

# Package ‘EBSEA’

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**Type** Package

**Title** Exon Based Strategy for Expression Analysis of genes

**Version** 1.35.0

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**Description** Calculates differential expression of genes based on exon counts of genes obtained from RNA-seq sequencing data.

**License** GPL-2

**biocViews** Software, DifferentialExpression, GeneExpression, Sequencing

**Imports** DESeq2, graphics, stats, EmpiricalBrownsMethod

**RoxygenNote** 7.1.1

**Encoding** UTF-8

**Suggests** knitr, rmarkdown

**VignetteBuilder** knitr

**Depends** R (>= 4.0.0)

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EBSEA

*Exon Based Strategy for Expression Analysis of genes***Description**

EBSEA takes the filtered raw exon-level read counts as input, normalizes and performs a two-group statistical comparison with DESeq2. The exon-level results are aggregated to the gene-level using empirical Brown's method. The samples in the two groups can be paired.

**Usage**

```
EBSEA(data, columnData, design, test = "Wald", contrasts = NULL, plot = FALSE)
```

**Arguments**

|                         |  |
|-------------------------|--|
| <code>data</code>       | A dataframe of raw exon-counts   |
| <code>columnData</code> | A dataframe indicated the groups of the samples.   |
| <code>design</code>     | Design matrix (see more information od design matrixes in DESeq2 reference manual)   |
| <code>test</code>       | The statistical test to be carried out. It can be either Wald or Likelihood Ratio Test. For further details about the methods you can look into DESeq2 refernce manual. Default: Wald                                    |
| <code>contrasts</code>  | a character vector with exactly three elements: the name of a factor in the design formula, the name of the numerator level for the fold change, and the name of the denominator level for the fold change Default: NULL |
| <code>plot</code>       | A logical value indicating a volcano plot is produced. Default: FALSE  |

**Value**

The function returns a list containing containing exon and gene-level results. ExonTable is a data frame containing an average expression, log2 fold-change, p-value and adjusted p-value. GeneTable is a data frame containing the gene-level p-value, and adjusted-value. Other returned elements include the raw and normalised exon-level read counts, group information and design matrix used.

**References**

Laiho, A., & Elo, L. L. (2014). A note on an exon-based strategy to identify differentially expressed genes in RNA-seq experiments. *PLoS One*, 9(12), e115964.

**Examples**

```
# The exon-based analysis for unpaired samples can be performed as follows:
data(exonCounts)
group <- data.frame('group' = as.factor(c('G1', 'G1', 'G1', 'G2', 'G2', 'G2', 'G2')))
row.names(group) <- colnames(exonCounts)
design <- ~group
```

```
ebsea.out <- EBSEA(exonCounts, group, design)
# The exon-based analysis for paired samples with contrast provided can be performed as follows:
data(exonCounts)
group <- data.frame('group' = as.factor(c('G1', 'G1', 'G1', 'G2', 'G2', 'G2', 'G2')),
  'paired' = as.factor(c(1,2,3,1,2,3,3)))
row.names(group) <- colnames(exonCounts)
design <- ~group
contrastInfo <- c('group', 'G2', 'G1')
ebsea.out <- EBSEA(exonCounts, group, design, contrasts = contrastInfo)
```

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|            |                                  |
|------------|----------------------------------|
| exonCounts | <i>Subset of Pasilla Dataset</i> |
|------------|----------------------------------|

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### Description

exonCounts consists of a subset of the exon counts from the pasilla dataset.

### Usage

```
data("exonCounts")
```

### Format

A data frame with 1000 rows and 7 variables

### Source

Exoncounts from Pasilla package <https://bioconductor.org/packages/release/data/experiment/html/pasilla.html>

### References

Huber W, Reyes A (2020). pasilla: Data package with per-exon and per-gene read counts of RNA-seq samples of Pasilla knock-down by Brooks et al., Genome Research 2011

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|              |                          |
|--------------|--------------------------|
| filterCounts | <i>Filter Count Data</i> |
|--------------|--------------------------|

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### Description

Filtering of exons based on their expression levels

### Usage

```
filterCounts(x, mean = 1, exonCount = 1)
```

**Arguments**

|           |  |
|-----------|--|
| x         | A numeric dataframe of exon counts across the samples. Exon number in format GeneName:Exonnumber should be indicated in the row name and sample names as column names. |
| mean      | Exons with average count value across the dataset less than mean are filtered out. Default: 1  |
| exonCount | After filtering the individual exons, only genes with at least the given number of exons remaining will be retained. Default: 1  |

**Value**

A dataframe of filtered counts of exons

**Examples**

```
data(exonCounts)
res <- filterCounts(exonCounts)
```

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visualizeGenes

*Visualize gene*

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**Description**

Produces a visualization summarizing the normalized read count in each sample group and expression difference across the expressed exons. Top panel contains the log<sub>2</sub> fold-change for each expressed exon. Asterisk denotes the significance level (\*: < 0.05, \*\*: < 0.01). Bottom panel shows the averaged normalized read count for each sample group. The title of the figure shows the gene name and the adjusted gene-level p-value (fdr)

**Usage**

```
visualizeGenes(gene, ebsea.out)
```

**Arguments**

|           |   |
|-----------|---|
| gene      | Gene name matching the input data.                                    |
| ebsea.out | Plots the mean count and fold-change the exons of the specified gene. |

**Value**

Plots the mean count and fold-change across the exons of the specified gene.

**Examples**

```
data(exonCounts)
group <- data.frame('group' = as.factor(c('G1', 'G1', 'G1', 'G2', 'G2', 'G2', 'G2')))
row.names(group) <- colnames(exonCounts)
design <- ~group
ebsea.out <- EBSEA(exonCounts, group, design)
visualizeGenes('FBgn000017', ebsea.out)
```

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## \* **datasets**

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