

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](https://doi.org/10.1038/s41598-022-06500-4)>).

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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

URL <https://github.com/duct317/scISR>

BugReports <https://github.com/duct317/scISR/issues>

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Repository CRAN

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Goolam	<i>Goolam</i>
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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	<i>scISR: Single-cell Imputation using Subspace Regression</i>
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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 samples
ncores	Number of cores that the algorithm should use. Default value is 1.
force_impute	Always perform imputation.
do_fast	Use fast imputation implementation.
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.

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 generalized 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), ]
label <- Goolam$label

# Perform the imputation
imputed <- scISR(data = raw)

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)$x
  cluster_raw <- kmeans(pca_raw, length(unique(label)),
    nstart = 2000, iter.max = 2000)$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)$x
  cluster_imputed <- kmeans(pca_imputed, length(unique(label)),
    nstart = 2000, iter.max = 2000)$cluster
  print(paste('ARI of clusters using imputed data:',
    round(adjustedRandIndex(cluster_imputed, label),3)))
}
}
```

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