Package: ASSISTant (via r-universe)

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Type Package Title Adaptive Subgroup Selection in Group Sequential Trials Version 1.4.3 Date 2019-11-21 VignetteBuilder knitr URL https://github.com/bnaras/ASSISTant BugReports https://github.com/bnaras/ASSISTant/issues Description Clinical trial design for subgroup selection in three-stage group sequential trial. Includes facilities for design, exploration and analysis of such trials. An implementation of the initial DEFUSE-3 trial is also provided as a vignette. License GPL(>=2) **Encoding** UTF-8 **Roxygen** list(markdown = TRUE) RoxygenNote 6.1.1 Imports R6, mvtnorm, knitr, magrittr, dplyr Suggests rmarkdown Repository https://bnaras.r-universe.dev RemoteUrl https://github.com/bnaras/assistant RemoteRef HEAD RemoteSha b6598c72ffdc0433b4130f498e55d819f44cbffd

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ASSISTant

Three stage group sequential adaptive design with subgroup selection

Description

ASSISTant is a package that implements a three-stage adaptive clinical trial design with provision for subgroup selection where the treatment may be effective. The main design object is an R6 class that can be instantiated and manipulated to obtain the operating characteristics. A vignette is provided showing the use of this package for designing the DEFUSE-3 trial, described in the paper by Lai, Lavori and Liao. The package contains everything necessary to reproduce the results of the paper.

References

Adaptive Choice of Patient Subgroup for Comparing Two Treatments by Tze Leung Lai and Philip W. Lavori and Olivia Yueh-Wen Liao. Contemporary Clinical Trials, Vol. 39, No. 2, pp 191-200 (2014). http://www.sciencedirect.com/science/article/pii/S1551714414001311

Adaptive design of confirmatory trials: Advances and challenges, http://www.sciencedirect. com/science/article/pii/S1551714415300239 by Tze Leung Lai and Philip W. Lavori and Ka Wai Tsang. Contemporary Clinical Trials, Vol. 45, Part A, pp 93-102 (2015).

ASSISTDesign

A class to encapsulate the adaptive clinical trial design of Lai, Lavori and Liao

Description

ASSISTDesign objects are used to design, simulate and analyze adaptive group sequential clinical trial with three stages.

ASSISTDesign

Usage

design <- ASSISTDesign\$new(trialParameters, designParameters)</pre>

Format

An R6Class generator object

Methods

- ASSISTDesign\$new(designParameters, trialParameters, discreteData = FALSE, boundaries) Create a new ASSISTDesign instance object using the parameters specified. If discreteData is TRUE use a discrete distribution for the Rankin scores and designParameters must contain the appropriate distributions to sample from. If boundaries is specified, it used.
- getDesignParameters,getTrialParameters, getBoundaries Accessor methods for (obvious)
 object fields
- setBoundaries Modifier method for boundaries a named vector of double values with names btilde, b, and c, in that order
- print() Print the object in a human readable form
- computeCriticalValues() Compute the critical boundary values \tilde{b} , b and c for futility, efficacy and final efficacy decisions; saved in field boundaries
- explore(numberOfSimulations = 5000, rngSeed = 12345) Explore the design using the specified number of simulations and random number seed. There are a number of further parameters. By default trueParameters = self\$getDesignParameters() as would be the case for a Type I error calculation. If changed, would yield power. Also recordStats = TRUE/FALSE, showProgress = TRUE/FALSE, saveRawData = TRUE/FALSE control recording statistics, raw data saves, display of progress. Fixed sample size (fixedSampleSize = TRUE/FALSE) can be specified to ensure that patients lost after a futile overall look are not made up. Returns a list of results
- analyze(trialExploration) Analyze the design given the trialExploration which is the result of a call to explore to simulate the design. Return a list of summary quantities
- summary(analysis) Print the operating characteristics of the design, using the analysis result from the analyze call

References

Adaptive Choice of Patient Subgroup for Comparing Two Treatments by Tze Leung Lai and Philip W. Lavori and Olivia Yueh-Wen Liao. Contemporary Clinical Trials, Vol. 39, No. 2, pp 191-200 (2014). doi:10.1016/j.cct.2014.09.001g

See Also

LLL. SETTINGS for an explanation of trial parameters

Examples

```
## Not run:
data(LLL.SETTINGS)
prevalence <- LLL.SETTINGS$prevalences$table1</pre>
scenario <- LLL.SETTINGS$scenarios$S0</pre>
designParameters <- list(prevalence = prevalence,</pre>
                        mean = scenario$mean,
                        sd = scenario$sd)
designA <- ASSISTDesign$new(trialParameters = LLL.SETTINGS$trialParameters,</pre>
                              designParameters = designParameters)
print(designA)
## A realistic design uses 5000 simulations or more!
result <- designA$explore(showProgress = interactive())</pre>
analysis <- designA$analyze(result)</pre>
designA$summary(analysis)
## End(Not run)
## For full examples, try:
## browseURL(system.file("full_doc/ASSISTant.html", package="ASSISTant"))
```

ASSISTDesignB

A fixed sample design to compare against the adaptive clinical trial design of Lai, Lavori and Liao.

Description

ASSISTDesignB objects are used to design a trial with certain characteristics provided in the object instantiation method. This design differs from ASSISTDesign in only how it computes the critical boundaries, how it performs the interim look, and what quantities are computed in a trial run.

Usage

design <- ASSISTDesignB\$new(trialParameters, designParameters, discreteData)</p>

Format

An R6Class generator object

Methods

- ASSISTDesignB\$new(designParameters, trialParameters, discreteData = FALSE, boundaries) Create a new ASSISTDesign instance object using the parameters specified. If discreteData is TRUE use a discrete distribution for the Rankin scores and designParameters must contain the appropriate distributions to sample from. If boundaries is specified, it is used
- getDesignParameters,getTrialParameters, getBoundaries Accessor methods for (obvious)
 object slots
- setBoundaries Modifier method for boundaries a named vector of double values with names btilde, b, and c, in that order

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ASSISTDesignB

print() Print the object in a human readable form

computeCriticalValues() Compute the critical boundary value c_{α}

- explore(numberOfSimulations = 5000, rngSeed = 12345) Explore the design using the specified number of simulations and random number seed. There are further parameters. By default trueParameters = self\$getDesignParameters() as would be the case for a Type I error calculation. If changed, would yield power. Also showProgress = TRUE/FALSE, saveRawData = TRUE/FALSE control raw data saves and display of progress. Returns a list of results
- analyze(trialExploration) Analyze the design given the trialExploration which is the result of a call to explore to simulate the design. Return a list of summary quantities
- summary(analysis) Print the operating characteristics of the design, using the analysis result from the analyze call

References

Adaptive Choice of Patient Subgroup for Comparing Two Treatments by Tze Leung Lai and Philip W. Lavori and Olivia Yueh-Wen Liao. Contemporary Clinical Trials, Vol. 39, No. 2, pp 191-200 (2014). doi:10.1016/j.cct.2014.09.001g

See Also

ASSISTDesign which is a superclass of this object

Examples

```
## Not run:
data(LLL.SETTINGS)
prevalence <- LLL.SETTINGS$prevalences$table1</pre>
scenario <- LLL.SETTINGS$scenarios$S0</pre>
designParameters <- list(prevalence = prevalence,</pre>
                        mean = scenario$mean,
                        sd = scenario$sd)
designB <- ASSISTDesignB$new(trialParameters = LLL.SETTINGS$trialParameters,</pre>
                             designParameters = designParameters)
print(designB)
## A realistic design uses 5000 simulations or more!
result <- designB$explore(showProgress = interactive())</pre>
analysis <- designB$analyze(result)</pre>
designB$summary(analysis)
## End(Not run)
## For full examples, try:
## browseURL(system.file("full_doc/ASSISTant.html", package="ASSISTant"))
```

ASSISTDesignC

A fixed sample RCT design to compare against the adaptive clinical trial design of Lai, Lavori and Liao.

Description

ASSISTDesignC objects are used to design a trial with certain characteristics provided in the object instantiation method. This design differs from ASSISTDesign in only how it computes the critical boundaries, how it performs the interim look, and what quantities are computed in a trial run.

Usage

```
# design <- ASSISTDesignC$new(trialParameters, designParameters)</pre>
```

Format

An R6Class generator object

Methods

- ASSISTDesignC\$new(designParameters, trialParameters, discreteData = FALSE, boundaries) Create a new ASSISTDesign instance object using the parameters specified. If discreteData is TRUE use a discrete distribution for the Rankin scores and designParameters must contain the appropriate distributions to sample from. If 'boundaries is specified, it is used.
- getDesignameters,getTrialParameters, getBoundaries Accessor methods for (obvious) object slots
- setBoundaries Modifier method for boundaries a named vector of double values with names btilde, b, and c, in that order
- print() Print the object in a human readable form
- computeCriticalValues() Compute the critical boundary value c_{α}
- explore(numberOfSimulations = 5000, rngSeed = 12345 Explore the design using the specified number of simulations and random number seed. There are further parameters. By default trueParameters = self\$getDesignParameters() as would be the case for a Type I error calculation. If changed, would yield power. Also showProgress = TRUE/FALSE, saveRawData = TRUE/FALSE control raw data saves and display of progress. Returns a list of results
- analyze(trialExploration) Analyze the design given the trialExploration which is the result of a call to explore to simulate the design. Return a list of summary quantities
- summary(analysis) Print the operating characteristics of the design, using the analysis result from the analyze call

References

Adaptive Choice of Patient Subgroup for Comparing Two Treatments by Tze Leung Lai and Philip W. Lavori and Olivia Yueh-Wen Liao. Contemporary Clinical Trials, Vol. 39, No. 2, pp 191-200 (2014). doi:10.1016/j.cct.2014.09.001g

colNamesForStage

See Also

ASSISTDesignB which is a superclass of this object

Examples

colNamesForStage Return a vector of column names for statistics for a given stage

Description

Return a vector of column names for statistics for a given stage

Usage

```
colNamesForStage(stage, J)
```

Arguments

stage	the trial stage (1 to 3 inclusive).
J	the number of subgroups

Value

a character vector of the column names

computeMeanAndSD

Description

Compute the mean and sd of a discrete Rankin distribution

Usage

```
computeMeanAndSD(probVec = rep(1, 7L), support = 0L:6L)
```

Arguments

probVec	a probability vector of length equal to length of support, default is uniform
support	a vector of support values (default 0:6 for Rankin Scores)

Value

a named vector of mean and sd

computeMHPBoundaries Compute the three modified Haybittle-Peto boundaries

Description

Compute the three modified Haybittle-Peto boundaries

Usage

```
computeMHPBoundaries(prevalence, N, alpha, beta, eps,
futilityOnly = FALSE)
```

Arguments

prevalence	the vector of prevalences between 0 and 1 summing to 1. J , the number of groups, is implicitly the length of this vector and should be at least 2.
Ν	a three-vector of total sample size at each stage
alpha	the type I error
beta	the type II error
eps	the fraction (between 0 and 1) of the type 1 error to spend in the interim stages 1 and 2
futilityOnly	a logical value indicating only the futility boundary is to be computed; default FALSE

Value

a named vector of three values containing \tilde{b} , b, c

References

Adaptive Choice of Patient Subgroup for Comparing Two Treatments by Tze Leung Lai and Philip W. Lavori and Olivia Yueh-Wen Liao. Contemporary Clinical Trials, Vol. 39, No. 2, pp 191-200 (2014). http://www.sciencedirect.com/science/article/pii/S1551714414001311

computeMHPBoundaryITT Compute the three modified Haybittle-Peto boundaries and effect size

Description

Compute the three modified Haybittle-Peto boundaries and effect size

Usage

computeMHPBoundaryITT(prevalence, alpha)

Arguments

prevalence	the vector of prevalences between 0 and 1 summing to 1. J , the number of
	groups, is implicitly the length of this vector and should be at least 2.
alpha	the type I error

Value

a named vector of a single value containing the value for c

References

Adaptive Choice of Patient Subgroup for Comparing Two Treatments by Tze Leung Lai and Philip W. Lavori and Olivia Yueh-Wen Liao. Contemporary Clinical Trials, Vol. 39, No. 2, pp 191-200 (2014). http://www.sciencedirect.com/science/article/pii/S1551714414001311

DEFUSE3Design

Description

DEFUSE3Design is a slight variant of the the adaptive clinical trial design of Lai, Lavori and Liao. Simulation is used to compute the expected maximum sample size and the boundary for early futility is adjusted to account as well.

Usage

design <- DEFUSE3Design\$new(designParameters, trialParameters)</pre>

Format

An R6Class generator object

Methods

- DEFUSE3Design\$new(designParameters, trialParameters, discreteData = FALSE, numberOfSimulations = 5000, Create a new DEFUSE3Design instance object using the parameters specified. If discreteData is TRUE use a discrete distribution for the Rankin scores and designParameters must contain the appropriate distributions to sample from. If boundaries is specified, it is used.
- getDesignParameters,getTrialParameters, getBoundaries Accessor methods for (obvious)
 object slots
- setBoundaries Modifier method for boundaries a named vector of double values with names btilde, b, and c, in that order
- print() Print the object in a human readable form
- adjustCriticalValues(numberOfSimulations, rngSeed, showProgress) Adjust the critical values by performing simulations using the parameters provided
- computeCriticalValues() Compute the critical boundary value c_{α}
- explore(numberOfSimulations = 5000, rngSeed = 12345, trueParameters = self\$getDesignParameters(), record Explore the design using the specified number of simulations and random number seed. trueParameters is by default the same as designParameters as would be the case for a Type I error calculation. If changed, would yield power. Record statistics, save raw data and show progress if so desired. Returns a list of results
- analyze(trialHistory) Analyze the design given the trialHistory which is the result of a call to explore to simulate the design. Return a list of summary quantities
- summary(analysis) Print the operating characteristics of the design, using the analysis result from the analyze call

References

Adaptive design of confirmatory trials: Advances and challenges, 2015 45(Pt A):93-102, by Tze Leung Lai and Philip W. Lavori and Ka Wai Tsang. doi:10.1016/j.cct.2015.06.007

See Also

ASSISTDesign which is a superclass of this object

Examples

```
trialParameters <- list(N = c(200, 340, 476), type1Error = 0.025,
                        eps = 1/2, type2Error = 0.1)
designParameters <- list(</pre>
  nul0 = list(prevalence = rep(1/6, 6), mean = matrix(0, 2, 6),
               sd = matrix(1, 2, 6)),
  alt1 = list(prevalence = rep(1/6, 6), mean = rbind(rep(0, 6),
               c(0.5, 0.4, 0.3, 0, 0, 0)),
               sd = matrix(1, 2, 6)),
  alt2 = list(prevalence = rep(1/6, 6), mean = rbind(rep(0, 6)),
               c(0.5, 0.5, 0, 0, 0, 0)),
               sd = matrix(1, 2, 6)),
  alt3 = list(prevalence = rep(1/6, 6), mean = rbind(rep(0, 6), rep(0.36, 6)),
               sd = matrix(1, 2, 6)),
  alt4 = list(prevalence = rep(1/6, 6), mean = rbind(rep(0, 6), rep(0.30, 6)),
               sd = matrix(1, 2, 6)),
  alt5 = list(prevalence = rep(1/6, 6), mean = rbind(rep(0, 6)),
               c(0.4, 0.3, 0.2, 0, 0, 0)),
               sd = matrix(1, 2, 6)),
  alt6 = list(prevalence = rep(1/6, 6), mean = rbind(rep(0, 6),
               c(0.5, 0.5, 0.3, 0.3, 0.1, 0.1)),
               sd = matrix(1,2, 6)))
## Not run:
## A realistic design uses 5000 simulations or more!
defuse3 <- DEFUSE3Design$new(trialParameters = trialParameters,</pre>
                             numberOfSimulations = 25,
                             designParameters = designParameters$nul0,
                              showProgress = FALSE)
print(defuse3)
result <- defuse3$explore(showProgress = interactive())</pre>
analysis <- defuse3$analyze(result)</pre>
print(defuse3$summary(analysis))
## End(Not run)
## For full examples, try:
## browseURL(system.file("full_doc/defuse3.html", package="ASSISTant"))
```

generateDiscreteData A data generation function using a discrete distribution for Rankin score rather than a normal distribution

Description

A data generation function using a discrete distribution for Rankin score rather than a normal distribution generateDiscreteData(prevalence, N, support = 0L:6L, ctlDist, trtDist)

Arguments

prevalence	a vector of group prevalences (length denoted by J below)
Ν	the sample size to generate
support	the support values of the discrete distribution (length K), default 0:6
ctlDist	a probability vector of length K denoting the Rankin score distribution for con- trol.
trtDist	an K x J probability matrix with each column is the Rankin distribution for the associated group

Value

a three-column data frame of subGroup, trt (0 or 1), and score

Examples

```
# Simulate data from a discrete distribution for the Rankin scores,
# which are typically ordinal integers from 0 to 6 in the following
# simulations. So we define a few scenarios.
library(ASSISTant)
null.uniform <- rep(1, 7L) ## uniform on 7 support points</pre>
hourglass <- c(1, 2, 2, 1, 2, 2, 1)
inverted.hourglass <- c(2, 1, 1, 2, 1, 1, 2)
bottom.heavy <- c(2, 2, 2, 1, 1, 1, 1)
bottom.heavier <- c(3, 3, 2, 2, 1, 1, 1)
top.heavy <- c(1, 1, 1, 1, 2, 2, 2)
top.heavier <- c(1, 1, 1, 2, 2, 3, 3)
ctlDist <- null.uniform
trtDist <- cbind(null.uniform, null.uniform, hourglass, hourglass) ## 4 groups</pre>
generateDiscreteData(prevalence = rep(1, 4), N = 10, ctlDist = ctlDist,
                     trtDist = trtDist) ## default support is 0:6
trtDist <- cbind(bottom.heavy, bottom.heavy, top.heavy)</pre>
generateDiscreteData(prevalence = rep(1, 4), N = 10, ctlDist = ctlDist,
                     trtDist = trtDist)
support <- c(-2, -1, 0, 1, 2) ## Support of distribution</pre>
top.loaded <- c(1, 1, 1, 3, 3) ## Top is heavier
ctl.dist <- c(1, 1, 1, 1, 1) ## null on 5 support points
trt.dist <- cbind(ctl.dist, ctl.dist, top.loaded) ## 3 groups</pre>
generateDiscreteData(prevalence = rep(1, 3), N = 10, support = support,
                     ctlDist = ctl.dist, trtDist = trt.dist)
```

generateNormalData A data generation function along the lines of what was used in the Lai, Lavori, Liao paper. score rather than a normal distribution

Description

A data generation function along the lines of what was used in the Lai, Lavori, Liao paper. score rather than a normal distribution

Usage

generateNormalData(prevalence, N, mean, sd)

Arguments

prevalence	a vector of group prevalences (length denoted by J below)
Ν	the sample size to generate
mean	a 2 x J matrix of means under the null (first row) and alternative for each group
sd	a 2 x J matrix of standard deviations under the null (first row) and alternative for each group

Value

a three-column data frame of subGroup, trt (0 or 1), and score

groupSampleSize	Compute the sample size for any group at a stage assuming a nested
	structure as in the paper.

Description

In the three stage design under consideration, the groups are nested with assumed prevalences and fixed total sample size at each stage. This function returns the sample size for a specified group at a given stage, where the futility stage for the overall group test may be specified along with the chosen subgroup.

Usage

Arguments

prevalence	the vector of prevalence, will be normalized if not already so. The length of this vector implicitly indicates the number of groups J.
Ν	an integer vector of length 3 indicating total sample size at each of the three stages
stage	the stage of the trial
group	the group whose sample size is desired
HJFutileAtStag	e
	is the stage at which overall futility occured. Default NA indicating it did not occur. Also ignored if stage is 1.
chosenGroup	the selected group if HJFutilityAtStage is not NA. Ignored if stage is 1.

Value

the sample size for group

References

Adaptive Choice of Patient Subgroup for Comparing Two Treatments by Tze Leung Lai and Philip W. Lavori and Olivia Yueh-Wen Liao. Contemporary Clinical Trials, Vol. 39, No. 2, pp 191-200 (2014). http://www.sciencedirect.com/science/article/pii/S1551714414001311

LLL.SETTINGS	Design and trial settings used in the Lai, Lavori, Liao paper simula-
	tions

Description

A list of design and trial design settings used for analysis and simulations in the Lai, Lavori, Liao paper displayed in Tables 1 and 2. The elements of the list are the following

trialParameters N the sample size at each of three interim looks, the last being the final one; The length of this also determines the number of interim looks

type1Error the overall type I error

eps the fraction of type I error spent at each interim look

type2Error the type II error desired

- scenarios A list of the 10 settings used in the simulations named S0, S1, ..., S10 as in the paper, each with three elements
 - **mean** a $2 \times J$ matrix of means, the first row for the null setting, the second for the alternative **sd** a $2 \times J$ matrix of standard deviations, the first row for the null setting, the second for the alternative
- **prevalences** A list of two elements with prevalence vectors used in the paper; the lengths of these vectors implicitly define the number of groups.

table1 a vector of equal prevalences for six groups used in table 1

table2 a vector of prevalences used in table 2 of the paper

mHP.b

References

Adaptive Choice of Patient Subgroup for Comparing Two Treatments by Tze Leung Lai and Philip W. Lavori and Olivia Yueh-Wen Liao. Contemporary Clinical Trials, Vol. 39, No. 2, pp 191-200 (2014). http://www.sciencedirect.com/science/article/pii/S1551714414001311

mHP.b	Compute the efficacy boundary (modified Haybittle-Peto) for the first
	two stages

Description

Compute the efficacy boundary (modified Haybittle-Peto) for the first two stages

Usage

mHP.b(prevalence, N, cov.J, mu.prime, Sigma.prime, alpha, btilde, theta)

Arguments

prevalence	the vector of prevalences between 0 and 1 summing to 1. J , the number of groups, is implicitly the length of this vector and should be at least 2.
Ν	a three-vector of total sample size at each stage
cov.J	the 3 x 3 covariance matrix for Z_J at each of the three stages
mu.prime	a list of J mean vectors, each of length $J-1$ representing the conditional means of all the other Z_j given Z_i . This mean does not account for the conditioned value of Z_i and so has to be multiplied by that during use!
Sigma.prime	a list of J covariance matrices, each $J-1$ by $J-1$ representing the conditional covariances all the other Z_j given Z_i
alpha	the amount of type I error to spend
btilde	the futility boundary
theta	the effect size on the probability scale

References

Adaptive Choice of Patient Subgroup for Comparing Two Treatments by Tze Leung Lai and Philip W. Lavori and Olivia Yueh-Wen Liao. Contemporary Clinical Trials, Vol. 39, No. 2, pp 191-200 (2014). http://www.sciencedirect.com/science/article/pii/S1551714414001311

mHP.btilde

Description

The futility boundary \tilde{b} is computed by solving (under the alternative)

Usage

mHP.btilde(beta, cov.J)

Arguments

beta	the type II error
cov.J	the 3 x 3 covariance matrix

Details

 $P(\tilde{Z}_J^1 \le \tilde{b}or\tilde{Z}_J^2 \le \tilde{b}) = \epsilon\beta$

where the superscripts denote the stage and ϵ is the fraction of the type I error (α) spent and β is the type II error. We make use of the joint normal density of Z_J (the overall group) at each of the three stages and the fact that the \tilde{Z}_J is merely a translation of Z_J . So here the calculation is based on a mean of zero and has to be translated during use!

References

Adaptive Choice of Patient Subgroup for Comparing Two Treatments by Tze Leung Lai and Philip W. Lavori and Olivia Yueh-Wen Liao. Contemporary Clinical Trials, Vol. 39, No. 2, pp 191-200 (2014). http://www.sciencedirect.com/science/article/pii/S1551714414001311

mHP.c	<i>Compute the efficacy boundary (modified Haybittle-Peto) for the final (third) stage</i>

Description

Compute the efficacy boundary (modified Haybittle-Peto) for the final (third) stage

Usage

mHP.c(prevalence, N, cov.J, mu.prime, Sigma.prime, alpha, btilde, b, theta)

wilcoxon

Arguments

prevalence	the vector of prevalences between 0 and 1 summing to 1. J , the number of groups, is implicitly the length of this vector and should be at least 2.
Ν	a three-vector of total sample size at each stage
cov.J	the 3 x 3 covariance matrix for Z_J at each of the three stages
mu.prime	a list of J mean vectors, each of length $J-1$ representing the conditional means of all the other Z_j given Z_i . This mean does not account for the conditioned value of Z_i and so has to be multiplied by that during use!
Sigma.prime	a list of J covariance matrices, each $J-1$ by $J-1$ representing the conditional covariances all the other Z_j given Z_i
alpha	the amount of type I error to spend
btilde	the futility boundary
b	the efficacy boundary for the first two stages
theta	the effect size on the probability scale

References

Adaptive Choice of Patient Subgroup for Comparing Two Treatments by Tze Leung Lai and Philip W. Lavori and Olivia Yueh-Wen Liao. Contemporary Clinical Trials, Vol. 39, No. 2, pp 191-200 (2014). http://www.sciencedirect.com/science/article/pii/S1551714414001311

wilcoxon

Compute the standardized Wilcoxon test statistic for two samples

Description

We compute the standardized Wilcoxon test statistic with mean 0 and and standard deviation 1 for samples x and y. The R function stats::wilcox.test() returns the statistic

Usage

wilcoxon(x, y, theta = 0)

Arguments

x	a sample numeric vector
У	a sample numeric vector
theta	a value > 0 but < $1/2$.

Details

$$U = \sum_{i} R_i - \frac{m(m+1)}{2}$$

where R_i are the ranks of the first sample x of size m. We compute

$$\frac{(U - mn(1/2 + \theta))}{\sqrt{mn(m + n + 1)/12}}$$

where θ is the alternative hypothesis shift on the probability scale, i.e. $P(X > Y) = 1/2 + \theta$.

Value

the standardized Wilcoxon statistic

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