

Package ‘scRNAseq’

January 21, 2021

Title Collection of Public Single-Cell RNA-Seq Datasets

Version 2.4.0

Date 2020-10-12

Description Gene-level counts for a collection of public scRNA-seq datasets, provided as SingleCellExperiment objects with cell- and gene-level metadata.

License CC0

NeedsCompilation no

Depends SingleCellExperiment

Imports utils, methods, BiocGenerics, S4Vectors, GenomicRanges, SummarizedExperiment, ExperimentHub, AnnotationHub, AnnotationDbi, ensemblDb, GenomicFeatures

Suggests BiocStyle, knitr, rmarkdown, BiocFileCache, testthat

VignetteBuilder knitr

Encoding UTF-8

biocViews ExperimentHub, ExperimentData, ExpressionData, SequencingData, RNASeqData, SingleCellData

BuildResaveData no

RoxygenNote 7.1.1

git_url <https://git.bioconductor.org/packages/scRNAseq>

git_branch RELEASE_3_12

git_last_commit 61f2c38

git_last_commit_date 2020-10-27

Date/Publication 2021-01-21

Author Davide Risso [aut, cph],
Michael Cole [aut],
Aaron Lun [ctb, cre],
Alan O'Callaghan [ctb],
Jens Preussner [ctb],
Charlotte Soneson [ctb],
Stephany Orjuela [ctb]

Maintainer Aaron Lun <infinite.monkeys.with.keyboards@gmail.com>

R topics documented:

scRNAseq-package	2
AztekinTailData	3
BachMammaryData	4
BaronPancreasData	5
BuettnerESCData	6
CampbellBrainData	7
ChenBrainData	8
ERCCSpikeInConcentrations	9
GrunHSCData	10
GrunPancreasData	11
HermannSpermatogenesisData	12
HuCortexData	13
KolodziejczykESCData	14
KotliarovPBMCData	15
LaMannoBrainData	16
LawlorPancreasData	17
LengESCData	18
listDatasets	19
LunSpikeInData	20
MacoskoRetinaData	21
MairPBMCData	22
MarquesBrainData	23
MessmerESCData	24
MuraroPancreasData	25
NestorowaHSCData	26
PaulHSCData	27
ReprocessedAllenData	28
RichardTCellData	29
RomanovBrainData	30
SegerstolpePancreasData	31
ShekharRetinaData	32
StoeckiusHashingData	33
TasicBrainData	35
UsoskinBrainData	36
WuKidneyData	37
XinPancreasData	38
ZeiselBrainData	39
ZilionisLungData	40
Index	42

scRNAseq-package

*Collection of Public Single-Cell RNA-Seq Datasets***Description**

Gene-level counts for a collection of public scRNA-seq datasets, provided as SingleCellExperiment objects with cell- and gene-level metadata.

Details

This package contains a collection of three publicly available single-cell RNA-seq datasets.

The dataset `fluidigm` contains 65 cells from Pollen et al. (2014), each sequenced at high and low coverage.

The dataset `th2` contains 96 T helper cells from Mahata et al. (2014).

The dataset `allen` contains 379 cells from the mouse visual cortex. This is a subset of the data published in Tasic et al. (2016).

See the package vignette for details on the pre-processing of the data.

Author(s)

NA

Maintainer: NA

References

Pollen, Nowakowski, Shuga, Wang, Leyrat, Lui, Li, Szpankowski, Fowler, Chen, Ramalingam, Sun, Thu, Norris, Lebofsky, Toppani, Kemp II, Wong, Clerkson, Jones, Wu, Knutsson, Alvarado, Wang, Weaver, May, Jones, Unger, Kriegstein, West. Low-coverage single-cell mRNA sequencing reveals cellular heterogeneity and activated signaling pathways in developing cerebral cortex. *Nature Biotechnology*, 32, 1053-1058 (2014).

Mahata, Zhang, Kolodziejczyk, Proserpio, Haim-Vilmovsky, Taylor, Hebenstreit, Dingler, Moignard, Gottgens, Arlt, McKenzie, Teichmann. Single-Cell RNA Sequencing Reveals T Helper Cells Synthesizing Steroids De Novo to Contribute to Immune Homeostasis. *Cell Reports*, 7(4): 1130–1142 (2014).

Tasic, Menon, Nguyen, Kim, Jarsky, Yao, Levi, Gray, Sorensen, Dolbeare, Bertagnolli, Goldy, Shapovalova, Parry, Lee, Smith, Bernard, Madisen, Sunkin, Hawrylycz, Koch, Zeng. Adult mouse cortical cell taxonomy revealed by single cell transcriptomics. *Nature Neuroscience*, 19, 335–346 (2016).

AztekinTailData

Obtain the Aztekin tail data

Description

Obtain the *Xenopus* tail single-cell RNA-seq data from Aztekin et al. (2019).

Usage

```
AztekinTailData()
```

Details

Column metadata is provided in the same form as supplied in E-MTAB-7761. This contains information such as the treatment condition, batch, putative cell type, putative cell cycle phase.

The UMAP results are available as the "UMAP" entry in the `reducedDims`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scrNAseq/aztekin-tail`.

Value

A [SingleCellExperiment](#) object with a single matrix of UMI counts.

Author(s)

Aaron Lun

References

Aztekin C et al. (2019). Identification of a regeneration-organizing cell in the *Xenopus* tail. *Science* 364(6441), 653-658

Examples

```
sce <- AztekinTailData()
```

BachMammaryData	<i>Obtain the Bach mammary data</i>
-----------------	-------------------------------------

Description

Obtain the mouse mammary gland single-cell RNA-seq data from Bach et al. (2017).

Usage

```
BachMammaryData(
  samples = c("NP_1", "NP_2", "G_1", "G_2", "L_1", "L_2", "PI_1", "PI_2"),
  location = TRUE
)
```

Arguments

samples	A character vector with at least one element, specifying which samples(s) to retrieve.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

Column metadata is extracted from the sample annotation in GSE106273, and refers to the developmental stage of the mammary gland.

If multiple samples are specified in `samples`, the count matrices will be cbinded together. Cells originating from different samples are identifiable by the "Sample" field in the column metadata.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the [rowRanges](#) of the output.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/bach-mammary`.

Value

A [SingleCellExperiment](#) object with a single matrix of UMI counts.

Author(s)

Aaron Lun

References

Bach K et al. (2017). Differentiation dynamics of mammary epithelial cells revealed by single-cell RNA sequencing. *Nat Commun.* 8(1), 2128

Examples

```
sce <- BachMammaryData(samples="NP_1")
```

BaronPancreasData	<i>Obtain the Baron pancreas data</i>
-------------------	---------------------------------------

Description

Obtain the human/mouse pancreas single-cell RNA-seq data from Baron et al. (2017).

Usage

```
BaronPancreasData(  
  which = c("human", "mouse"),  
  ensembl = FALSE,  
  location = TRUE  
)
```

Arguments

<code>which</code>	String specifying the species to get data for.
<code>ensembl</code>	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
<code>location</code>	Logical scalar indicating whether genomic coordinates should be returned.

Details

Column metadata is provided in the same form as supplied in GSE84133. This contains information such as the cell type labels and donor ID (for humans) or strain (for mouse).

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the `rowRanges` of the output. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/baron-pancreas`.

Value

A [SingleCellExperiment](#) object with a single matrix of read counts.

Author(s)

Aaron Lun

References

Baron M et al. (2017). Single-cell transcriptomic map of the human and mouse pancreas reveals inter- and intra-cell population structure. *Cell Syst.* 3(4), 346-360.

Examples

```
sce.human <- BaronPancreasData()
sce.mouse <- BaronPancreasData("mouse")
```

BuettnerESData	<i>Obtain the Buettner ESC data</i>
----------------	-------------------------------------

Description

Obtain the mouse embryonic stem cell single-cell RNA-seq data from Buettner et al. (2015).

Usage

```
BuettnerESData(remove.htseq = TRUE, location = TRUE)
```

Arguments

`remove.htseq` Logical scalar indicating whether HT-seq alignment statistics should be removed.
`location` Logical scalar indicating whether genomic coordinates should be returned.

Details

Rows corresponding to HT-seq's alignment statistics are removed by default. These can be retained by setting `remove.htseq=FALSE`.

Column metadata contains the experimentally determined cell cycle phase for each cell.

Counts for ERCC spike-ins are stored in the "ERCC" entry in the [altExps](#).

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the [rowRanges](#) of the output.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/buettner-esc`.

Value

A [SingleCellExperiment](#) object with a single matrix of read counts.

Author(s)

Aaron Lun

References

Buettner F et al. (2015). Computational analysis of cell-to-cell heterogeneity in single-cell RNA-sequencing data reveals hidden subpopulations of cells. *Nat. Biotechnol.* 33(2), 155-160.

Examples

```
sce <- BuettnerESData()
```

CampbellBrainData	<i>Obtain the Campbell brain data</i>
-------------------	---------------------------------------

Description

Obtain the mouse brain single-cell RNA-seq data from Campbell et al. (2017).

Usage

```
CampbellBrainData(ensembl = FALSE, location = TRUE)
```

Arguments

ensembl	Logical scalar indicating whether the row names of the returned object should contain Ensembl identifiers.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

Column metadata is provided in the same form as supplied in GSE93374. This contains information such as the diet of the mice, sex and proposed cell type for each cell.

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the `rowRanges` of the output. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scrnaseq/campbell-brain`.

Value

A [SingleCellExperiment](#) object with a single matrix of UMI counts.

Author(s)

Aaron Lun

References

Campbell R et al. (2017). A molecular census of arcuate hypothalamus and median eminence cell types. *Nat. Neurosci.* 20, 484-496.

Examples

```
sce <- CampbellBrainData()
```

ChenBrainData	<i>Obtain the Chen brain data</i>
---------------	-----------------------------------

Description

Obtain the mouse brain single-cell RNA-seq data from Chen et al. (2017).

Usage

```
ChenBrainData(ensembl = FALSE, location = TRUE)
```

Arguments

ensembl	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

Column metadata is provided in the same form as supplied in GSE87544. This contains the putative cell type assigned by the original authors.

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the `rowRanges` of the output. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/chen-brain`.

Value

A [SingleCellExperiment](#) object with a single matrix of UMI counts.

Author(s)

Aaron Lun

References

Chen R et al. (2017). Single-Cell RNA-Seq reveals hypothalamic cell diversity. *Cell Rep.* 18, 3227-3241.

Examples

```
sce <- ChenBrainData()
```

ERCCSpikeInConcentrations

Obtain ERCC concentrations

Description

Obtain ERCC spike-in concentrations from the Thermo Fisher Scientific website.

Usage

```
ERCCSpikeInConcentrations(volume = NULL, dilution = NULL, mix = c("1", "2"))
```

Arguments

volume	Numeric scalar specifying the added volume (in nanoliters) of ERCC spike-in mixture. Only used if dilution is specified.
dilution	Numeric scalar specifying the dilution factor used for the added volume of the spike-in mixture. Only used if volume is specified.
mix	String specifying whether to compute the number of molecules for mix 1 or 2. Only used if both dilution and volume are specified.

Details

If volume and dilution are specified, an additional column is added to the output specifying the number of molecules of spike-in transcript for the specified mix.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/ercc-concentrations`.

Value

A [DataFrame](#) object with one row per ERCC spike-in transcript. This contains information such as the spike-in concentration in each mix.

Author(s)

Alan O'Callaghan

Examples

```
df <- ERCCSpikeInConcentrations()
```

GrunHSCData

Obtain the Grun HSC data

Description

Obtain the mouse haematopoietic stem cell single-cell RNA-seq data from Grun et al. (2016).

Usage

```
GrunHSCData(ensembl = FALSE, location = TRUE)
```

Arguments

ensembl	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

Row metadata contains the symbol and chromosomal location for each gene. Column metadata contains the extraction protocol used for each sample, as described in GSE76983.

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the `rowRanges` of the output. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scrnaseq/grun-hsc`.

Value

A [SingleCellExperiment](#) object with a single matrix of UMI counts.

Author(s)

Aaron Lun

References

Grun D et al. (2016). De novo prediction of stem cell identity using single-cell transcriptome data. *Cell Stem Cell* 19(2), 266-277.

Examples

```
sce <- GrunHSCData()
```

GrunPancreasData	<i>Obtain the Grun pancreas data</i>
------------------	--------------------------------------

Description

Obtain the human pancreas single-cell RNA-seq data from Grun et al. (2016).

Usage

```
GrunPancreasData(ensembl = FALSE, location = TRUE)
```

Arguments

ensembl	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

Row metadata contains fields for the symbol and chromosomal location of each gene, as derived from the row names.

Column metadata is derived from the column names of the count matrix with the sample annotations in GSE81076. This includes the donor identity for each cell and the type of sample.

The "ERCC" entry in the [altExps](#) contains count data for the ERCC spike-in transcripts.

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the [rowRanges](#) of the output. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/grun-pancreas`.

Value

A [SingleCellExperiment](#) object with a single matrix of UMI counts.

Author(s)

Aaron Lun, using additional metadata obtained by Vladimir Kiselev.

References

Grun D et al. (2016). De novo prediction of stem cell identity using single-cell transcriptome data. *Cell Stem Cell* 19(2), 266-277.

Examples

```
sce <- GrunPancreasData()
```

HermannSpermatogenesisData

Obtain the Hermann spermatogenesis data

Description

Obtain the mouse spermatogenesis single-cell RNA-seq data from Hermann et al. (2018).

Usage

```
HermannSpermatogenesisData(strip = FALSE, location = TRUE)
```

Arguments

<code>strip</code>	Logical scalar indicating whether to strip the <code>.X</code> notation from the row names.
<code>location</code>	Logical scalar indicating whether genomic coordinates should be returned. Only used if <code>strip=TRUE</code> .

Details

Column metadata contains cell types provided by the data generators at <https://data.mendeley.com/datasets/kxd5f8vpt4/1#file-fe79c10b-c42e-472e-9c7e-9a9873d9b3d8>.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/hermann-spermatogenesis`.

Value

A `SingleCellExperiment` object with two matrices, containing spliced and unspliced counts, respectively.

Author(s)

Charlotte Soneson

References

Hermann B.P. et al. (2018). The Mammalian Spermatogenesis Single-Cell Transcriptome, from Spermatogonial Stem Cells to Spermatids. *Cell Rep.* 25(6), 1650-1667.e8.

Examples

```
sce <- HermannSpermatogenesisData()
```

HuCortexData	<i>Obtain the Hu cortex data</i>
--------------	----------------------------------

Description

Obtain the mouse cortex single-nuclei RNA-seq data from Hu et al. (2017).

Usage

```
HuCortexData(  
  mode = c("ctx", "3T3"),  
  samples = NULL,  
  ensembl = FALSE,  
  location = TRUE  
)
```

Arguments

mode	Character vector indicating whether to return data for the 3T3 cells or the mouse cortex.
samples	Character vector indicating whether to return data for specific samples, see Details. If specified, this overrides mode.
ensembl	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

Column metadata includes the mode and sample corresponding to each cell/nuclei. Available samples are:

- "cell-3T3" and "nuclei-3T3", generated from the 3T3 cell line.
- "nuclei-ctx-X", nuclei generated from the cortex of animal number X (from 1 to 13).
- "nuclei-ctx-salineX" or "nuclei-ctx-PTZX", nuclei generated from the cortex of saline- or PTZ-treated mice. X represents the replicate number and can be 1 or 2.

If multiple modes are requested, counts are only reported for the intersection of genes across all modes. This is because the gene annotation in the original count matrices differs across modes.

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the `rowRanges` of the output. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/wu-kidney`.

Value

A [SingleCellExperiment](#) object with a single matrix of read counts.

Author(s)

Aaron Lun

References

Hu P et al. (2017). Dissecting cell-type composition and activity-dependent transcriptional state in mammalian brains by massively parallel single-nucleus RNA-seq. *Mol. cell* 68, 1006-1015.

Examples

```
sce <- HuCortexData("3T3")
```

KolodziejczykESCData *Obtain the Kolodziejczyk ESC data*

Description

Obtain the mouse embryonic stem cell single-cell RNA-seq data from Kolodziejczyk et al. (2015).

Usage

```
KolodziejczykESCData(remove.htseq = TRUE, location = TRUE)
```

Arguments

<code>remove.htseq</code>	Logical scalar indicating whether HT-seq alignment statistics should be removed.
<code>location</code>	Logical scalar indicating whether genomic coordinates should be returned.

Details

Column metadata is generated from the column names, and contains the culture conditions and the plate of origin for each cell.

Count data for ERCC spike-ins are stored in the "ERCC" entry in the [altExps](#).

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the [rowRanges](#) of the output.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/kolodziejczyk-esc`.

Value

A [SingleCellExperiment](#) object with a single matrix of read counts.

Author(s)

Aaron Lun

References

Messmer T et al. (2019). Transcriptional heterogeneity in naive and primed human pluripotent stem cells at single-cell resolution. *Cell Rep* 26(4), 815-824.e4

Examples

```
sce <- KolodziejczykESData()
```

KotliarovPBMCData *Obtain the Kotliarov CITE-seq data*

Description

Obtain the Kotliarov PBMC CITE-seq data from Kotliarov et al. (2020).

Usage

```
KotliarovPBMCData(mode = c("rna", "adt"), ensembl = FALSE, location = TRUE)
```

Arguments

mode	Character vector specifying whether to return either or both the RNA and ADT counts.
ensembl	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

This dataset contains 20 samples from 2 experimental batches, where each batch contains 5 high and 5 low responders. The 10 samples per batch were mixed and distributed across the 6 lanes using a cell hashing approach.

The column metadata contains the following fields:

- `sample*`: identifiers for the sample of origin for each cell.
- `adjmfc.time`: type of responder for each sample.
- `tenx_lane`: 10X lane from which each cell was collected.
- `batch`: the batch of origin.
- `barcode_check`: barcode identifier.
- `hash_*` and `hto_*` columns: **HTOdemux** outputs.
- `DEMUXLET.*` columns: **demuxlet** outputs.
- `joint_classification_global`: **HTOdemux** and **demuxlet** joint classification.
- `nGene`: number of genes as defined from **Seurat**'s `CreateSeuratObject`.
- `nUMI`: number of UMIs as defined from **Seurat**'s `CreateSeuratObject`.
- `pctMT`: percent of mitochondrial reads as defined from **Seurat**'s `CreateSeuratObject`.

Note, no filtering has been performed based on the quality control metrics.

If `ensembl=TRUE`, the gene symbols in the RNA data are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the `rowRanges` for the RNA data. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scrnaSeq/kotliarov-pbmc`.

Value

A [SingleCellExperiment](#) object with a single matrix of UMI counts corresponding to the first mode, with an optional alternative Experiment if there is a second mode.

Author(s)

Stephany Orjuela, with modifications from Aaron Lun

References

Kotliarov et al. (2020). Broad immune activation underlies shared set point signatures for vaccine responsiveness in healthy individuals and disease activity in patients with lupus. *Nat. Med.* 26, 618–629

Examples

```
sce <- KotliarovPBMCData()
```

LaMannoBrainData *Obtain the La Manno brain data*

Description

Obtain the mouse/human brain scRNA-seq data from La Manno et al. (2016).

Usage

```
LaMannoBrainData(
  which = c("human-es", "human-embryo", "human-ips", "mouse-adult", "mouse-embryo"),
  ensembl = FALSE,
  location = TRUE
)
```

Arguments

which	A string specifying which dataset should be obtained.
ensembl	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

Column metadata is provided in the same form as supplied in the supplementary tables in GSE71585. This contains information such as the time point and cell type.

The various settings of which will obtain different data sets.

- "human-es", human embryonic stem cells.
- "human-embryo", human embryo midbrain.
- "human-ips", human induced pluripotent stem cells.

- "mouse-adult", mouse adult dopaminergic neurons.
- "mouse-embryo", mouse embryo midbrain.

Unfortunately, each of these datasets uses a different set of features. If multiple datasets are to be used simultaneously, users will have to decide how to merge them, e.g., by taking the intersection of common features across all datasets.

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the `rowRanges` of the output. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/lamanno-brain`.

Value

A `SingleCellExperiment` object with a single matrix of UMI counts.

Author(s)

Aaron Lun

References

La Manno A et al. (2016). Molecular diversity of midbrain development in mouse, human, and stem cells. *Cell* 167(2), 566-580.

Examples

```
sce.h.es <- LaMannoBrainData()
sce.h.em <- LaMannoBrainData("human-embryo")
sce.h.ip <- LaMannoBrainData("human-ips")
sce.m.ad <- LaMannoBrainData("mouse-adult")
sce.m.em <- LaMannoBrainData("mouse-embryo")
```

LawlorPancreasData *Obtain the Lawlor pancreas data*

Description

Provides the human pancreas single-cell RNA-seq data from Lawlor et al. (2017).

Usage

```
LawlorPancreasData()
```

Details

Column metadata is provided in the same form as supplied in GSE86469. This contains information such as the cell type labels and patient status.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for scRNAseq/Lawlor-pancreas.

Value

A [SingleCellExperiment](#) object with a single matrix of read counts.

Author(s)

Aaron Lun

References

Lawlor N et al. (2017). Single-cell transcriptomes identify human islet cell signatures and reveal cell-type-specific expression changes in type 2 diabetes. *Genome Res.* 27(2), 208-222.

Examples

```
sce <- LawlorPancreasData()
```

LengESCData

Obtain the Leng ESC data

Description

Obtain the human embryonic stem cell single-cell RNA-seq data from Leng et al. (2015).

Usage

```
LengESCData(ensembl = FALSE, location = TRUE)
```

Arguments

ensembl	Logical scalar indicating whether gene symbols should be converted to Ensembl annotation.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

Column metadata contains the cell line, experiment number and experimentally determined cell cycle phase for each cell.

If `ensembl=TRUE`, the gene symbols in the published annotation are converted to Ensembl. If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the [rowRanges](#) of the output.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for scRNAseq/leng-esc.

Value

A [SingleCellExperiment](#) object with a single matrix of normalized expected read counts.

Author(s)

Aaron Lun

References

Leng F et al. (2015). Oscope identifies oscillatory genes in unsynchronized single-cell RNA-seq experiments. *Nat. Methods* 12(10), 947-950.

Examples

```
sce <- LengESData()
```

listDatasets	<i>List all available datasets</i>
--------------	------------------------------------

Description

Summary information for all available datasets in the **scRNAseq** package.

Usage

```
listDatasets()
```

Details

A study may contribute multiple datasets if they cannot be reasonably combined (e.g., different species). The reported number of cells refers only to the dataset as it is stored in **scRNAseq**; this may be different to the number of cells used by the authors in their analysis, e.g., due to filtering.

Value

A [DataFrame](#) where each row corresponds to a dataset, containing the fields:

- Reference, a Markdown-formatted citation to `scripts/ref.bib` in the **scRNAseq** installation directory.
- Taxonomy, an identifier for the organism.
- Part, the part of the organism being studied.
- Number, the total number of cells in the dataset.
- Call, the relevant R call required to construct the dataset.

Author(s)

Aaron Lun

Examples

```
listDatasets()
```

LunSpikeInData	<i>Obtain the Lun spike-in data</i>
----------------	-------------------------------------

Description

Obtain the spike-in single-cell RNA-seq data from Lun et al. (2017).

Usage

```
LunSpikeInData(  
  which = c("416b", "tropho"),  
  split.oncogene = FALSE,  
  location = TRUE  
)
```

Arguments

which	String specifying whether the 416B or trophoblast data should be obtained.
split.oncogene	Logical scalar indicating whether the oncogene should be split to a separate altExp .
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

Row data contains a single "Length" field describing the total exonic length of each feature.

Column metadata is provided in the same form as supplied in E-MTAB-5522. This contains information such as the cell type, plate of origin, spike-in addition order and oncogene induction.

Two sets of spike-ins were added to each cell in each dataset. These are available as the "SIRV" and "ERCC" entries in the [altExps](#).

If `split.oncogene=TRUE` and `which="416b"`, the CFBF-MYH11-mcherry oncogene is moved to extra "oncogene" entry in the [altExps](#).

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the [rowRanges](#) of the output.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/lun-spikein`.

Value

A [SingleCellExperiment](#) object with a single matrix of read counts.

Author(s)

Aaron Lun

References

Lun ATL et al. (2017). Assessing the reliability of spike-in normalization for analyses of single-cell RNA sequencing data. *Genome Res.* 27(11), 1795-1806.

Examples

```
sce <- LunSpikeInData()

sce <- LunSpikeInData("tropho")
```

MacoskoRetinaData *Obtain the Macosko retina data*

Description

Obtain the mouse retina single-cell RNA-seq data from Macosko et al. (2016).

Usage

```
MacoskoRetinaData(ensembl = FALSE, location = TRUE)
```

Arguments

ensembl	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

Column metadata contains the cluster identity as reported in the paper. Note that some cells will have NA identities as they are present in the count matrix but not in the metadata file. These are presumably low-quality cells that were discarded prior to clustering.

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the `rowRanges` of the output. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scrnaseq/macosko-retina`.

Value

A [SingleCellExperiment](#) object with a single matrix of UMI counts.

Author(s)

Aaron Lun

References

Macosko E et al. (2016). Highly parallel genome-wide expression profiling of individual cells using nanoliter droplets. *Cell* 161(5), 1202-1214.

Examples

```
sce <- MacoskoRetinaData()
```

MairPBMCData *Obtain the Mair CITE-seq data*

Description

Obtain the Mair PBMC targeted CITE-seq data from Mair et al. (2020).

Usage

```
MairPBMCData(mode = c("rna", "adt"), ensembl = FALSE, location = TRUE)
```

Arguments

mode	Character vector specifying whether to return either or both the RNA and ADT counts.
ensembl	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

Column metadata contains the donor identity and cartridge of origin. Some libraries may also be classified as multiplets or have undeterminate origins after hash tag debarcoding.

If `ensembl=TRUE`, the gene symbols in the RNA data are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the `rowRanges` for the RNA data. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/mair-pbmc`.

Value

A `SingleCellExperiment` object with a single matrix of UMI counts corresponding to the first mode, with an optional alternative Experiment if there is a second mode.

Author(s)

Stephany Orjuela, with modifications from Aaron Lun

References

Mair C et al. (2020). A targeted multi-omic analysis approach measures protein expression and low-abundance transcripts on the single-cell level. *Cell Rep.* 31, 107499

Examples

```
sce <- MairPBMCData()
```

MarquesBrainData	<i>Obtain the Marques brain data</i>
------------------	--------------------------------------

Description

Obtain the mouse brain single-cell RNA-seq data from Marques et al. (2016).

Usage

```
MarquesBrainData(ensembl = FALSE, location = TRUE)
```

Arguments

ensembl	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

Column metadata is provided in the same form as supplied in GSE75330. This contains information such as the cell type and age/sex of the mouse of origin for each cell.

Note that some genes may be present in multiple rows corresponding to different genomic locations. These additional rows are identified by a `_loc[2-9]` suffix in their row names. Users may wish to consider either removing them or merging them, e.g., with `scater::sumCountsAcrossFeatures`.

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained. All searching is performed after removing the `_loc[2-9]` suffix.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the `rowRanges` of the output. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/marques-brain`.

Value

A [SingleCellExperiment](#) object with a single matrix of UMI counts.

Author(s)

Aaron Lun

References

Marques A et al. (2016). Oligodendrocyte heterogeneity in the mouse juvenile and adult central nervous system. *Science* 352(6291), 1326-1329.

Examples

```
sce <- MarquesBrainData()
```

MessmerESCData

Obtain the Messmer ESC data

Description

Obtain the human embryonic stem cell single-cell RNA-seq data from Messmer et al. (2019).

Usage

```
MessmerESCData(location = TRUE)
```

Arguments

`location` Logical scalar indicating whether genomic coordinates should be returned.

Details

Row data contains a single "Length" field describing the total exonic length of each feature.

Column metadata is provided in the same form as supplied in E-MTAB-6819. This contains information such as the cell phenotype (naive or primed) and the batch of origin. Note that counts for technical replicates have already been summed together.

Count data for ERCC spike-ins are stored in the "ERCC" entry of the [altExps](#).

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the [rowRanges](#) of the output.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/messmer-esc`.

Value

A [SingleCellExperiment](#) object with a single matrix of read counts.

Author(s)

Aaron Lun

References

Messmer T et al. (2019). Transcriptional heterogeneity in naive and primed human pluripotent stem cells at single-cell resolution. *Cell Rep* 26(4), 815-824.e4

Examples

```
sce <- MessmerESCData()
```

MuraroPancreasData *Obtain the Muraro pancreas data*

Description

Obtain the human pancreas single-cell RNA-seq data from Muraro et al. (2016).

Usage

```
MuraroPancreasData(ensembl = FALSE, location = TRUE)
```

Arguments

ensembl	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

Row data contains fields for the symbol and chromosomal location of each gene.

Column metadata is derived from the columns of the count matrix provided in GSE85241, with additional cell type labels obtained from the authors (indirectly, via the Hemberg group). Some cells have NA labels and were presumably removed prior to downstream analyses.

Count data for ERCC spike-ins are stored in the "ERCC" entry of the [altExps](#).

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the [rowRanges](#) of the output. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/muraro-pancreas`.

Value

A [SingleCellExperiment](#) object with a single matrix of UMI counts.

Author(s)

Aaron Lun, using additional metadata obtained by Vladimir Kiselev.

References

Muraro MJ et al. (2016). A single-cell transcriptome atlas of the human pancreas. *Cell Syst.* 3(4), 385-394.

Examples

```
sce <- MuraroPancreasData()
```

NestorowaHSCData	<i>Obtain the Nestorowa HSC data</i>
------------------	--------------------------------------

Description

Obtain the mouse haematopoietic stem cell single-cell RNA-seq data from Nestorowa et al. (2015).

Usage

```
NestorowaHSCData(remove.htseq = TRUE, location = TRUE)
```

Arguments

<code>remove.htseq</code>	Logical scalar indicating whether HT-seq alignment statistics should be removed.
<code>location</code>	Logical scalar indicating whether genomic coordinates should be returned.

Details

Rows corresponding to HT-seq's alignment statistics are removed by default. These can be retained by setting `remove.htseq=FALSE`.

Column metadata includes the cell type mapping, as described on the website (see References), and the FACS expression levels of selected markers. Note that these are stored as nested matrices within the `colData`.

Diffusion map components are provided as the "diffusion" entry in the `reducedDims`.

Counts for ERCC spike-ins are stored in the "ERCC" entry in the `altExps`.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the `rowRanges` of the output.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/nestorowa-hsc`.

Value

A `SingleCellExperiment` object with a single matrix of read counts.

Author(s)

Aaron Lun

References

Nestorowa S et al. (2016). A single-cell resolution map of mouse hematopoietic stem and progenitor cell differentiation *Blood* 128, e20-e31.

Gene and protein expression in adult haematopoiesis: Data. http://blood.stemcells.cam.ac.uk/single_cell_atlas.html#data.

Examples

```
sce <- NestorowaHSCData()
```

PaulHSCData

Obtain the Paul HSC data

Description

Obtain the mouse haematopoietic stem cell single-cell RNA-seq data from Paul et al. (2015).

Usage

```
PaulHSCData(ensembl = FALSE, discard.multiple = TRUE, location = TRUE)
```

Arguments

<code>ensembl</code>	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
<code>discard.multiple</code>	Logical scalar indicating whether ambiguous rows should be discarded.
<code>location</code>	Logical scalar indicating whether genomic coordinates should be returned.

Details

Column metadata includes the plate and the mouse of origin, fluorescence intensities from indexed sorting and the number of cells in each well.

Some of the original rownames are concatenated symbols from multiple genes. We consider these rows to represent ambiguously assigned counts and discard them if `discard.multiple=TRUE`.

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the `rowRanges` of the output. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/nestorowa-hsc`.

Value

A [SingleCellExperiment](#) object with a single matrix of read counts.

Author(s)

Aaron Lun

References

Paul F et al. (2015). Transcriptional heterogeneity and lineage commitment in myeloid progenitors. *Cell* 163, 1663-77.

Examples

```
sce <- PaulHSCData()
```

 ReprocessedAllenData *Reprocessed single-cell data sets*

Description

Obtain the legacy count matrices for three publicly available single-cell RNA-seq datasets. Raw sequencing data were downloaded from NCBI's SRA or from EBI's ArrayExpress, aligned to the relevant genome build and used to quantify gene expression.

Usage

```
ReprocessedAllenData(assays = NULL, ensembl = FALSE, location = TRUE)
```

```
ReprocessedTh2Data(assays = NULL, ensembl = FALSE, location = TRUE)
```

```
ReprocessedFluidigmData(assays = NULL, ensembl = FALSE, location = TRUE)
```

Arguments

assays	Character vector specifying one or more assays to return. Choices are "tophat_counts", "cufflinks_fpkms", "rsem_counts" and "rsem_tpm". If NULL, all assays are returned.
ensembl	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

ReprocessedFluidigmData returns a dataset of 65 human neural cells from Pollen et al. (2014), each sequenced at high and low coverage (SRA accession SRP041736).

ReprocessedTh2Data returns a dataset of 96 mouse T helper cells from Mahata et al. (2014), obtained from ArrayExpress accession E-MTAB-2512. Spike-in counts are stored in the "ERCC" entry of the `altExps`.

ReprocessedAllenData return a dataset of 379 mouse brain cells from Tasic et al. (2016). This is a re-processed subset of the data from [TasicBrainData](#), and contains spike-in information stored as in the `altExps`.

In each dataset, the first columns of the `colData` are sample quality metrics from FastQC and Picard. The remaining fields were obtained from the original study in their GEO/SRA submission and/or as Supplementary files in the associated publication. These two categories of `colData` are distinguished by a `which_qc` element in the `metadata`, which contains the names of the quality-related columns in each object.

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the `rowRanges` of the output. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scrnaseq/legacy-allen`, `scrnaseq/legacy-fluidigm` or `scrnaseq/legacy-th2`.

Value

A [SingleCellExperiment](#) object containing one or more expression matrices of counts and/or TPMs, depending on assays.

Pre-processing details

FASTQ files were either obtained directly from ArrayExpress, or converted from SRA files (downloaded from the Sequence Read Archive) using the SRA Toolkit.

Reads were aligned with TopHat (v. 2.0.11) to the appropriate reference genome (GRCh38 for human samples, GRCm38 for mouse). RefSeq mouse gene annotation (GCF_000001635.23_GRCm38.p3) was downloaded from NCBI on Dec. 28, 2014. RefSeq human gene annotation (GCF_000001405.28) was downloaded from NCBI on Jun. 22, 2015.

featureCounts (v. 1.4.6-p3) was used to compute gene-level read counts. Cufflinks (v. 2.2.0) was used to compute gene-level FPKMs. Reads were also mapped to the transcriptome using RSEM (v. 1.2.19) to compute read counts and TPM's.

FastQC (v. 0.10.1) and Picard (v. 1.128) were used to compute sample quality control (QC) metrics. However, no filtering on the QC metrics has been performed for any dataset.

References

Pollen AA et al. (2014). Low-coverage single-cell mRNA sequencing reveals cellular heterogeneity and activated signaling pathways in developing cerebral cortex. *Nat. Biotechnol.* 32(10), 1053-8.

Mahata B et al. (2014). Single-cell RNA sequencing reveals T helper cells synthesizing steroids de novo to contribute to immune homeostasis. *Cell Rep*, 7(4), 1130-42.

Tasic A et al. (2016). Adult mouse cortical cell taxonomy revealed by single cell transcriptomics. *Nat. Neurosci.* 19(2), 335-46.

Examples

```
sce <- ReprocessedAllenData()
```

RichardTCellData	<i>Obtain the Richard T cell data</i>
------------------	---------------------------------------

Description

Obtain the mouse CD8+ T cell single-cell RNA-seq data from Richard et al. (2018).

Usage

```
RichardTCellData(location = TRUE)
```

Arguments

location Logical scalar indicating whether genomic coordinates should be returned.

Details

Column metadata is provided in the same form as supplied in E-MTAB-6051. This contains information such as the stimulus, time after stimulation, age of the mice and sequencing batch.

Count data for ERCC spike-ins are stored in the "ERCC" entry of the [altExps](#).

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the [rowRanges](#) of the output.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/richard-tcell`.

Value

A [SingleCellExperiment](#) object with a single matrix of read counts.

Author(s)

Aaron Lun

References

Richard AC et al. (2018). T cell cytolytic capacity is independent of initial stimulation strength. *Nat. Immunol.* 19(8), 849-858.

Examples

```
sce <- RichardTCellData()
```

RomanovBrainData

Obtain the Romanov brain data

Description

Obtain the mouse brain single-cell RNA-seq dataset from Romanov et al. (2017).

Usage

```
RomanovBrainData(ensembl = FALSE, location = TRUE)
```

Arguments

`ensembl` Logical scalar indicating whether the output row names should contain Ensembl identifiers.

`location` Logical scalar indicating whether genomic coordinates should be returned.

Details

Column metadata is provided in the same form as supplied in GSE74672. This contains information such as the reporter gene expressed in each cell, the mouse line, dissection type and so on.

Counts for ERCC spike-ins are stored in the "ERCC" entry of the [altExps](#). Note that some of the spike-in rows have NA observations for some (but not all) cells.

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the [rowRanges](#) of the output. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/romanov-brain`.

Value

A [SingleCellExperiment](#) object with a single matrix of UMI counts.

Author(s)

Aaron Lun, based on code by Vladimir Kiselev and Tallulah Andrews.

References

Romanov RA et al. (2017). Molecular interrogation of hypothalamic organization reveals distinct dopamine neuronal subtypes. *Nat. Neurosci.* 20, 176-188.

Examples

```
sce <- RomanovBrainData()
```

SegerstolpePancreasData

Obtain the Segerstolpe pancreas data

Description

Download the human pancreas single-cell RNA-seq (scRNA-seq) dataset from Segerstolpe et al. (2016)

Usage

```
SegerstolpePancreasData(ensembl = FALSE, location = TRUE)
```

Arguments

<code>ensembl</code>	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
<code>location</code>	Logical scalar indicating whether genomic coordinates should be returned.

Details

Row data contains fields for the gene symbol and RefSeq transcript IDs corresponding to each gene. The rows of the output object are named with the symbol, but note that these are not unique.

Column metadata were extracted from the `Characteristics` fields of the SDRF file for ArrayExpress E-MTAB-5061. This contains information such as the cell type labels and patient status.

Count data for ERCC spike-ins are stored in the "ERCC" entry of the `altExps`. Estimated numbers of spike-in molecules are provided in the `rowData` of this entry. Note that these concentrations are incorrect for donor H1, as 100 uL of spike-in mixture were added for this donor, rather than 25 uL for all others.

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the `rowRanges` of the output. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/segerstolpe-pancreas`.

Value

A `SingleCellExperiment` object with a single matrix of read counts.

Author(s)

Aaron Lun

References

Segerstolpe A et al. (2016). Single-cell transcriptome profiling of human pancreatic islets in health and type 2 diabetes. *Cell Metab.* 24(4), 593-607.

Examples

```
sce <- SegerstolpePancreasData()
```

ShekharRetinaData *Obtain the Shekhar retina data*

Description

Obtain the mouse retina single-cell RNA-seq dataset from Shekhar et al. (2016).

Usage

```
ShekharRetinaData(ensembl = FALSE, location = TRUE)
```

Arguments

<code>ensembl</code>	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
<code>location</code>	Logical scalar indicating whether genomic coordinates should be returned.

Details

Column metadata contains the cluster identities as reported in the paper. Note that some cells will have NA identities as they are present in the count matrix but not in the metadata file. These are presumably low-quality cells that were discarded prior to clustering.

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the `rowRanges` of the output. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/shekhar-retina`.

Value

A `SingleCellExperiment` object with a single matrix of UMI counts.

Author(s)

Aaron Lun

References

Shekhar K et al. (2016). Comprehensive classification of retinal bipolar neurons by single-cell transcriptomics. *Cell* 166(5), 1308-1323.

Examples

```
sce <- ShekharRetinaData()
```

StoeckiusHashingData *Obtain the Stoeckius cell hashing data*

Description

Obtain the (mostly human) cell hashing single-cell RNA-seq data from Stoeckius et al. (2018).

Usage

```
StoeckiusHashingData(  
  type = c("pbmc", "mixed"),  
  mode = NULL,  
  ensembl = FALSE,  
  location = TRUE,  
  strip.metrics = TRUE  
)
```

Arguments

<code>type</code>	String specifying the dataset to obtain.
<code>mode</code>	String specifying the data modalities to obtain, see Details .
<code>ensembl</code>	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
<code>location</code>	Logical scalar indicating whether genomic coordinates should be returned.
<code>strip.metrics</code>	Logical scalar indicating whether quality control metrics should be removed from the HTO/ADT counts.

Details

When `type="pbmc"`, the mode can be one or more of:

- `"human"`, the RNA counts for human genes.
- `"mouse"`, the RNA counts for mouse genes. Present as the PBMC dataset is actually a mixture of human PBMCs and unlabelled mouse cells.
- `"hto"`, the HTO counts.
- `"adt1"`, counts for the first set of ADTs (immunoglobulin controls).
- `"adt2"`, counts for the second set of ADTs (cell type-specific markers).

When `type="mixed"`, the mode can be one or more of:

- `"rna"`, the RNA counts for the genes;
- `"hto"`, the HTO counts.

If `ensembl=TRUE`, gene symbols for the RNA counts are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the [rowRanges](#) of the output. Note that this is only performed if `ensembl=TRUE` and only for the RNA counts.

For the HTO and ADT matrices, some rows correspond to quality control metrics. If `strip.metrics=TRUE`, these rows are removed so that only data for actual HTOs or ADTs are present.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/nestorowa-hsc`.

Value

A [SingleCellExperiment](#) object with a matrix of UMI counts corresponding to the first mode, plus any number of alternative Experiments containing the remaining modes. If multiple modes are specified, the output object only contains the intersection of their column names.

Author(s)

Aaron Lun

References

Stoeckius et al. (2018). Cell Hashing with barcoded antibodies enables multiplexing and doublet detection for single cell genomics. *Genome Biol.* 19, 224.

Examples

```
sce.pbmc <- StoeckiusHashingData()
sce.pbmc

sce.mixed <- StoeckiusHashingData(type="mixed")
sce.mixed
```

TasicBrainData	<i>Obtain the Tasic brain data</i>
----------------	------------------------------------

Description

Obtain the mouse brain single-cell RNA-seq data from Tasic et al. (2015).

Usage

```
TasicBrainData(ensembl = FALSE, location = TRUE)
```

Arguments

ensembl	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

Column metadata is provided in the same form as supplied in GSE71585. This contains information such as the reporter gene expressed in each cell, the mouse line, dissection type and so on.

Count data for ERCC spike-ins are stored in the "ERCC" entry of the [altExps](#). Note that some of the spike-in rows have NA observations for some (but not all) cells.

The last 9 columns (containing `_CTX_` in their names) correspond to no-cell control libraries.

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the [rowRanges](#) of the output. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scrNaseq/tasic-brain`.

Value

A [SingleCellExperiment](#) object with a single matrix of read counts.

Author(s)

Aaron Lun

References

Tasic A et al. (2016). Adult mouse cortical cell taxonomy revealed by single cell transcriptomics. *Nat. Neurosci.* 19(2), 335-46.

Examples

```
sce <- TasicBrainData()
```

UsoskinBrainData	<i>Obtain the Usoskin brain data</i>
------------------	--------------------------------------

Description

Obtain the mouse brain single-cell RNA-seq dataset from Usoskin et al. (2015).

Usage

```
UsoskinBrainData(ensembl = FALSE, location = TRUE)
```

Arguments

ensembl	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

Column metadata is provided in the same form as supplied in External Table 2 of <http://linnarssonlab.org/drg/>. This contains information such as the library of origin and the cell type.

The count matrix contains information for repeats, marked with `r_` prefixes in the row names; as well as mitochondrial transcripts, marked with `mt-` prefixes.

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the `rowRanges` of the output. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scrnaseq/usoskin-brain`.

Value

A `SingleCellExperiment` object with a single matrix of RPMs.

Author(s)

Aaron Lun

References

Usoskin A et al. (2015). Unbiased classification of sensory neuron types by large-scale single-cell RNA sequencing. *Nat. Neurosci.* 18(1), 145-53.

Examples

```
sce <- UsoskinBrainData()
```

WuKidneyData	<i>Obtain the Wu kidney data</i>
--------------	----------------------------------

Description

Obtain the mouse kidney single-nuclei RNA-seq data from Wu et al. (2019).

Usage

```
WuKidneyData(mode = c("healthy", "disease"), ensembl = FALSE, location = TRUE)
```

Arguments

mode	String indicating whether to return data for healthy and/or diseased donors.
ensembl	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

Column metadata includes the single-cell technology and whether they came from a diseased or healthy individual.

If mode specifies both healthy and disease donors, counts are only reported for the intersection of genes that are present for both donors. This is because the original count matrices had differences in their annotation.

If ensembl=TRUE, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If location=TRUE, the coordinates of the Ensembl gene models are stored in the [rowRanges](#) of the output. Note that this is only performed if ensembl=TRUE.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for scRNAseq/wu-kidney.

Value

A [SingleCellExperiment](#) object with a single matrix of read counts.

Author(s)

Aaron Lun

References

Wu H et al. (2019). Advantages of single-nucleus over single-cell RNA sequencing of adult kidney: rare cell types and novel cell states revealed in fibrosis. *J. Am. Soc. Nephrol.* 30, 23-32.

Examples

```
sce <- WuKidneyData("disease")
```

XinPancreasData	<i>Obtain the Xin pancreas data</i>
-----------------	-------------------------------------

Description

Obtain the human pancreas single-cell RNA-seq dataset from Xin et al. (2016).

Usage

```
XinPancreasData(ensembl = FALSE, location = TRUE)
```

Arguments

ensembl	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

Row data contains fields for the Entrez ID and symbol for each gene. Column metadata was obtained from the authors (indirectly, via the Hemberg group) and contains information such as the cell type labels and donor status.

If `ensembl=TRUE`, the Entrez IDs are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the `rowRanges` of the output. Note that this is only performed if `ensembl=TRUE`.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/xin-pancreas`.

Value

A [SingleCellExperiment](#) object with a single matrix of RPKMs.

Author(s)

Aaron Lun, using additional metadata obtained by Vladimir Kiselev.

References

Xin A et al. (2016). RNA sequencing of single human islet cells reveals type 2 diabetes genes. *Cell Metab.* 24(4), 608-615.

Examples

```
sce <- XinPancreasData()
```

ZeiselBrainData	<i>Obtain the Zeisel brain data</i>
-----------------	-------------------------------------

Description

Obtain the mouse brain single-cell RNA-seq dataset from Zeisel et al. (2015).

Usage

```
ZeiselBrainData(ensembl = FALSE, location = TRUE)
```

Arguments

ensembl	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
location	Logical scalar indicating whether genomic coordinates should be returned.

Details

Row data contains a single "featureType" field describing the type of each feature (endogenous genes, mitochondrial genes, spike-in transcripts and repeats). Spike-ins and repeats are stored as separate entries in the [altExps](#).

Column metadata is provided in the same form as supplied in <http://linnarssonlab.org/cortex/>. This contains information such as the cell diameter and the published cell type annotations.

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the [rowRanges](#) of the output.

Spike-in metadata is added using [ERCCSpikeInConcentrations](#), with molecule counts computed using a volume of 9 nL per cell at a dilution of 1:20000.

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/zeisel-brain`.

Value

A [SingleCellExperiment](#) object with a single matrix of UMI counts.

Author(s)

Aaron Lun

References

Zeisel A et al. (2015). Brain structure. Cell types in the mouse cortex and hippocampus revealed by single-cell RNA-seq. *Science* 347(6226), 1138-42.

Examples

```
sce <- ZeiselBrainData()
```

ZilionisLungData	<i>Obtain the Zilionis lung cancer data</i>
------------------	---

Description

Obtain the human/mouse lung cancer single-cell RNA-seq data from Zilionis et al. (2019).

Usage

```
ZilionisLungData(  
  which = c("human", "mouse"),  
  ensembl = FALSE,  
  location = TRUE,  
  filter = FALSE  
)
```

Arguments

which	String specifying the species to get data for.
ensembl	Logical scalar indicating whether the output row names should contain Ensembl identifiers.
location	Logical scalar indicating whether genomic coordinates should be returned.
filter	Logical scalar indicating if the filtered subset should be returned.

Details

Column metadata is provided and contains information on the library, donor ID/animal ID, replicate and tissue.

If `ensembl=TRUE`, the gene symbols are converted to Ensembl IDs in the row names of the output object. Rows with missing Ensembl IDs are discarded, and only the first occurrence of duplicated IDs is retained.

If `location=TRUE`, the coordinates of the Ensembl gene models are stored in the `rowRanges` of the output. Note that this is only performed if `ensembl=TRUE`.

If `filter=TRUE`, only cells that have been used in the original analysis are returned. The cells used are specified in the `Used` column of the `colData`.

The `reducedDim` contains coordinates of SPRING representations. This may be filled with NAs for SPRING coordinates computed on a subset of cells (specified in `colData`).

All data are downloaded from ExperimentHub and cached for local re-use. Specific resources can be retrieved by searching for `scRNAseq/zilionis-lung`.

Value

A `SingleCellExperiment` object with a single matrix of read counts.

Author(s)

Jens Preussner

References

Zilionis R et al. (2019). Single-cell transcriptomics of human and mouse lung cancers reveals conserved myeloid populations across individuals and species. *Immunity* 50(5), 1317-1334.

Examples

```
sce.human <- ZilionisLungData()
```

```
sce.mouse <- ZilionisLungData("mouse")
```

Index

- allen (scRNAseq-package), 2
- altExp, 20
- altExps, 6, 11, 14, 20, 24–26, 28, 30–32, 35, 39
- AztekinTailData, 3
- BachMammaryData, 4
- BaronPancreasData, 5
- BuettnerESCDData, 6
- CampbellBrainData, 7
- ChenBrainData, 8
- colData, 26, 40
- DataFrame, 9, 19
- ERCCSpikeInConcentrations, 9, 39
- fluidigm (scRNAseq-package), 2
- GrunHSCData, 10
- GrunPancreasData, 11
- HermannSpermatogenesisData, 12
- HuCortexData, 13
- KolodziejczykESCDData, 14
- KotliarovPBMCData, 15
- LaMannoBrainData, 16
- LawlorPancreasData, 17
- LengESCDData, 18
- listDatasets, 19
- LunSpikeInData, 20
- MacoskoRetinaData, 21
- MairPBMCData, 22
- MarquesBrainData, 23
- MessmerESCDData, 24
- metadata, 28
- MuraroPancreasData, 25
- NestorowaHSCData, 26
- PaulHSCData, 27
- reducedDim, 40
- reducedDims, 3, 26
- ReprocessedAllenData, 28
- ReprocessedFluidigmData (ReprocessedAllenData), 28
- ReprocessedTh2Data (ReprocessedAllenData), 28
- RichardTCellData, 29
- RomanovBrainData, 30
- rowData, 32
- rowRanges, 4–8, 10, 11, 13–15, 17, 18, 20–28, 30–40
- scRNAseq (scRNAseq-package), 2
- scRNAseq-package, 2
- SegerstolpePancreasData, 31
- ShekharRetinaData, 32
- SingleCellExperiment, 4–8, 10–14, 16–27, 29–40
- StoeckiusHashingData, 33
- TasicBrainData, 28, 35
- th2 (scRNAseq-package), 2
- UsoskinBrainData, 36
- WuKidneyData, 37
- XinPancreasData, 38
- ZeiselBrainData, 39
- ZilionisLungData, 40