Package 'MOSim'

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Title Multi-Omics Simulation (MOSim)

Version 1.8.0

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

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License GPL-3

LazyData false

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MOSim-package MOSim

Description

Multiomics simulation package.

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</pre>
```

design_matrix <- experimentalDesign(rnaseq_simulation)</pre>

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.

	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.
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 associations. By default FALSE.

Value

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

Examples

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

omicData

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.
associationLis	t
	Only for regulatory omics, a data frame with 2 columns, the first called contain- ing the regulator ID and the second called Gene with the gene identifier.

Value

Initialized simulation object with the given data.

Examples

```
# 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
## ENSMUSG0000000001
                        6572
## ENSMUSG0000000003
                           0
## ENSMUSG000000028
                        4644
## ENSMUSG000000031
                           8
## ENSMUSG000000037
                           0
## ENSMUSG000000049
                           0
```

The helper 'omicData' returns an object with our custom data. rnaseq_customdata <- omicData("RNA-seq", data = custom_rnaseq)</pre> omicResults

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.

Examples

```
omic_list <- c("RNA-seq")
rnaseq_simulation <- mosim(omics = omic_list)
#' # 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	
# ENSMUSG0000073155	4539	5374	5808	
# ENSMUSG00000026251	0	0	0	
# ENSMUSG00000040472	2742	2714	2912	
# ENSMUSG0000021598	5256	4640	5130	
# ENSMUSG0000032348	421	348	492	
# ENSMUSG00000097226	16	14	9	
# ENSMUSG00000027857	0	0	0	
# ENSMUSG0000032081	1	0	0	
# ENSMUSG00000097164	794	822	965	
# ENSMUSG00000097871	0	0	0	

omicSettings

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 regula- tor.
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.

Examples

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

all_settings <- omicSettings(multi_simulation)</pre>

omicSim

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 appropiate structure to be given as options in mosim function.

Examples

plotProfile

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.

Examples

```
omic_list <- c("RNA-seq", "miRNA-seq")
rnaseq_simulation <- mosim(omics = omic_list)
plotProfile(rnaseq_simulation,
    omics = c("RNA-seq", "miRNA-seq"),
    featureIDS = list("RNA-seq"="ENSMUSG0000007682", "miRNA-seq"="mmu-miR-320-3p")
)</pre>
```

sampleData Default data

Description

Dataset with base counts and id-gene tables.

Usage

sampleData

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.

 $idToGene \ \ Data frame with \ region \ as \ "ID" \ column \ and \ gene \ name \ on \ "Gene" \ column.$

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" column.

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

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