Package 'cellscape'

October 14, 2021

Title Explores single cell copy number profiles in the context of a single cell tree

Version 1.16.0

Description CellScape facilitates interactive browsing of single cell clonal evolution datasets. The tool requires two main inputs:

(i) the genomic content of each single cell in the form of either copy number segments or targeted mutation values, and (ii) a single cell phylogeny. Phylogenetic formats can vary from dendrogram-like phylogenies with leaf nodes to evolutionary model-derived phylogenies with observed or latent internal nodes. The CellScape phylogeny is flexibly input as a table of source-target edges to support arbitrary representations, where each node may or may not have associated genomic data. The output of CellScape is an interactive interface displaying a single cell phylogeny and a cell-by-locus genomic heatmap representing the mutation status in each cell for each locus.

Depends R (>= 3.3)

Imports htmlwidgets (>= 0.5), jsonlite (>= 0.9.19), reshape2 (>= 1.4.1), stringr (>= 1.0.0), plyr (>= 1.8.3), dplyr (>= 0.4.3), gtools (>= 3.5.0)

biocViews Visualization

License GPL-3

LazyData true

RoxygenNote 6.0.1

Suggests knitr, rmarkdown

VignetteBuilder knitr

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R topics documented:

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Description

cellscape explores single cell copy number profiles in the context of a single cell phylogeny.

Usage

```
cellscape(cnv_data = NULL, mut_data = NULL, mut_data_matrix = NULL,
  mut_order = NULL, tree_edges, gtype_tree_edges = NULL, sc_annot = NULL,
  clone_colours = "NA", timepoint_title = "Timepoint",
  clone_title = "Clone", xaxis_title = "Time Point",
  yaxis_title = "Clonal Prevalence", phylogeny_title = "Clonal Phylogeny",
  value_type = NULL, node_type = "Cell", display_node_ids = FALSE,
  prop_of_clone_threshold = 0.2, vaf_threshold = 0.05,
  show_warnings = TRUE, width = 900, height = 800)
```

Arguments

cnv_data

data.frame (Required if not providing mut_data nor mut_data_matrix) Single cell copy number segments data. Note that every single cell id must be present in the tree_edges data frame. Required columns are:

single_cell_id: character() single cell id.
chr: character() chromosome number.

start: numeric() start position.
end: numeric() end position.

copy_number: numeric() copy number state.

 mut_data

data.frame (Required if not providing cnv_data nor mut_data_matrix) Single cell targeted mutation data frame. Note that every single cell id must be present in the tree_edges data frame. Required columns are:

single_cell_id: character() single cell id.
chr: character() chromosome number.
coord: numeric() genomic coordinate.

VAF: numeric() variant allele frequency [0, 1].

mut_data_matrix

mut_order

matrix (Required if not providing cnv_data nor mut_data) Single cell targeted mutation matrix. Rows are single cell IDs, columns are mutations. Rows and columns must be named, column names in the format "<chromosome>:<coordinate>". Note that the order of these rows and columns will not be preserved, unless mutation order is the same as that specified in the mut_order parameter. Also note

that every single cell id must be present in the tree_edges data frame.

vector (Optional) Mutation order for targeted mutation heatmap (each mutation should consist of a string in the form "chrom:coord"). Default will use a

clustering function to determine mutation order.

tree_edges data.frame Edges for the single cell phylogenetic tree. Required columns are:

source: character() edge source (single cell id). **target:** character() edge target (single cell id).

Optional columns are:

dist: numeric() edge distance.

gtype_tree_edges

data.frame (Required for TimeScape) Genotype tree edges of a rooted tree.

Required columns are:

source: character() source node id.

target: character() target node id.

sc_annot data.frame (Required for TimeScape) Annotations (genotype and sample id)

for each single cell. Required columns are:

single_cell_id: character() single cell id.

genotype: character() genotype assignment.

Optional columns are:

timepoint: character() id of the sampled time point. Note: time points will

be ordered alphabetically.

clone_colours data.frame (Optional) Clone ids and their corresponding colours (in hex for-

mat). Required columns are:

clone_id: character() clone id.

colour: character() the corresponding Hex colour for each clone id.

timepoint_title

character() (Optional) Legend title for timepoint groups. Default is "Time-

point".

clone_title character() (Optional) Legend title for clones. Default is "Clone".

xaxis_title character() (Optional) For TimeScape - x-axis title. Default is "Time Point".

yaxis_title character() (Optional) For TimeScape - y-axis title. Default is "Clonal Preva-

lence".

phylogeny_title

character() (Optional) For TimeScape - legend phylogeny title. Default is

"Clonal Phylogeny".

value_type character() (Optional) The type of value plotted in heatmap - will affect leg-

end and heatmap tooltips. Default is "VAF" for mutation data, and "CNV" for

copy number data.

node_type character() (Optional) The type of node plotted in single cell phylogeny - will affect phylogeny tooltips. Default is "Cell".

display_node_ids

logical() (Optional) Whether or not to display the single cell ID within the tree nodes. Default is FALSE.

prop_of_clone_threshold

numeric() (Optional) Used for the ordering of targeted mutations. The minimum proportion of a clone to have a mutation in order to consider the mutation as present within that clone. Default is 0.2.

vaf_threshold numeric() (Optional) Used for the ordering of targeted mutations. The mini-

mum variant allele frequency for a mutation to be considered as present within

a single cell. Default is 0.05.

show_warnings logical() (Optional) Whether or not to show any warnings. Default is TRUE.

width numeric() (Optional) Width of the plot. height numeric() (Optional) Height of the plot.

Details

Interactive components:

1. Mouseover any single cell in the phylogeny to view its corresponding genomic profile in the heatmap, and vice versa.

- 2. Mouseover any part of the heatmap to view the CNV or VAF value for that copy number segment or mutation site, respectively.
- 3. Mouseover any branch of the phylogeny to view downstream single cells, both in the phylogeny and heatmap.
- 4. Mouseover any clone to view its corresponding single cells in the phylogeny and heatmap.
- 5. Click any node in the phylogeny to flip the order of its descendant branches.
- 6. Use the selection tool in the tool bar to select single cell genomic profiles and view their corresponding single cells in the phylogeny.
- 7. Use the tree trimming tool in the tool bar to remove any branch of the phylogeny by clicking it.
- 8. Use the switch view tool in the tool bar to change the phylogeny view from force-directed to unidirectional, and vice versa.
- 9. Use the re-root phylogeny tool to root the phylogeny at a clicked node.
- 10. Use the flip branch tool to vertically rotate any branch by clicking its root node.
- 11. If present, use the scale tree/graph tool in the tool bar to scale the phylogeny by the provided edge distances.
- 12. If time-series information is present such that the TimeScape is displayed below the CellScape, clones and time points are interactively linked in both views on mouseover.
- 13. Click the download buttons to download a PNG or SVG of the view.

Note:

See TimeScape repo (https://bitbucket.org/MO_BCCRC/timescape) for more information about TimeScape.

Examples

```
library("cellscape")
# EXAMPLE 1 - TARGETED MUTATION DATA
# single cell tree edges
tree_edges <- read.csv(system.file("extdata", "targeted_tree_edges.csv",</pre>
    package = "cellscape"))
# targeted mutations
targeted_data <- read.csv(system.file("extdata", "targeted_muts.csv",</pre>
    package = "cellscape"))
# genotype tree edges
gtype_tree_edges <- data.frame("source"=c("Ancestral", "Ancestral", "B",</pre>
    "C", "D"), "target"=c("A", "B", "C", "D", "E"))
# annotations
sc_annot <- read.csv(system.file("extdata", "targeted_annots.csv",</pre>
    package = "cellscape"))
# mutation order
mut_order <- scan(system.file("extdata", "targeted_mut_order.txt",</pre>
    package = "cellscape"), what=character())
# run cellscape
cellscape(mut_data=targeted_data, tree_edges=tree_edges, sc_annot =
    sc_annot, gtype_tree_edges=gtype_tree_edges, mut_order=mut_order)
# EXAMPLE 2 - COPY NUMBER DATA
# single cell tree edges
tree_edges <- read.csv(system.file("extdata", "cnv_tree_edges.csv",</pre>
    package = "cellscape"))
# cnv segments data
cnv_data <- read.csv(system.file("extdata", "cnv_data.csv", package =</pre>
    "cellscape"))
# annotations
sc_annot <- read.csv(system.file("extdata", "cnv_annots.tsv", package =</pre>
    "cellscape"), sep="\t")
# custom clone colours
clone_colours <- data.frame( clone_id = c("1","2","3"),</pre>
    colour = c("7fc97f", "beaed4", "fdc086"))
# run cellscape
cellscape(cnv_data=cnv_data, tree_edges=tree_edges, sc_annot=sc_annot,
```

width=800, height=475, show_warnings=FALSE, clone_colours = clone_colours)

dfs_tree

Get depth first search of a tree

Description

Get depth first search of a tree

Widget output function for use in Shiny

Widget render function for use in Shiny

Function to get data frame of pixels

function to get min and max values for each chromosome

function to get chromosome box pixel info

function to get the genome length

function to get the number of base pairs per pixel

function to get information (chr, start, end, mode_cnv) for each pixel

function to get mutation order for targeted data

function to get targeted heatmap information

function to find the mode of a vector

Function to process the user data

Function to check minimum dimensions

Function to check required inputs are present

check alpha value input is correct

check clonal_prev parameter data

check tree_edges parameter data

check genotype_position parameter

check clone_colours parameter

check perturbations parameter

get mutation data

function to replace spaces with underscores in all data frames & keep maps of original names to space-replaced names

Usage

```
dfs_tree(edges, cur_root, dfs_arr)
cellscapeOutput(outputId, width = "100%", height = "400px")
renderCnvTree(expr, env = parent.frame(), quoted = FALSE)
getEmptyGrid(hm_sc_ids_ordered, ncols)
getChromBounds(chroms, cnv_data)
getChromBoxInfo(chrom_bounds, n_bp_per_pixel)
getGenomeLength(chrom_bounds)
getNBPPerPixel(ncols, chrom_bounds, genome_length)
getCNVHeatmapForEachSC(cnv_data, chrom_bounds, n_bp_per_pixel)
getMutOrder(mut_data)
getTargetedHeatmapForEachSC(mut_data, mut_order, heatmapWidth)
findMode(x)
processUserData(clonal_prev, tree_edges, mutations, clone_colours, xaxis_title,
  yaxis_title, phylogeny_title, alpha, genotype_position, perturbations, sort,
  show_warnings, width, height)
checkMinDims(mutations, height, width)
checkRequiredInputs(clonal_prev, tree_edges)
checkAlpha(alpha)
checkClonalPrev(clonal_prev)
checkTreeEdges(tree_edges)
checkGtypePositioning(genotype_position)
checkCloneColours(clone_colours)
checkPerts(perturbations)
getMutationsData(mutations, tree_edges, clonal_prev)
replaceSpaces(clonal_prev, tree_edges, clone_colours, mutation_info, mutations,
```

mutation_prevalences)

Arguments

edges – edges of tree

cur_root – current root of the tree

dfs_arr - array of depth first search results to be filled

outputId - id of output

width - width of output

height - height of output

expr - expression for Shiny

env - environment for Shiny

quoted - default is FALSE

hm_sc_ids_ordered

- array of single cell ids in order

ncols – number of columns in heatmap/grid

chroms – vector of chromosome names

cnv_data – copy number data

chrom_bounds – data frame of chromosome boundaries n_bp_per_pixel – integer of number of base pairs per pixel

genome_length - integer of length of the genome
mut_data - data frame of mutations data

mut_order – array of order of mutations for heatmap (chromosome:coordinate)

heatmapWidth — number for width of the heatmap (in pixels)

x – vector of numbers

clonal_prev - data frame of Clonal prevalence. Note: timepoints will be alphanumerically

sorted in the view. Format: columns are (1) character() "timepoint" - time point (2) character() "clone_id" - clone id (3) numeric() "clonal_prev" - clonal preva-

lence.

tree_edges — data frame of Tree edges of a rooted tree. Format: columns are (1) character()

"source" - source node id (2) character() "target" - target node id.

mutations - data frame (Optional) of Mutations occurring at each clone. Any additional

field will be shown in the mutation table. Format: columns are (1) character() "chrom" - chromosome number (2) numeric() "coord" - coordinate of mutation on chromosome (3) character() "clone_id" - clone id (4) character() "timepoint" - time point (5) numeric() "VAF" - variant allele frequency of the mutation in

the corresponding timepoint.

clone_colours - data frame (Optional) of Clone ids and their corresponding colours Format:

columns are (1) character() "clone_id" - the clone ids (2) character() "colour" -

the corresponding Hex colour for each clone id.

xaxis_title - String (Optional) of x-axis title. Default is "Time Point".

yaxis_title — String (Optional) of y-axis title. Default is "Clonal Prevalence". phylogeny_title

- String (Optional) of Legend phylogeny title. Default is "Clonal Phylogeny".

alpha – Number (Optional) of Alpha value for sweeps, range [0, 100].

genotype_position

String (Optional) of How to position the genotypes from ["centre", "stack", "space"] "centre" – genotypes are centred with respect to their ancestors "stack"
 genotypes are stacked such that no genotype is split at any time point "space"

- genotypes are stacked but with a bit of spacing at the bottom

perturbations - data frame (Optional) of any perturbations that occurred between two time

points. Format: columns are (1) character() "pert_name" - the perturbation name (2) character() "prev_tp" - the time point (as labelled in clonal prevalence data)

BEFORE perturbation.

sort – Boolean (Optional) of whether (TRUE) or not (FALSE) to vertically sort the

genotypes by their emergence values (descending). Default is FALSE. Note that genotype sorting will always retain the phylogenetic hierarchy, and this

parameter will only affect the ordering of siblings.

show_warnings - Boolean (Optional) of Whether or not to show any warnings. Default is TRUE.

mutation_info - processed mutation_info

mutation_prevalences

- mutation_prevalences data from user

chrom_bounds - data frame of chromosome boundaries
ncols - integer of number of columns (pixels) to fill

chrom_bounds - data frame of chromosome boundaries

cnv_data - data frame of copy number variant segments data

chrom_bounds – data frame of chromosome boundaries n_bp_per_pixel – integer of number of base pairs per pixel

mut_data — data frame of mutations data

width – Number (Optional) of width of the plot. Minimum width is 450.

height – Number (Optional) of height of the plot. Minimum height with and without

mutations is 500 and 260, respectively.

mutations — mutations provided by user
height — height provided by user
width — width provided by user
clonal_prev — clonal_prev provided by user
tree_edges — tree_edges provided by user
alpha — alpha provided by user

clonal_prev — clonal prevalence provided by user

tree_edges — tree edges provided by user

genotype_position

genotype_position provided by user

```
clone_colours

    clone_colours provided by user

perturbations
                  - perturbations provided by user
mutations
                  - mutations data from user
tree_edges
                  - tree edges data from user
clonal_prev
                  - clonal prevalence data from user
clonal_prev
                  – clonal_prev data from user
tree_edges
                  - tree edges data from user
clone_colours
                  - clone colours data from user
mutations
                  - mutations data from user
```

Examples

```
dfs_tree(data.frame(source = c("1","1","2","2","5","6"), target=c("2","5","3","4","6","7")), "1", c())
cellscapeOutput(1, '100%', '300px')
cellscapeOutput(1, '80%', '300px')
findMode(c(1,1,19,1))
checkMinDims(data.frame(chr = c("11"), coord = c(104043), VAF = c(0.1)), "700px", "700px")
checkRequiredInputs(data.frame(timepoint = c(rep("Diagnosis", 6), rep("Relapse", 1)), clone_id = c("1","2","3","4
data.frame(source = c("1","1","2","2","5","6"), target=c("2","5","3","4","6","7")))
checkRequiredInputs(data.frame(timepoint = c(rep("Diagnosis", 6), rep("Relapse", 1)), clone_id = c("1","2","3","4
data.frame(source = c("1","1","2","2","5","6"), target=c("2","5","3","4","6","7")))
checkAlpha(4)
checkAlpha(100)
checkClonalPrev(data.frame(timepoint=c(1), clone_id=c(2), clonal_prev=c(0.1)))
checkTreeEdges(data.frame(source = c("1","1","2","2","5","6"), target=c("2","5","3","4","6","7")))
checkGtypePositioning("centre")
checkCloneColours(data.frame(clone_id = c("1","2","3", "4"), colour = c("#beaed4", "#fdc086", "#beaed4", "#b
checkPerts(data.frame(pert_name = c("New Drug"), prev_tp = c("Diagnosis")))
getMutationsData(data.frame(chrom = c("11"), coord = c(104043), VAF = c(0.1), clone_id=c(1), timepoint=c("Relapse")
data.frame(source = c("1","1","2","2","5","6"), target=c("2","5","3","4","6","7")),
data.frame(timepoint = c(rep("Diagnosis", 6), rep("Relapse", 1)), clone_id = c("1","2","3","4","5","6","7"), clon
replaceSpaces(mutations = data.frame(chrom = c("11"), coord = c(104043), VAF = c(0.1), clone_id=c(1), timepoint=c(1)
clonal_prev = data.frame(timepoint = c(rep("Diagnosis", 6), rep("Relapse", 1)), clone_id = c("1","2","3","4","5","
mutation_prevalences = list("X:6154028" = data.frame(timepoint = c("Diagnosis"), VAF = c(0.5557))), mutation_info
clone_colours = data.frame(clone_id = c("1","2","3", "4"), colour = c("#beaed4", "#fdc086", "#beaed4", "#beaed4"))
```

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