

# Package ‘ThreeArmedTrials’

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**Type** Package

**Title** Design and Analysis of Clinical Non-Inferiority or Superiority Trials with Active and Placebo Control

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**Maintainer** Tobias Mütze <tobias.muetze@outlook.com>

**Description** Design and analyze three-arm non-inferiority or superiority trials which follow a gold-standard design, i.e. trials with an experimental treatment, an active, and a placebo control. Method for the following distributions are implemented: Poisson (Mielke and Munk (2009) <arXiv:0912.4169>), negative binomial (Muetze et al. (2016) <doi:10.1002/sim.6738>), normal (Pi-geot et al. (2003) <doi:10.1002/sim.1450>; Hasler et al. (2009) <doi:10.1002/sim.3052>), binary (Friede and Kieser (2007) <doi:10.1002/sim.2543>), nonparametric (Muetze et al. (2017) <doi:10.1002/sim.7176>), exponential (Mielke and Munk (2009) <arXiv:0912.4169>).

**Depends** R (>= 3.0.0)

**Imports** stats, MASS, numDeriv

**Suggests** testthat, knitr, rmarkdown

**License** GPL (>= 2)

**NeedsCompilation** yes

**URL** <https://github.com/tobiasmuetze/ThreeArmedTrials>

**BugReports** <https://github.com/tobiasmuetze/ThreeArmedTrials/issues>

**RxygenNote** 6.1.1

**Encoding** UTF-8

**VignetteBuilder** knitr

**LazyData** true

**Author** Tobias Mütze [aut, cre] (<<https://orcid.org/0000-0002-4111-1941>>),  
Tim Friede [ctb]

**Repository** CRAN

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check_missing	<i>check_missing</i>
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### Description

Check if all arguments are defined

### Usage

```
check_missing(args = NULL, envir = parent.frame())
```

### Arguments

args	Character vector of arguments to be checked for existence.
envir	Environment in which the arguments are defined.

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check_RET_arguments	<i>check_RET_arguments</i>
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### Description

Check arguments for their respective condition

### Usage

```
check_RET_arguments(sig.level, power, Delta, n, allocation)
```

**Arguments**

<code>sig.level</code>	A numeric value specifying the significance level (type I error probability)
<code>power</code>	A numeric value specifying the target power (1 - type II error probability)
<code>Delta</code>	A numeric value specifying the non-inferiority or superiority margin. Is between 0 and 1 in case of non-inferiority and larger than 1 in case of superiority.
<code>n</code>	The total sample size. Needs to be at least 7.
<code>allocation</code>	A (non-empty) vector specifying the sample size allocation (nExp/n, nRef/n, nPla/n)

`GElesions`*Total number of new galodinium-enhancing lesions.***Description**

A (fictional) dataset containing the total number of new galodinium-enhancing lesions for different treatments for multiple sclerosis.

**Usage**`GElesions`**Format**

A data frame with 50 rows and 3 variables:

**placebo** Placebo group  
**reference** Reference group  
**experimental** Experimental treatment group

`is.naturalnumber`*is.naturalnumber***Description**

check if input is natural number

**Usage**`is.naturalnumber(x, tol = .Machine$double.eps^0.5)`**Arguments**

<code>x</code>	numeric number to be checked
<code>tol</code>	maximum accepted tolerance when checking if natural

`loglikelihood_binary`    *loglikelihood\_binary*

### Description

log likelihood of Bernoulli function

### Usage

```
loglikelihood_binary(p, xExp, xRef, xPla)
```

### Arguments

<code>p</code>	numeric vector of probabilities with length 3
<code>xExp</code>	numeric vector of probabilities with length 3
<code>xRef</code>	numeric vector of probabilities with length 3
<code>xPla</code>	numeric vector of probabilities with length 3

<code>opt_alloc_RET</code>	<i>Optimal sample size for three-arm trials when analyzed with a Wald-type test</i>
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### Description

Calculate optimal sample size allocation for Wald-type test for superiority or non-inferiority of the experimental treatment versus reference treatment with respect to placebo

### Usage

```
opt_alloc_RET(experiment, reference, placebo, Delta, distribution,
             h = NULL)
```

### Arguments

<code>experiment</code>	a numeric vector specifying the parameters of the experimental treatment group in the alternative hypothesis
<code>reference</code>	a numeric vector specifying the parameters of the reference treatment group in the alternative hypothesis
<code>placebo</code>	a numeric vector specifying the parameters of the placebo treatment group in the alternative hypothesis
<code>Delta</code>	a numeric value specifying the non-inferiority/superiority margin
<code>distribution</code>	a character specifying the distribution of the endpoints. Must be either of "poisson", "negbin", "exponential", "normal"
<code>h</code>	Function measuring the efficacy; used to defined hypothesis

## Details

The arguments experiment, reference, and placebo define the parameters of the endpoint distribution for the respective groups:

`distribution = "poisson"`: experiment, reference, and placebo must have length one and define the means.

`distribution = "negbin"`: experiment, reference, and placebo must have length two and define the mean in the first entry and the shape parameter in the second entry.

`distribution = "exponential"`: experiment, reference, and placebo must have length two and define the mean in the first entry and the probability for an uncensored observation in the second entry.

`distribution = "normal"`: experiment, reference, and placebo must have length two and define the mean in the first entry and the variance in the second entry.

## Value

Vector with optimal sample size allocation in the order (experiment, reference, placebo)

## Examples

```
opt_alloc_RET(experiment = 1,
             reference = 1,
             placebo = 3,
             Delta = 0.8,
             distribution = "poisson")
```

power\_RET

*Power related calculations for three-arm clinical trials*

## Description

Compute power, sample size, or level of significance for Wald-type test for non-inferiority or superiority of the experimental treatment versus reference treatment with respect to placebo.

## Usage

```
power_RET(experiment, reference, placebo, Delta, sig_level = NULL,
          power = NULL, n = NULL, allocation = c(1/3, 1/3, 1/3),
          distribution = NULL, ...)
```

## Arguments

- |                         |  |
|-------------------------|--|
| <code>experiment</code> | a numeric vector specifying the parameters of the experimental treatment group in the alternative hypothesis |
| <code>reference</code>  | a numeric vector specifying the parameters of the reference treatment group in the alternative hypothesis    |

placebo	a numeric vector specifying the parameters of the placebo treatment group in the alternative hypothesis
Delta	a numeric value specifying the non-inferiority/superiority margin
sig_level	A numeric value specifying the significance level (type I error probability)
power	A numeric value specifying the target power (1 - type II error probability)
n	The total sample size. Needs to be at least 7.
allocation	A (non-empty) vector specifying the sample size allocation (nExp/n, nRef/n, nPla/n)
distribution	A character specifying the distribution of the endpoints. Must be either of "binary", "poisson", "negbin", "exponential", "normal"
...	Further arguments. See details.

## Details

If the individual group sample sizes, i.e.  $n * \text{allocation}$  are not natural number, the parameters  $n$  and  $\text{allocation}$  will be re-calculated.

The additional parameter `var_estimation` is a character string specifying how the variance for the Wald-type test statistic is estimated in the Poisson and negative binomial model. Must be *RML* for restricted maximum-likelihood, or *ML* for unrestricted maximum-likelihood

## Value

A list with class "power.htest" containing the following components:

n	The total sample size
power	A numeric value specifying the target power
Delta	A numeric value specifying the non-inferiority or superiority margin.
sig.level	A character string specifying the significance level
type	A character string indicating what type of Wald-type test will be performed
allocation	A vector with the sample size allocation (nExp/n, nRef/n, nPla/n)
sig.level	The significance level (Type I error probability)
nExp	A numeric value specifying the number of sample in the experimental treatment group
nRef	A numeric value specifying the number of sample in the reference treatment group
nPla	A numeric value specifying the number of sample in the placebo treatment group

## Examples

```
power_RET(experiment = 15, reference = 17, placebo = 20,
          Delta = 0.8, sig_level = 0.025, power = 0.8,
          allocation = c(1, 1, 1) / 3,
          var_estimation = "RML",
          distribution = "poisson")
```

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**remission***Remission in clinical trial in patients with depression.*

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**Description**

A dataset indicating whether a patient went into remission defined as a HAM-D total score of <= 7.

**Usage****remission****Format**

A data frame with 88 rows and 3 variables:

**placebo** Placebo group**reference** Reference group**experimental** Experimental treatment group

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**seizures***Number of seizures per patient.*

---

**Description**

A (fictional) dataset containing the number of seizures per patient for different add-on treatments evaluating an anti-epileptic drug.

**Usage****seizures****Format**

A data frame with 18 rows and 3 variables:

**pla** Placebo group**ref** Reference group**exp** Experimental treatment group

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T2lesions	<i>Number of new and enlarging T2 lesions per patient.</i>
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### Description

A (fictional) dataset containing the number of new and enlarging T2 lesions per patient for different treatments for multiple sclerosis.

### Usage

```
T2lesions
```

### Format

A data frame with 150 rows and 3 variables:

- pla** Placebo group
- ref** Reference group
- exp** Experimental treatment group

---

test_RET	<i>Wald-type test for three-arm trials</i>
----------	--

---

### Description

Wald-type test for superiority/non-inferiority of the experimental treatment versus reference treatment with respect to placebo.

### Usage

```
test_RET(xExp, xRef, xPla, Delta, ...)
```

### Arguments

- |       |   |
|-------|---|
| xExp  | A (non-empty) numeric vector of data values from the experimental treatment group.  |
| xRef  | A (non-empty) numeric vector of data values from the reference treatment group.   |
| xPla  | A (non-empty) numeric vector of data values from the placebo group.   |
| Delta | A numeric value specifying the non-inferiority or superiority margin. Is between 0 and 1 in case of non-inferiority and larger than 1 in case of superiority. |
| ...   | Other named arguments such as <i>distribution</i> , <i>var_estimation</i> . See details for more information.   |

## Details

Additional parameters include distribution and var\_estimation.

The parameter distribution is a character string and indicates whether a parametric model should be used. If not specified retention of effect hypothesis is tested using sample means and variances. The following options exist: "poisson" (Poisson distribution), "negbin" (negative binomial distribution), "normal" (normal distribution), "exponential" (censored exponential). "nonparametric" (non-parametric). If the parameter distribution is not specified the effect and the variance for the test statistic are estimated by the sample means and sample variances.

The parameter var\_estimation defines how the variance is estimated in the parametric models "poisson" and "negbin". The following options exist: RML for the restricted maximum-likelihood estimator and ML (default) for the unrestricted maximum-likelihood estimator.

## Value

A list with class "htest" containing the following components:

statistic	The value of the Wald-type test statistic.
p.value	The p-value for the Wald-type test.
method	A character string indicating what type of Wald-type-test was performed.
estimate	The estimated rates for each of the group as well as the maximum-likelihood estimator for the shape parameter.
sample.size	The total number of data points used for the Wald-type test.

## References

- I. Pigeot, J. Schaefer, J. Roehmel, D. Hauschke. (2008). *Assessing non-inferiority of a new treatment in a three-arm clinical trial including a placebo*. Statistics in Medicine. 30(6):883-99.
- M. Hasler, R. Vonk, and LA. Hothorn. (2008). *Assessing non-inferiority of a new treatment in a three-arm trial in the presence of heteroscedasticity*. Statistics in Medicine, 27(4):490-503.
- M. Mielke and A. Munk. (2009). *The assessment and planning of non-inferiority trials for retention of effect hypotheses-towards a general approach*. arXiv preprint arXiv:0912.4169.
- T. Muetze, A. Munk, and T. Friede. (2016). *Design and analysis of three-arm trials with negative binomially distributed endpoints*. Statistics in Medicine, 35(4):505-521.

## See Also

[power\\_RET](#)

## Examples

```
# Negative binomially distributed endpoints
# Test for non-inferiority test. lambda_P=8, lambda_R = 4, lambda_E = 5, and phi = 1
# Delta = (lambda_P-lambda_E)/(lambda_P-lambda_R)
xExp <- rnbnom(60, mu = 5, size = 1)
xRef <- rnbnom(40, mu = 4, size = 1)
xPla <- rnbnom(40, mu = 8, size = 1)
Delta <- (8-5) / (8-4)
test_RET(xExp, xRef, xPla, Delta, var_estimation = 'RML', distribution = "negbin")
```

```

test_RET(xExp, xRef, xPla, Delta, var_estimation = 'ML', distribution = "negbin")

# Poisson distributed endpoints
# Test for non-inferiority test. lambda_P=8, lambda_R = 4, lambda_E = 5
# Delta = (lambda_P-lambda_E)/(lambda_P-lambda_R)
xExp <- rpois(60, lambda = 5)
xRef <- rpois(40, lambda = 4)
xPla <- rpois(40, lambda = 8)
Delta <- (8-5) / (8-4)
test_RET(xExp, xRef, xPla, Delta, var_estimation = 'RML', distribution = "poisson")
test_RET(xExp, xRef, xPla, Delta, var_estimation = 'ML', distribution = "poisson")

# Censored exponential distributed endpoints
# Test for non-inferiority test. lambda_P=3, lambda_R = 1, lambda_E = 2
# Probability for uncensored observation: 0.9
# Delta = (lambda_P-lambda_E)/(lambda_P-lambda_R)
x_exp <- matrix(c(rexp(40, rate = 1/2), rbinom(40, size = 1, prob = 0.9)),
                  ncol = 2, byrow = FALSE)
x_ref <- matrix(c(rexp(40, rate = 1/1), rbinom(40, size = 1, prob = 0.9)),
                  ncol = 2, byrow = FALSE)
x_pla <- matrix(c(rexp(40, rate = 1/3), rbinom(40, size = 1, prob = 0.9)),
                  ncol = 2, byrow = FALSE)
Delta <- log(2/3) / log(1/3)
test_RET(xExp = x_exp,
        xRef = x_ref,
        xPla = x_pla,
        Delta = Delta,
        distribution = "exponential")

```

## Description

The package **ThreeArmedTrials** provides functions for designing and analyzing non-inferiority or superiority trials with an active and a placebo control. Non-inferiority and superiority are defined through the hypothesis  $(\lambda_P - \lambda_E)/(\lambda_P - \lambda_R) \leq \Delta$  with the alternative hypothesis  $(\lambda_P - \lambda_E)/(\lambda_P - \lambda_R) > \Delta$ . The parameters  $\lambda_E$ ,  $\lambda_R$ , and  $\lambda_P$  are associated with the distribution of the endpoints and smaller values of  $\lambda_E$ ,  $\lambda_R$ , and  $\lambda_P$  are considered to be desirable. A detailed description of these parameters can be found in the help file of the individual functions. The margin  $\Delta$  is between 0 and 1 for testing non-inferiority and larger than 1 for testing superiority.

A detailed discussion of the hypothesis can be found in Hauschke and Pigeot (2005).

The statistical theory for negative binomial distributed endpoint has been developed by Muetze et al. (2015).

## Author(s)

Tobias Muetze <tobias.muetze@outlook.com>

**References**

- Hauschke, D. and Pigeot, I. 2005. "Establishing efficacy of a new experimental treatment in the 'gold standard' design." *Biometrical Journal* 47, 782–786. Muetze, T. et al. 2015. "Design and analysis of three-arm trials with negative binomially distributed endpoints." *Submitted*.

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