

Package ‘seqcombo’

October 17, 2020

Title Visualization Tool for Sequence Recombination and Reassortment

Version 1.10.0

Description Provides useful functions for visualizing sequence recombination and virus reassortment events.

Depends R (>= 3.4.0)

Imports Biostrings, cowplot, dplyr, ggplot2, grid, igraph, magrittr, methods, rvcheck, utils

Suggests emojifont, knitr, prettydoc, tibble

VignetteBuilder knitr

ByteCompile true

License Artistic-2.0

Encoding UTF-8

LazyData true

BugReports <https://github.com/GuangchuangYu/seqcombo/issues>

biocViews Alignment, Software, Visualization

RoxygenNote 6.1.1

git_url <https://git.bioconductor.org/packages/seqcombo>

git_branch RELEASE_3_11

git_last_commit 5c29223

git_last_commit_date 2020-04-27

Date/Publication 2020-10-16

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| | |
|---------------|----------------------|
| geom_genotype | <i>geom_genotype</i> |
|---------------|----------------------|

Description

geom layer of genotype

Usage

```
geom_genotype(virus_info, v_color = "darkgreen", v_fill = "steelblue",
  v_shape = "ellipse", l_color = "black", asp = 1, g_height = 0.65,
  g_width = 0.65)
```

Arguments

| | |
|------------|---|
| virus_info | virus information |
| v_color | the color of outer boundary of virus; can use expression (e.g. v_color=~Host) to color virus by specific variable |
| v_fill | the color to fill viruses; can use expression (e.g. v_fill=~Host) to fill virus by specific variable |
| v_shape | one of 'hexagon' or 'ellipse' |
| l_color | color of the lines that indicate genetic flow |
| asp | aspect ratio of the plotting device |
| g_height | height of regions to plot gene segments relative to the virus |
| g_width | width of gene segment relative to width of the virus (the hexagon) |

Value

geom layer

Author(s)

guangchuang yu

Examples

```
library(tibble)
library(ggplot2)
n <- 8
virus_info <- tibble(id = 1:7,
  x = c(rep(1990, 4), rep(2000, 2), 2009),
  y = c(1,2,3,5, 1.5, 3, 4),
  segment_color = list(rep('purple', n),
    rep('red', n), rep('darkgreen', n), rep('lightgreen', n),
    c('darkgreen', 'darkgreen', 'red', 'darkgreen', 'red', 'purple', 'red', 'purple'),
    c('darkgreen', 'darkgreen', 'red', 'darkgreen', 'darkgreen', 'purple', 'red', 'purple'),
    c('darkgreen', 'lightgreen', 'lightgreen', 'darkgreen', 'darkgreen', 'purple', 'red', 'purple')))
ggplot() + geom_genotype(virus_info)
```

| | |
|-------------|--------------------|
| geom_hybrid | <i>geom_hybrid</i> |
|-------------|--------------------|

Description

geom layer for reassortment events

Usage

```
geom_hybrid(virus_info, flow_info, v_color = "darkgreen",
             v_fill = "steelblue", v_shape = "ellipse", l_color = "black",
             asp = 1, parse = FALSE, g_height = 0.65, g_width = 0.65,
             t_size = 3.88, t_color = "black")
```

Arguments

| | |
|------------|---|
| virus_info | virus information |
| flow_info | flow information |
| v_color | the color of outer boundary of virus; can use expression (e.g. v_color=~Host) to color virus by specific variable |
| v_fill | the color to fill viruses; can use expression (e.g. v_fill=~Host) to fill virus by specific variable |
| v_shape | one of 'hexagon' or 'ellipse' |
| l_color | color of the lines that indicate genetic flow |
| asp | aspect ratio of the plotting device |
| parse | whether parse label, only works if 'label' and 'label_position' exist |
| g_height | height of regions to plot gene segments relative to the virus |
| g_width | width of gene segment relative to width of the virus (the hexagon) |
| t_size | size of text label |
| t_color | color of text label |

Value

geom layer

Author(s)

Guangchuang Yu

Examples

```
library(tibble)
library(ggplot2)
n <- 8
virus_info <- tibble(id = 1:7,
x = c(rep(1990, 4), rep(2000, 2), 2009),
y = c(1,2,3,5, 1.5, 3, 4),
segment_color = list(rep('purple', n),
rep('red', n), rep('darkgreen', n), rep('lightgreen', n),
```

```
c('darkgreen', 'darkgreen', 'red', 'darkgreen', 'red', 'purple', 'red', 'purple'),
c('darkgreen', 'darkgreen', 'red', 'darkgreen', 'darkgreen', 'purple', 'red', 'purple'),
c('darkgreen', 'lightgreen', 'lightgreen', 'darkgreen', 'darkgreen', 'purple', 'red', 'purple')))

flow_info <- tibble(from = c(1,2,3,3,4,5,6), to = c(5,5,5,6,7,6,7))

ggplot() + geom_hybrid(virus_info, flow_info)
```

hybrid_plot*hyrid_plot***Description**

visualize virus reassortment events

Usage

```
hybrid_plot(virus_info, flow_info, v_color = "darkgreen",
            v_fill = "steelblue", v_shape = "ellipse", l_color = "black",
            asp = 1, parse = FALSE, g_height = 0.65, g_width = 0.65,
            t_size = 3.88, t_color = "black")
```

Arguments

| | |
|-------------------------|---|
| <code>virus_info</code> | virus information |
| <code>flow_info</code> | flow information |
| <code>v_color</code> | the color of outer boundary of virus; can use expression (e.g. <code>v_color=~Host</code>) to color virus by specific variable |
| <code>v_fill</code> | the color to fill viruses; can use expression (e.g. <code>v_fill=~Host</code>) to fill virus by specific variable |
| <code>v_shape</code> | one of 'hexagon' or 'ellipse' |
| <code>l_color</code> | color of the lines that indicate genetic flow |
| <code>asp</code> | aspect ratio of the plotting device |
| <code>parse</code> | whether parse label, only works if 'label' and 'label_position' exist |
| <code>g_height</code> | height of regions to plot gene segments relative to the virus |
| <code>g_width</code> | width of gene segment relative to width of the virus (the hexagon) |
| <code>t_size</code> | size of text label |
| <code>t_color</code> | color of text label |

Value

ggplot object

Author(s)

guangchuang yu

Examples

```
library(tibble)
n <- 8
virus_info <- tibble(id = 1:7,
x = c(rep(1990, 4), rep(2000, 2), 2009),
y = c(1,2,3,5, 1.5, 3, 4),
segment_color = list(rep('purple', n),
rep('red', n), rep('darkgreen', n), rep('lightgreen', n),
c('darkgreen', 'darkgreen', 'red', 'darkgreen', 'red', 'purple', 'red', 'purple'),
c('darkgreen', 'darkgreen', 'red', 'darkgreen', 'darkgreen', 'purple', 'red', 'purple'),
c('darkgreen', 'lightgreen', 'lightgreen', 'darkgreen', 'darkgreen', 'purple', 'red', 'purple')))

flow_info <- tibble(from = c(1,2,3,3,4,5,6), to = c(5,5,5,6,7,6,7))

hybrid_plot(virus_info, flow_info)
```

plot

plot method for SeqDiff object

Description

plot method for SeqDiff object

Usage

```
## S4 method for signature 'SeqDiff,ANY'
plot(x, width = 50, title = "auto",
      xlab = "Nucleotide Position", by = "bar", fill = "firebrick",
      colors = c(A = "#E495A5", C = "#ABB065", G = "#39BEB1", T = "#ACA4E2"),
      xlim = NULL)
```

Arguments

| | |
|--------|--------------------------------------|
| x | SeqDiff object |
| width | bin width |
| title | plot title |
| xlab | xlab |
| by | one of 'bar' and 'area' |
| fill | fill color of upper part of the plot |
| colors | color of lower part of the plot |
| xlim | limits of x-axis |

Value

plot

Author(s)

guangchuang yu

Examples

```
fas <- list.files(system.file("examples", "GVariation", package="seqcombo"), pattern="fas", full.names=TRUE)
x1 <- seqdiff(fas[1], reference=1)
plot(x1)
```

seqdiff

*seqdiff***Description**

calculate difference of two aligned sequences

Usage

```
seqdiff(fasta, reference = 1)
```

Arguments

| | |
|-----------|---|
| fasta | fasta file |
| reference | which sequence serve as reference, 1 or 2 |

Value

SeqDiff object

Author(s)

guangchuang yu

Examples

```
fas <- list.files(system.file("examples", "GVariation", package="seqcombo"), pattern="fas", full.names=TRUE)
seqdiff(fas[1], reference=1)
```

set_layout

*set_layout***Description**

set layout for reassortment plot

Usage

```
set_layout(virus_info, flow_info, layout = "layout.auto")
```

Arguments

| | |
|------------|-------------------|
| virus_info | virus information |
| flow_info | flow information |
| layout | layout method |

show

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Value

updated virus_info

Author(s)

guangchuang yu

show

show method

Description

show method

Usage

show(object)

Arguments

object SeqDiff object

Value

message

Examples

```
fas <- list.files(system.file("examples", "GVariation", package="seqcombo"), pattern="fas", full.names=TRUE)
x1 <- seqdiff(fas[1], reference=1)
x1
```

simplot

simplot

Description

Sequence similarity plot

Usage

```
simplot(file, query, window = 200, step = 20, group = FALSE, id, sep,
sd = FALSE)
```

Arguments

| | |
|--------|---|
| file | alignment fast file |
| query | query sequence |
| window | sliding window size (bp) |
| step | step size to slide the window (bp) |
| group | whether grouping sequence |
| id | position to extract id for grouping; only works if group = TRUE |
| sep | separator to split sequence name; only works if group = TRUE |
| sd | whether display standard deviation of similarity among each group; only works if group=TRUE |

Value

ggplot object

Author(s)

guangchuang yu

Examples

```
fas <- system.file("examples/GVariation/sample_alignment.fa", package="seqcombo")
simplot(fas, 'CF_YL21')
```

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