

Package ‘SIMLR’

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Title Single-cell Interpretation via Multi-kernel LeaRning (SIMLR)

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Imports parallel, Matrix, stats, methods, Rcpp, pracma, RcppAnnoy,
RSpectra

Suggests BiocGenerics, BiocStyle, testthat, knitr, igraph

Description Single-cell RNA-seq technologies enable high throughput gene expression measurement of individual cells, and allow the discovery of heterogeneity within cell populations. Measurement of cell-to-cell gene expression similarity is critical for the identification, visualization and analysis of cell populations. However, single-cell data introduce challenges to conventional measures of gene expression similarity because of the high level of noise, outliers and dropouts. We develop a novel similarity-learning framework, SIMLR (Single-cell Interpretation via Multi-kernel LeaRning), which learns an appropriate distance metric from the data for dimension reduction, clustering and visualization.

Encoding UTF-8

LazyData TRUE

License file LICENSE

URL <https://github.com/BatzogloulabSU/SIMLR>

BugReports <https://github.com/BatzogloulabSU/SIMLR>

biocViews ImmunoOncology, Clustering, GeneExpression, Sequencing,
SingleCell

RoxygenNote 7.1.0

LinkingTo Rcpp

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| BuettnerFlorian | <i>test dataset for SIMLR</i> |
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Description

example dataset to test SIMLR from the work by Buettner, Florian, et al.

Usage

```
data(BuettnerFlorian)
```

Format

gene expression measurements of individual cells

Value

list of 6: `in_X` = input dataset as an (m x n) gene expression measurements of individual cells, `n_clust` = number of clusters (number of distinct true labels), `true_labs` = ground true of cluster assignments for each of the `n_clust` clusters, `seed` = seed used to compute the results for the example, `results` = result by SIMLR for the inputs defined as described, `nmi` = normalized mutual information as a measure of the inferred clusters compared to the true labels

Source

Buettner, Florian, et al. "Computational analysis of cell-to-cell heterogeneity in single-cell RNA-sequencing data reveals hidden subpopulations of cells." *Nature biotechnology* 33.2 (2015): 155-160.

 CIMLR

please refer to <https://github.com/danro9685/CIMLR>

Description

perform the CIMLR clustering algorithm

Usage

```
CIMLR(X, c, no.dim = NA, k = 10, cores.ratio = 1)
```

Arguments

| | |
|--------------------------|--|
| <code>X</code> | a list of multi-omic data each of which is an (m x n) data matrix of measurements of cancer patients |
| <code>c</code> | number of clusters to be estimated over X |
| <code>no.dim</code> | number of dimensions |
| <code>k</code> | tuning parameter |
| <code>cores.ratio</code> | ratio of the number of cores to be used when computing the multi-kernel |

Value

clusters the patients based on CIMLR and their similarities

list of 8 elements describing the clusters obtained by CIMLR, of which y are the resulting clusters: y = results of k-means clusterings, S = similarities computed by CIMLR, F = results from network diffusion, ydata = data referring the the results by k-means, alphaK = clustering coefficients, execution.time = execution time of the present run, converge = iterative convergence values by T-SNE, LF = parameters of the clustering

Examples

```
clusters = 3
## Not run:
CIMLR(X = GliomasReduced$in_X, c = clusters, cores.ratio = 0)

## End(Not run)
```

 CIMLR_Estimate_Number_of_Clusters

please refer to <https://github.com/danro9685/CIMLR>

Description

estimate the number of clusters by means of two huristics as discussed in the CIMLR paper

Usage

```
CIMLR_Estimate_Number_of_Clusters(all_data, NUMC = 2:5, cores.ratio = 1)
```

Arguments

| | |
|--------------------------|---|
| <code>all_data</code> | is a list of multi-omic data each of which is an (m x n) data matrix of measurements of cancer patients |
| <code>NUMC</code> | vector of number of clusters to be considered |
| <code>cores.ratio</code> | ratio of the number of cores to be used when computing the multi-kernel |

Value

a list of 2 elements: K1 and K2 with an estimation of the best clusters (the lower values the better) as discussed in the original paper of SIMLR

Examples

```
clusters = 2:5
## Not run:
CIMLR_Estimate_Number_of_Clusters(GliomasReduced$in_X,
  NUMC = clusters,
  cores.ratio = 0)

## End(Not run)
```

SIMLR

*SIMLR***Description**

perform the SIMLR clustering algorithm

Usage

```
SIMLR(
  X,
  c,
  no.dim = NA,
  k = 10,
  if.impute = FALSE,
  normalize = FALSE,
  cores.ratio = 1
)
```

Arguments

| | |
|--------------------------|---|
| <code>X</code> | an (m x n) data matrix of gene expression measurements of individual cells or and object of class <code>SCESet</code> |
| <code>c</code> | number of clusters to be estimated over <code>X</code> |
| <code>no.dim</code> | number of dimensions |
| <code>k</code> | tuning parameter |
| <code>if.impute</code> | should I transpose the input data? |
| <code>normalize</code> | should I normalize the input data? |
| <code>cores.ratio</code> | ratio of the number of cores to be used when computing the multi-kernel |

Value

clusters the cells based on SIMLR and their similarities

list of 8 elements describing the clusters obtained by SIMLR, of which y are the resulting clusters:
 y = results of k-means clusterings, S = similarities computed by SIMLR, F = results from network diffusion, ydata = data referring the the results by k-means, alphaK = clustering coefficients, execution.time = execution time of the present run, converge = iterative convergence values by T-SNE, LF = parameters of the clustering

Examples

```
SIMLR(X = BuettnerFlorian$in_X, c = BuettnerFlorian$n_clust, cores.ratio = 0)
```

SIMLR_Estimate_Number_of_Clusters

SIMLR Estimate Number of Clusters

Description

estimate the number of clusters by means of two huristics as discussed in the SIMLR paper

Usage

```
SIMLR_Estimate_Number_of_Clusters(X, NUMC = 2:5, cores.ratio = 1)
```

Arguments

| | |
|-------------|--|
| X | an (m x n) data matrix of gene expression measurements of individual cells |
| NUMC | vector of number of clusters to be considered |
| cores.ratio | ratio of the number of cores to be used when computing the multi-kernel |

Value

a list of 2 elements: K1 and K2 with an estimation of the best clusters (the lower values the better) as discussed in the original paper of SIMLR

Examples

```
SIMLR_Estimate_Number_of_Clusters(BuettnerFlorian$in_X,
  NUMC = 2:5,
  cores.ratio = 0)
```

SIMLR_Feature_Ranking *SIMLR Feature Ranking*

Description

perform the SIMLR feature ranking algorithm. This takes as input the original input data and the corresponding similarity matrix computed by SIMLR

Usage

```
SIMLR_Feature_Ranking(A, X)
```

Arguments

| | |
|---|--|
| A | an (n x n) similarity matrix by SIMLR |
| X | an (m x n) data matrix of gene expression measurements of individual cells |

Value

a list of 2 elements: pvalues and ranking ordering over the n covariates as estimated by the method

Examples

```
SIMLR_Feature_Ranking(A = BuettnerFlorian$results$S, X = BuettnerFlorian$in_X)
```

SIMLR_Large_Scale *SIMLR Large Scale*

Description

perform the SIMLR clustering algorithm for large scale datasets

Usage

```
SIMLR_Large_Scale(X, c, k = 10, kk = 100, if.impute = FALSE, normalize = FALSE)
```

Arguments

| | |
|-----------|--|
| X | an (m x n) data matrix of gene expression measurements of individual cells or and object of class SCESet |
| c | number of clusters to be estimated over X |
| k | tuning parameter |
| kk | number of principal components to be assessed in the PCA |
| if.impute | should I traspose the input data? |
| normalize | should I normalize the input data? |

Value

clusters the cells based on SIMLR Large Scale and their similarities

list of 8 elements describing the clusters obtained by SIMLR, of which y are the resulting clusters: y = results of k-means clusterings, S0 = similarities computed by SIMLR, F = results from the large scale iterative procedure, ydata = data referring the the results by k-means, alphaK = clustering coefficients, val = distances from the k-nearest neighbour search, ind = indeces from the k-nearest neighbour search, execution.time = execution time of the present run

Examples

```
resized = ZeiselAmit$in_X[, 1:340]
## Not run:
SIMLR_Large_Scale(X = resized, c = ZeiselAmit$n_clust, k = 5, kk = 5)

## End(Not run)
```

ZeiselAmit

test dataset for SIMLR large scale

Description

example dataset to test SIMLR large scale. This is a reduced version of the dataset from the work by Zeisel, Amit, et al.

Usage

```
data(ZeiselAmit)
```

Format

gene expression measurements of individual cells

Value

list of 6: in_X = input dataset as an (m x n) gene expression measurements of individual cells, n_clust = number of clusters (number of distinct true labels), true_labs = ground true of cluster assignments for each of the n_clust clusters, seed = seed used to compute the results for the example, results = result by SIMLR for the inputs defined as described, nmi = normalized mutual information as a measure of the inferred clusters compared to the true labels

Source

Zeisel, Amit, et al. "Cell types in the mouse cortex and hippocampus revealed by single-cell RNA-seq." *Science* 347.6226 (2015): 1138-1142.

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