Package ‘BayesianPlatformDesignTimeTrend’

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Title Simulate and Analyse Bayesian Platform Trial with Time Trend

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Description Simulating the sequential multi-arm multi-stage or platform trial with Bayesian approach using the 'rstan' package, which provides the R interface for the Stan. This package supports fixed ratio and Bayesian adaptive randomization approaches for randomization. Additionally, it allows for the study of time trend problems in platform trials. There are demos available for a multi-arm multi-stage trial with two different null scenarios, as well as for Bayesian trial cutoff screening. The Bayesian adaptive randomisation approaches are described in:
The randomisation algorithm is described in:
Zhao W <doi:10.1016/j.cct.2015.06.008>.
The analysis methods of time trend effect in platform trial are described in:
Saville et al. (2022) <doi:10.1177/17407745221112013> and

URL https://github.com/ZXW834/BayesianPlatformDesignTimeTrend

BugReports https://github.com/ZXW834/BayesianPlatformDesignTimeTrend/issues

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BayesianPlatformDesignTimeTrend-package

Description

This package simulates the multi-arm multi-stage or platform trial with bayesian approach using the 'rstan' package, which provides the R interface for to the stan. The package uses Thall’s and Trippa’s randomisation approach for Bayesian adaptive randomisation. In addition, the time trend problem of platform trial can be studied in this package. There is a demo for multi-arm multi-stage trial for two different null scenario in this package.

References


AdaptiveRandomisation

Description

This is a function doing the randomisation process. This Function generates the Sequence for patient allocation to each arm, patient outcomes.
AdaptiveRandomisation

Usage

AdaptiveRandomisation(
  Fixratio,
  rand.algo,
  K,
  n.new,
  randomprob,
  treatmentindex,
  groupwise.response.probs,
  group,
  armleft,
  max.deviation,
  trend_add_or_multip,
  trend.function,
  trend.effect,
  ns,
  Fixratiocontrol
)

Arguments

Fixratio A indicator TRUE/FALSE
rand.algo Randomisation algorithm: "Coin": Biased coin; "Urn": Urn method
K Total number of arms at the beginning
n.new The cohort size
randomprob A named vector of randomisation probability to each arm
treatmentindex The vector of treatment arm index excluding the control arm whose index is 0
groupwise.response.probs A matrix of response probability of each arm
group The current stage
armleft The number of treatment left in the platform (>2)
max.deviation Tuning parameter of using urn randomisation method.
trend_add_or_multip How time trend affects the true response probability: "add" or "mult"
trend.function The function returns time trend effect regarding to different time trend pattern
trend.effect The strength of time trend effect as a parameter in trend.function()
ns A vector of accumulated number of patient at each stage
Fixratiocontrol A numeric value indicating the weight of control in randomisation. Eg. 1 means equal randomisation, 2 means the number of patients allocated to control is twice as large as other treatment arm.
Value

A list of patient allocation and patient outcome nstage: A vector of the number of patients allocated to each arm ystage: A vector of the patients outcome after treating with each arm znew: A vector of treatment index assigned to each patient in the current cohort ynew: A vector of outcome index record for each patient after treatment in the current cohort

Author(s)

Ziyan Wang

References


Examples

AdaptiveRandomisation(
   Fixratio = FALSE,
   rand.algo = "Urn",
   K = 2,
   n.new = 30,
   randomprob = matrix(c(0.5, 0.5), ncol = 2, dimnames = list(c(),c("1","2"))),
   treatmentindex = 1,
   groupwise.response.probs = matrix(rep(c(0.4, 0.4), 5), byrow = TRUE, ncol = 2, nrow = 5),
   group = 1,
   armleft = 2,
   max.deviation = 3,
   trend_add_or_multip = "mult",
   trend.function = function(ns, group, i, trend.effect) {delta = 0; return(delta)},
   trend.effect = c(0, 0),
   ns = c(30, 60, 90, 120, 150),
   Fixratiocontrol = NA)

AdaptiveRandomisation(
   Fixratio = TRUE,
   rand.algo = "Urn",
   K = 4,
   n.new = 30,
   randomprob = NA,
   treatmentindex = c(1,3),
   groupwise.response.probs = matrix(rep(c(0.4, 0.4,0.4, 0.4), 5), byrow = TRUE, ncol = 4, nrow = 5),
   group = 1,
   armleft = 3,
   max.deviation = 3,
   trend_add_or_multip = "mult",
   trend.function = function(ns, group, i, trend.effect) {delta = 0; return(delta)},
   trend.effect = c(0, 0),
   ns = c(30, 60, 90, 120, 150),
   Fixratiocontrol = 1)
**Description**

This function estimates the mean error rate spent at each interim analysis for a trial. Example usage:

1. `sapply(res = result, fun = alphaspending)` will generate list of the proportion of trial replicates are stopped at each stage for all scenarios in result where result is a list containing output data for different scenario.
2. `sapply(sapply(result,FUN = alphaspending),sum)` will generate the type I error rate or power for all scenario on the result list.
3. `alpha(result)` generate the proportion of trial replicates are stopped at each stage where result is the output data for one specific scenario.
4. `sum(alpha(result))` will generate the type I error rate or power for a specific scenario.

**Usage**

`alphaspending(res)`

**Arguments**

`res` A list of output matrix of a number of trial replicates.

**Value**

The error rate at each interim analysis.

**Author(s)**

Ziyan Wang.

**Examples**

```
## Not run: alphaspending(res)
```
Description

This function adjusts the posterior randomisation probability for each arm using many approaches. Currently Thall’s approach and Trippa’s approach are used. Double biased coin and other method will be added in the next version.

Usage

```r
ARmethod(
  BARmethod, 
  group, 
  stats, 
  post.prob.btcontrol, 
  K, 
  n, 
  tuningparameter = NA, 
  c = NA, 
  a = NA, 
  b = NA, 
  post.prob.best, 
  max.ar = NA, 
  armleft, 
  treatmentindex
)
```

Arguments

- **BARmethod**: The indicator of which adaptive randomisation method is used
- **group**: The current stage
- **stats**: The output matrix
- **post.prob.btcontrol**: The vector of posterior probability of each active treatment arm better than control
- **K**: Total number of arms at the beginning
- **n**: The vector of sample size for each arm
- **tuningparameter**: The tuning parameter indicator for Thall’s approach
- **c**: The tuning parameter for Thall’s approach
- **a**: The hyperparameter parameter for Trippa’s approach
- **b**: The hyperparameter parameter for Trippa’s approach
- **post.prob.best**: Posterior probability of each arm to be the best
max.ar The upper boundary for randomisation ratio for each arm, which is used in Thall’s approach since Trippa’s approach has protection on control arm.

armleft The number of treatment left in the platform (>2)

treatmentindex The vector of treatment arm index excluding the control arm whose index is 0

Value

randomprob: The vector of adjusted randomisation probability to each arm

Author(s)

Ziyan Wang

References


Examples

ARmethod(  
BARmethod = "Thall",  
group = 1,  
stats = matrix(rep(NA, 40), ncol = 8, nrow = 5),  
post.prob.btcontrol = 0.5,  
K = 2,  
n = c(30, 30),  
tuningparameter = "fixed",  
c = 1,  
post.prob.best = c(0.5, 0.5),  
max.ar = 0.75,  
armleft = 2,  
treatmentindex = 1)

ARmethod(  
BARmethod = "Trippa",  
group = 1,  
stats = matrix(rep(NA, 40), ncol = 8, nrow = 5),  
post.prob.bestcontrol = c(0.5, 0.6),  
K = 3,  
n = c(30, 30, 40),  
tuningparameter = NA,  
c = NA,  
a = 3,  
b = 0.75,  
post.prob.best = c(0.3, 0.3, 0.4),  
max.ar = NA,  
armleft = 3,  
post.prob.best = c(0.3, 0.3, 0.4),  
max.ar = NA,  
armleft = 3,  
post.prob.best = c(0.3, 0.3, 0.4),  
max.ar = NA,  
armleft = 3)
Boundaryconstruction

treatmentindex = c(1, 2))

Description
This function constructs the stopping boundary based on input information

Usage
Boundaryconstruction(Stopbound.inf = Stopbound.inf, ns = ns)

Arguments
Stopbound.inf The list of stop boundary information for more see Stopboundinf
ns A vector of accumulated number of patient at each stage

Value
A list of the futility boundary and the efficacy boundary

Author(s)
Ziyan Wang

Examples
Stopbound.inf=list(Stop.type="Early-Pocock",Boundary.type="Symmetric",cutoff=c(0.9928,0.0072))
ns=c(60,120,180,240,300)
Boundaryconstruction(Stopbound.inf, ns)

conjunctivepower_or_FWER

Description
This function reads in the output matrix of a number of trial replicates to calculate the Family wise error rate or Conjunctive power

Usage
conjunctivepower_or_FWER(res, scenario, test.type)
Arguments

res A list of output matrix of a number of trial replicates
scenario The true scenario used to generate the res list
test.type The indicator of whether using one side or two side testing. Please make sure that the input test.type does not conflicts to the data. Otherwise the conjunctive power calculation is wrong

Value

Family wise error rate or Conjunctive power

Examples

```r
## Not run: conjuncativepower_or_FWER(res)
```

dataloginformd Cutoff screening example: the details of grid

Description

Details of grid including family wise error rate of a cutoff value, the cutoff value and the square of cutoff value for modelling and prediction

Usage

dataloginformd

Format

A data frame with 24 rows and 3 variables:

<table>
<thead>
<tr>
<th>tpIE</th>
<th>FWER</th>
</tr>
</thead>
<tbody>
<tr>
<td>cutoff</td>
<td>Cutoff value</td>
</tr>
<tr>
<td>cutoff2</td>
<td>Square of cutoff value</td>
</tr>
</tbody>
</table>
Description

This function does a cutoff screening for trial simulation.

Usage

demo_Cutoffscreening(
  ntrials = 1000,
  trial.fun = simulatetrial,
  grid.inf = list(start = c(0.9, 0.95, 1), extendlength = 15),
  input.info = list(response.probs = c(0.4, 0.4), ns = c(30, 60, 90, 120, 150), max.ar =
  0.75, rand.algo = "Urn", max.deviation = 3, test.type = "Twoside", model.inf =
  list(model = "tlr", ibb.inf = list(pi.star = 0.5, pess = 2, betabinomialmodel =
  ibetabinomial.post), tlr.inf = list(betato_prior_mu = 0, betal_prior_mu = 0,
  betato_prior_sigma = 2.5, betal_prior_sigma = 2.5, beta0_df = 7, betal_df = 7, reg.inf
  = "main", variable.inf = "Fixeffect")), Stop.type = "Early-Pocock", Boundary.type =
  "Symmetric", Random.inf = list(Fixratio = FALSE, Fixratiocontrol = NA,
  BARmethod = "Thall", Thall.tuning.inf = list(tuningparameter = "Fixed", fixvalue =
  1)), trend.inf = list(trend.type = "step", trend.effect = c(0, 0),
  trend.add_or_multip = "mult"),
  cl = 2
)

Arguments

ntrials A numeric variable indicating how many trial replicates you want to run
trial.fun The function of trial simulation, related to MainFunction.R
grid.inf A list of grid information to create start grid and extend grid for cutoff screening.
input.info A list of input information including all information required for trial simulation.
cl A numeric variable indicating how many cores you want to use in parallel programming.

Value

A vector of recommended cutoff. The final value is the latest recommended value. A plot for all
tested cutoff and error rate

Author(s)

Ziyan Wang
A demo for cutoff screening using Bayesian optimisation

Examples

demo_Cutoffscreening(ntrials = 2, cl = 2,
    grid.inf = list(start = c(0.9, 0.95, 1), extendlength = 2))

demo_Cutoffscreening.GP

Description

This function does a cutoff screening for trial simulation using Bayesian optimisation.

Usage

demo_Cutoffscreening.GP(
    ntrials = 1000,
    trial.fun = simulatetrial,
    grid.inf = list(start.length = 10, grid.min = NULL, grid.max = NULL, confidence.level = 0.95, grid.length = 5000, change.scale = FALSE, noise = TRUE, errormrate = 0.1, simulationerror = 0.01, iter.max = 15, plotornot = FALSE),
    power.type = NA,
    response.probs.alt = NA,
    input.info = list(response.probs.null = c(0.4, 0.4, 0.4, 0.4), ns = c(120, 240, 360, 480, 600), max.ar = 0.85, rand.algo = "Urn", max.deviation = 3, test.type = "Twoside", model.inf = list(model = "tlr", ibb.inf = list(pi.star = 0.5, pess = 2, betabinomialmodel = ibetabinomial.post), tlr.inf = list(beta0_prior_mu = 0, betal_prior_mu = 0, beta0_prior_sigma = 2.5, betal_prior_sigma = 2.5, beta0_df = 7, betal_df = 7, reg.inf = "main", variable.inf = "Fixeffect")), Stop.type = "Early-OBF", Boundary.type = "Symmetric",
    Random.inf = list(Fixratio = FALSE, Fixratiocontrol = NA, BARmethod = "Thall", Thall.tuning.inf = list(tuningparameter = "Unfixed", fixvalue = 1)), trend.inf = list(trend.type = "step", trend.effect = c(0, 0, 0, 0), trend_add_or_multip = "mult")),
    cl = 2
)

Arguments

ntrials A numeric variable indicating how many trial replicates you want to run
trial.fun The function of trial simulation, related to MainFunction.R
grid.inf A list of grid information to create start grid and extend grid for cutoff screening. 'start.length' is the size of start grid. Default is 10. 'grid.min' A numeric value or vector (for asymmetric boundary) indicating the lower bound of the grid for screening. For asymmetric boundary, the first value is efficacy minimum value and the second value is futility minimum value. 'grid.max' A numeric value or vector (for asymmetric boundary) indicating the upper bound of the grid for
screening. For asymmetric boundary, the first value is efficacy maximum value and the second value is futility maximum value. 'errorrate' refers to the target of type I error rate or family-wise error rate. Default is 0.05. User can change it to 0.1 for FWER if they think 0.05 is too conservative. The per-hypothesis type I error equals errorrate / (K-1) where (K-1) is the number of treatment arms. 'confidence.level' is a numeric value indicating the confidence level of estimate. Default is 0.95. 'grid.length' A numeric value indicating the grid resolution. Default is 5000 for symmetric boundary. For asymmetric boundary, the length of grid is 101 for both efficacy grid and futility grid. A numeric value indicating the grid resolution. Default is 5000 for symmetric boundary. For asymmetric boundary, the length of grid is 101 for both efficacy grid and futility grid.

'change.scale' is a logic value indicating whether we want to change scale when doing Gaussian process. Default is FALSE. 'noise' is a logic value indicating whether the input x is noisy. Default is TRUE. 'simulationerror' is a numeric value indicating the tolerable error for simulated type I error rate. Default is 0.01, 'iter.max' is a numeric value indicating the maximum number of evaluations. Default is 15. 'plotornot' is a logic value indicating whether the errorrate vs grid plot needed to be generated at each iteration. Default is FALSE.

`power.type` A indicator of which type of power we need to optimise when tuning the cutoff value for asymmetric boundary. Default is NA (Symmetric boundary). The choice of power type is Conjunctive power ("Conjunctive") and Disconjunctive power ("Disconjunctive"). In a two arm trial design, these power type are the same.

`response.probs.alt` A vector of response probability of each arm under the alternative scenario. This is used for power optimisation when tuning the cutoff values for asymmetric boundary. Default is NA.

`input.info` A list of input information including all information required for trial simulation.

`cl` A numeric variable indicating how many cores you want to use in parallel programming.

**Value**

A vector of recommended cutoff. The final value is the latest recommended value. A plot for all tested cutoff and error rate.

**Author(s)**

Ziyan Wang

**Examples**

```r
#Two arm asymmetric boundary screening. Default is OBF boundary.
demo_Cutoffscreening.GP(ntrials = 2, cl = 2,
  power.type = NA,
  response.probs.alt = NA,
  grid.inf = list(
    start.length = 10,
  ))
```
# Four arm asymmetric OBF boundary screening where conjunctive power is optimised.

demo_Cutoffscreening.GP(ntrials = 2, cl = 2, power.type = "Conjunctive",
response.probs.alt = c(0.4, 0.6, 0.6, 0.4), grid.inf = list(
  start.length = 10,
  confidence.level = 0.95,
  grid.length = 101,
  change.scale = FALSE,
  noise = TRUE,
  errorrate = 0.1,
  simulationerror = 0.01,
  iter.max = 15,
  plotornot = FALSE))

input.info = list(
  response.probs.null = c(0.4, 0.4, 0.4, 0.4),
  ns = c(120, 240, 360, 480, 600),
  max.ar = 0.85,
  rand.algo = "Urn",
  max.deviation = 3,
  test.type = "Twoside",
  model.inf = list(
    model = "tlr",
    ibb.inf = list(
      pi.star = 0.5,
      pess = 2,
      betabinomialmodel = ibetabinomial.post
    ),
    tlr.inf = list(
      beta0_prior_mu = 0,
      beta1_prior_mu = 0,
      beta0_prior_sigma = 2.5,
      beta1_prior_sigma = 2.5,
      beta0_df = 7,
      beta1_df = 7,
      reg.inf = "main",
      variable.inf = "Fixeffect"
    )
  ),
  Stop.type = "Early-OBF",
  Boundary.type = "Asymmetric",
  Random.inf = list(
    Fixratio = FALSE,
    Fixratiocontrol = NA,
BARmethod = "Thall",
Thall.tuning.inf = list(tuningparameter = "Unfixed", fixvalue = 1)
),
trend.inf = list(
trend.type = "step",
trend.effect = c(0, 0, 0, 0),
trend_add_or_multip = "mult"
)
)

---

### Description

This is a demo function simulating multi-arm multi-stage design with two different null scenarios where the response probability of control is 0.15 and 0.4, respectively. The clinically meaningful increment on probability scale is 0.2. The stopping boundary is the OBF. The cutoff vector in the demo is tuned to keep Type I error rate to be 0.05. The output data can be saved as .RData file.

### Usage

```r
demo_multscenario(ntrials = 1000, cl = 2, save_data = FALSE)
```

### Arguments

- **ntrials**: A numeric value. The number of total trail replicates for each scenario.
- **cl**: A numeric variable indicating how many cores you want to use in parallel programming.
- **save_data**: An indicator of whether the output data need to be saved. Default is FALSE.

### Value

A list of data for plotting. One is results of trial replicates for all scenarios. The other one is a data frame containing all summarised evaluation metrics for all scenarios.

### Author(s)

Ziyan Wang

### Examples

```r
demo_multscenario(ntrials = 2, cl = 2, save_data = FALSE)
```
disconjunctivepowerfunc

Description

This function reads in the output matrix of a number of trial replicates to calculate the Family wise error rate or disconjunctive power.

Usage

disconjunctivepowerfunc(res)

Arguments

res

A list of output matrix of a number of trial replicates

Value

Disconjunctive power

Examples

## Not run: disconjunctivepowerfunc(res)

GP.optim

GP.optim: optimiser to give the next cutoff for evaluation

Description

A function to predict the next cutoff value for evaluation.

Usage

GP.optim(
  x,
  y.t1E,
  y.pow = NA,
  ESS = NA,
  error.rate = 0.05,
  confidence.level = 0.95,
  grid.length = 1000,
  change.scale = FALSE,
  noise = T,
  grid.min,
  grid.max,
  Boundary.type = "Symmetric"
)
Arguments

- **x**: A numeric vector of cutoff data
- **y.t1E**: A numeric vector of type I error rate data
- **y.pow**: A numeric vector of power data. You can input conjunctive, disjunctive and marginal power data. Default is NA. Only used when Boundary.type == "Asymmetric"
- **ESS**: A matrix of effective sample size. This is only called for asymmetric boundary cutoff screening. Default is NA for symmetric boundary. The first column is the ESS for different cutoff pair under the null scenario, the second column is the ESS for different cutoff pair under the alternative scenario.
- **errorrate**: 'errorrate' refers to the target of type I error rate or family-wise error rate. Default is 0.05. User can change it to 0.1 for FWER if they think 0.05 is too conservative. The per-hypothesis type I error equals errorrate / (K-1) where (K-1) is the number of treatment arms.
- **confidence.level**: A numeric value indicating the confidence level of estimate. Default is 0.95
- **grid.length**: A numeric value indicating the grid resolution. Default is 1000.
- **change.scale**: A logic value indicating whether we want to change scale when doing Gaussian process. Default is FALSE.
- **noise**: A logic value indicating whether the input x is noisy. Default is TRUE.
- **grid.min**: A numeric value or vector (for asymmetric boundary) indicating the lower bound of the grid for screening. For asymmetric boundary, the first value is efficacy minimum value and the second value is futility minimum value.
- **grid.max**: A numeric value or vector (for asymmetric boundary) indicating the upper bound of the grid for screening. For asymmetric boundary, the first value is efficacy maximum value and the second value is futility maximum value.
- **Boundary.type**: A text indicating what type of boundary used. Default is "Symmetric"

Value

A list including the next cutoff value for evaluation next.cutoff and a list of predictions for screening grid.

Author(s)

Ziyan Wang

References

Examples

```r
x = c(7.123968, 6.449631, 1.984406, 3.507463, 4.972510, 2.925768, 5.816682, 4.367796, 7.349160, 1.113648)
y.t1E = c(0.0396, 0.0450, 0.5116, 0.2172, 0.1840, 0.3058, 0.0592, 0.1384, 0.0296, 0.7936)
grid.min=1
grid.max=8
GP.res=GP.optim(x=x, y.t1E=y.t1E, errorrate = 0.1, grid.min = grid.min, grid.max = grid.max)
GP.res$next.cutoff
```

```r
x = data.frame(matrix(c(0.9563408, 0.006295626, 0.9669739, 0.014395030, 0.9959410, 0.034858339, 0.9635357, 0.048435579, 0.9794314, 0.021659226, 0.9552201, 0.018442535, 0.9655749, 0.035281833, 0.9837123, 0.010656442, 0.9974910, 0.047741842, 0.9989172, 0.012982826), byrow=TRUE, ncol = 2))
y.t1E = c(0.3044, 0.2938, 0.2573, 0.4780, 0.2923, 0.3733, 0.4263, 0.1962, 0.2941, 0.1131)
y.pow = c(0.8300, 0.8239, 0.7102, 0.7291, 0.8205, 0.7984, 0.7709, 0.8418, 0.6359, 0.5609
ESS = data.frame(matrix(c(594.672, 560.580, 596.148, 566.328, 597.840, 590.124, 590.052, 544.800, 597.024, 574.716, 593.952, 554.580, 593.676, 554.400, 598.500, 583.896, 595.740, 590.520, 599.580, 598.644),byrow=TRUE,ncol=2))
grid.min=c(0.95,0)
grid.max=c(1,0.05)
GP.res_asy=GP.optim(x=x, y.t1E=y.t1E, y.pow=y.pow, ESS=ESS, errorrate = 0.1, grid.min = grid.min, grid.max = grid.max, Boundary.type="Asymmetric")
GP.res_asy$next.cutoff
```
Description

This function calculates the posterior probability of each active treatment arm better than control using betabinomial model.

Usage

ibetabinomial.post(n, y, pi.star = 0.5, pess = 2)

Arguments

- **n**: A vector of treated patients for each arm (The first element is for control).
- **y**: A vector of treated patient outcomes for each arm (The first element is for control).
- **pi.star**: The prior response probability. The default is 0.5.
- **pess**: The effective sample size of beta prior. The default is 2.

Value

A vector posterior probability of each active treatment arm better than control.

Author(s)

Ziyan Wang

Examples

```r
n <- c(20,20,20,20)
y <- c(12,12,12,6)
ibetabinomial.post(n, y, pi.star = 0.5, pess = 2)
# [1] 0.5000000 0.5000000 0.0308018
```

Description

This function initialises the inner parameter used in simulate.trial function.

Usage

Initialize.trial.parameter(response.probs, ns)

Arguments

- **response.probs**: A vector of response probability of each arm.
- **ns**: A vector of accumulated number of patient at each stage.
Value

A list of initialised parameters including the number of arms for this trial 'K', the number of arm active 'armleft', the index of treatment arm 'treatmentindex', the vector of total number of patients allocated to each arm 'n' the number of total number of patients survived for each arm 'y1', the matrix for true response probability of each arm at each stage 'groupwise.response.probs' which is required for the time trend study, the vector of randomisation probability for each arm 'randomprob', the array of arm assignment for each patient 'z', the array of outcome for each patient 'y', the array of the stage index for each patient 'group_indicator', the matrix of the probability of each arm to be the best at each stage 'post.prob.best.mat'.

Author(s)

Ziyan Wang

Examples

Initializetrialparameter(response.probs = c(0.4,0.6), ns = c(15,30,45,60,75,90))

#$K
#[1] 2

#$armleft
#[1] 2

#$treatmentindex
#[1] 1

#$n
#[1] 0 0

#$y1
#[1] 0 0

#$groupwise.response.probs
# [,1] [,2]
# [1,] 0.4 0.6
# [2,] 0.4 0.6
# [3,] 0.4 0.6
# [4,] 0.4 0.6
# [5,] 0.4 0.6
# [6,] 0.4 0.6

#$randomprob
# 1 2
# [1,] 0.5 0.5

#$z
#numeric(0)

#$y
#numeric(0)
intbias

Description
This function estimates the mean bias of treatment - stage interaction effect

Usage
intbias(res)

Arguments
res A list of output matrix of a number of trial replicates

Value
A matrix of mean treatment - stage interaction effect bias

Examples
## Not run: intbias(res)

Meanfunc

Description
This function reads in the output matrix of a number of trial replicates to calculate mean treatment effect estimate.

Usage
Meanfunc(res)
modelinf.fun

Arguments

res A list of output matrix of a number of trial replicates

Value

Mean treatment effect estimates of each treatment arm

Examples

## Not run: Meanfunc(res)

---

modelinf.fun modelinf

Description

This function summarize the input parameters describing the model for analysis and transfer them into a list

Usage

modelinf.fun(
  model = "tlr",
  ibb.inf = list(pi.star = 0.5, pess = 2, betabinomialmodel = ibetabinomial.post),
  tlr.inf = list(beta0_prior_mu = 0, beta1_prior_mu = 0, beta0_prior_sigma = 2.5,
                  beta1_prior_sigma = 2.5, beta0_df = 7, betal_df = 7, reg.inf = "main", variable.inf =
                  "Fixeffect"
)

Arguments

model The statistical model. ibb: betabinomial model / tlr: logistic model
ibb.inf The list of information for betabinomial model including: betabinomialmodel: The betabinomial model, pi.star: prior response rate, pess: prior effective sample size
tlr.inf The list of information for logistic model including: The mean (mu), variance (sigma), degree of freedom (df) of the intercept and the main effect of the linear terms in logistic model. reg.inf: The type of linear function in logistic model. variable.inf: Fixeffect/Mixeffect. Indicating whether a mix effect model is used (for time trend effect modelling)

Value

A list of model information including model, ibb.inf and tlr.inf

Author(s)

Ziyan Wang
Examples

```r
modelinf.fun(model = "tlr",
             ibb.inf = list(pi.star = 0.5,
                            pess = 2,
                            betabinomialmodel = ibetabinomial.post),
             tlr.inf = list(beta0_prior_mu = 0,
                            beta1_prior_mu = 0,
                            beta0_prior_sigma = 2.5,
                            beta1_prior_sigma = 2.5,
                            beta0_df = 7,
                            beta1_df = 7,
                            reg.inf = "main",
                            variable.inf = "Fixeffect"
            )
)
```

Description

This function reads in the output matrix of a number of trial replicates to calculate mean estimate of total number of patients allocated to each arm

Usage

```r
Nfunc(res)
```

Arguments

- `res` A list of output matrix of a number of trial replicates

Value

The mean estimate of total number of patients allocated to each arm

Examples

```r
## Not run: Nfunc(res)
```
Description

Operation characteristic table for alternative scenario using main + continuous stage model. The time trend pattern is step. The strength of time trend is 0.1 equally for all arm. The effect of time trend on true response probability is multiplicative.

Usage

OPC_alt

Format

A data frame with 3 rows and 16 variables:

Type.I.Error.or.Power  Power
Bias.1  Treatment effect bias for treatment 1
Bias.2  Treatment effect bias for treatment 2
Bias.3  Treatment effect bias for treatment 3
rMSE.1  Rooted mean squared error for treatment 1
rMSE.2  Rooted mean squared error for treatment 2
rMSE.3  Rooted mean squared error for treatment 3
N.per.arm.1  Mean total number of patient allocated to control
N.per.arm.2  Mean total number of patient allocated to treatment 1
N.per.arm.3  Mean total number of patient allocated to treatment 2
N.per.arm.4  Mean total number of patient allocated to treatment 3
Survive.per.arm.1  Mean total number of patient survived when using control
Survive.per.arm.2  Mean total number of patient survived when using treatment 1
Survive.per.arm.3  Mean total number of patient survived when using treatment 2
Survive.per.arm.4  Mean total number of patient survived when using treatment 3
N  Mean total number of patient in a trial
Operation characteristic table for null scenario

Description

Operation characteristic table for null scenario using main and main + continuous stage model. The main effect model was run for a null scenario with and without time trend. The time trend pattern is step. The strength of time trend is 0.1 equally for all arm. The effect of time trend on true response probability is multiplicative.

Usage

OPC_null

Format

A data frame with 3 rows and 16 variables:

- Type.I.Error.or.Power: Family wise error rate
- Bias.1: Treatment effect bias for treatment 1
- Bias.2: Treatment effect bias for treatment 2
- Bias.3: Treatment effect bias for treatment 3
- rMSE.1: Rooted mean squared error for treatment 1
- rMSE.2: Rooted mean squared error for treatment 2
- rMSE.3: Rooted mean squared error for treatment 3
- N.per.arm.1: Mean total number of patient allocated to control
- N.per.arm.2: Mean total number of patient allocated to treatment 1
- N.per.arm.3: Mean total number of patient allocated to treatment 2
- N.per.arm.4: Mean total number of patient allocated to treatment 3
- Survive.per.arm.1: Mean total number of patient allocated to control
- Survive.per.arm.2: Mean total number of patient survived when using treatment 1
- Survive.per.arm.3: Mean total number of patient survived when using treatment 2
- Survive.per.arm.4: Mean total number of patient survived when using treatment 3
- N: Mean total number of patient in a trial
**OPC_Trial.simulation**  
*Operation characteristic table for Trial.simulation() null scenario*

**Description**

Operation characteristic table for null scenario using main model.

**Usage**

OPC_Trial.simulation

**Format**

A data frame with 1 rows and 8 variables:

- **Type.I.Error.or.Power**  
  Power
- **Bias**  
  Treatment effect bias for treatment 1
- **rMSE**  
  Rooted mean squared error for treatment effect 1
- **N.per.arm.1**  
  Mean total number of patient allocated to control
- **N.per.arm.2**  
  Mean total number of patient allocated to treatment 1
- **Survive.per.arm.1**  
  Mean total number of patient allocated to control
- **Survive.per.arm.2**  
  Mean total number of patient survived when using treatment 1
- **N**  
  Mean total number of patient in a trial

**optimdata_asy**  
*A list of data from Gaussian process and trial simulation for asymmetric cutoff screening.*

**Description**

A list of data from Gaussian process and trial simulation for asymmetric cutoff screening.

**Usage**

optimdata_asy
Format

A list with four element:

- `next.cutoff` The cutoff value for the next evaluation
- `prediction` A list of values from Gaussian process model
- `ESS` A two column twenty five rows matrix with the effective sample size for each cutoff pair under both null (first column) and alternative (second column) scenario
- `testeddata` A data frame containing each tested cutoff pair (column two and three for efficacy and futility, respectively), their FWER under null (column one) and conjunctive power under alternative (column four)

Description

A list of data from Gaussian process for symmetric cutoff screening.

Usage

`optimdata_sym`

Format

A list with two element:

- `next.cutoff` The cutoff value for the next evaluation
- `prediction` A list of values from Gaussian process model
- `tpIE` A vector of type I error rate data
- `cutoff` A vector of cutoff data

Description

This function initializes the output matrix including all evaluation metrics based on input information.

Usage

`OutputStats.initialising(variable.inf, reg.inf, ns, K)`
Arguments

variable.inf  The parameter information in the model
reg.inf  The model information. For the fixed effect model, the input of reg.inf can be main, main + stage_continuous, main * stage_continuous, main + stage_discrete, main * stage_discrete. For the mixed effect model, the reg.inf is invalid.
ns  A vector of accumulated number of patient at each stage
K  Total number of arm including control

Value

The empty output matrix including different evaluation metrics.

Author(s)

Ziyan Wang

Examples

OutputStats.initialising(
    variable.inf = "Fixeffect",
    reg.inf = "main",
    ns = c(15, 30, 45, 60, 75),
    K = 2)

perHtypeerror_marginalpowerfunc

Description

This function reads in the output matrix of a number of trial replicates to calculate the error rate

Usage

perHtypeerror_marginalpowerfunc(res)

Arguments

res  A list of output matrix of a number of trial replicates

Value

"per-hypothesis" Type I error rate or power (marginal power) for each treatment - control comparison

Author(s)

Ziyan Wang
Examples

```r
## Not run: perHtypeError_powerfunc(res)
```
Value

A list of input randomisation information

Author(s)

Ziyan Wang

Examples

Randomisation.inf(Random.inf = list(
  Fixratio = FALSE,
  Fixratiocontrol = NA,
  BARmethod = "Thall",
  Thall.tuning.inf = list(tuningparameter = "Fixed", fixvalue = 1),
  Trippa.tuning.inf = list(a = NA, b = NA)
))

<table>
<thead>
<tr>
<th>recommendloginformd</th>
<th>Cutoff screening example: the recommended grid value at each time point</th>
</tr>
</thead>
</table>

Description

The recommended grid value at each time point. There are 20 cutoff value explored

Usage

recommendloginformd

Format

A data frame with 20 rows and 1 variables:

recommendloginformd The cutoff value at each time point

<table>
<thead>
<tr>
<th>resultrtostats</th>
<th>resultrtostats</th>
</tr>
</thead>
</table>

Description

This is an inner function of the function resultstantoRfunc
Usage

```r
resultrtostats(
  trteff = NA,
  treatmentindex = NA,
  armleft,
  K,
  group,
  reg.inf,
  fit,
  ns
)
```

Arguments

- `trteff`: Stan posterior samples of treatment effect sample distribution
- `treatmentindex`: A vector of treatment index at the beginning of a trial
- `armleft`: The number of treatment left in the platform (>2)
- `K`: Total number of arms at the beginning
- `group`: The current stage
- `reg.inf`: The information of how much accumulated information will be used
- `fit`: The stan output
- `ns`: A vector of accumulated number of patient at each stage

Value

A list of stan result inference stats:
1. A vector of posterior probability for all treatment arms including dropped and active treatment arm
2. The mean treatment effect estimate of each treatment compared to control
3. The variance of treatment effect estimate of each treatment compared to control
4. A vector including Posterior probability of each active treatment arm better than control

Author(s)

Ziyan Wang

Examples

```r
## Not run: resultrtostats(trteff = NA, treatmentindex = NA, armleft, K, group, reg.inf, fit, ns)
```
Description

The inner function of function `resultstantoRfunc.rand`

Usage

```r
resultstostats.rand(
  trteff = NA,
  treatmentindex = NA,
  armleft,
  K,
  group,
  fit,
  ns
)
```

Arguments

- `trteff`: Stan posterior samples of treatment effect sample distribution
- `treatmentindex`: A vector of treatment index at the beginning of a trial
- `armleft`: The number of treatment left in the platform (>2)
- `K`: Total number of arms at the beginning
- `group`: The current stage
- `fit`: The stan output
- `ns`: A vector of accumulated number of patient at each stage

Value

A list of stan result inference stats:
- 1: A vector of posterior probability for all treatment arms including dropped and active treatment arm
- 4: The mean treatment effect estimate of each treatment compared to control
- 5: The variance of treatment effect estimate of each treatment compared to control
- `post.prob.btcontrol`: a vector including Posterior probability of each active treatment arm better than control

Author(s)

Ziyan Wang

Examples

```r
## Not run: resultstostats.rand(trteff = NA, treatmentindex = NA, armleft, K, group, fit, ns)
```
resultstantoRfunc

Description

This function summarise the fix effect stan output data to and transform them to be readable.

Usage

resultstantoRfunc(group, reg.inf, variable.inf, fit, armleft, treatmentindex, K, ns)

Arguments

group           The current stage
reg.inf         The information of how much accumulated information will be used
variable.inf    The information of whether to use random effect model or fix effect model.
fit             The stan output
armleft         The number of treatment left in the platform (>2)
treatmentindex  A vector of treatment index at the beginning of a trial
K               Total number of arms at the beginning
ns              A vector of accumulated number of patient at each stage

Value

A list of stan result inference stats1: A vector of posterior probability for all treatment arms including dropped and active treatment arm stats4: The mean treatment effect estimate of each treatment compared to control stats5: The variance of treatment effect estimate of each treatment compared to control post.prob.bcontrol: a vector including Posterior probability of each active treatment arm better than control stats6: A vector of mean stage (time trend) effect estimate stats7: A vector of mean treatment - stage (time trend) interaction effect estimate sampefftotal: The posterior samples of response probability of each active arm on logit scale. This can be transformed to probit scale by using inv.logit() function.

Author(s)

Ziyan Wang
resultstantoRfunc.rand

Description

This function summarise the mix effect stan output data to and transform them to be readable.

Usage

resultstantoRfunc.rand(group, fit, armleft, treatmentindex, K, ns)

Arguments

group The current stage
fit The stan output
armleft The number of treatment left in the platform (>2)
treatmentindex A vector of treatment index at the beginning of a trial
K Total number of arms at the beginning
ns A vector of accumulated number of patient at each stage

Value

A list of stan result inference stats1: A vector of posterior probability for all treatment arms including dropped and active treatment arm stats4: The mean treatment effect estimate of each treatment compared to control stats5: The variance of treatment effect estimate of each treatment compared to control post.prob.btc:control: a vector including Posterior probability of each active treatment arm better than control stats6: A vector of mean stage (time trend) effect estimate stats7: A vector of mean treatment - stage (time trend) interaction effect estimate sampfeftotal: The posterior samples of response probability of each active arm on logit scale. This can be transformed to probit scale by using inv.logit() function.

Author(s)

Ziyan Wang

Examples

## Not run: resultstantoRfunc.rand(group, fit, armleft, treatmentindex, K, ns)
Save.resulttoRDatafile

Description

This function generates the name of file for output table and dataset

Usage

Save.resulttoRDatafile(input.info = list(response.probs = c(0.4, 0.4), ns = c(30, 60, 90, 120, 150), max.ar = 0.75, rand.type = "Urn", max.deviation = 3, model.inf = list(model = "tlr", ibb.inf = list(betabinomialmodel = ibetabinomial.post), tlr.inf = list(beta0_prior_mu = 0, beta1_prior_mu = 0, beta0_prior_sigma = 2.5, beta1_prior_sigma = 2.5, beta0_df = 7, beta1_df = 7, reg.inf = "main", variable.inf = "Fixeffect")), Stopbound.inf = Stopboundinf(Stop.type = "Early-Pocock", Boundary.type = "Symmetric", cutoff = c(0.99, 0.01)), Random.inf = list(Fixratio = FALSE, Fixratiocontrol = NA, BARmethod = "Thall", Thall.tuning.inf = list(tuningparameter = "Fixed", fixvalue = 1)), trend.inf = list(trend.type = "step", trend.effect = c(0, 0), trend_add_or_multip = "mult")))

Arguments

input.info A list of input information require for trial simulation

Value

A list of name for table and dataset

Author(s)

Ziyan Wang

Examples

Save.resulttoRDatafile(input.info = list(response.probs = c(0.4, 0.4), ns = c(30, 60, 90, 120, 150), max.ar = 0.75, rand.type = "Urn", max.deviation = 3, model.inf = list(model = "tlr", ibb.inf = list(betabinomialmodel = ibetabinomial.post), tlr.inf = list(beta0_prior_mu = 0, beta1_prior_mu = 0, beta0_prior_sigma = 2.5, beta1_prior_sigma = 2.5, beta0_df = 7, beta1_df = 7, reg.inf = "main", variable.inf = "Fixeffect")), Stopbound.inf = Stopboundinf(Stop.type = "Early-Pocock", Boundary.type = "Symmetric", cutoff = c(0.99, 0.01)), Random.inf = list(Fixratio = FALSE, Fixratiocontrol = NA, BARmethod = "Thall", Thall.tuning.inf = list(tuningparameter = "Fixed", fixvalue = 1)), trend.inf = list(trend.type = "step", trend.effect = c(0, 0), trend_add_or_multip = "mult")))
Description

This function simulates a MAMS trial applying adaptive methods where the time trend effect can be studied.

Usage

```r
simulatetrial(
  ii,
  response.probs = c(0.4, 0.4),
  ns = c(30, 60, 90, 120, 150),
  test.type = "Twoside",
  max.ar = 0.75,
  rand.algo = "Urn",
  max.deviation = 3,
  model.inf = list(model = "tlr", ibb.inf = list(pi.star = 0.5, pess = 2,
pi.star = 0.5,
pess = 2,
betabinomialmodel = ibetabinomial.post
)
),
  tlr.inf = list(
    beta0_prior_mu = 0,
    beta1_prior_mu = 0,
    beta0_prior_sigma = 2.5,
    beta1_prior_sigma = 2.5,
    beta0_df = 7,
    beta1_df = 7,
    reg.inf = "main",
    variable.inf = "Fixefect"
  ),
  Stop.type = "Early-Pocock",
  Boundary.type = "Symmetric",
  Random.inf = list(
    Fixratio = FALSE,
    Fixratiocontrol = NA,
    BARmethod = "Thall",
    Thall.tuning.inf = list(tuningparameter = "Fixed", fixvalue = 1)
  ),
  trend.inf = list(
    trend.type = "step",
    trend.effect = c(0, 0),
    trend_add_or_multip = "mult"
  )
)
)
simulatetrial

```
betabinomialmodel = ibetabinomial.post, tlr.inf = list(bet0_prior_mu = 0, 
bet1_prior_mu = 0, beta0_prior_sigma = 2.5, beta1_prior_sigma = 2.5, beta0_df = 7, 
bet1_df = 7, reg.inf = "main", variable.inf = "Fixeffect")), 
Stopbound.inf = Stopbound.inf, 
Random.inf = Random.inf, 
trend.inf = trend.inf
)
```

Arguments

- **ii**: Meaning less parameter but required for foreach function in doParallel package
- **response.probs**: A vector of true response probability for each arm. Default `response.probs = c(0.4, 0.4)`.
- **ns**: A vector of accumulated number of patient at each stage. Default is `ns = c(30, 60, 90, 120, 150)`.
- **test.type**: A indicator of whether to use one side test or two side test for each treatment-control comparison.
- **max.ar**: The upper boundary for randomisation ratio for each arm. Default is 0.75 for a two arm trial. The minimum value depends on K where 1 - max.ar <= 1/K
- **rand.algo**: The method of applying patient allocation with a given randomisation probability vector. Default is "Urn".
- **max.deviation**: The tuning parameter for Urn randomisation method. Default is 3.
- **model.inf**: The list of interim data analysis model information for more see `modelinf.fun`
- **Stopbound.inf**: The list of stop boundary information for more see `Stopboundinf`
- **Random.inf**: The list of Adaptive randomisation information for more see `Randomisation.inf`
- **trend.inf**: The list of time trend information

Value

A matrix including all evaluation metrics

Author(s)

Ziyan Wang

Examples

```
set.seed(1)
simulatetrial(response.probs = c(0.4, 0.4), 
 ns = c(30, 60, 90, 120, 150), 
 max.ar = 0.75, 
 test.type = "Twoside", 
 rand.algo = "Urn", 
 max.deviation = 3, 
 model.inf = list( 
 model = "tlr", 
 ibb.inf = list(
```
Sperarmfunc

Description
This function reads in the output matrix of a number of trial replicates to calculate mean total number of survived patients of each arm

Usage
Sperarmfunc(res)

Arguments
res A list of output matrix of a number of trial replicates
Value

The mean total number of survived patients of each arm

Examples

## Not run: Sperarmfunc(res)

```r
stan.logisticmodeltrans
```

Description

This function transform the data in trial simulation to the data required for stan modelling

Usage

```r
stan.logisticmodeltrans(
  z,
  y,
  randomprob,
  group_indicator,
  armleft,
  group,
  variable.inf,
  reg.inf
)
```

Arguments

- `z`: A vector of all treatment index data from the beginning of a trial
- `y`: A vector of all outcome data from the beginning of a trial
- `randomprob`: A named vector of randomisation probability to each arm
- `group_indicator`: A vector for the stage at which each patient was treated
- `armleft`: The number of treatment left in the platform (>2)
- `group`: The current stage
- `variable.inf`: Fixeffect/Mixeffect for logistic model parameter
- `reg.inf`: The information of how much accumulated information will be used
Value

A list of information require for the stan model including: zdropped: The vector of treatment index for each patient whose treatment arm is active at current stage. ydropped: The vector of outcome index for each patient whose treatment arm is active at current stage. Ndropped: The total number of patients that are treated with active treatment arms at current stage. group_indicator_dropped: The vector of stage index for each patient whose treatment arm is active at current stage. zlevel: The active treatment arm index at current stage xdummy: A design matrix transformed from zdropped and group_indicator_dropped for modelling

Author(s)

Ziyan Wang

Examples

```r
stan.logisticmodeltrans(
  z = c(1,2,1,2,1,2,1,2),
  y = c(0,0,0,0,1,1,1,1),
  randomprob = matrix(c(0.5, 0.5), ncol = 2, dimnames = list(c("Stage1"), c("1", "2"))),
  group_indicator = c(1,1,1,1,1,1,1,1),
  armleft = 2,
  group = 1,
  variable.inf = "Fixeffect",
  reg.inf = "main")
```

Description

This function summaries and checks stopping boundary information.

Usage

```r
Stopboundinf(
  Stop.type = "Early-Pocock",
  Boundary.type = "Symmetric",
  cutoff = c(0.9928, 0.0072)
)
```

Arguments

**Stop.type** The type of stopping boundary should be "Early-Pocock", "Early-OBF" and "Noearly". Default is "Early-Pocock" which is the Pocock boundary with early stopping.
**Boundary.type**  Whether the futility boundary and the efficacy boundary are the same conservative. Default is "Symmetric" which means they are as conservative as each other. Boundary.type = "Asymmetric" means that the efficacy boundary and the futility boundary are not as conservative as each other.

**cutoff**  = c(cutoff1, cutoff2). A numerical vector of cutoff value for each boundary. The first element is the efficacy boundary cutoff. The second element is the futility boundary cutoff Pr(theta_1 > theta_0|D_n) > cutoff1. Should input the cutoff1 for efficacy boundary as the first element Pr(theta_1 < theta_0|D_n) < cutoff2. Should input the cutoff2 for futility boundary as the first element.

**Value**

The list of information required for boundary construction function 'Stopbound.inf'

**Author(s)**

Ziyan Wang

**Examples**

Stop.type = "Early-Pocock"  #(Pocock boundary is a flat boundary across time)
Boundary.type = "Symmetric"
cutoff = c(0.9928, 0.0072)

Stopbound.inf = Stopboundinf(Stop.type, Boundary.type, cutoff)
#Stopbound.inf
#$Stop.type
# [1] "Early-Pocock"
#$Boundary.type
# [1] "Symmetric"
#$cutoff
# [1] 0.9928 0.0072

**Description**

This function makes a decision on whether any active arm should be dropped based on posterior probability and return the vector of decision on each arm, the vector of active arms index and the number of arms left for further study.
Usage

testing_and_armdropping(
  K,
  armleft,
  post.prob.btcontrol,
  group,
  cutoffeff,
  cutoffful,
  treatmentindex,
  test.type
)

Arguments

K                   A numeric value indicating the total number of arm at the beginning of trial including both control and treatment.
armleft             A numeric vector indicating the number of active arms before this interim analysis;
post.prob.btcontrol A numeric vector of posterior probability of each treatment arm better than control
group               A numeric value. The current stage index.
cutoffeff           A numeric vector of the cutoff value at each stage for efficacy boundary.
cutoffful           A numeric vector of the cutoff value at each stage for futility boundary.
treatmentindex     A numeric vector of the current active treatment arm index
test.type           A character indicating which hypothesis testing we are use. "Oneside": H_0: \pi_k \leq \pi_0; H_0: \pi_k \geq \pi_0 "Twoside": H_0: \pi_k \equiv \pi_0; H_0: \pi_k \neq \pi_0

Value

A list of information including armleft: the number of active arms after this interim analysis; treatmentindex: the index vector of active arm after this interim analysis; stats3: the vector of conclusion on whether null hypothesis is rejected

Examples

testing_and_armdropping(
  K = 4,
  armleft = 4,
  post.prob.btcontrol = c(0.5, 0.99, 0.02),
  group = 3,
  cutoffeff = c(1, 0.99, 0.975, 0.96, 0.95),
  cutoffful = c(0, 0.01, 0.025, 0.04, 0.05),
  treatmentindex = c(1, 2, 3),
  test.type = "Oneside")
testing_and_armdropping(
K = 4,
armleft = 4,
post.prob.btcontrol = c(0.5, 0.99, 0.02),
group = 3,
cutoffeff = c(1, 0.99, 0.975, 0.96, 0.95),
cutoffful = c(0, 0.01, 0.025, 0.04, 0.05),
treatmentindex = c(1, 2, 3),
test.type = "Twoside")

Timetrend.fun

Description
This function generate the time trend function based on trend information. This function also check
the validity of the input time trend information.

Usage
Timetrend.fun(trend.inf)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>trend.inf</td>
<td>The list of information for time trend effect including 'trend.type', 'trend.effect', 'trend_add_or_mult'. 'trend.type' is the shape of time trend. Default is &quot;step&quot;. Other types are &quot;linear&quot;, &quot;inverse.U.linear&quot;, &quot;plateau&quot;. 'trend.effect' the vector of the strength of time trend for each arm. The first element is for the control arm. The value of time trend is the gap between the start of the trial and the end of the trial. The change between each interim or each patient is calculated in the function. For example, for linear time trend with trend.effect = c(0.2, 0.2). The trend effect increment in control group for patient $i$ is $0.2(i-1)/(N_{max}-1)$, for stage $j$ is $0.2(j-1)/(length(ns)-1)$. 'trend_add_or_mult' the pattern of time trend affecting the true response probability. Default is &quot;mult&quot;.</td>
</tr>
</tbody>
</table>

Value
A list containing the time trend function according to input trend.type variable, and a indicator of whether there is a time trend in data generation based on input trend information

Author(s)
Ziyan Wang
Examples

```r
Timetrend.fun(trend.inf = list(
    trend.type = "step",
    trend.effect = c(0, 0),
    trend_add_or_multip = "mult"
))
```

Description

This function simulates and does final analysis of a trial with one scenario. The time cost of this function depend on the cpu cores of the user’s computer.

Usage

```r
Trial.simulation(
    ntrials = 5000,
    trial.fun = simulatetrial,
    input.info = list(response.probs = c(0.4, 0.4), ns = c(30, 60, 90, 120, 150), max.ar = 0.75, test.type = "Twoside", rand.algo = "Urn", max.deviation = 3, model.inf = list(model = "tlr", ibb.inf = list(pi.star = 0.5, pess = 2, betabinomialmodel = ibetabinomial.post), tlr.inf = list(beta0_prior_mu = 0, beta1_prior_mu = 0, beta0_prior_sigma = 2.5, beta1_prior_sigma = 2.5, beta0_df = 7, beta1_df = 7, reg.inf = "main", variable.inf = "Fixeffect"), stopbound.inf = stopboundinf(Stop.type = "Early-Pocock", Boundary.type = "Symmetric", cutoff = c(0.99, 0.01)), Random.inf = list(Fixratio = TRUE, Fixratiocontrol = 1, BARmethod = NA, thall.tuning.inf = NA, trippa.tuning.inf = NA), trend.inf = list(trend.type = "step", trend.effect = c(0, 0), trend_add_or_multip = "mult")),
    cl = 2
)
```

Arguments

- **ntrials**: A numeric variable indicating how many trial replicates you want to run. Default is 5000.
- **trial.fun**: The function of trial simulation for more see `simulatetrial`
- **input.info**: A list of input information including all information required for trial simulation.
- **cl**: A numeric variable indicating how many cores you want to use in parallel programming.

Value

A list of output including the final output of each trial replicates called `result` The analysis result table of the specific trial called `OPC` and the file name for saving these output on the computer.
Author(s)

Ziyan Wang

Examples

```r
set.seed(1)
Trial.simulation(ntrials = 2, cl = 2)
```

---

trtbias

Description

This function estimates the mean bias of treatment effect

Usage

```
trtbias(res, trueeffect)
```

Arguments

- `res` A list of output matrix of a number of trial replicates
- `trueeffect` A vector of true treatment effect in each scenario

Value

A matrix of mean treatment effect bias

Examples

```r
## Not run: trtbias(res, trueeffect)
```

---

trteffect

Description

This function estimates the mean treatment effect bias and its rooted mean squared error

Usage

```
trteffect(res, trueeff)
```

Arguments

- `res` A list of output matrix of a number of trial replicates
- `trueeff` A vector of true treatment effect in each scenario

Examples

```r
## Not run: trteffect(res, trueeff)
```
Value

A vector of mean treatment effect bias and its rooted mean squared error

Examples

## Not run: trteffect(res, trueeff)

---

### Description

This function reads in the output matrix of a number of trial replicates to calculate the variance of treatment effect estimate.

### Usage

`varfunc(res)`

### Arguments

- `res` A list of output matrix of a number of trial replicates

### Value

The variance of Treatment effect estimates of each treatment arm

### Examples

## Not run: varfunc(res)
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