Package 'CancerEvolutionVisualization'

November 21, 2023

Title Publication Quality Phylogenetic Tree Plots
Version 2.0.1
Date 2023-11-17
Description Generates tree plots with precise branch lengths, gene annotations, and cellular prevalence. The package handles complex tree structures (angles, lengths, etc.) and can be further refined as needed by the user.
License GPL-2
${\bf URL}\ {\tt https://github.com/uclahs-cds/package-CancerEvolutionVisualization}$
<pre>BugReports https://github.com/uclahs-cds/package-CancerEvolutionVisualization/issues</pre>
Depends R (>= 3.5.0), graphics, grid, gridExtra, gtable,
Imports plyr, grDevices, utils, stringr, BoutrosLab.plotting.general
Suggests testthat, knitr
VignetteBuilder knitr
LazyLoad yes
LazyData yes
NeedsCompilation no
Author Paul Boutros [cre], Adriana Salcedo [aut], Dan Knight [aut], Helena Winata [ctb]
Maintainer Paul Boutros <pboutros@mednet.ucla.edu></pboutros@mednet.ucla.edu>
Repository CRAN
Date/Publication 2023-11-21 19:40:02 UTC
R topics documented:
colours 2 SRCGrob 2
Index 6

colours

Colour scheme vector

Description

Default colours used by CP polygons, etc.

Format

Character vector

SRCGrob

Subclone Tree Plot

Description

Creates a phylogenetic tree using subclonal reconstruction data. An optional node text data.frame can be used to annotate notable genes alonside branches.

Usage

```
SRCGrob(
    tree,
    node.text = NULL,
   main = NULL,
   horizontal.padding = 0.1,
    scale1 = 1,
    scale2 = 1,
    yat = NULL,
   yaxis1.label = NULL,
   yaxis2.label = NULL,
    xlab.cex = 1.55,
   ylab.cex = 1.55,
    xaxis.cex = 1.45,
    yaxis.cex = 1.45,
    xaxis.label = "CP",
    label.cex = NA,
    node.text.cex = 0.85,
   main.y = NULL,
   main.cex = 1.7,
    node.radius = 0.1,
    node.text.line.dist = 0.1,
    colour.scheme = CancerEvolutionVisualization::colours,
    draw.nodes = TRUE,
    add.normal = FALSE,
```

```
use.radians = FALSE,
normal.cex = 1,
sig.shape = 3,
label.nodes = TRUE,
disable.polygons = FALSE,
length.from.node.edge = TRUE,
size.units = "npc"
);
```

Arguments

tree Tree structure data.frame

node. text Dataframe for text labels to be displayed next to nodes

main Main plot title

horizontal.padding

Increase/reduce the plot's horizontal padding proportionally. A positive value

will expand the padding, and a negative value will reduce it.

scale1 Proportionally scale the values of the first branch length column in the tree input.

scale2 Proportionally scale the values of the second branch length column in the tree

input.

yat Specific values to be used for the y-axis ticks. A list is required, with each

element corresponding to an axis.

yaxis1.label Text label for the first, leftmost y-axis

yaxis2.label Text label for the second, rightmost y-axis

xlab.cex Font size for the x-axis label
ylab.cex Font size for the y-axis labels
xaxis.cex Font size for the x-axis tick labels
yaxis.cex Font size for the y-axis tick labels

xaxis.label Text label for the x-axis
label.cex Font size for the node labels
node.text.cex Font size for the node text

main.y Move the main plot title position up or down

main.cex Font size for the main plot title

node.radius Node size
node.text.line.dist

Distance between node text and tree branches (as a value between 0 and 1)

colour.scheme Vector of colour values to be used for CP polygons

draw.nodes Enable or disable drawing tree nodes

add.normal Adds a normal

use.radians Unit to be used for "angle" column (degrees or radians)

normal.cex Font size within the normal "box"

Value

A 'grob' of class "SRCGrob"

Author(s)

Dan Knight

Examples

```
# Simple Tree Plot
simple.tree <- data.frame(</pre>
    parent = c(NA, 1, 1)
    );
SRCGrob(simple.tree);
# Specify Branch Lengths
branch.lengths.tree <- data.frame(</pre>
    simple.tree,
    length1 = c(10, 20, 15)
    );
SRCGrob(branch.lengths.tree);
# Cellular Prevalence
CP.tree <- data.frame(</pre>
    branch.lengths.tree,
    CP = c(1, 0.3, 0.5)
    );
SRCGrob(CP.tree);
# Scaled Branch Lengths
SRCGrob(
    branch.lengths.tree,
    scale1 = 0.8
    );
```

```
# Override Branch Angles in Degrees
degrees.tree <- data.frame(</pre>
    parent = c(NA, 1, 2),
    angle = c(NA, NA, 90)
SRCGrob(degrees.tree);
# Override Branch Angles in Radians
radians.tree <- data.frame(</pre>
    parent = c(NA, 1, 2),
    angle = c(NA, NA, pi / 2)
    );
SRCGrob(
    radians.tree,
    use.radians = TRUE
    );
# Plot Title
SRCGrob(
    simple.tree,
    main = 'Example Plot'
    );
# Y-Axis Label
SRCGrob(
    branch.lengths.tree,
    yaxis1.label = 'SNVs'
    );
# Y-Axis Ticks
yaxis1.ticks <- seq(0, 10, 2);</pre>
SRCGrob(
    branch.lengths.tree,
    yat = list(yaxis1.ticks)
    );
# Normal
SRCGrob(
    simple.tree,
    add.normal = TRUE
    );
```

Index

colours, 2

SRCGrob, 2