# Package 'scMAGeCK'

October 14, 2021

| Type Package  |
|---|
| <b>Title</b> Identify genes associated with multiple expression phenotypes in single-cell CRISPR screening data   |
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| <b>Description</b><br>scMAGeCK is a computational model to identify genes associated with multiple expression phe-<br>notypes from CRISPR screening coupled with single-cell RNA sequencing data (CROP-seq) |
| License BSD_2_clause  |
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# **R** topics documented:

| scMAGeCK-package scmageck_lr |   |   |       |   |   |   |   |   |   |   |   |   |   |       |   |   |   |   |   |       |   |   |   |  |   |   |   |
|------------------------------|---|---|-------|---|---|---|---|---|---|---|---|---|---|-------|---|---|---|---|---|-------|---|---|---|--|---|---|---|
| scmageck_rra                 | • | • | <br>• | • | • | • | • | • | • | • | • | • | • | <br>• | • | • | • | • | • | <br>• | • | • | • |  | • | • | 4 |
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scMAGeCK-package

Identify genes associated with multiple expression phenotypes in single-cell CRISPR screening data

#### Description

scMAGeCK is a computational model to identify genes associated with multiple expression phenotypes from CRISPR screening coupled with single-cell RNA sequencing data (CROP-seq)

#### Details

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scMAGeCK is a computational model to identify genes associated with multiple expression phenotypes from CRISPR screening coupled with single-cell RNA sequencing data (CROP-seq).scMAGeCK is based on our previous MAGeCK and MAGeCK-VISPR models for pooled CRISPR screens.

The scMAGeCK manuscript can be found at bioRxiv(https://www.biorxiv.org/content/10.1101/658146v1/).

#### Author(s)

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#### Examples

```
### BARCODE file contains cell identity information, generated from
### the cell identity collection step
BARCODE <- system.file("extdata","barcode_rec.txt",package = "scMAGeCK")</pre>
### RDS can be a Seurat object or local RDS file path that contains
### the scRNA-seq dataset
RDS <- system.file("extdata", "singles_dox_mki67_v3.RDS", package = "scMAGeCK")
### Set RRA executable file path.
     You can generate RRA executable file by following commands:
###
###
      wget https://bitbucket.org/weililab/scmageck/downloads/RRA_0.5.9.zip
###
      unzip RRA_0.5.9.zip
     cd RRA_0.5.9
###
     make
###
RRAPATH <- "/Library/RRA_0.5.9/bin/RRA"</pre>
target_gene <- "MKI67"</pre>
rra_result <- scmageck_rra(BARCODE=BARCODE, RDS=RDS, GENE=target_gene,</pre>
                            RRAPATH=RRAPATH, LABEL='dox_mki67',
                            NEGCTRL=NULL, KEEPTMP=FALSE,
                            PATHWAY=FALSE, SAVEPATH=NULL)
head(rra_result)
```

# scmageck\_lr

scmageck\_lr

Use linear regression to test the association of gene knockout with all possible genes

# Description

echo "Use linear regression to test the association of gene knockout with all possible genes"

# Usage

scmageck\_lr(BARCODE, RDS, NEGCTRL, SELECT\_GENE=NULL, LABEL = NULL, PERMUTATION = NULL, SAVEPATH = "./",LAMBDA=0.01,GENE\_FRAC=0.01)

#### Arguments

| BARCODE     | A txt file to include cell identity information, generated from the cell identity collection step.  |
|-------------|---|
| RDS         | A Seurat object or local RDS file path that contains the scRNA-seq dataset. Note that the dataset has to be normalized and scaled.                      |
| NEGCTRL     | The name of the genes (separated by ",") served as negative controls.   |
| SELECT_GENE | The list of genes for regression. By default, all genes in the table are subject to regression.   |
| LABEL       | The label of the output file.   |
| PERMUTATION | The number of permutations for p value calculation.   |
| SAVEPATH    | The save path of result. Default save path is the current working directory. If you don't need save the result, set SAVEPATH as NULL.                   |
| LAMBDA      | A paramter for the LR model for ridge regression. Default: 0.01.  |
| GENE_FRAC   | A paramter for filtering low expressed genes. By default, only genes that have expressions in at least that fractions of cells are kept. Default: 0.01. |

#### Value

The result for object RDS

#### Examples

```
### BARCODE file contains cell identity information, generated from the cell identity collection step
BARCODE <- system.file("extdata","barcode_rec.txt",package = "scMAGeCK")
### RDS can be a Seurat object or local RDS file path that contains the scRNA-seq dataset
RDS <- system.file("extdata","singles_dox_mki67_v3.RDS",package = "scMAGeCK")
lr_result <- scmageck_lr(BARCODE=BARCODE, RDS=RDS, LABEL='dox_scmageck_lr',
NEGCTRL = 'NonTargetingControlGuideForHuman', PERMUTATION = 1000, SAVEPATH=NULL, LAMBDA=0.01)
lr_score <- lr_result[1]
lr_score_pval <- lr_result[2]
head(lr_score_pval)
```

| scmageck_rra | Use RRA to test the association of gene knockout with certain marker |
|--------------|--|
|              | expression   |

#### Description

echo "Use RRA to test the association of gene knockout with certain marker expression"

# Usage

```
scmageck_rra(BARCODE, RDS, GENE, RRAPATH = NULL, LABEL = NULL, NEGCTRL = NULL,
KEEPTMP = FALSE, PATHWAY = FALSE, SAVEPATH = "./")
```

#### Arguments

| BARCODE  | A txt file to include cell identity information, generated from the cell identity collection step.   |
|----------|--|
| RDS      | A Seurat object or local RDS file path that contains the scRNA-seq dataset. Note that the dataset has to be normalized and scaled.           |
| GENE     | Genes whose expressions are to be tested. Multiple genes can be provided, separated by ",". For example, "MKI67,TP53"                        |
| RRAPATH  | The path to the RRA program, if RRA cannot be found in the PATH environment variable.  |
| LABEL    | The label of the output file.  |
| NEGCTRL  | The name of the negative control gene. For example, "NonTargetingControl-GuideForHuman". Default is NULL (do not use any negative controls). |
| KEEPTMP  | Keep temporary files.  |
| PATHWAY  | Treat genes in –GENE option as a pathway. In other words, the averaged expression of these genes will be used for testing.                   |
| SAVEPATH | The save path of result. Default save path is the current working directory. If you don't need save the result, set SAVEPATH as NULL.        |

# scmageck\_rra

### Value

The result for object RDS

# Examples

### BARCODE file contains cell identity information, generated from the cell identity collection step BARCODE <- system.file("extdata","barcode\_rec.txt",package = "scMAGeCK")</pre>

### RDS can be a Seurat object or local RDS file path that contains the scRNA-seq dataset RDS <- system.file("extdata","singles\_dox\_mki67\_v3.RDS",package = "scMAGeCK")</pre>

```
target_gene <- "MKI67"</pre>
```

```
rra_result <- scmageck_rra(BARCODE=BARCODE, RDS=RDS, GENE=target_gene,
        LABEL='dox_mki67', NEGCTRL=NULL, KEEPTMP=FALSE, PATHWAY=FALSE, SAVEPATH=NULL)
head(rra_result)
```

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