Package 'blindrecalc'

July 6, 2021

Type Package

Title Blinded Sample Size Recalculation

Version 0.1.3

Description Computation of key characteristics and plots for blinded sample size recalculation. Continuous as well as binary endpoints are supported in superiority and non-inferiority trials. The implemented methods include the approaches by Lu, K. (2019) <doi:10.1002/pst.1737>,

Kieser, M. and Friede, T. (2000) <doi:10.1002/(SICI)1097-0258(20000415)19:7%3C901::AID-SIM405%3E3.0.CO;2-L>,

Friede, T. and Kieser, M. (2004) <doi:10.1002/pst.140>,

Friede, T., Mitchell, C., Mueller-Veltern, G. (2007) <doi:10.1002/bimj.200610373>, and Friede, T. and Kieser, M. (2011) <doi:10.3414/ME09-01-0063>.

License MIT + file LICENSE

URL https://github.com/imbi-heidelberg/blindrecalc

Encoding UTF-8

Suggests testthat, covr

Imports methods, Rcpp

Collate test_statistics.R methods.R Student.R blindrecalc.R ChiSquare.R ChiSquare_helper.R FarringtonManning.R FarringtonManning_helper.R RcppExports.R

RoxygenNote 7.1.1

LinkingTo Rcpp

NeedsCompilation yes

Author Lukas Baumann [aut, cre] (<https://orcid.org/0000-0001-7931-7470>), Maximilian Pilz [aut] (<https://orcid.org/0000-0002-9685-1613>), Institute of Medical Biometry and Informatics - University of Heidelberg [cph]

Maintainer Lukas Baumann <baumann@imbi.uni-heidelberg.de>

Repository CRAN

Date/Publication 2021-07-06 16:00:16 UTC

R topics documented:

adjusted_alpha	2
adjusted_alpha,ChiSquare-method	3
adjusted_alpha,FarringtonManning-method	4
adjusted_alpha,Student-method	6
blindrecalc.	7
ChiSquare-class	7
FarringtonManning-class	9
n_dist	10
n_dist,ChiSquare-method	11
n_dist,FarringtonManning-method	12
	13
n_fix	14
n_fix,ChiSquare-method	15
n_fix,FarringtonManning-method	16
n_fix,Student-method	17
pow	18
pow,ChiSquare-method	19
pow,FarringtonManning-method	20
pow,Student-method	21
simulation	22
Student-class	23
toer	25
toer,ChiSquare-method	26
toer,FarringtonManning-method	27
toer,Student-method	28
	30

Index

adjusted_alpha Adjusted level of significance

Description

This method returns an adjusted significance level that can be used such that the actual type I error rate is preserved.

Usage

```
adjusted_alpha(design, n1, nuisance, ...)
```

Arguments

design	object of class TestStatistic created by setup
n1	total number of patients that are recruited before the sample size is recalculated
nuisance	nuisance parameter that is estimated at the interim analysis
	Further optional arguments.

Details

The method is only vectorized in either nuisance or n1.

The method is implemented for the classes Student, ChiSquare, and FarringtonManning. Check the class-specific documentation for further parameters that have to be specified.

Value

Value of the adjusted significance level for every nuisance parameter and every value of n1.

Examples

```
d <- setupStudent(alpha = .025, beta = .2, r = 1, delta = 0, delta_NI = 1.5, n_max = 848)
sigma <- c(2, 5.5, 9)
adjusted_alpha(design = d, n1 = 20, nuisance = sigma, tol = 1e-4, iters = 1e3)</pre>
```

adjusted_alpha,ChiSquare-method

Adjusted level of significance

Description

This method returns an adjusted significance level that can be used such that the actual type I error rate is preserved.

Usage

```
## S4 method for signature 'ChiSquare'
adjusted_alpha(
    design,
    n1,
    nuisance,
    nuis_ass,
    precision = 0.001,
    gamma = 0,
    recalculation,
    allocation = c("exact", "approximate"),
    ...
)
```

Arguments

design	Object of class ChiSquare created by setupChiSquare.
n1	Either the sample size of the first stage (if recalculation = TRUE or the total sample size (if recalculation = FALSE).
nuisance	Value of the nuisance parameter in $(0,1)$. For the Chi-Squared test this is the overall response rate.

nuis_ass	If recalculation = FALSE this is the value for the overall response rate that is used to calculate the sample size for the adjusted significance level.
precision	Value by which the nominal type 1 error rate is reduced in each iteration until the nominal type 1 error rate is preserved.
gamma	If gamma > 0 then the significance level is adjusted such that the actual level is at most alpha -gamma. This is necessary to maintain the nomininal significance level if a confidence interval approach proposed by Friede & Kieser (2011) is used.
recalculation	Should the sample size be recalculated after n1 n1 patients are recruited?
allocation	Whether the allocation ratio should be preserved exactly (exact) or approxi- mately (approximate or kf_approx). approximate uses the unrounded calcu- lated sample size in the sample size recalculation, kf_approx rounds the sample size to the next integer.
	Further optional arguments.

adjusted_alpha,FarringtonManning-method

Details

The method is only vectorized in either nuisance or n1.

Value

Value of the adjusted significance level for every nuisance parameter and every value of n1.

Examples

```
d <- setupChiSquare(alpha = 0.025, beta = 0.2, r = 1, delta = 0.2)
adjusted_alpha(d, n1 = 10, nuisance = 0.3, gamma = 0.001,
    nuis_ass = 0.3, precision = 0.001, recalculation = TRUE)</pre>
```

adjusted_alpha,FarringtonManning-method Adjusted level of significance

Description

This method returns an adjusted significance level that can be used such that the actual type I error rate is preserved.

Usage

S4 method for signature 'FarringtonManning'
adjusted_alpha(
 design,
 n1,
 nuisance,

```
nuis_ass,
precision = 0.001,
gamma = 0,
recalculation,
allocation = c("exact", "approximate"),
...
```

Arguments

design	Object of class FarringtonManning created by setupFarringtonManning.
n1	Either the sample size of the first stage (if recalculation = TRUE or the total sample size (if recalculation = FALSE).
nuisance	Value of the nuisance parameter in $(0,1)$. For the Farrington-Manning test this is the overall response rate.
nuis_ass	If recalculation = FALSE this is the value for the overall response rate that is used to calculate the sample size for the adjusted significance level.
precision	Value by which the nominal type 1 error rate is reduced in each iteration until the nominal type 1 error rate is preserved.
gamma	If gamma > 0 then the significance level is adjusted such that the actual level is at most alpha -gamma. This is necessary to maintain the nomininal significance level if a confidence interval approach proposed by Friede & Kieser (2011) is used.
recalculation	Should the sample size be recalculated after n1 n1 patients are recruited?
allocation	Whether the allocation ratio should be preserved exactly (exact) or approximately (approximate).
	Further optional arguments.

Details

The method is only vectorized in either nuisance or n1.

Value

Value of the adjusted significance level for every nuisance parameter and every value of n1.

```
d <- setupFarringtonManning(alpha = 0.025, beta = 0.2, r = 1, delta = 0, delta_NI = 0.25)
adjusted_alpha(d, n1 = 20, nuisance = 0.5, recalculation = TRUE)</pre>
```

adjusted_alpha,Student-method

Adjusted level of significance

Description

This method returns an adjusted significance level that can be used such that the actual type I error rate is preserved.

Usage

```
## S4 method for signature 'Student'
adjusted_alpha(design, n1, nuisance, tol, iters = 10000, seed = NULL, ...)
```

Arguments

design	Object of class Student created by setupStudent.
n1	Either the sample size of the first stage (if recalculation = TRUE or the toal sample size (if recalculation = FALSE).
nuisance	Value of the nuisance parameter. For the Student's t-test this is the variance.
tol	desired absolute tolerance
iters	Number of simulation iterations.
seed	Random seed for simulation.
	Further optional arguments.

Details

The method is only vectorized in either nuisance or n1.

In the case of the Student's t-test, the adjusted alpha is calculated using the algorithm by Kieser and Friede (2000): "Re-calculating the sample size in internal pilot study designs with control of the type I error rate". Statistics in Medicine 19: 901-911.

Value

Value of the adjusted significance level for every nuisance parameter and every value of n1.

```
d <- setupStudent(alpha = .025, beta = .2, r = 1, delta = 0, delta_NI = 1.5, n_max = 848)
sigma <- c(2, 5.5, 9)
adjusted_alpha(design = d, n1 = 20, nuisance = sigma, tol = 1e-4, iters = 1e3)
```

blindrecalc

Description

The package **blindrecalc** provides characteristics and plots of trial designs with blinded sample size recalculation where a nuisance parameter is estimated at an blinded interim analysis.

Details

Currently, for continuous outcomes, a t-test is implemented for superiority and non-inferiority trials. For superiority trials with binary endpoint, the chi^2-test is implemented. The Farrington Manning test covers non-inferiority trials with binary endpoint.

ChiSquare-class Chi-squared test

Description

This class implements a chi-squared test for superiority trials. A trial with binary outcomes in two groups E and C is assumed. If alternative == "greater" the null and alternative hypotheses for the difference in response probabilities are

$$H_0: p_E \le p_C \text{ vs. } H_1: p_E > p_C.$$

If alternative == "smaller", the direction of the effect is changed.

The function setupChiSquare creates an object of class ChiSquare.

Usage

```
setupChiSquare(
   alpha,
   beta,
   r = 1,
   delta,
   alternative = c("greater", "smaller"),
   n_max = Inf,
   ...
)
```

Arguments

alpha	One-sided type I error rate.
beta	Type II error rate.
r	Allocation ratio between experimental and control group.
delta	Difference of effect size between alternative and null hypothesis.
alternative	Does the alternative hypothesis contain greater (greater) or smaller (smaller) values than the null hypothesis.
n_max	Maximal overall sample size. If the recalculated sample size is greater than n_max it is set to n_max .
	Further optional arguments.

Details

The nuisance parameter is the overall response probability p_0 . In the blinded sample size recalculation procedure it is blindly estimated by:

$$\hat{p}_0 := (X_{1,E} + X_{1,C})/(n_{1,E} + n_{1,C}),$$

where $X_{1,E}$ and $X_{1,C}$ are the numbers of responses and $n_{1,E}$ and $n_{1,C}$ are the sample sizes of the respective group after the first stage. The event rates in both groups under the alternative hypothesis can then be blindly estimated as:

$$\hat{p}_{C,A} := \hat{p}_0 - \Delta \cdot r/(1+r), \ \hat{p}_{E,A} := \hat{p}_0 + \Delta/(1+r),$$

where Δ is the difference in response probabilities under the alternative hypothesis and r is the allocation ratio of the sample sizes in the two groups. These blinded estimates can then be used to re-estimate the sample size.

The following methods are available for this class: toer, pow, n_dist, adjusted_alpha, and n_fix. Check the design specific documentation for details.

For non-inferiority trials use the function setupFarringtonManning.

Value

An object of class ChiSquare.

References

Friede, T., & Kieser, M. (2004). Sample size recalculation for binary data in internal pilot study designs. Pharmaceutical Statistics: The Journal of Applied Statistics in the Pharmaceutical Industry, 3(4), 269-279.

Kieser, M. (2020). Methods and applications of sample size calculation and recalculation in clinical trials. Springer.

```
design <- setupChiSquare(alpha = .025, beta = .2, r = 1, delta = 0.2,
alternative = "greater")
```

FarringtonManning-class

Farrington Manning test

Description

This class implements a Farrington-Manning test for non-inferiority trials. A trial with binary outcomes in two groups E and C is assumed. The null and alternative hypotheses for the non-inferiority of response probabilities are:

 $H_0: p_E - p_C \le -\delta$ vs. $H_1: p_E - p_C > -\delta$,

where δ denotes the non-inferiority margin.

The function setupFarringtonManning creates an object of FarringtonManning.

Usage

setupFarringtonManning(alpha, beta, r = 1, delta, delta_NI, n_max = Inf, ...)

Arguments

alpha	One-sided type I error rate.
beta	Type II error rate.
r	Allocation ratio between experimental and control group.
delta	Difference of effect size between alternative and null hypothesis.
delta_NI	Non-inferiority margin.
n_max	Maximal overall sample size. If the recalculated sample size is greater than n_max it is set to n_max .
	Further optional arguments.

Details

The nuisance parameter is the overall response probability p_0 . In the blinded sample size recalculation procedure it is blindly estimated by:

$$\hat{p}_0 := (X_{1,E} + X_{1,C}) / (n_{1,E} + n_{1,C}),$$

where $X_{1,E}$ and $X_{1,C}$ are the numbers of responses and $n_{1,E}$ and $n_{1,C}$ are the sample sizes of the respective group after the first stage. The event rates in both groups under the alternative hypothesis can then be blindly estimated as:

$$\hat{p}_{C,A} := \hat{p}_0 - \Delta \cdot r/(1+r), \, \hat{p}_{E,A} := \hat{p}_0 + \Delta/(1+r),$$

where Δ is the difference in response probabilities under the alternative hypothesis and r is the allocation ratio of the sample sizes in the two groups. These blinded estimates can then be used to re-estimate the sample size.

The following methods are available for this class: toer, pow, n_dist, adjusted_alpha, and n_fix. Check the design specific documentation for details.

Value

An object of class FarringtonManning.

References

Friede, T., Mitchell, C., & Mueller-Velten, G. (2007). Blinded sample size reestimation in noninferiority trials with binary endpoints. Biometrical Journal, 49(6), 903-916. Kieser, M. (2020). Methods and applications of sample size calculation and recalculation in clinical trials. Springer.

Examples

```
design <- setupFarringtonManning(alpha = .025, beta = .2, r = 1, delta = 0,
delta_NI = .15)
```

n_dist

Distribution of the Sample Size

Description

Calculates the distribution of the total sample sizes of designs with blinded sample size recalculation for different values of the nuisance parameter or of n1.

Usage

```
n_dist(design, n1, nuisance, summary = TRUE, plot = FALSE, ...)
```

Arguments

design	object of class TestStatistic created by setup
n1	total number of patients that are recruited before the sample size is recalculated
nuisance	nuisance parameter that is estimated at the interim analysis
summary	logical - is a summary of the sample size distribution desired? Otherwise, a vector with sample sizes is returned.
plot	Should a plot of the sample size distribution be drawn?
	Further optional arguments.

Details

The method is only vectorized in either nuisance or n1.

The method is implemented for the classes Student, ChiSquare, and FarringtonManning.

Value

Summary and/or plot of the sample size distribution for every nuisance parameter and every value of n1.

Examples

n_dist,ChiSquare-method

Distribution of the Sample Size

Description

Calculates the distribution of the total sample sizes of designs with blinded sample size recalculation for different values of the nuisance parameter or of n1.

Usage

```
## S4 method for signature 'ChiSquare'
n_dist(
    design,
    n1,
    nuisance,
    summary = TRUE,
    plot = FALSE,
    allocation = c("exact", "approximate"),
    ...
)
```

Arguments

design	Object of class ChiSquare created by setupChiSquare.
n1	Either the sample size of the first stage (if recalculation = TRUE or the total sample size (if recalculation = FALSE).
nuisance	Value of the nuisance parameter in $(0,1)$. For the Chi-Squared test this is the overall response rate.
summary	Is a summary of the sample size distribution desired? Otherwise, a vector with sample sizes is returned.
plot	Should a plot of the sample size distribution be drawn?
allocation	Whether the allocation ratio should be preserved exactly (exact) or approxi- mately (approximate or kf_approx). appproximate uses the unrounded calcu- lated sample size in the sample size recalculation, kf_approx rounds the sample size to the next integer.
	Further optional arguments.

Details

Only sample sizes that occur with a probability of at least 0.01 considered.

The method is only vectorized in either nuisance or n1.

Value

Summary and/or plot of the sample size distribution for every nuisance parameter and every value of n1.

Examples

```
d <- setupChiSquare(alpha = 0.025, beta = 0.2, r = 1, delta = 0.2)
n_dist(d, n1 = 20, nuisance = 0.25, summary = TRUE, plot = FALSE)
```

n_dist,FarringtonManning-method

Distribution of the Sample Size

Description

Calculates the distribution of the total sample sizes of designs with blinded sample size recalculation for different values of the nuisance parameter or of n1.

Usage

```
## S4 method for signature 'FarringtonManning'
n_dist(
    design,
    n1,
    nuisance,
    summary,
    plot,
    allocation = c("exact", "approximate"),
    ...
)
```

Arguments

design	Object of class FarringtonManning created by setupFarringtonManning.
n1	Either the sample size of the first stage (if recalculation = TRUE or the total sample size (if recalculation = FALSE).
nuisance	Value of the nuisance parameter in $(0,1)$. For the Farrington-Manning test this is the overall response rate.
summary	Is a summary of the sample size distribution desired? Otherwise, a vector with sample sizes is returned.

plot	Should a plot of the sample size distribution be drawn?
allocation	Whether the allocation ratio should be preserved exactly (exact) or approximately (approximate).
	Further optional arguments.

Details

Only sample sizes that occur with a probability of at least 0.01 considered.

The method is only vectorized in either nuisance or n1.

Value

Summary and/or plot of the sample size distribution for each nuisance parameter and every value of n1.

Examples

```
d <- setupFarringtonManning(alpha = 0.025, beta = 0.2, r = 1, delta = 0, delta_NI = 0.25)
n_dist(d, n1 = 30, nuisance = 0.2, summary = TRUE, plot = FALSE)</pre>
```

n_dist, Student-method Distribution of the Sample Size

Description

Calculates the distribution of the total sample sizes of designs with blinded sample size recalculation for different values of the nuisance parameter or of n1.

Usage

```
## S4 method for signature 'Student'
n_dist(
    design,
    n1,
    nuisance,
    summary = TRUE,
    plot = FALSE,
    iters = 10000,
    seed = NULL,
    range = 0,
    allocation = c("approximate", "exact"),
    ...
)
```

Arguments

design	Object of class Student created by setupStudent.
n1	Either the sample size of the first stage (if recalculation = TRUE or the toal sample size (if recalculation = FALSE).
nuisance	Value of the nuisance parameter. For the Student's t-test this is the variance.
summary	Is a summary of the sample size distribution desired? Otherwise, a vector with sample sizes is returned.
plot	Should a plot of the sample size distribution be drawn?
iters	Number of simulation iterations.
seed	Random seed for simulation.
range	this determines how far the plot whiskers extend out from the box. If range is positive, the whiskers extend to the most extreme data point which is no more than range times the interquartile range from the box. A value of zero causes the whiskers to extend to the data extremes.
allocation	Whether the allocation ratio should be preserved exactly (exact) or approximately (approximate).
	Further optional arguments.

Details

The method is only vectorized in either nuisance or n1.

Value

Summary and/or plot of the sample size distribution for every nuisance parameter and every value of n1.

Examples

n_fix	
-------	--

Fixed Sample Size

Description

Returns the total sample size of a fixed design without sample size recalculation.

Usage

n_fix(design, nuisance, ...)

Arguments

design	test statistic object created by setup
nuisance	nuisance parameter for the respective test problem
	Further optional arguments.

Details

The method is only vectorized in either nuisance or n1.

The method is implemented for the classes Student, ChiSquare, and FarringtonManning.

Value

One value of the fixed sample size for every nuisance parameter and every value of n1.

Examples

n_fix,ChiSquare-method

Fixed Sample Size

Description

Returns the sample size of a fixed design without sample size recalculation.

Usage

```
## S4 method for signature 'ChiSquare'
n_fix(
   design,
   nuisance,
   variance = c("heterogeneous", "homogeneous"),
   rounded = TRUE,
   ...
)
```

Arguments

design	Object of class ChiSquare created by setupChiSquare.
nuisance	Value of the nuisance parameter. For the Chi-Squared test this is the overall
	response rate.

variance	A character string indicating whether the "heterogenous" (default) or the "homogeneous" variance formula should be used.
rounded	Whether the calculated sample size should be rounded up such that the allocation ratio is preserved.
	Further optional arguments.

Details

The method is only vectorized in either nuisance or n1.

Value

One value of the fixed sample size for every nuisance parameter and every value of n1.

Examples

```
design1 <- setupChiSquare(alpha = 0.025, beta = 0.2, r = 1, delta = 0.2)
n_fix(design1, nuisance = c(0.2, 0.3))
```

n_fix,FarringtonManning-method

Fixed Sample Size

Description

Returns the sample size of a fixed design without sample size recalculation.

Usage

```
## S4 method for signature 'FarringtonManning'
n_fix(design, nuisance, rounded = TRUE, ...)
```

Arguments

design	Object of class FarringtonManning created by setupFarringtonManning.
nuisance	Value of the nuisance parameter. For the Farrington-Manning test this is the overall response rate.
rounded	Whether the calculated sample size should be rounded up such that the allocation ratio is preserved.
	Further optional arguments.

Details

The method is only vectorized in either nuisance or n1.

n_fix,Student-method

Value

One value of the fixed sample size for every nuisance parameter and every value of n1.

Examples

d <- setupFarringtonManning(alpha = 0.025, beta = 0.2, r = 1, delta = 0, delta_NI = 0.25)
n_fix(d, nuisance = 0.3)</pre>

n_fix, Student-method Fixed Sample Size

Description

Returns the sample size of a fixed design without sample size recalculation.

Usage

```
## S4 method for signature 'Student'
n_fix(design, nuisance, ...)
```

Arguments

design	test statistic object
nuisance	nuisance parameter
	Further optional arguments.

Details

The method is only vectorized in either nuisance or n1.

Value

One value of the fixed sample size for every nuisance parameter and every value of n1.

pow

Description

Calculates the power of designs with blinded sample size recalculation or of fixed designs for one or several values of the nuisance parameter.

Usage

pow(design, n1, nuisance, recalculation, ...)

Arguments

design	object of class TestStatistic created by setup
n1	total number of patients that are recruited before the sample size is recalculated
nuisance	nuisance parameter that is estimated at the interim analysis
recalculation	Should the sample size be recalculated after n1 n1 patients are recruited?
	Further optional arguments.

Details

The method is only vectorized in either nuisance or n1.

The method is implemented for the classes Student, ChiSquare, and FarringtonManning.

Value

One power value for every nuisance parameter and every value of n1.

pow,ChiSquare-method Power

Description

Calculates the power of designs with blinded sample size recalculation or of fixed designs for one or several values of the nuisance parameter.

Usage

```
## S4 method for signature 'ChiSquare'
pow(
    design,
    n1,
    nuisance,
    recalculation,
    allocation = c("exact", "approximate", "kf_approx"),
    ...
)
```

Arguments

design	Object of class ChiSquare created by setupChiSquare.
n1	Either the sample size of the first stage (if recalculation = TRUE or the total sample size (if recalculation = FALSE).
nuisance	Value of the nuisance parameter in $(0,1)$. For the Chi-Squared test this is the overall response rate.
recalculation	Should the sample size be recalculated after n1 n1 patients are recruited?
allocation	Whether the allocation ratio should be preserved exactly (exact) or approxi- mately (approximate or kf_approx). appproximate uses the unrounded calcu- lated sample size in the sample size recalculation, kf_approx rounds the sample size to the next integer.
•••	Further optional arguments.

Details

The method is only vectorized in either nuisance or n1.

Value

One power value for every nuisance parameter and every value of n1.

```
d <- setupChiSquare(alpha = 0.025, beta = 0.2, r = 1, delta = 0.2)
pow(d, n1 = 20, nuisance = c(0.2, 0.4), recalculation = TRUE)
```

Description

Calculates the power of designs with blinded sample size recalculation or of fixed designs for one or several values of the nuisance parameter.

Usage

```
## S4 method for signature 'FarringtonManning'
pow(
   design,
   n1,
   nuisance,
   recalculation,
   allocation = c("exact", "approximate"),
   ...
)
```

Arguments

design	Object of class FarringtonManning created by setupFarringtonManning.
n1	Either the sample size of the first stage (if recalculation = TRUE or the total sample size (if recalculation = FALSE).
nuisance	Value of the nuisance parameter in $(0,1)$. For the Farrington-Manning test this is the overall response rate.
recalculation	Should the sample size be recalculated after n1 n1 patients are recruited?
allocation	Whether the allocation ratio should be preserved exactly (exact) or approximately (approximate).
	Further optional arguments.

Details

The method is only vectorized in either nuisance or n1.

Value

One power value for every nuisance parameter and every value of n1.

```
d <- setupFarringtonManning(alpha = 0.025, beta = 0.2, r = 1, delta = 0, delta_NI = 0.25)
pow(d, n1 = 30, nuisance = 0.4, allocation = "approximate", recalculation = TRUE)</pre>
```

Description

Calculates the power of designs with blinded sample size recalculation or of fixed designs for one or several values of the nuisance parameter.

Usage

```
## S4 method for signature 'Student'
pow(
   design,
   n1,
   nuisance,
   recalculation = TRUE,
   iters = 10000,
   seed = NULL,
   allocation = c("approximate", "exact"),
   ...
)
```

Arguments

design	Object of class Student created by setupStudent.
n1	Either the sample size of the first stage (if recalculation = TRUE or the toal sample size (if recalculation = FALSE).
nuisance	Value of the nuisance parameter. For the Student's t-test this is the variance.
recalculation	Should the sample size be recalculated after n1 n1 patients are recruited?
iters	Number of simulation iterations.
seed	Random seed for simulation.
allocation	Whether the allocation ratio should be preserved exactly (exact) or approximately (approximate).
	Further optional arguments.

Details

The method is only vectorized in either nuisance or n1.

Value

One power value for every nuisance parameter and every value of n1.

Examples

simulation

Simulate Rejection Probability and Sample Size for Student's t-Test

Description

This function simulates the probability that a test defined by setupStudent rejects the null hypothesis. Note that here the nuisance parameter nuisance is the variance of the outcome variable sigma^2.

Usage

```
simulation(
  design,
  n1,
  nuisance,
  recalculation = TRUE,
  delta_true,
  iters = 1000,
  seed = NULL,
  allocation = c("approximate", "exact"),
  ...
)
```

Arguments

design	Object of class Student created by setupStudent.
n1	Either the sample size of the first stage (if recalculation = TRUE or the toal sample size (if recalculation = FALSE).
nuisance	Value of the nuisance parameter. For the Student's t-test this is the variance.
recalculation	Should the sample size be recalculated after n1 n1 patients are recruited?
delta_true	effect measure under which the rejection probabilities are computed
iters	Number of simulation iterations.
seed	Random seed for simulation.
allocation	Whether the allocation ratio should be preserved exactly (exact) or approximately (approximate).
	Further optional arguments.

22

Student-class

Details

The implementation follows the algorithm in Lu (2019): Distribution of the two-sample t-test statistic following blinded sample size re-estimation. Pharmaceutical Statistics 15: 208-215. Since Lu (2019) assumes negative non-inferiority margins, the non-inferiority margin of design is multiplied with -1 internally.

Value

Simulated rejection probabilities and sample sizes for each nuisance parameter.

Examples

Student-class Student's t test

Description

This class implements Student's t-test for superiority and non-inferiority tests. A trial with continuous outcomes of the two groups E and C is assumed. If alternative == "greater" the null hypothesis for the mean difference $\Delta = \mu_E - \mu_C$ is

$$H_0: \Delta \leq -\delta_{NI}$$
 vs. $H_1: \Delta > -\delta_{NI}$.

Here, $\delta_{NI} \ge 0$ denotes the non-inferiority margin. For superiority trials, δ_{NI} can be set to zero (default). If alternative=="smaller", the direction of the effect is changed.

The function setupStudent creates an object of class Student that can be used for sample size recalculation.

Usage

```
setupStudent(
    alpha,
    beta,
    r = 1,
    delta,
    delta_NI = 0,
    alternative = c("greater", "smaller"),
    n_max = Inf,
    ...
)
```

Arguments

alpha	One-sided type I error rate.
beta	Type II error rate.
r	Allocation ratio between experimental and control group.
delta	Difference of effect size between alternative and null hypothesis.
delta_NI	Non-inferiority margin.
alternative	Does the alternative hypothesis contain greater (greater) or smaller (smaller) values than the null hypothesis.
n_max	Maximal overall sample size. If the recalculated sample size is greater than n_max it is set to n_max .
•••	Further optional arguments.

Details

The nuisance parameter is the variance σ^2 . Within the blinded sample size recalculation procedure, it is re-estimated by the one-sample variance estimator that is defined by

$$\widehat{\sigma}^2 := \frac{1}{n_1 - 1} \sum_{j \in \{T, C\}} \sum_{k=1}^{n_{1,j}} (x_{j,k} - \overline{x})^2,$$

where $x_{j,k}$ is the outcome of patient k in group j, $n_{1,j}$ denotes the first-stage sample size in group j and \bar{x} equals the mean over all n_1 observations.

The following methods are available for this class: toer, pow, n_dist, adjusted_alpha, and n_fix. Check the design specific documentation for details.

Value

An object of class Student.

References

Lu, K. (2019). Distribution of the two-sample t-test statistic following blinded sample size reestimation. Pharmaceutical Statistics 15(3): 208-215.

toer

Description

Computes the type I error rate of designs with blinded sample size recalculation or of fixed designs for one or several values of the nuisance parameter.

Usage

toer(design, n1, nuisance, recalculation, ...)

Arguments

design	object of class TestStatistic created by setup
n1	total number of patients that are recruited before the sample size is recalculated
nuisance	nuisance parameter that is estimated at the interim analysis
recalculation	Should the sample size be recalculated after n1 n1 patients are recruited?
	Further optional arguments.

Details

The method is only vectorized in either nuisance or n1.

The method is implemented for the classes Student, ChiSquare, and FarringtonManning.

Value

One type I error rate value for every nuisance parameter and every value of n1.

toer, ChiSquare-method Type I Error Rate

Description

Computes the type I error rate of designs with blinded sample size recalculation or of fixed designs for one or several values of the nuisance parameter.

Usage

```
## S4 method for signature 'ChiSquare'
toer(
   design,
   n1,
   nuisance,
   recalculation,
   allocation = c("exact", "approximate", "kf_approx"),
   ...
)
```

Arguments

design	Object of class ChiSquare created by setupChiSquare.
n1	Either the sample size of the first stage (if recalculation = TRUE or the total sample size (if recalculation = FALSE).
nuisance	Value of the nuisance parameter in $(0,1)$. For the Chi-Squared test this is the overall response rate.
recalculation	Should the sample size be recalculated after n1 n1 patients are recruited?
allocation	Whether the allocation ratio should be preserved exactly (exact) or approxi- mately (approximate or kf_approx). appproximate uses the unrounded calcu- lated sample size in the sample size recalculation, kf_approx rounds the sample size to the next integer.
	Further optional arguments.

Details

The method is only vectorized in either nuisance or n1.

Value

One type I error rate value for every nuisance parameter and every value of n1.

```
d <- setupChiSquare(alpha = 0.025, beta = 0.2, r = 1, delta = 0.2)
toer(d, n1 = c(10, 20), nuisance = 0.25, recalculation = TRUE)
```

toer, FarringtonManning-method Type I Error Rate

Description

Computes the type I error rate of designs with blinded sample size recalculation or of fixed designs for one or several values of the nuisance parameter.

Usage

```
## S4 method for signature 'FarringtonManning'
toer(
   design,
   n1,
   nuisance,
   recalculation,
   allocation = c("exact", "approximate"),
   ...
)
```

Arguments

design	Object of class FarringtonManning created by setupFarringtonManning.
n1	Either the sample size of the first stage (if recalculation = TRUE or the total sample size (if recalculation = FALSE).
nuisance	Value of the nuisance parameter in $(0,1)$. For the Farrington-Manning test this is the overall response rate.
recalculation	Should the sample size be recalculated after n1 n1 patients are recruited?
allocation	Whether the allocation ratio should be preserved exactly (exact) or approximately (approximate).
	Further optional arguments.

Details

The method is only vectorized in either nuisance or n1.

Value

One type I error rate value for every nuisance parameter and every value of n1.

```
d <- setupFarringtonManning(alpha = 0.025, beta = 0.2, r = 1, delta = 0, delta_NI = 0.2)
toer(d, n1 = 20, nuisance = 0.25, recalculation = TRUE, allocation = "approximate")</pre>
```

toer, Student-method Type I Error Rate

Description

Computes the type I error rate of designs with blinded sample size recalculation or of fixed designs for one or several values of the nuisance parameter.

Usage

```
## S4 method for signature 'Student'
toer(
   design,
   n1,
   nuisance,
   recalculation = TRUE,
   iters = 10000,
   seed = NULL,
   allocation = c("approximate", "exact"),
   ...
)
```

Arguments

design	Object of class Student created by setupStudent.
n1	Either the sample size of the first stage (if recalculation = TRUE or the toal sample size (if recalculation = FALSE).
nuisance	Value of the nuisance parameter. For the Student's t-test this is the variance.
recalculation	Should the sample size be recalculated after n1 n1 patients are recruited?
iters	Number of simulation iterations.
seed	Random seed for simulation.
allocation	Whether the allocation ratio should be preserved exactly (exact) or approximately (approximate).
	Further optional arguments.

Details

The method is only vectorized in either nuisance or n1.

Value

One type I error rate value for every nuisance parameter and every value of n1.

Examples

Index

blindrecalc, 7

ChiSquare, *3*, *7*, *8*, *10*, *15*, *18*, *25* ChiSquare (ChiSquare-class), 7 ChiSquare-class, 7

FarringtonManning, 3, 9, 10, 15, 18, 25 FarringtonManning (FarringtonManning-class), 9 FarringtonManning-class, 9

n_dist, 8, 9, 10, 24 n_dist, ChiSquare-method, 11 n_dist, FarringtonManning-method, 12 n_dist, Student-method, 13 n_fix, 8, 9, 14, 24 n_fix, ChiSquare-method, 15 n_fix, FarringtonManning-method, 16 n_fix, Student-method, 17

pow, 8, 9, 18, 24
pow, ChiSquare-method, 19
pow, FarringtonManning-method, 20
pow, Student-method, 21

toer, 8, 9, 24, 25
toer, ChiSquare-method, 26
toer, FarringtonManning-method, 27
toer, Student-method, 28