

# Package ‘Single.mTEC.Transcriptomes’

April 13, 2017

**Type** Package

**Title** Single Cell Transcriptome Data and Analysis of Mouse mTEC cells

**Version** 1.2.1

**Date** 2015-01-30

**Author** Alejandro Reyes

**Maintainer** Alejandro Reyes <[alejandro.reyes@embl.de](mailto:alejandro.reyes@embl.de)>

**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

**RoxygenNote** 5.0.1

**NeedsCompilation** no

## R topics documented:

Single.mTec.Transcriptomes-package . . . . .	2
aireDependentSansom . . . . .	2
biotype . . . . .	3
biotypesHuman . . . . .	3
cea1Coexpression . . . . .	3
corMatsNoMarker . . . . .	4
deGenesNone . . . . .	4
deGenesSansom . . . . .	5
dxdATAC . . . . .	5
fantom . . . . .	5

geneNames . . . . .	6
geneNamesHuman . . . . .	6
geneRanges . . . . .	6
mTECdxd . . . . .	6
muc1Coexpression . . . . .	7
nomarkerCellsClustering . . . . .	7
percentsGG . . . . .	7
permutationResults . . . . .	8
scLVM_output . . . . .	8
tras . . . . .	8

**Index****9**

---

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    *List of Aire-dependent genes*

---

**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)
```

<i>mTECdx</i>	<i>Count data from the single-<i>mTEC</i> 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`      *Muc1 co-expression group data*

---

**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	<i>Permutation results</i>
--------------------	----------------------------

---

**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

\*Topic **datasets**

- aireDependentSansom, 2
- biotype, 3
- biotypesHuman, 3
- cea1Coexpression, 3
- corMatsNoMarker, 4
- deGenesNone, 4
- deGenesSansom, 5
- dxdATAC, 5
- fantom, 5
- geneNames, 6
- geneNamesHuman, 6
- geneRanges, 6
- mTECdx, 6
- muc1Coexpression, 7
- nomarkerCellsClustering, 7
- percentsGG, 7
- permutationResults, 8
- scLVM\_output, 8
- tras, 8

aireDependentSansom, 2

beta (scLVM\_output), 8

beta0 (scLVM\_output), 8

biotype, 3

biotypesHuman, 3

cea1Coexpression, 3

corMatsNoMarker, 4

corMatSp (corMatsNoMarker), 4

corMatSpNoMarker (corMatsNoMarker), 4

deGenesNone, 4

deGenesSansom, 5

dxd (mTECdx), 6

dxdATAC, 5

dxdFANTOM (fantom), 5

fantom, 5

geneNames, 6

geneNamesHuman, 6

geneRanges, 6

h5GeneNames (scLVM\_output), 8

heter (scLVM\_output), 8

mTECdx, 6

muc1Coexpression, 7

nomarkerCellsClustering, 7

percentsGG, 7

permsAllClusters (permutationResults), 8

permutationResults, 8

realAllClusters (permutationResults), 8

scLVM\_output, 8

Single.mTec.Transcriptomes-package, 2

tras, 8

vars (scLVM\_output), 8

Ycorr (scLVM\_output), 8