

Package ‘Single.mTEC.Transcriptomes’

August 6, 2024

Type Package

Title Single Cell Transcriptome Data and Analysis of Mouse mTEC cells

Version 1.32.0

Date 2021-11-21

Author Alejandro Reyes

Maintainer Alejandro Reyes <alejandro.reyes.ds@gmail.com>

Description This data package contains the code used to analyse the single-cell RNA-seq and the bulk ATAC-seq data from the manuscript titled: Single-cell transcriptome analysis reveals coordinated ectopic-gene expression patterns in medullary thymic epithelial cells. This paper was published in Nature Immunology 16,933-941(2015). The data objects provided in this package has been pre-processed: the raw data files can be downloaded from ArrayExpress under the accession identifiers E-MTAB-3346 and E-MTAB-3624. The vignette of this data package provides a documented and reproducible workflow that includes the code that was used to generate each statistic and figure from the manuscript.

License LGPL

biocViews ExperimentData

VignetteBuilder knitr

Suggests DESeq2, GenomicRanges, GenomicFeatures, genefilter, statmod, gdata, RColorBrewer, ggplot2, gplots, cluster, clue, grid, gridExtra, ggbio, Gviz, geneplotter, matrixStats, pheatmap, BiocStyle, knitr, BiocParallel

RoxygenNote 5.0.1

NeedsCompilation no

git_url <https://git.bioconductor.org/packages/Single.mTEC.Transcriptomes>

git_branch RELEASE_3_19

git_last_commit 5f3c26d

git_last_commit_date 2024-04-30

Repository Bioconductor 3.19

Date/Publication 2024-08-06

Contents

Single.mTec.Transcriptomes-package	2
aireDependentSansom	3
biotype	3
biotypesHuman	3
cealCoexpression	4
corMatsNoMarker	4
deGenesNone	5
deGenesSansom	5
dxATAC	5
fantom	6
geneNames	6
geneNamesHuman	6
geneRanges	7
mTECdx	7
muc1Coexpression	7
nomarkerCellsClustering	8
percentsGG	8
permutationResults	8
scLVM_output	9
tras	9
Index	10

Single.mTec.Transcriptomes-package
Single-cell transcriptome data of medullary thymic epithelial cells

Description

This document contains all the code used to analyse the single-cell RNA-seq and the bulk ATAC-seq data from the manuscript by Brennecke et al, 2015. The purpose of this package is to provide full reproducibility of the results presented in the manuscript. This package provides a documented and reproducible workflow of the code that was used to generate each number and figure from the manuscript.

References

Brennecke et al. Single-cell transcriptome analysis reveals coordinated ectopic gene-expression patterns in medullary thymic epithelial cells. *Nature Immunology* 16,933-941 (2015)

aireDependentSansom	<i>List of Aire-dependent genes</i>
---------------------	-------------------------------------

Description

A character vector of ensembl gene identifiers defined by Sansom et al, 2014. This list was downloaded from the supplementary material of the manuscript.

Usage

```
data(aireDependentSansom)
```

References

Sansom et al. Population and single-cell genomics reveal the Aire dependency, relief from Polycomb silencing, and distribution of self-antigen expression in thymic epithelia. *Genome Res.* 24, 1918–1931 (2014).

biotype	<i>Mouse biotypes</i>
---------	-----------------------

Description

A character vector defining biotype for each mouse ensembl gene identifier. Biotypes were queried using **biomaRt**.

Usage

```
data(biotypes)
```

biotypesHuman	<i>Human biotypes</i>
---------------	-----------------------

Description

A character vector defining biotype for each human ensembl gene identifier. Biotypes were queried using **biomaRt**.

Usage

```
data(biotypesHuman)
```

cea1Coexpression	<i>Cea1 co-expression group data</i>
------------------	--------------------------------------

Description

A data frame containing information about the Cea1 co-expression group defined in Pinto et al, 2013.

Usage

```
data(cea1Coexpression)
```

References

Pinto et al. Overlapping gene coexpression patterns in human medullary thymic epithelial cells generate self-antigen diversity. Proc. Natl. Acad. Sci. U.S.A. 110, E3497–3505, (2013).

corMatsNoMarker	<i>Gene-gene correlation network</i>
-----------------	--------------------------------------

Description

A matrix containing the gene-gene Spearman correlation across single unselected cells.

A matrix containing the gene-gene Spearman correlation across single unselected cells, using the data from Sansom et al, 2014.

Usage

```
data(corMatsNoMarker)
```

```
data(corMatsSansom)
```

References

Sansom et al. Population and single-cell genomics reveal the Aire dependency, relief from Polycomb silencing, and distribution of self-antigen expression in thymic epithelia. Genome Res. 24, 1918–1931 (2014).

deGenesNone	<i>Highly variable genes.</i>
-------------	-------------------------------

Description

List of highly variable genes according to the method by Brennecke et al, 2013.

Usage

```
data(deGenesNone)
```

References

Brennecke et al. Accounting for technical noise in single-cell RNA-seq experiments. Nat. Methods. 10, 1093-1095 (2013).

deGenesSansom	<i>Highly variable genes from Sansom et al.</i>
---------------	---

Description

List of highly variable genes according to the method by Brennecke et al, 2013, using the data by Santom et al.

Usage

```
data(deGenesSansom)
```

References

Brennecke et al. Accounting for technical noise in single-cell RNA-seq experiments. Nat. Methods. 10, 1093-1095 (2013). Sansom et al. Population and single-cell genomics reveal the Aire dependency, relief from Polycomb silencing, and distribution of self-antigen expression in thymic epithelia. Genome Res. 24, 1918–1931 (2014).

dxdATAC	<i>ATAC-seq summarized counts.</i>
---------	------------------------------------

Description

DESeqDataSet object summarizing the ATAC-seq data presented in the manuscript. This object contains the read counts of each sample over a window of 4Kb around transcription start sites.

Usage

```
data(dxdATAC)
```

fantom	<i>FANTOM dataset</i>
--------	-----------------------

Description

DESeqDataSet object of the read counts from selected tissues from the FANTOM dataset.

Usage

```
data(fantom)
```

References

Forrest et al. A promoter-level mammalian expression atlas. 24, 1918–1931. (2014).

geneNames	<i>Mouse gene names</i>
-----------	-------------------------

Description

Character vector of mouse gene names. The gene names were queried using **biomaRt**.

Usage

```
data(geneNames)
```

geneNamesHuman	<i>Human gene names</i>
----------------	-------------------------

Description

Character vector of human gene names. The gene names were queried using **biomaRt**.

Usage

```
data(geneNamesHuman)
```

geneRanges	<i>Genomic range coordinates.</i>
------------	-----------------------------------

Description

GenomicRanges object containing the coordinate ranges of mouse protein coding genes used in the manuscript.

Usage

```
data(geneRanges)
```

mTECdx	<i>Count data from the single-mTEC data.</i>
--------	--

Description

DESeqDataSet object containing the read counts from the single-cell RNA-seq dataset generated by Brennecke et al, 2015.

Usage

```
data(mTECdx)
```

References

Brennecke et al. Single-cell transcriptome analysis reveals coordinated ectopic gene-expression patterns in medullary thymic epithelial cells. *Nature Immunology* 16,933-941 (2015)

muc1Coexpression	<i>Muc1 co-expression group data</i>
------------------	--------------------------------------

Description

A data frame containing information about the Muc1 co-expression group defined in Pinto et al, 2013.

Usage

```
data(muc1Coexpression)
```

References

Pinto et al. Overlapping gene coexpression patterns in human medullary thymic epithelial cells generate self-antigen diversity. *Proc. Natl. Acad. Sci. U.S.A.* 110, E3497–3505, (2013).

nomarkerCellsClustering

Clustering results

Description

List containing the results from the clustering analysis from Figure 2 from the manuscript.

Usage

data(nomarkerCellsClustering)

percentsGG

Mapping statistics

Description

Data frame containing the mapping statistics from the single-cell RNA-seq data.

Usage

data(percentsGG)

permutationResults

Permutation results

Description

Result from the gene permutations from Figure 5 from the manuscript.

Usage

data(permutationResults)

scLVM_output	<i>Output from scLVM</i>
--------------	--------------------------

Description

Objects containing the output scLVM on the single-mTEC RNA-seq data.

Usage

```
data(scLVM_output)
```

References

Buettner et al. Computational analysis of cell-to-cell heterogeneity in single-cell RNA-sequencing data reveals hidden subpopulations of cells. Nat. Biotechnol. 2015.

tras	<i>Tissue restricted antigens lists.</i>
------	--

Description

Data frame containing the TRA classification from Pinto et al, 2013.

Usage

```
data(tras)
```

Index

* datasets

- aireDependentSansom, 3
 - biotype, 3
 - biotypesHuman, 3
 - cea1Coexpression, 4
 - corMatsNoMarker, 4
 - deGenesNone, 5
 - deGenesSansom, 5
 - dx dATAC, 5
 - fantom, 6
 - geneNames, 6
 - geneNamesHuman, 6
 - geneRanges, 7
 - mTECdx d, 7
 - muc1Coexpression, 7
 - nomarkerCellsClustering, 8
 - percentsGG, 8
 - permutationResults, 8
 - scLVM_output, 9
 - tras, 9
- aireDependentSansom, 3
- beta (scLVM_output), 9
- beta0 (scLVM_output), 9
- biotype, 3
- biotypesHuman, 3
- cea1Coexpression, 4
- corMatsNoMarker, 4
- corMatSp (corMatsNoMarker), 4
- corMatSpNoMarker (corMatsNoMarker), 4
- deGenesNone, 5
- deGenesSansom, 5
- dx d (mTECdx d), 7
- dx dATAC, 5
- dx dFANTOM (fantom), 6
- fantom, 6
- geneNames, 6
- geneNamesHuman, 6
- geneRanges, 7
- h5GeneNames (scLVM_output), 9
- heter (scLVM_output), 9
- mTECdx d, 7
- muc1Coexpression, 7
- nomarkerCellsClustering, 8
- percentsGG, 8
- permsAllClusters (permutationResults), 8
- permutationResults, 8
- realAllClusters (permutationResults), 8
- scLVM_output, 9
- Single.mTec.Transcriptomes-package, 2
- tras, 9
- vars (scLVM_output), 9
- Ycorr (scLVM_output), 9