

Package: INTRIGUE (via r-universe)

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Type Package

Title Quantify and Control Reproducibility in High-Throughput Experiments

Version 0.1.0

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Description Estimate the proportions of the null and the reproducibility and non-reproducibility of the signal group for the input data set. The Bayes factor calculation and EM (Expectation Maximization) algorithm procedures are also included.

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Encoding UTF-8

Imports SQUAREM, dplyr, rlist

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bf.approx	<i>Bayes Factor Approximation</i>
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Description

A function calculates the approximation for bayes factor, when the value of original bayes factor goes to infinity.

Usage

```
bf.approx(z, param, size, k2, oa2)
```

Arguments

z	The index for individual(i).
param	Input dataset.
size	Number of replicates(m).
k2	Grid value of k^2 .
oa2	Grid value of ω^2 .

Value

Approximation for bayes factor in log scale.

bf.cal.cefn	<i>Bayes Factor Calculation Scheme for CENF prior</i>
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Description

A function that calculates bayes factor for each data pair on each grid point in log scale.

Usage

```
bf.cal.cefn(data, hyperparam)
```

Arguments

- `data` A dataset which is constructed by pairs of coefficient values β and standard errors $se(\beta)$.
`hyperparam` A two-dimensional vector denoting all the grid points, namely, $k \times \omega$.

Value

A list records all the log scale bayes factor values.

bf.cal.meta

*Bayes Factor Calculation Scheme for META prior***Description**

A function that calculates bayes factor for each data pair on each grid point in log scale.

Usage

```
bf.cal.meta(data, hyperparam = NULL, bf.only = FALSE)
```

Arguments

- `data` A dataset which is constructed by pairs of coefficient values β and standard errors $se(\beta)$.
`hyperparam` A two-dimensional vector denoting all the grid points, namely, $\phi \times \omega$.
`bf.only` A boolean, denoting whether this function is called to calculate Bayes factor for META prior only. Usually used when publication bias issue is the target.

Value

A list records all the log scale bayes factor values or a list records log scale bayes factor for null, reproducible and irreproducible model (when `bf.only=TRUE`).

bf.em

*Bayes Factor EM Updating Scheme***Description**

A function that describes the updating process in E step and M step for EM algorithm. It will be used in SQUAREM package.

Usage

```
bf.em(w, bf)
```

Arguments

- w The weight vector in previous M step.
 bf A vector recording all the bayes factor values in log scale.

Value

The updated weight vector in current M step(wnew).

bf.loglik

*Bayes Factor Loglikelihood Function***Description**

Calculate the updated loglikelihood value in EM algorithm, and to evaluate whether converge or not.

Usage

```
bf.loglik(w, bf)
```

Arguments

- w The current weight vector
 bf A vector recording all the bayes factor values in log scale.

Value

Negative summation of loglikelihood values.

bf.weighted_sum

*Bayes Factor Weighted Summation***Description**

A function calculates the weighted summation of bayes factor.

Usage

```
bf.weighted_sum(w, bf, i)
```

Arguments

- w Input weight vector.
 bf Input bayes factor vector
 i Individual index.

Value

Weighted sum for bayes factor in log scale.

hetero	<i>Heterogeneity Evaluation</i>
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Description

Evaluating the overall and individually heterogeneity and reproducibility for the given individuals(units) shared in different replicates.

Usage

```
hetero(
  data,
  use_cefn = TRUE,
  rep = NULL,
  irre = NULL,
  phi_min = NULL,
  phi_max = NULL,
  sq_em_tol = 1e-04,
  fdr.level = NULL,
  sample_size = NULL
)
```

Arguments

data	A dataset which is constructed by pairs of coefficient values β and standard errors $se(\beta)$.
use_cefn	A boolean, denoting whether to use CEFN prior. If the value is TRUE, CEFN prior is used, else, META prior is applied. The default value is TRUE.
rep	A vector, denoting all the k^2 (under CEFN prior) or r (under META prior) values constructing the reproducible signals. If not specified, the default one is c(0.105,0.260,0.369), which corresponds to the several prior values satisfy that $Pr(\beta_{i,1}, \beta_{i,2} \text{ have a same sign}) = 0.99, 0.975, 0.95$ for CEFN prior.
irre	A vector, denoting all the k^2 or r values constructing the irreproducible signals. If not specified, the default one is c(2.198, 3.636, 6.735), which corresponds to the several prior values satisfy that $Pr(\beta_{i,1}, \beta_{i,2} \text{ have a same sign}) = 0.75, 0.70, 0.65$ for CEFN prior.
phi_min	A value which determines the maximum phi . If not specified, will be constructed from the input datasets.
phi_max	A value which determines the minimum phi . If not specified, will be constructed from the input datasets.
sq_em_tol	A small, positive scalar that determines when iterations should be terminated in squarem algorithm. The default value is $1e-4$.
fdr.level	The user-defined rejection level for false discovery rate.
sample_size	The user-defined sample size.

Value

A list with the following components:

<code>gridweight</code>	The final optimal weight vector evaluated on each grid point.
<code>ind_prob</code>	A matrix denoting the converged probability for each individual being inside the three different groups, namely, the null, the reproducible and the irreproducible group.
<code>est_prop</code>	The estimated proportion value for the three different groups, namely, the null, the reproducible and the irreproducible group.
<code>lfdr</code>	The local false discovery rate based on the null hypothesis of unit belonging to H_R , reproducible group. $lfdr = 1 - Pr(H_R)$
<code>significant</code>	If <code>fdr.level</code> is specified, a significant object recording True or False will be returned

Examples

```
data("heterodata")

hetero.out<-hetero(heterodata,fdr.level=0.05)
names(hetero.out)
print(hetero.out$est_prop)

## for CRAN check
hetero.out<-hetero(heterodata[1:100,],fdr.level=0.05)
```

hetero.lfdr*Local False Discovery Rate Evaluation***Description**

Local False Discovery Rate Evaluation

Usage

```
hetero.lfdr(cat, fdr.level)
```

Arguments

<code>cat</code>	The final individual-level probabilities of falling into three categories, separately.
<code>fdr.level</code>	Rejection level for Local false discovery rate, if not specified, the rejection decision procedure won't be run.

Value

A list that preserves local false discovery rate and 5tthe corresponding reject decision if called.

*heterodata**Simulation Dataset*

Description

This is a simulation dataset, containing n=5000 units and m=2 replicates. The true proportion for the null, the reproducible and the irreproducible group is 0.80, 0.18, 0.02 separately.

Usage

```
data("heterodata")
```

Format

An object of class `matrix` (inherits from `array`) with 1000 rows and 4 columns.

Examples

```
data("heterodata")
```

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