Package 'RaceID'

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Title Identification of Cell Types and Inference of Lineage Trees from Single-Cell RNA-Seq Data

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R topics documented:

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Description

A more detailed description of what the package does. A length of about one to five lines is recommended.

Details

This section should provide a more detailed overview of how to use the package, including the most important functions.

Author(s)

Your Name, email optional.

Maintainer: Your Name <your@email.com>

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References

This optional section can contain literature or other references for background information.

See Also

Optional links to other man pages

Examples

```
## Not run:
    ## Optional simple examples of the most important functions
    ## These can be in \dontrun{} and \donttest{} blocks.

## End(Not run)
```

barplotgene

Gene Expression Barplot

Description

This functions generates a barplot of gene expression across all clusters.

Usage

```
barplotgene(object, g, n = NULL, logsc = FALSE)
```

Arguments

object	SCseq class object.
g	Individual gene name or vector with a group of gene names corresponding to a subset of valid row names of the ndata slot of the SCseq object.
n	String of characters representing the title of the plot. Default is NULL and the first element of g is chosen.
logsc	logical. If TRUE, then gene expression values are log2-transformed after adding a pseudo-count of 0.1. Default is FALSE and untransformed values are shown.

Value

None

baseLineVar 5

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Baseline gene expression variability

Description

This function returns the base line variability as a function of the

Usage

```
baseLineVar(x, y)
```

Arguments

x mean expression. The corresponding corrected variance is returned.

y object returned by compNoise, noiseBaseFit, pruneKnn or fitBackVar. De-

pending on the input the funtion returns either the background variability (for

pruneKnn or fitBackVar) or the base line variability.

Value

Base line (or background) variability.

Examples

```
y <- noiseBaseFit(intestinalDataSmall,step=.01,thr=.05)
x <- apply(intestinalDataSmall,1,mean)
baseLineVar(x,y)</pre>
```

branchcells

Differential Gene Expression between Links

Description

This function computes expression z-score between groups of cells from the same cluster residing on different links

Usage

```
branchcells(object, br)
```

Arguments

object Ltree class object.

br List containing two branches, where each component has to be two valid cluster

numbers seperated by a . and with one common cluster in the two components. The lower number precedes the larger one, i.e. 1.3. For each component, the

cluster number need to be ordered in increasing order.

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Value

A list of four components:

n a vector with the number of significant links for each cluster.

scl a vector with the delta entropy for each cluster.k a vector with the StemID score for each cluster.diffgenes a vector with the StemID score for each cluster.

Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
sc <- clustexp(sc)
sc <- findoutliers(sc)
sc <- comptsne(sc)
ltr <- Ltree(sc)
ltr <- compentropy(ltr)
ltr <- projcells(ltr)
ltr <- lineagegraph(ltr)
ltr <- compovalue(ltr)
x <- branchcells(ltr,list("1.3","3.6"))
head(x$diffgenes$z)
plotmap(x$scl)
plotdiffgenes(x$diffgenes, names(x$diffgenes$z)[1])</pre>
```

CCcorrect

Dimensional Reduction by PCA or ICA

Description

This functions performs dimensional reduction by PCA or ICA and removes components enriched for particular gene sets, e.g. cell cycle related genes associated with technical batch effects.

Usage

```
CCcorrect(
  object,
  vset = NULL,
  CGenes = NULL,
  ccor = 0.4,
  pvalue = 0.01,
  quant = 0.01,
  nComp = NULL,
  dimR = FALSE,
  mode = "pca",
```

CCcorrect 7

```
logscale = FALSE,
FSelect = TRUE
)
```

Arguments

object	SCseq class object.
--------	---------------------

vset List of vectors with genes sets. The loadings of each component are tested

for enrichment in any of these gene sets and if the lower quant or upper 1 - quant fraction of genes ordered by loading is enriched at a p-value < pvalue

the component is discarded. Default is NULL.

CGenes Vector of gene names. If this argument is given, gene sets to be tested for en-

richment in PCA- or ICA-components are defined by all genes with a Pearson's correlation of >ccor to a gene in CGenes. The loadings of each component are tested for enrichment in any of these gene sets and if the lower quant or upper 1 - quant fraction of genes ordered by loading is enriched at a p-value < pvalue

the component is discarded. Default is NULL.

ccor Positive number between 0 and 1. Correlation threshold used to detrmine corre-

lating gene sets for all genes in CGenes. Default is 0.4.

pvalue Positive number between 0 and 1. P-value cutoff for determining enriched com-

ponents. See vset or CGenes. Default is 0.01.

quant Positive number between 0 and 1. Upper and lower fraction of gene loadings

used for determining enriched components. See vset or CGenes. Default is

0.01.

nComp Number of PCA- or ICA-components to use. Default is NULL and the maximal

number of components is computed.

dimR logical. If TRUE, then the number of principal components to use for down-

stream analysis is derived from a saturation criterion. See function plotdimsat.

Default is FALSE and all nComp components are used.

mode "pca" or "ica" to perform either principal component analysis or independent

component analysis. Default is pca.

logscale logical. If TRUE data are log-transformed prior to PCA or ICA. Default is FALSE.

FSelect logical. If TRUE, then PCA or ICA is performed on the filtered expression ma-

trix using only the features stored in slotcluster\$features as computed in the function filterdata. See FSelect for function filterdata. Default is TRUE.

Value

The function returns an updated SCseq object with the principal or independent component matrix written to the slot dimRed\$x of the SCseq object. Additional information on the PCA or ICA is stored in slot dimRed.

Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- CCcorrect(sc,dimR=TRUE,nComp=3)</pre>
```

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cc_genes

Cell cycle markers for Mus Muscuus

Description

This dataset contains official gene symbols for markers of the S phase and G2/M phase of the cell cycle in mouse.

Usage

cc_genes

Format

A list of two components with S phase marker (s) and G2M phase marker (g2m) gene symbols.

Value

None

cellsfromtree

Extract Cells on Differentiation Trajectory

Description

This function extracts a vector of cells on a given differentiation trajectory in pseudo-temporal order determined from the projection coordinates.

Usage

```
cellsfromtree(object, z)
```

Arguments

object Ltree class object.

z Vector of valid cluster numbers ordered along the trajectory.

Value

A list of four components:

f a vector of cells ids ordered along the trajectory defined by z.

g a vector of integer number. Number i indicates that a cell resides on the link

between the i-th and (i+1)-th cluster in z.

clustdiffgenes 9

Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
sc <- clustexp(sc)
sc <- findoutliers(sc)
sc <- comptsne(sc)
ltr <- Ltree(sc)
ltr <- compentropy(ltr)
ltr <- projcells(ltr)
ltr <- lineagegraph(ltr)
ltr <- comppvalue(ltr)
x <- cellsfromtree(ltr,c(1,3,6,2))</pre>
```

clustdiffgenes

Inference of differentially expressed genes in a cluster

Description

This functions computes differentially expressed genes in a (set of) cluster(s) by comparing to all remaining cells outside of the cluster (or a given background set of clusters) based on a negative binomial model of gene expression

Usage

```
clustdiffgenes(object, cl, bgr = NULL, pvalue = 0.01)
```

Arguments

object	SCseq class object.
cl	A valid set of cluster numbers from the final cluster partition stored in the cpart slot of the SCseq object.
bgr	Ordered vector of cluster numbers to be used as background set. If NULL then all clusters not in c1 are used as background set.
pvalue	Positive real number smaller than one. This is the p-value cutoff for the inference of differential gene expression. Default is 0.01.

Value

A list of two components. The first component dg contains a a data.frame of differentially expressed genes ordered by p-value in increasing order, with four columns:

mean.ncl	mean expression across cells outside of cluster c1.
mean.cl	mean expression across cells within cluster c1.
fc	fold-change of mean expression in cluster c1 versus the remaining cells.
DV	inferred p-value for differential expression.

10 clustexp

```
padj Benjamini-Hochberg corrected FDR.
```

The second component de contains the conventional output of diffexpnb, where set B corresponds to all clusters in cl and B to the background set (all clusters in bgr or not in cl). This component can be used for plotting by plotdiffgenesnb.

Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
sc <- clustexp(sc)
sc <- findoutliers(sc)
x <- clustdiffgenes(sc,1)
head(x$dg[x$dg$fc>1,])
```

clustexp

Clustering of single-cell transcriptome data

Description

This functions performs the initial clustering of the RaceID3 algorithm.

Usage

```
clustexp(
  object,
  sat = TRUE,
  samp = NULL,
  cln = NULL,
  clustnr = 30,
  bootnr = 50,
  rseed = 17000,
  FUNcluster = "kmedoids",
  verbose = TRUE
)
```

Arguments

object	SCseq class object.
sat	logical. If TRUE, then the number of clusters is determined based on finding the saturation point of the mean within-cluster dispersion as a function of the cluster number. Default is TRUE. If FALSE, then cluster number needs to be given as cln.
samp	Number of random sample of cells used for the inference of cluster number and for inferring Jaccard similarities. Default is 1000.
cln	Number of clusters to be used. Default is NULL and the cluster number is inferred by the saturation criterion.

clustheatmap 11

clustrr Maximum number of clusters for the derivation of the cluster number by the

saturation of mean within-cluster-dispersion. Default is 30.

bootnr Number of booststrapping runs for clusterboot. Default is 50.

rseed Integer number. Random seed to enforce reproducible clustering results. Default

is 17000.

FUNcluster Clustering method used by RaceID3. One of "kmedoids", "kmeans", "hclust".

Default is "kmedoids".

verbose logical. If FALSE then status output messages are disabled. Default is TRUE.

Value

SCseq object with clustering data stored in slot cluster and slot clusterpar. The clustering partition is stored in cluster\$kpart.

Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
sc <- clustexp(sc)</pre>
```

clustheatmap

Plotting a Heatmap of the Distance Matrix

Description

This functions plots a heatmap of the distance matrix grouped by clusters.

Usage

```
clustheatmap(object, final = TRUE, hmethod = "single")
```

Arguments

object SCseq class object.

final logical. If TRUE, then cells are grouped based on final clusters after outlier iden-

tification. If FALSE, then initial clusters prior to outlier identification are used

for grouping. Default is TRUE.

hmethod Agglomeration method used for determining the cluster order from hierarchical

clustering of the cluster medoids. See hclust function.

Value

Returns a vector of cluster numbers ordered as determined by herarchical clustering of cluster the cluster medoids as depicted in the heatmap.

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compdist

Computing a distance matrix for cell type inference

Description

This functions computes the distance matrix used for cell type inference by RaceID3.

Usage

```
compdist(
  object,
  metric = "pearson",
  FSelect = TRUE,
  knn = NULL,
  alpha = 1,
  no_cores = 1
)
```

Arguments

object

SCseq class object.

metric

Distances are computed from the filtered expression matrix after optional feature selection, dimensional reduction, and/or transformation (batch correction). Possible values for metric are spearman, pearson, logpearson, euclidean, rho, phi, kendall. Default is "pearson". In case of the correlation based methods, the distance is computed as 1- correlation. rho and phi are measures of proportionality computed on non-normalized counts, taken from the **propr** package.

FSelect

Logical parameter. If TRUE, then feature selection is performed prior to RaceID3

analysis. Default is TRUE.

knn

Positive integer number of nearest neighbours used for imputing gene expression

values. Default is NULL and no imputing is done.

alpha

Positive real number. Relative weight of a cell versus its k nearest neighbour applied for imputing gene expression. A cell receives a weight of alpha while the weight of its k nearest neighbours is determined by quadratic programming. The sum across all weights is normalized to one, and the wieghted mean expression is used for computing the joint probability of a cell and each of its k nearest neighbours. These probabilities are applied for the derivation of the imputed gene expression for each cell. Default is 1. Larger values give more weight to

the gene expression observed in a cell versus its neighbourhood.

no_cores

Positive integer number. Number of cores for multithreading during imputation. If set to NULL then the number of available cores minus two is used. Default is

1.

Value

SCseq object with the distance matrix in slot distances. If FSelect=TRUE, the genes used for computing the distance object are stored in slot cluster\$features.

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Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)</pre>
```

compentropy

Compute transcriptome entropy of each cell

Description

This function computes the transcriptome entropy for each cell.

Usage

```
compentropy(object)
```

Arguments

object

Ltree class object.

Value

An Ltree class object with a vector of entropies for each cell in the same order as column names in slot sc@ndata.

Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
sc <- clustexp(sc)
sc <- findoutliers(sc)
sc <- comptsne(sc)
ltr <- Ltree(sc)
ltr <- compentropy(ltr)</pre>
```

compfr

Computation of a two dimensional Fruchterman-Rheingold representation

Description

This functions performs the computation of a Fruchterman-Rheingold graph layout based on an adjacency matrix derived from the distance object in slot distances using the **igraph** package.

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Usage

```
compfr(object, knn = 10, rseed = 15555)
```

Arguments

object SCseq class object.

knn Positive integer number of nearest neighbours used for the inference of the

Fruchterman-Rheingold layout. Default is 10.

rseed Integer number. Random seed to enforce reproducible layouts.

Value

SCseq object with layout coordinates stored in slot fr.

Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
sc <- clustexp(sc)
sc <- findoutliers(sc)
sc <- compfr(sc)</pre>
```

compMean

Function for computing local gene expression averages

Description

This function performs computation of locally averaged gene expression across the pruned k nearest neighbours at given link probability cutoff.

Usage

```
compMean(
    x,
    res,
    pvalue = 0.01,
    genes = NULL,
    regNB = FALSE,
    batch = NULL,
    ngenes = NULL,
    regVar = NULL,
    span = 0.75,
    no_cores = NULL,
    seed = 12345
)
```

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Arguments

res

x Matrix of gene expression values with genes as rows and cells as columns. The matrix need to contain the same cell IDs as columns like the input matrix used to derive the pruned k nearest neighbours with the pruneKnn function. However, it may contain a different set of genes.

List object with k nearest neighbour information returned by pruneKnn function.

pvalue Positive real number between 0 and 1. All nearest neighbours with link proba-

bility < pvalue are discarded. Default is 0.01.

genes Vector of gene names corresponding to a subset of rownames of x. Only for

these genes local gene expression averages are computed. Default is NULL and

values for all genes are returned.

regNB logical. If TRUE then gene expression averages are computed from the pearson

residuals obtained from a negative binomial regression to eliminate the dependence of the expression variance on the mean. If FALSE then averages are com-

puted from raw UMI counts. Default is FALSE.

batch vector of batch variables. Component names need to correspond to valid cell

IDs, i.e. column names of expData. If regNB is TRUE, than the batch variable will be regressed out simultaneously with the $\log 10$ UMI count per cell.An interaction term is included for the $\log 10$ UMI count with the batch variable. Default

value is NULL.

ngenes Positive integer number. Randomly sampled number of genes (from rownames

of expData) used for predicting regression coefficients (if regNB=TRUE). Smoothed coefficients are derived for all genes. Default is NULL and all genes are used.

regVar data.frame with additional variables to be regressed out simultaneously with

the log10 UMI count and the batch variable (if batch is TRUE). Column names indicate variable names (name beta is reserved for the coefficient of the log10 UMI count), and rownames need to correspond to valid cell IDs, i.e. column names of expData. Interaction terms are included for each variable in regVar

with the batch variable (if batch is TRUE). Default value is NULL.

span Positive real number. Parameter for loess-regression (see regNB) controlling the

degree of smoothing. Default is 0.75.

no_cores Positive integer number. Number of cores for multithreading. If set to NULL then

the number of available cores minus two is used. Default is 1.

seed Integer number. Random number to initialize stochastic routines. Default is

12345.

Value

List object of three components:

mean matrix with local gene expression averages, computed from Pearson residuals (if

regNB=TRUE) or normalized UMI counts (if regNB=FALSE). In the latter case, the average UMI count for a local neighbourhood is normalized to one and rescaled

by the median UMI count across neighborhoods.

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regData

If regNB=TRUE this argument contains a list of four components: component pearsonRes contains a matrix of the Pearson Residual computed from the negative binomial regression, component nbRegr contains a matrix with the regression coefficients, component nbRegrSmooth contains a matrix with the smoothed regression coefficients, and log10_umi is a vector with the total log10 UMI count for each cell. The regression coefficients comprise the dispersion parameter theta, the intercept, the regression coefficient beta for the log10 UMI count, and the regression coefficients of the batches (if batch is not NULL).

Examples

```
res <- pruneKnn(intestinalDataSmall,metric="pearson",knn=10,alpha=1,no_cores=1,FSelect=FALSE)
mexp <- compMean(intestinalDataSmall,res,pvalue=0.01,genes = NULL,no_cores=1)</pre>
```

compmedoids

Computes Medoids from a Clustering Partition

Description

This functions computes cluster medoids given an SCseq object and a clustering partition. The medoids are either derived from the distance matrix or, if the slot distances is empty, from the dimensionally reduced feature matrix in slot dimRed\$x using the euclidean metric.

Usage

```
compmedoids(object, part)
```

Arguments

object SCseq class object.

part Clustering partition. A vector of cluster numbers for (a subset of) cells (i.e.

column names) of slot ndata from the SCseq object.

Value

Returns a list of medoids (column names of slot ndata from the SCseq object) ordered by increasing cluster number.

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compNoise

Function for computing local gene expression variability

Description

This function performs computation of the local gene expression variability across the pruned k nearest neighbours at given link probability cutoff. The estimated variance is corrected for the mean dependence utilizing the baseline model of gene expression variance.

Usage

```
compNoise(
    x,
    res,
    pvalue = 0.01,
    genes = NULL,
    regNB = FALSE,
    batch = NULL,
    ngenes = NULL,
    regVar = NULL,
    span = 0.75,
    step = 0.01,
    thr = 0.05,
    no_cores = NULL,
    seed = 12345
)
```

Arguments

v	
А	

Matrix of gene expression values with genes as rows and cells as columns. The matrix need to contain the same cell IDs as columns like the input matrix used to derive the pruned k nearest neighbours with the pruneKnn function. However, it may contain a different set of genes.

res

List object with k nearest neighbour information returned by pruneKnn function.

pvalue

Positive real number between 0 and 1. All nearest neighbours with link probability < pvalue are discarded. Default is 0.01.

genes

Vector of gene names corresponding to a subset of rownames of x. Only for these genes local gene expression variability is computed. Default is NULL and values for all genes are returned.

regNB

logical. If TRUE then gene expression variability is derived from the pearson residuals obtained from a negative binomial regression to eliminate the dependence of the expression variance on the mean. If FALSE then the mean dependence is regressed out from the raw variance using the baseline variance estimate. Default is FALSE.

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batch vector of batch variables. Component names need to correspond to valid cell

IDs, i.e. column names of expData. If regNB is TRUE, than the batch variable will be regressed out simultaneously with the $\log 10$ UMI count per cell. An interaction term is included for the $\log 10$ UMI count with the batch variable.

Default value is NULL.

ngenes Positive integer number. Randomly sampled number of genes (from rownames

of expData) used for predicting regression coefficients (if regNB=TRUE). Smoothed coefficients are derived for all genes. Default is NULL and all genes are used.

regVar data.frame with additional variables to be regressed out simultaneously with

the log10 UMI count and the batch variable (if batch is TRUE). Column names indicate variable names (name beta is reserved for the coefficient of the log10 UMI count), and rownames need to correspond to valid cell IDs, i.e. column names of expData. Interaction terms are included for each variable in regVar

with the batch variable (if batch is TRUE). Default value is NULL.

span Positive real number. Parameter for loess-regression (see regNB) controlling the

degree of smoothing. Default is 0.75.

step Positive real number between 0 and 1. See function noiseBaseFit. Default is

0.01.

thr Positive real number between 0 and 1. See function noiseBaseFit. Default is

0.05.

no_cores Positive integer number. Number of cores for multithreading. If set to NULL then

the number of available cores minus two is used. Default is 1.

seed Integer number. Random number to initialize stochastic routines. Default is

12345.

Value

List object of three components:

model the baseline noise model as computed by the noiseBaseFit function.

data matrix with local gene expression variability estimates, corrected for the mean

dependence.

regData If regNB=TRUE this argument contains a list of four components: component

pearsonRes contains a matrix of the Pearson Residual computed from the negative binomial regression, component nbRegr contains a matrix with the regression coefficients, component nbRegrSmooth contains a matrix with the smoothed regression coefficients, and $log10_umi$ is a vector with the total log10 UMI count for each cell. The regression coefficients comprise the dispersion parameter theta, the intercept, the regression coefficient beta for the log10 UMI count,

and the regression coefficients of the batches (if batch is not NULL).

Examples

res <- pruneKnn(intestinalDataSmall,metric="pearson",knn=10,alpha=1,no_cores=1,FSelect=FALSE)
noise <- compNoise(intestinalDataSmall,res,pvalue=0.01,genes = NULL,no_cores=1)</pre>

19 comppvalue

comppvalue Computing P-values for Link Significance	
---	--

Description

This function computes a p-value for the significance (i.e. over-representation of assigned cells) of each inter-cluster link.

Usage

```
comppvalue(object, pthr = 0.01, sensitive = FALSE)
```

Arguments

Ltree class object. object p-value cutoff for link significance. This threshold is applied for the calculation pthr of link scores reflecting how uniformly a link is occupied by cells. sensitive logical. Only relevant when nmode=TRUE in function projcell. If TRUE, then all cells on the most highly significant link are and the link itself are disregard

to test significance of the remaining links with a binomial p-value. Default is

FALSE.

Value

An Ltree class object with link p-value and occupancy data stored in slot cdata.

Examples

```
sc <- SCseq(intestinalDataSmall)</pre>
sc <- filterdata(sc)</pre>
sc <- compdist(sc)</pre>
sc <- clustexp(sc)</pre>
sc <- findoutliers(sc)</pre>
sc <- comptsne(sc)</pre>
ltr <- Ltree(sc)</pre>
ltr <- compentropy(ltr)</pre>
ltr <- projcells(ltr)</pre>
ltr <- lineagegraph(ltr)</pre>
ltr <- comppvalue(ltr)</pre>
```

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compscore	Compute StemID2 score	
mpscore	Compute Stem1D2 score	

Description

This function extracts the number of links connecting a given cluster to other cluster, the delta median entropy of each cluster (median entropy of a cluster after subtracting the minimum median entropy across all clusters), and the StemID2 score which is the product of both quantities for each cluster.

Usage

```
compscore(object, nn = 1, scthr = 0, show = TRUE)
```

Arguments

object Ltree class object.

nn Positive integer number. Number of higher order neighbors to be included for

the determination of links: indirect connections via n-1 intermittant neighbors

and determination of miks. Indirect connections via it is intermittal

are allowed. Default is 1.

scthr Real number between zero and one. Score threshold for links to be included in

the calculation. For scthr=0 all significant links are included. The maximum

score is one.

show logical. If TRUE, then plot heatmap of projections. Default is TRUE.

Value

A list of three components:

links a vector with the number of significant links for each cluster.

entropy a vector with the delta entropy for each cluster.

StemIDscore a vector with the StemID score for each cluster.

comptsne	Computation of a two dimensional t-SNE representation

Description

This functions performs the computation of a t-SNE map from the distance object in slot distances using the **Rtsne** package.

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Usage

```
comptsne(
  object,
  dimRed = FALSE,
  initial_cmd = TRUE,
  perplexity = 30,
  rseed = 15555
)
```

Arguments

object SCseq class object.

dimRed logical. If TRUE then the t-SNE is computed from the feature matrix in slot

dimRed\$x (if not equal to NULL). Default is FALSE and the t-SNE is computed from the distance matrix stored in slot distances. If slot distances equals

NULL dimRed is automatially set to TRUE.

initial_cmd logical. If TRUE, then the t-SNE map computation is initialized with a configu-

ration obtained by classical multidimensional scaling. Default is TRUE.

perplexity Positive number. Perplexity of the t-SNE map. Default is 30.

rseed Integer number. Random seed to enforce reproducible t-SNE map.

Value

SCseq object with t-SNE coordinates stored in slot tsne.

Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
sc <- clustexp(sc)
sc <- findoutliers(sc)
sc <- comptsne(sc)</pre>
```

compumap

Computation of a two dimensional umap representation

Description

This functions performs the computation of a two-dimensional umap representation based on the distance matrix in slot distances using the **umap** package.

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Usage

```
compumap(
  object,
  dimRed = FALSE,
  n_neighbors = 15,
  metric = "euclidean",
  n_epochs = 200,
  min_dist = 0.1,
  local_connectivity = 1,
  spread = 1
)
```

Arguments

object SCseq class object.

dimRed logical. If TRUE then the umap is computed from the feature matrix in slot

dimRed\$x (if not equal to NULL). Default is FALSE and the umap is computed from the distance matrix stored in slot distances. If slot distances equals

NULL dimRed is automatially set to TRUE.

n_neighbors Umap parameter. See help(umap.defaults) after loading package **umap**. De-

fault is 15.

metric Umap parameter. See help(umap.defaults) after loading package umap. De-

fault is "euclidean".

n_epochs Umap parameter. See help(umap.defaults) after loading package umap. De-

fault is 200.

min_dist Umap parameter. See help(umap.defaults) after loading package umap. De-

fault is 0.1.

local_connectivity

Umap parameter. See help(umap.defaults) after loading package umap. De-

fault is 1.

spread Umap parameter. See help(umap.defaults) after loading package umap. De-

fault is 1.

Value

SCseq object with umap coordinates stored in slot umap.

Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
sc <- clustexp(sc)
sc <- findoutliers(sc)
sc <- compumap(sc)</pre>
```

createKnnMatrix 23

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Function to create a knn matrix

Description

This creates an adjacency matrix, keeping only nearest neighbour with a link probability above a minimum probability

Usage

```
createKnnMatrix(res, pvalue = 0.01)
```

Arguments

res List object with k nearest neighbour information returned by pruneKnn function.

pvalue Positive real number between 0 and 1. All nearest neighbours with link proba-

bility < pvalue are discarded. Default is 0.01.

Value

Adjacency matrix in sparse matrix format (see package **Matrix**) with positive non-zero entries only for k nearest neighburs with link probability >= pvalue. The value of these entries equals the link probability.

Examples

```
res <- pruneKnn(intestinalDataSmall,metric="pearson",knn=10,alpha=1,no_cores=1,FSelect=FALSE)
y <- createKnnMatrix(res,pvalue=0.01)</pre>
```

diffexpnb

Function for differential expression analysis

Description

This function performs differential expression analysis between two sets of single cell transcriptomes. The inference is based on a noise model or relies on the DESeq2 approach.

Usage

```
diffexpnb(
    x,
    A,
    B,
    DESeq = FALSE,
    method = "pooled",
    norm = FALSE,
```

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```
vfit = NULL,
locreg = FALSE,
...
)
```

Arguments

Χ

expression data frame with genes as rows and cells as columns. Gene IDs should be given as row names and cell IDs should be given as column names. This can be a reduced expression table only including the features (genes) to be used in the analysis. This input has to be provided if g (see below) is given and corresponds to a valid gene ID, i. e. one of the rownames of x. The default value is NULL. In this case, cluster identities are highlighted in the plot.

A vector of cell IDs corresponding column names of x. Differential expression in

set A versus set B will be evaluated.

B vector of cell IDs corresponding column names of x. Differential expression in

set A versus set B will be evaluated.

DESeq logical value. If TRUE, then **DESeq2** is used for the inference of differentially

expressed genes. In this case, it is recommended to provide non-normalized input data x. The **DESeq2** package needs to be installed from bioconductor.

Default value is FALSE.

method either "per-condition" or "pooled". If DESeq is not used, this parameter deter-

mines, if the noise model is fitted for each set separately ("per-condition") or for

the pooled set comprising all cells in A and B. Default value is "pooled".

norm logical value. If TRUE then the total transcript count in each cell is normalized to

the minimum number of transcripts across all cells in set A and B. Default value

is FALSE.

vfit function describing the background noise model. Inference of differentially ex-

pressed genes can be performed with a user-specified noise model describing the expression variance as a function of the mean expression. Default value is

NULL.

locreg logical value. If FALSE then regression of a second order polynomial is performed

to determine the relation of variance and mean. If TRUE a local regression is

performed instead. Default value is FALSE.

... additional arguments to be passed to the low level function DESeqDataSetFromMatrix.

Value

If DESeq equals TRUE, the function returns the output of **DESeq2**. In this case list of the following two components is returned:

cds object returned by the **DESeq2** function DESeqDataSetFromMatrix.

res data frame containing the results of the **DESeq2** analysis.

Otherwise, a list of three components is returned:

vf1 a data frame of three columns, indicating the mean m, the variance v and the

fitted variance vm for set A.

diffgenes 25

vf2	a data frame of three columns, indicating the mean m , the variance ν and the fitted variance νm for set B.
res	a data frame with the results of the differential gene expression analysis with the structure of the DESeq output, displaying mean expression of the two sets, fold change and log2 fold change between the two sets, the p-value for differential expression (pval) and the Benjamini-Hochberg corrected false discovery rate (padj).

Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
sc <- clustexp(sc)
sc <- findoutliers(sc)
A <- names(sc@cpart)[sc@cpart %in% c(1,2)]
B <- names(sc@cpart)[sc@cpart %in% c(3)]
y <- diffexpnb(getfdata(sc,n=c(A,B)), A=A, B=B)</pre>
```

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Compute Expression Differences between Clusters

Description

This functions computes expression differences between clusters and ranks genes by z-score differences.

Usage

```
diffgenes(object, cl1, cl2, mincount = 1)
```

Arguments

object	SCseq class object.
cl1	A vector of valid cluster numbers (contained in the cpart slot of the SCseq object). Represents the first group of the comparison.
c12	A vector of valid cluster numbers (contained in the cpart slot of the SCseq object). Represents the second group of the comparison.
mincount	Minimal normalized expression level of a gene to be included into the analysis. A gene needs to be expressed at this level in at least a single cell.

Value

A list with four components:

z a vector of z-scores in decreasing order with genes up-regulated in cl1 appearing at the top of the list.

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cl1	a data.frame with expression values for cells in cl1.
c12	a data.frame with expression values for cells in c12.
cl1n	a vector of cluster numbers for cells in cl1.
cl2n	a vector of cluster numbers for cells in c12.

Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
sc <- clustexp(sc)
sc <- findoutliers(sc)
x <- diffgenes(sc,1,2)
head(x$z)
plotdiffgenes(x,names(x$z)[1])</pre>
```

diffNoisyGenes

Function for extracting genes with elevated variability in a cluster

Description

This function extracts genes with significantly elevated variability in a cluster on a basis of a Wilcoxon rank sum-test between cells in a cluster and all remaining cells.

Usage

```
diffNoisyGenes(noise, cl, set, bgr = NULL, no_cores = 1)
```

Arguments

noise	List object with the background noise model and a variability matrix, returned by the compNoise function.
cl	List object with Louvain clustering information, returned by the graphCluster function.
set	Postive integer number or vector of integers corresponding to valid cluster numbers. The function reports genes with elevated variability in all clusters contained in set.
bgr	Postive integer number or vector of integers corresponding to valid cluster numbers. Background set for comparison. The function reports genes with elevated variability in all clusters contained in set compared to clusters in bgr. Default is NULL and the comparison is against all clusters not in set.
no_cores	Positive integer number. Number of cores for multithreading. If set to NULL then the number of available cores minus two is used. Default is 1.

filterdata 27

Value

Data.frame reporting the log2 fold change between clusters in set and the remaining clusters and the p-value for elevated variability for each genes. Rows are ordered by decreasing log2 fold change.

Examples

```
res <- pruneKnn(intestinalDataSmall,metric="pearson",knn=10,alpha=1,no_cores=1,FSelect=FALSE)
noise <- compNoise(intestinalDataSmall,res,pvalue=0.01,genes = NULL,no_cores=1)
cl <- graphCluster(res,pvalue=0.01)
ngenes <- diffNoisyGenes(noise,cl,c(1,2),no_cores=1)</pre>
```

filterdata

Data filtering

Description

This function allows filtering of genes and cells to be used in the RaceID3 analysis. It also can perform batch effect correction using an internal method or a recently published alternative mnnCorrect from the **batchelor** package.

Usage

```
filterdata(
  object,
  mintotal = 3000,
  minexpr = 5,
  minnumber = 5,
  LBatch = NULL,
  knn = 10,
  CGenes = NULL,
  FGenes = NULL,
  ccor = 0.4,
  bmode = "RaceID",
  verbose = TRUE
)
```

Arguments

object SCseq class object.

minitotal minimum total transcript number required. Cells with less than mintotal tran-

scripts are filtered out. Default is 3000.

minexpr minimum required transcript count of a gene in at least minnumber cells. All

other genes are filtered out. Default is 5.

minnumber See minexpr. Default is 5.

LBatch List of experimental batches used for batch effect correction. Each list element

contains a vector with cell names (i.e. column names of the input expression

data) falling into this batch. Default is NULL, i.e. no batch correction.

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knn	Number of nearest neighbors used to infer corresponding cell types in different batches. Defult is 10.
CGenes	List of gene names. All genes with correlated expression to any of the genes in CGenes are filtered out for cell type inference. Default is NULL.
FGenes	List of gene names to be filtered out for cell type inference. Default is NULL.
ccor	Correlation coefficient used as a trehshold for determining genes correlated to genes in CGenes. Only genes correlating less than ccor to all genes in CGenes are retained for analysis. Default is 0.4.
bmode	Method used for batch effect correction. Any of "RaceID", "mnnCorrect". If mnnCorrect from the batchelor package is desired, this package needs to be installed from bioconductor. Default is "RaceID".
verbose	logical. If FALSE then status output messages are disabled. Default is TRUE.

Value

An SCseq class object with filtered and normalized expression data.

Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)</pre>
```

findoutliers

Inference of outlier cells and final clustering

Description

This functions performs the outlier identification based on the clusters infered with the clustexp function.

Usage

```
findoutliers(
  object,
  probthr = 0.001,
  outminc = 5,
  outlg = 2,
  outdistquant = 0.95,
  verbose = TRUE
)
```

fitBackVar 29

Arguments

object	SCseq class object.
probthr	outlier probability threshold for a minimum of outlg genes to be an outlier cell. This probability is computed from a negative binomial background model of expression in a cluster. Default is 0.001 .
outminc	minimal transcript count of a gene in a clusters to be tested for being an outlier gene. Default is 5.
outlg	Minimum number of outlier genes required for being an outlier cell. Default is 2.
outdistquant	Real number between zero and one. Outlier cells are merged to outlier clusters if their distance smaller than the outdistquant-quantile of the distance distribution of pairs of cells in the original clusters after outlier removal. Default is 0.95.
verbose	logical. If FALSE then status output messages are disabled. Default is TRUE.

Value

SCseq object with outlier data stored in slot out and slot outlierpar. The final clustering partition is stored in cpart.

Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
sc <- clustexp(sc)
sc <- findoutliers(sc)</pre>
```

fitBackVar	Function for computing a background model of gene expression variability

Description

This funtion fits a second order polynomial to the variance-mean dependence across all genes in log space.

Usage

```
fitBackVar(x, mthr = -1)
```

Arguments

Х	Matrix of gene expression values with genes as rows and cells as columns.
mthr	Real number. Threshold of log2 mean expression. Genes with mean expression
	< mthr are discarded prior to fitting the polynomial. Default is -1.

30 fractDotPlot

Value

List object of four components:

fit model fit as returned by the 1m function.

genes with expression variance greater than the polynomial fit.

m mean expression of all genesv expression variance of all genes

Examples

```
bg <- fitBackVar(intestinalDataSmall)</pre>
```

fractDotPlot

Dotplot of gene expression across clusters or samples

Description

This is a plotting function for visualizing gene expression across subsets of clusters or samples. The diameter of a dot reflects the fraction of cells expressing a gene, and the color indicates the expression z-score across all clusters or samples.

Usage

```
fractDotPlot(
  object,
  genes,
  cluster = NULL,
  samples = NULL,
  subset = NULL,
  zsc = FALSE,
  logscale = TRUE,
  cap = Inf,
  flo = -Inf
)
```

Arguments

object SCseq class object.

genes vector of valid gene names corresponding to row names of slot ndata. The

expression for this genes is shown.

cluster vector of valid cluster numbers contained in slot cpart. Default is NULL. If

not given, then the samples argument is expected. If both are given, only the samples argument is considered. If both are NULL, then cluster is initialized

with all clusters.

samples vector of sample names for all cells. Length and order has to correspond to

colnames of slot ndata. Default is NULL.

getExpData 31

subset	vector of unique sample names to show in the expression dotplot. Each sample names in subset has to occur in samples. Default is NULL. If not given and samples is not NULL, the subset is intialized with all sample names occuring in samples.
ZSC	logical. If TRUE then a z-score transformation is applied. Default is FALSE.
logscale	logical. If TRUE then a log2 transformation is applied. Default is TRUE.
сар	real number. Upper limit for the expression, log2 expression, or z-score. Values larges then cap are replaced by cap.
flo	real number. Lower limit for the expression, log2 expression, or z-score. Values smaller then flo are replaced by flo.

Value

None

getExpData	Function for extracting a filtered expression matrix from a RaceID SCseq object

Description

This function for extracts a filtered expression matrix from a **RaceID** SCseq object. The filterdata function from the **RaceID** package has to be run on the SCseq object before.

Usage

```
getExpData(object, genes = NULL)
```

Arguments

object RaceID SCseq object.

genes Vector of valid gene identifiers corresponding to valid rownames of the input

expression data. An expression matrix is returned only for these genes. Default is NULL and an expression matrix is returned for all genes retained after filtering

of the SCseq object, i.e. all genes in genes slot of the SCseq object.

Value

noise Sparse Matrix with genes as rows and cells as columns after filtering.

Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
d <- getExpData(sc)
res <- pruneKnn(d,distM=sc@distances,metric="pearson",knn=10,alpha=1,no_cores=1,FSelect=FALSE)</pre>
```

32 getproj

getfdata	Extracting filtered expression data	

Description

This functions allows the extraction of a filtered and normalized expression matrix

Usage

```
getfdata(object, g = NULL, n = NULL)
```

Arguments

object	SCseq class object.
g	Vector of gene names to be included corresponding to a subset of valid row names of the ndata slot of the SCseq object. Default is NULL and data for all genes remaining after filtering by the filterdata function are shown.
n	Vector of valid column names corresponding to a subset of valid column names of the ndata slot of the SCseq object. Default is NULL and data for all cells remaining after filtering by the filterdata function are shown.

Value

Matrix of filtered expression data with genes as rows and cells as columns.

getproj	Extract Projections of all Cells from a Cluster	

Description

This function extracts projections of all cells in a cluster and plots a heatmap of these hierarchically clustered projections (rows) to all other clusters (columns). A minimum spanning tree of the cluster centers is overlaid for comparison.

Usage

```
getproj(object, i, show = TRUE, zscore = FALSE)
```

Arguments

object	Ltree class object.
i	Cluster number. This number has to correspond to one of the RaceID3 clusters included for the StemID2 inference, i.e. to a number present in slot ldata\$1p.
show	logical. If TRUE, then plot heatmap of projections. Default is TRUE.
zscore	logical. If TRUE and show=TRUE, then plot z-score-transformed projections. If TRUE and show=FALSE, then plot untransformed projections. Default is FALSE.

graphCluster 33

Value

A list of two components:

pr a data frame of projections for all cells in cluster i (rows) onto all other clusters

(columns).

prz a data.frame of z-transformed projections for all cells in cluster i (rows) onto all

other clusters (columns).

graphCluster Function for infering Louvain clustering of the pruned k nearest neigh-

bour graph

Description

This function derives a graph object from the pruned nearest neighbours and infers clusters by the the Louvain clustering method on this graph. A Fruchterman-Rheingold graph layout is also derived from the pruned nearest neighbours.

Usage

```
graphCluster(res, pvalue = 0.01, use.weights = TRUE, rseed = 12345)
```

Arguments

res List object with k nearest neighbour information returned by pruneKnn function.

pvalue Positive real number between 0 and 1. All nearest neighbours with link proba-

hility < nual up are discarded Default is 0.01

bility < pvalue are discarded. Default is 0.01.

use.weights logical. If TRUE, then nearest-neighbor link probabilities are used to build

a graph as input for Louvain clustering. If FALSE, then all links have equal

weight. Default is TRUE.

rseed Integer number. Random seed to enforce reproducible clustering results. Default

is 12345.

Value

List object of three components:

graph graph derived from the pruned adjacency matrix computed with the igraph

package.

louvain Louvain clustering returned by the cluster_louvain function from the igraph

package.

fr Fruchterman-Rheingold graph layout derived from the pruned adjacency matrix.

Examples

```
res <- pruneKnn(intestinalDataSmall,metric="pearson",knn=10,alpha=1,no_cores=1,FSelect=FALSE)
cl <- graphCluster(res,pvalue=0.01)</pre>
```

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imputeexp

Imputed expression matrix

Description

This functions returns an imputed expression matrix based on the imputing computed with compdist.

Usage

```
imputeexp(object, genes = NULL)
```

Arguments

object

SCseq class object.

genes

vector of valid gene names corresponding to row names of slot ndata. Default

is NULL and imputing is done for all genes.

Value

An expression matrix with imputed expression values after size normalization. Genes are in rows and cells in columns.

intestinalData

Single-cell transcriptome data of intestinal epithelial cells

Description

This dataset contains gene expression values, i. e. transcript counts, of 278 intestinal epithelial cells

Usage

intestinalData

Format

A sparse matrix (using the **Matrix**) with cells as columns and genes as rows. Entries are raw transcript counts.

Value

None

References

Grün et al. (2016) Cell Stem Cell 19(2): 266-77 < DOI:10.1016/j.stem.2016.05.010 > (PubMed)

intestinalDataSmall 35

intestinalDataSmall

Single-cell transcriptome data of intestinal epithelial cells

Description

This dataset is a smaller subset of the original dataset, which contains gene expression values, i. e. transcript counts, of 278 intestinal epithelial cells. The dataset is included for quick testing and examples. Only cells with >10,000 transcripts per cell and only genes with >20 transcript counts in >10 cells were retained.

Usage

intestinalDataSmall

Format

A sparse matrix (using the **Matrix**) with cells as columns and genes as rows. Entries are raw transcript counts.

Value

None

References

Grün et al. (2016) Cell Stem Cell 19(2): 266-77 <DOI:10.1016/j.stem.2016.05.010> (PubMed)

lineagegraph

Inference of a Lineage Graph

Description

This function assembles a lineage graph based on the cell projections onto inter-cluster links.

Usage

```
lineagegraph(object, verbose = TRUE)
```

Arguments

object Ltree class object.

verbose logical. If FALSE then status output messages are disabled. Default is TRUE.

Value

An Ltree class object with lineage graph-related data stored in slots 1tcoord, prtree, and cdata.

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Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
sc <- clustexp(sc)
sc <- findoutliers(sc)
sc <- comptsne(sc)
ltr <- Ltree(sc)
ltr <- compentropy(ltr)
ltr <- projcells(ltr)
ltr <- lineagegraph(ltr)</pre>
```

Ltree-class

The Ltree Class

Description

The Ltree class is the central object storing all information generated during lineage tree inference by the StemID algorithm. It comprises a number of slots for a variety of objects.

Arguments

object

An Ltree object.

Slots

sc An SCseq object with the RaceID3 analysis of the single-cell RNA-seq data for which a lineage tree should be derived.

ldata List object storing information on the clustering partition, the distance matrix, and the cluster centers in dimensionally-reduced input space and in two-dimensional t-sne space. Elements: lp: vector with the filtered partition into clusters after discarding clusters with cthr cells or less. pdi:matrix with the coordinates of all cells in the embedded space. Clusters with cthr transcripts or less were discarded (see function projcells). Rows are medoids and columns are coordinates. cn: data.frame with the coordinates of the cluster medoids in the embedded space. Clusters with cthr transcripts or less were discarded. Rows are medoids and columns are coordinates. m: vector with the numbers of the clusters which survived the filtering. pdil: data.frame with coordinates of cells in the two-dimensional t-SNE representation computed by RaceID3. Clusters with cthr transcripts or less were discarded. Rows are cells and columns are coordinates. cnl: data.frame with the coordinates of the cluster medoids in the two-dimensional t-SNE representation computed by RaceID3. Clusters with cthr transcripts or less were discarded. Rows are medoids and columns are coordinates.

entropy Vector with transcriptome entropy computed for each cell.

trproj List containing two data.frames. Elements: res: data.frame with three columns for each cell. The first column o shows the cluster of a cell, the second column 1 shows the cluster number for the link the cell is assigned to, and the third column h shows the projection as a fraction of the length of the inter-cluster link. Parallel projections are positive, while antiparallel projections are negative. rma: data.frame with all projection coordinates for each cell.

maxNoisyGenes 37

Rows are cells and columns are clusters. Projections are given as a fraction of the length of the inter-cluster link. Parallel projections are positive, while anti-parallel projections are negative. The column corresponding to the originating cluster of a cell shows NA.

par List of parameters used for the StemID2 analysis.

prback data.frame of the same structure as the trproj\$res. In case randomizations are used to compute significant projections, the projections of all pdishuff randomizations are appended to this data.frame and therefore the number of rows corresponds to the number of cells multiplied by pdishuf. See function projback.

prbacka data.frame reporting the aggregated results of the randomizations with four columns. Column n denotes the number of the randomization sample, column o and 1 contain the numbers of the originating and the terminal cluster, respectively, for each inter-cluster link and column count shows the number of cells assigned to this link in randomization sample n. The discrete distribution for the computation of the link p-value is given by the data contained in this object (if nmode=FALSE).

1tcoord Matrix storing projection coordinates of all cells in the two-dimensional t-SNE space, used for visualization.

prtree List with two elements. The first element 1 stores a list with the projection coordinates for each link. The name of each element identifies the link and is composed of two cluster numbers separated by a dot. The second element n is a list of the same structure and contains the cell names corresponding to the projection coordinates stored in 1.

cdata list of data.frames, each with cluster ids as rows and columns: counts data.frame indicating the number of cells on the links connecting the cluster of origin (rows) to other clusters (columns). counts.br data.frame containing the cell counts on cluster connections averaged across the randomized background samples (if nmode = FALSE) or as derived from sampling statistics (if nmode = TRUE). pv.e matrix of enrichment p-values estimated from sampling statistics (if nmode = TRUE); entries are 0 if the observed number of cells on the respective link exceeds the (1 pethr)-quantile of the randomized background distribution and 0.5 otherwise (if nmode = FALSE), pv.d matrix of depletion p-values estimated from sampling statistics (if nmode = TRUE); entries are 0 if the observed number of cells on the respective link is lower than the pethr-quantile of the randomized background distribution and 0.5 otherwise (if nmode = FALSE). pvn.e matrix of enrichment p-values estimated from sampling statistics (if nmode = TRUE); 1- quantile, with the quantile estimated from the number of cells on a link as derived from the randomized background distribution (if nmode = FALSE). pvn.d matrix of depletion p-values estimated from sampling statistics (if nmode = TRUE); quantile estimated from the number of cells on a link as derived from the randomized background distribution (if nmode = FALSE).

maxNoisyGenes

Function for extracting genes maximal variability

Description

This function extracts genes with maximal variability in a cluster or in the entire data set.

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Usage

```
maxNoisyGenes(noise, cl = NULL, set = NULL)
```

Arguments

noise	List object with the background noise model and a variability matrix, returned by the compNoise function.
cl	List object with Louvain clustering information, returned by the graphCluster function. Default is NULL.
set	Postive integer number or vector of integers corresponding to valid cluster numbers. Noise levels are computed across all cells in this subset of clusters. Default is NULL and noise levels are computed across all cells.

Value

Vector with average gene expression variability in decreasing order, computed across all cells or only cells in a set of clusters (if cl and set are given.

Examples

```
res <- pruneKnn(intestinalDataSmall,metric="pearson",knn=10,alpha=1,no_cores=1,FSelect=FALSE)
noise <- compNoise(intestinalDataSmall,res,pvalue=0.01,genes = NULL,no_cores=1)
mgenes <- maxNoisyGenes(noise)</pre>
```

noiseBaseFit	Function for computing a fit to the baseline of gene expression variability

Description

This function fits a second order polynomial to the baseline variance-mean dependence across all genes in log space.

Usage

```
noiseBaseFit(x, step = 0.01, thr = 0.05)
```

Arguments

x	Matrix of gene expression values with genes as rows and cells as columns.
step	Positive real number between 0 and 1. Bin size for the computation. The interval of mean gene expression values is divided into bins with equal number of data points and step equals the fraction of data points in each bin. Default is 0.01.
thr	Positive real number between 0 and 1. In each mean expression bin defined by step the lowest thr-quantile of the gene expression variance distribution is selected. The selected data points from all bins are used for a second order polynomial fit of the variance-mean dependence in log space. Default is 0.05.

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Value

List object of three components:

nfit model fit as returned by the 1m function.

m mean expression of all genesv expression variance of all genes

Examples

```
x <- noiseBaseFit(intestinalDataSmall,step=.01,thr=.05)</pre>
```

plotbackground

Plot Background Model

Description

This functions produces a scatter plot showing the gene expression variance as a function of the mean and the inferred polynomial fit of the background model computed by RaceID3. It also shows a local regression.

Usage

```
plotbackground(object)
```

Arguments

object

SCseq class object.

Value

None

plotBackVar

Function for plottinhg the background model of gene expression variability

Description

This function plots the variance against mean expression across all genes and a second order polynomial to the variance-mean dependence in log space. It also plots a local regression.

Usage

```
plotBackVar(x)
```

40 plotdiffgenes

Arguments

Х

List object returned by function fitBackVar or list object returned by function pruneKnn.

Value

None

Examples

```
res <- pruneKnn(intestinalDataSmall,metric="pearson",knn=10,alpha=1,no\_cores=1,FSelect=FALSE) \\ plotBackVar(res)
```

plotdiffgenes

Barplot of differentially expressed genes

Description

This functions produces a barplot of differentially expressed genes derived by the function diffgenes

Usage

```
plotdiffgenes(z, gene)
```

Arguments

z Output of diffgenes

gene

Valid gene name. Has to correspond to one of the rownames of the ndata slot of the SCseq object.

Value

None

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
sc <- clustexp(sc)
sc <- findoutliers(sc)
x <- diffgenes(sc,1,2)
head(x$z)
plotdiffgenes(x,names(x$z)[1])</pre>
```

plotdiffgenesnb 41

Function for plotting differentially expressed genes

Description

This is a plotting function for visualizing the output of the diffexpnb or clustdiffgenes function as MA plot.

Usage

```
plotdiffgenesnb(
    X,
    pthr = 0.05,
    padj = TRUE,
    lthr = 0,
    mthr = -Inf,
    Aname = NULL,
    Bname = NULL,
    show_names = TRUE,
    ...
)
```

Arguments

X	output of the function diffexpnb.
pthr	real number between 0 and 1. This number represents the p-value cutoff applied for displaying differentially expressed genes. Default value is 0.05. The parameter padj (see below) determines if this cutoff is applied to the uncorrected p-value or to the Benjamini-Hochberg corrected false discovery rate.
padj	logical value. If TRUE, then genes with a Benjamini-Hochberg corrected false discovery rate lower than pthr are displayed. If FALSE, then genes with a p-value lower than pthr are displayed.
lthr	real number between 0 and Inf. Differentially expressed genes are displayed only for log2 fold-changes greater than 1thr. Default value is 0.
mthr	real number between -Inf and Inf. Differentially expressed genes are displayed only for log2 mean expression greater than mthr. Default value is -Inf.
Aname	name of expression set A, which was used as input to diffexpnb. If provided, this name is used in the axis labels. Default value is NULL.
Bname	name of expression set B, which was used as input to diffexpnb. If provided, this name is used in the axis labels. Default value is NULL.
show_names	logical value. If TRUE then gene names displayed for differentially expressed genes. Default value is FALSE.
	Additional arguments for function plot.

42 plotdimsat

Value

None

Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
sc <- clustexp(sc)
sc <- findoutliers(sc)
A <- names(sc@cpart)[sc@cpart %in% c(1,2)]
B <- names(sc@cpart)[sc@cpart %in% c(3)]
y <- diffexpnb(getfdata(sc,n=c(A,B)), A=A, B=B)
plotdiffgenesnb(y)</pre>
```

plotdimsat

Plotting the Saturation of Explained Variance

Description

This functions plots the explained variance as a function of PCA/ICA components computed by the function CCcorrect. The number of components where the change in explained variability upon adding further components approaches linear behaviour demarcates the saturation point and is highlighted in blue.

Usage

```
plotdimsat(object, change = TRUE, lim = NULL)
```

Arguments

object SCseq class object.

change logical. If TRUE then the change in explained variance is plotted. Default is

FALSE and the explained variance is shown.

lim Number of components included for he calculation and shown in the plot. De-

fault is NULL and all components are included.

Value

None

plotdistanceratio 43

plotdistanceratio

Histogram of Cell-to-Cell Distances in Real versus Embedded Space

Description

This function plots a histogram of the ratios of cell-to-cell distances in the original versus the high-dimensional embedded space used as input for the StemID2 inferences. The embedded space approximates correlation-based distances by Euclidean distances obtained by classical multi-dimensional scaling. A minimum spanning tree of the cluster centers is overlaid for comparison.

Usage

```
plotdistanceratio(object)
```

Arguments

object

Ltree class object.

Value

None.

plotexpmap

Highlighting gene expression in a dimensional reduction representation

Description

This functions highlights gene expression in a two-dimensional t-SNE map, UMAP, or a Fruchterman-Rheingold graph layout of the singe-cell transcriptome data.

Usage

```
plotexpmap(
  object,
  g,
  n = NULL,
  logsc = FALSE,
  imputed = FALSE,
  fr = FALSE,
  um = FALSE,
  cells = NULL,
  cex = 0.5,
  map = TRUE,
  leg = TRUE,
  noise = FALSE
)
```

44 plotgraph

Arguments

object	SCseq class object.
g	Individual gene name or vector with a group of gene names corresponding to a subset of valid row names of the ndata slot of the SCseq object.
n	String of characters representing the title of the plot. Default is NULL and the first element of g is chosen.
logsc	logical. If TRUE, then gene expression values are log2-transformed after adding a pseudo-count of 0.1. Default is FALSE and untransformed values are shown.
imputed	logical. If TRUE and imputing was done by calling compdist with knn > 0, then imputed expression values are shown. If FALSE, then raw counts are shown. Default is FALSE.
fr	logical. If TRUE then plot Fruchterman-Rheingold layout. Default is FALSE.
um	logical. If TRUE then plot umap dimensional reduction representation. Default is FALSE.
cells	Vector of valid cell names corresponding to column names of slot ndata of the SCseq object. Gene expression is ony shown for this subset.
cex	size of data points. Default value is 0.5.
map	logical. If TRUE then data points are shown. Default value is TRUE.
leg	logical. If TRUE then the legend is shown. Default value is TRUE.
noise	logical. If TRUE then display local gene expression variability instead of gene expression (requires VarID analysis)/ Default value is FALSE.

Value

None

	G. IDAI: G. I	
plotgraph	StemID2 Lineage Graph	

Description

This function plots a graph of lineage trajectories connecting RaceID3 cluster medoids as inferred by StemID2 to approximate the lineage tree. The plot highlights significant links, where colour indicates the level of significance and width indicates the link score. The node colour reflects the level of transcriptome entropy.

Usage

```
plotgraph(
  object,
  showCells = FALSE,
  showMap = TRUE,
  tp = 0.5,
  scthr = 0,
  cex = 1
)
```

plotjaccard 45

Arguments

object Ltree class object.

showCells logical. If TRUE, then projections of cells are shown in the plot. Default is FALSE.

showMap logical. Tf TRUE, then show transparent t-SNE map (with transparency tp) of

cells in the background. Default is TRUE.

tp Real number between zero and one. Level of transparency of the t-SNE map.

Deafault is 0.5. See showMap.

sethr Real number between zero and one. Score threshold for links to be shown in the

graph. For scthr=0 all significant links are shown. The maximum score is one.

Default is 0.

cex real positive number. Size of data points. Deault is 1.

Value

None.

plotjaccard	Plot Jaccard Similarities	

Description

This functions plots a barchart of Jaccard similarities for the RaceID3 clusters before outlier identification

Usage

```
plotjaccard(object)
```

Arguments

object SCseq class object.

Value

None

46 plotlinkpv

plotlabelsmap Plot labels in a dimensional reduction representation	
---	--

Description

This functions plots cell labels into a two-dimensional t-SNE map, UMAP, or a Fruchterman-Rheingold graph layout of the singe-cell transcriptome data.

Usage

```
plotlabelsmap(object, labels = NULL, fr = FALSE, um = FALSE, cex = 0.5)
```

Arguments

object	SCseq class object.
labels	Vector of labels for all cells to be highlighted in the t-SNE map. The order has to be the same as for the columns in slot ndata of the SCseq object. Default is NULL and cell names are highlighted.
fr	logical. If TRUE then plot Fruchterman-Rheingold layout. Default is FALSE.
um	logical. If TRUE then plot umap dimensional reduction representation. Default is FALSE.
cex	positive real number. Size of the labels. Default is 0.5.

Value

None

|--|

Description

This function plots a heatmap of link p-values.

Usage

```
plotlinkpv(object)
```

Arguments

object Ltree class object.

Value

None.

plotlinkscore 47

|--|

Description

This function plots a heatmap of link score.

Usage

```
plotlinkscore(object)
```

Arguments

object Ltree class object.

Value

None.

plotmap	Plotting a dimensional reduction representation

Description

This functions plots a two-dimensional t-SNE map, UMAP, or a Fruchterman-Rheingold graph layout of the singe-cell transcriptome data.

Usage

```
plotmap(object, final = TRUE, tp = 1, fr = FALSE, um = FALSE, cex = 0.5)
```

Arguments

object	SCseq class object.
final	logical. If TRUE, then highlight final clusters after outlier identification. If FALSE, then highlight initial clusters prior to outlier identification. Default is TRUE.
tp	Number between 0 and 1 to change transparency of dots in the map. Default is 1.
fr	logical. If TRUE then plot Fruchterman-Rheingold layout. Default is FALSE.
um	logical. If TRUE then plot umap dimensional reduction representation. Default is FALSE.
cex	size of data points. Default value is 0.5.

Value

None

48 plotmarkergenes

plotmarkergenes

Plotting a Heatmap of Marker Gene Expression

Description

This functions generates a heatmap of expression for defined group of genes and can highlight the clustering partition and another sample grouping, e.g. origin or cell type.

Usage

```
plotmarkergenes(
  object,
  genes,
  imputed = FALSE,
  cthr = 0,
  cl = NULL,
  cells = NULL,
  order.cells = FALSE,
  aggr = FALSE,
  norm = FALSE,
  cap = NULL,
  flo = NULL,
  samples = NULL,
  cluster_cols = FALSE,
  cluster_rows = TRUE,
  cluster_set = FALSE,
  samples_col = NULL,
  zsc = FALSE,
  logscale = TRUE,
  noise = FALSE,
  fontsize = 10
)
```

Arguments

object	SCseq class object.
genes	A vector with a group of gene names corresponding to a subset of valid row names of the ndata slot of the SCseq object.
imputed	logical. If TRUE and imputing was done by calling compdist with knn $>$ 0, then imputed expression values are shown. If FALSE, then raw counts are shown. Default is FALSE
cthr	Interger number greater or equal zero. Only clusters with $>$ cthr cells are included in the t-SNE map. Default is 0 .
cl	Vector of valid cluster numbers contained in slot cpart of the SCseq object. Default is NULL and all clusters with >cthr cells are included.

plotNoiseModel 49

cells	Vector of valid cell names corresponding to column names of slot ndata of the SCseq object. Gene expression is only shown for this subset. Default is NULL and all cells are included. The set of cells is intersected with the subset of clusters in cl if given.
order.cells	logical. If TRUE, then columns of the heatmap are ordered by cell name and not by cluster number. If cells are given, then columns are ordered as in cells.
aggr	logical. If TRUE, then only average expression is shown for each cluster. Default is FALSE and expression in individual cells is shown.
norm	logical. If TRUE, then expression of each gene across clusters is normalized to 1, in order to depict all genes on the same scale. Default is FALSE.
сар	Numeric. Upper bound for gene expression. All values larger then cap are replaced by cap. Default is NULL and no cap is applied.
flo	Numeric. Lower bound for gene expression. All values smaller then flo are replaced by flo. Default is NULL and no flo is applied.
samples	A vector with a group of sample names for each cell in the same order as the column names of the ndata slot of the SCseq object.
cluster_cols	logical. If TRUE, then columns are clustered. Default is FALSE.
cluster_rows	logical. If TRUE, then rows are clustered. Default is TRUE.
cluster_set	logical. If TRUE then clusters are ordered by hierarchical clustering of the cluster medoids.
samples_col	Vector of colors used for highlighting all samples contained in samples in the heatmap. Default is NULL.
zsc	logical. If TRUE then a z-score transformation is applied. Default is FALSE.
logscale	logical. If TRUE then a log2 transformation is applied. Default is TRUE.
noise	logical. If TRUE then display local gene expression variability instead of gene expression (requires VarID analysis)/ Default value is FALSE.
fontsize	postive real number. Font size of gene name labels. Default is 10.

Value

Object with clustering information for rows and columns returned by the function pheatmap from the package **pheatmap**.

plotNoiseModel	Function for plotting the baseline model of gene expression variability

Description

This function plots the variance against mean expression across all genes and a second order polynomial to the base line of the variance-mean dependence in log space.

Usage

```
plotNoiseModel(x, corrected = FALSE)
```

50 plotPearsonRes

Arguments

x List object returned by function noiseBaseFit or function compNoise.

corrected logical value. If TRUE, then the variance is plotted after regressing our the mean

dependence.

Value

None

Examples

```
x <- noiseBaseFit(intestinalDataSmall,step=.01,thr=.05)
plotNoiseModel(x)</pre>
```

plotoutlierprobs

Plot Outlier Probabilities

Description

This functions plots a barchart of outlier probabilities across all cells in each cluster.

Usage

```
plotoutlierprobs(object)
```

Arguments

object

SCseq class object.

Value

None

plotPearsonRes

Function for plotting the variance of Pearson residuals

Description

This function plots the variance versus the mean of the Pearson residuals obtained by the negative binomial regression computed by the function compNoise if regNB is TRUE. A local regression is also shown.

Usage

```
plotPearsonRes(noise, log = FALSE)
```

plotRegNB 51

Arguments

noise List object with the background noise model and a variability matrix, returned

by the compNoise function.

logical. If TRUE then the y-axis is log-transformed. Default is FALSE.

Value

None

Examples

```
res <- pruneKnn(intestinalDataSmall,metric="pearson",knn=10,alpha=1,no_cores=1,FSelect=FALSE)
noise <- compNoise(intestinalDataSmall,res,pvalue=0.01,genes = NULL,no_cores=1)
plotPearsonRes(noise,log=TRUE)</pre>
```

plotRegNB

Function for plotting negative binomial regression

Description

This function plots the parameters obtained by the negative binomial regression of the transcript counts on the total transcript count in each cells. Smoothed parameter estimates are also shown.

Usage

```
plotRegNB(expData, noise, par.nb = NULL)
```

Arguments

expData	Matrix of gene expression values with genes as rows and cells as columns. The
	matrix need to contain the same call IDs as solumns like the input matrix used

matrix need to contain the same cell IDs as columns like the input matrix used to derive the pruned k nearest neighbours with the pruneKnn function.

noise List object with the background noise model and a variability matrix, returned

by the compNoise function.

par.nb Parameter to be plotted, i.e. valid column of noise\$regData\$nbRegr. of the

log10 total UMI count. intercept is the intercept inferred by the regression.

Default is NULL and theta is shown.

Value

None

Examples

res <- pruneKnn(intestinalDataSmall,metric="pearson",knn=10,alpha=1,no_cores=1,FSelect=FALSE)
noise <- compNoise(intestinalDataSmall,res,regNB=TRUE,pvalue=0.01,genes = NULL,no_cores=1)
plotRegNB(intestinalDataSmall,noise,"theta")</pre>

52 plotsensitivity

plotsaturation

Plot Saturation of Within-Cluster Dispersion

Description

This functions plots the (change in the) mean within-cluster dispersion as a function of the cluster number and highlights the saturation point inferred based on the saturation criterion applied by RaceID3: The number of clusters where the change in within-cluster dispersion upon adding further clusters approaches linear behaviour demarcates the saturation point and is highlighted in blue.

Usage

```
plotsaturation(object, disp = FALSE)
```

Arguments

object SCseq class object.

disp logical. If FALSE, then the change of the within-cluster dispersion is plotted. if

TRUE the actual dispersion is plotted. Default is FALSE

Value

None

plotsensitivity

Plot Sensitivity

Description

This functions plots the number of outliers as a function of the outlier probability.

Usage

```
plotsensitivity(object)
```

Arguments

object

SCseq class object.

Value

None

plotsilhouette 53

Cluster Silhouette

Description

This functions produces a silhouette plot for RaceID3 clusters prior or post outlier identification.

Usage

```
plotsilhouette(object, final = FALSE)
```

Arguments

object SCseq class object.

final logical. If TRUE, then plot silhouette coefficients for final clusters after outlier

identification. Default is FALSE and silhouette coefficients are plotted for initial

clusters.

Value

None

plotspantree	Minimum Spanning Tree of RaceID3 clusters	

Description

This function plots a minimum spanning tree of the RaceID3 cluster medoids in a two-dimensional reduction representation.

Usage

```
plotspantree(object, tp = 0.5, cex = 1, projections = FALSE)
```

Arguments

object Ltree class object.

tp Real number between zero and one. Level of transparency of the t-SNE map.

Deafault is 0.5.

cex real positive number. Size of data points. Deault is 1.

projections logical. If TRUE, then the projections of the cells onto the inter-medoid links as

computed by StemID are shown. Default is FALSE

Value

None.

54 plotsymbolsmap

plotsymbolsmap Plotting groups as different symbols in a dimensional reduction representation

Description

This functions highlights groups of cells by different symbols in a two-dimensional t-SNE map, UMAP, or a Fruchterman-Rheingold graph layout of the singe-cell transcriptome data.

Usage

```
plotsymbolsmap(
  object,
  types,
  subset = NULL,
  samples_col = NULL,
  cex = 0.5,
  fr = FALSE,
  um = FALSE,
  leg = TRUE,
  map = TRUE
)
```

Arguments

object	SCseq class object.
types	Vector assigning each cell to a type to be highlighted in the t-SNE map. The order has to be the same as for the columns in slot ndata of the SCseq object. Default is NULL and each cell is highlighted by a different symbol.
subset	Vector containing a subset of types from types to be highlighted in the map. Default is NULL and all types are shown.
samples_col	Vector of colors used for highlighting all samples contained in samples in the map. Default is NULL.
cex	size of data points. Default value is 0.5.
fr	logical. If TRUE then plot Fruchterman-Rheingold layout. Default is FALSE.
um	logical. If TRUE then plot umap dimensional reduction representation. Default is FALSE.
leg	logical. If TRUE then the legend is shown. Default value is TRUE.
map	logical. If TRUE then data points are shown. Default value is TRUE.

Value

None

plotTrProbs 55

plotTrProbs	Function for plotting transition probabilities between clusters	

Description

This function plots the transitions probabilities in a dimensional reduction representation of a **RaceID** SCseq object updates with the updateSC function. in order to utilize **RaceID** functions for visualization.

Usage

```
plotTrProbs(
  object,
  probs,
  tp = 0.5,
  prthr = 0,
  cthr = 0,
  fr = FALSE,
  um = FALSE,
  cex = 1
)
```

Arguments

object	RaceID SCseq object, updated with the updateSC function.
probs	$Matrix\ of\ transition\ probabilities\ between\ clusters,\ returned\ by\ the\ transition Probs\ function.$
tp	Positive real number between 0 and 1. Transparency of the data points in the dimensional reduction map. Default is 0.5.
prthr	Positive real number between 0 and 1. Threshold of transition probabilities. Only transitions with probability >prthr are displayed in the map. Default is 0.
cthr	Integer number greater or equal 0 defining the minimum clusters size for inclusion into the map. Default is 0.
fr	logical. If TRUE, then a Fruchterman-Rheingold graph layout is shown (in case it has been computed for the RaceID bject), otherwise a t-SNE map is shown. Default is FALSE.
um	logical. If TRUE then plot umap dimensional reduction representation. Default is FALSE.
cex	real positive number. Size of data points. Default is 1.

Value

None

56 projback

Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
d <- getExpData(sc)
res <- pruneKnn(d,distM=sc@distances,metric="pearson",knn=10,alpha=1,no_cores=1,FSelect=FALSE)
cl <- graphCluster(res,pvalue=0.01)
sc <- updateSC(sc,res=res,cl=cl)
sc <- comptsne(sc)
probs <-transitionProbs(res,cl,pvalue=0.01)
plotTrProbs(sc,probs,tp=.5,prthr=0,cthr=0,fr=FALSE)</pre>
```

projback

Compute Cell Projections for Randomized Background Distribution

Description

This function computes the projections of cells onto inter-cluster links for randomized cell positions in a high-dimensional embedded space. Significance of link based on an increased number of cells on a link is inferred based on this background model.

Usage

```
projback(object, pdishuf = 500, fast = FALSE, rseed = 17000, verbose = TRUE)
```

Arguments

object	Ltree class object.
pdishuf	Number of randomizations of cell positions for which to compute projections of cells on inter-cluster links. Default is 2000. No randomizations are needed in this mode and the function will do nothing. Default is TRUE.
fast	logical. If TRUE and nmode=FALSE cells will still be assigned to links based on maximum projections but a fast approximate background model will be used to infer significance. The function will do nothing in this case. Default is FALSE.
rseed	Integer number used as seed to ensure reproducibility of randomizations. Defaut is 17000.
verbose	logical. If FALSE then status output messages are disabled. Default is TRUE.

Value

An Ltree class object with all information on randomized cell projections onto links stored in the prbacka slot.

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Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
sc <- clustexp(sc)
sc <- findoutliers(sc)
sc <- comptsne(sc)
ltr <- Ltree(sc)
ltr <- compentropy(ltr)
ltr <- projcells(ltr,nmode=FALSE)
ltr <- projback(ltr,pdishuf=50)</pre>
```

projcells

Compute transcriptome entropy of each cell

Description

This function computes the projections of cells onto inter-cluster links in a high-dimensional embedded space.

Usage

```
projcells(object, cthr = 5, nmode = TRUE, knn = 3, fr = FALSE, um = FALSE)
```

Arguments

object	Ltree class object.
cthr	Positive integer number. Clusters to be included into the StemID2 analysis must contain more than cthr cells. Default is 5.
nmode	logical. If TRUE, then a cell of given cluster is assigned to the link to the cluster with the smallest average distance of the knn nearest neighbours within this cluster. Default is TRUE.
knn	Positive integer number. See nmode. Default is 3.
fr	logical. Use Fruchterman-Rheingold layout instead of t-SNE for dimensional-reduction representation of the lineage graph. Default is FALSE.
um	logical. Use umap representation instead of t-SNE for dimensional-reduction representation of the lineage graph. Default is FALSE.

Value

An Ltree class object with all information on cell projections onto links stored in the 1data slot.

58 pruneKnn

Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
sc <- clustexp(sc)
sc <- findoutliers(sc)
sc <- comptsne(sc)
ltr <- Ltree(sc)
ltr <- compentropy(ltr)
ltr <- projeclls(ltr)</pre>
```

projenrichment

Enrichment of cells on inter-cluster links

Description

This function plots a heatmap of the enrichment ratios of cells on significant links.

Usage

```
projenrichment(object)
```

Arguments

object

Ltree class object.

Value

None.

pruneKnn

Function inferring a pruned knn matrix

Description

This function determines k nearest neighbours for each cell in gene expression space, and tests if the links are supported by a negative binomial joint distribution of gene expression. A probability is assigned to each link which is given by the minimum joint probability across all genes.

59 pruneKnn

Usage

```
pruneKnn(
  expData,
  distM = NULL,
  large = TRUE,
  regNB = TRUE,
  batch = NULL,
  regVar = NULL,
  ngenes = 2000,
  span = 0.75,
  pcaComp = 100,
  algorithm = "kd_tree",
 metric = "pearson",
  genes = NULL,
  knn = 10,
  alpha = NULL,
  no_cores = NULL,
  FSelect = FALSE,
  seed = 12345,
  res = NULL
)
```

Arguments

expData

Matrix of gene expression values with genes as rows and cells as columns. These values have to correspond to unique molecular identifier counts.

distM

Optional distance matrix used for determining k nearest neighbours. Default is NULL and the distance matrix is computed using a metric given by the parameter metric.

large

logical. If TRUE then no distance matrix is required and nearest neighbours are inferred by the FNN package based on a reduced feature matrix computed by a principle component analysis. Only the first pcaComp principle components are considered. Prior to principal component analysis a negative binomial regression is performed to eliminate the dependence on the total number of transcripts per cell. The pearson residuals of this regression serve as input for the principal component analysis after smoothing the parameter dependence on the mean by a loess regression. Deafult is TRUE. Recommended mode for very large datasets, where a distance matrix consumes too much memory. A distance matrix is no longer required, and if distM is initialized it will be ignored if large equals TRUE.

logical. If TRUE then gene a negative binomial regression is performed to prior to the principle component analysis if large = TRUE. See large. Default is TRUE.

batch

vector of batch variables. Component names need to correspond to valid cell IDs, i.e. column names of expData. If regNB is TRUE, than the batch variable will be regressed out simultaneously with the log10 UMI count per cell. An interaction term is included for the log10 UMI count with the batch variable. Default value is NULL.

regNB

60 pruneKnn

regVar data.frame with additional variables to be regressed out simultaneously with

the log10 UMI count and the batch variable (if batch is TRUE). Column names indicate variable names (name beta is reserved for the coefficient of the log10 UMI count), and rownames need to correspond to valid cell IDs, i.e. column names of expData. Interaction terms are included for each variable in regVar

with the batch variable (if batch is TRUE). Default value is NULL.

ngenes Positive integer number. Randomly sampled number of genes (from rownames

of expData) used for predicting regression coefficients (if regNB=TRUE). Smoothed

coefficients are derived for all genes. Default is 2000.

span Positive real number. Parameter for loess-regression (see large) controlling the

degree of smoothing. Default is 0.75.

pcaComp Positive integer number. Number of princple components to be included if

large is TRUE. Default is 100.

from the **FNN** package. See help(get.knn). Deafult is "kd_tree".

metric Distances are computed from the expression matrix x after optionally includ-

ing only genes given as argument genes or after optional feature selection (see FSelect). Possible values for metric are "pearson", "spearman", "logpearson", "euclidean". Default is "pearson". In case of the correlation based methods,

the distance is computed as 1 – correlation.

genes Vector of gene names corresponding to a subset of rownames of x. Only these

genes are used for the computation of a distance matrix and for the computation of joint probabilities of nearest neighbours. Default is NULL and all genes are

used.

knn Positive integer number. Number of nearest neighbours considered for each cell.

Default is 10.

alpha Positive real number. Relative weight of a cell versus its k nearest neigbour ap-

plied for the derivation of joint probabilities. A cell receives a weight of alpha while the weight of its k nearest neighbours is determined by quadratic programming. The sum across all weights is normalized to one, and the weighted mean expression is used for computing the joint probability of a cell and each of its k nearest neighbours. These probabilities are used for the derivation of of link probabilities. Larger values give more weight to the gene expression observed in a cell versus its neighbourhood. Typical values should be in the range of 0 to 10. Default is NULL. In this case, alpha is inferred by an optimization, i.e., alpha is minimized under the constraint that the gene expression in a cell does not deviate more then one standard deviation from the predicted weighted mean, where the standard deviation is calculated from the predicted mean using the background model (the average dependence of the variance on the mean

expression).

no_cores Positive integer number. Number of cores for multithreading. If set to NULL then

the number of available cores minus two is used. Default is 1.

FSelect Logical parameter. If TRUE, then feature selection is performed prior to distance

matrix calculation and VarID analysis. Default is FALSE.

seed Integer number. Random number to initialize stochastic routines. Default is

12345.

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res

Output object from pruneKnn. The rownames (genes) and colnames (cells) of the parameter expData have to be subsets on the input data used to produce this output. For example, the batch effects could have been corrected on the global dataset using the pruneKnn function, and using the output from the global run permits using regression parameters from the global analysis on specific subsets if expData contain a subset of genes and cells.

Value

List object of six components:

distM Distance matrix.

dimRed PCA transformation of expData including the first pcaComp principle compo-

nents, computed on including genes or variable genes only if Fselect equals

TRUE. Is is set to NULL if large equals FALSE.

pvM Matrix of link probabilities between a cell and each of its k nearest neighbours.

Column i shows the k nearest neighbour link probabilities for cell i in matrix x.

NN Matrix of column indices of k nearest neighbours for each cell according to input

matrix x. First entry corresponds to index of the cell itself. Column i shows the

k nearest neighbour indices for cell i in matrix x.

B List object with background model of gene expression as obtained by fitBackVar

function.

regData If regNB=TRUE this argument contains a list of four components: component

pearsonRes contains a matrix of the Pearson Residual computed from the negative binomial regression, component nbRegr contains a matrix with the regression coefficients, component nbRegrSmooth contains a matrix with the smoothed regression coefficients, and log10_umi is a vector with the total log10 UMI count for each cell. The regression coefficients comprise the dispersion parameter theta, the intercept, the regression coefficient beta for the log10 UMI count,

and the regression coefficients of the batches (if batch is not NULL).

Examples

res <- pruneKnn(intestinalDataSmall,metric="pearson",knn=10,alpha=1,no_cores=1,FSelect=FALSE)

rcpp_hello_world Simple function using Rcpp

Description

Simple function using Rcpp

Usage

rcpp_hello_world()

62 rfcorrect

Examples

```
## Not run:
rcpp_hello_world()
## End(Not run)
```

rfcorrect

Random Forests-based Reclassification

Description

This functions applies random forests-based reclassification of cell clusters to enhance robustness of the final clusters.

Usage

```
rfcorrect(
  object,
  rfseed = 12345,
  nbtree = NULL,
  final = TRUE,
  nbfactor = 5,
   ...
)
```

Arguments

object SCseq class object.

rfseed Seed for enforcing reproducible results. Default is 12345.

Number of trees to be built. Default is NULL and the number of tree is given by the number of cells times nbfactor.

final logical. If TRUE, then reclassification of cell types using out-of-bag analysis is performed based on the final clusters after outlier identification. If FALSE, then the cluster partition prior to outlier identification is used for reclassification.

Positive integer number. See nbtree.

additional input arguments to the randomForest function of the randomForest package.

Value

The function returns an updated SCseq object with random forests votes written to slot out\$rfvotes. The clustering partition prior or post outlier identification (slot cluster\$kpart or cpart, if parameter final equals FALSE or TRUE, respectively) is overwritten with the partition derived from the reclassification.

SCseq 63

Examples

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
sc <- clustexp(sc)
sc <- findoutliers(sc)
sc <- rfcorrect(sc)</pre>
```

SCseq

The SCseq Class

Description

The SCseq class is the central object storing all information generated during cell type identification with the RaceID3 algorithm. It comprises a number of slots for a variety of objects.

Arguments

object

An SCseq object.

Slots

expdata The raw expression data matrix with cells as columns and genes as rows in sparse matrix format.

ndata Filtered data with expression normalized to one for each cell.

noise Matrix with local gene expression noise estimates (used for VarID analysis)

counts Vector with total transcript counts for each cell in ndata remaining after filtering.

genes Vector with gene names of all genes in ndata remaining after filtering.

dimRed list object object storing information on a feature matrix obtained by dimensional reduction, batch effect correction etc. Component x stores the actual feature matrix.

distances distance (or dis-similarity) matrix computed by RaceID3.

imputed list with two matrices computed for imputing gene expression. The first matrix nn contains the cell indices of the knn nearest neighbours, the second matrix contains the probabilities at which each cell contributes to the imputed gene expression value for the cell correponding to the columns.

tsne data.frame with coordinates of two-dimensional tsne layout computed by RaceID3.

fr data.frame with coordinates of two-dimensional Fruchterman-Rheingold graphlayout computed by RaceID3.

umap data.frame with coordinates of two-dimensional umap representation computed by RaceID3.

cluster list storing information on the initial clustering step of the RaceID3 algorithm

background list storing the polynomial fit for the background model of gene expression variability computed by RaceID3, which is used for outlier identification.

out list storing information on outlier cells used for the prediction of rare cell types by RaceID3

64 transitionProbs

cpart vector containing the final clustering (i.e. cell type) partition computed by RaceID3 fcol vector containing the colour scheme for the RaceID3 clusters medoids vector containing the cell ids for th cluster medoids filterpar list containing the parameters used for cell and gene filterung clusterpar list containing the parameters used for clustering outlierpar list containing the parameters used for outlier identification

${\it transition Probs} \qquad {\it Function for the computation of transition probabilities between clusters}$		transitionProbs		
---	--	-----------------	--	--

Description

This function computes transition probabilities between clusters based on the link probabilities of the pruned k nearest neighbour graph.

Usage

```
transitionProbs(res, cl, pvalue = 0.01)
```

Arguments

res -	List object with k nearest neighbour information returned by pruneKnn function.
cl	List object with Louvain clustering information, returned by the graphCluster function.
pvalue	Positive real number between 0 and 1. All nearest neighbours with link probability < pvalue are discarded. Default is 0.01.

Value

Matrix of transition probabilities between clusters.

```
res <- pruneKnn(intestinalDataSmall,metric="pearson",knn=10,alpha=1,no_cores=1,FSelect=FALSE)
cl <- graphCluster(res,pvalue=0.01)
probs <-transitionProbs(res,cl,pvalue=0.01)</pre>
```

updateSC 65

VarID results
VarID results

Description

This function updates a **RaceID** SCseq object with a distance matrix or dimensionally reduced feature matrix, a clustering partition, and/or a matrix of gene expression variability, in order to utilize **RaceID** functions for visualization.

Usage

```
updateSC(object, res = NULL, cl = NULL, noise = NULL, flo = NULL)
```

Arguments

object	RaceID SCseq object.
res	List object returned by pruneKnn function to update SCseq with distance matrix and/or dimensionally reduced feature matrix in res. Default is NULL
cl	List object with Louvain clustering information, returned by the graphCluster function to update SCseq object with clustering partition and Fruchterman-Rheingold layout. Default is NULL.
noise	List object with the background noise model and a variability matrix, returned by the compNoise function, to update SCseq object with a noise matrix. Default is NULL.
flo	Real number. Lower cutoff for the gene expression variability. All values < flo in the variability matrix are set to this level. Default is NULL and values are not reset.

Value

SCseq object with a distance matrix (slot distances) and a dimensionally reduced feature matrix (slot dimRed\$x), or clustering partition (slot cpart and cluster\$kpart) derived from the VarID analysis, and/or with a gene expression variability matrix in slot noise.

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
sc <- compdist(sc)
d <- getExpData(sc)
res <- pruneKnn(d,distM=sc@distances,metric="pearson",knn=10,alpha=1,no_cores=1,FSelect=FALSE)
cl <- graphCluster(res,pvalue=0.01)
sc <- updateSC(sc,res=res,cl=cl)
sc <- comptsne(sc)
plotmap(sc)</pre>
```

varRegression

varRegression Linear Regression of Sources of Variability

Description

This functions regresses out variability associated with particular sources.

Usage

```
varRegression(object, vars = NULL, logscale = FALSE, Batch = FALSE)
```

Arguments

object SCseq class object.

vars data.frame of variables to be regressed out. Each column corresponds to a vari-

able and each variable corresponds to a cell. The object must contain all cells,

i.e. column names of the slot ndata from the SCseq object.

logical. If TRUE data are log-transformed prior to regression. Default is FALSE.

Batch logical. If TRUE, then the function will regress out batch-associated variability

based on genes stored in the filterpar\$BGenes slot of the SCseq object. This requires prior batch correction with the filterdata function using bmode="RaceID".

Value

The function returns an updated SCseq object with the corrected expression matrix written to the slot dimRed\$x of the SCseq object.

```
sc <- SCseq(intestinalDataSmall)
sc <- filterdata(sc)
b <- sub("(\\_\\d+)$","",colnames(intestinalData))
vars <- data.frame(row.names=colnames(intestinalData),batch=b)
sc <- varRegression(sc,vars)</pre>
```

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