Package ‘RCircos’

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Description A simple and flexible way to generate Circos 2D track plot images for genomic data visualization is implemented in this package. The types of plots include: heatmap, histogram, lines, scatterplot, tiles and plot items for further decorations include connector, link (lines and ribbons), and text (gene) label. All functions require only R graphics package that comes with R base installation.
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RCircos-package

Circos-like Plot of Genomic Data

Description

This is the R implement of basic Circos plots of genomic data.
Details
RCircos package provides a simple and flexible way to make Circos 2D track plots with R and could be easily integrated into other R data processing and graphic manipulation pipelines for presenting large-scale multi-sample genomic research data. It can also serve as a base tool to generate complex Circos images. Currently, RC circos supports genomic data plot for three species: human, mouse, and rat. Data plots include chromosome ideogram with cytoband, heatmap, histogram, lines, scatterplot, and tiles. Plot items for further decorations have connectors, links (lines and ribbons), and text (gene) labels. Each plot is implemented with a specific function and input data for all functions are data frames which can be objects read from text files or generated with other R pipelines.

Author(s)

Henry H Zhang
Maintainer: Henry H Zhang <hzhang.henry@gmail.com>

References

Hongen Zhang, Paul Meltzer, and Sean Davis. RCircos: an R package for Circos 2D track plots. BMC Bioinformatics, 2013, 14:244.

RCircos.Adjust.Scatter.Values

Adjust Scatter Plot Data to Fit the Data Track

Description

Adjust scatter plot values to fit the plot track. The data with lowest value will be plot on the centre of first subtrack and the one with highest value will be on the centre of top subtrack. After this adjustment, values could be added to inside position of plot track as final plot positions. This function is mainly for internal use.

Usage

RCircos.Adjust.Scatter.Values(scatter.values=NULL, min.value=NULL, max.value=NULL, track.height=NULL, subtrack=NULL)
### arguments

- **scatter.values**: Numeric vector, the data to be plotted.
- **min.value**: Numeric, minimum value of scatter data.
- **max.value**: Numeric, maximum value of scatter data.
- **track.height**: Non-negative numeric, height of plot track.
- **subtrack**: Non-negative numeric, the number of sub tracks.

### Author(s)

Henry Zhang

---

**RCircos.Area.Highlight**

*Highlight Plot Area Cross One or More Tracks*

### Description

Highlight a plot area with transparent color cross one or more tracks. RCircos core components and graphic device must be initialized first.

### Usage

```r
RCircos.Area.Highlight(highlight.area=NULL, track.num=NULL, side=c("in", "out"), highlight.color=rgb(0.5, 0.5, 0, 0.5), inside.pos=NULL, outside.pos=NULL)
```

### Arguments

- **highlight.area**: Vector with chromosome name, start and end position to be highlighted.
- **track.num**: Vector of non-negative integer, which track or track(s) to be highlighted.
- **side**: Character vector, location relative to the ideogram, either "in" or "out".
- **highlight.color**: An RGB color definition, alpha value must be defined.
- **inside.pos**: Non-negative numeric, inside position (relative to the centre of plot area) of the track.
- **outside.pos**: Non-negative numeric, outside position (relative to the centre of plot area) of the track.

### Author(s)

Henry Zhang
Examples

```r
## Not run:
library(RCircos)
data(UCSC.HG19.Human.CytoBandIdeogram)
data(RCircos.Gene.Label.Data)
  chr.exclude=NULL, tracks.inside=10, tracks.outside=0)
RCircos.Set.Plot.Area()
RCircos.Chromosome.Ideogram.Plot()

highlight.area <- c("chr1", 100000, 200000)
RCircos.Area.Highlight(highlight.area=c("chr1", 100000, 200000),
  track.num=c(1:3), side="in")
RCircos.Area.Highlight(highlight.areac="chr10", 100000, 200000),
  inside.pos=1.5, outside.pos=2)

## End(Not run)
```

---

**RCircos.Area.Plot**  
**Paint Areas on One Data Track**

**Description**

Paint area on one data track with different height and different locations inside of a track. Plot types includes "mountain", "curtain", and "band". Plot data should have genomic positions (chromosome names, start and end positions) as well as height values. For band type plot, two columns of height values are required.

**Usage**

```r
RCircos.Area.Plot(area.data=NULL, data.col=c(4,5),
  track.num=NULL, side=c("in", "out"),
  plot.type=c("mountain", "curtain", "band"),
  min.value=NULL, max.value=NULL, area.color="gray",
  border.col="black", inside.pos=NULL, outside.pos=NULL,
  genomic.columns=3, is.sorted=TRUE)
```

**Arguments**

- `area.data`: A data frame with two or three columns for genomic positions, one or more columns for heights of each data point, and an optional column for polygon colour.
- `data.col`: Non-negative integer, representing the ordinal number of the column in input data set that contains the data to be plotted.
- `track.num`: Non-negative integer, representing the ordinal number of the plot track where the lines will be plotted.
side Character vector, either "in" or "out", representing the position related to chromosome ideogram.

plot.type Character vector, either "mountain", "curtain", or "band"

min.value Numeric, minimum value in data column of polygon data.

max.value Numeric, maximum value in data column of polygon data.

area.color Color names for fill of the area.

border.col Color name for border color, default null.

inside.pos Non-negative numeric, inside position (relative to the centre of plot area) of the track.

outside.pos Non-negative numeric, outside position (relative to the centre of plot area) of the track.

genomic.columns Non-negative integer, total number of columns for genomic position in each row. Must be either 3 or 2.

is.sorted Logic, whether the data is sorted by chromosome names and start positions.

Author(s)

Henry Zhang

Examples

```R
## Not run:
library(RCircos);
data(UCSC.HG19.Human.CytoBandIdeogram);
data(RCircos.Polygon.Data);

RCircos.Set.Core.Components(
    cyto.info=UCSC.HG19.Human.CytoBandIdeogram,
    chr.exclude=NULL, tracks.inside=10, tracks.outside=5)
RCircos.Set.Plot.Area();
RCircos.Chromosome.Ideogram.Plot()

load("RCircos/data/RCircos.Histogram.Data.RData")

area.data <- RCircos.Histogram.Data;
adj.value <- runif(nrow(area.data), 0, 0.4)
area.data["DataT"] <- 0.5 + adj.value
area.data["DataB"] <- 0.5 - adj.value

RCircos.Area.Plot(area.data, data.col=4, plot.type="mountain",
    inside.pos=1.2, outside.pos=1.5, is.sorted=FALSE)

RCircos.Area.Plot(area.data, data.col=4, plot.type="curtain",
    inside.pos=0.9, outside.pos=1.1, is.sorted=FALSE)

RCircos.Area.Plot(area.data, data.col=c(5,6), plot.type="band",
    inside.pos=0.4, outside.pos=0.7, is.sorted=FALSE)
## End(Not run)
```
Description

Draw chromosome ideogram, chromosome highlights and chromosome names. Graphic device must be initialized first. The original function was split to three new ones in order to plot chromosome highlight or names in different area. RCircos core components and graphic device must be initialized first.

Usage

RCircos.Chromosome.Ideogram.Plot(tick.interval=0)
RCircos.Draw.Chromosome.Ideogram(ideo.pos=NULL, ideo.width=NULL)
RCircos.Highlight.Chromosome.Ideogram(highlight.pos=NULL, highlight.width=NULL)
RCircos.Ideogram.Tick.Plot(tick.interval=50, track.for.ticks=3)
RCircos.Label.Chromosome.Names(chr.name.pos=NULL)

Arguments

tick.interval Non-negative integer, the distance between two chromosome ideogram ticks in million base pairs. Set to 0 (zero) will ignore the ideogram ticks.
ideo.pos Non-negative numeric, the distance of chromosome ideogram position from center of plot area.
ideo.width Non-negative numeric, the width of chromosome ideogram.
highlight.pos Non-negative numeric, the distance of chromosome highlight line position from center of plot area.
highlight.width Non-negative integer, the width of chromosome highlight lines.
track.for.ticks Non-negative integer, total number of tracks for ideogram tick height.
chr.name.pos Non-negative numeric, the distance of chromosome names from center of plot area.

Details

RCircos.Chromosome.Ideogram.Plot() is the default method to plot chromosome ideogram, chromosome highlights, and chromosome names.
RCircos.Highlight.Chromosome.Ideogram() highlights chromosomes with different color lines only.
RCircos.Ideogram.Tick.Plot() draws chromosome ideogram ticks starting from chromosome highlight positions.
RCircos.Label.Chromosome.Names(chr.name.pos=NULL) add chromosome names for chromosome ideogram at default or customized location.
**Author(s)**

Henry Zhang

**Examples**

```r
## Not run:
library(RCircos)
data(UCSC.HG38.Human.CytoBandIdeogram)
cyto.info <- UCSC.HG38.Human.CytoBandIdeogram
RCircos.Set.Core.Components(cyto.info, chr.exclude=NULL,
    tracks.inside=10, tracks.outside=0)
RCircos.Set.Plot.Area()
RCircos.Chromosome.Ideogram.Plot()

## End(Not run)
```

---

**RCircos.Clear.Track**  
*Erase One or More Data Tracks*

**Description**

Erase one or more data track or all tracks from current track to the center of plot area. RCircos core components and graphic device must be already initialized.

**Usage**

```r
RCircos.Clear.Track(track.num=NULL, side=NULL, to.center=FALSE,
    inside.pos=NULL, outside.pos=NULL)
```

**Arguments**

- **track.num**: Non-negative integer, representing the ordinal number of the plot track to be erased.
- **side**: Character vector, either "in" or "out", representing the position related to chromosome ideogram. It could be NULL is both inside.pos and outside.pos are defined.
- **to.center**: Logic, FALSE for erasing current track only and TRUE for erasing all tracks from current track to the center of plot plot area.
- **inside.pos**: Non-negative numeric, inside position (relative to the centre of plot area) of the track.
- **outside.pos**: Non-negative numeric, outside position (relative to the centre of plot area) of the track.

**Author(s)**

Henry Zhang
Examples

```r
## Not run:
library(RCircos);
data(UCSC.HG19.Human.CytoBandIdeogram)
data(RCircos.Link.Data)
data(RCircos.Scatter.Data)

                           chr.exclude=c("chrX", "chrY"),
                           tracks.inside=10, tracks.outside=0 )

RCircos.Set.Plot.Area()
RCircos.Chromosome.Ideogram.Plot()

RCircos.Scatter.Plot(RCircos.Scatter.Data, data.col=5,
                      track.num=1, side="in", by.fold=1)
RCircos.Link.Plot(RCircos.Link.Data, track.num=3)

RCircos.Clear.Track(track.num=1, side="in", to.center=FALSE)
RCircos.Clear.Track(track.num=5, side="in", to.center=TRUE)

## End(Not run)
```

**RCircos.Clear.Zoom.Area**

_Erase Zoom-in Plot Area_

**Description**

Erase zoom-in plot area. RCircos core components and graphic device must be already initialized.

**Usage**

```
RCircos.Clear.Zoom.Area(zoom.pos=NULL, track.num=NULL,
                        inside.pos=NULL, outside.pos=NULL)
```

**Arguments**

- `zoom.pos`: Non-negative integer vector, representing the index of RCircos base plot positions.
- `track.num`: Non-negative integer, representing the ordinal number of the plot track to be erased.
- `inside.pos`: Non-negative numeric, inside position (relative to the centre of plot area) of the track.
- `outside.pos`: Non-negative numeric, outside position (relative to the centre of plot area) of the track.
Author(s)

Henry Zhang

Examples

```r
## Not run:
library(RCircos)
data(UCSC.HG38.Human.CytoBandIdeogram)
cyto.info <- UCSC.HG38.Human.CytoBandIdeogram
RCircos.Set.Core.Components(cyto.info, chr.exclude=NULL,
   tracks.inside=10, tracks.outside=0 )

RCircos.Set.Plot.Area()
RCircos.Clear.Zoom.Area(zoom.pos=c(1:10000), inside.pos=2, outside.pos=2.5)

## End(Not run)
```

Description

Draw connection lines (simple lines plot) between two sets of data points. One example of usage is to label genes at modified plot position and connect the gene label to its genomic position. The core components and graphics device must be initialized before drawing.

Usage

```r
RCircos.Customized.Connection.Plot(gene.data, label.data,
gene.pos=NULL, label.pos=NULL)
```

Arguments

gene.data A data frame with chromosome names and actual genomic positions for the genes to be labeled.

label.data A data frame with chromosome name and genomic positions that will be used to label gene names.

gene.pos Float numeric, scale factor relative to the center of plot area (0) for start points of connection lines.

label.pos Float numeric, scale factor relative to the center of plot area (0) for end points of connection lines.

Author(s)

Henry Zhang
Description

Plot one customized shape on a data track. RCircos core components and graphic device must be initialized first. The customized shape should be represented by coordinates of a polygon inside a circle with radius of 1 and default location is at 12 o’clock. When plotting, the polygon center will be scaled and transformed for new size and location. For example, following code will plot an arrow:

```
polygonX <- c(0, -0.7, -0.2, -0.2, 0.2, 0.2, 0.7, 0)
polygonY <- c(-1, 0.7, 0.4, 1, 1, 0.4, 0.7, -1)
plot(c(-2, 2), c(-2, 2))
polygon(polygonX, polygonY, col="red")
```

Usage

```
RCircos.Customized.Shape.Plot(shape.data=NULL, track.num=NULL,
   side=c("in", "out"), location=NULL, shape.color="red",
   inside.pos=NULL, outside.pos=NULL)
```

Arguments

- `shape.data`: A two dimensional numeric matrix for coordinates of a polygon.
- `track.num`: Non-negative integer, representing the ordinal number of the plot track where the lines will be plotted.
- `side`: Character vector, either "in" or "out", representing the position related to chromosome ideogram.
- `location`: Vector with chromosome name, start position, and end position where the shape to be drawn.
- `shape.color`: Character vector, color for the shape.
- `inside.pos`: Non-negative numeric, inside position (relative to the centre of plot area) of the track.
- `outside.pos`: Non-negative numeric, outside position (relative to the centre of plot area) of the track.

Author(s)

Henry Zhang

Examples

```
## Not run:
library(RCircos)

data(UCSC.HG19.Human.CytoBandIdeogram)
```
RCircos.Data.Point

**Description**

Calculate the index of Circos plot positions (a set of x- and y-coordinates for a circular line) for a genomic position (chromosome name and start position). This function is for internal use only.

**Usage**

RCircos.Data.Point(chromosome=NULL, start.posotion=NULL)

**Arguments**

- **chromosome**: Character vector, a chromosome name with prefix of "chr", e.g., "chr1".
- **start.posotion**: Non-negative integer, start position of a genomic interval (e.g. a gene or a band) on the chromosome.

**Value**

An integer representing the index of RCircos plot positions.

**Author(s)**

Henry Zhang
**RCircos.defaultBasePerUnits**

*The Default Base Pair Number of A Circos Unit*

**Description**

The default total number of base pairs a Circos unit represents. This is a read only parameter and the actual value in a RCircos session could be modified with get and reset methods for plot parameters.

**Format**

The format is: num 30000

---

**RCircos.defaultCharWidth**

*Default Character With in Circos Unit*

**Description**

Default Circos units of a character on RCircos plot. This is the character size for best resolution and is a read only parameter.

**Format**

The format is: num 500

---

**RCircos.defaultChromPadding**

*The Default Value of GAP in Circos (Chromosome) Unit between two Chromosome Ideogram*

**Description**

The default total unit of a gap bewtween the ends of two neighbor chromosome ideograms. This is a read only parameter.

**Format**

The format is: num 300
### Default Circos Units for RCircos

**Description**

The default Circos units for RCircos. A Circos unit is a point which represents an interval of genome length on the circumference of a circle line. By default an unit represents a point which equal to the length of 30000 base pairs and length of human genome is used to calculate the total number of Circos units. This is a read only parameter and is for internal use to modify plot parameters for RCircos plot with non-human genomes.

**Format**

The format is: `num 103190`

### Default Character Size of RCircos Plot

**Description**

The default character size for text plot of RCircos in best resolution. This is a read only parameter and the actual value in a RCircos session could be modified with get and reset methods for plot parameters.

**Format**

The format is: `num 0.4`

### RCircos Environment

**Description**

RCircos.Env is the user environment to hold RCircos core components and other objects for RCircos plot. Each object in the environment can be viewed with get method.

**Format**

The format is: `<environment: 0xd67930>"
RCircos.Gene.Connector.Plot

*Draw Connectors between Chromosome Ideogram and Gene Names*

**Description**

Draw a set of connectors (three lines for each connector) between chromosome ideogram or a data track and gene names. RCircos core components and graphics device must be initialized before drawing.

**Usage**

```
```

**Arguments**

- `genomic.data`: A data frame with the first four columns for chromosome names, start positions, end positions, and gene names.
- `track.num`: Non-negative integer, representing the ordinal number of the plot track where the connectors are plotted.
- `side`: Character vector, either "in" or "out" representing the position related to chromosome ideogram.
- `inside.pos`: Non-negative numeric, inside position (relative to the centre of plot area) of the track.
- `outside.pos`: Non-negative numeric, outside position (relative to the centre of plot area) of the track.
- `genomic.columns`: Non-negative integer, total number of columns for genomic position in each row. Must be either 3 or 2.
- `is.sorted`: Logic, whether the data is sorted by chromosome names and start positions.

**Author(s)**

Henry Zhang

**Examples**

```
## Not run:
library(RCircos)
data(RCircos.Gene.Label.Data)
data(UCSC.HG19.Human.CytoBandIdeogram)

RCircos.Set.Core.Components(cyto.info, chr.exclude=NULL,
```
tracks.inside=10, tracks.outside=0)

RCircos.Set.Plot.Area()
RCircos.Chromosome.Ideogram.Plot()
RCircos.Gene.Connector.Plot(gene.data, track.num=1, side="in")
   inside.pos=2.5, outside.pos=3)

## End(Not run)

---

\textbf{RCircos.Gene.Label.Data}

\textit{Sample Data for Gene Labels}

\section*{Description}

RCircos.Gene.Label.Data contains genomic position information for 192 genes. Each row of the
data contains genomic position and gene name for one gene in the order of chromosome name, start
position, end position, and gene name.

\section*{Usage}

data(RCircos.Gene.Label.Data)

\section*{Format}

A data frame with 192 observations on the following 4 variables.

\begin{verbatim}
Chromosome a factor with levels chr1 chr10 chr11 chr12 chr13 chr14 chr15 chr16
   chr17 chr18 chr19 chr20 chr21 chr22 chr3 chr4 chr5 chr6 chr7 chr8
   chr9 chrX
chromStart a numeric vector
chromEnd a numeric vector
Gene  a factor with levels ABL1 ACN9 ACVR1B AKT1 ALK APC ARHGEF2 ARID1A ASXL1 ATM ATRX
   BAG3 BA13 BAP1 BCAN BCL2 BRAF BRCA1 BRCA2 CA12 CA9 CALU CARD11 CBL
   CCN1 CD34 CDC7 CDC1 CDK4 CDK2A CDX2 CEACAM7 CEBPA CES3 CRLF2
   CSF1R CSF3 CTNNA1 CTNNB1 CUL2 CYLD CYP1A1 DAXX DCC DES DIRAS1
   DIRC2 DKK3 DLD DMBT1 DMNT3A DPYD EGF EGFR EGR3 EIF4G2 EML4
   ENO1 ENO2 ERBB2 ERBB3 ERBB4 ERG EZH2 F10 FAM12B FBXW7 FGFR1 FGFR2
   FGFR3 FH FHT1 FLCN FLT3 FOXL2 FZR1 GATA1 GATA2 GATA3 GFAP GNA11
   GNAQ GNAS GRP GSTM1 HIVEP3 HNF1A HRAS IDH1 IDH2 IFNA1 IL2 ITGB5
   JAK1 JAK2 JAK3 KDR KIT KRAS KRT20 MAGI1 MAP2K4 MEN1 MET MGMT
   MIB1 MKI67 MLH1 MPL MSH2 MSH6 MUC1 MUC17 MUC2 MUTYH MVP MYC
   MYD88 NAT1 NAT2 NES NF1 NF2 NOTCH1 NOTCH2 NPM1 NRAS PBRM1
   PCNA PDGFRA PDZD4 PGR PHOX2B PIK3CA PMS2 PPP2R1A PRCC PRKAR1A
   PTCH1 PTEN PTGS2 PTPN11 RB1 REEP5 RET RNF139 RNFL
   RUNX1 SDHA SDHAF1 SDHAF2 SDHB SDHD SELT SEF2 SLC38A1
   SLC6A2 SLTM SMDA4 SMARCA4 SMARCBI SMOK1 SOCS1 SRC SST STC1
   STK11 SUFU SYT TCF7L2 TET2 TFE3 TFE6 TME97 TNFAIP3 TNFSF13
   TP53 TPDS2L2 TPM4 TSC1 TSC2 TSHR TYK2 VHL VIM WT1 XRCC1
   ZNF135
\end{verbatim}
Source

Unpublished data.

---

**RCircos.Gene.Name.Plot**

*Draw Gene Names on a Data Track*

---

**Description**

Label gene names along chromosome ideogram or a data track. RCircos core components and graphic device must be initialized before drawing.

**Usage**

```r
RCircos.Gene.Name.Plot(gene.data=NULL, name.col=NULL, track.num=NULL, side="in", inside.pos=NULL, outside.pos=NULL, genomic.columns=3, is.sorted=FALSE)
```

**Arguments**

- **gene.data**
  A data frame. The first three columns should be chromosome name, start position, end position.

- **name.col**
  Non-negative integer, representing the ordinal number of the column in input data that contains gene names.

- **track.num**
  Non-negative integer, representing the ordinal number of the plot track where the gene names are plotted.

- **side**
  Character vector, either "in" or "out", representing the position related to chromosome ideogram.

- **inside.pos**
  Non-negative numeric, inside position (relative to the centre of plot area) of the track.

- **outside.pos**
  Non-negative numeric, outside position (relative to the centre of plot area) of the track.

- **genomic.columns**
  Non-negative integer, total number of columns for genomic position in each row. Must be either 3 or 2.

- **is.sorted**
  Logic, whether the data is sorted by chromosome names and start positions.

**Author(s)**

Henry Zhang
Examples

```r
## Not run:
library(RCircos)
data(UCSC.HG19.Human.CytoBandIdeogram)
data(RCircos.Gene.Label.Data)

cyto.info <- UCSC.HG19.Human.CytoBandIdeogram
RCircos.Set.Core.Components(cyto.info,
    chr.exclude=c("chrX", "chrY"),
    num.inside=10, num.outside=0)

RCircos.Set.Plot.Area()
RCircos.Chromosome.Ideoagram.Plot()

RCircos.Gene.Name.Plot(gene.data, name.col=4,
    track.num=2, side="in")
RCircos.Gene.Name.Plot(gene.data, name.col=4,
    track.num=2, side="out",
    inside.pos=2.5, outside.pos=3)

## End(Not run)
```

---

### RCircos.Get.Arrow.Shape

*Get Default Coordinates for An Arrow Shape*

**Description**

Get default coordinates for an arrow shape. The arrow is represented as a polygon inside of a circle with radius of 1. The coordinates will be different based which side of chromosome ideogram the arrow will be plotted.

**Usage**

```r
RCircos.Get.Arrow.Shape(side="in")
```

**Arguments**

- `side` Character vector, either "in" or "out".

**Value**

A two dimensional numeric matrix for x and y coordinates of a polygon.

**Author(s)**

Henry Zhang
Examples

    library(RCircos)
    arrow <- RCircos.Get.Arrow.Shape("in")
    arrow <- RCircos.Get.Arrow.Shape("out")

---


*Find the order of Chromosome Names*

Description

Generate an ordered chromosome names from input. For human and other mammalian animals, numeric names (integers or Roman numbers) will go first followed by chromosome X, Y, and M. If all chromosome names are character only, they will be in alphabetical order.

Usage

    RCircos.Get.Chromosome.Order(chromosomes=NULL)

Arguments

  chromosomes    Character vector, names of chromosomes.

Value

  Character vector, ordered chromosome name list.

Author(s)

  Henry Zhang

Examples

    library(RCircos)
    chromosomes <- paste0("chr", c(1:10, 21,22, 11:20, "X", "Y"))
RCircos.Get.Core.Components

Methods to Retrieve RCircos Core Components Used in Current RCircos Session

Description

Retrieve base plot positions, plot ideogram, and plot parameters used in current RCircos session. RCircos core components must be initialized first.

Usage

```
RCircos.Get.Plot.Parameters()
RCircos.Get.Plot.Positions()
RCircos.Get.Plot.Ideogram()
```

Value

- `RCircos.Get.Plot.Parameters()` returns a list that holding all parameters used in current RCircos session.
- `RCircos.Get.Plot.Positions()` returns a data frame with three columns for x, and y coordinates of a circular line with radius of 1 as well as the degrees used for text rotation at each point of the positions.
- `RCircos.Get.Plot.Ideogram()` returns a data frame containing all information for chromosome ideogram plot. The first five column are orginal cytoband data and following are chromosome highlight colors, and plot colors, length, number of chromosome unit, and relative location on the circular layout of each band.

Author(s)

Henry Zhang

RCircos.Get.Data.Point.Height

Calculate the Height of Data in A Track

Description

Calculate data point height inside a plot track such as scatter location, top or bottom location of a bar, layer of a tile or parallel link line.

Usage

```
RCircos.Get.Data.Point.Height(plot.values=NULL, min.value=NULL, max.value=NULL, plot.type=NULL, track.height=NULL)
```
Arguments

- **plot.values**: Numeric, the data to be plotted on a data track.
- **min.value**: Numeric, the minimum value of data range.
- **max.value**: Numeric, the maximum value of data range.
- **plot.type**: Character vector, plot type, valid values are "bar", "histogram", "uniform", or "points".
- **track.height**: Non-negative numeric, height of plot track.

Value

Numeric vector with values between 0 ~ 1

Author(s)

Henry Zhang

Examples

```r
data.values <- runif(1000, -4, 11)
library(RCircos)
data.height <- RCircos.Get.Data.Point.Height(data.values,
  min.value=-4, max.value=14, plot.type="points",
  track.height=NULL)
```

---

RCircos.Get.Default.Parameters

Retrieve Parameter Values Stored in RCircos Environment

Description

These methods are used to view the default parameters stored in RCircos Environments.

Usage

```r
RCircos.Get.Padding.Constant()
RCircos.Get.Supported.HeatmapColors()
RCircos.Get.Supported.Plot.Types()
```
**Value**

RCircos.Get.Default.Circos.Units() returns the default number of units (points) for a circular line. This value is derived from human genome length (3x10^9) and one unit represents length of 3000 base pairs.

RCircos.Get.Default.Base.Per.Units() returns the default number of base pairs that a unit (point) will represent.

RCircos.Get.Default.Chrom.Padding() returns the default length of gaps between two neighboring chromosomes in chromosome (Circos) units.

RCircos.Get.Padding.Constant() returns the default ratio of chromosome padding length and total default Circos (chromosome) units.

RCircos.Get.Supported.HeatmapColors() returns the types of colors for heatmap plot.

RCircos.Get.Supported.Plot.Types() returns the plot types supported by current RCircos version.

RCircos.Get.Default.Char.Width() returns the default number of units a character will span on the plot.


**Author(s)**

Henry Zhang

---


*Calculate Plot Positions for Gene Labels*

**Description**

This function calculates new plot positions from genomic positions for gene labeling. In case there are too many genes in a genomic interval, the labels may become overlapped so that correct numbers and new plot positions are often needed to make the labels readable. If all labels cannot be plotted for the given genomic interval extra labels will be ignored. This function is for internal use only.

**Usage**


**Arguments**

- **genomic.data** A data frame with leading columns for chromosome name, start position, and/or end position.
- **genomic.columns** Non-negative integer, total number of columns for genomic position in each row. Must be either 3 or 2.
- **is.sorted** Logic, whether the data is sorted by chromosome names and start positions.

Value

All or subset of input data frame with a new column for plot positions.

Author(s)

Henry Zhang

Description

Calculate maximum number of gene names to be plotted for each chromosome based on plot RCircos plot parameters. RCircos core components must be initialized first.

Usage


Value

A data frame with names, maximum number of labels could be plotted, start position, end position, and label width for each chromosome.

Author(s)

Henry Zhang

RCircos.Get.Heatmap.Color.Scale

Generate Color Scales for Heatmap Plot

Description

Create color scales for heatmap plot. Currently there are six color scales could be generated:

- **BlueWhiteRed**: colors from blue (lowest) to white then red (highest)
- **GreenWhiteRed**: colors from green (lowest) to white then red (highest)
- **GreenYellowRed**: colors from green (lowest) to yellow then red (highest)
- **GreenBlackRed**: colors from green (lowest) to black then red (highest)
- **YellowToRed**: colors from yellow (lowest) to red (highest)
- **BlackOnly**: black only
Usage

RCircos.Get.Heatmap.Color.Scale(heatmap.color=NULL)

Arguments

heatmap.color  Character vector, one of "BlueWhiteRed", "GreenWhiteRed", "GreenYellowRed", "GreenBlackRed", "YellowToRed", and "BlackOnly".

Value

A vector or two column matrix of rgb colors.

Author(s)

Henry Zhang

Examples

library(RCircos)


Calculate X and Y Coordinates for Color Scale

Description

Calculate X and Y coordinates for the topleft corner of heatmap color scale.

Usage

RCircos.Get.Heatmap.Color.Scale.Location(scale.location = 1)

Arguments

scale.location  Integer of 1 ~ 12, represents the plot location (default is 1):
  1: "bottom"
  2: "left"
  3: "top"
  4: "right"
  5: "bottomleft"
  6: "bottomright"
  7: "leftright"
  8: "leftbottom"
  9: "topleft"
 10: "topright"
 11: "righttop"
 12: "rightbottom"

Value

Numeric vector of length 2 for x and y coordinates

Author(s)

Henry Zhang

Examples

```r
library(RCircos)
location <- RCircos.Get.Heatmap.Color.Scale.Location(1)
```


*Set Colors for One Track of Heatmap*

Description

Assign colors to each value of a numeric vector for heatmap plot. The colour scale needs to be calculated from whole datasets when there are more than one data column so the minimum and maximum values for heatmap data must be provided.

Usage

```r
RCircos.Get.Heatmap.Data.Colors(heatmap.value=NULL, min.value=NULL, max.value=NULL)
```

Arguments

- `heatmap.value` A numeric vector.
- `min.value` Numeric, the minimum value for heatmap scale.
- `max.value` Numeric, the maximum value for heatmap scale.

Value

Character vector of R color names with length same as the length of data values.

Author(s)

Henry Zhang

Examples

```r
## Not run:
library(RCircos)
data(RCircos.Heatmap.Data)
## End(Not run)
```
RCircos.Get.Link.Colors

_Set up Colors for Link Lines or Ribbons_

**Description**

Assign colors for link lines or Ribbons. The default colors are rainbow with length of total link lines or ribbons. If `by.chromosome` is set to TRUE, red color will be used for links between positions of same chromosomes and blue color for links between different chromosomes. Users can also define their own colors by appending a column with color names to link data and give the column name as "PlotColor".

This function is for internal use only.

**Usage**

```r
circos.Get.Link.Colors(link.data, genomic.columns=3, by.chromosome=TRUE)
```

**Arguments**

- **link.data**
  A data frame with paired genomic positions in each row. The first three columns are for chromosome names, start and end position of one genomic position followed by three columns for chromosome names, start and end position of the second genomic position.

- **genomic.columns**
  Non-negative integer, total columns that define the positions of a genomic interval (chromosome name, start and optional end positions).

- **by.chromosome**
  Logic. If true, red color will be used for links between positions of same chromosomes and blue color for links between different chromosomes. Otherwise, user defined or random colors (rainbow) will be automatically assigned for each line.

**Value**

A vector of color names with length same as the rows of input data.

**Author(s)**

Henry Zhang

*Calculate Plot Positions for A Genomic Interval*

**Description**

Calculate x and y coordinates for a genomic interval which needs two plot positions (start and end) such as link lines/ribbons, tiles, parallel lines. As a set of points for a circular line is held in the RCircos session, we only need the index of the point for each genomic position.

**Usage**

```r
```

**Arguments**

- **genomic.data**: A data frame contains paired genomic positions (chromosome names, start and end positions). The data does not need to be sorted.
- **genomic.columns**: Non-negative integer, total number of columns for genomic position (chromosome name, start and/or end position).
- **plot.type**: Character vector, either "link", "ribbon", "pLink", "polygon", or "tile".

**Value**

A data frame same as input but with two new columns for index of plot positions on circular line.

**Author(s)**

Henry Zhang

**Examples**

```r
## Not run:
library(RCircos)
data(UCSC.HG19.Human.CytoBandIdeogram)
data(RCircos.Link.Data)
## End(Not run)
```
RCircos.Get.Plot.Boundary

Calculate Boundary of A Data Track

Description

Calculate plot track location based on track number, plot side, or user defined inside and outside position relative to plot area center. If all arguments are defined, user defined inside and outside position will be used. This function is mainly for internal use.

Usage

RCircos.Get.Plot.Boundary(track.num=NULL, side=NULL, inside.pos=NULL, outside.pos=NULL, erase.area=FALSE)

Arguments

track.num Non-negative integer, representing the ordinal number of the plot track where the lines will be plotted.
side Character vector, either "in" or "out", representing the position related to chromosome ideogram. It could be NULL if both inside.pos and outside.pos are defined.
inside.pos Non-negative numeric, inside position (relative to the centre of plot area) of the track.
outside.pos Non-negative numeric, outside position (relative to the centre of plot area) of the track.
erase.area Logic, if the track is going to be erased.

Value

Numeric vector of length 2, x and y coordinates of a data track.

Author(s)

Henry Zhang
**RCircos.Get.Plot.Colors**

*Assign Plot Colors for Non-Link Data Plot*

**Description**

Assign plot colors for each data point of genomic data except of link plot (link lines or ribbons). Users can define their own colors by appending a column with color names to plot data and give the column name as "PlotColor". Otherwise, default colors will be used. This function is for internal use only.

**Usage**

```r
RCircos.Get.Plot.Colors(plot.data, color)
```

**Arguments**

- `plot.data` A data frame with the first three columns as chromosome name, start and end position followed by data values for one or more samples.
- `color` Character vector of color names.

**Value**

A vector of color names with length same as the rows of input data.

**Author(s)**

Hongen Zhang

---

**RCircos.Get.Plot.Layers**

*Get the Layer Numbers for Tile Plot*

**Description**

Check out overlaps between different genomic positions on same chromosome and set up layer numbers for each line of the genomic data.

**Usage**

```r
RCircos.Get.Plot.Layers(genomic.data=NULL, genomic.columns=NULL)
```
Arguments

- genomic.data: A data frame with genomic positions (chromosomes, start and end positions) and the positions should be already validated and sorted by chromosome then start position.
- genomic.columns: Non-negative integer, total number of columns for genomic positions.

Value

A non-negative integer vector with length same as the total rows of input data.

Author(s)

Henry Zhang

Examples

```r
library(RCircos)
data(RCircos.Tile.Data)
```

Description

Calculate polygon heights (bottom and top location) inside a plot track. Polygon can be drawn from inside position to outside position, from outside position to inside position, or from the middle to both directions of the track. This is mainly for internal use.

Usage

```r
RCircos.Get.Polygon.Height(data.heights, min.value=NULL, max.value=NULL, inside.pos=NULL, outside.pos=NULL)
```

Arguments

- data.heights: Numeric vector, the data to be plotted on a data track.
- min.value: Numeric, the minimum value of data range.
- max.value: Numeric, the maximum value of data range.
- inside.pos: Non-negative numeric, the close position of a plot track relative to the center of plot area.
- outside.pos: Non-negative numeric, the far position of a plot track relative to the center of plot area.
Value
Non-negative numeric matrix of column 2 representing the bottom and top positions of polygons.

Author(s)
Henry Zhang

Description
Calculate x and y coordinates of one genomic position for each row of a genomic data set. As a set of points for a circular line is held in the RCircos session, we only need the index of the point for each genomic position.

Usage

Arguments
- genomic.data: A data frame contains genomic positions (at least two or three columns for chromosome names, start and/or end positions).
- genomic.columns: Non-negative integer, total number of columns for genomic position (chromosome name, start and/or end position).

Value
A data frame same as input but with a new column for index of plot positions on a circular line.

Author(s)
Henry Zhang

Examples
```r
## Not run:
library(RCircos)
data(RCircos.Heatmap.Data)

## End(Not run)
```
RCircos.Get.Start.End.Locations

*Calculate Start and End Positions for Plot Data*

**Description**

Generate start and end positions from single point for special plot types such as heatmap, histogram, or rectangle-like which need start and end positions along the track. This function is mainly for internal use.

**Usage**

```r
RCircos.Get.Start.End.Locations(plot.data, plot.width)
```

**Arguments**

- `plot.data`: Data frame returned from `RCircos.Get.Single.Point.Positions()` and the last column is the plot locations
- `plot.width`: Non-negative integer in number of base pairs, e.g., width or heatmap or histogram of each data points

**Author(s)**

Henry Zhang

---

RCircos.Get.Track.Positions

*Calculate Plot Positions for A Data Track*

**Description**

Calculate inner and outer plot positions for a data track. RCircos core components must be initialized first. This function is mainly for internal use.

**Usage**

```r
RCircos.Get.Track.Positions(side=NULL, track.num=NULL)
```

**Arguments**

- `side`: Character vector, either "in" or "out", representing the position related to chromosome ideogram.
- `track.num`: Integer, representing the ordinal number of the plot track
Value

Numeric vector of length 2. The outer and inner positions of a track.

Author(s)

Hongen Zhang

---

Retriev Subset of Plot Data for Zoom

Description

Retrieve subset of plot data based on gene name or row header.

Usage

```r
RCircos.Get.Zoom.Data(plot.data=NULL, name.col= NULL, genomic.columns=3, target.gene=NULL, neighbor.genes=5)
```

Arguments

- `plot.data`: A data frame containing genomic positions, gene names, and plot values. The data should be already sorted by chromosome names then start positions.
- `name.col`: Non-negative integer, which column is for gene names.
- `genomic.columns`: Non-negative integer, total number of columns for genomic position in each row. Must be either 3 or 2.
- `target.gene`: Character vector, target gene names.
- `neighbor.genes`: Non-negative integer, number of neighbor genes on each side of taget gene to be included in subset.

Value

Data frame, subset of input data.

Author(s)

Henry Zhang

Examples

```r
## Not run:
library(RCircos)
data(RCircos.Heatmap.Data)

## End(Not run)
```
RCircos.Get.Zoom.Range

*Retrieve Genomic Interval from Genomic Data for Zoom Plot*

**Description**

Extract zoom in range (chromosome name, start position, and end position) from zoom data with a gene name or row header. If using this function to get zoom-in range, there will be no need to call RCircos.Validate.Genomic.Info().

**Usage**

```
RCircos.Get.Zoom.Range(zoom.data=NULL, genomic.columns=3)
```

**Arguments**

- `zoom.data` A data frame with genomic positions, gene names, and plot values for genes to be zoomed-in only.
- `genomic.columns` Non-negative integer, total number of columns for genomic position in each row. Must be either 3 or 2.

**Value**

A list with chromosome names, start end end positions.

**Author(s)**

Henry Zhang

**Examples**

```r
## Not run:
library(RCircos)
data(RCircos.Heatmap.Data)
    name.col=4, target.gene="TP53")
zoom.info <- RCircos.Get.Zoom.Range(zoom.data, genomic.columns=3)

## End(Not run)
```
Sample Data for RCircos Heatmap Plot

Description
A dataset with the first three columns as chromosome names, start position, and end position followed by a column of gene names then columns of gene expression values.

Usage
data(RCircos.Heatmap.Data)

Format
A data frame with 6660 observations on the following 10 variables.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>chromosome</td>
<td>a factor with levels chr1 chr10 chr11 chr12 chr13 chr14 chr15 chr16 chr17 chr18 chr19 chr2 chr20 chr21 chr22 chr3 chr4 chr5 chr6 chr7 chr8 chr9 chrX chrY</td>
</tr>
<tr>
<td>chromStart</td>
<td>a numeric vector</td>
</tr>
<tr>
<td>chromEnd</td>
<td>a numeric vector</td>
</tr>
<tr>
<td>GeneName</td>
<td>a character vector</td>
</tr>
<tr>
<td>X786.0</td>
<td>a numeric vector</td>
</tr>
<tr>
<td>A498</td>
<td>a numeric vector</td>
</tr>
<tr>
<td>A549.ATCC</td>
<td>a numeric vector</td>
</tr>
<tr>
<td>ACHN</td>
<td>a numeric vector</td>
</tr>
<tr>
<td>BT.549</td>
<td>a numeric vector</td>
</tr>
<tr>
<td>CAKI.1</td>
<td>a numeric vector</td>
</tr>
</tbody>
</table>

Source
Modified from NCBI GEO dataset: GSE32474.
Usage

RCircos.Heatmap.Plot(heatmap.data=NULL, data.col=NULL, track.num=NULL, side=c("in", "out"), min.value=NULL, max.value=NULL, inside.pos=NULL, outside.pos=NULL, genomic.columns=3, is.sorted=TRUE)

Arguments

heatmap.data A data frame with the first three columns as chromosome names, start position and end position of each data point followed by data values for one or more samples.
data.col Non-negative integer, representing the ordinal number of the column in input data set that contains the data to be plotted.
track.num Non-negative Integer, representing the ordinal number of the plot track where the heatmap will be plotted.
side Character vector, either "in" or "out", representing the position related to chromosome ideogram.
min.value Numeric, minimum value for heatmap scale.
max.value Numeric, maximum value for heatmap scale
inside.pos Non-negative numeric, inside position (relative to the centre of plot area) of the track.
outside.pos Non-negative numeric, outside position (relative to the centre of plot area) of the track.
genomic.columns Non-negative integer, total number of columns for genomic position in each row. Must be either 3 or 2.
is.sorted Logic, whether the data is sorted by chromosome names and start positions.

Author(s)

Henry Zhang

Examples

```r
## Not run:
library(RCircos)
data(UCSC.HG19.Human.CytoBandIdeogram)
data(RCircos.Heatmap.Data)

RCircos.Set.Core.Components(
cyto.info=UCSC.HG19.Human.CytoBandIdeogram,
chr.exclude=c("chrX", "chrY"),
tracks.inside=10, tracks.outside=0)
RCircos.Set.Plot.Area()
RCircos.Chromosome.Ideogram.Plot()

RCircos.Heatmap.Plot(RCircos.Heatmap.Data, data.col=5,
```

**RCircos.heatmapColors**

```
track.num=4, side="in")
RCircos.Heatmap.Plot(RCircos.Heatmap.Data, data.col=5,
side="in", inside.pos=0.9, outside.pos=1.2)
## End(Not run)
```

---

**RCircos.heatmapColors  Heatmap Colors Supported by Current Version of RCircos**

**Description**

Default heatmap colors (total of 6) supported by current version of RCircos. This is a read only data and can be viewed with the method of `RCircos.Get.Supported.HeatmapColors()`.

**Format**

The format is: `chr [1:6] "BlueWhiteRed" "GreenWhiteRed" "GreenYellowRed" "GreenBlackRed"
"YellowToRed" "BlackOnly"`

---

**RCircos.Histogram.Data  Sample Data for RCircos Histogram Plot**

**Description**

A dataset with the first three columns as chromosome names, start position, and end position followed by a column of histogram data.

**Usage**

```
data(RCircos.Histogram.Data)
```

**Format**

A data frame with 324 observations on the following 4 variables.

- **Chromosome** a factor with levels `chr1 chr10 chr11 chr12 chr13 chr14 chr15 chr16 chr17 chr18
  chr19 chr2 chr20 chr21 chr22 chr3 chr4 chr5 chr6 chr7 chr8 chr9 chrX chrY
- **chromStart** a numeric vector
- **chromEnd** a numeric vector
- **Data** a numeric vector

**Source**

Unpublished data.


**RCircos.Histogram.Plot**

*Plot Histogram for One Data Track*

**Description**

Draw one track of histogram. RCircos core components and graphic device must be initialized before drawing.

**Usage**

```r
circos.histogram.plot(hist.data=NULL, data.col=4,
                      track.num=NULL, side=c("in", "out"),
                      min.value=NULL, max.value=NULL, inside.pos=NULL,
                      outside.pos=NULL, genomic.columns=3, is.sorted=TRUE)
```

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>hist.data</code></td>
<td>A data frame with the first three columns as chromosome names, start position and end position of each data point followed by data values for one or more samples.</td>
</tr>
<tr>
<td><code>data.col</code></td>
<td>Non-negative integer, representing the ordinal number of the column in input data that contains the data to be plotted.</td>
</tr>
<tr>
<td><code>track.num</code></td>
<td>Non-negative integer, representing the ordinal number of the plot track where the histogram will be plotted.</td>
</tr>
<tr>
<td><code>side</code></td>
<td>Character vector, either &quot;in&quot; or &quot;out&quot;, representing the position related to chromosome ideogram.</td>
</tr>
<tr>
<td><code>min.value</code></td>
<td>Numeric, minimum value for histogram</td>
</tr>
<tr>
<td><code>max.value</code></td>
<td>Numeric, maximum value for histogram</td>
</tr>
<tr>
<td><code>inside.pos</code></td>
<td>Non-negative numeric, inside position (relative to the centre of plot area) of the track.</td>
</tr>
<tr>
<td><code>outside.pos</code></td>
<td>Non-negative numeric, outside position (relative to the centre of plot area) of the track.</td>
</tr>
<tr>
<td><code>genomic.columns</code></td>
<td>Non-negative integer, total number of columns for genomic position in each row. Must be either 3 or 2.</td>
</tr>
<tr>
<td><code>is.sorted</code></td>
<td>Logic, whether the data is sorted by chromosome names and start positions.</td>
</tr>
</tbody>
</table>

**Author(s)**

Henry Zhang
Examples

```r
## Not run:
library(RCircos);
data(UCSC.HG19.Human.CytoBandIdeogram);
data(RCircos.Histogram.Data);
RCircos.Set.Core.Components(
cyto.info=UCSC.HG19.Human.CytoBandIdeogram,
chr.exclude=c("chrX", "chrY"),
tracks.inside=10, tracks.outside=5)
RCircos.Set.Plot.Area()
RCircos.Chromosome.Ideogram.Plot()

RCircos.Histogram.Plot(RCircos.Histogram.Data,
data.col=4, track.num=1, side="in")
RCircos.Histogram.Plot(RCircos.Histogram.Data,
data.col=4, side="in", inside.pos=1.7,
outside.pos=1.8)

## End(Not run)
```

RCircos.Initialize.Plot.Parameters

Initialize RCircos Plot Parameters

Description

Setup default values for RCircos plot parameters. All parameters are held in one list. This function is for internal use only.

Usage

```r
RCircos.Initialize.Plot.Parameters(tracks.inside, tracks.outside)
```

Arguments

- `tracks.inside`  Non-negative integer, how many tracks will be put inside of chromosome ideogram.
- `tracks.outside` Non-negative integer, how many tracks will be put outside of chromosome ideogram.

Author(s)

Henry Zhang
RCircos.Line.Data  Sample Data for RCircos Line Plot

Description
A data frame with the first three columns as chromosome name, start and end position followed by columns of DNA copy number variant data (num.mark and seg.mean).

Usage
data(RCircos.Line.Data)

Format
A data frame with 2037 observations on the following 5 variables.

- chromosome: a factor with levels `Q QP QQ QR QS QT QU QV QW QX QY R RP RQ RR S T U V W X Y x y`
- start: a numeric vector
- stop: a numeric vector
- num.mark: a numeric vector
- seg.mean: a numeric vector

Source
Unpublished data.

RCircos.Line.Plot  Draw Lines for One Data Track

Description
Draw one track of continue lines which connect each neighbor data points. RCircos core components and graphics device must be initialized before drawing.

Usage
RCircos.Line.Plot(line.data=NULL, data.col=4, track.num=NULL, side=c("in", "out"), min.value=NULL, max.value=NULL, inside.pos=NULL, outside.pos=NULL, genomic.columns=3, is.sorted=TRUE)
Arguments

- **line.data**: A data frame with the first three columns for chromosome names, start and end position of each data point followed by data values for one or more samples.
- **data.col**: Non-negative integer, representing the ordinal number of the column in input dataset that contains the data to be plotted.
- **track.num**: Non-negative integer, representing the ordinal number of the plot track where the lines will be plotted.
- **side**: Character vector, either "in" or "out", representing the position related to chromosome ideogram.
- **min.value**: Numeric, minimum value of line data.
- **max.value**: Numeric, maximum value of line data.
- **inside.pos**: Non-negative numeric, inside position (relative to the centre of plot area) of the track.
- **outside.pos**: Non-negative numeric, outside position (relative to the centre of plot area) of the track.
- **genomic.columns**: Non-negative integer, total number of columns for genomic position in each row. Must be either 3 or 2.
- **is.sorted**: Logic, whether the data is sorted by chromosome names and start positions.

Author(s)

Henry Zhang

Examples

```r
## Not run:
library(RCircos);
data(UCSC.HG19.Human.CytoBandIdeogram);
data(RCircos.Line.Data);

RCircos.Set.Core.Components(
    cyto.info=UCSC.HG19.Human.CytoBandIdeogram,
    chr.exclude=c("chrX", "chrY"),
    num.inside=10, num.outside=0)

RCircos.Set.Plot.Area()
RCircos.Chromosome.Ideogram.Plot()

RCircos.Line.Plot(RCircos.Line.Data, data.col=5,
    track.num=1, side="in")
RCircos.Line.Plot(RCircos.Line.Data, data.col=5,
    side="in", inside.pos=1.5, outside.pos=1.75)

## End(Not run)
```
RCircos.Link.Data  Sample Data for RCircos Link Plot

Description
A data frame containing paired genomic position data for each row. The content of each row must be in the order of name, start and end position of chromosome A then name, start and end position of chromosome B. This data frame is directly used to draw link lines without extra processing.

Usage
data(RCircos.Link.Data)

Format
A data frame with 71 observations on the following 6 variables.

Chromosome  a factor with levels chr1 chr10 chr11 chr12 chr14 chr15 chr17 chr19 chr2 chr21 chr22 chr3 chr4 chr5 chr6 chr7 chr8 chr9 chrX
chromStart  a numeric vector
chromEnd    a numeric vector
Chromosome.1 a factor with levels chr1 chr10 chr11 chr12 chr14 chr15 chr16 chr17 chr18 chr19 chr2 chr20 chr21 chr22 chr3 chr4 chr5 chr6 chr7 chr8 chr9 chrX
chromStart.1 a numeric vector
chromEnd.1  a numeric vector

Source
Unpublished data.

RCircos.Link.Line  Calculate x and y Coordinates for a Bezire Curve

Description
Calculate a quadratic Bezire curve between two points with (0,0) as control points. This function is for internal use.

Usage
RCircos.Link.Line(line.start=NULL, line.end=NULL)
RCircos.Link.Plot

Arguments

- `line.start`: The x and y coordinates of a point where the Bezier curve starts.
- `line.end`: The x and y coordinates of a point where the Bezier curve ends.

Author(s)

Hongen Zhang

---

Draw Link Lines between Two or More Genomic Positions

Description

Draw lines (quadratic Bezier curves) between paired genomic positions. RCircos core components and graphic device must be initialize before drawing. Link lines are always in the center of plot area.

Usage

```r
RCircos.Link.Plot(link.data=NULL, track.num=NULL, 
by.chromosome=FALSE, start.pos=NULL, genomic.columns=3, 
is.sorted=TRUE, linewidth=rep(1, nrow(link.data)))
```

Arguments

- `link.data`: A data frame with paired genomic positions in each row. The two sets of genomic positions must have same number of columns, e.g., chromosome names, start and end positions or only chromosome names and start positions.
- `track.num`: Non-negative integer, representing the ordinal number of the plot track where the link lines will start.
- `by.chromosome`: Logic. If true, red color will be used for links between positions of same chromosomes and blue color for links between different chromosomes. Otherwise, user defined or random colors (rainbow) will be automatically assigned for each line.
- `start.pos`: Non-negative numeric, scale factor relative to chromosome ideogram position. Must be smaller than 1.
- `genomic.columns`: Non-negative integer, total number of columns for genomic position in each row. Must be either 3 or 2.
- `is.sorted`: Logic, whether the data is sorted by chromosome names and start positions.
- `linewidth`: Non-negative integer vector, width for each link line.

Author(s)

Henry Zhang
Examples

```r
## Not run:
library(RCircos);
data(UCSC.HG19.Human.CytoBandIdeogram);
data(RCircos.Link.Data);

RCircos.Set.Core.Components(
cyto.info=UCSC.HG19.Human.CytoBandIdeogram,
chr.exclude=c("chrX", "chrY"),
um.n.inside=10, num.n.outside=0)

RCircos.Set.Plot.Area()
RCircos.Chromosome.Ideogram.Plot()
RCircos.Link.Plot(RCircos.Link.Data, track.num=4,
by.chromosome=TRUE)
RCircos.Link.Plot(RCircos.Link.Data, start.pos=0.75,
by.chromosome=TRUE)

## End(Not run)
```

---

**RCircos.List.Plot.Parameters**

*List RCircos Plot Parameters*

Description

Print out onto screen all plot parameters used by current RCircos session. RCircos core components must be already initialized.

Usage

```r
RCircos.List.Plot.Parameters()
```

Author(s)

Henry Zhang

Examples

```r
## Not run:
library(RCircos);
data(UCSC.HG19.Human.CytoBandIdeogram);

RCircos.Set.Core.Components(
cyto.info=UCSC.HG19.Human.CytoBandIdeogram,
chr.exclude=c("chrX", "chrY"),
tracks.inside=10, tracks.outside=0) RCircos.List.Plot.Parameters()

## End(Not run)
```
Sample Data of Mouse Gene Expression

Description

A data frame containing mouse genomic position information and gene expression values.

Usage

data(RCircos.Mouse.Expr.Data)

Format

A data frame with 16499 observations on the following 5 variables.

- chromosome: a factor with levels chr1 chr10 chr11 chr12 chr13 chr14 chr15 chr16 chr17 chr18 chr19 chr2 chr3 chr4 chr5 chr6 chr7 chr8 chr9 chrX chrY
- txStart: a numeric vector
- txEnd: a numeric vector
- Gene: a character vector
- Expr.Mean: a numeric vector

Source

Modified from GEO dataset: GSE42081.

Setup RCircos Core Components with Multiple Species Genomes

Description

Setup RCircos core components with chromosome ideograms of multiple species for visualization of similarity and difference of genomic data between different species.

Usage

RCircos.Multiple.Species.Core.Components(cyto.info.list, species.list, chr.exclude= NULL, tracks.inside, tracks.outside)
Arguments

cyto.info.list A List contains data frames for chromosome ideogram data of different species. All data frames in the list must have same columns.

species.list Character vector represent multiple species. Each one will be used as prefix of chromosome names for relavant species. The order of species must match the order of the species in cyto.info.list.

chr.exclude Character vector of chromosome names to be excluded from plot, e.g., chr.exclude <- c("chrX", "chrY");

tracks.inside Integer, total number of data tracks inside of chromosome ideogram.

tracks.outside Integer, total number of data tracks outside of chromosome ideogram.

Author(s)

Hongen Zhang

Examples

## Not run:
library(RCircos)
data(UCSC.Mouse.GRCm38.CytoBandIdeogram)
data(UCSC.Baylor.3.4.Rat.cytoBandIdeogram)

cyto.list <- list(UCSC.Mouse.GRCm38.CytoBandIdeogram, 
                  UCSC.Baylor.3.4.Rat.cytoBandIdeogram);
species <- c("M", "R")

RCircos.Multiple.Species.Core.Components(
  cyto.list, species, chr.exclude=typeof, 
  tracks.inside=10, tracks.outside=0)

## End(Not run)

RCircos.Multiple.Species.Dataset

Combine Multiple Data Sets form Different Genomes

Description

Combine and modify the chromosome names in multiple species datasets to match the chromosomes in multiple species cytoband data.

Usage

RCircos.Multiple.Species.Dataset(data.list, species)
Arguments

data.list  List of genomic data from multiple species.
species  Character vector for prefix of chromosome names to identify different species. The order of species in species list must match the species in data.list.

Value

A data frame same as the input data list but with modified chromosome names.

Author(s)

Henry Zhang

Examples

```r
## Not run:
library(RCircos)
data(RCircos.Mouse.Expr.Data)
data(RCircos.Rat.Expr.Data)
dataSets <- list(RCircos.Mouse.Expr.Data, RCircos.Rat_EXPR.Data)
dataset <- RCircos.Get.Multiple.Species.Dataset(dataSets,
  species.list=c("M", "L"))
```

RCircos.paddingConst  Padding Distance Constant between Two Chromosomes

Description

Default length of padding area between two chromosomes on chromosome ideogram track. This is a read only data and can be viewed with RCircos.Get.Padding.Constant() method.

Format

The format is: num 0.00291
RCircos.Parallel.Line.Plot

*Draw Lines between Two Genomic Positions on Same Chromosome*

**Description**

Plot link lines inside a data track between two genomic positions on same chromosome. RCircos core components and graphic device must be initialized first.

**Usage**

```r
RCircos.Parallel.Line.Plot(line.data=NULL, track.num=NULL, 
side=c("in", "out"), line.width=1, inside.pos=NULL, 
outside.pos=NULL, genomic.columns=3, is.sorted=TRUE)
```

**Arguments**

- `line.data`: A data frame containing chromosome, start and end position for each line.
- `track.num`: Non-negative integer, the number of track to plot
- `side`: Character vector, relative location to ideogram, either "in" or "out".
- `line.width`: Non-negative numeric, same as lwd in graphic package
- `inside.pos`: Non-negative numeric, inside position (relative to the centre of plot area) of the track.
- `outside.pos`: Non-negative numeric, outside position (relative to the centre of plot area) of the track.
- `genomic.columns`: Non-negative integer, total number of columns for genomic position in each row. Must be 3.
- `is.sorted`: Logic, whether the data is sorted by chromosome names and start positions.

**Author(s)**

Henry Zhang

**Examples**

```r
## Not run:
library(RCircos);
data(UCSC.HG19.Human.CytoBandIdeogram);
data(RCircos.Line.Data);
RCircos.Set.Core.Components(
   cyto.info=UCSC.HG19.Human.CytoBandIdeogram,
   chr.exclude=c("chrX", "chrY"),
   tracks.inside=10, tracks.outside=5)
RCircos.Set.Plot.Area();
```
Description

Plot a color scale showing the color ranges of heatmap. One or more tracks should be defined for plots in outside of chromosome ideogram.

Usage

```
RCircos.Plot.Heatmap.Color.Scale(max.value=NULL, min.value=NULL, 
    color.type="BlueWhiteRed", scale.location=1, 
    scale.width=0, scale.height=0)
```

Arguments

- **max.value**: Numeric, maximum value of heatmap scale
- **min.value**: Numeric, minimum value of heatmap scale
- **color.type**: Character vector for color specification, either "BlueWhiteRed", "GreenWhiteRed", "GreenYellowRed", "GreenBlackRed", "YellowToRed", or "BlackOnly". Default is "BlueWhiteRed".
- **scale.location**: Non negative integer representing color scale location, 1~4 for bottom, left, top, and right.
- **scale.width**: Non-negative numeric, width of color scale. if not defined, 1/2 of x axis will be used
- **scale.height**: Non-negative numeric, height of color scale. if not defined, 1/10 of scaleWidth axis will be used

Author(s)

Henry Zhang
Examples

## Not run:
```r
plot.new()
plot.window(c(-5, 5), c(-5, 5))
RCircos.Plot.Heatmap.Color.Scale(max.value=3, min.value=-3,
    scaleLocation=1, colorType="BlueWhiteRed",
    scaleWidth=0, scaleHeight=0)
```

## End(Not run)

---

**RCircos.plotTypes**  
*Plot Types Supported by Current RCircos Version*

### Description

All supported plot types saved in RCircos Environment. The values are read only through RCircos.Get.Supported.Plot.Types().

### Format


---

**RCircos.Point.Plot**  
*Point Plot for One Data Track*

### Description

Plot points on a track without track outlines and sub-track lines. The size and location of each point can be adjusted with the data values. RCircos core components and graphic device must be initialized first.

### Usage

```r
RCircos.Point.Plot(point.data=NULL, data.col=4,
    track.num=NULL, side=c("in", "out"),
    min.value=NULL, max.value=NULL,
    point.type=19, with.height=TRUE,
    with.size=FALSE, point.scale=1,
    inside.pos=NULL, outside.pos=NULL,
    genomic.columns=3, is.sorted=TRUE)
```
Arguments

point.data A data frame with leading columns of chromosome names, start position and/or end position of each data point followed by data values for one or more samples.
data.col Non-negative integer, representing the ordinal number of the column in input dataset that contains the data to be plotted.
track.num Non-negative integer, representing the ordinal number of the plot track where the lines will be plotted.
side Character vector, either "in" or "out", representing the position related to chromosome ideogram.
min.value Numeric, minimum value of point data.
max.value Numeric, maximum value of point data.
point.type Non-negative integer for pch. Default is 19.
with.height Logic, if TRUE, the location of each point will be adjusted according to the data value.
with.size Logic, if TRUE, the size of each point will be adjusted according to the data value.
point.scale Non-negative numeric, more scale for point size, must be greater than or equal to 1.
inside.pos Non-negative numeric, inside position (relative to the centre of plot area) of the track.
outside.pos Non-negative numeric, outside position (relative to the centre of plot area) of the track.
genomic.columns Non-negative integer, total number of columns for genomic position in each row. Must be either 3 or 2.
is.sorted Logic, whether the data is sorted by chromosome names and start positions.

Author(s)

Henry Zhang

Examples

## Not run:
library(RCircos)
data(UCSC.HG19.Human.CytoBandIdeogram)

RCircos.Set.Core.Components(
cyto.info=UCSC.HG19.Human.CytoBandIdeogram,
chr.exclude=c("chrX", "chrY"),
um.inside=10, num.outside=0)

RCircos.Set.Plot.Area()
RCircos.Chromosome.Ideogram.Plot()
data(RCircos.Scatter.Data)
RCircos.Scatter.Plot(RCircos.Scatter.Data, 
data.col=5, track.num=4, side="in", 
with.height=TRUE, with.size=FALSE)

## End(Not run)

RCircos.Polygon.Plot

**RCircos Polygon Demo Data**

**Description**

A data frame with four columns for chromosome names, start and end position for each genomic interval, and values for polygon heights.

**Usage**

data("RCircos.Polygon.Data")

**Format**

A data frame with 77 observations on the following 4 variables.

- **chromosome**: a factor with levels chr1 chr10 chr11 chr12 chr13 chr14 chr15 chr16 chr17 chr18 chr19 chr2 chr20 chr21 chr3 chr4 chr5 chr6 chr7 chr8 chr9 chrX chrY.
- **chromStart**: a numeric vector
- **chromEnd**: a numeric vector
- **Data**: a numeric vector

**RCircos.Polygon.Plot**

**Draw Polygons on One Data Track**

**Description**

Plot polygons with different height and different locations inside of a track. Polygon plot is an alternative bar plot that takes both positive and negative height values and genomic intervals with different lengths. Plot data should have genomic positions (chromosome names, start and end positions) as well as height values. Optional column for polygon colors may follow.

**Usage**

RCircos.Polygon.Plot(polygon.data=NULL, data.col=NULL, 
track.num=NULL, side=c("in", "out"), border.col=NULL, 
polygon.col="pink", min.value=NULL, max.value=NULL, 
inside.pos=NULL, outside.pos=NULL, genomic.columns=3, 
is.sorted = TRUE)
Arguments

- **polygon.data**: A data frame with three columns for genomic positions, one or columns for polygon heights, and an optional column for polygon colour.
- **data.col**: Non-negative integer, representing the ordinal number of the column in input data set that contains the data to be plotted.
- **track.num**: Non-negative integer, representing the ordinal number of the plot track where the lines will be plotted.
- **side**: Character vector, either "in" or "out", representing the position related to chromosome ideogram.
- **border.col**: Vector of color names for border color, default null.
- **polygon.col**: Color name for fill of polygon.
- **min.value**: Numeric, minimum value in data column of polygon data.
- **max.value**: Numeric, maximum value in data column of polygon data.
- **inside.pos**: Non-negative numeric, inside position (relative to the centre of plot area) of the track.
- **outside.pos**: Non-negative numeric, outside position (relative to the centre of plot area) of the track.
- **genomic.columns**: Non-negative integer, total number of columns for genomic position in each row. Must be either 3 or 2.
- **is.sorted**: Logic, whether the data is sorted by chromosome names and start positions.

Author(s)

Henry Zhang

Examples

```r
## Not run:
library(RCircos);
data(UCSC.HG19.Human.CytoBandIdeogram);
data(RCircos.Polygon.Data);
RCircos.Set.Core.Components(
  cyto.info=UCSC.HG19.Human.CytoBandIdeogram,
  chr.exclude=c("chrX", "chrY"),
  tracks.inside=10, tracks.outside=5)
RCircos.Set.Plot.Area();
RCircos.Chromosome.Ideogram.Plot()
RCircos.Polygon.Plot(RCircos.Polygon.Data,
  track.num=1, side="in")
## End(Not run)
```
RCircos.Pseudo.Ideogram.From.Labels

Generate A Pseudo Chromosome Ideogram from List of Labels

Description

Generate a pseudo ideogram from a character list for RCircos plot. Each chromosome will have the same length and no stain colors.

Usage

```r
RCircos.Pseudo.Ideogram.From.Labels(chromosomes=NULL)
```

Arguments

- `chromosomes` - A vector of labels (such as gene names) used for chromosome names and each element must be unique.

Value

A data frame as an ideogram data table.

Author(s)

Henry Zhang

Examples

```r
## Not run:
library(RCircos)
geneNames <- paste0("Gene", 1:20)

## End(Not run)
```

RCircos.Pseudo.Ideogram.From.Table

Generate Pseudo Chromosome Ideogram from Plot Data Table

Description

Generate a pseudo ideogram from plot data for Circos plot. Plot data must have at least two columns serving as chromosome names and band names. An optional gene length or locations can also be provided, e.g.,

```
Groups LocusName LocusPosition Group1 1 1.5 Group1 3 2.0 Group1 4 3.0
```

```r
```
Usage

RCircos.Pseudo.Ideogram.From.Table(plot.data=NULL, location.col=NULL, band.col=NULL)

Arguments

plot.data  A data frame with two or three columns, the first column is for chromosome names.
location.col  Non negative integer, column number in plot.data from which end positions will be derived for each chromosome or each band. This argument must be defined and could be either length of the locus or relative locus position.
band.col  Non negative integer, column number in plot.data which serves as band names, this argument is optional.

Value

A data frame with ideogram data.

Author(s)

Henry Zhang

Examples

## Not run:
library(RCircos)
data(RCircos.Gene.Label.Data);

## End(Not run)

---

RCircos.Rat.Expr.Data  Sample Data of Rat Gene Expression

Description

A data frame containing rat genes, genomic position and expression values of these genes.

Usage

data(RCircos.Rat.Expr.Data)
Format

A data frame with 11426 observations on the following 5 variables.

- **Chromosome** a factor with levels `chr1 chr10 chr11 chr12 chr13 chr14 chr15 chr16 chr17 chr18 chr19 chr2 chr20 chr3 chr4 chr5 chr6 chr7 chr8 chr9 chrX chrQ chrQP chrQQ chrQR chrQS chrQT chrQU chrQV chrQW chrQX
- **txStart** a numeric vector
- **txEnd** a numeric vector
- **Gene** a character vector
- **Expr.Mean** a numeric vector

Source

Modified from GEO dataset: GSE42081.

---

**RCircos.Reset.Core.Components**

*Reset Methods for Modifying RCircos Core Components*

---

Description

Reset RCircos core components: plot ideogram, plot position, and plot parameters.

Usage

```r
RCircos.Reset.Plot.Ideogram(chrom.ideo)
RCircos.Reset.Plot.Positions(plot.positions)
RCircos.Reset.Plot.Parameters(new.params)
```

Arguments

- **chrom.ideo** Data frame, object of RCircos cytoband data returned from `RCircos.Get.Plot.Ideogram()`.
- **plot.positions** Data frame, object of RCircos plot positions returned from `RCircos.Get.Plot.Positions()`.
- **new.params** List, object of RCircos plot positions returned from `RCircos.Get.Plot.Parameters()`

Author(s)

Henry Zhang
Examples

```r
## Not run:
library(RCircos);
data(UCSC.HG19.Human.CytoBandIdeogram);

RCircos.Set.Core.Components(
  cyto.info=UCSC.HG19.Human.CytoBandIdeogram,
  chr.exclude=c("chrX", "chrY"),
  tracks.inside=10, tracks.outside=5)

ideo <- RCircos.Get.Plot.Ideogram()
rows <- which(ideo$BandColor="red"
ideo$BandColor[rows] <- "green"
RCircos.Reset.Plot.Ideogram(ideo)

params <- RCircos.Get.Plot.Parameters()
params$base.per.unit <- 6000
RCircos.Reset.Plot.Parameters(params)

positions[1, 3] <- 0;
RCircos.Reset.Plot.Positions(positions);

## End(Not run)
```

---

**Sample Data for RCircos Ribbon Plot**

### Description

A data frame containing paired genomic position data for each row. The contents of each row are in the order of name, start and end positions of chromosome A then name, start and end positions of chromosome B. This data frame is directly used to draw ribbons without extra processing.

### Usage

```r
data(RCircos.Ribbon.Data)
```

### Format

A data frame with 4 observations on the following 6 variables.

- **chromA** a factor with levels chr1 chr5 chr8
- **chromStartA** a numeric vector
- **chromEndA** a numeric vector
- **chromB** a factor with levels chr10 chr13 chr17 chr18
- **chromStartB** a numeric vector
- **chromEndB** a numeric vector
Source

Unpublished data.

---

**RCircos.Ribbon.Plot**  
*Draw Ribbon between Two Genomic Regions*

### Description

Draw various ribbons instead of lines with uniform thickness. The thickness of ribbons are relative to the size of the genomic regions to be linked.

### Usage

```r
RCircos.Ribbon.Plot(ribbon.data=NULL, track.num=NULL,  
                    by.chromosome=FALSE, twist=FALSE, start.pos=NULL,  
                    genomic.columns=3, is.sorted=TRUE)
```

### Arguments

- **ribbon.data**
  - A data frame with paired genomic positions in each row. The first three columns of each row are for chromosome name, start and end positions of one genomic position followed by three columns for chromosome name, start and end positions of the second genomic position.

- **track.num**
  - Non negative-integer, representing the ordinal number of the plot track where the ribbons will start.

- **by.chromosome**
  - Logic, If true, red color will be used for ribbons between positions of same chromosomes and blue color for ribbons between different chromosomes. Otherwise, different colors will be automatically assigned for each ribbon.

- **twist**
  - Logic, TRUE for the ribbons with twisted shape and FALSE for no twist.

- **start.pos**
  - Non-negative numeric, scale factor relative to chromosome ideogram position. Must be smaller than 1.

- **genomic.columns**
  - Non-negative integer, total number of columns for genomic position in each row. Must be either 3 or 2.

- **is.sorted**
  - Logic, whether the data is sorted by chromosome names and start positions.

### Author(s)

Henry Zhang
Examples

```r
## Not run:
library(RCircos)
data(UCSC.HG19.Human.CytoBandIdeogram);
data(RCircos.Link.Data);
data(RCircos.Ribbon.Data);

cyto.info <- UCSC.HG19.Human.CytoBandIdeogram
chr.exclude <- c("chrX", "chrY")
RCircos.Set.Core.Components(
  cyto.info=UCSC.HG19.Human.CytoBandIdeogram,
  chr.exclude=c("chrX", "chrY"),
  tracks.inside=10, tracks.outside=0)
RCircos.Set.Plot.Area()
RCircos.Chromosome.Ideogram.Plot()

RCircos.Link.Plot(RCircos.Link.Data, track.num=4, twist=TRUE)
RCircos.Ribbon.Plot(RCircos.Ribbon.Data,
  track.num=4, twist=TRUE, by.chromosome=FALSE)

## End(Not run)
```

---

**RCircos.Scatter.Data**  
Sample Data for RCircos Scatter Plot

**Description**

A data frame with the first three columns as chromosome name, start and end position followed by columns of DNA copy number variant data (num.mark and seg.mean).

**Usage**

```r
data(RCircos.Scatter.Data)
```

**Format**

A data frame with 1757 observations on the following 5 variables.

- **chromosome**  
  a factor with levels 1 10 11 12 13 14 15 16 17 18 19 2 20 21 22 3 4 5 6 7 8 9 X Y
- **start**  
  a numeric vector
- **stop**  
  a numeric vector
- **num.mark**  
  a numeric vector
- **seg.mean**  
  a numeric vector
**Scater Plot for One Data Track**

**Description**

Draw one track of scatter plot. RCircos core components and graphics device must be initialized before drawing.

**Usage**

```r
RCircos.Scatter.Plot(scatter.data=NULL, data.col=4, track.num=NULL, side=c("in", "out"), by.fold=0, min.value=NULL, max.value=NULL, inside.pos=NULL, outside.pos=NULL, genomic.columns=3, is.sorted=TRUE)
```

**Arguments**

- `scatter.data`: A data frame with leading columns of chromosome names, start position and/or end position of each data point followed by data values for one or more samples.
- `data.col`: Non-negative integer, representing the ordinal number of the column in input dataset that contains the data to be plotted.
- `track.num`: Non-negative integer, representing the ordinal number of the plot track where the lines will be plotted.
- `side`: Character vector, either "in" or "out", representing the position related to chromosome ideogram.
- `by.fold`: Zero or a positive float number, if it is greater than zero, any data point with a value greater than or equal to the by.fold will be plot as red color and data points with values less or equal to the negative by.fold will be plot as blue color. Otherwise, black color is used.
- `min.value`: Numeric, minimum value of scatter plot data.
- `max.value`: Numeric, maximum value of scatter plot data.
- `inside.pos`: Non-negative numeric, inside position (relative to the centre of plot area) of the track.
- `outside.pos`: Non-negative numeric, outside position (relative to the centre of plot area) of the track.
- `genomic.columns`: Non-negative integer, total number of columns for genomic position in each row. Must be either 3 or 2.
- `is.sorted`: Logic, whether the data is sorted by chromosome names and start positions.

**Author(s)**

Henry Zhang
Setup Base Plot Positions

Description

Calculate x and y coordinates for a circular line that served as base plot positions to derive positions for all other plot items. Total number of coordinates are based on total chromosome units and padding. Degrees that text should be rotated at each point are also calculated. This function is for internal use.

Usage

RCircos.Set.Base.Plot.Positions(total.points=NULL)

Arguments

total.points Non-negative integer, total number of points to form a circular line.

Author(s)

Henry Zhang
RCircos.Set.Core.Components

*Initialize Core Components for RCircos Plot*

**Description**

Setup core components for RCircos Plot including plot parameters, cytoband data, and base plot positions. This function must be called before chromosome ideogram and any other data plot.

**Usage**

```r
RCircos.Set.Core.Components(cyto.info, chr.exclude = NULL, tracks.inside, tracks.outside)
```

**Arguments**

- `cyto.info`: A data frame with chromosome ideogram data and columns are ordered by chromosome names, start position, end position, band names, and stain intensity for each chromosome band.
- `chr.exclude`: Character vector of chromosome names to be excluded from plot, e.g., `chr.exclude <- c("chrX", "chrY");`
- `tracks.inside`: Non-negative integer, total number of data tracks inside of chromosome ideogram to be plotted.
- `tracks.outside`: Non-negative integer, total number of data tracks outside of chromosome ideogram to be plotted.

**Author(s)**

Hongen Zhang

**Examples**

```r
## Not run:
library(RCircos)
data(UCSC.HG19.Human.CytoBandIdeogram)
RCircos.Set.Core.Components(
cyto.info=UCSC.HG19.Human.CytoBandIdeogram,
chr.exclude=c("chrX", "chrY"),
tracks.inside=10, tracks.outside=0)
## End(Not run)
```
**RCircos.Set.Cytoband.Data**  

*Set Chromosome Cytoband Data for Circos Plot*

**Description**

Set chromosome cytoband data (one of three RCircos core components) for Circos plot. The cytoband data will be put into RCircos Environment. This function is for internal use and users may not directly call this function and use get and reset methods instead when need to modify ideogram data.

**Usage**

```r
RCircos.Set.Cytoband.Data(cyto.band.info=NULL)
```

**Arguments**

- `cyto.band.info` A data frame containing the chromosome ideogram data returned from RCircos.Validate.Cyto.Info() or a sorted ideogram table with correct chromosome names, start and end positions, band names, and staining status.

**Author(s)**

Henry Zhang

**RCircos.Set Plot Area**  

*Setup Plot Area for RCircos Plot*

**Description**

Open a new window for plotting. RCircos core components must be initialized before call this function. All four margins are set to 0.25 and window size are set to plot.radius in RCircos parameters. This function is optional and could be replaced by call `par()`, `plot.new()`, and `plot.window()` if users want control the plot window by themselves.

**Usage**

```r
RCircos.Set.Plot.Area(margins = 0.25, ...)
```

**Arguments**

- `margins` Non-negative numeric, value for margins around plot area.
- `...` Other arguments passed...
**Author(s)**

Henry Zhang

**Examples**

```r
## Not run:
library(RCircos);
data(UCSC.HG19.Human.CytoBandIdeogram);
RCircos.Set.Core.Components(
cyto.info=UCSC.HG19.Human.CytoBandIdeogram,
chr.exclude=c("chrX", "chrY"),
tracks.inside=10, tracks.outside=5)
RCircos.Set.Plot.Area()

## End(Not run)
```

---

**RCircos.Set.Zoom.Plot.Positions**

*Set Position for Zoom-in Plot*

**Description**

Set plot position for zooming-in plot by selecting index of default plot positions. The zoomed plot will be aligned with the original plot position such as gene position or a cytoband. The zoom area was defined as fraction of circle circumference, default is 1/8.

**Usage**

```r
RCircos.Set.Zoom.Plot.Positions(zoom.info=NULL, total.genes=11,
area.length=0.25, fixed.width=FALSE, gene.width=NULL)
```

**Arguments**

- `zoom.info` A vector with chromosome name, start and end position to be zoomed.
- `total.genes` Non-negative integer from 1 - maximum number of genes to be plotted. The maximum number of genes is calculated based on track height and will be adjusted inside the function.
- `area.length` Non-negative numeric, for better layout, it must be smaller than or equal to 1/4 (quarter of the circumference).
- `fixed.width` Logic, if the plot width for each data point is fixed or not.
- `gene.width` Non-negative integer, number of units for gene width.

**Value**

Numeric vector for index of RCircos.Base.Position
RCircos.Sort.Genomic.Data

Author(s)

Henry Zhang

Examples

```r
## Not run:
library(RCircos)
data(UCSC.HG19.Human.CytoBandIdeogram);

RCircos.Set.Core.Components(
  cyto.info=UCSC.HG19.Human.CytoBandIdeogram,
  chr.exclude=c("chrX", "chrY"),
  tracks.inside=10, tracks.outside=0)
zoom.info <- c("chr1", 10000, 50000)

## End(Not run)
```

RCircos.Sort.Genomic.Data

Sort Genomic or Ideogram Data

Description

Sort genomics/ideogram data. The order of chromosome names should be numeric names (integers or Roman numbers) first then character names. If chromosome names are all characters alphabets order will be used. This function could be used before making RCircos plot.

Usage

```r
RCircos.Sort.Genomic.Data(genomic.data=NULL, is.ideo=FALSE)
```

Arguments

- `genomic.data`: A data frame with the first two or three columns for chromosome names, start and end positions. If it is ideogram data, next two columns must be band names, and Giemsa stain status.
- `is.ideo`: Logic, whether the genomic data is ideogram or not.

Value

Data frame same as input data but ordered by chromosome names then start positions

Author(s)

Henry
Examples

```r
## Not run:
library(RCircos)

data(UCSC.Mouse.GRCm38.CytoBandIdeogram)
cyto <- UCSC.Mouse.GRCm38.CytoBandIdeogram
ideogram <- RCircos.Sort.Genomic.Data(
  genomic.data=UCSC.Mouse.GRCm38.CytoBandIdeogram,
  is.ideo=TRUE)
data(RCircos.Line.Data)
lineData <- RCircos.Sort.Genomic.Data( genomic.data=RCircos.Line.Data,
  is.ideo=FALSE)

## End(Not run)
```

**RCircos.Tile.Data**

*Sample Data for RCircos Tile Plot*

**Description**

A data frame containing genomic position data only (chromosome, start and end positions).

**Usage**

```r
data(RCircos.Tile.Data)
```

**Format**

A data frame with 152 observations on the following 3 variables.

- **chromosome** a factor with levels chr1 chr10 chr11 chr12 chr13 chr14 chr15 chr16 chr17 chr18 chr19 chr2 chr20 chr21 chr22 chr3 chr4 chr5 chr6 chr7 chr8 chr9 chrX chrY
- **chromStart** a numeric vector
- **chromEnd** a numeric vector

**Source**

Unpublished Data.
RCircos.Tile.Plot  
Plot Tiles for One Data Track

Description

Draw one track of Tiles. RCircos core components and graphics device must be initialized before drawing.

Usage

RCircos.Tile.Plot(tile.data=NULL, track.num=NULL, side= c("in", "out"), inside.pos=NULL, outside.pos=NULL, genomic.columns=3, is.sorted=TRUE)

Arguments

tile.data  A data frame with the three columns as chromosome names, start position and end position of each data point. Other column(s) is optional.
track.num  Non-negative integer, representing the ordinal number of the plot track where the lines will be plotted.
side  Character vector, either "in" or "out", representing the position related to chromosome ideogram.
inside.pos  Non-negative numeric, inside position (relative to the centre of plot area) of the track.
outside.pos  Non-negative numeric, outside position (relative to the centre of plot area) of the track.
genomic.columns  Non-negative integer. Must be 3 for tile plot.
is.sorted  Logic, whether the data is sorted by chromosome names and start positions.

Author(s)

Henry Zhang

Examples

```r
# Not run:
library(RCircos);
data(UCSC.HG19.Human.CytoBandIdeogram);

RCircos.Set.Core.Components(  
cyto.info=UCSC.HG19.Human.CytoBandIdeogram,  
chr.exclude=c("chrX", "chrY"),  
tracks.inside=10, tracks.outside=0  
)  
RCircos.Set.Plot.Area()  
RCircos.Chromosome.Ideogram.Plot()
```
RCircos.Track.Outline  Draw Outline for A Data Track

Description

Draw outline with subtracks for one data track. RCircos core components and graphic device must be initialized first. This function is mainly for internal use.

Usage

```r
RCircos.Track.Outline(inside.pos=NULL, outside.pos=NULL,
    num.layers=1, chrom.list=NULL, track.colors=NULL)
```

Arguments

- `inside.pos` Non-negative numeric, inside position (relative to the centre of plot area) of the track.
- `outside.pos` Non-negative numeric, outside position (relative to the centre of plot area) of the track.
- `num.layers` Non-negative integer, number of sub-tracks lines. 0 for no sub-track line.
- `chrom.list` List of chromosome names for which outline will be drawn.
- `track.colors` Vector of color names for outline of each chromosome.

Author(s)

Henry Zhang

Examples

```r
## Not run:
library(RCircos);
data(Ucsc.HG19.Human.Cytobandideogram);

RCircos.Set.Core.Components(
cyto.info=Ucsc.HG19.Human.Cytobandideogram,
    chr.exclude=c("chrX", "chrY"),
    tracks.inside=10, tracks.outside=5)
RCircos.Set.Area()
RCircos.Chromosome.Ideogram.Plot()

locations <- RCircos.Track.Positions(side="in", track.num=1);
out.pos <- locations[1];
```
**RCircos.Validate.Cyto.Info**

**Validate Chromosome Ideogram Information**

**Description**

Validate chromosome ideogram information for correct chromosome order including of correct order of chromosome names, chromosome start, and end positions. The prefix of "chr" in chromosome names is no longer required version 1.2.

**Usage**

```r
RCircos.Validate.Cyto.Info(cyto.info=NULL, chr.exclude=NULL, is.sorted=TRUE)
```

**Arguments**

- **cyto.info**: A data frame with chromosome ideogram data, e.g., an object returned by function of read.table() which read a file containing full information of cytobandIdeogram table from UCSC genome browser.
- **chr.exclude**: Character vector, name(s) of chromosomes to be excluded from RCircos plot.
- **is.sorted**: Logic, whether the cyto.info has been sorted.

**Value**

A data frame with validated chromosome ideogram table.

**Author(s)**

Henry Zhang

**Examples**

```r
## Not run:
library(RCircos)
data(UCSC.HG38.Human.CytobandIdeogram)
cyto.info <- RCircos.Validate.Cyto.Info(
  cyto.info=UCSC.HG38.Human.CytobandIdeogram,
  chr.exclude=NULL, is.sorted=FALSE)

## End(Not run)
```
RCircos.Validate.Genomic.Data

Validate Genomic Data

Description

Validate input dataset for correct chromosome names, chromosome start, and chromosome end positions. Chromosome names will be converted to character vectors if they are factor variables.

Usage

RCircos.Validate.Genomic.Data(genomic.data=NULL, plot.type=c("plot", "link"), genomic.columns=3)

Arguments

- genomic.data: Data frame with genomic position data.
- plot.type: Character vector, either "plot" or "link".
- genomic.columns: Non-negative integer, total number of columns for genomic position (chromosome name, start and/or end position).

Value

None. Function will exit with an error prompt.

Author(s)

Henry Zhang

Examples

```r
## Not run:
library(RCircos)
data(RCircos.Heatmap.Data)

data(RCircos.Link.Data)

## End(Not run)
```
**RCircos.Validate.Genomic.Info**

*Validate Genomic Information*

**Description**

Check out if the chromosome name, start and end positions in the range of ideogram data.

**Usage**

```r
RCircos.Validate.Genomic.Info(genomic.info=NULL)
```

**Arguments**

- `genomic.info`: A vector with a chromosome name, start and end position on the chromosome

**Author(s)**

Henry Zhang

**Examples**

```r
## Not run:
library(RCircos)
data(UCSC.HG38.Human.CytoBandIdeogram)

RCircos.Set.Core.Components(UCSC.HG38.Human.CytoBandIdeogram,
    chr.exclude=NULL, tracks.inside=10, tracks.outside=0 )
RCircos.Validate.Genomic.Info(c("chr1", 10000, 500000))

## End(Not run)
```

---

**RCircos.Validate.Plot.Parameters**

*Validate Plot Parameters*

**Description**

Validate values of plot parameters before reset them in RCircos Environment. This function is mainly for internal use.

**Usage**

```r
RCircos.Validate.Plot.Parameters(parameters=NULL)
```
**Arguments**


**Author(s)**

Henry Zhang

---

**RCircos.Validate.Track.Positions**

*Validate Track Positions*

---

**Description**

Validate if user defined plot locations (inside.pos and outside.pos) are inside of plot area and do not overlap with chromosome ideogram. RCircos core components must be initialized first. This function is mainly for internal use.

**Usage**

```r
circos.Validate.Track.Positions(inside.pos=0, outside.pos=0, erase.area=FALSE)
```

**Arguments**

- `inside.pos`: Non-negative numeric, inside position (relative to the centre of plot area) of the track.
- `outside.pos`: Non-negative numeric, outside position (relative to the centre of plot area) of the track.
- `erase.area`: Logic, if the track is going to be erased.

**Value**

A numeric vector with length of 2 for the real plot position of a customized track.

**Author(s)**

Henry Zhang
RCircos.Vertical.Line.Plot

**Draw Vertical Lines on One Data Track**

**Description**

Plot vertical lines on a data track for each genomic position without track outlines and sub-track lines. RCircos core components and graphic devices must be initialized first.

**Usage**

```r
RCircos.Vertical.Line.Plot(line.data=NULL, track.num=NULL, side=c("in", "out"), line.width=1, inside.pos=NULL, outside.pos=NULL, genomic.columns=3, is.sorted=TRUE)
```

**Arguments**

- **line.data** A data frame with leading columns as chromosome names, start position and/or end position of each data point followed by data values for one or more samples.
- **track.num** Non-negative integer, representing the ordinal number of the plot track where the lines will be plotted.
- **side** Character vector, either "in" or "out", representing the position related to chromosome ideogram.
- **line.width** Non-negative integer, width of vertical lines
- **inside.pos** Non-negative numeric, inside position (relative to the centre of plot area) of the track.
- **outside.pos** Non-negative numeric, outside position (relative to the centre of plot area) of the track.
- **genomic.columns** Non-negative integer, total number of columns for genomic position in each row. Must be either 3 or 2.
- **is.sorted** Logic, whether the data is sorted by chromosome names and start positions.

**Author(s)**

Henry Zhang

**Examples**

```r
## Not run:
library(RCircos);
data(UCSC.HG19.Human.CytoBandIdeogram);
data(RCircos.Line.Data);

RCircos.Set.Core.Components(
    cyto.info=UCSC.HG19.Human.CytoBandIdeogram,
```
RCircos.Zoom.Area.Outline

A Simple RCircos Plot Workflow

Description
Print out on screen to list procedures for a basic RCircos plot.

Usage
RCircos.Workflow()

Author(s)
Henry Zhang

Examples
## Not run:
library(RCircos)
RCircos.Workflow()

## End(Not run)

---

RCircos.Zoom.Area.Outline

Draw Outline for Zoom Plot Area

Description
Draw outline and subtrack lines for zoomed-in area of a track in outside of chromosome ideogram. RCircos core components must be initialized first. This function is mainly for internal use.

Usage
RCircos.Zoom.Area.Outline(zoom.pos=NULL, inside.pos=NULL, outside.pos=NULL, num.layers=5, fill.col="white")
Arguments

- **zoom.pos**: Non-negative numeric vector, the index of RCircos plot position.
- **inside.pos**: Non-negative numeric, inside position (relative to the centre of plot area) of the track.
- **outside.pos**: Non-negative numeric, outside position (relative to the centre of plot area) of the track.
- **num.layers**: Non-negative integer, number of sub-tracks in a data track.
- **fill.col**: Character vector for color name to fill the polygon.

Author(s)

Henry Zhang

Examples

```r
## Not run:
library(RCircos);
data(UCSC.HG19.Human.CytoBandIdeogram)

RCircos.Set.Core.Components(
cyto.info=UCSC.HG19.Human.CytoBandIdeogram,
chr.exclude=c("chrX", "chrY"),
tracks.inside=10, tracks.outside=5)
RCircos.Set.Plot.Area()
RCircos.Chromosome.Ideogram.Plot()

data(RCircos.Heatmap.Data)
name.col=4, genomic.columns=3,
target.gene="SP5", neighbor.genes=5)
zoom.range <- RCircos.Get.Zoom.Range(zoom.data, 3)
zoom.pos <- RCircos.Set.Zoom.Plot.Positions(zoom.range,
total.genes=11, area.length=0.25, gene.width=NULL)

RCircos.Zoom.Area.Outline(zoom.pos=NULL, inside.pos=2.5,
outside.pos=3, num.layers=5, fill.col="white")
```

Description

Scale up the index of Circos plot positions of a small range for zoom-in plot with data that require two positions to plot e.g., tiles and parallele lines. This function is mainly for internal use.
Usage


Arguments

zoom.data  Data frame, data to be plotted onto zoom area.
zoom.pos   Non-negative integer, index of RCircos base plot positions.

Value

A two columns integer matrix representing the index of RCircos plot positions.

Author(s)

Henry Zhang

---


*Calculate Zoomed Index of Circos Plot Positions for Single Point Genomic Data*

Description

Scale up the index of Circos plot positions of a small range for zoom-in plot with data that require only one position to plot e.g., scatter plot. This function is mainly for internal use.

Usage


Arguments

zoom.data  Data frame, data to be plotted onto zoom area.
zoom.pos   Non-negative integer, index of RCircos base plot positions.

Value

An integer representing the index of RCircos plot positions.

Author(s)

Henry Zhang
RCircos.ZoomIn.Chromosome

Zoom in One or Partial Chromosome Ideogram

Description

Increase plot length for one or partial chromosome in order to plot more data points. This works on original chromosome ideogram data and should be done before set RCircos core components. Also, this function only work with one region each time. If there are more than one regions to be zoomed in, repeat this function for each region.

Usage

RCircos.ZoomIn.Chromosome(ideogram=NULL, chromosome=NULL, from=NULL, to=NULL, zoom.in=NULL)

Arguments

ideogram Data frame, chromosome ideogram data with chromosome names, start position, end position, band name and stains for each band or each chromosome.

chromosome Character vector, name of chromosome to be zoomed in.

from Non-negative integer, genomic coordinates of start position of chromosome region to be zoomed in.

to Non-negative integer, genomic coordinates of end position of chromosome region to be zoomed in.

zoom.in Non-negative number, fold to zoom in.

Value

Data frame, ideogram with new zoomed in region.

Author(s)

Henry Zhang

Examples

```r
## Not run:
library(RCircos)
data(UCSC.HG19.Human.CytoBandIdeogram)
zoomedIdeo <- RCircos.ZoomIn.Chromosome(
   ideogram=UCSC.HG19.Human.CytoBandIdeogram, 
   chromosome="chr17", from=10000, to=20000, 
   zoom.in=1000)

## End(Not run)
```
RCircos.ZoomIn.Plot  Zoomed-in Plot of Small Genomic Region

Description

Zoom-in a small area on outside of chromosome ideogram to show the details for: heatmap, histogram, gene connector, gene labels, continue lines, vertical lines, parallel lines, scatters(points), areas, and tiles.

Usage

```r
RCircos.Plot.Zoomed.Area(zoom.data=NULL,
    plot.type="mountain", data.col=NULL, track.num=NULL,
    zoom.pos=NULL, min.value=NULL, max.value=NULL,
    area.color="gray", border.col="black", inside.pos=NULL,
    outside.pos=NULL, outline=TRUE)
```

```r
RCircos.Plot.Zoomed.Gene.Connectors(zoom.data=NULL,
    track.num=NULL, zoom.pos=NULL, line.width=1,
    inside.pos=NULL, outside.pos=NULL)
```

```r
RCircos.Plot.Zoomed.Heatmap(zoom.data=NULL, data.col=NULL,
    track.num=NULL, zoom.pos=NULL, min.value=NULL,
    max.value=NULL, inside.pos=NULL, outside.pos=NULL)
```

```r
RCircos.Plot.Zoomed.Histogram(zoom.data=NULL, data.col=NULL,
    track.num=NULL, zoom.pos=NULL, min.value=NULL,
    max.value=NULL, inside.pos=NULL, outside.pos=NULL,
    outline=TRUE)
```

```r
RCircos.Plot.Zoomed.Ideogram.Ticks(zoom.info=NULL,
    track.num=NULL, zoom.pos=NULL, tick.interval=5,
    inside.pos=NULL, outside.pos=NULL)
```

```r
RCircos.Plot.Zoomed.Continue.Lines(zoom.data=NULL,
    data.col=NULL, track.num=NULL, zoom.pos=NULL,
    line.width=1, min.value=NULL, max.value=NULL,
    inside.pos=NULL, outside.pos=NULL, outline=TRUE)
```

```r
RCircos.Plot.Zoomed.Parallel.Lines(zoom.data=NULL,
    track.num=NULL, zoom.pos=NULL, genomic.cols=3,
    line.width=NULL,inside.pos=NULL, outside.pos=NULL,
    outline=FALSE)
```

```r
RCircos.Plot.Zoomed.Vertical.Lines(zoom.data=NULL,
    track.num=NULL, zoom.pos=NULL, line.width=1,
    inside.pos=NULL, outside.pos=NULL, outline=FALSE)
```
RCircos.ZoomIn.Plot

```r
RCircos.Plot.Zoomed.Scatters(zoom.data=NULL, data.col=NULL,
track.num=NULL, zoom.pos=NULL, min.value=NULL,
max.value=NULL, point.type=16, by.fold=0,
with.size=TRUE, with.height=FALSE, point.scale=1,
inside.pos=NULL, outside.pos=NULL, outline=TRUE)

RCircos.Plot.Zoomed.Tiles(zoom.data=NULL, track.num=NULL,
zoom.pos=NULL, genomic.cols=3, layers=5, border.col=NULL,
inside.pos=NULL, outside.pos=NULL, outline=TRUE)

RCircos.Plot.Zoomed.Polygons(zoom.data=NULL, data.col=4,
track.num=NULL, zoom.pos=NULL, genomic.cols=3,
min.value=NULL, max.value=NULL, border.col=NULL,
inside.pos=NULL, outside.pos=NULL, outline=TRUE)

RCircos.Label.Zoom.Region(zoom.data=NULL, name.col=NULL,
track.num=NULL, zoom.pos=NULL, text.size=0.75,
inside.pos=NULL, outside.pos=NULL)

RCircos.Mark.Zoom.Area(zoom.range=NULL, track.num=1,
zoom.pos=NULL, fill.color="yellow",
inside.pos=NULL, outside.pos=NULL)
```

**Arguments**

- `zoom.data`  A data frame containing genomic positions, gene names, and plot values for zoom-in genes/rows.
- `zoom.range`  Character vector, chromosome name, start and end positions of a genomic interval to be zoomed in.
- `data.col`  Non-negative integer/vector, which column(s) is plot data values.
- `track.num`  Non-negative integer, which track will be plotted.
- `zoom.pos`  Non-negative numeric vector, the index of RCircos plot position.
- `min.value`  Numeric, minimum value for heatmap scale.
- `max.value`  Numeric, maximum value for heatmap scale.
- `inside.pos`  Non-negative numeric, inside position (relative to the centre of plot area) of the track.
- `outside.pos`  Non-negative numeric, outside position (relative to the centre of plot area) of the track.
- `name.col`  Non-negative integer, which column is for labels.
- `genomic.cols`  Non-negative integer, total number of columns for genomic position information (chromosome, start and/or end position).
- `fill.color`  Character vector for name of a color to fill the polygon.
- `border.col`  Character vector for name of a color to draw the polygon border.
- `area.color`  Character vector for name of a color to fill the area.
text.size  Non-negative numeric, character size for text plot.
line.width  Non-negative numeric, width of lines.
point.type  Non-negative integer, type of character for point plot (Default 19).
plot.type   Character vector, either "mountain", "curtain", or "band", types of area plot.
by.fold     Zero or a positive float number, if it is greater than zero, any data point with a value greater than or equal to the by.fold will be plot as red color and data points with values less or equal to the negative by.fold will be plot as blue colr. Otherwise, black color is used.
with.size   Logic. If TRUE, the point size will match to the data value in scatter/point plot.
with.height Logic. If TRUE, point location will match to the data value in scatter/point plot.
point.scale Non-negative numeric, more scale to add for scatter/point size in scatter/point plot.
outline     Logic. If TRUE, outline will be drawn for zoomed area.
zoom.info   Character vector contains chromosome name, start and end position.
tick.interval Non-negative integer, length of genomic interval in million base pairs.
layers      Non-negative integer, number of subtrack in the data track.

Details
RCircos.Plot.Zoomed.Area() paints partial area inside the zoomed region (an irregular polygon ship). Plot types include "mountain", "curtain" (reversed mountain), and "band".
RCircos.Plot.Zoomed.Vertical.Lines() plot zoomed-in vertical lines from bottom to top of a track for small number of genomic intervals (default 11).
RCircos.Label.Zoom.Region() plot gene names for small number of genes (default 11).
RCircos.Mark.Zoom.Area() highlight the area between zoomed-in plot area and chromosome ideogram to mark the original genomic interval.
**RCircos.ZoomOut.Chromosome**

**Author(s)**

Henry Zhang

**Examples**

```r
## Not run:
library(RCircos);
data(UCSC.HG19.Human.CytoBandIdeogram)

cyto <- UCSC.HG19.Human.CytoBandIdeogram
RCircos.Set.Core.Components(cyto, NULL, 10, 5)

RCircos.Set.Plot.Area()
RCircos.Chromosome.Ideogram.Plot()

data(RCircos.Heatmap.Data)
min.value <- min(as.matrix(RCircos.Heatmap.Data[,5:10]))
max.value <- max(as.matrix(RCircos.Heatmap.Data[,5:10]))

name.col=4, genomic.columns=3,
target.gene="SP5", neighbor.genes=5)
zoom.range <- RCircos.Get.Zoom.Range(zoom.data, 3)
zoom.pos <- RCircos.Set.Zoom.Plot.Positions(zoom.range,
total.genes=11, area.length=0.25, gene.width=NULL)

RCircos.Plot.Zoomed.Heatmap(zoom.data, data.col=5,
track.num=2, zoom.pos=zoom.pos,
min.value=min.value, max.value=max.value,
inside.pos=NULL, outside.pos=NULL)

RCircos.Mark.Zoom.Area(zoom.pos, zoom.data, color="red",
track.num=2, inside.pos=NULL, outside.pos=NULL)

RCircos.Label.Zoom.Region(zoom.data, name.col=4,
track.num=3, zoom.pos=zoom.pos, text.size=0.75,
inside.pos=NULL, outside.pos=NULL)

## End(Not run)
```

**Description**

Zoom out chromosome ideogram to leave a small room between the first and the last chromosome (at 12 O’clock) so that track names could be added. This function works on the RCircos chromosome ideogram component so the core components must be set first.
Usage
RCircos.ZoomOut.Chromosome(zoom.out.ratio=NULL)

Arguments

zoom.out.ratio  A float number between 0 and 1, the ratio of target chromosome ideogram to be zoomed out.

Value
None. New ideogram object is set to RCircos environment.

Author(s)
Henry Zhang

Examples
```r
## Not run:
library(RCircos)
data(UCSC.HG19.Human.CytoBandIdeogram)
RCircos.Set.Core.Components(  
  cyto.info=UCSC.HG19.Human.CytoBandIdeogram,  
  chr.exclude=c("chrX", "chrY"),  
  tracks.inside=10, tracks.outside=0) RCircos.ZoomOut.Chromosome(zoom.out.ratio=0.95)
## End(Not run)
```

---

UCSC.Baylor.3.4.Rat.cytoBandIdeogram

*Cytoband Information for Rat Chromosome Ideogram*

Description
A data frame containing chromosome name, start and end position, band name, and intensity of Giemsa stains for each cytoband.

Usage
data(UCSC.Baylor.3.4.Rat.cytoBandIdeogram)

Format
A data frame with 246 observations on the following 5 variables.

- **Chromosome**  a factor with levels chr1 chr10 chr11 chr12 chr13 chr14 chr15 chr16 chr17 chr18 chr19 chr2 chr20 chr3 chr4 chr5 chr6 chr7 chr8 chr9 chrX
- **ChromStart**  a numeric vector
ChromEnd a numeric vector

Band a factor with levels p11 p12 p13 p14 p15 p16 p21 p22 q11 q12 q12.1 q12.2 q12.3 q12.4 q12.5 q13 q14 q15 q16 q21 q22 q23 q24 q25 q26 q27 q31 q32 q32.1 q32.2 q32.3 q33 q34 q35 q36 q37 q38 q41 q42 q43 q44 q45 q51 q52 q53 q54 q55

Stain a factor with levels gneg gpos gvar

Source

http://genome.ucsc.edu

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UCSC.HG19.Human.CytoBandIdeogram

**Cytoband Information for Human Chromosome Ideogram**

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**Description**

A data frame containing chromosome name, start and end position, band name, and intensity of Giemsa stains for each cytoband.

**Usage**

data(UCSC.HG19.Human.CytoBandIdeogram)

**Format**

A data frame with 862 observations on the following 5 variables.

Chromosome a factor with levels chr1 chr10 chr11 chr12 chr13 chr14 chr15 chr16 chr17 chr18 chr19 chr2 chr20 chr21 chr22 chr3 chr4 chr5 chr6 chr7 chr8 chr9 chrX chrY

ChromStart a numeric vector

ChromEnd a numeric vector

Stain: a factor with levels acen gneg gpos gposQPP gposRU gposUP gposWU gvar stalk

Source
http://genome.ucsc.edu

### Format
A data frame with 862 observations on the following 5 variables.

- **Chromosome**: a factor with levels chr1 chr10 chr11 chr12 chr13 chr14 chr15 chr16 chr17 chr18 chr19 chr20 chr21 chr22 chr3 chr4 chr5 chr6 chr7 chr8 chr9 chrX chrY
- **chromStart**: a numeric vector
- **chromEnd**: a numeric vector
- **Name**: a factor with levels p11 p11.1 p11.11 p11.12 p11.2 p11.21 p11.22 p11.23 p11.3 p11.31 p11.32 p11.4 p12.1 p12.2 p12.3 p12.31 p12.32 p12.33 p13.1 p13.11 p13.12 p13.13 p13.2 p13.3 p13.31 p13.32 p13.33 p14.1 p14.2 p14.3 p15.1 p15.2 p15.3 p15.31 p15.32 p15.33 p15.4 p15.5 p16.1 p16.2 p16.3 p21.1 p21.2 p21.3 p21.31 p21.32 p21.33 p22.1 p22.11 p22.12 p22.13 p22.2 p22.21 p22.22 p22.23 p22.24 p22.3 p22.31 p22.32 p22.33 p23.1 p23.2 p23.3 p24.1 p24.2 p24.3 p25.1 p25.2 p25.3 p26.1 p26.2 p26.3 p27.1 p27.2 p27.3 p27.4 p27.5 p27.6 p28.1 p28.2 p28.3 p28.4 p28.5 p28.6 p28.7 p28.8 p28.9 p29.1 p29.2 p29.3 p29.4 p29.5 p29.6 p29.7 p29.8 p29.9 p30.1 p30.2 p30.3 p30.4 p30.5 p30.6 p30.7 p30.8 p30.9 p31.1 p31.2 p31.3 p31.4 p31.5 p31.6 p31.7 p31.8 p31.9 p32.1 p32.2 p32.3 p32.4 p32.5 p32.6 p32.7 p32.8 p32.9 p33.1 p33.2 p33.3 p33.4 p33.5 p33.6 p33.7 p33.8 p33.9 p34.1 p34.2 p34.3 p34.4 p34.5 p34.6 p34.7 p34.8 p34.9 p35.1 p35.2 p35.3 p35.4 p35.5 p35.6 p35.7 p35.8 p35.9 p36.1 p36.2 p36.3 p36.4 p36.5 p36.6 p36.7 p36.8 p36.9 p37.1 p37.2 p37.3 p37.4 p37.5 p37.6 p37.7 p37.8 p37.9 p38.1 p38.2 p38.3 p38.4 p38.5 p38.6 p38.7 p38.8 p38.9 p39.1 p39.2 p39.3 p39.4 p39.5 p39.6 p39.7 p39.8 p39.9 p40.1 p40.2 p40.3 p40.4 p40.5 p40.6 p40.7 p40.8 p40.9 p41.1 p41.2 p41.3 p41.4 p41.5 p41.6 p41.7 p41.8 p41.9 p42.1 p42.2 p42.3 p42.4 p42.5 p42.6 p42.7 p42.8 p42.9 p43.1 p43.2 p43.3 p43.4 p43.5 p43.6 p43.7 p43.8 p43.9 p44.1 p44.2 p44.3 p44.4
UCSC.Mouse.GRCm38.CytoBandIdeogram

Stain a factor with levels acen gneg gpos100 gpos25 gpos50 gpos75 gvar stalk

Source

http://genome.ucsc.edu

UCSC.Mouse.GRCm38.CytoBandIdeogram

Cytoband Information for Mouse Chromosome Ideogram

Description

A data frame containing chromosome name, start and end position, band name, and intensity of Giemsa stains for each cytoband.

Usage

data(UCSC.Mouse.GRCm38.CytoBandIdeogram)

Format

A data frame with 403 observations on the following 5 variables.

Chromosome a factor with levels chr1 chr10 chr11 chr12 chr13 chr14 chr15 chr16 chr17 chr18 chr19 chr2 chr3 chr4 chr5 chr6 chr7 chr8 chr9 chrX chrY

ChromStart a numeric vector

ChromEnd a numeric vector

Band a factor with levels qa qa1.1 qa1.2 qa1.3 qa2 qa3 qa3.1 qa3.2 qa3.3 qa4 qa5 qa5.1 qa5.2 qa5.3 qa6 qa7.1 qa7.2 qa7.3 qb qb1 qb1.1 qb1.2 qb1.3 qb2 qb2.1 qb2.2 qb2.3 qb3 qb3.1 qb3.2 qb3.3 qb4 qb5 qb5.1 qb5.2 qb5.3 qc qc1 qc1.1 qc1.2 qc1.3 qc2 qc3 qc3.1 qc3.2 qc3.3 qc4 qc5 qc6 qc7 qd qd1 qd2 qd2.1 qd2.2 qd2.3 qd3 qe qe1 qe1.1 qe1.2 qe1.3 qe2 qe2.1 qe2.2 qe2.3 qe3 qe3.1 qe3.2 qe3.3 qe4 qe5 qf qf1 qf2 qf2.1 qf2.2 qf2.3 qf3 qf4 qf5 qg1 qg1.1 qg1.2 qg1.3 qg2 qg3 qh qh1 qh2 qh2.1 qh2.2 qh2.3 qh3 qh4 qh5 qh6

Stain a factor with levels acen gneg gpos100 gpos33 gpos66 gpos75

Source

http://genome.ucsc.edu
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