Package 'powertools'

July 24, 2024

Type Package

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|--|
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Description

A short description... THIS IS A TEST DESCRIPTION.

Usage

```
altprimary(
   K,
   n1 = NULL,
   n.ratio = 1,
   delta = NULL,
   Sigma,
   sd,
   rho,
   alpha = 0.025,
   power = NULL,
   tol = .Machine$double.eps^0.25,
   v = FALSE
)
```

Arguments

| K | The number of endpoints. |
|---------|--|
| n1 | The sample size for group 1. |
| n.ratio | The ratio $n2/n1$ between the sample sizes of two groups; defaults to 1 (equal group sizes). |
| delta | A vector of length K of the true mean differences mu1k - mu2k; must all be positive. |
| Sigma | The covariance matrix of the K outcomes, of dimension K x K. |
| sd | A vector of length K of the standard deviations of the K outcomes. |
| rho | A vector of length $0.5*K*(K-1)$ of the correlations among the K outcomes. |
| alpha | The significance level or type 1 error rate; defaults to 0.025. A one-sided test is assumed. |
| power | The specified level of power. |
| tol | The desired accuracy (convergence tolerance) for uniroot. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |
| | |

Value

A list of the arguments (including the computed one).

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Examples

```
altprimary(K = 2, n1 = 100, delta = c(0.4, 0.5), sd = c(1, 1), rho = 0.3, alpha = 0.025, power = NULL)
```

anova1way.c.bal

Power calculations for one-way balanced analysis of variance contrast test

Description

This is a test description.

Usage

```
anova1way.c.bal(
  n = NULL,
  mvec = NULL,
  cvec = NULL,
  sd = 1,
  Rsq = 0,
  ncov = 0,
  alpha = 0.05,
  power = NULL,
  v = FALSE
)
```

Arguments

| n | The sample size per group. |
|-------|--|
| mvec | A vector of group means c(mu1, mu2,). |
| cvec | A vector of contrast coefficients c(c1, c2,). |
| sd | The estimated standard deviation within each group; defaults to 1. |
| Rsq | The estimated R^2 for regressing the outcome on the covariates; defaults to 0. |
| ncov | The number of covariates adjusted for in the model; defaults to 0. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
anova1way.c.bal(n = 20, mvec = c(5, 10, 12), cvec = c(1, -1, 0), sd = 10, alpha = 0.025) anova1way.c.bal(n = 20, mvec = c(5, 10, 12), cvec = c(1, 0, -1), sd = 10, alpha = 0.025)
```

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| anova1way.c.unbal | Power calculations for one-way unbalanced analysis of variance contrast test | |
|-------------------|--|--|
| anovalway.c.unbal | , , , , , , , , , , , , , , , , , , , | |

Description

Power calculations for one-way unbalanced analysis of variance contrast test

Usage

```
anova1way.c.unbal(
  nvec = NULL,
  mvec = NULL,
  cvec = NULL,
  sd = NULL,
  Rsq = 0,
  ncov = 0,
  alpha = 0.05,
  v = FALSE
)
```

Arguments

| nvec | A vector of group sample sizes c(n1, n2,). |
|-------|--|
| mvec | A vector of group mvec c(mu1, mu2,). |
| cvec | A vector of contrast coefficients c(c1, c2,). |
| sd | The estimated standard deviation within each group. |
| Rsq | The estimated R^2 for regressing the outcome on the covariates; defaults to 0. |
| ncov | The number of covariates adjusted for in the model; defaults to 0. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed power).

```
anova1way.c.unbal(nvec = c(20, 20, 20), mvec = c(5, 10, 12), cvec = c(1, -1, 0), sd = 10, alpha = 0.025) anova1way.c.unbal(nvec = c(20, 20, 20), mvec = c(5, 10, 12), cvec = c(1, 0, -1), sd = 10, alpha = 0.025)
```

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anova1way.F.bal

Power calculation for balanced one-way ANOVA omnibus F test

Description

This function performs power and sample size calculations for the overall (omnibus) F test in a balanced (equal-sized groups) one-way analysis of variance (ANOVA). For an unbalanced one-way ANOVA F test (that is, unequal group sample sizes), use 'anovalway.F.unbal'. For contrast tests in a one-way ANOVA, see 'anovalway.c.bal' and 'anovalway.c.unbal'.

Usage

```
anova1way.F.bal(
  n = NULL,
  mvec = NULL,
  sd = 1,
  Rsq = 0,
  ncov = 0,
  alpha = 0.05,
  power = NULL,
  v = FALSE
)
```

Arguments

| n | The sample size per group. |
|-------|--|
| mvec | A vector of group means c(mu1, mu2,). |
| sd | The estimated standard deviation within each group; defaults to 1. |
| Rsq | The estimated R^2 for regressing the outcome on the covariates; defaults to 0. |
| ncov | The number of covariates adjusted for in the model; defaults to 0. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
anova1way.F.bal(n = 20, mvec = c(5, 10, 12), sd = 10) anova1way.F.bal(n = NULL, mvec = c(-0.25, 0.25), sd = 1, Rsq = 0.5^2, ncov = 1, power = 0.8)
```

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| anova1way.F.unbal | Power calculations for one-way unbalanced analysis of variance omnibus F test |
|-------------------|---|
| | nibus F test |

Description

Power calculations for one-way unbalanced analysis of variance omnibus F test

Usage

```
anova1way.F.unbal(
  nvec = NULL,
  mvec = NULL,
  sd = NULL,
  Rsq = 0,
  ncov = 0,
  alpha = 0.05,
  v = FALSE
)
```

Arguments

| nvec | A vector of group sample sizes c(n1, n2,). |
|-------|--|
| mvec | A vector of group mvec c(mu1, mu2,). |
| sd | The estimated standard deviation within each group. |
| Rsq | The estimated R^2 for regressing the outcome on the covariates; defaults to 0. |
| ncov | The number of covariates adjusted for in the model; defaults to 0. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed power).

Examples

```
anova1way.F.unbal(nvec = c(10, 20, 30), mvec = c(5, 10, 12), sd = 10)
```

| anova2way.c.bal | Power calculations for two-way balanced analysis of variance con- |
|-----------------|---|
| | trast test |

Description

Power calculations for two-way balanced analysis of variance contrast test

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Usage

```
anova2way.c.bal(
  n = NULL,
  mmatrix = NULL,
  cvec = NULL,
  factor = c("a", "b"),
  sd = 1,
  Rsq = 0,
  ncov = 0,
  alpha = 0.05,
  power = NULL,
  v = FALSE
)
```

Arguments

| n | The sample size per group. |
|---------|--|
| mmatrix | A matrix of group means (see example). |
| cvec | A vector of contrast coefficients c(c1, c2,). |
| factor | Either "a" for rows or "b" for columns depending on which factor the contrast test is being made on. |
| sd | The estimated standard deviation within each group; defaults to 1. |
| Rsq | The estimated R^2 for regressing the outcome on the covariates; defaults to 0. |
| ncov | The number of covariates adjusted for in the model; defaults to 0. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

Examples

```
mmatrix <- matrix(c(9.3, 8.9, 8.5, 8.7, 8.3, 7.3), nrow = 2, byrow = TRUE) anova2way.c.bal(n = 30, mmatrix = mmatrix, cvec = c(1, 0, -1), factor = "b", sd = 2, alpha = 0.05)
```

anova2way.c.unbal Power calculations for two-way unbalanced analysis of variance contrast test

Description

Power calculations for two-way unbalanced analysis of variance contrast test

anova2way.F.bal

Usage

```
anova2way.c.unbal(
  nmatrix = nmatrix,
  mmatrix = NULL,
  cvec = NULL,
  factor = c("a", "b"),
  sd = NULL,
  Rsq = 0,
  ncov = 0,
  alpha = 0.05,
  v = FALSE
)
```

Arguments

| nmatrix | A matrix of group sample sizes (see example). |
|---------|--|
| mmatrix | A matrix of group means (see example). |
| cvec | A vector of contrast coefficients c(c1, c2,). |
| factor | Either "a" (rows) or "b" (columns) depending on which factor the contrast test is being made on. |
| sd | The estimated standard deviation within each group. |
| Rsq | The estimated R^2 for regressing the outcome on the covariates; defaults to 0. |
| ncov | The number of covariates adjusted for in the model; defaults to 0. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

Examples

```
nmatrix <- matrix(c(30, 30, 30, 30, 30, 30), nrow = 2, byrow = TRUE) mmatrix <- matrix(c(9.3, 8.9, 8.5, 8.7, 8.3, 7.3), nrow = 2, byrow = TRUE) anova2way.c.unbal(nmatrix = nmatrix, mmatrix = mmatrix, cvec = c(1, 0, -1), factor = "b", sd = 2, alpha = 0.05)
```

anova2way.F.bal Power calculations for two-way balanced analysis of variance omnibus F test

Description

Power calculations for two-way balanced analysis of variance omnibus F test

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Usage

```
anova2way.F.bal(
  n = NULL,
  mmatrix = NULL,
  sd = 1,
  Rsq = 0,
  ncov = 0,
  alpha = 0.05,
  power = NULL,
  v = FALSE
)
```

Arguments

| n | The sample size per group. |
|---------|--|
| mmatrix | A matrix of group means (see example). |
| sd | The estimated standard deviation within each group; defaults to 1. |
| Rsq | The estimated R^2 for regressing the outcome on the covariates; defaults to 0. |
| ncov | The number of covariates adjusted for in the model; defaults to 0. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

Examples

anova2way.F.unbal Power calculations for two-way unbalanced analysis of variance omnibus F test

Description

Power calculations for two-way unbalanced analysis of variance omnibus F test

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Usage

```
anova2way.F.unbal(
  nmatrix = NULL,
  mmatrix = NULL,
  sd = NULL,
  Rsq = 0,
  ncov = 0,
  alpha = 0.05,
  v = FALSE
)
```

Arguments

| nmatrix | A matrix of group sample sizes (see example). |
|---------|--|
| mmatrix | A matrix of group means (see example). |
| sd | The estimated standard deviation within each group. |
| Rsq | The estimated R^2 for regressing the outcome on the covariates; defaults to 0. |
| ncov | The number of covariates adjusted for in the model; defaults to 0. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed power).

Examples

```
nmatrix <- matrix(c(30, 30, 30, 30, 30, 30), nrow = 2, byrow = TRUE) mmatrix <- matrix(c(9.3, 8.9, 8.5, 8.7, 8.3, 7.9), nrow = 2, byrow = TRUE) anova2way.F.unbal(nmatrix = nmatrix, mmatrix = mmatrix, sd = 2, alpha = 0.05) nmatrix <- matrix(c(30, 30, 30, 30, 30), nrow = 2, byrow = TRUE) mmatrix <- matrix(c(9.3, 8.9, 8.5, 8.7, 8.3, 7.3), nrow = 2, byrow = TRUE) anova2way.F.unbal(nmatrix = nmatrix, mmatrix = mmatrix, sd = 2, alpha = 0.05) nmatrix <- matrix(c(30, 30, 30, 30, 30, 30), nrow = 2, byrow = TRUE) mmatrix <- matrix(c(9.3, 8.9, 8.5, 8.7, 8.3, 7.9), nrow = 2, byrow = TRUE) anova2way.F.unbal(nmatrix = nmatrix, mmatrix = mmatrix, sd = 2, Rsq = 0.4^2, ncov = 1, alpha = 0.05)
```

anova2way.se.bal

Power calculations for two-way balanced analysis of variance simple effects test

Description

Power calculations for two-way balanced analysis of variance simple effects test

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Usage

```
anova2way.se.bal(
  n = NULL,
  mmatrix = NULL,
  cmatrix = NULL,
  sd = 1,
  Rsq = 0,
  ncov = 0,
  alpha = 0.05,
  power = NULL,
  v = FALSE
)
```

Arguments

| n | The sample size per group. |
|---------|--|
| mmatrix | A matrix of group means (see example). |
| cmatrix | A matrix of contrast coefficients (see example). |
| sd | The estimated standard deviation within each group; defaults to 1. |
| Rsq | The estimated R^2 for regressing the outcome on the covariates; defaults to θ . |
| ncov | The number of covariates adjusted for in the model; defaults to 0. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

Examples

anova2way.se.unbal $\begin{array}{c} \textit{Power calculations for two-way unbalanced analysis of variance simple effects test} \end{array}$

Description

Power calculations for two-way unbalanced analysis of variance simple effects test

chisq.gof

Usage

```
anova2way.se.unbal(
  nmatrix = NULL,
  mmatrix = NULL,
  cmatrix = NULL,
  sd = 0,
  Rsq = 0,
  ncov = 0,
  alpha = 0.05,
  v = FALSE
)
```

Arguments

```
nmatrix A matrix of sample sizes (see example).

A matrix of group means (see example).

A matrix of contrast coefficients (see example).

A matrix of contrast coefficients (see example).

The estimated standard deviation within each group.

Rsq The estimated R^2 for regressing the outcome on the covariates; defaults to 0.

The number of covariates adjusted for in the model; defaults to 0.

The significance level or type 1 error rate; defaults to 0.05.

V Either TRUE for verbose output or FALSE to output computed argument only.
```

Value

A list of the arguments (including the computed power).

Examples

```
nmatrix <- matrix(c(30, 30, 30, 30, 30, 30), nrow = 2, byrow = TRUE)
mmatrix <- matrix(c(9.3, 8.9, 8.5, 8.7, 8.3, 7.3), nrow = 2, byrow = TRUE)
cmatrix <- matrix(c(-1, 0, 0, 1, 0, 0), nrow = 2, byrow = TRUE)
anova2way.se.unbal(nmatrix = nmatrix, mmatrix = mmatrix, cmatrix = cmatrix,
sd = 2, alpha = 0.025)</pre>
```

chisq.gof

Power calculations for chi-square goodness-of-fit test

Description

Power calculations for chi-square goodness-of-fit test

```
chisq.gof(
  p0vec = NULL,
  p1vec = NULL,
  N = NULL,
  alpha = 0.05,
  power = NULL,
  v = FALSE
```

14 chisq.indep

Arguments

| povee the first vector of probabilities (under the fidil). | p0vec | The first vector of probabilities (under the null). |
|--|-------|---|
|--|-------|---|

p1vec The second vector of probabilities (under the alternative hypothesis).

N The number of total observations.

alpha The significance level or type 1 error rate; defaults to 0.05.

power The specified level of power.

v Either TRUE for verbose output or FALSE to output computed argument only.

Value

A list of the arguments (including the computed one).

Examples

```
chisq.gof(p0vec = c(0.5, 0.3, 0.2), p1vec = c(0.7, 0.2, 0.1), N = 50)
```

chisq.indep

Power calculations for chi-square test of independence

Description

Power calculations for chi-square test of independence

Usage

```
chisq.indep(pmatrix = NULL, N = NULL, alpha = 0.05, power = NULL, v = FALSE)
```

Arguments

pmatrix The two-way probability table under the alternative hypothesis.

N The number of total observations.

alpha The significance level or type 1 error rate; defaults to 0.05.

power The specified level of power.

v Either TRUE for verbose output or FALSE to output computed argument only.

Value

A list of the arguments (including the computed one).

```
chisq.indep(pmatrix = matrix(c(0.050, 0.350, 0.100, 0.075, 0.250, 0.175), nrow = 2, byrow = TRUE), N = 230) chisq.indep(pmatrix = matrix(c(0.3, 0.2, 0.4, 0.1), nrow = 2, byrow = TRUE), N = 200)
```

ci.mean 15

| ci.mean | Power calculations for | precision analysis for one mean |
|---------|------------------------|---------------------------------|
| | | |

Description

Power calculations for precision analysis for one mean

Usage

```
ci.mean(
  N = NULL,
  halfwidth = NULL,
  sd = 1,
  alpha = 0.05,
  power = NULL,
  cond = FALSE,
  v = FALSE
)
```

Arguments

| N | The sample size. |
|-----------|--|
| halfwidth | The desired halfwidth. |
| sd | The estimated standard deviation; defaults to 1. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| cond | Specify using unconditional or conditional probability. Defaults to FALSE. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
ci.mean(N = NULL, halfwidth = 0.25, power = 0.8)
ci.mean(N = 62, halfwidth = 0.25, power = NULL)
ci.mean(N = 73, halfwidth = 0.25, cond = TRUE)
```

16 ci.meandiff

| ci.meandiff Power calculations for precision analysis for a different means | ence between |
|--|--------------|
|--|--------------|

Description

Power calculations for precision analysis for a difference between means

Usage

```
ci.meandiff(
  n1 = NULL,
  n.ratio = 1,
  halfwidth = NULL,
  sd = 1,
  alpha = 0.05,
  power = NULL,
  cond = FALSE,
  v = FALSE
)
```

Arguments

| n1 | The sample size for group 1. |
|-----------|--|
| n.ratio | The ratio n2/n1 between the sample sizes of two groups; defaults to 1 (equal group sizes). |
| halfwidth | The desired halfwidth for the difference in means. |
| sd | The estimated standard deviation; defaults to 1. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| cond | Specify using unconditional or conditional probability. Defaults to FALSE. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
ci.meandiff(n1 = NULL, halfwidth = 0.25, power = 0.8) ci.meandiff(n1 = 134, halfwidth = 0.25, cond = TRUE)
```

coprimary.t 17

| coprimary.t | Power calculations for multiple co-primary continuous endpoints as- |
|-------------|---|
| | suming unknown covariance matrix |

Description

Power calculations for multiple co-primary continuous endpoints assuming unknown covariance matrix

Usage

```
coprimary.t(
  Κ,
  n1 = NULL,
  n.ratio = 1,
  delta = NULL,
  Sigma,
  sd,
  rho,
  alpha = 0.025,
  power = NULL,
  M = 10000,
  min.n = NULL,
  max.n = NULL,
  tol = .Machine$double.eps^0.25,
  use.uniroot = TRUE,
  v = FALSE
```

Arguments

| K | The number of endpoints. |
|-------------|--|
| n1 | The sample size for group 1. |
| n.ratio | The ratio n2/n1 between the sample sizes of two groups; defaults to 1 (equal group sizes). |
| delta | A vector of length K of the true mean differences mu1k - mu2k; must all be positive. |
| Sigma | The covariance matrix of the K outcomes, of dimension K x K. |
| sd | A vector of length K of the standard deviations of the K outcomes. |
| rho | A vector of length $0.5*K*(K-1)$ of the correlations among the K outcomes. |
| alpha | The significance level or type 1 error rate; defaults to 0.025. A one-sided test is assumed. |
| power | The specified level of power. |
| М | The number of simulation. |
| min.n | Minimum value of n1; used in search for n1 to achieve desired power. |
| max.n | Maximum value of n1; used in search for n1 to achieve desired power. |
| tol | The desired accuracy (convergence tolerance) for uniroot. |
| use.uniroot | Whether to use the uniroot function to calculate n1; defaults to TRUE. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

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Value

A list of the arguments (including the computed one).

Examples

```
coprimary.t(K = 2, n1 = 100, delta = c(0.4, 0.5), sd = c(1, 1), rho = 0.3, alpha = 0.025, power = NULL)
```

coprimary.z

Power calculations for multiple co-primary continuous endpoints assuming known covariance matrix

Description

Power calculations for multiple co-primary continuous endpoints assuming known covariance matrix

Usage

```
coprimary.z(
   K,
   n1 = NULL,
   n.ratio = 1,
   delta = NULL,
   Sigma,
   sd,
   rho,
   alpha = 0.025,
   power = NULL,
   tol = .Machine$double.eps^0.25,
   v = FALSE
)
```

Arguments

| К | The number of endpoints. |
|---------|---|
| n1 | The sample size for group 1. |
| n.ratio | The ratio n2/n1 between the sample sizes of two groups; defaults to 1 (equal group sizes). |
| delta | A vector of length K of the true mean differences $mu1k$ - $mu2k$; $must$ all be positive. |
| Sigma | The covariance matrix of the K outcomes, of dimension K x K. |
| sd | A vector of length K of the standard deviations of the K outcomes. |
| rho | A vector of length $0.5*K*(K-1)$ of the correlations among the K outcomes. |
| alpha | The significance level or type 1 error rate; defaults to 0.025. A one-sided test is assumed. |
| power | The specified level of power. |
| tol | The desired accuracy (convergence tolerance) for uniroot. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

corr.1samp

Value

A list of the arguments (including the computed one).

Examples

```
coprimary.z(K = 2, n1 = 100, delta = c(0.4, 0.5), sd = c(1, 1), rho = 0.3, alpha = 0.025, power = NULL)
```

corr.1samp

Power calculations for one correlation coefficient

Description

Power calculations for one correlation coefficient

Usage

```
corr.1samp(
  N = NULL,
  rho0 = 0,
  rhoA = NULL,
  alpha = 0.05,
  power = NULL,
  sides = 2,
  v = FALSE
)
```

Arguments

| N | The sample size. |
|-------|---|
| rho0 | The correlation coefficient under the null hypothesis; defaults to 0. |
| rhoA | The correlation coefficient under the alternative hypothesis. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
corr.1samp(N = 100, rhoA = 0.2, sides = 1)
corr.1samp(N = 100, rho0 = 0.2, rhoA = 0.4, sides = 1)
```

20 corr.2samp

Power calculations for comparing two correlation coefficients

Description

Power calculations for comparing two correlation coefficients

Usage

```
corr.2samp(
  n1 = NULL,
  n.ratio = 1,
  rho1 = NULL,
  rho2 = NULL,
  alpha = 0.05,
  power = NULL,
  sides = 2,
  v = FALSE
)
```

Arguments

| n1 | The sample size for group 1. |
|---------|--|
| n.ratio | The ratio $n2/n1$ between the sample sizes of two groups; defaults to 1 (equal group sizes). |
| rho1 | The correlation coefficient in group 1. |
| rho2 | The correlation coefficient in group 2. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
corr.2samp(n1 = 300, rho1 = 0.3, rho2 = 0.1, sides = 1)
```

crt.long.cont 21

| crt.long.cont | Power for test of treatment effect in longitudinal cluster randomized trial with baseline measurement |
|---------------|---|
| | |

Description

Power for test of treatment effect in longitudinal cluster randomized trial with baseline measurement

Usage

```
crt.long.cont(
    m = NULL,
    J1 = NULL,
    J.ratio = 1,
    delta = NULL,
    sd = 1,
    icc = 0,
    cac = 0,
    sac = 0,
    alpha = 0.05,
    power = NULL,
    sides = 2,
    v = FALSE
)
```

Arguments

| m | The number of subjects measured during each cluster-period. |
|---------|---|
| J1 | The number of clusters in arm 1. |
| J.ratio | The ratio J2/J1 between the number of clusters in the two arms; defaults to 1 (equal clusters per arm). |
| delta | The difference between the intervention and control means under the alternative minus the difference under the null hypothesis. |
| sd | The total standard deviation of the outcome variable; defaults to 1. |
| icc | The within-cluster, within-period intraclass correlation coefficient; defaults to 0. |
| cac | The cluster autocorrelation; defaults to 0. |
| sac | The subject autocorrelation; defaults to 0. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
crt.long.cont(m = 30, J1 = 8, delta = 0.3, icc = 0.05, cac = 0.4, sac = 0.5)
```

22 crt.parallel.bin

crt.means.r

Correlation between a cluster mean at baseline and follow up

Description

Correlation between a cluster mean at baseline and follow up

Usage

```
crt.means.r(m, icc, cac, sac)
```

Arguments

m The number of measurements in each cluster at baseline and follow up.
 icc The intraclass correlation coefficient.
 cac The cluster autocorrelation.
 sac The subject autocorrelation.

Value

The computed correlation.

Examples

```
crt.means.r(m = 30, icc = 0.05, cac = 0.4, sac = 0.5)
```

crt.parallel.bin

Power for a cluster randomized trial with a binary outcome

Description

Power for a cluster randomized trial with a binary outcome

```
crt.parallel.bin(
  m = NULL,
  m.sd = 0,
  J = NULL,
  pc = NULL,
  pt = NULL,
  sigma.u = NULL,
  alpha = 0.05,
  power = NULL,
  sides = 2,
  v = FALSE
)
```

crt.parallel.cont 23

Arguments

| m | The number of subjects per cluster. |
|---------|---|
| m.sd | The standard deviation of cluster sizes (provide if unequal number of participants per cluster); defaults to 0. |
| J | The number of clusters. |
| рс | The probability of the outcome in control clusters. |
| pt | The probability of the outcome in treatment clusters. |
| sigma.u | Standard deviation of the cluster random effect. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

Examples

```
crt.parallel.bin(m = 60, J = NULL, pc = 0.25, pt = 0.15, sigma.u = 0.3, power = 0.8)
crt.parallel.bin(m = 60, m.sd = 1, J = NULL, pc = 0.25, pt = 0.15, sigma.u = 0.3, power = 0.8)
```

crt.parallel.cont

Power for a cluster randomized trial with a continuous outcome

Description

Power for a cluster randomized trial with a continuous outcome

```
crt.parallel.cont(
  m = NULL,
  m.sd = 0,
  J1 = NULL,
  J.ratio = 1,
  delta = NULL,
  sd = 1,
  icc1 = 0,
  icc2 = 0,
  ncov = 0,
  RsqB = 0,
  RsqW = 0,
  alpha = 0.05,
  power = NULL,
  sides = 2,
  v = FALSE
```

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Arguments

| m | The number of subjects per cluster or the mean cluster size (if unequal number of participants per cluster). |
|---------|---|
| m.sd | The standard deviation of cluster sizes (provide if unequal number of participants per cluster); defaults to 0 . |
| J1 | The number of clusters in arm 1. |
| J.ratio | The ratio J2/J1 between the number of clusters in the two arms; defaults to 1 (equal clusters per arm). |
| delta | The difference between the intervention and control means under the alternative minus the difference under the null hypothesis. |
| sd | The total standard deviation of the outcome variable; defaults to 1. |
| icc1 | The intraclass correlation coefficient in arm 1; defaults to 0. |
| icc2 | The intraclass correlation coefficient in arm 2; defaults to 0. |
| ncov | The number of cluster-level and individual-level covariates; defaults to 0. |
| RsqB | The estimated proportion of total variance explained by cluster-level covariates; defaults to 0. |
| RsqW | The estimated proportion of total variance explained by individual-level covariates; defaults to 0. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

Examples

```
crt.parallel.cont(m = 30, J1 = 8, delta = 0.4, icc1 = 0.05, icc2 = 0.05)
crt.parallel.cont(m = NULL, J1 = 6, delta = 0.5, icc1 = 0.05, icc2 = 0.05, power = 0.8)
crt.parallel.cont(m = 25, m.sd = 15, J1 = NULL, delta = 0.3, icc1 = 0.05,
icc2 = 0.05, power = 0.8)
crt.parallel.cont(m = 20, J1 = 15, delta = 0.3, icc1 = 0.05, icc2 = 0.05,
RsqB = 0.1, ncov = 1, sides = 1)
crt.parallel.cont(m = 20, J1 = 15, delta = 0.3, icc1 = 0.05, icc2 = 0.05,
RsqW = 0.5^2, ncov = 1, sides = 1)
crt.parallel.cont(m = NULL, J1 = 6, delta = 0.5, icc1 = 0.05, icc2 = 0.05, power = 0.8)
```

crt.varexplore Variance exploration for cluster randomized trials with binary outcomes

Description

Variance exploration for cluster randomized trials with binary outcomes

crt.xo.cont 25

Usage

```
crt.varexplore(pc, pt)
```

Arguments

pc The probability of the outcome in control clusters.

pt The probability of the outcome in treatment clusters.

Value

A list of the arguments and a dataframe of outputs.

Examples

```
crt.varexplore(pc = 0.25, pt = 0.15)
```

crt.xo.cont

Power for test of treatment effect in 2x2 crossover cluster randomized trial

Description

Power for test of treatment effect in 2x2 crossover cluster randomized trial

Usage

```
crt.xo.cont(
    m = NULL,
    J.arm = NULL,
    delta = NULL,
    sd = 1,
    icc = 0,
    icca = 0,
    iccb = NULL,
    cac = NULL,
    sac = 0,
    alpha = 0.05,
    power = NULL,
    sides = 2,
    v = FALSE
)
```

Arguments

| m | The number of subjects measured during each cluster-period. |
|-------|---|
| J.arm | The number of clusters in each arm. |
| delta | The difference between the intervention and control means under the alternative minus the difference under the null hypothesis. |
| sd | The total standard deviation of the outcome variable; defaults to 1. |
| icc | The within-cluster, within-period intraclass correlation coefficient; defaults to 0. |

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| icca | The within-cluster, within-subject correlation (correlation between two measurements within the same subject); defaults to 0. |
|-------|---|
| iccb | The within-cluster, between-period intraclass correlation coefficient. Either iccb OR cac must be specified. |
| cac | The cluster autocorrelation. Either iccb OR cac must be specified. |
| sac | The subject autocorrelation; defaults to 0. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

Examples

```
crt.xo.cont(m = 30, J.arm = 4, delta = 0.3, icc = 0.05, cac = 0.8, sac = 0.4) crt.xo.cont(m = 30, J.arm = 4, delta = 0.3, icc = 0.05, icca = 0.42, iccb = 0.04) crt.xo.cont(m = 30, J.arm = 4, delta = 0.3, icc = 0.05, cac = 0.5)
```

es.anova.f *Cohen's f effect size calculation for one- or two- way analysis of variance*

Description

Cohen's f effect size calculation for one- or two- way analysis of variance

Usage

```
es.anova.f(means = NULL, sd = NULL, v = TRUE)
```

Arguments

means A vector or matrix of group means.

sd The estimated standard deviation within each group.

v Either TRUE for verbose output or FALSE to output computed argument only.

Value

Various calculated f effect sizes.

```
es.anova.f(means = c(5, 10, 12), sd = 10)
mmatrix <- matrix(c(9.3, 8.9, 8.5, 8.7, 8.3, 7.9), nrow = 2, byrow = TRUE)
es.anova.f(means = mmatrix, sd = 2)
```

es.d 27

es.d

Cohen's d effect size calculation for one or two means

Description

Cohen's d effect size calculation for one or two means

Usage

```
es.d(delta = NULL, sd = 1)
```

Arguments

delta

If one mean: muA (the true mean) - mu0 (the mean under the null). If two means: DeltaA (the true difference mu1 - mu2) - Delta0 (the difference under

the null)

sd

The estimated standard deviation; defaults to 1.

Value

A list of the arguments and the d effect size.

Examples

```
es.d(delta = 6.5 - 5.7, sd = 0.4)
```

es.fsq

Cohen's f^2 effect size calculation for overall F test

Description

Cohen's f^2 effect size calculation for overall F test

Usage

```
es.fsq(Rsq = 0.02)
```

Arguments

Rsq

The squared sample multiple correlation coefficient.

Value

A list of the arguments and the f^2 effect size.

```
es.fsq(Rsq = 0.02)
```

es.q

es.h

Cohen's h effect size calculation for two proportions

Description

Cohen's h effect size calculation for two proportions

Usage

```
es.h(p1 = NULL, p2 = NULL)
```

Arguments

p1 The proportion in group 1. p2 The proportion in group 2.

Value

A list of the arguments and the h effect size.

Examples

```
es.h(p1 = 0.8, p2 = 0.6)
```

es.q

Cohen's q effect size calculation for two correlation coefficients

Description

Cohen's q effect size calculation for two correlation coefficients

Usage

```
es.q(rho1 = NULL, rho2 = NULL)
```

Arguments

rho1 The correlation coefficient in group 1. rho2 The correlation coefficient in group 2.

Value

A list of the arguments and the q effect size.

```
es.q(rho1 = 0.3, rho2 = 0.1)
```

es.w 29

es.w

Cohen's w effect size calculation for chi-square tests

Description

Cohen's w effect size calculation for chi-square tests

Usage

```
es.w(p0vec = NULL, p1vec = NULL, pmatrix = NULL)
```

Arguments

povec The first vector of probabilities. Both povec and plvec, or pmatrix must be

specified.

p1vec The second vector of probabilities. Both p0vec and p1vec, or pmatrix must be

specified.

pmatrix The two-way probability table. Both p0vec and p1vec, or pmatrix must be spec-

ified.

Value

A list of the arguments and the w effect size.

Examples

```
es.w(p0vec = c(0.5, 0.3, 0.2), p1vec = c(0.7, 0.2, 0.1))
es.w(pmatrix = matrix(c(0.050, 0.350, 0.100, 0.075, 0.250, 0.175), nrow = 2, byrow = TRUE))
```

irgtt.bin

Power for an individual randomized group treatment trial with binary outcomes

Description

Power for an individual randomized group treatment trial with binary outcomes

```
irgtt.bin(
  m = NULL,
  J = NULL,
  n = NULL,
  p1 = NULL,
  icc = 0,
  alpha = 0.05,
  power = NULL,
  sides = 2,
  v = FALSE
```

irgtt.cont

Arguments

| m | The number of subjects per cluster in the treatment group. |
|-------|---|
| J | The number of clusters in the treatment group. |
| n | The number of total participants in the control group. |
| p1 | The probability of the outcome in the control group. |
| p2 | The probability of the outcome in the intervention group. |
| icc | The intraclass correlation coefficient in the treatment group; defaults to 0. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

Examples

```
irgtt.bin(m = 20, J = 5, n = 100, p1 = 0.8, p2 = 0.6, icc = 0.04, sides = 2) irgtt.bin(m = 20, J = 6, n = 120, p1 = 0.8, p2 = 0.6, icc = 0.04, sides = 2)
```

irgtt.cont

Power for individually randomized group treatment trial with continuous outcome

Description

Power for individually randomized group treatment trial with continuous outcome

```
irgtt.cont(
  m = NULL,
  J = NULL,
  n = NULL,
  delta = NULL,
  sd = 1,
  icc = 0,
  Theta = 1,
  alpha = 0.05,
  power = NULL,
  sides = 2,
  tol = .Machine$double.eps^0.25,
  v = FALSE
)
```

margin.sign 31

Arguments

| m | The number of subjects per cluster in the treatment group. |
|-------|---|
| J | The number of clusters in the treatment group. |
| n | The number of total participants in the control group. |
| delta | The difference between the intervention and control means under the alternative minus the difference under the null hypothesis. |
| sd | The total standard deviation of the outcome variable in the control group; defaults to $1. $ |
| icc | The intraclass correlation coefficient in the treatment group; defaults to 0. |
| Theta | The ratio of the total variance between intervention and control groups; defaults to $1. $ |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| tol | The desired accuracy (convergence tolerance) for uniroot. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

Examples

```
irgtt.cont(m = 10, J = 12, n = 120, delta = 0.4, icc = 0.05, Theta = 1, power = NULL) irgtt.cont(m = 10, J = 12, n = NULL, delta = 0.4, icc = 0.05, Theta = 1, power = 0.8)
```

margin.sign

Margin Sign Table

Description

A table documenting the sign of margin for tests comparing two parameters.

Usage

```
{\tt margin.sign}
```

Format

'margin.sign' A data frame with 3 rows and 3 columns:

columns Whether the test is for noninferiority of superirority by a marginrows Whether a higher or lower parameter value is betterxij The sign of delta given the row & column

32 mlrF.overall

| mlrF.overall |
|--------------|
|--------------|

Power calculations for a multiple linear regression overall F test

Description

Power calculations for a multiple linear regression overall F test

Usage

```
mlrF.overall(
  N = NULL,
  p = NULL,
  Rsq = NULL,
  fsq = NULL,
  alpha = 0.05,
  power = NULL,
  random = FALSE,
  v = FALSE
)
```

Arguments

| N | The sample size. |
|--------|---|
| р | The number of predictors. |
| Rsq | The squared sample multiple correlation coefficient. |
| fsq | The squared f effect size. Either Rsq OR fsq must be specified. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| random | Whether the values of the predictors are random; defaults to FALSE. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
mlrF.overall(N = 400, p = 2, Rsq = 0.02)
mlrF.overall(N = 400, p = 2, fsq = 0.02 / (1 - 0.02))
mlrF.overall(N = 109, p = 1, Rsq = 0.3^2)
mlrF.overall(N = 50, p = 1, Rsq = 0.2)
mlrF.overall(N = 50, p = 3, Rsq = 0.2)
mlrF.overall(N = 50, p = 5, Rsq = 0.2)
mlrF.overall(N = 400, p = 2, Rsq = 0.02, random = TRUE)
```

mlrF.partial 33

Power calculations for a multiple linear regression partial F test

Description

Power calculations for a multiple linear regression partial F test

Usage

```
mlrF.partial(
  N = NULL,
  p = NULL,
  q = NULL,
  pc = NULL,
  Rsq.red = NULL,
  Rsq.full = NULL,
  alpha = 0.05,
  power = NULL,
  v = FALSE
)
```

Arguments

| N | The sample size. |
|----------|--|
| р | The number of control predictors. |
| q | The number of test predictors. |
| pc | The partial correlation coefficient. Either both Rsq terms OR pc must be specified. |
| Rsq.red | The squared sample multiple correlation coefficient in the reduced model. Either both Rsq terms OR pc must be specified. |
| Rsq.full | The squared sample multiple correlation coefficient in the full model. Either both Rsq terms OR pc must be specified. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
mlrF.partial(N = 80, p = 3, q = 2, Rsq.red = 0.25, Rsq.full = 0.35) mlrF.partial(N = 150, p = 4, pc = 0.2)
```

34 multisite.bin

ms.varexplore

Variance exploration for multisite trials with binary outcomes

Description

Variance exploration for multisite trials with binary outcomes

Usage

```
ms.varexplore(pc, pt)
```

Arguments

pc The probability of the outcome in control clusters.

pt The probability of the outcome in treatment clusters.

Value

A list of the arguments and a dataframe of outputs.

Examples

```
ms.varexplore(pc = 0.1, pt = 0.2)
```

multisite.bin

Power for multisite trials with binary outcomes

Description

Power for multisite trials with binary outcomes

```
multisite.bin(
  m = NULL,
  alloc.ratio = 1,
  J = NULL,
  pc = NULL,
  pt = NULL,
  sigma.u = NULL,
  alpha = 0.05,
  power = NULL,
  sides = 2,
  v = FALSE
)
```

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Arguments

| m | The total number of subjects per site. |
|-------------|---|
| alloc.ratio | The allocation ratio of intervention/control per site; defaults to 1. |
| J | The number of sites. |
| рс | The probability of the outcome in the control condition. |
| pt | The probability of the outcome in the treatment condition. |
| sigma.u | Standard deviation of the treatment effect across sites. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

Examples

```
multisite.bin(m = 30, J = 25, pc = 0.1, pt = 0.2, sigma.u = 0.4, power = NULL) multisite.bin(m = 30, J = NULL, pc = 0.1, pt = 0.2, sigma.u = 0.4, power = 0.9)
```

multisite.cont

Power for test of average treatment effect in a multisite trial

Description

Power for test of average treatment effect in a multisite trial

```
multisite.cont(
   m = NULL,
   m.sd = 0,
   alloc.ratio = 1,
   J = NULL,
   delta = NULL,
   sd = 1,
   icc0 = NULL,
   icc1 = NULL,
   Rsq = 0,
   alpha = 0.05,
   power = NULL,
   sides = 2,
   v = FALSE
)
```

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Arguments

| m | The total number of subjects per site or the mean cluster size (if unequal number of participants per site). |
|-------------|---|
| m.sd | The standard deviation of cluster sizes (provide if unequal number of participants per site); defaults to 0. |
| alloc.ratio | The allocation ratio of intervention/control per site; defaults to 1. |
| J | The number of sites. |
| delta | The difference between the intervention and control means under the alternative minus the difference under the null hypothesis. |
| sd | The total standard deviation of the outcome variable; defaults to 1. |
| icc0 | The proportion of total variance of the outcome attributable to variation in site- level means. |
| icc1 | The proportion of total variance of the outcome attributable to variation in the treatment effect across sites. |
| Rsq | The estimated R^2 for regressing the outcome on the covariates; defaults to 0. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

Examples

multisite.hte

Power for test of heterogeneity of treatment effect in multisite trials

Description

Power for test of heterogeneity of treatment effect in multisite trials

```
multisite.hte(
  m = NULL,
  alloc.ratio = 1,
  J = NULL,
  VR = NULL,
  alpha = 0.05,
  v = FALSE
```

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Arguments

m The total number of subjects per site.

alloc.ratio The allocation ratio of intervention/control per site; defaults to 1.

J The number of sites.

VR The variance ratio (site-level treatment effects / observations within sites) the under the alternative.

alpha The significance level or type 1 error rate; defaults to 0.05.

v Either TRUE for verbose output or FALSE to output computed argument only.

Value

A list of the arguments (including the computed power).

Examples

```
multisite.hte(m = 10, J = 30, VR = 8 / 36)
```

prop.1samp Power calculations for one sample proportion tests

Description

Power calculations for one sample proportion tests

Usage

```
prop.1samp(
  N = NULL,
  p0 = NULL,
  pA = NULL,
  alpha = 0.05,
  power = NULL,
  sides = 2,
  v = FALSE
)
```

Arguments

| N | The sample size. |
|-------|---|
| p0 | The proportion under the null. |
| pA | The true proportion. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

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Examples

```
prop.1samp(N = NULL, p0 = 0.2, pA = 0.3, power = 0.8, sides = 1) prop.1samp(N = NULL, p0 = 0.4, pA = 0.5, power = 0.8, sides = 1)
```

prop.2samp

Power calculations for two sample proportion tests

Description

Power calculations for two sample proportion tests

Usage

```
prop.2samp(
  n1 = NULL,
  n.ratio = 1,
  p1 = NULL,
  p2 = NULL,
  margin = 0,
  alpha = 0.05,
  power = NULL,
  sides = 2,
  v = FALSE
)
```

Arguments

| n1 | The sample size for group 1. |
|---------|---|
| n.ratio | The ratio n2/n1 between the sample sizes of two groups; defaults to 1 (equal group sizes). |
| p1 | The proportion in group 1. |
| p2 | The proportion in group 2. |
| margin | The margin of noninferiority or superiority; defaults to 0. See margin.sign for guidance on the sign of margin. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
prop.2samp(n1 = NULL, p1 = 0.6, p2 = 0.8, alpha = 0.025, power = 0.9, sides = 1) prop.2samp(n1 = NULL, p1 = 0.25, p2 = 0.25, margin = 0.1, alpha = 0.025, power = 0.8, sides = 1)
```

prop.paired 39

| prop.paired | Power approximation for McNemar's test for two correlated proportions |
|-------------|---|
| prop.paired | |

Description

Power approximation for McNemar's test for two correlated proportions

Usage

```
prop.paired(
  N = NULL,
  p1 = NULL,
  p2 = NULL,
  phi = NULL,
  dpr = NULL,
  alpha = 0.05,
  power = NULL,
  sides = 2,
  v = FALSE
)
```

Arguments

| N | The sample size; the number of pairs. |
|-------|---|
| p1 | The proportion in condition 1. |
| p2 | The proportion in condition 2. |
| phi | The estimated correlation between the two conditions. |
| paid | The smaller of the two discordant probabilities. Either p1, p2, and phi, OR paid and dpr must be specified. |
| dpr | The discordant proportion ratio. Either p1, p2, and phi, OR paid and dpr must be specified. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
prop.paired(N = NULL, p1 = 0.8, p2 = 0.9, phi = 0, power = 0.9, sides = 2) prop.paired(N = NULL, paid = 0.08, dpr = 0.18 / 0.08, power = 0.9, sides = 2)
```

40 prop.test.equiv

| prop. | test | Palli V |
|-------|------|---------|

Power calculations for test of equivalence of two proportions

Description

Power calculations for test of equivalence of two proportions

Usage

```
prop.test.equiv(
  n1 = NULL,
  n.ratio = 1,
  p1 = NULL,
  p2 = NULL,
  margin = NULL,
  alpha = 0.05,
  power = NULL,
  sides = 2,
  v = FALSE
)
```

Arguments

| n1 | The sample size for group 1. |
|---------|--|
| n.ratio | The ratio n2/n1 between the sample sizes of two groups; defaults to 1 (equal group sizes). |
| p1 | The outcome proportion in group 1. |
| p2 | The outcome proportion in group 2. |
| margin | The equivalence margin. See margin.sign for guidance on the sign of margin. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
prop.test.equiv(n1 = NULL, p1 = 0.5, p2 = 0.5, margin = 0.1, alpha = 0.05, power = 0.8, sides = 1)
```

propodds 41

| propodds | Power calculations for ordinal categorical variable under proportional odds assumption |
|----------|--|
| | • |

Description

Power calculations for ordinal categorical variable under proportional odds assumption

Usage

```
propodds(pC, OR, n1, n.ratio = 1, alpha = 0.05, power = NULL, v = FALSE)
```

Arguments

| pC | Vector of response probabilities in control group (group 1). Must sum to 1. Categories are ordered from best to worst. |
|---------|--|
| OR | Odds ratio when the alternative is true. Must be greater than 1. |
| n1 | Sample size for group 1 (control group). |
| n.ratio | The ratio n2/n1 between the sample sizes of two groups; defaults to 1 (equal group sizes). |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power; defaults to 0.8. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

Examples

```
library(Hmisc) 
 pC <- c(0.2, 0.5, 0.2, 0.1) 
 propodds(pC = pC, OR = 2.5, n1 = 65, n.ratio = 1, alpha = 0.05)
```

| ranksum | Power calculations for rank-sum test |
|---------|--------------------------------------|
| | |

Description

Power calculations for rank-sum test

re.clustsize.bin

Usage

```
ranksum(
  n1 = NULL,
  n.ratio = 1,
  p = NULL,
  alpha = 0.05,
  power = NULL,
  sides = 2,
  v = FALSE
)
```

Arguments

| n1 | The sample size in group 1. |
|---------|--|
| n.ratio | The ratio n2/n1 between the sample sizes of two groups; defaults to 1 (equal group sizes). |
| р | The probability that an observation in group 2 is greater than an observation in group 1 ($P(Y>X)$). |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two-sided hypothesis test. |
| ٧ | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

Examples

```
ranksum(n1 = 10, n.ratio = 1, p = 0.8, alpha = 0.05, power = NULL, sides = 2)
```

| re.clustsize.bin | Relative efficiency for cluster randomized trials with binary outcomes |
|------------------|--|
| | due to varying cluster sizes |

Description

Relative efficiency for cluster randomized trials with binary outcomes due to varying cluster sizes

Usage

```
re.clustsize.bin(m, m.sd, pc, pt, sigma.u)
```

re.clustsize.cont 43

Arguments

| m | The number of subjects per cluster or the mean cluster size (if unequal number of participants per cluster). |
|---------|--|
| m.sd | The standard deviation of cluster sizes (provide if unequal number of participants per cluster). |
| рс | The probability of the outcome in control clusters. |
| pt | The probability of the outcome in treatment clusters. |
| sigma.u | Standard deviation of the cluster random effect. |

Value

The computed RE.

Examples

```
re.clustsize.bin(m = 60, m.sd = 45, pc = 0.25, pt = 0.15, sigma.u = 0.3)
```

| re.clustsize.cont | Relative efficiency for cluster randomized or multisite trials due to |
|-------------------|---|
| | varying cluster sizes |

Description

Relative efficiency for cluster randomized or multisite trials due to varying cluster sizes

Usage

```
re.clustsize.cont(m, m.sd, icc)
```

Arguments

| m | The number of subjects per cluster or the mean cluster size (if unequal number of participants per cluster). |
|------|---|
| m.sd | The standard deviation of cluster sizes (provide if unequal number of participants per cluster). |
| icc | The intraclass correlation coefficient. For a multisite trial this is icc1. For a CRT this is the average of the 2 icc's. |

Value

The computed RE.

```
re.clustsize.cont(m = 25, m.sd = 15, icc = 0.05)
```

44 relrisk

relrisk

 $Power\ calculations\ for\ relative\ risk$

Description

Power calculations for relative risk

Usage

```
relrisk(
  n1 = NULL,
  n.ratio = 1,
  p1 = NULL,
  p2 = NULL,
  RR0 = 1,
  alpha = 0.05,
  power = NULL,
  sides = 2,
  v = FALSE
)
```

Arguments

| n1 | The sample size for group 1. |
|---------|--|
| n.ratio | The ratio $n2/n1$ between the sample sizes of two groups; defaults to 1 (equal group sizes). |
| p1 | The proportion in group 1. |
| p2 | The proportion in group 2. |
| RRØ | The relative risk under the null; defaults to 1. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
relrisk(n1 = NULL, n.ratio = 1/6, p1 = 0.1, p2 = 0.1 * 2, power = 0.8)
```

signedrank 45

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| | | | |

Power calculation for signed-rank test

Description

The signed-rank test is a nonparametric alternative to a one-sample or paired t test. This function performs power and sample size calculations for the signed-rank test using Noether's approximation; see Noether (1987) Sample size determination for some common nonparametric tests. JASA 82(398):645-647.

Usage

```
signedrank(
  N = NULL,
  ps = NULL,
  alpha = 0.05,
  power = NULL,
  sides = 2,
  v = FALSE
)
```

Arguments

| N | The sample size; number of observations or paired differences. |
|-------|--|
| ps | The probability that the sum of two values exceeds zero when the alternative hypothesis is true. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two-sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

Examples

```
signedrank(N = 20, ps = 0.87, power = NULL, sides = 2)
```

signtest

Power calculation for sign test (one-sample test for a median)

Description

The sign test is a one-sample test of location, specifically, a test of whether the median equals (or is less than or greater than) zero. It is used in place of a one-sample or paired t test. This function performs power and sample size calculations for a sign test.

46 slr

Usage

```
signtest(N = NULL, p = NULL, alpha = 0.05, power = NULL, sides = 2, v = FALSE)
```

Arguments

| N | The sample size. |
|-------|---|
| р | The probability of a positive difference when the alternative hypothesis is true. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two-sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

Examples

```
signtest(N = 40, p = 0.7, power = NULL, alpha = 0.05, sides = 1)
```

slr

Power calculations for a simple linear regression

Description

Power calculations for a simple linear regression

Usage

```
slr(
  N = NULL,
  beta10 = 0,
  beta1A = NULL,
  sd.x.sq = NULL,
  sigma.e = NULL,
  alpha = 0.05,
  power = NULL,
  sides = 2,
  v = FALSE
)
```

Arguments

| N | The sample size. |
|---------|--|
| beta10 | The slope regression coefficient under the null hypothesis; defaults to 0. |
| beta1A | The slope regression coefficient under the alternative hypothesis. |
| sd.x.sq | The sample variance of the covariate X. |
| sigma.e | The estimated standard deviation of the error terms. |

swd_1trt_power 47

| alpha | The significance level or type 1 error rate; defaults to 0.05. |
|-------|---|
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

Examples

```
# Yi = beta0 + beta1 * Xi + ei, i = 1,...,N slr(N = 100, beta10 = 1, beta1A = 1.5, sd.x.sq = 25, sigma.e = 10, sides = 1)
```

 swd_1trt_power

Stepped Wedge Design Power Calculation with 1 Treatment

Description

Calculates power for stepped wedge design with one treatment and one control condition.

Usage

```
swd_1trt_power(alpha = 0.05, J, m, K, b, d, rho, rho_c, rho_s)
```

Arguments

| alpha | The significance level or type 1 error rate |
|-------|---|
| J | Number of clusters |
| m | The number of individuals in each cluster at each time period |
| K | The number of steps (if one baseline period, then this is periods - 1) |
| b | The number of baseline periods |
| d | The standardized effect size |
| rho | Intraclass correlation coefficient (ICC) |
| rho_c | Cluster autocorrelation (proportion of cluster-level variance that is time invariant) |
| rho_s | Subject autocorrelation (This is 0 in repeated cross sectional designs because people are measured only once) |

Value

power

48 swd_1trt_ss

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|--------|------|----|
| swd 1 | ltrt | SS |

Stepped Wedge Design Sample Size Calculation with 1 Treatment

Description

Calculates sample size requirements for stepped wedge design with one treatment and one control condition.

Usage

```
swd_1trt_ss(alpha, power, m, K, b, d, rho, rho_c, rho_s)
```

Arguments

| alpha | The significance level or type 1 error rate |
|-------|---|
| power | The specified level of power |
| m | The number of individuals in each cluster at each time period |
| K | The number of steps (if one baseline period, then this is periods - 1) |
| b | The number of baseline periods |
| d | The standardized effect size |
| rho | Intraclass correlation coefficient (ICC) |
| rho_c | Cluster autocorrelation (proportion of cluster-level variance that is time invariant) |
| rho_s | Subject autocorrelation (This is 0 in repeated cross sectional designs because people are measured only once) |

Value

Calculated number of clusters required, adjusted number of clusters, and suggested number of clusters that includes the value of \boldsymbol{K}

```
swd_2trt_additive_power
```

Additive Treatments Stepped Wedge Design Power Calculation

Description

Calculate power for detecting treatment effects in a two-treatment stepped wedge design with additive treatment effects. We only consider complete designs. This function is based on Power Analysis for Stepped Wedge Trials with Two Treatments published in Stat Med in 2022 by Phillip Sundin and Catherine Crespi. This manuscript uses the normal distribution and the Wald Test to derive power.

Usage

```
swd_2trt_additive_power(
  RhoW,
  ModelChoice,
  n.individuals,
  n.clusters,
  n.periods,
  delta_1,
  delta_2,
  RhoA,
  IAC,
  Sequence_Tx1,
  Sequence_Tx2,
  typeIerror
)
```

Arguments

| RhoW The intraclass correlation coefficient, which re | efers to the correlation between |
|---|----------------------------------|
|---|----------------------------------|

outcomes of two different individuals in the same cluster at the same time.

ModelChoice A string indicating either a repeated cross sectional design ("RCS"), nested ex-

changeable ("NEC"), Cohort ("Cohort"), or any one of these three designs in their standardized form. Only one of "RCS Standardized", "RCS", "Cohort Standardized", "Cohort", "NEC Standardized", or "NEC" can be specified. The standardized design refers to dividing by the standard deviation of y (the out-

come) in the covariance specification.

n.individuals The number of individuals in each condition at one time period.

n.clustersn.periodsThe number of clustersn.periods

delta_1 The standardized effect size for treatment 1 delta_2 The standardized effect size for treatment 2

RhoA The correlation between two observations in the same individual in the same

cluster but different time periods. RCS models assume RhoW = RhoA, and

Cohort models assume RhoA >= RhoW.

IAC The individual auto-correlation, which is the proportion of the individual-level

variance that is time-invariant. IAC = 0 in a RCS.

Sequence_Tx1 A vector representing the time points (periods) at which each cluster transitions from control to treatment 1 (ie the first occurence).

Sequence_Tx2 A vector representing the time points (periods) at which each cluster transitions from control to treatment 2 (ie the first occurence).

typeIerror The significance level.

Value

A list with power for detecting treatment effect for treatment 1 and 2

Examples

```
## For example, the 3 cluster, 4 period SWD,
##
      0 1 1 1
##
      0 0 1+2 1+2
##
      0 0 2 1+2
## where 0 denotes control condition, 1 denotes the condition with only
## treatment 1, 2 denote the condition with only treatment 2, and 1+2 denotes the condition
## where a cluster receives both treatment 1 and 2
## would have Sequence1 <- c(2,3,4) and Sequence2 <- c(NA,3,3)
## notice sequence 1 cluster 3 has time period 4 even though time period 3 had treatment 2,
## so it is the first occurrence of trt 1 in this cluster.
# 12-cluster concurrent repeated cross sectional design
# Sequencing has 2 clusters transition to treatment 1 at time 2,
# 2 clusters transition to treatment 1 at time 3, and 2 clusters transition
# to treatment 1 at time 3. These clusters never receive treatment 2
Sequence_Tx1 <- c(2, 2, 3, 3, 4, 4, NA, NA, NA, NA, NA, NA, NA)
# similar sequencing for treatment 2
Sequence_Tx2 <- c(NA, NA, NA, NA, NA, NA, 4, 4, 3, 3, 2, 2)
swd_2trt_additive_power(RhoW = 0.05, ModelChoice = "RCS", RhoA = 0.2, IAC = 0,
          n.individuals = 30, n.clusters = 12, n.periods = 4, delta_1 = 0.4, delta_2 = 0.1,
               Sequence_Tx1 = Sequence_Tx1, Sequence_Tx2 = Sequence_Tx2,
               typeIerror = 0.05)
```

swd_2trt_interaction_power

Stepped Wedge Design Power Calculation for Interaction Term

Description

Calculate power for detecting interaction effects in a two-treatment stepped wedge design. We only consider complete designs. This function is based on Power Analysis for Stepped Wedge Trials with Two Treatments published in Stat Med in 2022 by Phillip Sundin and Catherine Crespi. This manuscript uses the normal distribution and the Wald Test to derive power.

Usage

```
swd_2trt_interaction_power(
  RhoW,
  ModelChoice,
  n.individuals,
```

```
n.clusters,
n.periods,
delta_1,
delta_2,
delta_3,
RhoA,
IAC,
typeIerror,
Sequence_Tx1,
Sequence_Tx2
```

Arguments

RhoW The intraclass correlation coefficient, which refers to the correlation between

outcomes of two different individuals in the same cluster at the same time.

ModelChoice A string indicating either a repeated cross sectional design ("RCS"), nested ex-

changeable ("NEC"), Cohort ("Cohort"), or any one of these three designs in their standardized form. Only one of "RCS Standardized", "RCS", "Cohort Standardized", "Cohort", "NEC Standardized", or "NEC" can be specified. The standardized design refers to dividing by the standard deviation of y (the out-

come) in the covariance specification.

n.individuals The number of individuals in each condition at one time period.

n.clusters The number of clusters

n.periods The number of time periods.

delta_1 The standardized effect size for treatment 1 delta_2 The standardized effect size for treatment 2

delta_3 The standardized effect size for the interaction effect.

RhoA The correlation between two observations in the same individual in the same

cluster but different time periods. RCS models assume RhoW = RhoA, and

Cohort models assume RhoA >= RhoW.

IAC The individual auto-correlation, which is the proportion of the individual-level

variance that is time-invariant. IAC = 0 in a RCS.

typeIerror The significance level.

Sequence_Tx1 A vector representing the time points (periods) at which each cluster transitions

from control to treatment 1 (ie the first occurence).

Sequence_Tx2 A vector representing the time points (periods) at which each cluster transitions

from control to treatment 2 (ie the first occurence).

Value

A matrix with the standard error, statistic, and power for detecting effects for treatment 1, treatment 2, and interaction effects.

```
delta_2 = 0.6, delta_3 = 0.6, typeIerror = 0.05/3,
Sequence_Tx1 = c(2,3,2,3,4,5,NA,NA), Sequence_Tx2 = c(NA,NA,2,3,4,5,3,2))
```

```
swd_2trt_linearcontrast_power
```

Stepped Wedge Design Power Calculation for Linear Contrasts of Treatment Effects

Description

Calculate the power for detecting linear contrasts of two treatment effects in a two-treatment stepped wedge design (SWD) with additive treatment effects. We only consider complete designs. This function is based on Power Analysis for Stepped Wedge Trials with Two Treatments published in Stat Med in 2022 by Phillip Sundin and Catherine Crespi. This manuscript uses the normal distribution and the Wald Test to derive power.

Usage

```
swd_2trt_linearcontrast_power(
  RhoW,
  ModelChoice,
  n.individuals,
  n.clusters,
  n.periods,
  delta_12,
  RhoA,
  IAC,
  Sequence_Tx1,
  Sequence_Tx2,
  typeIerror
)
```

Arguments

RhoW The intraclass correlation coefficient, which refers to the correlation between

outcomes of two different individuals in the same cluster at the same time.

ModelChoice A string indicating either a repeated cross sectional design ("RCS"), nested ex-

changeable ("NEC"), Cohort ("Cohort"), or any one of these three designs in their standardized form. Only one of "RCS Standardized", "RCS", "Cohort Standardized", "Cohort", "NEC Standardized", or "NEC" can be specified. The standardized design refers to dividing by the standard deviation of y (the out-

come) in the covariance specification.

n.individuals The number of individuals in each condition at one time period.

n.clusters The number of clustersn.periods The number of time periods.

delta_12 The standardized difference between the two treatment effects to be powered on.

RhoA The correlation between two observations in the same individual in the same

cluster but different time periods. RCS models assume RhoW = RhoA, and

Cohort models assume RhoA >= RhoW.

| IAC | The individual auto-correlation, which is the proportion of the individual-level variance that is time-invariant. IAC = 0 in a RCS. |
|--------------|---|
| Sequence_Tx1 | A vector representing the time points (periods) at which each cluster transitions from control to treatment 1 (ie the first occurence). |
| Sequence_Tx2 | A vector representing the time points (periods) at which each cluster transitions from control to treatment 2 (ie the first occurence). |
| typeIerror | The significance level. |

Value

Power for detecting the difference of treatment effects

```
# Late factorial SWD design with 12 clusters
swd_2trt_linearcontrast_power(RhoW = 0.2, ModelChoice="RCS", IAC = 0,
                      RhoA = 0.2, n.individuals = 15, delta_12 = 0.4,
                      n.clusters = 12, n.periods = 4,
                      Sequence_Tx1 = c(2,2,3,3,4,4,4,4,4,4,4,4),
                      Sequence_Tx2 = c(4,4,4,4,4,4,2,2,3,3,4,4),
                      typeIerror = 0.05)
\# Example where there are different potential designs to be considered \#
#creating data frame for power as a function of rho_w
PowerTable <- data.frame(RhoW = c(rep(seq(0,0.4,by=0.01),times=3)))
PowerTable$DesignChoice <- c(rep('"Late" Factorial Design, 12 clusters',times=41),</pre>
                             rep('"Early" Factorial Design, 10 clusters',times=41),
                              rep("Concurrent Design, 12 clusters",times=41))
PowerTable$n.clusters <- c(rep(12,times=41), rep(10,times=41), rep(12,times=41))
PowerTableSequence1 \leftarrow c(rep(list(c(2,2,3,3,4,4,4,4,4,4,4,4)),times=41),
                          rep(list(c(2,2,2,3,4,4,3,4,4,4)),times=41),
                          rep(list(c(2,2,3,3,4,4,NA,NA,NA,NA,NA,NA)),times=41))
PowerTableSequence2 <- c(rep(list(c(4,4,4,4,4,4,2,2,3,3,4,4)),times=41),
                          rep(list(c(4,4,4,3,4,4,3,2,2,2)),times=41),
                          rep(list(c(NA,NA,NA,NA,NA,4,4,3,3,2,2)),times=41))
PowerTable$Power <- mapply(swd_2trt_linearcontrast_power,</pre>
                        RhoW = PowerTable$RhoW,
                        ModelChoice="RCS",
                        IAC = 0, RhoA = PowerTable$RhoW,
                        n.individuals = 15, delta_12 = 0.4,
                        n.clusters = PowerTable$n.clusters,
                        n.periods = 4, typeIerror = 0.05,
                        Sequence_Tx1 = PowerTable$Sequence1,
                        Sequence_Tx2 = PowerTable$Sequence2)
##removing unnecessary columns of clusters
PowerTable <- PowerTable[ ,-which(names(PowerTable) %in%</pre>
            c("Sequence1", "Sequence2", "n.clusters"))]
PowerTable
```

54 t.test.1samp

t.test.1samp

Power calculations for one sample t tests

Description

Power calculations for one sample t tests

Usage

```
## S3 method for class 'test.1samp'
t(
   N = NULL,
   delta = NULL,
   sd = 1,
   alpha = 0.05,
   power = NULL,
   sides = 2,
   v = FALSE
)
```

Arguments

| N | The sample size. |
|-------|---|
| delta | muA (the true mean) - mu0 (the mean under the null). |
| sd | The estimated standard deviation; defaults to 1. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
t.test.1samp(N = 36, delta = 4.9 - 5.7, sd = 2, sides = 1)
t.test.1samp(N = 36, delta = 6.3 - 5.7, sd = 2, sides = 1)
t.test.1samp(N = 36, delta = 4.9 - 5.7, sd = 2, sides = 2)
t.test.1samp(delta = 0.6, sd = 1, power = 0.8, sides = 1)
```

t.test.2samp 55

| t.test.2samp | Power calculations for two sample t tests allowing for unequal sample sizes and/or variances |
|--------------|--|
| | sizes and or variances |

Description

Power calculations for two sample t tests allowing for unequal sample sizes and/or variances

Usage

```
## S3 method for class 'test.2samp'
t(
    n1 = NULL,
    n.ratio = 1,
    delta = NULL,
    sd1 = 1,
    sd.ratio = 1,
    df.method = c("welch", "classical"),
    alpha = 0.05,
    power = NULL,
    sides = 2,
    v = FALSE
)
```

Arguments

| n1 | The sample size for group 1. |
|-----------|---|
| n.ratio | The ratio n2/n1 between the sample sizes of two groups; defaults to 1 (equal group sizes). |
| delta | DeltaA (the true difference mu1 - mu2) - Delta0 (the difference under the null) - margin. See margin.sign for guidance on the sign of the margin. |
| sd1 | The estimated standard deviation for group 1; defaults to 1 (equal standard deviations in the two groups). |
| sd.ratio | The ratio sd2/sd1 between the standard deviations of the two groups. |
| df.method | Method for calculating the degrees of freedom: "welch" (default) or "classical". |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
t.test.2samp(n1 = 50, delta = 2, sd1 = 5, sides = 1)
t.test.2samp(n1 = NULL, n.ratio = 2, delta = 0.5, sd1 = 1, power = 0.8, sides = 2)
t.test.2samp(n1 = 49, n.ratio = 2, delta = 0.5, sd1 = 1, power = NULL, sides = 2)
t.test.2samp(n1 = 25, n.ratio = 3, delta = 3, sd1 = 4, sd.ratio = 1.5, alpha = 0.025, sides = 1)
t.test.2samp(n1 = NULL, delta = 0.5, sd1 = 1, power = 0.8, sides = 2)
```

56 t.test.paired

t.test.paired

Power calculations for paired t tests

Description

Power calculations for paired t tests

Usage

```
## S3 method for class 'test.paired'
t(
   N = NULL,
   delta = NULL,
   sd1 = 1,
   sd2 = 1,
   rho = NULL,
   alpha = 0.05,
   power = NULL,
   sides = 2,
   v = FALSE
)
```

Arguments

| N | The sample size; the number of pairs. |
|-------|--|
| delta | $Delta A \ (the \ true \ mean \ difference) - Delta 0 \ (the \ mean \ difference \ under \ the \ null).$ |
| sd1 | The estimated pre standard deviation; defaults to 1. |
| sd2 | The estimated post standard deviation; defaults to 1. |
| rho | The estimated correlation between pre and post measurements on the same individual. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
t.test.paired(N = NULL, delta = 4, sd1 = 10, sd2 = 10, rho = 0.4, power = 0.8, sides = 2)
```

z.test.1samp 57

z.test.1samp

Power calculation for one sample z test

Description

This function performs power and sample size calculations for a one-sample z test, which is analogous to a one-sample t test with the variance assumed to be known.

Usage

```
z.test.1samp(
  N = NULL,
  delta = NULL,
  sd = 1,
  alpha = 0.05,
  power = NULL,
  sides = 2,
  v = FALSE
)
```

Arguments

| N | The sample size. |
|-------|---|
| delta | muA (the true mean) - mu0 (the mean under the null). |
| sd | The estimated standard deviation; defaults to 1. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
z.test.1samp(N = NULL, delta = 6.5 - 5.7, sd = 2, power = 0.8, sides = 2) z.test.1samp(N = 40, delta = NULL, sd = 1, power = 0.9, sides = 2) z.test.1samp(N = NULL, delta = 0.6, sd = 1, power = 0.8, sides = 1)
```

z.test.2samp

z.test.2samp

Power calculation for two-sample z test

Description

This function performs power and sample size calculations for a two-sample z test, which is analogous to a two-sample t test but with the variances assumed to be known. The function allows for unequal sample sizes and/or variances in the two groups.

Usage

```
z.test.2samp(
  n1 = NULL,
  n.ratio = 1,
  delta = NULL,
  sd1 = 1,
  sd.ratio = 1,
  alpha = 0.05,
  power = NULL,
  sides = 2,
  v = FALSE
)
```

Arguments

| n1 | The sample size for group 1. |
|----------|--|
| n.ratio | The ratio $n2/n1$ between the sample sizes of two groups; defaults to 1 (equal group sizes). |
| delta | $\label{eq:DeltaA} DeltaA \ (the \ true \ difference \ mu1 - mu2) - DeltaO \ (the \ difference \ under \ the \ null) - margin See \ margin.sign \ for \ guidance \ on \ the \ sign \ of \ the \ margin.$ |
| sd1 | The estimated standard deviation for group 1; defaults to 1 (equal standard deviations in the two groups). |
| sd.ratio | The ratio sd2/sd1 between the standard deviations of the two groups. |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
z.test.2samp(n1 = NULL, n.ratio = 1, delta = 0.5, sd1 = 1, power = 0.8, sides = 2)
```

z.test.paired 59

| z.test.paired Power calculation for paired z test | z.test.paired | Power calculation for paired z test | |
|---|---------------|-------------------------------------|--|
|---|---------------|-------------------------------------|--|

Description

This function performs power and sample size calculations for a paired z test, which is analogous to a paired t test with variance assumed to be known.

Usage

```
z.test.paired(
  N = NULL,
  delta = NULL,
  sd1 = 1,
  sd2 = 1,
  rho = NULL,
  alpha = 0.05,
  power = NULL,
  sides = 2,
  v = FALSE
)
```

Arguments

| N | The sample size; the number of pairs. |
|-------|--|
| delta | $Delta A \ (the \ true \ mean \ difference) - Delta 0 \ (the \ mean \ difference \ under \ the \ null).$ |
| sd1 | The estimated pre standard deviation; defaults to 1. |
| sd2 | The estimated post standard deviation; defaults to 1. |
| rho | The estimated correlation between pre and post measurements on the same individual; defaults to 0 . |
| alpha | The significance level or type 1 error rate; defaults to 0.05. |
| power | The specified level of power. |
| sides | Either 1 or 2 (default) to specify a one- or two- sided hypothesis test. |
| V | Either TRUE for verbose output or FALSE to output computed argument only. |

Value

A list of the arguments (including the computed one).

```
z.test.paired(N = NULL, delta = 4, sd1 = 10, sd2 = 10, rho = 0.4, power = 0.8, sides = 2)
```

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