation
(design1 <- getDesignMeanRatioXOEquiv(
  beta = 0.1, n = NA, meanRatioLower = 0.8, meanRatioUpper = 1.25,
  meanRatio = 1, CV = 0.35,
  kMax = 4, alpha = 0.05, typeAlphaSpending = "sfOF"))

# Example 2: sample size calculation for t-test
(design2 <- getDesignMeanRatioXOEquiv(
  beta = 0.1, n = NA, meanRatioLower = 0.8, meanRatioUpper = 1.25,
  meanRatio = 1, CV = 0.35,
  normalApproximation = FALSE, alpha = 0.05))
```

getDesignOddsRatio	<i>Group Sequential Design for Two-Sample Odds Ratio</i>
--------------------	--

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for two-sample odds ratio.

Usage

```
getDesignOddsRatio(
  beta = NA_real_,
  n = NA_real_,
  oddsRatioH0 = 1,
  pi1 = NA_real_,
  pi2 = NA_real_,
  nullVariance = FALSE,
  allocationRatioPlanned = 1,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NA_real_,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NA_real_,
  futilityCP = NA_real_,
  futilityOddsRatio = NA_real_,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
```

```

    userBetaSpending = NA_real_,
    spendingTime = NA_real_
)

```

Arguments

beta	The type II error.
n	The total sample size.
oddsRatioH0	The odds ratio under the null hypothesis. Defaults to 1.
pi1	The assumed probability for the active treatment group.
pi2	The assumed probability for the control group.
nullVariance	Whether to use the variance under the null or the empirical variance under the alternative.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
kMax	The maximum number of stages.
informationRates	The information rates. Fixed prior to the trial. Defaults to $(1:kMax) / kMax$ if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.

futilityCP	The futility bounds on the conditional power scale.
futilityOddsRatio	The futility bounds on the odds ratio scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Details

Consider a group sequential design for two-sample odds ratio. The parameter of interest is

$$\psi = \frac{\pi_1(1 - \pi_2)}{(1 - \pi_1)\pi_2}$$

where π_1 is the response probability for the active treatment group and π_2 is the response probability for the control group. For statistical inference, the parameter is often transformed to the log scale:

$$\theta = \log(\psi) = \log(\pi_1/(1 - \pi_1)) - \log(\pi_2/(1 - \pi_2))$$

The variance of the estimator $\hat{\theta}$ can be derived from the binomial distributions as follows:

$$Var(\hat{\theta}) = \frac{1}{n} \left\{ \frac{1}{\pi_1(1 - \pi_1)r} + \frac{1}{\pi_2(1 - \pi_2)(1 - r)} \right\}$$

where n is the total number of subjects and r is the randomization probability for the active treatment group. When nullVariance = TRUE, the variance is computed under the null hypothesis. In this case, the values of π_1 and π_2 in the variance formula are replaced with their restricted maximum likelihood counterparts, subject to the constraint

$$\frac{\pi_1(1 - \pi_2)}{(1 - \pi_1)\pi_2} = \psi_0$$

Value

An S3 class designOddsRatio object with three components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The overall significance level.

- attainedAlpha: The attained significance level, which is different from the overall significance level in the presence of futility stopping.
- kMax: The number of stages.
- theta: The parameter value.
- information: The maximum information.
- expectedInformationH1: The expected information under H1.
- expectedInformationH0: The expected information under H0.
- drift: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
- inflationFactor: The inflation factor (relative to the fixed design).
- numberOfSubjects: The maximum number of subjects.
- expectedNumberOfSubjectsH1: The expected number of subjects under H1.
- expectedNumberOfSubjectsH0: The expected number of subjects under H0.
- oddsRatioH0: The odds ratio under the null hypothesis.
- pi1: The assumed probability for the active treatment group.
- pi2: The assumed probability for the control group.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale.
 - futilityBounds: The futility boundaries on the Z-scale.
 - rejectPerStage: The probability for efficacy stopping.
 - futilityPerStage: The probability for futility stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeFutility: The cumulative probability for futility stopping.
 - cumulativeAlphaSpent: The cumulative alpha spent.
 - efficacyP: The efficacy boundaries on the p-value scale.
 - futilityP: The futility boundaries on the p-value scale.
 - information: The cumulative information.
 - efficacyStopping: Whether to allow efficacy stopping.
 - futilityStopping: Whether to allow futility stopping.
 - rejectPerStageH0: The probability for efficacy stopping under H0.
 - futilityPerStageH0: The probability for futility stopping under H0.
 - cumulativeRejectionH0: The cumulative probability for efficacy stopping under H0.
 - cumulativeFutilityH0: The cumulative probability for futility stopping under H0.
 - efficacyOddsRatio: The efficacy boundaries on the odds ratio scale.
 - futilityOddsRatio: The futility boundaries on the odds ratio scale.
 - numberOfSubjects: The number of subjects.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - typeBetaSpending: The type of beta spending.
 - parameterBetaSpending: The parameter value for beta spending.

- userBetaSpending: The user defined beta spending.
- spendingTime: The error spending time at each analysis.
- varianceRatio: The ratio of the variance under H0 to the variance under H1.
- nullVariance: Whether to use the variance under the null or the empirical variance under the alternative.
- allocationRatioPlanned: Allocation ratio for the active treatment versus control.
- rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignOddsRatio(
  beta = 0.1, n = NA, pi1 = 0.5, pi2 = 0.3,
  alpha = 0.05))
```

getDesignOddsRatioEquiv

Group Sequential Design for Equivalence in Two-Sample Odds Ratio

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for equivalence in two-sample odds ratio.

Usage

```
getDesignOddsRatioEquiv(
  beta = NA_real_,
  n = NA_real_,
  oddsRatioLower = NA_real_,
  oddsRatioUpper = NA_real_,
  pi1 = NA_real_,
  pi2 = NA_real_,
  allocationRatioPlanned = 1,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  criticalValues = NA_real_,
  alpha = 0.05,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  spendingTime = NA_real_
)
```

Arguments

beta	The type II error.
n	The total sample size.
oddsRatioLower	The lower equivalence limit of odds ratio.
oddsRatioUpper	The upper equivalence limit of odds ratio.
pi1	The assumed probability for the active treatment group.
pi2	The assumed probability for the control group.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
kMax	The maximum number of stages.
informationRates	The information rates. Fixed prior to the trial. Defaults to (1:kMax) / kMax if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level for each of the two one-sided tests. Defaults to 0.05.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Value

An S3 class designOddsRatioEquiv object with three components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The significance level for each of the two one-sided tests. Defaults to 0.05.
 - attainedAlphaH10: The attained significance level under H10.
 - attainedAlphaH20: The attained significance level under H20.
 - kMax: The number of stages.
 - information: The maximum information.

- expectedInformationH1: The expected information under H1.
- expectedInformationH10: The expected information under H10.
- expectedInformationH20: The expected information under H20.
- numberOfSubjects: The maximum number of subjects.
- expectedNumberOfSubjectsH1: The expected number of subjects under H1.
- expectedNumberOfSubjectsH10: The expected number of subjects under H10.
- expectedNumberOfSubjectsH20: The expected number of subjects under H20.
- oddsRatioLower: The lower equivalence limit of odds ratio.
- oddsRatioUpper: The upper equivalence limit of odds ratio.
- pi1: The assumed probability for the active treatment group.
- pi2: The assumed probability for the control group.
- oddsRatio: The odds ratio.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale for each of the two one-sided tests.
 - rejectPerStage: The probability for efficacy stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeAlphaSpent: The cumulative alpha for each of the two one-sided tests.
 - cumulativeAttainedAlphaH10: The cumulative alpha attained under H10.
 - cumulativeAttainedAlphaH20: The cumulative alpha attained under H20.
 - efficacyOddsRatioLower: The efficacy boundaries on the odds ratio scale for the one-sided null hypothesis on the lower equivalence limit.
 - efficacyOddsRatioUpper: The efficacy boundaries on the odds ratio scale for the one-sided null hypothesis on the upper equivalence limit.
 - efficacyP: The efficacy bounds on the p-value scale for each of the two one-sided tests.
 - information: The cumulative information.
 - numberOfSubjects: The number of subjects.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - spendingTime: The error spending time at each analysis.
 - allocationRatioPlanned: Allocation ratio for the active treatment versus control.
 - rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignOddsRatioEquiv(
  beta = 0.2, n = NA, oddsRatioLower = 0.8,
  oddsRatioUpper = 1.25, pi1 = 0.12, pi2 = 0.12,
  kMax = 3, alpha = 0.05, typeAlphaSpending = "sfOF"))
```

getDesignOneMean	<i>Group Sequential Design for One-Sample Mean</i>
------------------	--

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for one-sample mean.

Usage

```
getDesignOneMean(
  beta = NA_real_,
  n = NA_real_,
  meanH0 = 0,
  mean = 0.5,
  stDev = 1,
  normalApproximation = TRUE,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NA_real_,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NA_real_,
  futilityCP = NA_real_,
  futilityMean = NA_real_,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
  spendingTime = NA_real_
)
```

Arguments

beta	The type II error.
n	The total sample size.

meanH0	The mean under the null hypothesis. Defaults to 0.
mean	The mean under the alternative hypothesis.
stDev	The standard deviation.
normalApproximation	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
kMax	The maximum number of stages.
informationRates	The information rates. Fixed prior to the trial. Defaults to $(1:kMax) / kMax$ if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	The futility boundary on the conditional power scale.
futilityMean	The futility boundary on the mean scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".

parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Value

An S3 class designOneMean object with three components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The overall significance level.
 - attainedAlpha: The attained significance level, which is different from the overall significance level in the presence of futility stopping.
 - kMax: The number of stages.
 - theta: The parameter value.
 - information: The maximum information.
 - expectedInformationH1: The expected information under H1.
 - expectedInformationH0: The expected information under H0.
 - drift: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
 - inflationFactor: The inflation factor (relative to the fixed design).
 - numberOfSubjects: The maximum number of subjects.
 - expectedNumberOfSubjectsH1: The expected number of subjects under H1.
 - expectedNumberOfSubjectsH0: The expected number of subjects under H0.
 - meanH0: The mean under the null hypothesis.
 - mean: The mean under the alternative hypothesis.
 - stDev: The standard deviation.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale.
 - futilityBounds: The futility boundaries on the Z-scale.
 - rejectPerStage: The probability for efficacy stopping.
 - futilityPerStage: The probability for futility stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeFutility: The cumulative probability for futility stopping.
 - cumulativeAlphaSpent: The cumulative alpha spent.
 - efficacyP: The efficacy boundaries on the p-value scale.
 - futilityP: The futility boundaries on the p-value scale.
 - information: The cumulative information.
 - efficacyStopping: Whether to allow efficacy stopping.
 - futilityStopping: Whether to allow futility stopping.

- rejectPerStageH0: The probability for efficacy stopping under H0.
- futilityPerStageH0: The probability for futility stopping under H0.
- cumulativeRejectionH0: The cumulative probability for efficacy stopping under H0.
- cumulativeFutilityH0: The cumulative probability for futility stopping under H0.
- efficacyMean: The efficacy boundaries on the mean scale.
- futilityMean: The futility boundaries on the mean scale.
- numberOfSubjects: The number of subjects.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - typeBetaSpending: The type of beta spending.
 - parameterBetaSpending: The parameter value for beta spending.
 - userBetaSpending: The user defined beta spending.
 - spendingTime: The error spending time at each analysis.
 - normalApproximation: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution.
 - rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
# Example 1: group sequential trial power calculation
(design1 <- getDesignOneMean(
  beta = 0.1, n = NA, meanH0 = 7, mean = 6, stDev = 2.5,
  kMax = 5, alpha = 0.025, typeAlphaSpending = "sfOF",
  typeBetaSpending = "sfP"))

# Example 2: sample size calculation for one-sample t-test
(design2 <- getDesignOneMean(
  beta = 0.1, n = NA, meanH0 = 7, mean = 6, stDev = 2.5,
  normalApproximation = FALSE, alpha = 0.025))
```

getDesignOneMultinom *Power and Sample Size for One-Sample Multinomial Response*

Description

Obtains the power given sample size or obtains the sample size given power for one-sample multinomial response.

Usage

```
getDesignOneMultinom(
  beta = NA_real_,
  n = NA_real_,
  ncats = NA_integer_,
  piH0 = NA_real_,
  pi = NA_real_,
  rounding = TRUE,
  alpha = 0.05
)
```

Arguments

beta	The type II error.
n	The total sample size.
ncats	The number of categories of the multinomial response.
piH0	The prevalence of each category under the null hypothesis. Only need to provide the values for the first ncats-1 categories.
pi	The prevalence of each category. Only need to provide the values for the first ncats-1 categories.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
alpha	The two-sided significance level. Defaults to 0.05.

Details

A single-arm multinomial response design is used to test whether the prevalence of each category is different from the null hypothesis prevalence. The null hypothesis is that the prevalence of each category is equal to π_{0i} , while the alternative hypothesis is that the prevalence of each category is equal to π_i , for $i = 1, \dots, C$, where C is the number of categories.

The sample size is calculated based on the chi-square test for multinomial response. The test statistic is given by

$$X^2 = \sum_{i=1}^C \frac{(n_i - n\pi_{0i})^2}{n\pi_{0i}}$$

where n_i is the number of subjects in category i , and n is the total sample size.

- Under the null hypothesis, X^2 follows a chi-square distribution with $C - 1$ degrees of freedom.
- Under the alternative hypothesis, X^2 follows a non-central chi-square distribution with non-centrality parameter

$$\lambda = n \sum_{i=1}^C \frac{(\pi_i - \pi_{0i})^2}{\pi_{0i}}$$

The sample size is chosen such that the power to reject the null hypothesis is at least $1 - \beta$ for a given significance level α .

Value

An S3 class designOneMultinom object with the following components:

- power: The power to reject the null hypothesis.
- alpha: The two-sided significance level.
- n: The maximum number of subjects.
- ncats: The number of categories of the multinomial response.
- piH0: The prevalence of each category under the null hypothesis.
- pi: The prevalence of each category.
- effectsize: The effect size for the chi-square test.
- rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignOneMultinom(
  beta = 0.1, ncats = 3, piH0 = c(0.25, 0.25),
  pi = c(0.3, 0.4), alpha = 0.05))
```

getDesignOneProportion

Group Sequential Design for One-Sample Proportion

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for one-sample proportion.

Usage

```
getDesignOneProportion(
  beta = NA_real_,
  n = NA_real_,
  piH0 = 0.1,
  pi = 0.2,
  nullVariance = TRUE,
  normalApproximation = TRUE,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
```

```

    futilityStopping = NA_integer_,
    criticalValues = NA_real_,
    alpha = 0.025,
    typeAlphaSpending = "sfOF",
    parameterAlphaSpending = NA_real_,
    userAlphaSpending = NA_real_,
    futilityBounds = NA_real_,
    futilityCP = NA_real_,
    futilitypi = NA_real_,
    typeBetaSpending = "none",
    parameterBetaSpending = NA_real_,
    userBetaSpending = NA_real_,
    spendingTime = NA_real_
  )

```

Arguments

beta	The type II error.
n	The total sample size.
piH0	The response probability under the null hypothesis.
pi	The response probability under the alternative hypothesis.
nullVariance	Whether to use the variance under the null or the variance under the alternative.
normalApproximation	The type of computation of the p-values. If TRUE, the normal approximation will be used, otherwise the calculations are performed with the binomial distribution. The exact calculation using the binomial distribution is only implemented for the fixed design.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
kMax	The maximum number of stages.
informationRates	The information rates. Fixed prior to the trial. Defaults to (1:kMax) / kMax if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".

parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	The futility bounds on the conditional power scale.
futilitypi	The futility bounds on the response probability scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Value

An S3 class designOneProportion object with three components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The overall significance level.
 - attainedAlpha: The attained significance level, which is different from the overall significance level in the presence of futility stopping as well as for the binomial exact test in a fixed design.
 - kMax: The number of stages.
 - theta: The parameter value.
 - information: The maximum information.
 - expectedInformationH1: The expected information under H1.
 - expectedInformationH0: The expected information under H0.
 - drift: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
 - inflationFactor: The inflation factor (relative to the fixed design).
 - numberOfSubjects: The maximum number of subjects.
 - expectedNumberOfSubjectsH1: The expected number of subjects under H1.
 - expectedNumberOfSubjectsH0: The expected number of subjects under H0.
 - piH0: The response probability under the null hypothesis.

- pi: The response probability under the alternative hypothesis.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale.
 - futilityBounds: The futility boundaries on the Z-scale.
 - rejectPerStage: The probability for efficacy stopping.
 - futilityPerStage: The probability for futility stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeFutility: The cumulative probability for futility stopping.
 - cumulativeAlphaSpent: The cumulative alpha spent.
 - efficacyP: The efficacy boundaries on the p-value scale.
 - futilityP: The futility boundaries on the p-value scale.
 - information: The cumulative information.
 - efficacyStopping: Whether to allow efficacy stopping.
 - futilityStopping: Whether to allow futility stopping.
 - rejectPerStageH0: The probability for efficacy stopping under H0.
 - futilityPerStageH0: The probability for futility stopping under H0.
 - cumulativeRejectionH0: The cumulative probability for efficacy stopping under H0.
 - cumulativeFutilityH0: The cumulative probability for futility stopping under H0.
 - efficacyResponses: The efficacy boundaries on the number of responses scale.
 - futilityResponses: The futility boundaries on the number of responses scale.
 - numberOfSubjects: The number of subjects.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - typeBetaSpending: The type of beta spending.
 - parameterBetaSpending: The parameter value for beta spending.
 - userBetaSpending: The user defined beta spending.
 - spendingTime: The error spending time at each analysis.
 - varianceRatio: The ratio of the variance under H0 to the variance under H1.
 - nullVariance: Whether to use the variance under the null or the empirical variance under the alternative.
 - normalApproximation: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the binomial distribution.
 - rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Example 1: group sequential trial power calculation
(design1 <- getDesignOneProportion(
  beta = 0.2, n = NA, piH0 = 0.15, pi = 0.25,
  kMax = 3, alpha = 0.05, typeAlphaSpending = "sf0F"))

# Example 2: sample size calculation for one-sample binomial exact test
(design2 <- getDesignOneProportion(
  beta = 0.2, n = NA, piH0 = 0.15, pi = 0.25,
  normalApproximation = FALSE, alpha = 0.05))
```

getDesignOneRateExact *Power and Sample Size for One-Sample Poisson Rate Exact Test*

Description

Obtains the power given sample size or obtains the sample size given power for one-sample Poisson rate.

Usage

```
getDesignOneRateExact(
  beta = NA_real_,
  n = NA_real_,
  lambdaH0 = NA_real_,
  lambda = NA_real_,
  D = 1,
  alpha = 0.025
)
```

Arguments

beta	The type II error.
n	The total sample size.
lambdaH0	The Poisson rate under the null hypothesis.
lambda	The Poisson rate under the alternative hypothesis.
D	The average exposure per subject.
alpha	The one-sided significance level. Defaults to 0.025.

Value

A data frame containing the following variables:

- alpha: The specified significance level.
- attainedAlpha: The attained type I error of the exact test.

- power: The actual power of the exact test.
- n: The sample size.
- lambdaH0: The Poisson rate under the null hypothesis.
- lambda: The Poisson rate under the alternative hypothesis.
- D: The average exposure per subject.
- r: The critical value of the number of events for rejecting the null hypothesis. Reject H0 if $Y \geq r$ for upper-tailed test, and reject H0 if $Y \leq r$ for lower-tailed test.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
# Example 1: power calculation
(design1 <- getDesignOneRateExact(
  n = 525, lambdaH0 = 0.049, lambda = 0.012,
  D = 0.5, alpha = 0.025))

# Example 2: sample size calculation
(design2 <- getDesignOneRateExact(
  beta = 0.2, lambdaH0 = 0.2, lambda = 0.3,
  D = 1, alpha = 0.05))
```

getDesignOneSlope

Group Sequential Design for One-Sample Slope

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for one-sample slope.

Usage

```
getDesignOneSlope(
  beta = NA_real_,
  n = NA_real_,
  slopeH0 = 0,
  slope = 0.5,
  stDev = 1,
  stDevCovariate = 1,
  normalApproximation = TRUE,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
```

```

futilityStopping = NA_integer_,
criticalValues = NA_real_,
alpha = 0.025,
typeAlphaSpending = "sfOF",
parameterAlphaSpending = NA_real_,
userAlphaSpending = NA_real_,
futilityBounds = NA_real_,
futilityCP = NA_real_,
futilitySlope = NA_real_,
typeBetaSpending = "none",
parameterBetaSpending = NA_real_,
userBetaSpending = NA_real_,
spendingTime = NA_real_
)

```

Arguments

beta	The type II error.
n	The total sample size.
slopeH0	The slope under the null hypothesis. Defaults to 0.
slope	The slope under the alternative hypothesis.
stDev	The standard deviation of the residual.
stDevCovariate	The standard deviation of the covariate.
normalApproximation	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
kMax	The maximum number of stages.
informationRates	The information rates. Fixed prior to the trial. Defaults to (1:kMax) / kMax if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for

	Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	The futility bounds on the conditional power scale.
futilitySlope	The futility bounds on the slope scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Details

We assume a simple linear regression of the form

$$y_i = \alpha + \beta x_i + \epsilon_i$$

where ϵ_i is the residual error, which is assumed to be normally distributed with mean 0 and standard deviation σ_ϵ . The covariate x_i is assumed to be normally distributed with mean 0 and standard deviation σ_x . The slope under the null hypothesis is β_0 , and the slope under the alternative hypothesis is β . Since

$$\hat{\beta} = \frac{\sum_{i=1}^n (x_i - \bar{x}) y_i}{\sum_{i=1}^n (x_i - \bar{x})^2}$$

it follows that

$$\hat{\beta} \sim N\left(\beta, \frac{\sigma_\epsilon^2}{\sum_{i=1}^n (x_i - \bar{x})^2}\right).$$

Since the variance of $\hat{\beta}$ is

$$\frac{\sigma_\epsilon^2}{n\sigma_x^2}$$

we can use it to calculate the power and sample size for the group sequential design.

Value

An S3 class `designOneSlope` object with three components:

- `overallResults`: A data frame containing the following variables:
 - `overallReject`: The overall rejection probability.
 - `alpha`: The overall significance level.
 - `attainedAlpha`: The attained significance level, which is different from the overall significance level in the presence of futility stopping.
 - `kMax`: The number of stages.
 - `theta`: The parameter value.
 - `information`: The maximum information.
 - `expectedInformationH1`: The expected information under H1.
 - `expectedInformationH0`: The expected information under H0.
 - `drift`: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
 - `inflationFactor`: The inflation factor (relative to the fixed design).
 - `numberOfSubjects`: The maximum number of subjects.
 - `expectedNumberOfSubjectsH1`: The expected number of subjects under H1.
 - `expectedNumberOfSubjectsH0`: The expected number of subjects under H0.
 - `slopeH0`: The slope under the null hypothesis.
 - `slope`: The slope under the alternative hypothesis.
 - `stDev`: The standard deviation of the residual.
 - `stDevCovariate`: The standard deviation of the covariate.
- `byStageResults`: A data frame containing the following variables:
 - `informationRates`: The information rates.
 - `efficacyBounds`: The efficacy boundaries on the Z-scale.
 - `futilityBounds`: The futility boundaries on the Z-scale.
 - `rejectPerStage`: The probability for efficacy stopping.
 - `futilityPerStage`: The probability for futility stopping.
 - `cumulativeRejection`: The cumulative probability for efficacy stopping.
 - `cumulativeFutility`: The cumulative probability for futility stopping.
 - `cumulativeAlphaSpent`: The cumulative alpha spent.
 - `efficacyP`: The efficacy boundaries on the p-value scale.
 - `futilityP`: The futility boundaries on the p-value scale.
 - `information`: The cumulative information.
 - `efficacyStopping`: Whether to allow efficacy stopping.
 - `futilityStopping`: Whether to allow futility stopping.
 - `rejectPerStageH0`: The probability for efficacy stopping under H0.
 - `futilityPerStageH0`: The probability for futility stopping under H0.
 - `cumulativeRejectionH0`: The cumulative probability for efficacy stopping under H0.
 - `cumulativeFutilityH0`: The cumulative probability for futility stopping under H0.
 - `efficacySlope`: The efficacy boundaries on the slope scale.
 - `futilitySlope`: The futility boundaries on the slope scale.

- numberOfSubjects: The number of subjects.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - typeBetaSpending: The type of beta spending.
 - parameterBetaSpending: The parameter value for beta spending.
 - userBetaSpending: The user defined beta spending.
 - spendingTime: The error spending time at each analysis.
 - normalApproximation: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution.
 - rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignOneSlope(
  beta = 0.1, n = NA, slope = 0.5,
  stDev = 15, stDevCovariate = 9,
  normalApproximation = FALSE,
  alpha = 0.025))
```

getDesignOrderedBinom *Power and Sample Size for Cochran-Armitage Trend Test for Ordered Multi-Sample Binomial Response*

Description

Obtains the power given sample size or obtains the sample size given power for the Cochran-Armitage trend test for ordered multi-sample binomial response.

Usage

```
getDesignOrderedBinom(
  beta = NA_real_,
  n = NA_real_,
  ngroups = NA_integer_,
  pi = NA_real_,
  w = NA_real_,
  allocationRatioPlanned = NA_integer_,
  rounding = TRUE,
  alpha = 0.05
)
```

Arguments

beta	The type II error.
n	The total sample size.
ngroups	The number of treatment groups.
pi	The response probabilities for the treatment groups.
w	The scores assigned to the treatment groups. This should reflect the ordinal nature of the treatment groups, e.g. dose levels. Defaults to equally spaced scores.
allocationRatioPlanned	Allocation ratio for the treatment groups.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
alpha	The two-sided significance level. Defaults to 0.05.

Details

An ordered multi-sample binomial response design is used to test whether the response probabilities differ across multiple treatment groups. The null hypothesis is that the response probabilities are equal across all treatment groups, while the alternative hypothesis is that the response probabilities are ordered, i.e. the response probability increases with the treatment group index. The Cochran-Armitage trend test is used to test this hypothesis. This test effectively regresses the response probabilities against treatment group scores, and test whether the slope of the regression line is significantly different from zero.

The trend parameter is defined as

$$\theta = \sum_{g=1}^G r_g (w_g - \bar{w}) \pi_g$$

where G is the number of treatment groups, r_g is the randomization probability for treatment group g , w_g is the score assigned to treatment group g , π_g is the response probability for treatment group g , and $\bar{w} = \sum_{g=1}^G r_g w_g$ is the weighted average score across all treatment groups.

Since $\hat{\theta}$ is a linear combination of the estimated response probabilities, its variance is given by

$$Var(\hat{\theta}) = \frac{1}{n} \sum_{g=1}^G r_g (w_g - \bar{w})^2 \pi_g (1 - \pi_g)$$

where n is the total sample size.

The sample size is chosen such that the power to reject the null hypothesis is at least $1 - \beta$ for a given significance level α .

Value

An S3 class `designOrderedBinom` object with the following components:

- power: The power to reject the null hypothesis.
- alpha: The two-sided significance level.

- n: The maximum number of subjects.
- ngroups: The number of treatment groups.
- pi: The response probabilities for the treatment groups.
- w: The scores assigned to the treatment groups.
- trendstat: The Cochran-Armitage trend test statistic.
- allocationRatioPlanned: Allocation ratio for the treatment groups.
- rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignOrderedBinom(
  beta = 0.1, ngroups = 3, pi = c(0.1, 0.25, 0.5), alpha = 0.05))
```

```
getDesignPairedMeanDiff
```

Group Sequential Design for Paired Mean Difference

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for paired mean difference.

Usage

```
getDesignPairedMeanDiff(
  beta = NA_real_,
  n = NA_real_,
  pairedDiffH0 = 0,
  pairedDiff = 0.5,
  stDev = 1,
  normalApproximation = TRUE,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NA_real_,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
```

```

    futilityBounds = NA_real_,
    futilityCP = NA_real_,
    futilityPairedDiff = NA_real_,
    typeBetaSpending = "none",
    parameterBetaSpending = NA_real_,
    userBetaSpending = NA_real_,
    spendingTime = NA_real_
  )

```

Arguments

beta	The type II error.
n	The total sample size.
pairedDiffH0	The paired difference under the null hypothesis. Defaults to 0.
pairedDiff	The paired difference under the alternative hypothesis.
stDev	The standard deviation for paired difference.
normalApproximation	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
kMax	The maximum number of stages.
informationRates	The information rates. Fixed prior to the trial. Defaults to $(1:kMax) / kMax$ if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.

futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	The futility boundary on the conditional power scale.
futilityPairedDiff	The futility boundary on the paired difference scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Value

An S3 class designPairedMeanDiff object with three components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The overall significance level.
 - attainedAlpha: The attained significance level, which is different from the overall significance level in the presence of futility stopping.
 - kMax: The number of stages.
 - theta: The parameter value.
 - information: The maximum information.
 - expectedInformationH1: The expected information under H1.
 - expectedInformationH0: The expected information under H0.
 - drift: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
 - inflationFactor: The inflation factor (relative to the fixed design).
 - numberOfSubjects: The maximum number of subjects.
 - expectedNumberOfSubjectsH1: The expected number of subjects under H1.
 - expectedNumberOfSubjectsH0: The expected number of subjects under H0.
 - pairedDiffH0: The paired difference under the null hypothesis.
 - pairedDiff: The paired difference under the alternative hypothesis.
 - stDev: The standard deviation for paired difference.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.

- efficacyBounds: The efficacy boundaries on the Z-scale.
- futilityBounds: The futility boundaries on the Z-scale.
- rejectPerStage: The probability for efficacy stopping.
- futilityPerStage: The probability for futility stopping.
- cumulativeRejection: The cumulative probability for efficacy stopping.
- cumulativeFutility: The cumulative probability for futility stopping.
- cumulativeAlphaSpent: The cumulative alpha spent.
- efficacyP: The efficacy boundaries on the p-value scale.
- futilityP: The futility boundaries on the p-value scale.
- information: The cumulative information.
- efficacyStopping: Whether to allow efficacy stopping.
- futilityStopping: Whether to allow futility stopping.
- rejectPerStageH0: The probability for efficacy stopping under H0.
- futilityPerStageH0: The probability for futility stopping under H0.
- cumulativeRejectionH0: The cumulative probability for efficacy stopping under H0.
- cumulativeFutilityH0: The cumulative probability for futility stopping under H0.
- efficacyPairedDiff: The efficacy boundaries on the paired difference scale.
- futilityPairedDiff: The futility boundaries on the paired difference scale.
- numberOfSubjects: The number of subjects.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - typeBetaSpending: The type of beta spending.
 - parameterBetaSpending: The parameter value for beta spending.
 - userBetaSpending: The user defined beta spending.
 - spendingTime: The error spending time at each analysis.
 - normalApproximation: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution.
 - rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Example 1: group sequential trial power calculation
(design1 <- getDesignPairedMeanDiff(
  beta = 0.1, n = NA, pairedDiffH0 = 0, pairedDiff = -2, stDev = 5,
  kMax = 5, alpha = 0.05, typeAlphaSpending = "sf0F"))

# Example 2: sample size calculation for one-sample t-test
(design2 <- getDesignPairedMeanDiff(
  beta = 0.1, n = NA, pairedDiffH0 = 0, pairedDiff = -2, stDev = 5,
  normalApproximation = FALSE, alpha = 0.025))
```

getDesignPairedMeanDiffEquiv

Group Sequential Design for Equivalence in Paired Mean Difference

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for equivalence in paired mean difference.

Usage

```
getDesignPairedMeanDiffEquiv(
  beta = NA_real_,
  n = NA_real_,
  pairedDiffLower = NA_real_,
  pairedDiffUpper = NA_real_,
  pairedDiff = 0,
  stDev = 1,
  normalApproximation = TRUE,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  alpha = 0.05,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  spendingTime = NA_real_
)
```

Arguments

beta	The type II error.
n	The total sample size.
pairedDiffLower	The lower equivalence limit of paired difference.
pairedDiffUpper	The upper equivalence limit of paired difference.
pairedDiff	The paired difference under the alternative hypothesis.
stDev	The standard deviation for paired difference.
normalApproximation	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
kMax	The maximum number of stages.

informationRates	The information rates. Fixed prior to the trial. Defaults to $(1:kMax) / kMax$ if left unspecified.
alpha	The significance level for each of the two one-sided tests. Defaults to 0.05.
typeAlphaSpending	The type of alpha spending. One of the following: "0F" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sf0F" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sf0F".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
spendingTime	A vector of length $kMax$ for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Value

An S3 class *designPairedMeanDiffEquiv* object with three components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The significance level for each of the two one-sided tests. Defaults to 0.05.
 - attainedAlpha: The attained significance level under H_0 .
 - kMax: The number of stages.
 - information: The maximum information.
 - expectedInformationH1: The expected information under H_1 .
 - expectedInformationH0: The expected information under H_0 .
 - numberOfSubjects: The maximum number of subjects.
 - expectedNumberOfSubjectsH1: The expected number of subjects under H_1 .
 - expectedNumberOfSubjectsH0: The expected number of subjects under H_0 .
 - pairedDiffLower: The lower equivalence limit of paired difference.
 - pairedDiffUpper: The upper equivalence limit of paired difference.
 - pairedDiff: The paired difference under the alternative hypothesis.
 - stDev: The standard deviation for paired difference.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale for each of the two one-sided tests.
 - rejectPerStage: The probability for efficacy stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.

- cumulativeAlphaSpent: The cumulative alpha for each of the two one-sided tests.
- cumulativeAttainedAlpha: The cumulative probability for efficacy stopping under H0.
- efficacyPairedDiffLower: The efficacy boundaries on the paired difference scale for the one-sided null hypothesis on the lower equivalence limit.
- efficacyPairedDiffUpper: The efficacy boundaries on the paired difference scale for the one-sided null hypothesis on the upper equivalence limit.
- efficacyP: The efficacy bounds on the p-value scale for each of the two one-sided tests.
- information: The cumulative information.
- numberOfSubjects: The number of subjects.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - spendingTime: The error spending time at each analysis.
 - normalApproximation: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
 - rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
# Example 1: group sequential trial power calculation
(design1 <- getDesignPairedMeanDiffEquiv(
  beta = 0.1, n = NA, pairedDiffLower = -1.3, pairedDiffUpper = 1.3,
  pairedDiff = 0, stDev = 2.2,
  kMax = 4, alpha = 0.05, typeAlphaSpending = "sf0F"))

# Example 2: sample size calculation for t-test
(design2 <- getDesignPairedMeanDiffEquiv(
  beta = 0.1, n = NA, pairedDiffLower = -1.3, pairedDiffUpper = 1.3,
  pairedDiff = 0, stDev = 2.2,
  normalApproximation = FALSE, alpha = 0.05))
```

getDesignPairedMeanRatio

Group Sequential Design for Paired Mean Ratio

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for paired mean ratio.

Usage

```

getDesignPairedMeanRatio(
  beta = NA_real_,
  n = NA_real_,
  pairedRatioH0 = 1,
  pairedRatio = 1.2,
  CV = 1,
  normalApproximation = TRUE,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NA_real_,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NA_real_,
  futilityCP = NA_real_,
  futilityPairedRatio = NA_real_,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
  spendingTime = NA_real_
)

```

Arguments

<code>beta</code>	The type II error.
<code>n</code>	The total sample size.
<code>pairedRatioH0</code>	The paired ratio under the null hypothesis.
<code>pairedRatio</code>	The paired ratio under the alternative hypothesis.
<code>CV</code>	The coefficient of variation for paired ratio.
<code>normalApproximation</code>	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
<code>rounding</code>	Whether to round up sample size. Defaults to 1 for sample size rounding.
<code>kMax</code>	The maximum number of stages.
<code>informationRates</code>	The information rates. Fixed prior to the trial. Defaults to (1:kMax) / kMax if left unspecified.
<code>efficacyStopping</code>	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.

futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	The futility boundary on the conditional power scale.
futilityPairedRatio	The futility boundary on the paired ratio scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Value

An S3 class designPairedMeanRatio object with three components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The overall significance level.
 - attainedAlpha: The attained significance level, which is different from the overall significance level in the presence of futility stopping.

- kMax: The number of stages.
- theta: The parameter value.
- information: The maximum information.
- expectedInformationH1: The expected information under H1.
- expectedInformationH0: The expected information under H0.
- drift: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
- inflationFactor: The inflation factor (relative to the fixed design).
- numberOfSubjects: The maximum number of subjects.
- expectedNumberOfSubjectsH1: The expected number of subjects under H1.
- expectedNumberOfSubjectsH0: The expected number of subjects under H0.
- pairedRatioH0: The paired ratio under the null hypothesis.
- pairedRatio: The paired ratio under the alternative hypothesis.
- CV: The coefficient of variation for paired ratio.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale.
 - futilityBounds: The futility boundaries on the Z-scale.
 - rejectPerStage: The probability for efficacy stopping.
 - futilityPerStage: The probability for futility stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeFutility: The cumulative probability for futility stopping.
 - cumulativeAlphaSpent: The cumulative alpha spent.
 - efficacyP: The efficacy boundaries on the p-value scale.
 - futilityP: The futility boundaries on the p-value scale.
 - information: The cumulative information.
 - efficacyStopping: Whether to allow efficacy stopping.
 - futilityStopping: Whether to allow futility stopping.
 - rejectPerStageH0: The probability for efficacy stopping under H0.
 - futilityPerStageH0: The probability for futility stopping under H0.
 - cumulativeRejectionH0: The cumulative probability for efficacy stopping under H0.
 - cumulativeFutilityH0: The cumulative probability for futility stopping under H0.
 - numberOfSubjects: The number of subjects.
 - efficacyPairedRatio: The efficacy boundaries on the paired ratio scale.
 - futilityPairedRatio: The futility boundaries on the paired ratio scale.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - typeBetaSpending: The type of beta spending.
 - parameterBetaSpending: The parameter value for beta spending.
 - userBetaSpending: The user defined beta spending.

- spendingTime: The error spending time at each analysis.
- normalApproximation: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution.
- rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Example 1: group sequential trial power calculation
(design1 <- getDesignPairedMeanRatio(
  beta = 0.1, n = NA, pairedRatio = 1.2, CV = 0.35,
  kMax = 5, alpha = 0.05, typeAlphaSpending = "sfOF"))

# Example 2: sample size calculation for one-sample t-test
(design2 <- getDesignPairedMeanRatio(
  beta = 0.1, n = NA, pairedRatio = 1.2, CV = 0.35,
  normalApproximation = FALSE, alpha = 0.05))
```

getDesignPairedMeanRatioEquiv

Group Sequential Design for Equivalence in Paired Mean Ratio

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for equivalence in paired mean ratio.

Usage

```
getDesignPairedMeanRatioEquiv(
  beta = NA_real_,
  n = NA_real_,
  pairedRatioLower = NA_real_,
  pairedRatioUpper = NA_real_,
  pairedRatio = 1,
  CV = 1,
  normalApproximation = TRUE,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  alpha = 0.05,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
```

```

    spendingTime = NA_real_
  )

```

Arguments

beta	The type II error.
n	The total sample size.
pairedRatioLower	The lower equivalence limit of paired ratio.
pairedRatioUpper	The upper equivalence limit of paired ratio.
pairedRatio	The paired ratio under the alternative hypothesis.
CV	The coefficient of variation for paired ratio.
normalApproximation	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
kMax	The maximum number of stages.
informationRates	The information rates. Fixed prior to the trial. Defaults to (1:kMax) / kMax if left unspecified.
alpha	The significance level for each of the two one-sided tests. Defaults to 0.05.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Value

An S3 class designPairedMeanRatioEquiv object with three components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The significance level for each of the two one-sided tests. Defaults to 0.05.

- attainedAlpha: The attained significance level under H0.
- kMax: The number of stages.
- information: The maximum information.
- expectedInformationH1: The expected information under H1.
- expectedInformationH0: The expected information under H0.
- numberOfSubjects: The maximum number of subjects.
- expectedNumberOfSubjectsH1: The expected number of subjects under H1.
- expectedNumberOfSubjectsH0: The expected number of subjects under H0.
- pairedRatioLower: The lower equivalence limit of paired ratio.
- pairedRatioUpper: The upper equivalence limit of paired ratio.
- pairedRatio: The paired ratio under the alternative hypothesis.
- CV: The coefficient of variation for paired ratios.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale for each of the two one-sided tests.
 - rejectPerStage: The probability for efficacy stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeAlphaSpent: The cumulative alpha for each of the two one-sided tests.
 - cumulativeAttainedAlpha: The cumulative alpha attained under H0.
 - efficacyP: The efficacy bounds on the p-value scale for each of the two one-sided tests.
 - information: The cumulative information.
 - numberOfSubjects: The number of subjects.
 - efficacyPairedRatioLower: The efficacy boundaries on the paired ratio scale for the one-sided null hypothesis on the lower equivalence limit.
 - efficacyPairedRatioUpper: The efficacy boundaries on the paired ratio scale for the one-sided null hypothesis on the upper equivalence limit.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - spendingTime: The error spending time at each analysis.
 - normalApproximation: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
 - rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
# Example 1: group sequential trial power calculation
(design1 <- getDesignPairedMeanRatioEquiv(
  beta = 0.1, n = NA, pairedRatioLower = 0.8, pairedRatioUpper = 1.25,
  pairedRatio = 1, CV = 0.35,
  kMax = 4, alpha = 0.05, typeAlphaSpending = "sf0F"))

# Example 2: sample size calculation for t-test
(design2 <- getDesignPairedMeanRatioEquiv(
  beta = 0.1, n = NA, pairedRatioLower = 0.8, pairedRatioUpper = 1.25,
  pairedRatio = 1, CV = 0.35,
  normalApproximation = FALSE, alpha = 0.05))
```

```
getDesignPairedPropMcNemar
```

Group Sequential Design for McNemar's Test for Paired Proportions

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for McNemar's test for paired proportions.

Usage

```
getDesignPairedPropMcNemar(
  beta = NA_real_,
  n = NA_real_,
  pDiscordant = NA_real_,
  riskDiff = NA_real_,
  nullVariance = TRUE,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NA_real_,
  alpha = 0.025,
  typeAlphaSpending = "sf0F",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NA_real_,
  futilityCP = NA_real_,
  futilityRiskDiff = NA_real_,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
  spendingTime = NA_real_
)
```

Arguments

beta	The type II error.
n	The total sample size.
pDiscordant	The proportion of discordant pairs ($\pi_{01} + \pi_{10}$).
riskDiff	The risk difference between the active and control treatments ($\delta = \pi_{1t} - \pi_{1c} = \pi_{01} - \pi_{10}$).
nullVariance	Whether to use the variance under the null or the variance under the alternative.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
kMax	The maximum number of stages.
informationRates	The information rates. Fixed prior to the trial. Defaults to $(1:kMax) / kMax$ if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsiatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	The futility bounds on the conditional power scale.
futilityRiskDiff	The futility bounds on the risk difference scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".

parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Details

Consider a group sequential design for McNemar's test for paired proportions. The table below shows joint probabilities for each cell (π_{ij} where i is for control group and j is for experimental group), with marginal totals.

	Experimental: No Response	Experimental: Response	Row Total
Control: No Response	π_{00}	π_{01}	$1 - \pi_c$
Control: Response	π_{10}	π_{11}	π_c
Column Total	$1 - \pi_t$	π_t	1

The parameters π_{01} and π_{10} are the discordant pairs (i.e., $\pi_{01} + \pi_{10} = \xi$) and the risk difference is $\pi_{01} - \pi_{10} = \delta$. The parameter π_t is the proportion of experimental group response, and π_c is the proportion of control group response. The parameter of interest is

$$\theta = \pi_t - \pi_c = \pi_{01} - \pi_{10} = \delta$$

The variance of $\hat{\theta}$ can be obtained from the multinomial distribution as follows:

$$Var(\hat{\theta}) = \frac{1}{n} \{ \pi_{01}(1 - \pi_{01}) + \pi_{10}(1 - \pi_{10}) + 2\pi_{01}\pi_{10} \}$$

which can be simplified to

$$Var(\hat{\theta}) = \frac{1}{n} (\xi - \delta^2)$$

Here, n is the total number of treatment pairs. This is the unconditional variance, which is used for the overall design.

Value

An S3 class designPairedPropMcNemar object with three components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The overall significance level.
 - attainedAlpha: The attained significance level, which is different from the overall significance level in the presence of futility stopping.
 - kMax: The number of stages.
 - theta: The parameter value.
 - information: The maximum information.
 - expectedInformationH1: The expected information under H1.

- expectedInformationH0: The expected information under H0.
- drift: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
- inflationFactor: The inflation factor (relative to the fixed design).
- numberOfSubjects: The maximum number of subjects.
- expectedNumberOfSubjectsH1: The expected number of subjects under H1.
- expectedNumberOfSubjectsH0: The expected number of subjects under H0.
- pDiscordant: The proportion of discordant pairs ($\pi_{10} + \pi_{01}$).
- riskDiff: The risk difference between the active and control treatments ($\delta = \pi_t - \pi_c = \pi_{01} - \pi_{10}$)
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale.
 - futilityBounds: The futility boundaries on the Z-scale.
 - rejectPerStage: The probability for efficacy stopping.
 - futilityPerStage: The probability for futility stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeFutility: The cumulative probability for futility stopping.
 - cumulativeAlphaSpent: The cumulative alpha spent.
 - efficacyP: The efficacy boundaries on the p-value scale.
 - futilityP: The futility boundaries on the p-value scale.
 - information: The cumulative information.
 - efficacyStopping: Whether to allow efficacy stopping.
 - futilityStopping: Whether to allow futility stopping.
 - rejectPerStageH0: The probability for efficacy stopping under H0.
 - futilityPerStageH0: The probability for futility stopping under H0.
 - cumulativeRejectionH0: The cumulative probability for efficacy stopping under H0.
 - cumulativeFutilityH0: The cumulative probability for futility stopping under H0.
 - efficacyRiskDiff: The efficacy boundaries on the risk difference scale.
 - futilityRiskDiff: The futility boundaries on the risk difference scale.
 - numberOfSubjects: The number of subjects.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - typeBetaSpending: The type of beta spending.
 - parameterBetaSpending: The parameter value for beta spending.
 - userBetaSpending: The user defined beta spending.
 - spendingTime: The error spending time at each analysis.
 - varianceRatio: The ratio of the variance under H0 to the variance under H1.
 - rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Example 1: fixed design
(design1 <- getDesignPairedPropMcNemar(
  beta = 0.1, n = NA, pDiscordant = 0.16, riskDiff = 0.1,
  alpha = 0.025))

# Example 2: group sequential design
(design2 <- getDesignPairedPropMcNemar(
  beta = 0.1, n = NA, pDiscordant = 0.16, riskDiff = 0.1,
  alpha = 0.025, kMax = 3, typeAlphaSpending = "sfOF"))
```

getDesignRepeatedANOVA

Power and Sample Size for Repeated-Measures ANOVA

Description

Obtains the power and sample size for one-way repeated measures analysis of variance. Each subject takes all treatments in the longitudinal study.

Usage

```
getDesignRepeatedANOVA(
  beta = NA_real_,
  n = NA_real_,
  ngroups = 2,
  means = NA_real_,
  stDev = 1,
  corr = 0,
  rounding = TRUE,
  alpha = 0.05
)
```

Arguments

beta	The type II error.
n	The total sample size.
ngroups	The number of treatment groups.
means	The treatment group means.
stDev	The total standard deviation.
corr	The correlation among the repeated measures.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
alpha	The two-sided significance level. Defaults to 0.05.

Details

Let y_{ij} denote the measurement under treatment condition j ($j = 1, \dots, k$) for subject i ($i = 1, \dots, n$). Then

$$y_{ij} = \alpha + \beta_j + b_i + e_{ij}$$

where b_i denotes the subject random effect, $b_i \sim N(0, \sigma_b^2)$ and $e_{ij} \sim N(0, \sigma_e^2)$ denotes the within-subject residual. If we set $\beta_k = 0$, then α is the mean of the last treatment (control), and β_j is the difference in means between the j th treatment and the control for $j = 1, \dots, k - 1$.

The repeated measures have a compound symmetry covariance structure. Let $\sigma^2 = \sigma_b^2 + \sigma_e^2$, and $\rho = \frac{\sigma_b^2}{\sigma_b^2 + \sigma_e^2}$. Then $\text{Var}(y_i) = \sigma^2\{(1 - \rho)I_k + \rho 1_k 1_k^T\}$. Let X_i denote the design matrix for subject i . Let $\theta = (\alpha, \beta_1, \dots, \beta_{k-1})^T$. It follows that

$$\text{Var}(\hat{\theta}) = \left(\sum_{i=1}^n X_i^T V_i^{-1} X_i \right)^{-1}.$$

It can be shown that

$$\text{Var}(\hat{\beta}) = \frac{\sigma^2(1 - \rho)}{n} (I_{k-1} + 1_{k-1} 1_{k-1}^T).$$

It follows that $\hat{\beta}^T \hat{V}_{\hat{\beta}}^{-1} \hat{\beta} \sim F_{k-1, (n-1)(k-1), \lambda}$ where the noncentrality parameter for the F distribution is

$$\lambda = \beta^T V_{\hat{\beta}}^{-1} \beta = \frac{n \sum_{j=1}^k (\mu_j - \bar{\mu})^2}{\sigma^2(1 - \rho)}.$$

Value

An S3 class `designRepeatedANOVA` object with the following components:

- `power`: The power to reject the null hypothesis that there is no difference among the treatment groups.
- `alpha`: The two-sided significance level.
- `n`: The number of subjects.
- `ngroups`: The number of treatment groups.
- `means`: The treatment group means.
- `stDev`: The total standard deviation.
- `corr`: The correlation among the repeated measures.
- `effectsize`: The effect size.
- `rounding`: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignRepeatedANOVA(
  beta = 0.1, ngroups = 4, means = c(1.5, 2.5, 2, 0),
  stDev = 5, corr = 0.2, alpha = 0.05))
```

getDesignRepeatedANOVAContrast

Power and Sample Size for One-Way Repeated Measures ANOVA Contrast

Description

Obtains the power and sample size for a single contrast in one-way repeated measures analysis of variance.

Usage

```
getDesignRepeatedANOVAContrast(
  beta = NA_real_,
  n = NA_real_,
  ngroups = 2,
  means = NA_real_,
  stDev = 1,
  corr = 0,
  contrast = NA_real_,
  meanContrastH0 = 0,
  rounding = TRUE,
  alpha = 0.025
)
```

Arguments

beta	The type II error.
n	The total sample size.
ngroups	The number of treatment groups.
means	The treatment group means.
stDev	The total standard deviation.
corr	The correlation among the repeated measures.
contrast	The coefficients for the single contrast.
meanContrastH0	The mean of the contrast under the null hypothesis.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
alpha	The one-sided significance level. Defaults to 0.025.

Value

An S3 class designRepeatedANOVAContrast object with the following components:

- power: The power to reject the null hypothesis for the treatment contrast.
- alpha: The one-sided significance level.

- n: The number of subjects.
- ngroups: The number of treatment groups.
- means: The treatment group means.
- stDev: The total standard deviation.
- corr: The correlation among the repeated measures.
- contrast: The coefficients for the single contrast.
- meanContrastH0: The mean of the contrast under the null hypothesis.
- meanContrast: The mean of the contrast under the alternative hypothesis.
- effectsize: The effect size.
- rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignRepeatedANOVAContrast(
  beta = 0.1, ngroups = 4, means = c(1.5, 2.5, 2, 0),
  stDev = 5, corr = 0.2, contrast = c(1, 1, 1, -3)/3,
  alpha = 0.025))
```

getDesignRiskDiff

Group Sequential Design for Two-Sample Risk Difference

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for two-sample risk difference.

Usage

```
getDesignRiskDiff(
  beta = NA_real_,
  n = NA_real_,
  riskDiffH0 = 0,
  pi1 = NA_real_,
  pi2 = NA_real_,
  nullVariance = TRUE,
  allocationRatioPlanned = 1,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
```

```

    futilityStopping = NA_integer_,
    criticalValues = NA_real_,
    alpha = 0.025,
    typeAlphaSpending = "sfOF",
    parameterAlphaSpending = NA_real_,
    userAlphaSpending = NA_real_,
    futilityBounds = NA_real_,
    futilityCP = NA_real_,
    futilityRiskDiff = NA_real_,
    typeBetaSpending = "none",
    parameterBetaSpending = NA_real_,
    userBetaSpending = NA_real_,
    spendingTime = NA_real_
  )

```

Arguments

beta	The type II error.
n	The total sample size.
riskDiffH0	The risk difference under the null hypothesis. Defaults to 0.
pi1	The assumed probability for the active treatment group.
pi2	The assumed probability for the control group.
nullVariance	Whether to use the variance under the null or the empirical variance under the alternative.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
kMax	The maximum number of stages.
informationRates	The information rates. Fixed prior to the trial. Defaults to (1:kMax) / kMax if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for

	Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	The futility bounds on the conditional power scale.
futilityRiskDiff	The futility bounds on the risk difference scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Details

Consider a group sequential design for two-sample risk difference. The parameter of interest is

$$\theta = \pi_1 - \pi_2$$

where π_1 is the response probability for the active treatment group and π_2 is the response probability for the control group. The variance of $\hat{\theta}$ can be obtained from the binomial distributions as follows:

$$Var(\hat{\theta}) = \frac{1}{n} \left\{ \frac{\pi_1(1 - \pi_1)}{r} + \frac{\pi_2(1 - \pi_2)}{1 - r} \right\}$$

where n is the total number of subjects and r is the randomization probability for the active treatment group. When nullVariance = TRUE, the variance is computed under the null hypothesis. In this case, the values of π_1 and π_2 in the variance formula are replaced with their restricted maximum likelihood counterparts, subject to the constraint

$$\pi_1 - \pi_2 = \theta_0$$

Value

An S3 class `designRiskDiff` object with three components:

- `overallResults`: A data frame containing the following variables:
 - `overallReject`: The overall rejection probability.
 - `alpha`: The overall significance level.
 - `attainedAlpha`: The attained significance level, which is different from the overall significance level in the presence of futility stopping.
 - `kMax`: The number of stages.
 - `theta`: The parameter value.
 - `information`: The maximum information.
 - `expectedInformationH1`: The expected information under H1.
 - `expectedInformationH0`: The expected information under H0.
 - `drift`: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
 - `inflationFactor`: The inflation factor (relative to the fixed design).
 - `numberOfSubjects`: The maximum number of subjects.
 - `expectedNumberOfSubjectsH1`: The expected number of subjects under H1.
 - `expectedNumberOfSubjectsH0`: The expected number of subjects under H0.
 - `riskDiffH0`: The risk difference under the null hypothesis.
 - `pi1`: The assumed probability for the active treatment group.
 - `pi2`: The assumed probability for the control group.
- `byStageResults`: A data frame containing the following variables:
 - `informationRates`: The information rates.
 - `efficacyBounds`: The efficacy boundaries on the Z-scale.
 - `futilityBounds`: The futility boundaries on the Z-scale.
 - `rejectPerStage`: The probability for efficacy stopping.
 - `futilityPerStage`: The probability for futility stopping.
 - `cumulativeRejection`: The cumulative probability for efficacy stopping.
 - `cumulativeFutility`: The cumulative probability for futility stopping.
 - `cumulativeAlphaSpent`: The cumulative alpha spent.
 - `efficacyRiskDiff`: The efficacy boundaries on the risk difference scale.
 - `futilityRiskDiff`: The futility boundaries on the risk difference scale.
 - `efficacyP`: The efficacy boundaries on the p-value scale.
 - `futilityP`: The futility boundaries on the p-value scale.
 - `information`: The cumulative information.
 - `efficacyStopping`: Whether to allow efficacy stopping.
 - `futilityStopping`: Whether to allow futility stopping.
 - `rejectPerStageH0`: The probability for efficacy stopping under H0.
 - `futilityPerStageH0`: The probability for futility stopping under H0.
 - `cumulativeRejectionH0`: The cumulative probability for efficacy stopping under H0.
 - `cumulativeFutilityH0`: The cumulative probability for futility stopping under H0.
 - `numberOfSubjects`: The number of subjects.

- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - typeBetaSpending: The type of beta spending.
 - parameterBetaSpending: The parameter value for beta spending.
 - userBetaSpending: The user defined beta spending.
 - spendingTime: The error spending time at each analysis.
 - varianceRatio: The ratio of the variance under H0 to the variance under H1.
 - nullVariance: Whether to use the variance under the null or the empirical variance under the alternative.
 - allocationRatioPlanned: Allocation ratio for the active treatment versus control.
 - rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignRiskDiff(
  beta = 0.2, n = NA, pi1 = 0.1, pi2 = 0.15,
  kMax = 3, alpha = 0.025, typeAlphaSpending = "sfOF",
  nullVariance = FALSE))
```

getDesignRiskDiffEquiv

Group Sequential Design for Equivalence in Two-Sample Risk Difference

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for equivalence in two-sample risk difference.

Usage

```
getDesignRiskDiffEquiv(
  beta = NA_real_,
  n = NA_real_,
  riskDiffLower = NA_real_,
  riskDiffUpper = NA_real_,
  pi1 = NA_real_,
  pi2 = NA_real_,
  allocationRatioPlanned = 1,
```

```

    rounding = TRUE,
    kMax = 1L,
    informationRates = NA_real_,
    criticalValues = NA_real_,
    alpha = 0.05,
    typeAlphaSpending = "sfOF",
    parameterAlphaSpending = NA_real_,
    userAlphaSpending = NA_real_,
    spendingTime = NA_real_
  )

```

Arguments

<code>beta</code>	The type II error.
<code>n</code>	The total sample size.
<code>riskDiffLower</code>	The lower equivalence limit of risk difference.
<code>riskDiffUpper</code>	The upper equivalence limit of risk difference.
<code>pi1</code>	The assumed probability for the active treatment group.
<code>pi2</code>	The assumed probability for the control group.
<code>allocationRatioPlanned</code>	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
<code>rounding</code>	Whether to round up sample size. Defaults to 1 for sample size rounding.
<code>kMax</code>	The maximum number of stages.
<code>informationRates</code>	The information rates. Fixed prior to the trial. Defaults to $(1:kMax) / kMax$ if left unspecified.
<code>criticalValues</code>	Upper boundaries on the z-test statistic scale for stopping for efficacy.
<code>alpha</code>	The significance level for each of the two one-sided tests. Defaults to 0.05.
<code>typeAlphaSpending</code>	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
<code>parameterAlphaSpending</code>	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
<code>userAlphaSpending</code>	The user defined alpha spending. Cumulative alpha spent up to each stage.
<code>spendingTime</code>	A vector of length <code>kMax</code> for the error spending time at each analysis. Defaults to missing, in which case, it is the same as <code>informationRates</code> .

Value

An S3 class `designRiskDiffEquiv` object with three components:

- `overallResults`: A data frame containing the following variables:
 - `overallReject`: The overall rejection probability.
 - `alpha`: The significance level for each of the two one-sided tests. Defaults to 0.05.
 - `attainedAlphaH10`: The attained significance level under H10.
 - `attainedAlphaH20`: The attained significance level under H20.
 - `kMax`: The number of stages.
 - `information`: The maximum information.
 - `expectedInformationH1`: The expected information under H1.
 - `expectedInformationH10`: The expected information under H10.
 - `expectedInformationH20`: The expected information under H20.
 - `numberOfSubjects`: The maximum number of subjects.
 - `expectedNumberOfSubjectsH1`: The expected number of subjects under H1.
 - `expectedNumberOfSubjectsH10`: The expected number of subjects under H10.
 - `expectedNumberOfSubjectsH20`: The expected number of subjects under H20.
 - `riskDiffLower`: The lower equivalence limit of risk difference.
 - `riskDiffUpper`: The upper equivalence limit of risk difference.
 - `pi1`: The assumed probability for the active treatment group.
 - `pi2`: The assumed probability for the control group.
 - `riskDiff`: The risk difference.
- `byStageResults`: A data frame containing the following variables:
 - `informationRates`: The information rates.
 - `efficacyBounds`: The efficacy boundaries on the Z-scale for each of the two one-sided tests.
 - `rejectPerStage`: The probability for efficacy stopping.
 - `cumulativeRejection`: The cumulative probability for efficacy stopping.
 - `cumulativeAlphaSpent`: The cumulative alpha for each of the two one-sided tests.
 - `cumulativeAttainedAlphaH10`: The cumulative alpha attained under H10.
 - `cumulativeAttainedAlphaH20`: The cumulative alpha attained under H20.
 - `efficacyP`: The efficacy bounds on the p-value scale for each of the two one-sided tests.
 - `information`: The cumulative information.
 - `efficacyRiskDiffLower`: The efficacy boundaries on the risk difference scale for the one-sided null hypothesis on the lower equivalence limit.
 - `efficacyRiskDiffUpper`: The efficacy boundaries on the risk difference scale for the one-sided null hypothesis on the upper equivalence limit.
 - `numberOfSubjects`: The number of subjects.
- `settings`: A list containing the following input parameters:
 - `typeAlphaSpending`: The type of alpha spending.
 - `parameterAlphaSpending`: The parameter value for alpha spending.
 - `userAlphaSpending`: The user defined alpha spending.
 - `spendingTime`: The error spending time at each analysis.
 - `allocationRatioPlanned`: Allocation ratio for the active treatment versus control.
 - `rounding`: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
(design1 <- getDesignRiskDiffEquiv(
  beta = 0.2, n = NA, riskDiffLower = -0.1,
  riskDiffUpper = 0.1, pi1 = 0.12, pi2 = 0.12,
  kMax = 3, alpha = 0.05, typeAlphaSpending = "sf0F"))
```

```
getDesignRiskDiffExact
```

Power and Sample Size for Exact Unconditional Test for Risk Difference

Description

Obtains the power given sample size or obtains the sample size given power for exact unconditional test of risk difference.

Usage

```
getDesignRiskDiffExact(
  beta = NA_real_,
  n = NA_real_,
  riskDiffH0 = 0,
  pi1 = NA_real_,
  pi2 = NA_real_,
  allocationRatioPlanned = 1,
  alpha = 0.025
)
```

Arguments

beta	The type II error.
n	The total sample size.
riskDiffH0	The risk difference under the null hypothesis. Defaults to 0.
pi1	The assumed probability for the active treatment group.
pi2	The assumed probability for the control group.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
alpha	The one-sided significance level. Defaults to 0.025.

Value

A data frame with the following variables:

- alpha: The specified one-sided significance level.
- attainedAlpha: The attained one-sided significance level.
- power: The power.
- n: The sample size.
- riskDiffH0: The risk difference under the null hypothesis.
- pi1: The assumed probability for the active treatment group.
- pi2: The assumed probability for the control group.
- allocationRatioPlanned: Allocation ratio for the active treatment versus control.
- zstatRiskDiffBound: The critical value on the scale of score test statistic for risk difference.
- pi2star: The response probability in the control group at which the critical value of the test statistic is attained.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Superiority test

getDesignRiskDiffExact(n = 50, pi1 = 0.6, pi2 = 0.25, alpha = 0.025)

# Non-inferiority test

getDesignRiskDiffExact(beta = 0.2, riskDiffH0 = -0.3,
                        pi1 = 0.9, pi2 = 0.9, alpha = 0.025)
```

```
getDesignRiskDiffExactEquiv
```

*Power and Sample Size for Exact Unconditional Test for Equivalence
in Risk Difference*

Description

Obtains the power given sample size or obtains the sample size given power for exact unconditional test of equivalence in risk difference.

Usage

```
getDesignRiskDiffExactEquiv(
  beta = NA_real_,
  n = NA_real_,
  riskDiffLower = NA_real_,
  riskDiffUpper = NA_real_,
  pi1 = NA_real_,
  pi2 = NA_real_,
  allocationRatioPlanned = 1,
  alpha = 0.05
)
```

Arguments

beta	The type II error.
n	The total sample size.
riskDiffLower	The lower equivalence limit of risk difference.
riskDiffUpper	The upper equivalence limit of risk difference.
pi1	The assumed probability for the active treatment group.
pi2	The assumed probability for the control group.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
alpha	The significance level for each of the two one-sided tests. Defaults to 0.05.

Value

A data frame with the following variables:

- alpha: The specified significance level for each of the two one-sided tests.
- attainedAlpha: The attained significance level.
- power: The power.
- n: The sample size.
- riskDiffLower: The lower equivalence limit of risk difference.
- riskDiffUpper: The upper equivalence limit of risk difference.
- pi1: The assumed probability for the active treatment group.
- pi2: The assumed probability for the control group.
- riskDiff: The risk difference.
- allocationRatioPlanned: Allocation ratio for the active treatment versus control.
- zstatRiskDiffLower: The efficacy boundaries on the z-test statistic scale for the one-sided null hypothesis on the lower equivalence limit.
- zstatRiskDiffUpper: The efficacy boundaries on the z-test statistic scale for the one-sided null hypothesis on the upper equivalence limit.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
getDesignRiskDiffExactEquiv(
  n = 200, riskDiffLower = -0.2, riskDiffUpper = 0.2,
  pi1 = 0.775, pi2 = 0.775, alpha = 0.05)
```

getDesignRiskRatio	<i>Group Sequential Design for Two-Sample Risk Ratio</i>
--------------------	--

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for two-sample risk ratio.

Usage

```
getDesignRiskRatio(
  beta = NA_real_,
  n = NA_real_,
  riskRatioH0 = 1,
  pi1 = NA_real_,
  pi2 = NA_real_,
  nullVariance = TRUE,
  allocationRatioPlanned = 1,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NA_real_,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NA_real_,
  futilityCP = NA_real_,
  futilityRiskRatio = NA_real_,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
  spendingTime = NA_real_
)
```

Arguments

<code>beta</code>	The type II error.
<code>n</code>	The total sample size.
<code>riskRatioH0</code>	The risk ratio under the null hypothesis. Defaults to 1.
<code>pi1</code>	The assumed probability for the active treatment group.
<code>pi2</code>	The assumed probability for the control group.
<code>nullVariance</code>	Whether to use the variance under the null or the empirical variance under the alternative.
<code>allocationRatioPlanned</code>	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
<code>rounding</code>	Whether to round up sample size. Defaults to 1 for sample size rounding.
<code>kMax</code>	The maximum number of stages.
<code>informationRates</code>	The information rates. Fixed prior to the trial. Defaults to $(1:kMax) / kMax$ if left unspecified.
<code>efficacyStopping</code>	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
<code>futilityStopping</code>	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
<code>criticalValues</code>	Upper boundaries on the z-test statistic scale for stopping for efficacy.
<code>alpha</code>	The significance level. Defaults to 0.025.
<code>typeAlphaSpending</code>	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
<code>parameterAlphaSpending</code>	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
<code>userAlphaSpending</code>	The user defined alpha spending. Cumulative alpha spent up to each stage.
<code>futilityBounds</code>	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., $kMax-1$. Defaults to <code>rep(-8, kMax-1)</code> if left unspecified. The futility bounds are non-binding for the calculation of critical values.
<code>futilityCP</code>	The futility bounds on the conditional power scale.
<code>futilityRiskRatio</code>	The futility bounds on the risk ratio scale.

typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Details

Consider a group sequential design for two-sample risk ratio. The parameter of interest is

$$\rho = \pi_1 / \pi_2$$

where π_1 is the response probability for the active treatment group and π_2 is the response probability for the control group. For statistical inference, the parameter is often transformed to the log scale:

$$\theta = \log(\rho) = \log(\pi_1) - \log(\pi_2)$$

The variance of the estimator $\hat{\theta}$ can be derived from the binomial distributions as follows:

$$Var(\hat{\theta}) = \frac{1}{n} \left\{ \frac{1 - \pi_1}{\pi_1 r} + \frac{1 - \pi_2}{\pi_2 (1 - r)} \right\}$$

where n is the total number of subjects and r is the randomization probability for the active treatment group. When nullVariance = TRUE, the variance is computed under the null hypothesis. In this case, the values of π_1 and π_2 in the variance formula are replaced with their restricted maximum likelihood counterparts, subject to the constraint

$$\pi_1 / \pi_2 = \rho_0$$

Value

An S3 class designRiskRatio object with three components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The overall significance level.
 - attainedAlpha: The attained significance level, which is different from the overall significance level in the presence of futility stopping.
 - kMax: The number of stages.
 - theta: The parameter value.
 - information: The maximum information.
 - expectedInformationH1: The expected information under H1.

- `expectedInformationH0`: The expected information under H_0 .
- `drift`: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
- `inflationFactor`: The inflation factor (relative to the fixed design).
- `numberOfSubjects`: The maximum number of subjects.
- `expectedNumberOfSubjectsH1`: The expected number of subjects under H_1 .
- `expectedNumberOfSubjectsH0`: The expected number of subjects under H_0 .
- `riskRatioH0`: The risk ratio under the null hypothesis.
- `pi1`: The assumed probability for the active treatment group.
- `pi2`: The assumed probability for the control group.
- `byStageResults`: A data frame containing the following variables:
 - `informationRates`: The information rates.
 - `efficacyBounds`: The efficacy boundaries on the Z-scale.
 - `futilityBounds`: The futility boundaries on the Z-scale.
 - `rejectPerStage`: The probability for efficacy stopping.
 - `futilityPerStage`: The probability for futility stopping.
 - `cumulativeRejection`: The cumulative probability for efficacy stopping.
 - `cumulativeFutility`: The cumulative probability for futility stopping.
 - `cumulativeAlphaSpent`: The cumulative alpha spent.
 - `efficacyP`: The efficacy boundaries on the p-value scale.
 - `futilityP`: The futility boundaries on the p-value scale.
 - `information`: The cumulative information.
 - `efficacyStopping`: Whether to allow efficacy stopping.
 - `futilityStopping`: Whether to allow futility stopping.
 - `rejectPerStageH0`: The probability for efficacy stopping under H_0 .
 - `futilityPerStageH0`: The probability for futility stopping under H_0 .
 - `cumulativeRejectionH0`: The cumulative probability for efficacy stopping under H_0 .
 - `cumulativeFutilityH0`: The cumulative probability for futility stopping under H_0 .
 - `efficacyRiskRatio`: The efficacy boundaries on the risk ratio scale.
 - `futilityRiskRatio`: The futility boundaries on the risk ratio scale.
 - `numberOfSubjects`: The number of subjects.
- `settings`: A list containing the following input parameters:
 - `typeAlphaSpending`: The type of alpha spending.
 - `parameterAlphaSpending`: The parameter value for alpha spending.
 - `userAlphaSpending`: The user defined alpha spending.
 - `typeBetaSpending`: The type of beta spending.
 - `parameterBetaSpending`: The parameter value for beta spending.
 - `userBetaSpending`: The user defined beta spending.
 - `spendingTime`: The error spending time at each analysis.
 - `varianceRatio`: The ratio of the variance under H_0 to the variance under H_1 .
 - `nullVariance`: Whether to use the variance under the null or the empirical variance under the alternative.
 - `allocationRatioPlanned`: Allocation ratio for the active treatment versus control.
 - `rounding`: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignRiskRatio(
  beta = 0.1, n = NA, pi1 = 0.5, pi2 = 0.3,
  alpha = 0.05))
```

getDesignRiskRatioEquiv

Group Sequential Design for Equivalence in Two-Sample Risk Ratio

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for equivalence in two-sample risk ratio.

Usage

```
getDesignRiskRatioEquiv(
  beta = NA_real_,
  n = NA_real_,
  riskRatioLower = NA_real_,
  riskRatioUpper = NA_real_,
  pi1 = NA_real_,
  pi2 = NA_real_,
  allocationRatioPlanned = 1,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  criticalValues = NA_real_,
  alpha = 0.05,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  spendingTime = NA_real_
)
```

Arguments

beta	The type II error.
n	The total sample size.
riskRatioLower	The lower equivalence limit of risk ratio.
riskRatioUpper	The upper equivalence limit of risk ratio.

<code>pi1</code>	The assumed probability for the active treatment group.
<code>pi2</code>	The assumed probability for the control group.
<code>allocationRatioPlanned</code>	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
<code>rounding</code>	Whether to round up sample size. Defaults to 1 for sample size rounding.
<code>kMax</code>	The maximum number of stages.
<code>informationRates</code>	The information rates. Fixed prior to the trial. Defaults to $(1:kMax) / kMax$ if left unspecified.
<code>criticalValues</code>	Upper boundaries on the z-test statistic scale for stopping for efficacy.
<code>alpha</code>	The significance level for each of the two one-sided tests. Defaults to 0.05.
<code>typeAlphaSpending</code>	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsiatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
<code>parameterAlphaSpending</code>	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
<code>userAlphaSpending</code>	The user defined alpha spending. Cumulative alpha spent up to each stage.
<code>spendingTime</code>	A vector of length <code>kMax</code> for the error spending time at each analysis. Defaults to missing, in which case, it is the same as <code>informationRates</code> .

Value

An S3 class `designRiskRatioEquiv` object with three components:

- `overallResults`: A data frame containing the following variables:
 - `overallReject`: The overall rejection probability.
 - `alpha`: The significance level for each of the two one-sided tests. Defaults to 0.05.
 - `attainedAlphaH10`: The attained significance level under H10.
 - `attainedAlphaH20`: The attained significance level under H20.
 - `kMax`: The number of stages.
 - `information`: The maximum information.
 - `expectedInformationH1`: The expected information under H1.
 - `expectedInformationH10`: The expected information under H10.
 - `expectedInformationH20`: The expected information under H20.
 - `numberOfSubjects`: The maximum number of subjects.
 - `expectedNumberOfSubjectsH1`: The expected number of subjects under H1.
 - `expectedNumberOfSubjectsH10`: The expected number of subjects under H10.

- expectedNumberOfSubjectsH20: The expected number of subjects under H20.
- riskRatioLower: The lower equivalence limit of risk ratio.
- riskRatioUpper: The upper equivalence limit of risk ratio.
- pi1: The assumed probability for the active treatment group.
- pi2: The assumed probability for the control group.
- riskRatio: The risk ratio.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale for each of the two one-sided tests.
 - rejectPerStage: The probability for efficacy stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeAlphaSpent: The cumulative alpha for each of the two one-sided tests.
 - cumulativeAttainedAlphaH10: The cumulative alpha attained under H10.
 - cumulativeAttainedAlphaH20: The cumulative alpha attained under H20.
 - efficacyRiskRatioLower: The efficacy boundaries on the risk ratio scale for the one-sided null hypothesis on the lower equivalence limit.
 - efficacyRiskRatioUpper: The efficacy boundaries on the risk ratio scale for the one-sided null hypothesis on the upper equivalence limit.
 - efficacyP: The efficacy bounds on the p-value scale for each of the two one-sided tests.
 - information: The cumulative information.
 - numberOfSubjects: The number of subjects.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - spendingTime: The error spending time at each analysis.
 - allocationRatioPlanned: Allocation ratio for the active treatment versus control.
 - rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
(design1 <- getDesignRiskRatioEquiv(
  beta = 0.2, n = NA, riskRatioLower = 0.8,
  riskRatioUpper = 1.25, pi1 = 0.12, pi2 = 0.12,
  kMax = 3, alpha = 0.05, typeAlphaSpending = "sf0F"))
```

getDesignRiskRatioExact

Power and Sample Size for Exact Unconditional Test for Risk Ratio

Description

Obtains the power given sample size or obtains the sample size given power for exact unconditional test of risk ratio.

Usage

```
getDesignRiskRatioExact(
  beta = NA_real_,
  n = NA_real_,
  riskRatioH0 = 1,
  pi1 = NA_real_,
  pi2 = NA_real_,
  allocationRatioPlanned = 1,
  alpha = 0.025
)
```

Arguments

beta	The type II error.
n	The total sample size.
riskRatioH0	The risk ratio under the null hypothesis. Defaults to 0.
pi1	The assumed probability for the active treatment group.
pi2	The assumed probability for the control group.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
alpha	The one-sided significance level. Defaults to 0.025.

Value

A data frame with the following variables:

- alpha: The specified one-sided significance level.
- attainedAlpha: The attained one-sided significance level.
- power: The power.
- n: The sample size.
- riskRatioH0: The risk ratio under the null hypothesis.
- pi1: The assumed probability for the active treatment group.
- pi2: The assumed probability for the control group.

- allocationRatioPlanned: Allocation ratio for the active treatment versus control.
- zstatRiskRatioBound: The critical value on the scale of score test statistic for risk ratio.
- pi2star: The response probability in the control group at which the critical value of the test statistic is attained.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Non-inferiority test

getDesignRiskRatioExact(beta = 0.2, riskRatioH0 = 0.7,
                        pi1 = 0.95, pi2 = 0.95, alpha = 0.025)
```

```
getDesignRiskRatioExactEquiv
```

*Power and Sample Size for Exact Unconditional Test for Equivalence
in Risk Ratio*

Description

Obtains the power given sample size or obtains the sample size given power for exact unconditional test of equivalence in risk ratio.

Usage

```
getDesignRiskRatioExactEquiv(
  beta = NA_real_,
  n = NA_real_,
  riskRatioLower = NA_real_,
  riskRatioUpper = NA_real_,
  pi1 = NA_real_,
  pi2 = NA_real_,
  allocationRatioPlanned = 1,
  alpha = 0.05
)
```

Arguments

beta	The type II error.
n	The total sample size.
riskRatioLower	The lower equivalence limit of risk ratio.
riskRatioUpper	The upper equivalence limit of risk ratio.

<code>pi1</code>	The assumed probability for the active treatment group.
<code>pi2</code>	The assumed probability for the control group.
<code>allocationRatioPlanned</code>	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
<code>alpha</code>	The significance level for each of the two one-sided tests. Defaults to 0.05.

Value

A data frame with the following variables:

- `alpha`: The specified significance level for each of the two one-sided tests.
- `attainedAlpha`: The attained significance level.
- `power`: The power.
- `n`: The sample size.
- `riskRatioLower`: The lower equivalence limit of risk ratio.
- `riskRatioUpper`: The upper equivalence limit of risk ratio.
- `pi1`: The assumed probability for the active treatment group.
- `pi2`: The assumed probability for the control group.
- `riskRatio`: The risk ratio.
- `allocationRatioPlanned`: Allocation ratio for the active treatment versus control.
- `zstatRiskRatioLower`: The efficacy boundaries on the z-test statistic scale for the one-sided null hypothesis on the lower equivalence limit.
- `zstatRiskRatioUpper`: The efficacy boundaries on the z-test statistic scale for the one-sided null hypothesis on the upper equivalence limit.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
getDesignRiskRatioExactEquiv(
  n = 200, riskRatioLower = 0.8, riskRatioUpper = 1.25,
  pi1 = 0.775, pi2 = 0.775, alpha = 0.05)
```

getDesignRiskRatioFM *Group Sequential Design for Two-Sample Risk Ratio Based on the Farrington-Manning Score Test*

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for two-sample risk ratio based on the Farrington-Manning score test

Usage

```
getDesignRiskRatioFM(
  beta = NA_real_,
  n = NA_real_,
  riskRatioH0 = 1,
  pi1 = NA_real_,
  pi2 = NA_real_,
  nullVariance = TRUE,
  allocationRatioPlanned = 1,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NA_real_,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NA_real_,
  futilityCP = NA_real_,
  futilityRiskRatio = NA_real_,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
  spendingTime = NA_real_
)
```

Arguments

beta	The type II error.
n	The total sample size.
riskRatioH0	The risk ratio under the null hypothesis. Defaults to 1.
pi1	The assumed probability for the active treatment group.
pi2	The assumed probability for the control group.

<code>nullVariance</code>	Whether to use the variance under the null or the empirical variance under the alternative.
<code>allocationRatioPlanned</code>	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
<code>rounding</code>	Whether to round up sample size. Defaults to 1 for sample size rounding.
<code>kMax</code>	The maximum number of stages.
<code>informationRates</code>	The information rates. Fixed prior to the trial. Defaults to $(1:kMax) / kMax$ if left unspecified.
<code>efficacyStopping</code>	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
<code>futilityStopping</code>	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
<code>criticalValues</code>	Upper boundaries on the z-test statistic scale for stopping for efficacy.
<code>alpha</code>	The significance level. Defaults to 0.025.
<code>typeAlphaSpending</code>	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
<code>parameterAlphaSpending</code>	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
<code>userAlphaSpending</code>	The user defined alpha spending. Cumulative alpha spent up to each stage.
<code>futilityBounds</code>	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., $kMax-1$. Defaults to <code>rep(-8, kMax-1)</code> if left unspecified. The futility bounds are non-binding for the calculation of critical values.
<code>futilityCP</code>	The futility bounds on the conditional power scale.
<code>futilityRiskRatio</code>	The futility bounds on the risk ratio scale.
<code>typeBetaSpending</code>	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
<code>parameterBetaSpending</code>	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".

userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Details

Consider a group sequential design for two-sample risk ratio. The parameter of interest is

$$\rho = \pi_1 / \pi_2$$

where π_1 is the response probability for the active treatment group and π_2 is the response probability for the control group. Let ρ_0 denote the risk ratio under the null hypothesis. The Farrington-Manning score test statistic is constructed as

$$Z = \frac{\hat{\pi}_1 - \rho_0 \hat{\pi}_2}{\sqrt{\text{Var}(\hat{\pi}_1 - \rho_0 \hat{\pi}_2)}}$$

The variance can be derived from the binomial distributions as follows:

$$\text{Var}(\hat{\pi}_1 - \rho_0 \hat{\pi}_2) = \frac{1}{n} \left\{ \frac{\pi_1(1 - \pi_1)}{r} + \frac{\rho_0^2 \pi_2(1 - \pi_2)}{1 - r} \right\}$$

where n is the total number of subjects and r is the randomization probability for the active treatment group. When nullVariance = TRUE, the variance is computed under the null hypothesis. In this case, the values of π_1 and π_2 in the variance formula are replaced with their restricted maximum likelihood counterparts, subject to the constraint

$$\pi_1 / \pi_2 = \rho_0$$

Value

An S3 class designRiskRatioFM object with three components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The overall significance level.
 - attainedAlpha: The attained significance level, which is different from the overall significance level in the presence of futility stopping.
 - kMax: The number of stages.
 - theta: The parameter value.
 - information: The maximum information.
 - expectedInformationH1: The expected information under H1.
 - expectedInformationH0: The expected information under H0.
 - drift: The drift parameter, equal to theta*sqrt(information).
 - inflationFactor: The inflation factor (relative to the fixed design).
 - numberOfSubjects: The maximum number of subjects.
 - expectedNumberOfSubjectsH1: The expected number of subjects under H1.
 - expectedNumberOfSubjectsH0: The expected number of subjects under H0.

- riskRatioH0: The risk ratio under the null hypothesis.
- pi1: The assumed probability for the active treatment group.
- pi2: The assumed probability for the control group.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale.
 - futilityBounds: The futility boundaries on the Z-scale.
 - rejectPerStage: The probability for efficacy stopping.
 - futilityPerStage: The probability for futility stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeFutility: The cumulative probability for futility stopping.
 - cumulativeAlphaSpent: The cumulative alpha spent.
 - efficacyP: The efficacy boundaries on the p-value scale.
 - futilityP: The futility boundaries on the p-value scale.
 - information: The cumulative information.
 - efficacyStopping: Whether to allow efficacy stopping.
 - futilityStopping: Whether to allow futility stopping.
 - rejectPerStageH0: The probability for efficacy stopping under H0.
 - futilityPerStageH0: The probability for futility stopping under H0.
 - cumulativeRejectionH0: The cumulative probability for efficacy stopping under H0.
 - cumulativeFutilityH0: The cumulative probability for futility stopping under H0.
 - efficacyRiskRatioScore: The efficacy boundaries on the score test $\pi_1 - \text{riskRatioH0} * \pi_2$ scale.
 - futilityRiskRatioScore: The futility boundaries on the score test $\pi_1 - \text{riskRatioH0} * \pi_2$ scale.
 - numberOfSubjects: The number of subjects.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - typeBetaSpending: The type of beta spending.
 - parameterBetaSpending: The parameter value for beta spending.
 - userBetaSpending: The user defined beta spending.
 - spendingTime: The error spending time at each analysis.
 - varianceRatio: The ratio of the variance under H0 to the variance under H1.
 - nullVariance: Whether to use the variance under the null or the empirical variance under the alternative.
 - allocationRatioPlanned: Allocation ratio for the active treatment versus control.
 - rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
(design1 <- getDesignRiskRatioFM(
  beta = 0.2, riskRatioH0 = 1.3, pi1 = 0.125, pi2 = 0.125,
  alpha = 0.05))
```

getDesignSlopeDiff	<i>Group Sequential Design for Two-Sample Slope Difference</i>
--------------------	--

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for two-sample slope difference.

Usage

```
getDesignSlopeDiff(
  beta = NA_real_,
  n = NA_real_,
  slopeDiffH0 = 0,
  slopeDiff = 0.5,
  stDev = 1,
  stDevCovariate = 1,
  allocationRatioPlanned = 1,
  normalApproximation = TRUE,
  rounding = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NA_real_,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NA_real_,
  futilityCP = NA_real_,
  futilitySlopeDiff = NA_real_,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
  spendingTime = NA_real_
)
```

Arguments

beta	The type II error.
n	The total sample size.
slopeDiffH0	The slope difference under the null hypothesis. Defaults to 0.
slopeDiff	The slope difference under the alternative hypothesis.
stDev	The standard deviation of the residual.
stDevCovariate	The standard deviation of the covariate.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
normalApproximation	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The exact calculation using the t distribution is only implemented for the fixed design.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
kMax	The maximum number of stages.
informationRates	The information rates. Fixed prior to the trial. Defaults to $(1:kMax) / kMax$ if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	The futility bounds on the conditional power scale.

futilitySlopeDiff	The futility bounds on the slope difference scale.
typeBetaSpending	The type of beta spending. One of the following: "sf0F" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Details

We assume a simple linear regression of the form

$$y_{g,i} = \alpha_g + \beta_g x_{g,i} + \epsilon_{g,i}$$

for treatment group g , where $\epsilon_{g,i}$ is the residual error for subject i in group g , which is assumed to be normally distributed with mean 0 and standard deviation σ_ϵ . The covariate $x_{g,i}$ is assumed to be normally distributed with mean 0 and standard deviation σ_x . Since

$$\hat{\beta}_g = \frac{\sum_{i=1}^{n_g} (x_{g,i} - \bar{x}_g) y_{g,i}}{\sum_{i=1}^{n_g} (x_{g,i} - \bar{x}_g)^2}$$

where n_g is the sample size for group g , it follows that

$$\hat{\beta}_g \sim N(\beta_g, \frac{\sigma_\epsilon^2}{\sum_{i=1}^{n_g} (x_{g,i} - \bar{x}_g)^2}).$$

The slope difference is defined as

$$\hat{\beta}_1 - \hat{\beta}_2$$

where $\hat{\beta}_1$ and $\hat{\beta}_2$ are the estimated slopes for treatment groups 1 and 2, respectively. Since the variance of $\hat{\beta}_g$ is

$$\frac{\sigma_\epsilon^2}{n_g \sigma_x^2}$$

we have

$$\hat{\beta}_1 - \hat{\beta}_2 \sim N(\beta_1 - \beta_2, (n_1^{-1} + n_2^{-1}) \sigma_\epsilon^2 / \sigma_x^2)$$

which can be used to calculate the power and sample size for the group sequential design.

Value

An S3 class `designSlopeDiff` object with three components:

- `overallResults`: A data frame containing the following variables:
 - `overallReject`: The overall rejection probability.
 - `alpha`: The overall significance level.
 - `attainedAlpha`: The attained significance level, which is different from the overall significance level in the presence of futility stopping.
 - `kMax`: The number of stages.
 - `theta`: The parameter value.
 - `information`: The maximum information.
 - `expectedInformationH1`: The expected information under H1.
 - `expectedInformationH0`: The expected information under H0.
 - `drift`: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
 - `inflationFactor`: The inflation factor (relative to the fixed design).
 - `numberOfSubjects`: The maximum number of subjects.
 - `expectedNumberOfSubjectsH1`: The expected number of subjects under H1.
 - `expectedNumberOfSubjectsH0`: The expected number of subjects under H0.
 - `slopeDiffH0`: The slope difference under the null hypothesis.
 - `slopeDiff`: The slope difference under the alternative hypothesis.
 - `stDev`: The standard deviation of the residual.
 - `stDevCovariate`: The standard deviation of the covariate.
- `byStageResults`: A data frame containing the following variables:
 - `informationRates`: The information rates.
 - `efficacyBounds`: The efficacy boundaries on the Z-scale.
 - `futilityBounds`: The futility boundaries on the Z-scale.
 - `rejectPerStage`: The probability for efficacy stopping.
 - `futilityPerStage`: The probability for futility stopping.
 - `cumulativeRejection`: The cumulative probability for efficacy stopping.
 - `cumulativeFutility`: The cumulative probability for futility stopping.
 - `cumulativeAlphaSpent`: The cumulative alpha spent.
 - `efficacyP`: The efficacy boundaries on the p-value scale.
 - `futilityP`: The futility boundaries on the p-value scale.
 - `information`: The cumulative information.
 - `efficacyStopping`: Whether to allow efficacy stopping.
 - `futilityStopping`: Whether to allow futility stopping.
 - `rejectPerStageH0`: The probability for efficacy stopping under H0.
 - `futilityPerStageH0`: The probability for futility stopping under H0.
 - `cumulativeRejectionH0`: The cumulative probability for efficacy stopping under H0.
 - `cumulativeFutilityH0`: The cumulative probability for futility stopping under H0.
 - `efficacySlopeDiff`: The efficacy boundaries on the slope difference scale.
 - `futilitySlopeDiff`: The futility boundaries on the slope difference scale.

- numberOfSubjects: The number of subjects.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - typeBetaSpending: The type of beta spending.
 - parameterBetaSpending: The parameter value for beta spending.
 - userBetaSpending: The user defined beta spending.
 - spendingTime: The error spending time at each analysis.
 - allocationRatioPlanned: Allocation ratio for the active treatment versus control.
 - normalApproximation: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution.
 - rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignSlopeDiff(
  beta = 0.1, n = NA, slopeDiff = -0.5,
  stDev = 10, stDevCovariate = 6,
  normalApproximation = FALSE, alpha = 0.025))
```

getDesignSlopeDiffMMRM

Group Sequential Design for Two-Sample Slope Difference From the MMRM Model

Description

Obtains the power given sample size or obtains the sample size given power for two-sample slope difference from the growth curve MMRM model.

Usage

```
getDesignSlopeDiffMMRM(
  beta = NA_real_,
  slopeDiffH0 = 0,
  slopeDiff = 0.5,
  stDev = 1,
  stDevIntercept = 1,
  stDevSlope = 1,
  corrInterceptSlope = 0.5,
```

```

w = NA_real_,
N = NA_real_,
accrualTime = 0,
accrualIntensity = NA_real_,
piecewiseSurvivalTime = 0,
gamma1 = 0,
gamma2 = 0,
accrualDuration = NA_real_,
followupTime = NA_real_,
fixedFollowup = FALSE,
allocationRatioPlanned = 1,
normalApproximation = TRUE,
rounding = TRUE,
kMax = 1L,
informationRates = NA_real_,
efficacyStopping = NA_integer_,
futilityStopping = NA_integer_,
criticalValues = NA_real_,
alpha = 0.025,
typeAlphaSpending = "sfOF",
parameterAlphaSpending = NA_real_,
userAlphaSpending = NA_real_,
futilityBounds = NA_real_,
futilityCP = NA_real_,
futilitySlopeDiff = NA_real_,
typeBetaSpending = "none",
parameterBetaSpending = NA_real_,
userBetaSpending = NA_real_,
spendingTime = NA_real_
)

```

Arguments

beta	The type II error.
slopeDiffH0	The slope difference under the null hypothesis. Defaults to 0.
slopeDiff	The slope difference under the alternative hypothesis.
stDev	The standard deviation of the residual.
stDevIntercept	The standard deviation of the random intercept.
stDevSlope	The standard deviation of the random slope.
corrInterceptSlope	The correlation between the random intercept and random slope.
w	The number of time units (e.g. weeks) per measurement visit in a period. In general, visits are more frequent in the beginning of the study and less frequent towards the end.
N	The number of measurement visits in a period. For example, $w = c(8, 16)$ and $N = c(2, \text{Inf})$ means that the response variable will be collected at baseline, week 8, week 16, and every 16 weeks thereafter.

accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
gamma1	The hazard rate for exponential dropout, or a vector of hazard rates for piecewise exponential dropout for the active treatment group.
gamma2	The hazard rate for exponential dropout, or a vector of hazard rates for piecewise exponential dropout for the control group.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
normalApproximation	The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution. The degrees of freedom for the t-distribution for testing the slope difference is calculated using the containment method, and is equal to the total number of observations minus two times the total number of subjects. The exact calculation using the t distribution is only implemented for the fixed design.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
kMax	The maximum number of stages.
informationRates	The information rates. Defaults to $(1:kMax) / kMax$ if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type

	spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	The futility bounds on the conditional power scale.
futilitySlopeDiff	The futility bounds on the slope difference scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Details

We use the following random-effects model to compare two slopes:

$$y_{ij} = \alpha + (\beta + \gamma x_i)t_j + a_i + b_i t_j + e_{ij}$$

where

- α : overall intercept common across treatment groups due to randomization
- β : slope for the control group
- γ : difference in slopes between the active treatment and control groups
- x_i : treatment indicator for subject i , 1 for the active treatment and 0 for the control
- t_j : time point j for repeated measurements, $t_1 = 0 < t_2 < \dots < t_k$
- (a_i, b_i) : random intercept and random slope for subject i , $Var(a_i) = \sigma_a^2$, $Var(b_i) = \sigma_b^2$, $Corr(a_i, b_i) = \rho$
- e_{ij} : within-subject residual with variance σ_e^2

By accounting for randomization, we improve the efficiency for estimating the difference in slopes. The model also accommodates unequally spaced time points and missing data. Specifically, given a calendar time τ for an interim or final analysis, let k be the number of scheduled time points up to and including τ , subject to the follow-up duration for fixed follow-up designs. Let the observed time points be $t_1, t_2, \dots, t_k$, where $t_1 = 0$ denotes baseline.

For a subject in treatment group g with observed data pattern j , the design matrix for the fixed effects $(\alpha, \beta, \gamma)'$ is given by

$$X_{g,j} = (\mathbf{1}_j, \vec{t}_j, I(g=1)\vec{t}_j)$$

where $\mathbf{1}_j$ is a j -vector of ones, and $\vec{t}_j = (t_1, \dots, t_j)'$ is the column vector of observed time points. The design matrix for the random effects $(a_i, b_i)'$ is

$$Z_j = (\mathbf{1}_j, \vec{t}_j)$$

The variance-covariance matrix of the random effects is

$$D = \begin{pmatrix} \sigma_a^2 & \rho\sigma_a\sigma_b \\ \rho\sigma_a\sigma_b & \sigma_b^2 \end{pmatrix}$$

Therefore, the variance-covariance matrix for the observed data for the subject is

$$V_j = Z_j D Z_j' + \sigma_e^2 I_j$$

where I_j is the $j \times j$ identity matrix. Let π_g denote the proportion of subjects in group g . The information matrix for the fixed effects is

$$I = nJ$$

where

$$J = \sum_{g=1}^2 \pi_g \sum_{j=1}^k p_{g,j} X_{g,j}' V_j^{-1} X_{g,j}$$

and $p_{g,j}$ is the proportion of subjects in group g with observed data pattern j .

The variance of the estimator for the slope difference $\hat{\gamma}$ is given by

$$\text{Var}(\hat{\gamma}) = \frac{1}{n} J^{-1} [3, 3]$$

which can be used to calculate the power and sample size for the group sequential design to detect a slope difference.

Value

An S3 class `designSlopeDiffMMRM` object with three components:

- `overallResults`: A data frame containing the following variables:
 - `overallReject`: The overall rejection probability.
 - `alpha`: The overall significance level.
 - `attainedAlpha`: The attained significance level, which is different from the overall significance level in the presence of futility stopping.
 - `kMax`: The number of stages.

- theta: The parameter value.
- information: The maximum information.
- expectedInformationH1: The expected information under H1.
- expectedInformationH0: The expected information under H0.
- drift: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
- inflationFactor: The inflation factor (relative to the fixed design).
- numberOfSubjects: The maximum number of subjects.
- studyDuration: The maximum study duration.
- expectedNumberOfSubjectsH1: The expected number of subjects under H1.
- expectedNumberOfSubjectsH0: The expected number of subjects under H0.
- expectedStudyDurationH1: The expected study duration under H1.
- expectedStudyDurationH0: The expected study duration under H0.
- accrualDuration: The accrual duration.
- followupTime: The follow-up time.
- fixedFollowup: Whether a fixed follow-up design is used.
- slopeDiffH0: The slope difference under H0.
- slopeDiff: The slope difference under H1.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale.
 - futilityBounds: The futility boundaries on the Z-scale.
 - rejectPerStage: The probability for efficacy stopping.
 - futilityPerStage: The probability for futility stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeFutility: The cumulative probability for futility stopping.
 - cumulativeAlphaSpent: The cumulative alpha spent.
 - efficacyP: The efficacy boundaries on the p-value scale.
 - futilityP: The futility boundaries on the p-value scale.
 - information: The cumulative information.
 - efficacyStopping: Whether to allow efficacy stopping.
 - futilityStopping: Whether to allow futility stopping.
 - rejectPerStageH0: The probability for efficacy stopping under H0.
 - futilityPerStageH0: The probability for futility stopping under H0.
 - cumulativeRejectionH0: The cumulative probability for efficacy stopping under H0.
 - cumulativeFutilityH0: The cumulative probability for futility stopping under H0.
 - efficacySlopeDiff: The efficacy boundaries on the slope difference scale.
 - futilitySlopeDiff: The futility boundaries on the slope difference scale.
 - numberOfSubjects: The number of subjects.
 - analysisTime: The average time since trial start.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.

- parameterAlphaSpending: The parameter value for alpha spending.
- userAlphaSpending: The user defined alpha spending.
- typeBetaSpending: The type of beta spending.
- parameterBetaSpending: The parameter value for beta spending.
- userBetaSpending: The user defined beta spending.
- spendingTime: The error spending time at each analysis.
- allocationRatioPlanned: The allocation ratio for the active treatment versus control.
- accrualTime: A vector that specifies the starting time of piecewise Poisson enrollment time intervals.
- accrualIntensity: A vector of accrual intensities. One for each accrual time interval.
- piecewiseSurvivalTime: A vector that specifies the starting time of piecewise exponential survival time intervals.
- gamma1: The hazard rate for exponential dropout or a vector of hazard rates for piecewise exponential dropout for the active treatment group.
- gamma2: The hazard rate for exponential dropout or a vector of hazard rates for piecewise exponential dropout for the control group.
- w: The number of time units per measurement visit in a period.
- N: The number of measurement visits in a period.
- stdDev: The standard deviation of the residual.
- G: The covariance matrix for the random intercept and random slope.
- normalApproximation: The type of computation of the p-values. If TRUE, the variance is assumed to be known, otherwise the calculations are performed with the t distribution.
- rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Daniel O. Scharfstein, Anastasios A. Tsiatis, and James M. Robins. Semiparametric efficiency and its implication on the design and analysis of group-sequential studies. *Journal of the American Statistical Association* 1997; 92:1342-1350.

Examples

```
(design1 <- getDesignSlopeDiffMMRM(
  beta = 0.2, slopeDiff = log(1.15)/52,
  stDev = sqrt(.182),
  stDevIntercept = sqrt(.238960),
  stDevSlope = sqrt(.000057),
  corrInterceptSlope = .003688/sqrt(.238960*.000057),
  w = 8,
  N = 10000,
  accrualIntensity = 15,
  gamma1 = 1/(4.48*52),
  gamma2 = 1/(4.48*52),
  accrualDuration = NA,
```

```
followupTime = 8,  
alpha = 0.025))
```

getDesignTwoMultinom	<i>Power and Sample Size for Difference in Two-Sample Multinomial Responses</i>
----------------------	---

Description

Obtains the power given sample size or obtains the sample size given power for difference in two-sample multinomial responses.

Usage

```
getDesignTwoMultinom(  
  beta = NA_real_,  
  n = NA_real_,  
  ncats = NA_integer_,  
  pi1 = NA_real_,  
  pi2 = NA_real_,  
  allocationRatioPlanned = 1,  
  rounding = TRUE,  
  alpha = 0.05  
)
```

Arguments

beta	The type II error.
n	The total sample size.
ncats	The number of categories of the multinomial response.
pi1	The prevalence of each category for the treatment group. Only need to specify the valued for the first ncats-1 categories.
pi2	The prevalence of each category for the control group. Only need to specify the valued for the first ncats-1 categories.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
alpha	The two-sided significance level. Defaults to 0.05.

Details

A two-arm multinomial response design is used to test whether the prevalence of each category differs between two treatment arms. Let π_{gi} denote the prevalence of category i in group g , where $g = 1$ for the treatment group and $g = 2$ for the control group. The chi-square test statistic is given by

$$X^2 = \sum_{g=1}^2 \sum_{i=1}^C \frac{(n_{gi} - n_{g+}n_{+i}/n)^2}{n_{g+}n_{+i}/n}$$

where n_{gi} is the number of subjects in category i for group g , n_{g+} is the total number of subjects in group g , and n_{+i} is the total number of subjects in category i across both groups, and n is the total sample size.

- Under the null hypothesis, X^2 follows a chi-square distribution with $C - 1$ degrees of freedom.
- Under the alternative hypothesis, X^2 follows a non-central chi-square distribution with non-centrality parameter

$$\lambda = nr(1 - r) \sum_{i=1}^C \frac{(\pi_{1i} - \pi_{2i})^2}{r\pi_{1i} + (1 - r)\pi_{2i}}$$

where r is the randomization probability for the active treatment.

The sample size is chosen such that the power to reject the null hypothesis is at least $1 - \beta$ for a given significance level α .

Value

An S3 class `designTwoMultinom` object with the following components:

- `power`: The power to reject the null hypothesis.
- `alpha`: The two-sided significance level.
- `n`: The maximum number of subjects.
- `ncats`: The number of categories of the multinomial response.
- `pi1`: The prevalence of each category for the treatment group.
- `pi2`: The prevalence of each category for the control group.
- `effectsize`: The effect size for the chi-square test.
- `allocationRatioPlanned`: Allocation ratio for the active treatment versus control.
- `rounding`: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignTwoMultinom(
  beta = 0.1, ncats = 3, pi1 = c(0.3, 0.35),
  pi2 = c(0.2, 0.3), alpha = 0.05))
```

getDesignTwoOrdinal *Power and Sample Size for the Wilcoxon Test for Two-Sample Ordinal Response*

Description

Obtains the power given sample size or obtains the sample size given power for the Wilcoxon test for two-sample ordinal response.

Usage

```
getDesignTwoOrdinal(
  beta = NA_real_,
  n = NA_real_,
  ncats = NA_integer_,
  pi1 = NA_real_,
  pi2 = NA_real_,
  allocationRatioPlanned = 1,
  rounding = TRUE,
  alpha = 0.05
)
```

Arguments

beta	The type II error.
n	The total sample size.
ncats	The number of categories of the ordinal response.
pi1	The prevalence of each category for the treatment group. Only need to specify the valued for the first ncats-1 categories.
pi2	The prevalence of each category for the control group. Only need to specify the valued for the first ncats-1 categories.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
alpha	The significance level. Defaults to 0.025.

Details

A two-sample ordinal response design is used to test whether the ordinal response distributions differ between two treatment arms. Let π_{gi} denote the prevalence of category i in group g , where $g = 1$ represents the treatment group and $g = 2$ represents the control group.

The parameter of interest is

$$\theta = \sum_{i=1}^C w_i (\pi_{1i} - \pi_{2i})$$

where w_i is the midrank score for category i . The Z-test statistic is given by

$$Z = \hat{\theta} / \sqrt{\text{Var}(\hat{\theta})}$$

where $\hat{\theta}$ is the estimate of θ .

The midrank score w_i for category i is calculated as:

$$w_i = \sum_{j=1}^i \pi_j - 0.5\pi_i$$

where $\pi_i = r\pi_{1i} + (1-r)\pi_{2i}$ denotes the average prevalence of category i across both groups, and r is the randomization probability for the active treatment.

To understand the midrank score, consider $n\pi_i$ subjects in category i . The midrank score is the average rank of these subjects:

$$s_i = \frac{1}{n\pi_i} \sum_{j=1}^{n\pi_i} (n\pi_1 + \dots + n\pi_{i-1} + j)$$

This simplifies to

$$s_i = n \left(\sum_{j=1}^i \pi_j - 0.5\pi_i \right) + \frac{1}{2}$$

By dividing by n and ignoring $\frac{1}{2n}$, we obtain the midrank score w_i .

The variance of $\hat{\theta}$ can be derived from the multinomial distributions and is given by

$$\text{Var}(\hat{\theta}) = \frac{1}{n} \sum_{g=1}^2 \frac{1}{r_g} \left\{ \sum_{i=1}^C w_i^2 \pi_{gi} - \left(\sum_{i=1}^C w_i \pi_{gi} \right)^2 \right\}$$

where r_g is the randomization probability for group g .

The sample size is chosen such that the power to reject the null hypothesis is at least $1 - \beta$ for a given significance level α .

Value

An S3 class `designTwoOrdinal` object with the following components:

- `power`: The power to reject the null hypothesis.
- `alpha`: The two-sided significance level.
- `n`: The maximum number of subjects.
- `ncats`: The number of categories of the ordinal response.
- `pi1`: The prevalence of each category for the treatment group.
- `pi2`: The prevalence of each category for the control group.
- `meanscore1`: The mean midrank score for the treatment group.
- `meanscore2`: The mean midrank score for the control group.
- `allocationRatioPlanned`: Allocation ratio for the active treatment versus control.
- `rounding`: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignTwoOrdinal(
  beta = 0.1, ncats = 4, pi1 = c(0.55, 0.3, 0.1),
  pi2 = c(0.214, 0.344, 0.251), alpha = 0.025))
```

getDesignTwoWayANOVA *Power and Sample Size for Two-Way ANOVA*

Description

Obtains the power and sample size for two-way analysis of variance.

Usage

```
getDesignTwoWayANOVA(
  beta = NA_real_,
  n = NA_real_,
  nlevelsA = 2,
  nlevelsB = 2,
  means = NA_real_,
  stDev = 1,
  rounding = TRUE,
  alpha = 0.05
)
```

Arguments

beta	The type II error.
n	The total sample size.
nlevelsA	The number of groups for Factor A.
nlevelsB	The number of levels for Factor B.
means	The matrix of treatment means for Factors A and B combination.
stDev	The common standard deviation.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
alpha	The two-sided significance level. Defaults to 0.05.

Value

An S3 class designTwoWayANOVA object with the following components:

- alpha: The two-sided significance level.
- nlevelsA: The number of levels for Factor A.
- nlevelsB: The number of levels for Factor B.
- means: The matrix of treatment group means.
- stDev: The common standard deviation.
- effectsizeA: The effect size for Factor A.
- effectsizeB: The effect size for Factor B.
- effectsizeAB: The effect size for Factor A and Factor B interaction.
- rounding: Whether to round up sample size.
- powerdf: The data frame containing the power and sample size results. It has the following variables:
 - n: The sample size.
 - powerA: The power to reject the null hypothesis that there is no difference among Factor A levels.
 - powerB: The power to reject the null hypothesis that there is no difference among Factor B levels.
 - powerAB: The power to reject the null hypothesis that there is no interaction between Factor A and Factor B.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignTwoWayANOVA(
  beta = 0.1, nlevelsA = 2, nlevelsB = 2,
  means = matrix(c(0.5, 4.7, 0.4, 6.9), 2, 2, byrow = TRUE),
  stDev = 2, alpha = 0.05))
```

getDesignUnorderedBinom

Power and Sample Size for Unordered Multi-Sample Binomial Response

Description

Obtains the power given sample size or obtains the sample size given power for the chi-square test for unordered multi-sample binomial response.

Usage

```
getDesignUnorderedBinom(
  beta = NA_real_,
  n = NA_real_,
  ngroups = NA_integer_,
  pi = NA_real_,
  allocationRatioPlanned = NA_integer_,
  rounding = TRUE,
  alpha = 0.05
)
```

Arguments

beta	The type II error.
n	The total sample size.
ngroups	The number of treatment groups.
pi	The response probabilities for the treatment groups.
allocationRatioPlanned	Allocation ratio for the treatment groups.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
alpha	The two-sided significance level. Defaults to 0.05.

Details

A multi-sample binomial response design is used to test whether the response probabilities differ among multiple treatment arms. Let π_g denote the response probability in group $g = 1, \dots, G$, where G is the total number of treatment groups.

The chi-square test statistic is given by

$$X^2 = \sum_{g=1}^G \sum_{i=1}^2 \frac{(n_{gi} - n_{g+}n_{+i}/n)^2}{n_{g+}n_{+i}/n}$$

where n_{gi} is the number of subjects in category i for group g , n_{g+} is the total number of subjects in group g , and n_{+i} is the total number of subjects in category i across all groups, and n is the total sample size.

Let r_g denote the randomization probability for group g , and define the weighted average response probability across all groups as

$$\bar{\pi} = \sum_{g=1}^G r_g \pi_g$$

- Under the null hypothesis, X^2 follows a chi-square distribution with $G-1$ degrees of freedom.
- Under the alternative hypothesis, X^2 follows a non-central chi-square distribution with non-centrality parameter

$$\lambda = n \sum_{g=1}^G \frac{r_g (\pi_g - \bar{\pi})^2}{\bar{\pi}(1 - \bar{\pi})}$$

The sample size is chosen such that the power to reject the null hypothesis is at least $1 - \beta$ for a given significance level α .

Value

An S3 class designUnorderedBinom object with the following components:

- power: The power to reject the null hypothesis.
- alpha: The two-sided significance level.
- n: The maximum number of subjects.
- ngroups: The number of treatment groups.
- pi: The response probabilities for the treatment groups.
- effectsize: The effect size for the chi-square test.
- allocationRatioPlanned: Allocation ratio for the treatment groups.
- rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignUnorderedBinom(
  beta = 0.1, ngroups = 3, pi = c(0.1, 0.25, 0.5), alpha = 0.05))
```

```
getDesignUnorderedMultinom
```

Power and Sample Size for Unordered Multi-Sample Multinomial Response

Description

Obtains the power given sample size or obtains the sample size given power for the chi-square test for unordered multi-sample multinomial response.

Usage

```
getDesignUnorderedMultinom(
  beta = NA_real_,
  n = NA_real_,
  ngroups = NA_integer_,
  ncats = NA_integer_,
  pi = NA_real_,
  allocationRatioPlanned = NA_integer_,
  rounding = TRUE,
  alpha = 0.05
)
```

Arguments

beta	The type II error.
n	The total sample size.
ngroups	The number of treatment groups.
ncats	The number of categories of the multinomial response.
pi	The matrix of response probabilities for the treatment groups. It should have ngroups rows and ncats-1 or ncats columns.
allocationRatioPlanned	Allocation ratio for the treatment groups.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
alpha	The two-sided significance level. Defaults to 0.05.

Details

A multi-sample multinomial response design is used to test whether the response probabilities differ among multiple treatment arms. Let π_{gi} denote the response probability for category $i = 1, \dots, C$ in group $g = 1, \dots, G$, where G is the total number of treatment groups, and C is the total number of categories for the response variable.

The chi-square test statistic is given by

$$X^2 = \sum_{g=1}^G \sum_{i=1}^C \frac{(n_{gi} - n_{g+}n_{+i}/n)^2}{n_{g+}n_{+i}/n}$$

where n_{gi} is the number of subjects in category i for group g , n_{g+} is the total number of subjects in group g , and n_{+i} is the total number of subjects in category i across all groups, and n is the total sample size.

Let r_g denote the randomization probability for group g , and define the weighted average response probability for category i across all groups as

$$\bar{\pi}_i = \sum_{g=1}^G r_g \pi_{gi}$$

- Under the null hypothesis, X^2 follows a chi-square distribution with $(G-1)(C-1)$ degrees of freedom.
- Under the alternative hypothesis, X^2 follows a non-central chi-square distribution with non-centrality parameter

$$\lambda = n \sum_{g=1}^G \sum_{i=1}^C \frac{r_g (\pi_{gi} - \bar{\pi}_i)^2}{\bar{\pi}_i}$$

The sample size is chosen such that the power to reject the null hypothesis is at least $1 - \beta$ for a given significance level α .

Value

An S3 class `designUnorderedMultinom` object with the following components:

- `power`: The power to reject the null hypothesis.
- `alpha`: The two-sided significance level.
- `n`: The maximum number of subjects.
- `ngroups`: The number of treatment groups.
- `ncats`: The number of categories of the multinomial response.
- `pi`: The response probabilities for the treatment groups.
- `effectsize`: The effect size for the chi-square test.
- `allocationRatioPlanned`: Allocation ratio for the treatment groups.
- `rounding`: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(design1 <- getDesignUnorderedMultinom(
  beta = 0.1, ngroups = 3, ncats = 4,
  pi = matrix(c(0.230, 0.320, 0.272,
               0.358, 0.442, 0.154,
               0.142, 0.036, 0.039),
             3, 3, byrow = TRUE),
  allocationRatioPlanned = c(2, 2, 1),
  alpha = 0.05))
```

getDesignWilcoxon

Group Sequential Design for Two-Sample Wilcoxon Test

Description

Obtains the power given sample size or obtains the sample size given power for a group sequential design for two-sample Wilcoxon test.

Usage

```
getDesignWilcoxon(
  beta = NA_real_,
  n = NA_real_,
  pLarger = 0.6,
  allocationRatioPlanned = 1,
  rounding = TRUE,
```

```

kMax = 1L,
informationRates = NA_real_,
efficacyStopping = NA_integer_,
futilityStopping = NA_integer_,
criticalValues = NA_real_,
alpha = 0.025,
typeAlphaSpending = "sfOF",
parameterAlphaSpending = NA_real_,
userAlphaSpending = NA_real_,
futilityBounds = NA_real_,
futilityCP = NA_real_,
futilitypLarger = NA_real_,
typeBetaSpending = "none",
parameterBetaSpending = NA_real_,
userBetaSpending = NA_real_,
spendingTime = NA_real_
)

```

Arguments

beta	The type II error.
n	The total sample size.
pLarger	The probability that a randomly chosen sample from the treatment group is larger than a randomly chosen sample from the control group under the alternative hypothesis.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
kMax	The maximum number of stages.
informationRates	The information rates. Fixed prior to the trial. Defaults to (1:kMax) / kMax if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for

	Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sf0F".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	The futility bounds on the conditional power scale.
futilitypLarger	The futility bounds on the pLarger scale.
typeBetaSpending	The type of beta spending. One of the following: "sf0F" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.

Details

The Mann-Whitney U test is a non-parametric test for the difference in location between two independent groups. It is also known as the Wilcoxon rank-sum test. The test is based on the ranks of the data rather than the actual values, making it robust to outliers and non-normal distributions. The test statistic is the number of times a randomly chosen sample from the treatment group is larger than a randomly chosen sample from the control group, i.e.,

$$W_{XY} = \sum_{i=1}^{n_1} \sum_{j=1}^{n_2} I(X_i > Y_j)$$

where X_i and Y_j are the samples from the treatment and control groups, respectively. The test is often used when the data do not meet the assumptions of the t-test, such as non-normality or unequal variances. The test is also applicable to ordinal data. The test is one-sided, meaning that it only tests whether the treatment group is larger than the control group. Asymptotically,

$$\frac{W_{XY} - n_1 n_2 / 2}{\sqrt{n_1 n_2 (n + 1) / 12}} \sim N(0, 1) \quad \text{under } H_0$$

where n_1 and n_2 are the sample sizes of the treatment and control groups, respectively, and $n = n_1 + n_2$. Let $\theta = P(X > Y)$, and $\hat{\theta} = \frac{1}{nm} W_{XY}$. It follows that

$$\sqrt{n}(\hat{\theta} - 1/2) \sim N\left(0, \frac{1}{12r(1-r)}\right) \quad \text{under } H_0$$

where $r = n_1/(n_1 + n_2)$ is the randomization probability for the active treatment group. This formulation allows for group sequential testing with futility stopping and efficacy stopping.

Value

An S3 class `designWilcoxon` object with three components:

- `overallResults`: A data frame containing the following variables:
 - `overallReject`: The overall rejection probability.
 - `alpha`: The overall significance level.
 - `attainedAlpha`: The attained significance level, which is different from the overall significance level in the presence of futility stopping..
 - `kMax`: The number of stages.
 - `theta`: The parameter value.
 - `information`: The maximum information.
 - `expectedInformationH1`: The expected information under H1.
 - `expectedInformationH0`: The expected information under H0.
 - `drift`: The drift parameter, equal to $\theta \cdot \sqrt{\text{information}}$.
 - `inflationFactor`: The inflation factor (relative to the fixed design).
 - `numberOfSubjects`: The maximum number of subjects.
 - `expectedNumberOfSubjectsH1`: The expected number of subjects under H1.
 - `expectedNumberOfSubjectsH0`: The expected number of subjects under H0.
 - `pLarger`: The probability that a randomly chosen sample from the treatment group is larger than a randomly chosen sample from the control group under the alternative hypothesis.
- `byStageResults`: A data frame containing the following variables:
 - `informationRates`: The information rates.
 - `efficacyBounds`: The efficacy boundaries on the Z-scale.
 - `futilityBounds`: The futility boundaries on the Z-scale.
 - `rejectPerStage`: The probability for efficacy stopping.
 - `futilityPerStage`: The probability for futility stopping.
 - `cumulativeRejection`: The cumulative probability for efficacy stopping.
 - `cumulativeFutility`: The cumulative probability for futility stopping.
 - `cumulativeAlphaSpent`: The cumulative alpha spent.
 - `efficacyP`: The efficacy boundaries on the p-value scale.
 - `futilityP`: The futility boundaries on the p-value scale.
 - `information`: The cumulative information.
 - `efficacyStopping`: Whether to allow efficacy stopping.
 - `futilityStopping`: Whether to allow futility stopping.

- rejectPerStageH0: The probability for efficacy stopping under H0.
- futilityPerStageH0: The probability for futility stopping under H0.
- cumulativeRejectionH0: The cumulative probability for efficacy stopping under H0.
- cumulativeFutilityH0: The cumulative probability for futility stopping under H0.
- efficacyPLarger: The efficacy boundaries on the proportion of pairs of samples from the two treatment groups with the sample from the treatment group greater than that from the control group.
- futilityPLarger: The futility boundaries on the proportion of pairs of samples from the two treatment groups with the sample from the treatment group greater than that from the control group.
- numberOfSubjects: The number of subjects.
- settings: A list containing the following input parameters:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for alpha spending.
 - userAlphaSpending: The user defined alpha spending.
 - typeBetaSpending: The type of beta spending.
 - parameterBetaSpending: The parameter value for beta spending.
 - userBetaSpending: The user defined beta spending.
 - spendingTime: The error spending time at each analysis.
 - allocationRatioPlanned: Allocation ratio for the active treatment versus control.
 - rounding: Whether to round up sample size.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Example 1: fixed design
(design1 <- getDesignWilcoxon(
  beta = 0.1, n = NA,
  pLarger = pnorm((8 - 2)/sqrt(2*25^2)), alpha = 0.025))

# Example 2: group sequential design
(design2 <- getDesignWilcoxon(
  beta = 0.1, n = NA,
  pLarger = pnorm((8 - 2)/sqrt(2*25^2)), alpha = 0.025,
  kMax = 3, typeAlphaSpending = "sfOF"))
```

getDesign_multiarm	<i>Power and Sample Size for Multi-Arm Multi-Stage Design</i>
--------------------	---

Description

Computes either the maximum information and stopping boundaries for a multi-arm multi-stage design, or the achieved power when the maximum information and stopping boundaries are provided.

Usage

```
getDesign_multiarm(
  beta = NA_real_,
  IMax = NA_real_,
  theta = NA_real_,
  M = NA_integer_,
  r = 1,
  corr_known = TRUE,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,
  futilityCP = NULL,
  futilityTheta = NULL,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
  spendingTime = NA_real_,
  nthreads = 0
)
```

Arguments

beta	Type II error rate. Provide either beta or IMax; the other should be missing.
IMax	Maximum information for any active arm versus the common control. Provide either IMax or beta; the other should be missing.
theta	A vector of length M representing the true treatment effects for each active arm versus the common control. The global null is $\theta_i = 0$ for all i , and alternatives are one-sided: $\theta_i > 0$ for at least one $i = 1, \dots, M$.
M	Number of active treatment arms.
r	Randomization ratio of each active arm to the common control.

corr_known	Logical. If TRUE, the correlation between Wald statistics is derived from the randomization ratio r as $r/(r + 1)$. If FALSE, a conservative correlation of 0 is used.
kMax	Number of sequential looks.
informationRates	A numeric vector of information rates fixed before the trial. If unspecified, defaults to $(1 : kMax)/kMax$.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	The matrix of by-level upper boundaries on the max z-test statistic scale for efficacy stopping. The first column is for level M , the second column is for level $M - 1$, and so on, with the last column for level 1. If left unspecified, the critical values will be computed based on the specified alpha spending function.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	A numeric vector of length $kMax - 1$ specifying the futility boundaries on the max z-test statistic scale for futility stopping.
futilityCP	A numeric vector of length $kMax - 1$ specifying the futility boundaries on the conditional power scale for futility stopping.
futilityTheta	A numeric vector of length $kMax - 1$ specifying the futility boundaries on the parameter scale for futility stopping.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".

userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A numeric vector of length kMax specifying the error spending time at each analysis. Values must be strictly increasing and ends at 1. If omitted, defaults to informationRates.
nthreads	The number of threads to use (0 leaves the RcppParallel setting unchanged).

Details

If `corr_known` is FALSE, critical boundaries are computed assuming independence among the Wald statistics in each stage (a conservative assumption). Power calculations, however, use the correlation implied by the randomization ratio r .

Value

An S3 object of class `multiarm` with the following components:

- `overallResults`: A data frame containing:
 - `overallReject`: Overall probability of rejecting the global null hypothesis.
 - `alpha`: Overall significance level.
 - `attainedAlpha`: The attained significance level, which is different from the overall significance level in the presence of futility stopping.
 - `M`: Number of active arms.
 - `r`: Randomization ratio per active arm versus control.
 - `corr_known`: Whether the correlation among Wald statistics was assumed known.
 - `kMax`: Number of stages.
 - `information`: Maximum information for any active arm versus control.
 - `expectedInformationH1`: The expected information under H1.
 - `expectedInformationH0`: The expected information under H0.
- `byStageResults`: A data frame containing:
 - `informationRates`: Information rates at each analysis.
 - `efficacyBounds`: Efficacy boundaries on the max Z-scale.
 - `futilityBounds`: Futility boundaries on the max Z-scale.
 - `rejectPerStage`: Probability of efficacy stopping at each stage.
 - `futilityPerStage`: Probability of futility stopping at each stage.
 - `cumulativeRejection`: Cumulative probability of efficacy stopping.
 - `cumulativeFutility`: Cumulative probability of futility stopping.
 - `cumulativeAlphaSpent`: Cumulative alpha spent.
 - `efficacyTheta`: Efficacy boundaries on the parameter scale.
 - `futilityTheta`: Futility boundaries on the parameter scale.
 - `efficacyP`: Efficacy boundaries on the p-value scale.
 - `futilityP`: Futility boundaries on the p-value scale.
 - `information`: Cumulative information for any active arm versus control at each analysis.
 - `efficacyStopping`: Indicator of whether efficacy stopping is permitted at each stage.

- futilityStopping: Indicator of whether futility stopping is permitted at each stage.
- rejectPerStageH0: The probability for efficacy stopping under H0.
- futilityPerStageH0: The probability for futility stopping under H0.
- cumulativeRejectionH0: The cumulative probability for efficacy stopping under H0.
- cumulativeFutilityH0: The cumulative probability for futility stopping under H0.
- settings: A list of input settings:
 - typeAlphaSpending: The type of alpha spending.
 - parameterAlphaSpending: The parameter value for the chosen alpha spending function.
 - userAlphaSpending: The user-specified alpha spending values.
 - typeBetaSpending: The type of beta spending.
 - parameterBetaSpending: The parameter value for the chosen beta spending function.
 - userBetaSpending: The user-specified beta spending values.
 - spendingTime: The error-spending time at each analysis.
- byLevelBounds: A data frame containing the efficacy boundaries for each level of testing (i.e., number of active arms remaining) and each stage. Columns include:
 - level: Number of active arms remaining (1 to M).
 - stage: Stage index (1 to $kMax$).
 - efficacyBounds: Efficacy boundaries on the max Z-scale for the given level and stage.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

References

Ping Gao, Yingqiu Li. Adaptive multiple comparison sequential design (AMCSD) for clinical trials. Journal of Biopharmaceutical Statistics, 2024, 34(3), 424-440.

Examples

```
# Example 1: obtain the maximum information given power
(design1 <- getDesign_multiarm(
  beta = 0.1, theta = c(0.3, 0.5), M = 2, r = 1.0,
  kMax = 3, informationRates = seq(1, 3)/3,
  alpha = 0.025, typeAlphaSpending = "OF", nthreads = 1))

# Example 2: obtain power given the maximum information
(design2 <- getDesign_multiarm(
  IMax = 110/(2*1^2), theta = c(0.3, 0.5), M = 2, r = 1.0,
  kMax = 3, informationRates = seq(1, 3)/3,
  alpha = 0.025, typeAlphaSpending = "OF", nthreads = 1))
```

getDesign_seamless	<i>Power and Sample Size for Phase 2/3 Seamless Design</i>
--------------------	--

Description

Computes either the maximum information and stopping boundaries for a phase 2/3 seamless design, or the achieved power when the maximum information and stopping boundaries are provided. Both efficacy and futility stopping can be incorporated.

Usage

```
getDesign_seamless(
  beta = NA_real_,
  IMax = NA_real_,
  theta = NA_real_,
  M = NA_integer_,
  r = 1,
  corr_known = TRUE,
  K = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,
  futilityCP = NULL,
  futilityTheta = NULL,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
  spendingTime = NA_real_,
  nthreads = 0
)
```

Arguments

beta	Type II error rate. Provide either beta or IMax; the other should be missing.
IMax	Maximum information for any active arm versus the common control. Provide either IMax or beta; the other should be missing.
theta	A vector of length M representing the true treatment effects for each active arm versus the common control. The global null is $\theta_i = 0$ for all i , and alternatives are one-sided: $\theta_i > 0$ for at least one $i = 1, \dots, M$.
M	Number of active treatment arms in Phase 2.

<code>r</code>	Randomization ratio of each active arm to the common control in Phase 2.
<code>corr_known</code>	Logical. If TRUE, the correlation between Wald statistics in Phase 2 is derived from the randomization ratio r as $r/(r + 1)$. If FALSE, a conservative correlation of 0 is used. This option is only used for critical value calculations; the correlation is always derived from r for power calculations.
<code>K</code>	Number of sequential looks in Phase 3.
<code>informationRates</code>	A numeric vector of information rates fixed before the trial. If unspecified, defaults to $(1 : (K + 1))/(K + 1)$.
<code>efficacyStopping</code>	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
<code>futilityStopping</code>	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
<code>criticalValues</code>	The upper boundaries on the Z-statistic scale for the most promising arm in Phase 2 and the Z statistics for the selected arm in Phase 3. If missing, boundaries will be computed based on the specified alpha spending function.
<code>alpha</code>	The significance level. Defaults to 0.025.
<code>typeAlphaSpending</code>	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
<code>parameterAlphaSpending</code>	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
<code>userAlphaSpending</code>	The user defined alpha spending. Cumulative alpha spent up to each stage.
<code>futilityBounds</code>	A numeric vector of length K specifying futility boundaries on the Z scale at the end of Phase 2 for the most promising arm and on the Z scale for the $K - 1$ analyses in Phase 3. The final analysis uses the efficacy boundary as the futility boundary.
<code>futilityCP</code>	A numeric vector of length K specifying futility boundaries on the conditional power scale.
<code>futilityTheta</code>	A numeric vector of length K specifying futility boundaries on the parameter scale.
<code>typeBetaSpending</code>	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".

parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A numeric vector of length $K + 1$ specifying the error spending time at each analysis. Values must be strictly increasing and end at 1. If omitted, defaults to informationRates.
nthreads	The number of threads to use (0 leaves the RcppParallel setting unchanged).

Details

If `corr_known` is FALSE, critical boundaries are computed assuming independence among the Phase-2 Wald statistics (a conservative assumption). Power calculations, however, use the correlation implied by the randomization ratio r .

Futility boundaries may be supplied directly on the Z scale, derived from conditional power, derived from parameter values, or computed from a beta spending function.

Value

An S3 object of class `seamless` with the following components:

- `overallResults`: A data frame containing:
 - `overallReject`: Overall probability of rejecting the null hypothesis.
 - `alpha`: Overall significance level.
 - `attainedAlpha`: The attained significance level, which may differ from `alpha` in the presence of futility stopping.
 - `M`: Number of active arms in Phase 2.
 - `r`: Randomization ratio per active arm versus control in Phase 2.
 - `corr_known`: Whether the phase-2 correlation was assumed known.
 - `K`: Number of looks in Phase 3.
 - `information`: Maximum information for any active arm versus control.
 - `expectedInformationH1`: Expected information under the alternative.
 - `expectedInformationH0`: Expected information under the null.
 - `informationOverall`: Maximum information for the overall study.
 - `expectedInformationH1`: Expected information under the alternative for the overall study.
 - `expectedInformationH0`: Expected information under the null for the overall study.
- `byStageResults`: A data frame containing:
 - `informationRates`: Information rates at each analysis.
 - `efficacyBounds`: Efficacy boundaries on the Z scale.
 - `futilityBounds`: Futility boundaries on the Z scale.
 - `rejectPerStage`: Probability of efficacy stopping at each stage.
 - `futilityPerStage`: Probability of futility stopping at each stage.
 - `cumulativeRejection`: Cumulative probability of efficacy stopping.

- cumulativeFutility: Cumulative probability of futility stopping.
- cumulativeAlphaSpent: Cumulative alpha spent.
- efficacyTheta: Efficacy boundaries on the parameter scale.
- futilityTheta: Futility boundaries on the parameter scale.
- efficacyP: Efficacy boundaries on the p-value scale.
- futilityP: Futility boundaries on the p-value scale.
- information: Cumulative information at each analysis.
- informationOverall: Cumulative information for the overall study at each analysis.
- efficacyStopping: Indicator of whether efficacy stopping is permitted.
- futilityStopping: Indicator of whether futility stopping is permitted.
- rejectPerStageH0: Probability of efficacy stopping under the global null.
- futilityPerStageH0: Probability of futility stopping under the global null.
- cumulativeRejectionH0: Cumulative probability of efficacy stopping under the global null.
- cumulativeFutilityH0: Cumulative probability of futility stopping under the global null.
- byArmResults: A data frame containing:
 - theta: Parameter values for the active arms.
 - selectionProb: Probability an arm is selected at the end of Phase 2.
 - powerByArm: Probability of rejecting the null for each arm by trial end.
 - condPowerByArm: Conditional power for each arm given it was selected as the most promising arm at the end of Phase 2.
- settings: A list of input settings:
 - typeAlphaSpending: Type of alpha spending function.
 - parameterAlphaSpending: Parameter value for the chosen alpha spending function.
 - userAlphaSpending: User-specified alpha spending values.
 - typeBetaSpending: Type of beta spending function.
 - parameterBetaSpending: Parameter value for the chosen beta spending function.
 - userBetaSpending: User-specified beta spending values.
 - spendingTime: Error-spending times at each analysis.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Ping Gao, Yingqiu Li. Adaptive two-stage seamless sequential design for clinical trials. Journal of Biopharmaceutical Statistics, 2025, 35(4), 565-587.

Examples

```
# Example 1: obtain the maximum information given power with no futility
(design1 <- getDesign_seamless(
  beta = 0.1, theta = c(0.3, 0.5), M = 2, r = 1.0,
  K = 2, informationRates = seq(1, 3)/3,
  alpha = 0.025, typeAlphaSpending = "OF", nthreads = 1))

# Example 2: obtain power given the maximum information and a futility rule
(design2 <- getDesign_seamless(
  IMax = 110/(2*1^2), theta = c(0.3, 0.5), M = 2, r = 1.0,
  K = 2, informationRates = seq(1, 3)/3,
  alpha = 0.025, typeAlphaSpending = "OF",
  futilityBounds = c(0.0, 0.5), nthreads = 1))

# Example 3: derive futility boundaries using beta spending
(design3 <- getDesign_seamless(
  beta = 0.1, theta = c(-log(0.5), -log(0.7)),
  M = 2, r = 1.0, corr_known = FALSE,
  K = 2, informationRates = seq(1, 3)/3,
  alpha = 0.025, typeAlphaSpending = "sfOF",
  typeBetaSpending = "sfHSD", parameterBetaSpending = -2,
  nthreads = 1))
```

getDurationFromNevents

Range of Accrual Duration for Target Number of Events

Description

Obtains a range of accrual duration to reach the target number of events.

Usage

```
getDurationFromNevents(
  nevents = NA_real_,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  followupTime = NA_real_,
  fixedFollowup = FALSE,
  npoints = 23L
)
```

Arguments

nevents	The target number of events.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
lambda1	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.
lambda2	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
followupTime	Follow-up time for the last enrolled subjects. Must be provided for fixed follow-up design.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
npoints	The number of accrual duration time points. Defaults to 23.

Value

A data frame of the following variables:

- nevents: The target number of events.
- fixedFollowup: Whether a fixed follow-up design is used.
- accrualDuration: The accrual duration.
- subjects: The total number of subjects.
- followupTime: The follow-up time for the last enrolled subject.
- studyDuration: The study duration.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Piecewise accrual, piecewise exponential survivals, and 5% dropout by
# the end of 1 year.
```

```
getDurationFromNevents(
  nevents = 80, allocationRatioPlanned = 1,
  accrualTime = seq(0, 8),
  accrualIntensity = 26/9*seq(1, 9),
  piecewiseSurvivalTime = c(0, 6),
  lambda1 = c(0.0533, 0.0309),
  lambda2 = c(0.0533, 0.0533),
  gamma1 = -log(1-0.05)/12,
  gamma2 = -log(1-0.05)/12,
  fixedFollowup = FALSE)
```

```
getNeventsFromHazardRatio
```

Required Number of Events Given Hazard Ratio

Description

Obtains the required number of events given the hazard ratios under the null and alternative hypotheses for a group sequential design.

Usage

```
getNeventsFromHazardRatio(
  beta = 0.2,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,
  futilityCP = NULL,
  futilityHR = NULL,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
```

```

    spendingTime = NA_real_,
    hazardRatioH0 = 1,
    hazardRatio = NA_real_,
    allocationRatioPlanned = 1,
    rounding = TRUE
)

```

Arguments

beta	Type II error. Defaults to 0.2.
kMax	The maximum number of stages.
informationRates	The information rates in terms of number of events. Fixed prior to the trial. Defaults to (1:kMax) / kMax if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsiatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	A vector of length kMax - 1 for the futility bounds on the conditional power scale.
futilityHR	A vector of length kMax - 1 for the futility bounds on the hazard ratio scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".

parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.
hazardRatioH0	Hazard ratio under the null hypothesis for the active treatment versus control. Defaults to 1 for superiority test.
hazardRatio	Hazard ratio under the alternative hypothesis for the active treatment versus control.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
rounding	Whether to round up the number of events. Defaults to 1 for rounding.

Value

The required number of events.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
getNeventsFromHazardRatio(
  beta = 0.2, kMax = 2,
  informationRates = c(0.5,1),
  alpha = 0.025, typeAlphaSpending = "sf0F",
  typeBetaSpending = "sfP",
  hazardRatio = 0.673)
```

getNFromNeventsAndFollowup

Accrual Duration for Target Number of Events and Follow-up Time

Description

Obtains the accrual duration and sample size to reach the target number of events given follow-up time.

Usage

```
getNFromNeventsAndFollowup(
  nevents = NA_real_,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  followupTime = NA_real_,
  fixedFollowup = FALSE
)
```

Arguments

<code>nevents</code>	The target number of events.
<code>allocationRatioPlanned</code>	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
<code>accrualTime</code>	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., <code>c(0, 3)</code> breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
<code>accrualIntensity</code>	A vector of accrual intensities. One for each accrual time interval.
<code>piecewiseSurvivalTime</code>	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., <code>c(0, 6)</code> breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
<code>stratumFraction</code>	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
<code>lambda1</code>	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.
<code>lambda2</code>	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
<code>gamma1</code>	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
<code>gamma2</code>	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
<code>followupTime</code>	Follow-up time for the last enrolled subjects. Must be provided.
<code>fixedFollowup</code>	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.

Value

A data frame of the following variables:

- nevents: The target number of events.
- fixedFollowup: Whether a fixed follow-up design is used.
- accrualDuration: The accrual duration.
- subjects: The total number of subjects.
- followupTime: The follow-up time for the last enrolled subject.
- studyDuration: The study duration.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Piecewise accrual, piecewise exponential survivals, and 5% dropout by
# the end of 1 year.
```

```
getNFromNeventsAndFollowup(
  nevents = 80, allocationRatioPlanned = 1,
  accrualTime = seq(0, 8),
  accrualIntensity = 26/9*seq(1, 9),
  piecewiseSurvivalTime = c(0, 6),
  lambda1 = c(0.0533, 0.0309),
  lambda2 = c(0.0533, 0.0533),
  gamma1 = -log(1-0.05)/12,
  gamma2 = -log(1-0.05)/12,
  followupTime = 18,
  fixedFollowup = FALSE)
```

getRCI

Repeated Confidence Interval for Group Sequential Design

Description

Obtains the repeated confidence interval for a group sequential trial.

Usage

```
getRCI(
  L = NA_integer_,
  zL = NA_real_,
  IMax = NA_real_,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
```

```

criticalValues = NA_real_,
alpha = 0.025,
typeAlphaSpending = "sfOF",
parameterAlphaSpending = NA_real_,
spendingTime = NA_real_
)

```

Arguments

<code>L</code>	The look of interest.
<code>zL</code>	The z-test statistic at the look.
<code>IMax</code>	The maximum information of the trial.
<code>informationRates</code>	The information rates up to look L.
<code>efficacyStopping</code>	Indicators of whether efficacy stopping is allowed at each stage up to look L. Defaults to true if left unspecified.
<code>criticalValues</code>	The upper boundaries on the z-test statistic scale for efficacy stopping up to look L.
<code>alpha</code>	The significance level. Defaults to 0.025.
<code>typeAlphaSpending</code>	The type of alpha spending for the trial. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsiatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early efficacy stopping. Defaults to "sfOF".
<code>parameterAlphaSpending</code>	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
<code>spendingTime</code>	The error spending time up to look L. Defaults to missing, in which case, it is the same as <code>informationRates</code> .

Value

A data frame with the following components:

- `pvalue`: Repeated p-value for rejecting the null hypothesis.
- `thetahat`: Point estimate of the parameter.
- `cilevel`: Confidence interval level.
- `lower`: Lower bound of repeated confidence interval.
- `upper`: Upper bound of repeated confidence interval.

If `typeAlphaSpending` is "OF", "P", "WT", or "none", then `informationRates`, `efficacyStopping`, and `spendingTime` must be of full length `kMax`, and `informationRates` and `spendingTime` must end with 1.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Christopher Jennison and Bruce W. Turnbull. Interim analyses: the repeated confidence interval approach (with discussion). J R Stat Soc Series B. 1989;51:305-361.

Examples

```
# group sequential design with 90% power to detect delta = 6
delta <- 6
sigma <- 17
n <- 282
(des1 <- getDesign(IMax = n/(4*sigma^2), theta = delta, kMax = 3,
                  alpha = 0.05, typeAlphaSpending = "sfHSD",
                  parameterAlphaSpending = -4))

# results at the second look
L <- 2
n1 <- n*2/3
delta1 <- 7
sigma1 <- 20
zL <- delta1/sqrt(4/n1*sigma1^2)

# repeated confidence interval
getRCI(L = L, zL = zL, IMax = n/(4*sigma1^2),
       informationRates = c(1/3, 2/3), alpha = 0.05,
       typeAlphaSpending = "sfHSD", parameterAlphaSpending = -4)
```

hazard_pd

Hazard Function for Progressive Disease (PD) Given Correlation Between PD and OS

Description

Computes the hazard function of a piecewise exponential distribution for progressive disease (PD), such that the resulting hazard function for progression-free survival (PFS) closely matches a given piecewise hazard for PFS.

Usage

```
hazard_pd(
  piecewiseSurvivalTime = 0L,
  hazard_pfs = NA_real_,
  hazard_os = NA_real_,
  rho_pd_os = 0.5
)
```

Arguments

piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
hazard_pfs	A scalar or numeric vector specifying the hazard(s) for PFS based on a piecewise exponential distribution.
hazard_os	A scalar or numeric vector specifying the hazard(s) for overall survival (OS) based on a piecewise exponential distribution.
rho_pd_os	A numeric value specifying the correlation between PD and OS times.

Details

This function determines the hazard vector λ_{pd} for the piecewise exponential distribution of PD, so that the implied survival function for PFS time, $T_{pfs} = \min(T_{pd}, T_{os})$, closely matches the specified piecewise exponential distribution for PFS with hazard vector λ_{pfs} .

To achieve this, we simulate (Z_{pd}, Z_{os}) from a standard bivariate normal distribution with correlation ρ . Then, $U_{pd} = \Phi(Z_{pd})$ and $U_{os} = \Phi(Z_{os})$ are generated, where Φ denotes the standard normal CDF.

The times to PD and OS are obtained via the inverse transform method using quantile functions of the piecewise exponential distribution:

$$T_{pd} = \text{qpwexp}(U_{pd}, u, \lambda_{pd})$$

$$T_{os} = \text{qpwexp}(U_{os}, u, \lambda_{os})$$

where $u = \text{piecewiseSurvivalTime}$.

The function solves for λ_{pd} such that the survival function of T_{pfs} closely matches that of a piecewise exponential distribution with hazard λ_{pfs} :

$$P(\min(T_{pd}, T_{os}) > t) = S_{pfs}(t)$$

Since

$$Z_{pd} = \Phi^{-1}(\text{ppwexp}(T_{pd}, u, \lambda_{pd}))$$

and

$$Z_{os} = \Phi^{-1}(\text{ppwexp}(T_{os}, u, \lambda_{os}))$$

we have

$$P(\min(T_{pd}, T_{os}) > t) = P(Z_{pd} > \Phi^{-1}(\text{ppwexp}(t, u, \lambda_{pd})), Z_{os} > \Phi^{-1}(\text{ppwexp}(t, u, \lambda_{os})))$$

while

$$S_{pfs}(t) = 1 - \text{ppwexp}(t, u, \lambda_{pfs})$$

Matching is performed sequentially at the internal cut points $u_2, \dots, u_J$ and at the point $u_J + \log(2)/\lambda_{pfs,J}$ for the final interval, as well as the percentile points at 10%, 20%, ..., 90%, and 95% to solve for $\lambda_{pd,1}, \dots, \lambda_{pd,K}$, where K is the total number of unique cut points.

Value

A list with the following components:

- `piecewiseSurvivalTime`: A vector that specifies the starting time points of the intervals for the piecewise exponential distribution for PD.
- `hazard_pd`: A numeric vector representing the calculated hazard rates for the piecewise exponential distribution of PD.
- `hazard_os`: A numeric vector representing the hazard rates for the piecewise exponential distribution of OS at the same time points as PD.
- `rho_pd_os`: The correlation between PD and OS times (as input).

Author(s)

Kaifeng Lu (<kaifengl@gmail.com>)

Examples

```
u <- c(0, 1, 3, 4)
lambda1 <- c(0.0151, 0.0403, 0.0501, 0.0558)
lambda2 <- 0.0145
rho_pd_os <- 0.5
hazard_pd(u, lambda1, lambda2, rho_pd_os)
```

<code>hazard_sub</code>	<i>Hazard Function for Sub Population</i>
-------------------------	---

Description

Computes the hazard function of a piecewise exponential distribution for the biomarker negative sub population, such that the resulting survival function for the ITT population closely matches a given piecewise survival function.

Usage

```
hazard_sub(
  piecewiseSurvivalTime = 0L,
  hazard_itt = NA_real_,
  hazard_pos = NA_real_,
  p_pos = NA_real_
)
```

Arguments

<code>piecewiseSurvivalTime</code>	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
<code>hazard_itt</code>	A scalar or numeric vector specifying the hazard(s) for the ITT population based on a piecewise exponential distribution.
<code>hazard_pos</code>	A scalar or numeric vector specifying the hazard(s) for the biomarker positive sub population based on a piecewise exponential distribution.
<code>p_pos</code>	A numeric value specifying the prevalence of the biomarker positive sub population.

Details

This function determines the hazard vector λ_{neg} for the piecewise exponential distribution of the biomarker negative sub population, so that the implied survival function for the ITT population closely matches the specified piecewise exponential distribution with hazard vector λ_{itt} .

Let p_{pos} be the prevalence of the biomarker positive sub population, then the survival function for the ITT population is given by

$$S_{\text{itt}}(t) = p_{\text{pos}}S_{\text{pos}}(t) + (1 - p_{\text{pos}})S_{\text{neg}}(t)$$

where $S_{\text{pos}}(t)$ and $S_{\text{neg}}(t)$ are the survival functions for the biomarker positive and biomarker negative sub populations, respectively.

Matching is performed sequentially at the internal cutpoints $u_2, \dots, u_J$ and at the point $u_J + \log(2)/\lambda_{\text{itt},J}$ for the final interval, as well as the percentile points at 10%, 20%, ..., 90%, and 95%, to solve for $\lambda_{\text{neg},1}, \dots, \lambda_{\text{neg},K}$, where K is the total number of unique cut points.

Value

A list with the following components:

- `piecewiseSurvivalTime`: A vector that specifies the starting time points of the intervals for the piecewise exponential distribution for the biomarker negative sub population.
- `hazard_pos`: A numeric vector representing the hazard rates for the piecewise exponential distribution of the biomarker positive sub population at the same time points as the biomarker negative sub population.
- `hazard_neg`: A numeric vector representing the estimated hazard rates for the piecewise exponential distribution of the biomarker negative sub population.
- `p_pos`: The prevalence of the biomarker positive sub population (as input).

Author(s)

Kaifeng Lu (<kaifenglu@gmail.com>)

Examples

```
u <- c(0, 1, 3, 4)
lambda_itt <- c(0.0151, 0.0403, 0.0501, 0.0558)
lambda_pos <- c(0.0115, 0.0302, 0.0351, 0.0404)
p_pos <- 0.3
hazard_sub(u, lambda_itt, lambda_pos, p_pos)
```

heart	<i>Stanford heart transplant data from the survival package</i>
-------	---

Description

Survival of patients on the waiting list for the Stanford heart transplant program.

start, stop, event entry and exit time and status for the time interval

age age-48 years

year year of acceptance (in years after Nov 1, 1967)

surgery prior bypass surgery 1=yes, 0=no

transplant received transplant 1=yes, 0=no

id patient id

Usage

heart

Format

An object of class `data.frame` with 172 rows and 8 columns.

<i>hedgesg</i>	<i>Hedges' g Effect Size</i>
----------------	------------------------------

Description

Obtains Hedges' g estimate and confidence interval of effect size.

Usage

```
hedgesg(tstat, m, ntilde, cilevel = 0.95)
```

Arguments

tstat	The value of the t-test statistic for comparing two treatment conditions.
m	The degrees of freedom for the t-test.
ntilde	The normalizing sample size to convert the standardized treatment difference to the t-test statistic, i.e., $tstat = \sqrt{ntilde} * meanDiff / stDev$.
cilevel	The confidence interval level. Defaults to 0.95.

Details

Hedges' g is an effect size measure commonly used in meta-analysis to quantify the difference between two groups. It's an improvement over Cohen's d , particularly when dealing with small sample sizes.

The formula for Hedges' g is

$$g = c(m)d$$

where d is Cohen's d effect size estimate, and $c(m)$ is the bias correction factor,

$$d = (\hat{\mu}_1 - \hat{\mu}_2) / \hat{\sigma}$$

$$c(m) = 1 - \frac{3}{4m - 1}.$$

Since $c(m) < 1$, Cohen's d overestimates the true effect size, $\delta = (\mu_1 - \mu_2) / \sigma$. Since

$$t = \sqrt{\tilde{n}}d$$

we have

$$g = \frac{c(m)}{\sqrt{\tilde{n}}}t$$

where t has a noncentral t distribution with m degrees of freedom and noncentrality parameter $\sqrt{\tilde{n}}\delta$.

The asymptotic variance of g can be approximated by

$$Var(g) = \frac{1}{\tilde{n}} + \frac{g^2}{2m}.$$

The confidence interval for δ can be constructed using normal approximation.

For two-sample mean difference with sample size n_1 for the treatment group and n_2 for the control group, we have $\tilde{n} = \frac{n_1 n_2}{n_1 + n_2}$ and $m = n_1 + n_2 - 2$ for pooled variance estimate.

Value

A data frame with the following variables:

- tstat: The value of the t test statistic.
- m: The degrees of freedom for the t-test.
- ntilde: The normalizing sample size to convert the standardized treatment difference to the t-test statistic.
- g: Hedges' g effect size estimate.

- varg: Variance of g.
- lower: The lower confidence limit for effect size.
- upper: The upper confidence limit for effect size.
- cilevel: The confidence interval level.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

References

Larry V. Hedges. Distribution theory for Glass's estimator of effect size and related estimators. *Journal of Educational Statistics* 1981; 6:107-128.

Examples

```
n1 = 7
n2 = 8
meanDiff = 0.444
stDev = 1.201
m = n1+n2-2
ntilde = n1*n2/(n1+n2)
tstat = sqrt(ntilde)*meanDiff/stDev

hedgesg(tstat, m, ntilde)
```

immdef

Simulated CONCORDE trial data from the rpsftm package

Description

Patients were randomly assigned to receive treatment immediately or deferred, and those in the deferred arm could cross over and receive treatment. The primary endpoint was time to disease progression.

id Patient identification number

def Indicator that the participant was assigned to the deferred treatment arm

imm Indicator that the participant was assigned to the immediate treatment arm

censyrs The censoring time, in years, corresponding to the close of study minus the time of entry for each patient

xo Indicator that crossover occurred

xoyrs The time, in years, from entry to switching, or 0 for patients in the immediate arm

prog Indicator of disease progression (1), or censoring (0)

progyrs Time, in years, from entry to disease progression or censoring

entry The time of entry into the study, measured in years from the date of randomisation

Usage

`immdef`

Format

An object of class `data.frame` with 1000 rows and 9 columns.

ingots	<i>The binary data from Cox and Snell (1989, pp. 10-11).</i>
--------	--

Description

The dataset consists of the number of ingots not ready for rolling and the number of ingots ready for rolling for a number of combinations of heating time and soaking time.

Usage

`ingots`

Format

An object of class `tbl_df` (inherits from `tbl`, `data.frame`) with 25 rows and 4 columns.

Details

- Heat The heating time
- Soak The soaking time
- NotReady Response indicator, with a value 1 for units not ready for rolling (event) and a value of 0 for units ready for rolling (nonevent)
- Freq The frequency of occurrence of each combination of Heat, Soak, and NotReady

kmdiff	<i>Estimate of Milestone Survival Difference</i>
--------	--

Description

Obtains the estimate of milestone survival difference between two treatment groups.

Usage

```
kmdiff(
  data,
  stratum = "",
  treat = "treat",
  time = "time",
  time2 = "",
  event = "event",
  weight = "",
  milestone = 0,
  survDiffH0 = 0,
  conflev = 0.95
)
```

Arguments

data	The input data frame that contains the following variables: • stratum: The stratum. • treat: The treatment. • time: The follow-up time for right censored data, or the left end of each interval for counting process data. • time2: The right end of each interval for counting process data. Intervals are assumed to be open on the left and closed on the right, and event indicates whether an event occurred at the right end of each interval. • event: The event indicator, 1=event, 0=no event. • weight: The weight for each observation.
stratum	The name of the stratum variable in the input data.
treat	The name of the treatment variable in the input data.
time	The name of the time variable or the left end of each interval for counting process data in the input data.
time2	The name of the right end of each interval for counting process data in the input data.
event	The name of the event variable in the input data.
weight	The name of the weight variable in the input data.
milestone	The milestone time at which to calculate the survival probability.
survDiffH0	The difference in milestone survival probabilities under the null hypothesis. Defaults to 0 for superiority test.
conflev	The level of the two-sided confidence interval for the difference in milestone survival probabilities. Defaults to 0.95.

Value

A data frame with the following variables:

- milestone: The milestone time relative to randomization.

- survDiffH0: The difference in milestone survival probabilities under the null hypothesis.
- surv1: The estimated milestone survival probability for the treatment group.
- surv2: The estimated milestone survival probability for the control group.
- survDiff: The estimated difference in milestone survival probabilities.
- vsurv1: The variance for surv1.
- vsurv2: The variance for surv2.
- sesurvDiff: The standard error for survDiff.
- survDiffZ: The Z-statistic value.
- survDiffPValue: The two-sided p-value.
- lower: The lower bound of confidence interval.
- upper: The upper bound of confidence interval.
- conflev: The level of confidence interval.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
kmest(data = rawdata[rawdata$iterationNumber == 1, ],
      stratum = "stratum", treat = "treatmentGroup",
      time = "timeUnderObservation", event = "event",
      milestone = 12)
```

kmest

Kaplan-Meier Estimates of Survival Curve

Description

Obtains the Kaplan-Meier estimates of the survival curve.

Usage

```
kmest(
  data,
  stratum = "",
  time = "time",
  time2 = "",
  event = "event",
  weight = "",
  conftype = "log-log",
  conflev = 0.95,
  keep_censor = FALSE
)
```

Arguments

data	The input data frame that contains the following variables: • stratum: The stratum. • time: The follow-up time for right censored data, or the left end of each interval for counting process data. • time2: The right end of each interval for counting process data. Intervals are assumed to be open on the left and closed on the right, and event indicates whether an event occurred at the right end of each interval. • event: The event indicator, 1=event, 0=no event. • weight: The weight for each observation.
stratum	The name(s) of the stratum variable(s) in the input data.
time	The name of the time variable or the left end of each interval for counting process data in the input data.
time2	The name of the right end of each interval for counting process data in the input data.
event	The name of the event variable in the input data.
weight	The name of the weight variable in the input data.
conftype	The type of the confidence interval. One of "none", "plain", "log", "log-log" (the default), or "arcsin". The arcsin option bases the intervals on $\text{asin}(\sqrt{\text{survival}})$.
conflev	The level of the two-sided confidence interval for the survival probabilities. Defaults to 0.95.
keep_censor	Whether to retain the censoring time in the output data frame.

Value

A data frame with the following variables:

- size: The number of subjects in the stratum.
- time: The event time.
- nrisk: The number of subjects at risk.
- nevent: The number of subjects having the event.
- ncensor: The number of censored subjects.
- surv: The Kaplan-Meier estimate of the survival probability.
- sesurv: The standard error of the estimated survival probability based on the Greenwood formula.
- lower: The lower bound of confidence interval if requested.
- upper: The upper bound of confidence interval if requested.
- conflev: The level of confidence interval if requested.
- conftype: The type of confidence interval if requested.
- stratum: The stratum.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
krmest(data = aml, stratum = "x", time = "time", event = "status")
```

kmpower

Power for Difference in Milestone Survival Probabilities

Description

Estimates the power for testing the difference in milestone survival probabilities in a two-sample survival design.

Usage

```
kmpower(
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,
  futilityCP = NULL,
  futilitySurvDiff = NULL,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  milestone = NA_real_,
  survDiffH0 = 0,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = FALSE,
```

```

    spendingTime = NA_real_,
    studyDuration = NA_real_
  )

```

Arguments

kMax	The maximum number of stages.
informationRates	The information rates. Defaults to $(1:kMax) / kMax$ if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	A vector of length kMax - 1 for the futility bounds on the conditional power scale.
futilitySurvDiff	A vector of length kMax - 1 for the futility bounds on the milestone survival difference scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
milestone	The milestone time at which to calculate the survival probability.

survDiffH0	The difference in milestone survival probabilities under the null hypothesis. Defaults to 0 for superiority test.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
lambda1	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.
lambda2	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.
studyDuration	Study duration for fixed follow-up design. Defaults to missing, which is to be replaced with the sum of accrualDuration and followupTime. If provided, the value is allowed to be less than the sum of accrualDuration and followupTime.

Value

An S3 class kmpower object with 4 components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.

- alpha: The overall significance level.
- numberOfEvents: The total number of events.
- numberOfDropouts: The total number of dropouts.
- numbeOfSubjects: The total number of subjects.
- numberOfMilestone: The total number of subjects reaching milestone.
- studyDuration: The total study duration.
- information: The maximum information.
- expectedNumberOfEvents: The expected number of events.
- expectedNumberOfDropouts: The expected number of dropouts.
- expectedNumberOfSubjects: The expected number of subjects.
- expectedNumberOfMilestone: The expected number of subjects reaching milestone.
- expectedStudyDuration: The expected study duration.
- expectedInformation: The expected information.
- accrualDuration: The accrual duration.
- followupTime: The follow-up duration.
- fixedFollowup: Whether a fixed follow-up design is used.
- kMax: The number of stages.
- milestone: The milestone time relative to randomization.
- survDiffH0: The difference in milestone survival probabilities under the null hypothesis.
- surv1: The milestone survival probability for the treatment group.
- surv2: The milestone survival probability for the control group.
- survDiff: The difference in milestone survival probabilities, equal to $\text{surv1} - \text{surv2}$.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale.
 - futilityBounds: The futility boundaries on the Z-scale.
 - rejectPerStage: The probability for efficacy stopping.
 - futilityPerStage: The probability for futility stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeFutility: The cumulative probability for futility stopping.
 - cumulativeAlphaSpent: The cumulative alpha spent.
 - numberOfEvents: The number of events.
 - numberOfDropouts: The number of dropouts.
 - numberOfSubjects: The number of subjects.
 - numberOfMilestone: The number of subjects reaching milestone.
 - analysisTime: The average time since trial start.
 - efficacySurvDiff: The efficacy boundaries on the survival difference scale.
 - futilitySurvDiff: The futility boundaries on the survival difference scale.
 - efficacyP: The efficacy boundaries on the p-value scale.
 - futilityP: The futility boundaries on the p-value scale.
 - information: The cumulative information.
 - efficacyStopping: Whether to allow efficacy stopping.

- `futilityStopping`: Whether to allow futility stopping.
- `settings`: A list containing the following input parameters: `typeAlphaSpending`, `parameterAlphaSpending`, `userAlphaSpending`, `typeBetaSpending`, `parameterBetaSpending`, `allocationRatioPlanned`, `accrualTime`, `accrualIntensity`, `piecewiseSurvivalTime`, `stratumFraction`, `lambda1`, `lambda2`, `gamma1`, `gamma2`, and `spendingTime`.
- `byTreatmentCounts`: A list containing the following counts by treatment group:
 - `numberOfEvents1`: The number of events by stage for the treatment group.
 - `numberOfDropouts1`: The number of dropouts by stage for the treatment group.
 - `numberOfSubjects1`: The number of subjects by stage for the treatment group.
 - `numberOfMilestone1`: The number of subjects reaching milestone by stage for the active treatment group.
 - `numberOfEvents2`: The number of events by stage for the control group.
 - `numberOfDropouts2`: The number of dropouts by stage for the control group.
 - `numberOfSubjects2`: The number of subjects by stage for the control group.
 - `numberOfMilestone2`: The number of subjects reaching milestone by stage for the control group.
 - `expectedNumberOfEvents1`: The expected number of events for the treatment group.
 - `expectedNumberOfDropouts1`: The expected number of dropouts for the active treatment group.
 - `expectedNumberOfSubjects1`: The expected number of subjects for the active treatment group.
 - `expectedNumberOfMilestone1`: The expected number of subjects reaching milestone for the active treatment group.
 - `expectedNumberOfEvents2`: The expected number of events for control group.
 - `expectedNumberOfDropouts2`: The expected number of dropouts for the control group.
 - `expectedNumberOfSubjects2`: The expected number of subjects for the control group.
 - `expectedNumberOfMilestone2`: The expected number of subjects reaching milestone for the control group.

Author(s)

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Examples

```
# Piecewise accrual, piecewise exponential survival, and 5% dropout by
# the end of 1 year.
```

```
kmpower(kMax = 2, informationRates = c(0.8, 1),
        alpha = 0.025, typeAlphaSpending = "sfOF",
        milestone = 18,
        allocationRatioPlanned = 1, accrualTime = seq(0, 8),
        accrualIntensity = 26/9*seq(1, 9),
        piecewiseSurvivalTime = c(0, 6),
        stratumFraction = c(0.2, 0.8),
        lambda1 = c(0.0533, 0.0309, 1.5*0.0533, 1.5*0.0309),
        lambda2 = c(0.0533, 0.0533, 1.5*0.0533, 1.5*0.0533),
```

```
gamma1 = -log(1-0.05)/12,
gamma2 = -log(1-0.05)/12, accrualDuration = 22,
followupTime = 18, fixedFollowup = FALSE)
```

kmpower1s

Power for One-Sample Milestone Survival Probability

Description

Estimates the power, stopping probabilities, and expected sample size in a one-group survival design.

Usage

```
kmpower1s(
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sf0F",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,
  futilityCP = NULL,
  futilitySurv = NULL,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  milestone = NA_real_,
  survH0 = NA_real_,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  lambda = NA_real_,
  gamma = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = FALSE,
  spendingTime = NA_real_,
  studyDuration = NA_real_
)
```

Arguments

kMax	The maximum number of stages.
informationRates	The information rates. Defaults to $(1:kMax) / kMax$ if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	A vector of length kMax - 1 for the futility bounds on the conditional power scale.
futilitySurv	A vector of length kMax - 1 for the futility bounds on the milestone survival scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
milestone	The milestone time at which to calculate the survival probability.
survH0	The milestone survival probability under the null hypothesis.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., c(0, 3) breaks the time axis into 2 accrual intervals: [0, 3) and [3, ∞).

accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
lambda	A vector of hazard rates for the event in each analysis time interval by stratum under the alternative hypothesis.
gamma	The hazard rate for exponential dropout or a vector of hazard rates for piecewise exponential dropout. Defaults to 0 for no dropout.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
spendingTime	A vector of length <code>kMax</code> for the error spending time at each analysis. Defaults to missing, in which case, it is the same as <code>informationRates</code> .
studyDuration	Study duration for fixed follow-up design. Defaults to missing, which is to be replaced with the sum of <code>accrualDuration</code> and <code>followupTime</code> . If provided, the value is allowed to be less than the sum of <code>accrualDuration</code> and <code>followupTime</code> .

Value

An S3 class `kmpower1s` object with 3 components:

- `overallResults`: A data frame containing the following variables:
 - `overallReject`: The overall rejection probability.
 - `alpha`: The overall significance level.
 - `numberOfEvents`: The total number of events.
 - `numbeOfSubjects`: The total number of subjects.
 - `numberOfMilestone`: The total number of subjects reaching milestone.
 - `studyDuration`: The total study duration.
 - `information`: The maximum information.
 - `expectedNumberOfEvents`: The expected number of events.
 - `expectedNumberOfSubjects`: The expected number of subjects.
 - `expectedNumberOfMilestone`: The expected number of subjects reaching milestone.
 - `expectedStudyDuration`: The expected study duration.
 - `expectedInformation`: The expected information.
 - `accrualDuration`: The accrual duration.
 - `followupTime`: The follow-up duration.
 - `fixedFollowup`: Whether a fixed follow-up design is used.

- kMax: The number of stages.
- milestone: The milestone time to calculate the survival probability.
- survH0: The milestone survival probability under the null hypothesis.
- surv: The milestone survival probability under the alternative hypothesis.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale.
 - futilityBounds: The futility boundaries on the Z-scale.
 - rejectPerStage: The probability for efficacy stopping.
 - futilityPerStage: The probability for futility stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeFutility: The cumulative probability for futility stopping.
 - cumulativeAlphaSpent: The cumulative alpha spent.
 - numberOfEvents: The number of events.
 - numberOfDropouts: The number of dropouts.
 - numberOfSubjects: The number of subjects.
 - numberOfMilestone: The number of subjects reaching milestone.
 - analysisTime: The average time since trial start.
 - efficacySurv: The efficacy boundaries on the milestone survival probability scale.
 - futilitySurv: The futility boundaries on the milestone survival probability scale.
 - efficacyP: The efficacy boundaries on the p-value scale.
 - futilityP: The futility boundaries on the p-value scale.
 - information: The cumulative information.
 - efficacyStopping: Whether to allow efficacy stopping.
 - futilityStopping: Whether to allow futility stopping.
- settings: A list containing the following input parameters: typeAlphaSpending, parameterAlphaSpending, userAlphaSpending, typeBetaSpending, parameterBetaSpending, accrualTime, accrualIntensity, piecewiseSurvivalTime, stratumFraction, lambda, gamma, and spendingTime.

Author(s)

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See Also

[kmstat](#)

Examples

```
kmpower1s(kMax = 2, informationRates = c(0.8, 1),
  alpha = 0.025, typeAlphaSpending = "sfOF",
  milestone = 18, survH0 = 0.30,
  accrualTime = seq(0, 8),
  accrualIntensity = 26/9*seq(1, 9),
  piecewiseSurvivalTime = c(0, 6),
```

```
stratumFraction = c(0.2, 0.8),
lambda = c(0.0533, 0.0309, 1.5*0.0533, 1.5*0.0309),
gamma = -log(1-0.05)/12, accrualDuration = 22,
followupTime = 18, fixedFollowup = FALSE)
```

kmpowerequiv

Power for Equivalence in Milestone Survival Probability Difference

Description

Obtains the power for equivalence in milestone survival probability difference.

Usage

```
kmpowerequiv(
  kMax = 1L,
  informationRates = NA_real_,
  criticalValues = NA_real_,
  alpha = 0.05,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  milestone = NA_real_,
  survDiffLower = NA_real_,
  survDiffUpper = NA_real_,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = FALSE,
  spendingTime = NA_real_,
  studyDuration = NA_real_
)
```

Arguments

kMax The maximum number of stages.

informationRates The information rates. Defaults to $(1:kMax) / kMax$ if left unspecified.

criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level for each of the two one-sided tests. Defaults to 0.05.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsiatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
milestone	The milestone time at which to calculate the survival probability.
survDiffLower	The lower equivalence limit of milestone survival probability difference.
survDiffUpper	The upper equivalence limit of milestone survival probability difference.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
lambda1	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.
lambda2	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
accrualDuration	Duration of the enrollment period.

<code>followupTime</code>	Follow-up time for the last enrolled subject.
<code>fixedFollowup</code>	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
<code>spendingTime</code>	A vector of length <code>kMax</code> for the error spending time at each analysis. Defaults to missing, in which case, it is the same as <code>informationRates</code> .
<code>studyDuration</code>	Study duration for fixed follow-up design. Defaults to missing, which is to be replaced with the sum of <code>accrualDuration</code> and <code>followupTime</code> . If provided, the value is allowed to be less than the sum of <code>accrualDuration</code> and <code>followupTime</code> .

Value

An S3 class `kmpowerequiv` object with 4 components:

- `overallResults`: A data frame containing the following variables:
 - `overallReject`: The overall rejection probability.
 - `alpha`: The overall significance level.
 - `numberOfEvents`: The total number of events.
 - `numberOfSubjects`: The total number of subjects.
 - `studyDuration`: The total study duration.
 - `information`: The maximum information.
 - `expectedNumberOfEvents`: The expected number of events.
 - `expectedNumberOfSubjects`: The expected number of subjects.
 - `expectedStudyDuration`: The expected study duration.
 - `expectedInformation`: The expected information.
 - `kMax`: The number of stages.
 - `milestone`: The milestone time relative to randomization.
 - `survDiffLower`: The lower equivalence limit of milestone survival probability difference.
 - `survDiffUpper`: The upper equivalence limit of milestone survival probability difference.
 - `surv1`: The milestone survival probability for the treatment group.
 - `surv2`: The milestone survival probability for the control group.
 - `survDiff`: The milestone survival probability difference.
 - `accrualDuration`: The accrual duration.
 - `followupTime`: The follow-up duration.
 - `fixedFollowup`: Whether a fixed follow-up design is used.
- `byStageResults`: A data frame containing the following variables:
 - `informationRates`: The information rates.
 - `efficacyBounds`: The efficacy boundaries on the Z-scale for each of the two one-sided tests.
 - `rejectPerStage`: The probability for efficacy stopping.
 - `cumulativeRejection`: The cumulative probability for efficacy stopping.
 - `cumulativeAlphaSpent`: The cumulative alpha for each of the two one-sided tests.

- cumulativeAttainedAlphaH10: The cumulative alpha attained under H10.
- cumulativeAttainedAlphaH20: The cumulative alpha attained under H20.
- numberOfEvents: The number of events.
- numberOfDropouts: The number of dropouts.
- numberOfSubjects: The number of subjects.
- numberOfMilestone: The number of subjects reaching milestone.
- analysisTime: The average time since trial start.
- efficacySurvDiffLower: The efficacy boundaries on the milestone survival probability difference scale for the one-sided null hypothesis at the lower equivalence limit.
- efficacySurvDiffUpper: The efficacy boundaries on the milestone survival probability difference scale for the one-sided null hypothesis at the upper equivalence limit.
- efficacyP: The efficacy bounds on the p-value scale for each of the two one-sided tests.
- information: The cumulative information.
- settings: A list containing the following input parameters: typeAlphaSpending, parameterAlphaSpending, userAlphaSpending, allocationRatioPlanned, accrualTime, accrualIntensity, piecewiseSurvivalTime, stratumFraction, lambda1, lambda2, gamma1, gamma2, and spendingTime.
- byTreatmentCounts: A list containing the following counts by treatment group:
 - numberOfEvents1: The number of events by stage for the treatment group.
 - numberOfDropouts1: The number of dropouts by stage for the treatment group.
 - numberOfSubjects1: The number of subjects by stage for the treatment group.
 - numberOfMilestone1: The number of subjects reaching milestone by stage for the active treatment group.
 - numberOfEvents2: The number of events by stage for the control group.
 - numberOfDropouts2: The number of dropouts by stage for the control group.
 - numberOfSubjects2: The number of subjects by stage for the control group.
 - numberOfMilestone2: The number of subjects reaching milestone by stage for the control group.
 - expectedNumberOfEvents1: The expected number of events for the treatment group.
 - expectedNumberOfDropouts1: The expected number of dropouts for the active treatment group.
 - expectedNumberOfSubjects1: The expected number of subjects for the active treatment group.
 - expectedNumberOfMilestone1: The expected number of subjects reaching milestone for the active treatment group.
 - expectedNumberOfEvents2: The expected number of events for control group.
 - expectedNumberOfDropouts2: The expected number of dropouts for the control group.
 - expectedNumberOfSubjects2: The expected number of subjects for the control group.
 - expectedNumberOfMilestone2: The expected number of subjects reaching milestone for the control group.

Author(s)

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See Also[kmstat](#)**Examples**

```

kmpowerequiv(kMax = 2, informationRates = c(0.5, 1),
             alpha = 0.05, typeAlphaSpending = "sfOF",
             milestone = 18,
             survDiffLower = -0.13, survDiffUpper = 0.13,
             allocationRatioPlanned = 1, accrualTime = seq(0, 8),
             accrualIntensity = 26/9*seq(1, 9),
             piecewiseSurvivalTime = c(0, 6),
             stratumFraction = c(0.2, 0.8),
             lambda1 = c(0.0533, 0.0533, 1.5*0.0533, 1.5*0.0533),
             lambda2 = c(0.0533, 0.0533, 1.5*0.0533, 1.5*0.0533),
             gamma1 = -log(1-0.05)/12,
             gamma2 = -log(1-0.05)/12, accrualDuration = 22,
             followupTime = 18, fixedFollowup = FALSE)

```

kmsamplesize

*Sample Size for Difference in Milestone Survival Probabilities***Description**

Obtains the needed accrual duration given power, accrual intensity, and follow-up time, the needed follow-up time given power, accrual intensity, and accrual duration, or the needed absolute accrual intensity given power, relative accrual intensity, accrual duration, and follow-up time in a two-group survival design.

Usage

```

kmsamplesize(
  beta = 0.2,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,
  futilityCP = NULL,
  futilitySurvDiff = NULL,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,

```

```

    userBetaSpending = NA_real_,
    milestone = NA_real_,
    survDiffH0 = 0,
    allocationRatioPlanned = 1,
    accrualTime = 0L,
    accrualIntensity = NA_real_,
    piecewiseSurvivalTime = 0L,
    stratumFraction = 1L,
    lambda1 = NA_real_,
    lambda2 = NA_real_,
    gamma1 = 0L,
    gamma2 = 0L,
    accrualDuration = NA_real_,
    followupTime = NA_real_,
    fixedFollowup = FALSE,
    spendingTime = NA_real_,
    rounding = TRUE
)

```

Arguments

beta Type II error. Defaults to 0.2.

kMax The maximum number of stages.

informationRates The information rates. Defaults to $(1:kMax) / kMax$ if left unspecified.

efficacyStopping Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.

futilityStopping Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.

criticalValues Upper boundaries on the z-test statistic scale for stopping for efficacy.

alpha The significance level. Defaults to 0.025.

typeAlphaSpending The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".

parameterAlphaSpending The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".

userAlphaSpending The user defined alpha spending. Cumulative alpha spent up to each stage.

futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	A vector of length kMax - 1 for the futility bounds on the conditional power scale.
futilitySurvDiff	A vector of length kMax - 1 for the futility bounds on the milestone survival difference scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
milestone	The milestone time at which to calculate the survival probability.
survDiffH0	The difference in milestone survival probabilities under the null hypothesis. Defaults to 0 for superiority test.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., c(0, 3) breaks the time axis into 2 accrual intervals: [0, 3) and [3, ∞).
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., c(0, 6) breaks the time axis into 2 event intervals: [0, 6) and [6, ∞). Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
lambda1	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.
lambda2	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.

gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.

Value

A list of two components:

- resultsUnderH1: An S3 class kmpower object under the alternative hypothesis.
- resultsUnderH0: An S3 class kmpower object under the null hypothesis.

Author(s)

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See Also

[kmpower](#)

Examples

```
# Example 1: Obtains follow-up time given power, accrual intensity,
# and accrual duration for variable follow-up. Of note, the power
# reaches the maximum when the follow-up time equals milestone.

kmsamplesize(beta = 0.25, kMax = 2, informationRates = c(0.8, 1),
              alpha = 0.025, typeAlphaSpending = "sf0F",
              milestone = 18,
              allocationRatioPlanned = 1, accrualTime = seq(0, 8),
              accrualIntensity = 26/9*seq(1, 9),
              piecewiseSurvivalTime = c(0, 6),
              stratumFraction = c(0.2, 0.8),
              lambda1 = c(0.0533, 0.0309, 1.5*0.0533, 1.5*0.0309),
              lambda2 = c(0.0533, 0.0533, 1.5*0.0533, 1.5*0.0533),
              gamma1 = -log(1-0.05)/12,
              gamma2 = -log(1-0.05)/12, accrualDuration = 22,
              followupTime = NA, fixedFollowup = FALSE)

# Example 2: Obtains accrual intensity given power, accrual duration, and
# follow-up time for variable follow-up

kmsamplesize(beta = 0.2, kMax = 2, informationRates = c(0.8, 1),
```

```

alpha = 0.025, typeAlphaSpending = "sf0F",
milestone = 18,
allocationRatioPlanned = 1, accrualTime = seq(0, 8),
accrualIntensity = 26/9*seq(1, 9),
piecewiseSurvivalTime = c(0, 6),
stratumFraction = c(0.2, 0.8),
lambda1 = c(0.0533, 0.0309, 1.5*0.0533, 1.5*0.0309),
lambda2 = c(0.0533, 0.0533, 1.5*0.0533, 1.5*0.0533),
gamma1 = -log(1-0.05)/12,
gamma2 = -log(1-0.05)/12, accrualDuration = 22,
followupTime = 18, fixedFollowup = FALSE)

# Example 3: Obtains accrual duration given power, accrual intensity, and
# follow-up time for fixed follow-up

kmsamplesize(beta = 0.2, kMax = 2, informationRates = c(0.8, 1),
alpha = 0.025, typeAlphaSpending = "sf0F",
milestone = 18,
allocationRatioPlanned = 1, accrualTime = seq(0, 8),
accrualIntensity = 26/9*seq(1, 9),
piecewiseSurvivalTime = c(0, 6),
stratumFraction = c(0.2, 0.8),
lambda1 = c(0.0533, 0.0309, 1.5*0.0533, 1.5*0.0309),
lambda2 = c(0.0533, 0.0533, 1.5*0.0533, 1.5*0.0533),
gamma1 = -log(1-0.05)/12,
gamma2 = -log(1-0.05)/12, accrualDuration = NA,
followupTime = 18, fixedFollowup = TRUE)

```

kmsamplesize1s

Sample Size for One-Sample Milestone Survival Probability

Description

Obtains the needed accrual duration given power and follow-up time, the needed follow-up time given power and accrual duration, or the needed absolute accrual rates given power, accrual duration, follow-up duration, and relative accrual rates in a one-group survival design.

Usage

```

kmsamplesize1s(
  beta = 0.2,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,

```

```

typeAlphaSpending = "sfOF",
parameterAlphaSpending = NA_real_,
userAlphaSpending = NA_real_,
futilityBounds = NULL,
futilityCP = NULL,
futilitySurv = NULL,
typeBetaSpending = "none",
parameterBetaSpending = NA_real_,
userBetaSpending = NA_real_,
milestone = NA_real_,
survH0 = NA_real_,
accrualTime = 0L,
accrualIntensity = NA_real_,
piecewiseSurvivalTime = 0L,
stratumFraction = 1L,
lambda = NA_real_,
gamma = 0L,
accrualDuration = NA_real_,
followupTime = NA_real_,
fixedFollowup = FALSE,
spendingTime = NA_real_,
rounding = TRUE
)

```

Arguments

beta	Type II error. Defaults to 0.2.
kMax	The maximum number of stages.
informationRates	The information rates. Defaults to (1:kMax) / kMax if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".

userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	A vector of length kMax - 1 for the futility bounds on the conditional power scale.
futilitySurv	A vector of length kMax - 1 for the futility bounds on the milestone survival scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
milestone	The milestone time at which to calculate the survival probability.
survH0	The milestone survival probability under the null hypothesis.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., c(0, 3) breaks the time axis into 2 accrual intervals: [0, 3) and [3, ∞).
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., c(0, 6) breaks the time axis into 2 event intervals: [0, 6) and [6, ∞). Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
lambda	A vector of hazard rates for the event in each analysis time interval by stratum under the alternative hypothesis.
gamma	The hazard rate for exponential dropout or a vector of hazard rates for piecewise exponential dropout. Defaults to 0 for no dropout.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.

Value

A list of two components:

- resultsUnderH1: An S3 class kmpower1s object under the alternative hypothesis.
- resultsUnderH0: An S3 class kmpower1s object under the null hypothesis.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

See Also

[kmpower1s](#)

Examples

```
# Example 1: Obtains follow-up duration given power, accrual intensity,
# and accrual duration for variable follow-up
```

```
kmsamplesize1s(beta = 0.2, kMax = 2,
  informationRates = c(0.8, 1),
  alpha = 0.025, typeAlphaSpending = "sfOF",
  milestone = 18, survH0 = 0.30,
  accrualTime = seq(0, 8),
  accrualIntensity = 26/9*seq(1, 9),
  piecewiseSurvivalTime = c(0, 6),
  stratumFraction = c(0.2, 0.8),
  lambda = c(0.0533, 0.0309, 1.5*0.0533, 1.5*0.0309),
  gamma = -log(1-0.05)/12, accrualDuration = 22,
  followupTime = NA, fixedFollowup = FALSE)
```

```
# Example 2: Obtains accrual intensity given power, accrual duration, and
# follow-up duration for variable follow-up
```

```
kmsamplesize1s(beta = 0.2, kMax = 2,
  informationRates = c(0.8, 1),
  alpha = 0.025, typeAlphaSpending = "sfOF",
  milestone = 18, survH0 = 0.30,
  accrualTime = seq(0, 8),
  accrualIntensity = 26/9*seq(1, 9),
  piecewiseSurvivalTime = c(0, 6),
  stratumFraction = c(0.2, 0.8),
  lambda = c(0.0533, 0.0309, 1.5*0.0533, 1.5*0.0309),
  gamma = -log(1-0.05)/12, accrualDuration = 22,
  followupTime = 18, fixedFollowup = FALSE)
```

```
# Example 3: Obtains accrual duration given power, accrual intensity, and
# follow-up duration for fixed follow-up
```

```
kmsamplesize1s(beta = 0.2, kMax = 2,
  informationRates = c(0.8, 1),
```

```

alpha = 0.025, typeAlphaSpending = "sfOF",
milestone = 18, survH0 = 0.30,
accrualTime = seq(0, 8),
accrualIntensity = 26/9*seq(1, 9),
piecewiseSurvivalTime = c(0, 6),
stratumFraction = c(0.2, 0.8),
lambda = c(0.0533, 0.0309, 1.5*0.0533, 1.5*0.0309),
gamma = -log(1-0.05)/12, accrualDuration = NA,
followupTime = 18, fixedFollowup = TRUE)

```

kmsamplesizeequiv	<i>Sample Size for Equivalence in Milestone Survival Probability Difference</i>
-------------------	---

Description

Obtains the sample size for equivalence in milestone survival probability difference.

Usage

```

kmsamplesizeequiv(
  beta = 0.2,
  kMax = 1L,
  informationRates = NA_real_,
  criticalValues = NULL,
  alpha = 0.05,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  milestone = NA_real_,
  survDiffLower = NA_real_,
  survDiffUpper = NA_real_,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = 0L,
  spendingTime = NA_real_,
  rounding = 1L
)

```

Arguments

beta	The type II error.
kMax	The maximum number of stages.
informationRates	The information rates. Defaults to $(1:kMax) / kMax$ if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level for each of the two one-sided tests. Defaults to 0.05.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsiatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
milestone	The milestone time at which to calculate the survival probability.
survDiffLower	The lower equivalence limit of milestone survival probability difference.
survDiffUpper	The upper equivalence limit of milestone survival probability difference.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
lambda1	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.
lambda2	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.

<code>gamma2</code>	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
<code>accrualDuration</code>	Duration of the enrollment period.
<code>followupTime</code>	Follow-up time for the last enrolled subject.
<code>fixedFollowup</code>	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
<code>spendingTime</code>	A vector of length <code>kMax</code> for the error spending time at each analysis. Defaults to missing, in which case, it is the same as <code>informationRates</code> .
<code>rounding</code>	Whether to round up sample size. Defaults to 1 for sample size rounding.

Value

An S3 class `kmpowerequiv` object

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

See Also

[kmpowerequiv](#)

Examples

```
kmsamplesizeequiv(beta = 0.1, kMax = 2, informationRates = c(0.5, 1),
  alpha = 0.05, typeAlphaSpending = "sfOF",
  milestone = 18,
  survDiffLower = -0.13, survDiffUpper = 0.13,
  allocationRatioPlanned = 1, accrualTime = seq(0, 8),
  accrualIntensity = 26/9*seq(1, 9),
  piecewiseSurvivalTime = c(0, 6),
  stratumFraction = c(0.2, 0.8),
  lambda1 = c(0.0533, 0.0533, 1.5*0.0533, 1.5*0.0533),
  lambda2 = c(0.0533, 0.0533, 1.5*0.0533, 1.5*0.0533),
  gamma1 = -log(1-0.05)/12,
  gamma2 = -log(1-0.05)/12, accrualDuration = NA,
  followupTime = 18, fixedFollowup = FALSE)
```

kmstat

*Stratified Difference in Milestone Survival Probabilities***Description**

Obtains the stratified milestone survival probabilities and difference in milestone survival probabilities at given calendar times.

Usage

```
kmstat(
  time = NA_real_,
  milestone = NA_real_,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = FALSE
)
```

Arguments

time	A vector of calendar times for data cut.
milestone	The milestone time at which to calculate the survival probability.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., <code>c(0, 3)</code> breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., <code>c(0, 6)</code> breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.

lambda1	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.
lambda2	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piece-wise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piece-wise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.

Value

A data frame containing the following variables:

- time: The calendar time since trial start.
- subjects: The number of enrolled subjects.
- nevents: The total number of events.
- nevents1: The number of events in the active treatment group.
- nevents2: The number of events in the control group.
- ndropouts: The total number of dropouts.
- ndropouts1: The number of dropouts in the active treatment group.
- ndropouts2: The number of dropouts in the control group.
- milestone: The milestone time relative to randomization.
- nmilestone: The total number of subjects reaching milestone.
- nmilestone1: The number of subjects reaching milestone in the active treatment group.
- nmilestone2: The number of subjects reaching milestone in the control group.
- surv1: The milestone survival probability for the treatment group.
- surv2: The milestone survival probability for the control group.
- survDiff: The difference in milestone survival probabilities, i.e., $\text{surv1} - \text{surv2}$.
- vsurv1: The variance for surv1.
- vsurv2: The variance for surv2.
- vsurvDiff: The variance for survDiff.
- information: The information for survDiff, equal to $1/\text{vsurvDiff}$.
- survDiffZ: The Z-statistic value, i.e., $\text{survDiff}/\sqrt{\text{vsurvDiff}}$.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Piecewise accrual, piecewise exponential survivals, and 5% dropout by
# the end of 1 year.
```

```
kmstat(time = c(22, 40),
        milestone = 18,
        allocationRatioPlanned = 1,
        accrualTime = seq(0, 8),
        accrualIntensity = 26/9*seq(1, 9),
        piecewiseSurvivalTime = c(0, 6),
        stratumFraction = c(0.2, 0.8),
        lambda1 = c(0.0533, 0.0309, 1.5*0.0533, 1.5*0.0309),
        lambda2 = c(0.0533, 0.0533, 1.5*0.0533, 1.5*0.0533),
        gamma1 = -log(1-0.05)/12,
        gamma2 = -log(1-0.05)/12,
        accrualDuration = 22,
        followupTime = 18, fixedFollowup = FALSE)
```

liferegr

Parametric Regression Models for Failure Time Data

Description

Obtains the parameter estimates from parametric regression models with uncensored, right censored, left censored, or interval censored data.

Usage

```
liferegr(
  data,
  stratum = "",
  time = "time",
  time2 = "",
  event = "event",
  covariates = "",
  weight = "",
  offset = "",
  id = "",
  dist = "weibull",
  init = NA_real_,
  robust = FALSE,
  plci = FALSE,
  alpha = 0.05,
```

```

    maxiter = 50,
    eps = 1e-09
)

```

Arguments

<code>data</code>	The input data frame that contains the following variables: • <code>stratum</code>: The stratum. • <code>time</code>: The follow-up time for right censored data, or the left end of each interval for interval censored data. • <code>time2</code>: The right end of each interval for interval censored data. • <code>event</code>: The event indicator, 1=event, 0=no event. • <code>covariates</code>: The values of baseline covariates. • <code>weight</code>: The weight for each observation. • <code>offset</code>: The offset for each observation. • <code>id</code>: The optional subject ID to group the score residuals in computing the robust sandwich variance.
<code>stratum</code>	The name(s) of the stratum variable(s) in the input data.
<code>time</code>	The name of the time variable or the left end of each interval for interval censored data in the input data.
<code>time2</code>	The name of the right end of each interval for interval censored data in the input data.
<code>event</code>	The name of the event variable in the input data for right censored data.
<code>covariates</code>	The vector of names of baseline covariates in the input data.
<code>weight</code>	The name of the weight variable in the input data.
<code>offset</code>	The name of the offset variable in the input data.
<code>id</code>	The name of the id variable in the input data.
<code>dist</code>	The assumed distribution for time to event. Options include "exponential", "weibull", "lognormal", and "loglogistic" to be modeled on the log-scale, and "normal" and "logistic" to be modeled on the original scale.
<code>init</code>	A vector of initial values for the model parameters, including regression coefficients and the log scale parameter. By default, initial values are derived from an intercept-only model. If this approach fails, ordinary least squares (OLS) estimates, ignoring censoring, are used instead.
<code>robust</code>	Whether a robust sandwich variance estimate should be computed. In the presence of the <code>id</code> variable, the score residuals will be aggregated for each <code>id</code> when computing the robust sandwich variance estimate.
<code>plci</code>	Whether to obtain profile likelihood confidence interval.
<code>alpha</code>	The two-sided significance level.
<code>maxiter</code>	The maximum number of iterations.
<code>eps</code>	The tolerance to declare convergence.

Details

There are two ways to specify the model, one for right censored data through the time and event variables, and the other for interval censored data through the time (lower) and time2 (upper) variables. For the second form, we follow the convention used in SAS PROC LIFEREG:

- If lower is not missing, upper is not missing, and lower is equal to upper, then there is no censoring and the event occurred at time lower.
- If lower is not missing, upper is not missing, and lower < upper, then the event time is censored within the interval (lower, upper).
- If lower is missing, but upper is not missing, then upper will be used as the left censoring value.
- If lower is not missing, but upper is missing, then lower will be used as the right censoring value.
- If lower is not missing, upper is not missing, but lower > upper, or if both lower and upper are missing, then the observation will not be used.

Value

A list with the following components:

- **sumstat**: The data frame of summary statistics of model fit with the following variables:
 - **n**: The number of observations.
 - **nevents**: The number of events.
 - **loglik0**: The log-likelihood under null.
 - **loglik1**: The maximum log-likelihood.
 - **niter**: The number of Newton-Raphson iterations.
 - **dist**: The assumed distribution.
 - **p**: The number of parameters, including the intercept, regression coefficients associated with the covariates, and the log scale parameters for the strata.
 - **nvar**: The number of regression coefficients associated with the covariates (excluding the intercept).
 - **robust**: Whether the robust sandwich variance estimate is requested.
 - **fail**: Whether the model fails to converge.
- **parest**: The data frame of parameter estimates with the following variables:
 - **param**: The name of the covariate for the parameter estimate.
 - **beta**: The parameter estimate.
 - **sebeta**: The standard error of parameter estimate.
 - **z**: The Wald test statistic for the parameter.
 - **expbeta**: The exponentiated parameter estimate.
 - **lower**: The lower limit of confidence interval.
 - **upper**: The upper limit of confidence interval.
 - **p**: The p-value from the chi-square test.
 - **method**: The method to compute the confidence interval and p-value.

- `sebeta_naive`: The naive standard error of parameter estimate if robust variance is requested.
- `linear_predictors`: The vector of linear predictors.
- `p`: The number of parameters.
- `nvar`: The number of columns of the design matrix excluding the intercept.
- `param`: The parameter names.
- `beta`: The parameter estimate.
- `vbeta`: The covariance matrix for parameter estimates.
- `vbeta_naive`: The naive covariance matrix for parameter estimates.
- `terms`: The terms object.
- `xlevels`: A record of the levels of the factors used in fitting.
- `settings`: A list containing the input parameter values.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

References

John D. Kalbfleisch and Ross L. Prentice. The Statistical Analysis of Failure Time Data. Wiley: New York, 1980.

Examples

```
library(dplyr)

# right censored data
(fit1 <- liferegr(
  data = rawdata %>% filter(iterationNumber == 1) %>%
    mutate(treat = (treatmentGroup == 1)),
  stratum = "stratum",
  time = "timeUnderObservation", event = "event",
  covariates = "treat", dist = "weibull"))

# tobit regression for left censored data
(fit2 <- liferegr(
  data = tobin %>% mutate(time = ifelse(durable>0, durable, NA)),
  time = "time", time2 = "durable",
  covariates = c("age", "quant"), dist = "normal"))
```

liver

*The liver data used in SAS PROC PHREG documentation examples.***Description**

This data set contains information on 418 patients with primary biliary cirrhosis.

Usage

```
liver
```

Format

An object of class `tbl_df` (inherits from `tbl`, `data.frame`) with 418 rows and 7 columns.

Details

Time The follow-up time in years from the time of registration. The event could be liver transplantation, death, or the end of the study, whichever came first

Status A censoring indicator, where a value of 1 indicating a death event, and 0 indicating a censored observation (the patient survived past the observation time)

Age The patient's age in years

Albumin Serum albumin level in g/dl

Bilirubin Serum bilirubin level in mg/dl

Edema Edema status, where a value of 0 indicates no edema, 0.5 indicates edema successfully treated with diuretics, and 1 indicates edema despite diuretic therapy

Protime Prothrombin time in seconds

logisregr

*Logistic Regression Models for Binary Data***Description**

Obtains the parameter estimates from logistic regression models with binary data.

Usage

```
logisregr(
  data,
  event = "event",
  covariates = "",
  freq = "",
  weight = "",
  offset = "",
```

```

id = "",
link = "logit",
init = NA_real_,
robust = FALSE,
firth = FALSE,
flic = FALSE,
plci = FALSE,
alpha = 0.05,
maxiter = 50,
eps = 1e-09
)

```

Arguments

data	The input data frame that contains the following variables: • event: The event indicator, 1=event, 0=no event. • covariates: The values of baseline covariates. • freq: The frequency for each observation. • weight: The weight for each observation. • offset: The offset for each observation. • id: The optional subject ID to group the score residuals in computing the robust sandwich variance.
event	The name of the event variable in the input data.
covariates	The vector of names of baseline covariates in the input data.
freq	The name of the frequency variable in the input data. The frequencies must be the same for all observations within each cluster as indicated by the id. Thus freq is the cluster frequency.
weight	The name of the weight variable in the input data.
offset	The name of the offset variable in the input data.
id	The name of the id variable in the input data.
link	The link function linking the response probabilities to the linear predictors. Options include "logit" (default), "probit", and "cloglog" (complementary log-log).
init	A vector of initial values for the model parameters. By default, initial values are derived from an intercept-only model.
robust	Whether a robust sandwich variance estimate should be computed. In the presence of the id variable, the score residuals will be aggregated for each id when computing the robust sandwich variance estimate.
firth	Whether the firth's bias reducing penalized likelihood should be used. The default is FALSE.
flic	Whether to apply intercept correction to obtain more accurate predicted probabilities. The default is FALSE.
plci	Whether to obtain profile likelihood confidence interval.
alpha	The two-sided significance level.
maxiter	The maximum number of iterations.
eps	The tolerance to declare convergence.

Details

Fitting a logistic regression model using Firth's bias reduction method is equivalent to penalization of the log-likelihood by the Jeffreys prior. Firth's penalized log-likelihood is given by

$$l(\beta) + \frac{1}{2} \log(\det(I(\beta)))$$

and the components of the gradient $g(\beta)$ are computed as

$$g(\beta_j) + \frac{1}{2} \text{trace} \left(I(\beta)^{-1} \frac{\partial I(\beta)}{\partial \beta_j} \right)$$

The Hessian matrix is not modified by this penalty.

Firth's method reduces bias in maximum likelihood estimates of coefficients, but it introduces a bias toward one-half in the predicted probabilities.

A straightforward modification to Firth's logistic regression to achieve unbiased average predicted probabilities involves a post hoc adjustment of the intercept. This approach, known as Firth's logistic regression with intercept correction (FLIC), preserves the bias-corrected effect estimates. By excluding the intercept from penalization, it ensures that we don't sacrifice the accuracy of effect estimates to improve the predictions.

Value

A list with the following components:

- **sumstat**: The data frame of summary statistics of model fit with the following variables:
 - **n**: The number of subjects.
 - **nevents**: The number of events.
 - **loglik0**: The (penalized) log-likelihood under null.
 - **loglik1**: The maximum (penalized) log-likelihood.
 - **niter**: The number of Newton-Raphson iterations.
 - **p**: The number of parameters, including the intercept, and regression coefficients associated with the covariates.
 - **link**: The link function.
 - **robust**: Whether a robust sandwich variance estimate should be computed.
 - **firth**: Whether the firth's penalized likelihood is used.
 - **flic**: Whether to apply intercept correction.
 - **fail**: Whether the model fails to converge.
 - **loglik0_unpenalized**: The unpenalized log-likelihood under null.
 - **loglik1_unpenalized**: The maximum unpenalized log-likelihood.
- **parest**: The data frame of parameter estimates with the following variables:
 - **param**: The name of the covariate for the parameter estimate.
 - **beta**: The parameter estimate.
 - **sebeta**: The standard error of parameter estimate.
 - **z**: The Wald test statistic for the parameter.
 - **expbeta**: The exponentiated parameter estimate.

- lower: The lower limit of confidence interval.
- upper: The upper limit of confidence interval.
- p: The p-value from the chi-square test.
- method: The method to compute the confidence interval and p-value.
- sebeta_naive: The naive standard error of parameter estimate.
- fitted: The data frame with the following variables:
 - linear_predictors: The linear fit on the link function scale.
 - fitted_values: The fitted probabilities of having an event, obtained by transforming the linear predictors by the inverse of the link function.
- p: The number of parameters.
- link: The link function.
- param: The parameter names.
- beta: The parameter estimate.
- vbeta: The covariance matrix for parameter estimates.
- vbeta_naive: The naive covariance matrix for parameter estimates.
- linear_predictors: The linear fit on the link function scale.
- fitted_values: The fitted probabilities of having an event.
- terms: The terms object.
- xlevels: A record of the levels of the factors used in fitting.
- settings: A list containing the input parameter values.

Author(s)

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References

David Firth. Bias Reduction of Maximum Likelihood Estimates. *Biometrika* 1993; 80:27–38.

Georg Heinze and Michael Schemper. A solution to the problem of separation in logistic regression. *Statistics in Medicine* 2002;21:2409–2419.

Rainer Puh, Georg Heinze, Mariana Nold, Lara Lusa, and Angelika Geroldinger. Firth's logistic regression with rare events: accurate effect estimates and predictions? *Statistics in Medicine* 2017; 36:2302-2317.

Examples

```
(fit1 <- logisregr(
  ingots, event = "NotReady", covariates = "Heat*Soak", freq = "Freq"))
```

lpower	<i>Log-Rank Test Power</i>
--------	----------------------------

Description

Estimates the power, stopping probabilities, and expected sample size in a two-group survival design.

Usage

```
lpower(
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,
  futilityCP = NULL,
  futilityHR = NULL,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  hazardRatioH0 = 1,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = FALSE,
  rho1 = 0,
  rho2 = 0,
  typeOfComputation = "",
  spendingTime = NA_real_,
  studyDuration = NA_real_
)
```

Arguments

kMax	The maximum number of stages.
------	-------------------------------

informationRates	The information rates in terms of number of events for the conventional log-rank test and in terms of the actual information for weighted log-rank tests. Defaults to $(1:kMax) / kMax$ if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., $kMax-1$. Defaults to $\text{rep}(-8, kMax-1)$ if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	A vector of length $kMax - 1$ for the futility bounds on the conditional power scale.
futilityHR	A vector of length $kMax - 1$ for the futility bounds on the hazard ratio scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
hazardRatioH0	Hazard ratio under the null hypothesis for the active treatment versus control. Defaults to 1 for superiority test.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.

accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
lambda1	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.
lambda2	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
rho1	The first parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
rho2	The second parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
typeOfComputation	The type of computation, either "direct" for the direct approximation method, or "schoenfeld" for the Schoenfeld method. Defaults to empty, which selects the Schoenfeld method under proportional hazards and ordinary log-rank test and the direct method otherwise.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.
studyDuration	Study duration for fixed follow-up design. Defaults to missing, which is to be replaced with the sum of accrualDuration and followupTime. If provided, the value is allowed to be less than the sum of accrualDuration and followupTime.

Value

An S3 class lpower object with 4 components:

- overallResults: A data frame containing the following variables:

- overallReject: The overall rejection probability.
- alpha: The overall significance level.
- numberOfEvents: The total number of events.
- numberOfDropouts: The total number of dropouts.
- numbeOfSubjects: The total number of subjects.
- studyDuration: The total study duration.
- information: The maximum information.
- expectedNumberOfEvents: The expected number of events.
- expectedNumberOfDropouts: The expected number of dropouts.
- expectedNumberOfSubjects: The expected number of subjects.
- expectedStudyDuration: The expected study duration.
- expectedInformation: The expected information.
- accrualDuration: The accrual duration.
- followupTime: The follow-up time.
- fixedFollowup: Whether a fixed follow-up design is used.
- rho1: The first parameter of the Fleming-Harrington family of weighted log-rank test.
- rho2: The second parameter of the Fleming-Harrington family of weighted log-rank test.
- kMax: The number of stages.
- hazardRatioH0: The hazard ratio under the null hypothesis.
- typeOfComputation: The type of computation, either "direct" for the direct approximation method, or "schoenfeld" for the Schoenfeld method.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale.
 - futilityBounds: The futility boundaries on the Z-scale.
 - rejectPerStage: The probability for efficacy stopping.
 - futilityPerStage: The probability for futility stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeFutility: The cumulative probability for futility stopping.
 - cumulativeAlphaSpent: The cumulative alpha spent.
 - numberOfEvents: The number of events.
 - numberOfDropouts: The number of dropouts.
 - numberOfSubjects: The number of subjects.
 - analysisTime: The average time since trial start.
 - efficacyHR: The efficacy boundaries on the hazard ratio scale.
 - futilityHR: The futility boundaries on the hazard ratio scale.
 - efficacyP: The efficacy boundaries on the p-value scale.
 - futilityP: The futility boundaries on the p-value scale.
 - information: The cumulative information.
 - HR: The average hazard ratio.
 - efficacyStopping: Whether to allow efficacy stopping.
 - futilityStopping: Whether to allow futility stopping.

- settings: A list containing the following input parameters: typeAlphaSpending, parameterAlphaSpending, userAlphaSpending, typeBetaSpending, parameterBetaSpending, allocationRatioPlanned, accrualTime, accrualIntensity, piecewiseSurvivalTime, stratumFraction, lambda1, lambda2, gamma1, gamma2, and spendingTime.
- byTreatmentCounts: A list containing the following counts by treatment group:
 - numberOfEvents1: The number of events by stage for the treatment group.
 - numberOfDropouts1: The number of dropouts by stage for the treatment group.
 - numberOfSubjects1: The number of subjects by stage for the treatment group.
 - numberOfEvents2: The number of events by stage for the control group.
 - numberOfDropouts2: The number of dropouts by stage for the control group.
 - numberOfSubjects2: The number of subjects by stage for the control group.
 - expectedNumberOfEvents1: The expected number of events for the treatment group.
 - expectedNumberOfDropouts1: The expected number of dropouts for the treatment group.
 - expectedNumberOfSubjects1: The expected number of subjects for the treatment group.
 - expectedNumberOfEvents2: The expected number of events for control group.
 - expectedNumberOfDropouts2: The expected number of dropouts for the control group.
 - expectedNumberOfSubjects2: The expected number of subjects for the control group.

Author(s)

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Examples

```
# Piecewise accrual, piecewise exponential survival, and 5% dropout by
# the end of 1 year.
```

```
lpower(kMax = 2, informationRates = c(0.8, 1),
      alpha = 0.025, typeAlphaSpending = "sfOF",
      allocationRatioPlanned = 1, accrualTime = seq(0, 8),
      accrualIntensity = 26/9*seq(1, 9),
      piecewiseSurvivalTime = c(0, 6),
      lambda1 = c(0.0533, 0.0309),
      lambda2 = c(0.0533, 0.0533),
      gamma1 = -log(1-0.05)/12,
      gamma2 = -log(1-0.05)/12, accrualDuration = 22,
      followupTime = 18, fixedFollowup = FALSE)
```

Description

Obtains the power for equivalence in hazard ratio.

Usage

```
lrpowerequiv(
  kMax = 1L,
  informationRates = NA_real_,
  criticalValues = NULL,
  alpha = 0.05,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  hazardRatioLower = NA_real_,
  hazardRatioUpper = NA_real_,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = FALSE,
  typeOfComputation = "direct",
  spendingTime = NA_real_,
  studyDuration = NA_real_
)
```

Arguments

kMax The maximum number of stages.

informationRates The information rates. Defaults to $(1:kMax) / kMax$ if left unspecified.

criticalValues Upper boundaries on the z-test statistic scale for stopping for efficacy.

alpha The significance level for each of the two one-sided tests. Defaults to 0.05.

typeAlphaSpending The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".

parameterAlphaSpending The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".

userAlphaSpending The user defined alpha spending. Cumulative alpha spent up to each stage.

hazardRatioLower	The lower equivalence limit of hazard ratio.
hazardRatioUpper	The upper equivalence limit of hazard ratio.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
lambda1	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.
lambda2	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
typeOfComputation	The type of computation, either "direct" for the direct approximation method, or "schoenfeld" for the Schoenfeld method. Defaults to empty, which selects the Schoenfeld method under proportional hazards and ordinary log-rank test and the direct method otherwise.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.
studyDuration	Study duration for fixed follow-up design. Defaults to missing, which is to be replaced with the sum of accrualDuration and followupTime. If provided, the value is allowed to be less than the sum of accrualDuration and followupTime.

Value

An S3 class *lrpowerequiv* object with 4 components:

- **overallResults**: A data frame containing the following variables:
 - **overallReject**: The overall rejection probability.
 - **alpha**: The overall significance level.
 - **numberOfEvents**: The total number of events.
 - **numberOfDropouts**: The total number of dropouts.
 - **numbeOfSubjects**: The total number of subjects.
 - **studyDuration**: The total study duration.
 - **information**: The maximum information.
 - **expectedNumberOfEvents**: The expected number of events.
 - **expectedNumberOfDropouts**: The expected number of dropouts.
 - **expectedNumberOfSubjects**: The expected number of subjects.
 - **expectedStudyDuration**: The expected study duration.
 - **expectedInformation**: The expected information.
 - **kMax**: The number of stages.
 - **hazardRatioLower**: The lower equivalence limit of hazard ratio.
 - **hazardRatioUpper**: The upper equivalence limit of hazard ratio.
 - **accrualDuration**: The accrual duration.
 - **followupTime**: The follow-up time.
 - **fixedFollowup**: Whether a fixed follow-up design is used.
- **byStageResults**: A data frame containing the following variables:
 - **informationRates**: The information rates.
 - **efficacyBounds**: The efficacy boundaries on the Z-scale for each of the two one-sided tests.
 - **rejectPerStage**: The probability for efficacy stopping.
 - **cumulativeRejection**: The cumulative probability for efficacy stopping.
 - **cumulativeAlphaSpent**: The cumulative alpha for each of the two one-sided tests.
 - **cumulativeAttainedAlphaH10**: The cumulative alpha attained under H10.
 - **cumulativeAttainedAlphaH20**: The cumulative alpha attained under H20.
 - **numberOfEvents**: The number of events.
 - **numberOfDropouts**: The number of dropouts.
 - **numberOfSubjects**: The number of subjects.
 - **analysisTime**: The average time since trial start.
 - **efficacyHRLower**: The efficacy boundaries on the hazard ratio scale for the one-sided null hypothesis at the lower equivalence limit.
 - **efficacyHRUpper**: The efficacy boundaries on the hazard ratio scale for the one-sided null hypothesis at the upper equivalence limit.
 - **efficacyP**: The efficacy bounds on the p-value scale for each of the two one-sided tests.
 - **information**: The cumulative information.
 - **HR**: The average hazard ratio.

- **settings:** A list containing the following input parameters: `typeAlphaSpending`, `parameterAlphaSpending`, `userAlphaSpending`, `allocationRatioPlanned`, `accrualTime`, `accrualIntensity`, `piecewiseSurvivalTime`, `stratumFraction`, `lambda1`, `lambda2`, `gamma1`, `gamma2`, `typeOfComputation`, and `spendingTime`.
- **byTreatmentCounts:** A list containing the following counts by treatment group:
 - `numberOfEvents1`: The number of events by stage for the treatment group.
 - `numberOfDropouts1`: The number of dropouts by stage for the treatment group.
 - `numberOfSubjects1`: The number of subjects by stage for the treatment group.
 - `numberOfEvents2`: The number of events by stage for the control group.
 - `numberOfDropouts2`: The number of dropouts by stage for the control group.
 - `numberOfSubjects2`: The number of subjects by stage for the control group.
 - `expectedNumberOfEvents1`: The expected number of events for the treatment group.
 - `expectedNumberOfDropouts1`: The expected number of dropouts for the treatment group.
 - `expectedNumberOfSubjects1`: The expected number of subjects for the treatment group.
 - `expectedNumberOfEvents2`: The expected number of events for control group.
 - `expectedNumberOfDropouts2`: The expected number of dropouts for the control group.
 - `expectedNumberOfSubjects2`: The expected number of subjects for the control group.

Author(s)

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See Also

[rmstat](#)

Examples

```
lrsamplesize(kMax = 2, informationRates = c(0.5, 1),
  alpha = 0.05, typeAlphaSpending = "sfOF",
  hazardRatioLower = 0.71, hazardRatioUpper = 1.4,
  allocationRatioPlanned = 1, accrualTime = seq(0, 8),
  accrualIntensity = 100/9*seq(1, 9),
  lambda1 = 0.0533,
  lambda2 = 0.0533,
  gamma1 = -log(1-0.05)/12,
  gamma2 = -log(1-0.05)/12, accrualDuration = 22,
  followupTime = 18, fixedFollowup = FALSE)
```

lrsamplesize

Log-Rank Test Sample Size

Description

Obtains the needed accrual duration given power and follow-up time, the needed follow-up time given power and accrual duration, or the needed absolute accrual rates given power, accrual duration, follow-up time, and relative accrual rates in a two-group survival design.

Usage

```

lrsamplesize(
  beta = 0.2,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,
  futilityCP = NULL,
  futilityHR = NULL,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
  hazardRatioH0 = 1,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = FALSE,
  rho1 = 0,
  rho2 = 0,
  typeOfComputation = "",
  spendingTime = NA_real_,
  rounding = TRUE
)

```

Arguments

<code>beta</code>	Type II error. Defaults to 0.2.
<code>kMax</code>	The maximum number of stages.
<code>informationRates</code>	The information rates in terms of number of events for the conventional log-rank test and in terms of the actual information for weighted log-rank tests. Defaults to $(1:kMax) / kMax$ if left unspecified.
<code>efficacyStopping</code>	Indicators of whether efficacy stopping is allowed at each stage. Defaults to

	TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	A vector of length kMax - 1 for the futility bounds on the conditional power scale.
futilityHR	A vector of length kMax - 1 for the futility bounds on the hazard ratio scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
hazardRatioH0	Hazard ratio under the null hypothesis for the active treatment versus control. Defaults to 1 for superiority test.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., c(0, 3) breaks the time axis into 2 accrual intervals: [0, 3) and [3, ∞).
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.

<code>piecewiseSurvivalTime</code>	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., <code>c(0, 6)</code> breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
<code>stratumFraction</code>	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
<code>lambda1</code>	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.
<code>lambda2</code>	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
<code>gamma1</code>	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
<code>gamma2</code>	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
<code>accrualDuration</code>	Duration of the enrollment period.
<code>followupTime</code>	Follow-up time for the last enrolled subject.
<code>fixedFollowup</code>	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
<code>rho1</code>	The first parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
<code>rho2</code>	The second parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
<code>typeOfComputation</code>	The type of computation, either "direct" for the direct approximation method, or "schoenfeld" for the Schoenfeld method. Defaults to empty, which selects the Schoenfeld method under proportional hazards and ordinary log-rank test and the direct method otherwise.
<code>spendingTime</code>	A vector of length <code>kMax</code> for the error spending time at each analysis. Defaults to missing, in which case, it is the same as <code>informationRates</code> .
<code>rounding</code>	Whether to round up sample size and events. Defaults to 1 for sample size rounding.

Value

A list of two components:

- `resultsUnderH1`: An S3 class `lrpower` object under the alternative hypothesis.
- `resultsUnderH0`: An S3 class `lrpower` object under the null hypothesis.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

See Also[lpower](#)**Examples**

```
# Piecewise accrual, piecewise exponential survival, and 5% dropout by
# the end of 1 year.
```

```
# Example 1: Obtains accrual duration given power and follow-up time
```

```
lrsamplesize(beta = 0.2, kMax = 2,
             informationRates = c(0.8, 1),
             alpha = 0.025, typeAlphaSpending = "sf0F",
             accrualTime = seq(0, 8),
             accrualIntensity = 26/9*seq(1, 9),
             piecewiseSurvivalTime = c(0, 6),
             lambda1 = c(0.0533, 0.0309),
             lambda2 = c(0.0533, 0.0533),
             gamma1 = -log(1-0.05)/12,
             gamma2 = -log(1-0.05)/12,
             accrualDuration = NA,
             followupTime = 18, fixedFollowup = FALSE)
```

```
# Example 2: Obtains follow-up time given power and accrual duration
```

```
lrsamplesize(beta = 0.2, kMax = 2,
             informationRates = c(0.8, 1),
             alpha = 0.025, typeAlphaSpending = "sf0F",
             accrualTime = seq(0, 8),
             accrualIntensity = 26/9*seq(1, 9),
             piecewiseSurvivalTime = c(0, 6),
             lambda1 = c(0.0533, 0.0309),
             lambda2 = c(0.0533, 0.0533),
             gamma1 = -log(1-0.05)/12,
             gamma2 = -log(1-0.05)/12,
             accrualDuration = 22,
             followupTime = NA, fixedFollowup = FALSE)
```

```
# Example 3: Obtains absolute accrual intensity given power,
# accrual duration, follow-up time, and relative accrual intensity
```

```
lrsamplesize(beta = 0.2, kMax = 2,
             informationRates = c(0.8, 1),
             alpha = 0.025, typeAlphaSpending = "sf0F",
             accrualTime = seq(0, 8),
             accrualIntensity = 26/9*seq(1, 9),
             piecewiseSurvivalTime = c(0, 6),
             lambda1 = c(0.0533, 0.0309),
             lambda2 = c(0.0533, 0.0533),
             gamma1 = -log(1-0.05)/12,
```

```
gamma2 = -log(1-0.05)/12,
accrualDuration = 22,
followupTime = 18, fixedFollowup = FALSE)
```

lrsamplesizeequiv	<i>Sample Size for Equivalence in Hazard Ratio</i>
-------------------	--

Description

Obtains the sample size for equivalence in hazard ratio.

Usage

```
lrsamplesizeequiv(
  beta = 0.2,
  kMax = 1L,
  informationRates = NA_real_,
  criticalValues = NULL,
  alpha = 0.05,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  hazardRatioLower = NA_real_,
  hazardRatioUpper = NA_real_,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = FALSE,
  typeOfComputation = "direct",
  spendingTime = NA_real_,
  rounding = TRUE
)
```

Arguments

- beta The type II error.
- kMax The maximum number of stages.

informationRates	The information rates. Defaults to $(1:kMax) / kMax$ if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level for each of the two one-sided tests. Defaults to 0.05.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
hazardRatioLower	The lower equivalence limit of hazard ratio.
hazardRatioUpper	The upper equivalence limit of hazard ratio.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
lambda1	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.
lambda2	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.

<code>accrualDuration</code>	Duration of the enrollment period.
<code>followupTime</code>	Follow-up time for the last enrolled subject.
<code>fixedFollowup</code>	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
<code>typeOfComputation</code>	The type of computation, either "direct" for the direct approximation method, or "schoenfeld" for the Schoenfeld method. Defaults to empty, which selects the Schoenfeld method under proportional hazards and ordinary log-rank test and the direct method otherwise.
<code>spendingTime</code>	A vector of length <code>kMax</code> for the error spending time at each analysis. Defaults to missing, in which case, it is the same as <code>informationRates</code> .
<code>rounding</code>	Whether to round up sample size. Defaults to 1 for sample size rounding.

Value

An S3 class `lrpowerequiv` object

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

See Also

[lrpowerequiv](#)

Examples

```
lrsamplesizeequiv(kMax = 2, informationRates = c(0.5, 1),
  alpha = 0.05, typeAlphaSpending = "sfOF",
  hazardRatioLower = 0.71, hazardRatioUpper = 1.4,
  allocationRatioPlanned = 1, accrualTime = seq(0, 8),
  accrualIntensity = 26/9*seq(1, 9),
  piecewiseSurvivalTime = c(0, 6),
  lambda1 = c(0.0533, 0.0533),
  lambda2 = c(0.0533, 0.0533),
  gamma1 = -log(1-0.05)/12,
  gamma2 = -log(1-0.05)/12, accrualDuration = NA,
  followupTime = 18, fixedFollowup = FALSE)
```

Irschoenfeld

*Schoenfeld Method for Log-Rank Test Sample Size Calculation***Description**

Obtains the sample size and study duration by calibrating the number of events calculated using the Schoenfeld formula under the proportional hazards assumption. The initial event target can be used directly or calibrated with simulation.

Usage

```
lrschoenfeld(
  beta = 0.2,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NA_real_,
  alpha = 0.025,
  typeAlphaSpending = "sf0F",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NA_real_,
  futilityCP = NA_real_,
  futilityHR = NA_real_,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
  hazardRatioH0 = 1,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  hazardRatio = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  followupTime = NA_real_,
  fixedFollowup = FALSE,
  spendingTime = NA_real_,
  rounding = TRUE,
  simulate = FALSE,
  calibrate = FALSE,
  maxNumberOfIterations = 10000L,
  maxNumberOfRawDatasetsPerStage = 0L,
  seed = NA_integer_
```

)

Arguments

beta	Type II error (one minus the target power). Defaults to 0.2.
kMax	The maximum number of stages.
informationRates	The information rates in terms of number of events for the conventional log-rank test and in terms of the actual information for weighted log-rank tests. Defaults to $(1:kMax) / kMax$ if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	A vector of length kMax - 1 for the futility bounds on the conditional power scale.
futilityHR	A vector of length kMax - 1 for the futility bounds on the hazard ratio scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".

userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
hazardRatioH0	Hazard ratio under the null hypothesis for the active treatment versus control. Defaults to 1 for superiority test.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
hazardRatio	Hazard ratio under the alternative hypothesis for the active treatment versus control.
lambda2	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.
rounding	Whether to round up sample size and events. Defaults to 1 for sample size rounding.
simulate	Whether to use simulation to estimate empirical power for the event target.
calibrate	Whether to use simulation to calibrate the number of events calculated using the Schoenfeld formula. If TRUE, simulate is also treated as TRUE.
maxNumberOfIterations	The number of simulation iterations. Defaults to 10000.
maxNumberOfRawDatasetsPerStage	The number of raw datasets per stage to extract.
seed	The seed to reproduce the simulation results.

Details

This function first calculates the required number of events using the group-sequential design and the Schoenfeld approximation. It then obtains the corresponding sample size and study duration from the accrual, event, and dropout assumptions. The optional simulation-based calibration is useful when the hazard ratio is far from one or the allocation is unequal, because the Schoenfeld approximation can then be inaccurate.

For a fixed design, the Schoenfeld formula for the required number of events is

$$D = \frac{(\Phi^{-1}(1 - \alpha) + \Phi^{-1}(1 - \beta))^2}{(\theta - \theta_0)^2 r(1 - r)}$$

where D is the total number of events required, α is the type I error rate, β is the type II error rate, r is the randomization probability for the active treatment group, θ_0 and θ are the log hazard ratios under the null and alternative hypotheses, respectively.

Let $D_{\text{schoenfeld}}$ be the initial number of events calculated by the Schoenfeld formula, and $D_{\text{calibrated}}$ be the calibrated number of events. A simulation estimates the empirical power $p_{\text{schoenfeld}}$ at $D_{\text{schoenfeld}}$. The calibrated number of events is then calculated as

$$D_{\text{calibrated}} = \frac{\{\Phi^{-1}(1 - \alpha) + \Phi^{-1}(1 - \beta)\}^2}{\{\Phi^{-1}(1 - \alpha) + \Phi^{-1}(p_{\text{schoenfeld}})\}^2} D_{\text{schoenfeld}}$$

Here $p_{\text{schoenfeld}}$ is the empirical rejection probability estimated by the first simulation. If `rounding = TRUE`, the calibrated total and cumulative stage event targets are rounded up or to the nearest integer as appropriate. A half-count correction is applied to the simulated rejection count if needed so that empirical power values of exactly 0 or 1 do not produce infinite normal quantiles.

A second simulation is performed to obtain empirical power using the calibrated number of events. With `simulate = FALSE`, only the analytical result is returned. With `simulate = TRUE` and `calibrate = FALSE`, the simulation uses the initial target. With `calibrate = TRUE`, the simulation uses the calibrated target.

When `simulate = FALSE`, the function returns the number of events calculated using the Schoenfeld formula and the corresponding sample size and study duration without simulation results for the empirical power.

When `simulate = TRUE`, the function returns the event target used for the simulation and the corresponding sample size and study duration along with simulation results for empirical power. The target is the Schoenfeld target when `calibrate = FALSE` and the calibrated target when `calibrate = TRUE`.

Value

A list of two components when `simulate = TRUE`:

- `analyticalResults`: An S3 class `lrmpower` object for the asymptotic power.
- `simulationResults`: An S3 class `lrsim` object for the empirical power.

A list of one component for `analyticalResults` when `simulate = FALSE`.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(lr1 <- lrschoenfeld(
  beta = 0.1, kMax = 2, alpha = 0.025,
  hazardRatioH0 = 1, allocationRatioPlanned = 1,
  accrualIntensity = 20, hazardRatio = 0.3,
  lambda2 = 1.9/12,
  gamma1 = -log(1-0.1)/24, gamma2 = -log(1-0.1)/24,
  fixedFollowup = FALSE, rounding = TRUE,
  simulate = TRUE, calibrate = FALSE,
  maxNumberOfIterations = 1000,
  seed = 12345))

(lr2 <- lrschoenfeld(
  beta = 0.1, kMax = 2, alpha = 0.025,
  hazardRatioH0 = 1, allocationRatioPlanned = 1,
  accrualIntensity = 20, hazardRatio = 0.3,
  lambda2 = 1.9/12,
  gamma1 = -log(1-0.1)/24, gamma2 = -log(1-0.1)/24,
  fixedFollowup = FALSE, rounding = TRUE,
  simulate = TRUE, calibrate = TRUE,
  maxNumberOfIterations = 1000,
  seed = 12345))
```

lrsim

*Log-Rank Test Simulation***Description**

Performs simulation for two-arm group sequential trials based on weighted log-rank test.

Usage

```
lrsim(
  kMax = 1,
  informationRates = NA,
  criticalValues = NA,
  futilityBounds = NA,
  hazardRatioH0 = 1,
  allocation1 = 1,
  allocation2 = 1,
  accrualTime = 0,
  accrualIntensity = NA,
  piecewiseSurvivalTime = 0,
  stratumFraction = 1,
  lambda1 = NA,
  lambda2 = NA,
  gamma1 = 0,
```

```

gamma2 = 0,
n = NA,
followupTime = NA,
fixedFollowup = FALSE,
rho1 = 0,
rho2 = 0,
plannedEvents = NA,
plannedTime = NA,
maxNumberOfIterations = 1000,
maxNumberOfRawDatasetsPerStage = 0,
seed = 0,
nthreads = 0
)

```

Arguments

kMax	The maximum number of stages.
informationRates	The information rates in terms of number of events for the conventional log-rank test and in terms of the actual information for weighted log-rank tests. Fixed prior to the trial. If left unspecified, it defaults to <code>plannedEvents / plannedEvents[kMax]</code> when <code>plannedEvents</code> is provided and to <code>plannedTime / plannedTime[kMax]</code> otherwise.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to <code>rep(-8, kMax-1)</code> if left unspecified. The futility bounds are non-binding for the calculation of critical values.
hazardRatioH0	Hazard ratio under the null hypothesis for the active treatment versus control. Defaults to 1 for superiority test.
allocation1	Number of subjects in the active treatment group in a randomization block. Defaults to 1 for equal randomization.
allocation2	Number of subjects in the control group in a randomization block. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., <code>c(0, 3)</code> breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., <code>c(0, 6)</code> breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
lambda1	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.

lambda2	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
n	Sample size.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
rho1	The first parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
rho2	The second parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
plannedEvents	The planned cumulative total number of events at each stage.
plannedTime	The calendar times for the analyses. To use calendar time to plan the analyses, plannedEvents should be missing.
maxNumberOfIterations	The number of simulation iterations. Defaults to 1000.
maxNumberOfRawDatasetsPerStage	The number of raw datasets per stage to extract.
seed	The seed to reproduce the simulation results.
nthreads	The number of threads to use in simulations (0 means the default RcppParallel behavior).

Value

An S3 class `lrsim` object with 3 components:

- overview: A list containing the following information:
 - rejectPerStage: The efficacy stopping probability by stage.
 - futilityPerStage: The futility stopping probability by stage.
 - cumulativeRejection: Cumulative efficacy stopping probability by stage.
 - cumulativeFutility: The cumulative futility stopping probability by stage.
 - numberOfEvents: The average number of events by stage.
 - numberOfDropouts: The average number of dropouts by stage.
 - numberOfSubjects: The average number of subjects by stage.
 - analysisTime: The average analysis time by stage.
 - overallReject: The overall rejection probability.
 - expectedNumberOfEvents: The expected number of events for the overall study.

- expectedNumberOfDropouts: The expected number of dropouts for the overall study.
- expectedNumberOfSubjects: The expected number of subjects for the overall study.
- expectedStudyDuration: The expected study duration.
- hazardRatioH0: Hazard ratio under the null hypothesis for the active treatment versus control.
- useEvents: whether the analyses are planned based on the number of events or calendar time.
- numberOfIterations: The number of simulation iterations.
- n: Sample size.
- fixedFollowup: Whether a fixed follow-up design is used.
- rho1: The first parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
- rho2: The second parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
- kMax: The maximum number of stages.
- sumdata: A data frame of summary data by iteration and stage:
 - iterationNumber: The iteration number.
 - eventsNotAchieved: Whether the final target number of events is not achieved for the iteration.
 - stopStage: The stage at which the trial stops.
 - stageNumber: The stage number, covering all stages even if the trial stops at an interim look.
 - analysisTime: The time for the stage since trial start.
 - accruals1: The number of subjects enrolled at the stage for the treatment group.
 - accruals2: The number of subjects enrolled at the stage for the control group.
 - totalAccruals: The total number of subjects enrolled at the stage.
 - events1: The number of events at the stage for the treatment group.
 - events2: The number of events at the stage for the control group.
 - totalEvents: The total number of events at the stage.
 - dropouts1: The number of dropouts at the stage for the treatment group.
 - dropouts2: The number of dropouts at the stage for the control group.
 - totalDropouts: The total number of dropouts at the stage.
 - uscore: The numerator of the log-rank test statistic.
 - vscore: The variance of the log-rank test statistic.
 - logRankStatistic: The log-rank test Z-statistic.
 - rejectPerStage: Whether to reject the null hypothesis at the stage.
 - futilityPerStage: Whether to stop the trial for futility at the stage.
- rawdata (exists if maxNumberOfRawDatasetsPerStage is a positive integer): A data frame for subject-level data for selected replications, containing the following variables:
 - iterationNumber: The iteration number.
 - stopStage: The stage at which the trial stops.
 - stageNumber: The stage number, covering all stages even if the trial stops at an interim look.

- analysisTime: The time for the stage since trial start.
- subjectId: The subject ID.
- arrivalTime: The enrollment time for the subject.
- stratum: The stratum for the subject.
- treatmentGroup: The treatment group (1 or 2) for the subject.
- survivalTime: The underlying survival time for the subject.
- dropoutTime: The underlying dropout time for the subject.
- timeUnderObservation: The time under observation since randomization.
- event: Whether the subject experienced the event.
- dropoutEvent: Whether the subject dropped out.

Author(s)

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Examples

Example 1: analyses based on number of events

```
sim1 <- lrsim(  
  kMax = 2, informationRates = c(0.5, 1),  
  criticalValues = c(2.797, 1.977),  
  accrualIntensity = 11,  
  lambda1 = 0.018, lambda2 = 0.030,  
  n = 132,  
  plannedEvents = c(60, 120),  
  maxNumberOfIterations = 1000,  
  maxNumberOfRawDatasetsPerStage = 1,  
  seed = 314159,  
  nthreads = 1)
```

```
# summary statistics  
sim1
```

```
# summary for each simulated data set  
head(sim1$sumdata)
```

```
# raw data for selected replication  
head(sim1$rawdata)
```

Example 2: analyses based on calendar time have similar power

```
sim2 <- lrsim(  
  kMax = 2, informationRates = c(0.5, 1),  
  criticalValues = c(2.797, 1.977),  
  accrualIntensity = 11,  
  lambda1 = 0.018, lambda2 = 0.030,  
  n = 132,  
  plannedTime = c(31.9, 113.2),
```

```

maxNumberOfIterations = 1000,
maxNumberOfRawDatasetsPerStage = 1,
seed = 314159,
nthreads = 1)

# summary statistics
sim2

# summary for each simulated data set
head(sim2$sumdata)

```

lrsim2e

Log-Rank Test Simulation for PFS and OS Endpoints

Description

Performs simulation for two-endpoint (PFS and OS) two-arm group sequential trials based on weighted log-rank test. The first kMaxpfs looks are driven by the total number of PFS events in two arms combined, and the subsequent looks are driven by the total number of OS events in two arms combined. Alternatively, the analyses can be planned to occur at specified calendar times.

Usage

```

lrsim2e(
  kMax = 1,
  kMaxpfs = 1,
  hazardRatioH0pfs = 1,
  hazardRatioH0os = 1,
  allocation1 = 1,
  allocation2 = 1,
  accrualTime = 0,
  accrualIntensity = NA,
  piecewiseSurvivalTime = 0,
  stratumFraction = 1,
  rho_pd_os = 0,
  lambda1pfs = NA,
  lambda2pfs = NA,
  lambda1os = NA,
  lambda2os = NA,
  gamma1pfs = 0,
  gamma2pfs = 0,
  gamma1os = 0,
  gamma2os = 0,
  n = NA,
  followupTime = NA,
  fixedFollowup = FALSE,
  rho1 = 0,

```

```

    rho2 = 0,
    plannedEvents = NA,
    plannedTime = NA,
    maxNumberOfIterations = 1000,
    maxNumberOfRawDatasetsPerStage = 0,
    seed = 0,
    nthreads = 0
)

```

Arguments

kMax	The maximum number of stages.
kMaxpfs	Number of stages with timing determined by PFS events. Ranges from 0 (none) to kMax.
hazardRatioH0pfs	Hazard ratio under the null hypothesis for the active treatment vs control for PFS. Defaults to 1 for superiority test.
hazardRatioH0os	Hazard ratio under the null hypothesis for the active treatment vs control for OS. Defaults to 1 for superiority test.
allocation1	Number of subjects in the treatment group in a randomization block. Defaults to 1 for equal randomization.
allocation2	Number of subjects in the control group in a randomization block. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
rho_pd_os	The correlation coefficient for the standard bivariate normal random variables used to generate time to disease progression (PD) and time to death using the inverse CDF method.
lambda1pfs	A vector of hazard rates for the event in each analysis time interval by stratum for the treatment group and PFS.
lambda2pfs	A vector of hazard rates for the event in each analysis time interval by stratum for the control group and PFS.
lambda1os	A vector of hazard rates for the event in each analysis time interval by stratum for the treatment group and OS.
lambda2os	A vector of hazard rates for the event in each analysis time interval by stratum for the control group and OS.

<code>gamma1pfs</code>	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the treatment group and PFS.
<code>gamma2pfs</code>	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group and PFS.
<code>gamma1os</code>	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the treatment group and OS.
<code>gamma2os</code>	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group and OS.
<code>n</code>	Sample size.
<code>followupTime</code>	Follow-up time for the last enrolled subject.
<code>fixedFollowup</code>	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
<code>rho1</code>	The first parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
<code>rho2</code>	The second parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
<code>plannedEvents</code>	The planned cumulative total number of PFS events at Look 1 to Look <code>kMaxpfs</code> and the planned cumulative total number of OS events at Look <code>kMaxpfs+1</code> to Look <code>kMax</code> .
<code>plannedTime</code>	The calendar times for the analyses. To use calendar time to plan the analyses, <code>plannedEvents</code> should be missing.
<code>maxNumberOfIterations</code>	The number of simulation iterations. Defaults to 1000.
<code>maxNumberOfRawDatasetsPerStage</code>	The number of raw datasets per stage to extract.
<code>seed</code>	The seed to reproduce the simulation results.
<code>nthreads</code>	The number of threads to use in simulations (0 means the default <code>RcppParallel</code> behavior).

Value

A list with 2 components:

- `sumdata`: A data frame of summary data by iteration and stage:
 - `iterationNumber`: The iteration number.
 - `eventsNotAchieved`: Whether the target number of events is not achieved for the iteration.

- stageNumber: The stage number, covering all stages even if the trial stops at an interim look.
- analysisTime: The time for the stage since trial start.
- accruals1: The number of subjects enrolled at the stage for the treatment group.
- accruals2: The number of subjects enrolled at the stage for the control group.
- totalAccruals: The total number of subjects enrolled at the stage.
- endpoint: The endpoint (1 for PFS or 2 for OS) under consideration.
- events1: The number of events at the stage for the treatment group.
- events2: The number of events at the stage for the control group.
- totalEvents: The total number of events at the stage.
- dropouts1: The number of dropouts at the stage for the treatment group.
- dropouts2: The number of dropouts at the stage for the control group.
- totalDropouts: The total number of dropouts at the stage.
- uscore: The numerator of the log-rank test statistic for the endpoint.
- vscore: The variance of the log-rank test statistic for the endpoint.
- logRankStatistic: The log-rank test Z-statistic for the endpoint.
- rawdata (exists if maxNumberOfRawDatasetsPerStage is a positive integer): A data frame for subject-level data for selected replications, containing the following variables:
 - iterationNumber: The iteration number.
 - stageNumber: The stage under consideration.
 - analysisTime: The time for the stage since trial start.
 - subjectId: The subject ID.
 - arrivalTime: The enrollment time for the subject.
 - stratum: The stratum for the subject.
 - treatmentGroup: The treatment group (1 or 2) for the subject.
 - endpoint: The endpoint (1 for PFS or 2 for OS) under consideration for the row. Each subject will have two rows, one for each endpoint.
 - survivalTime: The underlying survival time for the event endpoint for the subject.
 - dropoutTime: The underlying dropout time for the event endpoint for the subject.
 - timeUnderObservation: The time under observation since randomization for the event endpoint for the subject.
 - event: Whether the subject experienced the event endpoint.
 - dropoutEvent: Whether the subject dropped out for the endpoint.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
sim1 <- lrsim2e(
  kMax = 3,
  kMaxpfs = 2,
  allocation1 = 2,
  allocation2 = 1,
```

```

    accrualTime = c(0, 8),
    accrualIntensity = c(10, 28),
    piecewiseSurvivalTime = 0,
    rho_pd_os = 0,
    lambda1pfs = log(2)/12*0.60,
    lambda2pfs = log(2)/12,
    lambda1os = log(2)/30*0.65,
    lambda2os = log(2)/30,
    n = 420,
    plannedEvents = c(186, 259, 183),
    maxNumberOfIterations = 1000,
    maxNumberOfRawDatasetsPerStage = 1,
    seed = 314159,
    nthreads = 1)

head(sim1$sumdata)
head(sim1$rawdata)

```

lrsim2e3a

Log-Rank Test Simulation for Two Endpoints (PFS and OS) and Three Arms

Description

Performs simulation for two-endpoint (PFS and OS) three-arm group sequential trials based on weighted log-rank test. The first k_{Maxpfs} looks are driven by the total number of PFS events in Arm A and Arm C combined, and the subsequent looks are driven by the total number of OS events in Arm A and Arm C combined. Alternatively, the analyses can be planned to occur at specified calendar times.

Usage

```

lrsim2e3a(
  kMax = 1,
  kMaxpfs = 1,
  hazardRatioH013pfs = 1,
  hazardRatioH023pfs = 1,
  hazardRatioH012pfs = 1,
  hazardRatioH013os = 1,
  hazardRatioH023os = 1,
  hazardRatioH012os = 1,
  allocation1 = 1,
  allocation2 = 1,
  allocation3 = 1,
  accrualTime = 0,
  accrualIntensity = NA,
  piecewiseSurvivalTime = 0,
  stratumFraction = 1,

```

```

    rho_pd_os = 0,
    lambda1pfs = NA,
    lambda2pfs = NA,
    lambda3pfs = NA,
    lambda1os = NA,
    lambda2os = NA,
    lambda3os = NA,
    gamma1pfs = 0,
    gamma2pfs = 0,
    gamma3pfs = 0,
    gamma1os = 0,
    gamma2os = 0,
    gamma3os = 0,
    n = NA,
    followupTime = NA,
    fixedFollowup = FALSE,
    rho1 = 0,
    rho2 = 0,
    plannedEvents = NA,
    plannedTime = NA,
    maxNumberOfIterations = 1000,
    maxNumberOfRawDatasetsPerStage = 0,
    seed = 0,
    nthreads = 0
)

```

Arguments

kMax	The maximum number of stages.
kMaxpfs	Number of stages with timing determined by PFS events. Ranges from 0 (none) to kMax.
hazardRatioH013pfs	Hazard ratio under the null hypothesis for arm 1 vs arm 3 for PFS. Defaults to 1 for superiority test.
hazardRatioH023pfs	Hazard ratio under the null hypothesis for arm 2 vs arm 3 for PFS. Defaults to 1 for superiority test.
hazardRatioH012pfs	Hazard ratio under the null hypothesis for arm 1 vs arm 2 for PFS. Defaults to 1 for superiority test.
hazardRatioH013os	Hazard ratio under the null hypothesis for arm 1 vs arm 3 for OS. Defaults to 1 for superiority test.
hazardRatioH023os	Hazard ratio under the null hypothesis for arm 2 vs arm 3 for OS. Defaults to 1 for superiority test.

hazardRatioH012os	Hazard ratio under the null hypothesis for arm 1 vs arm 2 for OS. Defaults to 1 for superiority test.
allocation1	Number of subjects in Arm A in a randomization block. Defaults to 1 for equal randomization.
allocation2	Number of subjects in Arm B in a randomization block. Defaults to 1 for equal randomization.
allocation3	Number of subjects in Arm C in a randomization block. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
rho_pd_os	The correlation coefficient for the standard bivariate normal random variables used to generate time to disease progression and time to death using the inverse CDF method.
lambda1pfs	A vector of hazard rates for the event in each analysis time interval by stratum for arm 1 and PFS.
lambda2pfs	A vector of hazard rates for the event in each analysis time interval by stratum for arm 2 and PFS.
lambda3pfs	A vector of hazard rates for the event in each analysis time interval by stratum for arm 3 and PFS.
lambda1os	A vector of hazard rates for the event in each analysis time interval by stratum for arm 1 and OS.
lambda2os	A vector of hazard rates for the event in each analysis time interval by stratum for arm 2 and OS.
lambda3os	A vector of hazard rates for the event in each analysis time interval by stratum for arm 3 and OS.
gamma1pfs	The hazard rate for exponential dropout. A vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for arm 1 and PFS.
gamma2pfs	The hazard rate for exponential dropout. A vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for arm 2 and PFS.
gamma3pfs	The hazard rate for exponential dropout. A vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for arm 3 and PFS.

gamma1os	The hazard rate for exponential dropout. A vector of hazard rates for piece-wise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for arm 1 and OS.
gamma2os	The hazard rate for exponential dropout. A vector of hazard rates for piece-wise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for arm 2 and OS.
gamma3os	The hazard rate for exponential dropout. A vector of hazard rates for piece-wise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for arm 3 and OS.
n	Sample size.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
rho1	The first parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
rho2	The second parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
plannedEvents	The planned cumulative total number of PFS events at Look 1 to Look kMaxpfs for Arms A and C combined and the planned cumulative total number of OS events at Look kMaxpfs+1 to Look kMax for Arms A and C combined.
plannedTime	The calendar times for the analyses. To use calendar time to plan the analyses, plannedEvents should be missing.
maxNumberOfIterations	The number of simulation iterations. Defaults to 1000.
maxNumberOfRawDatasetsPerStage	The number of raw datasets per stage to extract.
seed	The seed to reproduce the simulation results.
nthreads	The number of threads to use in simulations (0 means the default RcppParallel behavior).

Value

A list with 2 components:

- **sumdata**: A data frame of summary data by iteration and stage:
 - **iterationNumber**: The iteration number.
 - **eventsNotAchieved**: Whether the target number of events is not achieved for the iteration.
 - **stageNumber**: The stage number, covering all stages even if the trial stops at an interim look.
 - **analysisTime**: The time for the stage since trial start.
 - **accruals1**: The number of subjects enrolled at the stage for the active treatment 1 group.
 - **accruals2**: The number of subjects enrolled at the stage for the active treatment 2 group.
 - **accruals3**: The number of subjects enrolled at the stage for the control group.

- totalAccruals: The total number of subjects enrolled at the stage.
- endpoint: The endpoint (1 for PFS or 2 for OS) under consideration.
- events1: The number of events at the stage for the active treatment 1 group.
- events2: The number of events at the stage for the active treatment 2 group.
- events3: The number of events at the stage for the control group.
- totalEvents: The total number of events at the stage.
- dropouts1: The number of dropouts at the stage for the active treatment 1 group.
- dropouts2: The number of dropouts at the stage for the active treatment 2 group.
- dropouts3: The number of dropouts at the stage for the control group.
- totalDropouts: The total number of dropouts at the stage.
- logRankStatistic13: The log-rank test Z-statistic comparing the active treatment 1 to the control for the endpoint.
- logRankStatistic23: The log-rank test Z-statistic comparing the active treatment 2 to the control for the endpoint.
- logRankStatistic12: The log-rank test Z-statistic comparing the active treatment 1 to the active treatment 2 for the endpoint.
- rawdata (exists if maxNumberOfRawDatasetsPerStage is a positive integer): A data frame for subject-level data for selected replications, containing the following variables:
 - iterationNumber: The iteration number.
 - stageNumber: The stage under consideration.
 - analysisTime: The time for the stage since trial start.
 - subjectId: The subject ID.
 - arrivalTime: The enrollment time for the subject.
 - stratum: The stratum for the subject.
 - treatmentGroup: The treatment group (1, 2, or 3) for the subject.
 - endpoint: The endpoint (1 for PFS or 2 for OS) under consideration.
 - survivalTime: The underlying survival time for the event endpoint for the subject.
 - dropoutTime: The underlying dropout time for the event endpoint for the subject.
 - timeUnderObservation: The time under observation since randomization for the event endpoint for the subject.
 - event: Whether the subject experienced the event endpoint.
 - dropoutEvent: Whether the subject dropped out for the endpoint.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
sim1 <- lrsim2e3a(
  kMax = 3,
  kMaxpfs = 2,
  allocation1 = 2,
  allocation2 = 2,
  allocation3 = 1,
```

```

    accrualTime = c(0, 8),
    accrualIntensity = c(10, 28),
    piecewiseSurvivalTime = 0,
    rho_pd_os = 0,
    lambda1pfs = log(2)/12*0.60,
    lambda2pfs = log(2)/12*0.70,
    lambda3pfs = log(2)/12,
    lambda1os = log(2)/30*0.65,
    lambda2os = log(2)/30*0.75,
    lambda3os = log(2)/30,
    n = 700,
    plannedEvents = c(186, 259, 183),
    maxNumberOfIterations = 1000,
    maxNumberOfRawDatasetsPerStage = 1,
    seed = 314159,
    nthreads = 1)

head(sim1$sumdata)
head(sim1$rawdata)

```

lrsim3a

Log-Rank Test Simulation for Three Arms

Description

Performs simulation for three-arm group sequential trials based on weighted log-rank test. The looks are driven by the total number of events in Arm A and Arm C combined. Alternatively, the analyses can be planned to occur at specified calendar times.

Usage

```

lrsim3a(
  kMax = 1,
  hazardRatioH013 = 1,
  hazardRatioH023 = 1,
  hazardRatioH012 = 1,
  allocation1 = 1,
  allocation2 = 1,
  allocation3 = 1,
  accrualTime = 0,
  accrualIntensity = NA,
  piecewiseSurvivalTime = 0,
  stratumFraction = 1,
  lambda1 = NA,
  lambda2 = NA,
  lambda3 = NA,
  gamma1 = 0,
  gamma2 = 0,

```

```

gamma3 = 0,
n = NA,
followupTime = NA,
fixedFollowup = FALSE,
rho1 = 0,
rho2 = 0,
plannedEvents = NA,
plannedTime = NA,
maxNumberOfIterations = 1000,
maxNumberOfRawDatasetsPerStage = 0,
seed = 0,
nthreads = 0
)

```

Arguments

kMax	The maximum number of stages.
hazardRatioH013	Hazard ratio under the null hypothesis for arm 1 versus arm 3. Defaults to 1 for superiority test.
hazardRatioH023	Hazard ratio under the null hypothesis for arm 2 versus arm 3. Defaults to 1 for superiority test.
hazardRatioH012	Hazard ratio under the null hypothesis for arm 1 versus arm 2. Defaults to 1 for superiority test.
allocation1	Number of subjects in Arm A in a randomization block. Defaults to 1 for equal randomization.
allocation2	Number of subjects in Arm B in a randomization block. Defaults to 1 for equal randomization.
allocation3	Number of subjects in Arm C in a randomization block. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
lambda1	A vector of hazard rates for the event in each analysis time interval by stratum for arm 1.

lambda2	A vector of hazard rates for the event in each analysis time interval by stratum for arm 2.
lambda3	A vector of hazard rates for the event in each analysis time interval by stratum for arm 3.
gamma1	The hazard rate for exponential dropout. A vector of hazard rates for piece-wise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for arm 1.
gamma2	The hazard rate for exponential dropout. A vector of hazard rates for piece-wise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for arm 2.
gamma3	The hazard rate for exponential dropout. A vector of hazard rates for piece-wise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for arm 3.
n	Sample size.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
rho1	The first parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
rho2	The second parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
plannedEvents	The planned cumulative total number of events at Look 1 to Look kMax for Arms A and C combined.
plannedTime	The calendar times for the analyses. To use calendar time to plan the analyses, plannedEvents should be missing.
maxNumberOfIterations	The number of simulation iterations. Defaults to 1000.
maxNumberOfRawDatasetsPerStage	The number of raw datasets per stage to extract.
seed	The seed to reproduce the simulation results.
nthreads	The number of threads to use in simulations (0 means the default RcppParallel behavior).

Value

A list with 2 components:

- sumdata: A data frame of summary data by iteration and stage:
 - iterationNumber: The iteration number.
 - eventsNotAchieved: Whether the target number of events is not achieved for the iteration.
 - stageNumber: The stage number, covering all stages even if the trial stops at an interim look.
 - analysisTime: The time for the stage since trial start.

- accruals1: The number of subjects enrolled at the stage for the active treatment 1 group.
- accruals2: The number of subjects enrolled at the stage for the active treatment 2 group.
- accruals3: The number of subjects enrolled at the stage for the control group.
- totalAccruals: The total number of subjects enrolled at the stage.
- events1: The number of events at the stage for the active treatment 1 group.
- events2: The number of events at the stage for the active treatment 2 group.
- events3: The number of events at the stage for the control group.
- totalEvents: The total number of events at the stage.
- dropouts1: The number of dropouts at the stage for the active treatment 1 group.
- dropouts2: The number of dropouts at the stage for the active treatment 2 group.
- dropouts3: The number of dropouts at the stage for the control group.
- totalDropouts: The total number of dropouts at the stage.
- uscore13: The log-rank test score statistic comparing the active treatment 1 to the control.
- vscore13: The log-rank test variance statistic comparing the active treatment 1 to the control.
- logRankStatistic13: The log-rank test Z-statistic comparing the active treatment 1 to the control.
- uscore23: The log-rank test score statistic comparing the active treatment 2 to the control.
- vscore23: The log-rank test variance statistic comparing the active treatment 2 to the control.
- logRankStatistic23: The log-rank test Z-statistic comparing the active treatment 2 to the control.
- uscore12: The log-rank test score statistic comparing the active treatment 1 to the active treatment 2.
- vscore12: The log-rank test variance statistic comparing the active treatment 1 to the active treatment 2.
- logRankStatistic12: The log-rank test Z-statistic comparing the active treatment 1 to the active treatment 2.
- rawdata (exists if maxNumberOfRawDatasetsPerStage is a positive integer): A data frame for subject-level data for selected replications, containing the following variables:
 - iterationNumber: The iteration number.
 - stageNumber: The stage under consideration.
 - analysisTime: The time for the stage since trial start.
 - subjectId: The subject ID.
 - arrivalTime: The enrollment time for the subject.
 - stratum: The stratum for the subject.
 - treatmentGroup: The treatment group (1, 2, or 3) for the subject.
 - survivalTime: The underlying survival time for the subject.
 - dropoutTime: The underlying dropout time for the subject.
 - timeUnderObservation: The time under observation since randomization for the subject.
 - event: Whether the subject experienced the event.
 - dropoutEvent: Whether the subject dropped out.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
sim1 <- lrsim3a(
  kMax = 3,
  allocation1 = 2,
  allocation2 = 2,
  allocation3 = 1,
  accrualTime = c(0, 8),
  accrualIntensity = c(10, 28),
  piecewiseSurvivalTime = 0,
  lambda1 = log(2)/12*0.60,
  lambda2 = log(2)/12*0.70,
  lambda3 = log(2)/12,
  n = 700,
  plannedEvents = c(186, 259, 295),
  maxNumberOfIterations = 1000,
  maxNumberOfRawDatasetsPerStage = 1,
  seed = 314159,
  nthreads = 1)

head(sim1$sumdata)
head(sim1$rawdata)
```

lrsimsub

Log-Rank Test Simulation for Enrichment Design

Description

Performs simulation for two-arm group sequential trials based on weighted log-rank test for a biomarker enrichment design. The looks are either driven by the total number of events in the ITT population or the biomarker positive sub population. Alternatively, the analyses can be planned to occur at specified calendar times.

Usage

```
lrsimsub(
  kMax = 1,
  kMaxitt = 1,
  hazardRatioH0itt = 1,
  hazardRatioH0pos = 1,
  hazardRatioH0neg = 1,
  allocation1 = 1,
  allocation2 = 1,
  accrualTime = 0,
```

```

    accrualIntensity = NA,
    piecewiseSurvivalTime = 0,
    stratumFraction = 1,
    p_pos = NA,
    lambda1itt = NA,
    lambda2itt = NA,
    lambda1pos = NA,
    lambda2pos = NA,
    gamma1itt = 0,
    gamma2itt = 0,
    gamma1pos = 0,
    gamma2pos = 0,
    n = NA,
    followupTime = NA,
    fixedFollowup = FALSE,
    rho1 = 0,
    rho2 = 0,
    plannedEvents = NA,
    plannedTime = NA,
    maxNumberOfIterations = 1000,
    maxNumberOfRawDatasetsPerStage = 0,
    seed = 0,
    nthreads = 0
)

```

Arguments

kMax	The maximum number of stages.
kMaxitt	Number of stages with timing determined by events in the ITT population. Ranges from 0 (none) to kMax.
hazardRatioH0itt	Hazard ratio under the null hypothesis for the ITT population. Defaults to 1 for superiority test.
hazardRatioH0pos	Hazard ratio under the null hypothesis for the biomarker positive sub population. Defaults to 1 for superiority test.
hazardRatioH0neg	Hazard ratio under the null hypothesis for the biomarker negative sub population. Defaults to 1 for superiority test.
allocation1	Number of subjects in the treatment group in a randomization block. Defaults to 1 for equal randomization.
allocation2	Number of subjects in the control group in a randomization block. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.

accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
p_pos	The prevalence of the biomarker positive sub population in each stratum.
lambda1itt	A vector of hazard rates for the event in each analysis time interval by stratum for the treatment group in the ITT population.
lambda2itt	A vector of hazard rates for the event in each analysis time interval by stratum for the control group in the ITT population.
lambda1pos	A vector of hazard rates for the event in each analysis time interval by stratum for the treatment group in the biomarker positive sub population.
lambda2pos	A vector of hazard rates for the event in each analysis time interval by stratum for the control group in the biomarker positive sub population.
gamma1itt	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the treatment group in the ITT population.
gamma2itt	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group in the ITT population.
gamma1pos	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the treatment group in the biomarker positive sub population.
gamma2pos	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group in the biomarker positive sub population.
n	Sample size.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
rho1	The first parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
rho2	The second parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
plannedEvents	The planned cumulative total number events in the ITT population at Look 1 to Look k_{Maxitt} and the planned cumulative total number of events at Look $k_{Maxitt}+1$ to Look k_{Max} in the biomarker positive sub population.

plannedTime	The calendar times for the analyses. To use calendar time to plan the analyses, plannedEvents should be missing.
maxNumberOfIterations	The number of simulation iterations. Defaults to 1000.
maxNumberOfRawDatasetsPerStage	The number of raw datasets per stage to extract.
seed	The seed to reproduce the simulation results.
nthreads	The number of threads to use in simulations (0 means the default RcppParallel behavior).

Value

A list with 2 components:

- **sumdata**: A data frame of summary data by iteration and stage:
 - **iterationNumber**: The iteration number.
 - **eventsNotAchieved**: Whether the target number of events is not achieved for the iteration.
 - **stageNumber**: The stage number, covering all stages even if the trial stops at an interim look.
 - **analysisTime**: The time for the stage since trial start.
 - **population**: The population ("ITT", "Biomarker Positive", "Biomarker Negative") under consideration.
 - **accruals1**: The number of subjects enrolled at the stage for the treatment group.
 - **accruals2**: The number of subjects enrolled at the stage for the control group.
 - **totalAccruals**: The total number of subjects enrolled at the stage.
 - **events1**: The number of events at the stage for the treatment group.
 - **events2**: The number of events at the stage for the control group.
 - **totalEvents**: The total number of events at the stage.
 - **dropouts1**: The number of dropouts at the stage for the treatment group.
 - **dropouts2**: The number of dropouts at the stage for the control group.
 - **totalDropouts**: The total number of dropouts at the stage.
 - **logRankStatistic**: The log-rank test Z-statistic for the population.
- **rawdata** (exists if **maxNumberOfRawDatasetsPerStage** is a positive integer): A data frame for subject-level data for selected replications, containing the following variables:
 - **iterationNumber**: The iteration number.
 - **stageNumber**: The stage under consideration.
 - **analysisTime**: The time for the stage since trial start.
 - **subjectId**: The subject ID.
 - **arrivalTime**: The enrollment time for the subject.
 - **stratum**: The stratum for the subject.
 - **biomarker**: The biomarker status for the subject (1 for positive, 0 for negative).
 - **treatmentGroup**: The treatment group (1 or 2) for the subject.
 - **survivalTime**: The underlying survival time for the subject.

- dropoutTime: The underlying dropout time for the subject.
- timeUnderObservation: The time under observation since randomization for the subject.
- event: Whether the subject experienced an event.
- dropoutEvent: Whether the subject dropped out.

Author(s)

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Examples

```
sim1 <- lrsimsub(
  kMax = 2,
  kMaxitt = 2,
  allocation1 = 1,
  allocation2 = 1,
  accrualTime = seq(0,9),
  accrualIntensity = c(seq(10,70,10),rep(70,3)),
  piecewiseSurvivalTime = c(0,12,24),
  p_pos = 0.6,
  lambda1itt = c(0.00256, 0.00383, 0.00700),
  lambda2itt = c(0.00427, 0.00638, 0.01167),
  lambda1pos = c(0.00299, 0.00430, 0.01064),
  lambda2pos = c(0.00516, 0.00741, 0.01835),
  gamma1itt = -log(1-0.04)/12,
  gamma2itt = -log(1-0.04)/12,
  gamma1pos = -log(1-0.04)/12,
  gamma2pos = -log(1-0.04)/12,
  n = 500,
  plannedEvents = c(108,144),
  maxNumberOfIterations = 1000,
  maxNumberOfRawDatasetsPerStage = 1,
  seed = 314159,
  nthreads = 1)

head(sim1$sumdata)
head(sim1$rawdata)
```

lrsim_bmTrtSel

*Simulation of a seamless phase II/III design with treatment selection
based on a short-term endpoint and toxicities*

Description

Simulates a two-stage seamless phase II/III trial in which several doses are compared with a common control. At the end of phase II a single dose is carried forward based on the posterior benefit-risk tradeoff of a binary short-term biomarker endpoint and a binary toxicity endpoint. The

confirmatory phase III analysis is performed on a time-to-event long-term endpoint linked to the biomarker through the copula correlation, and the type I error rate is protected by a closed testing procedure combined across the two stages.

Usage

```
lrsim_bmTrtSel(
  phase2SampleSizePerArm = NA_integer_,
  phase3SampleSizePerArmMin = NA_integer_,
  phase3SampleSizePerArmMax = NA_integer_,
  responseProbControl = NA_real_,
  responseProbTreatments = NA_real_,
  toxicityProbTreatments = NA_real_,
  corrEfficacyToxicity = 0,
  corrEfficacyTTE = 0,
  hazardRateControl = NA_real_,
  hazardRatioTreatments = NA_real_,
  totalNumberOfEvents = NA_integer_,
  studyDurationPhase3 = NA_real_,
  toxicityWeight = NA_real_,
  toxicityUpperLimit = NA_real_,
  efficacyThreshold = 0,
  safetyThreshold = 0,
  useUniformPrior = TRUE,
  methods = c("ctbonferroni", "ctdunnett", "ctsimes", "ctpooled", "cer", "tsssd.k",
    "tsssd.uk", "tsssd.k.rank", "tsssd.uk.rank", "tsssd.k.ce", "tsssd.uk.ce",
    "tsssd.k.rank.ce", "tsssd.uk.rank.ce", "bm.rank", "pe.rank", "naive", "ph3only"),
  accrualRatePhase2 = NA_real_,
  accrualRatePhase3 = NA_real_,
  followupTimePhase2 = 0,
  maxNumberOfIterations = 1000,
  maxNumberOfRawDatasets = 0,
  seed = 0,
  nthreads = 0
)
```

Arguments

phase2SampleSizePerArm
The number of subjects per arm enrolled in phase II (stage 1).

phase3SampleSizePerArmMin
The smallest number of subjects per arm enrolled in phase III (stage 2). Operating characteristics are reported for every stage 2 sample size from this value to **phase3SampleSizePerArmMax**.

phase3SampleSizePerArmMax
The largest number of subjects per arm enrolled in phase III (stage 2).

responseProbControl
The probability of a short-term response in the control arm.

responseProbTreatments	A vector of length M giving the probability of a short-term response for each of the M doses under investigation. Its length determines M.
toxicityProbTreatments	A vector of length M giving the probability of toxicity for each dose under investigation.
corrEfficacyToxicity	The correlation between the latent normal variable for the toxicity endpoint and z_B , where z_B is the latent normal variable that determines the short-term biomarker endpoint. Use 0 for independent biomarker and toxicity endpoints.
corrEfficacyTTE	The correlation between the latent normal variable for the long-term time-to-event endpoint and z_B . Use 0 for an independent biomarker and long-term endpoint.
hazardRateControl	The overall hazard rate for the long-term endpoint in the control arm. It must be positive.
hazardRatioTreatments	A vector of length M giving the hazard ratio for each treatment relative to the control arm. The hazard rate in treatment arm k is <code>hazardRateControl * hazardRatioTreatments[k]</code> ; values below 1 indicate effective treatments.
totalNumberOfEvents	The target total number of long-term endpoint events in the selected dose and control arms, including subjects enrolled in phases II and III. It must be a positive integer no greater than the combined sample size of those two arms at <code>phase3SampleSizePerArmMin</code> . When provided, the final analysis is event-driven and <code>studyDurationPhase3</code> is ignored.
studyDurationPhase3	The duration of phase III, measured from the opening of phase III enrollment to the final analysis. It must be provided when <code>totalNumberOfEvents</code> is missing.
toxicityWeight	The weight placed on the posterior mean toxicity rate in the benefit-risk tradeoff used for dose selection. Use 0 to select on efficacy alone.
toxicityUpperLimit	The prespecified upper limit for the toxicity rate used in the safety criterion. Use 1 when the safety criterion is not applied.
efficacyThreshold	The threshold for the posterior probability that a dose is superior to the control in short-term response. Use 0 when the efficacy criterion is not applied.
safetyThreshold	The threshold for the posterior probability that the toxicity rate of a dose is below <code>toxicityUpperLimit</code> . Use 0 when the safety criterion is not applied.
useUniformPrior	Whether to use the uniform Beta(1,1) prior (the default) or the Jeffreys Beta(0.5,0.5) prior for the beta-binomial posterior used in dose selection.
methods	A character vector naming the testing procedures to evaluate for the confirmatory analysis. Any subset of "ctbonferroni", "ctdunnett", "ctsimes",

	"ctpooled", "cer", "tsssd.k", "tsssd.uk", "tsssd.k.rank", "tsssd.uk.rank", "tsssd.k.ce", "tsssd.uk.ce", "tsssd.k.rank.ce", "tsssd.uk.rank.ce", "bm.rank", "pe.rank", "naive", and "ph3only". Restricting the set skips the corresponding computation entirely, which matters because the methods differ by orders of magnitude in cost.
accrualRatePhase2	The accrual rate per arm during phase II. Arrival times follow a homogeneous Poisson process.
accrualRatePhase3	The accrual rate per arm during phase III.
followupTimePhase2	The follow-up time after the last phase II enrollment across all arms. Phase III enrollment opens at that point. Use 0 when dose selection occurs immediately after the last phase II enrollment.
maxNumberOfIterations	The number of simulated trials.
maxNumberOfRawDatasets	The number of initial simulation iterations for which to retain subject-level raw data. Set to 0, the default, to skip raw-data retention.
seed	The seed for the random number generator.
nthreads	The number of threads to use. The default, 0, leaves the RcppParallel setting unchanged.

Details

Data generation uses a copula-based approach. For each subject, a base random variable $z_B \sim N(0, 1)$ drives the short-term biomarker endpoint, shorttv. Toxicity is generated from $z_S = \rho_{tox} z_B + \sqrt{1 - \rho_{tox}^2} z_{S, indep}$ and the toxicity indicator toxv is obtained by thresholding z_S at the specified toxicity probability. The long-term endpoint is generated from $z_E = \rho_{eff} z_B + \sqrt{1 - \rho_{eff}^2} z_{E, indep}$ as $v = -\log(\Phi(z_E))/\lambda$, where $\lambda = hazardRateControl$ in the control arm and $\lambda = hazardRateControl \times hazardRatioTreatments[k]$ in treatment arm k . Dose selection uses a beta-binomial model with independent priors, either uniform Beta(1,1) or Jeffreys Beta(0.5,0.5) depending on useUniformPrior. A dose enters the acceptable set when the posterior probability that its response rate exceeds that of the control is above efficacyThreshold and the posterior probability that its toxicity rate is below toxicityUpperLimit is above safetyThreshold. Among the acceptable doses, the one maximizing the posterior mean benefit-risk tradeoff is selected. When both thresholds are 0, all doses are acceptable.

Phase III enrollment opens followupTimePhase2 after the last phase II enrollment across all arms, and the final analysis occurs studyDurationPhase3 later. Subjects whose long-term endpoint has not occurred by then are censored at the analysis time.

Value

A list of operating characteristics. Let ngrid denote phase3SampleSizePerArmMax - phase3SampleSizePerArmMin + 1, the number of stage 2 sample sizes examined, and let M denote the number of doses. The list contains

- `n1`, `n2`, `totalNumberOfEvents`, `studyDurationPhase3`, `numberOfIterations`, `trueOBD`: The design inputs echoed back, with `n2` the vector of stage 2 sample sizes examined.
- `selectionProb`: A vector of length `M` giving the probability that each dose is selected at the end of phase II.
- `pcs`: The percentage of simulated trials selecting the dose with the largest true benefit-risk tradeoff $\text{responseProbTreatments} - \text{toxicityWeight} * \text{toxicityProbTreatments}$.
- `ave.event`: An `ngrid` by 3 matrix of the average number of events in the selected dose and the control arm combined, in stage 1, stage 2, and overall.
- `methods`: The testing procedures evaluated, in canonical order.
- `byMethod`: A named list with one element per evaluated method, each containing
 - `gpower`: A vector of length `ngrid` giving the generalized power, the probability of both selecting the true best dose and rejecting its null hypothesis.
 - `prob.rej.each`: An `ngrid` by `M` matrix of the probability of rejecting the null hypothesis for each dose conditional on that dose being selected.
 - `prob.rej.any`: A vector of length `ngrid` giving the probability of rejecting any null hypothesis.
 - `disjunctive.power`: A vector of length `ngrid` giving the probability of rejecting at least one true non-null hypothesis. It is missing when all treatment hazard ratios equal 1.
 - `conjunctive.power`: A vector of length `ngrid` giving the probability of rejecting all true non-null hypotheses. It is missing when all treatment hazard ratios equal 1.
 - `fwcr`: A vector of length `ngrid` giving the probability of rejecting any true null hypothesis.

The method names are `ctbonferroni`, `ctdunnett`, `ctsimes`, and `ctpooled` for the closed testing procedure with the inverse normal combination of stage 1 and stage 2 p-values, using the Bonferroni, Dunnett, Simes, and pooled log-rank local tests respectively; `cer` for the conditional error rate method; `tsssd.k` and `tsssd.uk` for the original two-stage seamless design boundaries with known and unknown correlation; `tsssd.k.rank` and `tsssd.uk.rank` for rank-based boundaries based on the effective number of less efficacious doses; and `tsssd.k.ce`, `tsssd.uk.ce`, `tsssd.k.rank.ce`, and `tsssd.uk.rank.ce` for conditional-error updates of the original and rank-based boundaries. These updates start from nominal boundaries based on $n1/(n1+n2)$ and then use the observed stage 1 z-statistic and observed information fraction; `bm.rank` for the Biomarker rank-based Dunnett adjustment for unknown biomarker-efficacy correlation in Wang et al. formula (3) for stage 1, along with the p-value combination test at the end of stage 2; `pe.rank` for the primary endpoint rank-based Dunnett adjustment for stage 1, along with the p-value combination test at the end of stage 2; `naive` for the unadjusted log-rank test on the combined stage 1 and stage 2 data; and `ph3only` for the unadjusted log-rank test on the stage 2 data only. Because dose selection uses stage 1 data only, `ph3only` is based on data independent of the selection and still controls the familywise error rate, at the cost of discarding the stage 1 information. `naive` reuses the selection data and is anticonservative; it is reported for reference.

- `sumdataBIN`: One row per iteration and phase II arm, containing `iterationNumber`, `treatmentGroup`, `responses`, `toxicities`, `zBiomarker`, and `selected`. Treatment group 0 is control; its toxicity count and biomarker Z statistic are missing. These rows reproduce the dose-selection operating characteristics.

- `sumdataTTE`: One row per iteration and phase III sample size, containing `iterationNumber`, `selectedDose`, `phase3SampleSize`, stage 1, stage 2, and total event counts, stage 1, stage 2, and cumulative log-rank Z statistics for the selected dose, as well as stage 1 log-rank Z statistics for all individual and pooled dose groups (e.g., `stage1LogRankZ1`, `stage1LogRankZ2`, `stage1LogRankZ12`). It also has one `reject.<method>` indicator column for every requested method. These rows reproduce `ave.event` and the operating characteristics in `byMethod`.
- `rawdataBIN` (present when `maxNumberOfRawDatasets` is positive): Subject-level phase II binary data with `iterationNumber`, `subjectId`, `treatmentGroup` (0 for control and 1 through M for doses), `response`, and `toxicity`. Toxicity is missing for the control arm because it is not simulated or used for dose selection.
- `rawdataTTE` (present when `maxNumberOfRawDatasets` is positive): Subject-level time-to-event data with `iterationNumber`, `subjectId`, `phase`, `treatmentGroup`, `response`, `arrivalTime`, `survivalTime`, `timeUnderObservation`, and `event`. Phase 2 includes all arms; phase 3 includes only the control arm and the selected dose.

Author(s)

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References

- Liyun Jiang and Ying Yuan. Seamless phase II/III design: a useful strategy to reduce the sample size for dose optimization. *Journal of the National Cancer Institute*. 2023, 115(9):1092-1098.
- Ping Gao and Yingqiu Li. Adaptive two-stage seamless sequential design for clinical trials. *Journal of Biopharmaceutical Statistics*. 2025, 35(4), 565-587.
- Cyrus Mehta, Ajoy Mukhopadhyay, and Martin Posch. Graph Based, Adaptive, Multiarm, Multiple Endpoint, Two-Stage Designs. *Statistics in Medicine*. 2025.

Examples

```
# Overall control-arm hazard rate for the long-term endpoint
hazardRateControl <- log(2) / 15.5

# response rates of the two doses under investigation
pe <- c(0.6, 0.5)

# Hazard ratio versus control for each treatment
hazardRatioTreatments <- c(0.65, 0.70)

sim <- lrsim_bmTrtSel(
  phase2SampleSizePerArm = 50,
  phase3SampleSizePerArmMin = 113,
  phase3SampleSizePerArmMax = 118,
  responseProbControl = 0.4,
  responseProbTreatments = pe,
  toxicityProbTreatments = c(0, 0),
  corrEfficacyToxicity = 0,
  corrEfficacyTTE = 0.43,
  hazardRateControl = hazardRateControl,
  hazardRatioTreatments = hazardRatioTreatments,
```

```

    studyDurationPhase3 = 42.1,
    toxicityWeight = 0,
    toxicityUpperLimit = 1,
    efficacyThreshold = 0,
    safetyThreshold = 0,
    methods = c("ctbonferroni", "ctdunnett", "ctsimes", "ctpooled",
                "cer", "naive", "ph3only"),
    accrualRatePhase2 = 3,
    accrualRatePhase3 = 6,
    followupTimePhase2 = 6,
    maxNumberOfIterations = 100,
    seed = 314159,
    nthreads = 1)

sim$pcs
sim$byMethod$ctdunnett$gpower

```

lrsim_mcpmod

Log-Rank Test Simulation for MCPMod Design

Description

Simulate an MCPMod design using a Cox model. Analyses can be triggered either by the total number of events or by the pre-specified calendar time.

Usage

```

lrsim_mcpmod(
  M = 2,
  alpha = 0.05,
  hazardRatioH0s = 1,
  allocations = 1,
  accrualTime = 0,
  accrualIntensity = NA,
  piecewiseSurvivalTime = 0,
  stratumFraction = 1,
  lambdas = NULL,
  candidateHazardRatios = NULL,
  gammas = NULL,
  n = NA,
  followupTime = NA,
  fixedFollowup = FALSE,
  plannedEvents = NA,
  plannedTime = NA,
  maxNumberOfIterations = 1000,
  maxNumberOfRawDatasetsPerStage = 0,
  seed = 0,
  nthreads = 0
)

```

Arguments

<code>M</code>	Number of active treatment arms.
<code>alpha</code>	Significance level for the max test (1-sided).
<code>hazardRatioH0s</code>	Scalar or numeric vector of length M . Hazard ratios under H_0 for each active arm versus the common control. Defaults to 1 for superiority tests.
<code>allocations</code>	Integer or integer vector of length $M + 1$. Number of subjects per arm within a randomization block.
<code>accrualTime</code>	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
<code>accrualIntensity</code>	A vector of accrual intensities. One for each accrual time interval.
<code>piecewiseSurvivalTime</code>	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
<code>stratumFraction</code>	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
<code>lambdas</code>	List of length $M + 1$ (one element per arm). Each element is a scalar or a numeric vector of event hazard rates for the corresponding arm.
<code>candidateHazardRatios</code>	Numeric matrix of dimension $M \times T$ containing the candidate hazard ratios for the T candidate models.
<code>gammas</code>	List of length $M + 1$ (one element per arm). Each element is a scalar or a numeric vector of dropout hazard rates for the corresponding arm.
<code>n</code>	Planned total sample size across all active arms and control.
<code>followupTime</code>	Follow-up time for the last enrolled subject.
<code>fixedFollowup</code>	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
<code>plannedEvents</code>	Planned total number of events to trigger the analysis. Leave missing when using calendar-time planning.
<code>plannedTime</code>	Planned calendar time for the analysis. Leave missing when using event-based planning.
<code>maxNumberOfIterations</code>	Number of Monte Carlo replications.
<code>maxNumberOfRawDatasetsPerStage</code>	Number of subject-level raw datasets to retain.
<code>seed</code>	Random seed for reproducibility.
<code>nthreads</code>	Number of threads for parallel simulation. Use 0 to accept the default RcppParallel behavior.

Value

An S3 object of class "lrsim_mcpmod" with components overview, sumdata1, sumdata2, and optionally rawdata.

Examples

```
## Not run:
# Define candidate dose-response models
# (negative sign models decreasing dose-response for hazard)
f_emax <- function(d, ED50) { -d / (ED50 + d) }
f_exponential <- function(d, delta) { -exp(d / delta) + 1 }
f_linear <- function(d) { -d }
f_logistic <- function(d, ED50, delta) {
  -1 / (1 + exp((ED50 - d) / delta))
}
f_betamod <- function(d, delta1, delta2, D) {
  -dbeta(d / D, delta1 + 1, delta2 + 1)
}

# Log hazard parameters
b0 <- log(log(2) / 0.5) # placebo: median survival 0.5 years
b1 <- log(0.6) + b0     # optimal dose: HR = 0.6 vs. placebo

# Rescale candidate models to match hazard rates
f1 <- function(dose) {
  a0 <- f_emax(0, 50); a1 <- f_emax(100, 50)
  slope <- (b1 - b0) / (a1 - a0)
  intercept <- b0 - slope * a0
  intercept + slope * f_emax(dose, 50)
}
f2 <- function(dose) {
  a0 <- f_exponential(0, 22.756); a1 <- f_exponential(100, 22.756)
  slope <- (b1 - b0) / (a1 - a0)
  intercept <- b0 - slope * a0
  intercept + slope * f_exponential(dose, 22.756)
}
f3 <- function(dose) {
  a0 <- f_emax(0, 6.25); a1 <- f_emax(100, 6.25)
  slope <- (b1 - b0) / (a1 - a0)
  intercept <- b0 - slope * a0
  intercept + slope * f_emax(dose, 6.25)
}
f4 <- function(dose) {
  a0 <- f_logistic(0, 40.3287, 6.9764)
  a1 <- f_logistic(100, 40.3287, 6.9764)
  slope <- (b1 - b0) / (a1 - a0)
  intercept <- b0 - slope * a0
  intercept + slope * f_logistic(dose, 40.3287, 6.9764)
}
f5 <- function(dose) {
  a0 <- f_linear(0); a1 <- f_linear(100)
  slope <- (b1 - b0) / (a1 - a0)
  intercept <- b0 - slope * a0
  intercept + slope * f_linear(dose)
}
f6 <- function(dose) {
  a0 <- f_betamod(0, 0.7489, 1.0485, 120)
```

```

dstar <- 0.7489 * 120 / (0.7489 + 1.0485)
a1 <- f_betamod(dstar, 0.7489, 1.0485, 120)
slope <- (b1 - b0) / (a1 - a0)
intercept <- b0 - slope * a0
intercept + slope * f_betamod(dose, 0.7489, 1.0485, 120)
}

f <- list(f1, f2, f3, f4, f5, f6)
doselevels <- c(5, 25, 50, 100)

# Run the simulations (each candidate model is the true model in one scenario)
sims <- lapply(1:6, function(i) {
  lrsim_mcpmod(
    M = 4, alpha = 0.05, accrualIntensity = 360,
    lambdas = as.list(exp(f[[i]](c(doselevels, 0)))),
    candidateHazardRatios = exp(sapply(f, function(ff) ff(doselevels) - ff(0))),
    gammas = list(0, 0, 0, 0, 0),
    n = 300, plannedEvents = 242,
    maxNumberOfIterations = 1000,
    maxNumberOfRawDatasetsPerStage = 10,
    seed = 314159,
    nthreads = 1
  )
})

sapply(sims, function(s) s$overview$overallReject)

## End(Not run)

```

lrsim_multiarm

Log-Rank Test Simulation for Multi-Arm Multi-Stage Design

Description

Simulate a multi-arm multi-stage design using a weighted log-rank test. Analyses can be triggered either by the cumulative number of events (combined for an active arm and the common control) or by pre-specified calendar times.

Usage

```

lrsim_multiarm(
  M = 2,
  kMax = 1,
  criticalValues = NULL,
  futilityBounds = NULL,
  hazardRatioH0s = 1,
  allocations = 1,
  accrualTime = 0,

```

```

    accrualIntensity = NA,
    piecewiseSurvivalTime = 0,
    stratumFraction = 1,
    lambdas = NULL,
    gammas = NULL,
    n = NA,
    followupTime = NA,
    fixedFollowup = FALSE,
    rho1 = 0,
    rho2 = 0,
    plannedEvents = NA,
    plannedTime = NA,
    maxNumberOfIterations = 1000,
    maxNumberOfRawDatasetsPerStage = 0,
    seed = 0,
    nthreads = 0
)

```

Arguments

M	Number of active treatment arms.
kMax	Number of sequential looks.
criticalValues	Numeric matrix of dimension $kMax \times M$ giving the by-look critical values for the closed testing procedure. The first column is used for the level-M test and the last column for the level-1 test.
futilityBounds	Numeric vector of length $kMax - 1$ giving the futility boundaries on the Wald-statistic scale for the first $kMax - 1$ looks. At an interim look, the study stops for futility if all active treatment arms fall below the futility boundary. If omitted, no interim futility stopping is applied.
hazardRatioH0s	Scalar or numeric vector of length M . Hazard ratios under H_0 for each active arm versus the common control. Defaults to 1 for superiority tests.
allocations	Integer or integer vector of length $M + 1$. Number of subjects per arm within a randomization block. A single value implies equal allocation; defaults to 1. The first M elements refer to the active arms and the last element refers to the common control.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.

lambdas	List of length $M + 1$ (one element per arm). Each element is a scalar or a numeric vector of event hazard rates for the corresponding arm, given by analysis interval and stratum as required by the simulation. The first M elements refer to the active arms and the last element refers to the common control.
gammas	List of length $M + 1$ (one element per arm). Each element is a scalar or a numeric vector of dropout hazard rates for the corresponding arm, by analysis interval and stratum. The first M elements refer to the active arms and the last element refers to the common control.
n	Planned total sample size across all active arms and control.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
rho1	The first parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
rho2	The second parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
plannedEvents	Numeric vector of length $K + 1$ giving the planned cumulative number of events to trigger Look 1 through Look $K + 1$. Each entry refers to the combined events for the first active arm and the common control. Use plannedEvents to schedule event-driven looks.
plannedTime	Numeric vector of calendar times for the analyses. If plannedTime is supplied, analyses are scheduled by calendar time and plannedEvents should be left missing.
maxNumberOfIterations	Number of Monte Carlo replications. Defaults to 1000.
maxNumberOfRawDatasetsPerStage	Number of subject-level raw datasets to retain per stage (for selected replications).
seed	Random seed for reproducibility.
nthreads	Number of threads for parallel simulation. Use 0 to accept the default RcppParallel behavior.

Value

An S3 object of class "lrsim_seamless" with these components:

- overview: A list summarizing trial-level results and settings:
 - overallReject: Overall probability of rejecting the null by trial end.
 - overallFutility: Overall probability of stopping for futility by trial end.
 - rejectPerStage: Probability of rejecting the null for each active arm at each stage.
 - futilityPerStage: Probability of futility stopping for each arm at each stage.
 - cumulativeRejection: Cumulative probability of rejection for each active arm by stage.
 - cumulativeFutility: Cumulative probability of futility stopping for each active arm by stage.

- numberOfEvents: Cumulative event counts by stage and arm.
- numberOfDropouts: Cumulative dropouts by stage and arm.
- numberOfSubjects: Cumulative enrollments by stage and arm.
- analysisTime: Average calendar time for each stage by arm among replications that reached that stage.
- expectedNumberOfEvents: Expected cumulative events at trial end.
- expectedNumberOfDropouts: Expected cumulative dropouts at trial end.
- expectedNumberOfSubjects: Expected cumulative enrollments at trial end.
- expectedStudyDuration: Expected study duration.
- criticalValues: The input matrix of by-level critical boundaries.
- futilityBounds: The input vector of futility boundaries.
- hazardRatioH0s: The input hazard ratios under H_0 .
- useEvents: Logical indicating whether analyses were event-driven.
- numberOfIterations: Number of simulation iterations performed.
- n: Planned total sample size.
- allocations: The input allocation ratios.
- fixedFollowup: Logical indicating whether fixed follow-up was used.
- rho1, rho2: Fleming–Harrington weighting parameters used.
- M: Number of active arms in Phase 2.
- K: Number of sequential looks in Phase 3.
- sumdata1: Data frame summarizing each iteration, stage, and treatment group:
 - iterationNumber, eventsNotAchieved, stopStage, stageNumber, analysisTime, treatmentGroup, accruals, events, dropouts.
 - For each stage the final row summarizes the overall study (all arms combined).
- sumdata2: Data frame summarizing log-rank statistics by iteration, stage, and active arm:
 - iterationNumber, stopStage, stageNumber, analysisTime, activeArm, totalAccruals, totalEvents, totalDropouts, uscore, vscore, logRankStatistic, reject, futility.
 - For each active arm, total accruals, events, and dropouts refer to the combined counts for that arm and the common control at that stage.
- rawdata (present when maxNumberOfRawDatasetsPerStage > 0): Subject-level data for selected replications with variables:
 - iterationNumber, stopStage, stageNumber, analysisTime, subjectId, arrivalTime, stratum, treatmentGroup, survivalTime, dropoutTime, timeUnderObservation, event, dropoutEvent.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(sim1 <- lrsim_multiarm(
  M = 2,
  kMax = 3,
```

```
criticalValues = matrix(c(3.880, 2.747, 2.275,
                        3.710, 2.511, 1.993), 3, 2),
futilityBounds = c(0.074, 1.207),
accrualTime = c(0, 8),
accrualIntensity = c(10, 28),
piecewiseSurvivalTime = 0,
lambdas = list(log(2)/12*0.5, log(2)/12*0.75, log(2)/12),
n = 700,
plannedEvents = c(36, 72, 108),
maxNumberOfIterations = 10000,
maxNumberOfRawDatasetsPerStage = 1,
seed = 314159,
nthreads = 1))
```

lrsim_seamless

Log-Rank Test Simulation for Phase 2/3 Seamless Design

Description

Simulate phase 2/3 seamless design using a weighted log-rank test. Analyses can be triggered either by the cumulative number of events (combined for an active arm and the common control) or by pre-specified calendar times.

Usage

```
lrsim_seamless(
  M = 2,
  K = 1,
  criticalValues = NA,
  futilityBounds = NULL,
  hazardRatioH0s = 1,
  allocations = 1,
  accrualTime = 0,
  accrualIntensity = NA,
  piecewiseSurvivalTime = 0,
  stratumFraction = 1,
  lambdas = NULL,
  gammas = NULL,
  n = NA,
  followupTime = NA,
  fixedFollowup = FALSE,
  rho1 = 0,
  rho2 = 0,
  plannedEvents = NA,
  plannedTime = NA,
  maxNumberOfIterations = 1000,
  maxNumberOfRawDatasetsPerStage = 0,
```

```

    seed = 0,
    nthreads = 0
)

```

Arguments

M	Number of active treatment arms in Phase 2.
K	Number of sequential looks in Phase 3.
criticalValues	Numeric vector of length $K + 1$ giving the critical value for the Wald statistic at each look (Look 1 through Look $K + 1$). Decision rule: • At Look 1, compute the Wald statistic (which equals the negative of the logrank statistic) for each active arm versus the common control. If the largest test statistic exceeds the Look 1 critical value, stop for efficacy. • If the Look 1 stopping rule is not met, select the active arm with the largest Wald statistic and continue with that arm only versus control at subsequent looks. • For each look $j = 2, \dots, K + 1$, compare the selected arm to control; if its Wald statistic exceeds the Look j critical value, stop for efficacy; otherwise continue. • If no critical value is exceeded by Look $K + 1$, the procedure ends without rejection.
futilityBounds	Numeric vector of length K giving the futility boundaries for Phase 2 and the first $K - 1$ looks in Phase 3. The study stops for futility: • in Phase 2 if the selected treatment arm crosses the phase-2 futility boundary; • in Phase 3 if the selected arm crosses the futility boundary at an interim look; If omitted, no interim futility stopping is applied.
hazardRatioH0s	Numeric vector of length M . Hazard ratios under H_0 for each active arm versus the common control. Defaults to 1 for superiority tests.
allocations	Integer or integer vector of length $M + 1$. Number of subjects per arm within a randomization block. A single value implies equal allocation; defaults to 1. The first M elements refer to the active arms and the last element refers to the common control.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.

lambdas	List of length $M + 1$ (one element per arm). Each element is a scalar or a numeric vector of event hazard rates for the corresponding arm, given by analysis interval and stratum as required by the simulation. The first M elements refer to the active arms and the last element refers to the common control.
gammas	List of length $M + 1$ (one element per arm). Each element is a scalar or a numeric vector of dropout hazard rates for the corresponding arm, by analysis interval and stratum. The first M elements refer to the active arms and the last element refers to the common control.
n	Planned total sample size across all active arms and control.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
rho1	The first parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
rho2	The second parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
plannedEvents	Numeric vector of length $K + 1$ giving the planned cumulative number of events to trigger Look 1 through Look $K + 1$. Each entry refers to the combined events for the first active arm and the common control. Use plannedEvents to schedule event-driven looks.
plannedTime	Numeric vector of calendar times for the analyses. If plannedTime is supplied, analyses are scheduled by calendar time and plannedEvents should be left missing.
maxNumberOfIterations	Number of Monte Carlo replications. Defaults to 1000.
maxNumberOfRawDatasetsPerStage	Number of subject-level raw datasets to retain per stage (for selected replications).
seed	Random seed for reproducibility.
nthreads	Number of threads for parallel simulation. Use 0 to accept the default RcppParallel behavior.

Value

An S3 object of class "lrsim_seamless" with these components:

- overview: A list summarizing trial-level results and settings:
 - selectionProb: Probability of selecting each active arm at the end of phase 2.
 - selectToStage2: Probability of selecting each active arm to enter stage 2.
 - selectAnyToStage2: Probability of selecting any active arm to enter stage 2.
 - rejectPerStage: Probability of rejecting the null for each active arm at each stage.
 - futilityPerStage: Probability of futility stopping for each active arm at each stage.
 - cumulativeRejection: Cumulative probability of rejection by stage.
 - cumulativeFutility: Cumulative futility stopping probabilities by stage.

- numberOfEvents: Cumulative event counts by stage, including events from all arms in stage 1 and events from the selected arm and control in later stages.
- numberOfDropouts: Cumulative dropouts by stage.
- numberOfSubjects: Cumulative enrollments by stage.
- analysisTime: Average calendar time for each stage among replications that reached that stage.
- overallReject: Overall probability of rejecting the null by trial end.
- overallFutility: Overall probability of stopping for futility by trial end.
- expectedNumberOfEvents: Expected cumulative events at trial end.
- expectedNumberOfDropouts: Expected cumulative dropouts at trial end.
- expectedNumberOfSubjects: Expected cumulative enrollments at trial end.
- expectedStudyDuration: Expected study duration.
- criticalValues: The input critical values for each stage.
- futilityBounds: The input futility boundaries for each stage.
- hazardRatioH0s: The input hazard ratios under H_0 .
- useEvents: Logical indicating whether analyses were event-driven.
- numberOfIterations: Number of simulation iterations performed.
- n: Planned total sample size.
- allocations: The input allocation ratios.
- fixedFollowup: Logical indicating whether fixed follow-up was used.
- rho1, rho2: Fleming–Harrington weighting parameters used.
- M: Number of active arms in Phase 2.
- K: Number of sequential looks in Phase 3.
- sumdata1: Data frame summarizing each iteration, stage, and treatment group:
 - iterationNumber, eventsNotAchieved, stopStage, stageNumber, analysisTime, treatmentGroup, accruals, events, dropouts.
 - For each stage the final row summarizes the overall study (all arms combined).
- sumdata2: Data frame summarizing log-rank statistics by iteration, stage, and active arm:
 - iterationNumber, selectedArm, stopStage, stageNumber, analysisTime, activeArm, totalAccruals, totalEvents, totalDropouts, uscore, vscore, logRankStatistic, reject, futility.
 - For each active arm, total accruals, events, and dropouts refer to the combined counts for that arm and the common control at that stage.
- rawdata (present when maxNumberOfRawDatasetsPerStage > 0): Subject-level data for selected replications with variables:
 - iterationNumber, stopStage, stageNumber, analysisTime, subjectId, arrivalTime, stratum, treatmentGroup, survivalTime, dropoutTime, timeUnderObservation, event, dropoutEvent.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
(sim1 <- lrstat_seamless(
  M = 2,
  K = 2,
  criticalValues = c(3.882, 2.733, 2.222),
  futilityBounds = c(0.259, 1.201),
  accrualTime = c(0, 8),
  accrualIntensity = c(10, 28),
  piecewiseSurvivalTime = 0,
  lambdas = list(log(2)/12*0.5, log(2)/12*0.7, log(2)/12),
  n = 700,
  plannedEvents = c(42, 84, 126),
  maxNumberOfIterations = 10000,
  maxNumberOfRawDatasetsPerStage = 1,
  seed = 314159,
  nthreads = 1))
```

lrstat

Number of Subjects Having an Event and Log-Rank Statistics

Description

Obtains the number of subjects accrued, number of events, number of dropouts, and number of subjects reaching the maximum follow-up in each group, mean and variance of weighted log-rank score statistic, estimated hazard ratio from weighted Cox regression and variance of log hazard ratio estimate at given calendar times.

Usage

```
lrstat(
  time = NA_real_,
  hazardRatioH0 = 1,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = FALSE,
  rho1 = 0,
  rho2 = 0,
  predictTarget = 2L
)
```

Arguments

<code>time</code>	A vector of calendar times at which to calculate the number of events and the mean and variance of log-rank test score statistic.
<code>hazardRatioH0</code>	Hazard ratio under the null hypothesis for the active treatment versus control. Defaults to 1 for superiority test.
<code>allocationRatioPlanned</code>	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
<code>accrualTime</code>	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
<code>accrualIntensity</code>	A vector of accrual intensities. One for each accrual time interval.
<code>piecewiseSurvivalTime</code>	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
<code>stratumFraction</code>	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
<code>lambda1</code>	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.
<code>lambda2</code>	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
<code>gamma1</code>	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
<code>gamma2</code>	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
<code>accrualDuration</code>	Duration of the enrollment period.
<code>followupTime</code>	Follow-up time for the last enrolled subject.
<code>fixedFollowup</code>	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
<code>rho1</code>	The first parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
<code>rho2</code>	The second parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
<code>predictTarget</code>	The target of prediction. Set <code>predictTarget = 1</code> to predict the number of events only. Set <code>predictTarget = 2</code> (default) to predict the number of events and log-rank score statistic mean and variance. Set <code>predictTarget = 3</code> to predict the number of events, log-rank score statistic mean and variance, and hazard ratio and variance of log hazard ratio.

Value

A data frame containing the following variables if predictTarget = 1:

- time: The analysis time since trial start.
- subjects: The number of enrolled subjects.
- nevents: The total number of events.
- nevents1: The number of events in the active treatment group.
- nevents2: The number of events in the control group.
- ndropouts: The total number of dropouts.
- ndropouts1: The number of dropouts in the active treatment group.
- ndropouts2: The number of dropouts in the control group.
- nfmaz: The total number of subjects reaching maximum follow-up.
- nfmaz1: The number of subjects reaching maximum follow-up in the active treatment group.
- nfmaz2: The number of subjects reaching maximum follow-up in the control group.

If predictTarget = 2, the following variables will also be included:

- uscore: The numerator of the log-rank test statistic.
- vscore: The variance of the log-rank score test statistic.
- logRankZ: The log-rank test statistic on the Z-scale.
- hazardRatioH0: The hazard ratio under the null hypothesis.

Furthermore, if predictTarget = 3, the following additional variables will also be included:

- HR: The average hazard ratio from weighted Cox regression.
- vlogHR: The variance of log hazard ratio.
- zlogHR: The Z-statistic for log hazard ratio.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Piecewise accrual, piecewise exponential survivals, and 5% dropout by
# the end of 1 year.
```

```
lrstat(time = c(22, 40), allocationRatioPlanned = 1,
       accrualTime = seq(0, 8),
       accrualIntensity = 26/9*seq(1, 9),
       piecewiseSurvivalTime = c(0, 6),
       lambda1 = c(0.0533, 0.0309),
       lambda2 = c(0.0533, 0.0533),
       gamma1 = -log(1-0.05)/12,
       gamma2 = -log(1-0.05)/12,
       accrualDuration = 22,
       followupTime = 18, fixedFollowup = FALSE)
```

lrtest

*Log-Rank Test of Survival Curve Difference***Description**

Obtains the log-rank test using the Fleming-Harrington family of weights.

Usage

```
lrtest(
  data,
  stratum = "",
  treat = "treat",
  time = "time",
  time2 = "",
  event = "event",
  weight = "",
  weight_readj = FALSE,
  rho1 = 0,
  rho2 = 0
)
```

Arguments

data	The input data frame or list of data frames that contains the following variables: • stratum: The stratum. • treat: The treatment. • time: The follow-up time for right censored data, or the left end of each interval for counting process data. • time2: The right end of each interval for counting process data. Intervals are assumed to be open on the left and closed on the right, and event indicates whether an event occurred at the right end of each interval. • event: The event indicator, 1=event, 0=no event. • weight: The weight for each observation.
stratum	The name(s) of the stratum variable(s) in the input data.
treat	The name of the treatment variable in the input data.
time	The name of the time variable or the left end of each interval for counting process data in the input data.
time2	The name of the right end of each interval for counting process data in the input data.
event	The name of the event variable in the input data.
weight	The name of the weight variable in the input data.
weight_readj	Whether the weight variable at each event time will be readjusted to be proportional to the number at risk by treatment group. Defaults to FALSE.

- rho1 The first parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.
- rho2 The second parameter of the Fleming-Harrington family of weighted log-rank test. Defaults to 0 for conventional log-rank test.

Value

A data frame with the following variables:

- uscore: The numerator of the log-rank test statistic.
- vscore: The variance of the log-rank score test statistic.
- logRankZ: The Z-statistic value.
- logRankPValue: The two-sided p-value.
- weight_readj: Whether the weight variable will be readjusted.
- rho1: The first parameter of the Fleming-Harrington weights.
- rho2: The second parameter of the Fleming-Harrington weights.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
lrtest(rawdata[rawdata$iterationNumber == 1, ],
       stratum = "stratum", treat = "treatmentGroup",
       time = "timeUnderObservation", event = "event",
       rho1 = 0.5, rho2 = 0)
```

mnOddsRatioCI	<i>Miettinen-Nurminen Score Confidence Interval for Two-Sample Odds Ratio</i>
---------------	---

Description

Obtains the Miettinen-Nurminen score confidence interval for two-sample odds ratio possibly with stratification.

Usage

```
mnOddsRatioCI(n1, y1, n2, y2, cilevel = 0.95)
```

Arguments

n1	The sample size for the active treatment group.
y1	The number of responses for the active treatment group.
n2	The sample size for the control group.
y2	The number of responses for the control group.
cilevel	The confidence interval level.

Details

The Mantel-Haenszel sample size weights are used for stratified samples.

Value

A list with two components:

- data: A data frame containing the input sample size and number of responses for each treatment group. It has the following variables:
 - n1: The sample size for the active treatment group.
 - y1: The number of responses for the active treatment group.
 - n2: The sample size for the control group.
 - y2: The number of responses for the control group.
- estimates: A data frame containing the point estimate and confidence interval for odds ratio. It has the following variables:
 - scale: The scale of treatment effect.
 - estimate: The point estimate.
 - lower: The lower limit of the confidence interval.
 - upper: The upper limit of the confidence interval.
 - cilevel: The confidence interval level.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
mnOddsRatioCI(n1 = c(10,10), y1 = c(4,3), n2 = c(20,10), y2 = c(2,0))
```

mnRateDiffCI	<i>Miettinen-Nurminen Score Confidence Interval for Two-Sample Rate Difference</i>
--------------	--

Description

Obtains the Miettinen-Nurminen score confidence interval for two-sample rate difference possibly with stratification.

Usage

```
mnRateDiffCI(t1, y1, t2, y2, cilevel = 0.95)
```

Arguments

t1	The exposure for the active treatment group.
y1	The number of events for the active treatment group.
t2	The exposure for the control group.
y2	The number of events for the control group.
cilevel	The confidence interval level.

Details

The Mantel-Haenszel weights are used for stratified samples.

Value

A list with two components:

- data A data frame containing the input exposure and number of events for each treatment group. It has the following variables:
 - t1: The exposure for the active treatment group.
 - y1: The number of events for the active treatment group.
 - t2: The exposure for the control group.
 - y2: The number of events for the control group.
- estimates: A data frame containing the point estimate and confidence interval for rate difference. It has the following variables:
 - scale: The scale of treatment effect.
 - estimate: The point estimate.
 - lower: The lower limit of the confidence interval.
 - upper: The upper limit of the confidence interval.
 - cilevel: The confidence interval level.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
mnRateDiffCI(t1 = c(10,10), y1 = c(4,3), t2 = c(20,10), y2 = c(2,0))
```

mnRateRatioCI	<i>Miettinen-Nurminen Score Confidence Interval for Two-Sample Rate Ratio</i>
---------------	---

Description

Obtains the Miettinen-Nurminen score confidence interval for two-sample rate ratio possibly with stratification.

Usage

```
mnRateRatioCI(t1, y1, t2, y2, cilevel = 0.95)
```

Arguments

t1	The exposure for the active treatment group.
y1	The number of events for the active treatment group.
t2	The exposure for the control group.
y2	The number of events for the control group.
cilevel	The confidence interval level.

Details

The Mantel-Haenszel weights are used for stratified samples.

Value

A list with two components:

- data A data frame containing the input exposure and number of events for each treatment group. It has the following variables:
 - t1: The exposure for the active treatment group.
 - y1: The number of events for the active treatment group.
 - t2: The exposure for the control group.
 - y2: The number of events for the control group.
- estimates: A data frame containing the point estimate and confidence interval for rate ratio. It has the following variables:

- scale: The scale of treatment effect.
- estimate: The point estimate.
- lower: The lower limit of the confidence interval.
- upper: The upper limit of the confidence interval.
- cilevel: The confidence interval level.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
mnRateRatioCI(t1 = c(10,10), y1 = c(4,3), t2 = c(20,10), y2 = c(2,0))
```

mnRiskDiffCI	<i>Miettinen-Nurminen Score Confidence Interval for Two-Sample Risk Difference</i>
--------------	--

Description

Obtains the Miettinen-Nurminen score confidence interval for two-sample risk difference possibly with stratification.

Usage

```
mnRiskDiffCI(n1, y1, n2, y2, cilevel = 0.95)
```

Arguments

- | | |
|---------|---|
| n1 | The sample size for the active treatment group. |
| y1 | The number of responses for the active treatment group. |
| n2 | The sample size for the control group. |
| y2 | The number of responses for the control group. |
| cilevel | The confidence interval level. |

Details

The Mantel-Haenszel sample size weights are used for stratified samples.

Value

A list with two components:

- data: A data frame containing the input sample size and number of responses for each treatment group. It has the following variables:
 - n1: The sample size for the active treatment group.
 - y1: The number of responses for the active treatment group.
 - n2: The sample size for the control group.
 - y2: The number of responses for the control group.
- estimates: A data frame containing the point estimate and confidence interval for risk difference. It has the following variables:
 - scale: The scale of treatment effect.
 - estimate: The point estimate.
 - lower: The lower limit of the confidence interval.
 - upper: The upper limit of the confidence interval.
 - cilevel: The confidence interval level.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
mnRiskDiffCI(n1 = c(10, 10), y1 = c(4, 3),
              n2 = c(20, 10), y2 = c(2, 0))
```

mnRiskRatioCI	<i>Miettinen-Nurminen Score Confidence Interval for Two-Sample Risk Ratio</i>
---------------	---

Description

Obtains the Miettinen-Nurminen score confidence interval for two-sample risk ratio possibly with stratification.

Usage

```
mnRiskRatioCI(n1, y1, n2, y2, cilevel = 0.95)
```

Arguments

n1	The sample size for the active treatment group.
y1	The number of responses for the active treatment group.
n2	The sample size for the control group.
y2	The number of responses for the control group.
cilevel	The confidence interval level.

Details

The Mantel-Haenszel sample size weights are used for stratified samples.

Value

A list with two components:

- data A data frame containing the input sample size and number of responses for each treatment group. It has the following variables:
 - n1: The sample size for the active treatment group.
 - y1: The number of responses for the active treatment group.
 - n2: The sample size for the control group.
 - y2: The number of responses for the control group.
- estimates: A data frame containing the point estimate and confidence interval for risk ratio. It has the following variables:
 - scale: The scale of treatment effect.
 - estimate: The point estimate.
 - lower: The lower limit of the confidence interval.
 - upper: The upper limit of the confidence interval.
 - cilevel: The confidence interval level.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
mnRiskRatioCI(n1 = c(10, 10), y1 = c(4, 3),
               n2 = c(20, 10), y2 = c(2, 0))
```

mTPI2Table	<i>mTPI-2 Decision Table</i>
------------	------------------------------

Description

Obtains the decision table for the modified toxicity probability interval-2 (mTPI-2) design.

Usage

```
mTPI2Table(
  nMax = NA_integer_,
  pT = 0.3,
  epsilon1 = 0.05,
  epsilon2 = 0.05,
  a = 1,
```

```

    b = 1,
    pExcessTox = 0.95
  )

```

Arguments

nMax	The maximum number of subjects in a dose cohort.
pT	The target toxicity probability. Defaults to 0.3.
epsilon1	The lower equivalence margin from the target. Defaults to 0.05.
epsilon2	The upper equivalence margin from the target. Defaults to 0.05.
a	The prior toxicity parameter for the beta prior.
b	The prior non-toxicity parameter for the beta prior.
pExcessTox	The threshold for excessive toxicity, i.e., if $\text{Prob}(p > pT \mid \text{Data}) > p\text{ExcessTox}$, then the current and all higher doses will be excluded and never be used again in the remainder of the trial to avoid any other subjects receiving treatment at those doses. Defaults to 0.95.

Value

An S3 class `mTPI2Table` object with the following components:

- `settings`: The input settings data frame with the following variables:
 - `nMax`: The maximum number of subjects in a dose cohort.
 - `pT`: The target toxicity probability.
 - `epsilon1`: The lower equivalence margin from the target.
 - `epsilon2`: The upper equivalence margin from the target.
 - `a`: The prior toxicity parameter for the beta prior.
 - `b`: The prior non-toxicity parameter for the beta prior.
 - `pExcessTox`: The threshold for excessive toxicity.
- `subintervals`: The subintervals of equal length in the mTPI-2 design. It includes the following variables:
 - `lower`: The lower bound of the subinterval.
 - `upper`: The upper bound of the subinterval.
 - `decision`: The dosing decision for the subinterval.
- `decisionDataFrame`: The decision data frame for the mTPI-2 design. It includes the following variables:
 - `n`: The sample size.
 - `y`: The number of toxicities.
 - `decision`: The dosing decision.
- `decisionMatrix`: The decision matrix corresponding to the decision data frame.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

References

Guo, W., Wang, S. J., Yang, S., Lynn, H., & Ji, Y. (2017). A Bayesian interval dose-finding design addressing Ockham’s razor: mTPI-2. Contemporary Clinical Trials, 58, 23-33.

Examples

```
mTPI2Table(nMax = 18, pT = 0.3, epsilon1 = 0.05, epsilon2 = 0.05)
```

mtpwexp	<i>Mean and Variance of Truncated Piecewise Exponential Distribution</i>
---------	--

Description

Obtains the mean and variance from a truncated piecewise exponential distribution.

Usage

```
mtpwexp(piecewiseSurvivalTime = 0L, lambda = NA_real_, lowerBound = 0)
```

Arguments

- piecewiseSurvivalTime**
A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., `c(0, 6)` breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
- lambda**
A vector of hazard rates for the event. One for each analysis time interval.
- lowerBound**
The left truncation time point for the survival time. Defaults to 0 for no truncation.

Value

A list with two components, one for the mean, and the other for the variance of the truncated piecewise exponential distribution.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
mtpwexp(piecewiseSurvivalTime = c(0, 6, 9, 15),  
        lambda = c(0.025, 0.04, 0.015, 0.007))
```

natrisk	<i>Number of Subjects at Risk</i>
---------	-----------------------------------

Description

Obtains the number of subjects at risk at given analysis times for each treatment group.

Usage

```
natrisk(
  t = NA_real_,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  accrualDuration = NA_real_,
  maxFollowupTime = NA_real_,
  time = NA_real_
)
```

Arguments

<code>t</code>	A vector of analysis times at which to calculate the number of patients at risk.
<code>allocationRatioPlanned</code>	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
<code>accrualTime</code>	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., <code>c(0, 3)</code> breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
<code>accrualIntensity</code>	A vector of accrual intensities. One for each accrual time interval.
<code>piecewiseSurvivalTime</code>	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., <code>c(0, 6)</code> breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
<code>lambda1</code>	A vector of hazard rates for the event for the active treatment group. One for each analysis time interval.
<code>lambda2</code>	A vector of hazard rates for the event for the control group. One for each analysis time interval.
<code>gamma1</code>	The hazard rate for exponential dropout, or a vector of hazard rates for piecewise exponential dropout for the active treatment group.

gamma2	The hazard rate for exponential dropout, or a vector of hazard rates for piecewise exponential dropout for the control group.
accrualDuration	Duration of the enrollment period.
maxFollowupTime	Follow-up time for the first enrolled subject. For fixed follow-up, maxFollowupTime = minFollowupTime. For variable follow-up, maxFollowupTime = accrualDuration + minFollowupTime.
time	Calendar time for the analysis.

Details

For a given treatment group g and calendar time τ , the number of patients at risk at analysis time t is calculated as

$$\phi_g A(\tau - t) S_g(t) G_g(t),$$

where ϕ_g is the probability of randomization to treatment group g , $A(\tau - t)$ is the number of patients enrolled by calendar time $\tau - t$, $S_g(t) G_g(t)$ is the probability of being at risk at analysis time t for a patient in treatment group g after enrollment. Obviously, $t < \min(\tau, T_{\text{fmax}})$.

Value

A matrix of the number of patients at risk at the specified analysis times (row) for each treatment group (column).

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Piecewise accrual, piecewise exponential survivals, and 5% dropout by
# the end of 1 year.
```

```
natrisk(t = c(9, 24), allocationRatioPlanned = 1,
        accrualTime = c(0, 3), accrualIntensity = c(10, 20),
        piecewiseSurvivalTime = c(0, 6),
        lambda1 = c(0.0533, 0.0309), lambda2 = c(0.0533, 0.0533),
        gamma1 = -log(1-0.05)/12, gamma2 = -log(1-0.05)/12,
        accrualDuration = 12, maxFollowupTime = 30, time = 30)
```

nbpower

*Power for Negative Binomial Rate Ratio***Description**

Estimates the power for negative binomial rate ratio test.

Usage

```
nbpower(
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,
  futilityCP = NULL,
  futilityRateRatio = NULL,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  rateRatioH0 = 1,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  kappa1 = NA_real_,
  kappa2 = NA_real_,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = FALSE,
  spendingTime = NA_real_,
  studyDuration = NA_real_,
  nullVariance = FALSE
)
```

Arguments

kMax The maximum number of stages.

informationRates	The information rates. Defaults to $(1:kMax) / kMax$ if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., $kMax-1$. Defaults to $rep(-8, kMax-1)$ if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	A vector of length $kMax - 1$ for the futility bounds on the conditional power scale.
futilityRateRatio	A vector of length $kMax - 1$ for the futility bounds on the rate ratio scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
rateRatioH0	Rate ratio under the null hypothesis.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.

piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
kappa1	The dispersion parameter (reciprocal of the shape parameter of the gamma mixing distribution) for the active treatment group by stratum.
kappa2	The dispersion parameter (reciprocal of the shape parameter of the gamma mixing distribution) for the control group by stratum.
lambda1	The rate parameter of the negative binomial distribution for the active treatment group by stratum.
lambda2	The rate parameter of the negative binomial distribution for the control group by stratum.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.
studyDuration	Study duration for fixed follow-up design. Defaults to missing, which is to be replaced with the sum of accrualDuration and followupTime. If provided, the value is allowed to be less than the sum of accrualDuration and followupTime.
nullVariance	Whether to calculate the variance for log rate ratio under the null hypothesis.

Value

An S3 class nbpower object with 4 components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The overall significance level.
 - numberOfEvents: The total number of events.
 - numberOfDropouts: The total number of dropouts.
 - numbeOfSubjects: The total number of subjects.

- exposure: The total exposure.
- studyDuration: The total study duration.
- information: The maximum information.
- expectedNumberOfEvents: The expected number of events.
- expectedNumberOfDropouts: The expected number of dropouts.
- expectedNumberOfSubjects: The expected number of subjects.
- expectedExposure: The expected exposure.
- expectedStudyDuration: The expected study duration.
- expectedInformation: The expected information.
- accrualDuration: The accrual duration.
- followupTime: The follow-up duration.
- fixedFollowup: Whether a fixed follow-up design is used.
- kMax: The number of stages.
- rateRatioH0: The rate ratio under the null hypothesis.
- rateRatio: The rate ratio.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale.
 - futilityBounds: The futility boundaries on the Z-scale.
 - rejectPerStage: The probability for efficacy stopping.
 - futilityPerStage: The probability for futility stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeFutility: The cumulative probability for futility stopping.
 - cumulativeAlphaSpent: The cumulative alpha spent.
 - numberOfEvents: The number of events.
 - numberOfDropouts: The number of dropouts.
 - numberOfSubjects: The number of subjects.
 - exposure: The exposure.
 - analysisTime: The average time since trial start.
 - efficacyRateRatio: The efficacy boundaries on the rate ratio scale.
 - futilityRateRatio: The futility boundaries on the rate ratio scale.
 - efficacyP: The efficacy boundaries on the p-value scale.
 - futilityP: The futility boundaries on the p-value scale.
 - information: The cumulative information.
 - efficacyStopping: Whether to allow efficacy stopping.
 - futilityStopping: Whether to allow futility stopping.
- settings: A list containing the following input parameters: typeAlphaSpending, parameterAlphaSpending, userAlphaSpending, typeBetaSpending, parameterBetaSpending, allocationRatioPlanned, accrualTime, accrualIntensity, piecewiseSurvivalTime, kappa1, kappa2, lambda1, lambda2, gamma1, gamma2, spendingTime, and nullVariance.
- byTreatmentCounts: A list containing the following counts by treatment group:
 - numberOfEvents1: The number of events by stage for the treatment group.

- numberOfDropouts1: The number of dropouts by stage for the treatment group.
- numberOfSubjects1: The number of subjects by stage for the treatment group.
- exposure1: The exposure by stage for the treatment group.
- numberOfEvents2: The number of events by stage for the control group.
- numberOfDropouts2: The number of dropouts by stage for the control group.
- numberOfSubjects2: The number of subjects by stage for the control group.
- exposure2: The exposure by stage for the control group.
- expectedNumberOfEvents1: The expected number of events for the treatment group.
- expectedNumberOfDropouts1: The expected number of dropouts for the treatment group.
- expectedNumberOfSubjects1: The expected number of subjects for the treatment group.
- expectedExposure1: The expected exposure for the treatment group.
- expectedNumberOfEvents2: The expected number of events for control group.
- expectedNumberOfDropouts2: The expected number of dropouts for the control group.
- expectedNumberOfSubjects2: The expected number of subjects for the control group.
- expectedExposure2: The expected exposure for the control group.

Author(s)

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See Also

[nbstat](#)

Examples

Example 1: Variable follow-up design

```
nbpower(kMax = 2, informationRates = c(0.5, 1),
        alpha = 0.025, typeAlphaSpending = "sfOF",
        accrualIntensity = 1956/1.25,
        stratumFraction = c(0.2, 0.8),
        kappa1 = 5, kappa2 = 5,
        lambda1 = c(0.7*0.125, 0.75*0.25),
        lambda2 = c(0.125, 0.25),
        gamma1 = 0, gamma2 = 0,
        accrualDuration = 1.25,
        followupTime = 2.75, fixedFollowup = FALSE,
        nullVariance = 1)
```

Example 2: Fixed follow-up design

```
nbpower(kMax = 2, informationRates = c(0.5, 1),
        alpha = 0.025, typeAlphaSpending = "sfOF",
        accrualIntensity = 220/1.5,
        kappa1 = 3, kappa2 = 3,
        lambda1 = 0.5*8.4, lambda2 = 8.4,
        gamma1 = 0, gamma2 = 0,
        accrualDuration = 1.5,
```

```
followupTime = 0.5, fixedFollowup = TRUE)
```

nbpower1s

Power for One-Sample Negative Binomial Rate

Description

Estimates the power, stopping probabilities, and expected sample size in a one-group negative binomial design.

Usage

```
nbpower1s(
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,
  futilityCP = NULL,
  futilityRate = NULL,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  lambdaH0 = NA_real_,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  kappa = NA_real_,
  lambda = NA_real_,
  gamma = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = FALSE,
  spendingTime = NA_real_,
  studyDuration = NA_real_
)
```

Arguments

kMax The maximum number of stages.

informationRates	The information rates. Defaults to $(1:kMax) / kMax$ if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., $kMax-1$. Defaults to $rep(-8, kMax-1)$ if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	A vector of length $kMax - 1$ for the futility bounds on the conditional power scale.
futilityRate	A vector of length $kMax - 1$ for the futility bounds on the rate scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
lambdaH0	The rate parameter of the negative binomial distribution under the null hypothesis.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.

<code>piecewiseSurvivalTime</code>	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., <code>c(0, 6)</code> breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
<code>stratumFraction</code>	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
<code>kappa</code>	The dispersion parameter (reciprocal of the shape parameter of the gamma mixing distribution) of the negative binomial distribution by stratum.
<code>lambda</code>	The rate parameter of the negative binomial distribution under the alternative hypothesis by stratum.
<code>gamma</code>	The hazard rate for exponential dropout or a vector of hazard rates for piecewise exponential dropout by stratum. Defaults to 0 for no dropout.
<code>accrualDuration</code>	Duration of the enrollment period.
<code>followupTime</code>	Follow-up time for the last enrolled subject.
<code>fixedFollowup</code>	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
<code>spendingTime</code>	A vector of length <code>kMax</code> for the error spending time at each analysis. Defaults to missing, in which case, it is the same as <code>informationRates</code> .
<code>studyDuration</code>	Study duration for fixed follow-up design. Defaults to missing, which is to be replaced with the sum of <code>accrualDuration</code> and <code>followupTime</code> . If provided, the value is allowed to be less than the sum of <code>accrualDuration</code> and <code>followupTime</code> .

Value

An S3 class `nbpower1s` object with 3 components:

- `overallResults`: A data frame containing the following variables:
 - `overallReject`: The overall rejection probability.
 - `alpha`: The overall significance level.
 - `numberOfEvents`: The total number of events.
 - `numberOfDropouts`: The total number of dropouts.
 - `numbeOfSubjects`: The total number of subjects.
 - `exposure`: The total exposure.
 - `studyDuration`: The total study duration.
 - `information`: The maximum information.
 - `expectedNumberOfEvents`: The expected number of events.
 - `expectedNumberOfDropouts`: The expected number of dropouts.
 - `expectedNumberOfSubjects`: The expected number of subjects.
 - `expectedExposure`: The expected exposure.
 - `expectedStudyDuration`: The expected study duration.
 - `expectedInformation`: The expected information.
 - `accrualDuration`: The accrual duration.

- followupTime: The follow-up duration.
- fixedFollowup: Whether a fixed follow-up design is used.
- kMax: The number of stages.
- lambdaH0: The rate parameter of the negative binomial distribution under the null hypothesis.
- lambda: The overall rate parameter of the negative binomial distribution under the alternative hypothesis.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale.
 - futilityBounds: The futility boundaries on the Z-scale.
 - rejectPerStage: The probability for efficacy stopping.
 - futilityPerStage: The probability for futility stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeFutility: The cumulative probability for futility stopping.
 - cumulativeAlphaSpent: The cumulative alpha spent.
 - numberOfEvents: The number of events.
 - numberOfDropouts: The number of dropouts.
 - numberOfSubjects: The number of subjects.
 - exposure: The exposure.
 - analysisTime: The average time since trial start.
 - efficacyRate: The efficacy boundaries on the rate scale.
 - futilityRate: The futility boundaries on the rate scale.
 - efficacyP: The efficacy boundaries on the p-value scale.
 - futilityP: The futility boundaries on the p-value scale.
 - information: The cumulative information.
 - efficacyStopping: Whether to allow efficacy stopping.
 - futilityStopping: Whether to allow futility stopping.
- settings: A list containing the following input parameters: typeAlphaSpending, parameterAlphaSpending, userAlphaSpending, typeBetaSpending, parameterBetaSpending, accrualTime, accrualIntensity, piecewiseSurvivalTime, stratumFraction, kappa, lambda, gamma, and spendingTime.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

See Also

[nbstat](#)

Examples

```
# Example 1: Variable follow-up design

nbpower1s(kMax = 2, informationRates = c(0.5, 1),
```

```

alpha = 0.025, typeAlphaSpending = "sf0F",
lambdaH0 = 0.125, accrualIntensity = 500,
stratumFraction = c(0.2, 0.8),
kappa = c(3, 5), lambda = c(0.0875, 0.085),
gamma = 0, accrualDuration = 1.25,
followupTime = 2.75, fixedFollowup = FALSE)

# Example 2: Fixed follow-up design

nbpower1s(kMax = 2, informationRates = c(0.5, 1),
alpha = 0.025, typeAlphaSpending = "sf0F",
lambdaH0 = 8.4, accrualIntensity = 40,
kappa = 3, lambda = 0.5*8.4,
gamma = 0, accrualDuration = 1.5,
followupTime = 0.5, fixedFollowup = TRUE)

```

nbpowerequiv

*Power for Equivalence in Negative Binomial Rate Ratio***Description**

Obtains the power for equivalence in negative binomial rate ratio.

Usage

```

nbpowerequiv(
  kMax = 1L,
  informationRates = NA_real_,
  criticalValues = NULL,
  alpha = 0.05,
  typeAlphaSpending = "sf0F",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  rateRatioLower = NA_real_,
  rateRatioUpper = NA_real_,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  kappa1 = NA_real_,
  kappa2 = NA_real_,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  accrualDuration = NA_real_,

```

```

    followupTime = NA_real_,
    fixedFollowup = FALSE,
    spendingTime = NA_real_,
    studyDuration = NA_real_
  )

```

Arguments

kMax	The maximum number of stages.
informationRates	The information rates. Defaults to $(1:kMax) / kMax$ if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level for each of the two one-sided tests. Defaults to 0.05.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsiatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
rateRatioLower	The lower equivalence limit of rate ratio.
rateRatioUpper	The upper equivalence limit of rate ratio.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
kappa1	The dispersion parameter (reciprocal of the shape parameter of the gamma mixing distribution) for the active treatment group by stratum.
kappa2	The dispersion parameter (reciprocal of the shape parameter of the gamma mixing distribution) for the control group by stratum.

lambda1	The rate parameter of the negative binomial distribution for the active treatment group by stratum.
lambda2	The rate parameter of the negative binomial distribution for the control group by stratum.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.
studyDuration	Study duration for fixed follow-up design. Defaults to missing, which is to be replaced with the sum of accrualDuration and followupTime. If provided, the value is allowed to be less than the sum of accrualDuration and followupTime.

Value

An S3 class `nbpowerequiv` object with 4 components:

- `overallResults`: A data frame containing the following variables:
 - `overallReject`: The overall rejection probability.
 - `alpha`: The overall significance level.
 - `numberOfEvents`: The total number of events.
 - `numbeOfSubjects`: The total number of subjects.
 - `exposure`: The total exposure.
 - `studyDuration`: The total study duration.
 - `information`: The maximum information.
 - `expectedNumberOfEvents`: The expected number of events.
 - `expectedNumberOfSubjects`: The expected number of subjects.
 - `expectedExposure`: The expected exposure.
 - `expectedStudyDuration`: The expected study duration.
 - `expectedInformation`: The expected information.
 - `kMax`: The number of stages.
 - `rateRatioLower`: The lower equivalence limit of rate ratio.
 - `rateRatioUpper`: The upper equivalence limit of rate ratio.
 - `rateRatio`: The rate ratio.

- accrualDuration: The accrual duration.
- followupTime: The follow-up duration.
- fixedFollowup: Whether a fixed follow-up design is used.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale for each of the two one-sided tests.
 - rejectPerStage: The probability for efficacy stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeAlphaSpent: The cumulative alpha for each of the two one-sided tests.
 - cumulativeAttainedAlphaH10: The cumulative alpha attained under H10.
 - cumulativeAttainedAlphaH20: The cumulative alpha attained under H20.
 - numberOfEvents: The number of events.
 - numberOfDropouts: The number of dropouts.
 - numberOfSubjects: The number of subjects.
 - exposure: The exposure.
 - analysisTime: The average time since trial start.
 - efficacyRateRatioLower: The efficacy boundaries on the rate ratio scale for the one-sided null hypothesis at the lower equivalence limit.
 - efficacyRateRatioUpper: The efficacy boundaries on the rate ratio scale for the one-sided null hypothesis at the upper equivalence limit.
 - efficacyP: The efficacy bounds on the p-value scale for each of the two one-sided tests.
 - information: The cumulative information.
- settings: A list containing the following input parameters: typeAlphaSpending, parameterAlphaSpending, userAlphaSpending, allocationRatioPlanned, accrualTime, accrualIntensity, piecewiseSurvivalTime, stratumFraction, kappa1, kappa2, lambda1, lambda2, gamma1, gamma2, spendingTime.
- byTreatmentCounts: A list containing the following counts by treatment group:
 - numberOfEvents1: The number of events by stage for the treatment group.
 - numberOfDropouts1: The number of dropouts by stage for the treatment group.
 - numberOfSubjects1: The number of subjects by stage for the treatment group.
 - exposure1: The exposure by stage for the treatment group.
 - numberOfEvents2: The number of events by stage for the control group.
 - numberOfDropouts2: The number of dropouts by stage for the control group.
 - numberOfSubjects2: The number of subjects by stage for the control group.
 - exposure2: The exposure by stage for the control group.
 - expectedNumberOfEvents1: The expected number of events for the treatment group.
 - expectedNumberOfDropouts1: The expected number of dropouts for the treatment group.
 - expectedNumberOfSubjects1: The expected number of subjects for the treatment group.
 - expectedExposure1: The expected exposure for the treatment group.
 - expectedNumberOfEvents2: The expected number of events for control group.
 - expectedNumberOfDropouts2: The expected number of dropouts for the control group.
 - expectedNumberOfSubjects2: The expected number of subjects for the control group.
 - expectedExposure2: The expected exposure for the control group.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

See Also

[nbstat](#)

Examples

```
# Example 1: Variable follow-up design
nbpowerequiv(kMax = 2, informationRates = c(0.5, 1),
  alpha = 0.05, typeAlphaSpending = "sfOF",
  rateRatioLower = 2/3, rateRatioUpper = 3/2,
  accrualIntensity = 1956/1.25,
  kappa1 = 5, kappa2 = 5,
  lambda1 = 0.125, lambda2 = 0.125,
  gamma1 = 0, gamma2 = 0,
  accrualDuration = 1.25,
  followupTime = 2.75, fixedFollowup = FALSE)

# Example 2: Fixed follow-up design
nbpowerequiv(kMax = 2, informationRates = c(0.5, 1),
  alpha = 0.05, typeAlphaSpending = "sfOF",
  rateRatioLower = 0.5, rateRatioUpper = 2,
  accrualIntensity = 220/1.5,
  stratumFraction = c(0.2, 0.8),
  kappa1 = 3, kappa2 = 3,
  lambda1 = c(8.4, 10.2),
  lambda2 = c(8.0, 11.5),
  gamma1 = 0, gamma2 = 0,
  accrualDuration = 1.5,
  followupTime = 0.5, fixedFollowup = TRUE)
```

nbsamplesize

Sample Size for Negative Binomial Rate Ratio

Description

Obtains the needed accrual duration given power and follow-up time, the needed follow-up time given power and accrual duration, or the needed absolute accrual rates given power, accrual duration, follow-up duration, and relative accrual rates in a two-group negative binomial design.

Usage

```
nbsamplesize(
  beta = 0.2,
  kMax = 1L,
  informationRates = NA_real_,
```

```

efficacyStopping = NA_integer_,
futilityStopping = NA_integer_,
criticalValues = NULL,
alpha = 0.025,
typeAlphaSpending = "sfOF",
parameterAlphaSpending = NA_real_,
userAlphaSpending = NA_real_,
futilityBounds = NULL,
futilityCP = NULL,
futilityRateRatio = NULL,
typeBetaSpending = "none",
parameterBetaSpending = NA_real_,
userBetaSpending = NA_real_,
rateRatioH0 = 1,
allocationRatioPlanned = 1,
accrualTime = 0L,
accrualIntensity = NA_real_,
piecewiseSurvivalTime = 0L,
stratumFraction = 1L,
kappa1 = NA_real_,
kappa2 = NA_real_,
lambda1 = NA_real_,
lambda2 = NA_real_,
gamma1 = 0L,
gamma2 = 0L,
accrualDuration = NA_real_,
followupTime = NA_real_,
fixedFollowup = FALSE,
spendingTime = NA_real_,
rounding = TRUE,
nullVariance = FALSE
)

```

Arguments

beta	Type II error. Defaults to 0.2.
kMax	The maximum number of stages.
informationRates	The information rates. Defaults to (1:kMax) / kMax if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.

<code>typeAlphaSpending</code>	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
<code>parameterAlphaSpending</code>	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
<code>userAlphaSpending</code>	The user defined alpha spending. Cumulative alpha spent up to each stage.
<code>futilityBounds</code>	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., $k_{\text{Max}}-1$. Defaults to <code>rep(-8, $k_{\text{Max}}-1$)</code> if left unspecified. The futility bounds are non-binding for the calculation of critical values.
<code>futilityCP</code>	A vector of length $k_{\text{Max}} - 1$ for the futility bounds on the conditional power scale.
<code>futilityRateRatio</code>	A vector of length $k_{\text{Max}} - 1$ for the futility bounds on the rate ratio scale.
<code>typeBetaSpending</code>	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
<code>parameterBetaSpending</code>	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
<code>userBetaSpending</code>	The user defined beta spending. Cumulative beta spent up to each stage.
<code>rateRatioH0</code>	Rate ratio under the null hypothesis.
<code>allocationRatioPlanned</code>	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
<code>accrualTime</code>	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., <code>c(0, 3)</code> breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
<code>accrualIntensity</code>	A vector of accrual intensities. One for each accrual time interval.
<code>piecewiseSurvivalTime</code>	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., <code>c(0, 6)</code> breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
<code>stratumFraction</code>	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.

kappa1	The dispersion parameter (reciprocal of the shape parameter of the gamma mixing distribution) for the active treatment group by stratum.
kappa2	The dispersion parameter (reciprocal of the shape parameter of the gamma mixing distribution) for the control group by stratum.
lambda1	The rate parameter of the negative binomial distribution for the active treatment group by stratum.
lambda2	The rate parameter of the negative binomial distribution for the control group by stratum.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.
nullVariance	Whether to calculate the variance for log rate ratio under the null hypothesis.

Value

A list of two components:

- resultsUnderH1: An S3 class nbpower object under the alternative hypothesis.
- resultsUnderH0: An S3 class nbpower object under the null hypothesis.

Author(s)

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See Also

[nbpower](#)

Examples

```
# Example 1: Obtains follow-up duration given power, accrual intensity,
# and accrual duration for variable follow-up

nbsamplesize(beta = 0.2, kMax = 2,
```

```

        informationRates = c(0.5, 1),
        alpha = 0.025, typeAlphaSpending = "sfOF",
        accrualIntensity = 1956/1.25,
        kappa1 = 5, kappa2 = 5,
        lambda1 = 0.0875, lambda2 = 0.125,
        gamma1 = 0, gamma2 = 0,
        accrualDuration = 1.25,
        followupTime = NA, fixedFollowup = FALSE)

# Example 2: Obtains accrual intensity given power, accrual duration, and
# follow-up duration for variable follow-up

nbsamplesize(beta = 0.2, kMax = 2,
             informationRates = c(0.5, 1),
             alpha = 0.025, typeAlphaSpending = "sfOF",
             accrualIntensity = 100,
             kappa1 = 5, kappa2 = 5,
             lambda1 = 0.0875, lambda2 = 0.125,
             gamma1 = 0, gamma2 = 0,
             accrualDuration = 1.25,
             followupTime = 2.25, fixedFollowup = FALSE)

# Example 3: Obtains accrual duration given power, accrual intensity, and
# follow-up duration for fixed follow-up

nbsamplesize(beta = 0.2, kMax = 2,
             informationRates = c(0.5, 1),
             alpha = 0.025, typeAlphaSpending = "sfOF",
             accrualIntensity = 1667,
             stratumFraction = c(0.2, 0.8),
             kappa1 = 5, kappa2 = 5,
             lambda1 = c(0.7*0.125, 0.75*0.25),
             lambda2 = c(0.125, 0.25),
             gamma1 = 0, gamma2 = 0,
             accrualDuration = NA,
             followupTime = 0.5, fixedFollowup = TRUE)

```

nbsamplesize1s

Sample Size for One-Sample Negative Binomial Rate

Description

Obtains the needed accrual duration given power and follow-up time, the needed follow-up time given power and accrual duration, or the needed absolute accrual rates given power, accrual duration, follow-up duration, and relative accrual rates in a one-group negative binomial design.

Usage

```
nbsamplesize1s(
  beta = 0.2,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,
  futilityCP = NULL,
  futilityRate = NULL,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
  lambdaH0 = NA_real_,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  kappa = NA_real_,
  lambda = NA_real_,
  gamma = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = FALSE,
  spendingTime = NA_real_,
  rounding = TRUE
)
```

Arguments

beta	Type II error. Defaults to 0.2.
kMax	The maximum number of stages.
informationRates	The information rates. Defaults to (1:kMax) / kMax if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.

typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsiatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	A vector of length kMax - 1 for the futility bounds on the conditional power scale.
futilityRate	A vector of length kMax - 1 for the futility bounds on the rate scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
lambdaH0	The rate parameter of the negative binomial distribution under the null hypothesis.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., c(0, 3) breaks the time axis into 2 accrual intervals: [0, 3) and [3, ∞).
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., c(0, 6) breaks the time axis into 2 event intervals: [0, 6) and [6, ∞). Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
kappa	The dispersion parameter (reciprocal of the shape parameter of the gamma mixing distribution) of the negative binomial distribution by stratum.
lambda	The rate parameter of the negative binomial distribution under the alternative hypothesis by stratum.

gamma	The hazard rate for exponential dropout or a vector of hazard rates for piecewise exponential dropout by stratum. Defaults to 0 for no dropout.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.

Value

A list of two components:

- resultsUnderH1: An S3 class nbpower1s object under the alternative hypothesis.
- resultsUnderH0: An S3 class nbpower1s object under the null hypothesis.

Author(s)

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See Also

[nbpower1s](#)

Examples

```
# Example 1: Obtains follow-up duration given power, accrual intensity,
# and accrual duration for variable follow-up
```

```
nbsamplesize1s(beta = 0.2, kMax = 2,
  informationRates = c(0.5, 1),
  alpha = 0.025, typeAlphaSpending = "sfOF",
  lambdaH0 = 0.125, accrualIntensity = 500,
  stratumFraction = c(0.2, 0.8),
  kappa = c(3, 5), lambda = c(0.0875, 0.085),
  gamma = 0, accrualDuration = 1.25,
  followupTime = NA, fixedFollowup = FALSE)
```

```
# Example 2: Obtains accrual intensity given power, accrual duration, and
# follow-up duration for variable follow-up
```

```
nbsamplesize1s(beta = 0.2, kMax = 2,
  informationRates = c(0.5, 1),
  alpha = 0.025, typeAlphaSpending = "sfOF",
  lambdaH0 = 0.125, accrualIntensity = 100,
  kappa = 5, lambda = 0.0875,
  gamma = 0, accrualDuration = 1.25,
  followupTime = 2.25, fixedFollowup = FALSE)
```

```
# Example 3: Obtains accrual duration given power, accrual intensity, and
# follow-up duration for fixed follow-up
```

```
nbsamplesize1s(beta = 0.2, kMax = 2,
               informationRates = c(0.5, 1),
               alpha = 0.025, typeAlphaSpending = "sfOF",
               lambdaH0 = 8.4, accrualIntensity = 40,
               kappa = 3, lambda = 4.2,
               gamma = 0, accrualDuration = NA,
               followupTime = 0.5, fixedFollowup = TRUE)
```

nbsamplesizeequiv

Sample Size for Equivalence in Negative Binomial Rate Ratio

Description

Obtains the sample size for equivalence in negative binomial rate ratio.

Usage

```
nbsamplesizeequiv(
  beta = 0.2,
  kMax = 1L,
  informationRates = NA_real_,
  criticalValues = NULL,
  alpha = 0.05,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  rateRatioLower = NA_real_,
  rateRatioUpper = NA_real_,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  kappa1 = NA_real_,
  kappa2 = NA_real_,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = FALSE,
```

```

    spendingTime = NA_real_,
    rounding = TRUE
)

```

Arguments

beta	The type II error.
kMax	The maximum number of stages.
informationRates	The information rates. Defaults to $(1:kMax) / kMax$ if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level for each of the two one-sided tests. Defaults to 0.05.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
rateRatioLower	The lower equivalence limit of rate ratio.
rateRatioUpper	The upper equivalence limit of rate ratio.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
kappa1	The dispersion parameter (reciprocal of the shape parameter of the gamma mixing distribution) for the active treatment group by stratum.
kappa2	The dispersion parameter (reciprocal of the shape parameter of the gamma mixing distribution) for the control group by stratum.

lambda1	The rate parameter of the negative binomial distribution for the active treatment group by stratum.
lambda2	The rate parameter of the negative binomial distribution for the control group by stratum.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piece-wise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piece-wise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.

Value

An S3 class nbpowerequiv object

Author(s)

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See Also

[nbpowerequiv](#)

Examples

```
# Example 1: Variable follow-up design and solve for follow-up time
nbsamplesizeequiv(beta = 0.1, kMax = 2, informationRates = c(0.5, 1),
  alpha = 0.05, typeAlphaSpending = "sfOF",
  rateRatioLower = 2/3, rateRatioUpper = 3/2,
  accrualIntensity = 1956/1.25,
  stratumFraction = c(0.2, 0.8),
  kappa1 = c(3, 5),
  kappa2 = c(2, 3),
  lambda1 = c(0.125, 0.165),
  lambda2 = c(0.135, 0.175),
  gamma1 = -log(1-0.05),
  gamma2 = -log(1-0.10),
  accrualDuration = 1.25,
  followupTime = NA, fixedFollowup = FALSE)
```

```
# Example 2: Fixed follow-up design and solve for accrual duration
nbsamplesizeequiv(beta = 0.2, kMax = 2, informationRates = c(0.5, 1),
  alpha = 0.05, typeAlphaSpending = "sfOF",
  rateRatioLower = 0.5, rateRatioUpper = 2,
  accrualIntensity = 220/1.5,
  kappa1 = 3, kappa2 = 3,
  lambda1 = 8.4, lambda2 = 8.4,
  gamma1 = 0, gamma2 = 0,
  accrualDuration = NA,
  followupTime = 0.5, fixedFollowup = TRUE)
```

nbstat

Negative Binomial Rate Ratio

Description

Obtains the number of subjects accrued, number of events, number of dropouts, number of subjects reaching the maximum follow-up, total exposure, and variance for log rate in each group, rate ratio, variance, and Wald test statistic of log rate ratio at given calendar times.

Usage

```
nbstat(
  time = NA_real_,
  rateRatioH0 = 1,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  kappa1 = NA_real_,
  kappa2 = NA_real_,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = FALSE,
  nullVariance = FALSE
)
```

Arguments

time	A vector of calendar times for data cut.
rateRatioH0	Rate ratio under the null hypothesis.

allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
kappa1	The dispersion parameter (reciprocal of the shape parameter of the gamma mixing distribution) for the active treatment group by stratum.
kappa2	The dispersion parameter (reciprocal of the shape parameter of the gamma mixing distribution) for the control group by stratum.
lambda1	The rate parameter of the negative binomial distribution for the active treatment group by stratum.
lambda2	The rate parameter of the negative binomial distribution for the control group by stratum.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
nullVariance	Whether to calculate the variance for log rate ratio under the null hypothesis.

Details

The probability mass function for a negative binomial distribution with dispersion parameter κ_i and rate parameter λ_i is given by

$$P(Y_{ij} = y) = \frac{\Gamma(y + 1/\kappa_i)}{\Gamma(1/\kappa_i)y!} \left(\frac{1}{1 + \kappa_i \lambda_i t_{ij}} \right)^{1/\kappa_i} \left(\frac{\kappa_i \lambda_i t_{ij}}{1 + \kappa_i \lambda_i t_{ij}} \right)^y,$$

where Y_{ij} is the event count for subject j in treatment group i , and t_{ij} is the exposure time for the subject. If $\kappa_i = 0$, the negative binomial distribution reduces to the Poisson distribution.

For treatment group i , let $\beta_i = \log(\lambda_i)$. The log-likelihood for $\{(\kappa_i, \beta_i) : i = 1, 2\}$ can be written as

$$l = \sum_{i=1}^2 \sum_{j=1}^{n_i} \{ \log \Gamma(y_{ij} + 1/\kappa_i) - \log \Gamma(1/\kappa_i) + y_{ij}(\log(\kappa_i) + \beta_i) - (y_{ij} + 1/\kappa_i) \log(1 + \kappa_i \exp(\beta_i) t_{ij}) \}.$$

It follows that

$$\frac{\partial l}{\partial \beta_i} = \sum_{j=1}^{n_i} \left\{ y_{ij} - (y_{ij} + 1/\kappa_i) \frac{\kappa_i \exp(\beta_i) t_{ij}}{1 + \kappa_i \exp(\beta_i) t_{ij}} \right\},$$

and

$$-\frac{\partial^2 l}{\partial \beta_i^2} = \sum_{j=1}^{n_i} (y_{ij} + 1/\kappa_i) \frac{\kappa_i \lambda_i t_{ij}}{(1 + \kappa_i \lambda_i t_{ij})^2}.$$

The Fisher information for β_i is

$$E \left(-\frac{\partial^2 l}{\partial \beta_i^2} \right) = n_i E \left(\frac{\lambda_i t_{ij}}{1 + \kappa_i \lambda_i t_{ij}} \right).$$

In addition, we can show that

$$E \left(-\frac{\partial^2 l}{\partial \beta_i \partial \kappa_i} \right) = 0.$$

Therefore, the variance of $\hat{\beta}_i$ is

$$\text{Var}(\hat{\beta}_i) = \frac{1}{n_i} \left\{ E \left(\frac{\lambda_i t_{ij}}{1 + \kappa_i \lambda_i t_{ij}} \right) \right\}^{-1}.$$

To evaluate the integral, we need to obtain the distribution of the exposure time,

$$t_{ij} = \min(\tau - W_{ij}, C_{ij}, T_{fmax}),$$

where τ denotes the calendar time since trial start, W_{ij} denotes the enrollment time for subject j in treatment group i , C_{ij} denotes the time to dropout after enrollment for subject j in treatment group i , and T_{fmax} denotes the maximum follow-up time for all subjects. Therefore,

$$P(t_{ij} \geq t) = P(W_{ij} \leq \tau - t) P(C_{ij} \geq t) I(t \leq T_{fmax}).$$

Let H denote the distribution function of the enrollment time, and G_i denote the survival function of the dropout time for treatment group i . By the change of variables, we have

$$E \left(\frac{\lambda_i t_{ij}}{1 + \kappa_i \lambda_i t_{ij}} \right) = \int_0^{\tau \wedge T_{fmax}} \frac{\lambda_i}{(1 + \kappa_i \lambda_i t)^2} H(\tau - t) G_i(t) dt.$$

A numerical integration algorithm for a univariate function can be used to evaluate the above integral.

For the restricted maximum likelihood (reml) estimate of (β_1, β_2) subject to the constraint that $\beta_1 - \beta_2 = \Delta$, we express the log-likelihood in terms of $(\beta_2, \Delta, \kappa_1, \kappa_2)$, and takes the derivative of the log-likelihood function with respect to β_2 . The resulting score equation has asymptotic limit

$$E \left(\frac{\partial l}{\partial \beta_2} \right) = s_1 + s_2,$$

where

$$s_1 = nrE \left\{ \lambda_1 t_{1j} - \left(\lambda_1 t_{1j} + \frac{1}{\kappa_1} \right) \frac{\kappa_1 e^{\tilde{\beta}_2 + \Delta t_{1j}}}{1 + \kappa_1 e^{\tilde{\beta}_2 + \Delta t_{1j}}} \right\},$$

and

$$s_2 = n(1-r)E \left\{ \lambda_2 t_{2j} - \left(\lambda_2 t_{2j} + \frac{1}{\kappa_2} \right) \frac{\kappa_2 e^{\tilde{\beta}_2 t_{2j}}}{1 + \kappa_2 e^{\tilde{\beta}_2 t_{2j}}} \right\}.$$

Here r is the randomization probability for the active treatment group. The asymptotic limit of the reml of β_2 is the solution $\tilde{\beta}_2$ to $E \left(\frac{\partial l}{\partial \beta_2} \right) = 0$.

Value

A list with two components:

- **resultsUnderH1**: A data frame containing the following variables:
 - **time**: The analysis time since trial start.
 - **subjects**: The number of enrolled subjects.
 - **nevents**: The total number of events.
 - **nevents1**: The number of events in the active treatment group.
 - **nevents2**: The number of events in the control group.
 - **ndropouts**: The total number of dropouts.
 - **ndropouts1**: The number of dropouts in the active treatment group.
 - **ndropouts2**: The number of dropouts in the control group.
 - **nfmax**: The total number of subjects reaching maximum follow-up.
 - **nfmax1**: The number of subjects reaching maximum follow-up in the active treatment group.
 - **nfmax2**: The number of subjects reaching maximum follow-up in the control group.
 - **exposure**: The total exposure time.
 - **exposure1**: The exposure time for the active treatment group.
 - **exposure2**: The exposure time for the control group.
 - **rateRatio**: The rate ratio of the active treatment group versus the control group.
 - **vlogRate1**: The variance for the log rate parameter for the active treatment group.
 - **vlogRate2**: The variance for the log rate parameter for the control group.
 - **vlogRR**: The variance of log rate ratio.
 - **information**: The information of log rate ratio.
 - **zlogRR**: The Z-statistic for log rate ratio.
- **resultsUnderH0** when **nullVariance = TRUE**: A data frame with the following variables:
 - **time**: The analysis time since trial start.
 - **lambda1H0**: The restricted maximum likelihood estimate of the event rate for the active treatment group.
 - **lambda2H0**: The restricted maximum likelihood estimate of the event rate for the control group.
 - **rateRatioH0**: The rate ratio under H0.

- vlogRate1H0: The variance for the log rate parameter for the active treatment group under H0.
- vlogRate2H0: The variance for the log rate parameter for the control group under H0.
- vlogRRH0: The variance of log rate ratio under H0.
- informationH0: The information of log rate ratio under H0.
- zlogRRH0: The Z-statistic for log rate ratio with variance evaluated under H0.
- varianceRatio: The ratio of the variance under H0 versus the variance under H1.
- lambda1: The true event rate for the active treatment group.
- lambda2: The true event rate for the control group.
- rateRatio: The true rate ratio.
- resultsUnderH0 when nullVariance = FALSE: A data frame with the following variables:
 - time: The analysis time since trial start.
 - rateRatioH0: The rate ratio under H0.
 - varianceRatio: Equal to 1.
 - lambda1: The true event rate for the active treatment group.
 - lambda2: The true event rate for the control group.
 - rateRatio: The true rate ratio.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

Example 1: Variable follow-up design

```
nbstat(time = c(1, 1.25, 2, 3, 4),
        accrualIntensity = 1956/1.25,
        kappa1 = 5,
        kappa2 = 5,
        lambda1 = 0.7*0.125,
        lambda2 = 0.125,
        gamma1 = 0,
        gamma2 = 0,
        accrualDuration = 1.25,
        followupTime = 2.75)
```

Example 2: Fixed follow-up design

```
nbstat(time = c(0.5, 1, 1.5, 2),
        accrualIntensity = 220/1.5,
        stratumFraction = c(0.2, 0.8),
        kappa1 = 3,
        kappa2 = 3,
        lambda1 = c(0.5*8.4, 0.6*10.5),
        lambda2 = c(8.4, 10.5),
        gamma1 = 0,
        gamma2 = 0,
```

```
accrualDuration = 1.5,  
followupTime = 0.5,  
fixedFollowup = 1,  
nullVariance = 1)
```

nevent	<i>Number of Subjects Having an Event by Calendar Time</i>
--------	--

Description

Obtains the number of subjects having an event by given calendar times for each treatment group.

Usage

```
nevent(  
  time = NA_real_,  
  allocationRatioPlanned = 1,  
  accrualTime = 0L,  
  accrualIntensity = NA_real_,  
  piecewiseSurvivalTime = 0L,  
  lambda1 = NA_real_,  
  lambda2 = NA_real_,  
  gamma1 = 0L,  
  gamma2 = 0L,  
  accrualDuration = NA_real_,  
  maxFollowupTime = NA_real_  
)
```

Arguments

time	A vector of calendar times at which to calculate the number of patients having an event.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., <code>c(0, 3)</code> breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., <code>c(0, 6)</code> breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
lambda1	A vector of hazard rates for the event for the active treatment group. One for each analysis time interval.

lambda2	A vector of hazard rates for the event for the control group. One for each analysis time interval.
gamma1	The hazard rate for exponential dropout, or a vector of hazard rates for piecewise exponential dropout for the active treatment group.
gamma2	The hazard rate for exponential dropout, or a vector of hazard rates for piecewise exponential dropout for the control group.
accrualDuration	Duration of the enrollment period.
maxFollowupTime	Follow-up time for the first enrolled subject. For fixed follow-up, maxFollowupTime = minFollowupTime. For variable follow-up, maxFollowupTime = accrualDuration + minFollowupTime.

Details

For a given treatment group g and calendar time τ , the number of patients having an event by calendar time τ is calculated as $I_1 + I_2$, where

$$I_1 = \phi_g A(\tau - T_{\text{fmax}}) P_g(T_{\text{fmax}}),$$

and

$$I_2 = \phi_g \int_{\tau - T_{\text{fmax}}}^{\tau} a(u) P_g(\tau - u) du,$$

where ϕ_g is the probability of randomization to treatment group g , $A(\tau - T_{\text{fmax}})$ is the number of patients enrolled by calendar time $\tau - T_{\text{fmax}}$, $P_g(T_{\text{fmax}})$ is the probability of having an event by the maximum follow-up time T_{fmax} for a patient in treatment group g after enrollment, $a(u)$ is the accrual intensity at calendar time u , and $P_g(\tau - u)$ is the probability of having an event by calendar time τ for a patient in treatment group g enrolled at calendar time u .

Value

A matrix of the number of patients having an event at the specified calendar times (row) for each treatment group (column).

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
# Piecewise accrual, piecewise exponential survivals, and 5% dropout by
# the end of 1 year.
nevent(time = c(9, 24), allocationRatioPlanned = 1,
        accrualTime = c(0, 3), accrualIntensity = c(10, 20),
        piecewiseSurvivalTime = c(0, 6),
        lambda1 = c(0.0533, 0.0309), lambda2 = c(0.0533, 0.0533),
        gamma1 = -log(1-0.05)/12, gamma2 = -log(1-0.05)/12,
        accrualDuration = 12, maxFollowupTime = 30)
```

patrisk

Probability of Being at Risk

Description

Obtains the probability of being at risk at given analysis times.

Usage

```
patrisk(time, piecewiseSurvivalTime, lambda, gamma)
```

Arguments

time	A vector of analysis times at which to calculate the probability of being at risk.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
lambda	A vector of hazard rates for the event. One for each analysis time interval.
gamma	The hazard rate for exponential dropout, or a vector of hazard rates for piecewise exponential dropout.

Value

A vector of probabilities of being at risk at the specified analysis times after enrollment for a patient in a treatment group with specified piecewise exponential survival and dropout distributions.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Piecewise exponential survival with hazard 0.0533 in the first 6
# months, and hazard 0.0309 thereafter, and 5% dropout by the end of
# 1 year.

patrisk(time = c(3, 9), piecewiseSurvivalTime = c(0, 6),
        lambda = c(0.0533, 0.0309), gamma = -log(1-0.05)/12)
```

pbvnorm

Distribution Function of the Standard Bivariate Normal

Description

Computes the cumulative distribution function (CDF) of the standard bivariate normal distribution with specified lower and upper integration limits and correlation coefficient.

Usage

```
pbvnorm(lower = NULL, upper = NULL, rho = 0)
```

Arguments

lower	A numeric vector of length 2 specifying the lower limits of integration.
upper	A numeric vector of length 2 specifying the upper limits of integration.
rho	A numeric value specifying the correlation coefficient of the standard bivariate normal distribution.

Details

This function evaluates the probability $P(\text{lower}[1] < X < \text{upper}[1], \text{lower}[2] < Y < \text{upper}[2])$ where (X, Y) follows a standard bivariate normal distribution with correlation `corr`.

Value

A numeric value representing the probability that a standard bivariate normal vector falls within the specified rectangular region.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
pbvnorm(c(-1, -1), c(1, 1), 0.5)
```

pevent

Probability of Having an Event

Description

Obtains the probability of having an event at given analysis times.

Usage

```
pevent(time, piecewiseSurvivalTime, lambda, gamma)
```

Arguments

time	A vector of analysis times at which to calculate the probability of having an event.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., <code>c(0, 6)</code> breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
lambda	A vector of hazard rates for the event. One for each analysis time interval.
gamma	The hazard rate for exponential dropout, or a vector of hazard rates for piecewise exponential dropout.

Value

A vector of probabilities of having an event at the specified analysis times after enrollment for a patient in a treatment group with specified piecewise exponential survival and dropout distributions.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Piecewise exponential survival with hazard 0.0533 in the first 6
# months, and hazard 0.0309 thereafter, and 5% dropout by the end of
# 1 year.

pevent(time = c(3, 9), piecewiseSurvivalTime = c(0, 6),
       lambda = c(0.0533, 0.0309), gamma = -log(1-0.05)/12)
```

phregr*Proportional Hazards Regression Models*

Description

Obtains the hazard ratio estimates from the proportional hazards regression model with right censored or counting process data.

Usage

```
phregr(  
  data,  
  stratum = "",  
  time = "time",  
  time2 = "",  
  event = "event",  
  covariates = "",  
  weight = "",  
  offset = "",  
  id = "",  
  ties = "efron",  
  init = NA_real_,  
  robust = FALSE,  
  est_basehaz = TRUE,  
  est_resid = TRUE,  
  firth = FALSE,  
  plci = FALSE,  
  alpha = 0.05,  
  maxiter = 50,  
  eps = 1e-09  
)
```

Arguments

data	The input data frame that contains the following variables: • stratum: The stratum. • time: The follow-up time for right censored data, or the left end of each interval for counting process data. • time2: The right end of each interval for counting process data. Intervals are assumed to be open on the left and closed on the right, and event indicates whether an event occurred at the right end of each interval. • event: The event indicator, 1=event, 0=no event. • covariates: The values of baseline covariates (and time-dependent covariates in each interval for counting process data). • weight: The weight for each observation. • offset: The offset for each observation.
------	--

	• <code>id</code>: The optional subject ID for counting process data with time-dependent covariates.
<code>stratum</code>	The name(s) of the stratum variable(s) in the input data.
<code>time</code>	The name of the time variable or the left end of each interval for counting process data in the input data.
<code>time2</code>	The name of the right end of each interval for counting process data in the input data.
<code>event</code>	The name of the event variable in the input data.
<code>covariates</code>	The vector of names of baseline and time-dependent covariates in the input data.
<code>weight</code>	The name of the weight variable in the input data.
<code>offset</code>	The name of the offset variable in the input data.
<code>id</code>	The name of the id variable in the input data.
<code>ties</code>	The method for handling ties, either "breslow" or "efron" (default).
<code>init</code>	The vector of initial values. Defaults to zero for all variables.
<code>robust</code>	Whether a robust sandwich variance estimate should be computed. In the presence of the id variable, the score residuals will be aggregated for each id when computing the robust sandwich variance estimate.
<code>est_basehaz</code>	Whether to estimate the baseline hazards. Defaults to TRUE.
<code>est_resid</code>	Whether to estimate the martingale residuals. Defaults to TRUE.
<code>firth</code>	Whether to use Firth's penalized likelihood method. Defaults to FALSE.
<code>plci</code>	Whether to obtain profile likelihood confidence interval.
<code>alpha</code>	The two-sided significance level.
<code>maxiter</code>	The maximum number of iterations.
<code>eps</code>	The tolerance to declare convergence.

Value

A list with the following components:

- `sumstat`: The data frame of summary statistics of model fit with the following variables:
 - `n`: The number of observations.
 - `nevents`: The number of events.
 - `loglik0`: The (penalized) log-likelihood under null.
 - `loglik1`: The maximum (penalized) log-likelihood.
 - `scoretest`: The score test statistic.
 - `niter`: The number of Newton-Raphson iterations.
 - `ties`: The method for handling ties, either "breslow" or "efron".
 - `p`: The number of columns of the Cox model design matrix.
 - `robust`: Whether to use the robust variance estimate.
 - `firth`: Whether to use Firth's penalized likelihood method.
 - `fail`: Whether the model fails to converge.

- loglik0_unpenalized: The unpenalized log-likelihood under null.
- loglik1_unpenalized: The maximum unpenalized log-likelihood.
- parest: The data frame of parameter estimates with the following variables:
 - param: The name of the covariate for the parameter estimate.
 - beta: The log hazard ratio estimate.
 - sebeta: The standard error of log hazard ratio estimate.
 - z: The Wald test statistic for log hazard ratio.
 - expbeta: The hazard ratio estimate.
 - lower: The lower limit of confidence interval.
 - upper: The upper limit of confidence interval.
 - p: The p-value from the chi-square test.
 - method: The method to compute the confidence interval and p-value.
 - sebeta_naive: The naive standard error of log hazard ratio estimate if robust variance is requested.
- basehaz: The data frame of baseline hazards with the following variables (if est_basehaz is TRUE):
 - time: The observed event time.
 - nrisk: The number of patients at risk at the time point.
 - nevent: The number of events at the time point.
 - haz: The baseline hazard at the time point.
 - varhaz: The variance of the baseline hazard at the time point assuming the parameter beta is known.
 - gradhaz: The gradient of the baseline hazard with respect to beta at the time point (in the presence of covariates).
 - stratum: The stratum.
- residuals: The martingale residuals.
- linear_predictors: The vector of linear predictors.
- p: The number of parameters.
- param: The parameter names.
- beta: The parameter estimate.
- vbeta: The covariance matrix for parameter estimates.
- vbeta_naive: The naive covariance matrix for parameter estimates.
- terms: The terms object.
- xlevels: A record of the levels of the factors used in fitting.
- settings: A list containing the input parameter values.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Per K. Anderson and Richard D. Gill. Cox's regression model for counting processes, a large sample study. *Annals of Statistics* 1982; 10:1100-1120.

Terry M. Therneau and Patricia M. Grambsch. *Modeling Survival Data: Extending the Cox Model*. Springer-Verlag, 2000.

Examples

```
library(dplyr)

# Example 1 with right-censored data
(fit1 <- phregr(
  data = rawdata %>% filter(iterationNumber == 1) %>%
    mutate(treat = 1*(treatmentGroup == 1)),
  stratum = "stratum",
  time = "timeUnderObservation", event = "event",
  covariates = "treat", est_basehaz = FALSE, est_resid = FALSE))

# Example 2 with counting process data and robust variance estimate
(fit2 <- phregr(
  data = heart %>% mutate(rx = as.numeric(transplant) - 1),
  time = "start", time2 = "stop", event = "event",
  covariates = c("rx", "age"), id = "id",
  robust = TRUE, est_basehaz = TRUE, est_resid = TRUE))
```

pmvnormr

Multivariate Normal Probabilities over a Hyperrectangle

Description

Computes the probability that a multivariate normal random vector falls within a rectangular region defined by lower and upper bounds.

Usage

```
pmvnormr(
  lower = NULL,
  upper = NULL,
  mean = NULL,
  sigma,
  n0 = 1024,
  n_max = 16384,
  R = 8,
  abseps = 1e-04,
  releps = 0,
  seed = 314159,
  parallel = TRUE,
```

```

    nthreads = 0
  )

```

Arguments

lower	A numeric vector of lower integration limits.
upper	A numeric vector of upper integration limits.
mean	The mean vector. If NULL (default), a zero vector of appropriate length is used.
sigma	The covariance (or correlation) matrix of the distribution.
n0	Initial number of samples per replication for the Monte Carlo integration.
n_max	Maximum number of samples allowed per replication.
R	Number of independent replications used to estimate the error.
abseps	Absolute error tolerance for the probability calculation.
releps	Relative error tolerance for the probability calculation.
seed	Random seed for reproducibility. If 0, a seed is generated from the computer clock.
parallel	Logical; if TRUE, computations are performed in parallel.
nthreads	Number of threads for parallel execution. If 0, the default RcppParallel behavior is used.

Details

The function automatically selects the most efficient computation method based on the input:

- Analytic Methods: Used for univariate cases or multivariate distributions with a compound symmetry correlation structure and non-negative correlations.
- Monte Carlo Estimation: Used for all other cases. The algorithm employs a randomized quasi-Monte Carlo (QMC) approach using generalized Halton sequences.

To improve efficiency and accuracy, the QMC approach incorporates:

- Sequential Conditioning: Mimics the standardization and transformation approach used in `mvtnorm::lpmvnorm`, reducing the J -dimensional integral to a $(J - 1)$ -dimensional problem over a hypercube.
- Adaptive Sampling: The number of samples per replication increases dynamically until the estimated error falls below `abseps` or `releps`, or until `n_max` is reached.

The standard error is derived from R independent replications. For high-dimensional problems, computations can be accelerated by setting `parallel = TRUE`, which distributes the replications across multiple CPU threads via `nthreads`.

Positive semidefinite sigma matrices (including singular cases) are supported in the general covariance branch via minimal diagonal stabilization during factorization.

Value

The estimated probability with the following attributes:

- method: "exact" for analytic methods or "qmc" for the Monte Carlo approach.
- error: The estimated error (half-width of 95% confidence interval).
- nsamples: The total number of samples used across all replications.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Example 1: Compound symmetry covariance structure and analytic method
n <- 5
mean <- rep(0, n)
lower <- rep(-1, n)
upper <- rep(3, n)
sigma <- matrix(0.5, n, n)
diag(sigma) <- 1
pdmnormr(lower, upper, mean, sigma)

# Example 2: General covariance structure and Monte Carlo method
n <- 5
mean <- rep(0, n)
lower <- rep(-1, n)
upper <- rep(3, n)
sigma <- matrix(c(1, 0.5, 0.3, 0.2, 0.1,
                  0.5, 1, 0.4, 0.3, 0.2,
                  0.3, 0.4, 1, 0.5, 0.3,
                  0.2, 0.3, 0.5, 1, 0.4,
                  0.1, 0.2, 0.3, 0.4, 1), nrow = n)
pdmnormr(lower, upper, mean, sigma, seed = 314159, nthreads = 1)
```

Description

Computes lower- or upper-tail probabilities for the r th order statistic of a multivariate normal random vector.

Usage

```
pordmvnormr(
  z,
  r,
  mean = NULL,
  sigma,
  lower.tail = TRUE,
  n0 = 1024,
  n_max = 16384,
  R = 8,
  abseps = 1e-04,
  releps = 0,
  seed = 314159,
  parallel = TRUE,
  nthreads = 0
)
```

Arguments

<code>z</code>	The threshold value.
<code>r</code>	The order statistic index, with $1 \leq r \leq m$.
<code>mean</code>	The mean vector. If <code>NULL</code> (default), a zero vector of appropriate length is used.
<code>sigma</code>	The covariance (or correlation) matrix of the distribution.
<code>lower.tail</code>	Logical; if <code>TRUE</code> (default), probabilities are $P(Z_{(r)} \leq z)$, otherwise $P(Z_{(r)} \geq z)$.
<code>n0</code>	Initial number of samples per replication for the Monte Carlo integration.
<code>n_max</code>	Maximum number of samples allowed per replication.
<code>R</code>	Number of independent replications used to estimate the error.
<code>abseps</code>	Absolute error tolerance for the probability calculation.
<code>releps</code>	Relative error tolerance for the probability calculation.
<code>seed</code>	Random seed for reproducibility. If 0, a seed is generated from the computer clock.
<code>parallel</code>	Logical; if <code>TRUE</code> , computations are performed in parallel.
<code>nthreads</code>	Number of threads for parallel execution. If 0, the default <code>RcppParallel</code> behavior is used.

Details

Let $Z_{(1)} \leq \dots \leq Z_{(m)}$ denote the ordered components of Z . The lower-tail probability is evaluated as $P(Z_{(r)} \leq z) = P(\sum_i 1\{Z_i \leq z\} \geq r)$. The upper-tail probability is evaluated as $P(Z_{(r)} \geq z) = P(\sum_i 1\{Z_i \geq z\} \geq m - r + 1)$.

For compound-symmetry covariance matrices with non-negative correlation, the function uses a one-dimensional conditioning integral. If the component means are equal, the conditional tail is an ordinary binomial tail; otherwise it is a Poisson-binomial tail.

For general covariance matrices, the function uses the equivalent inclusion-exclusion representation over upper-orthant multivariate normal probabilities. This reduces each rectangle probability term to a lower dimensional subvector probability.

Value

The estimated probability with the following attributes:

- **method**: "analytic" for analytic or one-dimensional integration methods, or "qmc" if randomized quasi-Monte Carlo was used for at least one inclusion-exclusion term.
- **error**: The accumulated estimated error from the multivariate normal probability calculations.
- **nsamples**: The total number of samples used across all randomized quasi-Monte Carlo terms.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
n <- 5
mean <- rep(0, n)
sigma <- matrix(0.5, n, n)
diag(sigma) <- 1
pordmvnormr(z = 1, r = 4, mean = mean, sigma = sigma, nthreads = 1)
pordmvnormr(z = 1, r = 4, mean = mean, sigma = sigma,
             lower.tail = FALSE, nthreads = 1)
```

ptpwexp

Distribution Function of Truncated Piecewise Exponential Distribution

Description

Obtains the probability of a truncated piecewise exponential distribution.

Usage

```
ptpwexp(
  q,
  piecewiseSurvivalTime = 0,
  lambda = 0.0578,
  lowerBound = 0,
  lower.tail = TRUE,
  log.p = FALSE
)
```

Arguments

<code>q</code>	The vector of quantiles.
<code>piecewiseSurvivalTime</code>	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., <code>c(0, 6)</code> breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
<code>lambda</code>	A vector of hazard rates for the event. One for each analysis time interval.
<code>lowerBound</code>	The left truncation time point for the survival time. Defaults to 0 for no truncation.
<code>lower.tail</code>	Logical; if TRUE (default), probabilities are $P(X \leq x)$, otherwise, $P(X > x)$.
<code>log.p</code>	Logical; if TRUE, probabilities p are given as $\log(p)$.

Value

The probability p such that $P(X > q \mid X > \text{lowerBound}) = 1 - p$.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
ptpwexp(q = c(8, 18), piecewiseSurvivalTime = c(0, 6, 9, 15),
        lambda = c(0.025, 0.04, 0.015, 0.007))
```

pwexpcuts

Piecewise Exponential Approximation to a Survival Distribution

Description

Obtains the piecewise exponential distribution that approximates a survival distribution.

Usage

```
pwexpcuts(S, ..., tol = 1e-04)
```

Arguments

<code>S</code>	The survival function of a univariate survival time.
<code>...</code>	Additional arguments to be passed to <code>S</code> .
<code>tol</code>	The tolerance for convergence of the profile log-likelihood. Defaults to 0.0001.

Details

This function computes the piecewise exponential approximation to a survival distribution. The piecewise exponential model divides the time axis into J intervals defined by the change points, where each interval $[t_j, t_{j+1})$ has a constant hazard rate λ_j . The time intervals are specified as:

$$[t_1, t_2), [t_2, t_3), \dots, [t_J, t_{J+1})$$

where $t_1 = 0$, $t_{J+1} = \infty$, and $t_j = \tau_{j-1}$ for $j = 2, \dots, J$. The function starts with $J = 2$ (1 change point) and gradually increases J by adding one change point at a time until the maximized profile log-likelihood for change points stabilizes, i.e., the relative increase in the maximum of the profile log-likelihood function is less than `tol`. If the relative change in the hazard rate is also less than `tol`, the function stops and returns the results.

Value

A list with three components:

- `piecewiseSurvivalTime`: A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., `c(0, 6)` breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$.
- `lambda`: A vector of hazard rates for the event. One for each analysis time interval.
- `loglik`: The sequence of the asymptotic limit of the piecewise exponential log-likelihood for an increasing number of change points.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Example 1: Piecewise exponential
pwexpcuts(ptpwexp, piecewiseSurvivalTime = c(0, 3.4, 5.5),
           lambda = c(0.0168, 0.0833, 0.0431), lowerBound = 0,
           lower.tail = FALSE)

# Example 2: Weibull
pwexpcuts(pweibull, shape = 1.37, scale = 1/0.818, lower.tail = FALSE)
```

pwexploglik

Profile Log-Likelihood Function for Change Points in Piecewise Exponential Approximation

Description

Obtains the profile log-likelihood function for change points in the piecewise exponential approximation to a survival function.

Usage

```
pwexploglik(tau, S, ...)
```

Arguments

tau	The numeric vector of change points.
S	The survival function of a univariate survival time.
...	Additional arguments to be passed to S.

Details

This function computes the profile log-likelihood for change points in a piecewise exponential survival model.

Let $S(t)$ denote the survival function of a univariate survival time, and τ be a vector of $J - 1$ change points. The piecewise exponential survival model divides the time axis into J intervals defined by the change points τ , where each interval $[t_j, t_{j+1})$ has a constant hazard rate λ_j . The time intervals are specified as:

$$[t_1, t_2), [t_2, t_3), \dots, [t_J, t_{J+1})$$

where $t_1 = 0$, $t_{J+1} = \infty$, and $t_j = \tau_{j-1}$ for $j = 2, \dots, J$.

For each subject, the expected number of events occurring in the j -th interval is

$$d_j = E\{I(t_j < Y \leq t_{j+1})\} = S(t_j) - S(t_{j+1})$$

The expected exposure in the j -th interval is:

$$e_j = E\{(Y - t_j)I(t_j < Y \leq t_{j+1}) + (t_{j+1} - t_j)I(Y > t_{j+1})\}$$

which can be shown to be equivalent to

$$e_j = \int_{t_j}^{t_{j+1}} S(t) dt$$

The log-likelihood for the piecewise exponential model is:

$$\ell(\tau, \lambda) = \sum_{j=1}^J \{d_j \log(\lambda_j) - e_j \lambda_j\}$$

The profile log-likelihood for τ is obtained by maximizing $\ell(\tau, \lambda)$ with respect to λ for fixed τ . The maximum likelihood estimate of the hazard rate in the j -th interval is

$$\lambda_j = \frac{d_j}{e_j}$$

Substituting back, the profile log-likelihood is

$$\ell(\tau) = \sum_{j=1}^J d_j \log(d_j / e_j) - 1$$

where we use the fact that $\sum_{j=1}^J d_j = 1$.

Value

A list with the following three components:

- `piecewiseSurvivalTime`: A vector that specifies the starting time of piecewise exponential survival time intervals.
- `lambda`: A vector of hazard rates for the event. One for each analysis time interval.
- `loglik`: The value of the profile log-likelihood.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
pwexploglik(tau = c(0.5, 1.2, 2.8), pweibull,
            shape = 1.37, scale = 1/0.818, lower.tail = FALSE)
```

qmvnormr

Equicoordinate Quantiles of the Multivariate Normal Distribution

Description

Computes the equicoordinate quantile q such that $P(X_1 \leq q, X_2 \leq q, \dots, X_k \leq q) = p$ for a multivariate normal random vector X .

Usage

```
qmvnormr(
  p,
  mean = NULL,
  sigma,
  n0 = 1024,
  n_max = 16384,
  R = 8,
  abseps = 1e-04,
  releps = 0,
  seed = 314159,
  parallel = TRUE,
  nthreads = 0
)
```

Arguments

<code>p</code>	The probability level (cumulative probability).
<code>mean</code>	The mean vector. If NULL (default), a zero vector of appropriate length is used.
<code>sigma</code>	The covariance (or correlation) matrix of the distribution.
<code>n0</code>	Initial number of samples per replication for the Monte Carlo integration.
<code>n_max</code>	Maximum number of samples allowed per replication.
<code>R</code>	Number of independent replications used to estimate the error.
<code>abseps</code>	Absolute error tolerance for the probability calculation.
<code>releps</code>	Relative error tolerance for the probability calculation.
<code>seed</code>	Random seed for reproducibility. If 0, a seed is generated from the computer clock.
<code>parallel</code>	Logical; if TRUE, computations are performed in parallel.
<code>nthreads</code>	Number of threads for parallel execution. If 0, the default RcppParallel behavior is used.

Details

This function finds the value q using a root-finding algorithm applied to the `pmvnormr` function. It solves for the value where the multivariate normal cumulative distribution function equals the target probability p .

Positive semidefinite sigma matrices (including singular cases) are supported in the general covariance branch via minimal diagonal stabilization during factorization.

Value

A numeric value representing the calculated equicoordinate quantile.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
n <- 5
mean <- rep(0, n)
sigma <- matrix(0.5, n, n)
diag(sigma) <- 1
qmvnormr(0.5, mean = mean, sigma = sigma, nthreads = 1)
```

qtpwexp

*Quantile Function of Truncated Piecewise Exponential Distribution***Description**

Obtains the quantile of a truncated piecewise exponential distribution.

Usage

```
qtpwexp(
  p,
  piecewiseSurvivalTime = 0,
  lambda = 0.0578,
  lowerBound = 0,
  lower.tail = TRUE,
  log.p = FALSE
)
```

Arguments

p	The vector of probabilities.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
lambda	A vector of hazard rates for the event. One for each analysis time interval.
lowerBound	The left truncation time point for the survival time. Defaults to 0 for no truncation.
lower.tail	Logical; if TRUE (default), probabilities are $P(X \leq x)$, otherwise, $P(X > x)$.
log.p	Logical; if TRUE, probabilities p are given as $\log(p)$.

Value

The quantile q such that $P(X > q \mid X > \text{lowerBound}) = 1 - p$.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
qtpwexp(p = c(0.205, 0.317), piecewiseSurvivalTime = c(0, 6, 9, 15),
  lambda = c(0.025, 0.04, 0.015, 0.007))
```

rawdata	<i>A simulated time-to-event data set with 10 replications</i>
---------	--

Description

A simulated data set with stratification and delayed treatment effect:

iterationNumber The iteration number
 arrivalTime The enrollment time for the subject
 stratum The stratum for the subject
 treatmentGroup The treatment group for the subject
 timeUnderObservation The time under observation since randomization
 event Whether the subject experienced the event
 dropoutEvent Whether the subject dropped out

Usage

```
rawdata
```

Format

An object of class `data.frame` with 4910 rows and 7 columns.

rdsim_multiarm	<i>Risk Difference Simulation for Multi-Arm Multi-Stage Design</i>
----------------	--

Description

Simulate a multi-arm multi-stage design for binary responses using risk-difference Wald statistics with closed testing.

Usage

```
rdsim_multiarm(
  M = 2,
  kMax = 1,
  criticalValues = NA,
  futilityBounds = NULL,
  riskDiffH0s = 0,
  allocations = 1,
  pis = NULL,
  nullVariance = TRUE,
  n = NA,
  plannedSubjects = NA,
```

```

    maxNumberOfIterations = 1000,
    seed = 0,
    nthreads = 0
)

```

Arguments

<code>M</code>	Number of active treatment arms.
<code>kMax</code>	Number of sequential looks.
<code>criticalValues</code>	Numeric matrix of dimension $kMax \times M$ giving the by-look critical values for the closed testing procedure. The first column is used for the level-M test and the last column for the level-1 test.
<code>futilityBounds</code>	Numeric vector of length $kMax - 1$ giving the futility boundaries on the Wald-statistic scale for the first $kMax - 1$ looks. At an interim look, the study stops for futility if all active treatment arms fall below the futility boundary. If omitted, no interim futility stopping is applied.
<code>riskDiffH0s</code>	Scalar or numeric vector of length M . Risk differences under H_0 for each active arm versus the common control. Defaults to 0.
<code>allocations</code>	Integer or integer vector of length $M + 1$. Number of subjects per arm within a randomization block. A single value implies equal allocation; defaults to 1. The first M elements refer to the active arms and the last element refers to the common control.
<code>pis</code>	Numeric vector of length $M + 1$. Each element corresponds to the response rate for a treatment arm. The first M elements refer to the active arms and the last element refers to the common control.
<code>nullVariance</code>	Whether to use the variance under the null or the empirical variance under the alternative.
<code>n</code>	Planned total sample size across all active arms and control.
<code>plannedSubjects</code>	Numeric vector of length $kMax$ giving the planned cumulative sample size at each look for the first active arm and the common control combined.
<code>maxNumberOfIterations</code>	Number of Monte Carlo replications. Defaults to 1000.
<code>seed</code>	Random seed for reproducibility.
<code>nthreads</code>	Number of threads for parallel simulation. Use 0 to accept the default RcppParallel behavior.

Value

An S3 object of class "rdsim_multiarm" with these components:

- overview: A list summarizing trial-level results and settings:
 - overallReject: Overall probability of rejecting the null by trial end.
 - overallFutility: Overall probability of stopping for futility by trial end.
 - rejectPerStage: Probability of rejecting the null for each active arm at each stage.

- `futilityPerStage`: Probability of futility stopping for each active arm at each stage.
- `cumulativeRejection`: Cumulative probability of rejection by stage.
- `cumulativeFutility`: Cumulative futility stopping probability by stage.
- `numberOfEvents`: Cumulative event counts by stage and arm.
- `numberOfSubjects`: Cumulative enrollments by stage and arm.
- `expectedNumberOfEvents`: Expected cumulative events at trial end.
- `expectedNumberOfSubjects`: Expected cumulative enrollments at trial end.
- `criticalValues`: The input matrix of by-level critical values.
- `futilityBounds`: The input futility boundaries for each stage.
- `riskDiffH0s`: The input risk differences under H_0 .
- `nullVariance`: Whether to use variance under H_0 .
- `numberOfIterations`: Number of simulation iterations performed.
- `n`: Planned total sample size.
- `allocations`: The input allocation ratios.
- `responseRates`: The input response rates for each arm.
- `plannedSubjects`: The input planned cumulative sample size at each look for the first active arm and the common control combined.
- `M`: Number of active arms.
- `kMax`: Number of sequential looks.
- `sumdata1`: Data frame summarizing each iteration, stage, and treatment group:
 - `iterationNumber`, `stopStage`, `stageNumber`, `treatmentGroup`, `accruals`, `events`, `phat`.
 - For each stage the final row summarizes the overall study (all arms combined).
- `sumdata2`: Data frame summarizing test statistics by iteration, stage, and active arm:
 - `iterationNumber`, `stopStage`, `stageNumber`, `activeArm`, `totalAccruals`, `totalEvents`, `riskDiff`, `vriskDiff`, `riskDiffZ`, `reject`, `futility`.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(sim1 <- rdsim_multiarm(
  M = 2,
  kMax = 3,
  criticalValues = matrix(c(3.880, 2.747, 2.275,
                           3.710, 2.511, 1.993), 3, 2),
  futilityBounds = c(0.043, 1.194),
  pis = c(0.25, 0.30, 0.20),
  n = 486,
  plannedSubjects = c(146, 292, 324),
  maxNumberOfIterations = 10000,
  seed = 314159,
  nthreads = 1))
```

rdsim_seamless

*Risk Difference Simulation for Phase 2/3 Seamless Design***Description**

Simulate phase 2/3 seamless design testing for risk difference.

Usage

```
rdsim_seamless(
  M = 2,
  K = 1,
  criticalValues = NA,
  futilityBounds = NULL,
  riskDiffH0s = 0,
  allocations = 1,
  pis = NULL,
  nullVariance = TRUE,
  n = NA,
  plannedSubjects = NA,
  maxNumberOfIterations = 1000,
  seed = 0,
  nthreads = 0
)
```

Arguments

- | | |
|----------------|--|
| M | Number of active treatment arms in Phase 2. |
| K | Number of sequential looks in Phase 3. |
| criticalValues | <p>Numeric vector of length $K + 1$ giving the critical value for the Wald statistic at each look (Look 1 through Look $K + 1$). Decision rule:</p> <ul style="list-style-type: none"> • At Look 1, compute the Wald statistic for each active arm versus the common control. If the largest test statistic exceeds the Look 1 critical value, stop for efficacy. • If the Look 1 stopping rule is not met, select the active arm with the largest Wald statistic and continue with that arm only versus control at subsequent looks. • For each look $j = 2, \dots, K + 1$, compare the selected arm to control; if its Wald statistic exceeds the Look j critical value, stop for efficacy; otherwise continue. • If no critical value is exceeded by Look $K + 1$, the procedure ends without rejection. |
| futilityBounds | Numeric vector of length K giving the futility boundaries for Phase 2 and the first $K - 1$ looks in Phase 3. The study stops for futility: |

	• in Phase 2 if the selected treatment arm crosses the phase-2 futility boundary; • in Phase 3 if the selected arm crosses the futility boundary at an interim look; If omitted, no interim futility stopping is applied.
riskDiffH0s	Numeric vector of length M . Risk differences under H_0 for each active arm versus the common control. Defaults to 0 for superiority tests.
allocations	Integer or integer vector of length $M + 1$. Number of subjects per arm within a randomization block. A single value implies equal allocation; defaults to 1. The first M elements refer to the active arms and the last element refers to the common control.
pis	Numeric vector of length $M + 1$. Each element corresponds to the response rate for a treatment arm. The first M elements refer to the active arms and the last element refers to the common control.
nullVariance	Whether to use the variance under the null or the empirical variance under the alternative.
n	Planned total sample size across all active arms and control.
plannedSubjects	Numeric vector of length $K + 1$ giving the planned cumulative sample size at each look. Each entry refers to the combined number of patients for the first active arm and the common control.
maxNumberOfIterations	Number of Monte Carlo replications. Defaults to 1000.
seed	Random seed for reproducibility.
nthreads	Number of threads for parallel simulation. Use 0 to accept the default RcppParallel behavior.

Value

An S3 object of class "rdsim_seamless" with these components:

- overview: A list summarizing trial-level results and settings:
 - selectionProb: Probability of selecting each active arm at the end of phase 2.
 - selectToStage2: Probability of selecting each active arm to enter stage 2.
 - selectAnyToStage2: Probability of selecting any active arm to enter stage 2.
 - rejectPerStage: Probability of rejecting the null for each active arm at each stage.
 - futilityPerStage: Probability of futility stopping for each active arm at each stage.
 - cumulativeRejection: Cumulative probability of rejection by stage.
 - cumulativeFutility: Cumulative futility stopping probabilities by stage.
 - numberOfEvents: Cumulative event counts by stage, including events from all arms in stage 1 and events from the selected arm and control in later stages.
 - numberOfSubjects: Cumulative enrollments by stage. replications that reached that stage.
 - overallReject: Overall probability of rejecting the null by trial end.
 - overallFutility: Overall probability of stopping for futility by trial end.

- expectedNumberOfEvents: Expected cumulative events at trial end.
- expectedNumberOfSubjects: Expected cumulative enrollments at trial end.
- criticalValues: The input critical values for each stage.
- futilityBounds: The input futility boundaries for each stage.
- riskDiffH0s: The input risk differences under H_0 .
- nullVariance: Whether to use variance under H_0 .
- numberOfIterations: Number of simulation iterations performed.
- n: Planned total sample size.
- allocations: The input allocation ratios.
- responseRates: The input response rates for each arm.
- plannedSubjects: The input planned cumulative sample size at each look for the first active arm and the common control combined.
- M: Number of active arms in Phase 2.
- K: Number of sequential looks in Phase 3.
- sumdata1: Data frame summarizing each iteration, stage, and treatment group:
 - iterationNumber, stopStage, stageNumber, treatmentGroup, accruals, events, phat.
 - For each stage the final row summarizes the overall study (all arms combined).
- sumdata2: Data frame summarizing test statistics by iteration, stage, and active arm:
 - iterationNumber, selectedArm, stopStage, stageNumber, activeArm, totalAccruals, totalEvents, riskDiff, vriskDiff, riskDiffZ, reject, futility.
 - For each active arm, total accruals and events refer to the combined counts for that arm and the common control at that stage.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
(sim1 <- rdsim_seamless(
  M = 3,
  K = 1,
  criticalValues = c(8, 2.349),
  futilityBounds = 1.036,
  pis = c(0.22, 0.25, 0.35, 0.20),
  n = 1120,
  plannedSubjects = c(280, 560),
  maxNumberOfIterations = 10000,
  seed = 314159,
  nthreads = 1))
```

remlOddsRatio	<i>REML Estimates of Individual Proportions With Specified Odds Ratio</i>
---------------	---

Description

Obtains the restricted maximum likelihood estimates of individual proportions with specified odds ratio.

Usage

```
remlOddsRatio(n1, y1, n2, y2, oddsRatioH0 = 1)
```

Arguments

n1	The sample size for the active treatment group.
y1	The number of responses for the active treatment group.
n2	The sample size for the control group.
y2	The number of responses for the control group.
oddsRatioH0	The specified odds ratio.

Value

A vector of the restricted maximum likelihood estimates of the response probabilities for the two treatment groups.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
remlOddsRatio(n1 = 10, y1 = 4, n2 = 20, y2 = 2, oddsRatioH0 = 1.25)
```

remlRateDiff	<i>REML Estimates of Individual Rates With Specified Rate Difference</i>
--------------	--

Description

Obtains the restricted maximum likelihood estimates of individual proportions with specified rate difference.

Usage

```
remlRateDiff(t1, y1, t2, y2, rateDiffH0 = 0)
```

Arguments

- t1 The exposure for the active treatment group.
- y1 The number of events for the active treatment group.
- t2 The exposure for the control group.
- y2 The number of events for the control group.
- rateDiffH0 The specified rate difference.

Value

A vector of the restricted maximum likelihood estimates of the incidence rates for the two treatment groups.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
remlRateDiff(t1 = 10, y1 = 4, t2 = 20, y2 = 2, rateDiffH0 = 0.1)
```

remlRateRatio	<i>REML Estimates of Individual Rates With Specified Rate Ratio</i>
---------------	---

Description

Obtains the restricted maximum likelihood estimates of individual proportions with specified rate ratio.

Usage

```
remlRateRatio(t1, y1, t2, y2, rateRatioH0 = 1)
```

Arguments

- t1 The exposure for the active treatment group.
- y1 The number of events for the active treatment group.
- t2 The exposure for the control group.
- y2 The number of events for the control group.
- rateRatioH0 The specified rate ratio.

Value

A vector of the restricted maximum likelihood estimates of the incidence rates for the two treatment groups.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
remlRateRatio(t1 = 10, y1 = 4, t2 = 20, y2 = 2, rateRatioH0 = 1.1)
```

remlRiskDiff	<i>REML Estimates of Individual Proportions With Specified Risk difference</i>
--------------	--

Description

Obtains the restricted maximum likelihood estimates of individual proportions with specified risk difference.

Usage

```
remlRiskDiff(n1, y1, n2, y2, riskDiffH0 = 0)
```

Arguments

n1	The sample size for the active treatment group.
y1	The number of responses for the active treatment group.
n2	The sample size for the control group.
y2	The number of responses for the control group.
riskDiffH0	The specified risk difference.

Value

A vector of the restricted maximum likelihood estimates of the response probabilities for the two treatment groups.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
remlRiskDiff(n1 = 10, y1 = 4, n2 = 20, y2 = 0, riskDiffH0 = 0.1)
```

remlRiskRatio

REML Estimates of Individual Proportions With Specified Risk Ratio

Description

Obtains the restricted maximum likelihood estimates of individual proportions with specified risk ratio.

Usage

```
remlRiskRatio(n1, y1, n2, y2, riskRatioH0 = 1)
```

Arguments

n1	The sample size for the active treatment group.
y1	The number of responses for the active treatment group.
n2	The sample size for the control group.
y2	The number of responses for the control group.
riskRatioH0	The specified risk ratio.

Value

A vector of the restricted maximum likelihood estimates of the response probabilities for the two treatment groups.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
remlRiskRatio(n1 = 10, y1 = 4, n2 = 20, y2 = 2, riskRatioH0 = 1.2)
```

repeatedPValue

Repeated p-Values for Group Sequential Design

Description

Obtains the repeated p-values for a group sequential design.

Usage

```
repeatedPValue(
  kMax,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA,
  maxInformation = 1,
  p,
  information,
  spendingTime = NULL,
  nthreads = 0
)
```

Arguments

kMax	The maximum number of stages.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
maxInformation	The target maximum information. Defaults to 1, in which case, information represents informationRates.
p	The raw p-values at look 1 to look k. It can be a matrix with k columns for $k \leq kMax$.
information	The observed information by look. It can be a matrix with k columns.
spendingTime	The error spending time at each analysis, must be increasing and less than or equal to 1. Defaults to NULL, in which case, it is the same as informationRates derived from information and maxInformation. It can be a matrix with k columns.
nthreads	The number of threads to use in simulations (0 means the default RcppParallel behavior).

Value

The repeated p-values at look 1 to look k.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Example 1: informationRates different from spendingTime
repeatedPValue(kMax = 3, typeAlphaSpending = "sf0F",
               maxInformation = 800,
               p = c(0.2, 0.15, 0.1),
               information = c(529, 700, 800),
               spendingTime = c(0.6271186, 0.8305085, 1))

# Example 2: Maurer & Bretz (2013), current look is not the last look
repeatedPValue(kMax = 3, typeAlphaSpending = "sf0F",
               p = matrix(c(0.0062, 0.017,
                           0.009, 0.13,
                           0.0002, 0.0035,
                           0.002, 0.06),
                           nrow=4, ncol=2),
               information = c(1/3, 2/3),
               nthreads = 1)
```

residuals_liferegr	<i>Residuals for Parametric Regression Models for Failure Time Data</i>
--------------------	---

Description

Obtains the response, martingale, deviance, dfbeta, and likelihood displacement residuals for a parametric regression model for failure time data.

Usage

```
residuals_liferegr(
  object,
  type = c("response", "martingale", "deviance", "dfbeta", "dfbetas", "working",
           "ldcase", "ldresp", "ldshape", "matrix"),
  collapse = FALSE,
  weighted = (type %in% c("dfbeta", "dfbetas"))
)
```

Arguments

object	The output from the phregr call.
type	The type of residuals desired, with options including "response", "martingale", "deviance", "dfbeta", "dfbetas", "working", "ldcase", "ldresp", "ldshape", and "matrix".
collapse	Whether to collapse the residuals by id.
weighted	Whether to compute weighted residuals.

Details

The algorithms follow the `residuals.survreg` function in the `survival` package, except for martingale residuals, which are defined only for event or right-censored data for exponential, weibull, lognormal, and loglogistic distributions.

Value

Either a vector or a matrix of residuals, depending on the specified type:

- response residuals are on the scale of the original data.
- martingale residuals are event indicators minus the cumulative hazards for event or right-censored data.
- working residuals are on the scale of the linear predictor.
- deviance residuals are on the log-likelihood scale.
- `dfbeta` residuals are returned as a matrix, where the i -th row represents the approximate change in the model coefficients resulting from the inclusion of subject i .
- `dfbetas` residuals are similar to `dfbeta` residuals, but each column is scaled by the standard deviation of the corresponding coefficient.
- matrix residuals are a matrix of derivatives of the log-likelihood function. Let L be the log-likelihood, p be the linear predictor ($X\beta$), and s be $\log(\sigma)$. Then the resulting matrix contains six columns: L , $\partial L/\partial p$, $\partial^2 L/\partial p^2$, $\partial L/\partial s$, $\partial^2 L/\partial s^2$, and $\partial L^2/\partial p \partial s$.
- `ldcase` residuals are likelihood displacement for case weight perturbation.
- `ldresp` residuals are likelihood displacement for response value perturbation.
- `ldshape` residuals are likelihood displacement related to the shape parameter.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Escobar, L. A. and Meeker, W. Q. Assessing influence in regression analysis with censored data. *Biometrics* 1992; 48:507-528.

Examples

```
library(dplyr)

fit1 <- liferegr(
  data = tobin %>% mutate(time = ifelse(durable>0, durable, NA)),
  time = "time", time2 = "durable",
  covariates = c("age", "quant"), dist = "normal")

resid <- residuals_liferegr(fit1, type = "response")
head(resid)
```

residuals_phregr

Residuals for Proportional Hazards Regression Models

Description

Obtains the martingale, deviance, score, or Schoenfeld residuals for a proportional hazards regression model.

Usage

```
residuals_phregr(
  object,
  type = c("martingale", "deviance", "score", "schoenfeld", "dfbeta", "dfbetas",
    "scaledsch"),
  collapse = FALSE,
  weighted = (type %in% c("dfbeta", "dfbetas"))
)
```

Arguments

object	The output from the phregr call.
type	The type of residuals desired, with options including "martingale", "deviance", "score", "schoenfeld", "dfbeta", "dfbetas", and "scaledsch".
collapse	Whether to collapse the residuals by id. This is not applicable for Schoenfeld type residuals.
weighted	Whether to compute weighted residuals.

Details

For score and Schoenfeld type residuals, the proportional hazards model must include at least one covariate. The algorithms for deviance, dfbeta, dfbetas, and scaledsch residuals follow the residuals.coxph function in the survival package.

Value

For martingale and deviance residuals, the result is a vector with one element corresponding to each subject (without collapse). For score residuals, the result is a matrix where each row represents a subject and each column corresponds to a variable. The row order aligns with the input data used in the original fit. For Schoenfeld residuals, the result is a matrix with one row for each event and one column per variable. These rows are sorted by time within strata, with the attributes stratum and time included.

Score residuals represent each individual's contribution to the score vector. Two commonly used transformations of this are dfbeta, which represents the approximate change in the coefficient vector if the observation is excluded, and dfbetas, which gives the approximate change in the coefficients scaled by the standard error of the coefficients.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Terry M. Therneau, Patricia M. Grambsch, and Thomas M. Fleming. Martingale based residuals for survival models. *Biometrika* 1990; 77:147-160.

Patricia M. Grambsch and Terry M. Therneau. Proportional hazards tests and diagnostics based on weighted residuals. *Biometrika* 1994; 81:515-26.

Examples

```
library(dplyr)

# Example 1 with right-censored data
fit1 <- phregr(data = rawdata %>% filter(iterationNumber == 1) %>%
  mutate(treat = 1*(treatmentGroup == 1)),
  stratum = "stratum",
  time = "timeUnderObservation", event = "event",
  covariates = "treat")

ressco <- residuals_phregr(fit1, type = "score")
head(ressco)

# Example 2 with counting process data
fit2 <- phregr(data = heart %>% mutate(rx = as.numeric(transplant) - 1),
  time = "start", time2 = "stop", event = "event",
  covariates = c("rx", "age"), id = "id", robust = TRUE)

resssch <- residuals_phregr(fit2, type = "scaledsch")
head(resssch)
```

riskDiffExactCI

Exact Unconditional Confidence Interval for Risk Difference

Description

Obtains the exact unconditional confidence interval for risk difference based on the standardized score statistic.

Usage

```
riskDiffExactCI(
  n1 = NA_integer_,
  y1 = NA_integer_,
  n2 = NA_integer_,
  y2 = NA_integer_,
  cilevel = 0.95
)
```

Arguments

n1	The sample size for the active treatment group.
y1	The number of responses for the active treatment group.
n2	The sample size for the control group.
y2	The number of responses for the control group.
cilevel	The confidence interval level.

Value

A data frame containing the following variables:

- scale: The scale of treatment effect.
- estimate: The point estimate.
- lower: The lower limit of the confidence interval.
- upper: The upper limit of the confidence interval.
- cilevel: The confidence interval level.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
riskDiffExactCI(n1 = 30, y1 = 2, n2 = 30, y2 = 1, cilevel = 0.95)
```

riskDiffExactPValue	<i>P-Value for Exact Unconditional Test of Risk Difference</i>
---------------------	--

Description

Obtains the p-value for exact unconditional test of risk difference.

Usage

```
riskDiffExactPValue(
  n1 = NA_integer_,
  y1 = NA_integer_,
  n2 = NA_integer_,
  y2 = NA_integer_,
  riskDiffH0 = 0,
  directionUpper = TRUE
)
```

Arguments

n1	The sample size for the active treatment group.
y1	The number of responses for the active treatment group.
n2	The sample size for the control group.
y2	The number of responses for the control group.
riskDiffH0	The risk difference under the null hypothesis. Defaults to 0.
directionUpper	Whether larger values represent better responses.

Value

A data frame containing the following variables:

- riskDiffH0: The risk difference under the null hypothesis.
- directionUpper: Whether larger values represent better responses.
- riskDiff: The observed risk difference.
- zstat: The observed value of the Z test statistic.
- pvalue: The one-sided p-value for the unconditional exact test.
- pi2star: The value of pi2 that yields the p-value.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
riskDiffExactPValue(riskDiffH0 = 0, directionUpper = 1,
                     n1 = 68, y1 = 2, n2 = 65, y2 = 1)
```

riskRatioExactCI	<i>Exact Unconditional Confidence Interval for Risk Ratio</i>
------------------	---

Description

Obtains the exact unconditional confidence interval for risk ratio based on the standardized score statistic.

Usage

```
riskRatioExactCI(
  n1 = NA_integer_,
  y1 = NA_integer_,
  n2 = NA_integer_,
  y2 = NA_integer_,
  cilevel = 0.95
)
```

Arguments

n1	The sample size for the active treatment group.
y1	The number of responses for the active treatment group.
n2	The sample size for the control group.
y2	The number of responses for the control group.
cilevel	The confidence interval level.

Value

A data frame containing the following variables:

- scale: The scale of treatment effect.
- estimate: The point estimate.
- lower: The lower limit of the confidence interval.
- upper: The upper limit of the confidence interval.
- cilevel: The confidence interval level.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
riskRatioExactCI(n1 = 30, y1 = 2, n2 = 30, y2 = 1, cilevel = 0.95)
```

riskRatioExactPValue *P-Value for Exact Unconditional Test of Risk Ratio*

Description

Obtains the p-value for exact unconditional test of risk ratio.

Usage

```
riskRatioExactPValue(
  n1 = NA_integer_,
  y1 = NA_integer_,
  n2 = NA_integer_,
  y2 = NA_integer_,
  riskRatioH0 = 1,
  directionUpper = TRUE
)
```

Arguments

n1	The sample size for the active treatment group.
y1	The number of responses for the active treatment group.
n2	The sample size for the control group.
y2	The number of responses for the control group.
riskRatioH0	The risk ratio under the null hypothesis. Defaults to 1.
directionUpper	Whether larger values represent better responses.

Value

A data frame containing the following variables:

- riskRatioH0: The risk ratio under the null hypothesis.
- directionUpper: Whether larger values represent better responses.
- riskRatio: The observed risk ratio.
- zstat: The observed value of the Z test statistic.
- pvalue: The one-sided p-value for the unconditional exact test.
- pi2star: The value of π_2 that yields the p-value.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
riskRatioExactPValue(riskRatioH0 = 1, directionUpper = 1,
                      n1 = 68, y1 = 2, n2 = 65, y2 = 1)
```

rmdiff

Estimate of Restricted Mean Survival Time Difference

Description

Obtains the estimate of restricted mean survival time difference between two treatment groups.

Usage

```
rmdiff(
  data,
  stratum = "",
  treat = "treat",
  time = "time",
  event = "event",
  milestone = 0,
```

```

rmstDiffH0 = 0,
conflev = 0.95,
biascorrection = FALSE
)

```

Arguments

<code>data</code>	The input data frame that contains the following variables: • <code>stratum</code>: The stratum. • <code>treat</code>: The treatment. • <code>time</code>: The possibly right-censored survival time. • <code>event</code>: The event indicator.
<code>stratum</code>	The name of the stratum variable in the input data.
<code>treat</code>	The name of the treatment variable in the input data.
<code>time</code>	The name of the time variable in the input data.
<code>event</code>	The name of the event variable in the input data.
<code>milestone</code>	The milestone time at which to calculate the restricted mean survival time.
<code>rmstDiffH0</code>	The difference in restricted mean survival times under the null hypothesis. Defaults to 0 for superiority test.
<code>conflev</code>	The level of the two-sided confidence interval for the difference in restricted mean survival times. Defaults to 0.95.
<code>biascorrection</code>	Whether to apply bias correction for the variance estimate of individual restricted mean survival times. Defaults to no bias correction.

Value

A data frame with the following variables:

- `milestone`: The milestone time relative to randomization.
- `rmstDiffH0`: The difference in restricted mean survival times under the null hypothesis.
- `rmst1`: The estimated restricted mean survival time for the treatment group.
- `rmst2`: The estimated restricted mean survival time for the control group.
- `rmstDiff`: The estimated difference in restricted mean survival times.
- `vrms1`: The variance for `rmst1`.
- `vrms2`: The variance for `rmst2`.
- `sermsDiff`: The standard error for `rmstDiff`.
- `rmstDiffZ`: The Z-statistic value.
- `rmstDiffPValue`: The two-sided p-value.
- `lower`: The lower bound of confidence interval.
- `upper`: The upper bound of confidence interval.
- `conflev`: The level of confidence interval.
- `biascorrection`: Whether to apply bias correction for the variance estimate of individual restricted mean survival times.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
rmdiff(data = rawdata[rawdata$iterationNumber == 1, ],
       stratum = "stratum", treat = "treatmentGroup",
       time = "timeUnderObservation", event = "event",
       milestone = 12)
```

rmest

Estimate of Restricted Mean Survival Time

Description

Obtains the estimate of restricted means survival time for each stratum.

Usage

```
rmest(
  data,
  stratum = "",
  time = "time",
  event = "event",
  milestone = 0,
  conflev = 0.95,
  biascorrection = FALSE
)
```

Arguments

data	The input data frame that contains the following variables: • stratum: The stratum. • time: The possibly right-censored survival time. • event: The event indicator.
stratum	The name of the stratum variable in the input data.
time	The name of the time variable in the input data.
event	The name of the event variable in the input data.
milestone	The milestone time at which to calculate the restricted mean survival time.
conflev	The level of the two-sided confidence interval for the survival probabilities. Defaults to 0.95.
biascorrection	Whether to apply bias correction for the variance estimate. Defaults to no bias correction.

Value

A data frame with the following variables:

- stratum: The stratum variable.
- size: The number of subjects in the stratum.
- milestone: The milestone time relative to randomization.
- rmst: The estimate of restricted mean survival time.
- stderr: The standard error of the estimated rmst.
- lower: The lower bound of confidence interval if requested.
- upper: The upper bound of confidence interval if requested.
- conflev: The level of confidence interval if requested.
- biascorrection: Whether to apply bias correction for the variance estimate.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
rmmest(data = aml, stratum = "x",
        time = "time", event = "status", milestone = 24)
```

rmpower

Power for Difference in Restricted Mean Survival Times

Description

Estimates the power for testing the difference in restricted mean survival times in a two-sample survival design.

Usage

```
rmpower(
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,
  futilityCP = NULL,
```

```

    futilityRmstDiff = NULL,
    typeBetaSpending = "none",
    parameterBetaSpending = NA_real_,
    milestone = NA_real_,
    rmstDiffH0 = 0,
    allocationRatioPlanned = 1,
    accrualTime = 0L,
    accrualIntensity = NA_real_,
    piecewiseSurvivalTime = 0L,
    stratumFraction = 1L,
    lambda1 = NA_real_,
    lambda2 = NA_real_,
    gamma1 = 0L,
    gamma2 = 0L,
    accrualDuration = NA_real_,
    followupTime = NA_real_,
    fixedFollowup = FALSE,
    spendingTime = NA_real_,
    studyDuration = NA_real_
)

```

Arguments

kMax The maximum number of stages.

informationRates The information rates. Defaults to $(1:kMax) / kMax$ if left unspecified.

efficacyStopping Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.

futilityStopping Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.

criticalValues Upper boundaries on the z-test statistic scale for stopping for efficacy.

alpha The significance level. Defaults to 0.025.

typeAlphaSpending The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".

parameterAlphaSpending The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".

userAlphaSpending The user defined alpha spending. Cumulative alpha spent up to each stage.

futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	A vector of length kMax - 1 for the futility bounds on the conditional power scale.
futilityRmstDiff	A vector of length kMax - 1 for the futility bounds on the restricted mean survival time difference scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
milestone	The milestone time at which to calculate the restricted mean survival time.
rmstDiffH0	The difference in restricted mean survival times under the null hypothesis. Defaults to 0 for superiority test.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., c(0, 3) breaks the time axis into 2 accrual intervals: [0, 3) and [3, ∞).
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., c(0, 6) breaks the time axis into 2 event intervals: [0, 6) and [6, ∞). Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
lambda1	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.
lambda2	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.

accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.
studyDuration	Study duration for fixed follow-up design. Defaults to missing, which is to be replaced with the sum of accrualDuration and followupTime. If provided, the value is allowed to be less than the sum of accrualDuration and followupTime.

Value

An S3 class rmpower object with 4 components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The overall significance level.
 - numberOfEvents: The total number of events.
 - numberOfDropouts: The total number of dropouts.
 - numbeOfSubjects: The total number of subjects.
 - numberOfMilestone: The total number of subjects reaching milestone.
 - studyDuration: The total study duration.
 - information: The maximum information.
 - expectedNumberOfEvents: The expected number of events.
 - expectedNumberOfDropouts: The expected number of dropouts.
 - expectedNumberOfSubjects: The expected number of subjects.
 - expectedNumberOfMilestone: The expected number of subjects reaching milestone.
 - expectedStudyDuration: The expected study duration.
 - expectedInformation: The expected information.
 - accrualDuration: The accrual duration.
 - followupTime: The follow-up duration.
 - fixedFollowup: Whether a fixed follow-up design is used.
 - kMax: The number of stages.
 - milestone: The milestone time relative to randomization.
 - rmstDiffH0: The difference in restricted mean survival times under the null hypothesis.
 - rmst1: The restricted mean survival time for the treatment group.
 - rmst2: The restricted mean survival time for the control group.
 - rmstDiff: The difference in restricted mean survival times, equal to $\text{rmst1} - \text{rmst2}$.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale.

- `futilityBounds`: The futility boundaries on the Z-scale.
- `rejectPerStage`: The probability for efficacy stopping.
- `futilityPerStage`: The probability for futility stopping.
- `cumulativeRejection`: The cumulative probability for efficacy stopping.
- `cumulativeFutility`: The cumulative probability for futility stopping.
- `cumulativeAlphaSpent`: The cumulative alpha spent.
- `numberOfEvents`: The number of events.
- `numberOfDropouts`: The number of dropouts.
- `numberOfSubjects`: The number of subjects.
- `numberOfMilestone`: The number of subjects reaching milestone.
- `analysisTime`: The average time since trial start.
- `efficacyRmstDiff`: The efficacy boundaries on the restricted mean survival time difference scale.
- `futilityRmstDiff`: The futility boundaries on the restricted mean survival time difference scale.
- `efficacyP`: The efficacy boundaries on the p-value scale.
- `futilityP`: The futility boundaries on the p-value scale.
- `information`: The cumulative information.
- `efficacyStopping`: Whether to allow efficacy stopping.
- `futilityStopping`: Whether to allow futility stopping.
- `settings`: A list containing the following input parameters: `typeAlphaSpending`, `parameterAlphaSpending`, `userAlphaSpending`, `typeBetaSpending`, `parameterBetaSpending`, `allocationRatioPlanned`, `accrualTime`, `accrualIntensity`, `piecewiseSurvivalTime`, `stratumFraction`, `lambda1`, `lambda2`, `gamma1`, `gamma2`, and `spendingTime`.
- `byTreatmentCounts`: A list containing the following counts by treatment group:
 - `numberOfEvents1`: The number of events by stage for the treatment group.
 - `numberOfDropouts1`: The number of dropouts by stage for the treatment group.
 - `numberOfSubjects1`: The number of subjects by stage for the treatment group.
 - `numberOfMilestone1`: The number of subjects reaching milestone by stage for the active treatment group.
 - `numberOfEvents2`: The number of events by stage for the control group.
 - `numberOfDropouts2`: The number of dropouts by stage for the control group.
 - `numberOfSubjects2`: The number of subjects by stage for the control group.
 - `numberOfMilestone2`: The number of subjects reaching milestone by stage for the control group.
 - `expectedNumberOfEvents1`: The expected number of events for the treatment group.
 - `expectedNumberOfDropouts1`: The expected number of dropouts for the active treatment group.
 - `expectedNumberOfSubjects1`: The expected number of subjects for the active treatment group.
 - `expectedNumberOfMilestone1`: The expected number of subjects reaching milestone for the active treatment group.
 - `expectedNumberOfEvents2`: The expected number of events for control group.

- expectedNumberOfDropouts2: The expected number of dropouts for the control group.
- expectedNumberOfSubjects2: The expected number of subjects for the control group.
- expectedNumberOfMilestone2: The expected number of subjects reaching milestone for the control group.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Piecewise accrual, piecewise exponential survival, and 5% dropout by
# the end of 1 year.
```

```
rmpower(kMax = 2, informationRates = c(0.8, 1),
        alpha = 0.025, typeAlphaSpending = "sfOF",
        milestone = 18,
        allocationRatioPlanned = 1, accrualTime = seq(0, 8),
        accrualIntensity = 26/9*seq(1, 9),
        piecewiseSurvivalTime = c(0, 6),
        stratumFraction = c(0.2, 0.8),
        lambda1 = c(0.0533, 0.0309, 1.5*0.0533, 1.5*0.0309),
        lambda2 = c(0.0533, 0.0533, 1.5*0.0533, 1.5*0.0533),
        gamma1 = -log(1-0.05)/12,
        gamma2 = -log(1-0.05)/12, accrualDuration = 22,
        followupTime = 18, fixedFollowup = FALSE)
```

rmpower1s

Power for One-Sample Restricted Mean Survival Time

Description

Estimates the power, stopping probabilities, and expected sample size in a one-group survival design.

Usage

```
rmpower1s(
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
```

```

futilityBounds = NULL,
futilityCP = NULL,
futilityRmst = NULL,
typeBetaSpending = "none",
parameterBetaSpending = NA_real_,
milestone = NA_real_,
rmstH0 = NA_real_,
accrualTime = 0L,
accrualIntensity = NA_real_,
piecewiseSurvivalTime = 0L,
stratumFraction = 1L,
lambda = NA_real_,
gamma = 0L,
accrualDuration = NA_real_,
followupTime = NA_real_,
fixedFollowup = FALSE,
spendingTime = NA_real_,
studyDuration = NA_real_
)

```

Arguments

kMax	The maximum number of stages.
informationRates	The information rates. Defaults to $(1:kMax) / kMax$ if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsiatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.

futilityCP	A vector of length $kMax - 1$ for the futility bounds on the conditional power scale.
futilityRmst	A vector of length $kMax - 1$ for the futility bounds on the restricted mean survival time scale.
typeBetaSpending	The type of beta spending. One of the following: "sf0F" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
milestone	The milestone time at which to calculate the restricted mean survival time.
rmsH0	The restricted mean survival time under the null hypothesis.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
lambda	A vector of hazard rates for the event in each analysis time interval by stratum under the alternative hypothesis.
gamma	The hazard rate for exponential dropout or a vector of hazard rates for piecewise exponential dropout. Defaults to 0 for no dropout.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
spendingTime	A vector of length $kMax$ for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.
studyDuration	Study duration for fixed follow-up design. Defaults to missing, which is to be replaced with the sum of accrualDuration and followupTime. If provided, the value is allowed to be less than the sum of accrualDuration and followupTime.

Value

An S3 class rmpower1s object with 3 components:

- overallResults: A data frame containing the following variables:
 - overallReject: The overall rejection probability.
 - alpha: The overall significance level.
 - numberOfEvents: The total number of events.
 - numbeOfSubjects: The total number of subjects.
 - numberOfMilestone: The total number of subjects reaching milestone.
 - studyDuration: The total study duration.
 - information: The maximum information.
 - expectedNumberOfEvents: The expected number of events.
 - expectedNumberOfSubjects: The expected number of subjects.
 - expectedNumberOfMilestone: The expected number of subjects reaching milestone.
 - expectedStudyDuration: The expected study duration.
 - expectedInformation: The expected information.
 - accrualDuration: The accrual duration.
 - followupTime: The follow-up duration.
 - fixedFollowup: Whether a fixed follow-up design is used.
 - kMax: The number of stages.
 - milestone: The milestone time to calculate the restricted mean survival time.
 - rmstH0: The restricted mean survival time under the null hypothesis.
 - rmst: The restricted mean survival time under the alternative hypothesis.
- byStageResults: A data frame containing the following variables:
 - informationRates: The information rates.
 - efficacyBounds: The efficacy boundaries on the Z-scale.
 - futilityBounds: The futility boundaries on the Z-scale.
 - rejectPerStage: The probability for efficacy stopping.
 - futilityPerStage: The probability for futility stopping.
 - cumulativeRejection: The cumulative probability for efficacy stopping.
 - cumulativeFutility: The cumulative probability for futility stopping.
 - cumulativeAlphaSpent: The cumulative alpha spent.
 - numberOfEvents: The number of events.
 - numberOfDropouts: The number of dropouts.
 - numberOfSubjects: The number of subjects.
 - numberOfMilestone: The number of subjects reaching milestone.
 - analysisTime: The average time since trial start.
 - efficacyRmst: The efficacy boundaries on the restricted mean survival time.
 - futilityRmst: The futility boundaries on the restricted mean survival time.
 - efficacyP: The efficacy boundaries on the p-value scale.
 - futilityP: The futility boundaries on the p-value scale.
 - information: The cumulative information.
 - efficacyStopping: Whether to allow efficacy stopping.
 - futilityStopping: Whether to allow futility stopping.
- settings: A list containing the following input parameters: typeAlphaSpending, parameterAlphaSpending, userAlphaSpending, typeBetaSpending, parameterBetaSpending, accrualTime, accuralIntensity, piecewiseSurvivalTime, stratumFraction, lambda, gamma, and spendingTime.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

See Also

[rmstat](#)

Examples

```
rmpower1s(kMax = 2, informationRates = c(0.8, 1),
  alpha = 0.025, typeAlphaSpending = "sf0F",
  milestone = 18, rmstH0 = 10,
  accrualTime = seq(0, 8),
  accrualIntensity = 26/9*seq(1, 9),
  piecewiseSurvivalTime = c(0, 6),
  stratumFraction = c(0.2, 0.8),
  lambda = c(0.0533, 0.0309, 1.5*0.0533, 1.5*0.0309),
  gamma = -log(1-0.05)/12, accrualDuration = 22,
  followupTime = 18, fixedFollowup = FALSE)
```

rmpowerequiv

Power for Equivalence in Restricted Mean Survival Time Difference

Description

Obtains the power for equivalence in restricted mean survival time difference.

Usage

```
rmpowerequiv(
  kMax = 1L,
  informationRates = NA_real_,
  criticalValues = NULL,
  alpha = 0.05,
  typeAlphaSpending = "sf0F",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  milestone = NA_real_,
  rmstDiffLower = NA_real_,
  rmstDiffUpper = NA_real_,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
```

```

gamma1 = 0L,
gamma2 = 0L,
accrualDuration = NA_real_,
followupTime = NA_real_,
fixedFollowup = FALSE,
spendingTime = NA_real_,
studyDuration = NA_real_
)

```

Arguments

kMax	The maximum number of stages.
informationRates	The information rates. Defaults to $(1:kMax) / kMax$ if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level for each of the two one-sided tests. Defaults to 0.05.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
milestone	The milestone time at which to calculate the restricted mean survival time.
rmstDiffLower	The lower equivalence limit of restricted mean survival time difference.
rmstDiffUpper	The upper equivalence limit of restricted mean survival time difference.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., $c(0, 3)$ breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., $c(0, 6)$ breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.

lambda1	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.
lambda2	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piece-wise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piece-wise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.
studyDuration	Study duration for fixed follow-up design. Defaults to missing, which is to be replaced with the sum of accrualDuration and followupTime. If provided, the value is allowed to be less than the sum of accrualDuration and followupTime.

Value

An S3 class `rmpowerequiv` object with 4 components:

- `overallResults`: A data frame containing the following variables:
 - `overallReject`: The overall rejection probability.
 - `alpha`: The overall significance level.
 - `numberOfEvents`: The total number of events.
 - `numberOfSubjects`: The total number of subjects.
 - `studyDuration`: The total study duration.
 - `information`: The maximum information.
 - `expectedNumberOfEvents`: The expected number of events.
 - `expectedNumberOfSubjects`: The expected number of subjects.
 - `expectedStudyDuration`: The expected study duration.
 - `expectedInformation`: The expected information.
 - `kMax`: The number of stages.
 - `milestone`: The milestone time relative to randomization.
 - `rmstDiffLower`: The lower equivalence limit of restricted mean survival time difference.
 - `rmstDiffUpper`: The upper equivalence limit of restricted mean survival time difference.
 - `rmst1`: The restricted mean survival time for the treatment group.
 - `rmst2`: The restricted mean survival time for the control group.

- `rmstDiff`: The restricted mean survival time difference.
- `accrualDuration`: The accrual duration.
- `followupTime`: The follow-up duration.
- `fixedFollowup`: Whether a fixed follow-up design is used.
- `byStageResults`: A data frame containing the following variables:
 - `informationRates`: The information rates.
 - `efficacyBounds`: The efficacy boundaries on the Z-scale for each of the two one-sided tests.
 - `rejectPerStage`: The probability for efficacy stopping.
 - `cumulativeRejection`: The cumulative probability for efficacy stopping.
 - `cumulativeAlphaSpent`: The cumulative alpha for each of the two one-sided tests.
 - `cumulativeAttainedAlphaH10`: The cumulative alpha attained under H_{10} .
 - `cumulativeAttainedAlphaH20`: The cumulative alpha attained under H_{20} .
 - `numberOfEvents`: The number of events.
 - `numberOfDropouts`: The number of dropouts.
 - `numberOfSubjects`: The number of subjects.
 - `numberOfMilestone`: The number of subjects reaching milestone.
 - `analysisTime`: The average time since trial start.
 - `efficacyRmstDiffLower`: The efficacy boundaries on the restricted mean survival time difference scale for the one-sided null hypothesis at the lower equivalence limit.
 - `efficacyRmstDiffUpper`: The efficacy boundaries on the restricted mean survival time difference scale for the one-sided null hypothesis at the upper equivalence limit.
 - `efficacyP`: The efficacy bounds on the p-value scale for each of the two one-sided tests.
 - `information`: The cumulative information.
- `settings`: A list containing the following input parameters: `typeAlphaSpending`, `parameterAlphaSpending`, `userAlphaSpending`, `allocationRatioPlanned`, `accrualTime`, `accrualIntensity`, `piecewiseSurvivalTime`, `stratumFraction`, `lambda1`, `lambda2`, `gamma1`, `gamma2`, and `spendingTime`.
- `byTreatmentCounts`: A list containing the following counts by treatment group:
 - `numberOfEvents1`: The number of events by stage for the treatment group.
 - `numberOfDropouts1`: The number of dropouts by stage for the treatment group.
 - `numberOfSubjects1`: The number of subjects by stage for the treatment group.
 - `numberOfMilestone1`: The number of subjects reaching milestone by stage for the active treatment group.
 - `numberOfEvents2`: The number of events by stage for the control group.
 - `numberOfDropouts2`: The number of dropouts by stage for the control group.
 - `numberOfSubjects2`: The number of subjects by stage for the control group.
 - `numberOfMilestone2`: The number of subjects reaching milestone by stage for the control group.
 - `expectedNumberOfEvents1`: The expected number of events for the treatment group.
 - `expectedNumberOfDropouts1`: The expected number of dropouts for the active treatment group.
 - `expectedNumberOfSubjects1`: The expected number of subjects for the active treatment group.

- expectedNumberOfMilestone1: The expected number of subjects reaching milestone for the active treatment group.
- expectedNumberOfEvents2: The expected number of events for control group.
- expectedNumberOfDropouts2: The expected number of dropouts for the control group.
- expectedNumberOfSubjects2: The expected number of subjects for the control group.
- expectedNumberOfMilestone2: The expected number of subjects reaching milestone for the control group.

Author(s)

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See Also

[rmstat](#)

Examples

```
rmpowerequiv(kMax = 2, informationRates = c(0.5, 1),
  alpha = 0.05, typeAlphaSpending = "sfOF",
  milestone = 18,
  rmstDiffLower = -2, rmstDiffUpper = 2,
  allocationRatioPlanned = 1, accrualTime = seq(0, 8),
  accrualIntensity = 29/9*seq(1, 9),
  piecewiseSurvivalTime = c(0, 6),
  stratumFraction = c(0.2, 0.8),
  lambda1 = c(0.0533, 0.0533, 1.5*0.0533, 1.5*0.0533),
  lambda2 = c(0.0533, 0.0533, 1.5*0.0533, 1.5*0.0533),
  gamma1 = -log(1-0.05)/12,
  gamma2 = -log(1-0.05)/12, accrualDuration = 22,
  followupTime = 18, fixedFollowup = FALSE)
```

rmsamplesize

Sample Size for Difference in Restricted Mean Survival Times

Description

Obtains the needed accrual duration given power, accrual intensity, and follow-up time, the needed follow-up time given power, accrual intensity, and accrual duration, or the needed absolute accrual intensity given power, relative accrual intensity, accrual duration, and follow-up time in a two-group survival design.

Usage

```

rmsamplesize(
  beta = 0.2,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,
  futilityCP = NULL,
  futilityRmstDiff = NULL,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
  milestone = NA_real_,
  rmstDiffH0 = 0,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = FALSE,
  spendingTime = NA_real_,
  rounding = TRUE
)

```

Arguments

beta	Type II error. Defaults to 0.2.
kMax	The maximum number of stages.
informationRates	The information rates. Defaults to $(1:kMax) / kMax$ if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.
futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.

criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	A vector of length kMax - 1 for the futility bounds on the conditional power scale.
futilityRmstDiff	A vector of length kMax - 1 for the futility bounds on the restricted mean survival time difference scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
milestone	The milestone time at which to calculate the restricted mean survival time.
rmstDiffH0	The difference in restricted mean survival times under the null hypothesis. Defaults to 0 for superiority test.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., c(0, 3) breaks the time axis into 2 accrual intervals: [0, 3) and [3, ∞).
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.

<code>piecewiseSurvivalTime</code>	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., <code>c(0, 6)</code> breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
<code>stratumFraction</code>	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
<code>lambda1</code>	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.
<code>lambda2</code>	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
<code>gamma1</code>	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
<code>gamma2</code>	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
<code>accrualDuration</code>	Duration of the enrollment period.
<code>followupTime</code>	Follow-up time for the last enrolled subject.
<code>fixedFollowup</code>	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
<code>spendingTime</code>	A vector of length <code>kMax</code> for the error spending time at each analysis. Defaults to missing, in which case, it is the same as <code>informationRates</code> .
<code>rounding</code>	Whether to round up sample size. Defaults to 1 for sample size rounding.

Value

A list of two components:

- `resultsUnderH1`: An S3 class `rmpower` object under the alternative hypothesis.
- `resultsUnderH0`: An S3 class `rmpower` object under the null hypothesis.

Author(s)

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See Also

[rmpower](#)

Examples

```
# Example 1: Obtains follow-up time given power, accrual intensity,
# and accrual duration for variable follow-up. Of note, the power
# reaches the maximum when the follow-up time equals milestone.
```

```

rmsamplesize(beta = 0.2, kMax = 2, informationRates = c(0.8, 1),
             alpha = 0.025, typeAlphaSpending = "sfOF",
             milestone = 18,
             allocationRatioPlanned = 1, accrualTime = seq(0, 8),
             accrualIntensity = 100/9*seq(1, 9),
             piecewiseSurvivalTime = c(0, 6),
             stratumFraction = c(0.2, 0.8),
             lambda1 = c(0.0533, 0.0309, 1.5*0.0533, 1.5*0.0309),
             lambda2 = c(0.0533, 0.0533, 1.5*0.0533, 1.5*0.0533),
             gamma1 = -log(1-0.05)/12,
             gamma2 = -log(1-0.05)/12, accrualDuration = 22,
             followupTime = NA, fixedFollowup = FALSE)

# Example 2: Obtains accrual intensity given power, accrual duration, and
# follow-up time for variable follow-up

rmsamplesize(beta = 0.2, kMax = 2, informationRates = c(0.8, 1),
             alpha = 0.025, typeAlphaSpending = "sfOF",
             milestone = 18,
             allocationRatioPlanned = 1, accrualTime = seq(0, 8),
             accrualIntensity = 100/9*seq(1, 9),
             piecewiseSurvivalTime = c(0, 6),
             stratumFraction = c(0.2, 0.8),
             lambda1 = c(0.0533, 0.0309, 1.5*0.0533, 1.5*0.0309),
             lambda2 = c(0.0533, 0.0533, 1.5*0.0533, 1.5*0.0533),
             gamma1 = -log(1-0.05)/12,
             gamma2 = -log(1-0.05)/12, accrualDuration = 22,
             followupTime = 18, fixedFollowup = FALSE)

# Example 3: Obtains accrual duration given power, accrual intensity, and
# follow-up time for fixed follow-up

rmsamplesize(beta = 0.2, kMax = 2, informationRates = c(0.8, 1),
             alpha = 0.025, typeAlphaSpending = "sfOF",
             milestone = 18,
             allocationRatioPlanned = 1, accrualTime = seq(0, 8),
             accrualIntensity = 100/9*seq(1, 9),
             piecewiseSurvivalTime = c(0, 6),
             stratumFraction = c(0.2, 0.8),
             lambda1 = c(0.0533, 0.0309, 1.5*0.0533, 1.5*0.0309),
             lambda2 = c(0.0533, 0.0533, 1.5*0.0533, 1.5*0.0533),
             gamma1 = -log(1-0.05)/12,
             gamma2 = -log(1-0.05)/12, accrualDuration = NA,
             followupTime = 18, fixedFollowup = TRUE)

```

Description

Obtains the needed accrual duration given power and follow-up time, the needed follow-up time given power and accrual duration, or the needed absolute accrual rates given power, accrual duration, follow-up duration, and relative accrual rates in a one-group survival design.

Usage

```
rmsamplesize1s(
  beta = 0.2,
  kMax = 1L,
  informationRates = NA_real_,
  efficacyStopping = NA_integer_,
  futilityStopping = NA_integer_,
  criticalValues = NULL,
  alpha = 0.025,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,
  futilityBounds = NULL,
  futilityCP = NULL,
  futilityRmst = NULL,
  typeBetaSpending = "none",
  parameterBetaSpending = NA_real_,
  userBetaSpending = NA_real_,
  milestone = NA_real_,
  rmstH0 = NA_real_,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  lambda = NA_real_,
  gamma = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = FALSE,
  spendingTime = NA_real_,
  rounding = TRUE
)
```

Arguments

beta	Type II error. Defaults to 0.2.
kMax	The maximum number of stages.
informationRates	The information rates. Defaults to (1:kMax) / kMax if left unspecified.
efficacyStopping	Indicators of whether efficacy stopping is allowed at each stage. Defaults to TRUE if left unspecified.

futilityStopping	Indicators of whether futility stopping is allowed at each stage. Defaults to TRUE if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level. Defaults to 0.025.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
futilityBounds	Lower boundaries on the z-test statistic scale for stopping for futility at stages 1, ..., kMax-1. Defaults to rep(-8, kMax-1) if left unspecified. The futility bounds are non-binding for the calculation of critical values.
futilityCP	A vector of length kMax - 1 for the futility bounds on the conditional power scale.
futilityRmst	A vector of length kMax - 1 for the futility bounds on the restricted mean survival time scale.
typeBetaSpending	The type of beta spending. One of the following: "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early futility stopping. Defaults to "none".
parameterBetaSpending	The parameter value for the beta spending. Corresponds to ρ for "sfKD", and γ for "sfHSD".
userBetaSpending	The user defined beta spending. Cumulative beta spent up to each stage.
milestone	The milestone time at which to calculate the restricted survival time.
rmstH0	The restricted mean survival time under the null hypothesis.
accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., c(0, 3) breaks the time axis into 2 accrual intervals: [0, 3) and [3, ∞).
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., c(0, 6) breaks the time axis into 2 event intervals: [0, 6) and [6, ∞). Defaults to 0 for exponential distribution.

stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
lambda	A vector of hazard rates for the event in each analysis time interval by stratum under the alternative hypothesis.
gamma	The hazard rate for exponential dropout or a vector of hazard rates for piecewise exponential dropout. Defaults to 0 for no dropout.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
spendingTime	A vector of length kMax for the error spending time at each analysis. Defaults to missing, in which case, it is the same as informationRates.
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.

Value

A list of two components:

- resultsUnderH1: An S3 class `rmpower1s` object under the alternative hypothesis.
- resultsUnderH0: An S3 class `rmpower1s` object under the null hypothesis.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

See Also

[rmpower1s](#)

Examples

```
# Example 1: Obtains follow-up duration given power, accrual intensity,
# and accrual duration for variable follow-up
```

```
rmsamplesize1s(beta = 0.2, kMax = 2,
  informationRates = c(0.8, 1),
  alpha = 0.025, typeAlphaSpending = "sfOF",
  milestone = 18, rmstH0 = 10,
  accrualTime = seq(0, 8),
  accrualIntensity = 26/9*seq(1, 9),
  piecewiseSurvivalTime = c(0, 6),
  stratumFraction = c(0.2, 0.8),
  lambda = c(0.0533, 0.0309, 1.5*0.0533, 1.5*0.0309),
  gamma = -log(1-0.05)/12, accrualDuration = 22,
  followupTime = NA, fixedFollowup = FALSE)
```

```
# Example 2: Obtains accrual intensity given power, accrual duration, and
# follow-up duration for variable follow-up
```

```

rmsamplesize1s(beta = 0.2, kMax = 2,
  informationRates = c(0.8, 1),
  alpha = 0.025, typeAlphaSpending = "sfOF",
  milestone = 18, rmstH0 = 10,
  accrualTime = seq(0, 8),
  accrualIntensity = 26/9*seq(1, 9),
  piecewiseSurvivalTime = c(0, 6),
  stratumFraction = c(0.2, 0.8),
  lambda = c(0.0533, 0.0309, 1.5*0.0533, 1.5*0.0309),
  gamma = -log(1-0.05)/12, accrualDuration = 22,
  followupTime = 18, fixedFollowup = FALSE)

# Example 3: Obtains accrual duration given power, accrual intensity, and
# follow-up duration for fixed follow-up

rmsamplesize1s(beta = 0.2, kMax = 2,
  informationRates = c(0.8, 1),
  alpha = 0.025, typeAlphaSpending = "sfOF",
  milestone = 18, rmstH0 = 10,
  accrualTime = seq(0, 8),
  accrualIntensity = 26/9*seq(1, 9),
  piecewiseSurvivalTime = c(0, 6),
  stratumFraction = c(0.2, 0.8),
  lambda = c(0.0533, 0.0309, 1.5*0.0533, 1.5*0.0309),
  gamma = -log(1-0.05)/12, accrualDuration = NA,
  followupTime = 18, fixedFollowup = TRUE)

```

rmsamplesizeequiv	<i>Sample Size for Equivalence in Restricted Mean Survival Time Difference</i>
-------------------	--

Description

Obtains the sample size for equivalence in restricted mean survival time difference.

Usage

```

rmsamplesizeequiv(
  beta = 0.2,
  kMax = 1L,
  informationRates = NA_real_,
  criticalValues = NULL,
  alpha = 0.05,
  typeAlphaSpending = "sfOF",
  parameterAlphaSpending = NA_real_,
  userAlphaSpending = NA_real_,

```

```

milestone = NA_real_,
rmstDiffLower = NA_real_,
rmstDiffUpper = NA_real_,
allocationRatioPlanned = 1,
accrualTime = 0L,
accrualIntensity = NA_real_,
piecewiseSurvivalTime = 0L,
stratumFraction = 1L,
lambda1 = NA_real_,
lambda2 = NA_real_,
gamma1 = 0L,
gamma2 = 0L,
accrualDuration = NA_real_,
followupTime = NA_real_,
fixedFollowup = 0L,
spendingTime = NA_real_,
rounding = 1L
)

```

Arguments

beta	The type II error.
kMax	The maximum number of stages.
informationRates	The information rates. Defaults to $(1:kMax) / kMax$ if left unspecified.
criticalValues	Upper boundaries on the z-test statistic scale for stopping for efficacy.
alpha	The significance level for each of the two one-sided tests. Defaults to 0.05.
typeAlphaSpending	The type of alpha spending. One of the following: "OF" for O'Brien-Fleming boundaries, "P" for Pocock boundaries, "WT" for Wang & Tsatis boundaries, "sfOF" for O'Brien-Fleming type spending function, "sfP" for Pocock type spending function, "sfKD" for Kim & DeMets spending function, "sfHSD" for Hwang, Shi & DeCani spending function, "user" for user defined spending, and "none" for no early efficacy stopping. Defaults to "sfOF".
parameterAlphaSpending	The parameter value for the alpha spending. Corresponds to Δ for "WT", ρ for "sfKD", and γ for "sfHSD".
userAlphaSpending	The user defined alpha spending. Cumulative alpha spent up to each stage.
milestone	The milestone time at which to calculate the restricted mean survival time.
rmstDiffLower	The lower equivalence limit of restricted mean survival time difference.
rmstDiffUpper	The upper equivalence limit of restricted mean survival time difference.
allocationRatioPlanned	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.

accrualTime	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., <code>c(0, 3)</code> breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
accrualIntensity	A vector of accrual intensities. One for each accrual time interval.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., <code>c(0, 6)</code> breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
stratumFraction	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
lambda1	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.
lambda2	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piecewise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.
spendingTime	A vector of length <code>kMax</code> for the error spending time at each analysis. Defaults to missing, in which case, it is the same as <code>informationRates</code> .
rounding	Whether to round up sample size. Defaults to 1 for sample size rounding.

Value

An S3 class `rmpowerequiv` object

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

See Also

[rmpowerequiv](#)

Examples

```
rmsamplesizeequiv(beta = 0.1, kMax = 2, informationRates = c(0.5, 1),
  alpha = 0.05, typeAlphaSpending = "sfOF",
  milestone = 18,
  rmstDiffLower = -2, rmstDiffUpper = 2,
  allocationRatioPlanned = 1, accrualTime = seq(0, 8),
  accrualIntensity = 26/9*seq(1, 9),
  piecewiseSurvivalTime = c(0, 6),
  stratumFraction = c(0.2, 0.8),
  lambda1 = c(0.0533, 0.0533, 1.5*0.0533, 1.5*0.0533),
  lambda2 = c(0.0533, 0.0533, 1.5*0.0533, 1.5*0.0533),
  gamma1 = -log(1-0.05)/12,
  gamma2 = -log(1-0.05)/12, accrualDuration = NA,
  followupTime = 18, fixedFollowup = FALSE)
```

rmst	<i>Restricted Mean Survival Time</i>
------	--------------------------------------

Description

Obtains the restricted mean survival time over an interval.

Usage

```
rmst(t1 = 0, t2 = NA_real_, piecewiseSurvivalTime = 0L, lambda = NA_real_)
```

Arguments

- t1 Lower bound of the analysis time interval.
- t2 Upper bound of the analysis time interval.
- piecewiseSurvivalTime A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., c(0, 6) breaks the time axis into 2 event intervals: [0, 6) and [6, ∞). Defaults to 0 for exponential distribution.
- lambda A vector of hazard rates for the event. One for each analysis time interval.

Value

The integral of the survival function from t1 to t2

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
rmst(t1 = 0, t2 = 7, piecewiseSurvivalTime = c(0, 6),
  lambda = c(0.0533, 0.0309))
```

rmstat

*Stratified Difference in Restricted Mean Survival Times***Description**

Obtains the stratified restricted mean survival times and difference in restricted mean survival times at given calendar times.

Usage

```
rmstat(
  time = NA_real_,
  milestone = NA_real_,
  allocationRatioPlanned = 1,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  piecewiseSurvivalTime = 0L,
  stratumFraction = 1L,
  lambda1 = NA_real_,
  lambda2 = NA_real_,
  gamma1 = 0L,
  gamma2 = 0L,
  accrualDuration = NA_real_,
  followupTime = NA_real_,
  fixedFollowup = FALSE
)
```

Arguments

<code>time</code>	A vector of calendar times for data cut.
<code>milestone</code>	The milestone time at which to calculate the restricted mean survival time.
<code>allocationRatioPlanned</code>	Allocation ratio for the active treatment versus control. Defaults to 1 for equal randomization.
<code>accrualTime</code>	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., <code>c(0, 3)</code> breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
<code>accrualIntensity</code>	A vector of accrual intensities. One for each accrual time interval.
<code>piecewiseSurvivalTime</code>	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., <code>c(0, 6)</code> breaks the time axis into 2 event intervals: $[0, 6)$ and $[6, \infty)$. Defaults to 0 for exponential distribution.
<code>stratumFraction</code>	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.

lambda1	A vector of hazard rates for the event in each analysis time interval by stratum for the active treatment group.
lambda2	A vector of hazard rates for the event in each analysis time interval by stratum for the control group.
gamma1	The hazard rate for exponential dropout, a vector of hazard rates for piece-wise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the active treatment group.
gamma2	The hazard rate for exponential dropout, a vector of hazard rates for piece-wise exponential dropout applicable for all strata, or a vector of hazard rates for dropout in each analysis time interval by stratum for the control group.
accrualDuration	Duration of the enrollment period.
followupTime	Follow-up time for the last enrolled subject.
fixedFollowup	Whether a fixed follow-up design is used. Defaults to FALSE for variable follow-up.

Value

A data frame containing the following variables:

- time: The calendar time since trial start.
- subjects: The number of enrolled subjects.
- nevents: The total number of events.
- nevents1: The number of events in the active treatment group.
- nevents2: The number of events in the control group.
- ndropouts: The total number of dropouts.
- ndropouts1: The number of dropouts in the active treatment group.
- ndropouts2: The number of dropouts in the control group.
- milestone: The milestone time relative to randomization.
- nmilestone: The total number of subjects reaching milestone.
- nmilestone1: The number of subjects reaching milestone in the active treatment group.
- nmilestone2: The number of subjects reaching milestone in the control group.
- rmst1: The restricted mean survival time for the treatment group.
- rmst2: The restricted mean survival time for the control group.
- rmstDiff: The difference in restricted mean survival times, i.e., $rmst1 - rmst2$.
- vrmst1: The variance for $rmst1$.
- vrmst2: The variance for $rmst2$.
- vrmstDiff: The variance for $rmstDiff$.
- information: The information for $rmstDiff$, equal to $1/vrmstDiff$.
- rmstDiffZ: The Z-statistic value, i.e., $rmstDiff/\sqrt{vrmstDiff}$.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
# Piecewise accrual, piecewise exponential survivals, and 5% dropout by
# the end of 1 year.
```

```
rmstat(time = c(22, 40),
        milestone = 18,
        allocationRatioPlanned = 1,
        accrualTime = seq(0, 8),
        accrualIntensity = 26/9*seq(1, 9),
        piecewiseSurvivalTime = c(0, 6),
        stratumFraction = c(0.2, 0.8),
        lambda1 = c(0.0533, 0.0309, 1.5*0.0533, 1.5*0.0309),
        lambda2 = c(0.0533, 0.0533, 1.5*0.0533, 1.5*0.0533),
        gamma1 = -log(1-0.05)/12,
        gamma2 = -log(1-0.05)/12,
        accrualDuration = 22,
        followupTime = 18, fixedFollowup = FALSE)
```

rtpwexp

Random Number Generation Function of Truncated Piecewise Exponential Distribution

Description

Obtains random samples from a truncated piecewise exponential distribution.

Usage

```
rtpwexp(n, piecewiseSurvivalTime = 0, lambda = 0.0578, lowerBound = 0)
```

Arguments

n	The number of observations.
piecewiseSurvivalTime	A vector that specifies the starting time of piecewise exponential survival time intervals. Must start with 0, e.g., c(0, 6) breaks the time axis into 2 event intervals: [0, 6) and [6, ∞). Defaults to 0 for exponential distribution.
lambda	A vector of hazard rates for the event. One for each analysis time interval.
lowerBound	The left truncation time point for the survival time. Defaults to 0 for no truncation.

Value

The random numbers from truncated piecewise exponential distribution.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
rtpwexp(n = 10, piecewiseSurvivalTime = c(0, 6, 9, 15),
        lambda = c(0.025, 0.04, 0.015, 0.007))
```

runShinyApp_lrstat	<i>Run Shiny App</i>
--------------------	----------------------

Description

Runs the log-rank test power and sample size calculation Shiny app.

Usage

```
runShinyApp_lrstat()
```

Value

No return value, called for side effects.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

sexagg	<i>Urinary tract infection data from the logistf package</i>
--------	--

Description

This data set deals with urinary tract infection in sexually active college women, along with covariate information on age and contraceptive use. The variables are all binary and coded in 1 (condition is present) and 0 (condition is absent).

Usage

```
sexagg
```

Format

An object of class `data.frame` with 36 rows and 9 columns.

Details

case urinary tract infection, the study outcome variable
 age ≥ 24 years
 dia use of diaphragm
 oc use of oral contraceptive
 vic use of condom
 vicl use of lubricated condom
 vis use of spermicide

shilong

The randomized clinical trial SHIVA data in long format from the ipcswitch package

Description

The original SHIdat data set contains an anonymized excerpt of data from the SHIVA01 trial. This was the first randomized clinical trial that aimed at comparing molecularly targeted therapy based on tumor profiling (MTA) versus conventional therapy (CT) for advanced cancer. Patients were randomly assigned to receive the active or control treatment and may switch to the other arm or subsequent anti-cancer therapy upon disease progression. The restructured data is in the long format.

id The patient's identifier
 tstart The start of the time interval
 tstop The end of the time interval
 event Whether the patient died at the end of the interval
 agerand The patient's age (in years) at randomization
 sex.f The patients' gender, either Male or Female
 tt_Lnum The number of previous lines of treatment
 rmh_alea.c The Royal Marsden Hospital score segregated into two categories
 pathway.f The molecular pathway altered (the hormone receptors pathway, the PI3K/ AKT/mTOR pathway, and the RAF/MEK pathway)
 bras.f The patient's randomized arm, either MTA or CT
 ps The ECOG performance status
 ttc The presence of concomitant treatments
 tran The use of platelet transfusions
 dpd The relative day of a potential progression
 dco The relative day of treatment switching
 ady The relative day of the latest news
 dcut The relative day of administrative cutoff
 pd Whether the patient had disease progression
 co Whether the patient switched treatment

Usage

shilong

Format

An object of class data.frame with 602 rows and 19 columns.

simon2stage	<i>Simon's Two-Stage Design</i>
-------------	---------------------------------

Description

Obtains Simon's two-stage minimax, admissible, and optimal designs.

Usage

```
simon2stage(  
  alpha = NA_real_,  
  beta = NA_real_,  
  piH0 = NA_real_,  
  pi = NA_real_,  
  n_max = 110L  
)
```

Arguments

alpha	Type I error rate (one-sided).
beta	Type II error rate (1-power).
piH0	Response probability under the null hypothesis.
pi	Response probability under the alternative hypothesis.
n_max	Upper limit for sample size, defaults to 110.

Value

A data frame containing the following variables:

- piH0: Response probability under the null hypothesis.
- pi: Response probability under the alternative hypothesis.
- alpha: The specified one-sided significance level.
- beta: The specified type II error.
- n: Total sample size.
- n1: Stage 1 sample size.
- r1: Futility boundary for stage 1.
- r: Futility boundary for stage 2.

- EN_0 : Expected sample size under the null hypothesis.
- attainedAlpha: Attained type 1 error.
- power: Attained power.
- PET_0 : Probability of early stopping under the null hypothesis.
- w_lower: Lower bound of the interval for w.
- w_upper: Upper bound of the interval for w.
- design: Description of the design, e.g., minimax, admissible, or optimal.

Here w is the weight in the objective function: $w \cdot n + (1-w) \cdot EN_0$.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
simon2stage(0.05, 0.2, 0.1, 0.3)
```

simonBayesAnalysis *Analysis of Simon's Bayesian Basket Trials*

Description

Obtains the prior and posterior probabilities for Simon's Bayesian basket discovery trials.

Usage

```
simonBayesAnalysis(
  nstrata = NA_integer_,
  r = NA_real_,
  n = NA_real_,
  lambda = NA_real_,
  gamma = NA_real_,
  phi = NA_real_,
  plo = NA_real_
)
```

Arguments

nstrata	The number of strata.
r	The vector of number of responders across strata.
n	The vector of number of subjects across strata.
lambda	The prior probability that the drug activity is homogeneous across strata.
gamma	The prior probability that the drug is active in a stratum.
phi	The response probability for an active drug.
plo	The response probability for an inactive drug.

Value

A list containing the following five components:

- `case`: The matrix with each row corresponding to a combination of drug activity over strata represented by the columns.
- `prior_case`: The vector of joint prior probabilities for the stratum-specific response rates.
- `prior_stratum`: The vector of marginal prior probabilities for the stratum-specific response rates.
- `post_case`: The vector of joint posterior probabilities for the stratum-specific response rates.
- `post_stratum`: The vector of marginal posterior probabilities for the stratum-specific response rates.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
a <- simonBayesAnalysis(
  nstrata = 10,
  r = c(8,0,1,1,6,2,0,0,3,3),
  n = c(19,10,26,8,14,7,8,5,4,14),
  lambda = 0.5, gamma = 0.33,
  phi = 0.35, plo = 0.15)

a$post_stratum
```

simonBayesSim

Simulation of Simon's Bayesian Basket Trials

Description

Obtains the simulated raw and summary data for Simon's Bayesian basket discovery trials.

Usage

```
simonBayesSim(
  p = NA_real_,
  accrualTime = 0L,
  accrualIntensity = NA_real_,
  stratumFraction = 1L,
  lambda = NA_real_,
  gamma = NA_real_,
  phi = NA_real_,
  plo = NA_real_,
  T = NA_real_,
```

```

    maxSubjects = NA_integer_,
    plannedSubjects = NA_integer_,
    maxNumberOfIterations = 1000L,
    maxNumberOfRawDatasets = 1L,
    seed = 0L
  )

```

Arguments

<code>p</code>	The vector of true response probabilities across strata.
<code>accrualTime</code>	A vector that specifies the starting time of piecewise Poisson enrollment time intervals. Must start with 0, e.g., <code>c(0, 3)</code> breaks the time axis into 2 accrual intervals: $[0, 3)$ and $[3, \infty)$.
<code>accrualIntensity</code>	A vector of accrual intensities. One for each accrual time interval.
<code>stratumFraction</code>	A vector of stratum fractions that sum to 1. Defaults to 1 for no stratification.
<code>lambda</code>	The prior probability that the drug activity is homogeneous across strata.
<code>gamma</code>	The prior probability that the drug is active in a stratum.
<code>phi</code>	The response probability for an active drug.
<code>plo</code>	The response probability for an inactive drug.
<code>T</code>	The threshold for a conclusive posterior probability to stop enrollment.
<code>maxSubjects</code>	The maximum total sample size.
<code>plannedSubjects</code>	The planned cumulative number of subjects at each stage.
<code>maxNumberOfIterations</code>	The number of simulation iterations. Defaults to 1000.
<code>maxNumberOfRawDatasets</code>	The number of raw datasets to extract.
<code>seed</code>	The seed to reproduce the simulation results.

Value

A list containing the following four components:

- `rawdata`: A data frame for subject-level data, containing the following variables:
 - `iterationNumber`: The iteration number.
 - `stageNumber`: The stage number.
 - `subjectId`: The subject ID.
 - `arrivalTime`: The enrollment time for the subject.
 - `stratum`: The stratum for the subject.
 - `y`: Whether the subject was a responder (1) or nonresponder (0).
- `sumdata1`: A data frame for simulation and stratum-level summary data, containing the following variables:

- iterationNumber: The iteration number.
- stageNumber: The stage number.
- stratum: The stratum number.
- active: Whether the drug is active in the stratum.
- n: The number of subjects in the stratum.
- r: The number of responders in the stratum.
- posterior: The posterior probability that the drug is active in the stratum.
- open: Whether the stratum is still open for enrollment.
- positive: Whether the stratum has been determined to be a positive stratum.
- negative: Whether the stratum has been determined to be a negative stratum.
- sumdata2: A data frame for the simulation level summary data, containing the following variables:
 - iterationNumber: The iteration number.
 - numberOfStrata: The total number of strata.
 - n_active_strata: The number of active strata.
 - true_positive: The number of true positive strata.
 - false_negative: The number of false negative strata.
 - false_positive: The number of false positive strata.
 - true_negative: The number of true negative strata.
 - n_indet_strata: The number of indeterminate strata.
 - numberOfSubjects: The number of subjects.
- overview: A data frame for the summary across simulations, containing the following variables:
 - numberOfStrata: The total number of strata.
 - n_active_strata: The average number of active strata.
 - true_positive: The average number of true positive strata.
 - false_negative: The average number of false negative strata.
 - false_positive: The average number of false positive strata.
 - true_negative: The average number of true negative strata.
 - n_indet_strata: The average number of indeterminate strata.
 - numberOfSubjects: The average number of subjects.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
sim1 <- simonBayesSim(
  p = c(0.25, 0.25, 0.05),
  accrualIntensity = 5,
  stratumFraction = c(1/3, 1/3, 1/3),
  lambda = 0.33, gamma = 0.5,
  phi = 0.25, plo = 0.05,
  T = 0.8, maxSubjects = 50,
```

```

plannedSubjects = seq(5, 50, 5),
maxNumberOfIterations = 1000,
maxNumberOfRawDatasets = 1,
seed = 314159)

sim1$overview

```

six

The repeated measures data from the "Six Cities" study of the health effects of air pollution (Ware et al. 1984).

Description

The data analyzed are the 16 selected cases in Lipsitz et al. (1994). The binary response is the wheezing status of 16 children at ages 9, 10, 11, and 12 years. A value of 1 of wheezing status indicates the occurrence of wheezing. The explanatory variables city of residence, age, and maternal smoking status at the particular age.

Usage

```
six
```

Format

An object of class `tbl_df` (inherits from `tbl`, `data.frame`) with 64 rows and 6 columns.

Details

```

case case id
city city of residence
age age of the child
smoke maternal smoking status
wheeze wheezing status

```

survfit_phreg

*Survival Curve for Proportional Hazards Regression Models***Description**

Obtains the predicted survivor function for a proportional hazards regression model.

Usage

```
survfit_phreg(
  object,
  newdata,
  sefit = TRUE,
  conftype = "log-log",
  conflev = 0.95
)
```

Arguments

object	The output from the phreg call.
newdata	A data frame with the same variable names as those that appear in the phreg call. For right-censored data, one curve is produced per row to represent a cohort whose covariates correspond to the values in newdata. For counting-process data, one curve is produced per id in newdata to present the survival curve along the path of time-dependent covariates at the observed event times in the data used to fit phreg.
sefit	Whether to compute the standard error of the survival estimates.
conftype	The type of the confidence interval. One of "none", "plain", "log", "log-log" (the default), or "arcsin". The arcsin option bases the intervals on <code>asin(sqrt(surv))</code> .
conflev	The level of the two-sided confidence interval for the survival probabilities. Defaults to 0.95.

Details

If newdata is not provided and there is no covariate, survival curves based on the basehaz data frame will be produced.

Value

A data frame with the following variables:

- id: The id of the subject for counting-process data with time-dependent covariates.
- time: The observed times in the data used to fit phreg.
- nrisk: The number of patients at risk at the time point in the data used to fit phreg.
- nevent: The number of patients having event at the time point in the data used to fit phreg.

- cumhaz: The cumulative hazard at the time point.
- surv: The estimated survival probability at the time point.
- sesurv: The standard error of the estimated survival probability.
- lower: The lower confidence limit for survival probability.
- upper: The upper confidence limit for survival probability.
- conflev: The level of the two-sided confidence interval.
- conftype: The type of the confidence interval.
- covariates: The values of covariates based on newdata.
- stratum: The stratum of the subject.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Terry M. Therneau and Patricia M. Grambsch. Modeling Survival Data: Extending the Cox Model. Springer-Verlag, 2000.

Examples

```
library(dplyr)

# Example 1 with right-censored data
fit1 <- phregr(data = rawdata %>% filter(iterationNumber == 1) %>%
  mutate(treat = 1*(treatmentGroup == 1)),
  stratum = "stratum",
  time = "timeUnderObservation", event = "event",
  covariates = "treat")

surv1 <- survfit_phregr(fit1,
  newdata = data.frame(
    stratum = as.integer(c(1,1,2,2)),
    treat = c(1,0,1,0)))

head(surv1)

# Example 2 with counting process data and robust variance estimate
fit2 <- phregr(data = heart %>% mutate(rx = as.numeric(transplant) - 1),
  time = "start", time2 = "stop", event = "event",
  covariates = c("rx", "age"), id = "id", robust = TRUE)

surv2 <- survfit_phregr(fit2,
  newdata = data.frame(
    id = c(4,4,11,11),
    age = c(-7.737,-7.737,-0.019,-0.019),
    start = c(0,36,0,26),
    stop = c(36,39,26,153),
    rx = c(0,1,0,1)))

head(surv2)
```

survQuantile	<i>Brookmeyer-Crowley Confidence Interval for Quantiles of Right-Censored Time-to-Event Data</i>
--------------	--

Description

Obtains the Brookmeyer-Crowley confidence interval for quantiles of right-censored time-to-event data.

Usage

```
survQuantile(
  time,
  event,
  cilevel = 0.95,
  transform = "loglog",
  probs = as.numeric(c(0.25, 0.5, 0.75))
)
```

Arguments

time	The vector of possibly right-censored survival times.
event	The vector of event indicators.
cilevel	The confidence interval level. Defaults to 0.95.
transform	The transformation of the survival function to use to construct the confidence interval. Options include "linear" (alternatively "plain"), "log", "loglog" (alternatively "log-log" or "cloglog"), "asinsqrt" (alternatively "asin" or "arcsin"), and "logit". Defaults to "loglog".
probs	The vector of probabilities to calculate the quantiles. Defaults to c(0.25, 0.5, 0.75).

Value

A data frame containing the estimated quantile and confidence interval corresponding to each specified probability. It includes the following variables:

- prob: The probability to calculate the quantile.
- quantile: The estimated quantile.
- lower: The lower limit of the confidence interval.
- upper: The upper limit of the confidence interval.
- cilevel: The confidence interval level.
- transform: The transformation of the survival function to use to construct the confidence interval.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
survQuantile(
  time = c(33.7, 3.9, 10.5, 5.4, 19.5, 23.8, 7.9, 16.9, 16.6,
           33.7, 17.1, 7.9, 10.5, 38),
  event = c(0, 1, 1, 1, 1, 0, 1, 0, 0, 0, 0, 0, 1, 1),
  probs = c(0.25, 0.5, 0.75))
```

svdcpp

*Singular Value Decomposition of a Matrix***Description**

Computes the singular-value decomposition of a rectangular matrix.

Usage

```
svdcpp(X, outtransform = TRUE, decreasing = TRUE)
```

Arguments

<code>X</code>	A numeric matrix whose SVD decomposition is to be computed.
<code>outtransform</code>	Whether the orthogonal matrices composing of the left and right singular vectors are to be computed.
<code>decreasing</code>	Whether the singular values should be sorted in decreasing order and the corresponding singular vectors rearranged accordingly.

Details

Given $A \in R^{m \times n} (m \geq n)$, the following algorithm overwrites A with $U^T A V = D$, where $U \in R^{m \times m}$ is orthogonal, $V \in R^{n \times n}$ is orthogonal, and $D \in R^{m \times n}$ is diagonal.

Value

A list with the following components:

- `d`: A vector containing the singular values of X .
- `U`: A matrix whose columns contain the left singular vectors of X .
- `V`: A matrix whose columns contain the right singular vectors of X .

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

References

Gene N. Golub and Charles F. Van Loan. Matrix Computations, second edition. Baltimore, Maryland: The John Hopkins University Press, 1989, p.434.

Examples

```
A <- matrix(c(1,0,0,0, 1,2,0,0, 0,1,3,0, 0,0,1,4), 4, 4)
svdcp(A)
```

tobin	<i>Tobin's tobit data from the survival package</i>
-------	---

Description

Data from Tobin's original paper.
durable Durable goods purchase
age Age in years
quant Liquidity ratio (x 1000)

Usage

```
tobin
```

Format

An object of class `data.frame` with 20 rows and 3 columns.

updateGraph	<i>Update Graph for Graphical Approaches</i>
-------------	--

Description

Updates the weights and transition matrix for graphical approaches.

Usage

```
updateGraph(w, G, I, j)
```

Arguments

- | | |
|---|--|
| w | The current vector of weights for elementary hypotheses. |
| G | The current transition matrix. |
| I | The set of indices for yet to be rejected hypotheses. |
| j | The hypothesis to remove from index set I. |

Value

A list containing the new vector of weights, the new transition matrix for the graph, and the new set of indices of yet to be rejected hypotheses.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
updateGraph(w = c(0.5, 0.5, 0, 0),
            G = matrix(c(0, 0.5, 0.5, 0, 0.5, 0, 0, 0.5,
                        0, 1, 0, 0, 1, 0, 0, 0),
                      nrow=4, ncol=4, byrow=TRUE),
            I = c(1, 2, 3, 4),
            j = 1)
```

zph_phregr	<i>Assess Proportional Hazards Assumption Based on Scaled Schoenfeld Residuals</i>
------------	--

Description

Obtains the scaled Schoenfeld residuals and tests the proportional hazards assumption using a score test for the interaction between each covariate and a transformed time variable.

Usage

```
zph_phregr(object, transform = "km")
```

Arguments

object	The output from the phregr call.
transform	A character string indicating how survival times should be transformed before the test is performed. Supported values include "identity", "log", "rank", and "km" (default).

Details

This corresponds to the `cox.zph` function from the `survival` package with `terms = FALSE` and `global = TRUE`.

Value

- A list with the following components:
- table A matrix with one row for each parameter and a final row for the global test. The columns contain the score test for adding the time-dependent term, the degrees of freedom, and the two-sided p-value.
 - x The transformed time values.
 - time The original (untransformed) event times, with tied event times repeated.
 - strata The stratum index for each event.
 - y The matrix of scaled Schoenfeld residuals, with one column for each parameter and one row for each event. Column names correspond to the parameter names.
 - var An approximate covariance matrix of the scaled Schoenfeld residuals, used to construct an approximate standard error band for plots.
 - transform the transformation applied to the time values.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

References

Patricia M. Grambsch and Terry M. Therneau. Proportional hazards tests and diagnostics based on weighted residuals. *Biometrika* 1994; 81:515-26.

Examples

```
fit <- phregr(data = liver, time = "Time", event = "Status",
             covariates = c("log(Bilirubin)", "log(Prottime)",
                           "log(Albumin)", "Age", "Edema"),
             ties = "breslow")

zph <- zph_phregr(fit, transform = "log")

zph$table
```

zstatOddsRatio	<i>Miettinen-Nurminen Score Test Statistic for Two-Sample Odds Ratio</i>
----------------	--

Description

Obtains the Miettinen-Nurminen score test statistic for two-sample odds ratio possibly with stratification.

Usage

```
zstatOddsRatio(n1, y1, n2, y2, oddsRatioH0 = 1)
```

Arguments

n1	The sample size for the active treatment group.
y1	The number of responses for the active treatment group.
n2	The sample size for the control group.
y2	The number of responses for the control group.
oddsRatioH0	The odds ratio under the null hypothesis. Defaults to 1.

Details

The Mantel-Haenszel sample size weights are used for stratified samples.

Value

The value of the score test statistic.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
zstatOddsRatio(n1 = c(10, 10), y1 = c(4, 3),
               n2 = c(20, 10), y2 = c(2, 0), oddsRatioH0 = 1)
```

zstatRateDiff	<i>Miettinen-Nurminen Score Test Statistic for Two-Sample Rate Difference</i>
---------------	---

Description

Obtains the Miettinen-Nurminen score test statistic for two-sample rate difference possibly with stratification.

Usage

```
zstatRateDiff(t1, y1, t2, y2, rateDiffH0 = 0)
```

Arguments

t1	The exposure for the active treatment group.
y1	The number of events for the active treatment group.
t2	The exposure for the control group.
y2	The number of events for the control group.
rateDiffH0	The rate difference under the null hypothesis. Defaults to 0.

Details

The Mantel-Haenszel weights are used for stratified samples.

Value

The value of the score test statistic.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
zstatRateDiff(t1 = c(10, 10), y1 = c(4, 3),
              t2 = c(20, 10), y2 = c(2, 0), rateDiffH0 = 0)
```

<code>zstatRateRatio</code>	<i>Miettinen-Nurminen Score Test Statistic for Two-Sample Rate Ratio</i>
-----------------------------	--

Description

Obtains the Miettinen-Nurminen score test statistic for two-sample rate ratio possibly with stratification.

Usage

```
zstatRateRatio(t1, y1, t2, y2, rateRatioH0 = 1)
```

Arguments

- `t1` The exposure for the active treatment group.
- `y1` The number of events for the active treatment group.
- `t2` The exposure for the control group.
- `y2` The number of events for the control group.
- `rateRatioH0` The rate ratio under the null hypothesis. Defaults to 1.

Details

The Mantel-Haenszel weights are used for stratified samples.

Value

The value of the score test statistic.

Author(s)

Kaifeng Lu, <kaifenglu@gmail.com>

Examples

```
zstatRateRatio(t1 = c(10, 10), y1 = c(4, 3),  
               t2 = c(20, 10), y2 = c(2, 0), rateRatioH0 = 1)
```

zstatRiskDiff	<i>Miettinen-Nurminen Score Test Statistic for Two-Sample Risk difference</i>
---------------	---

Description

Obtains the Miettinen-Nurminen score test statistic for two-sample risk difference possibly with stratification.

Usage

```
zstatRiskDiff(n1, y1, n2, y2, riskDiffH0 = 0)
```

Arguments

n1	The sample size for the active treatment group.
y1	The number of responses for the active treatment group.
n2	The sample size for the control group.
y2	The number of responses for the control group.
riskDiffH0	The risk difference under the null hypothesis. Defaults to 0.

Details

The Mantel-Haenszel sample size weights are used for stratified samples.

Value

The value of the score test statistic.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
zstatRiskDiff(n1 = c(10, 10), y1 = c(4, 3),  
              n2 = c(20, 10), y2 = c(2, 0), riskDiffH0 = 0)
```

zstatRiskRatio	<i>Miettinen-Nurminen Score Test Statistic for Two-Sample Risk Ratio</i>
----------------	--

Description

Obtains the Miettinen-Nurminen score test statistic for two-sample risk ratio possibly with stratification.

Usage

```
zstatRiskRatio(n1, y1, n2, y2, riskRatioH0 = 1)
```

Arguments

n1	The sample size for the active treatment group.
y1	The number of responses for the active treatment group.
n2	The sample size for the control group.
y2	The number of responses for the control group.
riskRatioH0	The risk ratio under the null hypothesis. Defaults to 1.

Details

The Mantel-Haenszel sample size weights are used for stratified samples.

Value

The value of the score test statistic.

Author(s)

Kaifeng Lu, <kaifengl@gmail.com>

Examples

```
zstatRiskRatio(n1 = c(10, 10), y1 = c(4, 3),  
               n2 = c(20, 10), y2 = c(2, 0), riskRatioH0 = 1)
```

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