Package 'trackViewer'

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```
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      interactive tracks or lollipop plot to facilitate integrated
      analysis of multi-omics data
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```

Type Package

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 ${\tt track Viewer-package} \qquad {\tt \textit{Minimal designed plotting tool for genomic data}$

Description

A package that plot data and annotation information along genomic coordinates in an elegance style. This tool is based on Gviz but want to draw figures in minimal style for publication.

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Examples

```
library(TxDb.Hsapiens.UCSC.hg19.knownGene)
library(org.Hs.eg.db)
trs <- geneModelFromTxdb(TxDb.Hsapiens.UCSC.hg19.knownGene,</pre>
                           org.Hs.eg.db,
                           chrom="chr11"
                           start=122929275,
                           end=122930122)
extdata <- system.file("extdata", package="trackViewer",</pre>
                         mustWork=TRUE)
repA <- importScore(paste(extdata, "cpsf160.repA_+.wig", sep="/"),</pre>
                     paste(extdata, "cpsf160.repA_-.wig", sep="/"),
                     format="WIG")
strand(repA@dat) <- "+"</pre>
strand(repA@dat2) <- "-"</pre>
fox2 <- importScore(paste(extdata, "fox2.bed", sep="/"), format="BED")</pre>
dat <- coverageGR(fox2@dat)</pre>
fox2@dat <- dat[strand(dat)=="+"]</pre>
fox2@dat2 <- dat[strand(dat)=="-"]</pre>
gr \leftarrow GRanges("chr11", IRanges(122929275, 122930122), strand="-")
vp <- viewTracks(trackList(repA, fox2, trs), gr=gr, autoOptimizeStyle=TRUE)</pre>
addGuideLine(c(122929767, 122929969), vp=vp)
addArrowMark(list(x=unit(.5, "npc"),
                   y=unit(.39, "npc")),
              col="blue")
```

addArrowMark

Add arrow mark to the figure at a given position

Description

A function to add arrow mark for emphasizing peaks

Usage

```
addArrowMark(pos = grid.locator(), label = NULL, angle = 15,
  length = unit(0.25, "inches"), col = "red", cex = 1,
  quadrant = 4, type = "closed", vp = NULL)
```

Arguments

pos	A unit object representing the location of arrow mark to be placed at current viewport. Default is the value of grid.locator, which will get the location of the mouse click.
label	A character or expression vector.
angle	A parameter passed into grid::arrow function. The angle of arrow head in degrees (smaller numbers produce narrower, pointier arrows). Essentially describes the width of the arrow head.
length	A parameter passed into grid::arrow function. Aunit specifying the length of the arrow head.
col	color of the arrow

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cex Multiplier applied to fontsize

quadrant the direction of arrow, 1: to bottomleft, 2: to bottomright, 3: to topright, 4: to

opleft

type A parameter passed into grid::arrow function. One of "open" or "closed" indi-

cating whether the arrow head should be a closed triangle.

vp A Grid viewport object. It must be output of viewTracks

Value

invisible x, y position value.

See Also

See Also as addGuideLine, arrow

Examples

addGuideLine

Add guide lines to the tracks

Description

A function to add lines for emphasizing the positions

Usage

```
addGuideLine(guideLine, col = "gray", lty = "dashed", lwd = 1,
    vp = NULL)
```

Arguments

guideLine The genomic coordinates to draw the lines col A vector for the line color

1ty A vector for the line type 1wd A vector for the line width

vp A Grid viewport object. It must be output of viewTracks

See Also

See Also as getCurTrackViewport, addArrowMark, viewTracks

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Examples

```
vp \leftarrow getCurTrackViewport(trackViewerStyle(), 10000, 10200) addGuideLine(c(10010, 10025, 10150), vp=vp)
```

browseTracks

browse tracks

Description

browse tracks by a web browser.

Usage

```
browseTracks(trackList, gr = GRanges(), ignore.strand = TRUE,
  width = NULL, height = NULL, ...)
```

Arguments

```
trackList an object of trackList
gr an object of GRanges
ignore.strand ignore the strand or not when do filter. default TRUE
width width of the figure
height height of the figure
... parameters not used
```

Value

An object of class htmlwidget that will intelligently print itself into HTML in a variety of contexts including the R console, within R Markdown documents, and within Shiny output bindings.

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browseTracks-shiny Shiny bindings for browseTracks

Description

Output and render functions for using browseTracks within Shiny applications and interactive Rmd documents.

Usage

```
browseTracksOutput(outputId, width = "100%", height = "600px")
renderbrowseTracks(expr, env = parent.frame(), quoted = FALSE)
```

Arguments

outputId output variable to read from

width, height Must be a valid CSS unit (like '100%', '400px', 'auto') or a number, which

will be coerced to a string and have 'px' appended.

expr An expression that generates a browseTracks env The environment in which to evaluate expr.

quoted Is expr a quoted expression (with quote())? This is useful if you want to save

an expression in a variable.

coverageGR calculate coverage

Description

calculate coverage for GRanges, GAlignments or GAlignmentPairs

Usage

```
coverageGR(gr)
```

Arguments

gr an object of RGanges, GAlignments or GAlignmentPairs

Value

an object of GRanges

See Also

See Also as coverage, coverage-methods

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Examples

dandelion.plot

dandelion.plots

Description

Plot variants and somatic mutations

Usage

```
dandelion.plot(SNP.gr, features = NULL, ranges = NULL,
  type = c("fan", "circle", "pie", "pin"), newpage = TRUE,
  ylab = TRUE, ylab.gp = gpar(col = "black"), xaxis = TRUE,
  xaxis.gp = gpar(col = "black"), yaxis = FALSE, yaxis.gp = gpar(col
  = "black"), legend = NULL, cex = 1, maxgaps = 1/50,
  heightMethod = NULL, ...)
```

Arguments

SNP.gr A object of GRanges or GRangesList. All the width of GRanges must be 1.

features A object of GRanges or GRangesList.
ranges A object of GRanges or GRangesList.

type Character. Could be fan, circle, pie or pin.

newpage plot in the new page or not.

ylab plot ylab or not. If it is a character vector, the vector will be used as ylab.

ylab.gp, xaxis.gp, yaxis.gp

An object of class gpar for ylab, xaxis or yaxis.

xaxis, yaxis plot xaxis/yaxis or not. If it is a numeric vector with length greater than 1, the

vector will be used as the points at which tick-marks are to be drawn. And the names of the vector will be used to as labels to be placed at the tick points if it

has names.

legend If it is a list with named color vectors, a legend will be added.

cex cex will control the size of circle.

maxgaps maxgaps between the stem of dandelions. It is calculated by the width of plot

region devided by maxgaps. If a GRanges object is set, the dandelions stem will

be clusted in each genomic range.

heightMethod A function used to determine the height of stem of dandelion. eg. Mean. Default

is length.

... not used.

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Details

In SNP.gr and features, metadata of the GRanges object will be used to control thecolor, fill, border, height, data source of pie if the type is pie.

Examples

geneModelFromTxdb

Prepare gene model from an object of TxDb

Description

Generate an object of track for viewTracks by given parameters.

Usage

```
geneModelFromTxdb(txdb, orgDb, gr, chrom, start, end, strand = c("*",
   "+", "-"), txdump = NULL)
```

Arguments

txdb	An object of TxDb
orgDb	An object of "OrgDb"
gr	An object of GRanges.
chrom	chromosome name, must be a seqname of txdb
start	start position
end	end position
strand	strand
txdump	output of as.list(txdb), a list of data frames that can be used to make the db again with no loss of information.

Value

An object of track

See Also

See Also as importScore, importBam, viewTracks

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Examples

geneTrack

track from TxDb

Description

Generate a track object from TxDb by given gene ids

Usage

```
geneTrack(ids, txdb, type = c("gene", "transcript"))
```

Arguments

ids Gene IDs. A vector of character. It should be keys in txdb.

txdb An object of TxDb

type Output type of track, "gene" or "transcript".

Value

An object of track

Examples

```
library(TxDb.Hsapiens.UCSC.hg19.knownGene)
geneTrack(c("3312", "3313"), TxDb.Hsapiens.UCSC.hg19.knownGene)
```

getCurTrackViewport

Get current track viewport

Description

Get current track viewport for addGuideLine

Usage

```
getCurTrackViewport(curViewerStyle, start, end)
```

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Arguments

```
curViewerStyle an object of trackViewerStyle
start start position of current track
end end position of current track
```

Value

```
an object of viewport
```

See Also

See Also as addGuideLine

Examples

```
vp \leftarrow getCurTrackViewport(trackViewerStyle(), 10000, 10200) addGuideLine(c(10010, 10025, 10150), vp=vp)
```

getLocation

get genomic location by gene symbol

Description

given a gene name, get the genomic coordinates.

Usage

```
getLocation(symbol, txdb, org)
```

Arguments

symbol Gene symbol

txdb will be used to extract the genes

org org package name

```
library(TxDb.Hsapiens.UCSC.hg19.knownGene)
library(org.Hs.eg.db)
getLocation("HSPA8", TxDb.Hsapiens.UCSC.hg19.knownGene, "org.Hs.eg.db")
```

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gieStain

color sheme for the schema for Chromosome Band (Ideogram)

Description

Describe the colors of giemsa stain results

Usage

```
gieStain()
```

Value

A character vector of colors

Examples

```
gieStain()
```

gridPlot

plot GRanges metadata

Description

plot GRanges metadata for different types

Usage

```
gridPlot(gr, gp, type, xscale)
```

Arguments

gr an object of GRanges with metadata. All metadata must be numeric.

gp an object of gpar

type type of the figure, could be barplot, line, point and heatmap

xscale x scale of the viewport

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GRo	per	ato	r

GRanges operator

Description

GRanges operations (add, aubtract, multiply, divide)

Usage

```
GRoperator(A, B, col = "score", operator = c("+", "-", "*", "/", "^", "%"), ignore.strand = TRUE)
```

Arguments

```
A an object of GRanges

B an object of GRanges

col colname of A and B to be calculated

operator operator, "+" means A + B, and so on. User-defined function also could be used.

ignore.strand When set to TRUE, the strand information is ignored in the overlap calculations.
```

Value

an object of GRanges

Examples

ideogramPlot

plot ideogram with data

Description

plot ideogram with data for multiple chromosomes

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Usage

```
ideogramPlot(ideo, dataList, layout = NULL, horiz = TRUE,
  parameterList = list(vp = plotViewport(margins = c(0.1, 4.1, 0.3,
  0.1)), ideoHeight = unit(1/(1 + length(dataList)), "npc"), vgap =
  unit(0.3, "lines"), ylabs = "auto", ylabsRot = ifelse(horiz, 0, 90),
  ylabsPos = unit(2.5, "lines"), xaxis = FALSE, yaxis = FALSE, xlab = "",
  types = "barplot", heights = NULL, dataColumn = "score", gps = gpar(col =
  "black", fill = "gray")), colorSheme = gieStain(), gp = gpar(fill =
  NA, lwd = 2), ...)
```

Arguments

ideo output of loadIdeogram. dataList a GRangesList of data to plot. layout The layout of chromosomes. Could be a list with chromosome names as its elements. horiz a logical value. If FALSE, the ideograms are drawn vertically to the left. If TRUE, the ideograms are drawn horizontally at the bottom. a list of parameters for each dataset in the dataList. The elements of the paramparameterList eters could be xlabs, ylabs, etc. type could be barplot, line, point, heatmap. colorSheme A character vector of giemsa stain colors. parameters used for grid.roundrect. gp parameters not used.

```
## Not run:
ideo <- loadIdeogram("hg38")</pre>
library(rtracklayer)
library(grid)
dataList <- ideo
dataList$score <- as.numeric(dataList$gieStain)</pre>
dataList <- dataList[dataList$gieStain!="gneg"]</pre>
dataList <- GRangesList(dataList)</pre>
grid.newpage()
ideogramPlot(ideo, dataList,
              layout=list("chr1", "chr2", c("chr3", "chr22"),
                          c("chr4", "chr21"), c("chr5", "chr20"),
                          c("chr6", "chr19"), c("chr7", "chr18"),
                          c("chr8", "chr17"), c("chr9", "chr16"),
                          c("chr10", "chr15"), c("chr11", "chr14"),
                          c("chr12", "chr13"), c("chrX", "chrY")).
              parameterList = list(types="heatmap", colorKeyTitle="sample1"))
## End(Not run)
```

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importBam

Reading data from a BAM file

Description

Read a track object from a BAM file

Usage

```
importBam(file, file2, ranges = GRanges(), pairs = FALSE)
```

Arguments

file The path to the BAM file to read.

file2 The path to the second BAM file to read.

ranges An object of GRanges to indicate the range to be imported

pairs logical object to indicate the BAM is paired or not. See readGAlignments

Value

a track object

See Also

See Also as importScore, track, viewTracks

Examples

```
bamfile <- system.file("extdata", "ex1.bam", package="Rsamtools",
mustWork=TRUE)
dat <- importBam(file=bamfile, ranges=GRanges("seq1", IRanges(1, 50), strand="+"))</pre>
```

importData

Reading data from a BED or WIG file to RleList

Description

Read a track object from a BED, bedGraph, WIG or BigWig file to RleList

Usage

```
importData(files, format = NA, ranges = GRanges())
```

Arguments

files The path to the files to read.

format The format of import file. Could be BAM, BED, bedGraph, WIG or BigWig

ranges An object of GRanges to indicate the range to be imported

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Value

```
a list of RleList.
```

Examples

```
#import a BED file
bedfile <- system.file("tests", "test.bed", package="rtracklayer",</pre>
                        mustWork=TRUE)
dat <- importData(files=bedfile, format="BED",</pre>
                   ranges=GRanges("chr7", IRanges(127471197, 127474697)))
##import a WIG file
wigfile <- system.file("tests", "step.wig", package = "rtracklayer",</pre>
                        mustWork=TRUE)
dat <- importData(files=wigfile, format="WIG",</pre>
                   ranges=GRanges("chr19",
                                   IRanges(59104701, 59110920)))
##import a BigWig file
if(.Platform$OS.type!="windows"){
  ##this is because we are using rtracklayer::import
 bwfile <- system.file("tests", "test.bw", package = "rtracklayer",</pre>
                         mustWork=TRUE)
 dat <- importData(files=bwfile, format="BigWig",</pre>
                    ranges=GRanges("chr19", IRanges(1500, 2700)))
}
```

importScore

Reading data from a BED or WIG file

Description

Read a track object from a BED, bedGraph, WIG or BigWig file

Usage

```
importScore(file, file2, format = c("BED", "bedGraph", "WIG", "BigWig"),
  ranges = GRanges(), ignore.strand = TRUE)
```

Arguments

file	The path to the file to read.
file2	The path to the second file to read.
format	The format of import file. Could be BED, bedGraph, WIG or BigWig
ranges	An object of GRanges to indicate the range to be imported
ignore.strand	ignore the strand or not when do filter. default TRUE

Value

```
a track object
```

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See Also

See Also as importBam, track, viewTracks

Examples

```
#import a BED file
bedfile <- system.file("tests", "test.bed", package="rtracklayer",</pre>
                        mustWork=TRUE)
dat <- importScore(file=bedfile, format="BED",</pre>
                    ranges=GRanges("chr7", IRanges(127471197, 127474697)))
##import a WIG file
wigfile <- system.file("tests", "step.wig", package = "rtracklayer",</pre>
                        mustWork=TRUE)
dat <- importScore(file=wigfile, format="WIG")</pre>
##import a BigWig file
if(.Platform$OS.type!="windows"){##this is because we are using rtracklayer::import
 bwfile <- system.file("tests", "test.bw", package = "rtracklayer",</pre>
                         mustWork=TRUE)
 dat <- importScore(file=bwfile, format="BigWig")</pre>
}
##import 2 file
wigfile1 <- system.file("extdata", "cpsf160.repA_+.wig", package="trackViewer",</pre>
                         mustWork=TRUE)
wigfile2 <- system.file("extdata", "cpsf160.repA_-.wig", package="trackViewer",</pre>
                         mustWork=TRUE)
dat <- importScore(wigfile1, wigfile2, format="WIG",</pre>
                    ranges=GRanges("chr11", IRanges(122817703, 122889073)))
```

loadIdeogram

load ideogram from UCSC

Description

Download ideogram table from UCSC

Usage

```
loadIdeogram(genome, chrom = NULL, ranges = NULL, ...)
```

Arguments

genome Assembly name assigned by UCSC, such as hg38, mm10.

chrom A character vector of chromosome names, or NULL.

ranges A Ranges object with the intervals.

Additional arguments to pass to the GRanges constructor.

Value

A GRanges object.

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See Also

See Also as ideogramPlot

Examples

```
## Not run:
head(loadIdeogram("hg38"))
## End(Not run)
```

lolliplot

Lolliplots

Description

Plot variants and somatic mutations

Usage

```
lolliplot(SNP.gr, features = NULL, ranges = NULL, type = "circle",
  newpage = TRUE, ylab = TRUE, ylab.gp = gpar(col = "black"),
  yaxis = TRUE, yaxis.gp = gpar(col = "black"), xaxis = TRUE,
  xaxis.gp = gpar(col = "black"), legend = NULL, cex = 1,
  dashline.col = "gray80", jitter = c("node", "label"),
  rescale = FALSE, ...)
```

Arguments

SNP.gr A object of GRanges, GRangesList or a list of GRanges. All the width of

GRanges must be 1.

features A object of GRanges, GRangesList or a list of GRanges. The metadata 'fea-

tureLayerID' are used for drawing features in different layers. See details in

vignette.

ranges A object of GRanges or GRangesList.

type character. Could be circle, pie, pin, pie.stack or flag.

newpage Plot in the new page or not.

ylab Plot ylab or not. If it is a character vector, the vector will be used as ylab.

ylab.gp, xaxis.gp, yaxis.gp

An object of class gpar for ylab, xaxis or yaxis.

yaxis Plot yaxis or not.

xaxis Plot xaxis or not. If it is a numeric vector with length greater than 1, the vector

will be used as the points at which tick-marks are to be drawn. And the names of the vector will be used to as labels to be placed at the tick points if it has names.

legend If it is a list with named color vectors, a legend will be added.

cex cex will control the size of circle.

dashline.col color for the dashed line.

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jitter jitter the position of nodes or labels.

rescale logical(1) or a dataframe with rescale from and to. Recalse the x-axis or not. if dataframe is used, colnames must be from.start, from.end, to.start, to.end.

... not used.

Details

In SNP.gr and features, metadata of the GRanges object will be used to control the color, fill, border, height, cex, dashline.col, data source of pie if the type is pie. And also the controls for labels by name the metadata start as label.parameter.properties> such as label.parameter.rot, label.parameter.gp. The parameter is used for grid.text. The metadata 'featureLayerID' for features are used for drawing features in different layers. The metadata 'SNPsideID' for SNP.gr are used for determining the side of lollipops. And the 'SNPsideID' could only be 'top' or 'bottom'.

Examples

 ${\tt optimizeStyle}$

Optimize the style of plot

Description

Automatic optimize the stlye of trackViewer

Usage

```
optimizeStyle(trackList, viewerStyle = trackViewerStyle(),
  theme = NULL)
```

Arguments

trackList An object of trackList
viewerStyle An object of trackViewerStyle
theme A character string. Could be "bw", "col" or "safe".

Value

```
a list of a trackList and a trackViewerStyle
```

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See Also

See Also as viewTracks

Examples

parse2GRanges

parse text into GRanges

Description

```
parse text like "chr13:99,443,451-99,848,821:-" into GRanges
```

Usage

```
parse2GRanges(text)
```

Arguments

text

character vector like "chr13:99,443,451-99,848,821:-" or "chr13:99,443,451-99,848,821"

Value

an object of GRanges

Examples

```
parse2GRanges("chr13:99,443,451-99,848,821:-")
```

parseWIG

convert WIG format track to BED format track

Description

convert WIG format track to BED format track for a given range

Usage

```
parseWIG(trackScore, chrom, from, to)
```

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Arguments

trackScore an object of track with WIG format chrom sequence name of the chromosome

from start coordinate to end coordinate

Value

```
an object of track
```

Examples

plotGInteractions

plot GInteractions

Description

```
plot graph for GInteractions
```

Usage

```
\verb|plotGInteractions(gi, range, feature.gr, ...)|\\
```

Arguments

```
gi an object of GInteractions
```

range the region to plot. an object of GRanges

feature.gr to be added. an object of GRanges

... Not used.

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|--|

Description

A function to plot GRanges data for given range

Usage

```
plotGRanges(..., range = GRanges(), viewerStyle = trackViewerStyle(),
  autoOptimizeStyle = FALSE, newpage = TRUE)
```

Arguments

```
one or more objects of GRanges

range an object of GRanges

viewerStyle an object of trackViewerStyle

autoOptimizeStyle should use optimizeStyle to optimize style

newpage should be draw on a new page?
```

Value

An object of viewport for addGuideLine

See Also

See Also as addGuideLine, addArrowMark

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plotIdeo

plot ideogram

Description

plot ideogram for one chromosome

Usage

```
plotIdeo(ideo, chrom = seqlevels(ideo)[1], colorSheme = gieStain(),
   gp = gpar(fill = NA), ...)
```

Arguments

ideo output of loadIdeogram.

chrom A length 1 character vector of chromosome name.

colorSheme A character vector of giemsa stain colors.

gp parameters used for grid.roundrect.

... parameters not used.

Examples

```
## Not run:
ideo <- loadIdeogram("hg38")
library(grid)
grid.newpage()
plotIdeo(ideo)
## End(Not run)</pre>
```

plot0neIdeo

plot ideogram with data for one chromosome

Description

plot ideogram with data for one chromosome

Usage

```
plotOneIdeo(ideo, dataList, parameterList = list(vp =
  plotViewport(margins = c(0.1, 4.1, 1.1, 0.1)), ideoHeight = unit(1/(1 +
  length(dataList)), "npc"), vgap = unit(1, "lines"), ylabs =
  seqlevels(ideo)[1], ylabsRot = 90, ylabsPos = unit(2.5, "lines"), xaxis =
  FALSE, yaxis = FALSE, xlab = "", types = "barplot", heights = NULL,
  dataColumn = "score", gps = gpar(col = "black", fill = "gray")),
  chrom = seqlevels(ideo)[1], colorSheme = gieStain(), gp = gpar(fill
  = NA, lwd = 2), ...)
```

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Arguments

ideo output of loadIdeogram.

dataList a GRangesList of data to plot.

parameterList a list of parameters for each dataset in the dataList. The elements of the param-

eters could be xlabs, ylabs, etc. type could be barplot, line, point, heatmap.

chrom A length 1 character vector of chromosome name.

colorSheme A character vector of giemsa stain colors.

gp parameters used for grid.roundrect.

... parameters not used.

Examples

```
## Not run:
ideo <- loadIdeogram("hg38")
library(rtracklayer)
library(grid)
dataList <- ideo[seqnames(ideo) %in% "chr1"]
dataList$score <- as.numeric(dataList$gieStain)
dataList <- dataList[dataList$gieStain!="gneg"]
dataList <- GRangesList(dataList, dataList)
grid.newpage()
plotOneIdeo(ideo, dataList, chrom="chr1")
## End(Not run)</pre>
```

pos-class Class "pos"

Description

An object of class "pos" represents a point location

Slots

```
x A numeric value, indicates the x positiony A numeric value, indicates the y positionunit "character" apecifying the units for the corresponding numeric values. See unit
```

24 trackStyle-class

trackList-class

List of tracks

Description

An extension of List that holds only track objects.

Usage

```
trackList(..., heightDist = NA)
```

Arguments

... Each tracks in ... becomes an element in the new trackList, in the same order.

This is analogous to the list constructor, except every argument in ... must be

derived from track.

heightDist A vector or NA to define the height of each track.

See Also

track.

trackStyle-class

Class "trackStyle"

Description

```
An object of class "trackStyle" represents track style.

An object of class "track" represents scores of a given track.
```

Usage

```
## $4 method for signature 'track'
show(object)

## $4 method for signature 'track'
x$name

## $4 replacement method for signature 'track'
x$name <- value

setTrackStyleParam(ts, attr, value)

## $4 method for signature 'track,character'
setTrackStyleParam(ts, attr, value)

setTrackXscaleParam(ts, attr, value)

## $4 method for signature 'track,character'</pre>
```

trackStyle-class 25

```
setTrackXscaleParam(ts, attr, value)
setTrackYaxisParam(ts, attr, value)
## S4 method for signature 'track,character'
setTrackYaxisParam(ts, attr, value)
```

Arguments

object an object of trackStyle.

x an object of trackStyle

name slot name of trackStyle

value values to be assigned.

ts An object of track.

attr the name of slot of trackStyle object to be changed.

Slots

tracktype "character" track type, could be peak or cluster. Default is "peak". "cluster" is not supported yet. #' @slot color "character" track color. If the track has dat and dat2 slot, it should have two values.

```
height "numeric" track height. It should be a value between 0 and 1 marginTop "numeric" track top margin marginBottom "numeric" track bottom margin xscale object of xscale, describe the details of x-scale yaxis object of yaxisStyle, describe the details of y-axis ylim "numeric" y-axis range
```

ylabpos "character", ylable postion, ylabpos should be 'left', 'right', 'topleft', 'bottomleft', 'topright', 'bottomright', 'abovebaseline' or 'underbaseline'. For gene type track, it also could be 'upstream' or 'downstream'

ylablas "numeric" y lable direction. It should be a integer 0-3. See par:las

ylabgp A "list" object, It will convert to an object of class gpar. This is basically a list of graphical parameter settings of y-label.

dat Object of class GRanges the scores of a given track. It should contain score metadata.

dat2 Object of class GRanges the scores of a given track. It should contain score metadata. When dat2 and dat is paired, dat will be drawn as positive value where dat2 will be drawn as negative value (-1 * score)

type The type of track. It could be 'data', 'gene', 'transcript' or 'lollipopData'.

format The format of the input. It could be "BED", "bedGraph", "WIG", "BigWig" or "BAM" style Object of class trackStyle

name unused yet

See Also

Please try to use importScore and importBam to generate the object.

Examples

```
extdata <- system.file("extdata", package="trackViewer",
mustWork=TRUE)
fox2 <- importScore(file.path(extdata, "fox2.bed"), format="BED")
setTrackStyleParam(fox2, "color", c("red","green"))
setTrackXscaleParam(fox2, "gp", list(cex=.5))
setTrackYaxisParam(fox2, "gp", list(col="blue"))
fox2$dat <- GRanges(score=numeric(0))</pre>
```

```
trackViewerStyle-class
```

Class "trackViewerStyle"

Description

An object of class "trackViewerStyle" represents track viewer style.

Usage

```
trackViewerStyle(...)
setTrackViewerStyleParam(tvs, attr, value)
## S4 method for signature 'trackViewerStyle,character'
setTrackViewerStyleParam(tvs, attr,
    value)
```

Arguments

... Each argument in ... becomes an slot in the new trackViewerStyle.

tvs An object of trackViewerStyle. attr the name of slot to be changed.

value values to be assigned.

Slots

```
margin "numeric", specify the bottom, left, top and right margin.
xlas "numeric", label direction of x-axis mark. It should be a integer 0-3. See par:las
xgp A "list", object, It will convert to an object of class gpar. This is basically a list of graphical
    parameter settings of x-axis. For y-axis, see yaxisStyle
xaxis "logical", draw x-axis or not
autolas "logical" automatic determine y label direction
flip "logical" flip the x-axis or not, default FALSE
```

```
tvs <- trackViewerStyle()
setTrackViewerStyleParam(tvs, "xaxis", TRUE)</pre>
```

viewGene 27

	viewGene	plot tracks based on gene name	
--	----------	--------------------------------	--

Description

given a gene name, plot the tracks.

Usage

```
viewGene(symbol, filenames, format, txdb, org, upstream = 1000,
downstream = 1000, anchor = c("gene", "TSS"), plot = FALSE)
```

Arguments

symbol Gene symbol filenames files used to generate tracks format file format used to generate tracks txdb txdb will be used to extract the genes org package name org upstream from anchor upstream downstream from anchor downstream anchor TSS, or gene

plot the tracks or not.

Value

plot

an invisible list of a trackList, a trackViewerStyle and a GRanges

28 viewTracks

viewTracks pl	ot	the	tracks
-----------------	----	-----	--------

Description

A function to plot the data for given range

Usage

```
viewTracks(trackList, chromosome, start, end, strand, gr = GRanges(),
  ignore.strand = TRUE, viewerStyle = trackViewerStyle(),
  autoOptimizeStyle = FALSE, newpage = TRUE, operator = NULL,
  smooth = FALSE)
```

Arguments

```
an object of trackList
trackList
chromosome
                  chromosome
start
                  start position
end
                  end position
strand
                  strand
                  an object of GRanges
gr
                  ignore the strand or not when do filter. default TRUE
ignore.strand
viewerStyle
                  an object of trackViewerStyle
autoOptimizeStyle
                  should use optimizeStyle to optimize style
                  should be draw on a new page?
newpage
                  operator, could be +, -, *, /, ^{\wedge}, \%\%. "-" means dat - dat2, and so on.
operator
                  logical(1) or numeric(1). Smooth the curve or not. If it is numeric, eg n, mean
smooth
```

Value

An object of viewport for addGuideLine

See Also

See Also as addGuideLine, addArrowMark

Examples

of nearby n points will be used for plot.

xscale-class 29

```
fox2@dat <- dat[strand(dat)=="+"]
fox2@dat2 <- dat[strand(dat)=="-"]
gr <- GRanges("chr11", IRanges(122929275, 122930122), strand="-")
viewTracks(trackList(track=tracks, fox2=fox2), gr=gr, autoOptimizeStyle=TRUE)</pre>
```

xscale-class

Class "xscale"

Description

An object of class "xscale" represents x-scale style.

Slots

from A pos class, indicates the start point postion of x-scale.

to A pos class, indicates the end point postion of x-scale.

label "character" the label of x-scale

gp A "list" object, It will convert to an object of class gpar. This is basically a list of graphical parameter settings of x-scale.

draw A "logical" value indicating whether the x-scale should be draw.

yaxisStyle-class

Class "yaxisStyle"

Description

An object of class "yaxisStyle" represents y-axis style.

Slots

at "numeric" vector of y-value locations for the tick marks

label "logical" value indicating whether to draw the labels on the tick marks.

gp A "list" object, It will convert to an object of class gpar. This is basically a list of graphical parameter settings of y-axis.

draw A "logical" value indicating whether the y-axis should be draw.

main A "logical" value indicating whether the y-axis should be draw in left (TRUE) or right (FALSE).

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