# An introduction of making genomic plots with circlize

Zuguang Gu <z.gu@dkfz.de>
April 23, 2016

Since circos plots are mostly used in genomic research, the **circlize** package particularly provides functions which focus on genomic plots. These functions are synonymous to the basic circos graphic functions but expect special format of input data:

- circos.genomicTrackPlotRegion: create a new track and add graphics.
- circos.genomicPoints: low-level function, add points
- circos.genomicLines: low-level function, add lines
- circos.genomicRect: low-level function, add rectangles
- circos.genomicText: low-level function, add text
- circos.genomicLink: add links

The genomic functions are implemented by basic circos functions (e.g. circos.trackPlotRegion, circos.points), thus, you can customize your own plots by both genomic functions and basic circos functions.

# 1 Input data

Genomic circos functions expect input data as a data frame or a list of data frames in which there are at least three columns. The first column is genomic category (e.g. chromosome), the second column is the start positions in the genomic category and the third column is the end positions. Following columns are optional where numeric values or other related values are stored. Such data structure is known as *BED* format and is broadly used in genomic research.

circlize provides a simple function generateRandomBed which can generate random genomic data. Positions are uniformly generated from human genome and the number of regions on chromosomes approximately proportional to the length of chromosomes. In the function, nr and nc are number of rows and numeric columns that users want. Please note nr are not exactly the same as the number of rows which are returned by the function. fun is a self-defined function to generate random values.

```
library(circlize)
set.seed(999)
bed = generateRandomBed()
head(bed)
##
      chr start
                    end
                             value1
## 1 chr1 39485 159163 -0.1887635
## 2 chr1 897195 1041959 -0.2435220
## 3 chr1 1161957 1177159 0.3749953
## 4 chr1 1201513 1481406 -0.2600839
## 5 chr1 1487402 1531773 -0.4633990
## 6 chr1 1769949 2736215 -0.8159909
bed = generateRandomBed(nr = 200, nc = 4)
nrow(bed)
```

```
## [1] 205
bed = generateRandomBed(nc = 2, fun = function(k) runif(k))
head(bed)

## chr start end value1 value2
## 1 chr1 98740 566688 0.18303240 0.1725320
## 2 chr1 769960 887938 0.04453375 0.6378507
## 3 chr1 906851 933021 0.80237105 0.5433349
## 4 chr1 1241911 1243537 0.41103676 0.2260829
## 5 chr1 1385344 1410947 0.83473785 0.9218159
## 6 chr1 1498302 1585389 0.77238964 0.8390309
```

# 2 Initialize the layout

#### 2.1 Initialize with cytoband data

#### 2.1.1 Basic usage

Similar as general circos plots, the first step is to initialize the plot with genomic categories. In most situations, genomic categories are measured by chromosomes. The easiest way is to used circos.initializeWithIdeogram (figure 1 A):

```
par(mar = c(1, 1, 1, 1))
circos.initializeWithIdeogram()
```

By default, the function will initialize the plot with cytoband data of hg19. You can also use your own cytoband data by specifying the path of your cytoband file (no matter compressed or not) or providing your cytoband data as a data frame. An example for cytoband file is http://hgdownload.soe.ucsc.edu/goldenpath/hg19/database/cytoBand.txt.gz.

If you want to read cytoband data from file, please explicitly specify colClasses arguments and set the class of position columns as numeric. The reason is since positions are represented as integers, read.table would treat those numbers as integer by default. In initialization of circos plot, circlize needs to calculate the summation of all chromosome lengths. The sumation of such large integers would throw error of data overflow.

For simple use, users can also specify abbreviation of the species and the function will download cytoband file from UCSC server automatically (If it exists in UCSC. As you can guess, the URL template we use is http://hgdownload.soe.ucsc.edu/goldenpath/\$species/database/cytoBand.txt.gz).

```
circos.initializeWithIdeogram(species = "hg18")
circos.initializeWithIdeogram(species = "mm10")
```

By default, the function will use all chromosomes which are available in cytoband data to initialize the circos plot. Users can also choose a subset of chromosomes by specifying chromosome.index. This argument is also for ordering chromosomes (figure 1 B).

```
circos.initializeWithIdeogram(chromosome.index = paste0("chr", 10:1))
```

#### 2.1.2 Order chromosomes

Initialization step is important for circos plot. It controls orders of chromosomes which are going to be put on the circle. There are several ways to control the order:

- If chromosome.index is set, the order of chromosome.index is taken as order of chromosomes.
- If chromosome.index is not set and sort.chr is set to TRUE, chromosomes will be sorted first by numbers then by letters (figure 1 C).
- If chromosome.index and sort.chr are not set neither:
  - If cytoband is provided as a data frame, and if the first column is not a factor, the order of chromosomes would be unique(cytoband[[1]]) (figure 1 D).
  - If cytoband is provided as a data frame, and if the first column is a factor, the order of chromosome would be levels(cytoband[[1]]) (figure 1 E).
  - If cytoband is specified as a file path, or species is specified, the order of chromosomes depends on the original order in the source file.

```
cytoband = cytoband.df
circos.initializeWithIdeogram(cytoband, sort.chr = TRUE)
cytoband = cytoband.df
circos.initializeWithIdeogram(cytoband, sort.chr = FALSE)
cytoband[[1]] = factor(cytoband[[1]], levels = pasteO("chr", c(22:1, "X", "Y")))
circos.initializeWithIdeogram(cytoband, sort.chr = FALSE)
```

circlize provides a function read.cytoband which can read/download and process cytoband data. In fact, circos.initializeWithIdeogram calls read.cytoband internally (actually, if there is no cytoband available, read.chromInfo will be called later). Please refer to the help page of the function for more details.

```
cytoband = read.cytoband()
cytoband = read.cytoband(file)
cytoband = read.cytoband(df)
cytoband = read.cytoband(species)
```

#### 2.1.3 Pre-defined tracks

After the initialization of the circos plot, the function will additionally create a track where there are genomic axes and chromosome names, and create another track where there is an ideogram (depends on whether cytoband data is available). plotType can be used to control which kind of graphics need to be added (figure 1 F, G).

```
circos.initializeWithIdeogram(plotType = c("axis", "labels"))
circos.initializeWithIdeogram(plotType = NULL)
circos.clear()
```

#### 2.1.4 Other general settings

Similar as general circos plot, the layout of circos plot can be controlled by circos.par (figure 1 H, I).

```
circos.par("start.degree" = 90)
circos.initializeWithIdeogram()
circos.clear()
circos.par("gap.degree" = rep(c(2, 4), 12))
circos.initializeWithIdeogram()
circos.clear()
```

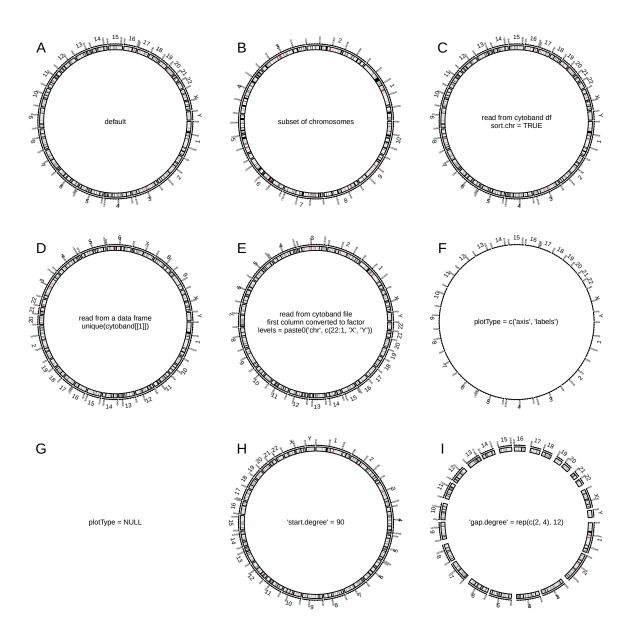


Figure 1: Different ways to initialize genomic circos plot. A) default; B) select subset of chromosomes; C) read from a data frame; D) the first column of the data frame is a factor; E) sort chromosomes; F) do not add ideogram; G) initialize the plot while plot nothing; H) set start degree of the plot; I) set gap degree.

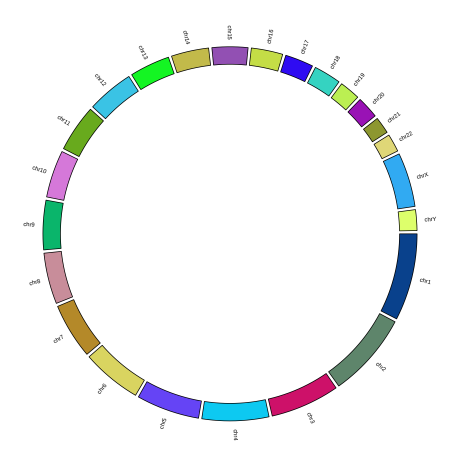


Figure 2: Customize ideogram.

## 2.2 Customize ideogram

The default behaviour of circos.initializeWithIdeogram is both initializing the layout and adding two tracks. But if plotType is set to NULL, the circular layout is only initialized but nothing is added. It provides possibility for users to completely design their own style of plots. In the following example, we use different colors to represent chromosomes and change the style of chromosome names (figure 2).

```
circos.clear()
```

## 2.3 Initialize with general genomic category

Cytoband data is just a special case of genomic category. circos.genomicInitialize can initialize circular layout with any kind of genomic categories. In fact, circos.initializeWithIdeogram is implemented by circos.genomicInitialize. The input data for the function is also a data frame with at least three columns. The first column is genomic category (for cytoband data, it is chromosome name), and the next two columns are genomic positions in each genomic category. The range of each category will be inferred from the minimum position and the maximum position in corresponding category. In the following example, a circos plot is initialized with three genes.

```
df = data.frame(
   name = c("TP53", "TP63", "TP73"),
   start = c(7565097, 189349205, 3569084),
   end = c(7590856, 189615068, 3652765))
circos.genomicInitialize(df)
```

Note it is not necessary that the record for each gene is one row.

As explained in previous section, the order of genomic categories is controlled by the first column of df which depends on whether it is a factor or a simple vector. Alternative names can be assigned to each category and the order of alternative names correspond to the order of genomic categories.

```
circos.genomicInitialize(df)
circos.genomicInitialize(df, sector.names = c("tp53", "tp63", "tp73"))
circos.genomicInitialize(df, plotType)

circos.par(gap.degree = 2)
circos.genomicInitialize(df)
```

In following example, we plot the transcripts for TP53, TP63 and TP73 in a circular layout (figure 3). The object tp\_family is a list of data frames which contain positions of exons for each transcript.

```
load(paste0(system.file(package = "circlize"), "/extdata/tp_family.RData"))
names(tp_family)
## [1] "TP73" "TP63" "TP53"
names(tp_family[["TP53"]])
   [1] "ENST00000413465.2" "ENST00000359597.4" "ENST00000504937.1"
   [4] "ENST00000269305.4" "ENST00000510385.1" "ENST00000504290.1"
  [7] "ENST00000455263.2" "ENST00000420246.2" "ENST00000445888.2"
## [10] "ENST00000576024.1" "ENST00000509690.1" "ENST00000514944.1"
## [13] "ENST00000574684.1" "ENST00000505014.1" "ENST00000508793.1"
## [16] "ENST00000604348.1" "ENST00000503591.1"
head(tp_family[["TP53"]][[1]])
           start
## exon_7 7565097 7565332
## exon_6 7577499 7577608
## exon_5 7578177 7578289
## exon_4 7578371 7578554
## exon_3 7579312 7579590
## exon_2 7579700 7579721
df = data.frame(gene = names(tp_family),
                start = sapply(tp_family, function(x) min(unlist(x))),
                end = sapply(tp_family, function(x) max(unlist(x))))
df
```

```
## gene start end

## TP73 TP73 3569084 3652765

## TP63 TP63 189349205 189615068

## TP53 TP53 7565097 7590856
```

In the following code, we first create a track which identifies three genes.

```
circos.genomicInitialize(df)
circos.genomicTrackPlotRegion(ylim = c(0, 1),
   bg.col = c("#FF000040", "#00FF0040", "#0000FF40"),
   bg.border = NA, track.height = 0.05)
```

Next, we put each transcript line by line. It is simply adding lines and rectangles. The usage of circos.genomicTrackPlotRegion will be introduced in later sections.

```
n = max(sapply(tp_family, length))
circos.genomicTrackPlotRegion(ylim = c(0.5, n + 0.5),
    panel.fun = function(region, value, ...) {
        gn = get.cell.meta.data("sector.index")
        tr = tp_family[[gn]] # all transcripts for this gene
        for(i in seq_along(tr)) {
            # for each transcript
            current_tr_start = min(tr[[i]]$start)
            current_tr_end = max(tr[[i]]$end)
            circos.lines(c(current_tr_start, current_tr_end),
                c(n - i, n - i), col = "#CCCCCC")
            circos.genomicRect(tr[[i]], ytop = n - i + 0.4,
                ybottom = n - i - 0.4, col = "orange", border = NA)
        }
}, bg.border = NA, track.height = 0.3)
circos.clear()
```

## 2.4 Zooming

circos.genomicInitialize is implemented by the general circos.initialize function, so same strategy can be applied to zoom chromosomes (figure 4). Just be careful with the order of chromosomes.

Here we can only use circos.genomicInitialize to deal with zoomed chromosomes because chromosome with name "zoom\_chr1" is not a normal chromosome.

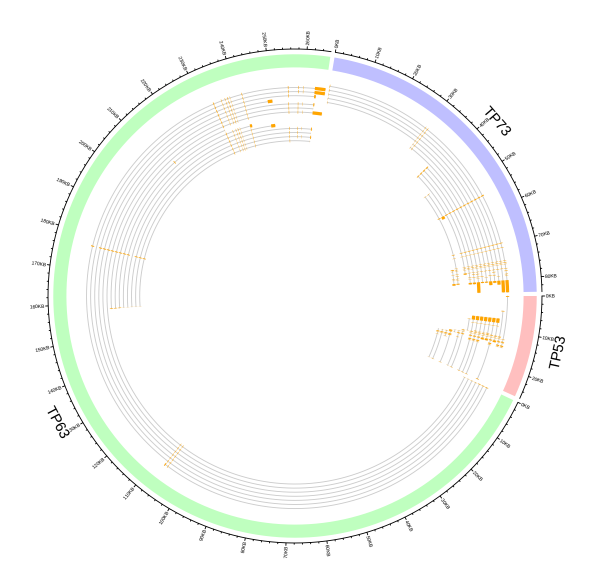


Figure 3: Alternative transcripts for genes.

```
par(mar = c(1, 1, 1, 1))
circos.par(start.degree = 90)
circos.genomicInitialize(df2, sector.width = sector.width)
```

In following tracks, data frames should also be extended with zoomed chromosomes. We can write a simple function to do the extension.

```
extend_zoomed_chromosome_in_bed = function(bed, chromosome, prefix = "zoom_") {
   zoom_bed = bed[bed[[1]] %in% chromosome, , drop = FALSE]
   zoom_bed[[1]] = pasteO(prefix, zoom_bed[[1]])
   rbind(bed, zoom_bed)
}
```

Then add a new track:

```
bed = generateRandomBed(100)
circos.genomicTrackPlotRegion(extend_zoomed_chromosome_in_bed(bed, chromosome[1:2]),
    panel.fun = function(region, value, ...) {
        circos.genomicPoints(region, value, pch = 16, cex = 0.5)
})
```

Also add link from original chromosome to the zoomed chromosome.

```
circos.link("chr1", get.cell.meta.data("cell.xlim", sector.index = "chr1"),
    "zoom_chr1", get.cell.meta.data("cell.xlim", sector.index = "zoom_chr1"),
    col = "#00000020", border = NA)
circos.clear()
```

Same strategy can be applied if you only want to zoom sub regions inside a certain chromosome (just make them as pseudo chromosomes).

# 3 Create plotting regions

In following sections, chromosome will be used as the type of genomic category. In this section, we assume data is simply a data frame. For more complex situations and behaviour of the functions, we will introduce in the next section.

Similar as circos.trackPlotRegion, circos.genomicTrackPlotRegion also accepts a self-defined function panel.fun which is applied in every cell but with different form.

```
circos.genomicTrackPlotRegion(data, panel.fun = function(region, value, ...) {
    circos.genomicPoints(region, value, ...)
})
```

Inside panel.fun, users can use low-level genomic graphic functions to add basic graphics in each cell. panel.fun expects two arguments region and value. region is a data frame containing start position and end position in the current chromosome which is extracted from data, and you can think it corresponds to values on x-axes. value is also a data frame which contains other columns in data, and you can think it corresponds to values on y-axes. Besides, there should be a third arguments ... which is mandatory and is used to pass user-invisible variables to inner functions (which we will explain in next sections).

Following codes demonstrate values for region and value when used inside panel.fun.

```
bed = generateRandomBed(nc = 1)
head(bed)

## chr start end value1
## 1 chr1 15804 90247 0.36038334
```

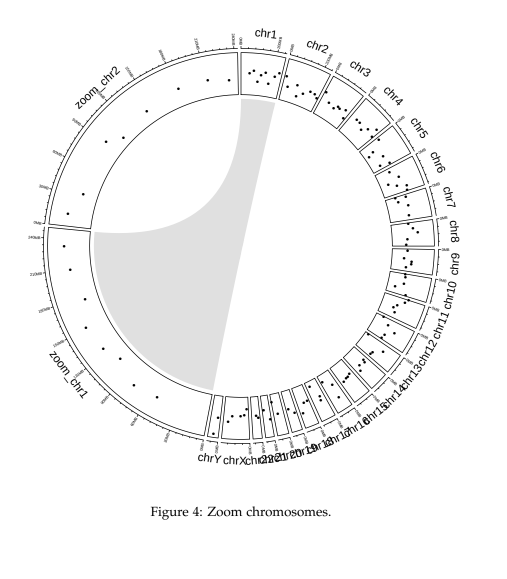


Figure 4: Zoom chromosomes.

```
## 2 chr1 122503 440777 -0.23412573
## 3 chr1 745473 1175215 -0.26368581
## 4 chr1 1662962 1781365 -0.94756937
## 5 chr1 1812844 1907062 -0.05621953
## 6 chr1 2060894 2353914 0.03344915
circos.initializeWithIdeogram(plotType = NULL)
circos.genomicTrackPlotRegion(bed, panel.fun = function(region, value, ...) {
    if(get.cell.meta.data("sector.index") == "chr1") {
        print(head(region))
        print(head(value))
    }
})
##
      start
                 end
             90247
## 1
      15804
## 2 122503 440777
## 3 745473 1175215
## 4 1662962 1781365
## 5 1812844 1907062
## 6 2060894 2353914
##
         value1
## 1 0.36038334
## 2 -0.23412573
## 3 -0.26368581
## 4 -0.94756937
## 5 -0.05621953
## 6 0.03344915
circos.clear()
```

Since circos.genomicTrackPlotRegion will create a new track, it needs values to calculate range of y-values to arrange data points. Users can either specify the index of numeric columns in data by numeric.column (named index or numeric index) or set ylim. If none of them are set, the function will try to look for all numeric columns in data (of course, excluding the first three columns), and set them as numeric.column.

```
circos.genomicTrackPlotRegion(data, ylim = c(0, 1),
    panel.fun = function(region, value, ...) {
        circos.genomicPoints(region, value, ...)
})
circos.genomicTrackPlotRegion(data, numeric.column,
    panel.fun = function(region, value, ...) {
        circos.genomicPoints(region, value, ...)
})
```

Since genomic functions are implemented by basic circos functions, you can use circos.info anywhere to get information of sectors and tracks.

## 3.1 Points

circos.genomicPoints is similar as circos.points. The difference is circos.genomicPoints expects a data frame containing genomic regions and a data frame containing values. The data column for plotting should be indicated by numeric.column. If the function is called inside circos.genomicTrackPlotRegion and users have been already set numeric.column in circos.genomicTrackPlotRegion, proper value of numeric.column will be passed to circos.genomicPoints through . . . in panel.fun. Which means, you need to add . . . as the final argument in circos.genomicPoints to get such information. If numeric.column is not set in both place, circos.genomicPoints will use all numeric columns detected in value.

Note here numeric.column is measured in value while numeric.column in circos.genomicTrackPlotRegion is measured in the complete data frame. There is a difference of 3! When numeric.column is passed to circos.genomicPoints internally, 3 is subtracted automatically. If you use character index instead of numeric index, you do not need to worry about it.

```
circos.genomicPoints(region, value, ...)
circos.genomicPoints(region, value, numeric.column = c(1, 2))
circos.genomicPoints(region, value, cex, pch)
circos.genomicPoints(region, value, sector.index, track.index)
```

If there is only one numeric column, graphical parameters such as pch, cex can be of length one or number of rows of region. If there are more than one numeric columns specified, points for each numeric column will be added iteratively, and the graphical parameters should be either length one or number of numeric columns specified.

circos.genomicPoints is implemented by circos.points. The basic idea of the implementation is like follows.

```
circos.genomicPoints = function(region, value, numeric.column = 1, ...) {
    x = (region[[2]] + region[[1]])/2
    y = value[[numeric.column]]
    circos.points(x, y, ...)
}
```

As shown above, circos.genomicPoints also provides arguments sector.index and track.index. The default values are current sector and current track. With these two arguments, you can add points by circos.genomicPoints outside circos.genomicPlotTrackRegion.

#### 3.2 Lines

circos.genomicLines is similar as circos.lines. The setting of graphical parameters is similar as circos.genomicPoints.

```
circos.genomicLines(region, value, ...)
circos.genomicLines(region, value, numeric.column = c(1, 2))
circos.genomicLines(region, value, lwd, lty = "segment")
circos.genomicLines(region, value, area, baseline, border)
circos.genomicLines(region, value, sector.index, track.index)
```

For 1ty, we additionally provide a new option segment by which each genomic interval will represent as a 'horizontal' line.

#### 3.3 Text

For circos.genomicText, the position of text can be specified either by numeric.column (index) or a separated vector y. The labels of text can be specified either by labels.column (index) or a vector labels.

```
circos.genomicText(region, value, ...)
circos.genomicText(region, value, y, labels)
circos.genomicText(region, value, numeric.column, labels.column)
circos.genomicText(region, value, facing, niceFacing, adj)
circos.genomicText(region, value, sector.index, track.index)
```

#### 3.4 Rectangle

For circos.genomicRect, the positions of top and bottom of the rectangles can be specified by ytop, ybottom or ytop.column, ybottom.column (index).

```
circos.genomicRect(region, value, ytop = 1, ybottom = 0)
circos.genomicRect(region, value, ytop.column = 2, ybottom = 0)
circos.genomicRect(region, value, col, border)
```

One of the usage of circos.genomicRect is to plot heatmap on the circle. **circlize** provides a simple function colorRamp2 to interpolate colors. The arguments of colorRamp2 are break points and colors which correspond to the break points. colorRamp2 returns a new function which can be used to generate new colors.

## 4 More details on circos.genomicTrackPlotRegion

The behaviour of circos.genomicTrackPlotRegion and panel.fun will be different according to different input data (e.g. is it a simple data frame or a list of data frames? If it is a data frame, how many numeric columns it has?) and different settings.

#### 4.1 Normal mode

#### 4.1.1 Input is a data frame

If input data is a simple data frame, region in panel.fun would be a data frame containing start position and end position in the current chromosome which is extracted from input data. value is also a data frame which contains columns in input data excluding the first three columns. Index of proper numeric columns will be passed by ... if it is set in circos.genomicTrackPlotRegion. So if users want to use such information, they need to pass ... to low-level genomic function such as circos.genoimcPoints as well.

If there are more than one numeric columns, all columns that are specified will be added to the plot.

```
bed = generateRandomBed(nc = 2)
# just note `numeric.column` is measured in `bed`
circos.genomicTrackPlotRegion(bed, numeric.column = 4,
    panel.fun = function(region, value, ...) {
        circos.genomicPoints(region, value, ...)
        circos.genomicPoints(region, value)
        # here `numeric.column` is measured in `value`
        circos.genomicPoints(region, value, numeric.column = 1)
})
```

## 4.1.2 Input is a list of data frames

If input data is a list of data frames, panel.fun is applied on each data frame iteratively. Under such situation, region and value will contain corresponding data in the current data frame. The index for the current data frame can be get by getI(...). For numeric.column argument, if input data is a list

of data frame, the length can only be one or the number of data frames, which means, there is only one numeric column that will be used in each data frame.

```
bedlist = list(generateRandomBed(), generateRandomBed())
circos.genomicTrackPlotRegion(bedlist,
    panel.fun = function(region, value, ...) {
        i = getI(...)
        circos.genomicPoints(region, value, col = i, ...)
})

# column 4 in the first bed and column 5 in the second bed
circos.genomicTrackPlotRegion(bedlist, numeric.column = c(4, 5),
    panel.fun = function(region, value, ...) {
        i = getI(...)
        circos.genomicPoints(region, value, col = i, ...)
})
```

#### 4.2 stack mode

circos.genomicTrackPlotRegion also supports a stack mode. Under stack mode, ylim is re-defined inside the function. The y-axis will be splitted into several layers with equal height and graphics are put on 'horizontal' layers (y = 1, 2, ...).

#### 4.2.1 Input is a data frame

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

```
bed = generateRandomBed(nc = 2)
circos.genomicTrackPlotRegion(bed, stack = TRUE,
    panel.fun = function(region, value, ...) {
        i = getI(...)
        circos.genomicPoints(region, value, col = i, ...)
})
```

## 4.2.2 Input is a list of data frame

When input data is a list containing data frames, each data frame will be treated as a single unit. The situation is quite similar as described previously. ylim is re-defined to c(0.5, n+0.5) in which n is number of data frames. panel.fun will be applied iteratively on each data frame. In each iteration, in panel.fun, region is still the genomic regions in current chromosome, and value contains columns in current data frame excluding the first three columns. Still, graphics by low-level genomic functions will be added on the 'horizontal' lines.

```
bedlist = list(generateRandomBed(), generateRandomBed())
circos.genomicTrackPlotRegion(bedlist, stack = TRUE,
    panel.fun = function(region, value, ...) {
        i = getI(...)
        circos.genomicPoints(region, value, ...)
})
```

Under stack mode, the differentce between using a data frame with multiple numeric columns and using a list of data frames is if using a data frame, all layers share the same genomic positions while if using a list of data frames, the genomic positions for each layer can be different.

# 5 Applications

In this section, we will show several real examples of making genomic plots under different modes.

#### 5.1 Points

To make plots more clear to look at, we only add graphics in the 1/4 of the circle and initialize the plot only with chromosome 1 (figure 5).

In track A, it is the most simple way to add points.

```
### track A
bed = generateRandomBed(nr = 300)
circos.genomicTrackPlotRegion(bed, panel.fun = function(region, value, ...) {
    circos.genomicPoints(region, value, pch = 16, cex = 0.5, ...)
})
```

In track B, under stack mode, points are added in one horizontal line (here we additionally add the dashed line) because there is only one numeric column in bed.

```
### track B
bed = generateRandomBed(nr = 300)
circos.genomicTrackPlotRegion(bed, stack = TRUE,
    panel.fun = function(region, value, ...) {
        circos.genomicPoints(region, value, pch = 16, cex = 0.5, ...)

        i = getI(...)
        cell.xlim = get.cell.meta.data("cell.xlim")
        circos.lines(cell.xlim, c(i, i), lty = 2, col = "#00000040")
})
```

In track C, the input data is a list of two data frames. In the plot, sizes of the points correspond to the values of regions, and colors correspond to different data frames.

```
### track C
bed1 = generateRandomBed(nr = 300)
bed2 = generateRandomBed(nr = 300)
bed_list = list(bed1, bed2)
circos.genomicTrackPlotRegion(bed_list,
    panel.fun = function(region, value, ...) {
        cex = (value[[1]] - min(value[[1]]))/(max(value[[1]]) - min(value[[1]]))
        i = getI(...)
        circos.genomicPoints(region, value, cex = cex, pch = 16, col = i, ...)
})
```

In track D, plot the data frame list under stack mode.

```
### track D
circos.genomicTrackPlotRegion(bed_list, stack = TRUE,
   panel.fun = function(region, value, ...) {
     cex = (value[[1]] - min(value[[1]]))/(max(value[[1]]) - min(value[[1]]))
     i = getI(...)
```

```
circos.genomicPoints(region, value, cex = cex, pch = 16, col = i, ...)
cell.xlim = get.cell.meta.data("cell.xlim")
circos.lines(cell.xlim, c(i, i), lty = 2, col = "#00000040")
})
```

In track E, the data frame has four numeric columns. Different colors are assigned to different columns. Note value is a data frame with four columns.

```
### track E
bed = generateRandomBed(nr = 300, nc = 4)
circos.genomicTrackPlotRegion(bed,
        panel.fun = function(region, value, ...) {
        circos.genomicPoints(region, value, cex = 0.5, pch = 16, col = 1:4, ...)
})
```

In track F, the data frame has four columns but is plotted under stack mode. Note here value is a data frame with only one column (but it will be exexuted 4 times in each cell).

```
### track F
bed = generateRandomBed(nr = 300, nc = 4)
circos.genomicTrackPlotRegion(bed, stack = TRUE,
    panel.fun = function(region, value, ...) {
        cex = (value[[1]] - min(value[[1]]))/(max(value[[1]]) - min(value[[1]]))
        i = getI(...)
        circos.genomicPoints(region, value, cex = cex, pch = 16, col = i, ...)

        cell.xlim = get.cell.meta.data("cell.xlim")
        circos.lines(cell.xlim, c(i, i), lty = 2, col = "#000000040")
})
circos.clear()
```

#### 5.2 Lines

Initialize the plot with one quarter of the circle (figure 6).

In track A, it is the most simple way to add lines.

```
### track A
bed = generateRandomBed(nr = 500)
circos.genomicTrackPlotRegion(bed,
    panel.fun = function(region, value, ...) {
        circos.genomicLines(region, value, type = "l", ...)
})
```

In track B, add lines for a list of data frames. Different colors refers to different data frames.

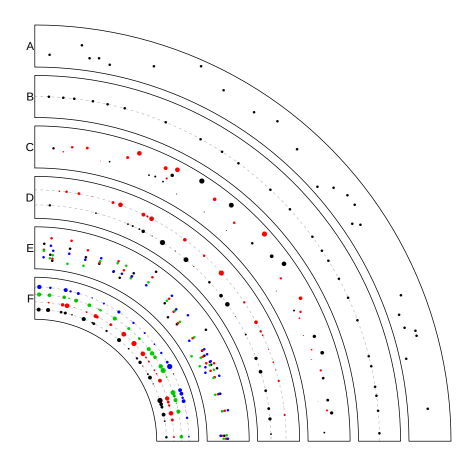


Figure 5: Add points under different modes.

```
i = getI(...)
circos.genomicLines(region, value, col = i, ...)
})
```

In track C, it is the stack mode of adding lines. Here the width of each line corresponds to the width of each genomic interval.

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

In track D, lines for the four numeric columns are added with different colors.

```
### track D
bed = generateRandomBed(nr = 500, nc = 4)
circos.genomicTrackPlotRegion(bed,
    panel.fun = function(region, value, ...) {
        circos.genomicLines(region, value, col = 1:4, ...)
})
```

In track E, it is the stack mode for a data frame with more than one numeric columns.

```
### track E
bed = generateRandomBed(nr = 500, nc = 4)
circos.genomicTrackPlotRegion(bed, stack = TRUE,
    panel.fun = function(region, value, ...) {
        i = getI(...)
        circos.genomicLines(region, value, col = i, ...)
})
```

In track F, we specify type to segment and use different colors for segments. Note each segment is located at its corresponding y-value.

## 5.3 Rectangles

Again, initialize the plot with one quarter of the circle (figure 7). Also, we want to map the values to colors by f.

In track A, bed has four numeric columns and stack mode is used to arrange the heatmap.

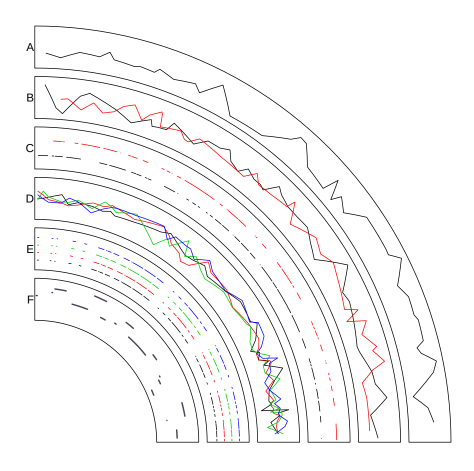


Figure 6: Add lines under different modes.

```
### track A
bed = generateRandomBed(nr = 100, nc = 4)
circos.genomicTrackPlotRegion(bed, stack = TRUE,
    panel.fun = function(region, value, ...) {
        circos.genomicRect(region, value, col = f(value[[1]]), border = NA, ...)
})
```

In track B, add rectangles for a list of data frames. Comparing to track A, here genomic regions for each data frame are different, so the positions of rectangles in the first layer are different from that in the second layer.

Under stack mode, ytop and ybottom can be set to adjust the height of rectangles. It would be straightforward because for each layer, the position is y = i and the height of each rectangle is 1.

```
### track B
bed1 = generateRandomBed(nr = 100)
bed2 = generateRandomBed(nr = 100)
bed_list = list(bed1, bed2)
circos.genomicTrackPlotRegion(bed_list, stack = TRUE,
    panel.fun = function(region, value, ...) {
        i = getI(...)
        circos.genomicRect(region, value, ytop = i + 0.4, ybottom = i - 0.4,
        col = f(value[[1]]), ...)
})
```

In track C, the plot are the same as track B, but without stack mode. Note here we explicitly specify ylim.

In track D, bars are added to the base line (y = 0).

#### 5.4 Mixed use of general circos functions

panel.fun is applied on each cell, which means, besides genomic functions, you can also use general circos functions to add more graphics. For example, some horizontal lines and texts are added to each cell and axes are put on top of each cell:

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

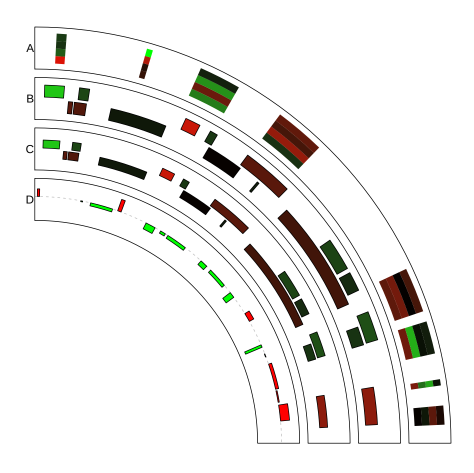


Figure 7: Add rectangles under different modes.

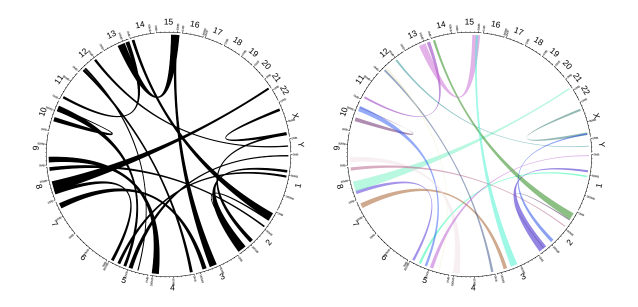


Figure 8: Add links from two sets of genomic regions.

```
circos.genomicPoints(region, value, ...)

cell.xlim = get.cell.meta.data("cell.xlim")
for(h in c(-1, -0.5, 0, 0.5, 1)) {
      circos.lines(cell.xlim, c(0, 0), lty = 2, col = "grey")
}
circos.text(x, y, labels)
circos.axis("top")
})
```

#### 6 Links

circos.genomicLink expects two data frames and it will add links from genomic intervals in the first data frame to corresponding genomic intervals in the second data frame (figure 8). You can set graphical parameters as a single scalar or a vector.

# 7 Highlight chromosomes

highlight.chromosome provides a simple way to highlight chromosomes. Just remember to use transparent filled colors. The position of the highlighted regions can be fine-tuned by padding argument which are percent of corresponding height and width in the highlighted regions.

In following examples, we first create five more tracks with random points.

```
circos.par("track.height" = 0.1)
circos.initializeWithIdeogram(plotType = c("axis", "labels"))

for(i in 1:5) {
    bed = generateRandomBed(nr = 100)
    circos.genomicTrackPlotRegion(bed, panel.fun = function(region, value, ...) {
        circos.genomicPoints(region, value, pch = 16, cex = 0.5, ...)
    })
}
```

Following are several ways of highlighting. we can highlight individual tracks by setting track.index. If two tracks or two chromosomes are neighbours, they will be taken as a union (figure 9).

circos.info would be helpful for you to find the correct track index. And if you only want to highlight some sub regions in one chromosome, use draw.sector directly.

# 8 High-level genomic functions

circlize implements several high-level functions which may help to visualize genomic data more effectively.

#### 8.1 Position transformation

There is one representative situation when genomic position transformation needs to be applied. For example, there are regions which are quite densely located on the chromosome. If heatmap or text for those dense regions are to be plotted. They would be overlapped and hard to identify, also ugly to visualize. Thus, a way to transform original positions to new positions would help for the visualization.

Low-level genomic functions such as circos.genomicPoints all accept an argument posTransform to apply user-defined position transformation. Value for posTransform is a self-defined function which only accepts at least one argument: a data frame with two columns (start position and end position). Because posTransform is bound to low-level graphic functions, it is not necessary to add chromosome information in the input data. There is only one requirement for position transformation: Number of rows of regions should be the same before and after the transformation.

## 8.1.1 The default position transformation function

In **circlize**, there already provides a position transformation function posTransform.default which distributes positions uniformly in the current chromosome.

Following code does the transformation. The points are plotted with the new transformed regions.

```
circos.genomicTrackPlotRegion(data, panel.fun = function(region, value, ...) {
    circos.genomicPoints(region, value, posTransform = posTransform.default, ...)
})
```

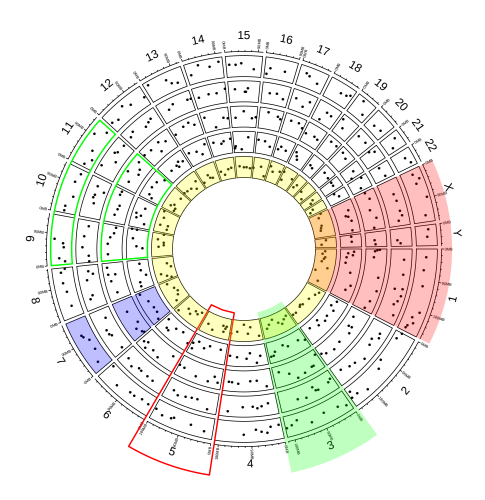


Figure 9: Highlight different chromosomes and tracks.

Of course we need identification lines to connect the untransformed regions to the transformed regions. There is a function called circos.genomicPosTransformLines which does the job. Same value for posTransform which was used before should be passed to circos.genomicPosTransformLines. Note circos.genomicPosTransformLines will create a new track to put such identification lines. In the function, direction controls whether the position transformed track is inside or outside the identification line track, and horizontalLine controls whether add lines to identify the regions.

```
circos.genomicPosTransformLines(data, posTransform = posTransform.default)
circos.genomicPosTransformLines(data, posTransform = posTransform.default,
    horizontalLine = "top")
circos.genomicPosTransformLines(data, posTransform = posTransform.default,
    direction = "outside")
```

In following example, we make heatmap for selected regions in the genome (figure 10 A).

For the second example, the heatmap is plotted outside the identification lines, so here direction is set to outside (figure 10 B). Also note how we add the ideogram inside the circle.

```
circos.par(cell.padding = c(0, 0, 0, 0))
circos.initializeWithIdeogram(plotType = NULL)
circos.genomicTrackPlotRegion(bed, stack = TRUE,
    panel.fun = function(region, value, ...) {
        circos.genomicRect(region, value, col = f(value[[1]]),
            border = f(value[[1]]), posTransform = posTransform.default, ...)
}, bg.border = NA)
circos.genomicPosTransformLines(bed, posTransform = posTransform.default,
   direction = "outside", horizontalLine = "bottom", track.height = 0.1)
cytoband = read.cytoband()$df
circos.genomicTrackPlotRegion(cytoband, stack = TRUE,
   panel.fun = function(region, value, ...) {
        circos.genomicRect(region, value, col = cytoband.col(value[[2]]), border = NA)
       xlim = get.cell.meta.data("xlim")
       ylim = get.cell.meta.data("ylim")
        circos.rect(xlim[1], ylim[1], xlim[2], ylim[2], border = "black")
}, track.height = 0.05, bg.border = NA)
circos.clear()
```

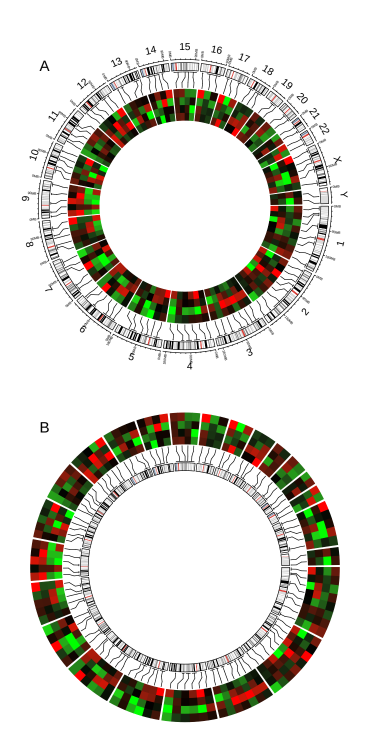


Figure 10: Position transformation with posTransform.default. A) transformation is inside; B) transformation is outside.

#### 8.1.2 Position transformation for text

There is another position transformation function posTransform.text provided in circlize that can smartly position text on the circle. Normally, we don't want the text too away from the original position and also we want to get avoid of text overlapping. posTransform.text calculates the height of text and transform positions properly. Since such text position transformation relies on font size of text and which track the text is in, the usage of posTransform.text is a little bit complex. Currently, posTransform.text only makes sense if it is called inside circos.genomicText and only works when facing is set to clockwise or reverse.clockwise.

In following example code, using posTransform.text is quite similar as posTransform.default (but note this function is quit experimental):

For the next track where position transformation lines are plotted. Code will be more complex. Since the calculation of the position of the line ends which corresponds to the transformed track relies on text settings (e.g. font size, font family) and track for the text (i.e. the track will affect the position of text), we need to go back to the track where text position transformation happened and use the same settings for the text. Such information should be collected when plotting transformation lines. The solution by **circlize** is to wrap posTransform.text with such information, then text information will be recovered and the transformation will be calculated in the correct track.

As we mentioned before, posTransform expect at least one argument, so the argument value is completely fine here.

```
i_track = get.cell.meta.data("track.index") # the nearest track
# we put `y`, `labels`, ... into a self-defined function
# because these parameters will affect the text position
circos.genomicPosTransformLines(bed, direction = "outside",
    posTransform = function(region, value)
        posTransform.text(region, y = 0, labels = value[[1]],
        cex = 0.8, track.index = i_track)
)
```

If the transformation line track is plotted before the track where text position is transformed, things will be more complicated. Since circos.genomicPosTransformLines needs information of the text, but at that moment, track which transforms text positions has not be created. The solution is first creating an empty track for position transformation lines, then adding the position transformation track. Finally go back to the transformation line track to add transformation lines.

```
posTransform = function(region, value)
    posTransform.text(region, y = 1, labels = value[[1]],
        cex = 0.8, track.index = tr_track),
    direction = "inside", track.index = i_track
)
```

Padding of text after transformation can be set through padding argument to adjust the space between two neighbouring text. Of course, when add transformation lines, padding should also be wrapped in.

For examples and comparisons between posTransform.default and posTransform.text when visualizing text, please refer to figure 11. Source code is available in the help page of posTransform.text.

#### 8.1.3 How about if I have a lot of labels?

Position transformation is applied inside each chromosome, which means, the transformed new positions are still located in the original chromosome. This would cause a problem that if you have a lot of labels to put, they will overlap with each other (figure 12 B). Unfortunately there is no simple solution to extend the positions outside of the chromosome.

Anyway, we provide a 'not-too-perfect' solution but the code is a little bit lengthy (figure 12 A). The idea is to put labels and transformation lines in one same track. Here we use posTransform.text directly to calculate the transformed positions.

To make a clear demonstration, we only add graphics in the first quarter of the circle, only on one chromosome.

The steps inside panel.fun are:

- 1. Original positions are plotted as dots.
- 2. In order to get rid of text overlapping, we separate the text into two groups. One group of text are plotted lower than the other group. In following code, the x-axis is splitted into 6 intervals and region and value are separated into groups by looking at whether the middle points of corresponding regions are in the odd intervals (i.e., the first interval, the third interval, ...) or even intervals. Of course, in this step, users should determine their own rule to separate the groups.
- 3. Text in different groups are assigned with different heights (y).

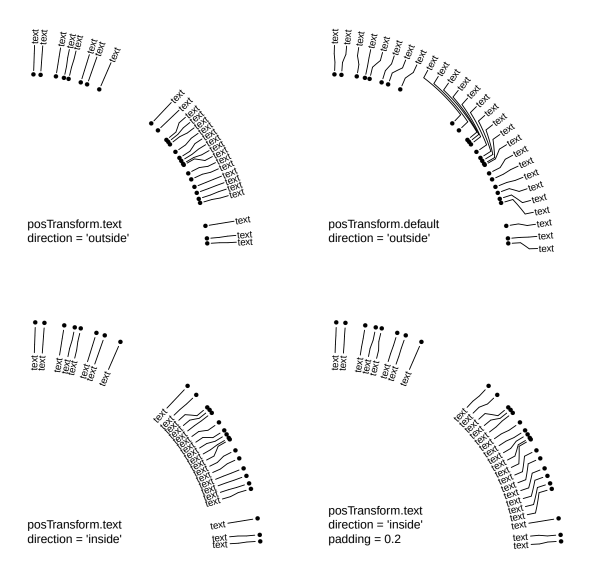


Figure 11: Transformation of text positions.

- 4. Do the position transformation on text and add labels.
- 5. Add the position transformation lines manually. Note the line for lower text and higher text are different. We do something more to make sure lines which points to higher text do not overlap with lower text too much.

```
circos.genomicTrackPlotRegion(bed, ylim = c(0, 1),
    panel.fun = function(region, value, ...) {
    # original positions
    circos.genomicPoints(region, data.frame(rep(0, nrow(region))), pch = 16)
   xlim = get.cell.meta.data("xlim")
   breaks = seq(xlim[1], xlim[2], length.out = 7)
   midpoints = (region[[1]] + region[[2]])/2
   for(i in seq_along(breaks)[-1]) {
        # index for current interval
        l = midpoints >= breaks[i - 1] & midpoints < breaks[i]</pre>
        # if there is no data in this interval
        if(sum(1) == 0) next
        # sub-regions in this interval
        sub_region = region[1, , drop = FALSE]
        sub_value = value[1, , drop = FALSE]
        # note here i == 2, 4, 6, \ldots corresponds to odd intervals
        # text in odd intervals are in lower position
        if(i %% 2 == 0) {
            y = 0.2
        } else {
            y = 0.8
        }
        # get the transformed position and add text with new positions
        tr_region = posTransform.text(sub_region, y = y, labels = sub_value[[1]],
            cex = 0.8, adj = c(0, 0.5))
        circos.genomicText(tr_region, sub_value, labels.column = 1, y = y,
            adj = c(0, 0.5), facing = "clockwise", niceFacing = TRUE, cex = 0.8)
        # add position transformation lines for odd intervals
        if(i %% 2 == 0) {
            for(i in seq_len(nrow(sub_region))) {
                x = c((sub\_region[i, 1] + sub\_region[i, 2])/2,
                       (sub_region[i, 1] + sub_region[i, 2])/2,
                       (tr_region[i, 1] + tr_region[i, 2])/2,
                       (tr_region[i, 1] + tr_region[i, 2])/2)
                y = c(0, 0.2/3, 0.2/3*2, 0.2)
                circos.lines(x, y)
            }
        } else { # add position transformation lines for even intervals
            median_sub_region_midpoint = median(midpoints[1])
            sub_region_width = max(midpoints[1]) - min(midpoints[1])
            for(i in seq_len(nrow(sub_region))) {
                x = c((sub\_region[i, 1] + sub\_region[i, 2])/2,
                       (sub_region[i, 1] + sub_region[i, 2])/2,
                       median_sub_region_midpoint +
                           sub_region_width*(i - nrow(sub_region))/nrow(sub_region) * 0.2,
                       median_sub_region_midpoint +
```

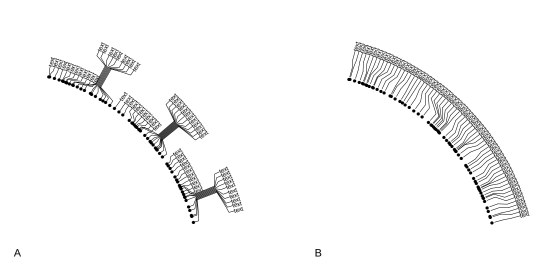


Figure 12: Position transformation for a lot of text. A) put text on two layers; B) put text on one layer.

You can self-define positions of the 'necks' of the long transformation lines so that they will not overlap with the lower text.

#### 8.2 Genomic density and Rainfall plot

circos.genomicDensity calculates how much a genomic window is covered by regions in bed. The input data can be a single data frame or a list of data frames.

```
circos.genomicDensity(bed)
circos.genomicDensity(bed, baseline = 0)
circos.genomicDensity(bed, window.size = 1e6)
circos.genomicDensity(bedlist, col = c("#FF000080", "#0000FF80"))
```

Rainfall plot can be used to visualize distribution of regions. On the plot, y-axis corresponds to the distance to neighbour regions (log10-based). So if there is a drop-down on the plot, it means there is a cluster of regions at that area (figure 13). The input data can be a single data frame or a list of data frames.

```
circos.genoimcRainfall(bed)
circos.genoimcRainfall(bedlist, col = c("red", "green"))
```

Following example makes a rainfall plot for differentially methylated regions (DMR) and their genomic densities. In the plot, red corresponds to hyper-methylated DMRs (gain of methylation) and blue corresponds to hypo-methylated DMRs (loss of methylation) (figure 13).

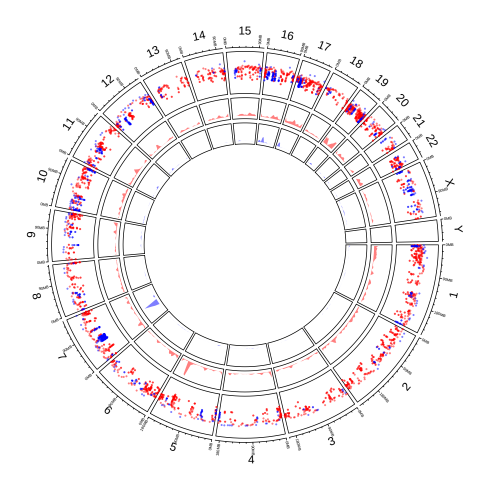


Figure 13: Genomic rainfall plot and densities.

```
load(paste0(system.file(package = "circlize"), "/extdata/DMR.RData"))
par(mar = c(1, 1, 1, 1))
circos.initializeWithIdeogram(plotType = c("axis", "labels"))

bed_list = list(DMR_hyper, DMR_hypo)
circos.genomicRainfall(bed_list, pch = 16, cex = 0.4, col = c("#FF000080", "#0000FF80"))
circos.genomicDensity(DMR_hyper, col = c("#FF000080"), track.height = 0.1)
circos.genomicDensity(DMR_hypo, col = c("#0000FF80"), track.height = 0.1)
```

circos.clear()