

Package ‘NBAMSeq’

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Type Package

Title Negative Binomial Additive Model for RNA-Seq Data

Version 1.18.0

Description High-throughput sequencing experiments followed by differential expression analysis is a widely used approach to detect genomic biomarkers. A fundamental step in differential expression analysis is to model the association between gene counts and covariates of interest. NBAMSeq a flexible statistical model based on the generalized additive model and allows for information sharing across genes in variance estimation.

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URL <https://github.com/reese3928/NBAMSeq>

BugReports <https://github.com/reese3928/NBAMSeq/issues>

Encoding UTF-8

Imports DESeq2, mgcv(>= 1.8-24), BiocParallel, genefilter, methods, stats,

Depends R (>= 3.6), SummarizedExperiment, S4Vectors

Suggests knitr, rmarkdown, testthat, ggplot2

RoxygenNote 6.1.0

VignetteBuilder knitr

biocViews RNASeq, DifferentialExpression, GeneExpression, Sequencing, Coverage

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R topics documented:

makeExample	2
makeplot	3
NBAMSeq	3
NBAMSeq-methods	4
NBAMSeqDataSet	5
NBAMSeqDataSet-class	6
results	7
Index	8

makeExample	<i>Make an example NBAMSeqDataSet</i>
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Description

This function makes an example NBAMSeqDataSet

Usage

```
makeExample(n = 200, m = 30)
```

Arguments

- n number of genes
- m number of samples

Value

a NBAMSeqDataSet object

References

Love, M.I., Huber, W., Anders, S. (2014) Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biology, 15:550. <https://doi.org/10.1186/s13059-014-0550-8>

Examples

```
gsd = makeExample()
```

makeplot*Making plots to visualize nonlinear associations*

Description

This function makes plots to visualize nonlinear associations.

Usage

```
makeplot(object, phenoname, genename, ...)
```

Arguments

object	a NBAMSeqDataSet object
phenoname	the name of nonlinear variable to be visualized
genename	the name of gene to be visualized
...	additional arguments provided to plot.gam

Value

the plot made by plot.gam() function

Examples

```
gsd = makeExample(n = 3, m = 10)
gsd = NBAMSeq(gsd)
makeplot(gsd, "pheno", "gene3", main = "gene10")
```

NBAMSeq*Differential expression analysis based on negative binomial additive model*

Description

This function performs differential expression analysis based on negative binomial additive model.

Usage

```
NBAMSeq(object, gamma = 2.5, parallel = FALSE, fitlin = FALSE,
  BPPARAM = bpparam(), ...)
```

Arguments

<code>object</code>	a NBAMSeqDataSet object
<code>gamma</code>	a number greater or equal to 1. Increase gamma to create smoother models. Default gamma is 2.5. See gam for details.
<code>parallel</code>	either TRUE or FALSE indicating whether parallel should be used. Default is FALSE
<code>fitlin</code>	either TRUE or FALSE indicating whether linear model should be fitted. Default is FALSE
<code>BPPARAM</code>	an argument provided to bplapply . See register for details.
<code>...</code>	additional arguments provided to gam

Value

a NBAMSeqDataSet object

References

Love, M.I., Huber, W., Anders, S. (2014) Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biology, 15:550. <https://doi.org/10.1186/s13059-014-0550-8>

Examples

```
gsd = makeExample(n = 3, m = 10)
gsd = NBAMSeq(gsd)
```

NBAMSeq-methods

Accessor functions and replace methods for NBAMSeqDataSet object

Description

Accessor functions and replace methods for NBAMSeqDataSet object

For `getDesign()`: accessor to the design formula

For `getsf()`: accessor to the size factors

Replace methods for NBAMSeqDataSet object

For `setsf()`: replace size factors

Usage

```
getDesign(theObject)
```

```
## S4 method for signature 'NBAMSeqDataSet'
getDesign(theObject)
```

```
getsf(theObject)
```

```
## S4 method for signature 'NBAMSeqDataSet'
getsf(theObject)

setsf(theObject) <- value

## S4 replacement method for signature 'NBAMSeqDataSet,numeric'
setsf(theObject) <- value
```

Arguments

theObject	a NBAMSeqDataSet object
value	the values to be included in the object

Value

For getDesign(): design formula
 For getsf(): size factor
 For setsf(): NBAMSeq object

References

Love, M.I., Huber, W., Anders, S. (2014) Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biology, 15:550. <https://doi.org/10.1186/s13059-014-0550-8>

Examples

```
## For getDesign() ##
gsd = makeExample()
design_gsd = getDesign(gsd)
## For getsf() ##
gsd = makeExample()
sf = getsf(gsd)
## For setsf() ##
n = 100
m = 50
gsd = makeExample(n = n, m = m)
sf = sample(1:5, m, replace = TRUE)
setsf(gsd) = sf
```

NBAMSeqDataSet

NBAMSeqDataSet constructor

Description

NBAMSeqDataSet constructor

Usage

```
NBAMSeqDataSet(countData, colData, design, ...)
```

Arguments

countData	a matrix or data frame contains gene count
colData	a DataFrame or data.frame
design	a mgcv type design. e.g. $\sim s(\text{pheno})$ or $\sim s(\text{pheno}) + \text{var1} + \text{var2}$
...	optional arguments passed to SummarizedExperiment

Value

a NBAMSeqDataSet object

Examples

```
n = 100 ## n stands for number of genes
m = 20  ## m stands for sample size
countData = matrix(rnbinom(n*m, mu=100, size=1/3), ncol = m)
mode(countData) = "integer"
colnames(countData) = paste0("sample", 1:m)
rownames(countData) = paste0("gene", 1:n)
pheno = runif(m, 20, 80)
colData = data.frame(pheno = pheno)
rownames(colData) = paste0("sample", 1:m)
gsd = NBAMSeqDataSet(countData = countData,
colData = colData, design = ~s(pheno))
```

NBAMSeqDataSet-class *NBAMSeqDataSet class*

Description

NBAMSeqDataSet is a class inherited from [SummarizedExperiment](#). It is used to store the count matrix, colData, and design formula in differential expression analysis.

Slots

design a mgcv-type design formula

References

Martin Morgan, Valerie Obenchain, Jim Hester and Hervé Pagès (2018). SummarizedExperiment: SummarizedExperiment container. R package version 1.12.0.

results	<i>Pulling out result</i>
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Description

This function pulls out result from NBAMSeqDataSet object returned by [NBAMSeq](#)

Usage

```
results(object, name, contrast, indepfilter = TRUE, alpha = 0.1,  
        pAdjustMethod = "BH", parallel = FALSE, BPPARAM = bpparam(), ...)
```

Arguments

object	a NBAMSeqDataSet object returned by NBAMSeq
name	the name of nonlinear variable or continuous linear variable
contrast	a character of length 3. 1st element: name of factor variable; 2nd element: name of numerator level; 3rd element: name of denominator level. contrast = c("group", "treatment", "control") means comparing treatment vs control for group variable.
indepfilter	either TRUE or FALSE indicating whether independent filtering should be performed. Default is TRUE.
alpha	significant threshold for declaring genes as differentially expressed. Default is 0.1.
pAdjustMethod	pvalue adjustment method. Default is "BH". See p.adjust for details.
parallel	either TRUE or FALSE indicating whether parallel should be used. Default is FALSE.
BPPARAM	an argument provided to bplapply . See register for details.
...	additional arguments provided to pvalueAdjustment function in DESeq2. See results for details.

Value

a DataFrame which contains the result

References

Love, M.I., Huber, W., Anders, S. (2014) Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. Genome Biology, 15:550. <https://doi.org/10.1186/s13059-014-0550-8>

Examples

```
gsd = makeExample(n = 3, m = 10)  
gsd = NBAMSeq(gsd)  
res = results(gsd, name = "pheno")
```

Index

bplapply, [4](#), [7](#)

gam, [4](#)

getDesign (NBAMSeq-methods), [4](#)

getDesign, NBAMSeqDataSet-method
(NBAMSeq-methods), [4](#)

getsf (NBAMSeq-methods), [4](#)

getsf, NBAMSeqDataSet-method
(NBAMSeq-methods), [4](#)

makeExample, [2](#)

makeplot, [3](#)

NBAMSeq, [3](#), [7](#)

NBAMSeq-methods, [4](#)

NBAMSeqDataSet, [5](#)

NBAMSeqDataSet-class, [6](#)

p.adjust, [7](#)

plot.gam, [3](#)

register, [4](#), [7](#)

results, [7](#), [7](#)

setsf<- (NBAMSeq-methods), [4](#)

setsf<- , NBAMSeqDataSet, numeric-method
(NBAMSeq-methods), [4](#)

SummarizedExperiment, [6](#)