# Package 'tiltmod'

June 11, 2018

Title Exponential Tilting Method for Reproducible Screening of Large

Type Package

Scale Testing Problem

Version 0.0.1
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<b>Description</b> Develop an exponential tilting method by conditioning on the false discovery rate to tilt a two component mixture of Beta distributions, yielding an tilted mixture model. Use a Boosted EM algorithm to fit a two-component mixture of Beta distributions for p-values or left tail areas of test statistics. The Boosted EM algorithm for the mixture model fitting is built in C++, which is quite fast and stable. The package also includes two utility functions to generate tilted false discovery rates and frequency network.
<b>Depends</b> R (>= $3.3.1$ )
License GPL (>= 2)
LinkingTo Rcpp, BH
<b>Imports</b> Rcpp (>= 0.12.15), stats, foreach, doParallel, plyr, utils, igraph, edgeR, limma
Encoding UTF-8
LazyData true
RoxygenNote 6.0.1
<pre>URL https://github.com/chongma1989/tiltmod</pre>
NeedsCompilation yes
<b>Archs</b> i386, x64
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etilt

The exponential tilting function

# Description

This function tilts the mixture model fitted from the training tail-areas (or p-values) by conditioning on the average of local fdr's from the testing tail-areas (or p-values)

## Usage

```
etilt(xl, xt, f = NULL, h = NULL, m = NULL, interval = NULL, ...)
```

# **Arguments**

xl	The training left-tail areas (or p-values)
xt	The testing left-tail areas (or p-values)
f	The objective function is tilted. If either $xl$ or $f$ is NULL, $f$ is fitted by UBMM. Default is NULL.
h	The conditioning function. If h is NULL, $h = -\log(f(x))$ . Default is NULL.
m	The constant is used to find the optimal theta such that $E(h(x))=m$ . If m is NULL, $m=mean(h(xt))$ . Default is NULL.
interval	The interval is used to search the optimal theta. Default is (-0.5,10).
	Arguments to be passed to uniroot.

## Value

A list includes theta, tau, tilt\_tau, tilt\_f, tilt\_f0, tilt\_f1, respectively.

# Examples

```
xl=c(rbeta(100,0.5,0.5),runif(900))
xt=c(rbeta(300,2,3),runif(700))
## Not run:
etilt(xl,xt)
## End(Not run)
```

fnet

Frequency network

# Description

This function displays the frequency network of discovered differentially expressed genes.

# Usage

```
fnet(x, Simplify = FALSE, threshold = 0.05, max.ew = 2,
   directed = FALSE, ...)
```

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## **Arguments**

X	A list of discovered differentially expressed genes
Simplify	logical indicating whether to discard the genes with lower relative frequency than the threshold. Default is FALSE.
threshold	a numeric value determining the cutoff point, where the genes are discarded with lower relative frequency than it. Default is $0.05$ .
max.ew	a numeric value. The maximum edge width in the network plot.
directed	logical indicating whether the edges are shown in directions. Default is FALSE.
	Arguments to be passed to plot.

## Value

A network plot.

# **Examples**

```
x=list(c(1,3,4),c(2,4,5),c(3,5,1),c(4,1,2),c(5,2,3))
## Not run:
fnet(x)
fnet(x,layout=layout.fruchterman.reingold,vertex.color="grey60")
## End(Not run)
```

MBM

Fit a two-point mixture of Beta distributions

# Description

Fit a two-point mixture of Beta distributions

## Usage

```
MBM(x, w = as.numeric(c()), a0 = as.numeric(c()), a1 = as.numeric(c()), precision = 1e-06, MaxIter = 10000L)
```

# Arguments

X	A vector of numeric values
W	A vector of two numeric values, representing the weights of two Beta distributions. Default values are 0.5, respectively.
a0	Initial values of the alpha and beta for Beta distribution f0. Default values are 1 and 1, respectively.
a1	Initial values of the alpha and beta for Beta distribution f1. Default values are 0.5 and 0.5, respectively.
precision	The tolerance for convergence. Default value is 1e-6.
MaxIter	The maximum iteration for the EM algorithm. Default value is 10000L.

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#### Value

A list of four components, including the converged weight, parameters for Beta distribution f0, parameters for Beta distribution f1, and the convergence iteration, respectively.

## **Examples**

```
x0=rbeta(900,0.8,0.8)
x1=rbeta(100,0.2,0.2)
## Not run:
    MBM(c(x0,x1),w=c(0.8,0.2),a0=c(1,1),a1=c(0.5,0.5))
## End(Not run)
```

tqvalue

The exponential tilting mixture model

# Description

This function tilts the mixture model fitted from the training tail-areas (or p-values) by conditioning on the average of local fdr's from the testing tail-areas (or p-values)

# Usage

```
tqvalue(x1, xt, w = NULL, a = NULL, precision = 1e-08, MaxIter = 10000,
  interval = NULL, adjust = TRUE, method = c("m1", "m2"),
  type = c("left tail area", "pvalue"), alpha = 0.9, q = 0.1,
  ncores = 1, rel.tol = .Machine$double.eps^0.6, tol = 1e-06)
```

#### **Arguments**

xt The testing left-tail areas (or p-values)  w A vector of two numeric values, representing the Beta distributions. See UBMM.  a A vector of two initial parameter values for Beta di precision The precision for convergence. Default value is 1e-MaxIter The maximum iteration for the EM algorithm.	
Beta distributions. See UBMM.  a A vector of two initial parameter values for Beta di precision The precision for convergence. Default value is 1e-	
precision The precision for convergence. Default value is 1e-	weights of the uniform and
-	istribution. See UBMM.
MaxIter The maximum iteration for the EM algorithm	-8.
The maximum iteration for the Entrangomann.	
interval A vector of two numeric values, which determines mal theta. Default is c(-100L,100L).	the range to search the opti-
adjust Whether or not to do the model adjustment. Defaul	lt is TRUE.
method A character chosen from m1, m2. Default is m1.	
type A character value, chosen from left tail area and pv	value. Default is left tail area.
alpha A numeric value. Used in method "m1" to determ Default is 0.9.	ine the probably null region.
q A numeric value. The global false discovery rate us mine the probable null region. Default is 0.1.	sed in method "m2", to deter-
ncores The number of cpus used for implementing this fur	nction.
rel.tol the accuracy used in integrate.	
tol the accuracy used in uniroot.	

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#### Value

A dataframe includes xl, xt, fdr, FDR, tfdr, and tFDR, respectively. fdr and FDR are the local and global false discovery rate for each value of xt. tfdr and tFDR are the corresponding tilted local and global false discovery rate, respectively.

The optimal theta calculated by solving log(E(exp(thetah(x))))-ctheta, where c=mean(h(xt)).

## **Examples**

```
xl=c(rbeta(50,0.2,0.2),runif(950))
xt=c(rbeta(50,0.1,0.1),runif(950))
## Not run:
tqvalue(xl,xt,ncores=4,adjust=FALSE,type="left tail area")
## End(Not run)
```

UBMM

Fit a mixture of uniform and Beta distribution

#### **Description**

Fit a mixture of uniform and Beta distribution

#### Usage

```
UBMM(x, w = as.numeric(c()), a = as.numeric(c()), precision = 1e-08, MaxIter = 10000L)
```

## **Arguments**

x	A vector of numeric values
W	A vector of two numeric values, representing the weights of the uniform and Beta distributions. Default values are 0.5, respectively.
a	Initial values of the alpha and beta for the Beta distribution. Defaults are obtained from MOM estimators.
precision	The tolerance for convergence. Default value is 1e-8.
MaxIter	The maximum iteration for the EM algorithm. Default value is 10000L.

### Value

A list of three components, including the converged weight, parameters for Beta distribution, and the convergence iteration, respectively.

## **Examples**

```
x0=runif(900)
x1=rbeta(100,0.5,0.5)
UBMM(c(x0,x1),w=c(0.8,0.2),a=c(0.7,0.8))
```

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