

# Package ‘switchde’

October 17, 2020

**Type** Package

**Title** Switch-like differential expression across single-cell  
trajectories

**Version** 1.14.0

**Date** 2017-10-24

**Description** Inference and detection of switch-like differential  
expression across single-cell RNA-seq trajectories.

**License** GPL (>= 2)

**LazyData** TRUE

**biocViews** ImmunoOncology, Software, Transcriptomics, GeneExpression,  
RNASeq, Regression, DifferentialExpression, SingleCell

**Depends** R (>= 3.4), SingleCellExperiment

**Imports** SummarizedExperiment, dplyr, ggplot2, methods, stats

**Suggests** knitr, rmarkdown, BiocStyle, testthat, numDeriv, tidyR

**VignetteBuilder** knitr

**RoxygenNote** 6.0.1

**URL** <https://github.com/kieranrcampbell/switchde>

**BugReports** <https://github.com/kieranrcampbell/switchde>

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<i>example_sigmoid</i>	<i>Example sigmoid plot</i>
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**Description**

Plot an example sigmoid function. For demonstration and documentation.

**Usage**

```
example_sigmoid()
```

**Value**

An object of class ggplot

**Examples**

```
example_sigmoid()
```

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<i>extract_pars</i>	<i>Extract parameters from fitted model</i>
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**Description**

Extract maximum likelihood parameter estimates from a call to *switchde*.

**Usage**

```
extract_pars(sde, gene)
```

**Arguments**

<i>sde</i>	The <code>data.frame</code> returned by <i>switchde</i>
<i>gene</i>	The gene for which to extract parameters

**Value**

A vector of length 3 corresponding to the parameters  $\mu_0$ ,  $k$  and  $t_0$

**Examples**

```
data(synth_gex)
data(ex_pseudotime)
sde <- switchde(synth_gex, ex_pseudotime)
pars <- extract_pars(sde, "Gene1")
```

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ex_pseudotime	<i>Synthetic gene pseudotimes</i>
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**Description**

A vector with example pseudotimes for the synthetic gene expression data in example\_gex

**Usage**

```
ex_pseudotime
```

**Format**

An object of class `array` of length 100.

**Value**

A vector of length 100

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fit_nzi_model	<i>Fit a (non-zero-inflated) model for a single gene</i>
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**Description**

Fits a sigmoidal expression model for a single gene vector, returning MLE model parameters and p-value.

**Usage**

```
fit_nzi_model(y, pst)
```

**Arguments**

y	Vector of gene expression values
pst	Pseudotime vector, of same length as y

**Value**

A vector with 5 entries: maximum likelihood estimates for  $\mu_0$ ,  $k$   $t_0$ ,  $\sigma^2$  and a p-value

**Examples**

```
data(synth_gex)
data(ex_pseudotime)
y <- synth_gex[1, ]
fit <- fit_nzi_model(y, ex_pseudotime)
```

<code>fit_zi_model</code>	<i>Fit a zero-inflated model for a single gene</i>
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## Description

Fits a zero-inflated sigmoidal model for a single gene vector, returning MLE model parameters and p-value.

## Usage

```
fit_zi_model(y, pst, maxiter = 10000, log_lik_tol = 0.001,
             verbose = FALSE)
```

## Arguments

<code>y</code>	Vector of gene expression values
<code>pst</code>	Pseudotime vector, of same length as <code>y</code>
<code>maxiter</code>	Maximum number of iterations for EM algorithm if zero inflation enabled. Default 100
<code>log_lik_tol</code>	If the change in the log-likelihood falls below this for zero inflated EM the algorithm is assumed to have converged
<code>verbose</code>	Print convergence update for EM algorithm

## Value

A vector with 6 entries: maximum likelihood estimates for  $\mu_0$ ,  $k$   $t_0$ ,  $\lambda$ ,  $\sigma^2$  and a p-value

## Examples

```
data(synth_gex)
data(ex_pseudotime)
y <- synth_gex[1, ]
fit <- fit_zi_model(y, ex_pseudotime)
```

<code>switchde</code>	<i>Switch-like model fitting and differential expression test</i>
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## Description

Fit sigmoidal differential expression models to gene expression across pseudotime. Parameter estimates are returned along with a p-value for switch-like differential expression over a null model (constant expression).

## Usage

```
switchde(object, pseudotime = NULL, zero_inflated = FALSE,
         lower_threshold = 0.01, maxiter = 1000, log_lik_tol = 0.01,
         verbose = FALSE, sce_assay = "exprs")
```

**Arguments**

object	Gene expression data that is either <ul style="list-style-type: none"> <li>• A vector of length number of cells for a single gene</li> <li>• A matrix of dimension number of genes x number of cells</li> <li>• An object of class <code>SingleCellExperiment</code> from package <code>SingleCellExperiment</code></li> </ul>
pseudotime	A pseudotime vector with a pseudotime corresponding to every cell. Can be <code>NULL</code> if <code>object</code> is of class <code>SCESet</code> and <code>colData(sce)\$pseudotime</code> is defined.
zero_inflated	Logical. Should zero inflation be implemented? Default <code>FALSE</code>
lower_threshold	The minimum threshold below which to set expression to zero to avoid numerical issues. Default is <code>0.01</code>
maxiter	Maximum number of iterations for EM algorithm if zero inflation enabled. Default <code>100</code>
log_lik_tol	If the change in the log-likelihood falls below this for zero inflated EM the algorithm is assumed to have converged
verbose	Print convergence update for EM algorithm
sce_assay	The assay from the <code>SingleCellExperiment</code> to be used as expression, defaulting to "exprs"

**Value**

A matrix where each column corresponds to a gene, the first row is the p-value for that gene and subsequent rows are model parameters.

**Examples**

```
data(synth_gex)
data(ex_pseudotime)
sde <- switchde(synth_gex, ex_pseudotime)
```

switchplot

*Plot gene behaviour***Description**

Plot gene behaviour and MLE sigmoid as a function of pseudotime.

**Usage**

```
switchplot(x, pseudotime, pars)
```

**Arguments**

x	Gene expression vector
pseudotime	Pseudotime vector (of same length as x)
pars	Fitted model parameters

**Details**

This plots expression of a single gene. Fitted model parameters can either be specified manually or can be extracted from the `data.frame` returned by `switchde` using the function `extract_pars`.

**Value**

A `ggplot2` plot of gene expression and MLE sigmoid

**Examples**

```
data(synth_gex)
data(ex_pseudotime)
sde <- switchde(synth_gex, ex_pseudotime)
switchplot(synth_gex[1, ], ex_pseudotime, extract_pars(sde, "Gene1"))
```

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`synth_gex`

*Synthetic gene expression matrix*

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**Description**

A matrix containing some synthetic gene expression data for 100 cells and 12 genes

**Usage**

`synth_gex`

**Format**

An object of class `matrix` with 12 rows and 100 columns.

**Value**

A 12 by 100 matrix

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