## Package ‘RTSA’

November 23, 2023

<table>
<thead>
<tr>
<th><strong>Type</strong></th>
<th>Package</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Title</strong></td>
<td>'Trial Sequential Analysis' for Error Control and Inference in Sequential Meta-Analyses</td>
</tr>
<tr>
<td><strong>Version</strong></td>
<td>0.2.2</td>
</tr>
<tr>
<td><strong>Description</strong></td>
<td>Frequentist sequential meta-analysis based on 'Trial Sequential Analysis' (TSA) in programmed in Java by the Copenhagen Trial Unit (CTU). The primary function is the calculation of group sequential designs for meta-analysis to be used for planning and analysis of both prospective and retrospective sequential meta-analyses to preserve type-I-error control under sequential testing. 'RTSA' includes tools for sample size and trial size calculation for meta-analysis and core meta-analyses methods such as fixed-effect and random-effects models and forest plots. TSA is described in Wetterslev et. al (2008) <a href="">doi:10.1016/j.jclinepi.2007.03.013</a>. The methods for deriving the group sequential designs are based on Jennison and Turnbull (1999, ISBN:9780849303166).</td>
</tr>
<tr>
<td><strong>License</strong></td>
<td>GPL (&gt;= 2)</td>
</tr>
<tr>
<td><strong>URL</strong></td>
<td><a href="https://github.com/AnneLyng/RTSA">https://github.com/AnneLyng/RTSA</a></td>
</tr>
<tr>
<td><strong>BugReports</strong></td>
<td><a href="https://github.com/AnneLyng/RTSA/issues">https://github.com/AnneLyng/RTSA/issues</a></td>
</tr>
<tr>
<td><strong>Imports</strong></td>
<td>stats, metafor, ggplot2, scales, Rcpp (&gt;= 0.11.0)</td>
</tr>
<tr>
<td><strong>LinkingTo</strong></td>
<td>Rcpp</td>
</tr>
<tr>
<td><strong>Encoding</strong></td>
<td>UTF-8</td>
</tr>
<tr>
<td><strong>Depends</strong></td>
<td>R (&gt;= 3.5.0)</td>
</tr>
<tr>
<td><strong>LazyData</strong></td>
<td>true</td>
</tr>
<tr>
<td><strong>RoxygenNote</strong></td>
<td>7.2.3</td>
</tr>
<tr>
<td><strong>Suggests</strong></td>
<td>gsDesign, CompQuadForm, dplyr, kableExtra, rmarkdown, knitr, bookdown, gridExtra, testthat (&gt;= 3.0.0)</td>
</tr>
<tr>
<td><strong>VignetteBuilder</strong></td>
<td>knitr, bookdown</td>
</tr>
<tr>
<td><strong>Config/testthat/edition</strong></td>
<td>3</td>
</tr>
<tr>
<td><strong>NeedsCompilation</strong></td>
<td>yes</td>
</tr>
</tbody>
</table>
boundaries

Boundaries for group sequential designs

Description

Calculates alpha- and potentially beta-spending boundaries for group sequential designs for meta-analysis. Should be used for exploring how the different arguments affect the sequential design. The function is not intended to be used individually for Trial Sequential Analysis. For this purpose, we recommend RTSA().

Usage

boundaries(
  timing,
  alpha = 0.05,
  beta = 0.1,
  side = 2,
  futility = "none",
  es_alpha = "esOF",
  es_beta = NULL,
  type = "design",
)
boundaries

```
design_R = NULL,
tol = 1e-09
```

Arguments

- **timing**
  Expected timings of interim analyses and final analysis as a vector consisting of values from 0 to 1.

- **alpha**
  The level of type I error as a percentage, the default is 0.05 corresponding to 5%.

- **beta**
  The level of type II error as a percentage, the default is 0.1 corresponding to 10%.

- **side**
  Whether a 1- or 2-sided hypothesis test is used. Defaults to 2. Options are 1 or 2.

- **futility**
  Futility boundaries added to design. Options are: none, non-binding and binding. Default is “none”.

- **es_alpha**
  The error spending function for alpha-spending. Options are: "esOF" (Lan & DeMets version of O’Brien-Fleming boundaries), "esPoc" (Lan & DeMets version of Pocock boundaries), "HSDC" (Hwang Sihi and DeCani) and "rho" (rho family). Defaults to "esOF".

- **es_beta**
  The error spending function for beta-spending. For options see es_alpha. Defaults to NULL.

- **type**
  Whether the boundaries are used for design or analysis. We recommend only to use the boundaries() function with type equal to design. Defaults to NULL.

- **design_R**
  If type is analysis, a scalar for achieving the right amount of power is required. It is recommended not to use the boundaries() function with the setting type equal to analysis. Defaults to NULL.

- **tol**
  Tolerance level for numerical integration. Defaults to 1e-09.

Value

A boundaries object which includes:

- **inf_frac**
  Timing of interim analyses and final analysis. Potentially modified if type = "analysis".

- **org_inf_frac**
  Original timing. If type = "design".

- **alpha_ubound**
  Upper alpha-spending boundaries

- **alpha_lbound**
  Lower alpha-spending boundaries

- **alpha**
  As input

- **alpha_spend**
  List of cumulative and incremental spending

- **delta**
  Drift parameter

- **design_R**
  If type = "analysis" it is the scalar for correct power in the design. Else NULL.
coronary

info List of the information as the squareroot of the information increments and the squareroot of the cumulative information
beta_ubound Upper beta-spending boundaries
beta_lbound Lower beta-spending boundaries
root Scalar for achieving correct power
beta_spend List of cumulative and incremental spending
pwr List of probabilities for rejecting the null under the sample size settings being true at each analysis and the sum.
tle List of probabilities for type-I-error at each analysis and the sum
side As input
beta As input
es_alpha As input
es_beta As input
type As input
futility As input

Examples

boundaries(timing = c(0.25, 0.5, 0.75, 1), alpha = 0.05, beta = 0.1,
side = 2, futility = "non-binding", es_alpha = "esOF", es_beta = "esOF")

coronary Dataset of trials investigating the intensity of statin therapy on the risk of myocardial infarction or coronary death

Description

A dataset containing trials investigating myocardial infarction or coronary death among patients with acute coronary syndromes or chronic coronary artery disease of statin therapy intensity. The trials compared low intensities of statin to higher intensities.

Usage

coronary

Format

A data frame with 4 rows and 5 variables:

study Name of first author of the trial
eI Number of events in the intervention group
nI Number of participants in the intervention group
eC Number of events in the control group
nC Number of participants in the control group
Description

A dataset containing trials investigating on the length of hospital stay when receiving early supported discharge (ESD) service versus conventional care. The outcome is length of initial hospital stay counted in days.

Usage

eds

Format

A data frame with 9 studies and 8 variables:

Details

- study. Name of the city of the study
- year. Year of the trial
- mI. Mean duration at hospital in intervention (ESD) group
- mC. Mean duration at hospital in control group
- sdI. Standard deviation of intervention (ESD) estimate
- sdC. Standard deviation of control estimate
- nI. Number of participants in the intervention (ESD) group
- nC. Number of participants in the control group

References

inference

Inference calculations for sequential meta-analysis

Description
Calculates point-estimates, p-values and confidence intervals. Computes naive inference and TSA-adjusted confidence intervals. If the meta-analysis crosses a alpha-spending boundary, a binding beta-spending boundary or reached the sequential RIS, stage-wise ordered inference is also calculated. This function is not supposed to be used individually for Trial Sequential Analysis (TSA). RTSA() is recommended for TSA.

Usage

inference(
  bounds,
  timing,
  ana_times,
  ma,
  fixed,
  org_timing,
  inf_type = "sw",
  conf_level = 0.95,
  final_analysis = FALSE,
  tol = 1e-15
)

Arguments

- **bounds**: The boundaries for the analysis as calculated by the boundaries() function in RTSA.
- **timing**: The timing of the studies relative to the sequential RIS. A vector consisting of values equal to the proportion of study participants out of the sequential RIS.
- **ana_times**: The analysis times presented as a vector. Describes at which studies the meta-analyses were performed. If one expects that the meta-analysis was updated per study a vector from 1 to the number of studies included can be used.
- **ma**: A metaanalysis object from the metaanalysis function.
- **fixed**: Whether the analysis is for fixed-effect or random-effects meta-analysis. Options are TRUE (meta-analysis is fixed-effect) or FALSE (meta-analysis is random-effects).
- **org_timing**: The timing of all included studies as a proportion of RIS and not sequential RIS.
- **inf_type**: For now only option is "sw" (stage-wise). Type of inference used for point estimates, confidence intervals and p-values.
- **conf_level**: The confidence interval level. Defaults to 0.95 which is 95%.
- **final_analysis**: Whether or not the this analysis is considered the final analysis.
- **tol**: The tolerance level. Set to 1e+09.
Value

A data.frame of cumulative meta-analysis results including stopping boundaries and a list of conditional sequential inference to be parsed to RTSA

results_df

A data.frame containing information about: Cumulative test values, cumulative outcomes, timing of trials, stopping boundaries (alpha_upper, alpha_lower, beta_upper, beta_lower), naive confidence intervals, TSA-adjusted confidence intervals, cumulative p-values and standard deviations.

seq_inf

If the meta-analysis crosses an alpha-spending boundary, a binding beta-spending boundary or reaches the required information size inference conditional on stopping is provided. A median unbiased estimate, lower and upper confidence interval, and p-value is calculated based on stage-wise ordering.

Examples

ma <- metaanalysis(data = perioOxy, outcome = "RR", mc = 0.8)
sts <- ma$res$NR_D2$NR_D2_full
timing <- cumsum(perioOxy$nI + perioOxy$nC)/sts
bound_oxy <- boundaries(timing = timing, alpha = 0.05, beta = 0.2, side = 2, futility = "none", es_alpha = "esOF")
inference(timing = bound_oxy$inf_frac, bounds = bound_oxy, ma = ma, fixed = FALSE, ana_times = 1:length(timing), org_timing = timing)

---

metaanalysis

**Fixed-effect or random-effects meta-analysis**

Description

Computes a fixed-effect or random-effects meta-analysis including heterogeneity statistics. If mc is specified, a retrospective sample and trial size is calculated.

Usage

metaanalysis(
    outcome,
    data,
    side = 2,
    alpha = 0.05,
    beta = 0.1,
    weights = "IV",
    re_method = "DL_HKSJ",
    tau_ci_method = "BJ",
    cont_vartype = "equal",
    mc = NULL,
    RRR = NULL,
    sd_mc = NULL,
metaanalysis

```r
study = NULL,
conf_level = 0.95,
zero_adj = 0.5,
...
)
```

**Arguments**

- `outcome` Outcome metric for the studies. Choose between: MD (mean difference), RR (relative risk), RD (risk difference), or OR (odds ratio).

- `data` A data.frame containing the study results. The data set must containing a specific set of columns. These are respectively `eI` (events in intervention group), `eC` (events in control group), `nC` (participants intervention group) or `nI` (participants control group) for discrete data, or, `mI` (mean intervention group), `mC` (mean control group), `sdI` (standard error intervention group), `sdC` (standard error control group), `nC` (participants intervention group) and `nI` (participants control group) for continuous outcomes. Preferable also a `study` column as an indicator of study.

- `side` Whether a 1- or 2-sided hypothesis test is used. Options are 1 or 2. Default is 2.

- `alpha` The level of type I error as a percentage, the default is 0.05 corresponding to 5%.

- `beta` The level of type II error as a percentage, the default is 0.1 corresponding to 10%. Not used unless a sample and trial size calculation is wanted.

- `weights` Method for calculating weights. Options are "MH" (Mantel-Haenzel and only optional for binary data) or "IV" (Inverse variance weighting). Default is "IV".

- `re_method` Methods are "DL" for DerSimonian-Laird or "DL_HKSJ" for DerSimonian-Laird with Hartung-Knapp-Sidik-Jonkman adjustment. Default is "DL_HKSJ".

- `tau_ci_method` Methods for computation of confidence interval for heterogeneity estimate tau. Calls rma.uni from the metafor package. Options are "BJ" and "QP". Default is "BJ"

- `cont_vartype` Variance type for continuous outcomes. Choices are "equal" (homogeneity of treatment group variances) or "non-equal" (heterogeneity of treatment group variances). Default is "equal".

- `mc` Minimum clinically relevant value. Used for sample and trial size calculation.

- `RRR` Relative risk reduction. Used for binary outcomes with outcome metric RR. Argument mc can be used instead. Must be a value between 0 and 1.

- `sd_mc` The expected standard deviation. Used for sample and trial size calculation for mean differences.

- `study` Optional vector of study IDs. If no study indicator is provided in `data`, a vector of study indicators e.g. names.

- `conf_level` Confidence interval coverage

- `zero_adj` Zero adjustment for null events in binary data. Options for now is 0.5. Default is 0.5.

- `...` Additional variables.
### Value

A `metaanalysis` object which is a list with 6 or 7 elements.

- **study_results**: A data.frame containing study results which is information about the individual studies.
- **meta_results**: A data.frame containing the results of the meta-analysis such as the pooled estimate, its standard error, confidence interval and p-value.
- **hete_results**: A list containing statistics about heterogeneity.
- **metaPrepare**: A list containing the elements used for calculating the study results.
- **synthesize**: A list containing the elements used for calculating the meta-analysis results.
- **settings**: A list containing the arguments used in the `metaanalysis` call.
- **ris**: (Only when `mc` has been specified or meta-analysis is created as part of RTSA). List of sample size and trial size calculation. See documentation for `ris`.

### Examples

#### ### Basic uses

```r
# Use perioOxy data from package and run meta-analysis with default settings
data(perioOxy)
metaanalysis(outcome = "RR", data = perioOxy, study = perioOxy$trial)
```

```r
# Run same meta-analysis but with odds ratio as outcome metric, Mantel-Haenzel
# weights and DerSimonian-Laird for the variance estimate
metaanalysis(outcome = "OR", data = perioOxy, study = perioOxy$trial,
weights = "MH", re_method = "DL")
```

```r
# Run meta-analysis with mean difference as outcome metric
data(eds)
metaanalysis(outcome = "MD", data = eds)
```

#### ### Retrospective sample size calculation

```r
# minimal clinical relevant difference set to an odds ratio of 0.7.
ma <- metaanalysis(outcome = "OR", data = perioOxy, mc = 0.7)
ma$ris
```

---

### minTrial

*Minimum number of trials needed for a specific level of power*

#### Description

Calculates minimum number of trials needed to achieve power in a meta-analysis with heterogeneity.
Usage

```r
minTrial(
  outcome,
  mc,
  tau2,
  alpha,
  beta,
  side,
  pC = NULL,
  p1 = NULL,
  var.mc = NULL,
  var.random = NULL,
  trials = NULL
)
```

Arguments

- **outcome**: Metric of interest, options include "RR" (relative risk), "OR" (odds ratio), "RD" (risk difference) and "MD" (mean difference).
- **mc**: Minimal clinical relevant value provided as a numeric value. Such as 0.8 for e.g. an odds ratio of 0.8.
- **tau2**: Heterogeneity estimate. Can be extracted from the metaanalysis() function.
- **alpha**: The level of type I error as a percentage, the default is 0.05 corresponding to 5%.
- **beta**: The level of type II error as a percentage, the default is 0.1 corresponding to 10%.
- **side**: Whether a 1- or 2-sided hypothesis test is used. Options are 1 or 2.
- **pC**: Probability of event in control group. Only used for outcomes "RR", "OR" and "RD".
- **p1**: Probability of event in treatment group. Only used for outcome "RD".
- **var.mc**: Variance of the estimated effect when outcome is "MD". Not required for outcome types "OR", "RR" or "RD".
- **var.random**: Estimated variance from the random-effects meta-analysis. Used then a meta-analysis have already been made previously.
- **trials**: Optional argument. Number of trials of interest for to provide the number of participants needed for that exact number of trials.

Value

Either a number (minimum required trials) or the minimum required required trials together with a matrix of required participants per trial given different number of trials.
Examples

# Minimum number of trials for a prospective meta-analysis
minTrial(outcome = "RR", pC = 0.5, mc = 0.7, tau2 = 0.05, alpha = 0.05, beta = 0.1, side = 2)

# Minimum number of trials still needed for a retrospective meta-analysis
# Note that retrospective sample size calculations are prone to bias
ma <- metaanalysis(outcome = "RR", data = perioOxy)
ris(outcome = "RR", mc = 0.80, ma = ma, type = "retrospective", fixed = FALSE, beta = 0.1, alpha = 0.05, side = 2)

perioOxy

Dataset of RCTs investigating the effect of 80% perioperative oxygen vs. 30-35% perioperative oxygen on surgical site infection.

Description

A dataset containing data on seven trials which includes their number of events per treatment group, where intervention is 80% oxygen and control is 30-35% oxygen, number of participants in each treatment group and the year of the trial.

Usage

perioOxy

Format

A data frame with 7 rows and 6 variables:

study  Name of first author of the trial
eI     Number of events in the intervention group (80% oxygen)
nI     Number of participants in the intervention group (80% oxygen)
eC     Number of events in the control group (30-35% oxygen)
nC     Number of participants in the control group (30-35% oxygen)
plot.boundaries  
Plot of boundaries for group sequential designs

Description
Plot of boundaries for group sequential designs

Usage
## S3 method for class 'boundaries'
plot(x, theme = "classic", ...)

Arguments
x  boundaries object
theme  Whether the theme is "classic" or "aussie"
...  Other arguments to plot.boundaries

Value
Plot. Either a plot for two- or one-sided testing.

Examples
bounds <- boundaries(timing = c(0.5, 0.75, 1), alpha = 0.025, beta = 0.2,
side = 1, futility = "none", es_alpha = "esOF")
plot(x = bounds)

plot.metaanalysis  
Forestplot for metaanalysis object.

Description
Forestplot for metaanalysis object.

Usage
## S3 method for class 'metaanalysis'
plot(x, type = "both", xlims = NULL, ...)

Arguments

- **x** metaanalysis object from the RTSA package.
- **type** Define whether or not both fixed-effect and random-effects meta-analysis results should be printed on the plot. Options are: "fixed", "random" or "both". Default is "both".
- **xlims** Set default limits on the outcome scale. Default is NULL.
- **...** Additional arguments

Examples

```r
# Example with OR
ma <- metaanalysis(data = coronary, outcome = "OR")
plot(ma)

# Example with RR
ma <- metaanalysis(data = perioOxy, outcome = "RR")
plot(ma)

# Example with MD
ma <- metaanalysis(data = eds, outcome = "MD")
plot(ma, type = "random")
```

plot.RTSA

Plot RTSA object. Returns the R version of the original TSA plot.

Description

Plot RTSA object. Returns the R version of the original TSA plot.

Usage

```r
## S3 method for class 'RTSA'
plot(x, model = "random", type = "classic", theme = "classic", ...)
```

Arguments

- **x** RTSA object
- **model** Whether a fixed- or random-effects meta-analysis should be used. Defaults to random.
- **type** Should Z-scores (classic) or outcome values (outcome) be plotted.
- **theme** Whether the theme is traditional TSA (classic) or modern (modern)
- **...** Other arguments to plot.RTSA

Value

Plot. Either a plot for two sided testing or one-sided
Examples

```r
data(perioOxy)
outRTSA <- RTSA(type = "analysis", data = perioOxy, outcome = "RR", mc = 0.8,
side = 2, alpha = 0.05, beta = 0.2, fixed = FALSE, es_alpha = "esOF", design = NULL)
plot(x = outRTSA)
```

---

### ris

**Calculate required sample and trials size.**

**Description**

Calculate required sample and trials size.

**Usage**

```r
ris(
  outcome,
  mc,
  side = 2,
  alpha = 0.05,
  beta = 0.1,
  fixed = TRUE,
  sd_mc = NULL,
  pC = NULL,
  p1 = NULL,
  ma = NULL,
  tau2 = NULL,
  I2 = NULL,
  D2 = NULL,
  type = "prospective",
  trials = NULL,
  RTSA = FALSE,
  ...
)
```

**Arguments**

- **outcome**: Choose between "MD" (mean difference), "RR" (relative risk), "OR" (odds ratio) or "RD" (risk difference).
- **mc**: Minimum clinical relevant effect. For "OR" or "RR" set to natural scale, not log scale.
- **side**: Test type. Set to 1 or 2 depending on the test being 1- or 2-sided.
- **alpha**: The level of type I error as a percentage, the default is 0.05 corresponding to 5%.
beta
The level of type II error as a percentage, the default is 0.1 corresponding to 10%.

fixed
Should sample size be based on a fixed-effect (TRUE) or random-effects (FALSE) model. Defaults to TRUE.

sd_mc
Standard deviation of estimated effect. Only needed when outcome type is "MD".

pC
Probability of event in control group. Only needed when outcome type is "OR", "RR" or "RD".

p1
Probability of event in treatment group. Only needed when outcome type is "RD".

ma
An optional metaanalysis object. Required for retrospective sample size calculations.

tau2
The value of the heterogeneity. Use when estimating the sample size under a random effects model. If data is provided, the estimated heterogeneity is used instead.

I2
Optional argument. Inconsistency.

D2
Optional argument. Diversity.

type
Whether the type of calculation is for "prospective" meta-analysis or "retrospective" meta-analysis. If the type is retrospective, one should add a meta-analysis object to the function. See argument ma.

trials
Optional numeric argument. If one is interested in a specific number of trials.

RTSA
Whether the ris function was called via the RTSA function. Purely operational argument.

...
additional arguments

Value
A list of up to 6 elements:

settings
A list containing the arguments provided to the ris function.

NF
The total number of required participants in a fixed-effect meta-analysis if type is prospective. Contains a list if the type is retrospective, where NF is the additional required number of participants and NF_full is the total required number of participants.

NR_tau
A list containing: minTrial the minimum number of trials. nPax a matrix containing four possible number of trials with the number of participants per trial and total number of participants. tau2 the estimate used for the calculation. Might contain NR_tau_l1 and NR_tau_u1 which contain the same three elements. NR_tau_l1 is based on the lower value in the confidence interval of tau2. NR_tau_u1 is based on the upper value in the confidence interval for tau2. If the type is prospective the numbers are the total required. If the type is retrospective the numbers are the additional required.

NR_D2
The total number of required participants in a random-effects meta-analysis adjusted by diversity (D2) if type is prospective. Contains a list if the type is retrospective, where NR_D2 is the additional required number of participants and NR_D2_full is the total required number of participants.
The total number of required participants in a random-effects meta-analysis adjusted by inconsistency (I^2) if type is prospective. Contains a list if the type is retrospective, where NR_I2 is the additional required number of participants and NR_I2_full is the total required number of participants.

Examples

```r
# Sample and trial size calculation for prospective meta-analysis
ris(outcome = "RR", mc = 0.8, pC = 0.12, fixed = TRUE, alpha = 0.05, beta = 0.1, side = 2)

# Additional sample and trial size calculation for retrospective meta-analysis
# It is calculated directly from the metaanalysis() function
data("perioOxy")
ma <- metaanalysis(outcome = "RR", data = perioOxy, mc = 0.8, beta = 0.2)
ma$ris
# Or by using the two functions in sequence
ma <- metaanalysis(outcome = "RR", data = perioOxy)
ris(outcome = "RR", mc = 0.8, ma = ma, type = "retrospective", fixed = FALSE, beta = 0.2, alpha = 0.05, side = 2)
```

RTSA

*R version of Trial Sequential Analysis. Used for designing and analysing sequential meta-analyses.*

Description

R version of Trial Sequential Analysis. Used for designing and analysing sequential meta-analyses.

Usage

```r
RTSA(
  type = "design",
  outcome = NULL,
  side = 2,
  alpha = 0.05,
  beta = 0.1,
  futility = "none",
  es_alpha = "esOF",
  es_beta = NULL,
  timing = NULL,
  data = NULL,
  design = NULL,
  ana_times = NULL,
  fixed = FALSE,
  mc = NULL,
  RRR = NULL,
  sd_mc = NULL,
)```
pC = NULL,
weights = "MH",
re_method = "DL_HKSJ",
tau_ci_method = "BJ",
gamma = NULL,
rho = NULL,
study = NULL,
cont_vartype = "equal",
zero_adj = 0.5,
tau2 = NULL,
I2 = NULL,
D2 = NULL,
trials = NULL,
final_analysis = NULL,
inf_type = "sw",
conf_level = 0.95,
random_adj = "tau2",
power_adj = TRUE,
...)

Arguments

type
Type of RTSA. Options are "design" or "analysis".

outcome
Outcome metric. Options are: RR (risk ratio/relative risk), OR (odds ratio), RD (risk difference) and MD (mean difference).

side
Whether a 1- or 2-sided hypothesis test is used. Options are 1 or 2. Default is 2.

alpha
The level of type I error as a percentage, the default is 0.05 corresponding to 5%.

beta
The level of type II error as a percentage, the default is 0.1 corresponding to 10%.

futility
Futility boundaries added to design. Options are: none, non-binding and binding. Default is "none".

es_alpha
The spending function for alpha-spending. Options are: esOF (Lan & DeMets version of O'Brien-Fleming), esPoc (Lan & DeMets version of Pocock), HSDC (Hwang Sihi and DeCani) and rho (rho family).

es_beta
The spending function for beta-spending. For options see es_alpha.

timing
Expected timings of interim analyses when type = "design". Defaults to NULL.

data
A data.frame containing the study results. The data set must containing a specific set of columns. These are respectively 'eI' (events in intervention group), 'eC' (events in control group), 'nC' (participants intervention group) or 'nI' (participants control group) for discrete data, or, 'mI' (mean intervention group), 'mC' (mean control group), 'sdI' (standard error intervention group), 'sdC' (standard error control group), 'nC' (participants intervention group) and 'nI' (participants control group) for continuous outcomes. Preferable also a 'study' column as an indicator of study.
design  RTSA object where type is design.
ana_times An optional vector of analysis times. Used if the sequential analysis is not done for all studies included in the meta-analysis.
fixed Should only a fixed-effect meta-analysis be computed. Default is FALSE.
mc Minimal clinical relevant outcome value
RRR Relative risk reduction. Used for binary outcomes with outcome metric RR. Argument mc can be used instead. Must be a value between 0 and 1.
sd_mc The expected standard deviation. Used for sample size calculation for mean differences.
pC The expected probability of event in the control group. Used for sample size calculation for binary outcomes.
weights Weighting method options include IV (inverse-variance) and MH (Mantel-Haenszel). Defaults to MH.
tau_ci_method Method for calculating confidence intervals for the estimated heterogeneity tau^2. Options are "QP" for Q-profiling and "BJ" for Biggelstaff ....
gamma Parameter for the HSDC error spending function.
rho Parameter for the rho family error spending function.
study An optional vector of study names and perhaps year of study. Defaults to NULL.
cont_vartype For mean difference outcomes, do we expect the variance in the different groups to be "equal" or "non-equal".
zero_adj Zero adjustment. Options for now is 0.5.
tau2 Heterogeneity estimate. Used for sample and trial size calculation. Defaults to NULL.
I2 Inconsistency estimate. Used for sample and trial size calculation. Defaults to NULL.
D2 Diversity estimate. Used for sample and trial size calculation. Defaults to NULL.
trials Number of anticipated extra trials. Used for heterogeneity adjustment by tau2.
final_analysis Whether or not the current analysis is the final analysis.
inf_type Stopping time confidence interval. Options for now is sw (stage-wise).
conf_level Confidence level on stopping time confidence interval.
random_adj The sample size adjustment based on presence of heterogeneity. Options are "D2" (Diversity), "I2" (Inconsistency) and "tau2" (the heterogeneity estimate). Default is "tau2".
power_adj Whether the sample size should be adjusted by the sequential design. Defaults to TRUE.

... other arguments
RTSA

Value

A RTSA object, a list of five elements:

- **settings** A list containing all of the settings used in the RTSA call. See Arguments.
- **ris** List containing sample and trial size calculations for a non-sequential meta-analysis. See documentation for ris function.
- **bounds** List of stopping boundaries, timing of trials and more. See documentation for boundaries function.
- **results** List of 3 to 7 elements. AIS Achieved information size. RIS Fixed-effect required information size for a non-sequential meta-analysis. SMA_RIS RIS adjusted for sequential analysis. HARIS Heterogeneity adjusted required information size for a non-sequential meta-analysis. SMA_HARIS HARIS adjusted for sequential analysis. results_df a data.frame of inference, see documentation for inference function. seq_inf a list of conditional inference, see documentation for inference function. metaanalysis A metaanalysis object, see documentation for metaanalysis function. design_df a data.frame containing the stopping boundaries and timings from the design.
- **warnings** List of warnings

Examples

```r
## Not run:
### Retrospective sequential meta-analysis:
# A RRR of 20% is expected which gives mc = 1 - RRR = 0.8.
# No futility boundaries
data(perioOxy)
RTSA(type = "analysis", data = perioOxy, outcome = "RR", mc = 0.8, side = 2,
       alpha = 0.05, beta = 0.2, es_alpha = "esOF")

# Set binding futility boundaries
# And use Lan and DeMets' version of Pocock stopping boundaries
RTSA(type = "analysis", data = perioOxy, outcome = "RR", mc = 0.8, side = 2,
       alpha = 0.05, beta = 0.2, es_alpha = "esOF", futility = "binding",
       es_beta = "esPoc", random_adj = "D2")

# Set non-binding futility boundaries
RTSA(type = "analysis", data = perioOxy, outcome = "RR", mc = 0.8, side = 2,
       alpha = 0.05, beta = 0.2, es_alpha = "esOF", futility = "non-binding",
       es_beta = "esOF")

### Design a prospective sequential meta-analysis
# For continuous data without expected heterogeneity
RTSA(type = "design", outcome = "MD", mc = 5, sd_mc = 10, side = 1,
       timing = c(0.33, 0.66, 1), fixed = TRUE,
       alpha = 0.025, beta = 0.1, es_alpha = "esOF", futility = "non-binding",
       es_beta = "esPoc")

# For binary outcome
RTSA(type = "design", outcome = "RR", mc = 0.75, side = 1,
       timing = c(0.33, 0.66, 1), pC = 0.1, D2 = 0.1,
```
alpha = 0.025, beta = 0.2, es_alpha = "esOF", futility = "non-binding",
es_beta = "esOF")

# extract sample size calculation
out_rtsa <- RTSA(type = "design", outcome = "RR", mc = 0.75, side = 1,
timing = c(0.33, 0.66, 1), pC = 0.1, D2 = 0.1,
alpha = 0.025, beta = 0.2, es_alpha = "esOF", futility = "non-binding",
es_beta = "esOF")
out_rtsa$ris

# plot the design
plot(out_rtsa)

# update the design with data as it accumulates (here toy-data)
fake_data <- data.frame(eI = c(10,10), eC = c(13, 11), nI = c(750, 750),
nC = c(750,750))
RTSA(type = "analysis", design = out_rtsa, data = fake_data)

# plot the analysis
an_rtsa <- RTSA(type = "analysis", design = out_rtsa, data = fake_data)
plot(an_rtsa)

## End(Not run)
Index

* datasets
  - coronary, 4
  - eds, 5
  - perioOxy, 11

boundaries, 2

coronary, 4

datasets, 4

eds, 5

inference, 6

metaanalysis, 7

minTrial, 9

perioOxy, 11

plot.boundaries, 12

plot.metaanalysis, 12

plot.RTSA, 13

print.metaanalysis (metaanalysis), 7

print.ris (ris), 14

print.RTSA (RTSA), 16

ris, 14

RTSA, 16