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Chapter 1

Getting started

1.1 Citation

```
citation("ggbio")
## To cite package 'ggbio' in publications use:
##
     Tengfei Yin, Dianne Cook and Michael Lawrence (2012): ggbio: an R
##
     package for extending the grammar of graphics for genomic data Genome
     Biology 13:R77
##
##
## A BibTeX entry for LaTeX users is
##
##
     @Article{,
       title = {ggbio: an R package for extending the grammar of graphics for genomic data},
##
##
       author = {Tengfei Yin and Dianne Cook and Michael Lawrence},
       journal = {Genome Biology},
       volume = \{13\},
##
       number = \{8\},
       pages = \{R77\},
       year = \{2012\},\
##
       publisher = {BioMed Central Ltd},
```

1.2 Introduction

ggbio is a Bioconductor package building on top of ggplot2(), leveraging the rich objects defined by Bioconductor and its statistical and computational power, it provides a flexible genomic visualization framework, extends the grammar of graphics into genomic data, try to delivers high quality, highly customizable graphics to the users.

What it features

autoplot function provides ready-to-use template for Bioconductor objects and different types of data.

- flexible low level components to use grammar of graphics to build you graphics layer by layer.
- layout transformation, so you could generate circular plot, grandlinear plot, stacked overview more easily.
- flexible tracks function to bind any ggplot2(), ggbio based plots.

Chapter 2

Case study: building your first tracks

In this chapter, you will learn

- how to add ideogram track.
- How to add gene model track.
- how to add track for bam files to visualize coverage and mismatch summary.
- how to add track for vcf file to visualize the variants.

2.1 Add an ideogram track

Ideogram provides functionality to construct ideogram, check the manual for more flexible methods. We build genome *hg19*, *hg18*, *mm10*, *mm9* inside, so you don't have download it on the fly. When embed with tracks, ideogram show zoomed region highlights automatically. xlim has special function here, is too changed highlighted zoomed region on the ideogram.

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

chr1

library(GenomicRanges)
## special highlights instead of zoomin!
p.ideo + xlim(GRanges("chr2", IRanges(1e8, 1e8+10000000)))</pre>
```

2.2 Add a gene model track

2.2.1 Introduction

Gene model track is one of the most frequently used track in genome browser, it is composed of genetic features CDS, UTR, introns, exons and non-genetic region. In *ggbio* we support three methods to make gene model track:

- OrganismDb object: recommended, support gene symbols and other combination of columns as label.
- TxDb object: don't support gene symbol labeling.
- *GRangesList* object: flexible, if you don't have annotation package available for the first two methods, you could prepare a data set parsed from gtf file, you can simply use it and plot it as gene model track.
- EnsDb object: supports gene symbol labeling, filtering etc.

2.2.2 Make gene model from OrganismDb object

OrganismDb object has a simpler API to retrieve data from different annotation resources, so we could label our transcripts in different ways

```
library(ggbio)
library(Homo.sapiens)
class(Homo.sapiens)

## [1] "OrganismDb"

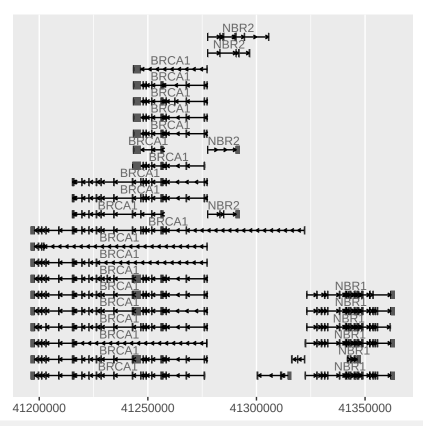
## attr(,"package")

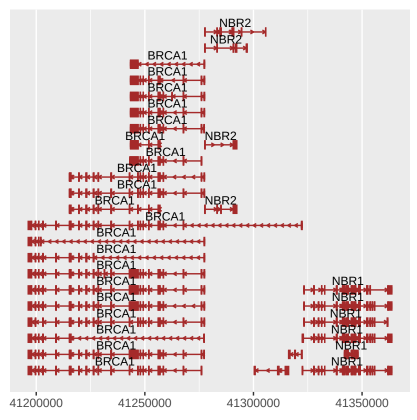
## [1] "OrganismDbi"

##

data(genesymbol, package = "biovizBase")
wh <- genesymbol[c("BRCA1", "NBR1")]
wh <- range(wh, ignore.strand = TRUE)

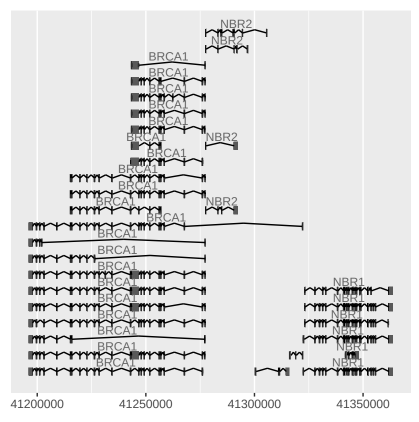
p.txdb <- autoplot(Homo.sapiens, which = wh)
p.txdb</pre>
```





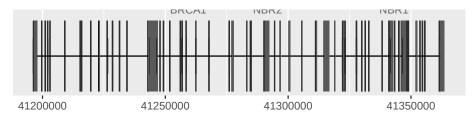
To change the intron geometry, use geom_alignment for more control parameters.

autoplot(Homo.sapiens, which = wh, gap.geom = "chevron")



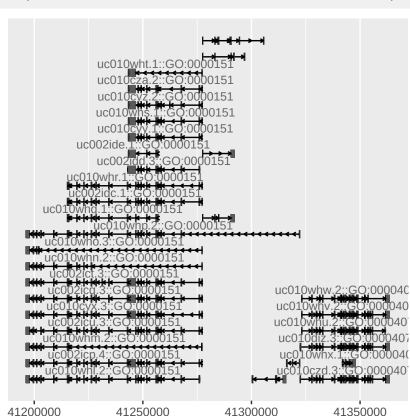
To collapse all features, use stat 'reduce'

```
autoplot(Homo.sapiens, which = wh, stat = "reduce")
```



Label could be turned off by setting it to FALSE, you could also use expression to make a flexible label combination from column names.

col	<pre>columns(Homo.sapiens)</pre>								
##	[1]	"ACCNUM"	"ALIAS"	"CDSCHROM"	"CDSEND"	"CDSID"			
##	[6]	"CDSNAME"	"CDSSTART"	"CDSSTRAND"	"DEFINITION"	"ENSEMBL"			
##	[11]	"ENSEMBLPROT"	"ENSEMBLTRANS"	"ENTREZID"	"ENZYME"	"EVIDENCE"			
##	[16]	"EVIDENCEALL"	"EXONCHROM"	"EXONEND"	"EXONID"	"EXONNAME"			
##	[21]	"EXONRANK"	"EXONSTART"	"EXONSTRAND"	"GENEID"	"GENENAME"			
##	[26]	"GENETYPE"	"G0"	"GOALL"	"GOID"	"IPI"			
##	[31]	"MAP"	"OMIM"	"ONTOLOGY"	"ONTOLOGYALL"	"PATH"			
##	[36]	"PFAM"	"PMID"	"PROSITE"	"REFSEQ"	"SYMBOL"			
##	[41]	"TERM"	"TXCHROM"	"TXEND"	"TXID"	"TXNAME"			
##	[46]	"TXSTART"	"TXSTRAND"	"TXTYPE"	"UCSCKG"	"UNIPROT"			

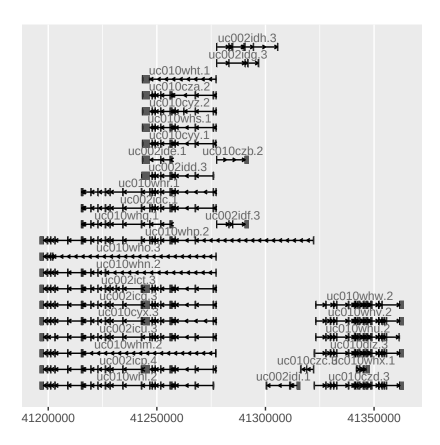


autoplot(Homo.sapiens, which = wh, columns = c("TXNAME", "GO"), names.expr = "TXNAME::GO")

2.2.3 Make gene model from *TxDb* object

TxDb doesn't contain any gene symbol information, so we use tx_id as default for label.

```
library(TxDb.Hsapiens.UCSC.hg19.knownGene)
txdb <- TxDb.Hsapiens.UCSC.hg19.knownGene
autoplot(txdb, which = wh)</pre>
```

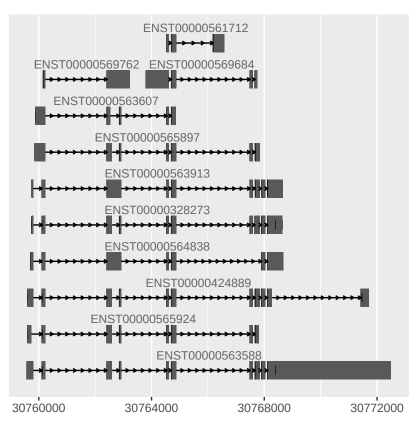


2.2.4 Make gene model from EnsDb object

An alternative source for gene models are the *EnsDb* objects from the *ensembldb* package that provide gene annotations provided from Ensembl. The *ensembldb* package provides a rich filtering system that allows to easily fetch specific information (genes/transcripts) from an *EnsDb*. The *EnsDb* objects provide gene symbol annotations in the column gene_name. Alternatively, we could use tx_id to label transcripts.

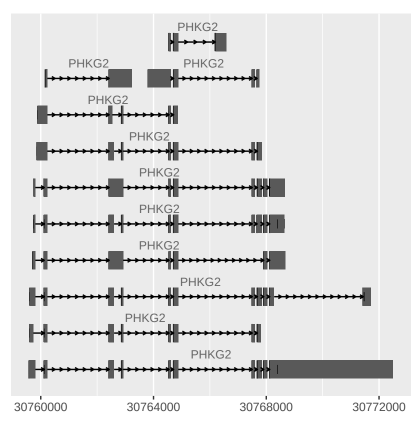
In the example below we plot the gene model of the gene PHKG2. We use a *GenenameFilter* to specify which gene we want to plot.

```
library(EnsDb.Hsapiens.v75)
ensdb <- EnsDb.Hsapiens.v75
autoplot(ensdb, GeneNameFilter("PHKG2"))</pre>
```



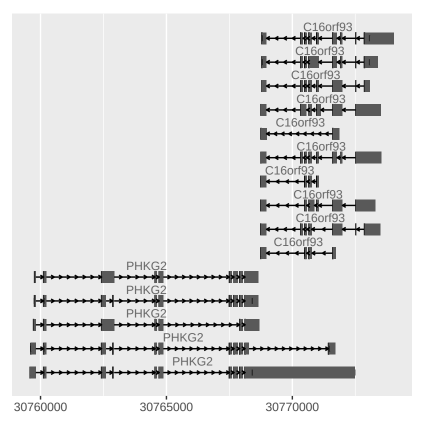
We can pass any filter class defined in the *AnnotationFilter* package with argument *which*. Alternatively we can combine filter classes using an *AnnotationFilterList* or we can pass a filter expression in form of a *formula*. Below we pass such a filter expression to the function.

```
autoplot(ensdb, ~ symbol == "PHKG2", names.expr="gene_name")
```



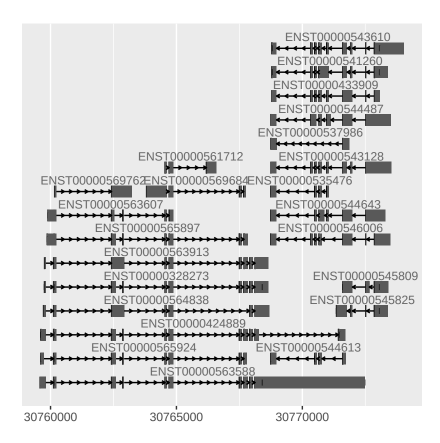
We could also specify a genomic region and fetch all transcripts overlapping that region (also partially, i.e. with a part of an intron or an exon).

```
## We specify "*" as strand, thus we query for genes encoded on both strands
gr <- GRanges(seqnames = 16, IRanges(30768000, 30770000), strand = "*")
autoplot(ensdb, GRangesFilter(gr), names.expr = "gene_name")</pre>
```



Also, we can spefify directly the gene ids and plot all transcripts of these genes (not only those overlapping with the region)

autoplot(ensdb, GeneIdFilter(c("ENSG00000196118", "ENSG00000156873")))



2.2.5 Make gene model from GRangesList object

Sometimes your gene model is not available as none of *OrganismDb* or *TxDb* object, it's may be stored in a table, you could simple parse it into a *GRangeList* object.

- each group indicate one transcripts
- names of group are shown as labels
- this object must has a column contains following key word: cds, exon, intron, and it's
 not case senstitive. use type to map this column. By default, we will try to parse 'type'
 column.

Let's make a sample GRangesList object which contains all information, and fake some labels.

```
library(biovizBase)
gr.txdb <- crunch(txdb, which = wh)
## change column to 'model'
colnames(values(gr.txdb))[4] <- "model"
grl <- split(gr.txdb, gr.txdb$tx_id)
## fake some randome names
names(grl) <- sample(LETTERS, size = length(grl), replace = TRUE)
grl

## GRangesList object of length 32:
## $W
## GRanges object with 7 ranges and 4 metadata columns:</pre>
```

```
ranges strand | tx_id
        segnames
                                                        tx_name
          <Rle>
                      <IRanges> <Rle> | <character> <character> <character>
##
                                           61241 uc002idf.3
           chr17 41277600-41277787
##
    [1]
                                    + |
                                                                    10230
                                               61241 uc002idf.3
##
    [2]
          chr17 41283225-41283287
                                     + |
                                                                     10230
                                             61241 uc002idf.3
##
    [3]
        chr17 41284973-41285154
                                    + |
                                                                     10230
                                                                    10230
    [4]
        chr17 41290674-41292342
                                             61241 uc002idf.3
##
                                    + |
    61241 uc002idf.3
61241 uc002idf.3
##
                                                                     10230
##
                                                                     10230
                                            61241 uc002idf.3
                                                                    10230
##
          model
##
     <factor>
##
    [1] exon
##
    [2]
          exon
    [3]
##
          exon
##
    [4]
          exon
    [5]
##
        gap
##
    [6]
           gap
##
    [7]
           gap
##
##
    seqinfo: 1 sequence from hg19 genome
##
## $I
## GRanges object with 3 ranges and 4 metadata columns:
        segnames
                    ranges strand | tx_id tx_name
          <Rle>
##
                      <IRanges> <Rle> | <character> <character> <character>
##
        chr17 41277600-41277787
                                 + | 61242 uc010czb.2
    [1]
                                            61242 uc010czb.2
61242 uc010czb.2
##
    [2] chr17 41290674-41292342
                                                                     10230
                                     + |
    [3] chr17 41277788-41290673
                                    * |
                                                                     10230
##
          model
##
       <factor>
##
    [1] exon
##
    [2]
           exon
    [3]
##
            gap
##
##
    seqinfo: 1 sequence from hg19 genome
##
## $H
## GRanges object with 9 ranges and 4 metadata columns:
        seqnames ranges strand | tx_id tx_name
                      <IRanges> <Rle> | <character> <character> <character>
##
          <Rle>
                                   + | 61243 uc002idg.3
##
    [1]
          chr17 41277600-41277787
                                                                   10230
                                            61243 uc002idg.3
##
    [2]
        chr17 41283225-41283287
                                                                     10230
                                     + |
    [3]
        chr17 41290674-41290939
                                             61243 uc002idg.3
                                                                     10230
                                    + |
    [4] chr17 41291833-41292300 + |

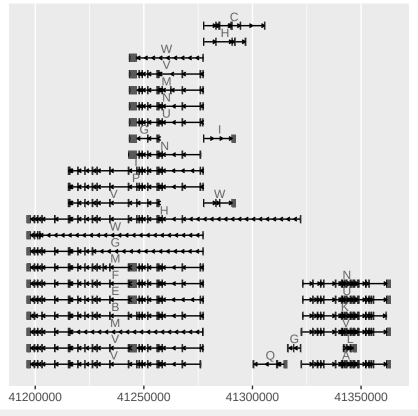
[5] chr17 41296745-41297125 + |

[6] chr17 41277788-41283224 * |
                                            61243 uc002idg.3
61243 uc002idg.3
61243 uc002idg.3
##
                                                                     10230
##
                                                                     10230
##
                                                                     10230
    [7] chr17 41283288-41290673
                                    * |
                                             61243 uc002idg.3
##
                                                                     10230
                                          61243 uc002idg.3
61243 uc002idg.3
##
    [8]
         chr17 41290940-41291832
                                    *
                                                                     10230
           chr17 41292301-41296744
##
    [9]
                                    * |
                                                                     10230
##
          model
##
        <factor>
```

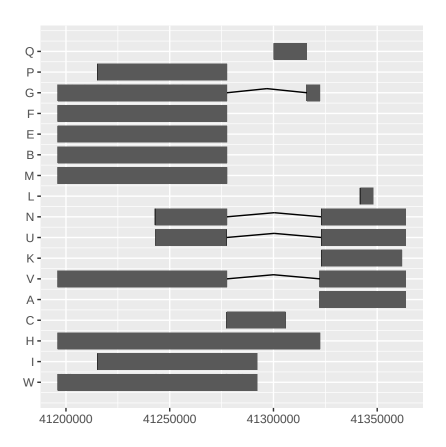
```
[1]
##
              exon
##
     [2]
              exon
##
     [3]
              exon
##
     [4]
              exon
##
     [5]
              exon
##
     [6]
              gap
##
     [7]
              gap
##
     [8]
              gap
##
     [9]
              gap
##
     seqinfo: 1 sequence from hg19 genome
##
##
##
## <29 more elements>
```

We get our example data ready, it meets all requirements, to make it a gene model track it's pretty simple to use autoplot, but don't forget mapping because we changed our column names, asssume you store you model key words in column 'model'.

```
autoplot(grl, aes(type = model))
```



ggplot() + geom_alignment(grl, type = "model")



2.3 Add a reference track

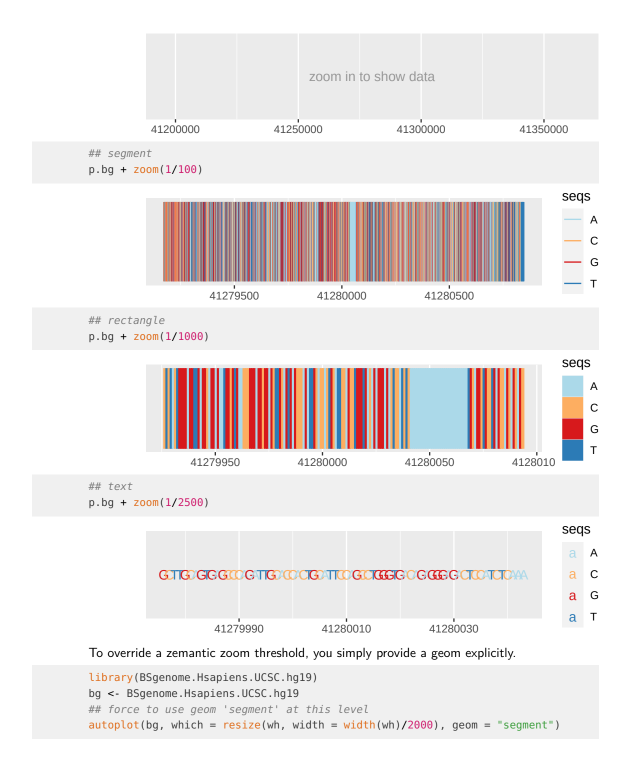
To add a reference track, we need to load a *BSgenome* object from the annotation package. You can choose to plot the sequence as *text, rect, segment*.

2.3.1 Semantic zoom

Here we introduce semantic zoom in *ggbio*, for some plots like reference sequence, we use pre-defined zoom level threshold to automatically assign geom to the track, unless the geom is explicitly specified. In the example below, when your region is too wide we show text 'zoom in to see text', when you zoom into different level, it shows you different details. zoom is a function we will introduce more in chapter 3 when we introduce more about navigation.

You can pass a zoom in factor into **zoom** function, if it's over 1 it's zooming out, if it's smaller than 1 it's zooming in.

```
library(BSgenome.Hsapiens.UCSC.hg19)
bg <- BSgenome.Hsapiens.UCSC.hg19
p.bg <- autoplot(bg, which = wh)
## no geom
p.bg</pre>
```



2.4 Add an alignment track

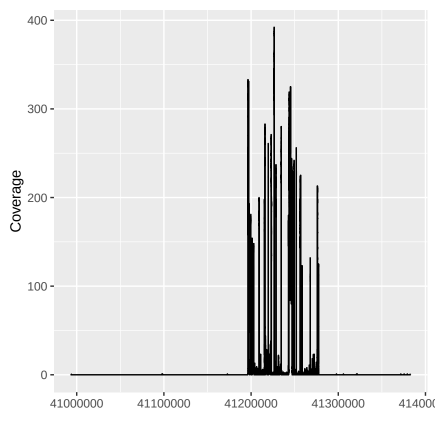
ggbio supports visuaization of alignemnts file stored in bam, autoplot method accepts

bam file path (indexed)

- BamFile object
- GappedAlignemnt object

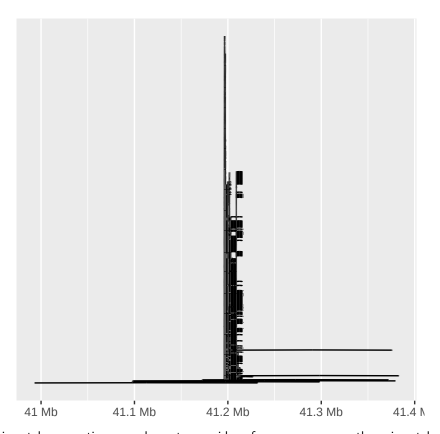
It's simple to just pass a file path to autoplot function, you can stream a chunk of region by providing 'which' parameter. Otherwise please use method 'estiamte' to show overall estiamted coverage.

```
fl.bam <- system.file("extdata", "wg-brcal.sorted.bam", package = "biovizBase")
wh <- keepSeqlevels(wh, "chr17")
autoplot(fl.bam, which = wh)</pre>
```



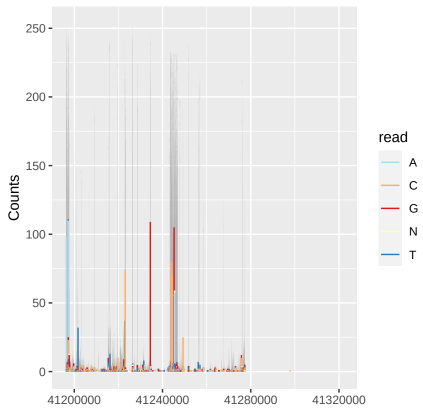
geom 'gapped pair' will show you alignments.

```
fl.bam <- system.file("extdata", "wg-brcal.sorted.bam", package = "biovizBase")
wh <- keepSeqlevels(wh, "chr17")
autoplot(fl.bam, which = resize(wh, width = width(wh)/10), geom = "gapped.pair")</pre>
```



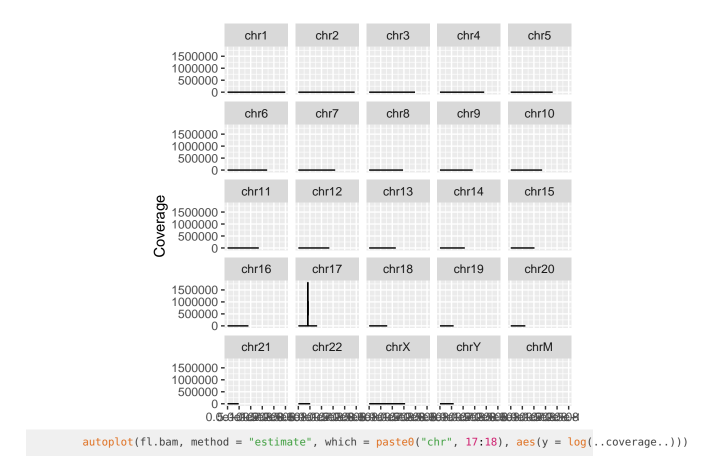
To show mismatch proportion, you have to provide reference sequence, the mismatched proportion is color coded in the bar chart.

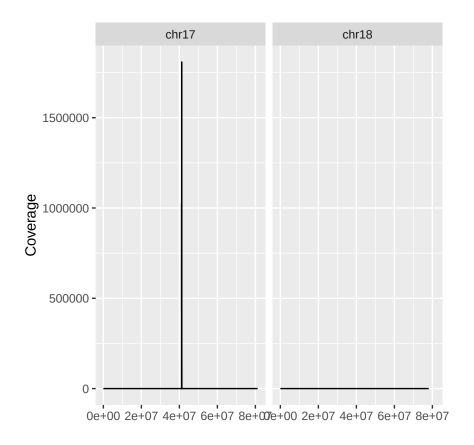
```
library(BSgenome.Hsapiens.UCSC.hg19)
bg <- BSgenome.Hsapiens.UCSC.hg19
p.mis <- autoplot(fl.bam, bsgenome = bg, which = wh, stat = "mismatch")
p.mis</pre>
```



To view overall estimated coverage distribution, please use method 'estiamte'. 'which' parameter also accept characters. And there is a hidden value called '..coverage..' to let you do simple transformation in aes().

autoplot(fl.bam, method = "estimate")





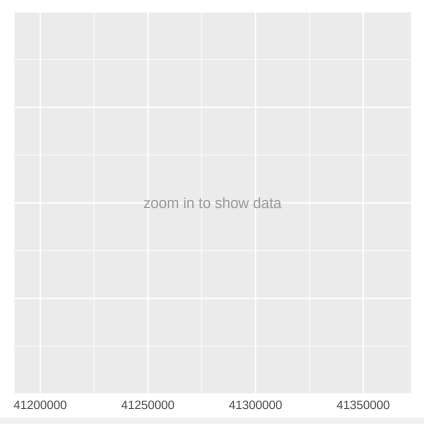
2.5 Add a variants track

This track is supported by semantic zoom.

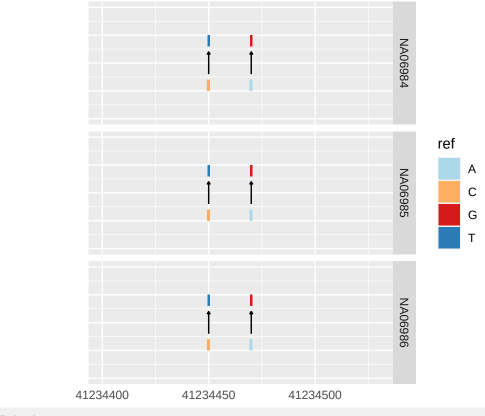
To view your variants file, you could

- Import it using package VariantAnntoation as VCF object, then use autoplot
- Convert it into *VRanges* object and use autoplot.
- Simply provide vcf file path in autoplot().

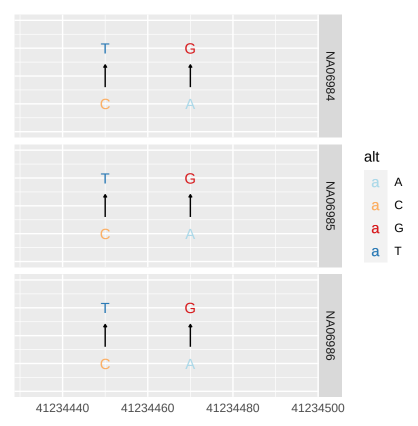
```
library(VariantAnnotation)
fl.vcf <- system.file("extdata", "17-1409-CEU-brca1.vcf.bgz", package="biovizBase")
vcf <- readVcf(fl.vcf, "hg19")
vr <- as(vcf[, 1:3], "VRanges")
vr <- renameSeqlevels(vr, value = c("17" = "chr17"))
## small region contains data
gr17 <- GRanges("chr17", IRanges(41234400, 41234530))
p.vr <- autoplot(vr, which = wh)
## none geom
p.vr</pre>
```



rect geom
p.vr + xlim(gr17)



text geom
p.vr + xlim(gr17) + zoom()

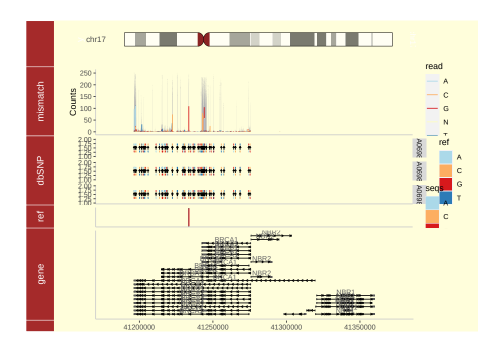


You can simply overide geom

```
autoplot(vr, which = wh, geom = "rect", arrow = FALSE)
```

2.6 Building your tracks

```
## tks <- tracks(p.ideo, mismatch = p.mis, dbSNP = p.vr, ref = p.bs, gene = p.txdb)
## tks <- tracks(fl.bam, fl.vcf, bs, Homo.sapiens) ## default ideo = FALSE, turned on
## tks <- tracks(fl.bam, fl.vcf, bs, Homo.sapiens, ideo = TRUE)
## tks + xlim(gr17)
gr17 <- GRanges("chr17", IRanges(41234415, 41234569))
tks <- tracks(p.ideo, mismatch = p.mis, dbSNP = p.vr, ref = p.bg, gene = p.txdb,
heights = c(2, 3, 3, 1, 4)) + xlim(gr17) + theme_tracks_sunset()
tks</pre>
```



Chapter 3

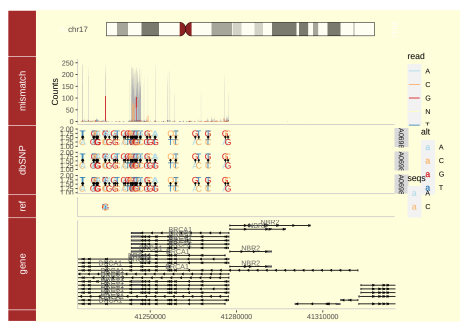
Simple navigation

We try to provide a simple navigation API for your plot, so you could zoom in and zoom out, or go through view chunks one by one.

- zoom: put a factor inside and you can zoom in or zoom out
- nextView: switch to next view
- prevView: switch to previous view

Navigation function also works for tracks plot too.

```
## zoom in
tks + zoom()
```



Try following command yourself.

```
## zoom in with scale
p.txdb + zoom(1/8)
## zoom out
p.txdb + zoom(2)
## next view page
p.txdb + nextView()
## previous view page
p.txdb + prevView()
```

Don't forget xlim accept *GRanges* object (single row), so you could simply prepare a *GRanges* to store the region of interests and go through them one by one.

Chapter 4

Overview plots

Overview is a good way to show all events at the same time, give overall summary statics for the whole genome.

In this chapter, we will introcue three different layouts that are used a lots in genomic data visualization.

4.1 how to make circular plots

4.1.1 Introduction

Circular view is a special layout in *ggbio*, this idea has been implemented in many different software, for example, the *Circos* project. However, we keep the grammar of graphics for users, so mapping variables to aesthetics is very easy, *ggbio* leverage the data structure defiend in *Bioconductor* to make this process as simple as possible.

4.1.2 Buidling circular plot layer by layer

Ok, let's start to process some raw data to the format we want. The data used in this study is from this a paper¹. In this tutorial, We are going to

¹http://www.nature.com/ng/journ

- 1. Visualize somatic mutation as segment.
- 2. Visualize inter, intro-chromosome rearrangement as links.
- 3. Visualize mutation score as point tracks with grid-background.
- 4. Add scale and ticks and labels.
- 5. To arrange multiple plots and legend. create multiple sample comparison.

All the raw data processed and stored in GRanges ready for use, you can simply load the sample data from biovizBase

```
data("CRC", package = "biovizBase")
```

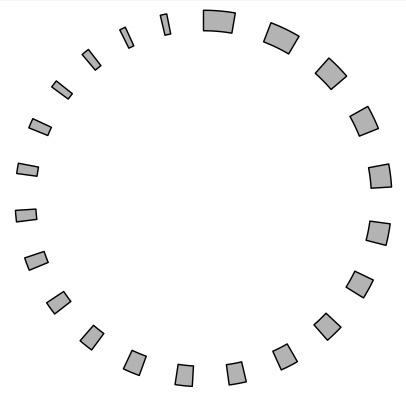
layout_circle is depreicated, because you have to set up radius and trackWidth manually with this function for creating circular plot.

We now present the new circle function, it accepts Granges object, and users don't have to specify radius, track width, you just add them one by one, it will be automatically created from innter circle to outside, unless you specify trackWidth and radius manually. To change default radius and trackWidth for all tracks, you simply put them in ggbio function.

- rule of thumb seqlengths, seqlevels and chromosomes names should be exactly the same
- to use circle, you have to use ggbio constructor at the beginning instead of ggplot.

You can use autoplot to create single track easily like

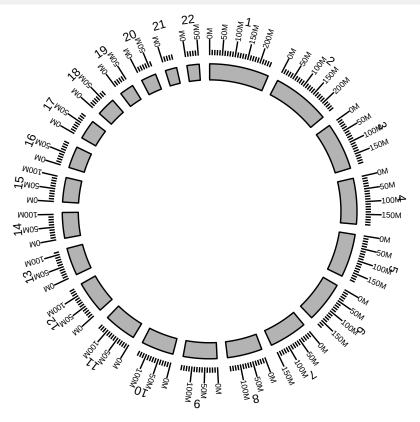
```
head(hg19sub)
## GRanges object with 6 ranges and 0 metadata columns:
         seqnames
                       ranges strand
##
            <Rle>
                    <IRanges> <Rle>
##
     [1]
                1 1-249250621
     [2]
                2 1-243199373
##
##
     [3]
                3 1-198022430
##
     [4]
                4 1-191154276
##
     [5]
                5 1-180915260
##
                6 1-171115067
##
     seginfo: 22 seguences from hg19 genome
autoplot(hg19sub, layout = "circle", fill = "gray70")
```



Hoever, the low level circle function leave you more flexibility to build circular plot one by one. Let's start to add tracks one by one.

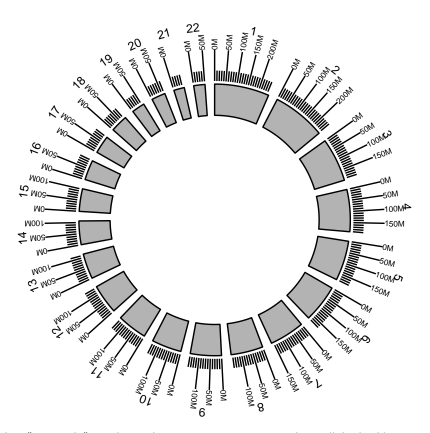
Let's use the same data to create ideogram, label and scale track, it layouts the circle by the order you created from inside to outside.

```
p <- ggbio() + circle(hg19sub, geom = "ideo", fill = "gray70") +
    circle(hg19sub, geom = "scale", size = 2) +
    circle(hg19sub, geom = "text", aes(label = seqnames), vjust = 0, size = 3)
p</pre>
```



To simply override the setting, you can do it globally in ggbio function or individually circle function by specifying parametters trackWidth and radius, you can also specify the global settin for buffer in between in ggbio like example below.

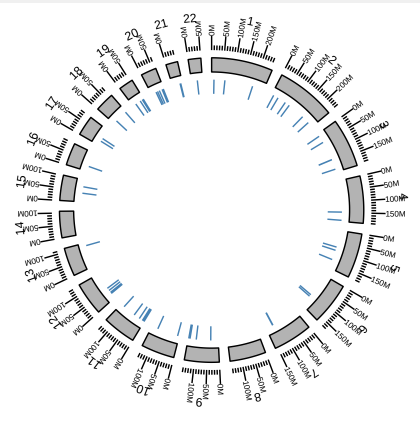
```
p <- ggbio(trackWidth = 10, buffer = 0, radius = 10) + circle(hg19sub, geom = "ideo", fill = "gray70") +
        circle(hg19sub, geom = "scale", size = 2) +
        circle(hg19sub, geom = "text", aes(label = seqnames), vjust = 0, size = 3)
p</pre>
```



Then we add a "rectangle" track to show somatic mutation, this will looks like vertical segments.

```
head(mut.gr)
## GRanges object with 6 ranges and 10 metadata columns:
##
         seqnames
                      ranges strand | Hugo_Symbol Entrez_Gene_Id
                                                                      Center
##
                                          <factor>
                                                          <integer> <factor>
             <Rle> <IRanges> <Rle>
##
                 1 11003085
                                           TARDBP
                                                              23435
                                                                       Broad
     [1]
                                                              10207
##
     [2]
                 1 62352395
                                            INADL
                                                                       Broad
##
     [3]
                 1 194960885
                                            CFH
                                                               3075
                                                                       Broad
##
     [4]
                 2 10116508
                                            CYS1
                                                             192668
                                                                       Broad
                 2
##
     [5]
                    33617747
                                            RASGRP3
                                                              25780
                                                                       Broad
##
                 2 73894280
                                                             388960
                                                                       Broad
                                            C2orf78
                                   + |
         NCBI\_Build
##
                       Strand Variant_Classification Variant_Type Reference_Allele
##
           <integer> <factor>
                                                                              <factor>
                                              <factor>
                                                            <factor>
##
     [1]
                  36
                                              Missense
                                                                 SNP
                                                                                     G
##
     [2]
                  36
                                              Missense
                                                                 SNP
                                                                                     T.
##
     [3]
                  36
                                              Missense
                                                                 SNP
                                                                                     G
##
     [4]
                  36
                                                                 SNP
                                                                                     C
                                              Missense
##
     [5]
                  36
                                              Missense
                                                                 SNP
                                                                                     C
##
                  36
                                              Missense
                                                                 SNP
                                                                                     T.
     [6]
##
         Tumor_Seq_Allele1 Tumor_Seq_Allele2
##
                   <factor>
                                      <factor>
##
     [1]
                          G
                                              Α
                          Τ
                                              G
##
     [2]
                          G
##
     [3]
                                              Α
```

```
##
     [4]
                          C
##
     [5]
                          C
                                             Т
                          Т
                                             C
##
     [6]
##
     seginfo: 22 seguences from an unspecified genome
p <- ggbio() + circle(mut.gr, geom = "rect", color = "steelblue") +</pre>
    circle(hg19sub, geom = "ideo", fill = "gray70") +
    circle(hg19sub, geom = "scale", size = 2) +
  circle(hg19sub, geom = "text", aes(label = seqnames), vjust = 0, size = 3)
р
```



Next, we need to add some "links" to show the rearrangement, of course, links can be used to map any kind of association between two or more different locations to indicate relationships like copies or fusions. To create a suitable structure to plot, please use another *GRanges* to represent the end of the links, and stored as elementMetadata for the "start point" *GRanges*. Here we named it as "to.gr" and will be used later.

```
head(crc.gr)
## GRanges object with 6 ranges and 17 metadata columns:
##
                      ranges strand | individual
                                                                 class
##
            <Rle> <IRanges>
                              <Rle>
                                        <factor> <integer>
                                                              <factor> <numeric>
##
     [1]
               18 56258628
                                           CRC-4
                                                          1 long_range
                                                                          2104165
##
     [2]
               18 44496014
                                            CRC-4
                                                          1 long_range
                                                                        12947165
     [3]
               18
                   45023683
                                            CRC-4
                                                          0 long_range
                                                                        13356670
                   52186319
                                            CRC-4
                                                          0 deletion
     [4]
                                                                              268
```

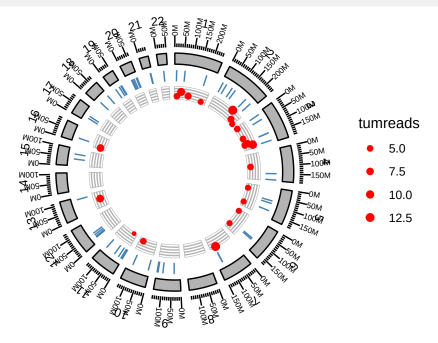
```
##
     [5]
                8 37328910
                                           CRC-4
                                                         0 inter_chr
                                                                             NaN
                8 35575394
                                           CRC-4
                                                         0 inter_chr
##
     [6]
                                                                             NaN
##
          tumreads normreads
                                gene1
                                          gene2
##
         <integer> <integer> <factor> <factor>
##
     [1]
               491
                           2 MC4R
                                        ZCCHC2
                           0 KIAA0427 CDH20
##
     [2]
               265
##
     [3]
               238
                           0 DYM
                                        ZCCHC2
##
                94
                           0 PXDNL
                                        PXDNL
     [4]
##
     [5]
                56
                           0 ZNF703
                                        PAK7
##
     [6]
                53
                           0 UNC5D
                                        RALGAPB
##
                                            site1
##
                                         <factor>
##
    [1] IGR: 69Kb before MC4R(-)
##
     [2] Intron of KIAA0427(+): 4Kb after exon 8
##
     [3] Intron of DYM(-): 14Kb after exon 13
     [4] IGR: 208Kb before PXDNL(-)
##
##
     [5] IGR: 344Kb before ZNF703(+)
##
     [6] Intron of UNC5D(+): 3Kb after exon 4
##
                                              site2
##
                                           <factor>
##
     [1] Intron of ZCCHC2(+): 222bp before exon 4
##
     [2] IGR: 134Kb before CDH20(+)
##
    [3] Intron of ZCCHC2(+): 854bp before exon 9
##
    [4] IGR: 208Kb before PXDNL(-)
##
     [5] Intron of PAK7(-): 11Kb after exon 4
     [6] Intron of RALGAPB(+): 839bp after exon 15
##
                                                              score BPresult
##
                                         fusion
                                                  quality
##
                                       <factor> <numeric> <numeric> <integer>
##
     [1] -
                                                 1.000000 491.0000
                                                                            - 1
##
     [2]
                                                 0.994412
                                                           263.5191
                                                                             1
##
     [3] Protein fusion: in frame (ZCCHC2-DYM) 1.000000 238.0000
                                                                             1
##
                                                 1.000000
                                                            94.0000
                                                                            - 1
##
     [5] -
                                                 0.974021
                                                            54.5452
                                                                             1
##
     [6] Antisense fusion
                                                 1.000000
                                                            53.0000
                                                                             1
##
                   validation_result
                                            to.gr
                                                    rearrangements
##
                            <factor>
                                        <GRanges>
                                                       <character>
     [1] not_subjected_to_validation 18:58362793 intrachromosomal
##
##
     [2] not_subjected_to_validation 18:57443167 intrachromosomal
##
     [3] somatic
                                     18:58380361 intrachromosomal
     [4] not_subjected_to_validation 8:52186587 intrachromosomal
##
##
     [5] somatic
                                       20:9561906 interchromosomal
##
     [6] not_subjected_to_validation 20:36595752 interchromosomal
##
##
     seqinfo: 22 sequences from an unspecified genome
```

Here in this example, we use "intrachromosomal" to label rearrangement within the same chromosomes and use "interchromosomal" to label rearrangement in different chromosomes.

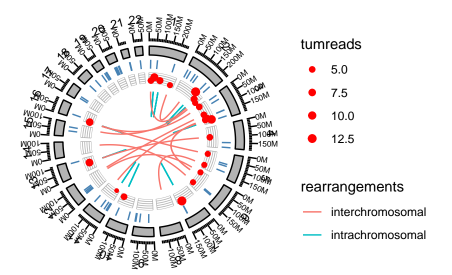
Get subset of links data for only one sample "CRC1"

```
gr.crc1 <- crc.gr[values(crc.gr)$individual == "CRC-1"]</pre>
```

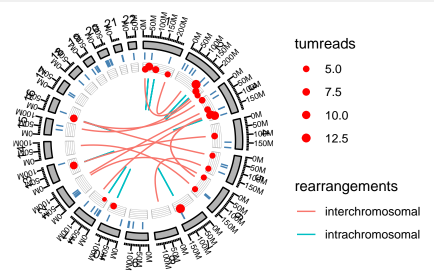
Ok, add a "point" track with grid background for rearrangement data and map 'y' to variable "score", map 'size' to variable "tumreads", rescale the size to a proper size range.



Finally, let's add links and map color to rearrangement types. Remember you need to specify 'linked.to' parameter to the column that contain end point of the data.



All those code could be simply constructed by following code



4.1.3 Complex arragnment of plots

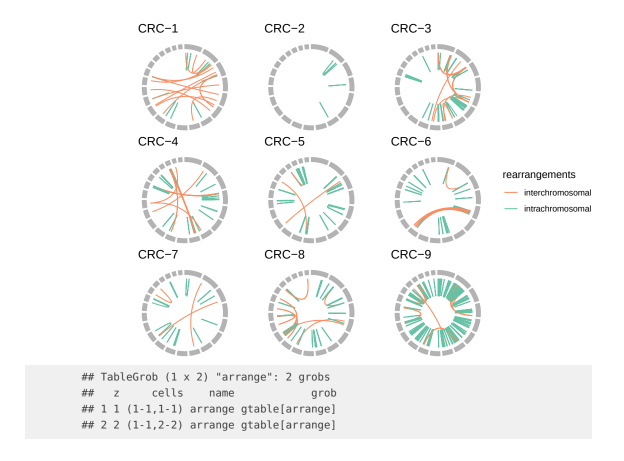
In this step, we are going to make multiple sample comparison, this may require some knowledge about package *grid* and *gridExtra*. We will introduce a more easy way to combine your graphics later after this.

We just want 9 single circular plots put together in one page, since we cannot keep too many tracks, we only keep ideogram and links. Here is one sample.

```
grl <- split(crc.gr, values(crc.gr)$individual)</pre>
## need "unit", load grid
library(grid)
crc.lst <- lapply(grl, function(gr.cur){</pre>
  print(unique(as.character(values(gr.cur)$individual)))
  cols <- RColorBrewer::brewer.pal(3, "Set2")[2:1]</pre>
  names(cols) <- c("interchromosomal", "intrachromosomal")</pre>
  p <- ggbio() + circle(gr.cur, geom = "link", linked.to = "to.gr",</pre>
                          aes(color = rearrangements)) +
                   circle(hg19sub, geom = "ideo",
                          color = "gray70", fill = "gray70") +
                   scale_color_manual(values = cols) +
                   labs(title = (unique(values(gr.cur)$individual))) +
                   theme(plot.margin = unit(rep(0, 4), "lines"))
})
## [1] "CRC-1"
## [1] "CRC-2"
## [1] "CRC-3"
## [1] "CRC-4"
## [1] "CRC-5"
## [1] "CRC-6"
## [1] "CRC-7"
## [1] "CRC-8"
## [1] "CRC-9"
```

We wrap the function in grid level to a more user-friendly high level function, called arrange GrobByParsingLegend. You can pass your ggplot2 graphics to this function, specify the legend you want to keep on the right, you can also specify the column/row numbers. Here we assume all plots we have passed follows the same color scale and have the same legend, so we only have to keep one legend on the right.

```
arrangeGrobByParsingLegend(crc.lst, widths = c(4, 1), legend.idx = 1, ncol = 3)
```



4.2 How to make grandlinear plots

4.2.1 Introduction

Let's use a subset of *PLINK* output (https://github.com/stephenturner/qqman/blob/master/plink.assoc.txt.gz) as our example test data.

```
snp <- read.table(system.file("extdata", "plink.assoc.sub.txt", package = "biovizBase"),</pre>
                   header = TRUE)
require(biovizBase)
gr.snp <- transformDfToGr(snp, seqnames = "CHR", start = "BP", width = 1)</pre>
head(gr.snp)
## GRanges object with 6 ranges and 10 metadata columns:
##
         seqnames
                               ranges strand |
                                                       CHR
                                                                   SNP
##
                           <IRanges>
                                      <Rle> |
                                                <integer> <character> <integer>
##
                 4 10794096-10794099
     [1]
                                                        4
                                                             rs9291494 10794096
     [2]
                14 55853742-55853755
                                                        14
                                                             rs1152481
                                                                         55853742
##
     [3]
                 6 55188853-55188858
                                                        6
                                                             rs3134708 55188853
     [4]
               17
                     4146033-4146049
                                                       17
                                                             rs2325988
                                                                          4146033
                19 46089501-46089519
##
     [5]
                                                       19
                                                             rs8103444 46089501
                            107051695
                                                            rs12072065 107051695
                             F_A
##
                   Α1
                                                             CHISQ
                                                                                      0R
                                       \mathsf{F}_{-}\mathsf{U}
                                                      A2
```

```
<character> <numeric> <numeric> <numeric> <numeric> <numeric> <numeric> <numeric>
##
    [1]
                 G
                      0.3061
                               0.1341
                                                   7.5070 0.006147
                                                                     2.8480
                                              Α
                                                   1.1030 0.293600
                 G
                      0.3542
                               0.2805
                                                                     1.4070
##
    [2]
                                              Α
##
    [3]
                 C
                     0.2500
                               0.2875
                                              Α
                                                   0.3135 0.575500
                                                                     0.8261
                               0.2317
                                                   0.1323 0.716100
##
    [4]
                 G 0.2551
                                              Α
                                                                     1.1360
    [5]
                 C 0.3980
                               0.2927
                                              Α
                                                   2.1750 0.140300
                                                                     1.5970
##
##
    [6]
                 0
                      0.0000
                               0.0000
                                              C
                                                       NA
                                                                NA
##
    seqinfo: 22 sequences from an unspecified genome; no seqlengths
## change the segname order
require(GenomicRanges)
gr.snp <- keepSeqlevels(gr.snp, as.character(1:22))</pre>
seqlengths(gr.snp)
## 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22
## need to assign seglengths
data(ideoCyto, package = "biovizBase")
seqlengths(gr.snp) <- as.numeric(seqlengths(ideoCyto$hg18)[1:22])</pre>
## remove missing
gr.snp <- gr.snp[!is.na(gr.snp$P)]</pre>
## transform pvalue
values(gr.snp)$pvalue <- -log10(values(gr.snp)$P)</pre>
head(gr.snp)
## GRanges object with 6 ranges and 11 metadata columns:
##
        segnames
                         ranges strand | CHR
                                                         SNP
##
           <Rle>
                       <IRanges> <Rle> | <integer> <character> <integer>
##
    [1]
             4 10794096-10794099
                                   * | 4 rs9291494 10794096
##
    [2]
             14 55853742-55853755
                                              14 rs1152481 55853742
                                     * |
##
    [3]
             6 55188853-55188858
                                    * |
                                               6 rs3134708 55188853
##
    [4]
             17 4146033-4146049
                                     *
                                               17
                                                    rs2325988
                                                              4146033
             19 46089501-46089519
                                              19
                                                    rs8103444 46089501
##
    [5]
                                     *
              9 81517907-81517915
##
                                    *
                                              9
                                                       rs2591 81517907
                        F_A
                                F_{-}U
                                                                 Р
##
                A1
                                             A2
                                                    CHISQ
##
        <character> <numeric> <numeric> <numeric> <numeric> <numeric> <numeric> <numeric>
                     0.3061 0.13410 A
                                                   7.5070 0.006147
##
    [1]
                 G
                                                                     2.8480
##
    [2]
                 G 0.3542 0.28050
                                              Α
                                                   1.1030 0.293600
                                                                     1.4070
                      0.2500 0.28750
##
    [3]
                 C
                                              Α
                                                   0.3135 0.575500
                                                                     0.8261
                 G 0.2551 0.23170
##
    [4]
                                              A 0.1323 0.716100
                                                                     1.1360
                 C
                     0.3980 0.29270
                                             A 2.1750 0.140300
                                                                     1.5970
##
    [5]
##
    [6]
                 C
                     0.1042 0.04878
                                             T 1.8720 0.171200
                                                                     2.2670
##
           pvalue
##
        <numeric>
##
    [1] 2.211337
    [2] 0.532244
##
    [3] 0.239955
##
##
    [4] 0.145026
    [5] 0.852942
    [6] 0.766496
##
```

```
## seqinfo: 22 sequences from an unspecified genome
## done
```

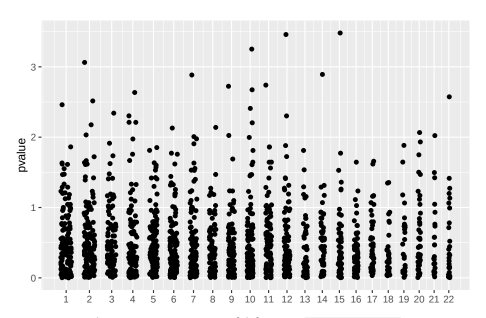
The data is ready, we need to pay attention

- if seqlengths is missing, we use data range, so the chromosome length is not accurate
- use seglevel to control order of chromosome

4.2.2 Corrdinate genome

In autoplot, argument coord is just used to transform the data, after that, you can use it as common GRanges, all other geom/stat works for it.

```
autoplot(gr.snp, geom = "point", coord = "genome", aes(y = pvalue))
```



However, we recommend you to use more powerful function plotGrandLinear to generate manhattan plot introduced in next section.

4.2.3 Convenient plotGrandLinear function

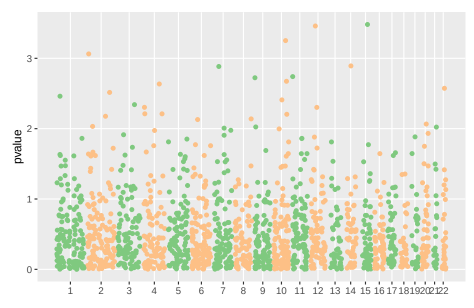
For *Manhattan plot*, we have a function called plotGrandLinear. aes(y =) is required to indicate the y value, e.g. p-value.

Color mapping is automatically figured out by ggbio following the rules

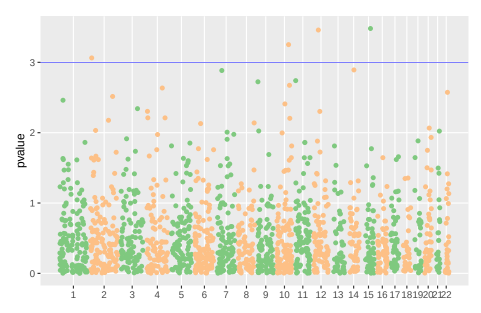
- if color present in aes(), like aes(color = seqnames), it will assume it's mapping to data column called 'seqnames'.
- if color is not wrapped in aes(), then this function will **recylcle** them to all chromosomes.
- if color is single character representing color, then just use one arbitrary color.

Let's test some examples for controling colors.

```
plotGrandLinear(gr.snp, aes(y = pvalue), color = c("#7fc97f", "#fdc086"))
```

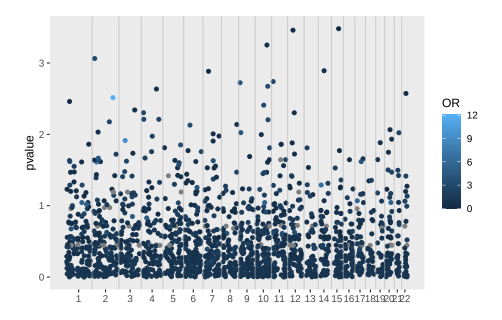


Let's add a cutoff line



Sometimes you use color to mapping other varibles so you may need a different to separate chromosomes.

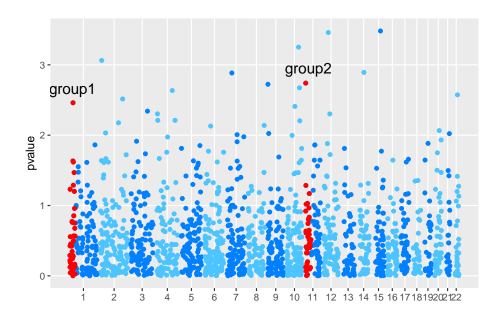
```
plotGrandLinear(gr.snp, aes(y = pvalue, color = OR), spaceline = TRUE, legend = TRUE)
```



4.2.4 How to highlight some points?

You can provide a highlight *GRanges*, and each row highlights a set of overlaped snps, and labeled by rownames or certain columns, there is more control in the function as parameters, with prefix highlight.*, so you could control color, label size and color, etc.

```
gro <- GRanges(c("1", "11"), IRanges(c(100, 2e6), width = 5e7))
names(gro) <- c("group1", "group2")
plotGrandLinear(gr.snp, aes(y = pvalue), highlight.gr = gro)</pre>
```



4.3 How to make stacked karyogram overview plots

4.3.1 Introduction

A karyotype is the number and appearance of chromosomes in the nucleus of a eukaryotic cell². It's one kind of overview when we want to show distribution of certain events on the genome, for example, binding sites for certain protein, even compare them across samples as example shows in this section.

²http://en.wikipedia.org/wiki/Kar

GRanges and Seqinfo objects are an ideal container for storing data needed for karyogram plot. Here is the strategy we used for generating ideogram templates.

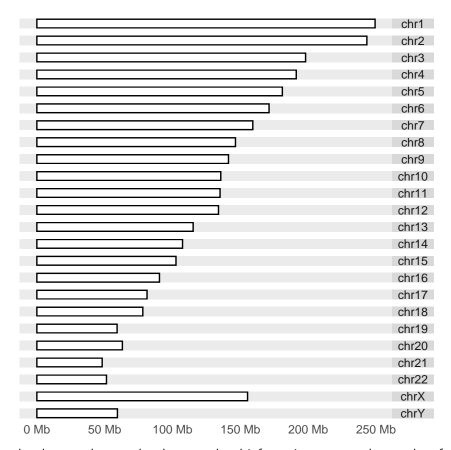
- Althouth seqlengths is not required, it's highly recommended for plotting karyogram.
 If a GRanges object contains seqlengths, we know exactly how long each chromosome is, and will use this information to plot genome space, particularly we plot all levels included in it, NOT JUST data space.
- If a GRanges has no seqlengths, we will issue a warning and try to estimate the chromosome lengths from data included. This is NOT accurate most time, so please pay attention to what you are going to visualize and make sure set seqlengths before hand.

4.3.2 Create karyogram temlate

Let's first introduce how to use autoplot to generate karyogram graphic.

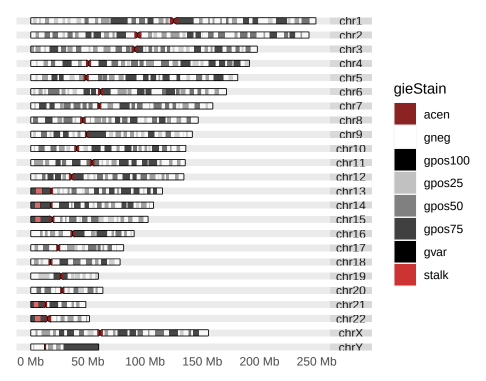
The most easy one is to just plot Seqinfo by using autoplot, if your *GRanges* object has seqinfo with seqlengths information. Then you add data layer later.

```
data(ideoCyto, package = "biovizBase")
autoplot(seqinfo(ideoCyto$hg19), layout = "karyogram")
```



To show cytobands, your data need to have cytoband information, we stored some data for you, including hg19, hg18, mm10, mm9.

```
## turn on cytobands if present
biovizBase::isIdeogram(ideoCyto$hg19)
## [1] TRUE
autoplot(ideoCyto$hg19, layout = "karyogram", cytobands = TRUE)
```



To change order or only show a subset of the karyogram, you have to manipulate seqlevels, please check out manual for keepSeqlevels, seqlevels in *GenomicRanges* package for more information. Or you could read the example below.

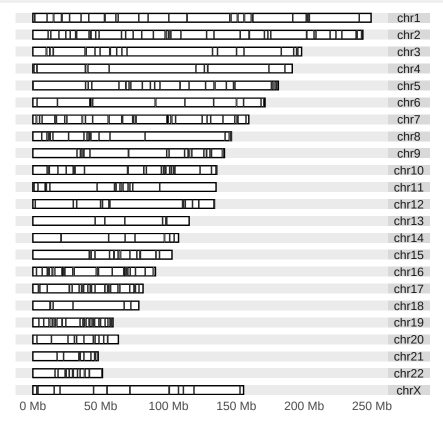
4.3.3 Add data on karyogram layout

If you have single data set stored as *GRanges* to show on a karyogram layout, autoplot function is enough for you to plot the data on it.

We use a default data in package *biovizBase*, which is a subset of RNA editing set in human. The data involved in this GRanges is sparse, so we cannot simply use it to make karyogram template, otherwise, the estimated chromosome lengths will be very rough and inaccurate. So what we need to do first is to *add seglength information to this object*.

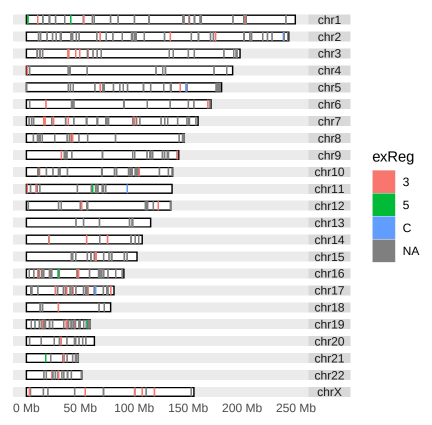
```
data(darned_hg19_subset500, package = "biovizBase")
dn <- darned_hg19_subset500</pre>
library(GenomicRanges)
seqlengths(dn)
    chr1 chr10 chr11 chr12 chr13 chr14 chr15 chr16 chr17 chr18 chr19
                                                                           chr2 chr20
##
                                                                             NA
      NA
            NA
                   NA
                         NA
                                NA
                                      NA
                                             NA
                                                   NA
                                                          NA
                                                                NA
                                                                       NA
                                                                                    NA
## chr21 chr22 chr3
                              chr5
                                          chr7
                                                 chr8
                                                        chr9
                       chr4
                                    chr6
                                                              chrX
##
      NA
            NA
                   NA
                         NA
                                NA
                                      NA
                                             NA
                                                   NA
                                                          NA
                                                                NA
## add seglengths
## we have seglegaths information in another data set
seqlengths(dn) <- seqlengths(ideoCyto$hg19)[names(seqlengths(dn))]</pre>
## then we change order
dn <- keepSeqlevels(dn, paste0("chr", c(1:22, "X")))</pre>
```

```
seqlengths(dn)
##
                   chr2
                             chr3
                                        chr4
                                                  chr5
                                                             chr6
                                                                       chr7
                                                                                  chr8
        chr1
## 249250621 243199373 198022430 191154276 180915260 171115067 159138663 146364022
                  chr10
        chr9
                            chr11
                                       chr12
                                                 chr13
                                                            chr14
                                                                      chr15
                                                                                 chr16
  141213431 135534747 135006516 133851895 115169878 107349540 102531392
                                                                              90354753
##
       chr17
                  chr18
                            chr19
                                       chr20
                                                 chr21
                                                            chr22
                                                                       chrX
##
    81195210
             78077248
                        59128983
                                   63025520
                                              48129895
                                                        51304566 155270560
autoplot(dn, layout = "karyogram")
```



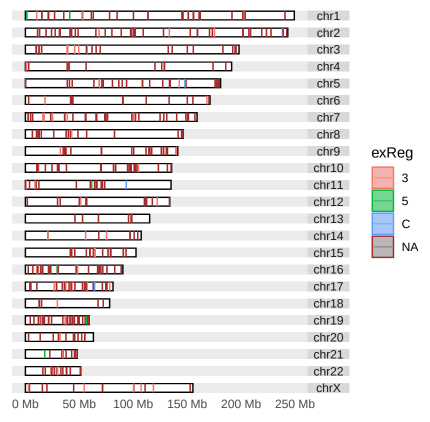
Then we take one step further, the power of *ggplot2* or *ggbio* is the flexible multivariate data mapping ability in graphics, make data exploration much more convenient. In the following example, we are trying to map a categorical variable 'exReg' to color, this variable is included in the data, and have three levels, '3' indicate 3' utr, '5' means 5' utr and 'C' means coding region. We have some missing values indicated as NA, in default, it's going to be shown in gray color, and keep in mind, since the basic geom(geometric object) is rectangle, and genome space is very large, so change both color/fill color of the rectangle to specify both border and filled color is necessary to get the data shown as different color, otherwise if the region is too small, border color is going to override the fill color.

```
## since default is geom rectangle, even though it's looks like segment
## we still use both fill/color to map colors
autoplot(dn, layout = "karyogram", aes(color = exReg, fill = exReg))
```



Or you can set the missing value to particular color yo u want (NA values is not shown on the legend).

```
## since default is geom rectangle, even though it's looks like segment
## we still use both fill/color to map colors
autoplot(dn, layout = "karyogram", aes(color = exReg, fill = exReg), alpha = 0.5) +
scale_color_discrete(na.value = "brown")
```



Well, sometimes we have too many values, we want to separate them by groups and show them at diffent height, below is a hack for that purpose and in next section, we will introduce a more flexible and general way to add data layer by layer.

Template chromosome y limits is [0, 10], that's why this hack works

4.3.4 Add more data using layout_karyogram function

In this section, a lower level function layout_karyogram is going to be introduced. This is convenient API for constructing karyogram plot and adding more data layer by layer. Function ggplot is just to create blank object to add layer on.

You need to pay attention to

• when you add plots layer by layer, seqnames of different data must be the same to make sure the data are mapped to the same chromosome. For example, if you name chromosome following schema like chr1 and use just number 1 to name other data, they will be treated as different chromosomes.

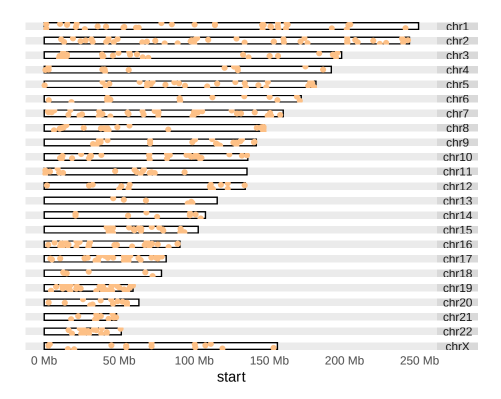
- cannot use the same aesthetics mapping multiple time for different data. For example, if you have used aes(color =), for one data, you cannot use aes(color =) anymore for mapping variables from other add-on data, this is currently not allowed in ggplot2, even though you expect multiple color legend shows up, this is going to confuse people which is which. HOWEVER, color or fill without aes() wrap around, is allowed for any track, it's set single arbitrary color.
- Default rectangle y range is [0, 10], so when you add on more data layer by layer on existing graphics, you can use ylim to control how to normalize your data and plot it relative to chromosome space. For example, with default, chromosome space is plotted between y [0, 10], if you use ylim = c(10, 20), you will stack data right above each chromosomes and with equal width. For geom like 'point', which you need to specify 'y' value in aes(), we will add 5% margin on top and at bottom of that track.

Many times we overlay different datas sets, so let's break down the previous samples into 4 groups and treat them as different data and build them layer by layer, assign the color by hand. You could use ylim to control where they are ploted.

```
## prepare the data
dn3 <- dn.nona[dn.nona$exReg == '3']
dn5 <- dn.nona[dn.nona$exReg == '5']
dnC <- dn.nona[dn.nona$exReg == 'C']
dn.na <- dn[is.na(dn$exReg)]
## now we have 4 different data sets
autoplot(seqinfo(dn3), layout = "karyogram") +
    layout_karyogram(data = dn3, geom = "rect", ylim = c(0, 10/3), color = "#7fc97f") +
    layout_karyogram(data = dn5, geom = "rect", ylim = c(10/3, 10/3*2), color = "#beaed4") +
    layout_karyogram(data = dnC, geom = "rect", ylim = c(10/3*2, 10), color = "#fdc086") +
    layout_karyogram(data = dn.na, geom = "rect", ylim = c(10, 10/3*4), color = "brown")</pre>
```

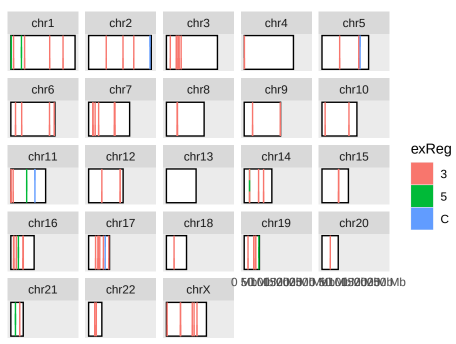
```
chr1
                                                      chr2
                                                      chr3
                                                      chr4
                                                      chr5
                                                      chr6
                                                      chr7
                                                      chr8
                                                      chr9
                                                      chr10
                                                      chr11
                                                      chr12
                                                      chr13
                                                      chr14
                                                      chr15
                                                      chr16
                                                      chr17
                                                      chr18
                                                      chr19
                                                      chr20
                                                      chr21
                                                      chr22
                                                      chrX
         50 Mb
                                      200 Mb
0 Mb
                  100 Mb
                            150 Mb
                                                250 Mb
                        start
```

What's more, you could even chagne the geom for those data



4.3.5 More flexible layout of karyogram

p.ylim + facet_wrap(~seqnames)



Chapter 5

Link ranges to your data

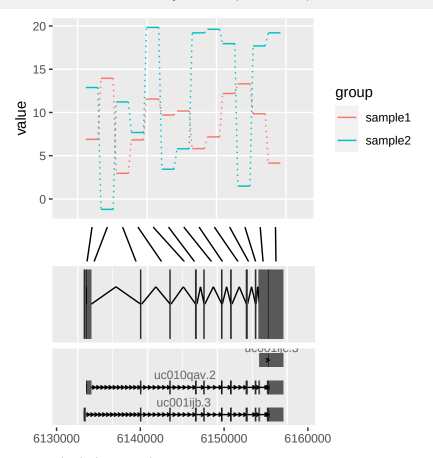
Plot GRanges object structure and linked to a even spaced paralell coordinates plot which represting the data in elementeMetadata.

```
library(TxDb.Hsapiens.UCSC.hg19.knownGene)
library(ggbio)
data(genesymbol, package = "biovizBase")
txdb <- TxDb.Hsapiens.UCSC.hg19.knownGene
model <- exonsBy(txdb, by = "tx")</pre>
model17 <- subsetByOverlaps(model, genesymbol["RBM17"])</pre>
exons <- exons(txdb)
exon17 <- subsetByOverlaps(exons, genesymbol["RBM17"])</pre>
## reduce to make sure there is no overlap
## just for example
exon.new <- reduce(exon17)
## suppose
values(exon.new)$sample1 <- rnorm(length(exon.new), 10, 3)</pre>
values(exon.new)$sample2 <- rnorm(length(exon.new), 10, 10)</pre>
values(exon.new)$score <- rnorm(length(exon.new))</pre>
values(exon.new)$significant <- sample(c(TRUE,FALSE), size = length(exon.new),replace = TRUE)</pre>
## data ready
exon.new
## GRanges object with 13 ranges and 4 metadata columns:
##
          seqnames ranges strand | sample1 sample2
                     <IRanges> <Rle> | <numeric> <numeric> <numeric>
##
             <Rle>
      [1] chr10 6130949-6131156 + | 6.89613 12.88536 -1.024697
##
      [2] chr10 6131309-6131934
                                       + | 13.95211 -1.20274 -0.394437
     [3] chr10 6139011-6139151 + | 2.96625 11.22116 0.353247  
[4] chr10 6143234-6143350 + | 6.83044 7.68600 1.224819
##
          chr10 6146894-6147060 + | 11.55350 19.83990 0.167351
     [5]
           chr10 6154173-6154324 + |
##
                                                   . . .
                                             7.17512 19.61853 -1.291514
     [9]
                                       + | 12.19564 17.95885 0.405627
##
     [10]
          chr10 6155471-6155544
                                       [11] chr10 6156012-6156110
            chr10 6156126-6157274 + | 9.85528 17.69333 0.726902
chr10 6157416-6159422 + | 4.14673 19.20935 -0.554322
##
    [12]
    [13]
```

```
##
          significant
##
             <logical>
                  TRUE
##
      [1]
##
                  TRUE
      [2]
##
                 FALSE
      [3]
                 FALSE
##
      [4]
##
                  TRUE
      [5]
##
##
      [9]
                  TRUE
##
     [10]
                 FALSE
##
     [11]
                 FALSE
##
                  TRUE
     [12]
##
     [13]
                  TRUE
##
     seqinfo: 93 sequences (1 circular) from hg19 genome
```

Make the plots, you can pass a list of annotation tracks too.

```
p17 <- autoplot(txdb, genesymbol["RBM17"])
plotRangesLinkedToData(exon.new, stat.y = c("sample1", "sample2"), annotation = list(p17))</pre>
```



For more information, check the manual.

Chapter 6

Miscellaneous

Every plot object produced by *ggplot2* is essentially a *ggplot2* object, so you could use all the tricks you know with *ggplot2* on *ggbio* plots too, including scales, colors, themes, etc.

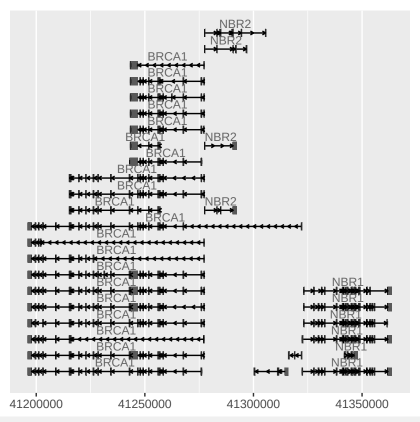
6.1 Themes

In ggbio, we developed some more themes to make things easier.

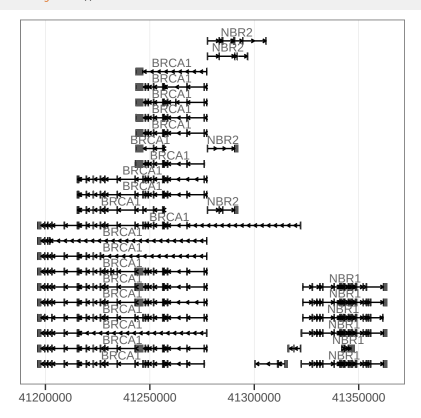
6.1.1 Plot theme

Plot level themes are like any other themes defined in ggplot2, simply apply it to a plot.

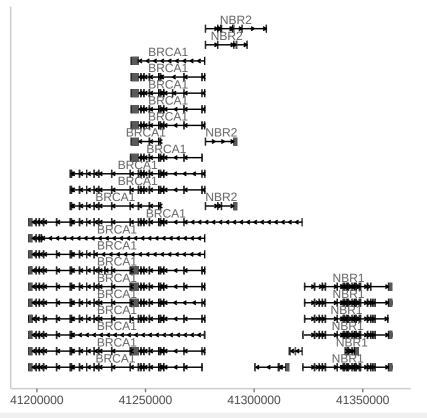
p.txdb



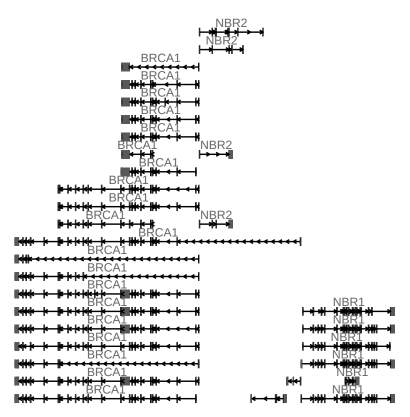
p.txdb + theme_alignment()



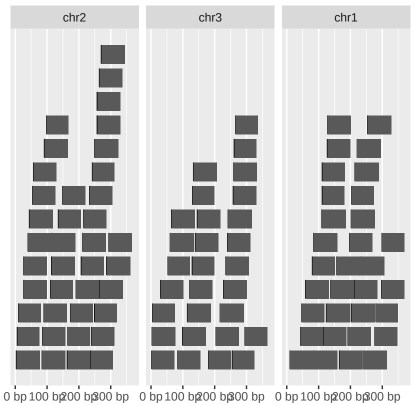




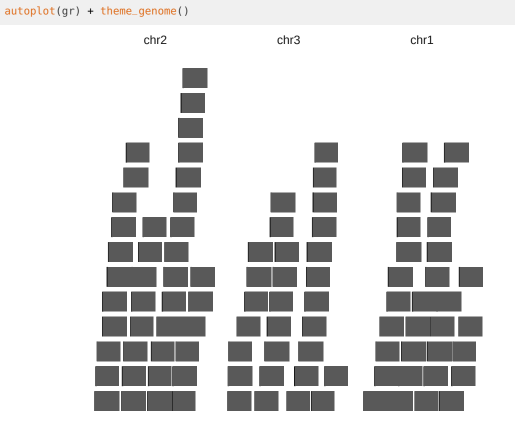
p.txdb + theme_null()



When you have multiple chromosomes encoded in seqnames, you could use theme_genome to make a 'fake' linear view of genome coordinates quickly by applying this theme, because it's not equal to chromosome lengths, it's simply



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6.1.2 Track theme

Track level themes are more complex, it controls whole looking of the tracks, it's essentially a theme object with some attributes controlling the tracks appearance.

See how we make a template, you could customize in the same way

```
theme_tracks_sunset

## function (bg = "#fffedb", alpha = 1, ...)

## {

## res <- theme_clear(grid.x.major = FALSE, ...)

## attr(res, "track.plot.color") <- sapply(bg, scales::alpha,

## alpha)

## attr(res, "track.bg.color") <- bg

## attr(res, "label.text.color") <- "white"

## attr(res, "label.bg.fill") <- "#a52a2a"

## res

## }

## <bytecode: 0x56526f398f18>

## <environment: namespace:ggbio>
```

The attributes you could control is basically passed to tracks() constructor, including

label.bg.color	character	
label.bg.fill	character	
label.text.color	character	
label.text.cex	numeric	
label.text.angle	numeric	
track.plot.color	character_OR_NULL	
track.bg.color	character_OR_NULL	
label.width	unit	

Table 6.1: tracks attributes

Chapter 7

Session Information

```
sessionInfo()
## R version 4.3.1 (2023-06-16)
## Platform: x86_64-pc-linux-gnu (64-bit)
## Running under: Ubuntu 22.04.3 LTS
##
## Matrix products: default
## BLAS: /home/biocbuild/bbs-3.18-bioc/R/lib/libRblas.so
## LAPACK: /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.10.0
## locale:
## [3] LC_TIME=en_GB
## [5] LC_MONETARY=en_US.UTF-8 LC_MESSAGES=en_US.UTF-8
## [7] LC_PAPER=en_US.UTF-8 LC_NAME=C
## [9] LC_ADDRESS=C
                                LC_TELEPHONE=C
## [11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
## time zone: America/New_York
## tzcode source: system (glibc)
## attached base packages:
## [1] grid stats4 stats graphics grDevices utils datasets
## [8] methods base
##
## other attached packages:
## [1] VariantAnnotation_1.48.0
## [2] Rsamtools_2.18.0
## [3] SummarizedExperiment_1.32.0
## [4] MatrixGenerics_1.14.0
## [5] matrixStats_1.0.0
## [6] BSgenome.Hsapiens.UCSC.hg19_1.4.3
## [7] BSgenome_1.70.1
## [8] rtracklayer_1.62.0
## [9] BiocIO_1.12.0
```

```
## [10] Biostrings_2.70.1
## [11] XVector_0.42.0
## [12] biovizBase_1.50.0
## [13] EnsDb.Hsapiens.v75_2.99.0
## [14] ensembldb_2.26.0
## [15] AnnotationFilter_1.26.0
## [16] Homo.sapiens_1.3.1
## [17] TxDb.Hsapiens.UCSC.hg19.knownGene_3.2.2
## [18] org.Hs.eg.db_3.18.0
## [19] GO.db_3.18.0
## [20] OrganismDbi_1.44.0
## [21] GenomicFeatures_1.54.1
## [22] AnnotationDbi_1.64.0
## [23] Biobase_2.62.0
## [24] GenomicRanges_1.54.1
## [25] GenomeInfoDb_1.38.0
## [26] IRanges_2.36.0
## [27] S4Vectors_0.40.1
## [28] ggbio_1.50.0
## [29] ggplot2_3.4.4
## [30] BiocGenerics_0.48.1
## [31] knitr_1.45
##
## loaded via a namespace (and not attached):
## [1] RColorBrewer_1.1-3
                                 rstudioapi_0.15.0
                                                           magrittr_2.0.3
## [4] farver_2.1.1
                                                           zlibbioc_1.48.0
                                 rmarkdown_2.25
                                                           RCurl_1.98-1.12
## [7] vctrs_0.6.4
                                 memoise_2.0.1
## [10] base64enc_0.1-3
                                 htmltools_0.5.6.1
                                                          S4Arrays_1.2.0
## [13] progress_1.2.2
                                 curl_5.1.0
                                                           SparseArray_1.2.0
## [16] Formula_1.2-5
                                 htmlwidgets_1.6.2
                                                           plyr_1.8.9
## [19] cachem_1.0.8
                                 GenomicAlignments_1.38.0 lifecycle_1.0.3
## [22] pkgconfig_2.0.3
                                 Matrix_1.6-1.1
                                                          R6_2.5.1
## [25] fastmap_1.1.1
                                 GenomeInfoDbData_1.2.11 digest_0.6.33
## [28] colorspace_2.1-0
                                 GGally_2.1.2
                                                           reshape_0.8.9
## [31] Hmisc_5.1-1
                                 RSQLite_2.3.2
                                                           labeling_0.4.3
## [34] filelock_1.0.2
                                 fansi_1.0.5
                                                           httr_1.4.7
## [37] abind_1.4-5
                                                           bit64_4.0.5
                                 compiler_4.3.1
## [40] withr_2.5.2
                                 htmlTable_2.4.2
                                                           backports_1.4.1
## [43] BiocParallel_1.36.0
                                 DBI_1.1.3
                                                          highr_0.10
## [46] biomaRt_2.58.0
                                 rappdirs_0.3.3
                                                           DelayedArray_0.28.0
## [49] rjson_0.2.21
                                 tools_4.3.1
                                                           foreign_0.8-85
## [52] nnet_7.3-19
                                                           restfulr_0.0.15
                                 glue_1.6.2
## [55] checkmate_2.3.0
                                 cluster_2.1.4
                                                           reshape2_1.4.4
## [58] generics_0.1.3
                                 gtable_0.3.4
                                                           data.table_1.14.8
## [61] hms_1.1.3
                                 xml2_1.3.5
                                                           utf8_1.2.4
## [64] pillar_1.9.0
                                 stringr_1.5.0
                                                           dplyr_1.1.3
## [67] BiocFileCache_2.10.1
                                 lattice_0.22-5
                                                          bit_4.0.5
## [70] tidyselect_1.2.0
                                 RBGL_1.78.0
                                                           gridExtra_2.3
## [73] ProtGenerics_1.34.0
                                 xfun_0.41
                                                           stringi_1.7.12
## [76] lazyeval_0.2.2
                                 yaml_2.3.7
                                                           evaluate_0.23
                                                           BiocManager_1.30.22
## [79] codetools_0.2-19
                                 tibble_3.2.1
```

## [82] graph_1.80.0	cli_3.6.1	$rpart_4.1.21$
## [85] munsell_0.5.0	$dichromat_2.0-0.1$	Rcpp_1.0.11
## [88] dbplyr_2.4.0	png_0.1-8	XML_3.99-0.14
## [91] parallel_4.3.1	$blob_1.2.4$	prettyunits $_{-}1.2.0$
## [94] bitops_1.0-7	scales_1.2.1	crayon_1.5.2
## [97] BiocStyle_2.30.0	rlang_1.1.1	KEGGREST_1.42.0