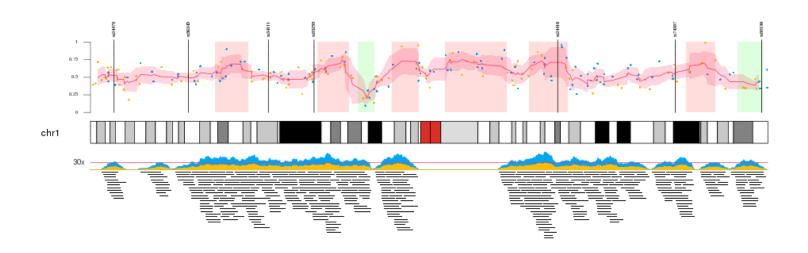
# Visualizing genomic data with karyoploteR

## Quick disclaimer

- R = several ways to get the same result
- Figures arent perfect
- Brief overview

#### Introduction

- · karyoploteR allows you to easily *plot* data across the genome
- built upon base R (sorry Tidyverse people!)



## Installing karyoploteR

Current version:

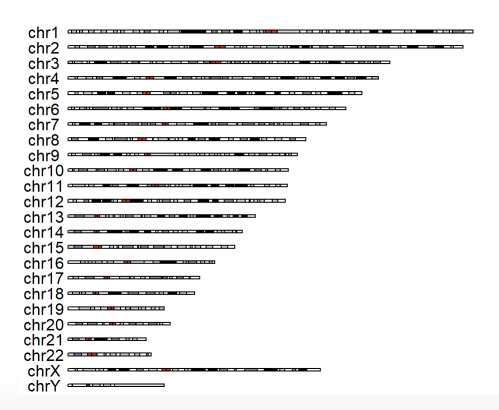
```
if (!requireNamespace("BiocManager", quietly = TRUE))
  install.packages("BiocManager")
  BiocManager::install("karyoploteR")
```

Developmental version can also be downloaded from github using devtools::install github()

#### **Getting started**

library(karyoploteR)

kp <- plotKaryotype()</pre>



#### Layouts

- plot.type = 1
  - data panel above ideogram
- plot.type = 2
  - two data panels one above and one below ideogram
- plot.type = 3
  - plot type 2 but arranged in a row
- plot.type = 4
  - plot type 1 but arranged in a row
- plot.type = 5
  - ideograms in a row with a data panel below
- plot.type = 6
  - no ideogram
- plot.type = 7
  - ideograms only in a row (caution!)

## Adding layers

#### Base R

```
kp <- plotKaryotype(chromosomes="chr17")
kpAddCytobandLabels(kp, force.all=TRUE, srt=90, col="purple", cex=1)</pre>
```

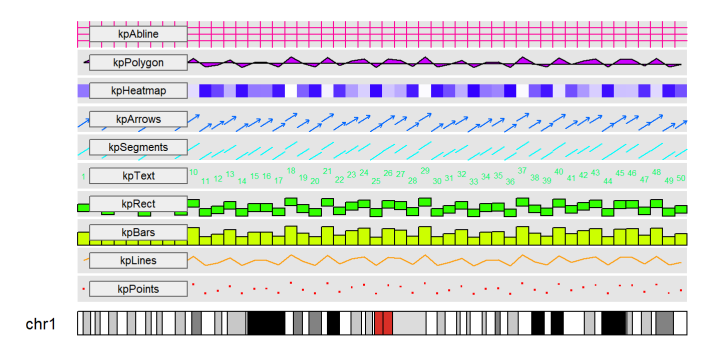
#### magrittr

```
library(magrittr)

kp <- plotKaryotype(chromosomes="chr17") %>%
   kpAddCytobandLabels(force.all=TRUE, srt=90, col="purple", cex=1)
```

cht12 p 13.3 p 1

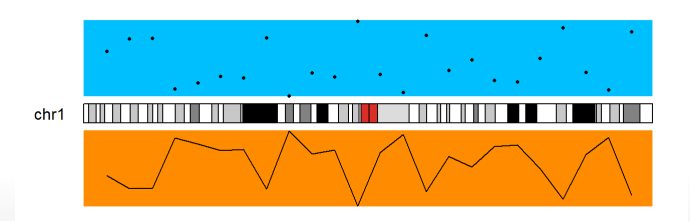
## Basic data types



#### Adding data

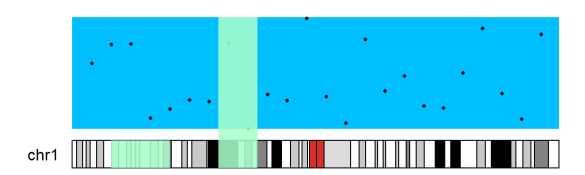
```
x <- 1:24*10e6 # genomic location
y <- runif(n = 24, min = 0, max = 1) #random values for plotting</pre>
```

```
plotKaryotype(plot.type=2, chromosomes = "chr1") %>%
  kpDataBackground(data.panel = 1, col="deepskyblue") %>%
  kpDataBackground(data.panel = 2, col="darkorange") %>%
  kpPoints(chr="chr1", x=x, y=y, data.panel = 1) %>%
  kpLines(chr="chr1", x=x, y=y, data.panel = 2)
```



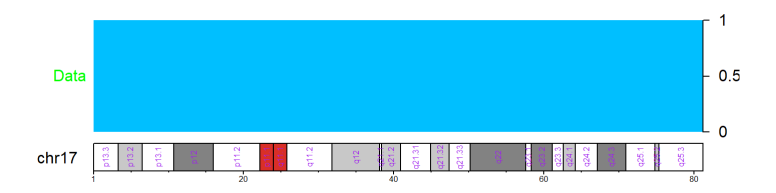
#### **Data Panels**

```
plotKaryotype(plot.type=1, chromosomes = "chr1") %>%
   kpDataBackground(data.panel = 1, col="deepskyblue") %>%
   kpPoints(chr="chr1", x=x, y=y, data.panel = 1) %>%
   kpRect(chr="chr1", x0=20e6, x1=50e6, y0=0, y1=1, col="#AAFFCBDD",
        data.panel="ideogram", border=NA) %>%
   kpRect(chr="chr1", x0=75e6, x1=95e6, y0=0, y1=1, col="#AAFFCBDD",
        data.panel="all", border=NA)
```



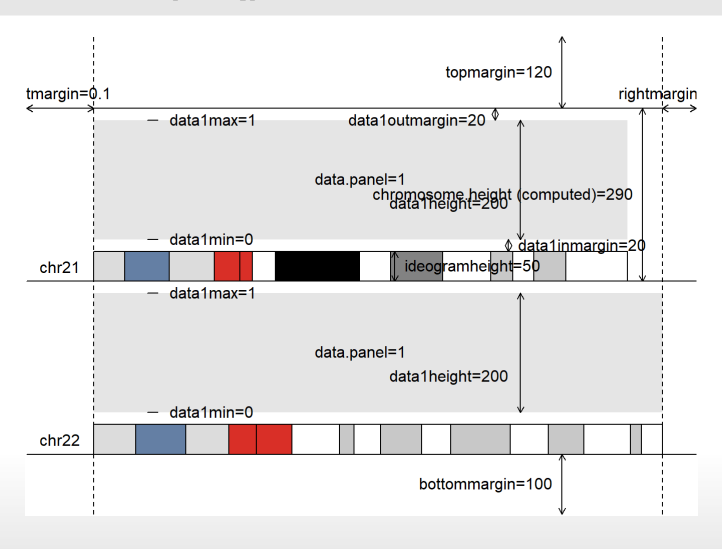
## Adding plot features

```
plotKaryotype(chromosomes="chr17", plot.type = 1) %>%
    kpDataBackground(data.panel = 1, col="deepskyblue") %>%
    kpAddCytobandLabels(force.all=TRUE, srt=90, col="purple", cex=0.5) %>%
    kpAddBaseNumbers() %>%
    kpAddLabels(labels="Data", data.panel = 1, col="green") %>%
    kpAxis(data.panel=1, side=2)
```



## Viewing plotting parameters

plotDefaultPlotParams(plot.type=1)



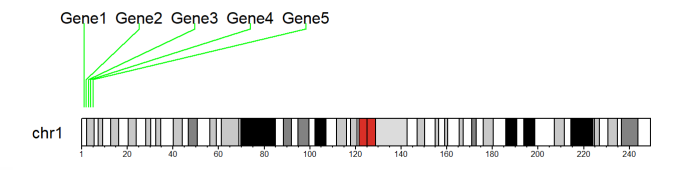
#### Changing plotting parameters

```
pp <- getDefaultPlotParams(plot.type=1)
pp$ideogramheight <- 100 ; pp$data1height <- 100
plotKaryotype(chromosomes="chr1", plot.type=2, plot.params = pp) %>%
   kpDataBackground(color = "deepskyblue")
```



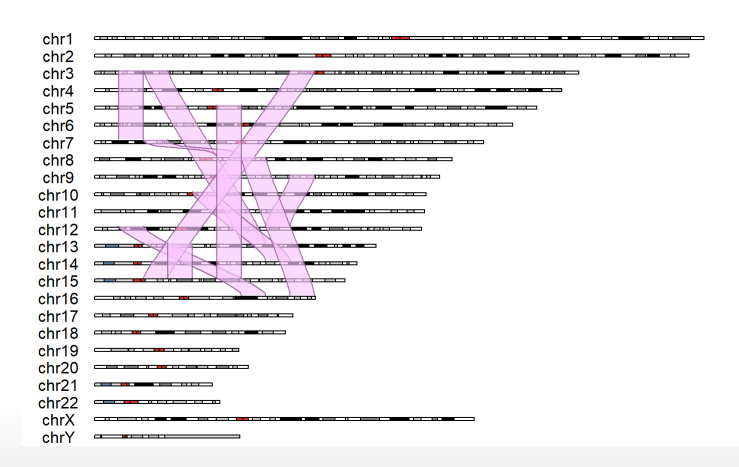


#### Gene markers



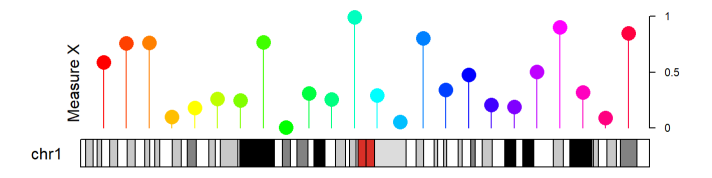
## Linking regions

```
plotKaryotype() %>%
kpPlotLinks(data=start.regs, data2=end.regs, col="#fac7ffaa")
```



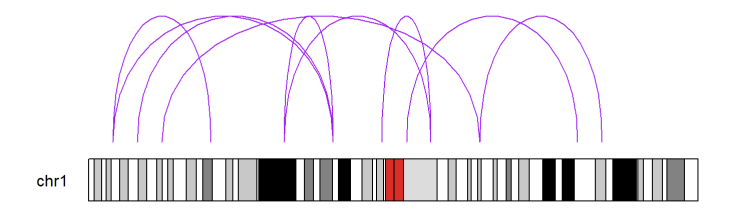
# Examples

#### **Lollipop Plot**



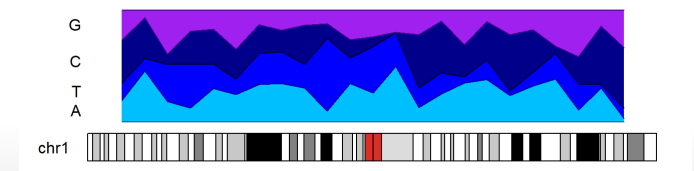
#### **5C**

```
plotKaryotype(chromosomes = "chr1") %>%
kpPlotLinks(data=start.regs, data2=end.regs, border="purple")
```



#### Nucleotide coverage

```
plotKaryotype(chromosomes = "chr1") %>%
   kpPlotRibbon(data=tmp, y0=0, y1=tmp$A, col="deepskyblue") %>%
   kpPlotRibbon(data=tmp, y0=tmp$A, y1=tmp$T, col="blue") %>%
   kpPlotRibbon(data=tmp, y0=tmp$T, y1=tmp$C, col="darkblue") %>%
   kpPlotRibbon(data=tmp, y0=tmp$C, y1=tmp$G, col="purple") %>%
   kpAddLabels(r0 = 0, r1=tmp$A[1], labels = "A") %>%
   kpAddLabels(r0 = tmp$A[1], r1=tmp$T[1], labels = "T") %>%
   kpAddLabels(r0 = tmp$T[1], r1=tmp$C[1], labels = "C") %>%
   kpAddLabels(r0 = tmp$C[1], r1=tmp$G[1], labels = "G")
```



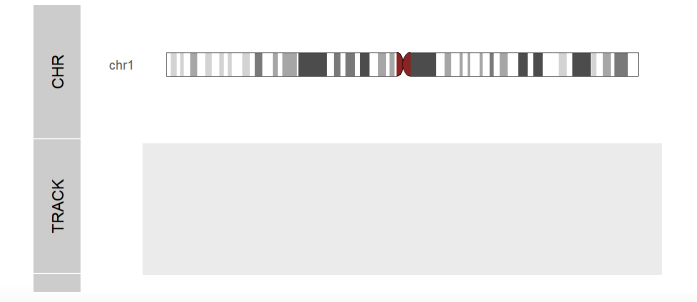
## Other ideogram options

## ggbio

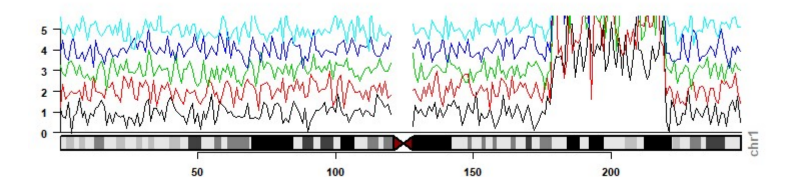
```
library(ggbio)
p.ideo <- Ideogram(genome = "hg19", subchr = "chr1")

track1 <- ggplot() + geom_blank()

tracks(CHR = p.ideo, TRACK = track1)</pre>
```



#### IdeoViz



#### Additional resources

- bioconductor vignettes
- github tutorial

