

Package ‘bacon’

April 10, 2023

Type Package

Title Controlling bias and inflation in association studies using the empirical null distribution

Version 1.26.0

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Description Bacon can be used to remove inflation and bias often observed in epigenome- and transcriptome-wide association studies. To this end bacon constructs an empirical null distribution using a Gibbs Sampling algorithm by fitting a three-component normal mixture on z-scores.

License GPL (>= 2)

Depends R (>= 3.3), methods, stats, ggplot2, graphics, BiocParallel, ellipse

Suggests BiocStyle, knitr, rmarkdown, testthat, roxygen2

biocViews ImmunoOncology, StatisticalMethod, Bayesian, Regression, GenomeWideAssociation, Transcriptomics, RNASeq, MethylationArray, BatchEffect, MultipleComparison

RoxygenNote 6.1.1

Collate 'BaconClass.R' 'BaconMethods.R' 'bacon.R' 'normmmixture.R'

VignetteBuilder knitr

NeedsCompilation yes

git_url <https://git.bioconductor.org/packages/bacon>

git_branch RELEASE_3_16

git_last_commit 7eec762

git_last_commit_date 2022-11-01

Date/Publication 2023-04-10

R topics documented:

| | |
|-------------------|----|
| bacon | 2 |
| Bacon-class | 4 |
| bias | 4 |
| dnormmmix | 5 |
| es | 6 |
| estimates | 6 |
| fit | 7 |
| inflation | 8 |
| meta | 9 |
| plot,Bacon-method | 10 |
| plotnormmmix | 10 |
| posteriors | 11 |
| pval | 12 |
| rnormmmix | 13 |
| se | 13 |
| topTable | 14 |
| traces | 15 |
| tstat | 16 |

Index

17

| | |
|-------|----------------------|
| bacon | <i>Gibbs sampler</i> |
|-------|----------------------|

Description

Gibbs Sampler Algorithm to fit a three component normal mixture to z-scores

Usage

```
bacon(teststatistics = NULL, effectsizes = NULL,
      standarderrors = NULL, niter = 5000L, nburnin = 2000L,
      nbins = 1000, trim = 0.999, level = 0.05, na.exclude = FALSE,
      verbose = FALSE, priors = list(sigma = list(alpha = 1.28, beta =
      0.36), mu = list(lambda = c(0, 3, -3), tau = c(1000, 100, 100)), epsilon
      = list(gamma = c(90, 5, 5))))
```

Arguments

| | |
|----------------|---|
| teststatistics | numeric vector or matrix of test-statistics |
| effectsizes | numeric vector or matrix of effect-sizes |
| standarderrors | numeric vector or matrix of standard errors |
| niter | number of iterations |
| nburnin | length of the burnin period |
| nbins | default 1000 else bin test-statistics |

| | |
|------------|--|
| trim | default 0.999 trimming test-statistics |
| level | significance leve used to determine prop. null for starting values |
| na.exclude | see ?na.exclude |
| verbose | default FALSE |
| priors | list of parameters of for the prior distributions |

Value

object of class-Bacon

Author(s)

mvaniterson

References

Implementation is based on a version from Zhihui Liu <https://macsphere.mcmaster.ca/handle/11375/9368>

Examples

```
##simulate some test-statistic from a normal mixture
##and run bacon
y <- rnormmmix(2000, c(0.9, 0, 1, 0, 4, 1))
bc <- bacon(y)
##extract all estimated mixture parameters
estimates(bc)
##extract inflation
inflation(bc)
##extract bias
bias(bc)

##extract bias and inflation corrected test-statistics
head(tstat(bc))

##inspect the Gibbs Sampling output
traces(bc)
posteriors(bc)
fit(bc)

##simulate multiple sets of test-statistic from a normal mixture
##and run bacon
y <- matrix(rnormmmix(10*2000, c(0.9, 0, 1, 0, 4, 1)), ncol=10)
bc <- bacon(y)
##extract all estimated mixture parameters
estimates(bc)
##extract only the inflation
inflation(bc)
##extract only the bias
bias(bc)
##extract bias and inflation corrected P-values
```

```
head(pval(bc))
##extract bias and inflation corrected test-statistics
head(tstat(bc))
```

Bacon-class*An S4 class container for storing Gibbs Sampler input and output***Description**

An S4 class container for storing Gibbs Sampler input and output

Slots

- `teststatistics` numeric vector or matrix of test-statistics
- `effectsizes` numeric vector or matrix of effect-sizes
- `standarderrors` numeric vector or matrix of standard errors
- `traces` array of Gibbs Sampler traces
- `estimates` vector or matrix of parameter estimates
- `priors` list of parameters of for the prior distributions
- `niter` number of iterations
- `nburnin` length of the burnin period

bias*Method to extract the estimated bias from the 'bacon'-object***Description**

Method to extract the estimated bias from the 'bacon'-object

Usage

```
bias(object)

## S4 method for signature 'Bacon'
bias(object)
```

Arguments

| | |
|---------------------|----------------|
| <code>object</code> | 'bacon'-object |
|---------------------|----------------|

Value

vector or matrix of inflation

See Also[bacon](#)**Examples**

```
y <- rnormmix(2000, c(0.9, 0, 1, 0, 4, 1))
##nbins = 100 to speed up the calculations
bc <- bacon(y, nbins=100)
bias(bc)
```

dnormmix*density of a k-component normal mixture*

Description

density of a k-component normal mixture

Usage

```
dnormmix(x, theta)
```

Arguments

| | |
|-------|---|
| x | x like dnorm(x, ...) |
| theta | parameters of the mixture proportion, mean and sd |

Details

details follow

Value

density of a k-component normal mixture

Author(s)

mvaniterson

Examples

```
n <- 2000
theta <- c(0.8, 0, 1, 0, 4, 1)
x <- rnormmix(n, theta)
hist(x, freq=FALSE, n=100)
curve(dnormmix(x, theta), add=TRUE, lwd=2)
```

es*Method to extract inflation- and bias-corrected effect-sizes***Description**

Method to extract inflation- and bias-corrected effect-sizes

Usage

```
es(object, corrected = TRUE)

## S4 method for signature 'Bacon'
es(object, corrected = TRUE)
```

Arguments

| | |
|-----------|-----------------------------|
| object | 'bacon'-object |
| corrected | optional return uncorrected |

Value

vector or matrix of effect-sizes

See Also

[bacon](#)

Examples

```
es <- replicate(6, rnormmmix(2000, c(0.9, 0, 1, 0, 4, 1)))
se <- replicate(6, 0.8*sqrt(4/rchisq(2000,df=4)))
bc <- bacon(NULL, es, se)
head(es(bc))
```

estimates*Method to extract the estimated parameters from the 'bacon'-object***Description**

Method to extract the estimated parameters from the 'bacon'-object

Usage

```
estimates(object)

## S4 method for signature 'Bacon'
estimates(object)
```

Arguments

object 'bacon'-object

Value

vector or matrix of estimates

See Also

[bacon](#)

Examples

```
y <- rnormmmix(2000, c(0.9, 0, 1, 0, 4, 1))
##nbins = 100 to speed up the calculations
bc <- bacon(y, nbins=100)
estimates(bc)
```

fit

Method to plot mixture fit

Description

Method to plot mixture fit

Usage

```
fit(object, index = 1, ...)
## S4 method for signature 'Bacon'
fit(object, index, col = "grey75", border = "grey75",
...)
```

Arguments

| | |
|---------------|---|
| object | 'bacon'-object |
| index | if multiple sets of test-statsistics where provided |
| ... | additional plotting parameters |
| col | line color default 'grey75' |
| border | border color 'grey75' |

Value

plot of the Gibbs Sampler mixture fit

See Also[bacon](#)**Examples**

```
y <- rnormmmix(2000, c(0.9, 0, 1, 0, 4, 1))
##nbins = 100 to speed up the calculations
bc <- bacon(y, nbins=100)
fit(bc)
```

inflation

*Method to extract the estimated inflation from the 'bacon'-object***Description**

Method to extract the estimated inflation from the 'bacon'-object

Usage

```
inflation(object)

## S4 method for signature 'Bacon'
inflation(object)
```

Arguments

| | |
|--------|----------------|
| object | 'bacon'-object |
|--------|----------------|

Value

vector or matrix of inflation

See Also[bacon](#)**Examples**

```
y <- rnormmmix(2000, c(0.9, 0, 1, 0, 4, 1))
##nbins = 100 to speed up the calculations
bc <- bacon(y, nbins=100)
inflation(bc)
```

| | |
|------|----------------------------|
| meta | <i>fixed meta-analysis</i> |
|------|----------------------------|

Description

Perform fixed meta-analysis using inflation and bias corrected effect-sizes and standard errors

Usage

```
meta(object, corrected = TRUE, ...)

## S4 method for signature 'Bacon'
meta(object, corrected = TRUE, ...)
```

Arguments

| | |
|-----------|-----------------------------|
| object | 'bacon'-object |
| corrected | optional return uncorrected |
| ... | additional arguments |

Details

TODO maybe add idea's from http://www.netstorm.be/home/meta_analysis#metaAnalysisU

Value

object of class 'bacon' with added fixed-effect meta-analysis test-statistics, effect-sizes and standard-errors

See Also

[bacon](#)

Examples

```
es <- replicate(6, rnormmmix(2000, c(0.9, 0, 1, 0, 4, 1)))
se <- replicate(6, 0.8*sqrt(4/rchisq(2000,df=4)))
bc <- bacon(NULL, es, se)
mbc <- meta(bc)
```

`plot`,`Bacon-method` *plot hist or qq*

Description

simple ggplot2 plotting function for 'bacon'-object

Usage

```
## S4 method for signature 'Bacon'
plot(x, y, type = c("hist", "qq"))
```

Arguments

| | |
|-------------------|----------------|
| <code>x</code> | 'bacon'-object |
| <code>y</code> | NULL |
| <code>type</code> | hist or qq |

Value

either qq-plot of P-values or histogram of Test-statistics

`plotnormmix` *plot normal mixtures*

Description

plot normal mixtures

Usage

```
plotnormmix(x, theta, ...)
```

Arguments

| | |
|--------------------|--|
| <code>x</code> | vector of test statistics |
| <code>theta</code> | parameters describing the mixture components |
| <code>...</code> | arguments passed to hist |

Details

details follow

Value

return plot with histogram of the data and mixture and individual components

Author(s)

mvaniterson

Examples

```
n <- 2000
theta <- c(0.8, 0, 1, 0, 4, 1)
x <- rnormmmix(n, theta)
plotnormmmix(x, theta)
```

posteriors*Method to plot posterior distribution*

Description

Method to plot posterior distribution

Usage

```
posteriors(object, thetas = c("sigma.0", "p.0"), index = 1,
           alphas = c(0.95, 0.9, 0.75), xlab = "", ylab = "", ...)
## S4 method for signature 'Bacon'
posteriors(object, thetas = c("sigma.0", "p.0"),
            index = 1, alphas = c(0.95, 0.9, 0.75), xlab = "", ylab = "", ...)
```

Arguments

| | |
|--------|---|
| object | 'bacon'-object |
| thetas | which thetas to plot |
| index | if multiple sets of test-statsistics where provided |
| alphas | significance level confidence ellipses |
| xlab | optional xlab |
| ylab | optional ylab |
| ... | additional plotting parameters |

Value

plot of the Gibbs Sampler posterior probabilities

See Also

[bacon](#)

Examples

```
y <- rnormmmix(2000, c(0.9, 0, 1, 0, 4, 1))
##nbins = 100 to speed up the calculations
bc <- bacon(y, nbins=100)
posterior(bc)
```

pval

Method to extract inflation- and bias-corrected P-values

Description

Method to extract inflation- and bias-corrected P-values

Usage

```
pval(object, corrected = TRUE)

## S4 method for signature 'Bacon'
pval(object, corrected = TRUE)
```

Arguments

| | |
|-----------|-----------------------------|
| object | 'bacon'-object |
| corrected | optional return uncorrected |

Value

vector or matrix of P-values

See Also

[bacon](#)

Examples

```
y <- rnormmmix(2000, c(0.9, 0, 1, 0, 4, 1))
bc <- bacon(y, nbins=100) #nbins = 100 to speed up the calculations
head(pval(bc))
```

| | |
|----------|-------------------------------------|
| rnormmix | <i>sample from a normal mixture</i> |
|----------|-------------------------------------|

Description

sample from a normal mixture

Usage

```
rnormmix(n, theta, shuffle = TRUE)
```

Arguments

| | |
|---------|---|
| n | size |
| theta | parameters |
| shuffle | shuffle return vectors or keep nulls and alternative ordered (null, alts) |

Details

details follow

Value

n samples from a normal mixture with parameters theta

Author(s)

mvaniterson

Examples

```
n <- 2000
theta <- c(0.8, 0, 1, 0, 4, 1)
x <- rnormmix(n, theta)
```

| | |
|----|--|
| se | <i>Method to extract inflation- and bias-corrected standard errors</i> |
|----|--|

Description

Method to extract inflation- and bias-corrected standard errors

Usage

```
se(object, corrected = TRUE)

## S4 method for signature 'Bacon'
se(object, corrected = TRUE)
```

Arguments

| | |
|-----------|-----------------------------|
| object | 'bacon'-object |
| corrected | optional return uncorrected |

Value

vector or matrix of standard-errors

See Also

[bacon](#)

Examples

```
es <- replicate(6, rnormmmix(2000, c(0.9, 0, 1, 0, 4, 1)))
se <- replicate(6, 0.8*sqrt(4/rchisq(2000,df=4)))
bc <- bacon(NULL, es, se)
head(se(bc))
```

topTable

topTable

Description

Extract top features after meta analysis

Usage

```
topTable(object, number = 10, adjust.method = "bonf",
         sort.by = c("pval", "eff.size"))

## S4 method for signature 'Bacon'
topTable(object, number = 10, adjust.method = "bonf",
         sort.by = c("pval", "eff.size"))
```

Arguments

| | |
|---------------|---|
| object | 'bacon'-object |
| number | return specified number of top features, n=-1 return all features |
| adjust.method | P-value multiple testing adjustment method default bonferroni |
| sort.by | order results by pval or eff.size |

Value

table with top features

See Also

[bacon](#)

Examples

```
es <- replicate(6, rnormmmix(2000, c(0.9, 0, 1, 0, 4, 1)))
se <- replicate(6, 0.8*sqrt(4/rchisq(2000,df=4)))
bc <- bacon(NULL, es, se)
mbc <- meta(bc)
topTable(mbc)
```

traces

Method to plot Gibbs sampling traces

Description

Method to plot Gibbs sampling traces

Usage

```
traces(object, burnin = TRUE, index = 1)

## S4 method for signature 'Bacon'
traces(object, burnin = TRUE, index = 1)
```

Arguments

| | |
|--------|---|
| object | 'bacon'-object |
| burnin | include burnin period default true |
| index | if multiple sets of test-statsistics where provided |

Value

plot of the Gibbs Sampler traces

See Also

[bacon](#)

Examples

```
y <- rnormmmix(2000, c(0.9, 0, 1, 0, 4, 1))
##nbins = 100 to speed up the calculations
bc <- bacon(y, nbins=100)
traces(bc)
```

tstat*Method to extract inflation- and bias-corrected test-statistics***Description**

Method to extract inflation- and bias-corrected test-statistics

Usage

```
tstat(object, corrected = TRUE)

## S4 method for signature 'Bacon'
tstat(object, corrected = TRUE)
```

Arguments

| | |
|-----------|-----------------------------|
| object | 'bacon'-object |
| corrected | optional return uncorrected |

Value

vector or matrix of test-statistics

See Also

[bacon](#)

Examples

```
y <- rnormmmix(2000, c(0.9, 0, 1, 0, 4, 1))
##nbins = 100 to speed up the calculations
bc <- bacon(y, nbins=100)
head(tstat(bc))
```

Index

bacon, 2, 5–9, 11, 12, 14–16
Bacon-class, 4
bias, 4
bias,Bacon-method (bias), 4

dnormmmix, 5

es, 6
es,Bacon-method (es), 6
estimates, 6
estimates,Bacon-method (estimates), 6

fit, 7
fit,Bacon-method (fit), 7

inflation, 8
inflation,Bacon-method (inflation), 8

meta, 9
meta,Bacon-method (meta), 9

plot,Bacon-method, 10
plotnormmmix, 10
posteriors, 11
posteriors,Bacon-method (posteriors), 11
pval, 12
pval,Bacon-method (pval), 12

rnormmmix, 13

se, 13
se,Bacon-method (se), 13

topTable, 14
topTable,Bacon-method (topTable), 14
traces, 15
traces,Bacon-method (traces), 15
tstat, 16
tstat,Bacon-method (tstat), 16