

# Package ‘rescue’

October 14, 2022

**Type** Package

**Title** Bootstrap Imputation for Single-Cell RNA-Seq Data

**Version** 1.0.3

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**Description** Given a log-transformed expression matrix and list of informative genes: subsample informative genes, cluster samples using shared nearest neighbors clustering, estimate missing expression values with the distribution mean of means extrapolated from these cell clusterings, and return an imputed expression matrix. See Tracy, S., Yuan, G.C. and Dries, R. (2019) <[doi:10.1186/s12859-019-2977-0](https://doi.org/10.1186/s12859-019-2977-0)> for more details.

**Config/reticulate** `list( packages = list( list(package = ``pandas"), list(package = ``networkx"), list(package = ``python-louvain") ) )`

**Depends** R (>= 3.4.0), utils

**Imports** data.table, dbscan (>= 1.1-3), igraph (>= 1.2.4.1), irlba, Matrix, methods, parallel, reticulate (>= 1.14)

**License** GPL-2 | GPL-3

**LazyData** FALSE

**URL** <https://github.com/seasamgo/rescue>

**BugReports** <http://github.com/seasamgo/rescue/issues>

**RoxygenNote** 7.1.1

**Encoding** UTF-8

**Suggests** knitr, rmarkdown

**NeedsCompilation** no

**Repository** CRAN

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**R topics documented:**

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bootstrapImputation	<i>Bootstrap Imputation</i>
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**Description**

Subsample informative genes, cluster cells using SNN, estimate missing expression values with the distribution mean of means extrapolated from these cell clusterings

**Usage**

```
bootstrapImputation(
  expression_matrix,
  select_cells = NULL,
  select_genes = NULL,
  log_transformed = TRUE,
  log_base = exp(1),
  proportion_genes = 0.6,
  bootstrap_samples = 100,
  number_pcs = 8,
  k_neighbors = 30,
  snn_resolution = 0.9,
  impute_index = NULL,
  use_mclapply = FALSE,
  cores = 2,
  return_individual_results = FALSE,
  python_path = NULL,
  verbose = FALSE
)
```

**Arguments**

expression_matrix	Row by column log-normalized expression matrix
select_cells	Subset cells if desired
select_genes	A vector of highly variable of differentially expressed gene names, defaults to the most variable
log_transformed	Whether the expression matrix has been log-transformed

log_base	If log-transformed, log-base used
proportion_genes	Proportion of informative genes to sample
bootstrap_samples	Number of samples for the bootstrap
number_pcs	Number of dimensions to inform SNN clustering
k_neighbors	Number of k neighbors to use for NN network
snn_resolution	Resolution parameter for SNN
impute_index	Index to impute, will default to all zeroes
use_mclapply	Run in parallel, default FALSE
cores	Number of cores for parallelization
return_individual_results	Return a list of subsampled means
python_path	path to your python binary (default = system path)
verbose	Print progress output to the console

**Value**

Returns a list with the imputed and original expression matrices

**Examples**

```

set.seed(0)
requireNamespace("Matrix")

## generate (meaningless) counts
c1 <- stats::rpois(5e3, 1)
c2 <- stats::rpois(5e3, 2)
m <- t(
  rbind(
    matrix(c1, nrow = 20),
    matrix(c2, nrow = 20)
  )
)

## construct an expression matrix m
colnames(m) <- paste0('cell', 1:ncol(m))
rownames(m) <- paste0('gene', 1:nrow(m))
m <- log(m/colSums(m)*1e4 + 1)
m <- methods::as(m, 'dgCMatrix')

## impute

m_imputed <- rescue::bootstrapImputation(
  expression_matrix = m,
  proportion_genes = .9,
  bootstrap_samples = 2,
  k_neighbors = 10

```

)

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clusterLouvain	<i>Cluster Cells via Louvain Algorithm</i>
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**Description**

Cluster cells using a NN-network and the Louvain algorithm from the community module in Python

**Usage**

```
clusterLouvain(
  nn_network,
  python_path = NULL,
  resolution = 1,
  weight_col = NULL,
  louv_random = F,
  set_seed = T,
  seed_number = 0,
  ...
)
```

**Arguments**

nn_network	Constructed nearest neighbor network to use
python_path	Specify specific path to python if required
resolution	Resolution
weight_col	Weight column
louv_random	Random
set_seed	Set seed
seed_number	Number for seed
...	Additional parameters

**Value**

A character vector of cluster labels

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`computeHVG`*Compute Highly Variable Genes*

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**Description**

Compute Highly Variable Genes

**Usage**

```
computeHVG(  
  expression_matrix,  
  reverse_log_scale = T,  
  log_base = exp(1),  
  expression_threshold = 0,  
  nr_expression_groups = 20,  
  zscore_threshold = 1.5  
)
```

**Arguments**

<code>expression_matrix</code>	Expression matrix
<code>reverse_log_scale</code>	Reverse log-scale of expression values
<code>log_base</code>	If <code>reverse_log_scale</code> is TRUE, which log base was used?
<code>expression_threshold</code>	Expression threshold to consider a gene detected
<code>nr_expression_groups</code>	Number of expression groups for <code>cov_groups</code>
<code>zscore_threshold</code>	Z-score to select hvg for <code>cov_groups</code>

**Value**

Character vector of highly variable genes

**Examples**

```
set.seed(0)  
requireNamespace("Matrix")  
  
## generate (meaningless) counts  
c1 <- stats::rpois(5e3, 1)  
c2 <- stats::rpois(5e3, 2)  
m <- t(  
  rbind(  
    matrix(c1, nrow = 20),
```

```

    matrix(c2, nrow = 20)
  )
)

## construct an expression matrix m
colnames(m) <- paste0('cell', 1:ncol(m))
rownames(m) <- paste0('gene', 1:nrow(m))
m <- log(m/colSums(m)*1e4 + 1)
m <- methods::as(m, 'dgCMatrix')

## calculate HVGs
hvgs <- computeHVG(m)

```

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constructNN

*Nearest Network*


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## Description

Construct a nearest neighbour network based on previously computed PCs

## Usage

```

constructNN(
  reduced_object,
  k_neighbors = 30,
  minimum_shared = 5,
  top_shared = 3,
  verbose = F,
  ...
)

```

## Arguments

reduced_object	PC reduction matrix
k_neighbors	Number of k neighbors to use
minimum_shared	Minimum shared neighbors
top_shared	Keep at ...
verbose	Be verbose
...	Additional parameters

## Value

NN network as igraph object

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sampleImputation	<i>Sample-mean Estimation</i>
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## Description

Cluster cells using SNN and a list of given genes, estimate missing expression values for each cell-gene combination with the within-cluster non-zero expression mean

## Usage

```
sampleImputation(  
  expression_matrix,  
  subset_genes = NULL,  
  scale_data = TRUE,  
  number_pcs = 8,  
  k_neighbors = 30,  
  snn_resolution = 0.9,  
  impute_index = NULL,  
  pseudo_zero = NULL,  
  python_path = NULL,  
  verbose = FALSE  
)
```

## Arguments

<code>expression_matrix</code>	Row by column log-normalized expression matrix
<code>subset_genes</code>	A vector of informative gene names, defaults to all genes
<code>scale_data</code>	Whether to standardize expression by gene, default TRUE
<code>number_pcs</code>	Number of dimensions to inform SNN clustering
<code>k_neighbors</code>	Number of k neighbors to use for NN network
<code>snn_resolution</code>	Resolution parameter for SNN
<code>impute_index</code>	Index to impute, will default to all zeroes
<code>pseudo_zero</code>	Pseudo-zero expression value
<code>python_path</code>	path to your python binary (default = system path)
<code>verbose</code>	Print progress output to the console

## Value

Returns a sparse matrix of class 'dgCMatrix'

**Examples**

```
set.seed(0)
requireNamespace("Matrix")

## generate (meaningless) counts
c1 <- stats::rpois(5e3, 1)
c2 <- stats::rpois(5e3, 2)
m <- t(
  rbind(
    matrix(c1, nrow = 20),
    matrix(c2, nrow = 20)
  )
)

## construct an expression matrix m
colnames(m) <- paste0('cell', 1:ncol(m))
rownames(m) <- paste0('gene', 1:nrow(m))
m <- log(m/colSums(m)*1e4 + 1)
m <- methods::as(m, 'dgCMatrix')

## impute

m_imputed <- rescue::sampleImputation(
  expression_matrix = m,
  k_neighbors = 10
)
```



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