# Package 'MOSim'

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**Title** Multi-Omics Simulation (MOSim)

Version 2.4.0

**Description** MOSim package simulates multi-omic experiments that mimic regulatory mechanisms within the cell, allowing flexible experimental design including time course and multiple groups.

**Encoding** UTF-8

**Depends** R (>= 4.2.0)

License GPL-3

LazyData false

biocViews Software, TimeCourse, ExperimentalDesign, RNASeq

BugReports https://github.com/ConesaLab/MOSim/issues

URL https://github.com/ConesaLab/MOSim

**Imports** HiddenMarkov, zoo, IRanges, S4Vectors, dplyr, ggplot2, lazyeval, matrixStats, methods, rlang, stringi, stringr, scran, Seurat, Signac, edgeR, Rcpp

**Suggests** testthat, knitr, rmarkdown, codetools, BiocStyle, stats, utils, purrr, scales, tibble, tidyr, Biobase, scater, SingleCellExperiment, decor, markdown, Rsamtools, igraph, leiden, bluster

Collate 'AllClass.R' 'AllGeneric.R' 'Simulator.R' 'SimulatorRegion.R' 'ChIP-seq.R' 'DNase-seq.R' 'MOSim-package.R' 'functions.R' 'Simulation.R' 'MOSim.R' 'RNA-seq.R' 'data.R' 'simulate\_WGBS\_functions.R' 'methyl-seq.R' 'miRNA-seq.R' 'sc\_MOSim.R' 'sc\_coexpression.R' 'sparsim\_functions.R' 'zzz.R'

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# **Description**

Multiomics simulation package.

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# See Also

Useful links:

- https://github.com/ConesaLab/MOSim
- Report bugs at https://github.com/ConesaLab/MOSim/issues

associationList

Data to showcase scRNA and scATAC-seq association

# Description

Data to showcase scRNA and scATAC-seq association

# Usage

```
data("associationList")
```

# **Format**

A dataframe with two columns and rows according to gene/feature relationships

Peak\_ID ATAC chromosomic positions associated to genes

Gene\_ID RNA genes associated to peaks

@source Created in-house to serve as an example

4 check\_patterns

#### **Description**

Helper function to calculate mean expression per celltype

# Usage

```
calculate_mean_per_list_df(df, named_lists)
```

# Arguments

df dataframe of expression where columns are cells named\_lists list of which cells belong to each celltype

# **Examples**

```
rna <- data.frame(c1 = c(1.5, 15.5, 3.5, 20.5), c2 = c(2, 15, 4, 20), c3 = c(10, 1, 12, 13), c4 = c(11, 1, 13, 14)) cell_types <- list("ct1" = c(1,2), "ct2" = c(3, 4)) calculate_mean_per_list_df(rna, cell_types)
```

check\_patterns

check\_patterns

# Description

Function to check if the TRUE FALSE patterns have at least two rows that are opposite, we need this to be able to generate repressor regulators

# Usage

```
check_patterns(patterns)
```

# **Arguments**

patterns

tibble of TRUE FALSE values

#### Value

list of indices where the rows are opposite

discretize 5

#### **Examples**

discretize

Discretize ChIP-Seq counts to simulate a binary dataset

# Description

Discretize ChIP-Seq counts to simulate a binary dataset

#### Usage

```
discretize(df, omic)
```

# **Arguments**

df A MOSimulated object

omic Character string of the omic to transform into binary data

# Value

A regulator dataframe of 0 and 1

```
omic_list <- c("RNA-seq", "ChIP-seq")
rnaseq_simulation <- mosim(omics = omic_list,
    omicsOptions = c(omicSim("ChIP-seq", totalFeatures = 2500)))
rnaseq_simulated <- omicResults(rnaseq_simulation, omic_list)
discrete_ChIP <- discretize(rnaseq_simulated, "ChIP-seq")</pre>
```

6 is.declared

experimentalDesign

Retrieves the experimental design

# **Description**

Retrieves the experimental design

# Usage

```
experimentalDesign(simulation)
```

# **Arguments**

simulation

A MOSimulation object

#### Value

A data frame containing the experimental design used to simulate the data.

# **Examples**

```
omic_list <- c("RNA-seq")
rnaseq_simulation <- mosim(omics = omic_list)
# This will be a data frame with RNA-seq counts
design_matrix <- experimentalDesign(rnaseq_simulation)</pre>
```

is.declared

Check if a variable is declared.

#### **Description**

Check if a variable is declared.

# Usage

```
is.declared(object, key = NULL)
```

# Arguments

object

Variable name to check

key

Optional key to check inside object.

# Value

TRUE or FALSE indicating if the variable is initialized & non-empty.

# **Description**

This function generates a dataframe containing the information of the relationship between ATAC and RNA, based on the cluster groups, and then tells the order the genes and peaks should be in the simulated dataframe of the group

# Usage

```
make_association_dataframe(
   group,
   genereggroup,
   numtotalgenes,
   numtotalpeaks,
   minFC,
   maxFC
)
```

#### Arguments

group Group from which we are generating the association dataframe

genereggroup list of elements to generate the association dataframe such as clusters of each

omic, indices of opposite clusters, which genes are activated, repressed, behav-

ior of the features etc.

numtotalgenes total number of genes numtotalpeaks total number of peaks

minFC FC below which is downregulated

maxFC FC above which is upregulated

#### Value

a dataframe with all the information the user needs about each gene and the order of gene and peak names to rename them in the simulated datasets of the group

```
make_cluster_patterns make_cluster_patterns
```

#### **Description**

Function to make the tibble with cluster combinations for the gene expression patterns along the cells This function is a slightly modified copy of the 'make\_cluster\_patterns' function from the 'Acorde' package (v1.0.0), originally developed by Arzalluz-Luque A, Salguero P, Tarazona S, Conesa A. (2022). acorde unravels functionally interpretable networks of isoform co-usage from single cell data. Nature communications 1828. DOI: 10.1038/s41467-022-29497-w. The original package is licensed under the GPL-3 license.

#### Usage

```
make_cluster_patterns(numcells = 4, clusters = 8)
```

#### **Arguments**

numcells Number of different celltypes we are simulating

clusters OPTIONAL. Number of co-expression patterns the user wants to simulate

#### Value

A tibble with number of columns equal to number of celltypes, rows according to the number of TRUE/FALSE combinations corresponding to the gene expression patterns along the cells

#### **Examples**

```
match_gene_regulator match_gene_regulator
```

#### **Description**

Helper function to make the most similar profiles possible between gene and regulator

#### Usage

```
match_gene_regulator(rna, atac, cell_types, associationList)
```

#### **Arguments**

rna dataframe of RNA expression

atac dataframe of ATAC expression

cell\_types list of which cells belong to each celltype

associationList

dataframe of two columns, Gene\_ID and Peak\_ID

# **Examples**

#### **Description**

```
match_gene_regulator_cluster
```

# Usage

```
match_gene_regulator_cluster(rna, atac, cell_types, associationMatrix)
```

#### **Arguments**

```
rna rna expression dataframe

atac atac expression dataframe

cell_types list of which cells belong to each celltype

associationMatrix

matrix of related genes and peaks
```

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#### **Examples**

mosim

mosim

#### **Description**

Performs a multiomic simulation by chaining two actions: 1) Creating the "MOSimulation" class with the provided params. 2) Calling "simulate" method on the initialized object.

# Usage

```
mosim(
  omics,
  omicsOptions,
  diffGenes,
  numberReps,
  numberGroups,
  times,
  depth,
  profileProbs,
  minMaxFC,
  TFtoGene
)
```

#### **Arguments**

omics

Character vector containing the names of the omics to simulate, which can be "RNA-seq", "miRNA-seq", "DNase-seq", "ChIP-seq" or "Methyl-seq" (e.g. c("RNA-seq", "miRNA-seq")). It can also be a list with the omic names as names and their options as values, but we recommend to use the argument omic-Sim to provide the options to simulated each omic.

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omicsOptions

List containing the options to simulate each omic. We recommend to apply the helper method omicSim to create this list in a friendly way, and the function omicData to provide custom data (see the related sections for more information). Each omic may have different configuration parameters, but the common ones are:

**simuData/idToGene** Seed sample and association tables for regulatory omics. The helper function omicData should be used to provide this information (see the following section).

**regulatorEffect** For regulatory omics. List containing the percentage of effect types (repressor, activator or no effect) over the total number of regulators. See vignette for more information.

**totalFeatures** Number of features to simulate. By default, the total number of features in the seed dataset.

**depth** Sequencing depth in millions of reads. If not provided, it takes the global parameter passed to mosim function.

**replicateParams** List with parameters *a* and *b* for adjusting the variability in the generation of replicates using the negative binomial. See vignette for more information.

diffGenes Number of differentially expressed genes to simulate, given in percentage (0 -

1) or in absolute number (> 1). By default 0.15

numberReps Number of replicates per experimetal condition (and time point, if time series

are to be generated). By default 3.

numberGroups Number of experimental groups or conditions to simulate.

times Vector of time points to consider in the experimental design.

depth Sequencing depth in millions of reads.

profileProbs Numeric vector with the probabilities to assign each of the patterns. Defaults to

0.2 for each.

minMaxFC Numeric vector of length 2 with minimum and maximum fold-change for dif-

ferentially expressed features, respectively.

TFtoGene A logical value indicating if default transcription factors data should be used

(TRUE) or not (FALSE), or a 3 column data frame containing custom associa-

tions. By default FALSE.

# Value

Instance of class "MOSimulation" containing the multiomic simulation data.

```
moSimulation <- mosim(
   omics = c("RNA-seq"),
   numberReps = 3,
   times = c(0, 2, 6, 12, 24)
)
# Retrieve simulated count matrix for RNA-seq</pre>
```

dataRNAseq <- omicResults(moSimulation, "RNA-seq")</pre>

MOSimulation-class

This class manages the global simulation process, like associating genes with gene classes, regulatory programs and other settings. Finally it will initialize the simulators with their options that will use the previously generated settings to simulate the data.

# **Description**

This class manages the global simulation process, like associating genes with gene classes, regulatory programs and other settings. Finally it will initialize the simulators with their options that will use the previously generated settings to simulate the data.

#### Slots

simulators Vector containing either S4 initialized classes of simulators or a list with the class name as keys, and its options as value, see example.

totalGenes A number with the total number of genes including not expressed. Overwritten if a genome reference is provided. Currently not used as we force to provide real data.

diffGenes A number with the total number of differential genes (if value > 1) or % or total genes (if value < 1).

numberReps Number of replicates of the experiment.

numberGroups Number of samples considered on the experiment.

times Numeric vector containing the measured times. If numberGroups < 2, the number of times must be at least 2.

geneNames Read only. List containing the IDs of the genes. Overwrited by the genome reference if provided. Currently not used as we force to provide real data.

simSettings List of settings that overrides initializing the configuration of the simulation by passing a previously generated list. This could be used to tweak by hand the assigned profiles, genes, regulatory programs, etc.

noiseFunction Noise function to apply when simulating counts. Must accept the parameter 'n' and return a vector of the same length. Defaults to 'rnorm'

profiles Named list containing the patterns with their coefficients.

profileProbs Numeric vector with the probabilities to assign each of the patterns. Defaults to 0.2 for each.

noiseParams Default noise parameters to be used with noise function.

depth Default depth to simulate.

TFtoGene Boolean (for default data) or 3 column data frame containing Symbol-TFGene-LinkedGene minMaxQuantile Numeric vector of length 2 indicating the quantiles to use in order to retrieve the absolute minimum and maximum value that a differentially expressed feature can have.

minMaxFC Numeric vector of length 2 indicating the minimum and maximum fold-change that a differentially expressed feature can have.

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MOSimulator-class

Virtual class containing common methods and slots for child classes.

# **Description**

Virtual class containing common methods and slots for child classes.

#### **Slots**

name Name of the simulator to be used in messages.

data Data frame containing the initial sample to be used, with the features IDs as rownames and only one column named "Counts".

regulator Boolean flag to indicate if the omic is a regulator or not.

regulatorEffect Possible regulation effects of the omic (enhancer, repressor or both).

idToGene Data frame with the association table between genes and other features. The structure must be 2 columns, one named "ID" and the other "Gene".

min Minimum value allowed in the omic.

max Maximum value allowed in the omic.

depth Sequencing depth to simulate.

depthRound Number of decimal places to round when adjusting depth.

depthAdjust Boolean indicating whether to adjust by sequencing depth or not.

totalFeatures Number of features to simulate. This will replace the data with a subset.

noiseFunction Noise function to apply when simulating counts. Must accept the parameter 'n' and return a vector of the same length. Defaults to 'rnorm'

increment Read-only. Minimum value to increase when simulating counts.

simData Contains the final simulated data.

pregenerated Indicates if the child class will generate the simulated data instead of the general process.

randData Auxiliary vector containing the original count data in random order with other adjustments.

noiseParams Noise parameters to be used with noise function.

roundDigits Number of digits to round the simulated count values.

minMaxQuantile Numeric vector of length 2 indicating the quantiles to use in order to retrieve the absolute minimum and maximum value that a differentially expressed feature can have.

minMaxFC Numeric vector of length 2 indicating the minimum and maximum fold-change that a differentially expressed feature can have.

minMaxDist Named list containing different minimum and maximum constraints values calculated at the beginning of the simulation process.

replicateParams Named list containing the parameters a and b to be used in the replicates generation process, see the vignette for more info.

MOSimulatorRegion-class

Virtual class containing general methods for simulators based on regions of the chromosomes, like DNase-seq, ChIP-seq or Methyl-seq

#### **Description**

Virtual class containing general methods for simulators based on regions of the chromosomes, like DNase-seq, ChIP-seq or Methyl-seq

Class to simulate RNA-seq data

Class to simulate transcription factor data

Class to simulate miRNA-seq

Class to simulate ChIP-seq data

Class to simulate DNase-seq data

Class to simulate Methyl-seq data.

#### Slots

locs Vector containing the list of locations of the sites.

locsName Type of the site to simulate, only for debug.

splitChar Character symbol used to split identifiers in chr/start/end

nCpG numeric. Number of CpG sites to simulate.

pSuccessMethReg numeric. Probability of success in methylated region.

pSuccessDemethReg numeric. Probability of success in non methylated region

errorMethReg numeric. Error rate in methylated region

errorDemethReg numeric. Error rate in methylated region

nReadsMethReg numeric. Mean number of reads in methylated region.

nReadsDemethReg numeric. Mean number of reads in non methylated regions.

 $phase \ Diff\ numeric.\ Phase\ difference\ in\ the\ differentially\ methylated\ regions\ between\ two\ samples$ 

balanceHypoHyper numeric. Balance of hypo/hyper methylation

ratesHMMMatrix numeric. Matrix of values that describes the exponential decay functions that define the distances between CpG values.

distType character. Distribution used to generate replicates:

transitionSize numeric.

PhiMeth matrix. Transition matrix for CpG locations.

PhiDemeth matrix. <Not used>

typesLocation numeric. <Not used>

returnValue character. Selected column:

betaThreshold numeric. Beta threshold value used to calculate M values.

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omicData

Set customized data for an omic.

#### **Description**

Set customized data for an omic.

#### Usage

```
omicData(omic, data = NULL, associationList = NULL)
```

# **Arguments**

omic The name of the omic to provide data.

data Data frame with the omic identifiers as row names and just one column named

Counts containing numeric values used as initial sample for the simulation.

associationList

Only for regulatory omics, a data frame with 2 columns, the first called containing the regulator ID and the second called Gene with the gene identifier.

#### Value

Initialized simulation object with the given data.

```
# Take a subset of the included dataset for illustration
# purposes. We could also load it from a csv file or RData,
# as long as we transform it to have 1 column named "Counts"
# and the identifiers as row names.
data(sampleData)
custom_rnaseq <- head(sampleData$SimRNAseq$data, 100)</pre>
# In this case, 'custom_rnaseq' is a data frame with
# the structure:
head(custom_rnaseq)
                      Counts
## ENSMUSG00000000001
## ENSMUSG00000000003
## ENSMUSG00000000028
                        4644
## ENSMUSG00000000031
                           8
## ENSMUSG0000000037
                           0
## ENSMUSG00000000049
                           0
# The helper 'omicData' returns an object with our custom data.
rnaseq_customdata <- omicData("RNA-seq", data = custom_rnaseq)</pre>
```

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omicResults	Retrieves the simulated data.

# **Description**

Retrieves the simulated data.

# Usage

```
omicResults(simulation, omics = NULL, format = "data.frame")
```

# Arguments

simulation A MOSimulation object.

omics List of the omics to retrieve the simulated data.

format Type of object to use for returning the results

#### Value

A list containing an element for every omic specifiec, with the simulation data in the format indicated, or a numeric matrix with simulated data if the omic name is directly provided.

```
omic_list <- c("RNA-seq")</pre>
rnaseq_simulation <- mosim(omics = omic_list)</pre>
#' # This will be a data frame with RNA-seq counts
rnaseq_simulated <- omicResults(rnaseq_simulation, "RNA-seq")</pre>
                     Group1.Time0.Rep1 Group1.Time0.Rep2 Group1.Time0.Rep3 ...
# ENSMUSG00000073155
                                   4539
                                                      5374
                                                                         5808 ...
# ENSMUSG00000026251
                                     0
                                                       0
                                                                            0 ...
# ENSMUSG00000040472
                                   2742
                                                      2714
                                                                         2912 ...
# ENSMUSG00000021598
                                   5256
                                                      4640
                                                                         5130 ...
# ENSMUSG00000032348
                                                       348
                                                                         492 ...
                                    421
                                                                           9 ...
# ENSMUSG00000097226
                                     16
                                                        14
                                                                           0 ...
# ENSMUSG00000027857
                                      0
                                                         0
# ENSMUSG00000032081
                                      1
                                                         0
                                                                            0 ...
                                                                          965 ...
# ENSMUSG00000097164
                                    794
                                                       822
# ENSMUSG00000097871
                                                                            0 ...
```

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omicSettings Retrieves the settings used in a simulation
--

# **Description**

Retrieves the settings used in a simulation

# Usage

```
omicSettings(
  simulation,
  omics = NULL,
  association = FALSE,
  reverse = FALSE,
  only.linked = FALSE,
  prefix = FALSE,
  include.lagged = TRUE
)
```

# Arguments

simulation	A MOSimulation object.
omics	List of omics to retrieve the settings.
association	A boolean indicating if the association must also be returned for the regulators.
reverse	A boolean, swap the column order in the association list in case we want to use the output directly and the program requires a different ordering.
only.linked	Return only the interactions that have an effect.
prefix	Logical indicating if the name of the omic should prefix the name of the regulator.
include.lagged	Logical indicating if interactions with transitory profile and different minimum/maximum time point between gene and regulator should be included or not.

# Value

A list containing a data frame with the settings used to simulate each of the indicated omics. If association is TRUE, it will be a list with 3 keys: 'associations', 'settings' and 'regulators', with the first two keys being a list containing the information for the selected omics and the last one a global data frame giving the merged information.

```
omic_list <- c("RNA-seq", "miRNA-seq")
multi_simulation <- mosim(omics = omic_list)

# This will be a data frame with RNA-seq settings (DE flag, profiles)
rnaseq_settings <- omicSettings(multi_simulation, "RNA-seq")</pre>
```

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```
# This will be a list containing all the simulated omics (RNA-seq
# and DNase-seq in this case)
all_settings <- omicSettings(multi_simulation)</pre>
```

omicSim

Set the simulation settings for an omic.

#### **Description**

Set the simulation settings for an omic.

#### Usage

```
omicSim(omic, depth = NULL, totalFeatures = NULL, regulatorEffect = NULL)
```

# **Arguments**

omic Name of the omic to set the settings.

depth Sequencing depth in millions of counts. If not provided will take the global

parameter passed to mosim function.

totalFeatures Limit the number of features to simulate. By default include all present in the

dataset.

regulatorEffect

only for regulatory omics. Associative list containing the percentage of effects over the total number of regulator, including repressor, association and no effect

(NE).

#### Value

A list with the appropriate structure to be given as options in mosim function.

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plotProfile	Generate a plot of a feature's profile for one or two omics.

# Description

Generate a plot of a feature's profile for one or two omics.

# Usage

```
plotProfile(simulation, omics, featureIDS, drawReps = FALSE, groups = NULL)
```

# Arguments

simulation	A MOSimulation object
omics	Character vector of the omics to simulate.
featureIDS	List containing the feature to show per omic. Must have the omics as the list names and the features as values.
drawReps	Logical to enable/disable the representation of the replicates inside the plot.
groups	Character vector indicating the groups to plot in the form "GroupX" (i.e. Group1)

# Value

A ggplot2 object.

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random\_unif\_interval

random\_unif\_interval Function to call the C code This function is a copy of the 'random\_unif\_interval' function from the 'SPARSim' package (v0.9.5), originally developed by Giacomo Baruzzo, Ilaria Patuzzi, Barbara Di Camillo (2020). The original package is licensed under the GPL-3 license.

# **Description**

random\_unif\_interval Function to call the C code This function is a copy of the 'random\_unif\_interval' function from the 'SPARSim' package (v0.9.5), originally developed by Giacomo Baruzzo, Ilaria Patuzzi, Barbara Di Camillo (2020). The original package is licensed under the GPL-3 license.

#### Usage

```
random_unif_interval(size, max_val)
```

#### **Arguments**

size from sparsim max\_val from sparsim

sampleData

Default data

#### **Description**

Dataset with base counts and id-gene tables.

#### Usage

```
data("sampleData")
```

#### **Format**

An object of class list of length 6.

# Details

List with 6 elements:

SimRNAseq data Dataframe with base counts with gene id as rownames.

geneLength Length of every gene.

**SimChIPseq data** Dataframe with base counts with regions as rownames.

idToGene Dataframe with region as "ID" column and gene name on "Gene" column.

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SimDNaseseq data Dataframe with base counts with regions as rownames.

idToGene Dataframe with region as "ID" column and gene name on "Gene" column.

SimMiRNAseq data Dataframe with base counts with miRNA id as rownames.

idToGene Dataframe with miRNA as "ID" column and gene name on "Gene" column.

SimMethylseq idToGene Dataframe with region as "ID" column and gene name on "Gene" col-

CpGisland Dataframe of CpG to be used as initialization data, located on "Region" column

scatac

Data to test scMOSim

# Description

Data to test scMOSim

# Usage

```
data("scatac")
```

#### **Format**

A seurat Object, subset from seuratData with ATAC

assays ATAC expression values

meta.data annotations of celltypes

@source https://github.com/satijalab/seurat-data, we took 11 cells from each of 4 celltypes

scrna

Data to test scMOSim

# Description

Data to test scMOSim

#### Usage

```
data("scrna")
```

sc\_mosim

#### **Format**

A seurat Object, subset from seuratData with RNA

```
assays RNA expression values
```

meta.data annotations of celltypes

@source https://github.com/satijalab/seurat-data, we took 11 cells from each of 4 celltypes This is how: dat <- pbmcMultiome.SeuratData::pbmc.rna dat <- subset(x = dat, subset = seurat\_annotations "cDC", "Memory B", "Treg")) unique\_cell\_types <- unique(datATmeta.data\$seurat\_annotations) extracted\_cells <- list() cellnames <- c() for (cell\_type in unique\_cell\_types) type\_cells <- subset(dat, subset = seurat\_annotations counts <- as.matrix(type\_cellsATassays[["RNA"]]ATcounts) extracted\_cells[[cell\_type]] <- counts[, 1:10] cellnames <- append(cellnames, replicate(11, cell\_type))

```
scrna <- Reduce(cbind, extracted_cells)</pre>
```

sc\_mosim

sc mosim

# **Description**

Performs multiomic simulation of single cell datasets

# Usage

```
sc_mosim(
 omics,
  cellTypes,
 numberReps = 1,
 numberGroups = 1,
 diffGenes = NULL,
 minFC = 0.25,
 maxFC = 4,
 numberCells = NULL,
 mean = NULL,
  sd = NULL,
  noiseRep = 0.1,
  noiseGroup = 0.5,
  regulatorEffect = NULL,
  associationList = NULL,
  feature_no = 8000,
  clusters = 3,
  cluster_size = NULL,
 TF = FALSE,
  TFdf = NULL
)
```

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#### **Arguments**

omics named list containing the omic to simulate as names, which can be "scRNA-seq"

or "scATAC-seq".

cellTypes list where the i-th element of the list contains the column indices for i-th exper-

imental conditions. List must be a named list.

numberReps OPTIONAL. Number of replicates per group numberGroups OPTIONAL. number of different groups

diffGenes OPTIONAL. If number groups > 1, Percentage DE genes to simulate. List of

vectors (one per group to compare to group 1) where the vector contains absolute number of genes for Up and Down ex: c(250, 500) or a percentage for up, down

ex: c(0.2, 0.2). The rest will be NE

minFC OPTIONAL. Threshold of FC below which are downregulated, by default 0.25

maxFC OPTIONAL. Threshold of FC above which are upregulated, by default 4

numberCells OPTIONAL. Vector of numbers. The numbers correspond to the number of cells

the user wants to simulate per each cell type. The length of the vector must be

the same as length of cellTypes.

mean OPTIONAL. Vector of numbers of mean depth per each cell type. Must be

specified just if numberCells is specified. The length of the vector must be the

same as length of cellTypes.

sd OPTIONAL. Vector of numbers of standard deviation per each cell type. Must

be specified just if numberCells is specified. The length of the vector must be

the same as length of cellTypes.

noiseRep OPTIONAL. Number indicating the desired standard deviation between biolog-

ical replicates.

noiseGroup OPTIONAL. Number indicating the desired standard deviation between treat-

ment groups

regulatorEffect

OPTIONAL. To simulate relationship scRNA-scATAC, list of vectors (one per group) where the vector contains absolute number of regulators for Activator and repressor ex: c(150, 200) or a percentage for Activator and repressor ex: c(0.2, 0.1). The rest will be NE. If not provided, no table of association between

scRNA and scATAC is outputted.

association List

REQUIRED A 2 columns dataframe reporting peak ids related to gene names.

If user doesnt have one, load from package data("associationList")

feature\_no OPTIONAL. If only scRNA-seq to simulate or scRNA and scATAC but no reg-

ulatory constraints, total number of features to be distributed between the coex-

pression clusters.

clusters OPTIONAL. Number of co-expression patterns the user wants to simulate

cluster\_size OPTIONAL. It may be inputted by the user. Recommended: by default, its the

number of features divided by the number of patterns to generate.

TF OPTIONAL default is FALSE, if true, extract TF dataframe

TFdf OPTIONAL, default is NULL. If an association matrix of TF and Target\_gene

is given the TF expression values are extracted. If no data.frame is given, using

the association of human TF from https://tflink.net/

24 sc\_omicData

# Value

a list of Seurat object, one per each omic.

# **Examples**

```
omic_list <- sc_omicData(list("scRNA-seq"))
cell_types <- list('Treg' = c(1:10),'cDC' = c(11:20),'CD4_TEM' = c(21:30),
'Memory_B' = c(31:40))
sim <- sc_mosim(omic_list, cell_types)</pre>
```

sc\_omicData

sc\_omicData

# **Description**

Checks if the user defined data is in the correct format, or loads the default multiomics pbmc dataset, a subset from SeuratData package

# Usage

```
sc_omicData(omics_types, data = NULL)
```

# **Arguments**

omics\_types A list of strings which can be either "scRNA-seq" or "scATAC-seq"

data A user input matrix with genes (peaks in case of scATAC-seq) as rows and cells

as columns. By default, it loads the example data. If a user input matrix is

included, cell columns must be sorted by cell t ype.

#### Value

a named list with omics type as name and the count matrix as value

```
# Simulate from PBMC
omicsList <- sc_omicData(list("scRNA-seq", "scATAC-seq"))</pre>
```

sc\_omicResults 25

sc\_omicResults

sc\_omicResults

# Description

```
sc\_omicResults
```

# Usage

```
sc_omicResults(sim)
```

# **Arguments**

sim

a simulated object from sc\_mosim function

# Value

list of seurat objects with simulated data

# **Examples**

```
cell_types <- list('Treg' = c(1:10),'cDC' = c(11:20),'CD4_TEM' = c(21:30),
'Memory_B' = c(31:40))
omicsList <- sc_omicData(list("scRNA-seq"))
sim <- sc_mosim(omicsList, cell_types)
res <- sc_omicResults(sim)</pre>
```

sc\_omicSettings

sc\_omicSettings

# Description

```
sc_omicSettings
```

#### Usage

```
sc_omicSettings(sim, TF = FALSE)
```

#### **Arguments**

sim a simulated object from sc\_mosim function

TF OPTIONAL default is FALSE, if true, extract TF association matrix

# Value

list of Association matrices explaining the effects of each regulator to each gene

26 sc\_param\_estimation

# **Examples**

```
 cell\_types <- list('Treg' = c(1:10), 'cDC' = c(11:20), 'CD4\_TEM' = c(21:30), 'Memory\_B' = c(31:40)) \\ omicsList <- sc\_omicData(list("scRNA-seq")) \\ sim <- sc\_mosim(omicsList, cell\_types) \\ res <- sc\_omicSettings(sim)
```

sc\_param\_estimation sc\_param\_estimation

# **Description**

Evaluate the users parameters for single cell simulation and use SPARSim to simulate the main dataset. Internal function

# Usage

```
sc_param_estimation(
  omics,
  cellTypes,
  diffGenes = list(c(0.2, 0.2)),
  minFC = 0.25,
  maxFC = 4,
  numberCells = NULL,
  mean = NULL,
  sd = NULL,
  noiseGroup = 0.5,
  group = 1,
  genereggroup
)
```

# Arguments

omics	named list containing the omics to simulate as names, which can be "scRNA-seq" or "scATAC-seq".
cellTypes	list where the i-th element of the list contains the column indices for i-th cell type. List must be a named list.
diffGenes	If number groups $>$ 1, Percentage DE genes to simulate. List of vectors (one per group to compare to group 1) where the vector contains absolute number of genes for Up and Down ex: c(250, 500) or a percentage for up, down ex: c(0.2, 0.2). The rest will be NE
minFC	Threshold of FC below which are downregulated, by default 0.25
maxFC	Threshold of FC above which are upregulated, by default 4
numberCells	vector of numbers. The numbers correspond to the number of cells the user wants to simulate per each cell type. The length of the vector must be the same

as length of cellTypes.

shuffle\_group\_matrix 27

mean	vector of numbers of	t mean depth per each cell type.	Must be specified just if
		'C 1	

numberCells is specified.

sd vector of numbers of standard deviation per each cell type. Must be specified

just if numberCells is specified.

noiseGroup OPTIONAL. Number indicating the desired standard deviation between treat-

ment groups

group Group for which to estimate parameters

genereggroup List with information of genes, clusters and regulators that must be related to

each other

#### Value

a list of Seurat object, one per each omic.

a named list with simulation parameters for each omics as values.

# **Examples**

```
omicsList <- sc_omicData(list("scRNA-seq"))
cell_types <- list('Treg' = c(1:10),'cDC' = c(11:20),'CD4_TEM' = c(21:30),
'Memory_B' = c(31:40))
#estimated_params <- sc_param_estimation(omicsList, cell_types)</pre>
```

shuffle\_group\_matrix

shuffle\_group\_matrix, Reorder cell type-specific expression matrix during co-expression simulation. Copied from ACORDE (https://github.com/ConesaLab/acorde) to facilitate stability and running within our scripts This function is a slightly modified copy of the 'shuffle\_group\_matrix' function from the 'Acorde' package (v1.0.0), originally developed by Arzalluz-Luque A, Salguero P, Tarazona S, Conesa A. (2022). acorde unravels functionally interpretable networks of isoform co-usage from single cell data. Nature communications 1828. DOI: 10.1038/s41467-022-29497-w. The original package is licensed under the GPL-3 license.

#### **Description**

This function is used internally by accorde to perform the shuffling of simulated features for an individual cell type, as part of the co-expression simulation process. The function is called recursively by simulate\_coexpression() to perform the simulation on a full scRNA-seq matrix.

#### Usage

```
shuffle_group_matrix(sim_data, feature_ids, group_pattern, ngroups)
```

#### **Arguments**

sim\_data A count matrix with features as rows and cells as columns. Feature IDs must be

included in an additional column named feature.

feature\_ids A two-column tibble containing top and bottom columns, each including the

feature IDs of features to be used as highly or lowly expressed when shuffling

by the indicated expression pattern.

group\_pattern A logical vector, containing TRUE to indicate that high expression in that cell

type is desired and FALSE if the opposite. The vector must be ordered as the cell

types in sim\_data.

ngroups An integer indicating the number of groups that top and bottom features should

be divided into. It is computed by dividing the number of features selected as

highly/lowly expressed by the size of the clusters that are to be generated.

#### Value

An expression matrix, with the same characteristics as sim\_data, and a number of features defined as the total amount of top/bottom features selected divided by the number of clusters for which co-expression patterns where supplied.

simulate\_coexpression simulate coexpression

#### **Description**

Adapted from ACORDE (https://github.com/ConesaLab/acorde) to adapt to our data input type. Simulates coexpression of genes along celltypes

#### Usage

```
simulate_coexpression(
   sim_matrix,
   feature_no,
   cellTypes,
   patterns,
   cluster_size = NULL
)
```

# **Arguments**

sim\_matrix Matrix with rows as features and columns as cells

feature\_no Total number of features to be distributed between the coexpression clusters cellTypes list where the i-th element of the list contains the column indices for i-th exper-

imental conditions. List must be a named list.

patterns Tibble with TRUE FALSE depicting the cluster patterns to simulate. Generated

by the user or by make\_cluster\_patterns.

cluster\_size OPTIONAL. It may be inputted by the user. By default, its the number of fea-

tures divided by the number of patterns to generate.

simulate\_hyper 29

#### **Details**

This function is a slightly modified copy of the 'simulate\_coexpression' function from the 'Acorde' package (v1.0.0), originally developed by Arzalluz-Luque A, Salguero P, Tarazona S, Conesa A. (2022). acorde unravels functionally interpretable networks of isoform co-usage from single cell data. Nature communications 1828. DOI: 10.1038/s41467-022-29497-w. The original package is licensed under the GPL-3 license.

#### Value

the simulated coexpression

late_hyper Simulate technical variabilit
--

# **Description**

Function to simulate the technical variability (i.e. a multivariate hypergeometric on a gamma expression value array)

#### Usage

```
simulate_hyper(avgAbund, seqdepth = NULL, digits, max_val)
```

#### **Arguments**

avgAbund	array containing the intensity values for each feature. It describes the intensity of a single sample
seqdepth	sequencing depth (i.e. sample size of the MH)
digits	number of digits for random number generation
max_val	max value for random number generation

# Details

This function is a copy of the 'simulate\_hyper' function from the 'SPARSim' package (v0.9.5), originally developed by Giacomo Baruzzo, Ilaria Patuzzi, Barbara Di Camillo (2020). The original package is licensed under the GPL-3 license.

#### Value

An array of length(avgAbund) elements representing the count values for the current sample

#### **Description**

Function to create a SPARSim simulation parameter. This function is a copy of the 'SPARSIM\_create\_simulation\_parameter' function from the 'SPARSim' package (v0.9.5), originally developed by Giacomo Baruzzo, Ilaria Patuzzi, Barbara Di Camillo (2020). The original package is licensed under the GPL-3 license. To simulate N feature (e.g. genes), user must specify N values of gene expression level and gene expression variability in the function input parameters intensity and variability, respectively. To simulate M samples (i.e. cells), user must specify M values of sample library size in the function input parameter library\_size.

# Usage

```
sparsim_create_simulation_parameter(
  intensity,
  variability,
  library_size,
  feature_names = NA,
  sample_names = NA,
  condition_name = NA,
  intensity_2 = NULL,
  variability_2 = NULL,
  p_bimod = NULL
)
```

# Arguments

intensity	Array of gene expression intensity values
variability	Array of gene expression variability values
library_size	Array of library size values
feature_names	Array of feature names. It must be of the same length of intensity array. If NA (default), feature will be automatically named "gene_1", "gene_2", "gene_ $<$ N>", where N = length(intensity)
sample_names	Array of sample names. It must be of the same length of library_size array. If NA (defatul), sample will be automatically named " <condition_name>_cell1", "<condition_name>_cell2",, "<condition_name>_cell<m>", where <math>M = length(library\_size)</math></m></condition_name></condition_name></condition_name>
condition_name	Name associated to the current experimental condition. If NA (default), it will be set to "cond<11><12>", where 11 and 12 are two random letters.
intensity_2	Array of gene expression intensity values for the second expression mode, if simulating genes with bimodal gene expression. Entries containing NAs will be ignored. If NULL (default), no bimodal gene expression is simulated.

variability\_2 Array of gene expression variability values for the second expression mode, if

simulating genes with bimodal gene expression. If NULL (default), no bimodal

gene expression is simulated.

p\_bimod Array of bimodal gene expression probabilities; the i-th value indicates the prob-

ability p of the i-th gene to be expressed in the first mode (i.e. the one specified in the i-th entries of parameters intensity and variability); with probability 1-p the i-th gene will be expressed in the second mode (i.e. the one specified in

the i-th entries of parameters intensity\_2 and variability\_2)

#### **Details**

User can optionally specify the names to assign at the single feature and sample to simulate (function input parameters feature\_names and sample\_names, respectively, as well as the name of the experimental condition (function input parameter condition\_name). If the user does not specify such information, the function will set some default values.

To simulate T different experimental conditions in a single count table, then T different simulation parameters must be created.

#### Value

SPARSim simulation parameter describing one experimental condition

sparsim\_estimate\_intensity

Estimate SPARSIm "intensity" parameter

#### **Description**

Function to estimate the intensity values from the genes in data. The intensity is computed as mean of normalized counts for each gene.

#### Usage

sparsim\_estimate\_intensity(data)

#### **Arguments**

data

normalized count data matrix (gene on rows, samples on columns). rownames(data) must contain gene names.

#### **Details**

This function is a copy of the 'SPARSIM\_estimate\_intensity' function from the 'SPARSim' package (v0.9.5), originally developed by Giacomo Baruzzo, Ilaria Patuzzi, Barbara Di Camillo (2020).

The original package is licensed under the GPL-3 license. This function is used in sparsim\_estimate\_parameter\_from\_date to compute SPARSim "intensity" parameter, given a real count table as input. If the count table contains more than one experimental condition, then the function is applied to each experimental conditions.

#### Value

An array of intensity values having N\_genes elements (N\_genes = nrow(data)). Array entries are named with gene names.

```
sparsim_estimate_library_size
```

Estimate SPARSim "library size" parameter

# **Description**

Function to estimate the library sizes from the samples in data.

#### Usage

```
sparsim_estimate_library_size(data)
```

# **Arguments**

data

raw count data matrix (gene on rows, samples on columns)

#### **Details**

This function is a copy of the 'SPARSIM\_estimate\_library\_size' function from the 'SPARSim' package (v0.9.5), originally developed by Giacomo Baruzzo, Ilaria Patuzzi, Barbara Di Camillo (2020). The original package is licensed under the GPL-3 license. This function is used in sparsim\_estimate\_parameter\_f to compute SPARSim "library size" parameter, given a real count table as input. If the count table contains more than one experimental condition, then the function is applied to each experimental conditions.

#### Value

An array of library size values having N\_samples elements (N\_samples = ncol(data))

```
sparsim_estimate_parameter_from_data
```

Estimate SPARSim simulation parameter from a given count table

# Description

Function to estimate SPARSim simulation parameters (intensity, variability and library sizes) from a real count table. If the real count table contains more than one experimental condition, it is possible to estimate the parameters for each experimental condition.

#### Usage

```
sparsim_estimate_parameter_from_data(raw_data, norm_data, conditions)
```

#### **Arguments**

norm\_data

raw\_data count matrix (gene on rows, samples on columns) containing raw count data

count matrix (gene on rows, samples on columns) containing normalized count

data

conditions list where the i-th element of the list contains the column indices for i-th exper-

imental conditions. List must be a named list.

#### **Details**

This function is a copy of the 'SPARSIM\_estimate\_parameter\_from\_data' function from the 'SPAR-Sim' package (v0.9.5), originally developed by Giacomo Baruzzo, Ilaria Patuzzi, Barbara Di Camillo (2020). The original package is licensed under the GPL-3 license.

#### Value

A SPARSim simulation parameters

sparsim\_estimate\_variability

Estimate SPARSim "variability" parameter

#### **Description**

Function to estimate the variability values from the genes in data.

# Usage

```
sparsim_estimate_variability(data)
```

# **Arguments**

data raw count data matrix (gene on rows, samples on columns)

#### **Details**

This function is a copy of the 'SPARSIM\_estimate\_variability' function from the 'SPARSim' package (v0.9.5), originally developed by Giacomo Baruzzo, Ilaria Patuzzi, Barbara Di Camillo (2020). The original package is licensed under the GPL-3 license. This function is used in sparsim\_estimate\_parameter\_from\_date to compute SPARSim "variability" parameter, given a real count table as input. If the count table contains more than one experimental condition, then the function is applied to each experimental conditions.

#### Value

An array of variability values having N\_genes elements (N\_genes = nrow(data))

34 sparsim\_simulation

sparsim\_simulation

Function to simulate a raw count table

#### **Description**

This function is a copy of the 'SPARSIM\_simulation' function from the 'SPARSim' package (v0.9.5), originally developed by Giacomo Baruzzo, Ilaria Patuzzi, Barbara Di Camillo (2020). The original package is licensed under the GPL-3 license.

# Usage

```
sparsim_simulation(
  dataset_parameter,
  output_sim_param_matrices = FALSE,
  output_batch_matrix = FALSE,
  count_data_simulation_seed = NULL
)
```

#### **Arguments**

dataset\_parameter

list containing, the intensity, variability and lib sizes of each experimental condition. It is the return value of "estimate\_parameter\_from\_data" or could be created by the users

output\_sim\_param\_matrices

boolean flag. If TRUE, the function will output two additional matrices, called abundance\_matrix and variability\_matrix, containing the gene intensities and gene variabilities used as simulation input. (Default: FALSE)

output\_batch\_matrix

boolean flag. If TRUE, the function will output an additional matrix, called batch\_factors\_matrix, containing the multiplicative factors used in batch effect simulation. (Default: FALSE)

count\_data\_simulation\_seed

inherited from sparsim

#### Value

A list of 5 elements:

- count\_matrix: the simulated count matrix (genes on rows, samples on columns)
- gene\_matrix: the simulated gene expression levels (genes on rows, samples on columns)
- abundance\_matrix: the input gene intensity values provided as input (genes on rows, samples on columns), if output\_sim\_param\_matrices = TRUE. NULL otherwise.
- variability\_matrix: the input gene variability values provided as input (genes on rows, samples on columns), if output\_sim\_param\_matrices = TRUE. NULL otherwise.
- batch\_factors\_matrix: the multiplicative factor used in batch generation (genes on rows, samples on columns), if output\_batch\_matrix = TRUE. NULL otherwise.

TF\_human 35

TF\_human

Data to extract human TF

# Description

Data to extract human TF

# Usage

```
data("TF_human")
```

# **Format**

vector of gene names

data.frame gene names corresponding to TF and to Target genes

@source https://tflink.net/

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