## Package 'BisqueRNA'

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Description Provides tools to accurately estimate cell type abundances from heterogeneous bulk expression. A reference-based method utilizes single-cell information to generate a signature matrix and transformation of bulk expression for accurate regression based estimates. A marker-based method utilizes known cell-specific marker genes to measure relative abundances across samples.
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## CalculateSCCellProportions

Calculate cell proportions based on single-cell data

## Description

Returns proportion of each cell type out of total cells for each individual in the single-cell Expression Set

## Usage

CalculateSCCellProportions(sc.eset, subject.names, cell.types)

## Arguments

sc.eset Expression Set with single-cell data
subject. names A character string. Name of phenoData attribute in sc.eset that indicates individual ID.
cell.types A character string. Name of phenoData attribute in sc.eset that indicates cell type

Value
sc.props Matrix. Cell proportions with number of cell types rows by number of individuals columns
CorTri Correlate columns of data frame

## Description

This function runs correlation between markers of a data frame or matrix, returning the values of the lower/upper triangular of the correlation matrix in a vector.

## Usage

CorTri(x, method = "pearson")

## Arguments

## X

Data frame or matrix. Column vectors are correlated
method Character string. Name of method passed to cor. Pearson by default.

## Value

cors Numeric vector. Correlation coefficients of pairs
CountsToCPM Convert counts data in Expression Set to counts per million (CPM)

## Description

Convert counts data in Expression Set to counts per million (CPM)

## Usage

CountsToCPM(eset)

## Arguments

eset Expression Set containing counts assay data.

## Value

eset Expression Set containing CPM assay data

## Description

Estimate cell type proportions using first PC of expression matrix

## Usage

EstimatePCACellTypeProportions(x, weighted = FALSE, w = NULL)

## Arguments

X
weighted
w

A sample by gene bulk expression matrix. Genes should be marker genes Boolean. If weighted=TRUE, multiply scaled gene expression by gene weights
Numeric vector. Weights of genes

## Value

ret List. Attribute pcs contains matrix of PCs, where PC1 should be used as estimates for cell type abundances Attribute sdev contains eigenvalues of eigendecomposition of var-covar matrix. The 1 st eigenvalue should explain most of the variance. Attribute genes contains names of genes.

FilterUnexpressedGenes
Remove genes in Expression Set with zero expression in all samples

## Description

Remove genes in Expression Set with zero expression in all samples

## Usage

FilterUnexpressedGenes(eset, verbose = TRUE)

## Arguments

$$
\begin{array}{ll}
\text { eset } & \text { Expression Set } \\
\text { verbose } & \text { Boolean. Print logging info }
\end{array}
$$

## Value

eset Expression Set with zero expression genes removed

FilterZeroVarianceGenes
Remove genes in Expression Set with zero variance across samples

## Description

Remove genes in Expression Set with zero variance across samples

## Usage

FilterZeroVarianceGenes(eset, verbose = TRUE)

## Arguments

eset
Expression Set
verbose
Boolean. Print logging info

## Value

eset Expression Set with zero variance genes removed

## Description

Averages expression within each cell type across all samples to use as reference profile.

## Usage

GenerateSCReference(sc.eset, cell.types)

## Arguments

$$
\begin{array}{ll}
\text { sc.eset } & \text { Expression Set with single-cell data } \\
\text { cell.types } & \text { A character string. Name of phenoData attribute in sc.eset that indicates cell } \\
\text { type }
\end{array}
$$

## Value

sc.ref Matrix. Reference profile with number of gene rows by number of cell types columns.
GetCTP Return cell type proportions from bulk

## Description

Calculate cell type proportions from a data frame containing bulk expression values. Uses PCA (weighted or regular) to estimate relative proportions within each cell type.

## Usage

GetCTP(
bulk,
cell_types,
markers,
ct_col,
gene_col,
min_gene,
max_gene,
weighted,
w_col,
verbose
)

## Arguments

| bulk | Expression Set containing bulk data |
| :--- | :--- |
| cell_types | Character vector. Names of cell types. <br> markers |
| Data frame with columns specifying cluster and gene, and optionally a column <br> for weights, typically the fold-change of the gene. Important that the genes for <br> each cell type are row-sorted by signficance. <br> Character string. Column name specifying cluster/cell type corresponding to <br> each marker gene in markers. |  |
| gene_col | Character string. Column name specifying gene names in markers. |
| min_gene | Numeric. Min number of genes to use for each cell type. |
| max_gene | Numeric. Max number of genes to use for each cell type. |
| weighted | Boolean. Whether to use weights for gene prioritization |
| w_col | Character string. Column name for weights, such as "avg_logFC", in markers <br> Boolean. Whether to print log info during decomposition. Errors will be printed <br> regardless. |
|  |  |

Value
A List. Slot cors contains list of vectors with correlation coefficients. Slot ctps contains list of CTP objects returned by GetCTP

## Description

Get number of genes to use with no weighted information

## Usage

GetNumGenes(x, min.gene = 25, max.gene = 200)

## Arguments

x
min.gene $\quad$ Numeric. Minimum number of genes to consider as markers.
max.gene $\quad$ Numeric. Maximum number of genes to consider as markers.

Value
best.n Numeric. Number of genes to use

GetNumGenesWeighted Get number of genes to use with weighted PCA

## Description

Get number of genes to use with weighted PCA

## Usage

GetNumGenesWeighted(x, w, min.gene = 25, max.gene = 200)

## Arguments

$x \quad$ Numeric Matrix. A sample by gene expression matrix containing the marker genes.
w Numeric Vector. The weights of the genes that correspond to the columns of x .
min.gene Numeric. Minimum number of genes to consider as markers.
max.gene Numeric. Maximum number of genes to consider as markers.

## Value

best.n Numeric. Number of genes to use

GetOverlappingGenes Find overlapping genes in single-cell data, bulk data, and marker genes

## Description

Find overlapping genes in single-cell data, bulk data, and marker genes

## Usage

GetOverlappingGenes(sc.eset, bulk.eset, markers, verbose)

## Arguments

| sc.eset | Expression Set with single-cell data |
| :--- | :--- |
| bulk.eset | Expression Set with bulk data |
| markers | Character vector. List of relevant marker genes |
| verbose | Boolean. Print logging info |

## Value

overlapping.genes Character vector. List of genes found in markers and both datasets.

GetOverlappingSamples Find overlapping samples in single-cell and bulk data

## Description

Find overlapping samples in single-cell and bulk data

## Usage

GetOverlappingSamples(sc.eset, bulk.eset, subject.names, verbose)

## Arguments

sc.eset Expression Set with single-cell data
bulk.eset Expression Set with bulk data
subject. names A character string. Name of phenoData attribute in sc.eset that indicates individual ID (that would be found in bulk.eset if overlapping)
verbose Boolean. Print logging info

## Value

samples A list with attributes overlapping and remaining. Each attribute refers to a character vector that lists the samples found in both datasets and samples found only in bulk, respectively

GetUniqueMarkers Get unique markers present in only 1 cell type

## Description

Given a data frame of marker genes for cell types, returns a new data frame with non-unique markers removed.

## Usage

GetUniqueMarkers(x, gene_col = "gene")

## Arguments

| $x$ | Data frame. Contains column with marker gene names |
| :--- | :--- |
| gene_col | Character string. Name of the column that contains the marker genes |

## Value

x Data frame. Markers with non-unique markers removed

## MarkerBasedDecomposition

Performs marker-based decomposition of bulk expression using marker genes

## Description

Estimates relative abundances of cell types from PCA-based decomposition. Uses a list of marker genes to subset the expression data, and returns the first PC of each sub-matrix as the cell type fraction estimates. Optionally, weights for each marker gene can be used to prioritize genes that are highly expressed in the given cell type.

## Usage

```
MarkerBasedDecomposition(
    bulk.eset,
    markers,
    ct_col = "cluster",
    gene_col = "gene",
    min_gene = 5,
    max_gene = 200,
    weighted = FALSE,
    w_col = "avg_logFC",
    unique_markers = TRUE,
    verbose = TRUE
)
```


## Arguments

| bulk.eset | Expression Set. Normalized bulk expression data. <br> markers |
| :--- | :--- |
| Data frame with columns specifying cluster and gene, and optionally a column <br> for weights, typically the fold-change of the gene. Important that the genes for <br> each cell type are row-sorted by signficance. |  |
| ct_col | Character string. Column name specifying cluster/cell type corresponding to <br> each marker gene in markers. |
| gene_col | Character string. Column name specifying gene names in markers. |
| min_gene | Numeric. Min number of genes to use for each cell type. |
| max_gene | Numeric. Max number of genes to use for each cell type. |
| weighted | Character string. Column name for weights, such as "avg_logFC", in markers |
| w_col | Boolean. If TRUE, subset markers to include only genes that are markers for <br> only one cell type |
| unique_markers weights for gene prioritization |  |

## Details

Note that this method expects the input bulk data to be normalized, unlike the reference-based method.

## Value

A List. Slot bulk.props contains estimated relative cell type abundances. Slot var.explained contains variance explained by first 20 PCs for cell type marker genes. Slot genes.used contains vector of genes used for decomposition.

## Examples

```
library(Biobase)
sim.data <- SimulateData(n.ind=10, n.genes=100, n.cells=100,
    cell.types=c("Neurons", "Astrocytes", "Microglia"),
    avg.props=c(.5, .3, .2))
res <- MarkerBasedDecomposition(sim.data$bulk.eset, sim.data$markers, weighted=FALSE)
estimated.cell.proportions <- res$bulk.props
```

```
ReferenceBasedDecomposition
    Performs reference-based decomposition of bulk expression using
    single-cell data
```


## Description

Generates a reference profile based on single-cell data. Learns a transformation of bulk expression based on observed single-cell proportions and performs NNLS regression on these transformed values to estimate cell proportions.

## Usage

ReferenceBasedDecomposition(
bulk.eset,
sc.eset,
markers = NULL,
cell.types = "cellType", subject.names = "SubjectName", use.overlap = TRUE, verbose = TRUE, old.cpm = TRUE
)

## Arguments

bulk.eset Expression Set containin bulk data. No PhenoData required but if overlapping option used, IDs returned by sampleNames(bulk.eset) should match those found in sc.eset phenoData individual labels.
sc.eset Expression Set containing single-cell data. PhenoData of this Expression Set should contain cell type and individual labels for each cell. Names of these fields specified by arguments below.
markers Structure, such as character vector, containing marker genes to be used in decomposition. 'base::unique(base::unlist(markers))' should return a simple vector containing each gene name. If no argument or NULL provided, the method will use all available genes for decomposition.
cell.types Character string. Name of phenoData attribute in sc.eset indicating cell type label for each cell
subject. names Character string. Name of phenoData attribute in sc.eset indicating individual label for each cell
use.overlap Boolean. Whether to use and expect overlapping samples in decomposition.
verbose Boolean. Whether to print log info during decomposition. Errors will be printed regardless.


#### Abstract

old.cpm Prior to version 1.0.4 (updated in July 2020), the package converted counts to CPM after subsetting the marker genes. Github user randel pointed out that the order of these operations should be switched. Thanks randel! This option is provided for replication of older BisqueRNA but should be enabled, especially for small marker gene sets. We briefly tested this change on the cortex and adipose datasets. The original and new order of operations produce estimates that have an average correlation of 0.87 for the cortex and 0.84 for the adipose within each cell type.


## Details

Expects read counts for both datasets, as they will be converted to counts per million (CPM). Two options available: Use overlapping indivudals found in both single-cell and bulk datasets to learn transformation or learn transformation from single-cell alone. The overlapping option is expected to have better performance.

## Value

A list. Slot bulk.props contains a matrix of cell type proportion estimates with cell types as rows and individuals as columns. Slot sc.props contains a matrix of cell type proportions estimated directly from counting single-cell data. Slot rnorm contains Euclidean norm of the residuals for each individual's proportion estimates. Slot genes.used contains vector of genes used in decomposition. Slot transformed.bulk contains the transformed bulk expression used for decomposition. These values are generated by applying a linear transformation to the CPM expression.

## Examples

```
library(Biobase)
sim.data <- SimulateData(n.ind=10, n.genes=100, n.cells=100,
    cell.types=c("Neurons", "Astrocytes", "Microglia"),
    avg.props=c(.5, .3, .2))
sim.data$sc.eset <- sim.data$sc.eset[,sim.data$sc.eset$SubjectName %in% as.character(6:10)]
res <- ReferenceBasedDecomposition(sim.data$bulk.eset, sim.data$sc.eset)
estimated.cell.proportions <- res$bulk.props
```

SemisupervisedTransformBulk

Transforms bulk expression of a gene using only single-cell data

## Description

For a specific gene, this function learns a transformation of the bulk expression to match the distribution produced by the single-cell based reference and observed single-cell based cell proportions.

## Usage

SemisupervisedTransformBulk(gene, Y.train, X.pred)

## Arguments

gene Character string. Gene name that corresponds to row in Y.train
Y.train Numeric Matrix. Number of gene rows by number of overlapping individuals columns. Contains weighted sum of reference profile by single-cell based cell proportion estimates for each individual
X.pred Numeric Matrix. Number of gene rows by number of remaining individuals columns. Contains observed bulk expression for each individual to be transformed.

## Value

Y.pred Numeric Matrix. One row for given gene by number of remaining individuals columns. Contains transformed bulk expression for each individual.

SeuratToExpressionSet Converts Seurat object to Expression Set

## Description

'SeuratToExpressionSet()' returns an Expression Set with phenotype data indicating cell type (cellType) and individual (SubjectName) for each cell in a Seurat object. Raw counts data is used for assay data.

## Usage

SeuratToExpressionSet (
seurat.object,
delimiter,
position,
version = c("v2", "v3")
)

## Arguments

seurat.object Seurat object with attributes raw.data, ident, and cell.names
delimiter $\quad$ Character to split cell names with to find individual ID.
position Integer indicating 1-indexed position of individual ID after splitting cell name with delimiter.
version Character string. Either "v2" or "v3. Seurat version used to create Seurat object.

## Details

Note that the Seurat and Biobase libraries should be attached before running this function. The delimiter and position arguments are used to infer the individual ID from the cell ID. For example, a delimiter of "-" and position of " 2 " indicates that the individual ID for the cell ID ACTG-2 would be 2 .

## Value

sc.eset Expression set containing relevant phenotype and individual data, cellType and SubjectName.

## Examples

```
library(Seurat)
library(Biobase)
# We make a class to emulate a Seurat v2 object for illustration only
setClass("testSeuratv2", representation(cell.names = "character",
                                    ident = "character",
                                    raw.data = "matrix"))
sc.counts <- matrix(0,nrow=3,ncol=3)
# These barcodes correspond to a delimiter of "-" and position 2 for individual id.
test.cell.names <- c("ATCG-1", "TAGC-2", "GTCA-3")
test.ident <- c("cell type a", "cell type b", "cell type c")
names(test.ident) <- test.cell.names
colnames(sc.counts) <- test.cell.names
test.seurat.obj <- new("testSeuratv2",
    cell.names=test.cell.names,
    ident=test.ident,
    raw.data=sc.counts)
single.cell.expression.set <- SeuratToExpressionSet(test.seurat.obj, delimiter='-',
                                    position=2, version="v2")
```


## Description

Generates a nucleotide barcode similar to those generated by 10 x chromium sequencing platforms for illustration purposes. Generates barcode and individual ID separated by '-' delimiter.

## Usage

SimulateBarcode(index, individual, barcode.length)

## Arguments

index Integer. Index of cell ID from 0 to barcode.length to the fourth power. Will generate a unique nucleotide barcode for each index.
individual Character. ID of individual that the cell is from.
barcode. length Integer. Length of nucleotide barcode.

## Value

Simulated barcode for cell from an individual

SimulateData Simulate data for decomposition illustration

## Description

Simulates bulk and single-cell expression, as well as marker genes and true proportions that can be used as an example of decomposition

## Usage

SimulateData(n.ind, n.genes, n.cells, cell.types, avg.props)

## Arguments

| n. ind | Integer. Number of individuals to simulate |
| :--- | :--- |
| n.genes | Integer. Number of genes to simulate |
| n.cells | Integer. Number of cells per individual for single-cell data |
| cell.types | Character vector. List of cell types to simulate |
| avg.props | Numeric vector. List of average proportions for given cell types. Should be <br> same length as cell.types and sum to 1 |

## Value

A list with simulated single-cell in slot 'sc.eset' and bulk in 'bulk.eset', as well as true proportions in 'props' and marker genes in 'markers'.

## Examples

```
library(Biobase)
sim.data <- SimulateData(n.ind=10, n.genes=100, n.cells=100,
    cell.types=c("Neurons", "Astrocytes", "Microglia"),
    avg.props=c(.5, .3, .2))
```

SupervisedTransformBulk

> Transforms bulk expression of a gene given overlapping data

## Description

For a specific gene, this function uses linear regression to learn a transformation of the bulk expression to match the values produced by the single-cell based reference and observed single-cell based cell proportions.

## Usage

SupervisedTransformBulk(gene, Y.train, X.train, X.pred)

## Arguments

gene Character string. Gene name that corresponds to row in Y.train
Y.train Numeric Matrix. Number of gene rows by number of overlapping individuals columns. Contains weighted sum of reference profile by single-cell based cell proportion estimates for each individual
X.train Numeric Matrix. Number of gene rows by number of overlapping individuals columns. Contains observed bulk expression for each individual
X.pred Numeric Matrix. Number of gene rows by number of remaining individuals columns. Contains observed bulk expression for each individual to be transformed.

## Details

If a linear transformation cannot be learned for a gene (zero variance in observed bulk or single-cell based weighted sums), a vector of NaNs will be returned of the expected length (length of X.pred)

## Value

Y.pred Numeric Matrix. One row for given gene by number of remaining individuals columns. Contains transformed bulk expression for each individual.

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