Package 'miRTalk'

June 9, 2023

```
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```

VignetteBuilder knitr

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create_miRTalk

Create miRTalk object

Description

create miRTalk object using single-cell transcriptomics data

```
create_miRTalk(sc_data, sc_celltype, species, if_normalize = TRUE)
```

demo_gene2go 3

Arguments

sc_data A data.frame or matrix or dgCMatrix containing raw counts of single-cell RNA-

seq data. see demo_sc_data

sc_celltype A character containing the cell type of the single-cell RNA-seq data

species A character meaning species of the single-cell transcriptomics data. 'Human',

'Mouse' or 'Rat'

been normalized.

Value

miRTalk object

demo_gene2go

Demo data of gene2go

Description

Demo data of gene2go

Usage

```
demo_gene2go()
```

Details

gene2go must be a data. frame object with three columns, namely 'symbol', 'GO_term', 'species'.

Examples

```
gene2go_demo <- demo_gene2go()</pre>
```

demo_geneinfo

Demo data of geneinfo

Description

Demo data of geneinfo

Usage

```
demo_geneinfo()
```

Details

geneinfo must be a data.frame object with three columns, namely 'symbol', 'synonyms', 'species'.

Examples

```
geneinfo_demo <- demo_geneinfo()</pre>
```

4 demo_mir_info

demo_mir2tar

Demo data of mir2tar

Description

Demo data of mir2tar

Usage

```
demo_mir2tar()
```

Details

```
mir2tar must be a data.frame object with four columns, namely 'miRNA', 'miRNA_mature',
'target_gene', 'species'
```

Examples

```
mir2tar_demo <- demo_mir2tar()</pre>
```

demo_mir_info

Demo data of mir_info

Description

Demo data of mir_info

Usage

```
demo_mir_info()
```

Details

```
mir\_info must be a data.frame object with four columns, namely 'miRNA', 'miRNA_mature', 'gene', 'species'
```

Examples

```
mir_info_demo <- demo_mir_info()</pre>
```

demo_pathways 5

demo_pathways

Demo data of pathways

Description

Demo data of pathways

Usage

```
demo_pathways()
```

Details

pathways must be a data.frame object with four columns, namely 'src', 'dest', 'pathway', 'species'

Examples

```
pathways_demo <- demo_pathways()</pre>
```

demo_sc_data

Demo data of sc_data

Description

Demo data of sc_data.

Usage

```
demo_sc_data()
```

Details

sc_data can be a data.frame, matrix, or dgCMatrix object, each column representing a cell, each row representing a gene.

Value

A dgCMatrix object.

Examples

```
sc_data_demo <- demo_sc_data()</pre>
```

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find_hvtg Find highly variable target genes

Description

Find highly variable target genes by excluding the cell-type-specific potential marker genes

Usage

```
find_hvtg(
  object,
  pvalue = 0.05,
  log2fc = 0.5,
  min_cell_num = 10,
  nfeatures = 3000
)
```

Arguments

object miRTalk object after find_miRNA pvalue Cutoff of p value. Default is 0.05

log2 fold change for identifying the highly expressed genes in each cell type.

Default is 0.5

nfeatures Number of features to select as top variable features. Default is 3000

Value

miRTalk object containing highly variable target genes without the cell-type-specific potential marker genes

find_miRNA

Find expressed miRNAs

Description

Find expressed miRNAs among all cells

Usage

```
find_miRNA(object, mir_info)
```

Arguments

object miRTalk object after create_miRTalk

mir_info A data.frame of the system data containing information of EV-derived miRNA

of 'Human', 'Mouse' or 'Rat'. see demo_mir_info

Value

miRTalk object containing the expressed miRNAs

find_miRTalk 7

find_miRTalk	Infer cell-cell communications mediated by EV-derived miRNAs

Description

Infer cell-cell communications mediated by exosomal miRNAs from senders to receivers

Usage

```
find_miRTalk(
  object,
  mir2tar,
  min_cell_num = 10,
  pvalue = 0.05,
  resolution = "mature",
  min_percent = 0.05,
  if_doParallel = TRUE,
  use_n_cores = 4
)
```

Arguments

object	miRTalk object after create_miRTalk
mir2tar	A data.frame of the system data containing relationship of miRNA and its target genes for 'Human', 'Mouse' or 'Rat'. see $demo_mir2tar$
min_cell_num	Min cell number for each cell type and expressed miRNA. Default is 10
pvalue	Cutoff of p value. Default is 0.05
resolution	Correct to precursor or mature miRNAs. Use 'precursor' or 'mature'. Default is 'mature'
min_percent	Min percent of expressed cells for target genes of miRNA. Default is 0.05
if_doParallel	Use doParallel. Default is TRUE.
use_n_cores	Number of CPU cores to use. Default is 4

Value

miRTalk object containing the inferred cell-cell communications mediated by EV-derived miRNAs

Description

A data.frame containing GO terms, i.e., process, function, and component, of "Human", "Mouse", and "Rat" updated on Jan, 02, 2022 for revising count matrix.

Usage

gene2go

get_gene2go

Format

An object of class data. frame with 1190681 rows and 7 columns.

Source

```
https://www.ncbi.nlm.nih.gov/gene
```

geneinfo

geneinfo

Description

A data.frame containing gene symbols of "Human", "Mouse", and "Rat" updated on Jan, 02, 2022 for revising count matrix.

Usage

geneinfo

Format

An object of class data. frame with 288559 rows and 3 columns.

Source

```
https://www.ncbi.nlm.nih.gov/gene
```

get_gene2go

Get GO terms

Description

Get GO terms for target genes

Usage

```
get_gene2go(target_genes, gene2go, species)
```

Arguments

target_genes Character of one or more target genes

gene2go A data.frame of the system data containing GO terms for 'Human', 'Mouse' or

'Rat'. see demo_gene2go

species A character meaning species of the target genes. 'Human', 'Mouse' or 'Rat'

Value

GO terms for one or more target genes

get_miRTalk_cci 9

Description

Get all miRNAs' activity for a given sender cell type

Usage

```
get_miRTalk_cci(object, simple = TRUE)
```

Arguments

object miRTalk object after find_miRTalk

simple Whether to show the simple results. Default is TRUE

Value

A data.frame containing all miRNA-target interactions.

get_pathways	Get pathways		
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Description

Get pathways for target genes

Usage

```
get_pathways(target_genes, pathways, species)
```

Arguments

target_genes Character of one or more target genes

pathways A data frame of the system data containing gene-gene interactions and pathways

from KEGG and Reactome for 'Human', 'Mouse' or 'Rat'. see demo_pathways

species A character meaning species of the target genes.'Human', 'Mouse' or 'Rat'

Value

Pathways for one or more target genes

10 miRTalk

mir2tar

mir2tar

Description

A data frame containing relationship of miRNA and its target genes for "Human", "Mouse", and "Rat".

Usage

mir2tar

Format

An object of class data. frame with 544329 rows and 8 columns.

Source

https://mirtarbase.cuhk.edu.cn/

miRTalk

Definition of 'miRTalk' class

Description

An S4 class containing the data, meta, and results of inferred cell-cell communications mediated by EV-derived miRNAs.

Slots

data A list containing the data and variable genes.

meta A data.frame containing the meta data.

species A character containing the species.

miR A data.frame containing expressed miRNA genes.

miR2tar A data.frame containing expressed miRNAs and their target genes.

type A character containing the type of miRNA.

cci A data.frame containing the significantly enriched EV-derived miRNAs and their target genes.

mir_info 11

mir_info

mir_info

Description

A data.frame containing infomation of exosomal miRNA of "Human", "Mouse", and "Rat".

Usage

mir_info

Format

An object of class data. frame with 3934 rows and 5 columns.

Source

```
http://bioinfo.life.hust.edu.cn/EVmiRNA#!/
https://mirtarbase.cuhk.edu.cn/
```

pathways

pathways

Description

```
KEGG pathways of "Human", "Mouse", and "Rat".
```

Usage

pathways

Format

An object of class data. frame with 222355 rows and 5 columns.

Source

```
https://www.genome.jp/kegg/pathway.html
```

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plot_miR2tar_chord Chord plot of EV-derived miRNAs and target genes

Description

Chord plot of EV-derived miRNAs and target genes from senders to receivers.

Usage

```
plot_miR2tar_chord(
  object,
  celltype_sender,
  celltype_receiver,
  celltype_color = NULL,
  miRNA = NULL,
  edge_color = NULL,
  edge_type = "circle",
  show_type = "prob",
  text_size = 0.5,
  y_scale = 1,
  ...
)
```

Arguments

miRTalk object after find_miRTalk object celltype_sender Name of celltype_sender. One or more cell types celltype_receiver Name of celltype_receiver. One or more cell types celltype_color Colors for the celltype_sender nodes and celltype_receiver nodes, or use "NO" to make it simple miRNA which miRNAs to use. Default is to plot all inferred miRNAs edge_color Colors for the edges from the sender cell type Types for the edges from the sender cell type. Default is "circle". "big.arrow" edge_type for big arrow, "triangle" for triangle, "ellipse" for ellipse, "curved" for curved. Details see chordDiagram which type of miRNAs to show, "prob" or "score" for inferred miRNAs-target show_type interactions. Default is "prob" Size of text labels. Default is 0.5 text_size y_scale to adjust the text. Default is 1 y_scale parameters pass to chordDiagram, e.g., link.arr.width, link.arr.length, link.arr.col

Value

Chord plot of EV-derived miRNAs and target genes

plot_miR2tar_circle 13

Description

Chord plot of EV-derived miRNAs and target genes from senders to receivers.

Usage

```
plot_miR2tar_circle(
  object,
  celltype_sender,
  celltype_receiver,
  celltype_color = NULL,
  miRNA = NULL,
  edge_color = NULL,
  edge_width = 0.5,
  text_size = 3
)
```

Arguments

```
object
                  miRTalk object after find_miRTalk
celltype_sender
                  Name of celltype_sender. One cell type
celltype_receiver
                  Name of celltype_receiver. One cell type
celltype_color Colors for the celltype_sender nodes and celltype_receiver nodes, or use "NO"
                  to make it simple
miRNA
                  which miRNAs to use. Default is to plot all inferred miRNAs
edge_color
                  Colors for the edges from the sender cell type
edge_width
                  Width of edge. Default is 0.5
text_size
                  Size of text labels. Default is 3
```

Value

ggplot2 object for Circle plot of EV-derived miRNAs and target genes

Description

Heatmap plot of EV-derived miRNAs and target genes from senders to receivers.

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Usage

```
plot_miR2tar_heatmap(
  object,
  celltype_sender,
  celltype_receiver,
  miRNA = NULL,
  show_type = "prob",
  text_size = 5,
  if_horizontal = TRUE,
  viridis_option = "D",
  ...
)
```

Arguments

```
object
                  miRTalk object after find_miRTalk
celltype_sender
                  Name of celltype_sender
celltype_receiver
                  Name of celltype_receiver
                  which miRNAs to use. Default is to plot all inferred miRNAs
miRNA
                  which type of miRNAs to show, "prob" or "score" for inferred miRNAs-target
show_type
                  interactions. Default is "prob"
text_size
                  Size of text labels. Default is 3
                  Whether to plot with the horizontal direction. Default is TRUE
if_horizontal
viridis_option option in scale_color_viridis, can be "A", "B", "C", "D", "E", "F", "G",
                  "H". Default is "D".
                  parameters pass to heatmaply, e.g., grid_color
. . .
```

Value

Heatmap plot of EV-derived miRNAs and target genes

```
plot_miRTalk_chord Chord plot of cell-cell communications
```

Description

Chord plot of cell-cell communications from senders to receivers with the sum of inferred miRNAs number, activity, score, or the max probability

```
plot_miRTalk_chord(
  object,
  celltype = NULL,
  celltype_color = NULL,
  miRNA = NULL,
  edge_color = NULL,
```

plot_miRTalk_circle 15

```
edge_type = "big.arrow",
    show_type = "number",
    if_show_autocrine = FALSE,
    text_size = 1.5,
    y_scale = 0.1,
    ...
)
```

Arguments

	object	miRTalk object after find_miRTalk
	celltype	which cell types to plot by order. Default is to plot all cell types
	celltype_color	Colors for the cell types, whose length must be equal to celltype
	miRNA	which miRNAs to use. Default is to plot all inferred miRNAs
	edge_color	Colors for the edges from the sender cell type, whose length must be equal to celltype
	edge_type	Types for the edges from the sender cell type. Default is "big.arrow". "ellipse" for ellipse, "triangle" for triangle, "curved" for curved. Details see chordDiagram
	show_type	which type of miRNAs to show, "number", "activity", or "score" for sum of inferred miRNAs number and activity, respectively, or "prob" for max probability. Default is "number"
if_show_autocrine		
		Whether to show autocrine. Default is FALSE
	text_size	Size of text labels. Default is 1.5
	y_scale	y_scale to adjust the text. Default is 0.1
		parameters pass to chordDiagram, e.g., link.arr.width, link.arr.length, link.arr.col

Value

Chord plot of cell-cell communications mediated by EV-derived miRNA

Description

Circle plot of cell-cell communications from senders to receivers with the sum of inferred miRNAs number, activity, score, or the max probability

```
plot_miRTalk_circle(
  object,
  celltype = NULL,
  miRNA = NULL,
  celltype_color = NULL,
  edge_color = NULL,
  edge_type = "fan",
```

```
show_type = "number",
if_show_autocrine = FALSE,
edge_alpha = 0.5,
node_size = 10,
text_size = 5
)
```

Arguments

object miRTalk object after find_miRTalk celltype which cell types to plot. Default is to plot all cell types miRNA which miRNAs to use. Default is to plot all inferred miRNAs celltype_color Colors for the cell types, whose length must be equal to celltype Colors for the edges from the sender cell type, whose length must be equal to edge_color celltype edge_type Types for the edges. "fan" by default, "link", "hive" which type of miRNAs to show, "number", "activity", or "score" for sum show_type of inferred miRNAs number and activity, respectively, or "prob" for max probability. Default is "number" if_show_autocrine Whether to show autocrine. Default is FALSE Transparency of edge. Default is 0.5 edge_alpha Size of node. Default is 10 node_size Size of text. Default is 5 text_size

Value

ggplot2 object for Circle plot of cell-cell communications mediated by EV-derived miRNA

```
plot_miRTalk_circle_simple
```

Circle plot of cell-cell communications by retaining all cell type nodes

Description

Circle plot of cell-cell communications from senders to receivers with the sum of inferred miRNAs number, activity, score, or the max probability by retaining all cell type nodes

```
plot_miRTalk_circle_simple(
  object,
  celltype,
  celltype_dir = "sender",
  miRNA = NULL,
  celltype_color = NULL,
  edge_color = NULL,
  edge_type = "fan",
```

```
show_type = "number",
if_show_autocrine = FALSE,
edge_alpha = 0.5,
node_size = 10,
text_size = 5
)
```

Arguments

	object	miRTalk object after find_miRTalk
	celltype	which cell types to plot. one or more cell types
	celltype_dir	which direction to plot, "sender" or "receiver". Default is as "sender".
	miRNA	which miRNAs to use. Default is to plot all inferred miRNAs
	celltype_color	Colors for the cell types, whose length must be equal to celltype
	edge_color	Colors for the edges from the sender cell type, whose length must be equal to $\ensuremath{\mbox{celltype}}$
	edge_type	Types for the edges. "fan" by default, "link", "hive"
	show_type	which type of miRNAs to show, "number", "activity", or "score" for sum of inferred miRNAs number and activity, respectively, or "prob" for max probability. Default is "number"
if_show_autocrine		
		Whether to show autocrine. Default is FALSE
	edge_alpha	Transparency of edge. Default is 0.5
	node_size	Size of node. Default is 10
	text_size	Size of text. Default is 5

Value

ggplot2 object for Circle plot of cell-cell communications mediated by EV-derived miRNA

```
\verb"plot_miRTalk_heatmap" \textit{ Heatmap plot of cell-cell communications}
```

Description

Heatmap plot of cell-cell communications from senders to receivers with the sum of inferred miR-NAs number, activity, score, or the max probability

```
plot_miRTalk_heatmap(
  object,
  celltype = NULL,
  miRNA = NULL,
  show_type = "number",
  text_size = 10,
  viridis_option = "D",
  ...
)
```

plot_miRTalk_sankey

Arguments

```
miRTalk\ object\ after\ \texttt{find\_miRTalk}
object
                  which cell types to plot by order. Default is to plot all cell types
celltype
miRNA
                  which miRNAs to use. Default is to plot all inferred miRNAs
                  which type of miRNAs to show, "number", "activity", or "score" for sum
show_type
                  of inferred miRNAs number and activity, respectively, or "prob" for max prob-
                  ability. Default is "number"
text_size
                  Size of text labels. Default is 10
viridis_option option in scale_color_viridis, can be "A", "B", "C", "D", "E", "F", "G",
                   "H". Default is "D".
                  parameters pass to heatmaply, e.g., grid_color, grid_width
. . .
```

Value

Heatmap plot of cell-cell communications mediated by EV-derived miRNA

```
plot_miRTalk_sankey Sankey plot of cell-cell communications
```

Description

Sankey plot of cell-cell communications from senders to receivers with the sum of inferred miRNAs number, activity, score, or the max probability

Usage

```
plot_miRTalk_sankey(
  object,
  celltype = NULL,
  miRNA = NULL,
  celltype_color = NULL,
  edge_color = NULL,
  show_type = "number",
  if_show_autocrine = FALSE,
  edge_alpha = 0.5,
  node_size = 40,
  text_size = 15,
  node_pad = 20,
  ...
)
```

Arguments

```
object miRTalk object after find_miRTalk

celltype which cell types to plot. Default is to plot all cell types

miRNA which miRNAs to use. Default is to plot all inferred miRNAs

celltype_color Colors for the cell types, whose length must be equal to celltype
```

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edge_color	Colors for the edges from the sender cell type, whose length must be equal to celltype, Or use "NO" to cancel it
show_type	which type of miRNAs to show, "number", "activity", or "score" for sum of inferred miRNAs number and activity, respectively, or "prob" for max probability. Default is "number"
if_show_autocri	ne
	Whether to show autocrine. Default is FALSE
edge_alpha	Transparency of edge. Default is 0.5
node_size	Size of node. Default is 40
text_size	Size of text. Default is 15
node_pad	Size of node padding. Numeric essentially influences the width height. Default is 20
	parameters pass to sankeyNetwork

Value

Sankey plot of cell-cell communications mediated by EV-derived miRNA

plot_miR_bubble Bubble plot of inferred miRNA

Description

Bubble plot of inferred miRNA from senders top receivers. Rows for cell pairs, and columns for miRNAs by default.

Usage

```
plot_miR_bubble(
  object,
  celltype = NULL,
  miRNA = NULL,
  if_show_autocrine = FALSE,
  if_horizontal = TRUE,
  viridis_option = "D"
)
```

Arguments

```
object miRTalk object after find_miRTalk

celltype which cell types to plot. Default is to plot all cell types

miRNA which miRNAs to plot. Default is to plot all inferred miRNAs

if_show_autocrine

Whether to show autocrine. Default is FALSE

if_horizontal Whether to plot with the horizontal direction. Default is TRUE

viridis_option option in scale_color_viridis, can be "A", "B", "C", "D", "E", "F", "G",

"H". Default is "D".
```

Value

ggplot2 object for Bubble plot of inferred miRNA

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plot_miR_heatmap

Heatmap plot of inferred miRNA

Description

heatmap plot of inferred miRNA for each sender. Rows for cell types, and columns for miRNAs by default

Usage

```
plot_miR_heatmap(
  object,
  celltype = NULL,
  miRNA = NULL,
  text_size = 10,
  if_horizontal = TRUE,
  viridis_option = "D",
  ...
)
```

Arguments

```
object miRTalk object after find_miRTalk

celltype which cell types to plot. Default is to plot all cell types

miRNA which miRNAs to plot. Default is to plot all inferred miRNAs

text_size Size of text labels. Default is 10

if_horizontal Whether to plot with the horizontal direction. Default is TRUE

viridis_option option in scale_color_viridis, can be "A", "B", "C", "D", "E", "F", "G",

"H". Default is "D".

... parameters pass to heatmaply, e.g., grid_color, grid_width
```

Value

Heatmap plot of inferred miRNA

rev_gene

Pre-processing step: revising gene symbols

Description

Revise genes according to NCBI Gene symbols updated in June 19, 2022 for count matrix, user-custom mir_info data.frame, and mir2tar data.frame

```
rev_gene(data = NULL, data_type = NULL, species = NULL, geneinfo = NULL)
```

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Arguments

data A data.frame or matrix or dgCMatrix

data_type A character to define the type of data, select 'count' for the data matrix,

'mir_info' for the mir_info data.frame, 'mir2tar' for the mir2tar data.frame, 'pathways' for the pathways data.frame, 'GO_BP' for the GO_BP data.frame

species Species of the data. 'Human', 'Mouse' or 'Rat'

geneinfo A data.frame of the system data containing gene symbols of 'Human', 'Mouse'

and 'Rat' updated on June 19, 2022 for revising gene symbols

Value

A new data.frame, matrix, or dgCMatrix.

show,miRTalk-method

Show miRTalk object

Description

Show miRTalk object

Usage

```
## S4 method for signature 'miRTalk'
show(object)
```

Arguments

object miRTalk object

Value

miRTalk object

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