

Package ‘GenTwoArmsTrialSize’

January 20, 2025

Title Generalized Two Arms Clinical Trial Sample Size Calculation

Version 0.0.5

Description Two arms clinical trials required sample size is calculated in the comprehensive parametric context. The calculation is based on the type of endpoints(continuous/binary/time-to-event/ordinal), design (parallel/crossover), hypothesis tests (equality/noninferiority/superiority/equivalence), trial arms noncompliance rates and expected loss of follow-up. Methods are described in: Chow SC, Shao J, Wang H, Lokhnygina Y (2017) <[doi:10.1201/9781315183084](https://doi.org/10.1201/9781315183084)>, Wittes, J (2002) <[doi:10.1093/epirev/24.1.39](https://doi.org/10.1093/epirev/24.1.39)>, Sato, T (2000) <[doi:10.1002/02582000101519:19%3C2689::aid-sim555%3E3.0.co;2-0](https://doi.org/10.1002/02582000101519:19%3C2689::aid-sim555%3E3.0.co;2-0)>, Lachin J M, Foulkes, M A (1986) <[doi:10.2307/2531201](https://doi.org/10.2307/2531201)>, Whitehead J (1993) <[doi:10.1002/sim.4780122404](https://doi.org/10.1002/sim.4780122404)>, Julius SA (2023) <[doi:10.1201/9780429503658](https://doi.org/10.1201/9780429503658)>.

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Suggests knitr, rmarkdown

VignetteBuilder knitr

NeedsCompilation no

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Description

This function computes the sample size required for two arms clinical trials with continuous outcome measure. Four hypothesis tests are available under two allocation designs.

Usage

```
getSizerMean(
  design = c("parallel", "crossover"),
  test = c("equality", "noninferiority", "superiority", "equivalence"),
  alpha = 0.05,
  beta = 0.2,
  sigma,
  k = 1,
  delta = 0,
  TTE,
  rho = c(0.05, 0.07),
  r = 0.1
)
```

Arguments

design	allocation method (parallel or crossover).
test	four hypothesis tests: equality, noninferiority, superiority, and equivalence.
alpha	level of significance.
beta	type II error.
sigma	pooled standard deviation of two groups.
k	ratio of control to treatment.
delta	delta margin in test hypothesis.
TTE	target treatment effect or effect size.
rho	vector of length 2, positive noncompliance rates of two arms.
r	projected proportion of trial uniform loss of follow-up.

Value

sample size per arm.

Examples

```
# Ex 1. (n_trt=91, n_ctl=91)
getSizeMean(design="parallel", test="equality", alpha=0.05, beta=0.20,
            sigma=0.10, k=1, delta=0, TTE=0.05, rho=c(0.05, 0.07), r=0.1)

getSizeMean(design="parallel", test="noninferiority", alpha=0.05,
            beta=0.20, sigma=0.10, k=1, delta=-0.05, TTE=0, rho=c(0.05, 0.07), r=0.1)

# Ex 3. (n_trt=1022, n_ctl=1022)
getSizeMean(design="parallel", test="superiority", alpha=0.05, beta=0.20,
            sigma=0.10, k=1, delta=0.05, TTE=0.07, rho=c(0.05, 0.07), r=0.1)

# Ex 4. (n_trt=113, n_ctl=113)
getSizeMean(design="parallel", test="equivalence", alpha=0.05, beta=0.20,
            sigma=0.10, k=1, delta=0.05, TTE=0.01, rho=c(0.05, 0.07), r=0.1)

# Ex 5. (n_trt=23, n_ctl=23)
getSizeMean(design="crossover", test="equality", alpha=0.05, beta=0.20,
            sigma=0.10, k=1, delta=0, TTE=0.05, rho=c(0.05, 0.07), r=0.1)

# Ex 6. (n_trt=14, n_ctl=14)
getSizeMean(design="crossover", test="noninferiority", alpha=0.05,
            beta=0.20, sigma=0.10, k=1, delta=-0.05, TTE=0, rho=c(0.05, 0.07), r=0.1)

# Ex 7. (n_trt=21, n_ctl=21)
getSizeMean(design="crossover", test="superiority", alpha=0.05, beta=0.20,
            sigma=0.10, k=1, delta=0.05, TTE=0.01, rho=c(0.05, 0.07), r=0.1)

# Ex 8. (n_trt=29, n_ctl=29)
getSizeMean(design="crossover", test="equivalence", alpha=0.05, beta=0.20,
            sigma=0.10, k=1, delta=0.05, TTE=0.01, rho=c(0.05, 0.07), r=0.1)
```

Description

This function computes the sample size required for two arms clinical trials with ordinal outcome measure. Four hypothesis tests are available under two allocation designs.

Usage

```
getSizeOrd(
  design = c("parallel", "crossover"),
  test = c("equality", "noninferiority", "superiority", "equivalence"),
  alpha = 0.05,
  beta = 0.2,
  varcatprob,
```

```

k = 1,
theta,
delta = 0,
rho = c(0.05, 0.07),
r = 0.1
)

```

Arguments

<code>design</code>	allocation method (parallel or crossover).
<code>test</code>	four hypothesis tests: equality, noninferiority, superiority, and equivalence.
<code>alpha</code>	level of significance.
<code>beta</code>	type II error.
<code>varcatprob</code>	list of two probability vectors per treatment arm
<code>k</code>	ratio of control to treatment.
<code>theta</code>	log odds ratio of outcome in treatment arm versus control arm
<code>delta</code>	delta margin in test hypothesis.
<code>rho</code>	vector of length 2, positive noncompliance rates of two arms.
<code>r</code>	projected proportion of trial uniform loss of follow-up.

Value

sample size per arm.

Examples

```

# Ex 1. (n_trt=135, n_ctl=135)
getSizeOrd(design="parallel", test="equality", alpha=0.05, beta=0.10,
varcatprob = list(c(0.2,0.5,0.2,0.1), c(0.378,0.472,0.106,0.044)),
k=1, theta=0.887, delta=0, rho=c(0.05, 0.07), r=0.1)

# Ex 2. (Check back next version)
getSizeOrd(design="crossover", test="equality", alpha=0.05, beta=0.10,
varcatprob = list(c(0.2,0.5,0.2,0.1), c(0.378,0.472,0.106,0.044)),
k=1, theta=0.887, delta=0, rho=c(0.05, 0.07), r=0.1)

```

Description

This function computes the sample size required for two arms clinical trials with binary outcome measure. Four hypothesis tests are available under two allocation designs.

Usage

```
getSizeProp(
  design = c("parallel", "crossover"),
  test = c("equality", "noninferiority", "superiority", "equivalence"),
  alpha = 0.05,
  beta = 0.2,
  varsigma,
  k = 1,
  seqnumber,
  delta = 0,
  TTE,
  rho = c(0.05, 0.07),
  r = 0.1
)
```

Arguments

<code>design</code>	allocation method (parallel or crossover).
<code>test</code>	four hypothesis tests: equality, noninferiority, superiority, and equivalence.
<code>alpha</code>	level of significance.
<code>beta</code>	type II error.
<code>varsigma</code>	(<code>varsigma1 > 0, varsigma2 > 0</code>) := (<code>p1, p2</code>) probability of mean response in control and treatment arms; (<code>varsigma1 > 0, varsigma2 > 0</code>) := (<code>sigma, sigma</code>) pooled standard deviation of two groups or their difference (<code>sigma>0</code>)
<code>k</code>	ratio of control to treatment.
<code>seqnumber</code>	Number of crossover sequences: 0 if parallel; 1+ if crossover (<code>seqnumber>=0</code>)
<code>delta</code>	delta margin in test hypothesis.
<code>TTE</code>	target treatment effect or effect size.
<code>rho</code>	vector of length 2, positive noncompliance rates of two arms.
<code>r</code>	projected proportion of trial uniform loss of follow-up.

Value

sample size per arm.

Examples

```
# Ex 1. (n_trt=102, n_ctl=102)
getSizeProp(design="parallel", test="equality", alpha=0.05, beta=0.20,
  varsigma=c(0.65, 0.85), k=1, seqnumber=0, delta=0, TTE=0,
  rho=c(0.05, 0.07), r=0.1)

# Ex 2. (n_trt=33, n_ctl=33)
getSizeProp(design="parallel", test="noninferiority", alpha=0.05, beta=0.20,
  varsigma=c(0.65, 0.85), k=1, seqnumber=0, delta=-0.10, TTE=0.20,
  rho=c(0.05, 0.07), r=0.1)
```

```

# Ex 3. (n_trt=157, n_ctl=157)
getSizeProp(design="parallel", test="superiority", alpha=0.05, beta=0.20,
  varsigma=c(0.65,0.85), k=1, seqnumber=0, delta=0.05, TTE=0.20,
  rho=c(0.05, 0.07), r=0.1)

# Ex 4. (n_trt=137, n_ctl=137)
getSizeProp(design="parallel", test="equivalence", alpha=0.05, beta=0.20,
  varsigma=c(0.75,0.80), k=1, seqnumber=0, delta=0.20, TTE=0.05,
  rho=c(0.05, 0.07), r=0.1)

# Ex 5. (n_trt=36, n_ctl=36)
getSizeProp(design="crossover", test="equality", alpha=0.05, beta=0.20,
  varsigma=c(0.5,0.5), k=1, seqnumber=2, delta=0, TTE=0.20,
  rho=c(0.05, 0.07), r=0.1)

# Ex 6. (n_trt=22, n_ctl=22)
getSizeProp(design="crossover", test="noninferiority", alpha=0.05,
  beta=0.20, varsigma=c(0.5,0.5), k=1, seqnumber=2, delta=-0.20, TTE=0,
  rho=c(0.05, 0.07), r=0.1)

# Ex 7. (n_trt=86, n_ctl=86)
getSizeProp(design="crossover", test="superiority", alpha=0.05, beta=0.20,
  varsigma=c(0.5,0.5), k=1, seqnumber=2, delta=0.10, TTE=0,
  rho=c(0.05, 0.07), r=0.1)

# Ex 8. (n_trt=30, n_ctl=30)
getSizeProp(design="crossover", test="equivalence", alpha=0.05, beta=0.20,
  varsigma=c(0.5,0.5), k=1, seqnumber=2, delta=0.20, TTE=0,
  rho=c(0.05, 0.07), r=0.1)

```

Description

This function computes the sample size required for two arms clinical trials with TTE outcome measure. Four hypothesis tests are available under two allocation designs.

Usage

```

getSizeTTE(
  design = c("parallel", "crossover"),
  test = c("equality", "noninferiority", "superiority", "equivalence"),
  alpha = 0.05,
  beta = 0.2,
  varlambda,
  k = 1,

```

```

    ttotal,
    taccrual,
    gamma,
    delta = 0,
    rho = c(0.05, 0.07),
    r = 0.1
)

```

Arguments

design	allocation method (parallel or crossover).
test	four hypothesis tests: equality, noninferiority, superiority, and equivalence.
alpha	level of significance.
beta	type II error.
varlambda	(varlambda1>0,varlambda2>0):=(lam1, lam2) hazard rates in control and treatment arms
k	ratio of control to treatment.
ttotal	total trial time (ttotal>0)
taccrual	accrual time period (taccrual>0)
gamma	parameter of exponential distribution (gamma>=0)
delta	delta margin in test hypothesis.
rho	vector of length 2, positive noncompliance rates of two arms.
r	projected proportion of trial uniform loss of follow-up.

Value

sample size per arm.

Examples

```

# Ex 1. (n_trt=56, n_ctl=56)
getSizerTTE(design="parallel", test="equality", alpha=0.05, beta=0.20,
            varlambda=c(1,2), k=1, ttotal=3, taccrual=1, gamma=0.00001, delta=0,
            rho=c(0.05, 0.07), r=0.1)

# Ex 2. (n_trt=30, n_ctl=30)
getSizerTTE(design="parallel", test="noninferiority", alpha=0.05, beta=0.20,
            varlambda=c(1,2), k=1, ttotal=3, taccrual=1, gamma=0.00001, delta= -0.2,
            rho=c(0.05, 0.07), r=0.1)

# Ex 3. (n_trt=74, n_ctl=74)
getSizerTTE(design="parallel", test="superiority", alpha=0.05, beta=0.20,
            varlambda=c(1,2), k=1, ttotal=3, taccrual=1, gamma=0.00001, delta=0.20,
            rho=c(0.05, 0.07), r=0.1)

# Ex 4. (n_trt=84, n_ctl=84)
getSizerTTE(design="parallel", test="equivalence", alpha=0.05, beta=0.20,

```

```
varlambda=c(1,1), k=1, ttotal=3, taccrual=1, gamma=0.00001, delta=0.5,
rho=c(0.05, 0.07), r=0.1)

# Ex 5. (Check back next version)
getSizeTTE(design="crossover", test="equality", alpha=0.05, beta=0.20,
varlambda=c(1,1), k=1, ttotal=3, taccrual=1, gamma=0.00001, delta=0.5,
rho=c(0.05, 0.07), r=0.1)

# Ex 6. (Check back next version)
getSizeTTE(design="crossover", test="noninferiority", alpha=0.05,
beta=0.20, varlambda=c(1,1), k=1, ttotal=3, taccrual=1, gamma=0.00001,
delta=0.5, rho=c(0.05, 0.07), r=0.1)

# Ex 7. (Check back next version)
getSizeTTE(design="crossover", test="superiority", alpha=0.05, beta=0.20,
varlambda=c(1,1), k=1, ttotal=3, taccrual=1, gamma=0.00001, delta=0.5,
rho=c(0.05, 0.07), r=0.1)

# Ex 8. (Check back next version)
getSizeTTE(design="crossover", test="equivalence", alpha=0.05, beta=0.20,
varlambda=c(1,1), k=1, ttotal=3, taccrual=1, gamma=0.00001, delta=0.5,
rho=c(0.05, 0.07), r=0.1)
```

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