Package 'FDRsamplesize2'

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Title Computing Power and Sample Size for the False Discovery Rate in

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Multiple Applications

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Description

Given the proportion pi0 of tests with a true null, find the p-value threshold that results in a desired FDR and average power.

```
alpha.power.fdr(fdr, pwr, pi0, method = "HH")
```

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Arguments

fdr	desired FDR	(scalar numeric)

pwr desired average power (scalar numeric)

pi0 the proportion of tests with a true null hypothesis

method method to estimate proportion pi0 of tests with true null, including: "HH" (p-

value histogram height), "HM" (p-value histogram mean), "BH" (Benjamini &

Hochberg 1995), "Jung" (Jung 2005)

Details

To get the fixed p-value threshold for multiple testing procedure, 4 approximation methods are provided, they are Benjamini & Hochberg procedure (1995), Jung's formula (2005), method of using p-value histogram height (HH) and method of using p-value histogram mean (HM). For last two methods' details, see Ni Y, Onar-Thomas A, Pounds S. "Computing Power and Sample Size for the False Discovery Rate in Multiple Applications"

Value

The fixed p-value threshold for multiple testing procedure

References

Pounds S and Cheng C, "Sample size determination for the false discovery rate." Bioinformatics 21.23 (2005): 4263-4271.

Gadbury GL, et al. (2004) Power and sample size estimation in high dimensional biology. Statistical Methods in Medical Research 13(4):325-38.

Jung, Sin-Ho. "Sample size for FDR-control in microarray data analysis." Bioinformatics 21.14 (2005): 3097-3104.

Ni Y, Seffernick A, Onar-Thomas A, Pounds S. "Computing Power and Sample Size for the False Discovery Rate in Multiple Applications", Manuscript.

Examples

```
alpha.power.fdr(fdr = 0.1, pwr = 0.9, pi0=0.9, method = "HH")
```

average.power.coxph

Compute the average power of many Cox regression models

Description

Compute the average power of many Cox regression models for a given number of events, p-value threshold, vector of effect sizes (log hazard ratio), and variance of predictor variables

```
average.power.coxph(n, alpha, logHR, v)
```

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Arguments

n	number of events (scalar)
alpha	p-value threshold (scalar)
logHR	log hazard ratio (vector)

v variance of predictor variable (vector)

Value

Average power estimate for multiple testing procedure

References

Hsieh, FY and Lavori, Philip W (2000) Sample-size calculations for the Cox proportional hazards regression model with non-binary covariates. Controlled Clinical Trials 21(6):552-560.

See Also

power.cox for more details about power calculation of single-predictor Cox regression model. The power calculation is based on asymptotic normal approximation.

Examples

```
logHR = log(rep(c(1, 2), c(900, 100)));
v = rep(1, 1000);
average.power.coxph(n = 50, alpha = 0.05, logHR = logHR, v = v)
```

average.power.fisher Compute average power of many Fisher's exact tests

Description

Compute average power of many Fisher's exact tests

Usage

```
average.power.fisher(p1, p2, n, alpha, alternative)
```

Arguments

p1 probability in one group (vector) p2 probability in other group (vector)

n per-group sample size
alpha p-value threshold
alternative one- or two-sided test

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Value

Average power estimate for multiple testing procedure

See Also

power. fisher for more details about power calculation of Fisher's exact test

Examples

```
set.seed(1234);
p1 = sample(seq(0,0.5,0.1),5,replace = TRUE);
p2 = sample(seq(0.5,1,0.1),5,replace = TRUE);
average.power.fisher(p1 = p1,p2 = p2,n = 20,alpha = 0.05,alternative = "two.sided")
```

average.power.hart

Compute average power for RNA-seq experiments assuming Negative Binomial distribution

Description

Compute average power for RNA-seq experiments assuming Negative Binomial distribution

Usage

```
average.power.hart(n, alpha, log.fc, mu, sig)
```

Arguments

n	per-group sample size (scalar)
alpha	p-value threshold (scalar)
log.fc	log fold-change (vector), usual null hypothesis is log.fc=0
mu	read depth per gene (vector, same length as log.fc)
sig	coefficient of variation (CV) per gene (vector, same length as log.fc)

Details

The power function is based on equation (1) of Hart et al (2013). It assumes a Negative Binomial model for RNA-seq read counts and equal sample size per group.

Value

Average power estimate for multiple testing procedure

References

SN Hart, TM Therneau, Y Zhang, GA Poland, and J-P Kocher (2013). Calculating Sample Size Estimates for RNA Sequencing Data. Journal of Computational Biology 20: 970-978.

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See Also

power.hart for more details about power calculation of data under Negative Binomial distribution. The power calculation is based on asymptotic normal approximation.

Examples

```
logFC = log(rep(c(1,2),c(900,100)));

mu = rep(5,1000);

sig = rep(0.6,1000);

average.power.hart(n = 50, alpha = 0.05,log.fc = logFC, mu = mu, sig = sig)
```

average.power.li

Compute average power for RNA-Seq experiments assuming Poisson distribution

Description

Use the formula of Li et al (2013) to compute power for comparing RNA-seq expression across two groups assuming the Poisson distribution.

Usage

```
average.power.li(n, alpha, rho, mu0, w, type)
```

Arguments

n	per-group sample size
alpha	p-value threshold (scalar)
rho	fold-change, usual null hypothesis is that rho=1 (vector)
mu0	average count in control group (vector)
W	ratio of the total number of reads mapped between the two groups (scalar or vector)
type	type of test: "w" for Wald, "s" for score, "lw" for log-transformed Wald, "ls" for

Details

This function computes the average power for a series of two-sided tests defined by the input parameters. The power is based on the sample size formulas in equations (10-13) of Li et al (2013). Also, note that the null effect is set to 1 in the examples because the usual null hypothesis is that the fold-change = 1.

Value

Average power estimate for multiple testing procedure

log-transformed score

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References

C-I Li, P-F Su, Y Guo, and Y Shyr (2013). Sample size calculation for differential expression analysis of RNA-seq data under Poisson distribution. Int J Comput Biol Drug Des 6(4).<doi:10.1504/IJCBDD.2013.056830>

See Also

power.li for more details about power calculation of data under Poisson distribution

Examples

```
rho = rep(c(1,1.25),c(900,100));
mu0 = rep(5,1000);
w = rep(0.5,1000);
average.power.li(n = 50, alpha = 0.05, rho = rho, mu0 = mu0, w = w, type = "w")
```

average.power.oneway

Compute average power of many one-way ANOVA tests

Description

Compute average power of many one-way ANOVA tests

Usage

```
average.power.oneway(n, alpha, theta, k)
```

Arguments

n per-group sample size (scalar) alpha p-value threshold (scalar)

theta sum of ((group mean - overall mean)/stdev)^2 across all groups for each hypoth-

esis test(vector)

k the number of groups to be compared

Value

Average power estimate for multiple testing procedure

See Also

power.oneway for more details about power calculation of one-way ANOVA

```
theta=rep(c(2,0),c(100,900));
average.power.oneway(n = 50, alpha = 0.05, theta = theta, k = 2)
```

average.power.ranksum Compute average power of rank-sum tests

Description

Compute average power of rank-sum tests

Usage

```
average.power.ranksum(n, alpha, p)
```

Arguments

n sample size (scalar)
alpha p-value threshold (scalar)
p Pr(Y>X), as in Noether (JASA 1987)

Value

Average power estimate for multiple testing procedure

See Also

power.ranksum for more details about power calculation of rank-sum test. The power calculation is based on asymptotic normal approximation.

Examples

```
p = rep(c(0.8,0.5),c(100,900));
average.power.ranksum(n = 50, alpha = 0.05, p=p)
```

average.power.signrank

Compute average power of many signed-rank tests

Description

Compute average power of many signed-rank tests

```
average.power.signrank(n, alpha, p1, p2)
```

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Arguments

n	sample size (scalar)
alpha	p-value threshold (scalar)
p1	Pr(X>0), as in Noether (JASA 1987)
p2	Pr(X+X'>0), as in Noether (JASA 1987)

Value

Average power estimate for multiple testing procedure

See Also

power.signrank for more details about power calculation of signed-rank test. The power calculation is based on asymptotic normal approximation.

Examples

```
p1 = rep(c(0.8,0.5),c(100,900));

p2 = rep(c(0.8,0.5),c(100,900));

average.power.signrank(n = 50, alpha = 0.05, p1 = p1, p2 = p2)
```

average.power.signtest

Compute average power of many sign tests

Description

Compute average power of many sign tests

Usage

```
average.power.signtest(n, alpha, p)
```

Arguments

```
n sample size (scalar)
alpha p-value threshold (scalar)
```

p Pr(Y>X), as in Noether (JASA 1987)

Value

Average power estimate for multiple testing procedure

See Also

power.signtest for more details about power calculation of sign test. The power calculation is based on asymptotic normal approximation.

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Examples

```
p = rep(c(0.8,0.5),c(100,900));
average.power.signtest(n = 50, alpha = 0.05, p=p)
```

```
average.power.t.test Compute average power of many t-tests
```

Description

Compute average power of many t-tests; Uses classical power formula for t-test; Assumes equal variance and sample size

Usage

```
average.power.t.test(
   n,
   alpha,
   delta,
   sigma = 1,
   type = "two.sample",
   alternative = "two.sided"
)
```

Arguments

```
n per-group sample size (scalar)
alpha p-value threshold (scalar)
delta difference of population means (vector)
sigma standard deviation (vector or scalar, default=1)
type type of t-test: "two.sample", "one.sample"
alternative one- or two-sided test
```

Value

Average power estimate for multiple testing procedure

```
d = rep(c(2,0),c(100,900));
average.power.t.test(n = 20, alpha = 0.05,delta = d)
```

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average.power.tcorr

Compute average power of many t-tests for non-zero correlation

Description

Compute average power of many t-tests for non-zero correlation

Usage

```
average.power.tcorr(n, alpha, rho)
```

Arguments

n sample size (scalar)
alpha p-value threshold (scalar)

rho population correlation coefficient (vector)

Details

For many applications, the null effect is rho = 0

Value

Average power estimate for multiple testing procedure

See Also

power.tcorr for more details about power calculation of t-test for non-zero correlation

Examples

```
rho = rep(c(0.3,0),c(100,900));
average.power.tcorr(n = 50, alpha = 0.05, rho = rho)
```

average.power.twoprop Computer average power of many two proportion z-tests

Description

Computer average power of many two proportion z-tests. The power calculation of two proportion z-test is based on asymptotic normal approximation.

```
average.power.twoprop(n, alpha, p1, p2, alternative)
```

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Arguments

n	per-group sample size (scalar)
alpha	p-value threshold (scalar)
p1	probability in one group (vector)
p2	probability in other group (vector)
alternative	one- or two-sided test

Value

Average power estimate for multiple testing procedure

Examples

```
set.seed(1234);

p1 = sample(seq(0,0.5,0.1),40,replace = TRUE);

p2 = sample(seq(0.5,1,0.1),40,replace = TRUE);

p2 = sample(seq(0.5,1,0.1),40,replace = TRUE);

p3 = sample(seq(0.5,1,0.1),40,replace = TRUE);

p4 = sample(seq(0.5,1,0.1),40,replace = TRUE);
```

fdr.avepow Compute FDR and average power for a given sample size and effect size vector

Description

For a given fixed sample size and effect size vector, compute FDR and average power as a function of the p-value threshold alpha.

Usage

```
fdr.avepow(n, avepow.func, null.hypo, alpha = 1:100/1000, method = "BH", ...)
```

Arguments

n	sample size
avepow.func	function to compute average power
null.hypo	string to evaluate null hypothesis
alpha	p-value threshold(s) to consider
method	method to estimate proportion pi0 of tests with a true null hypothesis, including: "HH" (p-value histogram height), "HM" (p-value histogram mean), "BH" (Benjamini & Hochberg 1995), "Jung" (Jung 2005)
	additional arguments, including effect size vector for average power function

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Value

A list with the following components:

n input sample size

avepow. func average power function null.hypo null hypothesis string pi0 computed value of pi0

method method to estimate proportion pi0 of tests with true null, including: "HH" (p-

value histogram height), "HM" (p-value histogram mean), "BH" (Benjamini &

Hochberg 1995), "Jung" (Jung 2005)

other.args additional arguments

res.tbl table of alpha, fdr, and average power

References

Pounds S and Cheng C, "Sample size determination for the false discovery rate." Bioinformatics 21.23 (2005): 4263-4271.

Gadbury GL, et al. (2004) Power and sample size estimation in high dimensional biology. Statistical Methods in Medical Research 13(4):325-38.

Jung, Sin-Ho. "Sample size for FDR-control in microarray data analysis." Bioinformatics 21.14 (2005): 3097-3104.

Ni Y, Seffernick A, Onar-Thomas A, Pounds S. "Computing Power and Sample Size for the False Discovery Rate in Multiple Applications", Manuscript.

Examples

```
n = 50; # number of events logHR = rep(c(0,0.5),c(950,50)); v = rep(1,length(logHR)); # variance of predictor variable (vector) res = fdr.avepow(n,average.power.coxph,"logHR==0",logHR=logHR,v=v); res$pi0; head(res$res.tbl)
```

fdr.power.alpha

Compute FDR for given p-value threshold, average power and proportion of tests with a true null

Description

Compute the FDR for given values of the p-value threshold alpha, average power, and proportion pi0 of tests with a true null hypothesis.

```
fdr.power.alpha(alpha, pwr, pi0, method = "HH")
```

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Arguments

alpha p-value threshold (vector)

pwr average power

pi0 actual proportion of tests with a true null hypothesis

method method to estimate proportion pi0 of tests with true null, including: "HH" (p-

value histogram height), "HM" (p-value histogram mean), "BH" (Benjamini &

Hochberg 1995), "Jung" (Jung 2005)

Value

FDR

References

Pounds S and Cheng C, "Sample size determination for the false discovery rate." Bioinformatics 21.23 (2005): 4263-4271.

Gadbury GL, et al. (2004) Power and sample size estimation in high dimensional biology. Statistical Methods in Medical Research 13(4):325-38.

Jung, Sin-Ho. "Sample size for FDR-control in microarray data analysis." Bioinformatics 21.14 (2005): 3097-3104.

Ni Y, Seffernick A, Onar-Thomas A, Pounds S. "Computing Power and Sample Size for the False Discovery Rate in Multiple Applications", Manuscript.

Examples

```
alpha = 1:100/1000;
pwr = rep(0.8,length(alpha));
pi0 = 0.95;
fdr.power.alpha(alpha,pwr,pi0,method="HH")
```

find.sample.size

Determines the sample size needed to achieve the desired FDR and average power

Description

Determines the sample size needed to achieve the desired FDR and average power by given the proportion of true null hypothesis.

```
find.sample.size(alpha, pwr, avepow.func, n0 = 3, n1 = 6, max.its = 50, ...)
```

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Arguments

alpha the fixed p-value threshold (scalar numeric)

pwr desired average power (scalar numeric)

avepow.func an R function to compute average power

no lower limit for initial sample size range

numeric)

upper limit for initial sample size range

max.its maximum number of iterations

additional arguments to average power function

Value

A list with the following components:

lower limit for initial sample size range

upper limit for initial sample size range

Note

n0

n1

For the test with power calculation based on asymptotic normal approximation, we suggest checking FDRsamplesize2 calculation by simulation.

```
#Here, calculating the sample size for the study involving many sign tests average.power.signtest; p.adj = 0.001; p = rep(c(0.8,0.5), c(100,9900)); find.sample.size(alpha = p.adj, pwr = 0.8, avepow.func = average.power.signtest, p = p)
```

n.fdr.coxph

n.fdr.coxph Sample size calculation for the Cox proportional hazards regression model	n.fdr.coxph		
---------------------------------------------------------------------------------------	-------------	--	--

Description

Find number of events needed to have a desired false discovery rate and average power for a large number of Cox regression models with non-binary covariates.

Usage

```
n.fdr.coxph(fdr, pwr, logHR, v, pi0.hat = "BH")
```

Arguments

fdr desired FDR (scalar numeric)

pwr desired average power (scalar numeric)

logHR log hazard ratio (vector)

v variance of predictor variable (vector)

pi0.hat method to estimate proportion pi0 of tests with true null, including: "HH" (p-

value histogram height), "HM" (p-value histogram mean), "BH" (Benjamini &

Hochberg 1995), "Jung" (Jung 2005)

Value

A list with the following components:

n number of events estimate

computed.avepow

average power

desired.avepow desired average power

desired.fdr desired FDR

input.pi0 proportion of tests with a true null hypothesis

alpha fixed p-value threshold for multiple testing procedure

n.its number of iteration

max.its maximum number of iteration, default is 50 no lower limit for initial sample size range n1 upper limit for initial sample size range

Note

For the test with power calculation based on asymptotic normal approximation, we suggest checking FDRsamplesize2 calculation by simulation.

n.fdr.fisher

References

Hsieh, FY and Lavori, Philip W (2000) Sample-size calculations for the Cox proportional hazards regression model with non-binary covariates. Controlled Clinical Trials 21(6):552-560.

Examples

```
\begin{split} &\log. \text{HR=log(rep(c(1,2),c(900,100)))} \\ &\textit{v=rep(1,1000)} \\ &\textit{n.fdr.coxph(fdr=0.1, pwr=0.8,logHR=log.HR, v=v, pi0.hat="BH")} \end{split}
```

n.fdr.fisher

Sample size calculation for Fisher's Exact tests

Description

Find the sample size needed to have a desired false discovery rate and average power for a large number of Fisher's exact tests.

Usage

```
n.fdr.fisher(fdr, pwr, p1, p2, alternative = "two.sided", pi0.hat = "BH")
```

Arguments

fdr desired FDR (scalar numeric)

pwr desired average power (scalar numeric)

p1 probability in one group (vector) p2 probability in other group (vector)

alternative one- or two-sided test

pi0.hat method to estimate proportion pi0 of tests with true null, including: "HH" (p-

value histogram height), "HM" (p-value histogram mean), "BH" (Benjamini &

Hochberg 1995), "Jung" (Jung 2005)

Value

A list with the following components:

n per-group sample size estimate

computed.avepow

average power

desired.avepow desired average power

desired.fdr desired FDR

input.pi0 proportion of tests with a true null hypothesis

alpha fixed p-value threshold for multiple testing procedure

n.fdr.negbin

n.its	number of iteration
max.its	maximum number of iteration, default is 50
n0	lower limit for initial sample size range
n1	upper limit for initial sample size range

Examples

```
set.seed(1234);
p1 = sample(seq(0,0.5,0.1),10,replace = TRUE);
p2 = sample(seq(0.5,1,0.1),10,replace = TRUE);
n.fdr.fisher(fdr = 0.1, pwr = 0.8, p1 = p1, p2 = p2, alternative = "two.sided", pi0.hat = "BH")
```

n.fdr.negbin

Sample size calculation for Negative Binomial data

Description

Find the sample size needed to have a desired false discovery rate and average power for a large number of Negative Binomial comparisons.

Usage

```
n.fdr.negbin(fdr, pwr, log.fc, mu, sig, pi0.hat = "BH")
```

Arguments

fdr	desired FDR (scalar numeric)
pwr	desired average power (scalar numeric)
log.fc	log fold-change (vector), usual null hypothesis is log.fc=0
mu	read depth per gene (vector, same length as log.fc)
sig	coefficient of variation (CV) per gene (vector, same length as log.fc)
pi0.hat	method to estimate proportion pi0 of tests with true null, including: "HH" (p-value histogram height), "HM" (p-value histogram mean), "BH" (Benjamini & Hochberg 1995), "Jung" (Jung 2005)

Value

A list with the following components:

```
n per-group sample size estimate
computed.avepow
average power

desired.avepow desired average power

desired.fdr desired FDR
input.pi0 proportion of tests with a true null hypothesis
```

n.fdr.oneway

alpha	fixed p-value threshold for multiple testing procedure
n.its	number of iteration
max.its	maximum number of iteration, default is 50
n0	lower limit for initial sample size range
n1	upper limit for initial sample size range

Note

For the test with power calculation based on asymptotic normal approximation, we suggest checking FDRsamplesize2 calculation by simulation.

References

SN Hart, TM Therneau, Y Zhang, GA Poland, and J-P Kocher (2013). Calculating Sample Size Estimates for RNA Sequencing Data. Journal of Computational Biology 20: 970-978.

Examples

```
\begin{split} \log & \text{FC = log(rep(c(1,2),c(900,100)));} \\ \text{mu = rep(5,1000);} \\ \text{sig = rep(0.6,1000);} \\ \text{n.fdr.negbin(fdr = 0.1, pwr = 0.8, log.fc = logFC, mu = mu, sig = sig, pi0.hat = "BH")} \end{split}
```

n.fdr.oneway

Sample size calculation for one-way ANOVA

Description

Find the sample size needed to have a desired false discovery rate and average power for a large number of one-way ANOVA tests.

Usage

```
n.fdr.oneway(fdr, pwr, theta, k, pi0.hat = "BH")
```

Arguments

fdr	desired FDR (scalar numeric)
pwr	desired average power (scalar numeric)
theta	$sum\ of\ ((group\ mean\ -\ overall\ mean)/stdev)^2\ across\ all\ groups\ for\ each\ hypothesis\ test\ (vector)$
k	the number of groups to be compared
pi0.hat	method to estimate proportion pi0 of tests with true null, including: "HH" (p-value histogram height), "HM" (p-value histogram mean), "BH" (Benjamini & Hochberg 1995), "Jung" (Jung 2005)

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Value

A list with the following components:

n per-group sample size estimate

computed.avepow

average power

desired.avepow desired average power

desired.fdr desired FDR

input.pi0 proportion of tests with a true null hypothesis

alpha fixed p-value threshold for multiple testing procedure

n.its number of iteration

max.its maximum number of iteration, default is 50 no lower limit for initial sample size range upper limit for initial sample size range

Examples

```
theta=rep(c(2,0),c(100,900));
n.fdr.oneway(fdr = 0.1, pwr = 0.8, theta = theta, k = 2, pi0.hat = "BH")
```

n.fdr.poisson

Sample size calculation for Poisson data

Description

Find the sample size needed to have a desired false discovery rate and average power for a large number of two-group comparisons under Poisson distribution.

Usage

```
n.fdr.poisson(fdr, pwr, rho, mu0, w, type, pi0.hat = "BH")
```

Arguments

fdr	desired FDR	(scalar numeric)
-----	-------------	------------------

pwr desired average power (scalar numeric)

rho fold-change, usual null hypothesis is that rho=1 (vector)

mu0 average count in control group (vector)

w ratio of the total number of reads mapped between the two groups

type type of test: "w" for Wald, "s" for score, "lw" for log-transformed Wald, "ls" for

log-transformed score.

pi0.hat method to estimate proportion pi0 of tests with true null, including: "HH" (p-

value histogram height), "HM" (p-value histogram mean), "BH" (Benjamini &

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Value

A list with the following components:

n per-group sample size estimate computed.avepow average power

desired.avepow desired average power

desired.fdr desired FDR

input.pi0 proportion of tests with a true null hypothesis

alpha fixed p-value threshold for multiple testing procedure

n.its number of iteration

max.its maximum number of iteration, default is 50 n0 lower limit for initial sample size range n1 upper limit for initial sample size range

References

C-I Li, P-F Su, Y Guo, and Y Shyr (2013). Sample size calculation for differential expression analysis of RNA-seq data under Poisson distribution. Int J Comput Biol Drug Des 6(4).doi:10.1504/IJCBDD.2013.056830

Examples

```
rho = rep(c(1,1.25),c(900,100));
mu0 = rep(5,1000);
w = rep(0.5,1000);
n.fdr.poisson(fdr = 0.1, pwr = 0.8, rho = rho, mu0 = mu0, w = w, type = "w", pi0.hat = "BH")
```

n.fdr.ranksum

Sample size calculation for rank-sum tests

Description

Find the sample size needed to have a desired false discovery rate and average power for a large number of rank-sum tests.

Usage

```
n.fdr.ranksum(fdr, pwr, p, pi0.hat = "BH")
```

Arguments

fdr desired FDR (scalar numeric)

pwr desired average power (scalar numeric)
p Pr(Y>X), as in Noether (JASA 1987)

pi0.hat method to estimate proportion pi0 of tests with true null, including: "HH" (p-

value histogram height), "HM" (p-value histogram mean), "BH" (Benjamini &

n.fdr.signrank

Value

A list with the following components:

n sample size estimate

computed.avepow

average power

desired.avepow desired average power

desired.fdr desired FDR

input.pi0 proportion of tests with a true null hypothesis

alpha fixed p-value threshold for multiple testing procedure

n.its number of iteration

max.its maximum number of iteration, default is 50 no lower limit for initial sample size range n1 upper limit for initial sample size range

References

Noether, Gottfried E (1987) Sample size determination for some common nonparametric tests. Journal of the American Statistical Association, 82:645-647.

Examples

```
p = rep(c(0.8, 0.5), c(100, 900));

n.fdr.ranksum(fdr = 0.1, pwr = 0.8, p = p, pi0.hat = "BH")
```

n.fdr.signrank

Sample size calculation for signed-rank tests

Description

Find the sample size needed to have a desired false discovery rate and average power for a large number of signed-rank tests.

Usage

```
n.fdr.signrank(fdr, pwr, p1, p2, pi0.hat = "BH")
```

Arguments

fdr	desired FDR (scalar numeric)
	` ,

pwr desired average power (scalar numeric)
p1 Pr(X>0), as in Noether (JASA 1987)
p2 Pr(X+X'>0), as in Noether (JASA 1987)

pi0.hat method to estimate proportion pi0 of tests with true null, including: "HH" (p-

value histogram height), "HM" (p-value histogram mean), "BH" (Benjamini &

n.fdr.signtest 23

Value

A list with the following components:

n sample size estimate

computed.avepow

average power

desired.avepow desired average power

desired.fdr desired FDR

input.pi0 proportion of tests with a true null hypothesis

alpha fixed p-value threshold for multiple testing procedure

n.its number of iteration

max.its maximum number of iteration, default is 50 n0 lower limit for initial sample size range n1 upper limit for initial sample size range

References

Noether, Gottfried E (1987) Sample size determination for some common nonparametric tests. Journal of the American Statistical Association, 82:645-647.

Examples

```
p1 = rep(c(0.8,0.5),c(100,900));

p2 = rep(c(0.8,0.5),c(100,900));

n.fdr.signrank(fdr = 0.1, pwr = 0.8, p1 = p1, p2 = p2, pi0.hat = "BH")
```

n.fdr.signtest

Sample size calculation for sign tests

Description

Find the sample size needed to have a desired false discovery rate and average power for a large number of sign tests.

Usage

```
n.fdr.signtest(fdr, pwr, p, pi0.hat = "BH")
```

Arguments

fdr desired FDR (scalar numeric)

pwr desired average power (scalar numeric)
p Pr(X>0), as in Noether (JASA 1987)

pi0.hat method to estimate proportion pi0 of tests with true null, including: "HH" (p-

value histogram height), "HM" (p-value histogram mean), "BH" (Benjamini &

24 n.fdr.tcorr

Value

A list with the following components:

n sample size estimate

computed.avepow

average power

desired.avepow desired average power

desired.fdr desired FDR

input.pi0 proportion of tests with a true null hypothesis

alpha fixed p-value threshold for multiple testing procedure

n.its number of iteration

max.its maximum number of iteration, default is 50 no lower limit for initial sample size range n1 upper limit for initial sample size range

Note

For the test with power calculation based on asymptotic normal approximation, we suggest checking FDRsamplesize2 calculation by simulation.

References

Noether, Gottfried E (1987) Sample size determination for some common nonparametric tests. Journal of the American Statistical Association, 82:645-647.

Examples

```
p = rep(c(0.8, 0.5), c(100, 900));
n.fdr.signtest(fdr = 0.1, pwr = 0.8, p = p, pi0.hat = "BH")
```

n.fdr.tcorr

Sample size calculation for t-tests for non-zero correlation

Description

Find the sample size needed to have a desired false discovery rate and average power for a large number of t-tests for non-zero correlation.

```
n.fdr.tcorr(fdr, pwr, rho, pi0.hat = "BH")
```

n.fdr.ttest 25

Arguments

fdr desired FDR (scalar numeric)

pwr desired average power (scalar numeric)

rho population correlation coefficient (vector)

pi0.hat method to estimate proportion pi0 of tests with true null, including: "HH" (p-

value histogram height), "HM" (p-value histogram mean), "BH" (Benjamini &

Hochberg 1995), "Jung" (Jung 2005)

Value

A list with the following components:

n sample size estimate

computed.avepow

average power

desired.avepow desired average power

desired.fdr desired FDR

input.pi0 proportion of tests with a true null hypothesis

alpha fixed p-value threshold for multiple testing procedure

n.its number of iteration

max.its maximum number of iteration, default is 50 no lower limit for initial sample size range

n1 upper limit for initial sample size range

Examples

```
rho = rep(c(0.3,0),c(100,900));
n.fdr.tcorr(fdr = 0.1, pwr = 0.8, rho = rho, pi0.hat="BH")
```

n.fdr.ttest

Sample size calculation for t-tests

Description

Find the sample size needed to have a desired false discovery rate and average power for a large number of t-tests.

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Usage

```
n.fdr.ttest(
  fdr,
  pwr,
  delta,
  sigma = 1,
  type = "two.sample",
  pi0.hat = "BH",
  alternative = "two.sided"
)
```

Arguments

fdr	desired FDR (scalar numeric)
pwr	desired average power (scalar numeric)
delta	difference of population means (vector)
sigma	standard deviation (vector or scalar)
type	type of t-test
pi0.hat	method to estimate proportion pi0 of tests with true null, including: "HH" (p-value histogram height), "HM" (p-value histogram mean), "BH" (Benjamini & Hochberg 1995), "Jung" (Jung 2005)
alternative	one- or two-sided test

Value

A list with the following components:

```
sample size (per group) estimate
computed.avepow
                  average power
desired.avepow desired average power
desired.fdr
                  desired FDR
input.pi0
                  proportion of tests with a true null hypothesis
alpha
                  fixed p-value threshold for multiple testing procedure
n.its
                  number of iteration
                  maximum number of iteration, default is 50
max.its
                  lower limit for initial sample size range
n0
                  upper limit for initial sample size range
n1
```

```
d = rep(c(2,0),c(100,900));

n.fdr.ttest(fdr = 0.1, pwr = 0.8, delta = d)
```

n.fdr.twoprop 27

n.fdr.twoprop Sample size calculation for comparing two proportions

Description

Find the sample size needed to have a desired false discovery rate and average power for a large number of two-group comparisons using the two proportion z-test.

Usage

```
n.fdr.twoprop(fdr, pwr, p1, p2, alternative = "two.sided", pi0.hat = "BH")
```

Arguments

fdr desired FDR (scalar numeric)

pwr desired average power (scalar numeric)

p1 probability in one group (vector)

p2 probability in other group (vector)

alternative one- or two-sided test

pi0.hat method to estimate proportion pi0 of tests with true null, including: "HH" (p-value histogram height), "HM" (p-value histogram mean), "BH" (Benjamini & Hochberg 1995), "Jung" (Jung 2005)

Value

A list with the following components:

per-group sample size estimate computed.avepow average power desired.avepow desired average power desired.fdr desired FDR input.pi0 proportion of tests with a true null hypothesis alpha fixed p-value threshold for multiple testing procedure number of iteration n.its maximum number of iteration, default is 50 max.its lower limit for initial sample size range n0 n1 upper limit for initial sample size range

Note

For the test with power calculation based on asymptotic normal approximation, we suggest checking FDRsamplesize2 calculation by simulation.

28 power.cox

Examples

```
set.seed(1234);
p1 = sample(seq(0,0.5,0.1),40,replace = TRUE);
p2 = sample(seq(0.5,1,0.1),40,replace = TRUE);
n.fdr.twoprop(fdr = 0.1, pwr = 0.8, p1 = p1, p2 = p2, alternative = "two.sided", pi0.hat = "BH")
```

power.cox

Compute the power of a single-predictor Cox regression model

Description

Use the formula of Hsieh and Lavori (2000) to compute the power of a single-predictor Cox model, which is based on asymptotic normal approximation.

Usage

```
power.cox(n, alpha, logHR, v)
```

Arguments

```
n number of events (scalar)
alpha p-value threshold (scalar)
logHR log hazard ratio (vector)
v variance of predictor variable (vector)
```

Value

Vector of power estimates for two-sided test

References

Hsieh, FY and Lavori, Philip W (2000) Sample-size calculations for the Cox proportional hazards regression model with non-binary covariates. Controlled Clinical Trials 21(6):552-560.

```
logHR = log(rep(c(1, 2),c(900, 100)));

v = rep(1, 1000);

res = power.cox(n = 50,alpha = 0.05,logHR = logHR, v = v)
```

power.fisher 29

|--|

Description

Compute power for Fisher's exact test

Usage

```
power.fisher(p1, p2, n, alpha, alternative)
```

Arguments

p1	probability in one group (scalar)
p2	probability in other group (scalar)
n	per-group sample size (scalar)
alpha	p-value threshold (scalar)
alternative	one- or two-sided test, must be one of "greater", "less", or "two.sided"

Value

Power estimate for one- or two-sided tests

Examples

```
power.fisher(p1 = 0.5, p2 = 0.9, n=20, alpha = 0.05, alternative = 'two.sided')
```

power.hart Compute power for RNA-seq experiments assuming Negative distribution

Description

Use the formula of Hart et al (2013) to compute power for comparing RNA-seq expression across two groups assuming a Negative Binomial distribution. The power calculation is based on asymptotic normal approximation.

```
power.hart(n, alpha, log.fc, mu, sig)
```

30 power.li

Arguments

n	per-group sample size (scalar)
alpha	p-value threshold (scalar)
log.fc	log fold-change (vector), usual null hypothesis is log.fc=0
mu	read depth per gene (vector, same length as log.fc)
sig	coefficient of variation (CV) per gene (vector, same length as log.fc)

Details

This function is based on equation (1) of Hart et al (2013). It assumes a Negative Binomial model for RNA-seq read counts and equal sample size per group.

Value

Vector of power estimates for the set of two-sided tests

tion

References

SN Hart, TM Therneau, Y Zhang, GA Poland, and J-P Kocher (2013). Calculating Sample Size Estimates for RNA Sequencing Data. Journal of Computational Biology 20: 970-978.

Examples

```
n.hart = 2*(qnorm(0.975)+qnorm(0.9))^2*(1/20+0.6^2)/(log(2)^2) # Equation (6) of Hart et al power.hart(n.hart,0.05,log(2),20,0.6) # Recapitulate 90% power

power.li Compute power for RNA-Seq experiments assuming Poisson distribu-
```

Description

Use the formula of Li et al (2013) to compute power for comparing RNA-seq expression across two groups assuming the Poisson distribution

Usage

```
power.li(n, alpha, rho, mu0, w, type)
```

Arguments

n	per-group sample size
alpha	p-value threshold (scalar)
rho	fold-change, usual null hypothesis is that rho=1 (vector)
mu0	average count in control group
W	ratio of the total number of reads mapped between the two groups
type	type of test: "w" for Wald, "s" for score, "lw" for log-transformed Wald, "ls" for log-transformed score

power.oneway 31

Details

This function computes the power for each of a series of two-sided tests defined by the input parameters. The power is based on the sample size formulas in equations (10-13) of Li et al (2013). Also, note that the null effect is set to 1 in the examples because the usual null hypothesis is that the fold-change = 1.

Value

Vector of power estimates for two-sided tests

References

C-I Li, P-F Su, Y Guo, and Y Shyr (2013). Sample size calculation for differential expression analysis of RNA-seq data under Poisson distribution. Int J Comput Biol Drug Des 6(4). <doi:10.1504/IJCBDD.2013.056830>

Examples

```
power.li(n = 88, alpha = 0.05, rho = 1.25, mu0 = 5, w = 0.5, type = "w") # recapitulate 80% power in Table 1 of Li et al (2013)
```

power.oneway

Compute power of one-way ANOVA

Description

Compute power of one-way ANOVA; Uses classical power formula for ANOVA; Assumes equal variance and sample size

Usage

```
power.oneway(n, alpha, theta, k = 2)
```

Arguments

n per-group sample size (scalar) alpha p-value threshold (scalar)

theta sum of ((group mean - overall mean)/stdev)^2 across all groups for each hypoth-

esis test(vector)

k the number of groups to be compared, default k=2

Details

For many applications, the null effect is zero for the parameter theta described above

Value

Vector of power estimates for test of equal means

32 power.ranksum

Examples

```
theta=rep(c(2,0),c(100,900));
res = power.oneway(n = 50, alpha = 0.05, theta = theta, k = 2)
```

power.ranksum

Compute power of the rank-sum test

Description

Compute power of rank-sum test; Uses formula of Noether (JASA 1987), which is based on asymptotic normal approximation.

Usage

```
power.ranksum(n, alpha, p)
```

Arguments

n sample size (scalar)
alpha p-value threshold (scalar)

p Pr(Y>X), as in Noether (JASA 1987)

Details

In most applications, the null effect size will be designated by p = 0.5

Value

Vector of power estimates for two-sided tests

References

Noether, Gottfried E (1987) Sample size determination for some common nonparametric tests. Journal of the American Statistical Association, 82:645-647.

```
p = rep(c(0.8, 0.5), c(100, 900))
res = power.ranksum(n = 50, alpha = 0.5, p=p)
```

power.signrank 33

power.signrank	Compute power of the signed-rank test

Description

Use the Noether (1987) formula to compute the power of the signed-rank test, which is based on asymptotic normal approximation.

Usage

```
power.signrank(n, alpha, p1, p2)
```

Arguments

n	sample size (scalar)
alpha	p-value threshold (scalar)
p1	Pr(X>0), as in Noether (JASA 1987)
p2	Pr(X+X'>0), as in Noether (JASA 1987)

Details

In most applications, the null effect size will be designated by p1 = p2 = 0.5

Value

Vector of power estimates for two-sided tests

References

Noether, Gottfried E (1987) Sample size determination for some common nonparametric tests. Journal of the American Statistical Association, 82:645-647.

```
\begin{array}{lll} p1 &=& rep(c(0.8,0.5),c(100,900));\\ p2 &=& rep(c(0.8,0.5),c(100,900));\\ res &=& power.signrank(n = 50, alpha = 0.05, p1 = p1, p2 = p2) \end{array}
```

power.signtest

power.signtest

Compute power of the sign test

Description

Use the Noether (1987) formula to compute the power of the sign test, which is based on asymptotic normal approximation.

Usage

```
power.signtest(n, alpha, p)
```

Arguments

n sample size (scalar)

alpha p-value threshold (scalar)

p Pr(X>0), as in Noether (JASA 1987)

Details

In most applications, the null effect size will be designated by p = 0.5

Value

Vector of power estimates for two-sided tests

References

Noether, Gottfried E (1987) Sample size determination for some common nonparametric tests. Journal of the American Statistical Association, 82:645-647.

```
p = rep(c(0.8,0.5),c(100,900));

res = power.signtest(n = 50, alpha = 0.05, p = p)
```

power.tcorr 35

power.tcorr

Compute power of the t-test for non-zero correlation

Description

Compute power of the t-test for non-zero correlation

Usage

```
power.tcorr(n, alpha, rho)
```

Arguments

n sample size (scalar) alpha p-value threshold (scalar)

rho population correlation coefficient (vector)

Details

For many applications, the null effect is rho = 0

Value

Vector of power estimates for two-sided tests

```
rho = rep(c(0.3,0),c(100,900));
res = power.tcorr(n = 50, alpha = 0.05, rho = rho)
```

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