Package 'TENxBUSData'

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Type Package

Title Single cell dataset from 10x in BUS format

Maintainer Lambda Moses <dlu2@caltech.edu>

```
Version 1.19.0
Description Download Barcode, UMI, and Set (BUS) format of 10x datasets from within R.
     This package accompanies the package BUSpaRse, which can load BUS format
     into R as a sparse matrix, and which has utility functions related to using
     the C++ command line package bustools.
License BSD_2_clause + file LICENSE
Encoding UTF-8
Imports ExperimentHub, AnnotationHub, utils, BiocGenerics
biocViews SingleCellData, SequencingData, ExperimentHub,
     ExperimentData, ExpressionData
BugReports https://github.com/BUStools/TENxBUSData/issues
URL https://github.com/BUStools/TENxBUSData
RoxygenNote 7.1.0
Roxygen list(markdown = TRUE)
Suggests knitr, rmarkdown, BiocStyle
VignetteBuilder knitr
git_url https://git.bioconductor.org/packages/TENxBUSData
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Author Lambda Moses [aut, cre] (<a href="https://orcid.org/0000-0002-7092-9427">https://orcid.org/0000-0002-7092-9427</a>)
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Description

This package provides 5 10x datasets in the BUS format, to be downloaded from within R. The files downloaded from this package are sufficient to generate a sparse matrix with package BUSpaRse to be used for downstream analysis with Seurat. Human and mouse transcriptomes from Ensembl version 99 were used to generate the BUS format from FASTQ files. This package server the following purposes: First, to demonstrate the kallisto bus workflow and downstream analyses. Second, for advanced users to experiment with other ways to collapse UMIs mapped to multiple genes and with other ways of barcode correction. The datasets are on ExperimentHub.

This function will download the 10x datasets, already processed and stored in the BUS format, from ExperimentHub. This function will decompress the downloaded file and return the directory where the files necessary to construct the sparse matrix with BUSpaRse are located.

Usage

```
TENxBUSData(
  file_path,
  dataset = c("hgmm100", "hgmm1k", "pbmc1k", "neuron10k"),
  force = FALSE,
  verbose = TRUE
)
```

Arguments

file_path	Character vector of length 1, specifying where to download the data.
dataset	Character, must be one of "hgmm100", "hgmm1k", "pbmc1k", "neuron10k".
force	Logical, whether to force redownload if the files are already present. Defaults to $\ensuremath{FALSE}.$
verbose	Whether to display progress of download.

Details

The following files will be downloaded:

matrix.ec Text file with 2 columns. The first column is the index of equivalence classes used in BUS files. The second column is the equivalence classes themselves, consisted of sets of transcript indices from the kallisto index.

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output.sorted Binary BUS file sorted by barcode, UMI, and equivalence classes. This can be directly parsedd by bustools to get the gene count matrix.

output.sorted.txt Sorted BUS file in text form. This is to be used with BUSpaRse to get the gene count matrix.

transcripts.txt Transcripts included in the BUS data, in the same order as in the kallisto index.

The gzipped file downloaded from ExperimentHub will be in a cache directory that can be retrieved by getExperimentHubOption("CACHE"). The cache will remain even if the decompressed files in the directory specified when calling this function are deleted. To delete cache, use removeCache.

Value

Character, directory to be used in BUSpaRse.

Datasets

```
hgmm100 100 1:1 Mixture of Fresh Frozen Human (HEK293T) and Mouse (NIH3T3) Cells. The
    raw data can be found here: https://support.10xgenomics.com/single-cell-gene-expression/
    datasets/2.1.0/hgmm_100

hgmm1k lk 1:1 Mixture of Fresh Frozen Human (HEK293T) and Mouse (NIH3T3) Cells (v3
        chemistry). The raw data can be found here: https://support.10xgenomics.com/single-cell-gene-expression
    datasets/3.0.0/hgmm_1k_v3

pbmc1k lk PBMCs from a Healthy Donor (v3 chemistry). The raw data can be found here: https:
    //support.10xgenomics.com/single-cell-gene-expression/datasets/3.0.0/pbmc_1k_
    v3

neuron10k 10k Brain Cells from an E18 Mouse (v3 chemistry). The raw data can be found
    here: https://support.10xgenomics.com/single-cell-gene-expression/datasets/
    3.0.0/neuron_10k_v3
```

Examples

```
TENxBUSData(".", dataset = "hgmm100")
```

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