Package 'ggtree'

April 23, 2023

```
Type Package
Title an R package for visualization of tree and annotation data
Version 3.7.2
Maintainer Guangchuang Yu <guangchuangyu@gmail.com>
Description 'ggtree' extends the 'ggplot2' plotting system which implemented
     the grammar of graphics. 'ggtree' is designed for visualization and annotation
     of phylogenetic trees and other tree-like structures with their annotation data.
Depends R (>= 3.5.0)
Imports ape, aplot, dplyr, ggplot2 (> 3.3.6), grid, magrittr, methods,
     purrr, rlang, ggfun (>= 0.0.9), yulab.utils, tidyr, tidytree
     (>= 0.3.9), treeio (>= 1.8.0), utils, scales, stats, cli
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BugReports https://github.com/YuLab-SMU/ggtree/issues
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${\sf R}$ topics documented:

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add_colorbar

 $add_colorbar$

Description

add colorbar legend

Usage

```
add_colorbar(p, color, x = NULL, ymin = NULL, ymax = NULL, font.size = 4)
```

Arguments

p tree view

color output of scale_color function

x x position
ymin ymin
ymax ymax
font.size font size

Value

ggplot2 object

Author(s)

applyLayoutDaylight 5

 $apply Layout Daylight \qquad apply Layout Daylight$

Description

Apply the daylight alorithm to adjust the spacing between the subtrees and tips of the specified node.

Usage

```
applyLayoutDaylight(df, node_id)
```

Arguments

df tree data.frame

node_id is id of the node from which daylight is measured to the other subtrees.

Value

list with tree data.frame with updated layout using daylight algorithm and max_change angle.

as.polytomy as.polytomy

Description

collapse binary tree to polytomy by applying 'fun' to 'feature'

Usage

```
as.polytomy(tree, feature, fun)
```

Arguments

tree object, 'phylo' object only

feature selected feature

fun function to select nodes to collapse

Value

polytomy tree

Author(s)

6 collapse.ggtree

collapse.ggtree

collapse-ggtree

Description

collapse a selected clade, which can later be expanded with the 'expand()' fuction if necessary

Usage

```
## S3 method for class 'ggtree'
collapse(x = NULL, node, mode = "none", clade_name = NULL, ...)
```

Arguments

x tree view (i.e. the ggtree object). If tree_view is NULL, the last ggplot will be

used.

node internal node number

mode one of 'none' (default), 'max', 'min' and 'mixed'. 'none' would simply collapse

the clade as 'tip' and the rest will display a triangle, whose shape is determined

by the farest/closest tip of the collapsed clade to indicate it

clade_name set a name for the collapsed clade. If clade_name = NULL, do nothing

... additional parameters to set the color or transparency of the triangle

Value

tree view

Author(s)

Guangchuang Yu

See Also

expand

Examples

```
x <- rtree(15)
p <- ggtree(x) + geom_tiplab()
p
p1 <- collapse(p, node = 17, mode = "mixed", clade_name = "cclade", alpha = 0.8, color = "grey", fill = "light blue")</pre>
```

Date2decimal 7

Date2decimal

Date2decimal

Description

convert Date to decimal format, eg "2014-05-05" to "2014.34"

Usage

Date2decimal(x)

Arguments

Χ

Date

Value

numeric

Author(s)

Guangchuang Yu

decimal2Date

decimal2Date

Description

convert decimal format to Date, eg "2014.34" to "2014-05-05"

Usage

```
decimal2Date(x)
```

Arguments

Χ

numerical number, eg 2014.34

Value

Date

Author(s)

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expand

expand

Description

expand collapsed clade

Usage

```
expand(tree_view = NULL, node)
```

Arguments

tree_view tree view (i.e. the ggtree object). If tree_view is NULL, the last ggplot object

will be used.

node internal node number to specify a clade. If NULL, using the whole tree

Value

tree view

Author(s)

Guangchuang Yu

See Also

collapse

Examples

```
x <- rtree(15)
p <- ggtree(x) + geom_tiplab()
p1 <- collapse(p, 17)
expand(p1, 17)</pre>
```

facet_data

facet_data

Description

```
extract data used in facet_plot or geom_facet
```

Usage

```
facet_data(tree_view, panel)
```

facet_labeller 9

Arguments

tree_view ggtree object

panel data plotted in specific panel. If only one dataset used in the panel, return the

data frame, else return a list of data frames.

Value

data frame or a list of data frames

Author(s)

Guangchuang Yu

References

G Yu, TTY Lam, H Zhu, Y Guan (2018). Two methods for mapping and visualizing associated data on phylogeny using ggtree. Molecular Biology and Evolution, 35(2):3041-3043. https://doi.org/10.1093/molbev/msy194

facet_labeller

facet_labeller

Description

function to relable selected panels created by 'geom_facet' or 'facet-plot'

Usage

```
facet_labeller(p, label)
```

Arguments

p facet_plot output

label new labels of facet panels

Value

ggplot object

Author(s)

10 facet_plot

Description

plot tree associated data in an additional panel

Usage

```
facet_plot(p, mapping = NULL, data, geom, panel, ...)
geom_facet(mapping = NULL, data, geom, panel, ...)
```

Arguments

p	tree view
mapping	aes mapping for 'geom'
data	data to plot by 'geom', first column should be matched with tip label of tree
geom	geom function to plot the data
panel	panel name for plot of input data
	additional parameters for 'geom'

Details

'facet_plot()' automatically re-arranges the input 'data' according to the tree structure, visualizes the 'data' on specific 'panel' using the 'geom' function with aesthetic 'mapping' and other parameters, and align the graph with the tree 'p' side by side. 'geom_facet' is a 'ggplot2' layer version of 'facet_plot'

Value

ggplot object

Author(s)

Guangchuang Yu

References

G Yu, TTY Lam, H Zhu, Y Guan (2018). Two methods for mapping and visualizing associated data on phylogeny using ggtree. Molecular Biology and Evolution, 35(2):3041-3043. https://doi.org/10.1093/molbev/msy194

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Examples

```
tr <- rtree(10)
dd = data.frame(id=tr$tip.label, value=abs(rnorm(10)))
p <- ggtree(tr)
facet_plot(p, 'Trait', data = dd, geom=geom_point, mapping=aes(x=value))</pre>
```

facet_widths

facet_widths

Description

set relative widths (for column only) of facet plots

Usage

```
facet_widths(p, widths)
```

Arguments

p ggplot or ggtree objectwidths relative widths of facet panels

Value

ggplot object by redrawing the figure (not a modified version of input object)

Author(s)

Guangchuang Yu

flip

flip

Description

exchange the position of 2 clades

Usage

```
flip(tree_view = NULL, node1, node2)
```

Arguments

tree_view tree view (i.e. the ggtree object). If tree_view is NULL, the last ggplot will be

used.

node number of clade 1. It should share a same parent node with node2 node number of clade 2. It should share a same parent node with node1 12 geom_aline

Value

ggplot object

Author(s)

Guangchuang Yu

Examples

```
set.seed(123)
x <- rtree(15)
p <- ggtree(x) + geom_tiplab() +
  geom_nodelab(aes(subset=!isTip, label=node), hjust = -.1, color = "red")
flip(p, 23, 24) ## Depends on the condition of your tree</pre>
```

geom_aline

geom_aline

Description

add horizontal align lines layer to a tree

Usage

```
geom_aline(mapping = NULL, linetype = "dotted", linewidth = 1, ...)
```

Arguments

mapping aes mapping

linetype set line type of the line, defaults to "dotted"

linewidth set width of the line, defaults to 1

... additional parameter

Details

'geom_aline'align all tips to the longest one by adding padding characters to the right side of the tip.

Value

aline layer

Author(s)

Yu Guangchuang

geom_balance 13

Description

highlights the two direct descendant clades of an internal node

Usage

```
geom_balance(
  node,
  fill = "steelblue",
  color = "white",
  alpha = 0.5,
  extend = 0,
  extendto = NULL
)
```

Arguments

fill color to fill in the highlight rectangle, default to "steelblue" color color to outline highlight rectangle and divide balance, defaults to "white" alpha alpha (transparency) for the highlight rectangle, defaults to 0.5 extend extend xmax of the highlight rectangle by the value of extend extendto extend xmax of the highlight rectangle to the value of extendto	node	selected node (balance) to highlight its two direct descendant
alpha alpha (transparency) for the highlight rectangle, defaults to 0.5 extend extend xmax of the highlight rectangle by the value of extend	fill	color to fill in the highlight rectangle, default to "steelblue"
extend extend xmax of the highlight rectangle by the value of extend	color	color to outline highlight rectangle and divide balance, defaults to "white"
	alpha	alpha (transparency) for the highlight rectangle, defaults to 0.5
extendto extend xmax of the highlight rectangle to the value of extendto	extend	extend xmax of the highlight rectangle by the value of extend
	extendto	extend xmax of the highlight rectangle to the value of extendto

Details

Particularly useful when studying neighboring clades. Note that balances that correspond to multichotomies will not be displayed.

Value

ggplot2

Author(s)

Justin Silverman and modified by Guangchuang Yu

References

J. Silverman, et al. *A phylogenetic transform enhances analysis of compositional microbiota data*. (in preparation)

For more detailed demonstration, please refer to chapter 5.2.2 of *Data Integration, Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

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Examples

```
library(ggtree)
set.seed(123)
tr<- rtree(15)
x <- ggtree(tr)
x + geom_balance(17)</pre>
```

geom_cladelab

geom cladelab

Description

annotate a clade with bar and text label or (image)

Usage

```
geom_cladelab(
  node = NULL,
  label = NULL,
  data = NULL,
  mapping = NULL,
  geom = "text",
  parse = FALSE,
  ...
)
```

Arguments

node selected node to annotate, when data and mapping is NULL, it is required. label character, character to be showed, when data and mapping is NULL, it is required. data data.frame, the data to be displayed in the annotation, defaults to NULL. Set of aesthetic mappings, defaults to NULL. The detail see the following exmapping planation. character, one of 'text', 'label', 'shadowtext', 'image' and 'phylopic', defaults geom to 'text', and the parameter see the Aesthetics For Specified Geom. logical, whether parse label to emoji font, defaults to FALSE. parse additional parameters, see also following section. . . . additional parameters can refer the following parameters.

- offset distance bar and tree, offset of bar and text from the clade, defaults to 0
- offset.text distance bar and text, offset of text from bar, defaults to 0.
- align logical, whether align clade lab, defaults to FALSE.

geom_cladelab 15

- extend numeric, extend the length of bar, defaults to 0.
- angle numeric or 'auto', if angle is auto, the angle of text will be calculated automatically, which is useful for the circular etc layout, defaults to 0.
- horizontal logical, whether set label to horizontal, defaults to TRUE.
- barsize the width of line, defaults to 0.5.
- barcolour the colour of line, defaults to 'black'.
- fontsize the size of text, defaults to 3.88.
- textcolour the colour of text, defaults to 'black'.
- imagesize the size of image, defaults to 0.05.
- imagecolor the colour of image, defaults to NULL, when geom="phylopic", it should be required.

The parameters also can be set in mapping, when data is provided. Note: the barsize, barcolour, fontsize, textcolour, imagesize and imagecolor should not be set in mapping (aesthetics). When the color and size are not be set in mapping, user can modify them to adjust the attributes of specified geom.

Aesthetics For Specified Geom

geom_cladelab() understands the following aesthetics for geom="text"(required aesthetics are in bold):

- node selected node to hight light, it is required.
- label labels to be shown, it is required.
- colour the colour of text, defaults to "black".
- size the size of text, defaults to 3.88.
- angle the angle of text, defaults to 0.
- hjust A numeric vector specifying horizontal justification, defaults to 0.
- vjust A numeric vector specifying vertical justification, defaults to 0.5.
- alpha the transparency of text, defaults to NA.
- family the family of text, defaults to 'sans'.
- font face the font face of text, defaults to 1 (plain), others are 2 (bold), 3 (italic), 4 (bold.italic).
- lineheight The height of a line as a multiple of the size of text, defaults to 1.2.

when the colour, size are not be set in mapping, and user want to modify the colour of text, they should use textcolour, fontsize to avoid the confusion with bar layer annotation.

geom_cladelab() understands the following aesthethics for geom="label" (required aesthetics are in bold):

- node selected node to hight light, it is required.
- label labels to be shown, it is required.
- colour the colour of text, defaults to "black".
- fill the background colour of the label, defaults to "white".
- size the size of text, defaults to 3.88.

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- angle the angle of text, defaults to 0.
- hjust A numeric vector specifying horizontal justification, defaults to 0.
- vjust A numeric vector specifying vertical justification, defaults to 0.5.
- alpha the transparency of text, defaults to NA.
- family the family of text, defaults to 'sans'.
- font face the font face of text, defaults to 1 (plain), others are 2 (bold), 3 (italic), 4 (bold.italic).
- lineheight The height of a line as a multiple of the size of text, defaults to 1.2.

when the colour, size are not be set in mapping, and user want to modify the colour of text, they should use textcolour, fontsize to avoid the confusion with bar layer annotation.

geom_cladelab() understands the following aesthethics for geom="shadowtext" (required aesthetics are in bold):

- node selected node to hight light, it is required.
- label labels to be shown, it is required.
- colour the colour of text, defaults to "black".
- bg.colour the background colour of text, defaults to 'black'.
- bg.r the width of background text, defaults to 0.1.
- size the size of text, defaults to 3.88.
- angle the angle of text, defaults to 0.
- hjust A numeric vector specifying horizontal justification, defaults to 0.
- vjust A numeric vector specifying vertical justification, defaults to 0.5.
- alpha the transparency of text, defaults to NA.
- family the family of text, defaults to 'sans'.
- fontface the font face of text, defaults to 1 (plain), others are 2 (bold), 3 (italic), 4 (bold.italic).
- lineheight The height of a line as a multiple of the size of text, defaults to 1.2.

when the colour, size are not be set in mapping, and user want to modify the colour of text, they should use textcolour, fontsize to avoid the confusion with bar layer annotation.

geom_cladelab() understands the following aesthethics for geom="image" or geom="phylopic" (required aesthetics are in bold):

- node selected node to hight light, it is required.
- label labels to be shown, it is required.
- image the image to be annotated, when geom="phylopic", the uid of phylopic databases, it is required.
- colour the color of image, defaults to NULL.
- size the size of image, defaults to 0.05.
- alpha the alpha of image, defaults to 0.8.

when the colour, size are not be set in mapping, and user want to modify the colour of image, they should use imagecolour, imagesize to avoid the confusion with bar layer annotation.

geom_cladelabel 17

Examples

```
set.seed(2015-12-21)
tree <- rtree(30)</pre>
data <- data.frame(id=c(34, 56),</pre>
                    annote=c("another clade", "long clade names"),
                    image=c("7fb9bea8-e758-4986-afb2-95a2c3bf983d",
                            "0174801d-15a6-4668-bfe0-4c421fbe51e8"),
                    group=c("A", "B"),
                    offset=c(0.1, 0.1),
                    offset.text=c(0.1, 0.2))
p <- ggtree(tree) + xlim(NA, 6)</pre>
p + geom_cladelab(node=45, label="test label") +
    geom_cladelab(node=34, label="another clade")
p2 <- p + geom_cladelab(data=data,</pre>
                         mapping=aes(
                              node=id,
                              label=annote,
                              image=image,
                              color=group,
                              offset=offset
                         ),
                         geom="shadowtext",
                         hjust=0.5,
                         align=TRUE,
                         horizontal=FALSE,
                         angle=90,
                         show.legend = FALSE
p2
```

geom_cladelabel

geom_cladelabel

Description

annotate a clade with bar and text label

Usage

```
geom_cladelabel(
  node,
  label,
  offset = 0,
  offset.text = 0,
  extend = 0,
  align = FALSE,
  barsize = 0.5,
```

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```
fontsize = 3.88,
  angle = 0,
  geom = "text",
  hjust = 0,
  color = NULL,
  fill = NA,
  family = "sans",
  parse = FALSE,
  horizontal = TRUE,
  ...
)
```

Arguments

node selected node label clade label

offset of bar and text from the clade

offset.text offset of text from bar extend extend bar height

align logical size of bar fontsize size of text angle angle of text

geom one of 'text' or 'label' hjust justify text horizontally

color color for clade & label, of length 1 or 2

fill label background, only work with geom='label'

family sans by default, can be any supported font

parse logical, whether parse label

horizontal logical, whether set label to horizontal, defaults to TRUE.

... additional parameter

Value

ggplot layers

Author(s)

Guangchuang Yu

See Also

geom_cladelabel2

geom_cladelabel2

geom_cladelabel2

 $geom_cladelabel2$

Description

annotate a clade with bar and text label

Usage

```
geom_cladelabel2(
  node,
  label,
 offset = 0,
  offset.text = 0,
  offset.bar = 0,
  align = FALSE,
 barsize = 0.5,
  fontsize = 3.88,
 hjust = 0,
  geom = "text",
  color = NULL,
  family = "sans",
  parse = FALSE,
 horizontal = TRUE,
)
```

Arguments

```
selected node
node
                   clade label
label
offset
                   offset of bar and text from the clade
offset.text
                   offset of text from bar
offset.bar
                   offset of bar from text
                  logical
align
                  size of bar
barsize
fontsize
                   size of text
                  justify text horizontally
hjust
                   one of 'text' or 'label'
geom
                  color for clade & label, of length 1 or 2
color
family
                   sans by default, can be any supported font
parse
                   logical, whether parse label
                  logical, whether set label to horizontal, defaults to TRUE.
horizontal
                   additional parameter
```

20 geom_hilight

Value

ggplot layers

Author(s)

JustGitting

See Also

geom_cladelabel

geom_hilight

geom_hilight

Description

layer of hilight clade

Usage

```
geom_hilight(
  data = NULL,
  mapping = NULL,
  node = NULL,
  type = "auto",
  to.bottom = FALSE,
  ...
)

geom_highlight(
  data = NULL,
  mapping = NULL,
  node = NULL,
  type = "auto",
  to.bottom = FALSE,
  ...
)
```

Arguments

data.frame, The data to be displayed in this layer, defaults to NULL.

mapping Set of aesthetic mappings, defaults to NULL.

node selected node to hilight, when data and mapping is NULL, it is required.

geom_hilight 21

type

the type of layer, defaults to auto, meaning rectangular, circular, slanted, fan, inward_circular, radial, equal_angle, ape layout tree will use rectangular layer, unrooted and daylight layout tree use will use encircle layer. You can specify this parameter to rect (rectangular layer) or encircle (encircle layer), 'gradient' (gradient color), 'roundrect' (round rectangular layer).

to.bottom

logical, whether set the high light layer to the bottom in all layers of 'ggtree' object, default is FALSE.

. . .

additional parameters, see also the below and Aesthetics section.

- align control the align direction of the edge of high light rectangular. Options is 'none' (default), 'left', 'right', 'both'. This argument only work when the 'geom_hilight' is plotting using geom_hilight(mapping=aes(...)).
- gradient.direction character, the direction of gradient color, defaults to 'rt' meaning the locations of gradient color is from root to tip, options are 'rt' and 'tr'.
- gradient.length.out integer, desired length of the sequence of gradient color, defaults to 2.
- roundrect.r numeric, the radius of the rounded corners, when roundrect=TRUE, defaults to 0.05.

Details

geom_hilight supports data.frame as input. And aesthetics of layer can be mapped. you can see the Aesthetics section to set parameters.

Value

a list object.

Aesthetics

geom_hilight() understands the following aesthetics for rectangular layer (required aesthetics are in bold):

- node selected node to hight light, it is required.
- colour the colour of margin, defaults to NA.
- fill the colour of fill, defaults to 'steelblue'.
- alpha the transparency of fill, defaults to 0.5.
- extend extend xmax of the rectangle, defaults to 0.
- extendto specify a value, meaning the rectangle extend to, defaults to NULL.
- linetype the line type of margin, defaults to 1.
- linewidth the width of line of margin, defaults to 0.5.

geom_hilight() understands the following aesthethics for encircle layer (required aesthetics are in bold):

• node selected node to hight light, it is required.

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- colour the colour of margin, defaults to 'black'.
- fill the colour of fill, defaults to 'steelblue'.
- alpha the transparency of fill, defaults to 0.5.
- expand expands the xspline clade region, defaults to 0.
- spread control the size, when only one point.
- linewidth the width of line of margin, defaults to 0.5.
- linetype the line type of margin, defaults to 1.
- s_shape the shape of the spline relative to the control points, defaults to 0.5.
- s_open whether the spline is a line or a closed shape, defaults to FALSE.

Author(s)

Guangchuang Yu and Shuangbin Xu

References

For more detailed demonstration, please refer to chapter 5.2.2 of *Data Integration, Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

Examples

```
library(ggplot2)
set.seed(102)
tree <- rtree(60)
p <- ggtree(tree)</pre>
p1 <- p + geom_hilight(node=62) + geom_hilight(node=88, fill="red")
dat <- data.frame(id=c(62, 88), type=c("A", "B"))</pre>
p2 <- p + geom_hilight(data=dat, mapping=aes(node=id, fill=type))</pre>
p3 <- p + geom_hilight(data=dat, mapping=aes(node=id, fill=type), align="left")
p4 <- p + geom_hilight(data=dat, mapping=aes(node=id, fill=type), align="right")
p5 <- p + geom_hilight(data=dat, mapping=aes(node=id, fill=type), align="both")
# display the high light layer with gradiental color rectangular.
p6 <- p + geom_hilight(data=dat, mapping=aes(node=id, fill=type), type = "gradient", alpha=0.68)
p7 <- p + geom_hilight(data=dat, mapping=aes(node=id, fill=type), type = "gradient", gradient.direction="tr", alph
# display the high light layer with round rectangular.
p8 <- p + geom_hilight(data=dat, mapping=aes(node=id, fill=type), type = "roundrect", alpha=0.68)
p2/ p3/ p4/ p5 / p6/ p7/ p8
```

geom_inset 23

geom_inset geom_inset

Description

gemo_inset can add subplots to tree by accepting a list of ggplot objects that are ancestral stats or data associated with selected nodes in the tree. These ggplot objects can be any kind of charts or hybrid of of these charts.

add subplots as insets in a tree

Usage

```
geom_inset(
  insets,
  width = 0.1,
  height = 0.1,
  hjust = 0,
  vjust = 0,
  x = "node",
  reverse_x = FALSE,
  reverse_y = FALSE
)
inset(
  tree_view,
  insets,
  width,
  height,
  hjust = 0,
  vjust = 0,
  x = "node",
  reverse_x = FALSE,
  reverse_y = FALSE
)
```

Arguments

insets	a list of ggplot objects, named by node number
width	width of the inset, relative to the range of x-axis, defaults to .1
height	height of the inset, relative to the range of y-axis, defaults to .1
hjust	adjust the horizontal position of the charts, charts will go left if hjust > 0
vjust	adjust the vertical position of the charts, charts will go down if vjust > 0
X	the position where users want to place the charts, one of 'node' (default) and 'branch'

24 geom_label2

reverse_x whether to reverse x axis of the charts by 'ggplot2::scale_x_reverse', defaults to 'FALSE'

reverse_y whether to reverse y axis of the charts by 'ggplot2::scale_y_reverse', defaults to 'FALSE'

tree_view tree view

Details

Users can also use

Value

inset layer tree view with insets

Author(s)

Guangchuang Yu

References

For demonstration of this function, please refer to chapter 8.3 of *Data Integration, Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

geom_label2

geom_label2

Description

geom_label2 support aes(subset) via setup_data

Usage

```
geom_label2(
  mapping = NULL,
  data = NULL,
  ...,
  stat = "identity",
  position = "identity",
  family = "sans",
  parse = FALSE,
  nudge_x = 0,
  nudge_y = 0,
  label.padding = unit(0.25, "lines"),
  label.r = unit(0.15, "lines"),
  label.size = 0.25,
```

geom_label2 25

```
na.rm = TRUE,
show.legend = NA,
inherit.aes = TRUE
)
```

Arguments

mapping Set of aesthetic mappings, defaults to NULL.

data A layer specific dataset - only needed if you want to override the plot defaults.

other arguments passed on to 'layer'.Name of the stat to modify data.

position The position adjustment to use for overlapping points on this layer.

family "sans" by default, can be any supported font.

parse if 'TRUE', the labels will be parsed as expressions, defaults to 'FALSE'.

nudge_x adjust the horizontal position of the labels.

nudge_y adjust the vertical position of the labels.

label.padding Amount of padding around label, defaults to 'unit(0.25, "lines")'.

label.r Use to set the radius of rounded corners of the label, defaults to 'unit(0.15,

"lines")'.

label.size Size of label border, in mm, defaults to 0.25.

na.rm If "FALSE" (default), missing values are removed with a warning. If "TRUE",

missing values are silently removed, logical.

show.legend Whether to show legend, logical, defaults to "NA".

inherit.aes Whether to inherit aesthetic mappings, logical, defaults to "TRUE".

Details

'geom_label2' is a modified version of geom_label, with subset aesthetic supported

Value

label layer

Author(s)

Guangchuang Yu

References

For more detailed demonstration of this function, please refer to chapter A.4.5 of *Data Integration*, *Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

See Also

```
geom_label
```

26 geom_motif

Examples

```
library(ggtree)
set.seed(123)
tr<- rtree(15)
x <- ggtree(tr)
x + geom_label2(aes(label = node, subset = isTip == FALSE))</pre>
```

geom_motif

geom_motif

Description

geom layer to draw aligned motif

Usage

```
geom_motif(mapping, data, on, label, align = "centre", ...)
```

Arguments

mapp	oing	aes mapping
data	1	data
on		gene to center (i.e. set middle position of the on gene to 0)
labe	·1	specify a column to be used to label genes
alig	ŗn	where to place gene label, default is 'centre' and can be set to 'left' and 'right'
		additional parameters

Value

geom layer

Author(s)

geom_nodelab 27

geom_nodelab

geom_nodelab

Description

add node label layer for a tree

Usage

```
geom_nodelab(
  mapping = NULL,
  nudge_x = 0,
  nudge_y = 0,
  geom = "text",
  hjust = 0.5,
  node = "internal",
  ...
)
```

Arguments

aesthetic mappings, defaults to NULL mapping nudge_x horizontal adjustment to nudge labels, defaults to 0 vertical adjustment to nudge labels, defaults to 0 nudge_y one of 'text', "shadowtext", 'label', 'image' and 'phylopic' geom hjust horizontal alignment, defaults to 0.5 node a character indicating which node labels will be displayed, it should be one of 'internal', 'external' and 'all'. If it is set to 'internal' will display internal node labels, 'external' will display the tip labels, and 'all' will display internal node and tip labels. additional parameters, see also the additional parameters of geom_tiplab().

Value

geom layer

Author(s)

Guangchuang Yu

References

For demonstration of this function, please refer to chapter A.4.5 of *Data Integration, Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

28 geom_nodelab2

See Also

```
geom_tiplab()
```

geom_nodelab2

geom_nodelab2

Description

add node label for circular layout

Usage

```
geom_nodelab2(
  mapping = NULL,
  nudge_x = 0,
  nudge_y = 0,
  geom = "text",
  hjust = 0.5,
  ...
)
```

Arguments

```
mapping aesthetic mappings, defaults to NULL

nudge_x horizontal adjustment to nudge labels, defaults to 0

nudge_y vertical adjustment to nudge labels, defaults to 0

geom one of 'text', "shadowtext", 'label', 'image' and 'phylopic'

hjust horizontal alignment, defaults to 0.5

... additional parameters, see also the additional parameters of geom_tiplab().
```

Value

node label layer

Author(s)

geom_nodepoint 29

geom_nodepoint

geom_nodepoint

Description

add node point layer to a tree

Usage

```
geom_nodepoint(
  mapping = NULL,
  data = NULL,
  position = "identity",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE,
  ...
)
```

Arguments

mapping	Set of aesthetic mapping created by aes(). If inherit.aes = TRUE, the mapping can be inherited from the plot mapping as specified in the call to ggplot().
data	The data to be displayed in this layer. If 'NULL' (the default), the data is inherited from the plot data as specified in the call to ggplot().
position	Position adjustment.
na.rm	logical. If 'FALSE' (default), missing values are removed with a warning. If 'TRUE', missing values are silently removed.
show.legend	logical. Should this layer be included in the legends? 'NA', the default, includes if any aesthetics are mapped. 'FALSE' never includes, and 'TRUE' always includes.
inherit.aes	logical (defaults to 'TRUE'). If 'FALSE', overrides the default aesthetics, rather then combining with them.
	additional parameters that passed on to this layer. These are often aesthetics, used to set an aesthetic to a fixed value, like colour = "red" or size = 3.

Value

node point layer

Author(s)

Guangchuang Yu library(ggtree) tr<- rtree(15) x <- ggtree(tr) x + geom_nodepoint()

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References

For more detailed demonstration, please refer to chapter 4.3.2 of *Data Integration, Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

geom_point2

geom_point2

Description

geom_point2 is a modified version of geom_point that supports aes(subset)

Usage

```
geom_point2(
  mapping = NULL,
  data = NULL,
  stat = "identity",
  position = "identity",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE,
  ...
)
```

Arguments

mapping	Set of aesthetic mapping created by aes(). If inherit.aes = TRUE, the mapping can be inherited from the plot mapping as specified in the call to ggplot().
data	The data to be displayed in this layer. If 'NULL' (the default), the data is inherited from the plot data as specified in the call to ggplot().
stat	Name of the statistical transformation to be used on the data for this layer.
position	Position adjustment.
na.rm	logical. If 'FALSE' (default), missing values are removed with a warning. If 'TRUE', missing values are silently removed.
show.legend	logical. Should this layer be included in the legends? 'NA', the default, includes if any aesthetics are mapped. 'FALSE' never includes, and 'TRUE' always includes.
inherit.aes	logical (defaults to 'TRUE'). If 'FALSE', overrides the default aesthetics, rather then combining with them.
	addtional parameters that passed on to this layer. These are often aesthetics, used to set an aesthetic to a fixed value, like colour = "red" or size = 3.

geom_point2 31

Details

geom_point2 creates scatterplots, just similar to ggplot2::geom_point. It extends the ggplot2::geom_point to support filtering via the subset aesthetic mapping (see Aesthetics section).

Value

point layer

Aesthetics

geom_point2() understands the following aesthetics

- subset logical expression indicating elements or rows to keep: missing values are taken as false; should be in aes().
- colour the colour of point, defaults to "black".
- fill the colour of fill, defaults to "black".
- alpha the transparency of fill, defaults to 1.
- size the size of point, defaults to 1.5.
- shape specify a shape, defaults to 19.
- stroke control point border thickness of point, defaults to 0.5.

Author(s)

Guangchuang Yu

References

- 1. G Yu, DK Smith, H Zhu, Y Guan, TTY Lam (2017). ggtree: an R package for visualization and annotation of phylogenetic trees with their covariates and other associated data. Methods in Ecology and Evolution, 8(1):28-36. https://doi.org/10.1111/2041-210X.12628
- 2. G Yu*, TTY Lam, H Zhu, Y Guan*. Two methods for mapping and visualizing associated data on phylogeny using ggtree. Molecular Biology and Evolution, 2018, 35(2):3041-3043. https://doi.org/10.1093/molbev/msy194
- 3. G Yu. Using ggtree to visualize data on tree-like structures. Current Protocols in Bioinformatics, 2020, 69:e96. https://doi.org/10.1002/cpbi.96

For more information about tree visualization, please refer to the online book https://yulab-smu.top/treedata-book/

See Also

geom_point; geom_rootpoint add point of root; geom_nodepoint add points of internal nodes; geom_tippoint add points of external nodes (also known as tips or leaves).

32 geom_range

Examples

```
library(ggtree)
## add point by aes(subset)
tr <- rtree(10)
# group tip and node
ggtree(tr) + geom_point2(aes(shape=isTip, color=isTip), size=3)
# specify a node to display
ggtree(tr) + geom_point2(aes(subset=(node==15)), shape=21, size=5, fill='green')
# specify a tip to display
ggtree(tr) + geom_point2(aes(subset=(label %in% c("t1", "t3"))), shape=21, size=5, fill='green')
## color point with continuous variables
library(ggtree)
library(treeio)
library(ggplot2)
beast_file <- system.file("examples/MCC_FluA_H3.tree", package="ggtree")</pre>
beast_tree <- read.beast(beast_file)</pre>
p <- ggtree(beast_tree) +</pre>
  geom\_tiplab(hjust = -.1)+
  geom_nodepoint(aes(fill = rate), shape = 21, size = 4) +
  scale_fill_continuous(low = 'blue', high = 'red') +
  theme_tree2() + theme(legend.position = 'right')
р
```

geom_range

geom_range

Description

horizontal bar of range (HPD, range etc) on nodes to present uncertainty of evolutionary inference

Usage

```
geom_range(range, center = "auto", ...)
```

Arguments

```
range range(interval) to be displayed, e.g. "height_0.95_HPD"

center center of the range, mean, median or auto (default, the center of the range)

additional parameter, e.g. color, linewidth, alpha
```

Value

ggplot layer

Author(s)

geom_rootedge 33

References

For demonstration of this function, please refer to chapter 5.2.4 of *Data Integration, Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

geom_rootedge

geom_rootedge

Description

display root edge layer for a tree

Usage

```
geom_rootedge(rootedge = NULL, ...)
```

Arguments

rootedge length of rootedge; use phylo\$root.edge if rootedge = NULL (default).
... additional parameters

Additional parameters can be referred to the following parameters:

- size control the width of rootedge, defaults to 0.5.
- colour color of rootedge, defaults to black.
- linetype the type of line, defaults to 1.
- alpha modify colour transparency, defaults to 1.

Details

geom_rootedge is used to create a rootedge as ggtree doesn't plot the root edge by default.

Value

ggtree rootedge layer

Author(s)

Guangchuang Yu

References

1. G Yu, DK Smith, H Zhu, Y Guan, TTY Lam (2017). ggtree: an R package for visualization and annotation of phylogenetic trees with their covariates and other associated data. Methods in Ecology and Evolution, 8(1):28-36. https://doi.org/10.1111/2041-210X.12628

34 geom_rootpoint

Examples

```
library(ggtree)
set.seed(123)
## with root edge = 1
tree1 <- read.tree(text='((A:1,B:2):3,C:2):1;')</pre>
ggtree(tree1) + geom_tiplab() + geom_rootedge()
## without root edge
tree2 <- read.tree(text='((A:1,B:2):3,C:2);')</pre>
ggtree(tree2) + geom_tiplab() + geom_rootedge()
## setting root edge
tree2$root.edge <- 2</pre>
ggtree(tree2) + geom_tiplab() + geom_rootedge()
## specify length of root edge for just plotting
## this will ignore tree$root.edge
ggtree(tree2) + geom_tiplab() + geom_rootedge(rootedge = 3)
## For more detailed demonstration of this function, please refer to chapter A.4.5 of
## *Data Integration, Manipulation and Visualization of Phylogenetic Trees*
## <http://yulab-smu.top/treedata-book/index.html> by Guangchuang Yu.
```

geom_rootpoint

geom_rootpoint

Description

geom_rootpoint is used to add root point layer to a tree

Usage

```
geom_rootpoint(
  mapping = NULL,
  data = NULL,
  position = "identity",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE,
  ...
)
```

Arguments

mapping

Set of aesthetic mapping created by aes(). If inherit.aes = TRUE, the mapping can be inherited from the plot mapping as specified in the call to ggplot().

geom_rootpoint 35

data	The data to be displayed in this layer. If 'NULL' (the default), the data is inherited from the plot data as specified in the call to ggplot().
position	Position adjustment.
na.rm	logical. If 'FALSE' (default), missing values are removed with a warning. If 'TRUE', missing values are silently removed.
show.legend	logical. Should this layer be included in the legends? 'NA', the default, includes if any aesthetics are mapped. 'FALSE' never includes, and 'TRUE' always includes.
inherit.aes	logical (defaults to 'TRUE'). If 'FALSE', overrides the default aesthetics, rather then combining with them.
	addtional parameters that passed on to this layer. These are often aesthetics, used to set an aesthetic to a fixed value, like colour = "red" or size = 3.

Details

geom_rootpoint inherit from geom_point2, and it is used to display and customize the points on the

Value

root point layer

Author(s)

Guangchuang Yu

References

- 1. Guangchuang Yu. Using ggtree to visualize data on tree-like structures. Current Protocols in Bioinformatics. 2020, 69:e96. doi:10.1002/cpbi.96
- 2. Guangchuang Yu, Tommy Tsan-Yuk Lam, Huachen Zhu, Yi Guan. Two methods for mapping and visualizing associated data on phylogeny using ggtree. Molecular Biology and Evolution. 2018, 35(12):3041-3043. doi:10.1093/molbev/msy194
- 3. Guangchuang Yu, David Smith, Huachen Zhu, Yi Guan, Tommy Tsan-Yuk Lam. ggtree: an R package for visualization and annotation of phylogenetic trees with their covariates and other associated data. Methods in Ecology and Evolution. 2017, 8(1):28-36. doi:10.1111/2041-210X.12628

For more information, please refer to the online book:Data Integration, Manipulation and Visualization of Phylogenetic Trees.http://yulab-smu.top/treedata-book/

See Also

geom_point; geom_rootpoint add point of root; geom_nodepoint add points of internal nodes; geom_tippoint add points of external nodes (also known as tips or leaves).

36 geom_segment2

Examples

```
library(ggtree)
tr <- rtree(10)
## add root point
ggtree(tr) + geom_rootpoint()
ggtree(tr) + geom_rootpoint(size=2,color="red",shape=2)</pre>
```

geom_segment2

geom_segment2

Description

geom_segment2 support aes(subset) via setup_data

Usage

```
geom_segment2(
  mapping = NULL,
  data = NULL,
  stat = "identity",
  position = "identity",
  lineend = "butt",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE,
  nudge_x = 0,
  arrow = NULL,
  arrow.fill = NULL,
  ...
)
```

Arguments

mapping	Set of aesthetic mappings, defaults to NULL
data	A layer specific dataset - only needed if you want to override the plot defaults.
stat	Name of stat to modify data.
position	The position adjustment to use for overlapping points on this layer.
lineend	Line end style, one of butt (default), round and square.
na.rm	If "FALSE" (default), missing values are removed with a warning. If "TRUE", missing values are silently removed, logical.
show.legend	Whether to show legend, logical.
inherit.aes	Whether to inherit aesthetic mappings, logical, defaults to "TRUE".
nudge_x	adjust the horizontal position of the segments.
arrow	specification for arrow heads, as created by arrow().
arrow.fill	fill color to usse for the arrow head (if closed). NULL means use colour aesthetic.
	additional parameter

geom_strip 37

Details

'geom_segment2' is a modified version of geom_segment, with subset aesthetic supported

Value

add segment layer

Author(s)

Guangchuang Yu

See Also

geom_segment

geom_strip

geom_strip

Description

annotate associated taxa (from taxa1 to taxa2, can be Monophyletic, Polyphyletic or Paraphyletc Taxa) with bar and (optional) text label

Usage

```
geom_strip(
  taxa1,
  taxa2,
  label,
  offset = 0,
  offset.text = 0,
  align = TRUE,
  barsize = 0.5,
  extend = 0,
  fontsize = 3.88,
  angle = 0,
  geom = "text",
  hjust = 0,
  color = "black",
  fill = NA,
  family = "sans",
  parse = FALSE,
)
```

38 geom_strip

Arguments

taxa1 taxa1 taxa2

label add label alongside the bar (optional)
offset offset of bar and text from the clade

offset.text offset of text from bar

align logical, whether to align bars to the most distant bar ,defaults to "TRUE" Note

that if "FALSE", the bars might cross the tree

barsize set size of the bar

extend extend bar length vertically

fontsize set size of the text
angle set the angle of text
geom one of 'text' or 'label'

hjust adjust the horizonal position of the bar

color set color for bar and label

fill set color to fill label background, only work with geom='label'

family "sans" by default, can be any supported font

parse logical, whether to parse labels, if "TRUE", the labels will be parsed into ex-

pressions, defaults to "FALSE"

... additional parameter

Value

ggplot layers

Author(s)

Guangchuang Yu

References

For more detailed demonstration of this function, please refer to chapter 5.2.1 of *Data Integration*, *Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

Examples

```
library(ggtree)
tr<- rtree(15)
x <- ggtree(tr)
x + geom_strip(13, 1, color = "red") + geom_strip(3, 7, color = "blue")</pre>
```

geom_striplab 39

geom_striplab geom_striplab

Description

annotate associated taxa (from taxa1 to taxa2, can be Monophyletic, Polyphyletic or Paraphyletc Taxa) with bar and (optional) text label or image

Usage

```
geom_striplab(
  taxa1 = NULL,
  taxa2 = NULL,
  label = NULL,
  data = NULL,
  mapping = NULL,
  geom = "text",
  parse = FALSE,
  ...
)
```

Arguments

taxa1	can be label or node number
taxa2	can be label or node number
label	character, character to be showed, when data and mapping is NULL, it is required.
data	data.frame, the data to be displayed in the annotation, default is NULL.
mapping	Set of aesthetic mappings, default is NULL. The detail see the following explanation.
geom	character, one of 'text', 'label', 'shadowtext', 'image' and 'phylopic', default is 'text', and the parameter see the Aesthetics For Specified Geom.
parse	logical, whether parse label to emoji font, default is FALSE.
	additional parameters, see also following section. additional parameters can refer the following parameters. ##'

- offset distance bar and tree, offset of bar and text from the clade, default is 0.
- offset.text distance bar and text, offset of text from bar, default is 0.
- align logical, whether align clade lab, default is FALSE.
- extend numeric, extend the length of bar, default is 0.
- angle numeric or 'auto', if angle is auto, the angle of text will be calculated automatically, which is useful for the circular etc layout, default is 0.
- horizontal logical, whether set label to horizontal, default is TRUE.

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- barsize the width of line, default is 0.5.
- barcolour the colour of line, default is 'black'.
- fontsize the size of text, default is 3.88.
- textcolour the colour of text, default is 'black'.
- imagesize the size of image, default is 0.05.
- imagecolor the colour of image, default is NULL, when geom="phylopic", it should be required.

The parameters also can be set in mapping, when data is provided. Note: the barsize, barcolour, fontsize, textcolour, imagesize and imagecolor should not be set in mapping (aesthetics). When the color and size are not be set in mapping, user can modify them to adjust the attributes of specified geom.

Aesthetics For Specified Geom

geom_striplab() understands the following aesthetics for geom="text"(required aesthetics are in bold):

- taxa1 selected tip label or tip node, it is required.
- taxa2 selected another tip label or tip node, it is required.
- label labels to be shown, it is required.
- colour the colour of text, default is "black".
- size the size of text, default is 3.88.
- angle the angle of text, default is 0.
- hjust A numeric vector specifying horizontal justification, default is 0.
- vjust A numeric vector specifying vertical justification, default is 0.5.
- alpha the transparency of text, default is NA.
- family the family of text, default is 'sans'.
- fontface the font face of text, default is 1 (plain), others are 2 (bold), 3 (italic), 4 (bold.italic).
- lineheight The height of a line as a multiple of the size of text, default is 1.2.

when the colour, size are not be set in mapping, and user want to modify the colour of text, they should use textcolour, fontsize to avoid the confusion with bar layer annotation.

geom_striplab() understands the following aesthethics for geom="label" (required aesthetics are in bold):

- taxa1 selected node to hight light, it is required.
- taxa2 selected another tip label or tip node, it is required.
- label labels to be shown, it is required.
- colour the colour of text, default is "black".
- fill the background colour of the label, default is "white".
- size the size of text, default is 3.88.
- angle the angle of text, default is 0.
- hjust A numeric vector specifying horizontal justification, default is 0.

geom_striplab 41

- vjust A numeric vector specifying vertical justification, default is 0.5.
- alpha the transparency of text, default is NA.
- family the family of text, default is 'sans'.
- fontface the font face of text, default is 1 (plain), others are 2 (bold), 3 (italic), 4 (bold.italic).
- lineheight The height of a line as a multiple of the size of text, default is 1.2.

when the colour, size are not be set in mapping, and user want to modify the colour of text, they should use textcolour, fontsize to avoid the confusion with bar layer annotation.

geom_striplab() understands the following aesthethics for geom="shadowtext" (required aesthetics are in bold):

- taxa1 selected node to hight light, it is required.
- taxa2 selected another tip label or tip node, it is required.
- label labels to be shown, it is required.
- colour the colour of text, default is "black".
- bg.colour the background colour of text, default is 'black'.
- bg.r the width of background text, default is 0.1.
- size the size of text, default is 3.88.
- angle the angle of text, default is 0.
- hjust A numeric vector specifying horizontal justification, default is 0.
- vjust A numeric vector specifying vertical justification, default is 0.5.
- alpha the transparency of text, default is NA.
- family the family of text, default is 'sans'.
- fontface the font face of text, default is 1 (plain), others are 2 (bold), 3 (italic), 4 (bold.italic).
- lineheight The height of a line as a multiple of the size of text, default is 1.2.

when the colour, size are not be set in mapping, and user want to modify the colour of text, they should use textcolour, fontsize to avoid the confusion with bar layer annotation.

geom_striplab() understands the following aesthethics for geom="image" or geom="phylopic" (required aesthetics are in bold):

- taxa1 selected node to hight light, it is required.
- taxa2 selected another tip label or tip node, it is required.
- label labels to be shown, it is required.
- image the image to be annotated, when geom="phylopic", the uid of phylopic databases, it is required.
- colour the color of image, default is NULL.
- size the size of image, default is 0.05.
- alpha the alpha of image, default is 0.8.

when the colour, size are not be set in mapping, and user want to modify the colour of image, they should use imagecolour, imagesize to avoid the confusion with bar layer annotation.

42 geom_taxalink

Examples

geom_taxalink

link between taxa

Description

geom_taxalink supports data.frame as input, the colour, linewidth, linetype and alpha can be mapped. When the data was provided, the mapping should be also provided, which taxal and taxal should be mapped created by aes, aes_ or aes_string. In addition, the hratio, control the height of curve line, when tree layout is cirular, default is 1. ncp, the number of control points used to draw the curve, more control points creates a smoother curve, default is 1. They also can be mapped to a column of data.

Usage

```
geom_taxalink(
  data = NULL,
  mapping = NULL,
  taxa1 = NULL,
  taxa2 = NULL,
  offset = NULL,
  outward = "auto",
  ...
)
```

Arguments

data data.frame, The data to be displayed in this layer, default is NULL.

mapping Set of aesthetic mappings, default is NULL.

taxa1 can be label or node number.

taxa2 can be label or node number.

offset numeric, control the shift of curve line (the ratio of axis value, range is "(0-1)"),

default is NULL.

geom_text2 43

outward

logical, control the orientation of curve when the layout of tree is circular, fan or other layout in polar coordinate, default is "auto", meaning It will automatically. additional parameter.

...,

Value

a list object.

Aesthetics

geom_taxalink() understands the following aesthethics (required aesthetics are in bold):

- taxa1 label or node number of tree.
- taxa2 label or node number of tree.
- group group category of link.
- colour control the color of line, default is black.
- linetype control the type of line, default is 1 (solid).
- linewidth control the width of line, default is 0.5.
- curvature control the curvature of line, default is 0.5, it will be created automatically in polar coordinate .
- hratio control the height of curve line, default is 1.
- ncp control the smooth of curve line, default is 1.

geom_text2

geom_text2

Description

geom_text2 support aes(subset) via setup_data

Usage

```
geom_text2(
  mapping = NULL,
  data = NULL,
  ...,
  stat = "identity",
  position = "identity",
  family = "sans",
  parse = FALSE,
  na.rm = TRUE,
  show.legend = NA,
  inherit.aes = TRUE,
  nudge_x = 0,
  nudge_y = 0,
  check_overlap = FALSE
)
```

geom_tiplab

Arguments

mapping the aesthetic mapping

data A layer specific dataset - only needed if you want to override he plot defaults.

... other arguments passed on to 'layer'

stat Name of stat to modify data

position The position adjustment to use for overlapping points on this layer

family sans by default, can be any supported font

parse if TRUE, the labels will be passd into expressions

na.rm logical show.legend logical inherit.aes logical

nudge_x horizontal adjustment nudge_y vertical adjustment

check_overlap if TRUE, text that overlaps previous text in the same layer will not be plotted

Value

text layer

Author(s)

Guangchuang Yu

See Also

geom_text

geom_tiplab

geom_tiplab

Description

add tip label layer for a tree

Usage

```
geom_tiplab(
  mapping = NULL,
  hjust = 0,
  align = FALSE,
  linetype = "dotted",
  linesize = 0.5,
  geom = "text",
  offset = 0,
  as_ylab = FALSE,
  ...
)
```

geom_tiplab 45

Arguments

mapping aes mapping

hjust horizontal adjustment, defaults to 0

align if TRUE, align all tip labels to the longest tip by adding padding characters to

the left side of tip labels, defaults to "FALSE" with a line connecting each tip

and its corresponding label, defaults to "FALSE"

linetype set linetype of the line if align = TRUE, defaults to "dotted"

linesize set line width if align = TRUE, defaults to 0.5

geom one of 'text', 'label', 'shadowtext', 'image' and 'phylopic'

offset tiplab offset, horizontal adjustment to nudge tip labels, defaults to 0

as_ylab display tip labels as y-axis label, only works for rectangular and dendrogram

layouts, defaults to "FALSE"

... additional parameter

additional parameters can refer the following parameters.

The following parameters for geom="text".

• size control the size of tip labels, defaults to 3.88.

- colour control the colour of tip labels, defaults to "black".
- angle control the angle of tip labels, defaults to 0.
- vjust A numeric vector specifying vertical justification, defaults to 0.5.
- alpha the transparency of text, defaults to NA.
- family the family of text, defaults to 'sans'.
- fontface the font face of text, defaults to 1 (plain), others are 2 (bold), 3 (italic), 4 (bold.italic).
- lineheight The height of a line as a multiple of the size of text, defaults to 1.2.
- nudge_x horizontal adjustment to nudge labels, defaults to 0.
- nudge_y vertical adjustment to nudge labels, defaults to 0.
- check.overlap if TRUE, text that overlaps previous text in the same layer will not be plotted.
- parse if TRUE, the labels will be parsed into expressions, if it is 'emoji', the labels will be parsed into emojifont.

The following parameters for geom="label".

- size the size of tip labels, defaults to 3.88.
- colour the colour of tip labels, defaults to "black".
- fill the colour of rectangular box of labels, defaults to "white".
- vjust numeric vector specifying vertical justification, defaults to 0.5.
- alpha the transparency of labels, defaults to NA.
- family the family of text, defaults to 'sans'.
- fontface the font face of text, defaults to 1 (plain), others are 2 (bold), 3 (italic), 4 (bold.italic).
- lineheight The height of a line as a multiple of the size of text, defaults to 1.2.

46 geom_tiplab

- nudge_x horizontal adjustment to nudge labels, defaults to 0.
- nudge_y vertical adjustment, defaults to 0.
- check.overlap if TRUE, text that overlaps previous text in the same layer will not be plotted.
- parse if TRUE, the labels will be parsed into expressions, if it is 'emoji', the labels will be parsed into emojifont.
- label.padding Amount of padding around label, defaults to 'unit(0.25, "lines")'.
- label.r Radius of rounded corners, defaults to 'unit(0.15, "lines")'.
- label.size Size of label border, in mm, defaults to 0.25.

The following parameters for geom="shadowtext", some parameters are like to geom="text".

- bg.colour the background colour of text, defaults to "black".
- bg.r the width of background of text, defaults to 0.1.

The following parameters for geom="image" or geom="phylopic".

- image the image file path for geom='image', but when geom='phylopic', it should be the uid of phylopic databases.
- size the image size, defaults to 0.05.
- colour the color of image, defaults to NULL.
- alpha the transparency of image, defaults to 0.8.

The following parameters for the line when align = TRUE.

- colour the colour of line, defaults to 'black'.
- alpha the transparency of line, defaults to NA.
- arrow specification for arrow heads, as created by arrow(), defaults to NULL.
- arrow.fill fill color to usse for the arrow head (if closed), defaults to 'NULL', meaning use 'colour' aesthetic.

Details

'geom_tiplab' not only supports using text or label geom to display tip labels, but also supports image geom to label tip with image files or phylopics.

For adding tip labels to a tree with circular layout, 'geom_tiplab' will automatically adjust the angle of the tip labels to the tree by internally calling 'geom_tiplab2'.

Value

tip label layer

Author(s)

Guangchuang Yu

References

For more detailed demonstration, please refer to chapter 4.3.3 of *Data Integration, Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

geom_tiplab2 47

Examples

```
require(ape)
tr <- rtree(10)
ggtree(tr) + geom_tiplab()</pre>
```

geom_tiplab2

geom_tiplab2

Description

add tip label for circular layout

Usage

```
geom_tiplab2(mapping = NULL, hjust = 0, ...)
```

Arguments

mapping aes mapping

hjust horizontal adjustment, defaults to 0
... additional parameter, see geom_tiplab

Details

'geom_tiplab2' will automatically adjust the angle of the tip labels to the tree with circular layout

Value

tip label layer

Author(s)

Guangchuang Yu

References

```
https://groups.google.com/forum/#!topic/bioc-ggtree/o35PV3iHO-0
```

See Also

```
geom_tiplab
```

Examples

```
library(ggtree)
set.seed(123)
tr <- rtree(10)
ggtree(tr, layout = "circular") + geom_tiplab2()</pre>
```

48 geom_tippoint

 ${\tt geom_tippoint}$

geom_tippoint

Description

add tip point layer for a tree

Usage

```
geom_tippoint(
  mapping = NULL,
  data = NULL,
  position = "identity",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE,
  ...
)
```

Arguments

mapping	Set of aesthetic mapping created by aes(). If inherit.aes = TRUE, the mapping can be inherited from the plot mapping as specified in the call to ggplot().
data	The data to be displayed in this layer. If 'NULL' (the default), the data is inherited from the plot data as specified in the call to ggplot().
position	Position adjustment.
na.rm	logical. If 'FALSE' (default), missing values are removed with a warning. If 'TRUE', missing values are silently removed.
show.legend	logical. Should this layer be included in the legends? 'NA', the default, includes if any aesthetics are mapped. 'FALSE' never includes, and 'TRUE' always includes.
inherit.aes	logical (defaults to 'TRUE'). If 'FALSE', overrides the default aesthetics, rather then combining with them.
• • •	additional parameters that passed on to this layer. These are often aesthetics, used to set an aesthetic to a fixed value, like colour = "red" or size = 3.

Value

tip point layer

Author(s)

Guangchuang Yu

geom_tree 49

References

For more detailed demonstration, please refer to chapter 4.3.2 of *Data Integration, Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

Examples

```
library(ggtree)
tr<- rtree(15)
x <- ggtree(tr)
x + geom_tippoint()</pre>
```

geom_tree

geom_tree

Description

```
add tree layer
```

Usage

```
geom_tree(
  mapping = NULL,
  data = NULL,
  layout = "rectangular",
  multiPhylo = FALSE,
  continuous = "none",
  position = "identity",
  ...
)
```

Arguments

```
mapping
                    aesthetic mapping
                    data of the tree
data
layout
                    one of 'rectangular', 'dendrogram', 'slanted', 'ellipse', 'roundrect', 'fan', 'cir-
                    cular', 'inward_circular', 'radial', 'equal_angle', 'daylight' or 'ape'
multiPhylo
                   logical, whether input data contains multiple phylo class, defaults to "FALSE".
continuous
                    character, continuous transition for selected aesthethic ('size' or 'color' ('colour')).
                    It should be one of 'color' (or 'colour'), 'size', 'all' and 'none', default is 'none'
                   Position adjustment, either as a string, or the result of a call to a position adjust-
position
                    ment function, default is "identity".
                    additional parameter
                   some dot arguments:
```

• nsplit integer, the number of branch blocks divided when 'continuous' is not "none", default is 200.

geom_tree2

Value

tree layer

Aesthetics

geom_tree() understands the following aesthethics:

- color character, control the color of line, default is black (continuous is "none").
- linetype control the type of line, default is 1 (solid).
- size numeric, control the width of line, default is 0.5 (continuous is "none").

Author(s)

Yu Guangchuang

References

For demonstration of this function, please refer to chapter 4.2.1 of *Data Integration, Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

Examples

```
tree <- rtree(10)
ggplot(tree) + geom_tree()</pre>
```

geom_tree2

geom_tree2

Description

add tree layer

Usage

```
geom_tree2(layout = "rectangular", ...)
```

Arguments

```
layout one of 'rectangular', 'slanted', 'circular', 'radial' or 'unrooted' additional parameter
```

Value

tree layer

Author(s)

Yu Guangchuang

geom_treescale 51

geom_treescale

geom_treescale

Description

add tree scale to a tree

Usage

```
geom_treescale(
  x = NULL,
  y = NULL,
  width = NULL,
  offset = NULL,
  offset.label = NULL,
  label = NULL,
  color = "black",
  linesize = 0.5,
  fontsize = 3.88,
  family = "sans"
)
```

Arguments

```
set x position of the scale
Χ
                    set y position of the scale
У
                    set the length of the tree scale
width
offset
                    set offset of text to line, defaults to NULL
                    set offset of the scale title to line.
offset.label
                    set the title of tree scale, defaults to NULL.
label
color
                    set color of the scale
linesize
                    set size of line
fontsize
                    set size of text
family
                    'sans' by default, can be any supported font
```

Details

'geom_treescale' automatically adds a scale bar for evolutionary distance

Value

ggplot layers

Author(s)

Guangchuang Yu

52 geom_zoom_clade

References

For demonstration of this function, please refer to chapter 4.3.1 of *Data Integration, Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

geom_zoom_clade

geom_zoom_clade

Description

zoom selected clade of a tree

Usage

```
geom_zoom_clade(node, xexpand = NULL)
```

Arguments

node internal node number to zoom in its corresponding clade

xexpand numeric, extend x, meaning the ratio of range of the xlim of the original tree,

defaults to NULL.

Details

'geom_zoom_clade' zooms in on a selected clade of a tree, while showing its on the full view of tree as a seperated panel for reference

Value

updated tree view

Author(s)

Guangchuang Yu

get.path 53

get.path get.path

Description

path from start node to end node

Usage

```
get.path(phylo, from, to)
```

Arguments

phylo phylo object from start node to end node

Value

node vectot

Author(s)

Guangchuang Yu

getNodeAngle.df

getNodeAngle.df

Description

Get the angle between the two nodes specified.

Usage

```
getNodeAngle.df(df, origin_node_id, node_id)
```

Arguments

```
df tree data.frame
origin_node_id origin node id number
node_id end node id number
```

Value

```
angle in range [-1, 1], i.e. degrees/180, radians/pi
```

54 getSubtree

```
getNodesBreadthFirst.df
```

getNodesBreadthFirst.df

Description

Get the nodes of tree from root in breadth-first order.

Usage

```
getNodesBreadthFirst.df(df)
```

Arguments

df tree data.frame

Value

list of node id's in breadth-first order.

getSubtree

getSubtree

Description

Get all children of node from tree, including start_node.

Usage

```
getSubtree(tree, node)
```

Arguments

tree ape phylo tree object

node is the tree node id from which the tree is derived.

Value

list of all child node id's from starting node.

getSubtree.df 55

getSubtree.df

getSubtree.df

Description

Get all children of node from df tree using breath-first.

Usage

```
getSubtree.df(df, node)
```

Arguments

df tree data.frame node id of starting node.

Value

list of all child node id's from starting node.

 ${\tt getSubtreeUnrooted}$

getSubtreeUnrooted

Description

Get all subtrees of specified node. This includes all ancestors and relatives of node and return named list of subtrees.

Usage

```
getSubtreeUnrooted(tree, node)
```

Arguments

tree ape phylo tree object

node is the tree node id from which the subtrees are derived.

Value

named list of subtrees with the root id of subtree and list of node id's making up subtree.

56 getTreeArcAngles

 ${\tt getSubtreeUnrooted.df} \ \ \textit{getSubtreeUnrooted}$

Description

Get all subtrees of node, as well as remaining branches of parent (ie, rest of tree structure as subtree) return named list of subtrees with list name as starting node id.

Usage

```
getSubtreeUnrooted.df(df, node)
```

Arguments

df tree data.frame

node is the tree node id from which the subtrees are derived.

Value

named list of subtrees with the root id of subtree and list of node id's making up subtree.

getTreeArcAngles getTreeArcAngles

Description

Find the right (clockwise rotation, angle from +ve x-axis to furthest subtree nodes) and left (anticlockwise angle from +ve x-axis to subtree) Returning arc angle in [0, 2] (0 to 360) domain.

Usage

```
getTreeArcAngles(df, origin_id, subtree)
```

Arguments

df tree data.frame

origin_id node id from which to calculate left and right hand angles of subtree.

subtree named list of root id of subtree (node) and list of node ids for given subtree

(subtree).

Value

named list with right and left angles in range [0, 2] i.e 1 = 180 degrees, 1.5 = 270 degrees.

get_clade_position 57

get_clade_position get_clade_position

Description

get position of clade (xmin, xmax, ymin, ymax)

Usage

```
get_clade_position(treeview, node)
```

Arguments

treeview tree view node selected node

Value

data.frame

Author(s)

Guangchuang Yu

```
{\tt get\_heatmap\_column\_position} \\ {\tt get\_heatmap\_column\_position}
```

Description

return a data.frame that contains position information for labeling column names of heatmap produced by gheatmap function

Usage

```
get_heatmap_column_position(treeview, by = "bottom")
```

Arguments

treeview output of gheatmap by one of 'bottom' or 'top'

Value

data.frame

58 get_taxa_name

Author(s)

Guangchuang Yu

get_taxa_name
get_taxa_name

Description

get taxa name of a selected node (or tree if node=NULL) sorted by their position in plotting

Usage

```
get_taxa_name(tree_view = NULL, node = NULL)
```

Arguments

tree_view tree view (i.e. the ggtree object). If tree_view is NULL, the last ggplot object

will be used.

node internal node number to specify a clade. If NULL, using the whole tree

Details

This function extract an ordered vector of the tips from selected clade or the whole tree based on the ggtree() plot.

Value

ordered taxa name vector

Author(s)

Guangchuang Yu

Examples

```
tree <- rtree(30)
p <- ggtree(tree)
get_taxa_name(p)</pre>
```

ggdensitree 59

ggdensitree ggden

Description

drawing phylogenetic trees from list of phylo objects

Usage

```
ggdensitree(
  data = NULL,
  mapping = NULL,
  layout = "slanted",
  tip.order = "mode",
  align.tips = TRUE,
  jitter = 0,
  ...
)
```

Arguments

data	a list of phylo objects or any object with an as.phylo and fortify method
mapping	aesthetic mapping
layout	one of 'slanted', 'rectangluar', 'fan', 'circular' or 'radial' (default: 'slanted')
tip.order	the order of the tips by a character vector of taxa names; or an integer, N, to order the tips by the order of the tips in the Nth tree; 'mode' to order the tips by the most common order; 'mds' to order the tips based on MDS of the path length between the tips; or 'mds_dist' to order the tips based on MDS of the distance between the tips (default: 'mode')
align.tips	TRUE (default) to align trees by their tips and FALSE to align trees by their root
jitter	deviation to jitter tips
	additional parameters passed to fortify, ggtree and geom_tree

Details

The trees plotted by 'ggdensitree()' will be stacked on top of each other and the structures of the trees will be rotated to ensure the consistency of the tip order.

Value

tree layer

Author(s)

Yu Guangchuang, Bradley R. Jones

60 ggtree

References

For more detailed demonstration of this function, please refer to chapter 4.4.2 of *Data Integration*, *Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

Examples

```
require(ape)
require(dplyr)
require(tidyr)
# Plot multiple trees with aligned tips
trees <- list(read.tree(text="((a:1,b:1):1.5,c:2.5);"), read.tree(text="((a:1,c:1):1,b:2);"));
ggdensitree(trees) + geom_tiplab()
# Plot multiple trees with aligned tips with tip labels and separate tree colors
trees.fort <- list(trees[[1]] %>% fortify %>% mutate(tree="a"), trees[[2]] %>% fortify %>% mutate(tree="b"));
ggdensitree(trees.fort, aes(colour=tree)) + geom_tiplab(colour='black')
# Generate example data
set.seed(1)
random.trees <- rmtree(5, 10)</pre>
time.trees <- lapply(seq_along(random.trees), function(i) {</pre>
 tree <- random.trees[[i]]</pre>
 tree$tip.label <- paste0("t", 1:10)</pre>
dates <- estimate.dates(tree, 1:10, mu=1, nsteps=1)</pre>
tree$edge.length <- dates[tree$edge[, 2]] - dates[tree$edge[, 1]]</pre>
fortify(tree) %>% mutate(tree=factor(i, levels=as.character(1:10)))
})
# Plot multiple trees with aligned tips from multiple time points
ggdensitree(time.trees, aes(colour=tree), tip.order=paste0("t", 1:10)) + geom_tiplab(colour='black')
# Read example data
example.trees <- read.tree(system.file("examples", "ggdensitree_example.tree", package="ggtree"))
# Compute OTU
grp <- list(A = c("a.t1", "a.t2", "a.t3", "a.t4"), B = c("b.t1", "b.t2", "b.t3", "b.t4"), C = c("c.t1", "c.t2", "c.t
otu.trees <- lapply(example.trees, groupOTU, grp)</pre>
# Plot multiple trees colored by OTU
ggdensitree(otu.trees, aes(colour=group), alpha=1/6, tip.order='mds') + scale_colour_manual(values=c("black", "r
```

ggtree

visualizing phylogenetic tree and heterogenous associated data based on grammar of graphics ggtree provides functions for visualizing phylogenetic tree and its associated data in R.

ggtree 61

Description

If you use ggtree in published research, please cite: Guangchuang Yu, David Smith, Huachen Zhu, Yi Guan, Tommy Tsan-Yuk Lam. ggtree: an R package for visualization and annotation of phylogenetic trees with their covariates and other associated data. Methods in Ecology and Evolution 2017, 8(1):28-36, doi:10.1111/2041-210X.12628

drawing phylogenetic tree from phylo object

Usage

```
ggtree(
  tr,
 mapping = NULL,
 layout = "rectangular",
 open.angle = 0,
 mrsd = NULL,
 as.Date = FALSE,
 yscale = "none",
 yscale_mapping = NULL,
  ladderize = TRUE,
  right = FALSE,
 branch.length = "branch.length",
  root.position = 0,
  xlim = NULL,
  layout.params = list(),
 hang = 0.1,
)
```

Arguments

tr	phylo object
mapping	aesthetic mapping
layout	one of 'rectangular', 'dendrogram', 'slanted', 'ellipse', 'roundrect', 'fan', 'circular', 'inward_circular', 'radial', 'equal_angle', 'daylight' or 'ape'
open.angle	open angle, only for 'fan' layout
mrsd	most recent sampling date
as.Date	logical whether using Date class in time tree
yscale	y scale
yscale_mapping	yscale mapping for category variable
ladderize	$logical \ (default \ TRUE). \ Should \ the \ tree \ be \ re-organized \ to \ have \ a \ 'ladder' \ aspect?$
right	logical. If ladderize = TRUE, should the ladder have the smallest clade on the right-hand side? See ape::ladderize() for more information.
branch.length	variable for scaling branch, if 'none' draw cladogram
root.position	position of the root node (default = 0)

62 ggtree

xlim x limits, only works for 'inward_circular' layout

layout.params list, the parameters of layout, when layout is a function.

hang numeric The fraction of the tree plot height by which labels should hang below

the rest of the plot. A negative value will cause the labels to hang down from 0. This parameter only work with the 'dendrogram' layout for 'hclust' like class,

default is 0.1.

. . . additional parameter

some dot arguments:

• nsplit integer, the number of branch blocks divided when 'continuous' is not "none", default is 200.

Value

tree

Author(s)

Yu Guangchuang

References

- 1. G Yu, TTY Lam, H Zhu, Y Guan (2018). Two methods for mapping and visualizing associated data on phylogeny using ggtree. Molecular Biology and Evolution, 35(2):3041-3043. https://doi.org/10.1093/molbev/msy194
- 2. G Yu, DK Smith, H Zhu, Y Guan, TTY Lam (2017). ggtree: an R package for visualization and annotation of phylogenetic trees with their covariates and other associated data. Methods in Ecology and Evolution, 8(1):28-36. https://doi.org/10.1111/2041-210X.12628

For more information, please refer to *Data Integration, Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

See Also

```
geom_tree()
```

Examples

```
require(ape)
tr <- rtree(10)
ggtree(tr)</pre>
```

63 gheatmap

Description

append a heatmap of a matrix to the right side of a phylogenetic tree

Usage

```
gheatmap(
 р,
  data,
 offset = 0,
 width = 1,
 low = "green",
 high = "red",
  color = "white",
  colnames = TRUE,
  colnames_position = "bottom",
  colnames_angle = 0,
  colnames_level = NULL,
  colnames_offset_x = 0,
  colnames_offset_y = 0,
  font.size = 4,
  family = "",
  hjust = 0.5,
  legend_title = "value",
  custom_column_labels = NULL
)
```

tree view

Arguments

```
р
data
                  matrix or data.frame
offset
                   set offset of the heatmap to tree
width
                  total width of heatmap, compare to width of tree, defaults to 1, which means
                  they are of the same length
low
                  set color of the lowest value, defaults to "green"
                   set color of the highest value, defaults to "red"
high
color
                   set color of heatmap cell border, defaults to "white"
                  logical, whether to add matrix colnames, defaults to "TRUE"
colnames
colnames\_position
                   set the position of the colnames, one of 'bottom' (default) or 'top'
colnames_angle set the angle of colnames
```

64 gzoom

```
colnames_level set levels of colnames
```

colnames_offset_x

set x offset for colnames

colnames_offset_y

set y offset for colnames

font.size set font size of matrix colnames

family font of matrix colnames, can be any supported font

hjust adjust horizonal position of column names (0: align left, 0.5: align center (de-

fault), 1: align righ)

legend_title title of fill legend

custom_column_labels

instead of using the colnames from the input matrix/data.frame, input a custom

vector to be set as column labels

Value

tree view

Author(s)

Guangchuang Yu

References

For demonstration of this function, please refer to chapter 7.3 of *Data Integration, Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

gzoom

gzoom method

Description

```
gzoom method
gzoom method
zoom selected subtree
```

Usage

```
gzoom(object, focus, subtree = FALSE, widths = c(0.3, 0.7), \ldots)
## S4 method for signature 'ggtree'
gzoom(object, focus, widths = c(0.3, 0.7), xmax_adjust = 0)
## S4 method for signature 'treedata'
```

gzoom.phylo 65

```
gzoom(object, focus, subtree = FALSE, widths = c(0.3, 0.7))
## S4 method for signature 'phylo'
gzoom(object, focus, subtree = FALSE, widths = c(0.3, 0.7))
```

Arguments

object supported tree objects

focus selected tips subtree logical widths widths

... additional parameter
xmax_adjust adjust xmax (xlim[2])

Value

figure

gzoom.phylo gzoom

Description

plots simultaneously a whole phylogenetic tree and a portion of it.

Usage

```
gzoom.phylo(phy, focus, subtree = FALSE, widths = c(0.3, 0.7))
```

Arguments

phy phylo object focus selected tips subtree logical widths widths

Value

a list of ggplot object

Author(s)

ygc

66 hexpand

hexpand hexpand

Description

hexpand

vexpand

expand xlim (ylim) by ratio of x (y) axis range

Usage

```
hexpand(ratio, direction = 1)
vexpand(ratio, direction = 1)
ggexpand(ratio, direction = 1, side = "hv")
```

Arguments

ratio expand x (y) axis limits by amount of xrange (yrange) * ratio

direction expand x axis limit at right hand side if direction is 1 (default), or left hand side

if direction is -1

side one of 'h' for horizontal and 'v' for vertical or 'hv' for both (default).

Value

ggexpand object

Author(s)

Guangchuang Yu

Examples

```
x <- rtree(20)
x$tip.label <- paste0('RRRRREEEEEAAAAALLLLLYYYYYY_Long_Lable_', x$tip.label)
p1 <- ggtree(x) + geom_tiplab()
p1 + ggexpand(1.5, side = "h")</pre>
```

label_pad 67

label_pad	Padding taxa labels

Description

This function adds padding characters to the left side of taxa labels, adjust their length to the longest label.

Usage

```
label_pad(label, justify = "right", pad = ".")
```

Arguments

label taxa label

justify should a character vector be right-justified (default), left-justified, centred or left

alone.

pad padding character (defaults to dots)

Value

Taxa labels with padding characters added

Author(s)

Guangchuang Yu and Yonghe Xia

References

```
https://groups.google.com/g/bioc-ggtree/c/INJ0Nfkq3b0/m/lXefnfV5AQAJ
```

Examples

```
library(ggtree)
set.seed(2015-12-21)
tree <- rtree(5)
tree$tip.label[2] <- "long string for test"
label_pad(tree$tip.label)</pre>
```

68 layoutEqualAngle

layoutDaylight

Equal daylight layout method for unrooted trees.

Description

#' @title

Usage

```
layoutDaylight(model, branch.length, MAX_COUNT = 5)
```

Arguments

model tree object, e.g. phylo or treedata

branch.length set to 'none' for edge length of 1. Otherwise the phylogenetic tree edge length

is used.

MAX_COUNT the maximum number of iterations to run (default 5)

Value

tree as data.frame with equal angle layout.

References

The following aglorithm aims to implement the vague description of the "Equal-daylight Algorithm" in "Inferring Phylogenies" pp 582-584 by Joseph Felsenstein.

```
Leafs are subtrees with no children
Initialise tree using equal angle algorithm
tree_df = equal_angle(tree)

nodes = get list of nodes in tree_df breadth-first
nodes = remove tip nodes.
```

layoutEqualAngle

layoutEqualAngle

Description

'Equal-angle layout algorithm for unrooted trees'

Usage

```
layoutEqualAngle(model, branch.length = "branch.length")
```

layout_rectangular 69

Arguments

model tree object, e.g. phylo or treedata

branch.length set to 'none' for edge length of 1. Otherwise the phylogenetic tree edge length

is used.

Value

tree as data.frame with equal angle layout.

References

"Inferring Phylogenies" by Joseph Felsenstein.

layout_rectangular layout_rectangular

Description

transform circular/fan layout to rectangular layout transform rectangular layout to circular layout transform rectangular/circular layout to inward circular layout transform rectangular/circular layout to fan layout transform rectangular layout to dendrogram layout

Usage

```
layout_rectangular()
layout_circular()
layout_inward_circular(xlim = NULL)
layout_fan(angle = 180)
layout_dendrogram()
```

Arguments

xlim setting x limits, which will affect the center space of the tree

angle open tree at specific angle

Author(s)

Guangchuang Yu

70 msaplot

Examples

```
tree <- rtree(20)
p <- ggtree(tree, layout = "circular") + layout_rectangular()
tree <- rtree(20)
p <- ggtree(tree)
p + layout_circular()
tree <- rtree(20)
p <- ggtree(tree)
p + layout_inward_circular(xlim=4) + geom_tiplab(hjust=1)
tree <- rtree(20)
p <- ggtree(tree)
p + layout_fan(angle=90)
tree <- rtree(20)
p <- ggtree(tree)
p + p + layout_dendrogram()</pre>
```

msaplot

msaplot

Description

visualize phylogenetic tree with multiple sequence alignment

Usage

```
msaplot(
  p,
  fasta,
  offset = 0,
  width = 1,
  color = NULL,
  window = NULL,
  bg_line = TRUE,
  height = 0.8
)
```

Arguments

р	tree view
fasta	fasta file that contains multiple sequence alignment information
offset	set the offset of MSA to tree
width	total width of alignment, compare to width of tree, defaults to 1 , which means they are of the same length
color	set color of the tree
window	specific a slice of alignment to display
bg_line	whether to add background line in alignment, defaults to "TRUE"
height	height ratio of sequence, defaults to 0.8

multiplot 71

Value

tree view

Author(s)

Guangchuang Yu

References

For demonstration of this function, please refer to chapter 7.4 of *Data Integration, Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

multiplot

multiplot

Description

plot multiple ggplot objects in one page

Usage

```
multiplot(
    ...,
    plotlist = NULL,
    ncol,
    widths = rep_len(1, ncol),
    labels = NULL,
    label_size = 5
)
```

Arguments

```
    plots
    plot list
    set the number of column to display the plots
    widths
    the width of each plot
```

labels set labels for labeling the plots label_size set font size of the label

Value

plot

Author(s)

Guangchuang Yu

72 nodepie

nodebar nodebar

Description

generate a list of bar charts for results of ancestral state reconstruction

Usage

```
nodebar(data, cols, color, alpha = 1, position = "stack")
```

Arguments

data a data.frame of stats with an additional column of node number named "node" cols columns of the data.frame that store the stats

color set color of bars

alpha set transparency of the charts

position position of bars, if 'stack' (default) make bars stacked atop one another, 'dodge'

make them dodged side-to-side

Value

list of ggplot objects

Author(s)

Guangchuang Yu

nodepie nodepie

Description

generate a list of pie charts for results of ancestral stat reconstruction

Usage

```
nodepie(
  data,
  cols,
  color,
  alpha = 1,
  outline.color = "transparent",
  outline.size = 0
)
```

open_tree 73

Arguments

data a data.frame of stats with an additional column of node number named "node"

cols columns of the data.frame that store the stats

color set color of bars

alpha set transparency of the charts

outline.color color of outline
outline.size size of outline

Value

list of ggplot objects

Author(s)

Guangchuang Yu

open_tree open_tree

Description

transform a tree in either rectangular or circular layout into the fan layout that opens with a specific angle

Usage

```
open_tree(treeview, angle)
```

Arguments

treeview tree view in rectangular/circular layout angle open the tree at a specific angle

Value

updated tree view

Author(s)

Guangchuang Yu

```
tree <- rtree(15)
p <- ggtree(tree) + geom_tiplab()
open_tree(p, 180)</pre>
```

74 reexports

range_format

range_format

Description

```
format a list of range (HPD, CI, etc that has length of 2)
```

Usage

```
range_format(x, trans = NULL)
```

Arguments

x input list

trans transformation function

Value

```
character vector of [lower, upper]
```

Author(s)

Guangchuang Yu

reexports

Objects exported from other packages

Description

These objects are imported from other packages. Follow the links below to see their documentation.

revts 75

revts

revts

Description

reverse timescle x-axis by setting the most recent tip to 0

Usage

```
revts(treeview)
```

Arguments

treeview

original tree view

Details

'scale_x_continuous(labels=abs)' is required if users want to set the x-axis lable to absolute value

Value

updated tree view

Author(s)

Guangchuang Yu

Examples

```
tr <- rtree(10)
p <- ggtree(tr) + theme_tree2()
p2 <- revts(p)
p2 + scale_x_continuous(labels=abs)</pre>
```

rotate

rotate

Description

rotate selected clade by 180 degree

Usage

```
rotate(tree_view = NULL, node)
```

76 rotateTreePoints.df

Arguments

tree_view tree view (i.e. the ggtree object). If tree_view is NULL, the last ggplot object

will be used.

node internal node number to specify a clade. If NULL, using the whole tree

Value

ggplot2 object

Author(s)

Guangchuang Yu

Examples

```
x <- rtree(15)
p <- ggtree(x) + geom_tiplab()
rotate(p, 17)</pre>
```

rotateTreePoints.df

rotate Tree Points. data. frame

Description

Rotate the points in a tree data.frame around a pivot node by the angle specified.

Usage

```
rotateTreePoints.df(df, pivot_node, nodes, angle)
```

Arguments

df tree data.frame

pivot_node is the id of the pivot node.

nodes list of node numbers that are to be rotated by angle around the pivot_node

angle in range [0, 2], ie degrees/180, radians/pi

Value

updated tree data.frame with points rotated by angle

rotate_tree 77

rotate_tree

rotate_tree

Description

rotate circular tree in a certain angle

Usage

```
rotate_tree(treeview, angle)
```

Arguments

treeview tree view in circular layout

angle the angle of rotation

Value

updated tree view

Author(s)

Guangchuang Yu

Examples

```
tree <- rtree(15)
p <- ggtree(tree) + geom_tiplab()
p2 <- open_tree(p, 180)
rotate_tree(p2, 180)</pre>
```

scaleClade

scaleClade

Description

zoom out/in a selected clade to emphasize or de-emphasize it

Usage

```
scaleClade(tree_view = NULL, node, scale = 1, vertical_only = TRUE)
```

78 scale_color

Arguments

tree_view tree view (i.e. the ggtree object). If tree_view is NULL, the last ggplot object

will be used.

node internal node number to specify a clade. If NULL, using the whole tree

scale the scale of the selected clade. The clade will be zoom in when scale > 1, and

will be zoom out when scale < 1

vertical_only logical. If TRUE (default), only vertical will be scaled. If FALSE, the clade will

be scaled vertical and horizontally.

Value

tree view

Author(s)

Guangchuang Yu

Examples

```
x <- rtree(15)
p <- ggtree(x) + geom_tiplab() +
  geom_nodelab(aes(subset=!isTip, label=node), hjust = -.1, color = "red")
scaleClade(p, 24, scale = .1)</pre>
```

scale_color

scale_color method

Description

```
scale_color method
scale color by a numerical tree attribute
```

Usage

```
scale_color(object, by, ...)
## S4 method for signature 'treedata'
scale_color(object, by, ...)
## S4 method for signature 'phylo'
scale_color(object, by, ...)
```

Arguments

object treedata object

by one of numerical attributes
... additional parameter

scale_color_subtree 79

Value

color vector

scale_color_subtree

 $scale_color_subtree$

Description

scale tree color by subtree (e.g., output of cutree, kmeans, or other clustering algorithm)

Usage

```
scale_color_subtree(group)
scale_colour_subtree(group)
```

Arguments

group

taxa group information

Value

updated tree view

Author(s)

Guangchuang Yu

scale_x_ggtree

scale_x_ggtree

Description

scale x for tree with gheatmap

Usage

```
scale_x_ggtree(breaks = waiver(), labels = waiver())
```

Arguments

breaks set breaks for tree

labels lables for corresponding breaks

scale_x_range

Details

Since setting x-axis for tree with gheatmap by using 'theme_tree2()' is quite tricky, 'scale_x_ggtree' can help set the x-axis more reasonably.

Value

updated tree view

Author(s)

Guangchuang Yu

References

For more detailed demonstration of this function, please refer to chapter 7.3 of *Data Integration*, *Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

scale_x_range

scale_x_range

Description

add second x-axis for geom_range

Usage

```
scale_x_range()
```

Details

notice that the first axis is disabled in the default theme thus users need to enable it first before using scale_x_range

Value

ggtree object

Author(s)

Guangchuang Yu

References

For demonstration of this function ,please refer to chapter 5.2.4 of *Data Integration*, *Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

set_hilight_legend 81

set_hilight_legend s	et_hilight_legend
----------------------	-------------------

Description

set legend for multiple geom_hilight layers

Usage

```
set_hilight_legend(p, color, label, alpha = 1)
```

Arguments

p ggtree objectcolor color vectorlabel label vector

alpha transparency of color

Value

updated ggtree object

Author(s)

Guangchuang Yu

StatBalance StatBalance

Description

StatBalance

82 td_filter

td_filter

td-filter

Description

filter data for tree annotation layer

Usage

```
td_filter(..., .f = NULL)
```

Arguments

... Expressions that return a logical value.

. f a function (if any, defaults to NULL) that pre-operate the data

Details

The 'td_filter()' function returns another function that can be used to subset ggtree() plot data. The function can be passed to the 'data' parameter of geom layer to perform subsetting. All rows that satisy your conditions will be retained.

Value

A function to filter ggtree plot data using conditions defined by '...'.

Author(s)

Guangchuang Yu

References

For more detailed demonstration of this function, please refer to chapter 12.5.1 of *Data Integration*, *Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

See Also

filter

```
tree <- rtree(30)
## similar to 'ggtree(tree) + geom_tippoint()'
ggtree(tree) + geom_point(data = td_filter(isTip))</pre>
```

td_mutate 83

td_mutate

td-mutate

Description

mutate data for tree annotation layer

Usage

```
td_mutate(..., .f = NULL)
```

Arguments

. . . additional parameters that pass to dplyr::mutate

.f a function (if any, defaults to NULL) that pre-operate the data

Details

The 'td_mutate()' function returns another function that can be used to mutate ggtree() plot data. The function can be passed to the 'data' parameter of geom layer to perform adding new variables and preserving existing ones.

Value

A function to mutate ggtree plot data

See Also

mutate

td_unnest

td-unnest

Description

flatterns a list-column of data frame

Usage

```
td_unnest(cols, ..., .f = NULL)
```

Arguments

cols columns to unnest

... additional parameters that pass to tidyr::unnest

.f a function (if any, defaults to NULL) that pre-operate the data

84 theme_dendrogram

Details

The 'td_unnest' function returns another function that can be used to unnest ggtree() plot data. The function can be passed to the 'data' parameter of a geom layer to flattern list-cloumn tree data.

Value

A function to unnest ggtree plot data

Author(s)

Guangchuang Yu

References

For demonstration of this function, please refer to chapter 12.5.2 of *Data Integration, Manipulation and Visualization of Phylogenetic Trees* http://yulab-smu.top/treedata-book/index.html by Guangchuang Yu.

See Also

unnest

theme_dendrogram

 $theme_dendrogram$

Description

dendrogram theme

Usage

```
theme_dendrogram(bgcolor = "white", fgcolor = "black", ...)
```

Arguments

bgcolor set background color, defaults to "white"

fgcolor set color of axis

... additional parameter

Author(s)

Guangchuang Yu

theme_inset 85

 ${\tt theme_inset}$

theme_inset

Description

inset theme

Usage

```
theme_inset(legend.position = "none", ...)
```

Arguments

```
legend.position
```

set the position of legend

.. additional parameter

Details

theme for inset function

Value

ggplot object

Author(s)

Guangchuang Yu

theme_tree

 $theme_tree$

Description

tree theme

Usage

```
theme_tree(bgcolor = "white", ...)
```

Arguments

```
bgcolor set background color, defaults to "white"
```

... additional parameter

theme_tree2

Details

'theme_tree' defines a blank background to display tree

Value

updated ggplot object with new theme

Author(s)

Guangchuang Yu

Examples

```
require(ape)
tr <- rtree(10)
ggtree(tr) + theme_tree()</pre>
```

theme_tree2

 $theme_tree2$

Description

tree2 theme

Usage

```
theme_tree2(bgcolor = "white", fgcolor = "black", ...)
```

Arguments

bgcolor set background color, defaults to "white" set foreground color, defaults to "black"

... additional parameter

Details

'theme_tree2' supports displaying phylogenetic distance by setting x-axis

Value

updated ggplot object with new theme

Author(s)

Guangchuang Yu

viewClade 87

Examples

```
require(ape)
tr <- rtree(10)
ggtree(tr) + theme_tree2()</pre>
```

viewClade

viewClade

Description

view a selected clade of tree, clade can be selected by specifying a node number or determined by the most recent common ancestor of selected tips

Usage

```
viewClade(tree_view = NULL, node, xmax_adjust = 0)
```

Arguments

tree_view tree view (i.e. the ggtree object). If tree_view is NULL, the last ggplot object

will be used.

node internal node number to specify a clade. If NULL, using the whole tree

xmax_adjust adjust the max range of x axis

Value

clade plot

Author(s)

Guangchuang Yu

```
x <- rtree(15)
p <- ggtree(x) + geom_tiplab()
viewClade(p, 18, xmax_adjust = 0.)</pre>
```

88 xlim_tree

xlim_expand

xlim_expand

Description

expand x axis limits for specific panel

Usage

```
xlim_expand(xlim, panel)
```

Arguments

xlim x axis limits

panel name of the panel to expand

Value

updated tree view

Author(s)

Guangchuang Yu

Examples

xlim_tree

xlim_tree

Description

set x axis limits specially for Tree panel

Usage

```
xlim_tree(xlim)
```

zoomClade 89

Arguments

xlim x axis limits

Value

updated tree view

Author(s)

Guangchuang Yu

Examples

zoomClade

zoomClade

Description

zoom in on a selected clade of a tree, while showing its on the full view of tree as a seperated panel for reference

Usage

```
zoomClade(tree_view = NULL, node, xexpand = NULL)
```

Arguments

tree_view tree view (i.e. the ggtree object). If tree_view is NULL, the last ggplot object

will be used.

node internal node number to specify a clade. If NULL, using the whole tree

numeric, expend the xlim of the zoom area. default is NULL.

Value

full tree with zoom in clade

Author(s)

Guangchuang Yu

90

Examples

```
## Not run:
x <- rtree(15)
p <- ggtree(x) + geom_tiplab() +
  geom_nodelab(aes(subset=!isTip, label=node), hjust = -.1, color = "red")
zoomClade(p, 21, xexpand = .2)
## End(Not run)</pre>
```

%+>%

%+>%

Description

update data with tree info (y coordination and panel)

Usage

```
p %+>% data
```

Arguments

p tree view data data.frame

Details

add tree information to an input data. This function will setup y coordination and panel info for data used in facet_plot and geom_faceet

Value

updated data.frame

Author(s)

Guangchuang Yu

References

G Yu, TTY Lam, H Zhu, Y Guan (2018). Two methods for mapping and visualizing associated data on phylogeny using ggtree. Molecular Biology and Evolution, 35(2):3041-3043. https://doi.org/10.1093/molbev/msy194

%<+%

%<+% %<+%

Description

add annotation data to a tree

Usage

```
pg %<+% data
```

Arguments

pg ggplot2 object

data annotation data that contains a column of "node", or the first column of taxa

labels

Details

This operator attaches annotation data to a ggtree graphic object

Value

ggplot object with annotation data added

Author(s)

Guangchuang Yu

References

G Yu, TTY Lam, H Zhu, Y Guan (2018). Two methods for mapping and visualizing associated data on phylogeny using ggtree. Molecular Biology and Evolution, 35(2):3041-3043. https://doi.org/10.1093/molbev/msy194

See Also

```
geom_facet
```

92

%<%

Description

update tree

Usage

```
pg %<% x
```

Arguments

pg ggtree object x tree object

Details

This operator apply the visualization directives in ggtree object (lhs) to visualize another tree object (rhs), that is similar to Format Painter.

Value

updated ggplot object

Author(s)

Guangchuang Yu

```
library("ggplot2")
nwk <- system.file("extdata", "sample.nwk", package="treeio")
tree <- read.tree(nwk)
p <- ggtree(tree) + geom_tippoint(color="#b5e521", alpha=1/4, size=10)
p %<% rtree(30)</pre>
```

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