

# Package ‘LPEadj’

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**Title** A correction of the local pooled error (LPE) method to replace the asymptotic variance adjustment with an unbiased adjustment based on sample size.

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**Depends** LPE

**Imports** LPE, stats

**Description** Two options are added to the LPE algorithm. The original LPE method sets all variances below the max variance in the ordered distribution of variances to the maximum variance. in LPEadj this option is turned off by default. The second option is to use a variance adjustment based on sample size rather than  $\pi/2$ . By default the LPEadj uses the sample size based variance adjustment.

**License** LGPL

**biocViews** Microarray, Proteomics

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## R topics documented:

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| adjBaseOlig.error | <i>Evaluates LPE variance function of M for quantiles of A within and experimental condition and then interpolates it for all genes.</i> |
|-------------------|------------------------------------------------------------------------------------------------------------------------------------------|

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**Description**

Calls adjBaseOlig.error.step1 and adjBaseOlig.error.step2 functions in order to calculate the base-line distribution.

**Usage**

```
adjBaseOlig.error(y, stats=median, q=0.01, min.genes.int=10,div.factor=1,
setMax1=FALSE)
```

**Arguments**

|               |                                                                                                                                                                              |
|---------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| y             | y is a preprocessed matrix or data frame of expression intensities in which columns are expression intensities for a particular experimental condition and rows are genes.   |
| stats         | It determines whether mean or median is to be used for the replicates                                                                                                        |
| q             | q is the quantile width; q=0.01 corresponds to 100 quantiles i.e. percentiles. Bins/quantiles have equal number of genes and are split according to the average intensity A. |
| min.genes.int | Determines the minimum number of genes in a subinterval for selecting the adaptive intervals.                                                                                |
| div.factor    | Determines the factor by which sigma needs to be divided for selecting adaptive intervals.                                                                                   |
| setMax1       | If T then all variances below the max variance in the ordered distribution of variances are set to the maximum variance. If F then variances are left as is (recommended)    |

**Value**

Returns object of class baseOlig comprising a data frame with 2 columns: A and var M, and rows for each quantile specified. The A column contains the median values of A for each quantile/bin and the M columns contains the pooled variance of the replicate chips for genes within each quantile/bin.

**Author(s)**

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## References

J.K. Lee and M.O.Connell(2003). *An S-Plus library for the analysis of differential expression*. In *The Analysis of Gene Expression Data: Methods and Software*. Edited by G. Parmigiani, ES Garrett, RA Irizarry and SL Zegar. Springer, New York.

Jain et. al. (2003) *Local pooled error test for identifying differentially expressed genes with a small number of replicated microarrays*, *Bioinformatics*, 1945-1951.

Jain et. al. (2005) *Rank-invariant resampling based estimation of false discovery rate for analysis of small sample microarray data*, *BMC Bioinformatics*, Vol 6, 187.

## See Also

[lpeAdj](#)

## Examples

```
# Loading the data from the LPE library
data(Ley)

dim(Ley)
# Gives 12488 by 7
Ley[1:3,]
# Returns
#      ID          c1  c2  c3   t1   t2   t3
# 1  AFFX-MurIL2_at 4.06 3.82 4.28 11.47 11.54 11.34
# 2  AFFX-MurIL10_at 4.56 2.79 4.83  4.25  3.72  2.94
# 3  AFFX-MurIL4_at 5.14 4.10 4.59  4.67  4.71  4.67

Ley[,2:7] <- preprocess(Ley[,2:7],data.type="MAS5")

subset <- 1:1000
Ley.subset <- Ley[subset,]

# Finding the baseline distribution of subset of the data
# condition one (3 replicates)
var.1 <- adjBaseOlig.error(Ley.subset[,2:4], q=0.01, setMax1=FALSE)
dim(var.1)
# Returns a matrix of 1000 by 2 (A,M) format, equal to the nrow(data)
```

---

adjBaseOlig.error.step1

*Evaluates LPE variance function of M for quantiles of A within and experimental condition by dividing the A in 100 intervals.*

---

## Description

Genes are placed in bins/quantiles according to their average expression intensity. The function adjBaseOlig.error calculates a pooled variance of M for genes within these bins/quantiles of A for the replicates of the experimental condition contained in y. Here the assumption is that variance of the genes in each interval is similar.

**Usage**

```
adjBaseOlig.error.step1(y, stats=median, setMax=FALSE, q=0.01, df=10)
```

**Arguments**

|        |                                                                                                                                                                              |
|--------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| y      | y is a preprocessed matrix or data frame of expression intensities in which columns are expression intensities for a particular experimental condition and rows are genes.   |
| stats  | It determines whether mean or median is to be used for the replicates                                                                                                        |
| setMax | If T then all variances below the max variance in the ordered distribution of variances are set to the maximum variance. If F then variances are left as is (recommended)    |
| q      | q is the quantile width; q=0.01 corresponds to 100 quantiles i.e. percentiles. Bins/quantiles have equal number of genes and are split according to the average intensity A. |
| df     | df stands for degrees of freedom. It is used in smooth.spline function to interpolate the variances of all genes. Default value is 10.                                       |

**Value**

Returns object of class baseOlig, comprising a data frame with 2 columns: A and var M. The A column contains the median values of each gene and the M columns contains the corresponding variance. Number of rows of the data-frame is same as that of the number of genes.

**Author(s)**

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**References**

J.K. Lee and M.O.Connell(2003). *An S-Plus library for the analysis of differential expression*. In The Analysis of Gene Expression Data: Methods and Software. Edited by G. Parmigiani, ES Garrett, RA Irizarry ad SL Zegar. Springer, NewYork.

Jain et. al. (2003) *Local pooled error test for identifying differentially expressed genes with a small number of replicated microarrays*, Bioinformatics, 1945-1951.

Jain et. al. (2005) *Rank-invariant resampling based estimation of false discovery rate for analysis of small sample microarray data*, BMC Bioinformatics, Vol 6, 187.

**See Also**

[lpeAdj](#)

**Examples**

```
# Loading the data from the LPE library
data(Ley)

dim(Ley)
```

```
# Gives 12488 by 7
Ley[1:3,]
# Returns
#      ID      c1  c2  c3  t1  t2  t3
# 1 AFFX-MurIL2_at 4.06 3.82 4.28 11.47 11.54 11.34
# 2 AFFX-MurIL10_at 4.56 2.79 4.83 4.25 3.72 2.94
# 3 AFFX-MurIL4_at 5.14 4.10 4.59 4.67 4.71 4.67

Ley[1:1000,2:7] <- preprocess(Ley[1:1000,2:7],data.type="MAS5")
# Finding the baseline distribution of subset of the data
# condition one (3 replicates)
var.1 <- adjBaseOlig.error.step1(Ley[1:1000,2:4], setMax=FALSE, q=0.01)
dim(var.1)
# Returns a matrix of 1000 by 2 (A,M) format
```

---

adjBaseOlig.error.step2

*Evaluates LPE variance function of M for quantiles of A within and experimental condition. It is based on the adaptive number of intervals.*

---

## Description

Similar to adjBaseOlig.error.step1 function, except that now the number of bins are chosen adaptively instead of fixed 100.

## Usage

```
adjBaseOlig.error.step2(y, baseOlig.error.step1.res, df=10, stats=median, setMax=FALSE, min.genes.int)
```

## Arguments

|                          |                                                                                                                                                                            |
|--------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| y                        | y is a preprocessed matrix or data frame of expression intensities in which columns are expression intensities for a particular experimental condition and rows are genes. |
| baseOlig.error.step1.res | It is the result obtained from adjBaseOlig.error.step1 function, in which number of bins are fixed=100                                                                     |
| df                       | df stands for degrees of freedom. It is used in smooth.spline function to interpolate the variances of all genes. Default value is 10.                                     |
| stats                    | It determines whether mean or median is to be used for the replicates                                                                                                      |
| setMax                   | If T then all variances below the max variance in the ordered distribution of variances are set to the maximum variance. If F then variances are left as is (recommended)  |
| min.genes.int            | Determines the minimum number of genes in a subinterval for selecting the adaptive intervals.                                                                              |
| div.factor               | Determines the factor by which sigma needs to be divided for selecting adaptive intervals.                                                                                 |

**Value**

Returns object of class baseOlig comprising a data frame with 2 columns: A and var M, and rows for each quantile specified. The A column contains the median values of A for each quantile/bin and the M columns contains the pooled variance of the replicate chips for genes within each quantile/bin.

**Author(s)**

Carl Murie <carl.murie@mcgill.ca>, Nitin Jain <nitin.jain@pfizer.com>

**References**

J.K. Lee and M.O.Connell(2003). *An S-Plus library for the analysis of differential expression*. In *The Analysis of Gene Expression Data: Methods and Software*. Edited by G. Parmigiani, ES Garrett, RA Irizarry and SL Zegar. Springer, New York.

Jain et. al. (2003) *Local pooled error test for identifying differentially expressed genes with a small number of replicated microarrays*, Bioinformatics, 1945-1951.

Jain et. al. (2005) *Rank-invariant resampling based estimation of false discovery rate for analysis of small sample microarray data*, BMC Bioinformatics, Vol 6, 187.

**See Also**

[lpeAdj](#)

**Examples**

```
# Loading the data from the LPE library
data(Ley)

dim(Ley)
# Gives 12488 by 7
Ley[1:3,]
# Returns
#      ID          c1  c2  c3   t1   t2   t3
# 1  AFX-MurIL2_at 4.06 3.82 4.28 11.47 11.54 11.34
# 2  AFX-MurIL10_at 4.56 2.79 4.83  4.25  3.72  2.94
# 3  AFX-MurIL4_at 5.14 4.10 4.59  4.67  4.71  4.67

Ley[1:1000,2:7] <- preprocess(Ley[1:1000,2:7],data.type="MAS5")
# Finding the baseline distribution of subset of the data
# condition one (3 replicates)
var.1 <- adjBaseOlig.error.step1(Ley[1:1000,2:4], q=0.01, df=10)
dim(var.1)
var.11 <- adjBaseOlig.error.step2(Ley[1:1000,2:4], var.1, df=10)
# Returns a matrix of 1000 by 2 (A,M) format
```

---

|                 |                                                                                              |
|-----------------|----------------------------------------------------------------------------------------------|
| calculateLpeAdj | <i>Evaluates local pooled error significance test with user chosen variance adjustments.</i> |
|-----------------|----------------------------------------------------------------------------------------------|

---

## Description

The local pooled error test attempts to reduce dependence on the within-gene estimates in tests for differential expression, by pooling error estimates within regions of similar intensity. Note that with the large number of genes there will be genes with low within-gene error estimates by chance, so that some signal-to-noise ratios will be large regardless of mean expression intensities and fold-change. The local pooled error attempts to avert this by combining within-gene error estimates with those of genes with similar expression intensity.

## Usage

```
calculateLpeAdj(x, y, basevar.x, basevar.y, df=10, array.type="olig",
  probe.set.name="OLIG.probe.name", trim.percent=5, adjust1=1.57, adjust2=1.57)
```

## Arguments

|                |                                                                                      |
|----------------|--------------------------------------------------------------------------------------|
| x              | Replicated data from first experimental condition (as matrix or data-frame)          |
| .              |                                                                                      |
| y              | Replicated data from second experimental condition (as matrix or data-frame)         |
| .              |                                                                                      |
| basevar.x      | Baseline distribution of first condition obtained from function baseOlig.error       |
| basevar.y      | Baseline distribution of second condition obtained from function baseOlig.error      |
| df             | Degrees of freedom used in fitting smooth.spline to estimates of var.M for bins in A |
| array.type     | Currently supports oligo arrays                                                      |
| probe.set.name | Gene IDs. By default if they are not provided then 1,2,3,... is assigned as GeneID   |
| trim.percent   | Percent of (A, var.M) estimates to trim from low end of A                            |
| adjust1        | adjustment factor of variance for first group                                        |
| adjust2        | adjustment factor of variance for second group                                       |

## Details

The LPE test statistic numerator is the difference in medians between the two experimental conditions. The test statistic denominator is the combined pooled standard error for the two experimental conditions obtained by looking up the var.M from each baseOlig.error variance function. The conversion to p-values is based on the Gaussian distribution for difference of order statistics (medians). The user may select both the smoother degrees of freedom (smaller is smoother) and the trim percent to obtain a variance function to suit particular issues i.e. variability of genes with low expression intensity. The default values for the adjustment of the variances of the two groups is the asymptotically correct value of  $\pi/2$ . This value is biased at small sample values and unbiased adjustment parameters based on sample size can be used instead. See documentation of lpeAdj for details.

**Value**

Data frame including x, median of x, y, median of y, median difference of (x,y), pooled standard deviation of difference, LPE p-value, outlier flag, probability of an outlier within x or y.

**Author(s)**

Carl Murie <carl.murie@mcgill.ca>, Nitin Jain <nitin.jain@pfizer.com>

**References**

J.K. Lee and M.O.Connell(2003). *An S-Plus library for the analysis of differential expression*. In *The Analysis of Gene Expression Data: Methods and Software*. Edited by G. Parmigiani, ES Garrett, RA Irizarry and SL Zegar. Springer, NewYork.

Jain et. al. (2003) *Local pooled error test for identifying differentially expressed genes with a small number of replicated microarrays*, Bioinformatics, 1945-1951.

Jain et. al. (2005) *Rank-invariant resampling based estimation of false discovery rate for analysis of small sample microarray data*, BMC Bioinformatics, Vol 6, 187.

**See Also**

[lpeAdj](#)

**Examples**

```
# Loading the data from the LPE library
data(Ley)

ADJ.VALUES <- c(1, 1, 1.34585905516761 ,1.19363228146169 ,1.436849413109
               ,1.289652132873 ,1.47658053092781 ,1.34382984852146
               ,1.49972130857404, 1.3835405678718)

dim(Ley)
# Gives 12488*7
# First column is ID.

# Subsetting the data
subset.Ley <- Ley[1:1000,]

subset.Ley[,2:7] <- preprocess(subset.Ley[,2:7],data.type="MAS5")

# Finding the baseline distribution of condition 1 and 2.
var.1 <- adjBaseOlig.error(subset.Ley[,2:4], q=0.01, setMax1=FALSE)
var.2 <- adjBaseOlig.error(subset.Ley[,5:7], q=0.01, setMax1=FALSE)

# Applying LPE
lpe.result <- calculateLpeAdj(subset.Ley[,2:4],subset.Ley[,5:7], var.1, var.2,
                             probe.set.name=subset.Ley[,1], adjust1=ADJ.VALUES[3],
                             adjust2=ADJ.VALUES[3])
```

---

|        |                                                                                                                                                                                          |
|--------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| lpeAdj | <i>High level lpeAdj function that executes the adjusted local pooled error significance test. If more control over parameters is needed then see documentation for calculateLpeAdj.</i> |
|--------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

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## Description

Applies the LPE algorithm with two additional options. The first is that the original LPE method sets all variances below the max variance in the ordered distribution of variances to the maximum variance. in LPEadj this option is turned off by default. The second option is to use a variance adjustment based on sample size rather than  $\pi/2$ . By default the LPEadj uses the sample size based variance adjustment. It is recommended to keep both of these options to the default.

## Usage

```
lpeAdj(dat, labels=NULL, doMax=FALSE, doAdj=TRUE, q=.01)
```

## Arguments

|        |                                                                                                                                                                                        |
|--------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| dat    | Replicated data of experiment containing two groups (as matrix or data-frame)                                                                                                          |
| .      |                                                                                                                                                                                        |
| labels | vector of group labels that correspond to the columns of dat. eg. labels=c(0,0,0,1,1,1) describes two groups with three replicates each                                                |
| .      |                                                                                                                                                                                        |
| doMax  | boolean: if T then all variances below the max variance in the ordered distribution of variances are set to the maximum variance. It is recommended to use the default value of False. |
| .      |                                                                                                                                                                                        |
| doAdj  | If T then run LPE with using variance adjustment value based on number of replicates (hardcoded in adjValues) rather than $\pi/2$ .                                                    |
| .      |                                                                                                                                                                                        |
| q      | is the quantile width; q=0.01 corresponds to 100 quantiles i.e. percentiles. Bins/quantiles have equal number of genes and are split according to the average intensity A.             |

## Details

The LPE test statistic numerator is the difference in medians between the two experimental conditions. The test statistic denominator is the combined pooled standard error for the two experimental conditions obtained by looking up the var.M from each baseOlig.error variance function. The conversion to p-values is based on the Gaussian distribution for difference of order statistics (medians).

**Value**

Data frame including x, median of x, y, median of y, median difference of (x,y), pooled standard deviation of difference, LPE p-value, outlier flag, probability of an outlier within x or y, .

**Author(s)**

Carl Murie <carl.murie@mcgill.ca>, Nitin Jain <nitin.jain@pfizer.com>

**References**

J.K. Lee and M.O.Connell(2003). *An S-Plus library for the analysis of differential expression*. In The Analysis of Gene Expression Data: Methods and Software. Edited by G. Parmigiani, ES Garrett, RA Irizarry and SL Zegar. Springer, New York.

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Jain et. al. (2005) *Rank-invariant resampling based estimation of false discovery rate for analysis of small sample microarray data*, BMC Bioinformatics, Vol 6, 187.

**Examples**

```
# Creating a null dataset (two groups with three
# replicates each)
dat <- matrix(rnorm(6000), ncol=6)

# Applying LPE
lpe.result <- lpeAdj(dat, labels=c(0,0,0,1,1,1))
```

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