

Package ‘CellTrails’

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Type Package

Title Reconstruction, visualization and analysis of branching trajectories

Version 1.28.0

Description CellTrails is an unsupervised algorithm for the de novo chronological ordering, visualization and analysis of single-cell expression data. CellTrails makes use of a geometrically motivated concept of lower-dimensional manifold learning, which exhibits a multitude of virtues that counteract intrinsic noise of single cell data caused by drop-outs, technical variance, and redundancy of predictive variables. CellTrails enables the reconstruction of branching trajectories and provides an intuitive graphical representation of expression patterns along all branches simultaneously. It allows the user to define and infer the expression dynamics of individual and multiple pathways towards distinct phenotypes.

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Encoding UTF-8

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Author Daniel Ellwanger [aut, cre, cph]

Maintainer Daniel Ellwanger <dc.ellwanger.dev@gmail.com>

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<i>.adjustLayoutByPtime</i>	<i>Adjusting the trajectory graph layout</i>
-----------------------------	--

Description

Adjusts the edge distances in the trajectory graph such that edge distances correlates to edge weights; pseudotime is stored in edge weights

Usage

```
.adjustLayoutByPtime(x, l)
```

Arguments

x	A SingleCellExperiment object
layout	Layout coordinates for each sample

Value

An adjusted layout

Author(s)

Daniel C. Ellwanger

```
.bhtsne      #' DEF: Truncate eigenbasis #' #' For details see reduceDimensions
             #' @keywords internal #' @author Daniel C. Ellwanger .reduceD-
             imensions_def <- function(x, s) CellTrails::latentSpace(x) <- Cell-
             Trails::latentSpace(x)[, seq_len(s@n)] CellTrails::eigenvalues(x) <-
             CellTrails::eigenvalues(x)[seq_len(s@n)] x t-Distributed Stochastic
             Neighbor Embedding
```

Description

Barnes-Hut implementation of t-Distributed Stochastic Neighbor Embedding

Usage

```
.bhtsne(x, dims = 2, perplexity = 30, theta = 0.5, max_iter = 1000)
```

Arguments

x	A numerical matrix
dims	Output dimensionality
perplexity	Perplexity parameter (default: 30)
theta	Speed/accuracy trade-off (increase for less accuracy), set to 0.0 for exact tSNE (default: .5)
max_iter	Number of iterations (default: 1000)

Value

A list with the following components:

Y Matrix containing the new representations for the objects

perplexity See above

Author(s)

Daniel C. Ellwanger

```
.capitalize      Capitalizes first character of string
```

Description

Capitalizes first character of string and sets the rest to lower case.

Usage

```
.capitalize(x)
```

Arguments

x A string

Details

Example: "abC" becoms "Abc".

Value

A string

Author(s)

Daniel C. Ellwanger

.color_hue

Color palette

Description

Generates equally spaced hues around the color wheel

Usage

```
.color_hue(n)
```

Arguments

n Number of colors to be generated

Value

Colors codes

Author(s)

Daniel C. Ellwanger

.color_ramp	Colors for vector
-------------	-------------------

Description

Generates colors for vector

Usage

```
.color_ramp(  
  x,  
  range = 3,  
  colPal = NULL,  
  min.val = 1e-10,  
  min.val.col = NA,  
  breaks = 25  
)
```

Arguments

x	A numeric vector
---	------------------

Value

Color codes

Author(s)

Daniel C. Ellwanger

.connectStates_def	DEF: Connect states
--------------------	---------------------

Description

For details see connectStates

Usage

```
.connectStates_def(dmat, cl, l = 10, sigma = 1)
```

Arguments

dmat	Distance matrix
cl	States

Author(s)

Daniel C. Ellwanger

.connect_ordi	<i>AUX Generate graph based on ordination</i>
---------------	---

Description

Generates 2D ordination from orthogonal projection

Usage

```
.connect_ordi(ordi)
```

Arguments

ordi	Ordination
------	------------

Value

An igraph object

Author(s)

Daniel C. Ellwanger

.connect_ortho	<i>AUX: Connect orthogonal projected samples</i>
----------------	--

Description

Connects samples based on their projected position on the trajectory.

Usage

```
.connect_ortho(Xorth)
```

Arguments

Xorth	Result from the orthogonal projection
-------	---------------------------------------

Value

An igraph object

Author(s)

Daniel C. Ellwanger

`.contrastExprTrail_def`*DEF: Differential expression between trails*

Description

DEF: Differential expression between trails

Usage

```
.contrastExprTrail_def(  
  ptime_1,  
  ptime_2,  
  feature_expr,  
  sts,  
  trail_names,  
  n = 250,  
  score,  
  dynamicFit_k = 5,  
  df = 10  
)
```

Arguments

<code>ptime_1</code>	Pseudotime trail 1
<code>ptime_2</code>	Pseudotime trail 2
<code>feature_expr</code>	Expression vector
<code>sts</code>	Samples vector
<code>n</code>	Number of fitted values
<code>score</code>	Score type
<code>dynamicFit_k</code>	Freedom param for dynamic fit
<code>df</code>	Freedom param for splines fit

Author(s)

Daniel C. Ellwanger

`.deleteMedianCentres` *AUX Remove median centres*

Description

Deletes median centres from trajectory graph if graph has not been simplified/has been expanded.

Usage

```
.deleteMedianCentres(g)
```

Arguments

`g` An object of class `igraph`

Value

An unupdated object of class `igraph`

Author(s)

Daniel C. Ellwanger

`.denoiseExpression` *Denoises expression*

Description

Blocks factors in expression matrix

Usage

`.denoiseExpression(x, design)`

Arguments

`x` A numeric expression vector
`design` A numeric matrix describing the factors that should be blocked

Value

A numeric denoised expression vector

Author(s)

Daniel C. Ellwanger

`.diffExpr` *Compute differential expression*

Description

Computes P-value and fold-change between two expression vectors.

Usage

`.diffExpr(x, y, lod = NULL, alternative = "two.sided")`

Arguments

`x` Feature expression in condition x
`y` Feature expression in condition y
`lod` Gene limit of detection

Details

For censored data a Peto-Peto test is performed, for non-censored data a Wilcoxon rank sum test. If limit of detection is provided, expectation maximization is used to compute the fold-change.

Value

A list containing the following components:

<code>p.value</code>	P-value
<code>fold</code>	Fold-change

Author(s)

Daniel C. Ellwanger

<code>.diffExprState_def</code>	<i>DEF: Differential expression between states</i>
---------------------------------	--

Description

DEF: Differential expression between states

Usage

```
.diffExprState_def(  
  x,  
  state1,  
  state2,  
  feature_name,  
  alternative = "two.sided",  
  lod = NULL  
)
```

Author(s)

Daniel C. Ellwanger

<code>.embedSamples_def</code>	<i>DEF: Spectral embedding of samples</i>
--------------------------------	---

Description

For details see `embedSamples`

Usage

```
.embedSamples_def(x, nbins = 10, sigma = 0.75, design = NULL)
```

Arguments

x	A numerical matrix
nbins	Cubic B-spline discretization is used to compute fuzzy mutual information between pairs of samples; nbins defines the number of intervals used for discretization of expression data. (default: 10)
sigma	The mutual information matrix is transformed to an adjacency matrix using a heat kernel; sigma defines the radius of heat kernel (in quantiles; default: sigma = 0.75 which is the third quantile of the mutual information matrix).
design	A numeric matrix describing the factors that should be blocked

Author(s)

Daniel C. Ellwanger

.exprs

GET expression matrix

Description

Retrieve numeric matrix of expression values for processing in CellTrails. This wrapper function ensures that all functions in the package receive the proper assay from the SingleCellExperiment object.

Usage

```
.exprs(object)
```

Arguments

object	An object of class SingleCellExperiment
--------	---

Value

A numeric matrix

Author(s)

Daniel C. Ellwanger

.featureNameExists	<i>Checks if feature exists</i>
--------------------	---------------------------------

Description

Checks if feature exists

Usage

```
.featureNameExists(x, feature_name)
```

Arguments

x	A SingleCellExperiment object
feature_name	Name of feature

Value

logical value

Author(s)

Daniel C. Ellwanger

.filterTrajFeaturesByCOV_def	<i>DEF: Filter features by coefficient of variation</i>
------------------------------	---

Description

For details see filterFeaturesByCOV

Usage

```
.filterTrajFeaturesByCOV_def(y, threshold, design = NULL, show_plot = TRUE)
```

Arguments

y	An expression vector
threshold	numeric cutoff value
design	Model matrix
show_plot	Show plot?

Author(s)

Daniel C. Ellwanger

```
.filterTrajFeaturesByDL_def
```

DEF: Filter feaures by Detection Level

Description

For details see filterFeaturesByPOD

Usage

```
.filterTrajFeaturesByDL_def(y, threshold, show_plot = TRUE)
```

Arguments

y	An expression matrix
threshold	numeric cutoff value
show_plot	Show plot?

Author(s)

Daniel C. Ellwanger

```
.filterTrajFeaturesByFF_def
```

DEF: Filter features by index of dispersion / fano factor

Description

For details see filterFeaturesByFF

Usage

```
.filterTrajFeaturesByFF_def(
  y,
  z,
  min_expr = 0,
  design = NULL,
  show_plot = TRUE
)
```

Arguments

y	An expression vector
z	A cutoff z-score
min_expr	Minimum expression level
design	Model matrix
show_plot	Show plot?

Author(s)

Daniel C. Ellwanger

.findSpectrum_def *DEF: Determine number of informative latent dimensions*

Description

For details see findSpectrum

Usage

```
.findSpectrum_def(D, frac = 100)
```

Author(s)

Daniel C. Ellwanger

.findStates_def *DEF: Find states*

Description

For details see findStates

Usage

```
.findStates_def(  
  X,  
  ordi,  
  link.method = "ward.D2",  
  min.size,  
  max.pval = 1e-04,  
  min.fc = 2,  
  min.g = 5,  
  reverse = FALSE,  
  verbose = FALSE  
)
```

Arguments

X	Expression matrix
ordi	Ordination of samples
link.method	Linkage criteria
min.size	Min size of initial clusters
max.pval	Pval threshold
min.fc	Fold-change threshold
reverse	Reverse order
verbose	For debug
min.g	Feature count threshold

Author(s)

Daniel C. Ellwanger

.fitDynamic_def

Fitting dynamics

Description

Fits expression as a function of pseudotime using generalized additive models

Usage

```
.fitDynamic_def(x, y, z, k = 5, n.out = NULL, x.out = NULL)
```

Arguments

x	Pseudotime values
y	Expression values
z	Sample states
k	Dimensions of the bases used to represent the smooth term
n.out	Number of predicted expression values within pseudotime range
x.out	Predicted expression values for given set of pseudotime values

Value

A list containing the following components:

x	Pseudotime
y	Fitted values
mod	GAM

Author(s)

Daniel C. Ellwanger

.fitTrajectory_def *DEF: Trajectory fitting*

Description

For details see `fitTrajectory`

Usage

```
.fitTrajectory_def(cl = cl, g = g, X = X, snames)
```

Arguments

<code>cl</code>	Trajectory states vector
<code>g</code>	Trajectory graph; <code>igraph</code> object
<code>X</code>	Lower-dimensional ordination of trajectory samples
<code>snames</code>	Sample names

Author(s)

Daniel C. Ellwanger

.fit_surface *Fitting the map surface*

Description

Fits the smooth surface of CellTrails maps

Usage

```
.fit_surface(X, y, npoints = 300, weights = NULL, knots = 10, rescale = FALSE)
```

Arguments

<code>X</code>	Ordination; numerical matrix
<code>y</code>	Expression values; numerical vector
<code>npoints</code>	Number of points along x and y axis
<code>weights</code>	Cluster states (defines weights based on fraction of non-detects per state)
<code>knots</code>	Number of knots in spline function

Value

A list containing the following components:

<code>fit</code>	GAM fit object
<code>grid</code>	Fitted values

Author(s)

Daniel C. Ellwanger

.fr_layout	<i>Computes state trajectory graph layout</i>
------------	---

Description

Uses the Fruchterman-Reingold layout algorithm

Usage

```
.fr_layout(g)
```

Arguments

g	An igraph graph object
---	------------------------

Value

numerical matrix with the layout coordinates

Author(s)

Daniel C. Ellwanger

.generate_ordination	<i>AUX: Orthogonal projection ordination</i>
----------------------	--

Description

Generates 2D ordination from orthogonal projection

Usage

```
.generate_ordination(g, error, factor = 7, rev = FALSE, only.ordi = FALSE)
```

Arguments

g	Graph from orthogonal projection, igraph object
error	Error from orthogonal projection; numeric vector
only.ordi	Compute only ordination?

Value

A list containing the following components:

ordi	Ordination
ordi.jitter	Jittered ordination
backbone	Backbone of trajectory

Author(s)

Daniel C. Ellwanger

.ihs	<i>Reverse inverse hyperbolic sine</i>
------	--

Description

Computes inverse hyperbolic sine

Usage

```
.ihs(x)
```

Arguments

x	A numeric value, vector or matrix
---	-----------------------------------

Details

In contrast to sqrt and log, ihs is defined for negative numbers.

Value

Transformed values

Author(s)

Daniel C. Ellwanger

.linear_fit	<i>Linear fit</i>
-------------	-------------------

Description

Perform simple linear fit

Usage

```
.linear_fit(x.in, y.in, x.out = NULL)
```

Arguments

x.in	Predictor variable, numeric vector
y.in	Response variable, numeric vector
x.out	Predictor variable for linear function, numeric vector

Value

A list containing the following components:

p.value	Anova P-value
r2	Adjusted R-squared
x.out	Predictor variable for linear function, numeric vector
y.out	Response from linear function

Author(s)

Daniel C. Ellwanger

.needsToBeExpanded	<i>AUX Check if projection needs to be expanded</i>
--------------------	---

Description

Simple graphs having only bifurcations along the backbone get simplified by a 2D projection. If a fork with more than two successors or a fork not located along the backbone, but along a side branch, the 2D simplification would collate the states and ignore the bifurcation. Therefore, the simplification step has to be skipped. This methods checks whether a simplification is possible or not.

Usage

```
.needsToBeExpanded(g, cl)
```

Arguments

g	Trajectory graph; igraph object
cl	Trajectory states vector; logical

Details

IF: any node has a degree > 3, return TRUE; OTHERWISE: Check all shortest paths from start node to any leaf in trajectory tree. IF any path passes a node with degree > 3, which is not located on the backbone, return TRUE; OTHERWISE: return FALSE.

Value

A logical value

Author(s)

Daniel C. Ellwanger

.nn_impute	<i>Nearest neighbor imputation</i>
------------	------------------------------------

Description

Nearest neighbor imputation

Usage

```
.nn_impute(y, D)
```

Arguments

<code>y</code>	Values; missing values need to be set to NA
<code>D</code>	distance matrix between samples

Value

Imputed values

Author(s)

Daniel C. Ellwanger

<code>.pca_def</code>	<i>DEF: PCA</i>
-----------------------	-----------------

Description

For details see `pca`

Usage

```
.pca_def(M, do_scaling = TRUE, design = NULL)
```

Arguments

<code>M</code>	expression matrix
<code>do_scaling</code>	use covariance/correlation matrix
<code>design</code>	Block factors

Author(s)

Daniel C. Ellwanger

<code>.pheno</code>	<i>GET phenotype values</i>
---------------------	-----------------------------

Description

Returns phenotype values from `SingleCellExperiment` object

Usage

```
.pheno(object, name)
```

Arguments

<code>object</code>	A <code>SingleCellExperiment</code> object
<code>name</code>	Name of phenotype

Details

Wrapper for colDat(object)[, name] which also accesses internal metadata (e.g., landmarks).

Value

A vector of any type

Author(s)

Daniel C. Ellwanger

.phenoNameExists	<i>Checks if phenotype exists</i>
------------------	-----------------------------------

Description

Checks if phenotype exists

Usage

```
.phenoNameExists(x, pheno_name)
```

Arguments

x	A SingleCellExperiment object
pheno_name	Name of phenotype

Value

logical value

Author(s)

Daniel C. Ellwanger

.plotDynamic	<i>DEF: Visualize expression dynamic</i>
--------------	--

Description

Shows feature expression as a function of pseudotime

Usage

```
.plotDynamic(x, Y, feature_name, trail_name, weights, k = 5)
```

Arguments

<code>x</code>	Pseudotime
<code>Y</code>	Expression matrix (samples in columns)
<code>weights</code>	Weight (states)
<code>k</code>	Dimensions of bases used to represent the smooth term

Author(s)

Daniel C. Ellwanger

<code>.plotManifold_def</code>	<i>DEF: Visualizing the manifold</i>
--------------------------------	--------------------------------------

Description

Plots approximation of sample embedding in latent space in two dimensions

Usage

```
.plotManifold_def(
  X,
  y,
  name,
  g = NULL,
  weights = NULL,
  axis_label = "",
  colors = c("pretty", "bw"),
  type = c("raw", "surface.fit", "surface.se"),
  samples_only = FALSE,
  setND = FALSE,
  alpha = 0.1
)
```

Arguments

<code>X</code>	Ordination of data points
<code>y</code>	Values to visualize
<code>name</code>	Label for the figure legend
<code>g</code>	The trajectory graph
<code>weights</code>	States along the trajectory
<code>axis_label</code>	Label prefix for each axis
<code>colors</code>	Either the color ramp is colorized or black/white
<code>type</code>	Type of visualization
<code>samples_only</code>	Does not show the whole surface, but colorizes only the samples
<code>setND</code>	Indicates if a '0' value should be set to 'ND' in the figure legend
<code>alpha</code>	Color param

Value

A ggplot object

Author(s)

Daniel C. Ellwanger

<code>.plotSpectrum_def</code>	<i>DEF: Visualizing the spectrum definition</i>
--------------------------------	---

Description

Plots Scree plot with eigengaps

Usage

```
.plotSpectrum_def(x)
```

Arguments

<code>x</code>	A list
----------------	--------

Value

A ggplot object

Author(s)

Daniel C. Ellwanger

<code>.plotStateExpression_def</code>	<i>DEF: Violine plots of genes</i>
---------------------------------------	------------------------------------

Description

Plot gene expression per state

Usage

```
.plotStateExpression_def(x, sts, label)
```

Arguments

<code>x</code>	A expression vector
<code>sts</code>	A states assignment vector
<code>label</code>	Label for legend

Value

A ggplot object

Author(s)

Daniel C. Ellwanger

`.plotStateSize_def` *DEF: Visualizing states sizes*

Description

Plots barplot of number of samples per state

Usage

```
.plotStateSize_def(x)
```

Arguments

x States

Value

A ggplot object

Author(s)

Daniel C. Ellwanger

`.plotStateTrajectory_def` *DEF: Visualizing the trajectory graph*

Description

Visualizes the trajectory spanning all states of a component

Usage

```
.plotStateTrajectory_def(  
  X,  
  g,  
  y,  
  all_sts,  
  name,  
  setND = FALSE,  
  point_size = 3,  
  label_offset = 2  
)
```

Arguments

<code>x</code>	Layout
<code>g</code>	State trajectory graph (igraph object)
<code>y</code>	Values; numeric or factor vector
<code>all_sts</code>	All state assignments
<code>setND</code>	If values of '0' are getting set to 'ND' in the figure legend
<code>point_size</code>	Point size parameter
<code>label_offset</code>	Label offset parameter
<code>label</code>	Label for plot legend

Value

A ggplot object

Author(s)

Daniel C. Ellwanger

.plotTrailblazing_def *DEF: Visualizing trailblazing*

Description

Shows trajectory graph and highlights landmarks

Usage

```
.plotTrailblazing_def(x, g, ltype, lid)
```

Arguments

<code>x</code>	The trajectory layout
<code>g</code>	The trajectory graph
<code>ltype</code>	Vector with landmark types
<code>lid</code>	Vector with landmark ids

Value

A ggplot object

Author(s)

Daniel C. Ellwanger

<code>.plotTrail_def</code>	<i>DEF: Visualize trail on map</i>
-----------------------------	------------------------------------

Description

Highlights a trail on map

Usage

```
.plotTrail_def(X, g, ptime, name)
```

Arguments

<code>X</code>	The trajectory layout
<code>g</code>	The trajectory graph
<code>ptime</code>	The trail pseudotime
<code>name</code>	Name of trail

Author(s)

Daniel C. Ellwanger

<code>.plot_trajectoryFit</code>	<i>DEF: Visualizing the trajectory fit</i>
----------------------------------	--

Description

Shows the residuals along the trajectory backbone

Usage

```
.plot_trajectoryFit(x, g, sts, factor = 7, rev = FALSE)
```

Arguments

<code>x</code>	The fitting residuals
<code>g</code>	The trajectory graph
<code>sts</code>	The states along the trajectory
<code>factor</code>	The jitter intensity correlates to the projection error
<code>rev</code>	Should the trajectory shown in reverse order?

Value

A ggplot object

Author(s)

Daniel C. Ellwanger

<code>.prettyColorRamp</code>	<i>Pretty color ramp</i>
-------------------------------	--------------------------

Description

Pretty color ramp

Usage

```
.prettyColorRamp(n, grayStart = TRUE)
```

Arguments

n	Number of colors
---	------------------

Value

Color codes

Author(s)

Daniel C. Ellwanger

<code>.prettyString</code>	<i>Pretty string from array</i>
----------------------------	---------------------------------

Description

Generates short representation of long character vectors

Usage

```
.prettyString(x, mmax = 4)
```

Arguments

x	A character vector
---	--------------------

Details

Returns pretty print of character vector

Value

String

Author(s)

Daniel C. Ellwanger

<code>.project_ortho</code>	<i>AUX: Orthogonal projection</i>
-----------------------------	-----------------------------------

Description

Orthogonally projects samples onto trajectory

Usage

```
.project_ortho(cl, g, X)
```

Arguments

<code>cl</code>	Trajectory states; factor vector
<code>g</code>	Trajectory tree; igraph object
<code>X</code>	Lower dimensional ordination of trajectory samples

Value

A list containing the following components:

<code>Y</code>	Projection
<code>X.c</code>	Mediancentres
<code>lambda</code>	Projection lambda (position: inside/outside of edge)
<code>adjmat</code>	Adjacency matrix with sample count per edge
<code>error</code>	Projection error per sample
<code>edgeId</code>	ID of each edge
<code>edgeId2Edge</code>	Mapping of edge to edgeId
<code>edge</code>	For each sample its assigned edge

Author(s)

Daniel C. Ellwanger

<code>.rbf</code>	<i>Radial basis function</i>
-------------------	------------------------------

Description

Computes radial basis function

Usage

```
.rbf(d, radius)
```

Arguments

x	A numeric value, vector or matrix
sigma	Radius of the kernel

Details

Also known as Heat kernel or Gaussian kernel

Value

Transformed values

Author(s)

Daniel C. Ellwanger

.rescale

Rescale vector

Description

Rescales vector to [ymin, ymax]

Usage

```
.rescale(x, ymin, ymax)
```

Arguments

x	X-values
ymin	New min
ymax	New max

Value

Rescaled vector

Author(s)

Daniel C. Ellwanger

.sampleNameExists	<i>Checks if sample exists</i>
-------------------	--------------------------------

Description

Checks if sample exists

Usage

```
.sampleNameExists(x, sample_name)
```

Arguments

x	A SingleCellExperiment object
sample_name	Name of sample

Value

logical value

Author(s)

Daniel C. Ellwanger

.spanForest	<i>GET state trajectory graph</i>
-------------	-----------------------------------

Description

Returns graph object spanning all states (spanning forest)

Usage

```
.spanForest(object)
```

Arguments

object	A SingleCellExperiment object
--------	-------------------------------

Value

A list object with an igraph object per component of the spanning forest

Author(s)

Daniel C. Ellwanger

```
.spanForest<-
```

SET state trajectory graph

Description

Sets graph object spanning all states (spanning forest) to SingleCellExperiment object

Usage

```
.spanForest(object) <- value
```

Arguments

object	A SingleCellExperiment object
value	A list object with an igraph object per component of the spanning forest

Value

An updated object of class SingleCellExperiment

Author(s)

Daniel C. Ellwanger

```
.spatmed
```

Spatial median

Description

Computes mediancentres

Usage

```
.spatmed(X, maxiter = 500, eps = 1e-06)
```

Arguments

x	A numeric matrix
---	------------------

Value

A numeric vector

Author(s)

Daniel C. Ellwanger

.stateTrajLayout	<i>GET state trajectory layout</i>
------------------	------------------------------------

Description

Gets layout of state trajectory from SingleCellExperiment object

Usage

```
.stateTrajLayout(object, component)
```

Arguments

object	A SingleCellExperiment object
--------	-------------------------------

Value

A numeric matrix

Author(s)

Daniel C. Ellwanger

.trailNameExists	<i>Checks if trail exists</i>
------------------	-------------------------------

Description

Checks if trail exists

Usage

```
.trailNameExists(x, trail_name)
```

Arguments

x	A SingleCellExperiment object
trail_name	Name of trail

Value

logical value

Author(s)

Daniel C. Ellwanger

<code>.trajGraph</code>	<i>GET trajectory graph</i>
-------------------------	-----------------------------

Description

Returns trajectory graph from a `SingleCellExperiment` object

Usage

```
.trajGraph(object)
```

Arguments

<code>object</code>	A <code>SingleCellExperiment</code> object
---------------------	--

Value

A `igraph` object

Author(s)

Daniel C. Ellwanger

<code>.trajGraph<-</code>	<i>SET trajectory graph</i>
------------------------------	-----------------------------

Description

Stores trajectory graph in a `SingleCellExperiment` object

Usage

```
.trajGraph(object) <- value
```

Arguments

<code>object</code>	A <code>SingleCellExperiment</code> object
<code>value</code>	A <code>igraph</code> object

Value

An updated object of class `SingleCellExperiment`

Author(s)

Daniel C. Ellwanger

.trajLandmark	<i>GET trajectory landmark annotation</i>
---------------	---

Description

Returns trajectory landmark information from SingleCellExperiment object

Usage

```
.trajLandmark(object, type = c("type", "id", "shape"))
```

Arguments

object	A SingleCellExperiment object
type	A character in "type", "id", "shape"

Value

A vector of any type

Author(s)

Daniel C. Ellwanger

.trajLandmark<-	<i>SET trajectory landmark annotation</i>
-----------------	---

Description

Stores information on trajectory landmarks in a SingleCellExperiment object

Usage

```
.trajLandmark(object, type = c("type", "id", "shape")) <- value
```

Arguments

object	A SingleCellExperiment object
type	A character in "type", "id", "shape"
value	A vector of any type

Value

An updated object of class SingleCellExperiment

Author(s)

Daniel C. Ellwanger

```
.trajResiduals<-      SET trajectory fitting residuals
```

Description

Stores trajectory fitting residuals in SingleCellExperiment object

Usage

```
.trajResiduals(object) <- value
```

Arguments

object	A SingleCellExperiment object
value	A numeric vector

Value

An updated object of class SingleCellExperiment

Author(s)

Daniel C. Ellwanger

```
.useFeature      GET trajectory features indicator
```

Description

Indicates if feature should be used for trajectory reconstruction. Spike-in controls are removed.

Usage

```
.useFeature(object)
```

Arguments

object	An object of class SingleCellExperiment
--------	---

Value

A logical vector

Author(s)

Daniel C. Ellwanger

<code>.useFeature<-</code>	<i>SET trajectory features indicator</i>
-------------------------------	--

Description

Sets indicator if feature should be used for trajectory reconstruction.

Usage

```
.useFeature(object) <- value
```

Arguments

<code>object</code>	An object of class <code>SingleCellExperiment</code>
<code>value</code>	A logical vector

Value

An updated object of class `SingleCellExperiment`

Author(s)

Daniel C. Ellwanger

<code>.useSample</code>	<i>GET trajectory samples indicator</i>
-------------------------	---

Description

Indicates if sample was used for trajectory reconstruction.

Usage

```
.useSample(object)
```

Arguments

<code>object</code>	An object of class <code>SingleCellExperiment</code>
---------------------	--

Value

A logical vector

Author(s)

Daniel C. Ellwanger

```
.useSample<-                    SET trajectory samples indicator
```

Description

Sets indicator if sample was used for trajectory reconstruction.

Usage

```
.useSample(object) <- value
```

Arguments

<code>object</code>	An object of class <code>SingleCellExperiment</code>
<code>value</code>	A logical vector

Value

An updated object of class `SingleCellExperiment`

Author(s)

Daniel C. Ellwanger

```
.validatePlotParams        Validates parameter settings for plot methods
```

Description

Validates parameter settings for plot methods

Usage

```
.validatePlotParams(x, color_by, name)
```

Arguments

<code>x</code>	A <code>SingleCellExperiment</code> object
<code>color_by</code>	Color_by parameter
<code>name</code>	Name parameter

Details

Parameter `color_by` needs to be in `'featureName'`, `'phenoName'`. Parameter `name` needs to be existent in `rownames(object)` or in `colnames(colData(object))`, `state`, `landmark`. If parameters were validated true, this function automatically extracts and returns the values. Character vectors get converted to factors.

Value

A list with the parameters and values

Author(s)

Daniel C. Ellwanger

<code>.write_ygraphml_def</code>	<i>DEF: Export trajectory graph</i>
----------------------------------	-------------------------------------

Description

For details see `write.ygraphml`

Usage

```
.write_ygraphml_def(g, X, file, col_values, lbl_values, shapes)
```

Arguments

<code>g</code>	Trajectory graph
<code>X</code>	Lower-dimensional ordination of trajectory samples
<code>col_values</code>	Color values
<code>lbl_values</code>	Label values
<code>shapes</code>	Trajectory landmark shapes
<code>tlayout</code>	Trajectory layout (if existent)

Author(s)

Daniel C. Ellwanger

<code>addTrail</code>	<i>ADD trail</i>
-----------------------	------------------

Description

Function to define a single trail on the trajectory.

Usage

```
addTrail(sce, from, to, name)
```

Arguments

<code>sce</code>	An object of class <code>SingleCellExperiment</code>
<code>from</code>	Start landmark
<code>to</code>	End landmark
<code>name</code>	Name of trail

Details

A trajectory can be composed of multiple single trails (e.g., developmental progression from a common start towards distinct terminal phenotypes). Start and endpoints of trails can be identified visually using the plot function `plotMap`. Here, start (=from) and end (=to) IDs of landmarks are starting with the character "B" (for branching points), "H" (for trail heads, i.e. terminal nodes), and "U" for user-defined landmarks.

Diagnostic messages

An error is thrown if the trajectory has not been fitted yet. Please, call `fitTrajectory` first. Further, an error is thrown if the provided start or end ID is unknown. A warning is shown if a trail with the same name already exists and gets re-defined.

Value

An updated object of class `SingleCellExperiment`

Author(s)

Daniel C. Ellwanger

See Also

`fitTrajectory` `landmarks` `plotMap`

Examples

```
# Example data
data(exSCE)

# Add trail
exSCE <- addTrail(exSCE, "H1", "H2", "Tr3")
trailNames(exSCE)
phenoNames(exSCE)
```

connectStates

Connect trajectory states

Description

Connects states using maximum interface scoring. For each state an interface score is defined by the relative distribution of states in its local `l`-neighborhood. A filter is applied to remove outliers (ie. false positive neighbors). States are spanned by maximizing the total interface score.

Usage

```
connectStates(sce, l = 10)
```

Arguments

<code>sce</code>	A <code>SingleCellExperiment</code> object
<code>l</code>	Neighborhood size (default: 10)

Details

CellTrails assumes that the arrangement of samples in the computed lower-dimensional latent space constitutes a trajectory. Therefore, CellTrails aims to place single samples along a maximum parsimony tree, which resembles a branching developmental continuum. Distances between samples in the latent space are computed using the Euclidean distance.

To avoid overfitting and to facilitate the accurate identification of bifurcations, CellTrails simplifies the problem. Analogous to the idea of a ‘broken-stick regression’, CellTrails groups the data and perform linear fits to separate trajectory segments, which are determined by the branching chronology of states. This leaves the optimization problem of finding the minimum number of associations between states while maximizing the total parsimony, which in theory can be solved by any minimum spanning tree algorithm. CellTrails adapts this concept by assuming that adjacent states should be located nearby and therefore share a relative high number of neighboring cells.

Each state defines a submatrix of samples that is composed of a distinct set of data vectors, i.e., each state is a distinct set of samples represented in the lower-dimensional space. For each state CellTrails identifies the l -nearest neighbors to each state’s data vector and takes note of their state memberships and distances. This results in two vectors of length l times the state size (i.e., a vector with memberships and a vector with distances).

CellTrails removes spurious neighbors (outliers), whose distance to a state is greater than or equal to

$$e^{\text{median}(\log(D)) + \text{MAD}(\log(D))}$$

where D is a matrix containing all collected l -nearest neighbor sample distances to any state in the latent space.

For each state CellTrails calculates the relative frequency on how often a state occurs in the neighborhood of a given state, which is referred to as the interface cardinality scores.

CellTrails implements a greedy algorithm to find the tree maximizing the total interface cardinality score, similar to a minimum spanning tree algorithm (Kruskal, 1956). In a nutshell, all interface cardinality scores are organized in a sorted linked list, and a graph with no edges, but k nodes (one for each state) is initialized. During each iteration the highest score is selected, removed from the list and its corresponding edge (connecting two states), if it is not introducing a cycle or is already existent, is added to the graph. The algorithm terminates if the size of the graph is $k-1$ (with k equals number of states) or the list is empty. A cycle is determined if nodes were revisited while traversing the graph using depth-first search. Its construction has a relaxed requirement (number of edges < number of nodes) compared to a tree (number of edges = number of nodes - 1), which may result in a graph (forest) having multiple tree components, i.e. several trajectories or isolated nodes.

Diagnostic messages

An error is thrown if the states have not been defined yet; function `findStates` needs to be called first.

Value

An updated `SingleCellExperiment` object

Author(s)

Daniel C. Ellwanger

References

Kruskal, J.B. (1956). On the shortest spanning subtree of a graph and the traveling salesman problem. Proc Amer Math Soc 7, 48-50.

See Also

findStates states

Examples

```
# Example data
data(exSCE)

# Connect states
exSCE <- connectStates(exSCE, l=30)
```

contrastTrailExpr	<i>Differential trail expression analysis</i>
-------------------	---

Description

Comparison of feature expression dynamic between two trails.

Usage

```
contrastTrailExpr(
  sce,
  feature_names = featureNames(sce),
  trail_names,
  score = "rmsd"
)
```

Arguments

sce	A SingleCellExperiment object
feature_names	Name of feature; can be multiple names
trail_names	Name of trails
score	Score type; one of {"rmsd", "tad", "abc", "cor"}

Details

Genes have non-uniform expression rates and each trail has a distinct set of upregulated genes, but also contains unequal numbers of cells. Because pseudotime is based on transcriptional change, its axis may be distorted, leading to stretched or compressed sections of longitudinal expression data that make comparison of trails challenging. To align different trails, despite these differences, CellTrails employs a dynamic programming based algorithm that has long been known in speech recognition, called dynamic time warping (Sakoe and Chiba, 1978). RNA expression rates are modeled analogous to speaking rates (Aach and Church, 2001); the latter accounts for innate non-linear variation in the length of individual phonemes (i.e., states) resulting in stretching and shrinking of word (i.e., trail) segments. This allows the computation of inter-trail alignment warps of individual expression time series that are similar but locally out of phase.

Univariate pairwise alignments are computed resulting in one warp per feature and per trail set. Similar to a (global) pairwise protein sequence alignment, monotonicity (i.e., no time loops) and continuity (i.e., no time leaps) constraints have to be imposed on the warping function to preserve temporal sequence ordering. To find the optimal warp, a recursion rule is applied which selects the local minimum of three moves through a dynamic programming matrix: suppose that query snapshot g and reference snapshot h have already been aligned, then the alignment of $h+1$ with $g+1$ is a (unit slope) diagonal move, h with $g+1$ denotes an expansion by repetition of h , and $h+2$ with $g+1$ contracts the query by dropping $h+1$.

The overall dissimilarity between two aligned expression time series x and y of length n is estimated by either the root-mean-square deviation $RMSD(x, y) = \sqrt{(\sum (x - y)^2 / n)}$, the total absolute deviation $TAD(x, y) = \sum (|x - y|)$, the area between the aligned dynamic curves (ABC), or Pearson's correlation coefficient (cor) over all aligned elements.

Value

Numeric value

Author(s)

Daniel C. Ellwanger

References

- Sakoe, H., and Chiba, S. (1978). Dynamic programming algorithm optimization for spoken word recognition. *IEEE Transactions on Acoustics, Speech, and Signaling Processing* 26, 43-49.
- Aach, J., and Church, G.M. (2001). Aligning gene expression time series with time warping algorithms. *Bioinformatics* 17, 495-508.

See Also

dtw

Examples

```
# Example data
data(exSCE)

# Differential expression between trails
contrastTrailExpr(exSCE, feature_name=c("feature_1", "feature_10"),
                  trail_names=c("Tr1", "Tr2"), score="rmsd")
```

embedSamples

Spectral embedding of biological samples

Description

Non-linear learning of a data representation that captures the intrinsic geometry of the trajectory. This function performs spectral decomposition of a graph encoding conditional entropy-based sample-to-sample similarities.

Usage

```
embedSamples(x, design = NULL)

## S4 method for signature 'matrix'
embedSamples(x, design = NULL)
```

Arguments

<code>x</code>	A <code>SingleCellExperiment</code> object or a numeric matrix with samples in columns and features in rows
<code>design</code>	A numeric matrix describing the factors that should be blocked

Details

Single-cell gene expression measurements comprise high-dimensional data of large volume, i.e. many features (e.g., genes) are measured in many samples (e.g., cells); or more formally, m samples can be described by the expression of n features (i.e., n dimensions). The cells' expression profiles are shaped by many distinct unobserved biological causes related to each cell's geno- and phenotype, such as developmental age, tissue region of origin, cell cycle stage, as well as extrinsic sources such as status of signaling receptors, and environmental stressors, but also technical noise. In other words, a single dimension, despite just containing gene expression information, represents an underlying combination of multiple dependent and independent, relevant and non-relevant factors, whereat each factors' individual contribution is non-uniform. To obtain a better resolution and to extract underlying information, CellTrails aims to find a meaningful low-dimensional structure - a manifold - that represents cells mainly by their temporal relation along a biological process.

This method assumes that the expression vectors are lying on or near a manifold with dimensionality d that is embedded in the n -dimensional space. By using spectral embedding CellTrails aims to amplify latent temporal information; it reduces noise (ie. truncates non-relevant dimensions) by transforming the expression matrix into a new dataset while retaining the geometry of the original dataset as much as possible. CellTrails captures overall cell-to-cell relations based on the statistical mutual dependency between any two data vectors. A high dependency between two samples should be represented by their close proximity in the lower-dimensional space.

First, the mutual dependency between samples is scored using mutual information. This entropy framework naturally requires discretization of data vectors by an indicator function, which assigns each continuous data point (expression value) to exactly one discrete interval (e.g. low, mid or high). However, measurement points located close to the interval borders may get wrongly assigned due to noise-induced fluctuations. Therefore, CellTrails fuzzifies the indicator function by using a piecewise polynomial function, i.e. the domain of each sample expression vector is divided into contiguous intervals (based on Daub *et al.*, 2004). Second, the computed mutual information matrix, which is left-bounded and composed of bits, is scaled to a generalized correlation coefficient. Third, CellTrails constructs a simple complete graph with m nodes, one for each data vector (ie. sample), and weights each edge between two nodes by a heat kernel function applied on the generalized correlation coefficient. Finally, nonlinear spectral embedding (ie. spectral decomposition of the graph's adjacency matrix) is performed (Belkin & Niyogi, 2003; Sussman *et al.*, 2012) unfolding the manifold. Please note that this methods only uses the set of defined trajectory features in a `SingleCellExperiment` object; spike-in controls are ignored and are not listed as trajectory features.

To account for systematic bias in the expression data (e.g., cell cycle effects), a design matrix can be provided for the learning process. It should list the factors that should be blocked and their

values per sample. It is suggested to construct a design matrix with `model.matrix`.

Diagnostic messages

The method throws an error if expression matrix contains samples with zero entropy (e.g., the samples exclusively contain non-detects, that is all expression values are zero).

Value

A list containing the following components:

eigenvectors	Ordered components of latent space
eigenvalues	Information content of latent components

Author(s)

Daniel C. Ellwanger

References

Daub, C.O., Steuer, R., Selbig, J., and Kloska, S. (2004). Estimating mutual information using B-spline functions – an improved similarity measure for analysing gene expression data. *BMC Bioinformatics* 5, 118.

Belkin, M., and Niyogi, P. (2003). Laplacian eigenmaps for dimensionality reduction and data representation. *Neural computation* 15, 1373-1396.

Sussman, D.L., Tang, M., Fishkind, D.E., and Priebe, C.E. (2012). A Consistent Adjacency Spectral Embedding for Stochastic Blockmodel Graphs. *J Am Stat Assoc* 107, 1119-1128.

See Also

`SingleCellExperiment` `trajectoryFeatureNames` `model.matrix`

Examples

```
# Example data
data(exSCE)

# Embed samples
res <- embedSamples(exSCE)
```

enrichment.test

Enrichment test

Description

Statistical enrichment analysis using either a Hypergeometric or Fisher's test

Usage

```
enrichment.test(  
  sample_true,  
  sample_size,  
  pop_true,  
  pop_size,  
  method = c("fisher", "hyper")  
)
```

Arguments

sample_true	Number of hits in sample
sample_size	Size of sample
pop_true	Number of hits in population
pop_size	Size of population
method	Statistical method that should be used

Details

Hypergeometric or one-tailed Fisher exact test is useful for enrichment analyses. For example, one needs to estimate which features are enriched among a set of instances sampled from a population.

Value

A list containing the following components:

p.value	P-value of the test
odds.ratio	Odds ratio
conf.int	Confidence interval for the odds ratio (only shown with method="fisher")
method	Used statistical test

Author(s)

Daniel C. Ellwanger

See Also

Hypergeometric and fisher.test

Examples

```
# Population has 13 of total 52 instances positive for a given feature  
# Sample has 1 of total 5 instances positive for a given feature  
# Test for significance of enrichment in sample  
enrichment.test(sample_true=1, sample_size=5,  
                 pop_true=13, pop_size=52, method="fisher")
```

exSCE

*Example single-cell expression data***Description**

This dataset contains simulated transcript expression profiles of 25 genes in expressed 100 cells. Simulation was performed using using the Negative Binomial Distribution. Distribution parameters for each feature were sampled from a Gamma distribution. The resulting expression matrix is log2-scaled and was stored in in an object of class 'SingleCellExperiment' (assay logcounts). The sample metainformation contains the underlying (discrete) simulated age of the cells.

Usage

```
data(exSCE)
```

Format

An object of class SingleCellExperiment

```
featureNames, SingleCellExperiment-method
```

GET feature names

Description

Retrieve feature names from a SingleCellExperiment object

Usage

```
## S4 method for signature 'SingleCellExperiment'
featureNames(object)
```

Arguments

object An object of class SingleCellExperiment

Details

Wrapper for rownames(object)

Value

A character vector

Author(s)

Daniel C. Ellwanger

See Also

SingleCellExperiment

Examples

```
# Example data
data(exSCE)

featureNames(exSCE)
```

```
filterTrajFeaturesByCOV
```

Filter features by Coefficient of Variation (COV)

Description

Filters trajectory features by their coefficient of variation.

Usage

```
filterTrajFeaturesByCOV(sce, threshold, design = NULL, show_plot = TRUE)
```

Arguments

sce	An SingleCellExperiment object
threshold	Minimum coefficient of variation; numeric value between 0 and 1
design	A numeric matrix describing the factors that should be blocked
show_plot	Indicates if plot should be shown (default: TRUE)

Details

For each trajectory feature x listed in the SingleCellExperiment object the coefficient of variation is computed by $CoV(x) = sd(x)/mean(x)$. Features with a $CoV(x)$ greater than threshold remain labeled as trajectory feature in the SingleCellExperiment object, otherwise they are not considered for dimensionality reduction, clustering and trajectory reconstruction. Please note that spike-in controls are ignored and are not listed as trajectory features.

To account for systematic bias in the expression data (e.g., cell cycle effects), a design matrix can be provided for the learning process. It should list the factors that should be blocked and their values per sample. It is suggested to construct a design matrix with `model.matrix`.

Value

A character vector

Author(s)

Daniel C. Ellwanger

See Also

trajFeatureNames `model.matrix`

Examples

```
# Simulate example data
set.seed(1101)
dat <- simulate_exprs(n_features=15000, n_samples=100)

# Create container
alist <- list(logcounts=dat)
sce <- SingleCellExperiment(assays=alist)

# Filter incrementally
trajFeatureNames(sce) <- filterTrajFeaturesByDL(sce, threshold=2)
trajFeatureNames(sce) <- filterTrajFeaturesByCOV(sce, threshold=0.5)

# Number of features
length(trajFeatureNames(sce)) #filtered
nrow(sce) #total
```

filterTrajFeaturesByDL

Filter trajectory features by Detection Level (DL)

Description

Filters trajectory features that are detected in a minimum number of samples.

Usage

```
filterTrajFeaturesByDL(sce, threshold, show_plot = TRUE)
```

Arguments

sce	An SingleCellExperiment object
threshold	Minimum number of samples; if value < 1 it is interpreted as fraction, otherwise as absolute sample count
show_plot	Indicates if plot should be shown (default: TRUE)

Details

The detection level denotes the fraction of samples in which a feature was detected. For each trajectory feature listed in the CellTrajectorySet object the relative number of samples having a feature expression value greater than 0 is counted. Features that are expressed in a fraction of all samples greater than threshold remain labeled as trajectory feature as listed in the SingleCellExperiment object, otherwise they may be not considered for dimensionality reduction, clustering, and trajectory reconstruction. If the parameter threshold fullfills threshold ≥ 1 it becomes converted to a relative fraction of the total sample count. Please note that spike-in controls are ignored and are not listed as trajectory features.

Value

A character vector

Author(s)

Daniel C. Ellwanger

See Also

trajFeatureNames

Examples

```
# Example data
set.seed(1101)
dat <- simulate_exprs(n_features=15000, n_samples=100)

# Create container
alist <- list(logcounts=dat)
sce <- SingleCellExperiment(assays=alist)

# Filter features
tfeat <- filterTrajFeaturesByDL(sce, threshold=2)
head(tfeat)

# Set trajectory features to object
trajFeatureNames(sce) <- tfeat

# Number of features
length(trajFeatureNames(sce)) #filtered
nrow(sce) #total
```

filterTrajFeaturesByFF

Filter features by Fano Factor

Description

Filters trajectory features that exhibit a significantly high fano factor (index of dispersion) by considering average expression levels.

Usage

```
filterTrajFeaturesByFF(
  sce,
  threshold = 1.7,
  min_expr = 0,
  design = NULL,
  show_plot = TRUE
)
```

Arguments

sce	An SingleCellExperiment object
threshold	A Z-score cutoff (default: 1.7)
min_expr	Minimum average expression of feature to be considered

design	A numeric matrix describing the factors that should be blocked
show_plot	Indicates if plot should be shown (default: TRUE)

Details

To identify the most variable features an unsupervised strategy that controls for the relationship between a features's average expression intensity and its expression variability is applied. Features are placed into 20 bins based on their mean expression. For each bin the fano factor (a windowed version of the index of dispersion, $IOD = \text{variance} / \text{mean}$) distribution is computed and standardized ($Z\text{-score}(x) = x/\text{sd}(x) - \text{mean}(x)/\text{sd}(x)$). Features with a Z-score greater than threshold remain labeled as trajectory feature in the `SingleCellExperiment` object. The parameter `min_expr` defines the minimum average expression level of a feature to be considered for this filter method. Please note that spike-in controls are ignored and are not listed as trajectory features.

To account for systematic bias in the expression data (e.g., cell cycle effects), a design matrix can be provided for the learning process. It should list the factors that should be blocked and their values per sample. It is suggested to construct a design matrix with `model.matrix`.

Value

A character vector

Author(s)

Daniel C. Ellwanger

See Also

`trajFeatureNames` `model.matrix`

Examples

```
# Simulate example data
set.seed(1101)
dat <- simulate_exprs(n_features=15000, n_samples=100)

# Create container
alist <- list(logcounts=dat)
sce <- SingleCellExperiment(assays=alist)

# Filter incrementally
trajFeatureNames(sce) <- filterTrajFeaturesByDL(sce, threshold=2)
trajFeatureNames(sce) <- filterTrajFeaturesByCOV(sce, threshold=0.5)
trajFeatureNames(sce) <- filterTrajFeaturesByFF(sce, threshold=1.7)

# Number of features
length(trajFeatureNames(sce)) #filtered
nrow(sce) #total
```

findSpectrum	<i>Determine number of informative latent dimensions</i>
--------------	--

Description

Identifies the dimensionality of the latent space

Usage

```
findSpectrum(x, frac = 100)
```

Arguments

x	A numeric vector with eigenvalues
frac	Fraction or number (if frac > 1) of eigengaps used to perform linear fit. (default: 100)

Details

Similar to a scree plot, this method generates a simple line segment plot showing the lagged differences between ordered eigenvalues (eigengaps). A linear fit is calculated on a fraction of top ranked values to identify informative eigenvectors.

Value

A numeric vector with indices of relevant dimensions

Author(s)

Daniel C. Ellwanger

See Also

pca embedSamples

Examples

```
# Example data
data(exSCE)

# Embedding
res <- embedSamples(exSCE)

# Find spectrum
d <- findSpectrum(res$eigenvalues, frac=30)
d
```

findStates	<i>Identify trajectory states</i>
------------	-----------------------------------

Description

Determines states using hierarchical spectral clustering with a *post-hoc* test.

Usage

```
findStates(sce, min_size = 0.01, min_feat = 5, max_pval = 1e-04, min_fc = 2)
```

Arguments

sce	A SingleCellExperiment object
min_size	The initial cluster dendrogram is cut at an height such that the minimum cluster size is at least min_size; if min_size < 1 than the fraction of total samples is used, otherwise it is used as absolute count (default: 0.01).
min_feat	Minimum number of differentially expressed features between siblings. If this number is not reached, two neighboring clusters (siblings) in the pruned dendrogram get joined. (default: 5)
max_pval	Maximum <i>P</i> -value for differential expression computation. (default: 1e-4)
min_fc	Minimum fold-change for differential expression computation (default: 2)

Details

To identify cellular subpopulations, CellTrails performs hierarchical clustering via minimization of a square error criterion (Ward, 1963) in the lower-dimensional space. To determine the cardinality of the clustering, CellTrails conducts an unsupervised *post-hoc* analysis. Here, it is assumed that differential expression of assayed features determines distinct cellular stages. First, Celltrails identifies the maximal fragmentation of the data space, i.e. the lowest cutting height in the clustering dendrogram that ensured that the resulting clusters contained at least a certain fraction of samples. Then, processing from this height towards the root, CellTrails iteratively joins siblings if they did not have at least a certain number of differentially expressed features. Statistical significance is tested by means of a two-sample non-parametric linear rank test accounting for censored values (Peto & Peto, 1972). The null hypothesis is rejected using the Benjamini-Hochberg (Benjamini & Hochberg, 1995) procedure for a given significance level.

Since this methods performs pairwise comparisons, the fold change threshold value is valid in both directions: higher and lower expressed than min_fc. Thus, input values < 0 are interpreted as a fold-change of 0. For example, min_fc=2 checks for features that are 2-fold differentially expressed in two given states (e.g., S1, S2). Thus, a feature can be either 2-fold higher expressed in state S1 or two-fold lower expressed in state S2 to be validated as differentially expressed.

Please note that this methods only uses the set of defined trajectory features in a SingleCellExperiment object; spike-in controls are ignored and are not listed as trajectory features.

Diagnostic messages

An error is thrown if the samples stored in the SingleCellExperiment object were not embedded yet (ie. the SingleCellExperiment object does not contain a latent space matrix object; latentSpace(object)is NULL).

Value

A factor vector

Author(s)

Daniel C. Ellwanger

References

Ward, J.H. (1963). Hierarchical Grouping to Optimize an Objective Function. *Journal of the American Statistical Association*, 58, 236-244.

Peto, R., and Peto, J. (1972). Asymptotically Efficient Rank Invariant Test Procedures (with Discussion). *Journal of the Royal Statistical Society of London, Series A* 135, 185–206.

Benjamini, Y., and Hochberg, Y. (1995). Controlling the false discovery rate: a practical and powerful approach to multiple testing. *Journal of the Royal Statistical Society Series B* 57, 289–300.

See Also

latentSpace trajectoryFeatureNames

Examples

```
# Example data
data(exSCE)

# Find states
cl <- findStates(exSCE, min_feat=2)
head(cl)
```

fitDynamic

Fit expression dynamic

Description

Fits feature expression as a function of pseudotime along a defined trail.

Usage

```
fitDynamic(sce, feature_name, trail_name)
```

Arguments

sce	A SingleCellExperiment object
feature_name	Name of feature
trail_name	Name of trail

Details

A trail is an induced subgraph of the trajectory graph. A trajectory graph is composed of samples (nodes) that are connected (by weighted edges) if they are chronologically related. A trail has to be defined by the user using `addTrail`. A pseudotime vector is extracted by computing the geodesic distance for each sample from the trail's start node. To infer the expression level of a feature as a function of pseudotime, CellTrails used generalized additive models with a single smoothing term with four basis dimensions. Here, for each feature CellTrails introduces prior weights for each observation to lower the confounding effect of drop-outs to the maximum-likelihood-based fitting process as follows. Each non-detect of feature j in state h is weighted by the relative fraction of non-detects of feature j in state h ; detected values are always assigned weight = 1.

Value

An object of type `list` with the following components

`pseudotime` The pseudotime along the trail

`expression` The fitted expression values for each value of pseudotime

`gam` A object of class `gamObject`

Author(s)

Daniel C. Ellwanger

See Also

`addTrail gamObject`

Examples

```
# Example data
data(exSCE)

# Fit dynamic
fit <- fitDynamic(exSCE, feature_name="feature_3", trail_name="Tr1")

summary(fit)
```

<code>fitTrajectory</code>	<i>Align samples to trajectory</i>
----------------------------	------------------------------------

Description

Orthogonal projection of each sample to the trajectory backbone.

Usage

```
fitTrajectory(sce)
```

Arguments

`sce` A `SingleCellExperiment` object

Details

The previously selected component (with k states) defines the trajectory backbone. With this function CellTrails embeds the trajectory structure in the latent space by computing $k-1$ straight lines passing through k mediancentres (Bedall & Zimmermann, 1979) of adjacent states. Then, a fitting function is learned. Each sample is projected to its most proximal straight line passing through the mediancentre of its assigned state. Here, whenever possible, projections on line segments *between* two mediancentres are preferred. Residuals (fitting deviations) are given by the Euclidean distance between the sample's location and the straight line. Finally, a weighted acyclic trajectory graph can be constructed based on each sample's position along its straight line. In addition, data vectors are connected to mediancentres to enable the proper determination of branching points. Each edge is weighted by the distance between each node (sample) after orthogonal projection.

Of note, the fitting function implies potential side branches in the trajectory graph; those could be caused due to technical variance or encompass samples that were statistically indistinguishable from the main trajectory given the selected genes used for trajectory reconstruction.

Diagnostic messages

An error is thrown if an trajectory graph component was not computed or selected yet; functions `connectStates` and `selectTrajectory` need to be run first.

Value

An updated `SingleCellExperiment` object

Author(s)

Daniel C. Ellwanger

References

Bedall, F.K., and Zimmermann, H. (1979). Algorithm AS143. The mediancentre. *Appl Statist* 28, 325-328.

See Also

`connectStates` `selectTrajectory`

Examples

```
# Example data
data(exSCE)

# Align samples to trajectory
exSCE <- fitTrajectory(exSCE)
```

landmarks	<i>GET landmarks</i>
-----------	----------------------

Description

Gets landmarks from a SingleCellExperiment object.

Usage

```
landmarks(object)
```

Arguments

object	A SingleCellExperiment object
--------	-------------------------------

Details

Trail branches (B) and heads (H) are automatically assigned; landmarks can also be defined on the trajectory by the user (U). Landmarks can be used to extract single trails from a trajectory.

Value

A character vector with sample names

Author(s)

Daniel C. Ellwanger

See Also

userLandmarks

Examples

```
# Example data
data(exSCE)

# Get landmarks
landmarks(exSCE)[seq_len(5)]
```

latentSpace	<i>GET CellTrails' latent space</i>
-------------	-------------------------------------

Description

Retrieve computed latent space from a SingleCellExperiment object.

Usage

```
latentSpace(object)
```

Arguments

object	A SingleCellExperiment object
--------	-------------------------------

Details

Returns the latent space set for a CellTrails analysis. The resulting matrix is numeric. Rows are samples and columns are d components. It is a wrapper for `reducedDim` to ensure that the proper matrix is received from a SingleCellExperiment object.

Value

An object of class `matrix`

Author(s)

Daniel C. Ellwanger

See Also

`SingleCellExperiment` `reducedDim`

Examples

```
# Example data
data(exSCE)

# Get latent space
latentSpace(exSCE)[seq_len(5), ]
```

latentSpace<-	<i>SET latent space</i>
---------------	-------------------------

Description

Set CellTrails' latent space to a SingleCellExperiment object.

Usage

```
latentSpace(object) <- value
```

Arguments

object	A SingleCellExperiment object
value	A numeric matrix with rows are samples and columns are components

Details

Rows need to be samples and columns to be d components (spanning the lower-dimensional latent space).

Value

An updated object of class SingleCellExperiment

Author(s)

Daniel C. Ellwanger

See Also

SingleCellExperiment reducedDim

Examples

```
# Example data
data(exSCE)

# Set latent space
latentSpace(exSCE) <- pca(exSCE)$components[, seq_len(10)]
```

manifold2D	<i>GET 2D manifold representation</i>
------------	---------------------------------------

Description

Returns 2D manifold representation of latent space from SingleCellExperiment object

Usage

```
manifold2D(object)
```

Arguments

object A SingleCellExperiment object

Value

A numeric matrix

Author(s)

Daniel C. Ellwanger

Examples

```
# Example data
data(exSCE)

manifold2D(exSCE)[seq_len(5), ]
```

manifold2D<-	<i>SET 2D manifold representation</i>
--------------	---------------------------------------

Description

Stores 2D manifold representation in SingleCellExperiment object

Usage

```
manifold2D(object) <- value
```

Arguments

object A SingleCellExperiment object
value A numeric matrix with one column per dimension

Value

An updated object of class SingleCellExperiment

Author(s)

Daniel C. Ellwanger

Examples

```
# Example data
data(exSCE)

gp <- plotManifold(exSCE, color_by="featureName", name="feature_10",
                   recalculate=TRUE)
manifold2D(exSCE) <- gp
```

pca

*Principal Component Analysis***Description**

Performs principal component analysis by spectral decomposition of a covariance or correlation matrix

Usage

```
pca(sce, do_scaling = TRUE, design = NULL)
```

Arguments

sce	SingleCellExperiment object
do_scaling	FALSE = covariance matrix, TRUE = correlation matrix
design	A numeric matrix describing the factors that should be blocked

Details

The calculation is done by a spectral decomposition of the (scaled) covariance matrix of the trajectory features as defined in the SingleCellExperiment object. Features with zero variance get automatically removed. Please note that this methods only uses the set of defined trajectory features in a SingleCellExperiment object; spike-in controls are ignored and are not listed as trajectory features.

To account for systematic bias in the expression data (e.g., cell cycle effects), a design matrix can be provided for the learning process. It should list the factors that should be blocked and their values per sample. It is suggested to construct a design matrix with `model.matrix`.

Value

A list object containing the following components:

components	Principal components
eigenvalues	Variance per component
variance	Fraction of variance explained by each component
loadings	Loading score for each feature

Author(s)

Daniel C. Ellwanger

See Also

SingleCellExperiment model.matrix

Examples

```
# Example data
data(exSCE)

# Principal component analysis
res <- pca(exSCE)

# Find relevant number of principal components
d <- findSpectrum(res$eigenvalues, frac=20)

barplot(res$variance[d] * 100, ylab="Variance (%)",
        names.arg=colnames(res$components)[d], las=2)
plot(res$component, xlab="PC1", ylab="PC2")
```

phenoNames

GET phenotype names

Description

Retrieve phenotype names from a SingleCellExperiment object

Usage

```
phenoNames(object)
```

Arguments

object An object of class SingleCellExperiment

Details

Wrapper for colnames(colData(object))

Value

A character vector

Author(s)

Daniel C. Ellwanger

See Also

SingleCellExperiment

Examples

```
# Example data
data(exSCE)

phenoNames(exSCE)
```

plotDynamic

Visualize dynamics

Description

Shows dynamics of one or multiple features along a given trail

Usage

```
plotDynamic(sce, feature_name, trail_name)
```

Arguments

sce	A SingleCellExperiment object
feature_name	Name of one or multiple features
trail_name	Name of trail

Details

An error is thrown if the trail_name or feature_name are unknown. The function is case-sensitive. All available trails can be listed by trailNames, all features with featureNames.

Value

A ggplot object

Author(s)

Daniel C. Ellwanger

See Also

addTrail trailNames featureNames

Examples

```
# Example data
data(exSCE)

# Plot dynamic of feature_10
plotDynamic(exSCE, trail_name="Tr1", feature_name="feature_1")
# Plot dynamic of feature_1 and feature_10
plotDynamic(exSCE, trail_name="Tr1",
            feature_name=c("feature_1", "feature_10"))
```

plotManifold	<i>Visualize the learned manifold</i>
--------------	---------------------------------------

Description

Method visualizes an approximation of the manifold in the latent space in two dimensions.

Usage

```
plotManifold(
  sce,
  color_by = c("phenoName", "featureName"),
  name,
  perplexity = 30,
  recalculate = FALSE
)
```

Arguments

sce	A SingleCellExperiment object
color_by	Indicates if nodes are colorized by a feature expression ('featureName') or phenotype label ('phenoName')
name	A character string specifying the featureName or phenoName
perplexity	Perplexity parameter for tSNE computation (default: 30)
recalculate	Indicates if tSNE should be recalculated and results returned (default: FALSE)

Details

Visualizes the learned lower-dimensional manifold in two dimensions using an approximation obtained by Barnes-Hut implementation of t-Distributed Stochastic Neighbor Embedding (tSNE; van der Maaten and Hinton 2008). Each point in this plot represents a sample. Points can be colorized according to feature expression or experimental metadata. The points' coloration can be defined via the attributes color_by and name, respectively. A previously computed tSNE visualization will be reused if it was set accordingly (see manifold2D<-). The parameter perplexity is used for the tSNE calculation.

Value

A ggplot object

Author(s)

Daniel C. Ellwanger

References

van der Maaten, L.J.P. & Hinton, G.E., 2008. Visualizing High-Dimensional Data Using t-SNE. Journal of Machine Learning Research, 9, pp.2579-2605.

See Also

Rtsne latentSpace manifold2D

Examples

```
# Example data
data(exSCE)

plotManifold(exSCE, color_by="featureName", name="feature_10")
gp <- plotManifold(exSCE, color_by="phenoName", name="age",
                    recalculate=TRUE)
manifold2D(exSCE) <- gp
```

plotMap	<i>Visualize expression maps</i>
---------	----------------------------------

Description

Method visualizes topographical expression maps in two dimensions.

Usage

```
plotMap(
  sce,
  color_by = c("phenoName", "featureName"),
  name,
  type = c("surface.fit", "surface.se", "raw"),
  samples_only = FALSE
)
```

Arguments

sce	A SingleCellExperiment object
color_by	Indicates if nodes are colorized by a feature expression
name	A character string specifying the featureName or phenoName
type	Type of map; one of "raw", "surface.fit", "surface.se"
samples_only	If only individual samples should be colorized rather than the whole surface (default: FALSE)

Details

Two-dimensional visualization of the trajectory. The red line represents the trajectory and individual points denote samples. This plot type can either show the topography of a given feature's expression landscape or colorizes individual samples by a metadata label. The feature is selected by setting the parameter `color_type` and the respective name. To show feature expression, a surface is fitted using isotropic (i.e., same parameters for both map dimensions) thin-plate spline smoothing in `gam`. It gives an overview of expression dynamics along all branches of the trajectory. The parameter `type` defines if either the raw/original expression data should be shown, the full fitted expression surface should be shown (`type="surface.fit"`) or the standard error of the surface prediction (`type="surface.se"`), or the expression values of single samples only (`type="surface.fit"` and

only_samples=TRUE).

To show all landmarks on the map, please use the parameters color_by="phenoName" and name="landmark".

Value

A ggplot object

Author(s)

Daniel C. Ellwanger

See Also

gam

Examples

```
# Example data
data(exSCE)

# Plot landmarks
plotMap(exSCE, color_by="phenoName", name="landmark")

# Plot phenotype
plotMap(exSCE, color_by="phenoName", name="age")

# Plot feature expression map
plotMap(exSCE, color_by="featureName", name="feature_10", type="surface.fit")
plotMap(exSCE, color_by="featureName", name="feature_10", type="surface.fit",
        samples_only=TRUE)

#Plot surface fit standard errors
plotMap(exSCE, color_by="featureName", name="feature_10", type="surface.se")
```

plotStateExpression	<i>Visualize feature expression distribution per state</i>
---------------------	--

Description

Violin plots showing the expression distribution of a feature per state.

Usage

```
plotStateExpression(sce, feature_name)
```

Arguments

sce	A SingleCellExperiment object
feature_name	The name of the feature to be visualized

Details

Each data point displays the feature's expression value in a single sample. A violine plot shows the density (mirrored on the y-axis) of the expression distribution per sample.

Value

A ggplot object

Author(s)

Daniel C. Ellwanger

See Also

ggplot states

Examples

```
# Example data
data(exSCE)

plotStateExpression(exSCE, feature_name="feature_1")
```

plotStateSize

Visualize the number of samples per state

Description

Shows barplot of state size distribution

Usage

```
plotStateSize(sce)
```

Arguments

sce A SingleCellExperiment object

Details

Barplot showing the absolute number of samples per state.

Value

A ggplot object

Author(s)

Daniel C. Ellwanger

See Also

ggplot states

Examples

```
# Example data
data(exSCE)

plotStateSize(exSCE)
```

plotStateTrajectory	<i>Visualize state trajectory graph</i>
---------------------	---

Description

Method visualizes the state-to-state relations delineating the trajectory backbone.

Usage

```
plotStateTrajectory(
  sce,
  color_by = c("phenoName", "featureName"),
  name,
  component = NULL,
  point_size = 3,
  label_offset = 2,
  recalculate = FALSE
)
```

Arguments

sce	A SingleCellExperiment object
color_by	Indicates if nodes are colorized by a feature expression ('featureName') or phenotype label ('phenoName')
name	A character string specifying the featureName or phenoName
component	Component of trajectory graph that should be shown (integer value)
point_size	Adjusts the point size of the data points shown
label_offset	Adjusts the offset of the data point labels
recalculate	If layout should be re-drawn (default: FALSE)

Details

Shows a single tree component of the computed trajectory graph. Each point in this plot represents a state and can be colorized according to feature expression (mean expression per state) or experimental metadata (arithmetic mean or percentage distribution of categorical values). The component is defined by parameter component. If the trajectory graph contains only a single component, then this parameter can be left undefined. The points' coloration can be defined via the attributes color_by and name, respectively. Missing sample labels are recovered using nearest neighbor learning. If the state trajectory graph layout was set with stateTrajLayout<- then the layout will be reused for visualization.

Value

A ggplot object

Author(s)

Daniel C. Ellwanger

See Also

connectStates

Examples

```
# Example data
data(exSCE)

plotStateTrajectory(exSCE, color_by="phenoName", name="age",
                    component=1, point_size = 1.5, label_offset = 4)

gp <- plotStateTrajectory(exSCE, color_by="featureName", name="feature_1",
                        component=1, recalculate=TRUE)
stateTrajLayout(exSCE) <- gp
```

plotTrail

*Visualize single trails***Description**

Method highlights a single trail on the trajectory map

Usage

```
plotTrail(sce, name)
```

Arguments

sce	A SingleCellExperiment object
name	Name of the trail

Details

A trail can be defined with the function `addTrail` between two landmarks. User-defined landmarks can be set with the function `userLandmarks`. This function visualizes the start and endpoints, and the pseudotime of a defined trail along the trajectory. The trail pseudotimes can be directly accessed via the trails.

An error is thrown if the `trail_name` is unknown. The function is case-sensitive. All available trails can be listed by `trailNames`.

Value

A ggplot object

Author(s)

Daniel C. Ellwanger

See Also

addTrail userLandmarks trailNames trails

Examples

```
# Example data
data(exSCE)

# Plot trail
plotTrail(exSCE, name="Tr1")
```

plotTrajectoryFit	<i>Visualize trajectory fit residuals</i>
-------------------	---

Description

Method visualizes the fitting residuals along the trajectory backbone.

Usage

```
plotTrajectoryFit(sce)
```

Arguments

sce A SingleCellExperiment object

Details

Shows the trajectory backbone (longest shortest path between two samples) and the fitting deviations of each sample indicated by the perpendicular jitter. Data points are colorized by state.

Value

A ggplot object

Author(s)

Daniel C. Ellwanger

See Also

fitTrajectory trajResiduals

Examples

```
# Example data
data(exSCE)

plotTrajectoryFit(exSCE)
```

read.ygraphml	<i>Reads trajectory graph layout</i>
---------------	--------------------------------------

Description

Reads ygraphml file containing the trajectory graph's layout

Usage

```
read.ygraphml(file)
```

Arguments

file	A character string naming a file
------	----------------------------------

Details

To visualize the trajectory graph, a proper graph layout has to be computed. Ideally, edges should not cross and nodes should not overlap. CellTrails enables the export and import of the trajectory graph structure using the graphml file format. This file format can be interpreted by most third-party graph analysis applications, allowing the user to subject the trajectory graph to a wide range of layout algorithms. Please note that the graphml file needs to contain layout information ("`<y:Geometry x=... y=... >`" entries) as provided by the 'ygraphml' file definition used by the Graph Visualization Software 'yEd' (freely available from yWorks GmbH, <http://www.yworks.com/products/yed>).

Value

An data.frame with coordinates of data points and visualization metadata

Author(s)

Daniel C. Ellwanger

See Also

`write.ygraphml`

Examples

```
# Example data
data(exSCE)

## Not run:
fn <- system.file("exdata", "exDat.graphml", package="CellTrails")
tl <- read.ygraphml(fn)

## End(Not run)
```

removeTrail	<i>REMOVE trail</i>
-------------	---------------------

Description

Removes trail from a SingleCellExperiment object.

Usage

```
removeTrail(sce, name)
```

Arguments

sce	An object of class SingleCellExperiment
name	Name of trail

Details

Diagnostic messages

An error is thrown if the trail name is unknown. All stored trail names can be shown using function `trailNames`.

Value

An updated object of class SingleCellExperiment

Author(s)

Daniel C. Ellwanger

See Also

`trailNames` `addTrail`

Examples

```
# Example data
data(exSCE)

# Remove trail
trailNames(exSCE)
exSCE <- removeTrail(exSCE, "Tr1")
trailNames(exSCE)
```

sampleNames, SingleCellExperiment-method
GET sample names

Description

Retrieve sample names from a SingleCellExperiment object

Usage

```
## S4 method for signature 'SingleCellExperiment'  
sampleNames(object)
```

Arguments

object An object of class SingleCellExperiment

Details

Wrapper for colnames(object)

Value

A character vector

Author(s)

Daniel C. Ellwanger

See Also

SingleCellExperiment

Examples

```
# Example data  
data(exSCE)  
  
sampleNames(exSCE)[seq_len(5)]
```

selectTrajectory	<i>Select component from trajectory graph</i>
------------------	---

Description

Retains a single component of a trajectory graph.

Usage

```
selectTrajectory(sce, component)
```

Arguments

sce	A SingleCellExperiment object
component	Number of component to be selected

Details

The construction of a trajectory graph may result in a forest having multiple tree components, which may represent individual trajectories or isolated nodes. This method should be used to extract a single component from the graph. A component is identified by its (integer) number.

Diagnostic messages

An error is thrown if the states have not been connected yet; function `connectStates` needs to be called first. An error is thrown if an unknown component (number) is selected.

Value

An updated `SingleCellExperiment` object

Author(s)

Daniel C. Ellwanger

See Also

`connectStates`
`findStates` `states`

Examples

```
# Example data
data(exSCE)

# Select trajectory
exSCE <- selectTrajectory(exSCE, component=1)
```

showTrajInfo	<i>Shows relevant content of a SingleCellExperiment object for a Cell-Trails analysis</i>
--------------	---

Description

Shows relevant content of a SingleCellExperiment object for a CellTrails analysis

Usage

```
showTrajInfo(object)
```

Arguments

object	A SingleCellExperiment object
--------	-------------------------------

Value

showTrajInfo returns an invisible NULL

Author(s)

Daniel C. Ellwanger

Examples

```
# Example data
data(exSCE)

showTrajInfo(exSCE)
```

simulate_exprs	<i>Simulation of RNA-Seq expression data</i>
----------------	--

Description

Simple simulation of RNA-Seq expression data estimating counts based on the negative binomial distribution

Usage

```
simulate_exprs(n_features, n_samples, prefix_sample = "")
```

Arguments

n_features	Number of genes
n_samples	Number of samples
prefix_sample	Prefix of sample name

Details

RNA-Seq counts are generated using the Negative Binomial Distribution. Distribution parameters for each feature are sampled from a Gamma distribution. The resulting expression matrix is log2-scaled.

Value

A numeric matrix with genes in rows and samples in columns

Author(s)

Daniel C. Ellwanger

See Also

NegBinomial and GammaDist

Examples

```
# Matrix with 100 genes and 50 cells
dat <- simulate_exprs(n_features=100, n_samples=50)
```

states

GET states

Description

Retrieve computed states from a SingleCellExperiment object

Usage

```
states(object)
```

Arguments

object An object of class SingleCellExperiment

Details

State information is extracted from colData; factor levels are alphanumerically ordered by ID.

Value

A factor vector

Author(s)

Daniel C. Ellwanger

See Also

SingleCellExperiment findStates

```
states<-
```

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Examples

```
# Example data
data(exSCE)

states(exSCE)[seq_len(5)]
```

```
states<-
```

SET states

Description

Sets states to a SingleCellExperiment object

Usage

```
states(object) <- value
```

Arguments

object	An object of class SingleCellExperiment
value	A numeric, character or factor vector

Details

State information is added to a SingleCellExperiment object via colData. If the vector containing the cluster assignments is numeric, the prefix "S" is added and the vector is converted to type factor.

Value

An updated object of class SingleCellExperiment

Author(s)

Daniel C. Ellwanger

See Also

colData

Examples

```
# Example data
data(exSCE)

# Assign clusters
cl <- kmeans(logcounts(exSCE), centers=10)$cluster
states(exSCE) <- cl
```

```
stateTrajLayout<-      SET state trajectory layout
```

Description

Stores layout of state trajectory in SingleCellExperiment object

Usage

```
stateTrajLayout(object) <- value
```

Arguments

object A SingleCellExperiment object
value A ggplot object

Value

An updated object of class SingleCellExperiment

Author(s)

Daniel C. Ellwanger

Examples

```
# Example data
data(exSCE)

gp <- plotStateTrajectory(exSCE, color_by="featureName",
                           name="feature_10", recalculate=TRUE)
stateTrajLayout(exSCE) <- gp
```

```
trailNames      GET trail names
```

Description

Function to extract trail names from SingleCellExperiment object.

Usage

```
trailNames(object)
```

Arguments

object An object of class SingleCellExperiment

Value

A character vector

```
trailNames<-
```

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Author(s)

Daniel C. Ellwanger

See Also

addTrail

Examples

```
# Example data
data(exSCE)

trailNames(exSCE)
```

```
trailNames<-
```

SET trail names

Description

Enables to rename trails stored in a SingleCellExperiment object.

Usage

```
trailNames(object) <- value
```

Arguments

object	An object of class SingleCellExperiment
value	A character vector with the trail names

Details

Diagnostic messages

An error is thrown if the number of names does not correspond to the number of trails stored in the object. Further, trail names are required to be unique.

Value

An updated object of class SingleCellExperiment

Author(s)

Daniel C. Ellwanger

See Also

addTrail

Examples

```
# Example data
data(exSCE)

trailNames(exSCE)
trailNames(exSCE) <- c("ABC", "DEF")
trailNames(exSCE)
```

trails	<i>GET trails</i>
--------	-------------------

Description

Function to extract trail pseudotimes from a SingleCellExperiment object.

Usage

```
trails(object)
```

Arguments

object	An object of class SingleCellExperiment
--------	---

Value

A DataFrame with numeric columns

Author(s)

Daniel C. Ellwanger

See Also

addTrail

Examples

```
# Example data
data(exSCE)

trails(exSCE)
```

trajComponents	<i>GET trajectory component states</i>
----------------	--

Description

Returns states of trajectory components SingleCellExperiment object

Usage

```
trajComponents(object)
```

Arguments

object	A SingleCellExperiment object
--------	-------------------------------

Value

A character vector

Author(s)

Daniel C. Ellwanger

Examples

```
# Example data
data(exSCE)

trajComponents(exSCE)
```

trajFeatureNames	<i>GET trajectory feature names</i>
------------------	-------------------------------------

Description

Retrieve names of features that were selected for trajectory reconstruction from a SingleCellExperiment object.

Usage

```
trajFeatureNames(object)
```

Arguments

object	An object of class SingleCellExperiment
--------	---

Details

Features can be selected prior to trajectory inference. This method retrieves the user-defined features from a SingleCellExperiment object. The return value is a character vector containing the feature names.

Value

An object of class character

Author(s)

Daniel C. Ellwanger

Examples

```
# Example data
data(exSCE)

# Get trajectory features
trajFeatureNames(exSCE)[seq_len(5)]
```

```
trajFeatureNames<-      SET trajectory features by name
```

Description

Function to set trajectory features by name

Usage

```
trajFeatureNames(object) <- value
```

Arguments

object	An object of class SingleCellExperiment
value	A character vector

Value

An updated object of class SingleCellExperiment

Author(s)

Daniel C. Ellwanger

Examples

```
# Example data
data(exSCE)

# Set trajectory features
trajFeatureNames(exSCE) <- rownames(exSCE)[seq_len(5)]
```

trajLayout	<i>GET trajectory layout</i>
------------	------------------------------

Description

Returns trajectory layout from SingleCellExperiment object

Usage

```
trajLayout(object)
```

Arguments

object	A SingleCellExperiment object
--------	-------------------------------

Value

A data.frame

Author(s)

Daniel C. Ellwanger

Examples

```
# Example data
data(exSCE)

trajLayout(exSCE)[seq_len(5), ]
```

trajLayout<-	<i>SET trajectory layout</i>
--------------	------------------------------

Description

Sets layout used for trajectory visualization to a SingleCellExperiment object.

Usage

```
trajLayout(object, adjust) <- value
```

Arguments

object	An object of class SingleCellExperiment
adjust	Indicates if layout has to be adjusted such that edge lengths correlate to pseudo-time (default: TRUE)
value	A data.frame with x- and y-coordinates for each sample (rows = samples, columns = coordinates)

Details

CellTrails implements a module which can incorporate pseudotime information into the the graph layout (activated via parameter `adjust`). Here, edge lengths between two nodes (samples) will then correspond to the inferred pseudotime that separates two samples along the trajectory.

Diagnostic messages

An error is thrown if the number of rows of the layout does not correspond to the number of trajectory samples or if the number of columns is less than 2, or if the row names do not correspond to `sampleNames`.

Value

An updated object of class `SingleCellExperiment`

Author(s)

Daniel C. Ellwanger

See Also

`write.ygraphml trajSampleNames`

Examples

```
# Example data
data(exSCE)
tl <- trajLayout(exSCE)

trajLayout(exSCE) <- tl
```

`trajResiduals`*GET trajectory fitting residuals*

Description

Returns trajectory fitting residuals from `SingleCellExperiment` object

Usage

```
trajResiduals(object)
```

Arguments

`object` A `SingleCellExperiment` object

Details

The trajectory fitting deviation is defined as the vector rejection from a sample in the latent space to the trajectory backbone. The trajectory backbone is defined by a tree spanning all relevant states. Samples get orthogonally projected onto straight lines connecting related states. This function quantifies the distance between the actual position of a sample in the latent space and its projected position on the trajectory backbone. In other words, the higher the distance, the higher its deviation (residual) from the trajectory fit. This function returns all residuals for each projected sample. Residuals of samples which were excluded for trajectory reconstruction are NA.

Value

A numeric vector

Author(s)

Daniel C. Ellwanger

See Also

fitTrajectory trajSampleNames

Examples

```
# Example data
data(exSCE)

trajResiduals(exSCE)[seq_len(5)]
```

trajSampleNames	<i>GET trajectory sample names</i>
-----------------	------------------------------------

Description

Retrieve names of samples that were aligned onto the trajectory from a SingleCellExperiment object.

Usage

```
trajSampleNames(object)
```

Arguments

object An object of class SingleCellExperiment

Details

A trajectory graph can be initially a forest. Trajectory fitting is performed on one component. This function returns the names of the samples which are member of the selected component.

Value

An object of class character

Author(s)

Daniel C. Ellwanger

Examples

```
# Example data
data(exSCE)

# Get trajectory samples
trajSampleNames(exSCE)[seq_len(5)]
```

userLandmarks	<i>GET user-defined landmarks</i>
---------------	-----------------------------------

Description

Gets user-defined landmarks from a SingleCellExperiment object.

Usage

```
userLandmarks(object)
```

Arguments

object	A SingleCellExperiment object
--------	-------------------------------

Details

Landmarks can be defined on the trajectory by the user with userLandmarks. Landmarks can be used to extract single trails from a trajectory.

Value

A character vector with sample names

Author(s)

Daniel C. Ellwanger

See Also

SingleCellExperiment

Examples

```
# Example data
data(exSCE)

# Get landmarks
userLandmarks(exSCE)
```

userLandmarks<-	<i>SET user-defined landmarks</i>
-----------------	-----------------------------------

Description

Set user-defined landmarks to a SingleCellExperiment object.

Usage

```
userLandmarks(object) <- value
```

Arguments

object	A SingleCellExperiment object
value	A character vector with sample names

Details

Landmarks can be defined on the trajectory and can be necessary to extract individual trails from a trajectory.

Diagnostic messages

An error is thrown if the trajectory has not been reconstructed yet.

Value

An updated SingleCellExperiment object

Author(s)

Daniel C. Ellwanger

See Also

SingleCellExperiment fitTrajectory

Examples

```
# Example data
data(exSCE)

# Set landmarks
userLandmarks(exSCE) <- colnames(exSCE)[5:7]
```

write.ygraphml	<i>Export trajectory graph</i>
----------------	--------------------------------

Description

Writes ygraphml file containing the trajectory graph's structure.

Usage

```
write.ygraphml(
  sce,
  file,
  color_by = c("phenoName", "featureName"),
  name,
  node_label = "state"
)
```

Arguments

sce	A SingleCellExperiment object
file	Character string naming a file
color_by	Indicates if nodes are colorized by a feature expression ('featureName') or phenotype label ('phenoName')
name	A character string specifying the featureName or phenoName
node_label	Defines the node label name (optional). Can be either set to the samples' states ('state') or the samples' names ('name').

Details

To visualize the trajectory graph, a proper graph layout has to be computed. Ideally, edges should not cross and nodes should not overlap (i.e., a planar embedding of the graph). CellTrails enables the export and import of the trajectory graph structure using the graphml file format. This file format can be interpreted by most third-party graph analysis applications, allowing the user to subject the trajectory graph to a wide range of (tree) layout algorithms. In particular, its format has additional ygraph attributes best suited to be used with the Graph Visualization Software 'yEd' which is freely available from yWorks GmbH (<http://www.yworks.com/products/yed>) for all major platforms.

The colors of the nodes can be defined by the parameters color_by and name. Please note that the trajectory landmarks are indicated by setting color_by='phenoName' and name='landmark'. States can be indicated by color_by='phenoName' and name='state'.

If a layout is already present in the provided CellTrailsSet object, the samples' coordinates will be listed in the graphml file.

Diagnostic messages

An error is thrown if the trajectory has not been computed yet; function fitTrajectory needs to be called first. Feature names and phenotype names get checked and will throw an error if not contained in the dataset. Please note, the parameter name is case-sensitive.

Value

write.ygraphml returns an invisible NULL

Author(s)

Daniel C. Ellwanger

See Also

fitTrajectory featureNames phenoNames

Examples

```
# Example data
data(exSCE)

## Not run:
# Export trajectory graph structure to graphml
# Color nodes by gene expression (e.g, feature_10)
write.ygraphml(sce, file="yourFilePath", color_by="featureName",
               name="feature_10")

# Color nodes by metadata (e.g., state) and
# label nodes by the (simulated) age of each sample
write.ygraphml(sce, file="yourFilePath", color_by="phenoName",
               name="state", node_label="age")

# Color and label nodes by landmark type and id
write.ygraphml(sce, file="yourFilePath", color_by="phenoName",
               name="landmark", node_label="landmark")
## End(Not run)
```

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