# NGS Analysis Basics

Author: Thomas Girke
Last update: 25 April, 2018

#### Overview

## Sequence Analysis in R and Bioconductor

#### R. Base

• Some basic string handling utilities. Wide spectrum of numeric data analysis tools.

#### Bioconductor

Bioconductor packages provide much more sophisticated string handling utilities for sequence analysis (Lawrence et al. 2013, Huber et al. (2015)).

- Biostrings: general sequence analysis environment
- ShortRead: pipeline for short read data
- IRanges: low-level infrastructure for range data
- GenomicRanges: high-level infrastructure for range data
- GenomicFeatures: managing transcript centric annotations
- GenomicAlignments: handling short genomic alignments
- Rsamtools: interface to samtools, bcftools and tabix
- BSgenome: genome annotation data
- biomaRt: interface to BioMart annotations
- rtracklayer: Annotation imports, interface to online genome browsers
- HelloRanges: Bedtools semantics in Bioc's Ranges infrastructure

## Package Requirements

Several Bioconductor packages are required for this tutorial. To install them, execute the following lines in the R console. Please also make sure that you have a recent R version installed on your system. R versions 3.3.x or higher are recommended.

```
source("https://bioconductor.org/biocLite.R")
biocLite(c("Biostrings", "GenomicRanges", "GenomicRanges", "rtracklayer", "systemPipeR", "seqLogo", "Sh
```

## Strings in R Base

## Basic String Matching and Parsing

## String matching

```
Generate sample sequence data set
```

```
myseq <- c("ATGCAGACATAGTG", "ATGAACATAGATCC", "GTACAGATCAC")
```

String searching with regular expression support

```
myseq[grep("ATG", myseq)]
## [1] "ATGCAGACATAGTG" "ATGAACATAGATCC"
Searches myseq for first match of pattern "AT"
pos1 <- regexpr("AT", myseq)</pre>
as.numeric(pos1); attributes(pos1) $match.length # Returns position information of matches
## [1] 1 1 7
## [1] 2 2 2
Searches myseq for all matches of pattern "AT"
pos2 <- gregexpr("AT", myseq)</pre>
as.numeric(pos2[[1]]); attributes(pos2[[1]]) $match.length # Returns positions of matches in first seque
## [1] 1 9
## [1] 2 2
String substitution with regular expression support
gsub("^ATG", "atg", myseq)
## [1] "atgCAGACATAGTG" "atgAACATAGATCC" "GTACAGATCAC"
Positional parsing
nchar(myseq) # Computes length of strings
## [1] 14 14 11
substring(myseq[1], c(1,3), c(2,5)) # Positional parsing of several fragments from one string
## [1] "AT" "GCA"
substring(myseq, c(1,4,7), c(2,6,10)) # Positional parsing of many strings
## [1] "AT"
              "AAC" "ATCA"
Random Sequence Generation
Random DNA sequences of any length
rand <- sapply(1:100, function(x) paste(sample(c("A", "T", "G", "C"), sample(10:20), replace=T), collapse=
rand[1:3]
## [1] "CAGTTCATGA"
                             "TATTTTCACCGCATAAAA" "TTGTAGTCGTGAAAT"
Count identical sequences
```

table(c(rand[1:4], rand[1]))

```
##
## CAGTTCATGA GATAGTACAC TATTTTCACCGCATAAAA TTGTAGTCGTGAAAT
## 2 1 1 1
```

#### Extract reads from reference

Note: this requires Biostrings package.

```
library(Biostrings)
ref <- DNAString(paste(sample(c("A","T","G","C"), 100000, replace=T), collapse=""))
randstart <- sample(1:(length(ref)-15), 1000)
randreads <- Views(ref, randstart, width=15)
rand_set <- DNAStringSet(randreads)
unlist(rand_set)</pre>
```

```
## 15000-letter "DNAString" instance
## seq: ACATGATTTCGTGCTTGGTCGCGTTGTCTCAATACGGGGCGTGTAT...AGTCGAAGGAGCTCAACGTCCTTAAGCTTTCAACAATCGTTATTTG
```

## Sequences in Bioconductor

## Important Data Objects of Biostrings

#### XString for single sequence

DNAString: for DNA
RNAString: for RNA
AAString: for amino acid
BString: for any string

## XStringSet for many sequences

'DNAStringSet": for DNA
RNAStringSet: for RNA
AAStringSet: for amino acid
BStringSet: for any string

#### QualityScaleXStringSet for sequences with quality data

QualityScaledDNAStringSet: for DNA
 QualityScaledRNAStringSet: for RNA
 QualityScaledAAStringSet: for amino acid
 QualityScaledBStringSet: for any string

#### Sequence Import and Export

Download the following sequences to your current working directory and then import them into R: ftp://ftp.ncbi.nlm.nih.gov/genomes/archive/old\_genbank/Bacteria/Halobacterium\_sp\_uid217/AE004437.ffn

```
dir.create("data", showWarnings = FALSE)
# system("wqet ftp://ftp.ncbi.nlm.nih.gov/qenomes/archive/old_qenbank/Bacteria/Halobacterium_sp_uid217/
download.file("ftp://ftp.ncbi.nlm.nih.gov/genomes/archive/old_genbank/Bacteria/Halobacterium_sp_uid217/
Import FASTA file with readDNAStringSet
myseq <- readDNAStringSet("data/AE004437.ffn")</pre>
myseq[1:3]
     A DNAStringSet instance of length 3
##
##
       width seq
## [1] 1206 ATGACTCGGCGGTCTCGTGTCGGTGCCGGCCTC...GTCGTCGTTGTTCGACGCTGGCGGAACCCATGA gi|12057215|gb|AE...
## [2]
         666 ATGAGCATCATCGAACTCGAAGGCGTGGTCAAA...GTCAACCTCGTCGATGGGGTGTTACACACGTGA gi|12057215|gb|AE...
## [3] 1110 ATGGCGTGGCGGAACCTCGGGCGGAACCGCGTG...AACGATCCGCCCGTCGAGGCGCTCGGCGAATGA gi|12057215|gb|AE...
Subset sequences with regular expression on sequence name field
sub <- myseq[grep("99.*", names(myseq))]</pre>
length(sub)
## [1] 170
Export subsetted sequences to FASTA file
writeXStringSet(sub, file="./data/AE004437sub.ffn", width=80)
```

Now inspect exported sequence file  ${\tt AE004437sub.ffn}$  in a text editor

## Working with XString Containers

The XString stores the different types of biosequences in dedicated containers

```
library(Biostrings)
d <- DNAString("GCATAT-TAC")</pre>
     10-letter "DNAString" instance
## seq: GCATAT-TAC
d[1:4]
     4-letter "DNAString" instance
## seq: GCAT
RNA sequences
r <- RNAString("GCAUAU-UAC")
r <- RNAString(d) # Converts d to RNAString object
     10-letter "RNAString" instance
## seq: GCAUAU-UAC
Protein sequences
p <- AAString("HCWYHH")</pre>
р
     6-letter "AAString" instance
## seq: HCWYHH
```

Any type of character strings

```
b <- BString("I store any set of characters. Other XString objects store only the IUPAC characters.")
b
## 85-letter "BString" instance
## seq: I store any set of characters. Other XString objects store only the IUPAC characters.</pre>
```

## Working with XStringSet Containers

XStringSet containers allow to store many biosequences in one object

```
dset <- DNAStringSet(c("GCATATTAC", "AATCGATCC", "GCATATTAC"))</pre>
names(dset) <- c("seq1", "seq2", "seq3") # Assigns names</pre>
dset[1:2]
##
     A DNAStringSet instance of length 2
##
       width seq
                                                                                        names
## [1]
           9 GCATATTAC
                                                                                        seq1
## [2]
           9 AATCGATCC
                                                                                        seq2
Important utilities for XStringSet containers
width(dset) # Returns the length of each sequences
## [1] 9 9 9
d <- dset[[1]] # The [[ subsetting operator returns a single entry as XString object
dset2 <- c(dset, dset) # Appends/concatenates two XStringSet objects</pre>
dsetchar <- as.character(dset) # Converts XStringSet to named vector</pre>
dsetone <- unlist(dset) # Collapses many sequences to a single one stored in a DNAString container
```

Sequence subsetting by positions:

```
DNAStringSet(dset, start=c(1,2,3), end=c(4,8,5))
```

## Multiple Alignment Class

The XMultipleAlignment class stores the different types of multiple sequence alignments:

```
## [6] ------ gi|148540149|ref|...
## [7] ------ CGGCTCCGCAGCGCCTCACTCG...----- gi|45383056|ref|N...
## [8] GGGGGAGACTTCAGAAGTTGTTCTCCTCTCCGCTGA...----- gi|213515133|ref|...
```

## **Basic Sequence Manipulations**

#### Reverse and Complement

```
randset <- DNAStringSet(rand)</pre>
complement(randset[1:2])
     A DNAStringSet instance of length 2
##
       width seq
## [1]
          10 GTCAAGTACT
## [2]
          18 ATAAAAGTGGCGTATTTT
reverse(randset[1:2])
##
     A DNAStringSet instance of length 2
##
       width seq
          10 AGTACTTGAC
## [1]
## [2]
          18 AAAATACGCCACTTTTAT
reverseComplement(randset[1:2])
     A DNAStringSet instance of length 2
##
       width seq
## [1]
          10 TCATGAACTG
## [2]
          18 TTTTATGCGGTGAAAATA
```

#### Translate DNA into Protein

```
translate(randset[1:2])

## Warning in .Call2("DNAStringSet_translate", x, skip_code, dna_codes[codon_alphabet], : in 'x[[1]]':

## last base was ignored

## A AAStringSet instance of length 2

## width seq

## [1] 3 QFM
```

### Pattern Matching

## [2]

#### Pattern matching with mismatches

6 YFHRIK

Find pattern matches in reference

```
myseq1 <- readDNAStringSet("./data/AE004437.ffn")
mypos <- matchPattern("ATGGTG", myseq1[[1]], max.mismatch=1)</pre>
```

Count only the corresponding matches

```
countPattern("ATGGCT", myseq1[[1]], max.mismatch=1)
## [1] 3
Count matches in many sequences
vcountPattern("ATGGCT", myseq1, max.mismatch=1)[1:20]
## [1] 3 0 5 4 1 2 2 1 4 3 0 0 1 2 0 1 4 0 0 1
Results shown in DNAStringSet object
tmp <- c(DNAStringSet("ATGGTG"), DNAStringSet(mypos))</pre>
Return a consensus matrix for query and hits
consensusMatrix(tmp)[1:4,]
     [,1] [,2] [,3] [,4] [,5] [,6]
##
## A
             0
                  0
                        0
## C
                   0
                        0
                             0
                                   0
        1
             1
## G
                        4
                                   4
        0
             0
                   4
                             1
## T
        0
             3
                   0
                        0
                                   0
Find all pattern matches in reference
myvpos <- vmatchPattern("ATGGCT", myseq1, max.mismatch=1)</pre>
myvpos # The results are stored as MIndex object.
## MIndex object of length 2058
## $`gi|12057215|gb|AE004437.1|:248-1453 Halobacterium sp. NRC-1, complete genome`
## IRanges object with 3 ranges and 0 metadata columns:
##
             start
                          end
                                  width
##
         <integer> <integer> <integer>
##
     [1]
                            6
                 1
     [2]
                383
                          388
                                       6
##
##
     [3]
                928
                          933
                                       6
##
## $`gi|12057215|gb|AE004437.1|:1450-2115 Halobacterium sp. NRC-1, complete genome`
## IRanges object with 0 ranges and 0 metadata columns:
##
          start
                       end
                               width
##
      <integer> <integer> <integer>
##
## $`gi|12057215|gb|AE004437.1|:2145-3254 Halobacterium sp. NRC-1, complete genome`
## IRanges object with 5 ranges and 0 metadata columns:
##
             start
                          end
                                  width
##
         <integer> <integer> <integer>
##
     [1]
                 1
                            6
                                       6
                94
##
     [2]
                           99
                                       6
##
     [3]
                221
                          226
                                       6
##
     [4]
                535
                                       6
                          540
##
     [5]
                601
                          606
                                       6
##
##
## <2055 more elements>
Views(myseq1[[1]], start(myvpos[[1]]), end(myvpos[[1]])) # Retrieves the result for single entry
```

Views on a 1206-letter DNAString subject

##

```
## subject: ATGACTCGGCGGTCTCGTGTCGGTGCCGGCCTCGCAGCCATTGT...TTGCGATCGTCGTCGTCGTTGTTCGACGCTGGCGGAACCCATGA
## views:
##
       start end width
           1 6
                     6 [ATGACT]
## [1]
## [2]
         383 388
                     6 [ATGGCA]
## [3]
         928 933
                     6 [ATGACT]
Return all matches
sapply(seq(along=myseq1), function(x)
       as.character(Views(myseq1[[x]], start(myvpos[[x]]), end(myvpos[[x]]))))[1:4]
```

#### Pattern matching with regular expression support

```
myseq <- DNAStringSet(c("ATGCAGACATAGTG", "ATGAACATAGATCC", "GTACAGATCAC"))</pre>
myseq[grep("ATG", myseq, perl=TRUE)] # String searching with regular expression support
     A DNAStringSet instance of length 2
##
       width seq
## [1]
         14 ATGCAGACATAGTG
## [2]
          14 ATGAACATAGATCC
pos1 <- regexpr("AT", myseq) # Searches 'myseq' for first match of pattern "AT"
as.numeric(pos1); attributes(pos1) $match.length # Returns position information of matches
## [1] 1 1 7
## [1] 2 2 2
pos2 <- gregexpr("AT", myseq) # Searches 'myseq' for all matches of pattern "AT"
as.numeric(pos2[[1]]); attributes(pos2[[1]]) $match.length # Match positions in first sequence
## [1] 1 9
## [1] 2 2
DNAStringSet(gsub("^ATG", "NNN", myseq)) # String substitution with regular expression support
##
     A DNAStringSet instance of length 3
##
       width seq
## [1]
          14 NNNCAGACATAGTG
## [2]
          14 NNNAACATAGATCC
## [3]
         11 GTACAGATCAC
```

#### PWM Viewing and Searching

#### Plot with seqLogo

```
library(seqLogo)

## Loading required package: grid

pwm <- PWM(DNAStringSet(c("GCT", "GGT", "GCA")))
pwm

## [,1] [,2] [,3]</pre>
```

## Plot with ggseqlogo

The ggseqlogo package (manual) provides many customization options for plotting sequence logos. It also supports various alphabets including sequence logos for amino acid sequences.

```
library(ggplot2); library(ggseqlogo)
pwm <- PWM(DNAStringSet(c("GCT", "GGT", "GCA")))
ggseqlogo(pwm)</pre>
```



Search sequence for PWM matches with score better than min.score

```
chr <- DNAString("AAAGCTAAAGGTAAAGCAAAA")
matchPWM(pwm, chr, min.score=0.9)

## Views on a 21-letter DNAString subject
## subject: AAAGCTAAAGGTAAAGCAAAA
## views:
## start end width</pre>
```

## [1] 4 6 3 [GCT] ## [2] 10 12 3 [GGT] ## [3] 16 18 3 [GCA]

## **NGS** Sequences

## Sequence and Quality Data: FASTQ Format

Four lines per sequence:

- 1. ID
- 2. Sequence
- 3 ID
- 4. Base call qualities (Phred scores) as ASCII characters

The following gives an example of 3 Illumina reads in a FASTQ file. The numbers at the beginning of each line are not part of the FASTQ format. They have been added solely for illustration purposes.

- 1. @SRR038845.3 HWI-EAS038:6:1:0:1938 length=36
- 2. CAACGAGTTCACACCTTGGCCGACAGGCCCGGGTAA
- 3. +SRR038845.3 HWI-EAS038:6:1:0:1938 length=36

```
4. BA@7>B=>:>>7@7@>>9=BAA?;>52;>:9=8.=A
1. @SRR038845.41 HWI-EAS038:6:1:0:1474 length=36
2. CCAATGATTTTTTCCGTGTTTCAGAATACGGTTAA
3. +SRR038845.41 HWI-EAS038:6:1:0:1474 length=36
4. BCCBA@BB@BBBBAB@B9B@=BABA@A:@693:@B=
1. @SRR038845.53 HWI-EAS038:6:1:1:360 length=36
2. GTTCAAAAAGAACTAAATTGTGTCAATAGAAAACTC
3. +SRR038845.53 HWI-EAS038:6:1:1:360 length=36
4. BBCBBBBBB@@BAB?BBBBCBC>BBBAA8>BBBAA@
```

## Sequence and Quality Data: QualityScaleXStringSet

Phred quality scores are integers from 0-50 that are stored as ASCII characters after adding 33. The basic R functions rawToChar and charToRaw can be used to interconvert among their representations.

Phred score interconversion

##

##

## [1]

## [2]

width seq

```
phred <- 1:9
phreda <- paste(sapply(as.raw((phred)+33), rawToChar), collapse="")</pre>
phreda
## [1] "\"#$%&'()*"
as.integer(charToRaw(phreda))-33
## [1] 1 2 3 4 5 6 7 8 9
Construct QualityScaledDNAStringSet from scratch
dset <- DNAStringSet(sapply(1:100, function(x) paste(sample(c("A", "T", "G", "C"), 20, replace=T), collaps
myqlist <- lapply(1:100, function(x) sample(1:40, 20, replace=T)) # Creates random Phred score list.
myqual <- sapply(myqlist, function(x) toString(PhredQuality(x))) # Converts integer scores into ASCII c
myqual <- PhredQuality(myqual) # Converts to a PhredQuality object.
dsetq1 <- QualityScaledDNAStringSet(dset, myqual) # Combines DNAStringSet and quality data in QualitySc
dsetq1[1:2]
##
     A QualityScaledDNAStringSet instance containing:
##
##
     A DNAStringSet instance of length 2
##
       width seq
## [1]
          20 CCTTATAAATAAAGACCGTA
          20 TTGCGACAAGCTATTGCCTC
## [2]
##
```

### Processing FASTQ Files with ShortRead

A PhredQuality instance of length 2

20 +>IFBGC\*#0I@##%,&OC<

20 3&D?\*@\*.9&\$I+H89,90%

The following expains the basic usage of ShortReadQ objects. To make the sample code work, download and unzip this file to your current working directory. The following code performs the download for you.

```
library(ShortRead)
download.file("http://faculty.ucr.edu/~tgirke/HTML_Presentations/Manuals/Workshop_Dec_6_10_2012/Rsequen
unzip("data.zip")
```

```
Important utilities for accessing FASTQ files
```

```
fastq <- list.files("data", "*.fastq$"); fastq <- paste("data/", fastq, sep="")</pre>
names(fastq) <- paste("flowcell6_lane", 1:length(fastq), sep="_")</pre>
(fq <- readFastq(fastq[1])) # Imports first FASTQ file</pre>
## class: ShortReadQ
## length: 1000 reads; width: 36 cycles
countLines(dirPath="./data", pattern=".fastq$")/4 # Counts numbers of reads in FASTQ files
## SRR038845.fastq SRR038846.fastq SRR038848.fastq SRR038850.fastq
##
              1000
                               1000
                                                1000
id(fq)[1] # Returns ID field
     A BStringSet instance of length 1
##
##
       width seq
          43 SRR038845.3 HWI-EAS038:6:1:0:1938 length=36
## [1]
sread(fq)[1] # Returns sequence
##
     A DNAStringSet instance of length 1
##
       width seq
## [1]
          36 CAACGAGTTCACACCTTGGCCGACAGGCCCGGGTAA
quality(fq)[1] # Returns Phred scores
## class: FastqQuality
## quality:
##
     A BStringSet instance of length 1
##
       width seq
          36 BA07>B=>:>>7070>>9=BAA?;>52;>:9=8.=A
as(quality(fq), "matrix")[1:4,1:12] # Coerces Phred scores to numeric matrix
##
        [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9] [,10] [,11] [,12]
## [1,]
          33
               32
                    31
                          22
                               29
                                    33
                                          28
                                               29
                                                    25
                                                           29
                                                                 29
                                                                       22
## [2,]
          33
               34
                    34
                          33
                               32
                                    31
                                          33
                                               33
                                                    31
                                                           33
                                                                 33
                                                                       33
                                                                       33
## [3,]
          33
               33
                    34
                          33
                               33
                                    33
                                          33
                                               33
                                                    33
                                                                 31
                                                           31
## [4,]
          33
                    33
                          33
                               31
                                    33
                                          28
                                               31
                                                           32
                                                                 33
                                                                       33
ShortReadQ(sread=sread(fq), quality=quality(fq), id=id(fq)) # Constructs a ShortReadQ from components
## class: ShortReadQ
## length: 1000 reads; width: 36 cycles
```

## FASTQ Quality Reports

#### Using systemPipeR

The following seeFastq and seeFastqPlot functions generate and plot a series of useful quality statistics for a set of FASTQ files.

```
library(systemPipeR)
fqlist <- seeFastq(fastq=fastq, batchsize=800, klength=8) # For real data set batchsize to at least 10^
seeFastqPlot(fqlist)</pre>
```



Handles many samples in one PDF file. For more details see here

#### Using ShortRead

The ShortRead package contains several FASTQ quality reporting functions.

## Filtering and Trimming FASTQ Files with ShortRead

## Adaptor trimming

```
fqtrim <- trimLRPatterns(Rpattern="GCCCGGGTAA", subject=fq)</pre>
sread(fq)[1:2] # Before trimming
##
     A DNAStringSet instance of length 2
##
       width seq
## [1]
          36 CAACGAGTTCACACCTTGGCCGACAGGCCCGGGTAA
## [2]
          36 CCAATGATTTTTTCCGTGTTTCAGAATACGGTTAA
sread(fqtrim)[1:2] # After trimming
##
     A DNAStringSet instance of length 2
##
       width seq
## [1]
          26 CAACGAGTTCACACCTTGGCCGACAG
## [2]
          36 CCAATGATTTTTTCCGTGTTTCAGAATACGGTTAA
```

#### Read counting and duplicate removal

```
tables(fq)$distribution # Counts read occurences
    nOccurrences nReads
##
## 1
                1
## 2
                2
                      26
sum(srduplicated(fq)) # Identifies duplicated reads
## [1] 26
fq[!srduplicated(fq)]
## class: ShortReadQ
## length: 974 reads; width: 36 cycles
Trimming low quality tails
cutoff <- 30
cutoff <- rawToChar(as.raw(cutoff+33))</pre>
sread(trimTails(fq, k=2, a=cutoff, successive=FALSE))[1:2]
##
     A DNAStringSet instance of length 2
##
       width seq
## [1]
           4 CAAC
## [2]
          20 CCAATGATTTTTTTCCGTGT
```

#### Removal of reads with Phred scores below a threshold value

```
cutoff <- 30
qcount <- rowSums(as(quality(fq), "matrix") <= 20)
fq[qcount == 0] # Number of reads where all Phred scores >= 20
```

```
## class: ShortReadQ
## length: 349 reads; width: 36 cycles
```

#### Removal of reads with x Ns and/or low complexity segments

```
filter1 <- nFilter(threshold=1) # Keeps only reads without Ns
filter2 <- polynFilter(threshold=20, nuc=c("A","T","G","C")) # Removes reads with >=20 of
filter <- compose(filter1, filter2)
fq[filter(fq)]

## class: ShortReadQ
## length: 989 reads; width: 36 cycles</pre>
```

## Memory Efficient FASTQ Processing

Streaming through FASTQ files with FastqStreamer and random sampling reads with FastqSampler

```
fq <- yield(FastqStreamer(fastq[1], 50)) # Imports first 50 reads
fq <- yield(FastqSampler(fastq[1], 50)) # Random samples 50 reads</pre>
```

Streaming through a FASTQ file while applying filtering/trimming functions and writing the results to a new file here SRR038845.fastq\_sub in data directory.

```
f <- FastqStreamer(fastq[1], 50)
while(length(fq <- yield(f))) {
   fqsub <- fq[grepl("^TT", sread(fq))]
   writeFastq(fqsub, paste(fastq[1], "sub", sep="_"), mode="a", compress=FALSE)
}
close(f)</pre>
```

## **Range Operations**

## Important Data Objects for Range Operations

- IRanges: stores range data only (IRanges library)
- GRanges: stores ranges and annotations (GenomicRanges library)
- GRangesList: list version of GRanges container (GenomicRanges library)

#### Range Data Are Stored in IRanges and GRanges Containers

#### Construct GRanges Object

```
library(GenomicRanges); library(rtracklayer)
gr <- GRanges(seqnames = Rle(c("chr1", "chr2", "chr1", "chr3"), c(1, 3, 2, 4)), ranges = IRanges(1:10, -</pre>
```

#### Import GFF into GRanges Object

```
gff <- import.gff("http://faculty.ucr.edu/~tgirke/Documents/R_BioCond/Samples/gff3.gff") # Imports a si
seqlengths(gff) <- end(ranges(gff[which(values(gff)[,"type"]=="chromosome"),]))</pre>
names(gff) <- 1:length(gff) # Assigns names to corresponding slot</pre>
gff[1:4,]
## GRanges object with 4 ranges and 10 metadata columns:
##
       seqnames
                           ranges strand |
                                                                                                          ID
                                               source
                                                                      score
                                                                                 phase
                                                             type
##
          <Rle>
                        <IRanges>
                                    <Rle> | <factor>
                                                         <factor> <numeric> <integer>
                                                                                                <character>
##
                     1, 30427671]
                                               TAIR10 chromosome
                                                                                  <NA>
                                                                                                        Chr1
     1
           Chr1 [
                                                                        <NA>
##
     2
           Chr1 [3631,
                            58991
                                               TAIR10
                                                             gene
                                                                        <NA>
                                                                                   <NA>
                                                                                                  AT1G01010
##
     3
           Chr1 [3631,
                            58991
                                               TAIR10
                                                             mRNA
                                                                        <NA>
                                                                                  <NA>
                                                                                                AT1G01010.1
##
           Chr1 [3760,
                            5630]
                                               TAIR10
                                                                        <NA>
                                                                                  <NA> AT1G01010.1-Protein
                                                         protein
##
              Name
                                                                 Index Derives_from
                                                   Parent
                                    Note
##
       <character>
                        <CharacterList> <CharacterList> <character>
##
               Chr1
                                                                  <NA>
                                                                                <NA>
     1
##
         AT1G01010 protein_coding_gene
                                                                  <NA>
                                                                                <NA>
##
     3 AT1G01010.1
                                                AT1G01010
                                                                     1
                                                                                <NA>
     4 AT1G01010.1
                                                                        AT1G01010.1
##
                                                                  <NA>
##
##
     seqinfo: 7 sequences from an unspecified genome
```

#### Coerce GRanges object to data.frame

```
as.data.frame(gff)[1:4, 1:7]
##
     seqnames start
                           end
                                  width strand source
                                                              type
## 1
         Chr1
                   1 30427671 30427671
                                              + TAIR10 chromosome
## 2
         Chr1
                3631
                          5899
                                   2269
                                              + TAIR10
                                                              gene
## 3
         Chr1
                3631
                          5899
                                   2269
                                              + TAIR10
                                                              mRNA
## 4
         Chr1 3760
                          5630
                                   1871
                                              + TAIR10
                                                           protein
```

#### Coerce GRanges to RangedData object and vice versa

```
gff_rd <- as(gff, "RangedData")
gff_gr <- as(gff_rd, "GRanges")</pre>
```

#### **Utilities for Range Containers**

#### Accessor and subsetting methods for GRanges objects

Subsetting and replacement

```
gff[1:4]
## GRanges object with 4 ranges and 10 metadata columns:
##
       seqnames
                            ranges strand |
                                               source
                                                                       score
                                                                                  phase
                                                                                                           ID
                                                             type
##
           <Rle>
                         <IRanges>
                                    <Rle> | <factor>
                                                         <factor> <numeric> <integer>
                                                                                                 <character>
##
           Chr1 [
                     1, 30427671]
                                               TAIR10 chromosome
                                                                        <NA>
                                                                                   <NA>
                                                                                                         Chr1
##
     2
           Chr1 [3631,
                             5899]
                                         + |
                                               TAIR10
                                                                        <NA>
                                                                                   <NA>
                                                                                                   AT1G01010
                                                             gene
                                                                        <NA>
##
     3
           Chr1 [3631,
                             5899]
                                         + |
                                               TAIR10
                                                             mRNA
                                                                                   <NA>
                                                                                                 AT1G01010.1
##
     4
           Chr1 [3760,
                             5630]
                                               TAIR10
                                                                        <NA>
                                                                                   <NA> AT1G01010.1-Protein
                                         + |
                                                          protein
```

```
##
              Name
                                   Note
                                                 Parent
                                                               Index Derives from
##
       <character>
                       <CharacterList> <CharacterList> <character>
                                                                      <character>
##
              Chr1
                                                                <NA>
         AT1G01010 protein_coding_gene
                                                                <NA>
                                                                             <NA>
##
##
     3 AT1G01010.1
                                              AT1G01010
                                                                   1
                                                                             <NA>
##
     4 AT1G01010.1
                                                                <NA>
                                                                      AT1G01010.1
##
##
     seqinfo: 7 sequences from an unspecified genome
gff[1:4, c("type", "ID")]
## GRanges object with 4 ranges and 2 metadata columns:
##
       segnames
                          ranges strand |
                                                                        ID
                                                 type
##
          <Rle>
                       <IRanges> <Rle> |
                                             <factor>
                                                               <character>
           Chr1 [
##
                    1, 30427671]
                                       + | chromosome
                                                                      Chr1
##
           Chr1 [3631,
                           5899]
                                       + |
                                                                 AT1G01010
     2
                                                 gene
##
           Chr1 [3631,
                           5899]
                                       + |
                                                 mRNA
                                                               AT1G01010.1
##
           Chr1 [3760,
                           5630]
                                       + |
                                              protein AT1G01010.1-Protein
##
     seqinfo: 7 sequences from an unspecified genome
gff[2] <- gff[3]
GRanges objects can be concatenated with the c function
c(gff[1:2], gff[401:402])
## GRanges object with 4 ranges and 10 metadata columns:
##
         segnames
                            ranges strand |
                                               source
                                                                 type
                                                                          score
                                                                                    phase
##
            <Rle>
                         <IRanges>
                                     <Rle> | <factor>
                                                            <factor> <numeric> <integer>
##
                      1, 30427671]
                                         + |
       1
             Chr1 [
                                               TAIR10
                                                          chromosome
                                                                           <NA>
                                                                                      <NA>
             Chr1 [3631,
##
       2
                             5899]
                                         + |
                                               TAIR10
                                                                mRNA
                                                                           <NA>
                                                                                      <NA>
##
             Chr5 [5516,
                             5769]
                                               TAIR10
                                                                           <NA>
                                                                                      <NA>
     401
                                         - |
                                                             protein
##
     402
             Chr5 [5770,
                             5801]
                                         - |
                                               TAIR10 five_prime_UTR
                                                                           <NA>
                                                                                      <NA>
##
                          ID
                                     Name
                                                     Note
                                                                    Parent
                                                                                  Index Derives from
##
                 <character> <character> <characterList> <character> <character> <character>
##
                        Chr1
                                                                                  <NA>
                                                                                                <NA>
                 AT1G01010.1 AT1G01010.1
##
                                                                 AT1G01010
                                                                                     1
                                                                                                <NA>
       2
##
     401 AT5G01015.2-Protein AT5G01015.2
                                                                                   <NA>
                                                                                        AT5G01015.2
                                                               AT5G01015.2
##
     402
                        <NA>
                                                                                  <NA>
                                                                                                <NA>
##
     seqinfo: 7 sequences from an unspecified genome
##
Acessor functions
seqnames(gff)
## factor-Rle of length 449 with 7 runs
               72
                    22
                          38 118 172
     Lengths:
                                          13
     Values : Chr1 Chr2 Chr3 Chr4 Chr5 ChrC ChrM
## Levels(7): Chr1 Chr2 Chr3 Chr4 Chr5 ChrC ChrM
ranges(gff)
\#\# IRanges object with 449 ranges and 0 metadata columns:
                                  width
             start
                         end
##
         <integer> <integer> <integer>
##
                 1 30427671 30427671
```

```
##
       2
               3631
                          5899
                                    2269
##
       3
               3631
                          5899
                                    2269
##
       4
               3760
                          5630
                                    1871
##
               3631
                                      283
       5
                          3913
##
                . . .
                                      . . .
                           . . .
     . . .
##
     445
              11918
                         12241
                                      324
##
     446
              11918
                        12241
                                      324
##
     447
              11918
                         12241
                                      324
##
     448
              11918
                        12241
                                      324
              11918
##
     449
                         12241
                                      324
strand(gff)
## factor-Rle of length 449 with 13 runs
##
     Lengths:
                18 54 28 21
                                12 117
                                           1 171
                                                       12
                                                                     5
##
     Values :
## Levels(3): + -
seqlengths(gff)
##
       Chr1
                 Chr2
                           Chr3
                                    Chr4
                                              Chr5
                                                        ChrC
                                                                  ChrM
## 30427671 19698289 23459830 18585056 26975502
                                                      154478
                                                                366924
start(gff[1:4])
## [1]
          1 3631 3631 3760
end(gff[1:4])
## [1] 30427671
                     5899
                               5899
                                         5630
width(gff[1:4])
## [1] 30427671
                     2269
                               2269
                                         1871
Accessing metadata component
values(gff) # or elementMetadata(qff)
## DataFrame with 449 rows and 10 columns
##
                                            phase
                                                                     ID
         source
                        type
                                 score
                                                                                Name
                                                                                                      Note
##
                                                           <character> <character>
       <factor>
                   <factor> <numeric> <integer>
                                                                                          <CharacterList>
## 1
         TAIR10 chromosome
                                    NA
                                               NA
                                                                   Chr1
## 2
         TAIR10
                                    NA
                                               NA
                                                           AT1G01010.1 AT1G01010.1
                       mRNA
## 3
         TAIR10
                       mRNA
                                    NA
                                                           AT1G01010.1 AT1G01010.1
## 4
         TAIR10
                                    NA
                                               NA AT1G01010.1-Protein AT1G01010.1
                    protein
## 5
         TAIR10
                        exon
                                    NA
                                               NA
                                                                     NA
                                                                                  NA
##
## 445
         TAIR10
                                    NA
                                               NA
                                                             ATMG00030
                                                                           ATMG00030 protein_coding_gene
                       gene
## 446
         TAIR10
                                                           ATMG00030.1 ATMG00030.1
                       mRNA
                                    NA
                                               NA
## 447
         TAIR10
                    protein
                                    NA
                                               NA ATMG00030.1-Protein ATMG00030.1
## 448
                                               NA
                                                                     NA
         TAIR10
                        exon
                                    NA
                                                                                  NA
##
  449
         TAIR10
                         CDS
                                    NA
                                                0
                                                                     NA
                                                                                  NA
##
                                  Parent
                                                Index Derives_from
##
                         <CharacterList> <character>
                                                        <character>
## 1
                                                    NA
                                                                  NA
## 2
                               AT1G01010
                                                     1
                                                                  NA
## 3
                               AT1G01010
                                                     1
                                                                  NA
                                                    NA
## 4
                                                        AT1G01010.1
```

```
## 5
                             AT1G01010.1
                                                                  NA
                                                    NA
## ...
                                                   . . .
                                                                  . . .
## 445
                                                    NA
                                                                  NA
## 446
                               ATMG00030
                                                                  NA
                                                     1
## 447
                                                    NA
                                                        ATMG00030.1
## 448
                             ATMG00030.1
                                                    NA
                                                                  NA
## 449 ATMG00030.1, ATMG00030.1-Protein
                                                    NA
                                                                  NA
values(gff)[, "type"][1:20]
    [1] chromosome
                          mRNA
                                           mRNA
                                                             protein
                                                                              exon
                                                                                                five prime UTR
##
    [7] CDS
                          exon
                                           CDS
                                                             exon
                                                                              CDS
                                                                                                exon
## [13] CDS
                          exon
                                           CDS
                                                                              CDS
                                                             exon
                                                                                                three_prime_UTR
## [19] gene
                          mRNA
## Levels: chromosome gene mRNA protein exon five_prime_UTR CDS three_prime_UTR rRNA tRNA
gff[values(gff)[ ,"type"] == "gene"]
## GRanges object with 21 ranges and 10 metadata columns:
##
                            ranges strand |
                                                                                                 ID
         segnames
                                                source
                                                                                phase
                                                            type
                                                                      score
##
             <Rle>
                         <IRanges>
                                     <Rle> | <factor> <factor> <numeric> <integer> <character>
##
      19
              Chr1 [ 5928, 8737]
                                                TAIR10
                                                                       <NA>
                                                                                  <NA>
                                                                                         AT1G01020
                                                            gene
##
      64
              Chr1 [11649, 13714]
                                         - 1
                                                TAIR10
                                                                       <NA>
                                                                                  <NA>
                                                                                         AT1G01030
                                                            gene
##
      74
                                                TAIR10
              Chr2 [ 1025,
                             2810]
                                         + |
                                                            gene
                                                                       <NA>
                                                                                  <NA>
                                                                                         AT2G01008
##
      84
              Chr2 [ 3706,
                             5513]
                                                TAIR10
                                                                       <NA>
                                                                                  <NA>
                                                                                         AT2G01010
                                         + |
                                                            gene
##
      87
              Chr2 [ 5782,
                             5945]
                                                TAIR10
                                                                       <NA>
                                                                                  <NA>
                                                                                         AT2G01020
                                         + |
                                                            gene
##
     . . .
               . . .
                                                             . . .
                                                                        . . .
                                                                                   . . .
##
     427
              ChrC [ 383,
                             1444]
                                         - |
                                                TAIR10
                                                                       <NA>
                                                                                  <NA>
                                                                                         ATCG00020
                                                            gene
##
     432
              ChrC [ 1717,
                             4347]
                                                TAIR10
                                                                       <NA>
                                                                                  <NA>
                                         - |
                                                            gene
                                                                                         ATCG00030
##
     437
              ChrM [ 273,
                              734]
                                                TAIR10
                                                                       <NA>
                                                                                  <NA>
                                                                                         ATMG00010
                                                            gene
##
              ChrM [ 8848, 11415]
                                                                       <NA>
                                                                                  <NA>
     442
                                                TAIR10
                                                                                         ATMG00020
                                                            gene
##
     445
              ChrM [11918, 12241]
                                                TAIR10
                                                                       <NA>
                                                                                  <NA>
                                                                                         ATMG00030
                                         + |
                                                            gene
##
                 Name
                                       Note
                                                      Parent
                                                                     Index Derives from
##
                           <CharacterList> <CharacterList> <character>
                                                                            <character>
         <character>
##
      19
           AT1G01020 protein_coding_gene
                                                                      <NA>
                                                                                    <NA>
##
            AT1G01030 protein_coding_gene
                                                                                    <NA>
      64
                                                                      <NA>
##
      74
            AT2G01008 protein_coding_gene
                                                                      <NA>
                                                                                    <NA>
##
      84
           AT2G01010
                                       rRNA
                                                                      <NA>
                                                                                    <NA>
##
      87
           AT2G01020
                                       rRNA
                                                                      <NA>
                                                                                    <NA>
##
                                                                       . . .
                                                                                     . . .
     . . .
##
     427
            ATCG00020 protein_coding_gene
                                                                                    <NA>
                                                                      <NA>
##
     432
           ATCG00030
                                                                      <NA>
                                                                                    <NA>
##
     437
                                                                      <NA>
                                                                                    <NA>
            ATMG00010 protein_coding_gene
##
     442
            ATMG00020
                                                                      < NA >
                                                                                    <NA>
##
     445
           ATMG00030 protein_coding_gene
                                                                      <NA>
                                                                                    <NA>
##
     seqinfo: 7 sequences from an unspecified genome
##
```

#### Useful utilities for GRanges objects

Remove chromosome ranges

```
gff <- gff[values(gff)$type != "chromosome"]</pre>
```

Erase the strand information

```
strand(gff) <- "*"</pre>
Collapses overlapping ranges to continuous ranges.
reduce(gff)
## GRanges object with 22 ranges and 0 metadata columns:
##
          seqnames
                             ranges strand
##
              <Rle>
                         <IRanges>
                                     <Rle>
##
               Chr1 [ 3631, 5899]
      [1]
##
      [2]
               Chr1 [ 5928, 8737]
      [3]
               Chr1 [11649, 13714]
##
##
      [4]
               Chr2 [ 1025,
                             28107
##
      [5]
               Chr2 [ 3706,
                             5513]
##
      . . .
##
     [18]
               ChrC [
                      383,
                              1444]
               ChrC [ 1717,
##
     [19]
                              4347]
##
     [20]
               ChrM [ 273,
                               734]
##
     [21]
               ChrM [ 8848, 11415]
##
     [22]
               ChrM [11918, 12241]
##
##
     seqinfo: 7 sequences from an unspecified genome
Return uncovered regions
gaps(gff)
## GRanges object with 43 ranges and 0 metadata columns:
##
          segnames
                               ranges strand
##
              <Rle>
                            <IRanges> <Rle>
##
      [1]
               Chr1 [
                        1, 30427671]
##
      [2]
               Chr1 [
                        1, 30427671]
##
      [3]
               Chr1 [
                        1,
                                3630]
      [4]
##
               Chr1 [5900,
                                5927]
##
      [5]
               Chr1 [8738,
                               116487
##
      . . .
     [39]
##
               ChrM [
                           1, 366924]
##
     [40]
               ChrM
                     1,
                                 272]
##
     [41]
               ChrM [
                        735,
                                8847]
##
     [42]
               ChrM [11416,
                               11917]
     [43]
##
               ChrM [12242, 366924]
##
##
     seqinfo: 7 sequences from an unspecified genome
More intuitive way to get uncovered regions
setdiff(as(seqinfo(gff), "GRanges"), gff)
## GRanges object with 29 ranges and 0 metadata columns:
##
          seqnames
                                ranges strand
##
              <Rle>
                             <IRanges>
                                         <Rle>
```

```
##
       [1]
                Chr1 [
                            1,
                                     3630]
##
       [2]
                Chr1 [ 5900,
                                    5927]
##
       [3]
                Chr1 [ 8738,
                                   11648]
##
       [4]
                Chr1 [13715, 30427671]
       [5]
                Chr2 [
##
                            1,
                                    1024]
##
       . . .
                 . . .
                                       . . .
```

```
[25]
              ChrC
                      [ 4348, 154478]
##
##
     [26]
              ChrM
                      1,
                                 272]
     [27]
##
              ChrM
                      [ 735,
                                 8847]
##
     [28]
              ChrM
                      [11416, 11917]
                      [12242, 366924]
##
     [29]
              ChrM
##
     seqinfo: 7 sequences from an unspecified genome
```

Return disjoint ranges

#### disjoin(gff)

```
## GRanges object with 211 ranges and 0 metadata columns:
           seqnames
                             ranges strand
##
               <Rle>
                          <IRanges>
                                      <Rle>
##
       [1]
                Chr1
                       [3631, 3759]
##
       [2]
                Chr1
                       [3760, 3913]
##
       [3]
                       [3914, 3995]
                Chr1
##
       [4]
               Chr1
                       [3996, 4276]
##
       [5]
               Chr1
                       [4277, 4485]
##
       . . .
                . . .
##
     [207]
               ChrC [ 1752,
                              4310]
##
     [208]
                ChrC [ 4311,
                              4347]
##
     [209]
                ChrM [ 273,
                                734]
##
     [210]
                ChrM [ 8848, 11415]
##
               ChrM [11918, 12241]
     [211]
##
##
     seqinfo: 7 sequences from an unspecified genome
```

## integer-Rle of length 26975502 with 64 runs

Returns coverage of ranges

#### coverage(gff)

```
## RleList of length 7
## $Chr1
## integer-Rle of length 30427671 with 45 runs
##
     Lengths:
                   3630
                             129
                                       154
                                                  82
                                                           281 ...
                                                                         233
                                                                                  161
                                                                                            380 30413957
##
     Values :
                      0
                                4
                                         5
                                                   3
                                                             5 ...
                                                                                    2
                                                                                              4
                                                                           4
##
## $Chr2
## integer-Rle of length 19698289 with 14 runs
     Lengths:
                   1024
                             248
                                       185
                                                  53
                                                           362 ...
                                                                         164
                                                                                  625
                                                                                            102 19691617
     Values :
                                5
                                                   5
##
                      0
                                         3
                                                             3 ...
                                                                           3
                                                                                    0
                                                                                              5
##
## $Chr3
## integer-Rle of length 23459830 with 29 runs
     Lengths:
                   1652
                             145
                                       139
                                                 111
                                                            95 ...
                                                                         155
                                                                                  148
                                                                                            156 23453781
##
     Values :
                      0
                                4
                                         5
                                                   3
                                                            5 ...
                                                                          3
                                                                                    5
                                                                                              4
## $Chr4
## integer-Rle of length 18585056 with 72 runs
                                                                         212
##
     Lengths:
                   1179
                             357
                                      1358
                                                 128
                                                           872 ...
                                                                                  114
                                                                                             74 18571697
##
     Values :
                      0
                                5
                                         0
                                                   5
                                                             3 ...
                                                                           3
                                                                                    5
                                                                                              4
##
```

```
72 ...
##
     Lengths:
                   1222
                              28
                                        28
                                                109
                                                                        76
                                                                                 55
                                                                                          174 26967058
##
     Values:
                               4
                                        7
                                                 13
                                                          16 ...
                                                                         3
                                                                                  5
                                                                                            4
                     0
                                                                                                     0
##
## ...
## <2 more elements>
Return the index pairings for overlapping ranges
findOverlaps(gff, gff[1:4])
## Hits object with 55 hits and 0 metadata columns:
##
          queryHits subjectHits
##
          <integer>
                       <integer>
##
      [1]
                  1
##
      [2]
                               2
                  1
##
      [3]
                  1
                               4
##
      [4]
                  1
                               3
##
      [5]
                  2
                               1
##
      . . .
                 . . .
                             . . .
##
     [51]
                 16
                               1
##
     [52]
                 16
                               2
##
     [53]
                 16
                               3
##
     [54]
                  17
                               1
##
     [55]
                 17
                               2
##
##
     queryLength: 442 / subjectLength: 4
Counts overlapping ranges
countOverlaps(gff, gff[1:4])[1:40]
                      8
                          9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34
                            ## 35 36 37 38 39 40 41
   0 0 0 0 0 0 0
Return only overlapping ranges
subsetByOverlaps(gff, gff[1:4])
## GRanges object with 17 ranges and 10 metadata columns:
##
        segnames
                        ranges strand |
                                           source
                                                             type
                                                                       score
                                                                                 phase
##
           <Rle>
                     <IRanges>
                                <Rle> | <factor>
                                                         <factor> <numeric> <integer>
      2
##
            Chr1 [3631, 5899]
                                    * |
                                           TAIR10
                                                             mRNA
                                                                        <NA>
                                                                                   <NA>
##
      3
            Chr1 [3631, 5899]
                                           TAIR10
                                                                        <NA>
                                                                                   <NA>
                                    * |
                                                             mRNA
##
      4
            Chr1 [3760, 5630]
                                    * |
                                          TAIR10
                                                          protein
                                                                        <NA>
                                                                                   <NA>
##
      5
            Chr1 [3631, 3913]
                                    * |
                                           TAIR10
                                                                        <NA>
                                                                                   <NA>
                                                             exon
##
      6
            Chr1 [3631, 3759]
                                    * |
                                           TAIR10
                                                                        <NA>
                                                                                   <NA>
                                                  five_prime_UTR
##
                                                              . . .
##
     14
            Chr1 [5174, 5326]
                                           TAIR10
                                                                        <NA>
                                                                                   <NA>
                                    * |
                                                             exon
##
                                           TAIR10
                                                              CDS
     15
            Chr1 [5174, 5326]
                                                                        <NA>
                                                                                     0
                                    * |
##
     16
            Chr1 [5439, 5899]
                                    * |
                                           TAIR10
                                                             exon
                                                                        <NA>
                                                                                   <NA>
##
            Chr1 [5439, 5630]
                                           TAIR10
                                                              CDS
                                                                        <NA>
     17
                                    * |
                                                                                     0
##
     18
                                                                        <NA>
                                                                                   <NA>
            Chr1 [5631, 5899]
                                    * |
                                           TAIR10 three_prime_UTR
##
                                    Name
                          ID
                                                     Note
                                                                                    Parent
                                                                                                  Index
##
                <character> <character> <CharacterList>
                                                                           <CharacterList> <character>
##
      2
                AT1G01010.1 AT1G01010.1
                                                                                 AT1G01010
```

AT1G01010

1

AT1G01010.1 AT1G01010.1

##

3

```
##
      4 AT1G01010.1-Protein AT1G01010.1
                                                                                                              <NA>
##
      5
                          <NA>
                                        <NA>
                                                                                       AT1G01010.1
                                                                                                              <NA>
                                        <NA>
##
      6
                          < NA >
                                                                                       AT1G01010.1
                                                                                                              <NA>
##
                            . . .
                                         . . .
                                                                                                               . . .
##
     14
                          <NA>
                                        <NA>
                                                                                       AT1G01010.1
                                                                                                              <NA>
##
                          <NA>
                                        <NA>
                                                                AT1G01010.1,AT1G01010.1-Protein
                                                                                                              <NA>
     15
##
     16
                          <NA>
                                        <NA>
                                                                                       AT1G01010.1
                                                                                                              <NA>
##
     17
                          <NA>
                                        <NA>
                                                                AT1G01010.1, AT1G01010.1-Protein
                                                                                                              <NA>
##
     18
                          <NA>
                                        <NA>
                                                                                       AT1G01010.1
                                                                                                              <NA>
##
         Derives_from
##
          <character>
##
      2
                  <NA>
##
      3
                  <NA>
##
      4
          AT1G01010.1
##
      5
                  <NA>
##
      6
                  <NA>
##
                   . . .
##
     14
                  <NA>
##
     15
                  <NA>
##
     16
                  <NA>
##
     17
                  <NA>
##
                  <NA>
##
     seginfo: 7 sequences from an unspecified genome
```

### GRangesList Objects

```
 sp \leftarrow split(gff, seq(along=gff)) \ \# \ Stores \ every \ range \ in \ separate \ component \ of \ a \ GRangesList \ object \\ split(gff, seqnames(gff)) \ \# \ Stores \ ranges \ of \ each \ chromosome \ in \ separate \ component.
```

```
## GRangesList object of length 7:
##
   GRanges object with 71 ranges and 10 metadata columns:
##
        seqnames
                           ranges strand |
                                               source
                                                                   type
                                                                             score
                                                                                        phase
                        <IRanges>
##
            <Rle>
                                    <Rle> |
                                             <factor>
                                                               <factor> <numeric> <integer>
##
      2
             Chr1
                     [3631, 5899]
                                                                              <NA>
                                               TAIR10
                                                                   mRNA
                                                                                         <NA>
##
      3
             Chr1
                     [3631, 5899]
                                               TAIR10
                                                                   mRNA
                                                                              <NA>
                                                                                         <NA>
##
      4
             Chr1
                     [3760, 5630]
                                                                              <NA>
                                                                                         <NA>
                                               TAIR10
                                                                protein
##
      5
             Chr1
                     [3631, 3913]
                                               TAIR10
                                                                   exon
                                                                              <NA>
                                                                                         <NA>
##
                     [3631, 3759]
                                                       five_prime_UTR
      6
             Chr1
                                        * |
                                               TAIR10
                                                                              <NA>
                                                                                         <NA>
##
                                                                    . . .
                                                                               . . .
                                                                                          . . .
                                       . . .
##
             Chr1 [13335, 13714]
                                               TAIR10
     68
                                        * |
                                                                   exon
                                                                              <NA>
                                                                                         <NA>
##
     69
             Chr1 [12941, 13173]
                                        * |
                                               TAIR10 five_prime_UTR
                                                                              <NA>
                                                                                         <NA>
             Chr1 [11864, 12940]
##
     70
                                        * |
                                               TAIR10
                                                                    CDS
                                                                              <NA>
                                                                                            0
##
     71
             Chr1 [11649, 11863]
                                        * |
                                               TAIR10 three_prime_UTR
                                                                              <NA>
                                                                                         <NA>
##
     72
             Chr1 [11649, 13173]
                                        * |
                                               TAIR10
                                                                              <NA>
                                                                                         <NA>
                                                                   exon
##
                                                                                         Parent
                                                                                                       Index
                                      Name
                                                        Note
##
                 <character> <character> <CharacterList>
                                                                               <CharacterList> <character>
##
      2
                 AT1G01010.1 AT1G01010.1
                                                                                      AT1G01010
                                                                                                            1
##
                 AT1G01010.1 AT1G01010.1
                                                                                      AT1G01010
                                                                                                            1
##
      4 AT1G01010.1-Protein AT1G01010.1
                                                                                                         <NA>
##
      5
                         <NA>
                                      <NA>
                                                                                    AT1G01010.1
                                                                                                         <NA>
##
      6
                         <NA>
                                      <NA>
                                                                                    AT1G01010.1
                                                                                                         <NA>
```

```
##
     . .
                           . . .
                                        . . .
                                                           . . .
                                                                                                            . . .
##
     68
                                                                                     AT1G01030.1
                          <NA>
                                       <NA>
                                                                                                           <NA>
##
     69
                                                                                     AT1G01030.1
                          <NA>
                                       <NA>
                                                                                                           <NA>
##
     70
                                       <NA>
                                                                                                           <NA>
                          <NA>
                                                               AT1G01030.1, AT1G01030.1-Protein
##
     71
                          <NA>
                                       <NA>
                                                                                     AT1G01030.1
                                                                                                           <NA>
##
     72
                          <NA>
                                       <NA>
                                                                                     AT1G01030.1
                                                                                                           <NA>
##
        Derives from
##
          <character>
##
      2
                  <NA>
##
      3
                  <NA>
##
          AT1G01010.1
##
      5
                  <NA>
      6
##
                  <NA>
##
     . .
                   . . .
##
     68
                  <NA>
##
     69
                  <NA>
##
     70
                  <NA>
##
     71
                  <NA>
##
     72
                  <NA>
##
##
## <6 more elements>
## -----
## seqinfo: 7 sequences from an unspecified genome
unlist(sp) # Returns data as GRanges object
   GRanges object with 442 ranges and 10 metadata columns:
##
              seqnames
                                 ranges strand |
                                                      source
                                                                         type
                                                                                   score
                                                                                              phase
##
                  <Rle>
                              <IRanges>
                                          <Rle> | <factor>
                                                                    <factor> <numeric> <integer>
##
                   Chr1
                           [3631, 5899]
                                                                                               <NA>
          1.2
                                                      TAIR10
                                                                         mRNA
                                                                                    <NA>
          2.3
##
                   Chr1
                           [3631, 5899]
                                                      TAIR10
                                                                         mRNA
                                                                                    <NA>
                                                                                               <NA>
##
          3.4
                   Chr1
                           [3760, 5630]
                                                      TAIR10
                                                                                    <NA>
                                                                                               <NA>
                                               *
                                                                      protein
##
          4.5
                   Chr1
                           [3631, 3913]
                                                      TAIR10
                                                                                    <NA>
                                                                                               <NA>
                                                                         exon
          5.6
                   Chr1
##
                           [3631, 3759]
                                               *
                                                      TAIR10 five_prime_UTR
                                                                                    <NA>
                                                                                               <NA>
##
                    . . .
                                                                                                . . .
          . . .
                                                         . . .
                                                                          . . .
                                                                                     . . .
##
     438.445
                   ChrM [11918, 12241]
                                                                                               <NA>
                                                      TAIR10
                                                                                    <NA>
                                               *
                                                                         gene
##
     439.446
                   ChrM [11918, 12241]
                                                      TAIR10
                                                                                               <NA>
                                               *
                                                                         mRNA
                                                                                    <NA>
##
     440.447
                   ChrM [11918, 12241]
                                                      TAIR10
                                                                                    <NA>
                                                                                               <NA>
                                               *
                                                                      protein
##
     441.448
                   ChrM [11918, 12241]
                                                      TAIR10
                                                                                    <NA>
                                                                                               <NA>
                                               * |
                                                                         exon
                   ChrM [11918, 12241]
                                               * |
##
     442.449
                                                                                    <NA>
                                                      TAIR10
                                                                          CDS
                                                                                                   0
##
                                 ID
                                             Name
                                                                   Note
                                                                                                     Parent
##
                       <character> <character>
                                                                                           <CharacterList>
                                                       <CharacterList>
                       AT1G01010.1 AT1G01010.1
##
          1.2
                                                                                                  AT1G01010
##
          2.3
                       AT1G01010.1 AT1G01010.1
                                                                                                  AT1G01010
##
          3.4 AT1G01010.1-Protein AT1G01010.1
##
          4.5
                               <NA>
                                             <NA>
                                                                                               AT1G01010.1
```

<NA>

. . .

<NA>

<NA>

ATMG00030 protein\_coding\_gene

AT1G01010.1

ATMG00030

ATMG00030.1

ATMG00030.1, ATMG00030.1-Protein

. . .

<NA>

. . .

<NA>

<NA>

Index Derives\_from

ATMG00030.1 ATMG00030.1

ATMG00030

440.447 ATMG00030.1-Protein ATMG00030.1

##

##

##

##

## ##

##

##

5.6

. . .

438.445

439.446

441.448

442.449

```
##
             <character> <character>
##
         1.2
                                  <NA>
                 1
                                  <NA>
##
         2.3
##
         3.4
                    <NA> AT1G01010.1
##
         4.5
                    <NA>
                                  <NA>
##
         5.6
                    <NA>
                                  <NA>
##
         . . .
                     . . .
                                  . . .
##
     438.445
                    <NA>
                                  <NA>
##
     439.446
                       1
                                  <NA>
##
     440.447
                    <NA> ATMG00030.1
##
     441.448
                    <NA>
                                  <NA>
     442.449
                    <NA>
                                  <NA>
##
     -----
##
     seqinfo: 7 sequences from an unspecified genome
sp[1:4, "type"] # Subsetting of GRangesList objects is similar to GRanges objects.
## GRangesList object of length 4:
## GRanges object with 1 range and 1 metadata column:
##
       seqnames
                      ranges strand |
##
          <Rle>
                   <IRanges> <Rle> | <factor>
##
     2
           Chr1 [3631, 5899]
                                   * |
##
  GRanges object with 1 range and 1 metadata column:
##
##
       segnames
                      ranges strand | type
##
           Chr1 [3631, 5899]
##
## $3
  GRanges object with 1 range and 1 metadata column:
##
       seqnames
                      ranges strand |
##
           Chr1 [3760, 5630]
                                 * | protein
##
## ...
## <1 more element>
## seqinfo: 7 sequences from an unspecified genome
lapply(sp[1:4], length) # Looping over GRangesList objects similar to lists
## $`1`
## [1] 1
##
## $\2\
## [1] 1
##
## $`3`
## [1] 1
## $`4`
## [1] 1
```

## Transcript Ranges

```
Storing annotation ranges in TranscriptDb databases makes many operations more robust and convenient.
library(GenomicFeatures)
download.file("http://faculty.ucr.edu/~tgirke/Documents/R_BioCond/Samples/gff3.gff", "data/gff3.gff")
txdb <- makeTxDbFromGFF(file="data/gff3.gff", format="gff", dataSource="TAIR", organism="Arabidopsis th
## Warning in .extract_exons_from_GRanges(cds_IDX, gr, ID, Name, Parent, feature = "cds", : The following
     seqid start end strand
                               ID
                                               Parent Name
## 1 Chr1 3760 3913
                           + <NA> AT1G01010.1-Protein <NA>
## 2 Chr1 3996 4276
                           + <NA> AT1G01010.1-Protein <NA>
## 3 Chr1 4486 4605
                          + <NA> AT1G01010.1-Protein <NA>
## 4 Chr1 4706 5095
                           + <NA> AT1G01010.1-Protein <NA>
## 5 Chr1 5174 5326
                           + <NA> AT1G01010.1-Protein <NA>
## 6 Chr1 5439 5630
                           + <NA> AT1G01010.1-Protein <NA>
saveDb(txdb, file="./data/TAIR10.sqlite")
## TxDb object:
## # Db type: TxDb
## # Supporting package: GenomicFeatures
## # Data source: TAIR
## # Organism: Arabidopsis thaliana
## # Taxonomy ID: 3702
## # miRBase build ID: NA
## # Genome: NA
## # transcript_nrow: 28
## # exon_nrow: 113
## # cds_nrow: 99
## # Db created by: GenomicFeatures package from Bioconductor
## # Creation time: 2018-04-25 16:29:04 -0700 (Wed, 25 Apr 2018)
## # GenomicFeatures version at creation time: 1.30.3
## # RSQLite version at creation time: 2.0
## # DBSCHEMAVERSION: 1.2
txdb <- loadDb("./data/TAIR10.sqlite")</pre>
transcripts(txdb)
## GRanges object with 28 ranges and 2 metadata columns:
##
          segnames
                           ranges strand |
                                               tx id
                                                          tx name
##
                        <IRanges> <Rle> | <integer> <character>
##
      [1]
              Chr1 [ 3631,
                           5899]
                                       + |
                                                    1 AT1G01010.1
##
      [2]
              Chr1 [ 5928, 8737]
                                                   2 AT1G01020.1
##
      [3]
              Chr1 [ 6790, 8737]
                                       - |
                                                   3 AT1G01020.2
```

```
##
      [4]
              Chr1 [11649, 13714]
                                         - 1
                                                      4 AT1G01030.1
      [5]
##
              Chr2 [ 1025, 2810]
                                         + |
                                                      5 AT2G01008.1
##
      . . .
               . . .
                                . . .
                                       . . . .
                                                    . . .
##
     [24]
              ChrC [ 383, 1444]
                                         - 1
                                                     24 ATCG00020.1
##
     [25]
              ChrC [ 1717, 4347]
                                         - 1
                                                     25 ATCG00030.1
     [26]
              ChrM [11918, 12241]
                                         + |
##
                                                     26 ATMG00030.1
                              734]
##
     [27]
              ChrM [ 273,
                                         - |
                                                     27 ATMG00010.1
##
     [28]
              ChrM [ 8848, 11415]
                                         - |
                                                     28 ATMG00020.1
##
##
     seqinfo: 7 sequences (2 circular) from an unspecified genome; no seqlengths
```

```
transcriptsBy(txdb, by = "gene")
## GRangesList object of length 22:
## $AT1G01010
## GRanges object with 1 range and 2 metadata columns:
##
         segnames
                        ranges strand |
                                           {\sf tx\_id}
                                                      tx_name
##
            <Rle>
                     <IRanges> <Rle> | <integer> <character>
##
             Chr1 [3631, 5899]
     [1]
                                    + |
                                                1 AT1G01010.1
##
## $AT1G01020
## GRanges object with 2 ranges and 2 metadata columns:
##
         segnames
                        ranges strand | tx_id
##
             Chr1 [5928, 8737]
                                    - |
                                           2 AT1G01020.1
     [1]
##
     [2]
             Chr1 [6790, 8737]
                                    - |
                                            3 AT1G01020.2
##
## $AT1G01030
## GRanges object with 1 range and 2 metadata columns:
                        seqnames
##
                                              4 AT1G01030.1
             Chr1 [11649, 13714]
                                    - |
     [1]
##
## ...
## <19 more elements>
## -----
## seqinfo: 7 sequences (2 circular) from an unspecified genome; no seqlengths
exonsBy(txdb, by = "gene")
## GRangesList object of length 22:
## $AT1G01010
## GRanges object with 6 ranges and 2 metadata columns:
##
         segnames
                        ranges strand |
                                          exon id
                                                    exon name
                     <IRanges> <Rle> | <integer> <character>
##
            <Rle>
##
             Chr1 [3631, 3913]
                                    + |
     [1]
                                                1
                                                          <NA>
             Chr1 [3996, 4276]
##
     [2]
                                    + |
                                                2
                                                          <NA>
##
     [3]
             Chr1 [4486, 4605]
                                    + |
                                                3
                                                          <NA>
##
     [4]
             Chr1 [4706, 5095]
                                    + |
                                                4
                                                          <NA>
##
                                                5
                                                          <NA>
     [5]
             Chr1 [5174, 5326]
                                    + |
##
     [6]
             Chr1 [5439, 5899]
                                    + |
                                                6
                                                          <NA>
##
## $AT1G01020
  GRanges object with 12 ranges and 2 metadata columns:
##
                         ranges strand | exon_id exon_name
          segnames
              Chr1 [5928, 6263]
##
      [1]
                                     - |
                                               7
                                                       <NA>
              Chr1 [6437, 7069]
##
      [2]
                                     - |
                                               8
                                                       <NA>
##
      [3]
              Chr1 [6790, 7069]
                                     - |
                                               9
                                                       <NA>
##
      [4]
              Chr1 [7157, 7232]
                                     - 1
                                              10
                                                       <NA>
              Chr1 [7157, 7450]
##
      [5]
                                     - |
                                              11
                                                       <NA>
##
      . . .
              . . .
##
      [8]
                                     - 1
              Chr1 [7762, 7835]
                                              14
                                                       <NA>
              Chr1 [7942, 7987]
                                     - |
##
      [9]
                                              15
                                                       <NA>
##
     Γ107
              Chr1 [8236, 8325]
                                     - |
                                              16
                                                      <NA>
##
     [11]
              Chr1 [8417, 8464]
                                     - |
                                              17
                                                      <NA>
              Chr1 [8571, 8737]
##
     [12]
                                     - |
                                              18
                                                       <NA>
```

##

```
## $AT1G01030
## GRanges object with 2 ranges and 2 metadata columns:
##
                          ranges strand | exon id exon name
             Chr1 [11649, 13173]
##
                                       - |
                                                19
                                                         <NA>
     [1]
##
     [2]
             Chr1 [13335, 13714]
                                       - |
                                                20
                                                         <NA>
##
## <19 more elements>
## seqinfo: 7 sequences (2 circular) from an unspecified genome; no seqlengths
```

#### txdb from BioMart

Alternative sources for creating txdb databases are BioMart, Bioc annotation packages, UCSC, etc. The following shows how to create a txdb from BioMart.

```
library(GenomicFeatures); library("biomaRt")
txdb <- makeTxDbFromBiomart(biomart = "plants_mart", dataset = "athaliana_eg_gene", host="plants.ensemb")</pre>
```

The following steps are useful to find out what is availble in BioMart.

```
listMarts() # Lists BioMart databases
listMarts(host="plants.ensembl.org")
mymart <- useMart("plants_mart", host="plants.ensembl.org") # Select one, here plants_mart_25
listDatasets(mymart) # List datasets available in the selected BioMart database
mymart <- useMart("plants_mart", dataset="athaliana_eg_gene", host="plants.ensembl.org")
listAttributes(mymart) # List available features
getBM(attributes=c("ensembl_gene_id", "description"), mart=mymart)[1:4,]</pre>
```

#### Efficient Sequence Parsing

#### getSeq

The following parses all annotation ranges provided by a GRanges object (e.g. gff) from a genome sequence stored in a local file.

```
gff <- gff[values(gff)$type != "chromosome"] # Remove chromosome ranges
rand <- DNAStringSet(sapply(unique(as.character(seqnames(gff))), function(x) paste(sample(c("A","T","G"
writeXStringSet(DNAStringSet(rand), "./data/test")
getSeq(FaFile("./data/test"), gff)</pre>
```

```
##
     A DNAStringSet instance of length 442
##
        width seq
##
     [1] 2269 ATGTGTAGGCAATCAGTCACCTTGCATCGATG...CTAGGATAGAGACCACACACAGCAAGCGACTTAG Chr1
     [2] 2269 ATGTGTAGGCAATCAGTCACCTTGCATCGATG...CTAGGATAGAGACCACACACAGCAAGCGACTTAG Chr1
##
     [3] 1871 GAGGCAATATCTTCATGGAGCAAATAAAAGCT...GATCCGATCGACCACCAAGGGATGGGGCCCCG Chr1
##
##
     [4]
          283 ATGTGTAGGCAATCAGTCACCTTGCATCGATG...GAGGCAGATCTTTCAGCGGACTACAGCATAAC Chr1
##
     [5]
          129 ATGTGTAGGCAATCAGTCACCTTGCATCGATG...CGATCTACCCATCAAGCTCTGGCTCACGCCAT Chr1
##
          324 CTGTTAGAGATTTCGGGCACCGACTGGGAGCT...CCACGGTGATCGAAAAGCTTTAGTTG ChrM
## [438]
          324 CTGTTAGAGATTTCGGGCACCGACTGGGAGCT...CCACGGTGATCGAAAAGCTTTAGTTG ChrM
## [439]
## [440]
          324 CTGTTAGAGATTTCGGGCACCGACTGGGAGCT...CCACGGTGATCGAAACTCGAAAAGCTTTAGTTG ChrM
          324 CTGTTAGAGATTTCGGGCACCGACTGGGAGCT...CCACGGTGATCGAAAAGCTTTAGTTG ChrM
## [441]
## [442]
          324 CTGTTAGAGATTTCGGGCACCGACTGGGAGCT...CCACGGTGATCGAACTCGAAAAGCTTTAGTTG ChrM
```

#### extractTranscriptSeqs

Sequences composed of several ranges, such as transcripts (or CDSs) with several exons, can be parsed with extractTranscriptSeqs. Note: the following expects the genome sequence in a file called mygenome.fasta and a valid txdb defining the ranges for that genome.

```
library(GenomicFeatures); library(Biostrings); library(Rsamtools)
extractTranscriptSeqs(FaFile("mygenome.fasta"), exonsBy(txdb, "tx", use.names=TRUE))
```

## Homework 6

## HW6a - Demultiplexing

Write a demultiplexing function that accepts any number of barcodes and splits a FASTQ file into as many subfiles as there are barcodes. At the same time the function should remove low quality tails from the reads. The following function accomplishes the first step. Expand this function so that it performs the second step as well.

#### HW6b - Sequence Parsing

- Download GFF from Halobacterium sp here
- Download genome sequence from *Halobacterium sp* here
- Task 1 Extract gene ranges, parse their sequences from genome and translate them into proteins
- Task 2 Reduce overlapping genes and parse their sequences from genome
- Task 3 Generate intergenic ranges and parse their sequences from genome

#### Useful commands

```
download.file("ftp://ftp.ncbi.nlm.nih.gov/genomes/archive/old_genbank/Bacteria/Halobacterium_sp_uid217/
download.file("ftp://ftp.ncbi.nlm.nih.gov/genomes/archive/old_genbank/Bacteria/Halobacterium_sp_uid217/
chr <- readDNAStringSet("data/AE004437.fna")
gff <- import("data/AE004437.gff")
gffgene <- gff[values(gff)[,"type"]=="gene"]
gene <- DNAStringSet(Views(chr[[1]], IRanges(start(gffgene), end(gffgene))))
names(gene) <- values(gffgene)[,"locus_tag"]
pos <- values(gffgene[strand(gffgene) == "+"])[,"locus_tag"]
p1 <- translate(gene[names(gene) %in% pos])
names(p1) <- names(gene[names(gene) %in% pos])</pre>
```

```
neg <- values(gffgene[strand(gffgene) == "-"])[,"locus_tag"]
p2 <- translate(reverseComplement(gene[names(gene) %in% neg]))
names(p2) <- names(gene[names(gene) %in% neg])
writeXStringSet(c(p1, p2), "./data/mypep.fasta")</pre>
```

#### Homework submission

Submit the homework results in one well structured and annotated R script to the instructor. The script should include instructions on how to use the functions.

#### Due date

This homework is due on Thu, May 4th at 6:00 PM.

#### **Homework Solutions**

See here.

## Session Info

```
sessionInfo()
## R version 3.4.4 (2018-03-15)
## Platform: x86_64-pc-linux-gnu (64-bit)
## Running under: Ubuntu 16.04.4 LTS
##
## Matrix products: default
## BLAS: /usr/lib/libblas/libblas.so.3.6.0
## LAPACK: /usr/lib/lapack/liblapack.so.3.6.0
##
## locale:
## [1] LC_CTYPE=en_US.UTF-8
                                   LC_NUMERIC=C
                                                               LC_TIME=en_US.UTF-8
   [4] LC_COLLATE=en_US.UTF-8
                                   LC_MONETARY=en_US.UTF-8
                                                               LC_MESSAGES=en_US.UTF-8
## [7] LC_PAPER=en_US.UTF-8
                                   LC_NAME=C
                                                               LC_ADDRESS=C
## [10] LC_TELEPHONE=C
                                   LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
##
## attached base packages:
## [1] grid
                  stats4
                            parallel methods
                                                           graphics grDevices utils
                                                stats
                                                                                         datasets
## [10] base
##
## other attached packages:
## [1] GenomicFeatures_1.30.3
                                   AnnotationDbi 1.40.0
                                                               rtracklayer_1.38.3
## [4] systemPipeR_1.12.0
                                   ShortRead_1.36.1
                                                               GenomicAlignments_1.14.2
## [7] SummarizedExperiment_1.8.1 DelayedArray_0.4.1
                                                               matrixStats_0.52.2
                                                               GenomicRanges_1.30.3
## [10] Biobase_2.38.0
                                   Rsamtools_1.30.0
## [13] GenomeInfoDb 1.14.0
                                   BiocParallel 1.12.0
                                                               ggseqlogo 0.1
                                                               XVector_0.18.0
## [16] seqLogo_1.44.0
                                   Biostrings_2.46.0
## [19] IRanges_2.12.0
                                   S4Vectors_0.16.0
                                                               BiocGenerics 0.24.0
                                   limma_3.34.1
                                                               BiocStyle_2.6.0
## [22] ggplot2_2.2.1
```

```
##
## loaded via a namespace (and not attached):
    [1] Category 2.44.0
                                bitops 1.0-6
                                                       bit64 0.9-7
                                                                               RColorBrewer 1.1-2
    [5] progress_1.1.2
                                httr_1.3.1
                                                       rprojroot_1.2
                                                                               Rgraphviz_2.22.0
##
##
    [9] tools_3.4.4
                                backports_1.1.1
                                                       R6_2.2.2
                                                                               DBI 0.7
## [13] lazyeval 0.2.1
                                colorspace_1.3-2
                                                                               RMySQL 0.10.14
                                                       prettyunits 1.0.2
  [17] bit 1.1-12
                                compiler 3.4.4
                                                       sendmailR_1.2-1
                                                                               graph_1.56.0
  [21] labeling_0.3
                                scales_0.5.0
                                                       checkmate_1.8.5
                                                                               BatchJobs_1.7
        genefilter_1.60.0
  [25]
                                RBGL_1.54.0
                                                       stringr_1.2.0
                                                                               digest_0.6.12
  [29]
        rmarkdown_1.8
                                AnnotationForge_1.20.0
                                                       base64enc_0.1-3
                                                                               pkgconfig_2.0.1
  [33] htmltools_0.3.6
                                rlang_0.2.0
                                                       RSQLite_2.0
                                                                               BBmisc_1.11
        GOstats_2.44.0
                                hwriter_1.3.2
                                                       RCurl_1.95-4.8
                                                                               magrittr_1.5
  [37]
       GO.db_3.5.0
  [41]
                                GenomeInfoDbData_1.0.0 Matrix_1.2-14
                                                                               Rcpp_0.12.13
##
        munsell_0.4.3
                                                       yaml_2.1.14
  [45]
                                stringi_1.1.6
                                                                                edgeR_3.20.1
## [49] zlibbioc_1.24.0
                                plyr_1.8.4
                                                       blob_1.1.0
                                                                               lattice_0.20-35
  [53] splines_3.4.4
                                annotate_1.56.1
                                                       locfit_1.5-9.1
                                                                               knitr_1.17
        pillar_1.2.1
                                                                               biomaRt_2.34.2
  [57]
                                rjson_0.2.15
                                                       codetools_0.2-15
  [61] XML 3.98-1.9
                                evaluate 0.10.1
                                                       latticeExtra_0.6-28
                                                                               data.table_1.10.4-3
  [65] gtable_0.2.0
                                                                               survival_2.42-3
                                assertthat_0.2.0
                                                       xtable_1.8-2
  [69] tibble 1.4.2
                                pheatmap_1.0.8
                                                       memoise_1.1.0
                                                                               brew_1.0-6
  [73] GSEABase_1.40.0
```

## References

Huber, Wolfgang, Vincent J Carey, Robert Gentleman, Simon Anders, Marc Carlson, Benilton S Carvalho, Hector Corrada Bravo, et al. 2015. "Orchestrating High-Throughput Genomic Analysis with Bioconductor." *Nat. Methods* 12 (2): 115–21. doi:10.1038/nmeth.3252.

Lawrence, Michael, Wolfgang Huber, Hervé Pagès, Patrick Aboyoun, Marc Carlson, Robert Gentleman, Martin T Morgan, and Vincent J Carey. 2013. "Software for Computing and Annotating Genomic Ranges." *PLoS Comput. Biol.* 9 (8): e1003118. doi:10.1371/journal.pcbi.1003118.