Package 'SCIBER'

May 2, 2023

Type Package
Title Single-Cell Integrator and Batch Effect Remover
Version 0.2.2
Description Remove batch effects by projecting query batches into the reference batch space.
License MIT + file LICENSE
Encoding UTF-8
RoxygenNote 7.2.1
LazyData true
<pre>URL https://github.com/RavenGan/SCIBER</pre>
<pre>BugReports https://github.com/RavenGan/SCIBER/issues</pre>
Imports dplyr, parallel, stats
Depends R (>= 2.10)
Suggests knitr, rmarkdown, testthat (>= 3.0.0), cowplot, ggplot2, ggthemes, Matrix, uwot
Config/testthat/edition 3
VignetteBuilder knitr
NeedsCompilation no
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R topics documented:
HumanDC
Index

2 SCIBER

HumanDC

Human dendritic cell gene expression and meta data

Description

A dataset human dendritic cells gene expression and corresponding meta data

Usage

 ${\it HumanDC}$

Format

An object of class list of length 2.

Source

```
https://pubmed.ncbi.nlm.nih.gov/28428369/
```

SCIBER

Batch effect removal with SCIBER

Description

Batch effect removal with SCIBER

Usage

```
SCIBER(
  input_batches,
  ref_index = NULL,
  batches_meta_data = NULL,
  omega = 0.5,
  h_fisher = 75,
  n_core = parallel::detectCores(),
  seed = 7,
  k = NULL
)
```

SCIBER 3

Arguments

input_batches A list contains all the pre-processed matrices with dimension of n_genes*n_cells.

ref_index The index of the reference batch in the object "input_batches"

batches_meta_data

A list contains the meta data for all the batches. The order should be consistent with that in "input_batches". Each meta data contains three columns, "cell_id", "cell_type", and "dataset". "dataset" indicates which batch the data comes from.

The row names of meta data should match the column names of batch.

omega A list of proportion of matched clusters or a single value between 0 and 1 applied

to all query batches.

h_fisher The number of marker genes used for Fisher exact test.

n_core Specify the number of cores otherwise use all the available cores.

seed random seed.

k Number of clusters used for K-means. If not provided, the default is k = the

square root of n_0, where n_0 is the number of cells in the reference batch.

Value

A list which contains the reference and batch-effect-corrected batches. The order is the same as that in input_batches.

Examples

```
data(HumanDC)
exp <- HumanDC[["exp"]]
meta <- HumanDC[["metadata"]]
omega <- c()
omega[[1]] <- 0.5
res <- SCIBER(input_batches = exp, ref_index = 1,
batches_meta_data = meta, omega = omega, n_core = 1)</pre>
```

Index

$*\ datasets$

HumanDC, 2

HumanDC, 2

SCIBER, 2