Package 'SingleCellStat'

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Type Package

Title A Toolkit for Statistical Analysis of Single-Cell Omics Data

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Description

A suite of statistical methods for analysis of single-cell omics data including linear modelbased methods for differential abundance analysis for individual level single-cell RNAseq data. For more details see Zhang, et al. (Submitted to Bioinformatics)<https: //github.com/Lujun995/DiSC_Replication_Code>.

Depends R (>= 4.3.0)

Suggests knitr, rmarkdown

Imports matrixStats, Matrix, stats, utils, vegan

License GPL-3

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LazyData true

LazyDataCompression xz

NeedsCompilation no

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Contents

DiSC	 	2
sim_data	 	4

6

Index

DiSC

DiSC: A statistical tool for differential expression analyis of individual level single-cell RNA-Seq data

Description

A statistical tool for differential expression analyis of individual level single-cell RNA-Seq data

Usage

```
DiSC(data.mat, cell.ind, metadata, outcome, covariates = NULL,
    cell.id = "cell_id",
    individual.id = "individual", perm.no = 999,
    features = c('prev', 'm', 'nzsd'), verbose = TRUE,
    sequencing.data = TRUE)
```

Arguments

data.mat	A data matrix for single-cell RNA sequencing data, or other single-cell data such as CyToF data. Rows - genes/features, columns - cells. Column names are cell ids.	
cell.ind	A data frame of cell-individual relationship. It includes two columns for cell ids and individual ids. It links cell ids to individual ids.	
metadata	A data frame of individual-level metadata. It includes a column for individual ids, a column for the outcome of interest and columns for other covariates if applicable.	
outcome	A character string for the column name of the outcome variable in metadata.	
covariates	A character string or vector of character strings for the covariates to be adjusted. Should be the column names in metadata. Default: NULL.	
cell.id	A character string for the column name of cell ids in cell.ind.	
individual.id	A character string for the column name of the individual ids in cell.ind and metadata.	
perm.no	An integer, number of permutations used. Default: 999. It can be reduced to 99 if adjusted P-values (false discovery rate) are the only interest.	
features	Features of the distribution used to test for the differentially expressed genes. Choose from "prev" (logit(non-zero proportion), "nzm" (sqrt(non-zero mean)), "nzsd" (sqrt(non-zero standard deviation)), "m" (overall mean)), "sd" (overall standard deviation), "nzm^1", "nzsd^1", "m^1", "sd^1" (non-sqrt-transformed versions). Default: "prev", "m" and "nzsd".	
verbose	Logical. Should the function print the processes? Default: TRUE.	
sequencing.data		
	Logical. Is the data.mat a sequencing data matrix (i.e., count data)? If TRUE, the total sum scaling will be used to normalize the count data. The users can normalize/transform the data themselves by setting it to be FALSE. Default: TRUE.	

DiSC

Value

call How was the function called?
R2 Description of R2
F0 Description of F0
RSS Description of RSS
df.model Description of df.model
df.residual Description of df.residual
coef.list Description of coef.list
p.raw Raw, unadjusted P-values.
p.adj.fdr P-values which have been adjusted for false discovery rate.
p.adj.fwer P-values which have been adjusted for family-wise error rate.

Author(s)

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References

Zhang, L., Yang, L., Ren, Y., Zhang, S., Guan, W., & Chen, J. (Bioinformatics): DiSC: a Statistical Tool for Fast Differential Expression Analysis of Individual-level Single-cell RNA-seq Data.

Examples

```
set.seed(seed = 1234556)
data(sim_data)
count_matrix <- sim_data$count_matrix</pre>
meta_cell <- sim_data$meta_cell</pre>
gene_index <- sim_data$gene_index</pre>
meta_ind <- sim_data$meta_ind</pre>
obj1 <- DiSC(data.mat = count_matrix, cell.ind = meta_cell,</pre>
              metadata = meta_ind, outcome = "phenotype",
              covariates = "RIN", cell.id = "cell_id",
              individual.id = "individual", perm.no = 999,
              features = c('prev', 'm', 'nzsd'), verbose = TRUE,
              sequencing.data = TRUE)
# Type I error (the nominal level: 0.05)
mean(obj1$p.raw[gene_index$EE_index] <= 0.05)</pre>
# True positive rate (based on raw P-values, the higher the better.)
mean(obj1$p.raw[gene_index$mean_index] <= 0.05)</pre>
mean(obj1$p.raw[gene_index$var_index] <= 0.05)</pre>
mean(obj1$p.raw[gene_index$mean_var_index] <= 0.05)</pre>
# False discovery rate (the nominal level: 0.10)
sum(obj1$p.adj.fdr[gene_index$EE_index] <= 0.10)/</pre>
  sum(obj1$p.adj.fdr <= 0.10)</pre>
# True positive rate (based on FDR-adjusted P-values, the higher the better.)
mean(obj1$p.adj.fdr[gene_index$mean_index] <= 0.10)</pre>
```

```
mean(obj1$p.adj.fdr[gene_index$var_index] <= 0.10)</pre>
mean(obj1$p.adj.fdr[gene_index$mean_var_index] <= 0.10)</pre>
# By default, DiSC normalizes the scRNA-seq data using TSS (total sum scaling),
# adjusted for log median sequencing depths
# Other user-specified normalization methods can also be used:
# log2 transformed, adjusted for log median sequencing depth
# data_mat_log <- log2(data_mat+1)</pre>
# inds <- unique(meta_cell[["individual"]])</pre>
# meta_ind <- meta_ind[base::match(inds, meta_ind[["individual"]]), ]</pre>
# data_mat <- count_matrix</pre>
# depth <- colSums(data_mat)</pre>
# cell.list <- list()</pre>
# for (ind in inds)
   cell.list[[ind]] <- meta_cell[meta_cell$individual == ind, ][["cell_id"]]</pre>
#
# log_md_depth <- numeric(length = length(inds))</pre>
# names(log_md_depth) <- inds</pre>
# for(ind in inds)
    log_md_depth[ind] <- log(median(depth[cell.list[[ind]]]))</pre>
#
# meta_ind$log_md_depth <- log_md_depth</pre>
# obj_log <-</pre>
    DiSC(data.mat = data_mat_log, cell.ind = meta_cell,
#
         metadata = meta_ind, outcome = "phenotype",
#
         covariates = c("RIN", "log_md_depth"),
#
         cell.id = "cell_id", individual.id = "individual",
#
#
         perm.no = 999, verbose = FALSE,
         sequencing.data = FALSE, # sequencing.data needs to be FALSE
#
         features = c('prev', 'm', 'nzsd'))
#
# Size factor: DESeq2, adjusted for log median sequencing depths
# require(DESeq2)
# colData <- data.frame(condition = rep(meta_ind$phenotype, each = 375),</pre>
                         row.names = colnames(data_mat))
#
# dds <- DESeq2::DESeqDataSetFromMatrix(countData = data_mat + 1,</pre>
                                        # avoid every gene contains at least one zero
                                           colData = colData, design = ~ condition)
# dds <- DESeq2::estimateSizeFactors(dds)</pre>
# data_mat_des <- sweep(data_mat, 2, DESeq2::sizeFactors(dds), FUN = "/")</pre>
# obj_des <-</pre>
#
   DiSC(data.mat = data_mat_des, cell.ind = meta_cell,
         metadata = meta_ind, outcome = "phenotype",
#
         covariates = c("RIN", "log_md_depth"),
#
#
         cell.id = "cell_id", individual.id = "individual",
#
         perm.no = 999, verbose = FALSE,
#
         sequencing.data = FALSE, # sequencing.data needs to be FALSE
#
         features = c('prev', 'm', 'nzsd'))
```

4

sim_data

Description

This dataset was generated in the "Generate Simulation Datasets" step in the Parametric_simulation.rmd (https://github.com/Lujun995/DiSC_Replication_Code)

Usage

data("sim_data")

Format

It contains 12 cases and 12 controls, each with 375 cell replicates. The read depths of each cell replicate are well-balanced. A covariate called RIN (RNA Integrity Number) at the individual level is included in the dataset.

The dataset comprises a total of 1,000 genes. The signal density was 15%, with differences in mean, variance, and mean+variance (each at 5%). The ground truth of differential/equally expression genes are indicated by gene_index, including mean_index (genes with a difference in mean), var_index (genes with a difference in variance), mean_var_index (genes with a difference in both mean and variance), EE_index (otherwise (to estimate type-I error)).

A list of elements:

count_matrix A numeric count matrix.

meta_cell A data.frame of the metadata at the cell level.

meta_ind A data.frame of the metadata at the individual level.

gene_index A list of 4 numeric vectors representing the ground truth of the IDs of the differentially or equally expressed genes.

Source

Simulated in the "Generate Simulation Datasets" step in the Parametric_simulation.rmd (https://github.com/Lujun995/DiSC_Replication_Code)

Examples

data(sim_data)
str(sim_data)

Index

* **datasets** sim_data, 4

DiSC, 2

sim_data,4