

Package ‘satuRn’

January 20, 2026

Type Package

Title Scalable Analysis of Differential Transcript Usage for Bulk and Single-Cell RNA-sequencing Applications

Date 2022-08-05

Version 1.18.0

Description satuRn provides a highly performant and scalable framework for performing differential transcript usage analyses. The package consists of three main functions. The first function, fitDTU, fits quasi-binomial generalized linear models that model transcript usage in different groups of interest. The second function, testDTU, tests for differential usage of transcripts between groups of interest. Finally, plotDTU visualizes the usage profiles of transcripts in groups of interest.

Depends R (>= 4.1)

Imports locfdr, SummarizedExperiment, BiocParallel, limma, pbapply, ggplot2, boot, Matrix, stats, methods, graphics

Suggests knitr, rmarkdown, testthat, covr, BiocStyle, AnnotationHub, ensemblDb, edgeR, DEXSeq, stageR, DelayedArray

VignetteBuilder knitr

Collate 'data.R' 'satuRn-framework.R' 'allGenerics.R' 'accessors.R' 'fitDTU.R' 'testDTU.R' 'plotDTU.R'

License Artistic-2.0

URL <https://github.com/statOmics/satuRn>

BugReports <https://github.com/statOmics/satuRn/issues>

Encoding UTF-8

LazyData false

RoxygenNote 7.2.0

biocViews Regression, ExperimentalDesign, DifferentialExpression, GeneExpression, RNASeq, Sequencing, Software, SingleCell, Transcriptomics, MultipleComparison, Visualization

git_url <https://git.bioconductor.org/packages/satuRn>

git_branch RELEASE_3_22

git_last_commit 20071e7

git_last_commit_date 2025-10-29

Repository Bioconductor 3.22

Date/Publication 2026-01-19

Author Jeroen Gilis [aut, cre],
 Kristoffer Vitting-Seerup [ctb],
 Koen Van den Berge [ctb],
 Lieven Clement [ctb]

Maintainer Jeroen Gilis <jeroen.gilis@ugent.be>

Contents

fitDTU	2
getModel,StatModel-method	3
plotDTU	4
StatModel	6
StatModel-class	7
sumExp_example	8
Tasic_counts_vignette	8
Tasic_metadata_vignette	9
testDTU	9

Index	11
--------------	-----------

fitDTU	<i>fitDTU</i>
--------	---------------

Description

Parameter estimation of quasi-binomial models.

Usage

```
fitDTU(object, ...)
```

```
## S4 method for signature 'SummarizedExperiment'
fitDTU(
  object,
  formula,
  parallel = FALSE,
  BPPARAM = BiocParallel::bpparam(),
  verbose = TRUE
)
```

Arguments

object	A ‘SummarizedExperiment’ instance generated with the SummarizedExperiment function of the SummarizedExperiment package. In the assay slot, provide the transcript-level expression counts as an ordinary ‘matrix’, ‘DataFrame’, a ‘sparseMatrix’ or a ‘DelayedMatrix’. The ‘rowData’ slot must be a ‘DataFrame’ object describing the rows, which must contain a column ‘isoform_id’ with the row names of the expression matrix and a column ‘gene_id’ with the corresponding gene identifiers of each transcript. ‘colData’ is a ‘DataFrame’ describing the samples or cells in the experiment. Finally, specify the experimental design as a
--------	---

	formula in the metadata slot. This formula must be based on the colData. See the documentation examples and the vignette for more details.
...	parameters including:
formula	Model formula. The model is built based on the covariates in the data object.
parallel	Logical, defaults to FALSE. Set to TRUE if you want to parallelize the fitting procedure.
BPPARAM	object of class bpparamClass that specifies the back-end to be used for computations. See bpparam in BiocParallel package for details.
verbose	Logical, should progress be printed?

Value

An updated ‘SummarizedExperiment’ instance. The instance now includes a new list of models ("fitDTUModels") in its rowData slot, which can be accessed by `rowData(object)[["fitDTUModels"]]`.

Author(s)

Jeroen Gilis

Examples

```
data(sumExp_example, package = "saturn")
sumExp <- fitDTU(
  object = sumExp_example,
  formula = ~ 0 + group,
  parallel = FALSE,
  BPPARAM = BiocParallel::bpparam(),
  verbose = TRUE
)
```

getModel, StatModel-method

Accessor functions for StatModel class

Description

Accessor functions for StatModel class

getModel(object) to get model

getDF(object) to get the residual degrees of freedom of the model

getDfPosterior(object) to get the degrees of freedom of the empirical Bayes variance estimator

getDispersion(object) to get the dispersion estimate of the model

getCoef(object) to get the parameter estimates of the mean model

Usage

```
## S4 method for signature 'StatModel'
getModel(object)

## S4 method for signature 'StatModel'
getDF(object)

## S4 method for signature 'StatModel'
getDfPosterior(object)

## S4 method for signature 'StatModel'
getDispersion(object)

## S4 method for signature 'StatModel'
getCoef(object)
```

Arguments

object StatModel object

Value

The requested parameter of the StatModel object

Examples

```
## A fully specified dummy model
myModel <- StatModel(
  type = "glm",
  params = list(x = 3, y = 7, b = 4),
  varPosterior = c(0.1, 0.2, 0.3),
  dfPosterior = c(6, 7, 8)
)
getModel(myModel)
```

plotDTU

Plot function to visualize differential transcript usage (DTU)

Description

Plot function to visualize differential transcript usage (DTU)

Usage

```
plotDTU(
  object,
  contrast,
  groups,
  coefficients,
  summaryStat = "model",
  transcripts = NULL,
  genes = NULL,
```

```
    top.n = 6
  )
```

Arguments

object	A ‘SummarizedExperiment’ containing the models and results of the DTU analysis as obtained by the ‘fitDTU’ and ‘testDTU’ function from this ‘satuRn’ package, respectively.
contrast	Specifies the specific contrast for which the visualization should be returned. This should be one of the column names of the contrast matrix that was provided to the ‘testDTU’ function.
groups	A ‘list’ containing two character vectors. Each character vector contains the names (sample names or cell names) of the respective groups in the target contrast.
coefficients	A ‘list’ containing two numeric vectors. Each numeric vector specifies the model coefficient of the corresponding groups in the selected contrast.
summaryStat	Which summary statistic for the relative usage of the transcript should be displayed. ‘Character’ or ‘character vector’, must be any of following summary statistics; model (default), mean or median.
transcripts	A ‘character’ or ‘character vector’ of transcript IDs, to specify which transcripts should be visualized. Can be used together with ‘genes’. If not specified, ‘plotDTU’ will check if the ‘genes’ slot is specified.
genes	A single ‘character’ or ‘character vector’ of gene IDs, to specify the genes for which the individual transcripts should be visualized. Can be used together with ‘transcripts’. If not specified, ‘plotDTU’ will check if the ‘transcripts’ slot is specified.
top.n	A ‘numeric’ value. If neither ‘transcripts’ nor ‘genes’ was specified, this argument leads to the visualization of the ‘n’ most significant DTU transcripts in the contrast. Defaults to 6 transcripts.

Value

A ggplot object that can be directly displayed in the current R session or stored in a list.

Author(s)

Jeroen Gilis

Examples

```
data(sumExp_example, package = "satuRn")
library(SummarizedExperiment)
sumExp <- fitDTU(
  object = sumExp_example,
  formula = ~ 0 + group,
  parallel = FALSE,
  BPPARAM = BiocParallel::bpparam(),
  verbose = TRUE
)
group <- as.factor(colData(sumExp)$group)
design <- model.matrix(~ 0 + group)
colnames(design) <- levels(group)
```

```

L <- matrix(0, ncol = 2, nrow = ncol(design))
rownames(L) <- colnames(design)
colnames(L) <- c("Contrast1", "Contrast2")
L[c("VISp.L5_IT_VISp_Hsd11b1_Endou", "ALM.L5_IT_ALM_Tnc"), 1] <- c(1, -1)
L[c("VISp.L5_IT_VISp_Hsd11b1_Endou",
    "ALM.L5_IT_ALM_Tmem163_Dmrtb1"), 2] <- c(1, -1)

sumExp <- testDTU(object = sumExp,
  contrasts = L,
  diagplot1 = FALSE,
  diagplot2 = FALSE,
  sort = FALSE)

group1 <- rownames(colData(sumExp))[colData(sumExp)$group ==
  "VISp.L5_IT_VISp_Hsd11b1_Endou"]
group2 <- rownames(colData(sumExp))[colData(sumExp)$group ==
  "ALM.L5_IT_ALM_Tnc"]

plots <- plotDTU(
  object = sumExp,
  contrast = "Contrast1",
  groups = list(group1, group2),
  coefficients = list(c(0, 0, 1), c(0, 1, 0)),
  summaryStat = "model",
  transcripts = c("ENSMUST00000165123",
    "ENSMUST00000165721",
    "ENSMUST0000005067"),
  genes = NULL,
  top.n = 6
)

```

StatModel

StatModel

Description

Function for constructing a new ‘StatModel’ object.

Usage

```

StatModel(
  type = "fitError",
  params = list(),
  varPosterior = NA_real_,
  dfPosterior = NA_real_
)

```

Arguments

<code>type</code>	default set to fiterror, can be a glm
<code>params</code>	A list containing the parameters of the fitted glm
<code>varPosterior</code>	Numeric, posterior variance of the glm, default is NA
<code>dfPosterior</code>	Numeric, posterior degrees of freedom of the glm, default is NA

Value

A StatModel object

Examples

```
## A fully specified dummy model
myModel <- StatModel(
  type = "glm",
  params = list(x = 3, y = 7, b = 4),
  varPosterior = c(0.1, 0.2, 0.3),
  dfPosterior = c(6, 7, 8)
)
myModel
```

StatModel-class

The StatModel class for satuRn

Description

The ‘StatModel’ class contains a statistical model generated by the DTU analysis.

Models are created by the dedicated user-level function ‘fitDTU()’ or manually, using the ‘StatModel()’ constructor. In the former case, each quantitative feature is assigned its statistical model and the models are stored as a variable in a ‘DataFrame’ object, that in turn will be stored in the ‘RowData’ slot of a ‘SummarizedExperiment’ object.

Slots

type ‘character(1)’ defining type of the used model. Default is “fitError”, i.e. an error model. If not an error, class type will be “glm”.

params A ‘list()’ containing the parameters of the fitted model.

varPosterior ‘numeric()’ of posterior variance.

dfPosterior ‘numeric()’ of posterior degrees of freedom.

Author(s)

Jeroen Gilis

Examples

```
## A fully specified dummy model
myModel <- StatModel(
  type = "glm",
  params = list(x = 3, y = 7, b = 4),
  varPosterior = c(0.1, 0.2, 0.3),
  dfPosterior = c(6, 7, 8)
)
myModel
```

sumExp_example	<i>A ‘SummarizedExperiment’ derived from our case study which builds on the dataset of Tasic et al. It contains the same cells as the data object used in the vignette (see ‘?Tasic_counts_vignette’ for more information). In this SummarizedExperiment, we performed a filtering with ‘filterByExpr’ of edgeR with more stringent than default parameter settings (min.count = 100,min.total.count = 200, large.n = 50, min.prop = 0.9) to reduced the number of retained transcripts. We used this object to create an executable example in the help files of satuRn.</i>
----------------	---

Description

A ‘SummarizedExperiment’ derived from our case study which builds on the dataset of Tasic et al. It contains the same cells as the data object used in the vignette (see ‘?Tasic_counts_vignette’ for more information). In this SummarizedExperiment, we performed a filtering with ‘filterByExpr’ of edgeR with more stringent than default parameter settings (min.count = 100,min.total.count = 200, large.n = 50, min.prop = 0.9) to reduced the number of retained transcripts. We used this object to create an executable example in the help files of satuRn.

Usage

```
data(sumExp_example)
```

Format

An object of class SummarizedExperiment with 1286 rows and 60 columns.

Tasic_counts_vignette	<i>A ‘Matrix’ with transcript-level counts derived from our case study which builds on the dataset of Tasic et al. We used Salmon (V1.1.0) to quantify all L5IT cells (both for ALM and VISp tissue) from mice with a normal eye condition. From these cells, we randomly sampled 20 cells of each of the following cell types to use for this vignette; L5_IT_VISp_Hsd11b1_Endou, L5_IT_ALM_Tmem163_Dmrtb1 and L5_IT_ALM_Tnc. The data has already been leniently filtered with the ‘filterByExpr’ function of edgeR.</i>
-----------------------	--

Description

A ‘Matrix’ with transcript-level counts derived from our case study which builds on the dataset of Tasic et al. We used Salmon (V1.1.0) to quantify all L5IT cells (both for ALM and VISp tissue) from mice with a normal eye condition. From these cells, we randomly sampled 20 cells of each of the following cell types to use for this vignette; L5_IT_VISp_Hsd11b1_Endou, L5_IT_ALM_Tmem163_Dmrtb1 and L5_IT_ALM_Tnc. The data has already been leniently filtered with the ‘filterByExpr’ function of edgeR.

Usage

```
data(Tasic_counts_vignette)
```


Format

An object of class matrix (inherits from array) with 22273 rows and 60 columns.

Tasic_metadata_vignette

Metadata associated with the expression matrix 'Tasic_counts_vignette'. See '?Tasic_counts_vignette' for more information on the dataset.

Description

Metadata associated with the expression matrix 'Tasic_counts_vignette'. See '?Tasic_counts_vignette' for more information on the dataset.

Usage

```
data(Tasic_metadata_vignette)
```

Format

An object of class data.frame with 60 rows and 3 columns.

testDTU

Test function to obtain a top list of transcripts that are differentially used in the contrast of interest

Description

Function to test for differential transcript usage (DTU)

Usage

```
testDTU(object, contrasts, diagplot1 = TRUE, diagplot2 = TRUE, sort = FALSE)
```

Arguments

object	A 'SummarizedExperiment' instance containing a list of objects of the 'StatModel' class as obtained by the 'fitDTU' function of the 'satuRn' package.
contrasts	'numeric' matrix specifying one or more contrasts of the linear model coefficients to be tested. The rownames of the matrix should be equal to the names of parameters of the model that are involved in the contrast. The column names of the matrix will be used to construct names to store the results in the rowData of the SummarizedExperiment.

diagplot1	'boolean(1)' Logical, defaults to TRUE. If set to TRUE, a plot of the histogram of the z-scores (computed from p-values) is displayed using the locfdr function of the 'locfdr' package. The blue dashed curve is fitted to the mid 50 to originate from null transcripts, thus representing the estimated empirical null component densities. The maximum likelihood estimates (MLE) and central matching estimates (CME) of this estimated empirical null distribution are given below the plot. If the values for delta and sigma deviate from 0 and 1 respectively, the downstream inference will be influenced by the empirical adjustment implemented in satuRn.
diagplot2	'boolean(1)' Logical, defaults to TRUE. If set to TRUE, a plot of the histogram of the "empirically adjusted" test statistics and the standard normal distribution will be displayed. Ideally, the majority (mid portion) of the adjusted test statistics should follow the standard normal.
sort	'boolean(1)' Logical, defaults to FALSE. If set to TRUE, the output of the topT-able test function is sorted according to the empirical p-values.

Value

An updated 'SummarizedExperiment' that contains the 'Dataframes' that display the significance of DTU for each transcript in each contrast of interest.

Author(s)

Jeroen Gilis

Examples

```
data(sumExp_example, package = "satuRn")
library(SummarizedExperiment)
sumExp <- fitDTU(
  object = sumExp_example,
  formula = ~ 0 + group,
  parallel = FALSE,
  BPPARAM = BiocParallel::bpparam(),
  verbose = TRUE
)
group <- as.factor(colData(sumExp)$group)
design <- model.matrix(~ 0 + group)
colnames(design) <- levels(group)
L <- matrix(0, ncol = 2, nrow = ncol(design))
rownames(L) <- colnames(design)
colnames(L) <- c("Contrast1", "Contrast2")
L[c("VISP.L5_IT_VISP_Hsd11b1_Endou",
    "ALM.L5_IT_ALM_Tnc"), 1] <- c(1, -1)
L[c("VISP.L5_IT_VISP_Hsd11b1_Endou",
    "ALM.L5_IT_ALM_Tmem163_Dmrtb1"), 2] <- c(1, -1)

sumExp <- testDTU(object = sumExp,
  contrasts = L,
  diagplot1 = FALSE,
  diagplot2 = FALSE,
  sort = FALSE)
```

Index

- * **datasets**
 - sumExp_example, [8](#)
 - Tasic_counts_vignette, [8](#)
 - Tasic_metadata_vignette, [9](#)
 - .StatModel (StatModel-class), [7](#)
- fitDTU, [2](#)
- fitDTU, SummarizedExperiment-method
 - (fitDTU), [2](#)
- getCoef (getModel, StatModel-method), [3](#)
- getCoef, StatModel-method
 - (getModel, StatModel-method), [3](#)
- getDF (getModel, StatModel-method), [3](#)
- getDF, StatModel-method
 - (getModel, StatModel-method), [3](#)
- getDfPosterior
 - (getModel, StatModel-method), [3](#)
- getDfPosterior, StatModel-method
 - (getModel, StatModel-method), [3](#)
- getDispersion
 - (getModel, StatModel-method), [3](#)
- getDispersion, StatModel-method
 - (getModel, StatModel-method), [3](#)
- getModel (getModel, StatModel-method), [3](#)
- getModel, StatModel-method, [3](#)
- plotDTU, [4](#)
- StatModel, [6](#)
- StatModel-class, [7](#)
- statModelAccessors
 - (getModel, StatModel-method), [3](#)
- sumExp_example, [8](#)
- Tasic_counts_vignette, [8](#)
- Tasic_metadata_vignette, [9](#)
- testDTU, [9](#)