Package ‘plotgardener’

May 6, 2024

Title Coordinate-Based Genomic Visualization Package for R

Version 1.10.0

Date 2024-04-12

Description Coordinate-based genomic visualization package for R. It grants users
the ability to programmatically produce complex, multi-paneled figures. Tailored for
genomics, plotgardener allows users to visualize large complex genomic datasets and
provides exquisite control over how plots are placed and arranged on a page.

Depends R (>= 4.1.0)

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URL https://phanstiellab.github.io/plotgardener,
     https://github.com/PhanstielLab/plotgardener

Encoding UTF-8

RoxygenNote 7.2.3

Imports curl, data.table, dplyr, GenomeInfoDb, GenomicRanges,
grDevices, grid, ggplotify, IRanges, methods, plyranges, purrr,
Rcpp, RColorBrewer, rlang, stats, strawr, tools, utils, withr

biocViews Visualization, GenomeAnnotation, FunctionalGenomics,
   GenomeAssembly, HiC

LinkingTo Rcpp

Suggests AnnotationDbi, AnnotationHub, BSgenome,
   BSgenome.Hsapiens.UCSC.hg19, ComplexHeatmap, GenomicFeatures,
ggplot2, InteractionSet, knitr, org.Hs.eg.db, rtracklayer,
   plotgardenerData, pdftools, png, rmarkdown, scales, showtext,
   testthat (>= 3.0.0), TxDb.Hsapiens.UCSC.hg19.knownGene

VignetteBuilder knitr

Config/testthat/edition 3

BugReports https://github.com/PhanstielLab/plotgardener/issues

git_url https://git.bioconductor.org/packages/plotgardener

git_branch RELEASE_3_19

git_last_commit 5955e9a
Contents

annoDomains ................................................................. 3
annoGenomeLabel .......................................................... 5
annoHeatmapLegend .......................................................... 8
annoHighlight ................................................................. 10
annoPixels ................................................................. 12
annoSegments ............................................................... 14
annoText ................................................................. 17
annoXaxis ................................................................. 18
annoYaxis ................................................................. 20
annoZoomLines ............................................................. 22
assembly ................................................................. 24
c ................................................................. 25
calcSignalRange ............................................................. 26
colorby ................................................................. 27
defaultPackages .............................................................. 28
genomes ................................................................. 29
mapColors ................................................................. 29
pageCreate ............................................................... 31
pageGuideHide ............................................................. 32
pageGuideHorizontal .......................................................... 33
pageGuideShow ............................................................. 34
pageGuideVertical ............................................................ 35
pageLayoutCol ............................................................. 36
pageLayoutRow ............................................................. 36
pagePlotPlace ............................................................. 37
pagePlotRemove ............................................................ 39
pgParams ................................................................. 40
plotCircle ................................................................. 44
plotgardener ............................................................... 45
plotGenes ................................................................. 46
plotGenomeLabel .......................................................... 49
plotGG ................................................................. 52
annoDomains

Annotate domains in a Hi-C plot

Description

Annotate domains in a Hi-C plot

Usage

annoDomains(
  plot,
  data,
  half = "inherit",
  linecolor = "black",
  params = NULL,
  ...
)

Arguments

plot          Hi-C plot object from plotHicSquare or plotHicTriangle on which to annotate pixels.
data          A string specifying the BED file path, a dataframe in BED format, or a GRanges object specifying domain ranges.
half

Character value specifying which half of hic plots to annotate. Triangle Hi-C plots will always default to the entirety of the triangular plot. Default value is half = "inherit". Options are:

- "inherit": Domains will be annotated on the half inherited by the input Hi-C plot.
- "both": Domains will be annotated on both halves of the diagonal of a square Hi-C plot.
- "top": Domains will be annotated on the upper diagonal half of a square Hi-C plot.
- "bottom": Domains will be annotated on the bottom diagonal half of a square Hi-C plot.

linecolor

A character value specifying the color of the domain annotations. Default value is linecolor = "black".

params

An optional pgParams object containing relevant function parameters.

Additional grid graphical parameters. See gpar.

Value

Returns a domain object containing relevant genomic region, placement, and grob information.

Examples

```r
## Define a GRanges object with TAD ranges
library(GenomicRanges)
library(IRanges)
domains <- GRanges("chr21",
  ranges = IRanges(
    start = c(28210000, 29085000, 29430000, 29700000),
    end = c(29085000, 29430000, 29700000, 30125000)
  )
)

## Load Hi-C data
library(plotgardenerData)
data("IMR90_HiC_10kb")

## Create page
pageCreate(width = 4.5, height = 4, default.units = "inches")

## Plot and place a square Hi-C plot
hicPlot <- plotHicSquare(
  data = IMR90_HiC_10kb, resolution = 10000,
  zrange = c(0, 70),
  chrom = "chr21",
  chromstart = 28000000, chromend = 30000000,
  assembly = "hg19",
  x = 0.5, y = 0.5, width = 3, height = 3,
  just = c("left", "top"),
  default.units = "inches"
)
```
## Annotate domains on bottom half of Hi-C plot

```r
annoDomains(
    plot = hicPlot, data = domains,
    half = "bottom", linecolor = "red"
)
```

## Annotate heatmap legend

```r
annoHeatmapLegend(
    plot = hicPlot,
    x = 3.6, y = 0.5, width = 0.12, height = 1.2,
    just = c("left", "top"), default.units = "inches"
)
```

## Annotate genome label

```r
annoGenomeLabel(
    plot = hicPlot, x = 0.5, y = 3.53, scale = "Mb",
    just = c("left", "top")
)
```

## Hide page guides

```r
pageGuideHide()
```

---

### annoGenomeLabel

**Annotate genomic coordinates along the x or y-axis of a plot**

**Description**

Annotate genomic coordinates along the x or y-axis of a plot

**Usage**

```r
annoGenomeLabel(
    plot, fontsize = 10,
    fontcolor = "black",
    linecolor = "black",
    margin = unit(1, "mm"),
    scale = "bp",
    commas = TRUE,
    sequence = TRUE,
    boxWidth = 0.5,
    axis = "x",
    at = NULL,
    tcl = 0.5,
    x,
    y,
    just = c("left", "top"),
    default.units = "inches",
```
annoGenomeLabel

```r
params = NULL,
...
)
```

**Arguments**

- `plot`: Input plot to annotate genomic coordinates. Genomic coordinates and assembly will be inherited from `plot`.
- `fontsize`: A numeric specifying text fontsize in points. Default value is `fontsize = 10`.
- `fontcolor`: A character value indicating the color for text. Default value is `fontcolor = "black"`.
- `linecolor`: A character value indicating the color of the genome label axis. Default value is `linecolor = "black"`.
- `margin`: A numeric or unit vector specifying space between axis and coordinate labels. Default value is `margin = unit(1, "mm")`.
- `scale`: A character value indicating the scale of the coordinates along the genome label. Default value is `scale = "bp"`. Options are:
  - "bp": base pairs.
  - "Kb": kilobase pairs. 1 kilobase pair is equal to 1000 base pairs.
  - "Mb": megabase pairs. 1 megabase pair is equal to 1000000 base pairs.
- `commas`: A logical value indicating whether to include commas in start and stop labels. Default value is `commas = TRUE`.
- `sequence`: A logical value indicating whether to include sequence information above the label of an x-axis (only at appropriate resolutions).
- `boxWidth`: A numeric value indicating the width of the boxes representing sequence information at appropriate resolutions. Default value is `boxWidth = 0.5`.
- `axis`: A character value indicating along which axis to add genome label. Sequence information will not be displayed along a y-axis. Default value is `axis = "x"`. Options are:
  - "x": Genome label will be plotted along the x-axis.
  - "y": Genome label will be plotted along the y-axis. This is typically used for a square Hi-C plot made with `plotHicSquare`.
- `at`: A numeric vector of x-value locations for tick marks.
- `tcl`: A numeric specifying the length of tickmarks as a fraction of text height. Default value is `tcl = 0.5`.
- `x`: A numeric or unit object specifying genome label x-location.
- `y`: A numeric, unit object, or character containing a "b" combined with a numeric value specifying genome label y-location. The character value will place the genome label y relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.
- `just`: Justification of genome label relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is `just = c("left", "top")`.
annoGenomeLabel

**default.units**
A string indicating the default units to use if \( x \) or \( y \) are only given as numerics. Default value is `default.units = "inches"`.

**params**
An optional `pgParams` object containing relevant function parameters.

... Additional grid graphical parameters or digit specifications. See `gpar` and `formatC`.

**Value**
Returns a `genomeLabel` object containing relevant genomic region, placement, and `grob` information.

**Examples**

```r
## Load hg19 genomic annotation packages
library("TxDb.Hsapiens.UCSC.hg19.knownGene")
library("org.Hs.eg.db")

## Create page
pageCreate(width = 5, height = 2, default.units = "inches")

## Plot and place gene track on page
genesPlot <- plotGenes(
  chrom = "chr8",
  chromstart = 1000000, chromend = 2000000,
  assembly = "hg19", fill = c("grey", "grey"),
  fontcolor = c("grey", "grey"),
  x = 0.5, y = 0.25, width = 4, height = 1,
  just = c("left", "top"),
  default.units = "inches"
)

## Annotate x-axis genome labels at different scales
annoGenomeLabel(
  plot = genesPlot, scale = "Mb",
  x = 0.5, y = 1.25, just = c("left", "top"),
  default.units = "inches"
)
annoGenomeLabel(
  plot = genesPlot, scale = "Kb",
  x = 0.5, y = 1.5, just = c("left", "top"),
  default.units = "inches"
)
annoGenomeLabel(
  plot = genesPlot, scale = "bp",
  x = 0.5, y = 1.75, just = c("left", "top"),
  default.units = "inches"
)

## Hide page guides
pageGuideHide()
```
annoHeatmapLegend  
Add a color scale legend for heatmap-style plots

Description
Add a color scale legend for heatmap-style plots

Usage
annoHeatmapLegend(
  plot,  
  orientation = "v", 
  fontsize = 8, 
  fontcolor = "dark grey", 
  scientific = FALSE, 
  digits = 1, 
  ticks = FALSE, 
  breaks = NULL, 
  border = FALSE, 
  x, 
  y, 
  width, 
  height, 
  just = c("left", "top"), 
  default.units = "inches", 
  params = NULL, 
  ...
)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>plot</td>
<td>Heatmap-style plot object to add heatmap legend for.</td>
</tr>
<tr>
<td>orientation</td>
<td>A string specifying legend orientation. Default value is orientation = &quot;v&quot;. Options are:</td>
</tr>
<tr>
<td>fontsize</td>
<td>A numeric specifying text fontsize in points. Default value is fontsize = 8.</td>
</tr>
<tr>
<td>fontcolor</td>
<td>Character value specifying text fontcolor. Default value is fontcolor = &quot;dark grey&quot;.</td>
</tr>
<tr>
<td>scientific</td>
<td>Logical value specifying if numeric color value labels should be encoded in scientific format. Default value is scientific = FALSE.</td>
</tr>
<tr>
<td>digits</td>
<td>Numeric specifying how many significant digits to include of numeric color value labels. Default value is digits = 1.</td>
</tr>
<tr>
<td>ticks</td>
<td>Logical value specifying if tick marks on the heatmap colorbar should be visible. Default value is ticks = FALSE.</td>
</tr>
</tbody>
</table>
annoHeatmapLegend

breaks A numeric vector specifying tick breaks. Default value is breaks = NULL.

border Logical value indicating whether to add a border around heatmap legend. Default value is border = FALSE.

x A numeric or unit object specifying x-location of legend.

y A numeric, unit object, or character containing a "b" combined with a numeric value specifying y-location of legend. The character value will place the legend y relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.

width A numeric or unit object specifying width of legend.

height A numeric or unit object specifying height of legend.

just Justification of heatmap legend relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is just = c("left", "top").

default.units A string indicating the default units to use if x, y, width, or height are only given as numerics. Default value is default.units = "inches".

params An optional pgParams object containing relevant function parameters.

... Additional grid graphical parameters. See gpar.

Value

Returns a heatmapLegend object with relevant color value, placement, and grob information.

Examples

```r
## Load Hi-C data
library(plotgardenerData)
data("IMR90_HiC_10kb")

## Create page
pageCreate(width = 4, height = 3.5, default.units = "inches")

## Plot and place a square Hi-C plot
hicPlot <- plotHicSquare(
  data = IMR90_HiC_10kb, resolution = 10000,
  zrange = c(0, 70),
  chrom = "chr21",
  chromstart = 28000000, chromend = 30300000,
  assembly = "hg19",
  x = 0.5, y = 0.5, width = 2.5, height = 2.5,
  just = c("left", "top"),
  default.units = "inches"
)

## Add heatmap legend
annoHeatmapLegend(
  plot = hicPlot,
)```

annoHighlight

Annotates a highlight box around a specified genomic region of a plot

Description

Annotates a highlight box around a specified genomic region of a plot

Usage

annoHighlight(
  plot,
  chrom,
  chromstart = NULL,
  chromend = NULL,
  fill = "grey",
  linecolor = NA,
  alpha = 0.4,
  y,
  height,
  just = c("left", "top"),
  default.units = "inches",
  params = NULL,
  ...
)

Arguments

plotInput plot on which to annotate genomic region.
chromChromosome of region to be highlighted, as a string.
chromstartInteger start position on chromosome to be highlighted.
chromendInteger end position on chromosome to be highlighted.
fillA character value specifying highlight box fill color. Default value is fill = "grey".
annoHighlight

**linecolor**
A character value specifying highlight box line color. Default value is `linecolor = NA`.

**alpha**
Numeric value specifying color transparency. Default value is `alpha = 0.4`.

**y**
A numeric, unit object, or character containing a "b" combined with a numeric value specifying square highlight box y-location. The character value will place the highlight box y relative to the bottom of the most recently plotted plot according to the units of the `plotgardener` package.

**height**
A numeric or unit object specifying highlight box height.

**just**
Justification of highlight box relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is `just = c("left", "top")`.

**default.units**
A string indicating the default units to use if `y` or `height` are only given as numerics or numeric vectors. Default value is `default.units = "inches"`.

**params**
An optional `pgParams` object containing relevant function parameters.

... Additional grid graphical parameters. See `gpar`.

**Value**

Returns a highlight object containing relevant genomic region, placement, and `grob` information.

**Examples**

```r
## Create a page
pageCreate(width = 7.5, height = 1.5, default.units = "inches")

## Plot and place a signal plot
library(plotgardenerData)
data("IMR90_ChIP_H3K27ac_signal")
region <- pgParams(
  chrom = "chr21",
  chromstart = 28000000, chromend = 30300000,
  assembly = "hg19",
  range = c(0, 45)
)
signalPlot <- plotSignal(
  data = IMR90_ChIP_H3K27ac_signal, params = region,
  x = 0.5, y = 0.25, width = 6.5, height = 0.65,
  just = c("left", "top"),
  default.units = "inches"
)

## Highlight genomic region on signal plot
annoHighlight(
  plot = signalPlot,
  chrom = "chr21",
  chromstart = 29000000, chromend = 29125000,
  y = 0.25, height = 1, just = c("left", "top"),
  default.units = "inches"
)
```
annoPixels

Annotate pixels in a Hi-C plot

Description
Annotate pixels in a Hi-C plot

Usage
annoPixels(
  plot, 
  data, 
  type = "box", 
  half = "inherit", 
  shift = 4, 
  params = NULL, 
  quiet = FALSE, 
  ... 
)

Arguments

plot    Hi-C plot object from plotHicSquare or plotHicTriangle on which to annotate pixels.
data    A string specifying the BEDPE file path, a dataframe in BEDPE format specifying pixel positions, or a GInteractions object specifying pixel positions.
type    Character value specifying type of annotation. Default value is type = "box". Options are:
        • "box": Boxes are drawn around each pixel.
annoPixels

- "circle": Circles are drawn around each pixel.
- "arrow": Arrows are drawn pointing to each pixel.

half

Character value specifying which half of hic plots to annotate. Triangle Hi-C plots will always default to the entirety of the triangular plot. Default value is half = "inherit". Options are:

- "inherit": Pixels will be annotated on the half inherited by the input Hi-C plot.
- "both": Pixels will be annotated on both halves of the diagonal of a square Hi-C plot.
- "top": Pixels will be annotated on the upper diagonal half of a square Hi-C plot.
- "bottom": Pixels will be annotated on the bottom diagonal half of a square Hi-C plot.

shift

Numeric specifying the number of pixels on either end of main pixel in a box or circle. Numeric specifying number of pixels for the length of an arrow.

params

An optional pgParams object containing relevant function parameters.

quiet

A logical indicating whether or not to print messages.

... Additional grid graphical parameters. See gpar.

Value

Returns a pixel object containing relevant genomic region, placement, and grob information.

Examples

```r
## Load Hi-C data and BEDPE data
library(plotgardenerData)
data("IMR90_HiC_10kb")
data("IMR90_DNAloops_pairs")

## Create page
pageCreate(width = 4.5, height = 4, default.units = "inches")

## Plot and place a square Hi-C plot
hicPlot <- plotHicSquare(
  data = IMR90_HiC_10kb, resolution = 10000,
  zrange = c(0, 70),
  chrom = "chr21",
  chromstart = 28000000, chromend = 30300000,
  assembly = "hg19",
  x = 0.5, y = 0.5, width = 3, height = 3,
  just = c("left", "top"),
  default.units = "inches"
)

## Annotate loops of both sides of Hi-C plot with squares
pixels <- annoPixels(
  plot = hicPlot, data = IMR90_DNAloops_pairs, type = "box",
)```

annoSegments

Annotates a line segment within a plot

Description

Annotates a line segment within a plot

Usage

annoSegments(
  x0,
  y0,
  x1,
  y1,
  plot,
  default.units = "native",
  linecolor = "black",
  lwd = 1,
annoSegments

lty = 1,
lineend = "butt",
linejoin = "mitre",
arrow = NULL,
params = NULL,
...
)

Arguments

x0 A numeric vector or unit object indicating the starting x-values of the line segments.
y0 A numeric vector or unit object indicating the starting y-values of the line segments.
x1 A numeric vector or unit object indicating the stopping x-values of the line segments.
y1 A numeric vector or unit object indicating the stopping y-values of the line segments.
plot Input plotgardener plot to internally plot line segments relative to.
default.units A string indicating the default units to use if x0, y0, x1, or y1 are only given as numeric vectors. Default value is default.units = "native".
linecolor A character value specifying segment line color. Default value is linecolor = "black".
lwd A numeric specifying segment line width. Default value is lwd = 1.
lty A numeric specifying segment line type. Default value is lty = 1.
lineend A character value specifying line end style. Default value is lineend = "butt". Options are:
  • "round": Segment ends are rounded.
  • "butt": Segment ends end exactly where ended.
  • "square": Segment ends are squared.
linejoin A character value specifying line join style. Default value is linejoin = "mitre". Options are:
  • "round": Line joins are rounded.
  • "mitre": Line joins are sharp corners.
  • "bevel": Line joins are flattened corners.
arrow A list describing arrow heads to place at either end of the line segments, as produced by the arrow function.
params An optional pgParams object containing relevant function parameters.
... Additional grid graphical parameters. See gpar.

Value

Returns a segments object containing relevant placement and grob information.
See Also

grid.segments, arrow

Examples

library(grid)
## Create a page
pageCreate(width = 7.5, height = 2.5, default.units = "inches")

## Plot a Manhattan plot
library(plotgardenerData)
library("TxDb.Hsapiens.UCSC.hg19.knownGene")
data("hg19_insulin_GWAS")
manhattanPlot <- plotManhattan(
data = hg19_insulin_GWAS, assembly = "hg19",
fill = c("grey", "#37a7db"),
sigLine = TRUE,
col = "grey", lty = 2, range = c(0, 14),
x = 0.5, y = 0, width = 6.5, height = 2,
just = c("left", "top"),
default.units = "inches"
)
## Annotate genome label
annoGenomeLabel(
plot = manhattanPlot, x = 0.5, y = 2, fontsize = 8,
just = c("left", "top"),
default.units = "inches"
)
plotText(
label = "Chromosome", fontsize = 8,
x = 3.75, y = 2.20, just = "center", default.units = "inches"
)
## Annotate y-axis
annoYaxis(
plot = manhattanPlot, at = c(0, 2, 4, 6, 8, 10, 12, 14),
axisLine = TRUE, fontsize = 8
)
## Annotate a line segment for an additional significance line of
## the Manhattan plot
annoSegments(
x0 = unit(0, "npc"), y0 = 10,
x1 = unit(1, "npc"), y1 = 10,
plot = manhattanPlot, default.units = "native",
linecolor = "red", lty = 2
)
## Plot y-axis label
plotText(
label = "-log10(p-value)", x = 0.15, y = 1, rot = 90,
annoText

annoText(  
  label,  
  fontcolor = "black",  
  fontsize = 12,  
  rot = 0,  
  check.overlap = FALSE,  
  plot,  
  x,  
  y,  
  just = "center",  
  default.units = "native",  
  params = NULL,  
  ...  
)

Arguments

- **label**: Character or expression of text to be plotted.
- **fontcolor**: A character value specifying text fontcolor. Default value is fontcolor = "black".
- **fontsize**: A numeric specifying text fontsize in points. Default value is fontsize = 12.
- **rot**: A numeric specifying the angle to rotate the text. Default value is rot = 0.
- **check.overlap**: A logical value to indicate whether to check for and omit overlapping text. Default value is check.overlap = FALSE.
- **plot**: Input plotgardener plot to internally place text relative to.
- **x**: A numeric vector or unit object specifying text x-location.
- **y**: A numeric vector or unit object specifying text y-location.
- **just**: Justification of text relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is just = "center".

Description

Annotates text within a plot

Usage

annoText(  
  label,  
  fontcolor = "black",  
  fontsize = 12,  
  rot = 0,  
  check.overlap = FALSE,  
  plot,  
  x,  
  y,  
  just = "center",  
  default.units = "native",  
  params = NULL,  
  ...  
)
default.units A string indicating the default units to use if x or y are only given as numerics or numeric vectors. Default value is default.units = "native".

params An optional pgParams object containing relevant function parameters.

... Additional grid graphical parameters. See gpar.

Value

Returns a text object containing relevant placement and grob information.

See Also

grid.text

Examples

## Create a page
pageCreate(width = 4, height = 4, default.units = "inches")

## Plot text relative to a plotgardener plot
library(plotgardenerData)
data("IMR90_HiC_10kb")
hicPlot <- plotHicSquare(
  data = IMR90_HiC_10kb, chrom = "chr21",
  chromstart = 28000000, chromend = 29500000,
  assembly = "hg19",
  zrange = c(0, 70),
  x = 0.5, y = 0.5, width = 3, height = 3,
  just = c("left", "top"),
  default.units = "inches"
)
annoGenomeLabel(
  plot = hicPlot, x = 0.5, y = 3.55, scale = "Mb",
  just = c("left", "top"), default.units = "inches"
)

annoText(
  label = "Loop", fontsize = 8, plot = hicPlot,
  x = 29075000, y = 28150000,
  just = "center", default.units = "native"
)

## Hide page guides
pageGuideHide()
Usage

annoXaxis(
  plot,
  at = NULL,
  label = TRUE,
  main = TRUE,
  scipen = 999,
  axisLine = FALSE,
  params = NULL,
  ...
)

Arguments

plot  Plot object to annotate with x-axis.
at A numeric vector of x-value locations for tick marks.
label A logical value indicating whether to draw the labels on the tick marks, or an
expression or character vector which specify the labels to use. If not logical, must be the same length as the at argument.
main A logical value indicating whether to draw the x-axis at the bottom of the plot.
Default value is main = TRUE. Options are:
  • TRUE: x-axis is drawn at the bottom of the plot.
  • FALSE: x-axis is drawn at the top of the plot.
scipen An integer indicating the penalty to be applied when deciding to print numeric
values in fixed or exponential notation. Default value is scipen = 999.
axisLine A logical value indicating whether to show the axis line. Default value is axisLine
  = FALSE.
params An optional pgParams object containing relevant function parameters.
... Additional grid graphical parameters. See gpar.

Value

Returns a xaxis object containing relevant grob information.

Examples

## Load transcript information
library("TxDb.Hsapiens.UCSC.hg19.knownGene")
library("org.Hs.eg.db")

## Create page
pageCreate(width = 7.5, height = 4.5, default.units = "inches")

## Plot gene transcripts
transcriptPlot <- plotTranscripts(
  chrom = "chr1",
  chromstart = 1000000,
annoYaxis

Add a y-axis to a plot

Description
Add a y-axis to a plot

Usage

annoYaxis(
  plot,
  at = NULL,
  label = TRUE,
  main = TRUE,
  scipen = 999,
  axisLine = FALSE,
  params = NULL,
  ...
)

Arguments

plot  Plot object to annotate with y-axis.
at   A numeric vector of y-value locations for tick marks.
label A logical value indicating whether to draw the labels on the tick marks, or an expression or character vector which specify the labels to use. If not logical, must be the same length as the at argument.
main

A logical value indicating whether to draw the y-axis at the left of the plot. Default value is main = TRUE. Options are:

- TRUE: y-axis is drawn at the left of the plot.
- FALSE: y-axis is drawn at the right of the plot.

scipen

An integer indicating the penalty to be applied when deciding to print numeric values in fixed or exponential notation. Default value is scipen = 999.

axisLine

A logical value indicating whether to show the axis line. Default value is axisLine = FALSE.

params

An optional pgParams object containing relevant function parameters.

Additional grid graphical parameters. See gpar.

Value

Returns a yaxis object containing relevant grob information.

Examples

```r
## Load Hi-C data
library(plotgardenerData)
data("IMR90_HiC_10kb")

## Create page
pageCreate(width = 4, height = 3.5, default.units = "inches")

## Plot and place a square Hi-C plot
hicPlot <- plotHicSquare(
  data = IMR90_HiC_10kb, resolution = 10000,
  zrange = c(0, 70),
  chrom = "chr21",
  chromstart = 28000000, chromend = 30300000,
  assembly = "hg19",
  x = 1, y = 0.5, width = 2.5, height = 2.5,
  just = c("left", "top"),
  default.units = "inches"
)

## Add standard y-axis to Hi-C plot
annoYaxis(
  plot = hicPlot, at = c(28000000, 29000000, 30300000),
  fontsize = 10
)

## Annotate genome label on x-axis
annoGenomeLabel(plot = hicPlot, x = 1, y = 3.03)

## Annotate heatmap legend
annoHeatmapLegend(
  plot = hicPlot,
  x = 3.6, y = 0.5, width = 0.12, height = 1.2
)
```
# Hide page guides
pageGuideHide()

## annoZoomLines
Annotates zoom lines for a specified genomic region of a plot

### Description
Annotates zoom lines for a specified genomic region of a plot

### Usage

```r
annoZoomLines(
  plot,
  chrom,
  chromstart = NULL,
  chromend = NULL,
  y0,
  x1 = NULL,
  y1,
  extend = 0,
  default.units = "inches",
  linecolor = "grey",
  lty = 2,
  params = NULL,
  ...
)
```

### Arguments

- **plot**: Input plot to annotate genomic region zoom lines from.
- **chrom**: Chromosome of region to draw zoom lines from, as a string.
- **chromstart**: Integer start position on chromosome to draw zoom lines from.
- **chromend**: Integer end position on chromosome to draw zoom lines from.
- **y0**: A numeric vector or unit object indicating the starting y-values of the zoom line segments. If two values are given, the first value will correspond to the left zoom line and the second value will correspond to the right zoom line.
- **x1**: A numeric vector or unit object indicating the stopping x-values of the zoom line segments. If two values are given, the first value will correspond to the left zoom line and the second value will correspond to the right zoom line. If NULL, straight lines from zoomed genomic region will be drawn.
- **y1**: A numeric vector or unit object indicating the stopping y-values of the zoom line segments. If two values are given, the first value will correspond to the left zoom line and the second value will correspond to the right zoom line.
extend  A numeric vector or unit object indicating the length to extend straight lines from each end of the zoom line segments. If two values are given, the first value will correspond to the top extension length and the second value will correspond to the bottom extension length. Default value is extend = 0.

default.units  A string indicating the default units to use if y0, x1, y1, or extend are only given as numerics or numeric vectors. Default value is default.units = "inches".

default.units  A string indicating the default units to use if y0, x1, y1, or extend are only given as numerics or numeric vectors. Default value is default.units = "inches".

default.units  A string indicating the default units to use if y0, x1, y1, or extend are only given as numerics or numeric vectors. Default value is default.units = "inches".

default.units  A string indicating the default units to use if y0, x1, y1, or extend are only given as numerics or numeric vectors. Default value is default.units = "inches".

Value

Returns a zoom object containing relevant genomic region, placement, and grob information.

Examples

```r
## Create a page
pageCreate(width = 7.5, height = 4.75, default.units = "inches")

## Plot and place a Manhattan plot
library(plotgardenerData)
library("TxDb.Hsapiens.UCSC.hg19.knownGene")
data("hg19_insulin_GWAS")
manhattanPlot <- plotManhattan(
  data = hg19_insulin_GWAS, assembly = "hg19",
  fill = c("grey", "#37a7db"),
  sigLine = FALSE,
  col = "grey", lty = 2, range = c(0, 14),
  x = 0.5, y = 0, width = 6.5, height = 2,
  just = c("left", "top"),
  default.units = "inches"
)
annoYaxis(
  plot = manhattanPlot, at = c(0, 2, 4, 6, 8, 10, 12, 14),
  axisLine = TRUE, fontsize = 8
)

## Annotate zoom lines for a region on chromosome 21
zoomRegion <- pgParams(
  chrom = "chr21",
  chromstart = 28000000, chromend = 30300000,
  assembly = "hg19"
)
annoZoomLines(
  plot = manhattanPlot, params = zoomRegion,
  y0 = 2, x1 = c(0.5, 7), y1 = 2.5, extend = c(0, 1.1),
  default.units = "inches",
  lty = 3
)
```
## Annotate highlight region for zoom region

```r
annoHighlight(
  plot = manhattanPlot, params = zoomRegion,
  y = 2, height = 2, just = c("left", "bottom"),
  default.units = "inches",
  fill = "red", alpha = 0.8
)
```

## Plot Manhattan plot data and signal track under zoom lines

```r
manhattanPlotZoom <- plotManhattan(
  data = hg19_insulin_GWAS,
  fill = "grey",
  sigLine = FALSE,
  baseline = TRUE,
  params = zoomRegion, range = c(0, 14),
  x = 0.5, y = 2.6,
  width = 6.5, height = 1
)
data("IMR90_ChIP_H3K27ac_signal")
signalPlot <- plotSignal(
  data = IMR90_ChIP_H3K27ac_signal, params = zoomRegion,
  range = c(0, 45),
  x = 0.5, y = "b0.1",
  width = 6.5, height = 0.65,
  just = c("left", "top"),
  default.units = "inches"
)
```

## Plot genome label

```r
plotGenomeLabel(
  chrom = "chr21",
  chromstart = 28000000, chromend = 30300000,
  assembly = "hg19",
  x = 0.5, y = 4.4, length = 6.5,
  default.units = "inches"
)
```

## Hide page guides

```r
pageGuideHide()
```

---

**assembly**

Make a assembly object for alternate TxDb, OrgDb, and BSgenome genomic annotation packages

**Description**

Make a assembly object for alternate TxDb, OrgDb, and BSgenome genomic annotation packages
Usage

```r
assembly(
  Genome,
  TxDb,
  OrgDb,
  gene.id.column = "ENTREZID",
  display.column = "SYMBOL",
  BSgenome = NULL
)
```

Arguments

- **Genome** String indicating the name of the genome assembly.
- **TxDb** String of existing TxDb package name or a TxDb object.
- **OrgDb** String of the desired OrgDb package name.
- **gene.id.column** String of the TxDb column name that refers to the given TxDb gene IDs. Default value is `gene.id.column = "ENTREZID"`.
- **display.column** String of the OrgDb column name that refers to the type of gene symbol to be displayed in plots. Default value is `display.column = "SYMBOL"`.
- **BSgenome** String of the desired BSgenome package name.

Value

Returns a `assembly` object containing all input parameters.

See Also

`TxDb`, `OrgDb-class`, `BSgenome`

Examples

```r
## Create a custom `assembly` object for hg38/GRCh38 packages
newAssembly <- assembly(
  Genome = "hg38_GRCh38",
  TxDb = "TxDb.Hsapiens.UCSC.hg38.knownGene",
  OrgDb = "org.Hs.eg.db",
  BSgenome = "BSgenome.Hsapiens.NCBI.GRCh38"
)
```

---

Combine multiple pgParams objects into a vector

Description

Combine multiple pgParams objects into a vector
Usage

c(..., recursive = FALSE)

Arguments

... pgParams objects to be concatenated.
recursive logical. If recursive = TRUE, the function recursively descends through lists (and pairlists) combining all their elements into a vector.

Value

NULL or an expression or a vector of an appropriate mode. (With no arguments the value is NULL.)

Examples

## Define parameters
p1 <- pgParams(chrom = "chr1", assembly = "hg19")

## Define another set of parameters
p2 <- pgParams(fontsize = 10, assembly = "hg19")

## Combine parameters into one `pgParams` object
pTotal <- c(p1, p2)

calcSignalRange Calculate a score range for multiple signals

Description

Calculate a score range for multiple signals

Usage

calcSignalRange(
  data,
  chrom = NULL,
  chromstart = 1,
  chromend = .Machine$integer.max,
  assembly = "hg38",
  negData = FALSE)

Arguments

data List of data to be plotted as character values specifying multiple bigwig file paths, dataframes in BED format, or GRanges objects with metadata column score.

chrom Chromosome of data region range as a string, if range for a specific chromosome is desired.
chromstart  Integer start position on chromosome to get data range.
chromend   Integer end position on chromosome to get data range.
assembly   Default genome assembly as a string or a assembly object. Default value is assembly = "hg38".
negData    A logical value indicating whether any of the data has both positive and negative scores and the signal range should be adjusted accordingly. Default value is negData = FALSE.

Value

Returns a vector of length 2 with the calculated c(min, max) range.

Examples

library("plotgardenerData")
data("GM12878_ChIP_CTCF_signal")
data("IMR90_ChIP_CTCF_signal")
data("GM12878_ChIP_H3K27ac_signal")
data("IMR90_ChIP_H3K27ac_signal")
calcSignalRange(data = list(GM12878_ChIP_CTCF_signal, 
    GM12878_ChIP_H3K27ac_signal, 
    IMR90_ChIP_CTCF_signal, 
    IMR90_ChIP_H3K27ac_signal),
    chrom = "chr21",
    chromstart = 28150000, chromend = 29150000,
    assembly = "hg38", negData = FALSE)

---

**colorby**  
Handle plotgardener color scaling parameters

**Description**

colorby should be used to create a set of parameters that specify color scaling for the functions plotPairs, plotPairsArches, and plotRanges.

**Usage**

colorby(column, palette = NULL, range = NULL, scalePerRegion = FALSE)

**Arguments**

column    String specifying name of data column to scale colors by.
palette    (optional) A function describing the color palette to use for color scaling.
range      (optional) A numeric vector specifying the range of values to apply a color scale to.
scalePerRegion  A logical value indicating whether to adjust NULL range of numerical `colorby` values to subset of data in a plotted genomic region. Default value is `scalePerRegion = FALSE`.

Value

Returns a "colorby" object.

Examples

```r
## Load paired ranges data in BEDPE format
library(plotgardenerData)
data("IMR90_DNAloops_pairs")

## Add a length column
IMR90_DNAloops_pairs$length <-
  (IMR90_DNAloops_pairs$start2 - IMR90_DNAloops_pairs$start1) / 1000

## Plot pairs with colorby object set for `length` column
bedpePlot <- plotPairs(
  data = IMR90_DNAloops_pairs,
  chrom = "chr21",
  chromstart = 27900000, chromend = 30700000,
  assembly = "hg19",
  fill = colorby("length", palette =
    colorRampPalette(c("dodgerblue2", "firebrick2"))),
  lwd = 2, spaceHeight = .7,
)
```

---

defaultPackages  

Display the default genomic annotation packages associated with a genome build

Description

Display the default genomic annotation packages associated with a genome build

Usage

```r
defaultPackages(Genome)
```

Arguments

- `Genome`  
  String indicating the name of the genome assembly.

Value

Returns a list of the default data packages for a genome build.
genomes

Examples

```r
## View default genomic annotation packages associated with "hg19"
defaultPackages(Genome = "hg19")

## View default genomic annotation packages associated with "mm9"
defaultPackages(Genome = "mm9")
```

genomes

Display the included available default genome assemblies

Description

Display the included available default genome assemblies

Usage

genomes()

Value

Returns the included available default genome assemblies

Examples

genomes()

mapColors

Maps a numeric or character vector to a color palette and returns the vector of colors

Description

Maps a numeric or character vector to a color palette and returns the vector of colors

Usage

mapColors(vector, palette, range = NULL)

Arguments

- **vector**: Vector to map to color.
- **palette**: Color palette function.
- **range**: Range of values to map for a numerical value.
mapColors

Details
This function allows for the manual mapping of a numerical or factor vector to a palette of colors. For a more automatic implementation of this functionality in plotgardener functions, colorby objects can be used.

Value
Returns a character vector of color values. If the input vector is numerical, this vector will have additional ‘palette’ and ‘range’ attributes.

See Also
colorby

Examples
```r
## Load paired ranges data in BEDPE format
library(plotgardenerData)
data("IMR90_DNAloops_pairs")

## Add a length column
IMR90_DNAloops_pairs$length <- (IMR90_DNAloops_pairs$start2 -
    IMR90_DNAloops_pairs$start1) / 1000

## Map length column to a vector of colors
colors <- mapColors(vector = IMR90_DNAloops_pairs$length,
    palette = colorRampPalette(c("dodgerblue2", "firebrick2")))

## Pass color vector into bbPlotPairsArches
heights <- IMR90_DNAloops_pairs$length / max(IMR90_DNAloops_pairs$length)
pageCreate(width = 7.5, height = 2.1, default.units = "inches",
    showGuides = FALSE, xgrid = 0, ygrid = 0)
params <- pgParams(chrom = "chr21",
    chromstart = 27900000, chromend = 30700000,
    assembly = "hg19",
    width = 7)

archPlot <- plotPairsArches(data = IMR90_DNAloops_pairs, params = params,
    fill = colors,
    linecolor = "fill",
    archHeight = heights, alpha = 1,
    x = 0.25, y = 0.25, height = 1.5,
    just = c("left", "top"),
    default.units = "inches")

annoGenomeLabel(plot = archPlot, x = 0.25, y = 1.78, scale = "Mb")
annoHeatmapLegend(plot = archPlot, fontcolor = "black",
    textcolor = "black", label = "Length")
```
```
Create a page for a plotgardener layout

Description

Create a page for a plotgardener layout

Usage

pageCreate(
  width = 8.5,
  height = 11,
  default.units = "inches",
  bg = NA,
  xgrid = 0.5,
  ygrid = 0.5,
  showGuides = TRUE,
  params = NULL
)

Arguments

- **width**: A numeric or unit object specifying page width. Default value is width = 8.
- **height**: A numeric or unit object specifying page height. Default value is height = 11.
- **default.units**: A string indicating the default units to use if width or height are only given as numerics. Default value is default.units = "inches".
- **bg**: Character value indicating page background color. Default value is bg = NA.
- **xgrid**: A numeric indicating the increment by which to place vertical gridlines. Default value is xgrid = 0.5.
- **ygrid**: A numeric indicating the increment by which to place horizontal gridlines. Default value is ygrid = 0.5.
- **showGuides**: A logical value indicating whether to draw a black border around the entire page and guiding rulers along the top and left side of the page. Default value is showGuides = TRUE.
- **params**: An optional pgParams object containing relevant function parameters.
Details

width and height must be specified in the same units.

Value

None.

Examples

```r
## Create a 6-inch wide, 4.5-inch high page
pageCreate(width = 6, height = 4.5, default.units = "inches")

## Create a 14-cm wide, 10-cm high page
pageCreate(width = 14, height = 10, default.units = "cm")
```

---

**Description**

Remove guides from a plotgardener page

**Usage**

```r
pageGuideHide()
```

**Value**

None.

**Note**

Please note that due to the implementation of `grid` removal functions, using `pageGuideHide` within a `pdf` call will result in the rendering of a separate, new page with the plot guides removed. To avoid this artifact, hide guides in the `pageCreate` function call with `showGuides = FALSE`.

**Examples**

```r
## Make a page
pageCreate(width = 7, height = 4, default.units = "inches")

## Hide page guides
pageGuideHide()
```
**pageGuideHorizontal**  
*Draw a horizontal guideline at a specified y-coordinate on a plotgardener page*

**Description**

Draw a horizontal guideline at a specified y-coordinate on a plotgardener page

**Usage**

```r
pageGuideHorizontal(
  y,
  default.units = "inches",
  linecolor = "grey55",
  params = NULL,
  ...
)
```

**Arguments**

- `y`  
  A numeric or unit object specifying y-coordinate of guide.
- `default.units`  
  A string indicating the default units to use if `y` is only given as a numeric. Default value is `default.units = "inches"`.
- `linecolor`  
  Character value indicating color of guideline. Default value is `linecolor = "grey55"`.
- `params`  
  An optional `pgParams` object containing relevant function parameters.
- `...`  
  Additional grid graphical parameters. See `gpar`.

**Value**

None.

**Examples**

```r
## Create a page
pageCreate(width = 6, height = 5, default.units = "inches")

## Add red horizontal guideline at y = 2.5 inches
pageGuideHorizontal(y = 2.5, linecolor = "red")
```
Description


Usage

```
pageGuideShow()
```

Value

None.

See Also

`pageCreate`, `pageGuideHorizontal`, `pageGuideVertical`

Examples

```r
## Load Hi-C data
library(plotgardenerData)
data("IMR90_HiC_10kb")

## Create a page
pageCreate(width = 3, height = 3, default.units = "inches")

## Add a page guide
pageGuideHorizontal(y = 0.5, default.units = "inches")

## Plot and place Hi-C plot
hicPlot <- plotHicSquare(
  data = IMR90_HiC_10kb, resolution = 10000,
  zrange = c(0, 70),
  chrom = "chr21",
  chromstart = 28000000, chromend = 30300000,
  x = 0.5, y = 0.5, width = 2, height = 2,
  just = c("left", "top"),
  default.units = "inches"
)

## Hide page guides
pageGuideHide()

## Re-show page guides
pageGuideShow()

## Annotate genome label
```
pageGuideVertical

Draw a vertical guideline at a specified x-coordinate on a plotgardener page

Description

Draw a vertical guideline at a specified x-coordinate on a plotgardener page

Usage

pageGuideVertical(
  x,
  default.units = "inches",
  linecolor = "grey55",
  params = NULL,
  ...
)

Arguments

x A numeric or unit object specifying x-coordinate of guide.
default.units A string indicating the default units to use if x is only given as a numeric. Default value is default.units = "inches".
linecolor Character value indicating color of guideline. Default value is linecolor = "grey55".
params An optional pgParams object containing relevant function parameters.
... Additional grid graphical parameters. See gpar.

Value

None.

Examples

## Create a page
pageCreate(width = 6, height = 5, default.units = "inches")

## Add blue vertical guideline at x = 1.7 inches
pageGuideVertical(x = 1.7, linecolor = "blue")
pageLayoutCol

Generate column positions for a number of plot elements with a specified width and space between them

Description

Generate column positions for a number of plot elements with a specified width and space between them

Usage

pageLayoutCol(x, width, space, n, default.units = "inches")

Arguments

x
A numeric or unit object specifying the starting column x-position.

width
A numeric or unit object specifying the width of columns.

space
A numeric or unit object specifying the space between columns.

n
An integer specifying the number of elements to generate column positions for.

default.units
A string indicating the default units to use if x, w, or s are only given as numerics. Default value is default.units = "inches"

Value

Returns a unit vector of page positions.

Examples

# Starting at 0.5 units, return a vector of positions for 3 objects that
# are 2 units in width with 0.1 units of space between them

pageLayoutCol(x = 0.5, width = 2, space = 0.1, n = 3,
               default.units = "inches")

pageLayoutRow

Generate row positions for a number of plot elements with a specified height and space between them

Description

Generate row positions for a number of plot elements with a specified height and space between them
Usage

pageLayoutRow(y, height, space, n, default.units = "inches")

Arguments

y A numeric or unit object specifying the starting row y-position.
height A numeric or unit object specifying the height of rows.
space A numeric or unit object specifying the space between rows.
n An integer specifying the number of elements to generate row positions for.
default.units A string indicating the default units to use if y, h, or s are only given as numerics. Default value is default.units = "inches".

Value

Returns a unit vector of page positions.

Examples

# Starting at 0.5 units, return a vector of positions for 3 objects that
# are 2 units in height with 0.1 units of space between them
pageLayoutRow(y = 0.5, height = 2, space = 0.1, n = 3,
   default.units = "inches")

pagePlotPlace

Place a plot that has been previously created but not drawn

Description

Place a plot that has been previously created but not drawn

Usage

pagePlotPlace(
   plot,
   x = NULL,
   y = NULL,
   width = NULL,
   height = NULL,
   just = c("left", "top"),
   default.units = "inches",
   draw = TRUE,
   params = NULL
)

Arguments

plot  Plot object to be placed, defined by the output of a plotgardener plotting function.

x  A numeric or unit object specifying plot x-location.

y  A numeric, unit object, or character containing a "b" combined with a numeric value specifying plot y-location. The character value will place the plot y relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.

width  A numeric or unit object specifying plot width.

height  A numeric or unit object specifying plot height.

just  Justification of plot relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is just = c("left", "top").

default.units  A string indicating the default units to use if x, y, width, or height are only given as numerics. Default value is default.units = "inches".

draw  A logical value indicating whether graphics output should be produced. Default value is draw = TRUE.

params  An optional pgParams object containing relevant function parameters.

Value

Function will update dimensions of an input plot and return an updated plot object.

Examples

```r
## Load Hi-C data
library(plotgardenerData)
data("IMR90_HiC_10kb")

## Create, but do not plot, square Hi-C plot
hicPlot <- plotHicSquare(
  data = IMR90_HiC_10kb, resolution = 10000,
  zrange = c(0, 70),
  chrom = "chr21",
  chromstart = 28000000, chromend = 30300000,
  draw = FALSE
)

## Create page
pageCreate(width = 3.75, height = 3.5, default.units = "inches")

## Place Hi-C plot on page
pagePlotPlace(
  plot = hicPlot,
  x = 0.25, y = 0.25, width = 3, height = 3,
  just = c("left", "top"),
  default.units = "inches", draw = TRUE
)```

## Annotate heatmap legend
annoHeatmapLegend(
    plot = hicPlot,
    x = 3.4, y = 0.25, width = 0.12, height = 1.2,
    just = c("left", "top"), default.units = "inches"
)

## Hide page guides
pageGuideHide()

---

**Description**

Remove plotgardener plots and annotations

**Usage**

pagePlotRemove(plot)

**Arguments**

- **plot**: Plot object to be removed, defined by the output of a plotgardener plotting function.

**Value**

None.

**Examples**

```r
## Load Hi-C data
library(plotgardenerData)
data("IMR90_HiC_10kb")

## Create page
pageCreate(width = 5.5, height = 4, default.units = "inches")

## Plot and place a square Hi-C plot
hicPlot <- plotHicSquare(
    data = IMR90_HiC_10kb, resolution = 10000,
    zrange = c(0, 70),
    chrom = "chr21",
    chromstart = 28000000, chromend = 30300000,
    x = 0.5, y = 0.5, width = 2.5, height = 2.5,
    just = c("left", "top"),
    default.units = "inches"
)
```
pgParams: plotgardener parameters object

Description

Creates an object of class "pgParams" that can be used by plotgardener functions. pgParams can be used to set a set of parameters to be shared across multiple functions.

Usage

pgParams(assembly = "hg38", gene = NULL, geneBuffer = NULL, ...)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>assembly</td>
<td>String defining the genome build. Default value is assembly = &quot;hg38&quot;.</td>
</tr>
<tr>
<td>gene</td>
<td>(optional) String naming a gene used to set the chrom, chromstart, and chromend arguments.</td>
</tr>
<tr>
<td>geneBuffer</td>
<td>(optional) Integer base-pairs to extend the start and end of a gene defined by argument gene. Can be one integer or a vector of length 2, where the first integer will extend the start of the gene and the second integer will extend the end of the gene.</td>
</tr>
<tr>
<td>...</td>
<td>This function will take any plotgardener function parameters and their values:</td>
</tr>
<tr>
<td></td>
<td>• alpha</td>
</tr>
<tr>
<td></td>
<td>• altchrom</td>
</tr>
<tr>
<td></td>
<td>• altchromend</td>
</tr>
<tr>
<td></td>
<td>• altchromstart</td>
</tr>
<tr>
<td></td>
<td>• archHeight</td>
</tr>
<tr>
<td></td>
<td>• arrow</td>
</tr>
<tr>
<td></td>
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<td>• boxHeight</td>
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## Remove square Hi-C plot from page
pagePlotRemove(plot = hicPlot)
• boxWidth
• breaks
• BSgenome
• cex
• check.overlap
• chrom
• chromend
• chromstart
• clip
• collapse
• colorbyStrand
• colorTrans
• column
• commas
• curvature
• data
• default.units
• digits
• display.column
• draw
• extend
• file
• fill
• flip
• fontcolor
• fontsize
• geneBackground
• geneHighlights
• gene.id.column
• geneOrder
• Genome
• half
• height
• id
• id.lengths
• image
• interpolate
• just
• label
• labels
• leadSNP
• legend
• length
• limitLabel
• linecolor
• lineend
• linejoin
• lty
• lwd
• main
• margin
• matrix
• negData
• norm
• OrgDb
• orientation
• palette
• pch
• plot
• quiet
• r
• range
• resolution
• res_scale
• rot
• scale
• scientific
• scipen
• sequence
• shift
• showBands
• showGuides
• sigCol
• sigLine
• sigVal
• spaceHeight
• spaceWidth
• strand
• strandLabels
• strandSplit
• stroke
• style
• tcl
• ticks
pgParams

- title
- TxDb
- type
- width
- x
- xgrid
- x0
- x1
- y
- ygrid
- ymax
- y0
- y1
- zrange

Details

pgParams generates arguments from exported plotgardener functions at loading time of the package. Arguments defined in a pgParams object can be passed into the params argument of plotgardener functions. params arguments can be overridden from within plotgardener functions.

pgParams also provides an alternative region definition mechanism. Given a gene name and genome assembly, pgParams returns the appropriate "chrom", "chromstart", and "chromend" with a default buffer of (gene length) / 2 added to the ends of the gene coordinates. The buffer amount can be set manually with the geneBuffer parameter. Buffer extending beyond the length of the chromosome will be trimmed.

Value

Returns an object of class pgParams containing plotgardener function arguments.

Examples

```r
## Load hg19 genomic annotation packages
library("TxDb.Hsapiens.UCSC.hg19.knownGene")
library("org.Hs.eg.db")

## Define parameters
p1 <- pgParams(gene = "IL1B", assembly = "hg19")

## Optionally add more parameters
p2 <- pgParams(fontsize = 10, assembly = "hg19")

## Combine parameters and pass them to a plotgardener function
plotGenes(params = c(p1, p2))
```
plotCircle

Plot a circle within a plotgardener layout

Description
Plot a circle within a plotgardener layout

Usage
plotCircle(
  x,
  y,
  r,
  default.units = "inches",
  linecolor = "black",
  lwd = 1,
  lty = 1,
  fill = NA,
  alpha = 1,
  params = NULL,
  ...
)

Arguments

x
A numeric vector or unit object specifying circle x-locations relative to center.

y
A numeric vector, unit object, or a character vector of values containing a "b" combined with a numeric value specifying circle y-locations relative to center. The character vector will place circle y-locations relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.

r
A numeric vector or unit object specifying radii.

default.units
A string indicating the default units to use if r, x, or y are only given as numerics or numeric vectors. Default value is default.units = "inches".

linecolor
A character value specifying circle line color. Default value is linecolor = "black".

lwd
A numeric specifying circle line width. Default value is lwd = 1.

lty
A numeric specifying circle line type. Default value is lty = 1.

fill
A character value specifying circle fill color. Default value is fill = NA.

alpha
Numeric value specifying color transparency. Default value is alpha = 1.

params
An optional pgParams object containing relevant function parameters.

...
Additional grid graphical parameters. See gpar.

Value
Returns a circle object containing relevant placement and grob information.
### plotgardener

**plotgardener: Coordinate-based Genomic Visualization Package for R**

**Description**

plotgardener is a coordinate-based genomic visualization package for R. It grants users the ability to programmatically produce complex, multi-paneled figures. Tailored for genomics, plotgardener allows users to visualize large complex genomic datasets and provides exquisite control over how plots are placed and arranged on a page.

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- Douglas H. Phanstiel <douglas_phanstiel@med.unc.edu> [copyright holder]

Other contributors:
- Sarah Parker <sarmae@live.unc.edu> [contributor]

### Examples

```r
## Create a page
pageCreate(width = 2, height = 2, default.units = "inches")

## Plot two circles, one at a time
plotCircle(
    x = 0.6, y = 0.5, r = 0.1, fill = "black",
    default.units = "inches"
)
plotCircle(
    x = 1.4, y = 0.5, r = 0.1, fill = "black",
    default.units = "inches"
)

## Plot a vector of circles
xVals <- 1 + (0.5 * cos(seq(0, pi, pi / 8)))
yVals <- 1 + (0.5 * sin(seq(0, pi, pi / 8)))
plotCircle(x = xVals, y = yVals, r = 0.05, default.units = "inches")

## Hide page guides
pageGuideHide()
```
PlotGenes

Plot a gene track for a specified genomic region

Description

Plot a gene track for a specified genomic region

Usage

plotGenes(
  chrom,
  chromstart = NULL,
  chromend = NULL,
  assembly = "hg38",
  fontsize = 8,
  fontcolor = c("#669fd9", "#abcc8e"),
  fill = c("#669fd9", "#abcc8e"),
  geneOrder = NULL,
  geneHighlights = NULL,
  geneBackground = "grey",
  strandLabels = TRUE,
  stroke = 0.1,
  bg = NA,
  x = NULL,
  y = NULL,
  width = NULL,
  height = unit(0.6, "inches"),
  just = c("left", "top"),
  default.units = "inches",
  draw = TRUE,
  params = NULL
)

See Also

Useful links:

- https://phanstiellab.github.io/plotgardener
- https://github.com/PhanstielLab/plotgardener
Arguments

chrom Chromosome of region to be plotted, as a string.
chromstart Integer start position on chromosome to be plotted.
chromend Integer end position on chromosome to be plotted.
assembly Default genome assembly as a string or a assembly object. Default value is assembly = "hg38".
fontsize A numeric specifying text fontsize in points. Default value is fontsize = 8.
fontcolor A character value or vector of length 2 indicating the fontcolors for the plus strand and minus strand gene labels. The first value will color the plus strand gene labels and the second value will color the minus strand gene labels. Default value is fontcolor = c("#669fd9", "#abcc8e").
fill A character value or vector of length 2 indicating the strand fill colors for the plus strand and minus strand plot elements. The first value will color the plus strand plot elements and the second label will color the minus strand plot elements. Default value is fill = c("#669fd9", "#abcc8e").
geneOrder An ordered character vector of gene names to prioritize when labeling genes.
geneHighlights A two-column dataframe with a column named "gene" containing gene names as strings to highlight and a named column "color" containing corresponding highlight colors.
geneBackground If geneHighlights is given, a character value indicating the color for genes that are not highlighted.
strandLabels A logical value indicating whether to include + and - strand labels to the left of the gene track.
stroke A numeric value indicating the stroke width for gene body outlines. Default value is stroke = 0.1.
bg Character value indicating background color. Default value is bg = NA.
x A numeric or unit object specifying genes plot x-location.
y A numeric, unit object, or character containing a "b" combined with a numeric value specifying genes plot y-location. The character value will place the genes plot y relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.
width A numeric or unit object specifying genes plot width.
height A numeric or unit object specifying genes plot height.
just Justification of genes plot relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is just = c("left", "top").
default.units A string indicating the default units to use if x, y, width, or height are only given as numerics. Default value is default.units = "inches".
draw A logical value indicating whether graphics output should be produced. Default value is draw = TRUE.
params An optional pgParams object containing relevant function parameters.
Details

A gene track can be placed on a page by providing plot placement parameters:

```r
plotGenes(chrom, chromstart = NULL, chromend = NULL,
        x, y, width, height, just = c("left", "top"),
        default.units = "inches")
```

This function can be used to quickly plot an unnannotated gene track by ignoring plot placement parameters:

```r
plotGenes(chrom, chromstart = NULL, chromend = NULL)
```

Genomic annotation information is acquired through `TxDb` and `OrgDb-class` packages, as determined through the `assembly` parameter. To avoid overcrowding of gene name labels, plotted gene labels are by default prioritized according to citation counts.

Value

Returns a `genes` object containing relevant genomic region, placement, and `grob` information.

See Also

`assembly`, `genomes`, `defaultPackages`

Examples

```r
## Load hg19 genomic annotation packages
library("TxDb.Hsapiens.UCSC.hg19.knownGene")
library("org.Hs.eg.db")

## Set genomic coordinates
paramssmall <- pgParams(
    chrom = "chr8",
    chromstart = 1, chromend = 3000000,
    assembly = "hg19", width = 7
)
paramsbig <- pgParams(
    chrom = "chr8",
    chromstart = 1, chromend = 146364022,
    assembly = "hg19", width = 7
)

## Set colors
cols <- c("#41B6C4", "#225EA8")

## Create page
pageCreate(width = 7.5, height = 3.5, default.units = "inches")

## Plot genes big
genesisPlot <- plotGenes(
    params = paramsbig, fill = cols,
    fontcolor = cols,
```
plotGenomeLabel

Plot genomic coordinates along the x or y-axis of a plotgardener plot

Description

Plot genomic coordinates along the x or y-axis of a plotgardener plot

Usage

plotGenomeLabel(
  chrom,
  chromstart = NULL,
  chromend = NULL,
  assembly = "hg38",
  fontsize = 10,
  fontcolor = "black",
  linecolor = "black",
  x = 0.25, y = 0.25, height = 0.75,
  just = c("left", "top"),
  default.units = "inches"
)

## Annotate genome label
annoGenomeLabel(
  plot = genesPlot, x = 0.25, y = 1.0,
  scale = "Mb", just = c("left", "top")
)

## Plot genes small
genesPlot <- plotGenes(
  params = paramssmall,
  geneHighlights = data.frame(
    "gene" = c("DLGAP2"),
    "color" = c("#225EA8")
  ),
  geneBackground = "grey",
  x = 0.25, y = 2.25, height = 0.75,
  just = c("left", "top"), default.units = "inches"
)

## Annotate genome label
annoGenomeLabel(
  plot = genesPlot, x = 0.25, y = 3.0, scale = "Mb",
  just = c("left", "top")
)

## Hide page guides
pageGuideHide()
margin = unit(1, "mm"),
scale = "bp",
comas = TRUE,
sequence = TRUE,
boxWidth = 0.5,
axis = "x",
at = NULL,
tcl = 0.5,
x,
y,
length,
just = c("left", "top"),
default.units = "inches",
params = NULL,
...
)

Arguments

chrom  Chromosome of genome label, as a string, or a character vector of chromosomes for a whole genome Manhattan plot.
chromstart  Integer start of genome label.
chromend  Integer end of genome label.
assembly  Default genome assembly as a string or a assembly object.
fontsize  A numeric specifying text fontsize in points. Default value is fontsize = 10.
fontcolor  A character value indicating the color for text. Default value is fontcolor = "black".
linecolor  A character value indicating the color of the genome label axis. Default value is linecolor = "black".
margin  A numeric or unit vector specifying space between axis and coordinate labels. Default value is margin = unit(1, "mm"),
scale  A character value indicating the scale of the coordinates along the genome label. Default value is scale = "bp". Options are:
  • "bp": base pairs.
  • "Kb": kilobase pairs. 1 kilobase pair is equal to 1000 base pairs.
  • "Mb": megabase pairs. 1 megabase pair is equal to 1000000 base pairs.
comas  A logical value indicating whether to include commas in start and stop labels. Default value is comas = TRUE.
sequence  A logical value indicating whether to include sequence information above the label of an x-axis (only at appropriate resolutions).
boxWidth  A numeric value indicating the width of the boxes representing sequence information at appropriate resolutions. Default value is boxWidth = 0.5.
axis  A character value indicating along which axis to add genome label. Sequence information will not be displayed along a y-axis. Default value is axis = "x". Options are:
• "x": Genome label will be plotted along the x-axis.
• "y": Genome label will be plotted along the y-axis. This is typically used for a square Hi-C plot made with plotHicSquare.

at
A numeric vector of x-value locations for tick marks.
tcl
A numeric specifying the length of tickmarks as a fraction of text height. Default value is \( tcl = 0.5 \).
x
A numeric or unit object specifying genome label x-location.
y
A numeric, unit object, or character containing a "b" combined with a numeric value specifying genome label y-location. The character value will place the genome label y relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.

length
A numeric or unit object specifying length of genome label axis.
just
Justification of genome label relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is just = c("left", "top").

default.units
A string indicating the default units to use if x, y, or length are only given as numerics. Default value is default.units = "inches".

params
An optional pgParams object containing relevant function parameters.
...
Additional grid graphical parameters or digit specifications. See gpar and formatC.

Value

Returns a genomeLabel object containing relevant genomic region, placement, and grob information.

Examples

```r
## Load hg19 genomic annotation packages
library("TxDb.Hsapiens.UCSC.hg19.knownGene")
library("org.Hs.eg.db")
library("BSgenome.Hsapiens.UCSC.hg19")

## Create page
tCreate(width = 5, height = 3, default.units = "inches")

ePlot and place gene track on page
genPlot <- plotGenes(
    chrom = "chr8",
    chromstart = 1000000, chromend = 2000000,
    assembly = "hg19", fill = c("grey", "grey"),
    fontcolor = c("grey", "grey"),
    x = 0.5, y = 0.25, width = 4, height = 1,
    just = c("left", "top"),
    default.units = "inches"
)"
```
## Plot x-axis genome labels at different scales

```r
plotGenomeLabel(
  chrom = "chr8",
  chromstart = 1000000, chromend = 2000000,
  assembly = "hg19",
  scale = "Mb",
  x = 0.5, y = 1.25, length = 4, just = c("left", "top"),
  default.units = "inches"
)
plotGenomeLabel(
  chrom = "chr8",
  chromstart = 1000000, chromend = 2000000,
  assembly = "hg19",
  scale = "Kb",
  x = 0.5, y = 1.5, length = 4, just = c("left", "top"),
  default.units = "inches"
)
plotGenomeLabel(
  chrom = "chr8",
  chromstart = 1000000, chromend = 2000000,
  assembly = "hg19",
  scale = "bp",
  x = 0.5, y = 1.75, length = 4, just = c("left", "top"),
  default.units = "inches"
)
```

## Plot a different genomic label region, zooming in enough ## to see base pairs

```r
plotGenomeLabel(
  chrom = "chr8",
  chromstart = 1000000, chromend = 1000050,
  assembly = "hg19",
  x = 0.25, y = 2.2, length = 4.5
)
plotGenomeLabel(
  chrom = "chr8",
  chromstart = 1000000, chromend = 1000020,
  assembly = "hg19",
  x = 0, y = 2.6, length = 5
)
```

## Hide page guides

```r
pageGuideHide()
```

---

**plotGG**

*Plot a ggplot2 plot, gtable, or grob object in a plotgardener layout*

---

**Description**

Plot a ggplot2 plot, gtable, or grob object in a plotgardener layout.
Usage

plotGG(
  plot,
  x,
  y,
  width,
  height,
  just = c("left", "top"),
  default.units = "inches",
  params = NULL
)

Arguments

plot ggplot, gtable, or grob object.
x A numeric or unit object specifying ggplot x-location.
y A numeric, unit object, or character containing a "b" combined with a numeric value specifying ggplot y-location. The character value will place the ggplot y relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.
width A numeric or unit object specifying ggplot width.
height A numeric or unit object specifying ggplot height.
just Justification of ggplot relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is just = c("left", "top").
default.units A string indicating the default units to use if x, y, width, or height are only given as numerics. Default value is default.units = "inches".
params An optional pgParams object containing relevant function parameters.

Value

Returns a pg_gg object containing relevant placement and grob information.

See Also

ggplot

Examples

## Create a plot using ggplot2
library(ggplot2)
p <- ggplot(mtcars) +
  geom_point(aes(mpg, disp))

## Create a page
pageCreate(width = 4, height = 4, default.units = "inches")
## Place ggplot in page
plotGG(
    plot = p, x = 0.5, y = 0.5, width = 3, height = 3,
    just = c("left", "top"), default.units = "inches"
)

## Add title
plotText(
    label = "mtcars", fontsize = 14, fontface = "bold",
    x = 1, y = 0.35
)

## Hide page guides
pageGuideHide()

---

**plotHicRectangle**: Plot a triangular Hi-C interaction matrix in a rectangular format

### Description
Plot a triangular Hi-C interaction matrix in a rectangular format

### Usage
```r
plotHicRectangle(
    data,
    resolution = "auto",
    zrange = NULL,
    norm = "KR",
    matrix = "observed",
    chrom,
    chromstart = NULL,
    chromend = NULL,
    assembly = "hg38",
    palette = colorRampPalette(brewer.pal(n = 9, "YlGnBu")),
    colorTrans = "linear",
    flip = FALSE,
    bg = NA,
    x = NULL,
    y = NULL,
    width = NULL,
    height = NULL,
    just = c("left", "top"),
    default.units = "inches",
    draw = TRUE,
    params = NULL,
    quiet = FALSE
)
```
**Arguments**

- **data**
  Path to `.hic` file as a string or a 3-column dataframe of interaction counts in sparse upper triangular format.

- **resolution**
  A numeric specifying the width in basepairs of each pixel. For hic files, "auto" will attempt to choose a resolution based on the size of the region. For dataframes, "auto" will attempt to detect the resolution the dataframe contains.

- **zrange**
  A numeric vector of length 2 specifying the range of interaction scores to plot, where extreme values will be set to the max or min.

- **norm**
  Character value specifying hic data normalization method, if giving `.hic` file. This value must be found in the `.hic` file. Default value is `norm = "KR"`.

- **matrix**
  Character value indicating the type of matrix to output. Default value is `matrix = "observed"`. Options are:
  - "observed": Observed counts.
  - "oe": Observed/expected counts.
  - "log2oe": Log2 transformed observed/expected counts.

- **chrom**
  Chromosome of region to be plotted, as a string.

- **chromstart**
  Integer start position on chromosome to be plotted.

- **chromend**
  Integer end position on chromosome to be plotted.

- **assembly**
  Default genome assembly as a string or a `assembly` object. Default value is `assembly = "hg38"`.

- **palette**
  A function describing the color palette to use for representing scale of interaction scores. Default value is `palette = colorRampPalette(brewer.pal(n = 9, "YlGnBu"))`.

- **colorTrans**
  A string specifying how to scale Hi-C colors. Options are "linear", "log", "log2", or "log10". Default value is `colorTrans = "linear"`.

- **flip**
  A logical indicating whether to flip the orientation of the Hi-C matrix over the x-axis. Default value is `flip = FALSE`.

- **bg**
  Character value indicating background color. Default value is `bg = NA`.

- **x**
  A numeric or unit object specifying rectangle Hi-C plot x-location.

- **y**
  A numeric, unit object, or character containing a "b" combined with a numeric value specifying rectangle Hi-C plot y-location. The character value will place the rectangle Hi-C plot y relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.

- **width**
  A numeric or unit object specifying the width of the Hi-C plot rectangle.

- **height**
  A numeric or unit object specifying the height of the Hi-C plot rectangle.

- **just**
  Justification of rectangle Hi-C plot relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is `just = c("left", "top")`.

- **default.units**
  A string indicating the default units to use if x, y, width, or height are only given as numerics. Default value is `default.units = "inches"`. 
plotHicRectangle

draw A logical value indicating whether graphics output should be produced. Default value is draw = TRUE.

params An optional pgParams object containing relevant function parameters.

quiet A logical indicating whether or not to print messages.

Details

This function is similar to plotHicTriangle but will fill in additional pixels around the triangular portion of the plot to make a rectangle. The x-axis represents the genomic coordinates and the y-axis corresponds to distance in Hi-C bins.

A rectangle Hi-C plot can be placed on a plotgardener coordinate page by providing plot placement parameters:

plotHicRectangle(data, chrom,
    chromstart = NULL, chromend = NULL,
    x, y, width, height, just = c("left", "top"),
    default.units = "inches")

This function can also be used to quickly plot an unannotated rectangle Hi-C plot by ignoring plot placement parameters:

plotHicRectangle(data, chrom,
    chromstart = NULL, chromend = NULL)

Value

Returns a hicRectangle object containing relevant genomic region, Hi-C data, placement, and grob information.

See Also

readHic, plotHicTriangle

Examples

## Load Hi-C data
library(plotgardenerData)
data("IMR90_HiC_10kb")

## Create a page
pageCreate(width = 6, height = 3.5, default.units = "inches")

## Plot and place rectangle Hi-C plot
hicPlot <- plotHicRectangle(
    data = IMR90_HiC_10kb, resolution = 10000,
    zrange = c(0, 70),
    chrom = "chr21",
    chromstart = 28950000, chromend = 29800000,
    assembly = "hg19",
    x = 0.5, y = 0.5, width = 5, height = 2.5,
plotHicSquare

Plot a Hi-C interaction matrix in a square format

Description

Plot a Hi-C interaction matrix in a square format

Usage

plotHicSquare(
  data,
  resolution = "auto",
  zrange = NULL,
  norm = "KR",
  matrix = "observed",
  chrom,
  chromstart = NULL,
  chromend = NULL,
  altchrom = NULL,
  altchromstart = NULL,
  altchromend = NULL,
  assembly = "hg38",
  palette = colorRampPalette(brewer.pal(n = 9, "YlGnBu")),
  colorTrans = "linear",
  half = "both",
  yaxisDir = "increase",
  bg = NA,
  x = NULL,
  just = c("left", "top"),
  default.units = "inches"
)
plotHicSquare

```r
y = NULL,
width = NULL,
height = NULL,
just = c("left", "top"),
default.units = "inches",
draw = TRUE,
params = NULL,
quiet = FALSE
)
```

**Arguments**

- **data**
  Path to .hic file as a string or a 3-column dataframe of interaction counts in sparse upper triangular format.

- **resolution**
  A numeric specifying the width in basepairs of each pixel. For hic files, "auto" will attempt to choose a resolution based on the size of the region. For dataframes, "auto" will attempt to detect the resolution the dataframe contains.

- **zrange**
  A numeric vector of length 2 specifying the range of interaction scores to plot, where extreme values will be set to the max or min.

- **norm**
  Character value specifying hic data normalization method, if giving .hic file. This value must be found in the .hic file. Default value is `norm = "KR"`.

- **matrix**
  Character value indicating the type of matrix to output. Default value is `matrix = "observed"`. Options are:
  - "observed": Observed counts.
  - "oe": Observed/expected counts.
  - "log2oe": Log2 transformed observed/expected counts.

- **chrom**
  Chromosome of region to be plotted, as a string.

- **chromstart**
  Integer start position on chromosome to be plotted.

- **chromend**
  Integer end position on chromosome to be plotted.

- **altchrom**
  Alternate chromosome for off-diagonal plotting or interchromosomal plotting, as a string.

- **altchromstart**
  Alternate chromosome integer start position for off-diagonal plotting or interchromosomal plotting.

- **altchromend**
  Alternate chromosome integer end position for off-diagonal plotting or interchromosomal plotting.

- **assembly**
  Default genome assembly as a string or a `assembly` object. Default value is `assembly = "hg38"`.

- **palette**
  A function describing the color palette to use for representing scale of interaction scores. Default value is `palette = colorRampPalette(brewer.pal(n = 9, "YlGnBu"))`.

- **colorTrans**
  A string specifying how to scale Hi-C colors. Options are "linear", "log", "log2", or "log10". Default value is `colorTrans = "linear"`. 

A character value indicating which diagonal regions to plot. For intrachromosomal plotting, options are "both", "top", or "bottom". For off-diagonal or interchromosomal plotting, options are "top" or "bottom". Default value is half = "both".
- "both": Both diagonal halves.
- "top": Half above the diagonal.
- "bottom": Half below the diagonal.

A string specifying the genomic direction of the y-axis. Options are "increase", where the bottom left corner of the Hi-C map indicates the origin, and "decrease", where the top left corner of the Hi-C map indicates the origin. This will change the direction of the diagonal for intrachromosomal maps. Default value is yaxisDir = "increase".

Character value indicating background color. Default value is bg = NA.

A numeric or unit object specifying square Hi-C plot x-location.

A numeric, unit object, or character containing a "b" combined with a numeric value specifying square Hi-C plot y-location. The character value will place the square Hi-C plot y relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.

A numeric or unit object specifying square Hi-C plot width.

A numeric or unit object specifying square Hi-C plot height.

Justification of square Hi-C plot relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is just = c("left", "top").

A string indicating the default units to use if x, y, width, or height are only given as numerics. Default value is default.units = "inches".

A logical value indicating whether graphics output should be produced. Default value is draw = TRUE.

An optional pgParams object containing relevant function parameters.

A logical indicating whether or not to print messages.

A square Hi-C plot can be placed on a plotgardener coordinate page by providing plot placement parameters:

```r
plotHicSquare(data, chrom,
             chromstart = NULL, chromend = NULL,
             x, y, width, height, just = c("left", "top"),
             default.units = "inches")
```

This function can be used to quickly plot an unannotated square Hi-C plot by ignoring plot placement parameters:

```r
plotHicSquare(data, chrom,
             chromstart = NULL, chromend = NULL)
```
Value

Returns a hicSquare object containing relevant genomic region, Hi-C data, placement, and grob information.

See Also

readHic

Examples

## Load Hi-C data
library(plotgardenerData)
data("IMR90_HiC_10kb")

## Create a page
pageCreate(width = 3, height = 3, default.units = "inches")

## Plot and place Hi-C plot
hicPlot <- plotHicSquare(
  data = IMR90_HiC_10kb, resolution = 10000,
  zrange = c(0, 70),
  chrom = "chr21",
  chromstart = 28000000, chromend = 30300000,
  assembly = "hg19",
  x = 0.5, y = 0.5, width = 2, height = 2,
  just = c("left", "top"),
  default.units = "inches", bg = "black"
)

## Annotate heatmap legend
annoHeatmapLegend(
  plot = hicPlot, x = 2.6, y = 0.5,
  width = 0.12, height = 1.2,
  just = c("left", "top"), default.units = "inches"
)

## Annotate x-axis and y-axis genome labels
annoGenomeLabel(
  plot = hicPlot, scale = "Mb", axis = "x",
  x = 0.5, y = 2.53, just = c("left", "top")
)
annoGenomeLabel(
  plot = hicPlot, scale = "Mb", axis = "y",
  x = 0.47, y = 0.5, just = c("right", "top")
)

## Hide page guides
pageGuideHide()
plotHicTriangle 

Plot a Hi-C interaction matrix in a triangular format

Description

Plot a Hi-C interaction matrix in a triangular format

Usage

plotHicTriangle(
  data,
  resolution = "auto",
  zrange = NULL,
  norm = "KR",
  matrix = "observed",
  chrom,
  chromstart = NULL,
  chromend = NULL,
  assembly = "hg38",
  palette = colorRampPalette(brewer.pal(n = 9, "YlGnBu")),
  colorTrans = "linear",
  flip = FALSE,
  bg = NA,
  x = NULL,
  y = NULL,
  width = NULL,
  height = NULL,
  just = c("left", "top"),
  default.units = "inches",
  draw = TRUE,
  params = NULL,
  quiet = FALSE
)

Arguments

- **data** Path to .hic file as a string or a 3-column dataframe of interaction counts in sparse upper triangular format.
- **resolution** A numeric specifying the width in basepairs of each pixel. For hic files, "auto" will attempt to choose a resolution based on the size of the region. For dataframes, "auto" will attempt to detect the resolution the dataframe contains.
- **zrange** A numeric vector of length 2 specifying the range of interaction scores to plot, where extreme values will be set to the max or min.
- **norm** Character value specifying hic data normalization method, if giving .hic file. This value must be found in the .hic file. Default value is norm = "KR".
matrix | Character value indicating the type of matrix to output. Default value is matrix = "observed". Options are:
| "observed": Observed counts.
| "oe": Observed/expected counts.
| "log2oe": Log2 transformed observed/expected counts.

chrom | Chromosome of region to be plotted, as a string.

chromstart | Integer start position on chromosome to be plotted.

chromend | Integer end position on chromosome to be plotted.

assembly | Default genome assembly as a string or a assembly object. Default value is assembly = "hg38".

palette | A function describing the color palette to use for representing scale of interaction scores. Default value is palette = colorRampPalette(brewer.pal(n = 9, "YlGnBu")).

colorTrans | A string specifying how to scale Hi-C colors. Options are "linear", "log", "log2", or "log10". Default value is colorTrans = "linear".

flip | A logical indicating whether to flip the orientation of the Hi-C matrix over the x-axis. Default value is flip = FALSE.

bg | Character value indicating background color. Default value is bg = NA.

x | A numeric or unit object specifying triangle Hi-C plot x-location.

y | A numeric, unit object, or character containing a "b" combined with a numeric value specifying triangle Hi-C plot y-location. The character value will place the triangle Hi-C plot y relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.

width | A numeric or unit object specifying the bottom width of the Hi-C plot triangle.

height | A numeric or unit object specifying the height of the Hi-C plot triangle.

just | Justification of triangle Hi-C plot relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is just = c("left", "top").

default.units | A string indicating the default units to use if x, y, width, or height are only given as numerics. Default value is default.units = "inches".

draw | A logical value indicating whether graphics output should be produced. Default value is draw = TRUE.

params | An optional pgParams object containing relevant function parameters.

quiet | A logical indicating whether or not to print messages.

Details

In this orientation, the x-axis represents the genomic coordinates and the y-axis corresponds to distance in Hi-C bins.

A triangle Hi-C plot can be placed on a plotgardener coordinate page by providing plot placement parameters:
plotHicTriangle(data, chrom,
chromstart = NULL, chromend = NULL,
x, y, width, height, just = c("left", "top"),
default.units = "inches")

This function can also be used to quickly plot an unannotated triangle Hi-C plot by ignoring plot placement parameters:

plotHicTriangle(data, chrom,
chromstart = NULL, chromend = NULL)

If height is $<$ 0.5 * width, the top of the triangle will be cropped to the given height.

Value

Returns a hicTriangle object containing relevant genomic region, Hi-C data, placement, and grob information.

See Also

readHic

Examples

## Load Hi-C data
library(plotgardenerData)
data("IMR90_HiC_10kb")

## Create a page
df <- pageCreate(width = 4, height = 2.5, default.units = "inches")

## Plot and place triangle Hi-C plot
df <- plotHicTriangle(
data = IMR90_HiC_10kb, resolution = 10000,
zrange = c(0, 70),
chrom = "chr21",
chromstart = 28000000, chromend = 30300000,
assembly = "hg19",
x = 2, y = 0.5, width = 3, height = 1.5,
just = "top", default.units = "inches"
)

## Annotate x-axis genome label
df <- annoGenomeLabel(
plot = df, scale = "Mb", x = 0.5, y = 2.03,
just = c("left", "top")
)

## Annotate heatmap legend
df <- annoHeatmapLegend(
plot = df, x = 3.5, y = 0.5,
width = 0.13, height = 1.2,
## Hide page guides
pageGuideHide()

---

**plotIdeogram**

*Plot a chromosome ideogram with or without cytobands*

### Description

Plot a chromosome ideogram with or without cytobands

### Usage

```r
plotIdeogram(
  chrom,
  assembly = "hg38",
  data = NULL,
  orientation = "h",
  flip = FALSE,
  showBands = TRUE,
  fill = NULL,
  x = NULL,
  y = NULL,
  width = NULL,
  height = NULL,
  just = c("left", "top"),
  default.units = "inches",
  draw = TRUE,
  params = NULL
)
```

### Arguments

- **chrom**: Chromosome to be plotted, as a string.
- **assembly**: Default genome assembly as a string or a `assembly` object. Default value is `assembly = "hg38"`. (default value is `"hg38"`)  
- **data**: Custom cytoband data, as a dataframe with the following columns: "seqnames", "start", "end", "width", "strand", "name", "gieStain". (default value is `NULL`)  
- **orientation**: Character value indicating the orientation of the ideogram. Default value is `orientation = "h"`. Options are:
  - "v": Vertical ideogram orientation.
  - "h": Horizontal ideogram orientation. (default value is `"h"`)
Logical value indicating whether to reverse the start and end positions of the ideogram. If `flip = FALSE` (default), the ideogram starts at the top/left and ends at the bottom/right. If `flip = TRUE`, the start and end positions are reversed, meaning the ideogram starts at the bottom/right and ends at the top/left.

**Arguments:**

- `flip`: Logical value indicating whether to reverse the start and end positions of the ideogram. Default value is `flip = FALSE`.

- `showBands`: Logical value indicating whether to draw colored cytobands within ideogram. Default value is `showBands = TRUE`.

- `fill`: (optional) A vector specifying alternate colors for cytoband stains. To change specific gieStain values (i.e. gneg, gpos, etc.) to specific colors, this vector can be named. This vector must have the same number of colors as there are gieStain values for each genome assembly.

- `x`: A numeric or unit object specifying ideogram x-location.

- `y`: A numeric, unit object, or character containing a "b" combined with a numeric value specifying ideogram y-location. The character value will place the ideogram y relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.

- `width`: A numeric or unit object specifying ideogram width.

- `height`: A numeric or unit object specifying ideogram height.

- `just`: Justification of ideogram relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is `just = c("left", "top")`.

- `default.units`: A string indicating the default units to use if x, y, width, or height are only given as numerics. Default value is `default.units = "inches"`.

- `draw`: A logical value indicating whether graphics output should be produced. Default value is `draw = TRUE`.

- `params`: An optional `pgParams` object containing relevant function parameters.

**Details**

An ideogram can be placed on a plotgardener coordinate page by providing plot placement parameters:

```r
plotIdeogram(chrom, 
  x, y, width, height, just = c("left", "top"), 
  default.units = "inches")
```

This function can also be used to quickly plot an unannotated ideogram by ignoring plot placement parameters:

```r
plotIdeogram(chrom)
```

If no data is provided, Giemsa stain band data will first try to fetch UCSC with AnnotationHub. The results are cached for faster access, but these cached items can be deleted. If no internet connection is available and AnnotationHub has not previously cached the data, custom Giemsa stain band data can be loaded with the `data` parameter.
Value

Returns a ideogram object containing relevant genomic region, placement, and grob information.

See Also

AnnotationHub

Examples

```r
## Load Giemsa stain band information and genomic
## annotation data for hg19 genome assembly
library(TxDb.Hsapiens.UCSC.hg19.knownGene)
library(AnnotationHub)

## Create page
pageCreate(width = 4.5, height = 1, default.units = "inches")

## Plot and place ideogram
ideogramPlot <- plotIdeogram(
  chrom = "chr2", assembly = "hg19",
  x = 0.25, y = 0.25, width = 4, height = 0.3,
  just = c("left", "top"),
  default.units = "inches"
)

## Plot text
plotText(
  label = "Chromosome 2", fontcolor = "dark grey",
  x = 4.25, y = 0.65, just = "right"
)

## Hide page guides
pageGuideHide()
```

---

**plotLegend**

Plot a legend

**Description**

Plot a legend

**Usage**

```r
plotLegend(
  legend,
  fill = NULL,
  pch = NULL,
  lty = NULL,
  orientation = "v",
```
Arguments

legend A character or expression vector to appear in the legend.

fill If specified, this argument will produce boxes filled with the specified colors to appear beside the legend text.

pch The plotting symbols appearing in the legend, as a numeric vector.

lty The line types for lines appearing in the legend.

orientation A string specifying legend orientation. Default value is orientation = "v". Options are:
  • "v": Vertical legend orientation.
  • "h": Horizontal legend orientation.

title A character value giving a title to be placed at the top of the legend.

fontsize A numeric specifying text fontsize in points. Default value is fontsize = 10.

border Logical value indicating whether to add a border around heatmap legend. Default value is border = TRUE.

bg Character value indicating background color. Default value is bg = NA.

x A numeric or unit object specifying legend x-location.

y A numeric, unit object, or character containing a "b" combined with a numeric value specifying legend y-location. The character value will place the legend y relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.

width A numeric or unit object specifying legend width.

height A numeric or unit object specifying legend height.

just Justification of legend relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is just = c("left", "top").

default.units A string indicating the default units to use if x, y, width, or height are only given as numerics. Default value is default.units = "inches".
A logical value indicating whether graphics output should be produced. Default value is `draw = TRUE`.

An optional `pgParams` object containing relevant function parameters.

Additional grid graphical parameters. See `gpar`.

Returns a legend object containing relevant placement and `grob` information.

Examples

```r
## Load BED data
library(plotgardenerData)
data("IMR90_ChIP_CTCF_reads")

## Create page
pageCreate(width = 7.5, height = 4, default.units = "inches")

## Plot a pileup plot, coloring elements by strand
pileupPlot <- plotRanges(
data = IMR90_ChIP_CTCF_reads, chrom = "chr21",
chromstart = 29072500, chromend = 29075000,
assembly = "hg19",
fill = colorby("strand", palette =
  colorRampPalette(c("steel blue", "light salmon"))),
x = 0.5, y = 3.5, width = 6.5, height = 3.5,
just = c("left", "bottom"),
default.units = "inches"
)

## Add a legend depicting strand colors
legendPlot <- plotLegend(
  legend = c("- strand", "+ strand"),
  fill = c("steel blue", "light salmon"),
  border = FALSE,
x = 5, y = 0.5, width = 1.5, height = 0.7,
just = c("left", "top"),
default.units = "inches"
)

## Annotate genome label
annoGenomeLabel(
  plot = pileupPlot, x = 0.5, y = 3.5,
  just = c("left", "top")
)

## Hide page guides
pageGuideHide()
```
plotManhattan

Plot a Manhattan plot

Description

Plot a Manhattan plot

Usage

plotManhattan(
  data,
  sigVal = 5e-08,
  chrom = NULL,
  chromStart = NULL,
  chromEnd = NULL,
  assembly = "hg38",
  fill = "black",
  pch = 19,
  cex = 0.25,
  leadSNP = NULL,
  sigLine = FALSE,
  sigCol = NULL,
  trans = "-log10",
  range = NULL,
  yscale_reverse = FALSE,
  space = 0.01,
  bg = NA,
  baseline = FALSE,
  baseline.color = "grey",
  baseline.lwd = 1,
  x = NULL,
  y = NULL,
  width = NULL,
  height = NULL,
  just = c("left", "top"),
  flip = FALSE,
  default.units = "inches",
  draw = TRUE,
  params = NULL,
  ...
)

Arguments

data          Data to be plotted, as a character value specifying a file path of GWAS data, a
dataframe, or a GRanges object. Each of these data types must have the following columns:
• "chrom": Chromosome names. This column must be a character.
• "pos": Chromosomal position. This column must be an integer or numeric.
• "p": p-value or similar measure. This column must be numeric. Values will be converted according to the `trans` parameter.
• "snp"(optional): SNP name or rsid. This column should be a character.

sigVal
A numeric specifying the significance level of p-values. Along with data p-values, this value will be converted according to the `trans` parameter. Default value is `sigVal = 5e-08`.

chrom
Chromosome of region to be plotted, as a string. If left `NULL`, all chromosomes found in data will be plotted.

chromstart
Integer start position on chromosome to be plotted.

chromend
Integer end position on chromosome to be plotted.

assembly
Default genome assembly as a string or a `assembly` object. Default value is `assembly = "hg38"`.

fill
A single character value, a vector, or a `colorby` object specifying fill colors of data points. For a Manhattan plot with multiple chromosomes, a vector of colors will be used to color points of different chromosomes. Default value is `fill = "black"`.

pch
A numeric value or numeric vector specifying point symbols. If `colorby` object is supplied for `fill`, point symbols will be mapped to `colorby` values. Default value is `pch = 19`.

cex
A numeric indicating the amount by which points should be scaled relative to the default. Default value is `cex = 0.25`.

leadSNP
A list specifying the lead SNP in the desired region and any associated aesthetic features of the lead SNP data point and text label. The lead SNP should be specified as a character with the name slot "snp" in the list. Accepted lead SNP aesthetic features in the list include `fill`, `pch`, `cex`, `fontcolor`, and `fontsize`.

sigLine
Logical value indicating whether to draw a line at the significance level indicated with `sigVal`. Default value is `sigLine = FALSE`.

sigCol
Single character value specifying the color of significant data points.

trans
Character value specifying the transformation to apply to the "p" column plotted along the y-axis. For no transformation, set value to the empty character "". Default value is `trans = "-log10"`.

range
A numeric vector of length 2 specifying the y-range of p-values to plot (c(min, max)).

yscale_reverse
Logical value indicating whether to reverse the y-scale and order points from max to min.

space
A numeric value indicating the space between each chromosome as a fraction of the width of the plot, if plotting multiple chromosomes. Default value is `space = 0.01`.

bg
Character value indicating background color. Default value is `bg = NA`.

baseline
Logical value indicating whether to include a baseline along the x-axis. Default value is `baseline = FALSE`.
plotManhattan

baseline.color  Baseline color. Default value is baseline.color = "grey".
baseline.lwd    Baseline line width. Default value is baseline.lwd = 1.
x             A numeric or unit object specifying Manhattan plot x-location.
y             A numeric, unit object, or character containing a "b" combined with a numeric value specifying Manhattan plot y-location. The character value will place the Manhattan plot y relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.
width          A numeric or unit object specifying Manhattan plot width.
height         A numeric or unit object specifying Manhattan plot height.
just           Justification of Manhattan plot relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is just = c("left", "top").
flip           Logical value indicating whether to reflect Manhattan plot over the x-axis. Default value is flip = FALSE.
default.units  A string indicating the default units to use if x, y, width, or height are only given as numerics. Default value is default.units = "inches".
draw           A logical value indicating whether graphics output should be produced. Default value is draw = TRUE.
params         An optional pgParams object containing relevant function parameters.
...            Additional grid graphical parameters. See gpar.

Details

A Manhattan plot can be placed on a plotgardener coordinate page by providing plot placement parameters:

```r
plotManhattan(data,
              chrom = NULL,
              chromstart = NULL, chromend = NULL,
              x, y, width, height, just = c("left", "top"),
              default.units = "inches")
```

This function can also be used to quickly plot an unannotated Manhattan plot by ignoring plot placement parameters:

```r
plotManhattan(data,
              chrom = NULL,
              chromstart = NULL, chromend = NULL)
```

Value

Returns a manhattan object containing relevant genomic region, placement, and grob information.
Examples

```
## Load genomic assembly information
library("TxDb.Hsapiens.UCSC.hg19.knownGene")
## Load GWAS data
library(plotgardenerData)
data("hg19_insulin_GWAS")

## Create a page
pageCreate(width = 7.5, height = 4.5, default.units = "inches")

## Plot all GWAS data
manhattanPlot <- plotManhattan(
data = hg19_insulin_GWAS, assembly = "hg19",
fill = c("grey", "#37a7db"),
sigLine = TRUE,
trans = "-log10",
col = "grey", lty = 2, range = c(0, 14),
x = 0.5, y = 0, width = 6.5, height = 2,
just = c("left", "top"),
default.units = "inches"
)
## Annotate genome label
annoGenomeLabel(
plot = manhattanPlot, x = 0.5, y = 2, fontsize = 8,
just = c("left", "top"),
default.units = "inches"
)

plotText(
label = "Chromosome", fontsize = 8,
x = 3.75, y = 2.20, just = "center", default.units = "inches"
)

## Annotate y-axis
annoYaxis(
plot = manhattanPlot, at = c(0, 2, 4, 6, 8, 10, 12, 14),
axisLine = TRUE, fontsize = 8
)

## Plot y-axis label
plotText(
label = "-log10(p-value)", x = 0.15, y = 1, rot = 90,
fontsize = 8, fontface = "bold", just = "center",
default.units = "inches"
)

## Plot GWAS data zooming in on chromosome 11
## highlighting a lead SNP, and coloring by LD score
hg19_insulin_GWAS$LD <- as.numeric(hg19_insulin_GWAS$LD)
## Group LD column into LD ranges
hg19_insulin_GWAS <- as.data.frame(dplyr::group_by(hg19_insulin_GWAS,
LDgrp = cut(
```
hg19_insulin_GWAS$LD, c(0, 0.2, 0.4, 0.6, 0.8, 1)))

hg19_insulin_GWAS$LDgrp <- addNA(hg19_insulin_GWAS$LD)
leadSNP_p <- min(hg19_insulin_GWAS[which(hg19_insulin_GWAS$chrom == "chr11"), ]$p)
leadSNP <- hg19_insulin_GWAS[which(hg19_insulin_GWAS$p == leadSNP_p), ]$snp

chr11_manhattanPlot <- plotManhattan(
  data = hg19_insulin_GWAS, chrom = "chr11",
  chromstart = 60000000,
  chromend = 130000000,
  assembly = "hg19",
  fill = colorby("LDgrp",
    palette = colorRampPalette(c("#1f4297",
                              "#37a7db", "green",
                              "orange", "red", "grey"))),
  trans = "-log10",
  sigLine = TRUE, col = "grey",
  lty = 2, range = c(0, 16),
  leadSNP = list(
    snp = leadSNP,
    pch = 18,
    cex = 0.75,
    fill = "#7ecdbb",
    fontsize = 8
  ),
  x = 0.5, y = 2.5, width = 6.5,
  height = 1.5,
  just = c("left", "top"),
  default.units = "inches"
)

## Plot legend for LD scores
plotLegend(
  legend = c("LD Ref Var",
             paste(""0.4", "">", "r^2",
                   "", "">=", "0.2"),
             paste(""0.2", "">", "r^2",
                   "", "">="", "0"),
             "no LD data"),
  fill = c("#7ecdbb", "#37a7db", "#1f4297", "grey"), cex = 0.75,
  pch = c(18, 19, 19, 19), border = FALSE, x = 7, y = 2.5,
  width = 1.5, height = 0.6, just = c("right", "top"),
  default.units = "inches"
)

## Annotate genome label
annoGenomeLabel(
  plot = chr11_manhattanPlot, x = 0.5, y = 4.01,
## Plot multiple signal tracks in line with each other

**plotMultiSignal**

**Description**

Plot multiple signal tracks in line with each other

**Usage**

```r
plotMultiSignal(
  data,
  binSize = NA,
  binCap = TRUE,
  negData = FALSE,
  chrom,
  chromstart = NULL,
  chromend = NULL,
  assembly = "hg38",
  linecolor = "#37a7db",
  fill = NA,
  ymax = 1,
  range = NULL,
  scale = FALSE,
  label = NULL,
  bg = NA,
  baseline = TRUE,
  baseline.color = "grey",
  baseline.lwd = 1,
)```
plotMultiSignal

```
orientation = "h",
x = NULL,
y = NULL,
width = NULL,
height = NULL,
just = c("left", "top"),
gapdistance = .2,
default.units = "inches",
draw = TRUE,
params = NULL, ...
```

**Arguments**

- **data**: List of data to be plotted as character values specifying multiple bigwig file paths, dataframes in BED format, or GRanges objects with metadata column score.
- **binSize**: A numeric specifying the length of each data bin in basepairs. Default value is binSize = NA.
- **binCap**: A logical value indicating whether the function will limit the number of data bins to 8,000. Default value is binCap = TRUE.
- **negData**: A logical value indicating whether any of the data has both positive and negative scores and the y-axis of each signal track should be split. Default value is negData = FALSE.
- **chrom**: Chromosome of region to be plotted, as a string.
- **chromstart**: Integer start position on chromosome to be plotted.
- **chromend**: Integer end position on chromosome to be plotted.
- **assembly**: Default genome assembly as a string or a assembly object. Default value is assembly = "hg38".
- **linecolor**: A character value or vector of character values specifying the line color(s) outlining the signal tracks. Default value is linecolor = "#37a7db".
- **fill**: A character value or vector specifying the fill color(s) of the signal tracks. Default value is fill = NA.
- **ymax**: A numeric specifying the fraction of the max y-value to set as the height of each plot. Default value is ymax = 1.
- **range**: A numeric vector of length 2 specifying the y-range of data to plot (c(min, max)) in each signal track. If range = NULL, an optimal range for all signal tracks will be calculated.
- **scale**: A logical value indicating whether to include a data scale label in the top left corner of each plot. Default value is scale = FALSE.
- **label**: An optional character vector to conveniently add text labels to signal tracks. If scale = TRUE, the labels will be drawn in the top right of the signal tracks. Otherwise, the label will be drawn in the top left of the plot. For more customizable labels, use plotText.
- **bg**: Character value indicating background color. Default value is bg = NA.
plotMultiSignal

baseline Logical value indicating whether to include a baseline along the x-axis. Default value is baseline = TRUE.

baseline.color Baseline color. Default value is baseline.color = "grey".

baseline.lwd Baseline line width. Default value is baseline.lwd = 1.

orientation A string specifying signal track orientations. Default value is orientation = "h". Options are:

  • "v": Vertical signal track orientations, where signal tracks will be stacked from left to right.
  • "h": Horizontal signal track orientations, where signal tracks will be stacked from top to bottom.

x A numeric vector or unit object specifying the overall multisignal x-location.

y A numeric, unit object, or character containing a "b" combined with a numeric value specifying overall multisignal plot y-location. The character value will place the multisignal plot y relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.

width A numeric or unit object specifying overall multisignal plot width.

height A numeric or unit object specifying overall multisignal plot height.

just Justification of overall multisignal plot relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is just = c("left", "top").

gapdistance A numeric or unit object specifying space between plots. Default value is gapdistance = 0.2.

default.units A string indicating the default units to use if x or y are only given as numerics. Default value is default.units = "inches".

draw A logical value indicating whether graphics output should be produced. Default value draw = TRUE.

params An optional pgParams object containing relevant function parameters.

... Additional grid graphical parameters. See gpar.

Value

Returns a list of signal objects containing relevant genomic region, placement, and grob information for each signal track.

Examples

library("plotgardenerData")
data("GM12878_ChIP_CTCF_signal")
data("IMR90_ChIP_CTCF_signal")
data("GM12878_ChIP_H3K27ac_signal")
data("IMR90_ChIP_H3K27ac_signal")
library("RColorBrewer")
## List of multiple signal datasets

```r
signalList <- list(GM12878_ChIP_CTCF_signal, GM12878_ChIP_H3K27ac_signal, IMR90_ChIP_CTCF_signal, IMR90_ChIP_H3K27ac_signal)
```

## Create page

```r
pageCreate(width = 6.9, height = 3.5, default.units = "inches")
```

## Plot multiple signals

```r
multisignal <- plotMultiSignal(signalList, chrom = "chr21", chromstart = 28150000, chromend = 29150000, linecolor = c(brewer.pal(n = 9, "YlGnBu")[4], brewer.pal(n = 9, "YlGnBu")[5], brewer.pal(n = 9, "YlGnBu")[6], brewer.pal(n = 9, "YlGnBu")[7]), label = c("GM12878 CTCF", "GM12878 H3K27ac", "IMR90 CTCF", "IMR90 H3K27ac"), assembly = "hg19", x = 0.2, y = 0.2, width = 6.5, height = 3, default.units = "inches", gapdistance = 0.1)
```

## Plot genome label

```r
plotGenomeLabel(
  chrom = "chr21", chromstart = 28150000, chromend = 29150000, assembly = "hg19", scale = "Kb", x = 0.2, y = 3.25, length = 6.5, default.units = "inches"
)
```

## Hide page guides

```r
pageGuideHide()
```

---

### plotPairs

**Plot paired-end genomic range elements**

#### Description

Plot paired-end genomic range elements

#### Usage

```r
plotPairs(
  data, chrom, chromstart = NULL, chromend = NULL, assembly = "hg38",
)```
plotPairs

fill = "#1f4297",
linecolor = NA,
bg = NA,
boxHeight = unit(2, "mm"),
spaceWidth = 0.02,
spaceHeight = 0.3,
limitLabel = TRUE,
baseline = FALSE,
baseline.color = "grey",
baseline.lwd = 1,
x = NULL,
y = NULL,
width = NULL,
height = NULL,
just = c("left", "top"),
default.units = "inches",
draw = TRUE,
params = NULL,
...
)

Arguments

data A string specifying the BEDPE file path, a dataframe in BEDPE format specifying data to be plotted, or a GInteractions object.

chrom Chromosome of region to be plotted, as a string.

chromstart Integer start position on chromosome to be plotted.

chromend Integer end position on chromosome to be plotted.

assembly Default genome assembly as a string or a assembly object. Default value is assembly = "hg38".

fill A single character value, a vector, or a colorby object specifying fill colors of paired range elements. Default value is fill = "#1f4297".

linecolor A single character value, a vector, or a colorby object specifying the color of the lines outlining paired range elements. Default value is linecolor = NA. Special options include:
   • NA: No line color.
   • "fill": Same color as fill.

bg Character value indicating background color. Default value is bg = NA.

boxHeight A numeric or unit object specifying height of boxes at either end of paired range elements. Default value is boxHeight = unit(2, "mm").

spaceWidth A numeric specifying the width of spacing between paired range elements, as a fraction of the plot’s genomic range. Default value is spaceWidth = 0.02.

spaceHeight A numeric specifying the height of space between boxes of paired range elements on different rows. Default value is spaceHeight = 0.3.
plotPairs

limitLabel A logical value indicating whether to draw a "+" when not all elements can be plotted in the plotting space. Default value is limitLabel = TRUE.

baseline Logical value indicating whether to include a baseline along the x-axis. Default value is baseline = FALSE.

baseline.color Baseline color. Default value is baseline.color = "grey".

baseline.lwd Baseline line width. Default value is baseline.lwd = 1.

x A numeric or unit object specifying paired range plot x-location.

y A numeric, unit object, or character containing a "b" combined with a numeric value specifying paired range plot y-location. The character value will place the paired range plot y relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.

width A numeric or unit object specifying paired range plot width.

height A numeric or unit object specifying paired range plot height.

just Justification of paired range plot relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is just = c("left", "top").

default.units A string indicating the default units to use if x, y, width, or height are only given as numerics. Default value is default.units = "inches".

draw A logical value indicating whether graphics output should be produced.

params An optional pgParams object containing relevant function parameters.

... Additional grid graphical parameters. See gpar.

Details

#' A paired ranges plot can be placed on a plotgardener coordinate page by providing plot placement parameters:

plotPairs(data, chrom,
           chromstart = NULL, chromend = NULL,
           x, y, width, height, just = c("left", "top"),
           default.units = "inches")

This function can also be used to quickly plot an unannotated paired ranges plot by ignoring plot placement parameters:

plotPairs(data, chrom,
           chromstart = NULL, chromend = NULL)

Value

Returns a pairs object containing relevant genomic region, placement, and grob information.
Examples

```r
## Load paired ranges data in BEDPE format
library(plotgardenerData)
data("IMR90_DNAloops_pairs")

## Set the coordinates
params <- pgParams(
  chrom = "chr21",
  chromstart = 27900000, chromend = 30700000,
  assembly = "hg19",
  width = 7
)

## Create a page
pageCreate(width = 7.5, height = 2.1, default.units = "inches")

## Add a length column
IMR90_DNAloops_pairs$length <-
  (IMR90_DNAloops_pairs$start2 - IMR90_DNAloops_pairs$start1) / 1000

## Plot the data
bedpePlot <- plotPairs(
  data = IMR90_DNAloops_pairs, params = params,
  fill = colorby("length", palette =
    colorRampPalette(c("dodgerblue2", "firebrick2"))),
  lwd = 2, spaceHeight = .7,
  x = 0.25, y = 0.25, height = 1.5,
  just = c("left", "top"), default.units = "inches"
)

## Annotate genome label
annoGenomeLabel(plot = bedpePlot, x = 0.25, y = 1.78, scale = "Mb")

## Add heatmap legend
annoHeatmapLegend(
  plot = bedpePlot, fontcolor = "black",
  x = 7.0, y = 0.25,
  width = 0.10, height = 1, fontsize = 10
)

## Add heatmap legend label
plotText(
  label = "Kb", rot = 90, x = 6.9, y = 0.75,
  just = c("center", "center"), fontsize = 10
)

## Hide page guides
pageGuideHide()
```

plotPairsArches  
Plot paired-end genomic range data in an arch style
**Description**

Plot paired-end genomic range data in an arch style

**Usage**

```r
plotPairsArches(
  data,
  chrom,
  chromstart = NULL,
  chromend = NULL,
  assembly = "hg38",
  style = "2D",
  flip = FALSE,
  curvature = 5,
  archHeight = NULL,
  fill = "#1f4297",
  linecolor = NA,
  alpha = 0.4,
  bg = NA,
  clip = FALSE,
  clip.noAnchor = TRUE,
  range = NULL,
  baseline = FALSE,
  baseline.color = "grey",
  baseline.lwd = 1,
  x = NULL,
  y = NULL,
  width = NULL,
  height = NULL,
  just = c("left", "top")
)
```

**Arguments**

- `data` A string specifying the BEDPE file path, a dataframe in BEDPE format specifying data to be plotted, or a `GInteractions` object.
- `chrom` Chromosome of region to be plotted, as a string.
- `chromstart` Integer start position on chromosome to be plotted.
- `chromend` Integer end position on chromosome to be plotted.
- `assembly` Default genome assembly as a string or a `assembly` object. Default value is `assembly = "hg38"`.
- `style` Character value describing the style of arches. Default value is `style = "2D"`. Options are:
plotPairsArches

- "2D": Arches will be drawn in a 2-dimensional style.
- "3D": Arches will be drawn in a 3-dimensional style.

flip

Logical value indicating whether to reflect arches over the x-axis. Default value is flip = FALSE.

curvature

Numeric indicating the number of points along the arch curvature. Default value is curvature = 5.

archHeight

Single numeric value, numeric vector, or column name in data specifying the arch heights. When NULL, all arches will be the same height, filling up the given plot area.

fill

A single character value, a vector, or a colorby object specifying fill colors of arches. Default value is fill = #1f4297.

linecolor

A single character value, a vector, or a colorby object specifying the color of the lines outlining arches. Default value is linecolor = NA. Special options include:
- NA: No line color.
- "fill": Same color as fill.

alpha

Numeric value specifying transparency. Default value is alpha = 0.4.

text

Character value indicating background color. Default value is bg = NA.

clip

A logical value indicating whether to clip any arches that get cutoff in the given genomic region. Default value is clip = FALSE.

clip.noAnchor

A logical value indicating whether to clip any arches that overlap the given genomic region but do not have an anchor in that region. Default value is clip.noAnchor = TRUE.

range

A numeric vector of length 2 specifying the y-range of archHeight to plot (c(min, max)).

baseline

Logical value indicating whether to include a baseline along the x-axis. Default value is baseline = FALSE.

baseline.color

Baseline color. Default value is baseline.color = "grey".

baseline.lwd

Baseline line width. Default value is baseline.lwd = 1.

x

A numeric or unit object specifying pair arches plot x-location.

y

A numeric, unit object, or character containing a "b" combined with a numeric value specifying BEDPE arches plot y-location. The character value will place the pair arches plot y relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.

width

A numeric or unit object specifying pair arches plot width.

height

A numeric or unit object specifying pair arches plot height.

just

Justification of pair arches plot relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is just = c("left", "top").

default.units

A string indicating the default units to use if x, y, width, or height are only given as numerics. Default value is default.units = "inches".
draw A logical value indicating whether graphics output should be produced. Default value is `draw = TRUE`.
params An optional `pgParams` object containing relevant function parameters.
... Additional grid graphical parameters. See `gpar`.

Details
A pair arches plot can be placed on a plotgardener coordinate page by providing plot placement parameters:

```r
plotPairsArches(data, chrom, chromstart = NULL, chromend = NULL, x, y, width, height, just = c("left", "top"), default.units = "inches")
```

This function can also be used to quickly plot an unannotated pair arches plot by ignoring plot placement parameters:

```r
plotPairsArches(data, chrom, chromstart = NULL, chromend = NULL)
```

Value
Returns a `arches` object containing relevant genomic region, placement, and `grob` information.

Examples
```r
## Load paired ranges data in BEDPE format
library(plotgardenerData)
data("IMR90_DNAloops_pairs")

## Set the coordinates
params <- pgParams(
    chrom = "chr21",
    chromstart = 27900000, chromend = 30700000,
    assembly = "hg19",
    width = 7
)

## Create a page
pageCreate(width = 7.5, height = 2.1, default.units = "inches")

## Add a length column to color by
IMR90_DNAloops_pairs$length <-
    (IMR90_DNAloops_pairs$start2 - IMR90_DNAloops_pairs$start1) / 1000

## Translate lengths into heights
IMR90_DNAloops_pairs$h <-
    IMR90_DNAloops_pairs$length / max(IMR90_DNAloops_pairs$length)

## Plot the data
archPlot <- plotPairsArches(
  data = IMR90_DNAloops_pairs, params = params,
  fill = colorby("length", palette =
    colorRampPalette(c("dodgerblue2", "firebrick2"))),
  linecolor = "fill",
  archHeight = "h", alpha = 1,
  x = 0.25, y = 0.25, height = 1.5,
  just = c("left", "top"),
  default.units = "inches"
)

## Annotate genome label
annoGenomeLabel(plot = archPlot, x = 0.25, y = 1.78, scale = "Mb")

## Annotate heatmap legend
annoHeatmapLegend(
  plot = archPlot, fontcolor = "black",
  x = 7.0, y = 0.25,
  width = 0.10, height = 1, fontsize = 10
)

## Add the heatmap legend title
plotText(
  label = "Kb", rot = 90, x = 6.9, y = 0.75,
  just = c("center", "center"),
  fontsize = 10
)

## Hide page guides
pageGuideHide()

---

**plotPolygon**

Plot a polygon within a plotgardener layout

---

**Description**

Plot a polygon within a plotgardener layout

**Usage**

```r
plotPolygon(
  x,
  y,
  default.units = "inches",
  linecolor = "black",
  lwd = 1,
  lty = 1,
  fill = NA,
```

---

```r
```
Arguments

- **x**: A numeric vector or unit object specifying polygon vertex x-locations.
- **y**: A numeric vector, unit object, or a character vector of values containing a "b" combined with a numeric value specifying polygon vertex y-locations. The character vector will place polygon vertex y-locations relative to the bottom of the most recently plotted plot according to the units of the plotgardeener page.
- **default.units**: A string indicating the default units to use if x or y are only given as numeric vectors. Default value is default.units = "inches".
- **linecolor**: A character value specifying polygon line color. Default value is linecolor = "black".
- **lwd**: A numeric specifying polygon line width. Default value is lwd = 1.
- **lty**: A numeric specifying polygon line type. Default value is lty = 1.
- **fill**: A character value specifying polygon fill color. Default value is fill = NA.
- **alpha**: Numeric value specifying color transparency. Default value is alpha = 1.
- **id**: A numeric vector used to separate locations in x and y into multiple polygons. All locations with the same id belong to the same polygon.
- **id.lengths**: A numeric vector used to separate locations in x and y into multiple polygons. Specifies consecutive blocks of locations which make up separate polygons.
- **params**: An optional pgParams object containing relevant function parameters.
- **...**: Additional grid graphical parameters. See gpar.

Value

Returns a polygon object containing relevant placement and grob information.

See Also

- grid polygon

Examples

```r
## Create a page
age = pageCreate(width = 7.5, height = 6, default.units = "inches")

## Plot complex polygons one at a time
plotPolygon(
  x = c(2.6, 4.65, 4.75, 6.05, 1.4, 1.3),
  y = c(2.5, 3.1, 3.5, 4, 3.15, 2.8),
  fill = "#4a168e", linecolor = NA
)
```
plotRanges

Plot genomic range elements in a pileup or collapsed format

Description

Plot genomic range elements in a pileup or collapsed format

Usage

plotRanges(
  data,
  chrom,
  chromstart = NULL,
  chromend = NULL,
  assembly = "hg38",
  fill = "#7ecdff",
  linecolor = NA,
  order = "width",
  collapse = FALSE,
  boxHeight = unit(2, "mm"),
)
Arguments

data      Data to be plotted; as a character value specifying a BED file path, a data frame in BED format, a character value specifying a .bam file path where a bam index file (.bam.bai) is in the same directory, or a GRanges object.

chrom     Chromosome of region to be plotted, as a string.

chromstart Integer start position on chromosome to be plotted.

chromend  Integer end position on chromosome to be plotted.

assembly  Default genome assembly as a string or a assembly object. Default value is assembly = "hg38".

fill      A single character value, a vector, or a colorby object specifying fill colors of range elements. Default value is fill = "#7ecdbb".

linecolor A single character value, a vector, or a colorby object specifying the color of the lines outlining range elements. Default value is linecolor = NA. Special options include:
• NA: No line color.
• "fill": Same color as fill.

order     A character value specifying how to order pileup data before assigning rows. Default value is order = "width". Options include:
• "width": Ordered by decreasing width of elements.
• "random": Ordered randomly in each function call.

collapse  A logical value indicating whether to collapse range elements into a single row, or into two rows if strandSplit = TRUE. If collapse = TRUE, boxHeight will be ignored and elements will be the height of the entire plot if strandSplit =
plotRanges

FALSE or be the height of half of the entire plot if strandSplit = TRUE. Default value is collapse = FALSE.

boxHeight A numeric or unit object specifying height of range element boxes. Default value is boxHeight = unit(2, "mm").

spaceWidth A numeric value specifying the width of minimum spacing between range element boxes, as a fraction of the plot’s genomic range. Default value is spaceWidth = 0.02.

spaceHeight A numeric value specifying the height of spacing between range element boxes on different rows, as a fraction of boxHeight. Default value is spaceHeight = 0.3.

limitLabel A logical value indicating whether to draw a "+" when not all elements can be plotted in the plotting space. Default value is limitLabel = TRUE.

strandSplit A logical value indicating whether plus and minus-stranded elements should be separated. Elements can only be split by strand if a strand column is found in data. Default value is strandSplit = FALSE.

bg Character value indicating background color. Default value is bg = NA.

baseline Logical value indicating whether to include a baseline along the x-axis. Default value is baseline = FALSE.

baseline.color Baseline color. Default value is baseline.color = "grey".

baseline.lwd Baseline line width. Default value is baseline.lwd = 1.

x A numeric or unit object specifying ranges plot x-location.

y A numeric, unit object, or character containing a "b" combined with a numeric value specifying ranges plot y-location. The character value will place the ranges plot y relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.

width A numeric or unit object specifying ranges plot width.

height A numeric or unit object specifying ranges plot height.

just Justification of ranges plot relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is just = c("left", "top").

default.units A string indicating the default units to use if x, y, width, or height are only given as numerics. Default value is default.units = "inches".

draw A logical value indicating whether graphics output should be produced. Default value draw = TRUE.

params An optional pgParams object containing relevant function parameters.

... Additional grid graphical parameters. See gpar.

Details

A ranges plot can be placed on a plotgardener coordinate page by providing plot placement parameters:
plotRanges(data, chrom,
    chromstart = NULL, chromend = NULL,
    x, y, width, height, just = c("left", "top"),
    default.units = "inches")

This function can also be used to quickly plot an unannotated BED plot by ignoring plot placement parameters:

plotRanges(data, chrom,
    chromstart = NULL, chromend = NULL)

Value

Returns a ranges object containing relevant genomic region, coloring data, placement, and grob information.

Examples

## Load ranges data in BED format
library(plotgardenerData)
data("IMR90_ChIP_CTCF_reads")

## Create page
pageCreate(width = 7.5, height = 5, default.units = "inches")

## Plot and place a pileup ranges plot
pileupPlot <- plotRanges(
    data = IMR90_ChIP_CTCF_reads, chrom = "chr21",
    chromstart = 29073000, chromend = 29074000,
    assembly = "hg19",
    order = "random",
    fill = colorby("strand", palette =
        colorRampPalette(c("#7ecdbb", "#37a7db")),
        strandSplit = TRUE,
    x = 0.5, y = 0.25, width = 6.5, height = 4.25,
    just = c("left", "top"), default.units = "inches"
)

## Annotate genome label
annoGenomeLabel(
    plot = pileupPlot, x = 0.5, y = 4.5,
    just = c("left", "top")
)

## Add text labels
plotText(
    label = "+ strand", fontcolor = "#37a7db", fontsize = 12,
    x = 0.5, y = 1.25, just = "left"
)
plotText(
    label = "- strand", fontcolor = "#7ecdbb", fontsize = 12,
    x = 0.5, y = 3.5, just = "left"
## Hide page guides
pageGuideHide()

---

plotRaster **Plot a raster object within a plotgardener layout**

### Description

Plot a raster object within a plotgardener layout

### Usage

```r
plotRaster(
  image,  
  x,      
  y,      
  width,  
  height, 
  just = "center", 
  default.units = "inches", 
  interpolate = TRUE, 
  params = NULL, 
  ...
)
```

### Arguments

- **image**: Any R object that can be coerced to a raster object.
- **x**: A numeric vector or unit object specifying raster x-locations.
- **y**: A numeric vector, unit object, or a character vector of values containing a "b" combined with a numeric value specifying raster y-locations. The character vector will place raster y relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.
- **width**: A numeric vector or unit object specifying raster widths.
- **height**: A numeric vector or unit object specifying raster heights.
- **just**: Justification of text relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is just = "center".
- **default.units**: A string indicating the default units to use if x, y, width, or height are only given as numerics or numeric vectors. Default value is default.units = "inches".
- **interpolate**: A logical value indicating whether to linearly interpolate the image. Default value is interpolate = TRUE.
- **params**: An optional `pgParams` object containing relevant function parameters.
- **...**: Additional grid graphical parameters. See `gpar`.
Value

Returns a raster object containing relevant placement and grob information.

See Also

grid.raster

Examples

library(png)

## Load images

pg_type <- readPNG(system.file("images", "pg-wordmark.png", package = "plotgardener"))

gene_gnome <- readPNG(system.file("images", "pg-gnome-hole-shadow.png", package = "plotgardener"))

rlogo <- readPNG(system.file("images", "Rlogo.png", package = "plotgardener"))

## Create page
pageCreate(width = 5, height = 6)

## Plot images
plotRaster(
  image = pg_type,
  x = 2.5, y = 0.25, width = 4, height = 1.5,
  just = "top"
)

plotRaster(
  image = gene_gnome,
  x = 2.5, y = 2.25, width = 3.5, height = 3.5,
  just = "top"
)

plotRaster(
  image = rlogo,
  x = 1, y = 1.5, width = 0.5, height = 0.45,
  just = c("left", "top")
)

## Hide page guies
pageGuideHide()
**Description**

Plot a rectangle within a plotgardener layout

**Usage**

```r
plotRect(
  x,  
  y,  
  width,  
  height,  
  just = "center",  
  default.units = "inches",  
  linecolor = "black",  
  lwd = 1,  
  lty = 1,  
  fill = NA,  
  alpha = 1,  
  params = NULL,  
  ...  
)
```

**Arguments**

- `x` A numeric vector or unit object specifying rectangle x-locations.
- `y` A numeric vector, unit object, or a character vector of values containing a "b" combined with a numeric value specifying rectangle y-locations. The character vector will place rectangle y-locations relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.
- `width` A numeric vector or unit object specifying rectangle widths.
- `height` A numeric vector or unit object specifying rectangle heights.
- `just` Justification of rectangle relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is `just = "center"`.
- `default.units` A string indicating the default units to use if `x`, `y`, `width`, and `height` are only given as numerics or numeric vectors. Default value is `default.units = "inches"`.
- `linecolor` A character value specifying rectangle line color. Default value is `linecolor = "black"`.
- `lwd` A numeric specifying rectangle line width. Default value is `lwd = 1`.
- `lty` A numeric specifying rectangle line type. Default value is `lty = 1`.
plotRect

fill  A character value specifying rectangle fill color. Default value is fill = NA.
alpha Numeric value specifying color transparency. Default value is alpha = 1.
params An optional pgParams object containing relevant function parameters.
... Additional grid graphical parameters. See gpar.

Value

Returns a rect object containing relevant placement and grob information.

See Also

grid.rect

Examples

## Create a page
pageCreate(width = 7.5, height = 6, default.units = "inches")

## Plot one rectangle with no fill
plotRect(
  x = 0.5, y = 0.5, width = 3, height = 3,
  just = c("left", "top"), default.units = "inches",
  lwd = 2, fill = NA
)

## Plot two rectangles with same width and height at different locations
plotRect(
  x = 4, y = c(0.5, 2.25), width = 3, height = 1.25,
  just = c("left", "top"), default.units = "inches",
  fill = "#7ecdbb"
)

## Plot two rectangles with different widths, heights, locations, and colors
## Plot two rectangles with different widths, heights, locations, and colors
plotRect(
  x = 3.75, y = c(4, 5.25), width = c(6.5, 4.5),
  height = c(1, 0.25),
  just = "top", default.units = "inches",
  fill = c("#7ecdbb", "#37a7db"), linecolor = NA, alpha = 0.4
)

## Hide page guides
pageGuideHide()
plotSegments

Draw a line segment within a plotgardener layout

Description

Draw a line segment within a plotgardener layout

Usage

```r
plotSegments(
  x0,
  y0,
  x1,
  y1,
  default.units = "inches",
  linecolor = "black",
  lwd = 1,
  lty = 1,
  lineend = "butt",
  linejoin = "mitre",
  arrow = NULL,
  params = NULL,
  ...
)
```

Arguments

- **x0**: A numeric vector or unit object indicating the starting x-values of the line segments.
- **y0**: A numeric vector, unit object, or a character vector of values containing a "b" combined with a numeric value specifying starting y-values of the line segments. The character vector will place starting y-values relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.
- **x1**: A numeric vector or unit object indicating the stopping x-values of the line segments.
- **y1**: A numeric vector, unit object, or a character vector of values containing a "b" combined with a numeric value specifying stopping y-values of the line segments. The character vector will place stopping y-values relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.
- **default.units**: A string indicating the default units to use if x0, y0, x1, or y1 are only given as numeric vectors. Default value is default.units = "inches".
- **linecolor**: A character value specifying segment line color. Default value is linecolor = "black".
- **lwd**: A numeric specifying segment line width. Default value is lwd = 1.
- **lty**: A numeric specifying segment line type. Default value is lty = 1.
plotSegments

lineend A character value specifying line end style. Default value is lineend = "butt". Options are:
  • "round": Segment ends are rounded.
  • "butt": Segment ends end exactly where ended.
  • "square": Segment ends are squared.

linejoin A character value specifying line join style. Default value is linejoin = "mitre". Options are:
  • "round": Line joins are rounded.
  • "mitre": Line joins are sharp corners.
  • "bevel": Line joins are flattened corners.

arrow A list describing arrow heads to place at either end of the line segments, as produced by the arrow function.

params An optional pgParams object containing relevant function parameters.

... Additional grid graphical parameters. See gpar.

Value

Returns a segments object containing relevant placement and grob information.

See Also

grid.segments, arrow

Examples

library(grid)
## Create a page
pageCreate(width = 7.5, height = 6, default.units = "inches")

## Plot one line segment
plotSegments(
  x0 = 3.75, y0 = 0.25, x1 = 3.75, y1 = 5.75,
  default.units = "inches",
  lwd = 3, lty = 2
)

## Plot multiple line segments at different locations in different colors
plotSegments(
  x0 = 0.5, y0 = c(1, 3, 5), x1 = 3.25, y1 = c(1, 3, 5),
  default.units = "inches",
  lwd = 2, linecolor = c("#7ecdbb", "#37a7db", "grey")
)

## Plot a line segment with an arrowhead
plotSegments(
  x0 = 4.5, y0 = 0.5, x1 = 7, y1 = 3,
  default.units = "inches",
  arrow = arrow(type = "closed"), fill = "black"
)
## Plot lines with round lineends

```r
plotSegments(
    x0 = c(4, 7), y0 = 3.5, x1 = 5.5, y1 = 4.5,
    default.units = "inches",
    lwd = 5, lineend = "round"
)
```

## Hide page guides

```r
pageGuideHide()
```

---

**plotSignal**  
*Plot any kind of signal track data for a single chromosome*

### Description

Plot any kind of signal track data for a single chromosome

### Usage

```r
plotSignal(
    data,
    binSize = NA,
    binCap = TRUE,
    negData = FALSE,
    chrom,
    chromstart = NULL,
    chromend = NULL,
    assembly = "hg38",
    linecolor = "#37a7db",
    fill = NA,
    ymax = 1,
    range = NULL,
    scale = FALSE,
    label = NULL,
    bg = NA,
    baseline = TRUE,
    baseline.color = "grey",
    baseline.lwd = 1,
    orientation = "h",
    x = NULL,
    y = NULL,
    width = NULL,
    height = NULL,
    just = c("left", "top"),
    default.units = "inches",
    draw = TRUE,
    params = NULL,
)```
Arguments

data  Data to be plotted as a character value specifying a bigwig file path, a dataframe in BED format, or a GRanges object with metadata column score. Either one data argument or a list of two can be provided, where the second data will be plotted below the x-axis if positive. The second data can also be negative data.

binSize  A numeric specifying the length of each data bin in basepairs. Default value is binSize = NA.

binCap  A logical value indicating whether the function will limit the number of data bins to 8,000. Default value is binCap = TRUE.

negData  A logical value indicating whether the provided data has negative scores. Default value is negData = FALSE.

chrom  Chromosome of region to be plotted, as a string.

chromstart  Integer start position on chromosome to be plotted.

chromend  Integer end position on chromosome to be plotted.

assembly  Default genome assembly as a string or a assembly object. Default value is assembly = "hg38".

linecolor  A character value or vector of length 2 specifying the line color(s) outlining the signal track(s). Default value is linecolor = "#37a7db".

fill  A character value or vector of length 2 specifying the fill color(s) of the signal track(s). Default value is fill = NA.

ymax  A numeric specifying the fraction of the max y-value to set as the height of the plot. Default value is ymax = 1.

range  A numeric vector of length 2 specifying the y-range of data to plot (c(min, max)).

scale  A logical value indicating whether to include a data scale label in the top left corner of the plot. Default value is scale = FALSE.

label  An optional character value to conveniently add a text label to the plot. If scale = TRUE, the label will be draw in the top right of the plot. Otherwise, the label will be drawn in the top left of the plot. For more customizable labels, use plotText. Default value is label = NULL.

bg  Character value indicating background color. Default value is bg = NA.

baseline  Logical value indicating whether to include a baseline along the x-axis. Default value is baseline = TRUE.

baseline.color  Baseline color. Default value is baseline.color = "grey".

baseline.lwd  Baseline line width. Default value is baseline.lwd = 1.

orientation  A string specifying signal track orientation. Default value is orientation = "h". Options are:
  • "v": Vertical signal track orientation.
  • "h": Horizontal signal track orientation.
A signal track can be placed on a plotgardener coordinate page by providing plot placement parameters:

```r
plotSignal(data, chrom, 
  chromstart = NULL, chromend = NULL,
  x, y, width, height, just = c("left", "top"),
  default.units = "inches")
```

This function can also be used to quickly plot an unannotated signal track by ignoring plot placement parameters:

```r
plotSignal(data, chrom, 
  chromstart = NULL, chromend = NULL)
```

Returns a signal object containing relevant genomic region, placement, and grob information.

### Examples

```r
## Load signal data
library(plotgardenerData)
data("IMR90_ChIP_H3K27ac_signal")
data("GM12878_ChIP_H3K27ac_signal")

## Create a page
pageCreate(width = 7.5, height = 2.1, default.units = "inches")
```
## Define region
region <- pgParams(
  chrom = "chr21",
  chromstart = 28000000, chromend = 30300000,
  assembly = "hg19",
  range = c(0, 45)
)

## Plot and place signal plots
signal1 <- plotSignal(
  data = IMR90_ChIP_H3K27ac_signal, params = region,
  x = 0.5, y = 0.25, width = 6.5, height = 0.65,
  just = c("left", "top"), default.units = "inches"
)

signal2 <- plotSignal(
  data = GM12878_ChIP_H3K27ac_signal, params = region,
  linecolor = "#7ecdbb",
  x = 0.5, y = 1, width = 6.5, height = 0.65,
  just = c("left", "top"), default.units = "inches"
)

## Plot genome label
plotGenomeLabel(
  chrom = "chr21",
  chromstart = 28000000, chromend = 30300000,
  assembly = "hg19",
  x = 0.5, y = 1.68, length = 6.5,
  default.units = "inches"
)

## Add text labels
plotText(
  label = "IMR90", fonsize = 10, fontcolor = "#37a7db",
  x = 0.5, y = 0.25, just = c("left", "top"),
  default.units = "inches"
)

plotText(
  label = "GM12878", fonsize = 10, fontcolor = "#7ecdbb",
  x = 0.5, y = 1, just = c("left", "top"),
  default.units = "inches"
)

## Hide page guides
pageGuideHide()
Description

Plot text within a plotgardener layout

Usage

plotText(
  label,
  fontcolor = "black",
  fontsize = 12,
  rot = 0,
  check.overlap = FALSE,
  x,
  y,
  just = "center",
  default.units = "inches",
  params = NULL,
  ...
)

Arguments

label Character or expression of text to be plotted.
fontcolor A character value specifying text fontcolor. Default value is fontcolor = "black".
fontsize A numeric specifying text fontsize in points. Default value is fontsize = 12.
rot A numeric specifying the angle to rotate the text. Default value is rot = 0.
check.overlap A logical value to indicate whether to check for and omit overlapping text. Default value is check.overlap = FALSE.
x A numeric vector or unit object specifying text x-location.
y A numeric vector, unit object, or a character vector of values containing a "b" combined with a numeric value specifying text y-locations. The character vector will place text y-locations relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.
just Justification of text relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is just = "center".
default.units A string indicating the default units to use if x or y are only given as numerics or numeric vectors. Default value is default.units = "inches".
params An optional pgParams object containing relevant function parameters.
... Additional grid graphical parameters. See gpar.

Value

Returns a text object containing relevant placement and grob information.
plotTranscripts

Plot gene transcripts in a pileup style for a single chromosome

Description

Plot gene transcripts in a pileup style for a single chromosome

Usage

plotTranscripts(
  chrom,
  chromstart = NULL,
  chromend = NULL,
```r
assembly = "hg38",
fill = c("#669fd9", "#abcc8e"),
colorbyStrand = TRUE,
strandSplit = FALSE,
boxHeight = unit(2, "mm"),
spaceWidth = 0.02,
spaceHeight = 0.3,
limitLabel = TRUE,
transcriptHighlights = NULL,
fontsize = 8,
labels = "transcript",
stroke = 0.1,
bg = NA,
x = NULL,
y = NULL,
width = NULL,
height = NULL,
just = c("left", "top"),
default.units = "inches",
draw = TRUE,
params = NULL
)
```

**Arguments**

- `chrom`  
  Chromosome of region to be plotted, as a string.

- `chromstart`  
  Integer start position on chromosome to be plotted.

- `chromend`  
  Integer end position on chromosome to be plotted.

- `assembly`  
  Default genome assembly as a string or a `assembly` object. Default value is `assembly = "hg38"`.

- `fill`  
  Character value(s) as a single value or vector specifying fill colors of transcripts. Default value is `fill = c("#669fd9", "#abcc8e")`.

- `colorbyStrand`  
  A logical value indicating whether to color plus and minus strands by the first two colors in a `fill` vector, where plus strand transcripts will be colored by the first `fill` color and minus strand transcripts will be colored by the second `fill` color. Default value is `colorbyStrand = TRUE`.

- `strandSplit`  
  A logical value indicating whether plus and minus-stranded transcripts should be separated, with plus strand transcripts plotted above the x-axis and minus strand transcripts plotted below the x-axis. Default value is `strandSplit = FALSE`.

- `boxHeight`  
  A numeric or unit object specifying height of transcripts. Default value is `boxHeight = unit(2, "mm")`.

- `spaceWidth`  
  A numeric value specifying the width of minimum spacing between transcripts, as a fraction of the plot’s genomic range. Default value is `spaceWidth = 0.02`.

- `spaceHeight`  
  A numeric value specifying the height of spacing between transcripts on different rows, as a fraction of `boxHeight`. Default value is `spaceHeight = 0.3`. 
limitLabel A logical value indicating whether to draw a "+" when not all elements can be plotted in the plotting space. Default value is limitLabel = TRUE.

transcriptHighlights A two-column dataframe with a column named "transcript" or "gene" containing transcript names or their associated gene names as strings to highlight and a column named "color" containing corresponding highlight colors.

fontsize A numeric specifying text fontsize in points. Default value is fontsize = 8.

labels A character value describing the format of transcript text labels. Default value is labels = "transcript". Options are:

- NULL: No labels.
- "transcript": Transcript name labels.
- "gene": Gene name labels.
- "both": Combined transcript and gene name labels with the format "gene name:transcript name".

stroke A numeric value indicating the stroke width for transcript body outlines. Default value is stroke = 0.1.

bg Character value indicating background color. Default value is bg = NA.

x A numeric or unit object specifying transcript plot x-location.

y A numeric, unit object, or character containing a "b" combined with a numeric value specifying transcript plot y-location. The character value will place the transcript plot y relative to the bottom of the most recently plotted plot according to the units of the plotgardener page.

width A numeric or unit object specifying transcript plot width.

height A numeric or unit object specifying transcript plot height.

just Justification of transcript plot relative to its (x, y) location. If there are two values, the first value specifies horizontal justification and the second value specifies vertical justification. Possible string values are: "left", "right", "centre", "center", "bottom", and "top". Default value is just = c("left", "top").

default.units A string indicating the default units to use if x, y, width, or height are only given as numerics. Default value is default.units = "inches".

draw A logical value indicating whether graphics output should be produced. Default value is draw = TRUE.

params An optional pgParams object containing relevant function parameters.

Details

A transcripts plot can be placed on a plotgardener coordinate page by providing plot placement parameters:

`plotTranscripts(chrom, chromstart = NULL, chromend = NULL, x, y, width, height, just = c("left", "top"), default.units = "inches")`
This function can also be used to quickly plot an unannotated transcripts plot by ignoring plot placement parameters:

```r
plotTranscripts(chrom, chromstart = NULL, chromend = NULL)
```

Genomic annotation information is acquired through `TxDb` and `OrgDb-class` packages, as determined through the assembly parameter.

**Value**

Returns a `transcripts` object containing relevant genomic region, placement, and `grob` information.

**See Also**

`assembly`, `genomes`, `defaultPackages`

**Examples**

```r
## Load hg19 genomic annotation packages
library("TxDb.Hsapiens.UCSC.hg19.knownGene")
library("org.Hs.eg.db")

## Create page
pageCreate(width = 7.5, height = 3.5, default.units = "inches")

## Plot and place transcripts
plotTranscripts(
  chrom = "chr8", chromstart = 1000000, chromend = 2000000,
  assembly = "hg19", labels = "gene",
  x = 0.5, y = 0.5, width = 6.5, height = 2.5,
  just = c("left", "top"), default.units = "inches"
)

## Plot genome label
plotGenomeLabel(
  chrom = "chr8", chromstart = 1000000, chromend = 2000000,
  assembly = "hg19",
  x = 0.5, y = 3.03, length = 6.5, default.units = "inches"
)

## Plot a legend
plotLegend(
  legend = c("+ strand", "- strand"),
  fill = c("#669fd9", "#abcc8e"), border = FALSE,
  x = 0.5, y = 1, width = 1, height = 0.5,
  just = c("left", "top")
)

## Hide page guides
pageGuideHide()
```
readBigwig  

Read a bigWig file and return it as a data frame

Description

Read a bigWig file and return it as a data frame

Usage

readBigwig(
  file,
  chrom = NULL,
  chromstart = 1,
  chromend = .Machine$integer.max,
  strand = "*",
  params = NULL
)

Arguments

file  A character value specifying the path to the bigwig file.
chrom  Chromosome of data as a string, if data for a specific chromosome is desired.
chromstart  Integer start position on chromosome.
chromend  Integer end position on chromosome.
strand  A character value specifying strand. Default value is strand = "*". Options are:
  • "+": Plus strand.
  • "-": Minus strand.
  • "*": Plus and minus strands.
params  An optional pgParams object containing relevant function parameters.

Details

This function does not work on Windows.

Value

Returns a 6-column dataframe of bigwig information.

See Also

import.bw
Examples

```r
if (.Platform$OS.type != "windows"){
  bwFile <- system.file("extdata/test.bw", package="plotgardenerData")

  ## Read in entire file
  bwData <- readBigwig(file = bwFile)

  ## Read in specified region
  bwRegion <- readBigwig(file = bwFile,
                         chrom = "chr2",
                         chromstart = 1,
                         chromend = 1500)
}
```

---

**readHic**

*Read a .hic file and return Hi-C data as a dataframe*

**Description**

Read a .hic file and return Hi-C data as a dataframe

**Usage**

```r
readHic(
  file,
  chrom,
  chromstart = NULL,
  chromend = NULL,
  altchrom = NULL,
  altchromstart = NULL,
  altchromend = NULL,
  assembly = "hg38",
  resolution = "auto",
  res_scale = "BP",
  zrange = NULL,
  norm = "KR",
  matrix = "observed",
  params = NULL,
  quiet = FALSE
)
```

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>file</td>
<td>A character value specifying the path to the .hic file.</td>
</tr>
<tr>
<td>chrom</td>
<td>Chromosome of data, as a string.</td>
</tr>
<tr>
<td>chromstart</td>
<td>Integer start position on chromosome.</td>
</tr>
</tbody>
</table>
readHic

chromend  Integer end position on chromosome.
altchrom   Alternate chromosome for interchromosomal data, as a string.
altchromstart  Alternate chromosome integer start position for interchromosomal data.
altchromend  Alternate chromosome integer end position for interchromosomal data.
assembly   Default genome assembly as a string or a assembly object. Default value is assembly = "hg38".
resolution A numeric specifying the width of each pixel. "auto" will attempt to choose a resolution in basepairs based on the size of the region.
res_scale  A character value specifying the resolution scale. Default value is res_scale = "BP". Options are:
  • "BP": Base pairs.
  • "FRAG": Fragments.
zrange    A numeric vector of length 2 specifying the range of interaction scores, where extreme values will be set to the max or min.
norm      Character value specifying hic data normalization method. This value must be found in the .hic file. Default value is norm = "KR".
matrix    Character value indicating the type of matrix to output. Default value is matrix = "observed". Options are:
  • "observed": Observed counts.
  • "oe": Observed/expected counts.
  • "log2oe": Log2 transformed observed/expected counts.
params    An optional pgParams object containing relevant function parameters.
quiet     A logical indicating whether or not to print messages.

Value

Returns a 3-column dataframe in sparse upper triangular format with the following columns: chrom, altchrom, counts.

See Also

straw

Examples

hicFile <- system.file("extdata/test_chr22.hic", package="plotgardenerData")

## Read in data for all chr22 file at 2.5Mb bp resolution
hicData <- readHic(file = hicFile, chrom = "22", assembly = "hg19", resolution = 2500000)

## Read in region "chr22:20000000-47500000" at 100 Kb resolution
hicData10Kb <- readHic(file = hicFile, chrom = "22", chromstart = 20000000, chromend = 47500000, assembly = "hg19"
resolution = 100000)
Index

annoDomains, 3
annoGenomeLabel, 5
annoHeatmapLegend, 8
annoHighlight, 10
annoPixels, 12
annoSegments, 14
AnnotationHub, 66
annoText, 17
annoXaxis, 18
annoYaxis, 20
annoZoomLines, 22
arrow, 15, 16, 95
assembly, 24, 27, 47, 48, 50, 55, 58, 62, 64, 70, 75, 78, 81, 87, 97, 102, 104, 107
BSgenome, 25
c, 25
calcSignalRange, 26
colorby, 27, 30, 70, 78, 82, 87
defaultPackages, 28, 48, 104
formatC, 7, 51
genomes, 29, 48, 104
ggplot, 53
GInteractions, 12, 78, 81
gpar, 4, 7, 9, 11, 13, 15, 18, 19, 21, 23, 33, 35, 44, 51, 68, 71, 76, 79, 83, 85, 88, 90, 93, 95, 98, 100
GRanges, 3, 26, 69, 75, 87, 97
grid.circle, 45
grid.polygon, 85
grid.raster, 91
grid.rect, 93
grid.segments, 16, 95
grid.text, 18, 101
grob, 4, 7, 9, 11, 13, 15, 18, 19, 21, 23, 44, 48, 51, 53, 56, 60, 63, 66, 68, 71, 76, 79, 83, 85, 91, 93, 95, 98, 100, 104
import.bw, 105
mapColors, 29
OrgDb-class, 25, 48, 104
pageCreate, 31, 34
pageGuideHide, 32
pageGuideHorizontal, 33, 34
pageGuideShow, 34
pageGuideVertical, 34, 35
pageLayoutCol, 36
pageLayoutRow, 36
pagePlotPlace, 37
pagePlotRemove, 39
plotCircle, 44
plotgardener, 45
plotgardener-package (plotgardener), 45
plotGenes, 46
plotGenomeLabel, 49
plotGG, 52
plotHicRectangle, 54
plotHicSquare, 57
plotHicTriangle, 56, 61
plotIdeogram, 64
plotLegend, 66
plotManhattan, 69
plotMultiSignal, 74
plotPairs, 77
plotPairsArches, 80
plotPolygon, 84
plotRanges, 86
plotRaster, 90
plotRect, 92
plotSegments, 94
plotSignal, 96

109
plotText, 75, 97, 99
plotTranscripts, 101
readBigwig, 105
readHic, 56, 60, 63, 106
straw, 107
TxDB, 25, 48, 104