Package 'scISR'

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Type Package
Title Single-Cell Imputation using Subspace Regression
Version 0.1.1
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Description Provides an imputation pipeline for single-cell RNA sequencing data. The 'scISR' method uses a hypothesis-testing technique to identify zero-valued entries that are most likely affected by dropout events and estimates the dropout values using a subspace regression model (Tran et.al. (2022) <doi:10.1038 s41598-022-06500-4="">).</doi:10.1038>
License LGPL
Depends R (>= 3.4)
Imports cluster, entropy, stats, utils, parallel, irlba, PINSPlus, matrixStats, markdown
Encoding UTF-8
LazyData true
RoxygenNote 7.1.1
NeedsCompilation no
Suggests testthat, knitr, mclust
VignetteBuilder knitr
<pre>URL https://github.com/duct317/scISR</pre>
<pre>BugReports https://github.com/duct317/scISR/issues</pre>
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Goolam Goolam

Description

Goolam dataset with data and cell types information. The number of genes is reduced to 10,000.

Usage

Goolam

Format

An object of class list of length 2.

scISR

scISR: Single-cell Imputation using Subspace Regression

Description

Perform single-cell Imputation using Subspace Regression

Usage

```
scISR(
  data,
  ncores = 1,
  force_impute = FALSE,
  do_fast = TRUE,
  preprocessing = TRUE,
  batch_impute = FALSE,
  seed = 1
)
```

Arguments

data Input matrix or data frame. Rows represent genes while columns represent sam-

ples

ncores Number of cores that the algorithm should use. Default value is 1.

force_impute Always perform imputation.

preprocessing Perform preprocessing on original data to filter out low quality features.

batch_impute Perform imputation in batches to reduce memory consumption.

seed Seed for reproducibility. Default value is 1.

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Details

scISR performs imputation for single-cell sequencing data. scISR identifies the true dropout values in the scRNA-seq dataset using hyper-geometric testing approach. Based on the result obtained from hyper-geometric testing, the original dataset is segregated into two subsets including training data and imputable data. Next, training data is used for constructing a generalize linear regression model that is used for imputation on the imputable data.

Value

scISR returns an imputed single-cell expression matrix where rows represent genes while columns represent samples.

Examples

```
# Load the package
library(scISR)
# Load Goolam dataset
data('Goolam');
# Use only 500 random genes for example
set.seed(1)
raw <- Goolam$data[sample(seq_len(nrow(Goolam$data)), 500), ]</pre>
label <- Goolam$label</pre>
# Perform the imputation
imputed <- scISR(data = raw)</pre>
if(requireNamespace('mclust'))
{
 library(mclust)
 # Perform PCA and k-means clustering on raw data
 set.seed(1)
 # Filter genes that have only zeros from raw data
 raw_filer <- raw[rowSums(raw != 0) > 0, ]
 pca_raw <- irlba::prcomp_irlba(t(raw_filer), n = 50)$x</pre>
 cluster_raw <- kmeans(pca_raw, length(unique(label)),</pre>
                         nstart = 2000, iter.max = 2000)$cluster
 print(paste('ARI of clusters using raw data:',
              round(adjustedRandIndex(cluster_raw, label),3)))
 # Perform PCA and k-means clustering on imputed data
 set.seed(1)
 pca_imputed <- irlba::prcomp_irlba(t(imputed), n = 50)$x</pre>
 cluster_imputed <- kmeans(pca_imputed, length(unique(label)),</pre>
                             nstart = 2000, iter.max = 2000)$cluster
 print(paste('ARI of clusters using imputed data:',
              round(adjustedRandIndex(cluster_imputed, label),3)))
}
}
```

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