Package 'circlize'

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Type Package

Title Circular Visualization

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Description Circular layout is an efficient way for the visualization of huge amounts of information. Here this package provides an implementation of circular layout generation in R as well as an enhancement of available software. The flexibility of the package is based on the usage of low-level graphics functions such that self-defined high-level graphics can be easily implemented by users for specific purposes. Together with the seamless connection between the powerful computational and visual environment in R, it gives users more convenience and freedom to design figures for better understanding complex patterns behind multiple dimensional data.

URL https://github.com/jokergoo/circlize,

http://jokergoo.github.io/circlize_book/book/

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circlize-package

Circular visualization in R

Description

Circular visualization in R

Details

This package aims to implement circular layout in R.

Since most of the figures are composed of points, lines and polygons, we just need to implement low-level functions for drawing points, lines and polygons.

Current there are following low-level graphic functions:

- circos.points
- circos.lines
- circos.rect
- circos.polygon
- circos.segments
- circos.text
- circos.axis, circos.xaxis, circos.yaxis
- circos.link

For drawing points, lines and text through the whole track (among several sectors), the following functions are available:

- circos.trackPoints
- circos.trackLines
- circos.trackText

Functions to arrange circular layout:

- circos.initialize
- circos.track
- circos.update
- circos.par
- circos.info
- circos.clear

Theoretically, you are able to draw most kinds of circular plots by the above functions.

For specific use in genomics, we also implement functions which add graphics in genome scale.

Functions to initialize circos plot with genomic coordinates:

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- circos.initializeWithIdeogram
- circos.genomicInitialize

Functions to arrange genomic circular layout:

• circos.genomicTrack

Functions to add basic graphics in genomic scale:

- circos.genomicPoints
- circos.genomicLines
- circos.genomicText
- circos.genomicRect
- circos.genomicLink

Functions with specific purpose:

- circos.genomicDensity
- circos.genomicRainfall
- circos.genomicIdeogram
- circos.genomicHeatmap
- circos.genomicLabels

Finally, function that draws Chord diagram:

• chordDiagram

Please refer to the vignettes (http://jokergoo.github.io/circlize_book/book/) to find out how to draw basic and advanced circular plots by this package.

Examples

There is no example NULL

add_transparency

Add transparency to colors

Description

Add transparency to colors

Usage

```
add_transparency(col, transparency = 0)
```

Arguments

col a vector of colors

transparency transparency, numeric value between 0 and 1

Value

A vector of colors

Examples

```
add_transparency("red", 0.5)
add_transparency(1, 0.5)
add_transparency("#FF000080", 0.2)
```

adjacencyList2Matrix

Convert adjacency list to adjacency matrix

Description

Convert adjacency list to adjacency matrix

Usage

```
adjacencyList2Matrix(lt, square = FALSE)
```

Arguments

1t a data frame which contains adjacency list. square is the returned matrix a square matrix?

Details

Convert adjacency list to adjacency matrix.

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Examples

```
lt = data.frame(letters[1:5], letters[6:10])
adjacencyList2Matrix(lt)

lt = data.frame(letters[1:5], letters[6:10], 1:5)
adjacencyList2Matrix(lt)

set.seed(123)
lt = data.frame(sample(letters, 4), sample(letters, 4), 1:4)
adjacencyList2Matrix(lt)
adjacencyList2Matrix(lt, square = TRUE)
```

calc_gap

Calculate gap to make two Chord diagram with same scale

Description

Calculate gap to make two Chord diagram with same scale

Usage

```
calc_gap(x1, x2, big.gap = 10, small.gap = 1)
```

Arguments

x1	The matrix or the data frame for the first Chord diagram.
x2	The matrix or the data frame for the second Chord diagram.
big.gap	big.gap for the first Chord diagram.
small.gap	small.gap for both Chord diagrams.

Details

There should be no overlap between the two sets of sectors.

Value

A numeric value which can be directly set to big.gap in the second Chord diagram.

```
\hbox{\tt\# There is no example}\\ \hbox{\tt NULL}
```

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CELL_META

Easy way to get meta data in the current cell

Description

Easy way to get meta data in the current cell

Usage

```
CELL_META
```

Details

The variable CELL_META can only be used to get meta data of the "current" cell. Basically you can simply replace e.g. get.cell.meta.data("sector.index") to CELL_META\$sector.index.

See Also

```
get.cell.meta.data
```

Examples

```
pdf(NULL)
circos.initialize("a", xlim = c(0, 1))
circos.track(ylim = c(0, 1), panel.fun = function(x, y) {
print(CELL_META$sector.index)
print(CELL_META$xlim)
})
dev.off()
```

chordDiagram

Plot Chord Diagram

Description

Plot Chord Diagram

Usage

```
chordDiagram(x, grid.col = NULL, grid.border = NA, transparency = 0.5,
    col = NULL, row.col = NULL, column.col = NULL,
    order = NULL, directional = 0, xmax = NULL,
    symmetric = FALSE, keep.diagonal = FALSE,
    direction.type = "diffHeight", diffHeight = convert_height(2, "mm"),
    reduce = 1e-5, self.link = 2,
    preAllocateTracks = NULL,
```

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```
annotationTrack = c("name", "grid", "axis"),
annotationTrackHeight = convert_height(c(3, 2), "mm"),
link.border = NA, link.lwd = par("lwd"), link.lty = par("lty"),
link.sort = FALSE, link.decreasing = TRUE,
link.arr.length = ifelse(link.arr.type == "big.arrow", 0.02, 0.4),
link.arr.width = link.arr.length/2,
link.arr.type = "triangle", link.arr.lty = par("lty"),
link.arr.lwd = par("lwd"), link.arr.col = par("col"),
link.largest.ontop = FALSE, link.visible = TRUE,
link.rank = NULL, link.overlap = FALSE,
scale = FALSE, big.gap = 10, small.gap = 1, ...)
```

Arguments

a matrix or a data frame. The function will pass all argument to chordDiagramFromMatrix Χ or chordDiagramFromDataFrame depending on the type of x, also format of other arguments depends of the type of x. If it is in the form of a matrix, it should be an adjacency matrix. If it is in the form of a data frame, it should be an adjacency list. grid.col pass to chordDiagramFromMatrix or chordDiagramFromDataFrame grid.border pass to chordDiagramFromMatrix or chordDiagramFromDataFrame transparency pass to chordDiagramFromMatrix or chordDiagramFromDataFrame col pass to chordDiagramFromMatrix or chordDiagramFromDataFrame row.col pass to chordDiagramFromMatrix column.col pass to chordDiagramFromMatrix order pass to chordDiagramFromMatrix or chordDiagramFromDataFrame directional pass to chordDiagramFromMatrix or chordDiagramFromDataFrame maximum value on x-axes, the value should be a named vector. xmax pass to chordDiagramFromMatrix symmetric pass to chordDiagramFromMatrix keep.diagonal direction.type pass to chordDiagramFromMatrix or chordDiagramFromDataFrame diffHeight pass to chordDiagramFromMatrix or chordDiagramFromDataFrame reduce pass to chordDiagramFromMatrix or chordDiagramFromDataFrame self.link pass to chordDiagramFromMatrix or chordDiagramFromDataFrame preAllocateTracks pass to chordDiagramFromMatrix or chordDiagramFromDataFrame annotationTrack pass to chordDiagramFromMatrix or chordDiagramFromDataFrame annotationTrackHeight pass to chordDiagramFromMatrix or chordDiagramFromDataFrame link.border pass to chordDiagramFromMatrix or chordDiagramFromDataFrame link.lwd pass to chordDiagramFromMatrix or chordDiagramFromDataFrame link.lty pass to chordDiagramFromMatrix or chordDiagramFromDataFrame

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link.sort link.decreasing	<pre>pass to chordDiagramFromMatrix or chordDiagramFromDataFrame g</pre>
link.arr.lengt	<pre>pass to chordDiagramFromMatrix or chordDiagramFromDataFrame h</pre>
	<pre>pass to chordDiagramFromMatrix or chordDiagramFromDataFrame</pre>
link.arr.width	<pre>pass to chordDiagramFromMatrix or chordDiagramFromDataFrame</pre>
link.arr.type	<pre>pass to chordDiagramFromMatrix or chordDiagramFromDataFrame</pre>
link.arr.lty	<pre>pass to chordDiagramFromMatrix or chordDiagramFromDataFrame</pre>
link.arr.lwd	<pre>pass to chordDiagramFromMatrix or chordDiagramFromDataFrame</pre>
link.arr.col	<pre>pass to chordDiagramFromMatrix or chordDiagramFromDataFrame</pre>
link.largest.o	ntop
	pass to chordDiagramFromMatrix or chordDiagramFromDataFrame
link.visible	pass to chordDiagramFromMatrix or chordDiagramFromDataFrame
link.rank	order to add links to the circle, a large value means to add it later.
link.overlap	<pre>pass to chordDiagramFromMatrix or chordDiagramFromDataFrame</pre>
scale	scale each sector to same width
big.gap	Gap between the two sets of sectors. If the input is a matrix, the two sets are row sectors and column sectors. If the input is a data frame, the two sets correspond to the first column and the second column. It only works when there is no intersection between the two sets.
small.gap	Small gap between sectors.
	pass to circos.link.

Details

Chord diagram is a way to visualize numeric tables (http://circos.ca/intro/tabular_visualization/), especially useful when the table represents information of directional relations. This function visualize tables in a circular way.

This function is flexible and contains some settings that may be a little difficult to understand. Please refer to vignette for better explanation.

Value

A data frame which contains positions of links, columns are:

- rn sector name corresponding to rows in the adjacency matrix or the first column in the adjacency list
- cn sector name corresponding to columns in the adjacency matrix or the second column in the adjacency list

value value for the interaction or relation

- o1 order of the link on the "from" sector
- o2 order of the link on the "to" sector
- x1 and position of the link on the "from" sector, the interval for the link on the "from" sector is c(x1-abs(value),x1)
- x2 and position of the link on the "to" sector, the interval for the link on the "from" sector is c(x2-abs(value),x2)

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

See Also

http://jokergoo.github.io/circlize_book/book/the-chorddiagram-function.html

Examples

```
set.seed(999)
mat = matrix(sample(18, 18), 3, 6)
rownames(mat) = paste0("S", 1:3)
colnames(mat) = paste0("E", 1:6)

df = data.frame(from = rep(rownames(mat), times = ncol(mat)),
    to = rep(colnames(mat), each = nrow(mat)),
    value = as.vector(mat),
    stringsAsFactors = FALSE)

chordDiagram(mat)
chordDiagram(df)
circos.clear()
```

chordDiagramFromDataFrame

Plot Chord Diagram from a data frame

Description

Plot Chord Diagram from a data frame

Usage

```
chordDiagramFromDataFrame(df, grid.col = NULL, grid.border = NA, transparency = 0.5,
    col = NULL, order = NULL, directional = 0, xmax = NULL,
    direction.type = "diffHeight", diffHeight = convert_height(2, "mm"),
    reduce = 1e-5, self.link = 2, preAllocateTracks = NULL,
    annotationTrack = c("name", "grid", "axis"),
    annotationTrackHeight = convert_height(c(3, 2), "mm"),
    link.border = NA, link.lwd = par("lwd"), link.lty = par("lty"),
    link.sort = FALSE, link.decreasing = TRUE,
    link.arr.length = ifelse(link.arr.type == "big.arrow", 0.02, 0.4),
    link.arr.width = link.arr.length/2,
    link.arr.type = "triangle", link.arr.lty = par("lty"),
    link.arr.lwd = par("lwd"), link.arr.col = par("col"),
    link.largest.ontop = FALSE, link.visible = TRUE,
    link.rank = seq_len(nrow(df)),
```

```
link.overlap = FALSE,
scale = FALSE, big.gap = 10, small.gap = 1,
...)
```

Arguments

grid.col

df A data frame with at least two columns. The first two columns specify the

connections and the third column (optional) contains numeric values which are mapped to the width of links as well as the colors if col is specified as a color mapping function. The sectors in the plot will be up in (df[[1]]] df[[2]])

mapping function. The sectors in the plot will be union(df[[1]],df[[2]]).

Grid colors which correspond to sectors. The length of the vector should be either 1 or the number of sectors. It's preferred that grid.col is a named vector of which names correspond to sectors. If it is not a named vector, the order of

grid.col corresponds to order of sectors.

grid.border border for grids. If it is NULL, the border color is same as grid color

transparency Transparency of link colors, 0 means no transparency and 1 means full trans-

parency. If transparency is already set in col or $\operatorname{row.col}$ or $\operatorname{column.col}$, this

argument will be ignored. NAalso ignores this argument.

col Colors for links. It can be a vector which corresponds to connections in df, or a function which generate colors according to values (the third column) in df.

a function which generate colors according to values (the third column) in df, or a single value which means colors for all links are the same. You may use

colorRamp2 to generate a function which maps values to colors.

order Order of sectors. Default order is union(df[[1]],df[[2]]).

directional Whether links have directions. 1 means the direction is from the first column

in df to the second column, -1 is the reverse, 0 is no direction, and 2 for two directional. The value can be a vector which has same length as number of rows

in df.

xmax maximum value on x-axes, the value should be a named vector.

direction.type type for representing directions. Can be one or two values in "diffHeight" and

"arrows". If the value contains "diffHeight", different heights of the links are used to represent the directions for which starting root has long height to give people feeling that something is comming out. If the value contains "arrows", users can customize arrows with following arguments. The value can be a vector which has same length as number of rows in df. Note if you want to set both diffHeight and arrows for certain links, you need to embed these two options

into one string such as "diffHeight+arrows".

diffHeight The difference of height between two 'roots' if directional is set to TRUE. If

the value is set to a positive value, start root is shorter than end root and if it is set to a negative value, start root is longer than the end root. The value can be a

vector which has same length as number of rows in df.

reduce if the ratio of the width of certain grid compared to the whole circle is less than

this value, the grid is removed on the plot. Set it to value less than zero if you

want to keep all tiny grid.

self.link if there is a self link in one sector, 1 means the link will be degenerated as a

'mountain' and the width corresponds to the value for this connection. 2 means

the width of the starting root and the ending root all have the same width that corresponds to the value for the connection.

preAllocateTracks

Pre-allocate empty tracks before drawing Chord diagram. It can be a single number indicating how many empty tracks needed to be created or a list containing settings for empty tracks. Please refer to vignette for details.

annotationTrack

Which annotation track should be plotted? By default, a track containing sector names and a track containing grid will be created.

annotationTrackHeight

Track height corresponding to values in annotationTrack.

link.border border for links, single scalar or a vector which has the same length as nrows of

df or a data frame

link.lwd width for link borders, single scalar or a vector which has the same length as

nrows of df or a data frame

link.lty style for link borders, single scalar or a vector which has the same length as

nrows of df or a data frame

link.sort whether sort links on every sector based on the width of the links on it. If it

is set to "overall", all links are sorted regardless whether they are from the first

column or the second column.

link.decreasing

for link.sort

link.arr.length

pass to circos.link. The format of this argument is same as link.lwd.

link.arr.width pass to Arrowhead. The format of this argument is same as link.lwd.

link.arr.type pass to circos.link, same settings as link.lwd. Default value is triangle.

link.arr.col color or the single line link which is put in the center of the belt. The format of

this argument is same as link.lwd.

link.arr.lwd line width ofthe single line link which is put in the center of the belt. The format

of this argument is same as link.lwd.

link.arr.lty line type of the single line link which is put in the center of the belt. The format

of this argument is same as link.lwd.

link.largest.ontop

controls the order of adding links, whether based on the absolute value?

link.visible whether plot the link. The value is logical, if it is set to FALSE, the corresponding

link will not plotted, but the space is still ocuppied. The format of this argument

is same as link.lwd

link.rank order to add links to the circle, a large value means to add it later.

link.overlap if it is a directional Chord Diagram, whether the links that come or end in a same

sector overlap?

scale scale each sector to same width

big.gap Gaps between the sectors in the first column of df and sectors in the second

column in df.

small.gap Small gap between sectors.

... pass to circos.link

Details

The data frame can have a column named "rank" which is used to control the order of adding links to the diagram.

Value

A data frame which contains positions of links, see explanation in chordDiagram.

Examples

```
# There is no example NULL
```

 ${\tt chordDiagramFromMatrix}$

Plot Chord Diagram from an adjacency matrix

Description

Plot Chord Diagram from an adjacency matrix

Usage

```
chordDiagramFromMatrix(mat, grid.col = NULL, grid.border = NA, transparency = 0.5,
   col = NULL, row.col = NULL, column.col = NULL, order = NULL, directional = 0,
   direction.type = "diffHeight", diffHeight = convert_height(2, "mm"),
   reduce = 1e-5, xmax = NULL, self.link = 2,
   symmetric = FALSE, keep.diagonal = FALSE, preAllocateTracks = NULL,
   annotationTrack = c("name", "grid", "axis"),
   annotationTrackHeight = convert_height(c(3, 2), "mm"),
   link.border = NA, link.lwd = par("lwd"), link.lty = par("lty"),
   link.sort = FALSE, link.decreasing = TRUE,
   link.arr.length = ifelse(link.arr.type == "big.arrow", 0.02, 0.4),
   link.arr.width = link.arr.length/2,
   link.arr.type = "triangle", link.arr.lty = par("lty"),
   link.arr.lwd = par("lwd"), link.arr.col = par("col"),
   link.largest.ontop = FALSE, link.visible = TRUE,
   link.rank = NULL, link.overlap = FALSE,
   scale = FALSE, big.gap = 10, small.gap = 1, ...)
```

Arguments

mat

A table which represents as a numeric matrix.

grid.col Grid colors which correspond to matrix rows/columns (or sectors). The length of

the vector should be either 1 or length(union(rownames(mat), colnames(mat))). It's preferred that grid.col is a named vector of which names correspond to sectors. If it is not a named vector, the order of grid.col corresponds to order

of sectors.

grid.border border for grids. If it is NULL, the border color is same as grid color

transparency Transparency of link colors, 0 means no transparency and 1 means full trans-

parency. If transparency is already set in col or $\operatorname{row.col}$ or $\operatorname{column.col}$, this

argument will be ignored. NAalso ignores this argument.

col Colors for links. It can be a matrix which corresponds to mat, or a function

which generate colors according to values in mat, or a single value which means colors for all links are the same, or a three-column data frame in which the first two columns correspond to row names and columns and the third column is colors. You may use colorRamp2 to generate a function which maps values to

colors.

row.col Colors for links. Links from the same row in mat will have the same color.

Length should be same as number of rows in mat. This argument only works

when col is set to NULL.

column.col Colors for links. Links from the same column in mat will have the same color.

Length should be same as number of columns in mat. This argument only works

when col and row.col is set to NULL.

order Order of sectors. Default order is union(df[[1]],df[[2]]).

directional Whether links have directions. 1 means the direction is from the first column

in df to the second column, -1 is the reverse, 0 is no direction, and 2 for two

directional. Same setting as link.border.

xmax maximum value on x-axes, the value should be a named vector.

direction.type type for representing directions. Can be one or two values in "diffHeight"

and "arrows". If the value contains "diffHeight", different heights of the links are used to represent the directions for which starting root has long height to give people feeling that something is comming out. If the value contains "arrows", users can customize arrows with following arguments. Same setting as link.border. Note if you want to set both diffHeight and arrows for certain links, you need to embed these two options into one string such as

"diffHeight+arrows".

diffHeight The difference of height between two 'roots' if directional is set to TRUE. If

the value is set to a positive value, start root is shorter than end root and if it is

set to a negative value, start root is longer than the end root.

reduce if the ratio of the width of certain grid compared to the whole circle is less than

this value, the grid is removed on the plot. Set it to value less than zero if you

want to keep all tiny grid.

self.link if there is a self link in one sector, 1 means the link will be degenerated as a

'mountain' and the width corresponds to the value for this connection. 2 means the width of the starting root and the ending root all have the width that corre-

sponds to the value for the connection.

symmetric Whether the matrix is symmetric. If the value is set to TRUE, only lower triangular matrix without the diagonal will be used.

keep.diagonal If the matrix is specified as symmetric, whether keep diagonal for visualization. preAllocateTracks

Pre-allocate empty tracks before drawing Chord diagram. It can be a single number indicating how many empty tracks needed to be created or a list containing settings for empty tracks. Please refer to vignette for details.

annotationTrack

Which annotation track should be plotted? By default, a track containing sector names and a track containing grid will be created.

annotation Track Height

Track height corresponding to values in annotationTrack.

link.border border for links, single scalar or a matrix with names or a data frame with three

olumns

link.lwd width for link borders, single scalar or a matrix with names or a data frame with

three columns

link.lty style for link borders, single scalar or a matrix with names or a data frame with

three columns

link.sort whether sort links on every sector based on the width of the links on it. If it

is set to "overall", all links are sorted regardless whether they are from rows or

columns.

link.decreasing

for link.sort

link.arr.length

pass to circos.link. The format of this argument is same as link.lwd.

 $\label{link.arr.width} {\it pass to Arrowhead}. \ {\it The format of this argument is same as link.lwd}.$

link.arr.type pass to circos.link, same format as link.lwd. Default value is triangle.

link.arr.col color or the single line link which is put in the center of the belt. The format of

this argument is same as link.lwd.

link arr.lwd line width of the single line link which is put in the center of the belt. The format

of this argument is same as link.lwd.

link.arr.lty line type of the single line link which is put in the center of the belt. The format

of this argument is same as link.lwd.

link.largest.ontop

controls the order of adding links, whether based on the absolute value?

link.visible whether plot the link. The value is logical, if it is set to FALSE, the corresponding

link will not plotted, but the space is still ocuppied. The format of this argument

is same as link.lwd

link.rank order to add links to the circle, a large value means to add it later.

link.overlap if it is a directional Chord Diagram, whether the links that come or end in a same

sector overlap?

scale scale each sector to same width

big.gap Gap between row sectors and column sectors.

small.gap Small gap between sectors.

... pass to circos.link

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Details

Internally, the matrix is transformed to a data frame and sent to chordDiagramFromDataFrame.

Value

A data frame which contains positions of links, see explanation in chordDiagram.

Examples

```
# There is no example NULL
```

circlize

Convert to polar coordinate system

Description

Convert to polar coordinate system

Usage

```
circlize(x, y, sector.index = get.current.sector.index(),
    track.index = get.current.track.index())
```

Arguments

Х	Data points on x-axis. The value can also be a two-column matrix/data frame if you put x and y data points into one variable.
У	Data points on y-axis.
sector.index	Index for the sector to convert the coordinates
track.index	Index for the track to convert the coordinates

Details

This is the core function in the package. It transform data points from data coordinate system (in a specific cell) to the polar coordinate system.

Value

A matrix with two columns (theta and rou). rou is measured in degree.

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Examples

```
pdf(NULL)
factors = c("a", "b")
circos.initialize(factors, xlim = c(0, 1))
circos.track(ylim = c(0, 1))
# x = 0.5, y = 0.5 in sector a and track 1
circlize(0.5, 0.5, sector.index = "a", track.index = 1)
circos.clear()
dev.off()
```

circos.arrow

Draw arrow which is paralle to the circle

Description

Draw arrow which is paralle to the circle

Usage

```
circos.arrow(x1, x2, y = get.cell.meta.data("ycenter", sector.index, track.index),
    width = get.cell.meta.data("yrange", sector.index, track.index)/2,
    sector.index = get.current.sector.index(), track.index = get.current.track.index(),
    arrow.head.length = convert_x(5, "mm", sector.index, track.index),
    arrow.head.width = width*2, arrow.position = c("end", "start"),
    tail = c("normal", "point"), border = "black", col = "white", lty = par("lty"), ...)
```

Arguments

col

lty

start position of the arrow on the x-axis. х1 end position of the arrow on the x-axis. x2 position of the arrow on the y-axis. Note this is the center of the arrow on y-axis. y width width of the arrow body. sector.index index of the sector. track.index index of the track. arrow.head.length length of the arrow head. Note the value should be smaller than the length of the arrow itself (which is x2 -x1). arrow.head.width width of the arrow head. arrow.position where is the arrow head on the arrow. tail the shape of the arrow tail (the opposite side of arrow head). border border color of the arrow.

filled color of the arrow.

line style of the arrow. pass to polygon.

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Details

Note all position values are measured in the data coordinate (the coordinate in each cell).

If you see points overflow warnings, you can set circos.par(points.overflow.warning = FALSE) to turn it off.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

```
circos.initialize(letters[1:4], x \lim = c(0, 1))
circos.track(ylim = c(0, 1), panel.fun = function(x, y) {
circos.arrow(0, 1, y = 0.5, width = 0.4, arrow.head.length = ux(1, "cm"),
col = "red", tail = ifelse(CELL_META$sector.index %in% c("a", "c"),
"point", "normal"))
}, bg.border = NA, track.height = 0.4)
######## cell cycle #########
cell_cycle = data.frame(phase = factor(c("G1", "S", "G2", "M"),
                                    levels = c("G1", "S", "G2", "M")),
                      hour = c(11, 8, 4, 1)
color = c("#66C2A5", "#FC8D62", "#8DA0CB", "#E78AC3")
circos.par(start.degree = 90)
circos.initialize(cell_cycle$phase, xlim = cbind(rep(0, 4), cell_cycle$hour))
circos.track(ylim = c(0, 1), panel.fun = function(x, y) {
circos.arrow(CELL_META$xlim[1], CELL_META$xlim[2],
arrow.head.width = CELL_META$yrange*0.8, arrow.head.length = ux(1, "cm"),
col = color[CELL_META$sector.numeric.index])
circos.text(CELL_META$xcenter, CELL_META$ycenter, CELL_META$sector.index,
facing = "downward")
}, bg.border = NA, track.height = 0.3)
circos.clear()
```

circos.axis

Draw x-axis

Description

Draw x-axis

Usage

```
circos.axis(h = "top", major.at = NULL, labels = TRUE, major.tick = TRUE,
    sector.index = get.cell.meta.data("sector.index"),
    track.index = get.cell.meta.data("track.index"),
    labels.font = par("font"), labels.cex = par("cex"),
    labels.facing = "inside", labels.direction = NULL, labels.niceFacing = TRUE,
```

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```
direction = c("outside", "inside"), minor.ticks = 4,
major.tick.percentage = 0.1, labels.away.percentage = major.tick.percentage/2,
major.tick.length = convert_y(1, "mm", sector.index, track.index),
lwd = par("lwd"), col = par("col"), labels.col = par("col"), labels.pos.adjust = TRUE)
```

Arguments

h	Position of the x-axis, can be "top", "bottom" or a numeric value
major.at	If it is numeric vector, it identifies the positions of the major ticks. It can exceed xlim value and the exceeding part would be trimmed automatically. If it is NULL, about every 10 degrees there is a major tick.
labels	labels of the major ticks. Also, the exceeding part would be trimmed automatically. The value can also be logical (either an atomic value or a vector) which represents which labels to show.
major.tick	Whether to draw major tick. If it is set to FALSE, there would be no minor ticks.
sector.index	Index for the sector
track.index	Index for the track
labels.font	font style for the axis labels
labels.cex labels.directi	font size for the axis labels on
	deprecated, use facing instead.
labels.facing labels.niceFac	facing of labels on axis, passing to circos.text
	Should facing of axis labels be human-easy
direction	whether the axis ticks point to the outside or inside of the circle.
minor.ticks	Number of minor ticks between two close major ticks.
major.tick.per	-
	not used. Length of the major ticks. It is the percentage to the height of the cell.
labels.away.pe	
	not used. The distance for the axis labels to the major ticks. It is the percentage to the height of the cell.
major.tick.length	
	length of the major ticks, measured in "current" data coordinate. convert_y can be used to convert an absolute unit to the data coordinate.
lwd	line width for ticks
col	color for the axes
labels.col labels.pos.adj	color for the labels
	whether to adjust the positions of the first label and the last label. The value can

be a vector of length two which correspond to the first label and the last label.

Details

It can only draw axes on x-direction.

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References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

See Also

circos. yaxis draws axes on y-direction.

```
factors = letters[1:8]
circos.par(points.overflow.warning = FALSE)
circos.initialize(factors = factors, xlim = c(0, 10))
circos.trackPlotRegion(factors = factors, ylim = c(0, 10), track.height = 0.1,
   bg.border = NA, panel.fun = function(x, y) {
        circos.text(5, 10, get.cell.meta.data("sector.index"))
})
circos.trackPlotRegion(factors = factors, ylim = c(0, 10))
circos.axis(sector.index = "a")
circos.axis(sector.index = "b", direction = "inside", labels.facing = "outside")
circos.axis(sector.index = "c", h = "bottom")
circos.axis(sector.index = "d", h = "bottom", direction = "inside",
    labels.facing = "reverse.clockwise")
circos.axis(sector.index = "e", h = 5, major.at = c(1, 3, 5, 7, 9))
circos.axis(sector.index = "f", h = 5, major.at = c(1, 3, 5, 7, 9),
    labels = c("a", "c", "e", "g", "f"), minor.ticks = 0)
circos.axis(sector.index = "g", h = 5, major.at = c(1, 3, 5, 7, 9),
    labels = c("a1", "c1", "e1", "g1", "f1"), major.tick = FALSE,
    labels.facing = "reverse.clockwise")
circos.axis(sector.index = "h", h = 2, major.at = c(1, 3, 5, 7, 9),
    labels = c("a1", "c1", "e1", "g1", "f1"), major.tick.percentage = 0.3,
    labels.away.percentage = 0.2, minor.ticks = 2, labels.facing = "clockwise")
circos.clear()
## Not run:
factors = letters[1]
circos.par("gap.degree" = 0, "cell.padding" = c(0, 0, 0, 0), "start.degree" = 90)
circos.initialize(factors = factors, x \lim = c(0, 12))
circos.trackPlotRegion(factors = factors, ylim = c(0, 1), bg.border = NA)
circos.axis(sector.index = "a", major.at = 0:12, labels = "",
    direction = "inside", major.tick.percentage = 0.3)
circos.text(1:12, rep(0.5, 12), 1:12, facing = "downward")
while(1) {
   current.time = as.POSIXlt(Sys.time())
    sec = ceiling(current.time$sec)
   min = current.time$min
   hour = current.time$hour
```

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```
# erase the clock hands
draw.sector(rou1 = 0.8, border = "white", col = "white")

sec.degree = 90 - sec/60 * 360
arrows(0, 0, cos(sec.degree/180*pi)*0.8, sin(sec.degree/180*pi)*0.8)

min.degree = 90 - min/60 * 360
arrows(0, 0, cos(min.degree/180*pi)*0.7, sin(min.degree/180*pi)*0.7, lwd = 2)

hour.degree = 90 - hour/12 * 360 - min/60 * 360/12
arrows(0, 0, cos(hour.degree/180*pi)*0.4, sin(hour.degree/180*pi)*0.4, lwd = 2)

Sys.sleep(1)
}
circos.clear()

## End(Not run)
```

circos.clear

Reset the circular layout parameters

Description

Reset the circular layout parameters

Usage

```
circos.clear()
```

Details

Because there are several parameters for the circular plot which can only be set before circos.initialize. So before you draw the next circular plot, you need to reset all these parameters.

If you meet some errors when re-drawing the circular plot, try running this function and it will solve most of the problems.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

```
# There is no example NULL
```

circos.dendrogram 23

circos.dendrogram Add circular dendrograms

Description

Add circular dendrograms

Usage

```
circos.dendrogram(dend, facing = c("outside", "inside"),
    max_height = NULL, use_x_attr = FALSE)
```

Arguments

dend A dendrogram object.

facing Is the dendromgrams facing inside to the circle or outside.

max_height Maximum height of the dendrogram. This is important if more than one den-

drograms are drawn in one track and making them comparable.

use_x_attr Whether use the x attribute to determine node positions in the dendrogram, used

internally.

Details

Assuming there are n nodes in the dendrogram, the positions for leaves on x-axis is 0.5,1.5,...,n -0.5. So you must be careful with xlim when you initialize the cirular layout.

You can use the dendextend package to render the dendrograms.

```
load(system.file(package = "circlize", "extdata", "bird.orders.RData"))
labels = hc$labels # name of birds
ct = cutree(hc, 6) # cut tree into 6 pieces
n = length(labels) # number of bird species
dend = as.dendrogram(hc)
circos.par(cell.padding = c(0, 0, 0, 0))
circos.initialize(factors = "a", x \lim = c(0, n)) # only one sector
max_height = attr(dend, "height") # maximum height of the trees
circos.trackPlotRegion(ylim = c(0, 1), bg.border = NA, track.height = 0.3,
    panel.fun = function(x, y) {
        for(i in seq_len(n)) {
            circos.text(i-0.5, 0, labels[i], adj = c(0, 0.5),
                facing = "clockwise", niceFacing = TRUE,
                col = ct[labels[i]], cex = 0.7)
        }
})
```

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circos.genomicAxis

Add genomic axes

Description

Add genomic axes

Usage

```
circos.genomicAxis(h = "top", major.at = NULL, labels = NULL,
    major.by = NULL, tickLabelsStartFromZero = TRUE,
    labels.cex = 0.4*par("cex"), sector.index = get.cell.meta.data("sector.index"),
    track.index = get.cell.meta.data("track.index"), ...)
```

Arguments

h	Position of the axes. "top" or "bottom".	
major.at	Major breaks. If major.at is set, major.by is ignored.	
labels	labels corresponding to major.at. If labels is set, major.at must be set.	
major.by	Increment of major ticks. It is calculated automatically if the value is not set (about every 10 degrees there is a major tick).	
tickLabelsStartFromZero		
	Whether axis tick labels start from 0? This will only affect the axis labels while not affect x-values in cells.	
labels.cex	the font size for the axis tick labels.	
sector.index	Index for the sector	
track.index	Index for the track	
	Other arguments pass to circos.axis.	

Details

It assigns proper tick labels under genomic coordinate.

```
circos.initializeWithIdeogram(plotType = NULL)
circos.track(ylim = c(0, 1), panel.fun = function(x, y) circos.genomicAxis())
circos.clear()
```

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circos.genomicDensity Calculate and add genomic density track

Description

Calculate and add genomic density track

Usage

```
circos.genomicDensity(data, ylim.force = FALSE, window.size = NULL, overlap = TRUE,
  col = ifelse(area, "grey", "black"), lwd = par("lwd"), lty = par("lty"), type = "l",
  area = TRUE, area.baseline = NULL, baseline = 0, border = NA, ...)
```

Arguments

data	A bed-file-like data frame or a list of data frames
ylim.force	Whether to force upper bound of ylim to be 1.
window.size	Pass to genomicDensity
overlap	Pass to genomicDensity
col	Colors. It should be length of one. If data is a list of data frames, the length of col can also be the length of the list.
lwd	Width of lines
lty	Style of lines
type	Type of lines, see circos.lines
area	See circos.lines
area.baseline	Deprecated, use baseline instead.
baseline	See circos.lines
border	See circos.lines
	Pass to circos.trackPlotRegion

Details

This function is a high-level graphical function, and it will create a new track.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

Examples

```
## Not run:
load(system.file(package = "circlize", "extdata", "DMR.RData"))
# rainfall
circos.initializeWithIdeogram(plotType = c("axis", "labels"))
bed_list = list(DMR_hyper, DMR_hypo)
circos.genomicRainfall(bed_list, pch = 16, cex = 0.4, col = c("#FF000080", "#0000FF80"))
circos.genomicDensity(bed_list[[1]], col = c("#FF000080"), track.height = 0.1)
circos.genomicDensity(bed_list[[2]], col = c("#0000FF80"), track.height = 0.1)
circos.clear()
## End(Not run)
```

circos.genomicHeatmap Add heatmaps for selected regions

Description

Add heatmaps for selected regions

Usage

```
circos.genomicHeatmap(bed, col, na_col = "grey",
   numeric.column = NULL, border = NA, border_lwd = par("lwd"),
   border_lty = par("lty"), connection_height = convert_height(5, "mm"),
   line_col = par("col"), line_lwd = par("lwd"), line_lty = par("lty"),
   heatmap_height = 0.15, side = c("inside", "outside"),
   track.margin = circos.par("track.margin"))
```

Arguments

bed a data frame in bed format, the matrix is stored from the fourth column.

col colors for the heatmaps. The value can be a matrix or a color mapping function

generated by colorRamp2.

na_col color for NA values.

numeric.column column index for the numeric columns. The values can be integer index or

character index

border border of the heatmap grids.

border_lwd line width for borders of heatmap grids

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```
border_lty line style for borders of heatmap grids connection_height height of the connection lines
```

line_col col of the connection line. The value can be a vector.

line_lwd line width of the connection lines.
line_lty line style of the connection lines.
heatmap_height height of the heatmap track

side side of the heatmaps. Is the heatmap facing inside or outside?

track.margin bottom and top margins

Details

The function visualizes heatmaps which correspond to a subset of regions in the genome. The correspondance between heatmaps and regions are identified by connection lines.

The function actually creates two tracks, one track for the connection lines and one track for the heamtaps. The heatmaps always fill the whole track.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

circos.genomicIdeogram

Add an ideogram track

Description

Add an ideogram track

Usage

```
circos.genomicIdeogram(cytoband = system.file(package = "circlize",
   "extdata", "cytoBand.txt"), species = NULL, track.height = convert_height(2, "mm"),
   track.margin = circos.par("track.margin"))
```

Arguments

cytoband a data frame or a file path, pass to read.cytoband species Abbreviations of species, pass to read.cytoband

track.height height of the ideogram track

track.margin margins for the track

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

```
# There is no example NULL
```

circos.genomicInitialize

Initialize circular plot with any genomic data

Description

Initialize circular plot with any genomic data

Usage

```
circos.genomicInitialize(data, sector.names = NULL, major.by = NULL,
    plotType = c("axis", "labels"), tickLabelsStartFromZero = TRUE,
    axis.labels.cex = 0.4*par("cex"), labels.cex = 0.8*par("cex"),
    track.height = NULL, ...)
```

Arguments

data A data frame containing genomic data.

sector.names Labels for each sectors which will be drawn along each sector. It will not modify

values of sector index.

major.by Increment of major ticks. It is calculated automatically if the value is not set

(about every 10 degrees there is a major tick).

plotType If it is not NULL, there will create a new track containing axis and names for

sectors. This argument controls which part should be drawn, axis for genomic

axis and labels for chromosome names

tickLabelsStartFromZero

Whether axis tick labels start from 0? This will only affect the axis labels while

not affect x-values in cells.

axis.labels.cex

the font size for the axis tick labels.

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```
labels.cex the font size for the labels.

track.height If PlotType is not NULL, height of the annotation track.

... Pass to circos.initialize
```

Details

The function will initialize circular plot from genomic data. If plotType is set with value in axis or labels, there will create a new track.

The order of sectors related to data structure of data. If the first column in data is a factor, the order of sectors is levels(data[[1]]); If the first column is just a simple vector, the order of sectors is unique(data[[1]]).

For more details on initializing genomic plot, please refer to the vignettes.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

```
## Not run:
df = read.cytoband()$df
circos.genomicInitialize(df)
df = data.frame(name = c("TP53", "TP63", "TP73"),
                 start = c(7565097, 189349205, 3569084),
                 end = c(7590856, 189615068, 3652765),
                 stringsAsFactors = FALSE)
circos.genomicInitialize(df)
circos.clear()
circos.genomicInitialize(df, major.by = 10000)
circos.clear()
circos.genomicInitialize(df, plotType = "labels")
circos.clear()
circos.genomicInitialize(df, sector.names = c("tp53", "tp63", "tp73"))
circos.clear()
circos.genomicInitialize(df, sector.names = c("tp53x", "tp63x", "tp73"))
circos.clear()
df[[1]] = factor(df[[1]], levels = c("TP73", "TP63", "TP53"))
circos.genomicInitialize(df)
circos.clear()
## End(Not run)
```

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circos.genomicLabels Add labels to specified genomic regions

Description

Add labels to specified genomic regions

Usage

```
circos.genomicLabels(bed, labels = NULL, labels.column = NULL,
    facing = "clockwise", niceFacing = TRUE,
    col = par("col"), cex = 0.8, font = par("font"), padding = 0.4,
    connection_height = convert_height(5, "mm"),
    line_col = par("col"), line_lwd = par("lwd"), line_lty = par("lty"),
    labels_height = min(c(convert_height(1.5, "cm"),
    max(strwidth(labels, cex = cex, font = font)))),
    side = c("inside", "outside"), track.margin = circos.par("track.margin"))
```

Arguments

bed a data frame in bed format

labels a vector of labels corresponding to rows in bed

labels.column if the label column is already in bed, the index for this column in bed

facing facing of the labels. The value can only be 'clockwise' or 'reverse.clockwise'.

niceFacing whether automatically adjust the facing of the labels.

col color for the labels
cex size of the labels
font font of the labels

padding padding of the labels, the value is the ratio to the height of the label

connection_height

height of the connection track

line_col color for the connection lines
line_lwd line width for the connection lines
line_lty line type for the connection lines

labels_height height of the labels track

side side of the labels track, is it in the inside of the track where the regions are

marked?

track.margin bottom and top margins

Details

The function adds labels for the specified regions. The positions of labels are arranged so that they are not overlapping to each other.

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Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

```
## Not run:
circos.initializeWithIdeogram(plotType = c("labels", "axis"))
bed = generateRandomBed(nr = 100, fun = function(k) sample(letters, k, replace = TRUE))
bed[1, 4] = "aaaaaaaa"
circos.genomicLabels(bed, labels.column = 4, side = "inside",
        col = as.numeric(factor(bed[[1]])))
circos.genomicLabels(bed, labels.column = 4, side = "outside",
        line_col = as.numeric(factor(bed[[1]])))
## End(Not run)
```

circos.genomicLines

Add lines to a plotting region, specifically for genomic graphics

Description

Add lines to a plotting region, specifically for genomic graphics

Usage

```
circos.genomicLines(region, value, numeric.column = NULL,
    sector.index = get.cell.meta.data("sector.index"),
    track.index = get.cell.meta.data("track.index"), posTransform = NULL,
    col = ifelse(area, "grey", "black"), lwd = par("lwd"),
    lty = par("lty"), type = "l",
    area = FALSE, area.baseline = NULL, border = "black", baseline = "bottom",
    pt.col = par("col"), cex = par("cex"), pch = par("pch"), ...)
```

Arguments

region A data frame contains 2 column which correspond to start position and end

position

value A data frame contains values and other information

numeric.column Which column in value data frame should be taken as y-value. If it is not

defined, the whole numeric columns in value will be taken.

sector.index Pass to circos.lines track.index Pass to circos.lines

posTransform Self-defined function to transform genomic positions, see posTransform.default

for explaination

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col	col of lines/areas. If there are more than one numeric column, the length of col can be either one or number of numeric columns. If there is only one numeric column and type is either segment or h, the length of col can be either one or number of rows of region. pass to circos.lines
lwd	Settings are similar as col. Pass to circos.lines
lty	Settings are similar as col. Pass to circos.lines
type	There is an additional option segment which plot segment lines from start position to end position. Settings are similar as col. Pass to circos.lines.
area	Settings are similar as col. Pass to circos.lines
area.baseline	Deprecated, use baseline instead.
baseline	Settings are similar as col. Pass to circos.lines
border	Settings are similar as col. Pass to circos.lines
pt.col	Settings are similar as col. Pass to circos.lines
cex	Settings are similar as col. Pass to circos.lines
pch	Settings are similar as col. Pass to circos.lines
	mysterious parameters

Details

The function is a low-level graphical function and usually is put in panel. fun when using circos.genomicTrackPlotRegio

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

```
### Not run:
### test bed
circos.par("track.height" = 0.1)
circos.initializeWithIdeogram(plotType = NULL)

bed = generateRandomBed(nr = 100)
circos.genomicTrackPlotRegion(bed, panel.fun = function(region, value, ...) {
    circos.genomicLines(region, value, type = "l", ...)
})

bed1 = generateRandomBed(nr = 100)
bed2 = generateRandomBed(nr = 100)
bed_list = list(bed1, bed2)

circos.genomicTrackPlotRegion(bed_list, panel.fun = function(region, value, ...) {
    i = getI(...)
    circos.genomicLines(region, value, col = i, ...)
})
```

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```
circos.genomicTrackPlotRegion(bed_list, stack = TRUE,
               panel.fun = function(region, value, ...) {
                i = getI(...)
               circos.genomicLines(region, value, col = i, ...)
})
bed = generateRandomBed(nr = 100, nc = 4)
circos.genomicTrackPlotRegion(bed, panel.fun = function(region, value, ...) {
                circos.genomicLines(region, value, col = 1:4, ...)
})
\verb|circos.genomicTrackPlotRegion| (bed, stack = TRUE, panel.fun = function(region, value, \ldots) \{ (beta = true, true
                i = getI(...)
               circos.genomicLines(region, value, col = i, ...)
})
bed = generateRandomBed(nr = 100)
circos.genomicTrackPlotRegion(bed, panel.fun = function(region, value, ...) {
               circos.genomicLines(region, value, type = "segment", lwd = 2, ...)
circos.clear()
## End(Not run)
```

circos.genomicLink

Add links from two sets of genomic positions

Description

Add links from two sets of genomic positions

Usage

```
circos.genomicLink(region1, region2,
    rou = get_most_inside_radius(), rou1 = rou, rou2 = rou,
    col = "black", lwd = par("lwd"), lty = par("lty"), border = col, ...)
```

Arguments

region1	A genomic data frame
region2	A genomic data frame
rou	Pass to circos.link
rou1	Pass to circos.link
rou2	Pass to circos.link
col	Pass to circos.link, length can be either one or nrow of region1

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lwd	Pass to circos.link, length can be either one or nrow of region1
lty	Pass to circos.link, length can be either one or nrow of region1
border	Pass to circos.link, length can be either one or nrow of region1
	Pass to circos.link

Details

Of course, number of rows should be same in region1 and region2.

If you want to have more controls on links, please use circos.link directly.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

Examples

```
## Not run:
set.seed(123)

bed1 = generateRandomBed(nr = 100)
bed1 = bed1[sample(nrow(bed1), 20), ]
bed2 = generateRandomBed(nr = 100)
bed2 = bed2[sample(nrow(bed2), 20), ]
circos.par("track.height" = 0.1, cell.padding = c(0, 0, 0, 0))
circos.initializeWithIdeogram()

circos.genomicLink(bed1, bed2, col = sample(1:5, 20, replace = TRUE), border = NA)
circos.clear()

## End(Not run)
```

circos.genomicPoints Add points to a plotting region, specifically for genomic graphics

Description

Add points to a plotting region, specifically for genomic graphics

Usage

```
circos.genomicPoints(region, value, numeric.column = NULL,
    sector.index = get.cell.meta.data("sector.index"),
    track.index = get.cell.meta.data("track.index"), posTransform = NULL,
    pch = par("pch"), col = par("col"), cex = par("cex"), bg = par("bg"), ...)
```

circos.genomicPoints 35

Arguments

region	A data frame contains 2 columns which correspond to start positions and end positions
value	A data frame contains values and other information
numeric.column	Which column in value data frame should be taken as y-value. If it is not defined, the whole numeric columns in value will be taken.
sector.index	Pass to circos.points
track.index	Pass to circos.points
posTransform	Self-defined function to transform genomic positions, see ${\tt posTransform.default}$ for explanation
col	color of points. If there is only one numeric column, the length of col can be either one or number of rows of region. If there are more than one numeric column, the length of col can be either one or number of numeric columns. Pass to circos.points
pch	Type of points. Settings are similar as col. Pass to circos.points
cex	Size of points. Settings are similar as col. Pass to circos.points
bg	background colors for points.
	Mysterious parameters

Details

The function is a low-level graphical function and usually is put in panel. fun when using circos.genomicTrackPlotRegio

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

```
## Not run:
circos.par("track.height" = 0.1)
circos.initializeWithIdeogram(plotType = NULL)

bed = generateRandomBed(nr = 100)
circos.genomicTrackPlotRegion(bed, panel.fun = function(region, value, ...) {
    circos.genomicPoints(region, value, pch = 16, cex = 0.5, ...)
})

circos.genomicTrackPlotRegion(bed, stack = TRUE, panel.fun = function(region, value, ...) {
    circos.genomicPoints(region, value, pch = 16, cex = 0.5, ...)
    i = getI(...)
    cell.xlim = get.cell.meta.data("cell.xlim")
    circos.lines(cell.xlim, c(i, i), lty = 2, col = "#000000040")
})
bed1 = generateRandomBed(nr = 100)
```

```
bed2 = generateRandomBed(nr = 100)
bed_list = list(bed1, bed2)
# data frame list
circos.genomicTrackPlotRegion(bed_list, panel.fun = function(region, value, ...) {
   cex = (value[[1]] - min(value[[1]]))/(max(value[[1]]) - min(value[[1]]))
   i = getI(...)
   circos.genomicPoints(region, value, cex = cex, pch = 16, col = i, ...)
})
circos.genomicTrackPlotRegion(bed_list, stack = TRUE,
   panel.fun = function(region, value, ...) {
   cex = (value[[1]] - min(value[[1]]))/(max(value[[1]]) - min(value[[1]]))
   i = getI(...)
   circos.genomicPoints(region, value, cex = cex, pch = 16, col = i, ...)
   cell.xlim = get.cell.meta.data("cell.xlim")
   circos.lines(cell.xlim, c(i, i), lty = 2, col = "#00000040")
})
bed = generateRandomBed(nr = 100, nc = 4)
circos.genomicTrackPlotRegion(bed, panel.fun = function(region, value, ...) {
   cex = (value[[1]] - min(value[[1]]))/(max(value[[1]]) - min(value[[1]]))
   circos.genomicPoints(region, value, cex = 0.5, pch = 16, col = 1:4, ...)
})
cex = (value[[1]] - min(value[[1]]))/(max(value[[1]]) - min(value[[1]]))
   i = getI(...)
   circos.genomicPoints(region, value, cex = cex, pch = 16, col = i, ...)
   cell.xlim = get.cell.meta.data("cell.xlim")
   circos.lines(cell.xlim, c(i, i), lty = 2, col = "#00000040")
})
circos.clear()
## End(Not run)
```

circos.genomicPosTransformLines

Add genomic position transformation lines between tracks

Description

Add genomic position transformation lines between tracks

Usage

```
circos.genomicPosTransformLines(data, track.height = 0.1, posTransform = NULL,
   horizontalLine = c("none", "top", "bottom", "both"), track.margin = c(0, 0),
   direction = c("inside", "outside"), col = "black", lwd = par("lwd"),
   lty = par("lty"), ...)
```

Arguments

data A data frame containing genomic data

track.height Height of the track

posTransform Genomic position transformation function, see posTransform.default for an

example.

horizontalLine Whether to draw horizontal lines which indicate region width

track.margin Margin of tracks

direction Type of the transformation. inside means position transformed track are lo-

cated inside and outside means position transformed track are located outside.

col Color of lines, can be length of one or nrow of data

lwd Width of lineslty Style of lines

... pass to circos.trackPlotRegion

Details

There is one representative situation when such position transformation needs to be applied. For example, there are two sets of regions in a chromosome in which regions in one set regions are quite densely to each other and regions in other set are far from others. Heatmap or text is going to be drawn on the next track. If there is no position transformation, heatmap or text for those dense regions would be overlapped and hard to identify, also ugly to visualize. Thus, a way to transform original positions to new positions would help for the visualization.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

```
f = colorRamp2(breaks = c(-1, 0, 1), colors = c("green", "black", "red"))
circos.genomicTrackPlotRegion(bed, stack = TRUE, panel.fun = function(region, value, ...) {
           circos.genomicRect(region, value, col = f(value[[1]]),
                      border = f(value[[1]]), posTransform = posTransform.default, ...)
}, bg.border = NA)
circos.clear()
circos.par(cell.padding = c(0, 0, 0, 0))
circos.initializeWithIdeogram(plotType = NULL)
bed = generateRandomBed(nr = 20, nc = 4)
circos.genomicTrackPlotRegion(bed, ylim = c(0, 1), panel.fun = function(region, value, ...) {
           circos.genomicText(region, value, y = 0, adj = c(1, 0.5),
                      labels = "gene", facing = "reverse.clockwise",
                      posTransform = posTransform.default)
}, bg.border = NA)
circos.genomicPosTransformLines(bed, posTransform = posTransform.default,
           horizontalLine = "bottom", direction = "outside", track.height = 0.1)
cytoband = read.cytoband()$df
 {\tt circos.genomicTrackPlotRegion(cytoband, stack = TRUE, panel.fun = function(region, value, \ldots)} \ \{ (in the constant of th
           circos.genomicRect(region, value, col = cytoband.col(value$V5), border = NA, ...)
}, track.height = 0.05)
circos.clear()
## End(Not run)
```

circos.genomicRainfall

Genomic rainfall plot

Description

Genomic rainfall plot

Usage

```
circos.genomicRainfall(data, mode = "min", ylim = NULL, col = "black",
    pch = par("pch"), cex = par("cex"), normalize_to_width = FALSE, ...)
```

circos.genomicRainfall 39

Arguments

data	A bed-file-like data frame or a list of data frames	
mode	how to calculate the distance of two neighbouring regions, pass to rainfallTransform	
ylim	ylim for rainfall plot track. If normalize_to_width is FALSE, the value should correspond to log10(dist+1), and if normalize_to_width is TRUE, the value should correspond to log2(rel_dist).	
col	Color of points. It should be length of one. If data is a list, the length of col can also be the length of the list.	
pch	Style of points	
cex	Size of points	
normalize_to_width		
	If it is TRUE, the value is the relative distance divided by the width of the region.	
	Pass to circos.trackPlotRegion	

Details

This is high-level graphical function, which mean, it will create a new track.

Rainfall plot can be used to visualize distribution of regions. On the plot, y-axis corresponds to the distance to neighbour regions (log-based). So if there is a drop-down on the plot, it means there is a cluster of regions at that area.

On the plot, y-axis are log10-transformed.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

```
## Not run:
load(system.file(package = "circlize", "extdata", "DMR.RData"))

# rainfall
circos.initializeWithIdeogram(plotType = c("axis", "labels"))

bed_list = list(DMR_hyper, DMR_hypo)
circos.genomicRainfall(bed_list, pch = 16, cex = 0.4, col = c("#FF000080", "#0000FF80"))

circos.genomicDensity(bed_list[[1]], col = c("#FF000080"), track.height = 0.1)
circos.genomicDensity(bed_list[[2]], col = c("#0000FF80"), track.height = 0.1)
circos.clear()

## End(Not run)
```

40 circos.genomicRect

circos.genomicRect Draw rectangle-like grid, specifically for genomic graphics

Description

Draw rectangle-like grid, specifically for genomic graphics

Usage

```
circos.genomicRect(region, value = NULL,
   ytop = NULL, ybottom = NULL, ytop.column = NULL, ybottom.column = NULL,
   sector.index = get.cell.meta.data("sector.index"),
   track.index = get.cell.meta.data("track.index"), posTransform = NULL,
   col = NA, border = "black", lty = par("lty"), ...)
```

Arguments

region	A data frame contains 2 column which correspond to start position and end position
value	A data frame contains values and other information
ytop	A vector or a single value indicating top position of rectangles
ybottom	A vector or a single value indicating bottom position of rectangles
ytop.column	If ytop is in value, the index of the column
ybottom.column	If ybottom is in value, the index of the column
sector.index	Pass to circos.rect
track.index	Pass to circos.rect
posTransform	Self-defined function to transform genomic positions, see ${\tt posTransform.default}$ for explaination
col	The length of col can be either one or number of rows of region. Pass to circos.rect
border	Settings are similar as col. Pass to circos.rect
lty	Settings are similar as col. Pass to circos.rect
	Mysterious parameters

Details

The function is a low-level graphical function and usually is put in panel. fun when using circos.genomicTrackPlotRegio

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

circos.genomicRect 41

```
## Not run:
### rect matrix
circos.par("track.height" = 0.1, cell.padding = c(0, 0, 0, 0))
circos.initializeWithIdeogram(plotType = NULL)
bed = generateRandomBed(nr = 100, nc = 4)
circos.genomicTrackPlotRegion(bed, stack = TRUE, panel.fun = function(region, value, ...) {
   circos.genomicRect(region, value, col = sample(1:10, nrow(region), replace = TRUE),
       border = NA, ...)
   i = getI(...)
   cell.xlim = get.cell.meta.data("cell.xlim")
   \#circos.lines(cell.xlim, c(i, i), lty = 2, col = "#00000040")
}, bg.border = NA)
circos.genomicPosTransformLines(bed, posTransform = posTransform.default,
   horizontalLine = "top")
circos.genomicTrackPlotRegion(bed, stack = TRUE, panel.fun = function(region, value, ...) {
   circos.genomicRect(region, value, col = sample(1:10, nrow(region), replace = TRUE),
       border = NA, posTransform = posTransform.default, ...)
   i = getI(...)
   cell.xlim = get.cell.meta.data("cell.xlim")
   \#circos.lines(cell.xlim, c(i, i), lty = 2, col = "#00000040")
}, bg.border = NA)
circos.genomicPosTransformLines(bed, posTransform = posTransform.default,
   direction = "outside", horizontalLine = "bottom")
circos.genomicTrackPlotRegion(bed, stack = TRUE, panel.fun = function(region, value, ...) {
   circos.genomicRect(region, value, col = sample(1:10, nrow(region), replace = TRUE),
       border = NA, ...)
   i = getI(...)
   cell.xlim = get.cell.meta.data("cell.xlim")
   \#circos.lines(cell.xlim, c(i, i), lty = 2, col = "#00000040")
}, bg.border = NA)
circos.clear()
### rect from bed list
circos.par("track.height" = 0.1, cell.padding = c(0, 0, 0, 0))
circos.initializeWithIdeogram(plotType = NULL)
bed1 = generateRandomBed(nr = 100)
bed2 = generateRandomBed(nr = 100)
bed_list = list(bed1, bed2)
f = colorRamp2(breaks = c(-1, 0, 1), colors = c("green", "black", "red"))
circos.genomicTrackPlotRegion(bed_list, stack = TRUE,
   panel.fun = function(region, value, ...) {
```

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```
circos.genomicRect(region, value, col = f(value[[1]]),
        border = NA, ...)
    i = getI(...)
    cell.xlim = get.cell.meta.data("cell.xlim")
    circos.lines(cell.xlim, c(i, i), lty = 2, col = "#000000")
})
circos.genomicTrackPlotRegion(bed_list, ylim = c(0, 3),
    panel.fun = function(region, value, ...) {
    i = getI(...)
   circos.genomicRect(region, value, ytop = i+0.4, ybottom = i-0.4, col = f(value[[1]]),
        border = NA, ...)
    cell.xlim = get.cell.meta.data("cell.xlim")
    circos.lines(cell.xlim, c(i, i), lty = 2, col = "#000000")
})
circos.genomicTrackPlotRegion(bed1, panel.fun = function(region, value, ...) {
    circos.genomicRect(region, value, col = "red", border = NA, ...)
})
circos.genomicTrackPlotRegion(bed_list, panel.fun = function(region, value, ...) {
    i = getI(...)
    circos.genomicRect(region, value, col = i, border = NA, ...)
})
circos.clear()
## End(Not run)
```

circos.genomicText

Draw text in a cell, specifically for genomic graphics

Description

Draw text in a cell, specifically for genomic graphics

Usage

```
circos.genomicText(region, value = NULL, y = NULL, labels = NULL, labels.column = NULL,
    numeric.column = NULL, sector.index = get.cell.meta.data("sector.index"),
    track.index = get.cell.meta.data("track.index"), posTransform = NULL,
    direction = NULL, facing = "inside", niceFacing = FALSE,
    adj = par("adj"), cex = 1, col = "black", font = par("font"), padding = 0,
    extend = 0, align_to = "region", ...)
```

circos.genomicText 43

Arguments

region	A data frame contains 2 column which correspond to start position and end position
value	A data frame contains values and other information
у	A vector or a single value indicating position of text.
labels	Labels of text corresponding to each genomic positions
labels.column	If labels are in value, index of column in value
numeric.column	Which column in value data frame should be taken as y-value. If it is not defined, only the first numeric columns in value will be taken.
sector.index	Pass to circos.rect
track.index	Pass to circos.rect
posTransform	Self-defined function to transform genomic positions, see ${\tt posTransform.default}$ for explanation
facing	Passing to circos.text. Settings are similar as col
niceFacing	Should the facing of text be adjusted to fit human eyes?
direction	Deprecated, use facing instead.
adj	Pass to circos.text. Settings are similar as col
cex	Pass to circos.text. Settings are similar as col
col	Pass to circos.text. The length of col can be either one or number of rows of region.
font	Pass to circos.text. Settings are similar as col
padding	pass to posTransform if it is set as posTransform.text
extend	pass to posTransform if it is set as posTransform.text
align_to	pass to posTransform if it is set as posTransform.text
	Mysterious parameters

Details

 $The function is a low-level graphical function and usually is put in panel. fun when using \verb|circos.genomicTrackPlotRegio|| and the panel of the function of the panel of th$

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

```
## Not run:
circos.par("track.height" = 0.1, cell.padding = c(0, 0, 0, 0))
circos.initializeWithIdeogram(plotType = NULL)

bed = generateRandomBed(nr = 20)

circos.genomicTrackPlotRegion(bed, ylim = c(0, 1), panel.fun = function(region, value, ...) {
```

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```
circos.genomicText(region, value, y = 0.5, labels = "text", ...)
})

bed = cbind(bed, sample(letters, nrow(bed), replace = TRUE))
circos.genomicTrackPlotRegion(bed, panel.fun = function(region, value, ...) {
    circos.genomicText(region, value, labels.column = 2, ...)
})

circos.clear()

## End(Not run)
```

circos.genomicTrack

Create a track for genomic graphics

Description

Create a track for genomic graphics

Usage

```
circos.genomicTrack(...)
```

Arguments

```
... pass to circos.genomicTrackPlotRegion
```

Details

 $shortcut\ function\ of\ \verb|circos.genomicTrackPlotRegion|.$

```
\# There is no example NULL
```

circos.genomicTrackPlotRegion

Create a track for genomic graphics

Description

Create a track for genomic graphics

Usage

```
circos.genomicTrackPlotRegion(data = NULL, ylim = NULL, stack = FALSE,
    numeric.column = NULL, jitter = 0,
    panel.fun = function(region, value, ...) {NULL}, ...)
```

Arguments

data A bed-file-like data frame or a list of data frames

ylim If it is NULL, the value will be calculated from data. If stack is set to TRUE, this

value is ignored.

stack whether to plot in a "stack" mode.

numeric.column Columns of numeric values in data that will be used for plotting. If data is

a data frame list, numeric.column should be either length of one or length of data. If value of numeric.column is not set, its value will depend on the structure of data. If data is a data frame, the default value for numeric.column is all the numeric column starting from the fourth column. If data is a list of data frame, the default value for numeric.column is a vector which have the same length as data and the value in default numeric.column is the index of the first

numeric column in corresponding data frame.

jitter Numeric. Only works for adding points in circos.genomicTrackPlotRegion

under stack mode

panel.fun Self-defined function which will be applied on each sector. Please not it is dif-

ferent from that in circos.trackPlotRegion. In this function, there are two arguments (region and value) plus In them, region is a two-column data frame with start positions and end positions in current genomic category (e.g. chromosome). value is a data frame which is derived from data but excluding the first three columns. Rows in value correspond to rows in region. ... is mandatory and is used to pass internal parameters to other functions. The definition of value will be different according to different input data (data frame or list of data frame) and different settings (stacked or not), please refer to 'details'

section and vignettes to detailed explanation.

... Pass to circos.trackPlotRegion.

Details

Similar as circos.trackPlotRegion, users can add customized graphics by panel.fun, but the behaviour of panel.fun will change depending on users' input data and stack setting.

When data is a single data frame, region in panel. fun is a data frame containing the second and third column in data in 'current' genomic category (e.g. current chromosome). value is also a data frame containing columns in data excluding the first three columns.

When data is a list containing data frames, panel.fun will be applied iteratively on each data frame, thus, region is extracted from the data frame which is in the current iteration. For example, if data contains two data frames, panel.fun will be applied with the first data frame in current chromosome and then applied with the second data frame in the same chromosome.

If stack is set to TRUE, ylim will be re-defined. in stack mode, the y-axis will be splitted into several part with equal height and graphics will be drawn on each 'horizontal' lines (y = 1, 2, ...). In this case:

When data is a single data frame containing one or more numeric columns, each numeric column defined in numeric.column will be treated as a single unit. ylim is re-defined to c(0.5,n+0.5) in which n is number of numeric columns. panel.fun will be applied iteratively on each numeric column. In each iteration, in panel.fun, region is still the genomic regions in current genomic category, but value contains current numeric column plus all non-numeric columns. Under stack mode, in panel.fun, all low-level genomic graphical functions will draw on the 'horizontal line' y = i in which i is the index of current numeric column and the value of i can be obtained by getI.

When data is a list containing data frames, each data frame will be treated as a single unit. The situation is quite similar as described in previous paragraph. ylim is re-defined to c(0.5,n+0.5) in which n is number of data frames. panel.fun will be applied iteratively on each data frame. In each iteration, in panel.fun, region is still the genomic regions in current genomic category, and value contains columns in current data frame excluding the first three columns. Under stack mode, in panel.fun, all low-level genomic graphical functions will draw on the 'horizontal line' y = i in which i is the index of current data frame.

Being different from panel.fun in circos.trackPlotRegion, there should be an additional argument ... in panel.fun. This additional argument is used to pass hidden values to low-level graphical functions. So if you are using functions like circos.genomicPoints, you should also add ... as an additional argument into circos.genomicPoints.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

Examples

There is no example NULL

circos.info 47

circos.info	Get information of the circular plot	

Description

Get information of the circular plot

Usage

```
circos.info(sector.index = NULL, track.index = NULL, plot = FALSE)
```

Arguments

sector.index Which sectors you want to look at? It can be a vector.
track.index Which tracks you want to look at? It can be a vector.
plot Whether to add information on the plot

Details

It tells you the basic parameters for sectors/tracks/cells. If both sector.index and track.index are set to NULL, the function would print index for all sectors and all tracks. If sector.index and/or track.index are set, the function would print xlim, ylim, cell.xlim, cell.ylim, xplot, yplot, track.margin and cell.padding for every cell in specified sectors and tracks. Also, the function will print index of your current sector and current track.

If plot is set to TRUE, the function will plot the index of the sector and the track for each cell on the figure.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

```
factors = letters[1:4]
circos.initialize(factors, xlim = c(0, 1))
circos.trackPlotRegion(ylim = c(0, 1))
circos.trackPlotRegion(ylim = c(0, 1))
circos.info(sector.index = "a", track.index = 1)
circos.info(sector.index = "a", track.index = 1:2)
circos.info(sector.index = c("a", "b"), track.index = 1)
circos.info(sector.index = "a")
circos.info(track.index = 1)
circos.info()
circos.info(plot = TRUE)
circos.clear()
```

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circos.initialize

Initialize the circular layout

Description

Initialize the circular layout

Usage

```
circos.initialize(factors, x = NULL, xlim = NULL, sector.width = NULL)
```

Arguments

factors A factor variable or a character vector which represent data categories

x Data on x-axes, a vector

xlim Ranges for values on x-axes, see "details" section for explanation of the format

sector. width Width for each sector. The length of the vector should be either 1 which means

all sectors have same width or as same as the number of sectors. Values for the vector are relative, and they will be scaled by dividing their summation. By default, it is NULL which means the width of sectors correspond to the data range

in sectors.

Details

The function allocates the sectors according to the values on x-axis. The number of sectors are determined by the factors and the order of sectors are determined by the levels of factors. In this function, the start and end position for each sector on the circle (measured by degree) are calculated according to the values on x-axis or by x1im.

If x is set, the length of x must be equal to the length of factors. Then the data range for each sector are calculated from x by splitting factors.

If xlim is set, it should be a vector containing two numbers or a matrix with 2 columns. If xlim is a 2-element vector, it means all sector share the same xlim. If xlim is a 2-column matrix, the number of rows should be equal to the number of categories identified by factors, then each row of xlim corresponds to the data range for each sector and the order of rows is corresponding to the order of levels of factors. If xlim is a matrix for which row names cover all sector names, xlim is automatically adjusted.

Normally, width of sectors will be calculated internally according to the data range in sectors. But you can still set the width manually. However, it is not always a good idea to change the default sector width since the width can reflect the range of data in sectors. However, in some cases, it is useful to manually set the width such as you want to zoom some part of the sectors.

The function finally calls plot with enforing aspect ratio to be 1 and be ready for adding graphics.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

See Also

http://jokergoo.github.io/circlize_book/book/circular-layout.html

Examples

```
circos.initialize(factors = sample(letters[1:4], 20, replace = TRUE), xlim = c(0, 1))
circos.info()
circos.clear()

circos.initialize(factors = sample(letters[1:4], 20, replace = TRUE), xlim = cbind(1:4, 1:4*2))
circos.info()
circos.clear()

circos.initialize(factors = sample(letters[1:4], 20, replace = TRUE), x = rnorm(20))
circos.info()
circos.clear()
```

circos.initializeWithIdeogram

Initialize the circular layout with an ideogram

Description

Initialize the circular layout with an ideogram

Usage

```
circos.initializeWithIdeogram(cytoband = system.file(package = "circlize",
    "extdata", "cytoBand.txt"), species = NULL, sort.chr = TRUE,
    chromosome.index = usable_chromosomes(species), major.by = NULL,
    plotType = c("ideogram", "axis", "labels"),
    track.height = NULL, ideogram.height = convert_height(2, "mm"),
    ...)
```

Arguments

cytoband A path of the cytoband file or a data frame that already contains cytoband data.

By default it is cytoband for hg19. Pass to read.cytoband.

species Abbreviations of species. e.g. hg19 for human, mm10 for mouse. If this value

is specified, the function will download cytoBand.txt.gz from UCSC website automatically. If there is no cytoband for user's species, it will keep on trying to

download chromInfo file. Pass to read.cytoband or read.chromInfo.

chromosome.index

subset of chromosomes, also used to reorder chromosomes.

sort.chr Whether chromosome names should be sorted (first sort by numbers then by

letters). If chromosome index is set, this argumeth is enforced to FALSE

major.by Increment of major ticks. Pass to circos.genomicInitialize.

plotType Which tracks should be drawn. ideogram for ideogram rectangle, axis for genomic axis and labels for chromosome names. If there is no ideogram for specified species, ideogram will be enforced to be excluded. If it is set to NULL, the function just initialize the plot but draw nothing.

track.height Height of the track which contains "axis" and "labels".

ideogram.height

Height of the ideogram track

Pass to circos.genomicInitialize.

Details

The function will initialize the circular plot in which each sector corresponds to a chromosome. You can control the order of chromosomes by chromosome.index or by sort.chr, or by setting a special format of cytoband (please refer to read.cytoband to find out how to control a proper cytoband).

The function finally pass data to circos.genomicInitialize to initialize the circular plot.

The style of ideogram is almost fixed, but you can customize it with your self-sefined code. Refer to vignette for demonstration.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

```
## Not run:
circos.initializeWithIdeogram()
cytoband.file = system.file(package = "circlize"),
    "extdata", "cytoBand.txt")
circos.initializeWithIdeogram(cytoband.file)
cytoband.df = read.table(cytoband.file, colClasses = c("character", "numeric",
    "numeric", "character", "character"), sep = "\t")
circos.initializeWithIdeogram(cytoband.df)
circos.initializeWithIdeogram(species = "hg18")
circos.initializeWithIdeogram(species = "mm10")
circos.initializeWithIdeogram(chromosome.index = c("chr1", "chr2"))
cytoband = read.table(cytoband.file, colClasses = c("character", "numeric",
    "numeric", "character", "character"), sep = "\t")
circos.initializeWithIdeogram(cytoband, sort.chr = FALSE)
cytoband[[1]] = factor(cytoband[[1]], levels = paste0("chr", c(22:1, "X", "Y")))
circos.initializeWithIdeogram(cytoband, sort.chr = FALSE)
```

circos.lines 51

circos.lines

Add lines to the plotting region

Description

Add lines to the plotting region

Usage

```
circos.lines(x, y, sector.index = get.cell.meta.data("sector.index"),
    track.index = get.cell.meta.data("track.index"),
    col = ifelse(area, "grey", par("col")), lwd = par("lwd"), lty = par("lty"),
    type = "l", straight = FALSE, area = FALSE, area.baseline = NULL,
    border = "black", baseline = "bottom", pt.col = par("col"), cex = par("cex"),
    pch = par("pch"))
```

Arguments

X	Data points on x-axis, measured in "current" data coordinate
у	Data points on y-axis, measured in "current" data coordinate
sector.index	Index for the sector
track.index	Index for the track
col	Line color
lwd	line width
lty	line style
type	line type, similar as type argument in lines, but only in c("1", "o", "h", "s")

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straight whether draw straight lines between points.

area whether to fill the area below the lines. If it is set to TRUE, col controls the filled color in the area and border controls color of the line.

area.baseline deprecated, use baseline instead.

baseline the base line to draw areas. By default it is the minimal of y-range (bottom). It

can be a string or a number. If a string, it should be one of bottom and top. This

argument also works if type is set to h.

border color for border of the area
pt.col if type is "o", point color
cex if type is "o", point size
pch if type is "o", point type

Details

Normally, straight lines in the Cartesian coordinate have to be transformed into curves in the circular layout. But if you do not want to do such transformation you can use this function just drawing straight lines between points by setting straight to TRUE.

Drawing areas below lines can help to identify the direction of y-axis in cells (since it is a circle). This can be done by specifying area to TURE.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

```
factors = letters[1:9]
circos.par(points.overflow.warning = FALSE)
circos.initialize(factors = factors, xlim = c(0, 10))
circos.trackPlotRegion(factors = factors, ylim = c(0, 10), track.height = 0.5)
circos.lines(sort(runif(10)*10), runif(10)*8, sector.index = "a")
circos.text(5, 9, "type = 'l'", sector.index = "a", facing = "outside")
circos.lines(sort(runif(10)*10), runif(10)*8, sector.index = "b", type = "o")
circos.text(5, 9, "type = 'o'", sector.index = "b", facing = "outside")
circos.lines(sort(runif(10)*10), runif(10)*8, sector.index = "c", type = "h")
circos.text(5, 9, "type = 'h'", sector.index = "c", facing = "outside")
circos.lines(sort(runif(10)*10), runif(10)*8, sector.index = "d", type = "h", baseline = 5)
circos.text(5, 9, "type = 'h', baseline = 5", sector.index = "d", facing = "outside")
circos.lines(sort(runif(10)*10), runif(10)*8, sector.index = "e", type = "s")
circos.text(5, 9, "type = 's'", sector.index = "e", facing = "outside")
circos.lines(sort(runif(10)*10), runif(10)*8, sector.index = "f", area = TRUE)
circos.text(5, 9, "type = 'l', area = TRUE", sector.index = "f")
```

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```
circos.lines(sort(runif(10)*10), runif(10)*8, sector.index = "g", type = "o", area = TRUE)
circos.text(5, 9, "type = 'o', area = TRUE", sector.index = "g")

circos.lines(sort(runif(10)*10), runif(10)*8, sector.index = "h", type = "s", area = TRUE)
circos.text(5, 9, "type = 's', area = TRUE", sector.index = "h")

circos.lines(sort(runif(10)*10), runif(10)*8, sector.index = "i", area = TRUE, baseline = "top")
circos.text(5, 9, "type = 'l', area = TRUE\nbaseline = 'top'", sector.index = "i")

circos.clear()
```

circos.link

Draw links between points or/and intervals

Description

Draw links between points or/and intervals

Usage

```
circos.link(sector.index1, point1, sector.index2, point2,
    rou = get_most_inside_radius(),
    rou1 = rou, rou2 = rou, h = NULL, h.ratio = 0.5, w = 1, h2 = h, w2 = w,
    col = "black", lwd = par("lwd"), lty = par("lty"), border = col,
    directional = 0, arr.length = ifelse(arr.type == "big.arrow", 0.02, 0.4),
    arr.width = arr.length/2, arr.type = "triangle", arr.lty = lty,
    arr.lwd = lwd, arr.col = col)
```

Arguments

sector.index1	Index for the first sector where one link end locates	
point1	A single value or a numeric vector of length 2. If it is a 2-elements vector, then the link would be a belt/ribbon.	
sector.index2	Index for the other sector where the other link end locates	
point2	A single value or a numeric vector of length 2. If it is a 2-elements vector, then the link would be a belt/ribbon.	
rou	The position of the the link ends (if rou1 and rou2 are not set). It is the percentage of the radius of the unit circle. By default its value is the position of bottom margin of the most inner track.	
rou1	The position of end 1 of the link.	
rou2	The position of end 2 of the link.	
h	Height of the link, measured as percent to the radius to the unit circle. By default it is automatically infered.	
h.ratio	systematically change the link height. The value is between 0 and 1.	

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W	Since the link is a Bezier curve, it controls the shape of Bezier curve.
h2	Height of the bottom edge of the link if it is a ribbon.
w2	Shape of the bottom edge of the link if it is a ribbon.
col	Color of the link. If the link is a ribbon, then it is the filled color for the ribbon.
lwd	Line (or border) width
lty	Line (or border) style
border	If the link is a ribbon, then it is the color for the ribbon border.
directional	0 for no direction, 1 for direction from point1 to point2, -1 for direction from point2 to point1. 2 for two directional. The direction is important when arrow heads are added.
arr.width	Width of the arrows, pass to Arrowhead.
arr.type	Type of the arrows, pass to Arrowhead. Default value is triangle. There is an additional option big.arrow.
arr.length	Length of the arrows, measured in 'cm', pass to Arrowhead. If arr. type is set to big.arrow, the value is percent to the radius of the unit circle.
arr.col	Color of the arrows, pass to Arrowhead.
arr.lwd	Line width of arrows, pass to Arrowhead.
arr.lty	Line type of arrows, pass to Arrowhead.

Details

Links are implemented as quadratic Bezier curves (https://en.wikipedia.org/wiki/B%C3%A9zier_curve#Rational_B.C3.A9zier_curves).

Drawing links does not create any track. So you can think it is independent of the tracks.

By default you only need to set sector.index1, point1, sector.index2 and point2. The links would look nice.

Please refer to the vignette for detailed explanation.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

See Also

http://jokergoo.github.io/circlize_book/book/graphics.html#links

```
factors = letters[1:8]
circos.par(points.overflow.warning = FALSE)
circos.initialize(factors = factors, xlim = c(0, 10))
circos.track(factors = factors, ylim = c(0, 1), bg.col = "grey",
    bg.border = NA, track.height = 0.05)
circos.link("a", 5, "c", 5, border = 1)
circos.link("b", 5, "d", c(4, 6), border = 1)
```

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```
circos.link("a", c(2, 3), "f", c(4, 6), border = 1)
circos.link("e", c(2, 3), "g", 5, border = 1)
circos.clear()

circos.par(points.overflow.warning = FALSE)
circos.initialize(factors = factors, xlim = c(0, 10))
circos.track(factors = factors, ylim = c(0, 1), bg.col = "grey",
    bg.border = NA, track.height = 0.05)
circos.link("a", 5, "b", 5, directional = 1, arr.length = 0.2)
circos.link("c", c(3, 7), "d", c(3, 7), directional = 1,
    arr.col = "white", arr.length = 0.2)
circos.link("e", c(4, 6), "f", c(4, 6), directional = 1,
    arr.type = "big.arrow", arr.length = 0.04)
circos.clear()
```

circos.nested

Nested zooming with two circular plots

Description

Nested zooming with two circular plots

Usage

```
circos.nested(f1, f2, correspondance, connection_height = convert_height(5, "mm"),
    connection_col = NA, connection_border = "black",
    connection_lty = par("lty"), connection_lwd = par("lwd"),
    adjust_start_degree = TRUE)
```

Arguments

f2

f1 a self-defined function for making the first circular plot. The function should have no argument.

a self-defined function for making the second circular plot. The function should have no argument.

correspondance a six-column data frame which contains correspondance between the coordinates in two circular plots

connection_height

the height of the connection track, measured as the percent to the radius of the unit circle. The value can be specified by uh or convert_height with absolute units.

connection_col filled color of the connection track. The value can be a vector with same length as number of rows of correspondance

connection_border

border color of the connection track.

connection_lty line style of the connection track borders

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```
connection_lwd line width of the connection track borders
adjust_start_degree
```

If circos.par(start.degree = ...) is not set in f2(), the start degree for the second circular plot will be adjusted to make the distance of sectors between the two plots to the minimal.

Details

The function visualizes zoomings by combining two circular plots into one page where one is the normal circular plot and the other one only contains regions that need to be zoomed. This function automatically arranges the two plots to make it easy to correspond between the original and the zoomed sectors.

Since the function needs to know the information of the two circular plots, please do not call circos.clear in either f1() or f2(). It will be called internally in circos.nested.

If adjust_start_degree is set to TRUE, start.degree should not be set in f2(). Also canvas.xlim and canvas.ylim are reset in f2(), they should not be set in f2() either.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

See Also

http://jokergoo.github.io/circlize_book/book/nested-zooming.html

```
## Not run:
#### simulate data ####
set.seed(123)
df = data.frame(cate = sample(letters[1:8], 400, replace = TRUE),
               x = runif(400),
               y = runif(400),
                stringsAsFactors = FALSE)
df = df[order(df[[1]], df[[2]]), ]
rownames(df) = NULL
df$interval_x = as.character(cut(df$x, c(0, 0.2, 0.4, 0.6, 0.8, 1.0)))
df$name = paste(df$cate, df$interval_x, sep = ":")
df$start = as.numeric(gsub("^\\((\\d(\\.\\d)?).*(\\d(\\.\\d)?)]", "\\1", df$interval_x))
dfend = as.numeric(gsub("^\\((\\d(\\.\\d)?),(\\d(\\.\\d)?)]$", "\\3", df$interval_x))
nm = sample(unique(df$name), 20)
df2 = df[df$name %in% nm, ]
correspondance = unique(df2[, c("cate", "start", "end", "name", "start", "end")])
zoom_sector = unique(df2[, c("name", "start", "end", "cate")])
zoom_data = df2[, c("name", "x", "y")]
data = df[, 1:3]
sector = data.frame(cate = letters[1:8], start = 0, end = 1, stringsAsFactors = FALSE)
sector_col = structure(rand_color(8, transparency = 0.5), names = letters[1:8])
```

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```
#### define two circular plots ####
f1 = function() {
   circos.par(gap.degree = 10)
    circos.initialize(sector[, 1], xlim = sector[, 2:3])
    circos.track(data[[1]], x = data[[2]], y = data[[3]], ylim = c(0, 1),
        panel.fun = function(x, y) {
            1 = correspondance[[1]] == CELL_META$sector.index
            if(sum(1)) {
                for(i in which(l)) {
                    circos.rect(correspondance[i, 2], CELL_META$cell.ylim[1],
                                correspondance[i, 3], CELL_META$cell.ylim[2],
                                col = sector_col[CELL_META$sector.index],
                                border = sector_col[CELL_META$sector.index])
                }
            circos.points(x, y, pch = 16, cex = 0.5)
            circos.text(CELL_META$xcenter, CELL_META$ylim[2] + uy(2, "mm"),
                CELL_META$sector.index, niceFacing = TRUE, adj = c(0.5, 0))
    })
}
f2 = function() {
    circos.par(gap.degree = 2, cell.padding = c(0, 0, 0, 0))
    circos.initialize(zoom_sector[[1]], xlim = as.matrix(zoom_sector[, 2:3]))
    circos.track(zoom_data[[1]], x = zoom_data[[2]], y = zoom_data[[3]],
        panel.fun = function(x, y) {
            circos.points(x, y, pch = 16, cex = 0.5)
        }, bg.col = sector_col[zoom_sector$cate],
        track.margin = c(0, 0))
circos.nested(f1, f2, correspondance, connection_col = sector_col[correspondance[[1]]])
## End(Not run)
```

circos.par

Parameters for the circular layout

Description

Parameters for the circular layout

Usage

```
circos.par(..., RESET = FALSE, READ.ONLY = NULL, LOCAL = FALSE, ADD = FALSE)
```

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Arguments

... Arguments for the parameters, see "details" section

RESET reset to default values

READ.ONLY please ignore
LOCAL please ignore
ADD please ignore

Details

Global parameters for the circular layout. Currently supported parameters are:

- start.degree The starting degree from which the circle begins to draw. Note this degree is measured in the standard polar coordinate which means it is always reverse-clockwise.
- gap.degree Gap between two neighbour sectors. It can be a single value or a vector. If it is a vector, the first value corresponds to the gap after the first sector.
- gap.after identical to gap.degree option, but a more understandable name. Modifying this option will also affect gap.degree.
- track.margin Like margin in Cascading Style Sheets (CSS), it is the blank area out of the plotting region, also outside of the borders. Since left and right margin are controlled by gap.degree, only bottom and top margin need to be set. And all cells in a same track share the same margins, and that's why this parameter is called track.margin. The value for the track.margin is the percentage according to the radius of the unit circle. convert_height can be used to set to an absolute unit (e.g cm/inche).
- unit.circle.segments Since curves are simulated by a series of straight lines, this parameter controls the amount of segments to represent a curve. The minimal length of the line segmentation is the length of the unit circle (2pi) divided by unit.circoe.segments. More segments means better approximation for the curves while larger size if you generate figures as PDF format.
- cell.padding Padding of the cell. Like padding in Cascading Style Sheets (CSS), it is the blank area around the plotting regions, but within the borders. The parameter has four values, which controls the bottom, left, top and right paddings respectively. The first and the third padding values are the percentages according to the radius of the unit circle and the second and fourth values are degrees. Similar as track.margin option, the first and the third value can be set by convert_height to an absolute unit.
- track.height The default height of tracks. It is the percentage according to the radius of the unit circle. The height includes the top and bottom cell paddings but not the margins. convert_height can be used to set the height to an absolute unit.
- points.overflow.warning Since each cell is in fact not a real plotting region but only an ordinary rectangle, it does not eliminate points that are plotted out of the region. So if some points are out of the plotting region, circlize would continue drawing the points and printing warnings. In some cases, draw something out of the plotting region is useful, such as draw some legend or text. Set this value to FALSE to turn off the warnings.
- canvas.xlim The coordinate for the canvas. Because circlize draws everything (or almost everything) inside the unit circle, the default canvas.xlim and canvas.ylim for the canvas would be all c(-1,1). However, you can set it to a more broad interval if you want to draw

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other things out of the circle. By choosing proper canvas.xlim and canvas.ylim, you can draw part of the circle. E.g. setting canvas.xlim to c(0,1) and canvas.ylim to c(0,1) would only draw circle in the region of (0, pi/2).

```
canvas. ylim The coordinate for the canvas. By default it is c(-1,1) clock.wise The direction for adding sectors. Default is TRUE.
```

Similar as par, you can get the parameter values by specifying the names of parameters and you can set the parameter values by specifying a named list which contains the new values.

gap.degree, start.degree, canvas.xlim, canvas.ylim and clock.wise only be set before the initialization of the circular layout (i.e. before calling circos.initialize) because these values will not be changed after adding sectors on the circle. The left and right padding for cell.padding will also be ignored after the initialization because all cells in a sector would share the same left and right paddings.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

Examples

```
# There is no example NULL
```

circos.points

Add points to a plotting region

Description

Add points to a plotting region

Usage

```
circos.points(x, y, sector.index = get.cell.meta.data("sector.index"),
    track.index = get.cell.meta.data("track.index"),
    pch = par("pch"), col = par("col"), cex = par("cex"), bg = par("bg"))
```

Arguments

X	Data points on x-axis, measured in "current" data coordinate
У	Data points on y-axis, measured in "current" data coordinate
sector.index	Index for the sector
track.index	Index for the track

pch Point type
col Point color
cex Point size

bg backgrond of points

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Details

This function can only add points in one specified cell. Pretending a low-level plotting function, it can only be applied in plotting region which has been created.

You can think the function similar as the normal points function, just adding points in the circular plotting region. The position of cell is identified by sector.index and track.index, if they are not specified, they are in 'current' sector and 'current' track.

Data points out of the plotting region will also be added, but with warning messages.

Other graphics parameters which are available in the function are pch, col and cex which have same meaning as those in the par.

It is recommended to use circos.points inside panel.fun in circos.trackPlotRegion so that it draws points directly on "curent" cell.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

Examples

```
circos.initialize(letters[1:8], xlim = c(0, 1))
circos.track(ylim = c(0, 1), panel.fun = function(x, y) {
    circos.points(runif(10), runif(10))
})
circos.points(runif(10), runif(10), sector.index = "c", pch = 16, col = "red")
circos.clear()
```

circos.polygon

Draw polygon

Description

Draw polygon

Usage

Arguments

```
x Data points on x-axis
y Data points on y-axis
sector.index Index for the sector
track.index Index for the track
pass to polygon
```

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Details

```
similar as polygon.
```

Note: start point should overlap with the end point,

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

Examples

```
set.seed(123)
factors = letters[1:4]
circos.initialize(factors, x = c(0, 1))
circos.trackPlotRegion(ylim = c(-3, 3), track.height = 0.4, panel.fun = function(x, y) {
    x1 = runif(20)
   y1 = x1 + rnorm(20)
   or = order(x1)
   x1 = x1[or]
    y1 = y1[or]
    loess.fit = loess(y1 \sim x1)
   loess.predict = predict(loess.fit, x1, se = TRUE)
   d1 = c(x1, rev(x1))
   d2 = c(loess.predict$fit + loess.predict$se.fit,
        rev(loess.predict$fit - loess.predict$se.fit))
   circos.polygon(d1, d2, col = "#CCCCCC", border = NA)
   circos.points(x1, y1, cex = 0.5)
    circos.lines(x1, loess.predict$fit)
})
circos.clear()
```

circos.raster

Add raster images

Description

Add raster images

Usage

```
circos.raster(image, x, y, width, height,
    facing = c("inside", "outside", "reverse.clockwise", "clockwise",
    "downward", "bending.inside", "bending.outside"),
    niceFacing = FALSE, sector.index = get.cell.meta.data("sector.index"),
    track.index = get.cell.meta.data("track.index"),
    scaling = 1)
```

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Arguments

image	a raster object, or an object that can be converted by as.raster
x	position of the center of the raster image, measued in the data coordinate in the cell
У	position of the center of the raster image, measued in the data coordinate in the cell
width	width of the raster image. When facing is one of "inside", "outside", "clockwise" and "reverse.clockwise", the image should have absolute size where the value of width should be specified like 20mm, 1cm or 0.5 inche. When facing is one of bending.inside and bending.outside, the value of width is measured in the data coordinate in the cell.
height	height of the raster image. Same format as width. If the value of height is omit, default height is calculated by taking the aspect ratio of the original image. But when facing is one of bending.inside and bending.outside, height is mandatory to set.
facing	facing of the raster image
niceFacing	facing of text. Please refer to vignette for different settings
sector.index	index for the sector
track.index	index for the track
scaling	scaling factor to resize the raster image.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

```
image = system.file("extdata", "Rlogo.png", package = "circlize")
image = as.raster(readPNG(image))
circos.initialize(letters[1:8], xlim = c(0, 1))
circos.track(ylim = c(0, 1), panel.fun = function(x, y) {
   circos.raster(image, CELL_META$xcenter, CELL_META$ycenter, width = "2cm",
        facing = "inside", niceFacing = TRUE)
})
circos.clear()
## Not run:
# NOTE: following takes quite a long time to run
load(system.file("extdata", "doodle.RData", package = "circlize"))
circos.par("cell.padding" = c(0, 0, 0, 0))
circos.initialize(letters[1:16], x \lim = c(0, 1))
circos.track(ylim = c(0, 1), panel.fun = function(x, y) {
    img = img_list[[CELL_META$sector.numeric.index]]
    circos.raster(img, CELL_META$xcenter, CELL_META$ycenter, width = 1,
       height = 1, facing = "bending.inside")
}, track.height = 0.25, bg.border = NA)
```

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```
circos.track(ylim = c(0, 1), panel.fun = function(x, y) {
   img = img_list[[CELL_META$sector.numeric.index + 16]]
   circos.raster(img, CELL_META$xcenter, CELL_META$ycenter, width = 1,
        height = 1, facing = "bending.inside")
}, track.height = 0.25, bg.border = NA)
circos.clear()
## End(Not run)
```

circos.rect

Draw rectangle-like grid

Description

Draw rectangle-like grid

Usage

```
circos.rect(xleft, ybottom, xright, ytop,
    sector.index = get.cell.meta.data("sector.index"),
    track.index = get.cell.meta.data("track.index"), ...)
```

Arguments

```
xleft x for the left bottom points
ybottom y for the left bottom points
xright x for the right top points
ytop y for the right top points
sector.index Index for the sector
track.index Index for the track
... pass to polygon
```

Details

The name for this function is circos.rect because if you imagine the plotting region as Cartesian coordinate, then it is rectangle. in the polar coordinate, the up and bottom edge become two arcs.

This function can be vectorized.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

```
# There is no example NULL
```

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ci	rcas	.segments
-	1 603	. 30 8 1110 11 03

Draw segments through pairwise of points

Description

Draw segments through pairwise of points

Usage

```
circos.segments(x0, y0, x1, y1, sector.index = get.cell.meta.data("sector.index"),
    track.index = get.cell.meta.data("track.index"), straight = FALSE,
    col = par("col"), lwd = par("lwd"), lty = par("lty"), ...)
```

Arguments

x0	x coordinates for starting points
y0	y coordinates for ending points
x1	x coordinates for starting points
y1	y coordinates for ending points

sector.index Index for the sector track.index Index for the track

straight whether the segment is a straight line

col color of the segments

lwd line width of the segments

lty line type of the segments

... pass to lines

Examples

```
# There is no example NULL
```

circos.text

Draw text in a cell

Description

Draw text in a cell

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Usage

```
circos.text(x, y, labels, sector.index = get.cell.meta.data("sector.index"),
    track.index = get.cell.meta.data("track.index"), direction = NULL,
    facing = c("inside", "outside", "reverse.clockwise", "clockwise",
    "downward", "bending", "bending.inside", "bending.outside"), niceFacing = FALSE,
    adj = par("adj"), cex = 1, col = par("col"), font = par("font"), ...)
```

Arguments

X	Data points on x-axis
у	Data points on y-axis
labels	Labels for each points
sector.index	Index for the sector
track.index	Index for the track
direction	deprecated, use facing instead.
facing	Facing of text. Please refer to vignette for different settings
niceFacing	Should the facing of text be adjusted to fit human eyes?
adj	offset for text. By default the text position adjustment is either horizontal or vertical in the canvas coordinate system. The "circular horizontal" offset can be set as a value in degree unit and the value should be wrapped by degree.
	Pass to text
cex	Font size
col	Font color
font	Font style

Details

The function is similar to text. All you need to note is the facing settings.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

See Also

```
http://jokergoo.github.io/circlize_book/book/graphics.html#text
```

```
factors = letters[1:4]
circos.par(points.overflow.warning = FALSE)
circos.initialize(factors = factors, xlim = c(0, 10))
circos.trackPlotRegion(factors = factors, ylim = c(0, 10),
    track.height = 0.5, panel.fun = function(x, y) {
    circos.text(3, 1, "inside", facing = "inside", cex = 0.8)
    circos.text(7, 1, "outside", facing = "outside", cex = 0.8)
```

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```
circos.text(0, 5, "reverse.clockwise", facing = "reverse.clockwise",
    adj = c(0.5, 0), cex = 0.8)
circos.text(10, 5, "clockwise", facing = "clockwise", adj = c(0.5, 0),
    cex = 0.8)
circos.text(5, 5, "downward", facing = "downward", cex = 0.8)
circos.text(3, 9, "====bending.inside====", facing = "bending.inside",
    cex = 0.8)
circos.text(7, 9, "====bending.outside====", facing = "bending.outside",
    cex = 0.8)
})
circos.clear()
```

circos.track

Create plotting regions for a whole track

Description

Create plotting regions for a whole track

Usage

```
circos.track(...)
```

Arguments

... pass to circos.trackPlotRegion

Details

Shortcut function of circos.trackPlotRegion.

```
\label{eq:continuous_problem} \mbox{\ensuremath{\mbox{\sc H}}} \mbox{\ensuremath{\mbox{\sc There}}} \mbox{\ensuremath{\mbox{\sc NULL}}} \mbox{\ensuremath{\mbox{\sc NULL}}}}
```

circos.trackHist 67

circos.trackHist	Draw histogram in cells among a whole track	
CIT COS. CT aCKITISC	Draw histogram in cetts among a whole track	

Description

Draw histogram in cells among a whole track

Usage

```
circos.trackHist(factors, x, track.height = circos.par("track.height"),
   track.index = NULL, force.ylim = TRUE, col = ifelse(draw.density, "black", NA),
   border = "black", lty = par("lty"), lwd = par("lwd"),
   bg.col = NA, bg.border = "black", bg.lty = par("lty"), bg.lwd = par("lwd"),
   breaks = "Sturges", include.lowest = TRUE, right = TRUE, draw.density = FALSE,
   bin.size = NULL, area = FALSE)
```

Arguments

bin.size

g	guments	
	factors	Factors which represent the categories of data
	x	Data on the x-axis
	track.index	Index for the track which is going to be updated. Setting it to NULL means creating the plotting regions in the next newest track.
	track.height	Height of the track. It is the percentage to the radius of the unit circle. If to update a track, this argument is disabled.
	force.ylim	Whether to force all cells in the track to share the same ylim. Btw, ylim is calculated automatically.
	col	Filled color for histogram
	border	Border color for histogram
	lty	Line style for histogram
	lwd	Line width for histogram
	bg.col	Background color for the plotting regions
	bg.border	Color for the border of the plotting regions
	bg.lty	Line style for the border of the plotting regions
	bg.lwd	Line width for the border of the plotting regions
	breaks	see hist
	$\verb"include.lowest"$	see hist
	right	see hist
	draw.density	whether draw density lines instead of histogram bars.
	area	whether to fill the area below the density lines. If it is set to TRUE, col controls

the filled color in the area and border controls color of the line.

size of the bins of the histogram

68 circos.trackLines

Details

It draw histogram in cells among a whole track. It is also an example to show how to add self-defined high-level graphics by this package.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

Examples

```
## Not run:
x = rnorm(1600)
factors = sample(letters[1:16], 1600, replace = TRUE)
circos.initialize(factors = factors, x = x)
circos.trackHist(factors = factors, x = x, col = "#999999",
   border = "#999999")
circos.trackHist(factors = factors, x = x, bin.size = 0.1,
   col = "#999999", border = "#999999")
circos.trackHist(factors = factors, x = x, draw.density = TRUE,
   col = "#999999", border = "#999999")
circos.clear()
## End(Not run)
```

circos.trackLines

Add lines to the plotting regions in a same track

Description

Add lines to the plotting regions in a same track

Usage

```
circos.trackLines(factors, x, y, track.index = get.cell.meta.data("track.index"),
  col = par("col"), lwd = par("lwd"), lty = par("lty"), type = "l", straight = FALSE,
  area = FALSE, area.baseline = NULL, border = "black", baseline = "bottom",
  pt.col = par("col"), cex = par("cex"), pch = par("pch"))
```

Arguments

factors	A factor or a character vector which represents the categories of data
x	Data points on x-axis
у	Data points on y-axis
track.index	Index for the track
col	Line color
lwd	line width

circos.trackPlotRegion 69

lty	line style
type	line type, similar as type argument in lines, but only in c("l", "o", "h", "s")
straight	whether draw straight lines between points
area	whether to fill the area below the lines. If it is set to TRUE, col controls the filled color in the area and border controls the color of the line.
area.baseline	deprecated, use baseline instead.
baseline	the base line to draw area, pass to circos.lines.
border	color for border of the area
pt.col	if type is "o", points color
cex	if type is "o", points size
pch	if type is "o", points type

Details

The function adds lines in multiple cells by first splitting data into several parts in which each part corresponds to one factor (sector index) and then add lines in cells by calling circos.lines.

This function can be replaced by a for loop containing circos.lines.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

Examples

```
\label{eq:continuous_problem} \mbox{\ensuremath{\mbox{\sc H}}} \mbox{\ensuremath{\mbox{\mbox{\sc H}}}} \mbox{\ensuremath{\mbox{\sc H}}} \mbox{\ensuremath{\mbox{\sc
```

```
circos.trackPlotRegion
```

Create plotting regions for a whole track

Description

Create plotting regions for a whole track

Usage

```
circos.trackPlotRegion(factors = NULL, x = NULL, y = NULL, ylim = NULL,
    force.ylim = TRUE, track.index = NULL,
    track.height = circos.par("track.height"),
    track.margin = circos.par("track.margin"),
    cell.padding = circos.par("cell.padding"),
    bg.col = NA, bg.border = "black", bg.lty = par("lty"), bg.lwd = par("lwd"),
    panel.fun = function(x, y) {NULL})
```

Arguments

factors	A factor or a character vector which represents categories of data, if it is NULL, then it uses all sector index.
X	Data on x-axis. It is only used if panel. fun is set.
у	Data on y-axis
ylim	Range of data on y-axis
force.ylim	Whether to force all cells in the track to share the same ylim. Normally, all cells on a same track should have same ylim.
track.index	Index for the track which is going to be created/updated. If the specified track has already been created, this function just updated corresponding track with new plot. If the specified track is NULL or has not been created, this function just creates it. Note the value for this argument should not exceed maximum track index plus 1.
track.height	Height of the track. It is the percentage to the radius of the unit circles. The value can be set by uh to an absolute unit. If updating a track (with proper track.index value), this argument is ignored.
track.margin	only affect current track
cell.padding	only affect current track
bg.col	Background color for the plotting regions. It can be vector which has the same length of sectors.
bg.border	Color for the border of the plotting regions. It can be vector which has the same length of sectors.
bg.lty	Line style for the border of the plotting regions. It can be vector which has the same length of sectors.
bg.lwd	Line width for the border of the plotting regions. It can be vector which has the same length of sectors.
panel.fun	Panel function to add graphics in each cell, see "details" section and vignette for explanation.

Details

This function tends to be a high-level plotting function, which means, you must first call this function to create plotting regions, then those low-level graphic function such as circos.points, circos.lines can be applied.

Currently, all the cells that are created in a same track sharing same height, which means, there is no cell has larger height than others.

Since ranges for values on x-axis has already been defined by circos.initialize, only ranges for values on y-axis should be specified in this function. There are two ways to identify the ranges for values on y-axes either by y or ylim. If y is set, it must has the same length as factors and the ylim for each cell is calculated from y values. Also, the ylim can be specified from ylim which can be a two-element vector or a matrix which has two columns and the number of rows is the same as the length of the levels of the factors.

If there is no enough space for the new track or the new track overlaps with other tracks, there will be an error.

circos.trackPoints 71

If factors does not cover all sectors, the cells in remaining unselected sectors would also be created but without drawing anything. The ylim for these cells are the same as that in the last created cell.

The function can also update a already-created track if the index for the track is specified. If updating an existed track, those parameters related to the position (such as track height and track margin) of the plotting region can not be changed.

Panel

panel.fun provides a convenient way to add graphics in each cell when initializing the tracks. The self-defined function needs two arguments: x and y which correspond to the data points in the current cell. When factors, x, and y are set in circos.trackPlotRegion, a subset of x and y are split by factors and are sent to panel.fun in the "current" cell. circos.trackPlotRegion creates plotting regions one by one on the track and panel.fun adds graphics in the 'current' cell after the plotting region for a certain cell has been created.

See vignette for examples of how to use this feature.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

See Also

http://jokergoo.github.io/circlize_book/book/circular-layout.html

Examples

circos.trackPoints

Add points to the plotting regions in a same track

Description

Add points to the plotting regions in a same track

Usage

```
circos.trackPoints(factors = NULL, x, y, track.index = get.cell.meta.data("track.index"),
    pch = par("pch"), col = par("col"), cex = par("cex"), bg = par("bg"))
```

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Arguments

factors	A factor or a character vector which represents the categories of data
х	Data points on x-axis
У	Data points on y-axis
track.index	Index for the track
pch	Point type
col	Point color
cex	Point size
bg	backgrond color

Details

The function adds points in multiple cells by first splitting data into several parts in which each part corresponds to one factor (sector index) and then adding points in each cell by calling circos.points.

Length of pch, col and cex can be one, length of levels of the factors or length of factors.

This function can be replaced by a for loop containing circos.points.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

Examples

circos.trackText

Draw text in cells among the whole track

Description

Draw text in cells among the whole track

Usage

```
circos.trackText(factors, x, y, labels, track.index = get.cell.meta.data("track.index"),
    direction = NULL, facing = c("inside", "outside", "reverse.clockwise", "clockwise",
    "downward", "bending", "bending.inside", "bending.outside"), niceFacing = FALSE,
    adj = par("adj"), cex = 1, col = par("col"), font = par("font"))
```

circos.update 73

Arguments

factors A factor or a character vector which represents the categories of data

x Data points on x-axis y Data points on y-axis

labels Labels

track.index Index for the track

direction deprecated, use facing instead.

facing Facing of text

niceFacing Should the facing of text be adjusted to fit human eyes?

adj Adjustment for text

cex Font size
col Font color
font Font style

Details

The function adds texts in multiple cells by first splitting data into several parts in which each part corresponds to one factor (sector index) and then add texts in cells by calling circos.text.

This function can be replaced by a for loop containing circos.text.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

Examples

```
# There is no example NULL
```

circos.update

Create plotting regions for a whole track

Description

Create plotting regions for a whole track

Usage

```
circos.update(...)
```

Arguments

... pass to circos.updatePlotRegion

Details

shortcut function of circos.updatePlotRegion.

Examples

```
# There is no example NULL
```

circos.updatePlotRegion

Update the plotting region in an existed cell

Description

Update the plotting region in an existed cell

Usage

Arguments

sector.index

track.index Index for the track
bg.col Background color for the plotting region
bg.border Color for the border of the plotting region
bg.lty Line style for the border of the plotting region
bg.lwd Line width for the border of the plotting region

Index for the sector

Details

You can update an existed cell by this function by erasing all the graphics. But the xlim and ylim inside the cell still remain unchanged.

Note if you use circos.track to update an already created track, you can re-define ylim in these cells.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

circos.xaxis 75

Examples

circos.xaxis

Draw x-axis

Description

Draw x-axis

Usage

```
circos.xaxis(...)
```

Arguments

... all pass to circos.axis

Examples

```
\hbox{\tt\# There is no example}\\ \hbox{\tt NULL}
```

circos.yaxis

Draw y-axis

Description

Draw y-axis

```
circos.yaxis(side = c("left", "right"), at = NULL, labels = TRUE, tick = TRUE,
    sector.index = get.cell.meta.data("sector.index"),
    track.index = get.cell.meta.data("track.index"),
    labels.font = par("font"), labels.cex = par("cex"),
    labels.niceFacing = TRUE,
    tick.length = convert_x(1, "mm", sector.index, track.index),
    lwd = par("lwd"), col = par("col"), labels.col = par("col"))
```

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Arguments

side add the y-axis on the left or right of the cell If it is numeric vector, it identifies the positions of the ticks. It can exceed ylim at value and the exceeding part would be trimmed automatically. labels labels of the ticks. The exceeding part would be trimmed automatically. The value can also be logical (either an atomic value or a vector) which represents which labels to show. tick Whether to draw ticks. sector.index Index for the sector track.index Index for the track labels.font font style for the axis labels font size for the axis labels labels.cex labels.niceFacing Should facing of axis labels be human-easy length of the tick tick.length

lwd line width for ticks col color for the axes labels.col

Details

Note, you need to set the gap between sectors manually by circos.par to make sure there is enough space for y-axis.

```
op = par(no.readonly = TRUE)

factors = letters[1:8]
circos.par(points.overflow.warning = FALSE)
circos.par(gap.degree = 8)
circos.initialize(factors = factors, xlim = c(0, 10))
circos.trackPlotRegion(factors = factors, ylim = c(0, 10), track.height = 0.5)
par(cex = 0.8)
for(a in letters[2:4]) {
    circos.yaxis(side = "left", sector.index = a)
}
for(a in letters[5:7]) {
    circos.yaxis(side = "right", sector.index = a)
}
circos.clear()
par(op)
```

col2value 77

Transform back from colors to values

Description

Transform back from colors to values

Usage

```
col2value(r, g, b, col_fun)
```

Arguments

r	red channel in sRGB color space, value should be between 0 and 1. The r, g and b argumentc can be wrapped into one variable which is either a three-column matrix or a vector of colors.
g	green channel in sRGB color space, value should be between 0 and 1.
b	blue channel in sRGB color space, value should be between 0 and 1.
col_fun	the color mapping function generated by colorRamp2.

Details

colorRamp2 transforms values to colors and this function does the reversed job. Note for some color spaces, it cannot transform back to the original value perfectly.

Value

A vector of original numeric values.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

```
x = seq(0, 1, length = 11)
col_fun = colorRamp2(c(0, 0.5, 1), c("blue", "white", "red"))
col = col_fun(x)
col2value(col, col_fun = col_fun)
col2value("red", col_fun = col_fun)

col_fun = colorRamp2(c(0, 0.5, 1), c("blue", "white", "red"), space = "sRGB")
col = col_fun(x)
col2value(col, col_fun = col_fun)
```

78 colorRamp2

-		
COL	orRamp2	

Color interpolation

Description

Color interpolation

Usage

```
colorRamp2(breaks, colors, transparency = 0, space = "LAB")
```

Arguments

breaks A vector indicating numeric breaks

colors A vector of colors which correspond to values in breaks

transparency A single value in [0,1]. O refers to no transparency and 1 refers to full trans-

parency

space color space in which colors are interpolated. Value should be one of "RGB",

"HSV", "HLS", "LAB", "XYZ", "sRGB", "LUV", see color-class for detail.

Details

Colors are linearly interpolated according to break values and corresponding colors through CIE Lab color space (LAB) by default. Values exceeding breaks will be assigned with corresponding maximum or minimum colors.

Value

It returns a function which accepts a vector of numeric values and returns interpolated colors.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

See Also

col2value converts back to the original values by providing the color mapping function generated by colorRamp2.

```
col_fun = colorRamp2(c(-1, 0, 1), c("green", "white", "red"))
col_fun(c(-2, -1, -0.5, 0, 0.5, 1, 2))
```

convert_height 79

convert_height

Convert units

Description

Convert units

Usage

```
convert_height(...)
```

Arguments

```
... pass to convert_length
```

Details

This function is same as convert_length. The reason for naming this function is convert_length is mostely used for defining the height of tracks and track margins.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

```
# see example in `convert_length` page
NULL
```

convert_length

Convert units

Description

Convert units

Usage

```
convert_length(x, unit = c("mm", "cm", "inches"))
```

Arguments

```
x a numeric vector
```

unit supported units, only "mm", "cm", "inches".

80 convert_x

Details

This function coverts mm/cm/inches units to units measured in the canvas coordinate, e.g. how much is it in the canvas coordinate for 1 mm/cm/inches.

Since in the circular plot, the aspect ratio is always 1, it does not matter this conversion is applied on x direction or y direction.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

See Also

convert_x and convert_y convert absolute units into a data coordinate in a specified cell.

Examples

```
fa = letters[1:10]
circos.par(cell.padding = c(0, 0, 0, 0), track.margin = c(0, 0))
circos.initialize(fa, xlim = cbind(rep(0, 10), runif(10, 0.5, 1.5)))
circos.track(ylim = c(0, 1), track.height = convert_length(5, "mm"))
circos.par(track.margin = c(0, convert_length(2, "mm")))
circos.track(ylim = c(0, 1), track.height = convert_length(1, "cm"))
circos.par(track.margin = c(0, convert_length(5, "mm")))
circos.track(ylim = c(0, 1), track.height = convert_length(1, "inches"))
circos.clear()
```

convert_x

Convert unit on x direction in data coordinate

Description

Convert unit on x direction in data coordinate

Usage

```
convert_x(x, unit = c("mm", "cm", "inches"),
    sector.index = get.cell.meta.data("sector.index"),
    track.index = get.cell.meta.data("track.index"),
    h = get.cell.meta.data("ycenter", sector.index = sector.index,
    track.index = track.index))
```

Arguments

```
x a numeric vector
unit supported units, only "mm", "cm", "inches"
sector.index index for the sector where the conversion is applied
track.index index for the track where the conversion is applied
```

convert_y 81

h

since the width of the cell is not identical from the top to the bottom in the cell, the position on y direction needs to be specified. By default it is at the middle point on y-axis

Value

A vector of numeric values which are measured in the specified data coordinate

Author(s)

Zuguang Gu <z.gu@dkfz.de>

See Also

convert_y converts on y direction.

Examples

```
fa = letters[1:10]
circos.par(cell.padding = c(0, 0, 0, 0), track.margin = c(0, 0))
circos.initialize(fa, xlim = cbind(rep(0, 10), runif(10, 0.5, 1.5)))
circos.track(ylim = c(0, 1), track.height = convert_height(5, "mm"),
    panel.fun = function(x, y) {
        circos.lines(c(0, 0 + convert_x(5, "mm")), c(0.5, 0.5), col = "blue")
circos.par(track.margin = c(0, convert_height(2, "mm")))
circos.track(ylim = c(0, 1), track.height = convert_height(1, "cm"),
    panel.fun = function(x, y) {
        xcenter = get.cell.meta.data("xcenter")
        circos.lines(c(xcenter, xcenter), c(0, convert_y(1, "cm")), col = "red")
    })
circos.par(track.margin = c(0, convert_height(5, "mm")))
circos.track(ylim = c(0, 1), track.height = convert_height(1, "inches"),
    panel.fun = function(x, y) {
        line_length_on_x = convert_x(1*sqrt(2)/2, "cm")
        line_length_on_y = convert_y(1*sqrt(2)/2, "cm")
        circos.lines(c(0, line_length_on_x), c(0, line_length_on_y), col = "orange")
    })
circos.clear()
```

convert_y

Convert unit on y direction in data coordinate

Description

Convert unit on y direction in data coordinate

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Usage

```
convert_y(x, unit = c("mm", "cm", "inches"),
    sector.index = get.current.sector.index(),
    track.index = get.current.track.index())
```

Arguments

x a numeric vector

unit supported units, only "mm", "cm", "inches"

sector.index index for the sector where the conversion is applied track.index index for the track where the conversion is applied

Value

A vector of numeric values which are measured in the specified data coordinate

Author(s)

Zuguang Gu <z.gu@dkfz.de>

See Also

convert_x converts on x direction.

Examples

```
# see example on `convert_x` page
NULL
```

cytoband.col

Assign colors to cytogenetic band (hg19) according to the Giemsa stain results

Description

Assign colors to cytogenetic band (hg19) according to the Giemsa stain results

Usage

```
cytoband.col(x)
```

Arguments

Х

A vector containing the Giemsa stain results

degree 83

Details

The color theme is from http://circos.ca/tutorials/course/slides/session-2.pdf, page 42.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

Examples

```
## Not run:
cytoband = read.cytoband()
cytoband.col(cytoband$df[[5]])
## End(Not run)
```

degree

Mark the value as a degree value

Description

Mark the value as a degree value

Usage

```
degree(x)
```

Arguments

Χ

degree value

Value

```
a degree object
```

```
\label{eq:continuous_problem} \mbox{\ensuremath{\mbox{\sc H}}} \mbox{\ensuremath{\mbox{\sc There}}} \mbox{\ensuremath{\mbox{\sc NULL}}} \mbox{\ensuremath{\mbox{\sc NULL}}}}
```

84 draw.sector

draw.sector	Draw sectors or rings in a circle	
araw.scctor	Draw sectors of rings in a circle	

Description

Draw sectors or rings in a circle

Usage

```
draw.sector(start.degree = 0, end.degree = 360, rou1 = 1, rou2 = NULL,
    center = c(0, 0), clock.wise = TRUE, col = NA, border = "black", lwd = par("lwd"),
    lty = par("lty"))
```

Arguments

start.degree start degree for the sector end.degree end degree for the sector

rou1 Radius for one of the arc in the sector rou2 Radius for the other arc in the sector

center Center of the circle

clock.wise The direction from start.degree to end.degree

col Filled color
border Border color
lwd Line width
lty Line style

Details

If the interval between start and end (larger or equal to 360 or smaller or equal to -360) it would draw a full circle or ring. If rou2 is set, it would draw part of a ring.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

```
plot(c(-1, 1), c(-1, 1), type = "n", axes = FALSE, ann = FALSE, asp = 1) draw.sector(20, 0) draw.sector(30, 60, rou1 = 0.8, rou2 = 0.5, clock.wise = FALSE, col = "#FF000080") draw.sector(350, 1000, col = "#00FF0080", border = NA) draw.sector(0, 180, rou1 = 0.25, center = c(-0.5, 0.5), border = 2, lwd = 2, lty = 2) draw.sector(0, 360, rou1 = 0.7, rou2 = 0.6, col = "#0000FF80") factors = letters[1:8]
```

fontsize 85

```
circos.initialize(factors, x \lim = c(0, 1))
for(i in 1:3) {
   circos.trackPlotRegion(ylim = c(0, 1))
circos.info(plot = TRUE)
draw.sector(get.cell.meta.data("cell.start.degree", sector.index = "a"),
            get.cell.meta.data("cell.end.degree", sector.index = "a"),
            rou1 = 1, col = "#FF000040")
draw.sector(0, 360,
    rou1 = get.cell.meta.data("cell.top.radius", track.index = 1),
    rou2 = get.cell.meta.data("cell.bottom.radius", track.index = 1),
    col = "#00FF0040")
draw.sector(get.cell.meta.data("cell.start.degree", sector.index = "e"),
            get.cell.meta.data("cell.end.degree", sector.index = "f"),
            get.cell.meta.data("cell.top.radius", track.index = 2),
            get.cell.meta.data("cell.bottom.radius", track.index = 3),
            col = "#0000FF40")
pos = circlize(c(0.2, 0.8), c(0.2, 0.8), sector.index = "h", track.index = 2)
draw.sector(pos[1, "theta"], pos[2, "theta"], pos[1, "rou"], pos[2, "rou"],
    clock.wise = TRUE, col = "#00FFFF40")
circos.clear()
```

fontsize

Convert fontsize to cex

Description

Convert fontsize to cex

Usage

fontsize(x)

Arguments

х

value for fontsize

Details

It just mark the value as a fontsize setting for circos.text

Author(s)

Zuguang Gu <z.gu@dkfz.de>

86 generateRandomBed

Examples

```
# There is no example NULL
```

generateRandomBed

Generate random genomic data

Description

Generate random genomic data

Usage

```
generateRandomBed(nr = 10000, nc = 1, fun = function(k) rnorm(k, 0, 0.5), species = NULL)
```

Arguments

nr Number of rows

nc Number of numeric columns / value columns

fun Function for generating random values

species species, pass to read.cytoband

Details

The function will uniformly sample positions from the genome. Chromosome names start with "chr" and positions are sorted. The final number of rows may not be exactly as same as nr.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

```
bed = generateRandomBed()
bed = generateRandomBed(nr = 200, nc = 4)
bed = generateRandomBed(fun = function(k) runif(k))
```

genomicDensity 87

genomicDe	nsity	Calculate genomic region density

Description

Calculate genomic region density

Usage

```
genomicDensity(region, window.size = 1e7, n.window = NULL, overlap = TRUE, chr.len = NULL)
```

Arguments

region	Genomic positions. It can be a data frame with two columns which are start positions and end positions on a single chromosome. It can also be a bed-format data frame which contains the chromosome column.
window.size	Window size to calculate genomic density
n.window	number of windows, if it is specified, window. size is ignored
overlap	Whether two neighbouring windows have half overlap
chr.len	the chromosome length. The value should be named vector

Details

It calculate the percent of each genomic windows that is covered by the input regions.

Value

If the input is a two-column data frame, the function returns a data frame with three columns: start position, end position and percent of overlapping. And if the input is a bed-format data frame, there will be an additionally chromosome name column.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

```
bed = generateRandomBed()
bed = subset(bed, chr == "chr1")
head(genomicDensity(bed))
```

88 get.all.track.index

```
get.all.sector.index Get index for all sectors
```

Description

Get index for all sectors

Usage

```
get.all.sector.index()
```

Details

It simply returns a vector of all sector index.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

Examples

```
## Not run:
library(circlize)
factors = letters[1:4]
circos.initialize(factors, xlim = c(0, 1))
circos.trackPlotRegion(ylim = c(0, 1))
get.all.sector.index()
circos.clear()
## End(Not run)
```

```
get.all.track.index Get index for all tracks
```

Description

Get index for all tracks

Usage

```
get.all.track.index()
```

Details

It simply returns a vector of all track index.

get.cell.meta.data 89

Examples

```
## Not run:
library(circlize)
factors = letters[1:4]
circos.initialize(factors, xlim = c(0, 1))
circos.trackPlotRegion(ylim = c(0, 1))
circos.trackPlotRegion(ylim = c(0, 1))
circos.trackPlotRegion(ylim = c(0, 1))
get.all.track.index()
circos.clear()
## End(Not run)
```

get.cell.meta.data

Get the meta data of a cell

Description

Get the meta data of a cell

Usage

Arguments

name Only support one name at a time, see "details" section

sector.index Index of the sector track.index Index of the track

Details

The following meta information for a cell can be obtained:

```
sector.index The name (index) for the sector sector.numeric.index Numeric index for the sector track.index Numeric index for the track xlim Minimal and maximal values on the x-axis ylim Minimal and maximal values on the y-axis xrange Range of xlim. It equals to xlim[2] -xlim[1] yrange Range of ylim xcenter Center of x-axis. It equals to (xlim[2] + xlim[1])/2 ycenter Center of y-axis
```

```
cell.xlim Minimal and maximal values on the x-axis extended by cell paddings cell.ylim Minimal and maximal values on the y-axis extended by cell paddings xplot Degrees for right and left borders of the cell.

yplot Radius for top and bottom borders of the cell.

cell.start.degree Same as xplot[1]

cell.end.degree Same as xplot[2]

cell.bottom.radius Same as yplot[1]

cell.top.radius Same as yplot[2]

track.margin Margin for the cell

cell.padding Padding for the cell
```

The function is useful when using panel. fun in circos. track to get detailed information of the current cell.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

See Also

```
CELL_META is a short version of get.cell.meta.data.
```

Examples

```
factors = letters[1:4]
circos.initialize(factors, xlim = c(0, 1))
circos.trackPlotRegion(ylim = c(0, 1), panel.fun = function(x, y) {
    print(get.cell.meta.data("xlim"))
})
print(get.cell.meta.data("xlim", sector.index = "a", track.index = 1))
circos.clear()
```

```
get.current.chromosome
```

Get current chromosome name

Description

Get current chromosome name

```
get.current.chromosome()
```

get.current.sector.index 91

Details

The function is same as get.current.sector.index and should only be put inside panel.fun when using circos.genomicTrackPlotRegion.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

Examples

```
## Not run:
library(circlize)
circos.initializeWithIdeogram()
circos.genomicTrackPlotRegion(ylim = c(0, 1), panel.fun = function(region, value, ...) {
    print(get.current.chromosome())
})
circos.clear()
## End(Not run)
```

```
get.current.sector.index
```

Get current sector index

Description

Get current sector index

Usage

```
get.current.sector.index()
```

Value

Simply returns the name of current sector

```
# There is no example
```

92 getI

```
get.current.track.index
```

Get current track index

Description

Get current track index

Usage

```
get.current.track.index()
```

Value

Simply returns the numeric index for the current track.

Examples

```
\# There is no example NULL
```

getI

Which data that panel. fun is using

Description

Which data that panel. fun is using

Usage

```
getI(...)
```

Arguments

... Invisible arguments that users do not need to care

Details

 $The function should only be put inside panel. fun when using {\tt circos.genomicTrackPlotRegion}.$

If stack is set to TRUE in circos.genomicTrackPlotRegion, the returned value indicates which stack the function will be applied to.

If data is a list of data frames, the value indicates which data frame is being used. Please see the vignette to get a more clear explanation.

highlight.chromosome 93

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

Examples

```
# There is no example NULL
```

highlight.chromosome Highlight chromosomes

Description

Highlight chromosomes

Usage

```
highlight.chromosome(...)
```

Arguments

```
... pass to highlight.sector
```

Details

This is only a shortcut function of highlight.sector.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

```
## Not run:

par(mar = c(1.5, 1.5, 1.5, 1.5))
# highlight
circos.par("track.height" = 0.1, cell.padding = c(0, 0, 0, 0))
circos.initializeWithIdeogram(plotType = c("axis", "labels"))

bed = generateRandomBed(nr = 100)
circos.genomicTrackPlotRegion(bed, panel.fun = function(region, value, ...) {
    circos.genomicPoints(region, value, pch = 16, cex = 0.5, ...)
})

bed = generateRandomBed(nr = 100)
circos.genomicTrackPlotRegion(bed, panel.fun = function(region, value, ...) {
    circos.genomicPoints(region, value, pch = 16, cex = 0.5, ...)
```

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```
})
bed = generateRandomBed(nr = 100)
circos.genomicTrackPlotRegion(bed, panel.fun = function(region, value, ...) {
    circos.genomicPoints(region, value, pch = 16, cex = 0.5, ...)
})
bed = generateRandomBed(nr = 100)
circos.genomicTrackPlotRegion(bed, panel.fun = function(region, value, ...) {
    circos.genomicPoints(region, value, pch = 16, cex = 0.5, ...)
})
\label{eq:highlight.chromosome} \mbox{\tt ("chr1", col = "\#FF000040", padding = c(0.05, 0.05, 0.15, 0.05))}
highlight.chromosome("chr3", col = NA, border = "red", lwd = 2,
    padding = c(0.05, 0.05, 0.15, 0.05))
highlight.chromosome("chr5", col = "#0000FF40", track.index = c(2, 4, 5))
highlight.chromosome("chr7", col = NA, border = "green", lwd = 2,
    track.index = c(2, 4, 5))
circos.clear()
## End(Not run)
```

highlight.sector

Highlight sectors and tracks

Description

Highlight sectors and tracks

Usage

```
highlight.sector(sector.index, track.index = get.all.track.index(),
    col = "#FF000040", border = NA, lwd = par("lwd"), lty = par("lty"),
    padding = c(0, 0, 0, 0), text = NULL, text.col = par("col"),
    text.vjust = 0.5, ...)
```

Arguments

sector.index A vector of sector index

track.index A vector of track index that you want to highlight

col Color for highlighting. Note the color should be semi-transparent.

border Border of the highlighted region

lwd Width of borderslty Style of borders

names.CELL_META 95

padding	Padding for the highlighted region. It should contain four values representing ratios of the width or height of the highlighted region
text	text added in the highlight region, only support plotting one string at a time
text.vjust	adjustment on 'vertical' (radical) direction. Besides to set it as numeric values, the value can also be a string contain absoute unit, e.g. "2.1mm", "-1 inche", but only "mm", "cm", "inches"/"inche" are allowed.
text.col	color for the text
	pass to circos.text

Details

You can use circos.info to find out index for all sectors and all tracks.

The function calls draw.sector.

Examples

```
factors = letters[1:8]
circos.initialize(factors, xlim = c(0, 1))
for(i in 1:4) {
      circos.trackPlotRegion(ylim = c(0, 1))
}
circos.info(plot = TRUE)

highlight.sector(c("a", "h"), track.index = 1)
highlight.sector("c", col = "#00FF0040")
highlight.sector("d", col = NA, border = "red", lwd = 2)
highlight.sector("e", col = "#0000FF40", track.index = c(2, 3))
highlight.sector(c("f", "g"), col = NA, border = "green",
      lwd = 2, track.index = c(2, 3))
highlight.sector(factors, col = "#FFFF0040", track.index = 4)
circos.clear()
```

names.CELL_META

Names of all meta data in the current cell

Description

Names of all meta data in the current cell

Usage

```
## S3 method for class 'CELL_META'
names(x)
```

Arguments

```
x use CELL_META.
```

96 posTransform.default

Details

The variable CELL_META can only be used to get meta data of the "current" cell. Basically you can simply replace e.g. get.cell.meta.data("sector.index") to CELL_META\$sector.index.

See Also

```
get.cell.meta.data
```

Examples

```
names(CELL_META)
```

posTransform.default Genomic position transformation function

Description

Genomic position transformation function

Usage

```
posTransform.default(region, ...)
```

Arguments

region Genomic positions at a single chromosome. It is a data frame with two columns

which are start position and end position.

... other arguments

Details

The default position transformation functions transforms position to be equally distributed along the chromosome. If users want to define their own transformation function, the requirement is that the returned value should be a data frame with two columns: transformed start position and transformed end position. The returned value should have same number of rows as the input one.

For details why need to use position transformation, please refer to circos.genomicPosTransformLines.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

```
## Not run:
par(mfrow = c(2, 1))
### rect matrix
circos.par(cell.padding = c(0, 0, 0, 0))
circos.initializeWithIdeogram()
bed = generateRandomBed(nr = 100, nc = 4)
circos.genomicPosTransformLines(bed, posTransform = posTransform.default,
    horizontalLine = "top", track.height = 0.1)
f = colorRamp2(breaks = c(-1, 0, 1), colors = c("green", "black", "red"))
circos.genomicTrackPlotRegion(bed, stack = TRUE, panel.fun = function(region, value, ...) {
    circos.genomicRect(region, value, col = f(value[[1]]),
        border = f(value[[1]]), posTransform = posTransform.default, ...)
}, bg.border = NA)
circos.clear()
circos.par(cell.padding = c(0, 0, 0, 0))
circos.initializeWithIdeogram(plotType = NULL)
bed = generateRandomBed(nr = 20, nc = 4)
circos.genomicTrackPlotRegion(bed, ylim = c(0, 1), panel.fun = function(region, value, ...) {
    circos.genomicText(region, value, y = 0, adj = c(1, 0.5), labels = "gene",
        facing = "reverse.clockwise", posTransform = posTransform.default)
}, bg.border = NA)
circos.genomicPosTransformLines(bed, posTransform = posTransform.default,
    horizontalLine = "bottom", direction = "outside", track.height = 0.1)
cytoband = read.cytoband()$df
circos.genomicTrackPlotRegion(cytoband, stack = TRUE, panel.fun = function(region, value, ...) {
    circos.genomicRect(region, value, col = cytoband.col(value$V5), border = NA, ...)
}, track.height = 0.05)
circos.clear()
## End(Not run)
```

Description

Genomic position transformation function specifically for text

Usage

```
posTransform.text(region, y, labels, cex = 1, font = par("font"),
    sector.index = get.cell.meta.data("sector.index"),
    track.index = get.cell.meta.data("track.index"), padding = 0,
    extend = 0, ...)
```

Arguments

region Genomic positions at a single chromosome. It is a data frame with two columns

which are start position and end position.

y positions of texts

labels text labels

cex text size

font text font style

sector.index sector index

track.index track index

padding padding of text

extend extend to allow labels to be put in an region which is wider than the current

chromosome. The value should be a proportion value and the length is either

one or two.

... other arguments

Details

This position transformation function is designed specifically for text. Under the transformation, texts will be as close as possible to the original positions.

```
circos.genomicTrackPlotRegion(bed, ylim = c(0, 1),
   panel.fun = function(region, value, ...) {
       circos.genomicText(region, value, y = 0, labels.column = 1,
           facing = "clockwise", adj = c(0, 0.5),
           posTransform = posTransform.text, cex = 0.8, niceFacing = F)
}, track.height = 0.1, bg.border = NA)
i_track = get.cell.meta.data("track.index")
circos.genomicPosTransformLines(bed,
   posTransform = function(region, value)
       posTransform.text(region, y = 0, labels = value[[1]],
           cex = 0.8, track.index = i_track),
   direction = "outside"
circos.genomicTrackPlotRegion(bed, ylim = c(0, 1),
   panel.fun = function(region, value, ...) {
       circos.points( (region[[1]] + region[[2]])/2, rep(0.5, nrow(region)), pch = 16)
}, track.height = 0.02, bg.border = NA)
circos.clear()
text(0, 0.05, "posTransform.text\ndirection = 'outside'", adj = c(0, 0))
circos.par("start.degree" = 90, canvas.xlim = c(0, 1), canvas.ylim = c(0, 1),
   gap.degree = 270, cell.padding = c(0, 0, 0, 0), track.margin = c(0.005, 0.005))
circos.initializeWithIdeogram(plotType = c("axis"), chromosome.index = "chr1")
circos.genomicTrackPlotRegion(bed, ylim = c(0, 1),
   panel.fun = function(region, value, ...) {
       circos.genomicText(region, value, y = 0, labels.column = 1,
           facing = "clockwise", adj = c(0, 0.5),
           posTransform = posTransform.default, cex = 0.8, niceFacing = F)
}, track.height = 0.1, bg.border = NA)
i_track = get.cell.meta.data("track.index")
circos.genomicPosTransformLines(bed, posTransform = posTransform.default,
   direction = "outside")
circos.genomicTrackPlotRegion(bed, ylim = c(0, 1),
   panel.fun = function(region, value, ...) {
       circos.points( (region[[1]] + region[[2]])/2, rep(0.5, nrow(region)), pch = 16)
}, track.height = 0.02, bg.border = NA)
circos.clear()
text(0, 0.05, "posTransform.default\ndirection = 'outside'", adj = c(0, 0))
circos.par("start.degree" = 90, canvas.xlim = c(0, 1), canvas.ylim = c(0, 1),
   gap.degree = 270, cell.padding = c(0, 0, 0, 0), track.margin = c(0.005, 0.005))
circos.initializeWithIdeogram(plotType = c("axis"), chromosome.index = "chr1")
circos.par(cell.padding = c(0, 0, 0, 0))
circos.genomicTrackPlotRegion(bed, ylim = c(0, 1),
```

```
panel.fun = function(region, value, ...) {
       circos.points( (region[[1]] + region[[2]])/2, rep(0.5, nrow(region)), pch = 16)
}, track.height = 0.02, bg.border = NA)
circos.genomicTrackPlotRegion(bed, ylim = c(0, 1), track.height = 0.1, bg.border = NA)
i_track = get.cell.meta.data("track.index")
circos.genomicTrackPlotRegion(bed, ylim = c(0, 1),
   panel.fun = function(region, value, ...) {
       circos.genomicText(region, value, y = 1, labels.column = 1,
           facing = "clockwise", adj = c(1, 0.5),
           posTransform = posTransform.text, cex = 0.8, niceFacing = F)
}, track.height = 0.1, bg.border = NA)
circos.genomicPosTransformLines(bed,
   posTransform = function(region, value)
       posTransform.text(region, y = 1, labels = value[[1]],
           cex = 0.8, track.index = i_track+1),
   direction = "inside", track.index = i_track
)
circos.clear()
text(0, 0.05, "posTransform.text\ndirection = 'inside'", adj = c(0, 0))
circos.par("start.degree" = 90, canvas.xlim = c(0, 1), canvas.ylim = c(0, 1),
   gap.degree = 270, cell.padding = c(0, 0, 0, 0), track.margin = c(0.005, 0.005))
circos.initializeWithIdeogram(plotType = c("axis"), chromosome.index = "chr1")
circos.par(cell.padding = c(0, 0, 0, 0))
circos.genomicTrackPlotRegion(bed, ylim = c(0, 1),
   panel.fun = function(region, value, ...) {
       circos.points( (region[[1]] + region[[2]])/2, rep(0.5, nrow(region)), pch = 16)
}, track.height = 0.02, bg.border = NA)
circos.genomicTrackPlotRegion(bed, ylim = c(0, 1), track.height = 0.1, bg.border = NA)
i_track = get.cell.meta.data("track.index")
circos.genomicTrackPlotRegion(bed, ylim = c(0, 1),
   panel.fun = function(region, value, ...) {
       circos.genomicText(region, value, y = 1, labels.column = 1, facing = "clockwise",
           adj = c(1, 0.5), posTransform = posTransform.text, cex = 0.8,
           niceFacing = F, padding = 0.2)
}, track.height = 0.1, bg.border = NA)
circos.genomicPosTransformLines(bed,
   posTransform = function(region, value)
       posTransform.text(region, y = 1, labels = value[[1]],
           cex = 0.8, track.index = i_track+1, padding = 0.2),
   direction = "inside", track.index = i_track
circos.clear()
text(0, 0.05, "posTransform.text\ndirection = 'inside'\npadding = 0.2", adj = c(0, 0))
```

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```
par(op)
## End(Not run)
```

print.CELL_META

Print CELL_META

Description

Print CELL_META

Usage

```
## S3 method for class 'CELL_META'
print(x, ...)
```

Arguments

x input

... additional parameters

Examples

```
\label{eq:continuous_problem} \mbox{\ensuremath{\mbox{\sc H}}} \mbox{\ensuremath{\mbox{\mbox{\sc H}}}} \mbox{\ensuremath{\mbox{\sc H}}} \mbox{\ensuremath{\mbox{\sc
```

rainfallTransform

Calculate inter-distance of genomic regions

Description

Calculate inter-distance of genomic regions

rand_color

Arguments

region Genomic positions. It can be a data frame with two columns which are start

positions and end positions on a single chromosome. It can also be a bed-format

data frame which contains the chromosome column.

mode How to calculate inter-distance. For a region, there is a distance to the prevous

region and also there is a distance to the next region. mode controls how to merge

these two distances into one value.

normalize_to_width

If it is TRUE, the value is the relative distance divided by the width of the region.

Value

If the input is a two-column data frame, the function returnes a data frame with three columns: start position, end position and distance. And if the input is a bed-format data frame, there will be the chromosome column added.

The row order of the returned data frame is as same as the input one.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

Examples

```
bed = generateRandomBed()
bed = subset(bed, chr == "chr1")
head(rainfallTransform(bed))
```

rand_color

Generate random colors

Description

Generate random colors

Usage

```
rand_color(n, hue = NULL, luminosity = "random", transparency = 0)
```

Arguments

n number of colors

hue the hue of the generated color. You can use following default color name: red,

orange, yellow, green, blue, purple, pink and monochrome. If the value is a hexidecimal color string such as #00FFFF, the function will extract its hue value

and use that to generate colors.

read.chromInfo

```
luminosity controls the luminosity of the generated color. The value should be a string containing bright, light, dark and random.

transparency transparency, numeric value between 0 and 1.
```

Details

```
The code is adapted from randomColor.js (https://github.com/davidmerfield/randomColor).
```

Value

a vector of colors

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

```
plot(NULL, xlim = c(1, 10), ylim = c(1, 8), axes = FALSE, ann = FALSE)
points(1:10, rep(1, 10), pch = 16, cex = 5,
   col = rand_color(10, luminosity = "random"))
points(1:10, rep(2, 10), pch = 16, cex = 5,
   col = rand_color(10, luminosity = "bright"))
points(1:10, rep(3, 10), pch = 16, cex = 5,
   col = rand_color(10, luminosity = "light"))
points(1:10, rep(4, 10), pch = 16, cex = 5,
   col = rand_color(10, luminosity = "dark"))
points(1:10, rep(5, 10), pch = 16, cex = 5,
   col = rand_color(10, hue = "red", luminosity = "bright"))
points(1:10, rep(6, 10), pch = 16, cex = 5,
    col = rand_color(10, hue = "green", luminosity = "bright"))
points(1:10, rep(7, 10), pch = 16, cex = 5,
    col = rand_color(10, hue = "blue", luminosity = "bright"))
points(1:10, rep(8, 10), pch = 16, cex = 5,
    col = rand_color(10, hue = "monochrome", luminosity = "bright"))
```

read.chromInfo

Read/parse chromInfo data from a data frame/file/UCSC database

Description

Read/parse chromInfo data from a data frame/file/UCSC database

```
read.chromInfo(chromInfo = system.file(package = "circlize",
    "extdata", "chromInfo.txt"), species = NULL,
    chromosome.index = usable_chromosomes(species),
    sort.chr = TRUE)
```

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Arguments

chromInfo Path of the chromInfo file or a data frame that already contains chromInfo data

species Abbreviations of species. e.g. hg19 for human, mm10 for mouse. If this value is

specified, the function will download chromInfo.txt.gz from UCSC website

automatically.

chromosome.index

subset of chromosomes, also used to reorder chromosomes.

sort.chr Whether chromosome names should be sorted (first sort by numbers then by

letters). If chromosome. index is set, this argument is enforced to FALSE

Details

The function read the chromInfo data, sort the chromosome names and calculate the length of each chromosome. By default, it is human hg19 chromInfo data.

You can find the data structure for the chromInfo data from http://hgdownload.cse.ucsc.edu/goldenpath/hg19/database/chromInfo.txt.gz

Value

```
df Data frame for chromInfo data (rows are sorted if sort.chr is set to TRUE) chromosome Sorted chromosome names chr.len Length of chromosomes. Order are same as chromosome
```

Examples

```
data = read.chromInfo(species = "hg19")
data = read.chromInfo(chromInfo = system.file(package = "circlize", "extdata", "chromInfo.txt"))
chromInfo = read.table(system.file(package = "circlize", "extdata", "chromInfo.txt"),
        colClasses = c("character", "numeric"), sep = "\t")
data = read.chromInfo(chromInfo = chromInfo)
```

read.cytoband

Read/parse cytoband data from a data frame/file/UCSC database

Description

Read/parse cytoband data from a data frame/file/UCSC database

```
read.cytoband(cytoband = system.file(package = "circlize",
    "extdata", "cytoBand.txt"), species = NULL,
    chromosome.index = usable_chromosomes(species),
    sort.chr = TRUE)
```

reverse.circlize 105

Arguments

cytoband Path of the cytoband file or a data frame that already contains cytoband data

species Abbreviations of species. e.g. hg19 for human, mm10 for mouse. If this value

is specified, the function will download cytoBand.txt.gz from UCSC website

automatically.

chromosome.index

subset of chromosomes, also used to reorder chromosomes.

sort.chr Whether chromosome names should be sorted (first sort by numbers then by

letters). If chromosome.index is set, this argument is enforced to FALSE

Details

The function read the cytoband data, sort the chromosome names and calculate the length of each chromosome. By default, it is human hg19 cytoband data.

You can find the data structure of the cytoband data from http://hgdownload.cse.ucsc.edu/goldenpath/hg19/database/cytoBand.txt.gz

Value

```
df Data frame for cytoband data (rows are sorted if sort.chr is set to TRUE) chromosome Sorted chromosome names chr.len Length of chromosomes. Orders are same as chromosome
```

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

Examples

```
data = read.cytoband(species = "hg19")
data = read.cytoband(cytoband = system.file(package = "circlize", "extdata", "cytoBand.txt"))
cytoband = read.table(system.file(package = "circlize", "extdata", "cytoBand.txt"),
    colClasses = c("character", "numeric", "numeric", "character", "character"), sep = "\t")
data = read.cytoband(cytoband = cytoband)
```

reverse.circlize

Convert to data coordinate system

Description

Convert to data coordinate system

```
reverse.circlize(x, y, sector.index = get.current.sector.index(),
    track.index = get.current.track.index())
```

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Arguments

X	degree values. The value can also be a two-column matrix/data frame if you put x and y data points into one variable.
у	distance to the circle center (the radius)
sector.index	Index for the sector where the data coordinate is used
track.index	Index for the track where the data coordinate is used

Details

This is the reverse function of circlize. It transform data points from polar coordinate system to a specified data coordinate system.

Value

A matrix with two columns (x and y)

Examples

```
pdf(NULL)
factors = letters[1:4]
circos.initialize(factors, xlim = c(0, 1))
circos.trackPlotRegion(ylim = c(0, 1))
reverse.circlize(c(30, 60), c(0.9, 0.8))
reverse.circlize(c(30, 60), c(0.9, 0.8), sector.index = "d", track.index = 1)
reverse.circlize(c(30, 60), c(0.9, 0.8), sector.index = "a", track.index = 1)
circos.clear()
dev.off()
```

set.current.cell

Set flag to current cell

Description

Set flag to current cell

Usage

```
set.current.cell(sector.index, track.index)
```

Arguments

```
sector.index sector index track.index track index
```

show.index 107

Details

After setting the current cell, all functions which need sector.index and track.index arguments and are applied to the current cell do not need to specify the two arguments explicitly.

Examples

```
pdf(NULL)
circos.initialize(letters[1:8], xlim = c(0, 1))
circos.track(ylim = c(0, 1))
circos.info()
set.current.cell("b", 1)
circos.info()
circos.clear()
dev.off()
```

show.index

Label the sector index and the track index on each cell

Description

Label the sector index and the track index on each cell

Usage

```
show.index()
```

Details

This function is deprecated, please use circos.info instead.

References

Gu, Z. (2014) circlize implements and enhances circular visualization in R. Bioinformatics.

```
\hbox{\tt\# There is no example}\\ \hbox{\tt NULL}
```

108 uh

 ${\it smartAlign}$

Adjust positions of text

Description

Adjust positions of text

Usage

```
smartAlign(x1, x2, xlim)
```

Arguments

x1 position which corresponds to the top of the text
 x2 position which corresponds to the bottom of the text
 x1im ranges on x-axis

Details

used internally

Examples

```
\# There is no example NULL
```

uh

Convert units

Description

Convert units

Usage

```
uh(...)
```

Arguments

... pass to convert_length

Details

This function is same as convert_length.

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Author(s)

```
Zuguang Gu <z.gu@dkfz.de>
```

Examples

```
# see example in `convert_length` page
NULL
```

ux

Convert unit on x direction in data coordinate

Description

Convert unit on x direction in data coordinate

Usage

```
ux(...)
```

Arguments

```
... pass to convert_x
```

Details

This function is same as convert_x.

Author(s)

```
Zuguang Gu <z.gu@dkfz.de>
```

```
# see example in `convert_x` page
NULL
```

\$.CELL_META

uy

Convert unit on y direction in data coordinate

Description

Convert unit on y direction in data coordinate

Usage

```
uy(...)
```

Arguments

```
... pass to convert_y
```

Details

This function is same as convert_y.

Author(s)

Zuguang Gu <z.gu@dkfz.de>

Examples

```
# see example in `convert_y` page
NULL
```

\$.CELL_META

Easy to way to get meta data in the current cell

Description

Easy to way to get meta data in the current cell

Usage

```
## S3 method for class 'CELL_META'
x$name
```

Arguments

x name of the variable should be "CELL_META"

name of the cell meta name

\$.CELL_META

Details

The variable CELL_META can only be used to get meta data of the "current" cell. Basically you can simply replace e.g. get.cell.meta.data("sector.index") to CELL_META\$sector.index.

See Also

```
get.cell.meta.data
```

Examples

 $\label{eq:continuous_problem} \mbox{\ensuremath{\mbox{\textbf{T}}}} \mbox{\ensuremath{\mbox{\textbf{T}}}} \mbox{\ensuremath{\mbox{\textbf{e}}}} \mbox{\ensuremath{\mbox{\textbf{e}}}$

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