Package 'WOTPLY'

October 12, 2022

Description It shows the connections between selected clusters from the latest time point and the clusters from all the previous time points. The transition matrices between time point t and t+1 are ob-

Title Plot Connectivity Between Cells from Different Time Points

Type Package

tained from Wad	dington-OT a	ınalysis <	nttps:	//git	hub.	com/	Scia	aldo	neL	.ab/	'WO	ΓPL	Y >.	
License GPL-3														
Depends R (>= 4.0)														
Imports network, GG	ally, sna													
Suggests testthat, knit	r													
VignetteBuilder knitr														
biocViews software														
Encoding UTF-8														
RoxygenNote 7.1.1														
NeedsCompilation no)													
Author Gabriele Luba	tti [aut, cre, c	ph]												
Maintainer Gabriele	Lubatti <gabr< td=""><td>riele.luk</td><td>oatti@</td><td>helmh</td><td>oltz</td><td>-mue</td><td>nche</td><td>en.d</td><td>e></td><td></td><td></td><td></td><td></td><td></td></gabr<>	riele.luk	oatti@	helmh	oltz	-mue	nche	en.d	e>					
Repository CRAN														
Date/Publication 202	2-09-12 07:4	3:01 UTC												
D														
R topics docum	iented:													
convert_name get_transition select_top_we WOTPLY .	_matrix eights					 								. 3
Index														6
			- 1											

2 convert_names

convert_names

convert_names

Description

```
convert_names
```

Usage

```
convert_names(new_row, new_col, transition_matrix)
```

Arguments

```
new_row Vector with the new row names to assign to transition_matrix

new_col Vector with the new column names to assign to transition_matrix

transition_matrix

Output from get_transition_matrix.
```

Value

A matrix with row names equal to new_row and column names equal to new_col.

Author(s)

Gabriele Lubatti <gabriele.lubatti@helmholtz-muenchen.de>

Examples

```
transition_1 <- matrix(1, ncol = 2, nrow = 2)
colnames(transition_1) <- c("Stage1", "Stage2")
row.names(transition_1) <- c("Stage1", "Stage2")
col_name_new <- c("Stage1_new", "Stage2_new")
row_name_new <- c("Stage1_new", "Stage2_new")
transition_1 <- convert_names(row_name_new, col_name_new, transition_1)</pre>
```

get_transition_matrix 3

```
{\tt get\_transition\_matrix} \ \ {\tt get\_transition\_matrix}
```

Description

The output of *compute_all_transport_maps* from pythonpackage *WOT* is a matrix. Each entry (i,j) describes the transition probability of cell i at time t towards cluster j at time t+1. From this matrix, the average of the transition probability for all the cells at time t belonging to the same cluster is computed. Finally only the entries of the resulting matrix with above *threshold* are kept. The row names of the final matrix are equal to *level_t_plus*, while the column names are equal to the levels of *cluster_t*.

Usage

```
get_transition_matrix(path, cluster_t, threshold, cells_t)
```

Arguments

path	Character string with the path to the folder with the output of the function compute_all_transport_maps from pythonpackage WOT.
cluster_t	Vector with cluster assignment for cells at time t . The length is equal to the length of $cells_t$.
threshold	Numeric value. Only entry of the transition matrix with weight equal or above <i>threshold</i> are kept.
cells_t	Character vector with the name of cells at time t for which we want to obtain the transition matrix.

Value

A matrix with row names equal to level_t_plus and column names equal to the levels of cluster_t.

Author(s)

Gabriele Lubatti <gabriele.lubatti@helmholtz-muenchen.de>

See Also

https://broadinstitute.github.io/wot/

4 WOTPLY

```
select_top_weights
```

select_top_weights

Description

```
select_top_weights
```

Usage

```
select_top_weights(transition_matrix, top_link = NULL)
```

Arguments

transition_matrix

Output from *get_transition_matrix*.

top_link

Integer.Maximum number of links to select between clusters at time t and clusters at time t+1. Links are sorted according to the weight and then only the *top_link* are kept. If *NULL* (default), all the links are kept.

Value

A matrix

Author(s)

Gabriele Lubatti <gabriele.lubatti@helmholtz-muenchen.de>

WOTPLY

WOTPLY

Description

A ggnet2 plot is generated showing the connections between *selected_stages* from the latest time point and the clusters from previous time points. The number of columns is equal to the numbers of time points. In each column, the cluster of the corresponding time point is shown as network node. The weight of the links between clusters at time points t and t+1 reflect the weight of the transition probabilities from *list_transition_matrices*.

Usage

```
WOTPLY(
   list_transition_matrices,
   selected_stages,
   cluster_label,
   legend_time,
   customize_color,
   top_link = NULL
)
```

WOTPLY 5

Arguments

list_transition_matrices

List of transition matrices. Each matrix contains the transition probabilities from the clusters at time t (on the columns) towards the clusters at time t+1 (on the rows). The matrices can be obtain from function *get_transition_matrix*

selected_stages

Vector with the name of the clusters related to the latest time point for which we want to know the connection to clusters at previous time points.

cluster_label Vector with the cluster information for all the cells from all time points.

legend_time Vector with time information with length equal to the number of time points.

customize_color

Character vector with the name of the colour for each cluster (node) in each time point.

top_link

Integer.Maximum number of links to select between clusters at time t and clusters at time t+1. Links are sorted according to the weight and then only the *top_link* are kept. If *NULL* (default), all the links are kept.

Value

A ggnet2 plot

Author(s)

Gabriele Lubatti <gabriele.lubatti@helmholtz-muenchen.de>

See Also

```
https://CRAN.R-project.org/package=GGally
```

Examples

```
transition_1 <- matrix(1,ncol = 2,nrow = 2)
colnames(transition_1) <- c("Stage1", "Stage2")
row.names(transition_1) <- c("Stage1", "Stage2")
l_t <- list((transition_1))
selected_stages <- c("Stage1")
cluster_label <- c("Stage1", "Stage2")
legend_time <- c("Day1", "Day2")
customize_color <- c("#F8766D", "#00BFC4")
WOTPLY(l_t, selected_stages, cluster_label, legend_time, customize_color)</pre>
```

Index

```
convert_names, 2
get_transition_matrix, 3
select_top_weights, 4
WOTPLY, 4
```