Basics

Contents

1	General terminology	2
2	Markers	2
3	Data	2
4	Public databases	3
5		3
	5.1 Python	3
	5.2 GRanges	6
	5.3 DNAStrings	16
	5.4 Data classes	18
	5.5 Expression array archives	23
	5.6 Obtaining the ExpressionSet for an EMBL series	23
	5.7 Storing data	24
	5.8 Reference genomes	
	5.9 Annotations	

1 General terminology

Term	Description
Genome	The entire DNA content of an individual's chromosome the the articular
Marker	A measurable variation (polymorphism) at the DNA level
Locus	A genomic region that explains phenotypic differences due to genetic polymorphisms
Gene	A basic unit of heredity and a sequence of nucleotides in DNA that encodes the synthesis of a gene product
Isoform	Each mRNA maps to a gene but one gene can be mapped by numerous mRNAs. A gene encodes different mRNA transcripts, or isoforms
Variant	A locus in the genome where there are differences between individuals
Allele	One of the possible bases/sequences that can occur at the variant
SNP	Single nucleotide polymorphism: only one nucleotide substitution within a short DNA sequence A type of variant
SNP array	Short DNA sequences, with their variant nucleotides at their ends, who constitute probes
VNTRs	Variable number of tandem repeats: multiple copies of the same sequence of DNA A type of variant
Haplotype	Blocks of correlated SNPs that tend to be inherited together (e.g., HLA alleles in the major histocompatibility complex)
QTL	A locus that correlates with the variation of a quantitative trait in the phenotype of a population
Linkage-	It is a measure of the statistical association between the allele frequencies at two different
Disequilibrium	loci in a population
Transcriptome	The complete set of RNA transcripts that are produced by the genome
Transcript	Refers to which exons are spliced to form the RNA molecule
Epigenome	Non-sequence modifications of DNA that are heritable (e.g., methylation)
Exposome	The measure of all the exposures of an individual in a lifetime

2 Markers

In general terms, the markers are used to identify regions harboring quantitative trait loci (QTL).

Types:

- Restriction fragment length polymorphism (RFLP)
- Random amplification of polymorphic DNA (RAPD)
- Amplified fragment length polymorphism (AFLP)
- Single-stranded conformation polymorphism (SSCP)
- Copy number variation (CNV)
- Microsatellites:
 - A type of VNTR where the length of the repeating unit is less than five base pairs
 - A microsatellite can have many alleles (10-30)
 - They are widely used in QTL (quantitative trait locus) mapping projects
- Single nucleotide polymorphism (SNP):
 - 0: homozygous (AA, for the minor allele A)
 - 1: heterozygous (AC/CA, for the minor allele A)
 - 2: variant homozygous (CC, for the minor allele A)

3 Data

- Transcriptomic data:
 - Generation methods:

- * Microarray method: mRNA is collected and converted back to cDNAs which are then dyed and hybridized on a chip containing probes
- * RNA-seq method: the application of high throughput sequencing to RNA
- Epigenomic data:
 - Methylomic data: bounded by the number of CpG sequences in the genome
 - Others
- Genomic data:
 - Strands * 5' to 3' (+) * 3' to 5' (-)

4 Public databases

- dbGaP:
 - genomic, epigenomic, somatic mutations, transcriptomic and microbiomic data, with associated phenotypes
 - Permission is required
- EGA: same as above
- GEO:
 - Only transcriptomic data from microarray and RNA-seq
 - Data can be directly retrieved with Bioconductor
- 1000 Genomes: genomic data from the sequencing of 2,504 individuals from 26 different ancestries
- GTEx: collected transcriptomic data for 714 donors, 635 of which were also genotyped
- TCGA: high-throughput DNA and RNA sequencing, SNP, DNA methylation, and reverse-protein arrays
 of 33 cancers

5 Data management

5.1 Python

5.1.1 Read, count, and reverse complement

Read genomic data:

GGGCGGCGACCTCGCGGGTT

Number of bases in the strand:

```
def count(genome):
    counts = {'A': 0, 'C': 0, 'G': 0, 'T': 0}
    for base in genome:
        counts[base] += 1
    print(counts)
```

```
count(lambda_virus_genome)

## {'A': 12334, 'C': 11362, 'G': 12820, 'T': 11986}

Get the antisense strand:

def reverseComplement(senseStrand):
    complement = {'A': 'T', 'T': 'A', 'C': 'G', 'G': 'C', 'N': 'N'}
    antisenseStrand = ''
    for base in senseStrand:
        antisenseStrand = complement[base] + antisenseStrand
    return antisenseStrand

lambda_virus_genomeAntisense = reverseComplement(lambda_virus_genome)
print(lambda_virus_genomeAntisense[:20])
```

CGTAACCTGTCGGATCACCG

5.1.2 Quality scores

Read genome reads (sequences and quality values):

```
def readReads(filename):
    sequences = []
   qualities = []
   with open(filename, 'r') as f:
        while True:
            f.readline()
            seq = f.readline().rstrip()
            f.readline()
            qual = f.readline().rstrip()
            if len(seq) == 0:
                break
            sequences.append(seq)
            qualities.append(qual)
   return sequences, qualities
human_seqs, human_quals = readReads('Data/SRR835775_1.first1000.fastq')
print(human_seqs[1])
```

TAACCCTAACCTAACCCTAACCCTAACCCTAACCCTAACCCTAACCCTAACCCTAACCCTAACCCTAACCCTAACCCTAACCCTAACCCTAACCCTAACCCTAAC

Convert quality values to numbers:

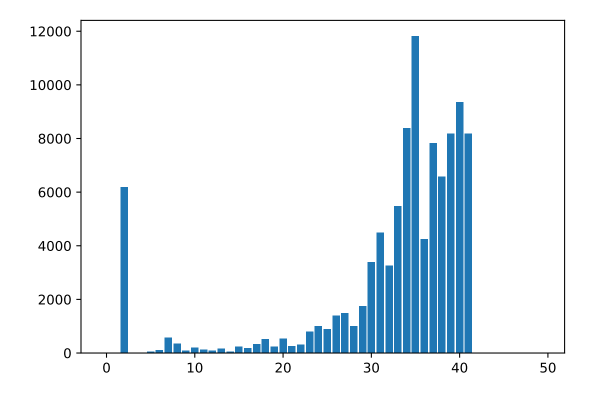
```
def phred33ToQ(qual):
    return ord(qual) - 33
```

Create a histogram of the frequency of qualities:

```
import matplotlib.pyplot as plt
def createHist(qualities):
    hist = [0] * 50
    for qual in qualities:
        for phred in qual:
```

```
q = phred33ToQ(phred)
    hist[q] += 1
    return hist
h = createHist(human_quals)
plt.bar(range(len(h)), h)

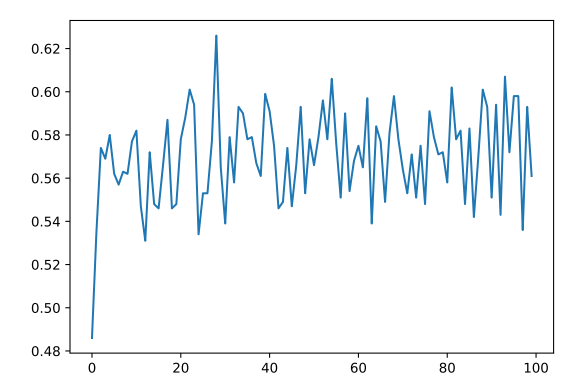
## <BarContainer object of 50 artists>
plt.show()
```



5.1.3 Find GCs

Find GCs by position:

```
gc = findGCByPos(human_seqs)
plt.plot(range(len(gc)), gc)
plt.show()
```



5.2 GRanges

Let's start by creating a sample gene:

```
# Import packages
library(GenomicRanges)

# Create an example gene
gen1 <- GRanges("chr1", IRanges(1000001, 1001000), strand = "+")

Now, let's check its specification:
# Get gene spec
## Start
start(gen1)

## [1] 1000001

## End
end(gen1)</pre>
```

[1] 1001000

```
## Width
width(gen1)
## [1] 1000
## Strand
strand(gen1)
## factor-Rle of length 1 with 1 run
    Lengths: 1
     Values : +
## Levels(3): + - *
## The 'metadata columns', any information stored alongside each range
mcols(gen1)
## DataFrame with 1 row and 0 columns
## IRanges
ranges(gen1)
## IRanges object with 1 range and 0 metadata columns:
##
             start
                         end
                                 width
         <integer> <integer> <integer>
##
          1000001 1001000
##
     [1]
                                  1000
## The chromosomes for each ranges
seqnames(gen1)
## factor-Rle of length 1 with 1 run
    Lengths:
##
    Values : chr1
## Levels(1): chr1
## The possible chromosomes
seqlevels(gen1)
## [1] "chr1"
## The lengths for each chromosome
seqlengths(gen1)
## chr1
##
    NA
```

5.2.1 Intra-range methods

Affects ranges independently.

function	description
shift	moves left/right
narrow	narrows by relative position within range
resize	resizes to width, fixing start for $+$, end for $-$
flank	returns flanking ranges to the left $+$, or right $-$
promoters	similar to flank
restrict	restricts ranges to a start and end position
trim	trims out of bound ranges
+/-	expands/contracts by adding/subtracting fixed amount

function	description	
*	zooms in (positive) or out (negative) by multiples	

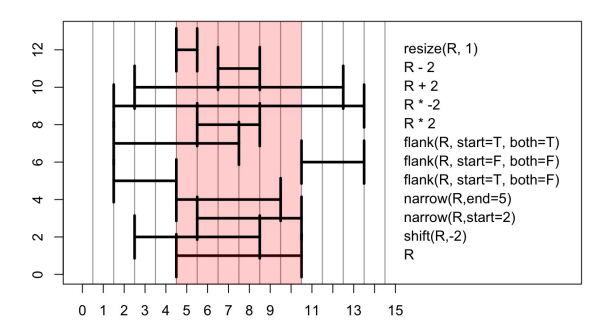


Figure 1: Intra-range methods 1

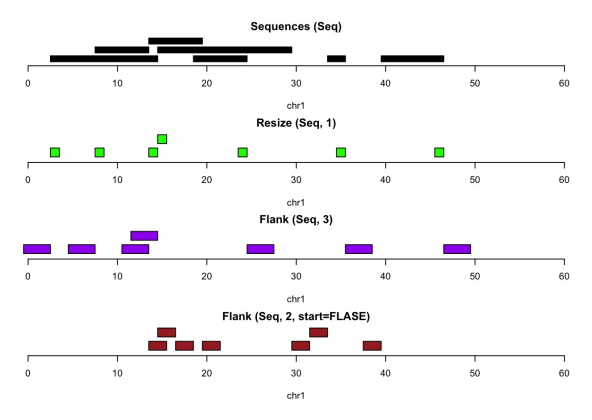


Figure 2: Intra-range methods 2

5.2.2 Inter-range methods

Affects ranges as a group.

function	description
range	one range, leftmost start to rightmost end
reduce	cover all positions with only one range
gaps	uncovered positions within range
disjoin	breaks into discrete ranges based on original starts/ends

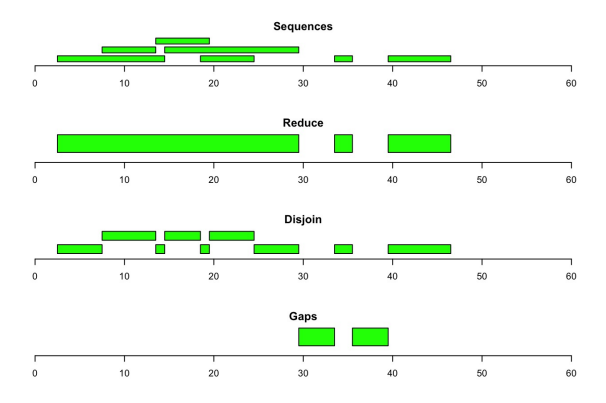
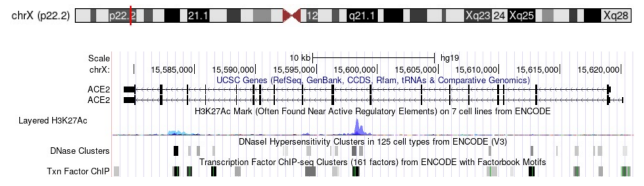


Figure 3: Inter-range methods

5.2.3 ENCODE project experiment

Scenario: A ChIP-seq experiment to identify the ESRRA nuclear protein binding sites in HepG2 cell line (of liver origin) and GM12878 cell line (of lymphoblastoid origin).

ESRRA: Estrogen Related Receptor Alpha



Let's start by importing and preprocessing the data:

```
# Load data
library(ERBS)
data(HepG2)
data(GM12878)

# Check data
HepG2[1:5, 4:5]
```

```
## GRanges object with 5 ranges and 2 metadata columns:
##
                             ranges strand | signalValue
                                                               pValue
        segnames
##
            <Rle>
                            <IRanges> <Rle> |
                                                 <numeric> <numeric>
##
             chr2 20335378-20335787
     [1]
                                           * |
                                                    58.251
                                                               75.899
##
     [2]
           chr20
                        328285-329145
                                           * |
                                                     10.808
                                                               69.685
##
            chr6 168135432-168136587
                                           * |
                                                     17.103
                                                               54.311
     [3]
##
           chr19 1244419-1245304
                                            * |
                                                     12.427
                                                               43.855
     ۲4٦
                    64071828-64073069
                                           * |
                                                     10.850
                                                               40.977
##
     [5]
           chr11
##
     seqinfo: 93 sequences (1 circular) from hg19 genome
# Extract data as Rle data
## Extract seqnames (chromosomes)
chr <- seqnames(HepG2)[1:5,]</pre>
## Extract ranges
r <- ranges(HepG2)[1:5,]
## Extract values
vals <- values(HepG2)[1:5,]</pre>
# Convert extracted data to character
chr <- as.character(chr)</pre>
chr
## [1] "chr2" "chr20" "chr6" "chr19" "chr11"
r <- as.data.frame(r)
         start
                     end width
## 1 20335378 20335787
                           410
       328285
                  329145
## 3 168135432 168136587 1156
     1244419
               1245304
                          886
## 5 64071828 64073069 1242
vals <- as.data.frame(vals)</pre>
vals[, 4:7]
     signalValue pValue
                              qValue peak
## 1
         58.251 75.899 6.143712e-72 195
## 2
         10.808 69.685 5.028065e-66 321
## 3
         17.103 54.311 7.930665e-51 930
## 4
          12.427 43.855 1.359756e-40
                                      604
          10.850 40.977 7.333863e-38
                                      492
# Cross-tab
table(chr)
## chr
## chr11 chr19 chr2 chr20 chr6
##
       1
           1
                   1
                         1
# Subset operation
chr20 <- HepG2[seqnames(HepG2) == "chr20", ]</pre>
chr20[1:5, 7]
## GRanges object with 5 ranges and 1 metadata column:
##
                             ranges strand |
         seqnames
```

```
##
            <Rle>
                           <IRanges> <Rle> | <integer>
##
            chr20
                      328285-329145
                                          * |
     Г17
                                                     321
##
            chr20 22410891-22411863
                                          * |
                                                     660
                                                     315
##
     [3]
            chr20 56039583-56040249
                                          * |
##
     [4]
            chr20 16455811-16456232
                                          * |
                                                     199
##
     [5]
            chr20
                    3140243-3140774
                                                     315
                                          * |
##
##
     seqinfo: 93 sequences (1 circular) from hg19 genome
# Re-order the data by segnames and ranges
HepG2_ordered = HepG2[order(HepG2),]
HepG2_ordered[1:5, 7]
## GRanges object with 5 ranges and 1 metadata column:
##
         seqnames
                              ranges strand |
##
            <Rle>
                           <IRanges>
                                     <Rle> | <integer>
##
                    9294157-9294957
     [1]
             chr1
                                          * |
##
     [2]
             chr1 15911325-15911873
                                          * |
                                                     264
##
     [3]
             chr1 16339152-16339862
                                          * |
                                                     332
##
     [4]
             chr1 26146200-26147004
                                          * |
                                                     337
##
             chr1 27264576-27265423
                                                     191
     [5]
##
##
     seqinfo: 93 sequences (1 circular) from hg19 genome
Now, we are going to find binding sites and extract their ranges:
# Find binding sites common to both HepG2 and GM12878
res = findOverlaps(HepG2, GM12878)
res[1]
## Hits object with 1 hit and 0 metadata columns:
##
         queryHits subjectHits
##
         <integer>
                     <integer>
##
     [1]
                 1
                             12
##
     queryLength: 303 / subjectLength: 1873
HepG2[1]
## GRanges object with 1 range and 7 metadata columns:
##
         seqnames
                              ranges strand |
                                                   name
                                                             score
                                                                          col
##
                           <IRanges> <Rle> | <numeric> <integer> <logical>
            <Rle>
##
             chr2 20335378-20335787
                                          * |
                                                      NA
                                                                 0
##
         signalValue
                        pValue
                                     qValue
                                                 peak
##
           <numeric> <numeric>
                                  <numeric> <integer>
##
     Г17
              58.251
                        75.899 6.14371e-72
##
##
     seqinfo: 93 sequences (1 circular) from hg19 genome
GM12878[12]
  GRanges object with 1 range and 7 metadata columns:
##
##
         seqnames
                              ranges strand |
                                                    name
                                                             score
##
            <Rle>
                           <IRanges> <Rle> | <numeric> <integer> <logical>
             chr2 20335155-20336108
##
     [1]
                                          * |
                                                      16
##
         signalValue
                        pValue
                                   qValue
                                               peak
##
           <numeric> <numeric> <integer>
##
     [1]
               55.95 223.815
                                                438
                                       32
```

```
##
##
     seqinfo: 93 sequences (1 circular) from hg19 genome
# Ranges from the guery for which we found a hit in the subject
index = queryHits(res)
erbs = HepG2[index,]
erbs[1:5]
## GRanges object with 5 ranges and 7 metadata columns:
##
         seqnames
                              ranges strand |
                                                                           col
                                                     name
                                                              score
##
            <Rle>
                           <IRanges>
                                       <Rle> | <numeric> <integer> <logical>
##
             chr2 20335378-20335787
     [1]
                                                                          <NA>
                                           * |
                                                       NA
##
     [2]
            chr20
                       328285-329145
                                                                   0
                                                       NA
                                                                          <NA>
##
     [3]
                                                       NA
                                                                   0
                                                                          <NA>
            chr19
                     1244419-1245304
##
     [4]
            chr11 64071828-64073069
                                                       NA
                                                                   0
                                                                          <NA>
##
     [5]
             chr2 16938364-16938840
                                                       NA
                                                                   0
                                                                          <NA>
##
         signalValue
                         pValue
                                      qValue
                                                   peak
##
           <numeric> <numeric>
                                   <numeric> <integer>
##
     [1]
              58.251
                         75.899 6.14371e-72
##
     [2]
              10.808
                         69.685 5.02806e-66
                                                    321
##
     [3]
              12.427
                         43.855 1.35976e-40
                                                    604
              10.850
                                                    492
##
     [4]
                         40.977 7.33386e-38
##
     [5]
              12.783
                         38.004 5.36029e-35
                                                    255
##
     seqinfo: 93 sequences (1 circular) from hg19 genome
# Extract only the ranges
erbs = granges(erbs)
erbs[1:5]
## GRanges object with 5 ranges and 0 metadata columns:
##
         seqnames
                              ranges strand
##
            <Rle>
                           <IRanges>
                                       <Rle>
##
     [1]
             chr2 20335378-20335787
##
     [2]
            chr20
                       328285-329145
##
     [3]
            chr19
                     1244419-1245304
##
     [4]
            chr11 64071828-64073069
##
             chr2 16938364-16938840
##
     seqinfo: 93 sequences (1 circular) from hg19 genome
```

5.2.3.1 Acquisition of gene transcription start sites Using the binding sites, we want to know which genes are preceded by each binding site.

Note: This information is strand-based, meaning that "preceding" depends on the direction of which we read the genome.

```
# Define human genes
library(Homo.sapiens)
ghs = genes(Homo.sapiens)
ghs[1:5]
  GRanges object with 5 ranges and 1 metadata column:
##
           seqnames
                                                            GENEID
                                  ranges strand |
##
              <Rle>
                               <IRanges> <Rle> | <CharacterList>
                                              - 1
##
         1
              chr19
                      58858172-58874214
                                                                 1
##
        10
               chr8
                      18248755-18258723
                                              + |
                                                                10
```

```
##
       100
              chr20
                      43248163-43280376
                                                               100
##
      1000
              chr18
                      25530930-25757445
                                              - 1
                                                              1000
                                                             10000
##
     10000
               chr1 243651535-244006886
                                              - 1
##
     seqinfo: 93 sequences (1 circular) from hg19 genome
# First three ERBS ranges
erbs[1:3]
## GRanges object with 3 ranges and 0 metadata columns:
##
         segnames
                             ranges strand
##
            <Rle>
                           <IRanges>
                                     <Rle>
##
     [1]
             chr2 20335378-20335787
            chr20
##
     [2]
                      328285-329145
            chr19
##
     [3]
                    1244419-1245304
##
     seqinfo: 93 sequences (1 circular) from hg19 genome
# For each range in ERBS, find the closest preceding genes in GHS
index = precede(erbs, ghs)
# First three closest preceding genes in GHS to ERBS ranges
ghs[index[1:3]]
## GRanges object with 3 ranges and 1 metadata column:
##
           seqnames
                               ranges strand |
                                                          GENEID
##
                            <IRanges> <Rle> | <CharacterList>
              <Rle>
##
      9741
               chr2 20232411-20251789
                                            - |
                                                           9741
##
     57761
              chr20
                        361308-378203
                                            + |
                                                          57761
##
     90007
              chr19
                      1248552-1259142
                                            + |
                                                          90007
##
     seqinfo: 93 sequences (1 circular) from hg19 genome
# Distance between binding sites and nearest preceding genes
distance(erbs, ghs[index])[1:3]
## [1] 83588 32162 3247
Now, let's find the transcription start site nearest to each binding site:
# Get all TSS of human genome
tssgr = resize(ghs, 1)
tssgr[1:3]
## GRanges object with 3 ranges and 1 metadata column:
##
         segnames
                     ranges strand |
##
            <Rle> <IRanges> <Rle> | <CharacterList>
##
       1
            chr19 58874214
                                  - 1
                                                    1
             chr8 18248755
##
      10
                                                   10
##
     100
            chr20 43280376
                                                  100
##
     seqinfo: 93 sequences (1 circular) from hg19 genome
# Distance between erbs binding sites and nearest TSS
d = distanceToNearest(erbs, tssgr)
d[1:3]
## Hits object with 3 hits and 1 metadata column:
         queryHits subjectHits | distance
##
```

```
##
         <integer>
                     <integer> | <integer>
##
     [1]
                          22817 |
                                      83588
                 1
##
     [2]
                 2
                          19883 |
                                        914
##
     [3]
                 3
                          13475 |
                                       2669
##
     queryLength: 75 / subjectLength: 23056
##
dists = values(d)$distance
# Get the TSS of genes that are less than 1000 bases away from the erbs BS
index = subjectHits(d)[dists < 1000]</pre>
index[1:3]
## [1] 19883 6316 18488
tssgr[index, ]
## GRanges object with 41 ranges and 1 metadata column:
##
            seqnames
                        ranges strand |
                                                   GENEID
##
               <Rle> <IRanges> <Rle> | <CharacterList>
##
      80023
               chr20
                        327370
                                     + |
                                                    80023
                                     + |
##
       2101
               chr11 64073044
                                                     2101
##
       7086
                chr3 53290130
                                     - 1
                                                     7086
##
       5478
                chr7 44836241
                                     + |
                                                     5478
##
     286262
                chr9 140095163
                                     - |
                                                   286262
##
                 . . .
        . . .
                            . . .
                                                      . . .
                                   . . . .
##
      55090
               chr17 17380300
                                                    55090
                                     + |
##
      11165
               chr6 34360457
                                     - 1
                                                    11165
##
                chr1 26146397
                                     + |
      56181
                                                    56181
##
     347734
                chr6 44225283
                                     - |
                                                   347734
                chr1 110198698
##
       2948
                                     + |
                                                     2948
##
     seqinfo: 93 sequences (1 circular) from hg19 genome
##
# Extract those genes IDs
keys = as.character(values(tssgr[index])$GENEID)
keys[1:3]
## [1] "80023" "2101" "7086"
# Extract the symbole and names of those IDs
res = select(
 Homo.sapiens,
 keys = keys,
  columns = c("SYMBOL", "GENENAME"),
 keytype = "GENEID"
)
res[1:3,]
    GENEID
                                    GENENAME SYMBOL
## 1 80023
                                 neurensin 2 NRSN2
## 2
       2101 estrogen related receptor alpha ESRRA
## 3
       7086
                               transketolase
```

These genes are the genes that are:

- common between both HepG2 and GM12878 cell lines;
- their TSS are close (<1000 bases far from) to the ESRRA nuclear protein BS;

• as a result, most probably activated by estrogen via Estrogen Related Receptor Alpha (ESRRA).

Conclusion: So basically, we went from the protein binding sites of a specific nuclear protein to the genes that are activated via it.

5.3 DNAStrings

Some other functions:

Let's start by creating a sample DNA string:

```
library(Biostrings)
# Define a DNAString
dna <- DNAString("TCGAGCAAT")</pre>
## 9-letter DNAString object
## seq: TCGAGCAAT
In DNAStrings there are the four bases (ACGT), the wild card or unknown base (N), and the dash representing
a gap (-).
# Create a set of DNAStrings (DNAStringSet)
set1 <- DNAStringSet(c("TCA", "AAATCG", "ACGTGCCTA", "CGCGCA", "GTT", "TCA"))</pre>
set1
## DNAStringSet object of length 6:
##
       width seq
## [1]
           3 TCA
           6 AAATCG
## [2]
## [3]
           9 ACGTGCCTA
           6 CGCGCA
## [4]
## [5]
           3 GTT
## [6]
           3 TCA
# Extract a base from a seq
set1[[1]][2]
## 1-letter DNAString object
## seq: C
\# Number of DNAStrings in the set
length(set1)
## [1] 6
# Width of each DNAString in the set
width(set1)
## [1] 3 6 9 6 3 3
```

function	description
duplicated	detect which sequences are duplicated
unique	keep only unique sequences
sort	sort sequences alphabetically
letterFrequency	find the freq of one character
dinucleotideFrequency	find the freq of all dinucleotides
trinucleotide Frequency	find the freq of all trinucleotides
countPattern	find the freq of a pattern

function	description
matchPattern	find the locations of a pattern
reverseComplement	find the reverse complement of a DNAString
translate	find the amino acid translation of a DNAString

Let's check a specific pattern in our DNAString sample:

subject: ATGCGCGCGCTA

3

5

start end width

5

7

views:

[1]

[2]

[3]

##

##

##

##

```
# Check for a pattern in a DNAString
dna_seq <- DNAString("ATGCGCGCGGCTA")</pre>
matchPattern("CGC", dna_seq)
## Views on a 13-letter DNAString subject
## subject: ATGCGCGCGCTA
## views:
##
         start end width
##
     [1]
             4
                  6
                        3 [CGC]
     [2]
                        3 [CGC]
##
             6
                  8
Because DNA is double-stranded, when checking for a pattern, you should also check for its complement:
# Check for a the reverse complement of a pattern in a DNAString
matchPattern(reverseComplement(DNAString("CGC")), dna_seq)
## Views on a 13-letter DNAString subject
```

A somehow similar process goes with DNAStringSets, only with a different command name:

```
# Check for a pattern in a DNAStringSet
set2 <- DNAStringSet(c("AACCGGTTTCGA", "CGATCGCGCCGG"))
vmatchPattern("CG", set2)</pre>
```

```
## MIndex object of length 2
## [[1]]
## IRanges object with 2 ranges and 0 metadata columns:
##
                           end
                                    width
              start
##
         <integer> <integer> <integer>
##
     [1]
                  4
                             5
                                        2
                                        2
##
     [2]
                 10
                            11
##
## IRanges object with 4 ranges and 0 metadata columns:
##
              start
                           end
                                    width
##
         <integer> <integer> <integer>
##
     [1]
                  1
                             2
                                        2
     [2]
                             6
                                        2
##
                  5
                                        2
     [3]
                  7
                             8
##
                                        2
     [4]
                 10
##
                            11
```

3 [GCG]

3 [GCG]

3 [GCG]

Now let's get back to the ENCODE project and find DNAStrings where ChIP-seq binding peaks are:

```
# Load human reference genome
library(BSgenome.Hsapiens.UCSC.hg19)
# Get DNAStrings of human genome where ChIP-seq binding peaks are
hepseq = getSeq(Hsapiens, HepG2)
hepseq[1:5]
## DNAStringSet object of length 5:
##
       width seq
         410 GAGACAGGGTTTCACCATGTTGGCCAGGCTGGTT...CCTTCCAGGAAGCAGAAATGTTCAAGGACTCTC
## [1]
## [2]
         861 TGGGAAGGACACAACTGAATGAGGCTGTGCAGAG...AGCAGAACCTCCAACCGTGTGTGTGTGTGTGT
       1156 GACACCTGCCACCCGGACCCCACAGAATGGGCA...GCTTCGTGTCTGCTTTCTTATGTGTTTTTGTTT
## [3]
## [4]
         886 GTGAAGGCCCTGGAGTAGGCGGTGCGTACCCGGT...AGTGTTTTTGGCACCTCCGTGGGCACCTAGGCT
       1242 CATCCTCCACCTTAACACTCAGCACCCTTAGAGA...TTTTGTGTCCTACAAGCAGCCGGCGGCGCCCCC
## [5]
```

Note: A DNA motif is defined as a nucleic acid sequence pattern that has some biological significance such as being DNA binding sites for a regulatory protein, i.e., a transcription factor.

Count occurrences of a motif in the gathered DNAStrings:

```
mot = "TCAAGGTCA"
sum(vcountPattern(mot, hepseq),
     vcountPattern(mot, reverseComplement(hepseq)))
## [1] 55
```

5.4 Data classes

In this section, we will work with different forms of genome data classes.

5.4.1 PLINK

PLINK format includes three files:

- .bed: Contains the genomic SNP data (Homozygous normal 00, Heterozygous 10, Homozygous variant 11, missing 01)
- .bim: Contains SNPs annotations
- .fam: Contains the subject's information

```
library(snpStats)
snps <- read.plink(bed = "Data/obesity.bed",
  bim = "Data/obesity.bim",
  fam = "Data/obesity.fam")
geno <- snps$genotypes
pheno <- snps$fam
annotation <- snps$map</pre>
```

5.4.1.1 Coordinating information from diverse tables We will use a dataset which has three objects, like those of the PLINK format: a SnpMatrix and two dataframes.

```
# Load sample data
library(GSE5859Subset)
data(GSE5859Subset)
```

Upon attachment and loading of data, we have three data elements:

- geneAnnotation
- geneExpression

• sampleInfo

```
# Check elements
geneExpression[1:2, 1:4]
             GSM136508.CEL.gz GSM136530.CEL.gz GSM136517.CEL.gz GSM136576.CEL.gz
##
                                                        6.298943
## 1007_s_at
                     6.543954
                                       6.401470
                                                                          6.837899
## 1053_at
                     7.546708
                                       7.263547
                                                         7.201699
                                                                          7.052761
geneAnnotation[1, ]
##
       PROBEID CHR
                      CHRLOC SYMBOL
## 1 1007_s_at chr6 30852327
                                DDR1
sampleInfo[1, ]
       ethnicity
##
                       date
                                     filename group
## 107
             ASN 2005-06-23 GSM136508.CEL.gz
Now to integrate data:
# Integrate data
cbind(sampleInfo[1:3,], colnames(geneExpression)[1:3], t(geneExpression)[1:3, 1:4])
                                     filename group colnames(geneExpression)[1:3]
##
       ethnicity
                       date
## 107
             ASN 2005-06-23 GSM136508.CEL.gz
                                                                  GSM136508.CEL.gz
                                                  1
## 122
             ASN 2005-06-27 GSM136530.CEL.gz
                                                                  GSM136530.CEL.gz
                                                  1
## 113
             ASN 2005-06-27 GSM136517.CEL.gz
                                                   1
                                                                  GSM136517.CEL.gz
                             117_at
##
       1007_s_at 1053_at
                                      121 at
## 107
       6.543954 7.546708 5.402622 7.892544
## 122
        6.401470 7.263547 5.050546 7.707754
       6.298943 7.201699 5.024917 7.461886
```

5.4.2 ExpressionSet

An ExpressionSet is another form of data container which is designed to combine several different sources of information into a single convenient structure. The data in an ExpressionSet consists of expression data from microarray experiments, 'meta-data' describing samples in the experiment, annotations and meta-data about the features on the chip and information related to the protocol used for processing each sample.

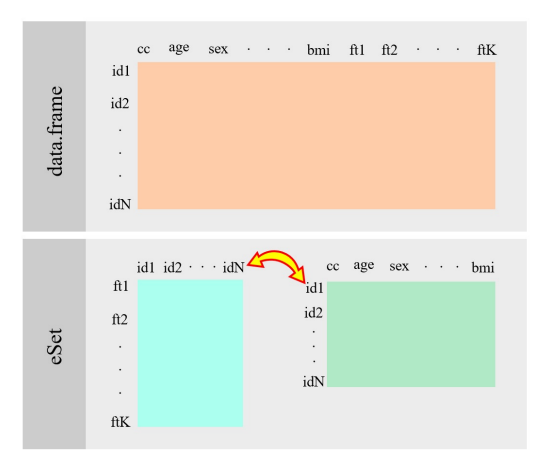


Figure 4: ExpressionSet data

```
# Import packages
library(Biobase)
```

Let's convert our previous PLINK data to an ExpressionSet:

```
# Bind the previous tables in an ExpressionSet
rownames(sampleInfo) = sampleInfo$filename
rownames(geneAnnotation) = geneAnnotation$PROBEID
es5859 = ExpressionSet(assayData = geneExpression)
pData(es5859) = sampleInfo
fData(es5859) = geneAnnotation
```

And we can easily subset data like the following:

```
# Subsetting data from an ExpressionSet
es5859_Y = es5859[which(fData(es5859)$CHR == "chrY"), ]
```

5.4.2.1 Annotation in ExpressionSet annotation argument in an ExpressionSet points to a character describing the platform on which the samples were assayed. This is often the name of a Bioconductor chip annotation package, which facilitated down-stream analysis.

```
# Set annotation for the ExpressionSet
annotation(es5859) = "hgfocus.db"
```

5.4.2.2 ExperimentData in ExpressionSet experimentData argument in an ExpressionSet points to an optional MIAME (Minimum Information About a Microarray Experiment) instance with meta-data (e.g., the lab and resulting publications from the analysis) about the experiment.

```
# Set experimentData for the ExpressionSet
library(annotate)
experimentData(es5859) = pmid2MIAME("17206142")
# Check experimentData
experimentData(es5859)
## Experiment data
     Experimenter name: Spielman RS
##
     Laboratory: NA
##
     Contact information:
##
     Title: Common genetic variants account for differences in gene expression among ethnic groups.
##
     URL:
##
     PMIDs: 17206142
##
##
     Abstract: A 145 word abstract is available. Use 'abstract' method.
# Check abstract
abstract(es5859)
```

[1] "Variation in DNA sequence contributes to individual differences in quantitative traits, but in

5.4.3 SummarizedExperiment

SummarizedExperiment is a comprehensive data structure that can be used to store expression and methylation data from microarrays or read counts from RNA-seq experiments. It can contain slots for one or more omic datasets, feature annotation (e.g. genes, transcripts, SNPs, CpGs), individual phenotypes and experimental details,

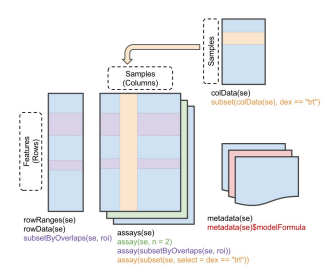


Figure 5: SummarizedExperiment data

Let's load a sample SummarizedExperiment data:

```
# Load sample data (RNA-seq study)
library(airway)
library(SummarizedExperiment)
data(airway)
Check the data:
# Get metadata about the experiment
metadata(airway)
## [[1]]
## Experiment data
##
    Experimenter name: Himes BE
##
    Laboratory: NA
##
     Contact information:
##
     Title: RNA-Seq transcriptome profiling identifies CRISPLD2 as a glucocorticoid responsive gene that
##
     URL: http://www.ncbi.nlm.nih.gov/pubmed/24926665
##
     PMIDs: 24926665
##
##
     Abstract: A 226 word abstract is available. Use 'abstract' method.
# Get the first four feature names
rownames(airway)[1:4]
## [1] "ENSG0000000003" "ENSG0000000005" "ENSG00000000419" "ENSG00000000457"
# Get the first four exon coordinates for a specific gene
rowRanges(airway) $ENSG00000172057[1:4]
##
  GRanges object with 4 ranges and 2 metadata columns:
##
         seqnames
                             ranges strand |
                                               exon_id
                                                              exon_name
##
            <Rle>
                          <IRanges> <Rle> | <integer>
                                                           <character>
##
     [1]
              17 38077294-38078938
                                         - 1
                                                549057 ENSE00001316037
                                         - |
##
     [2]
              17 38077296-38077570
                                                549058 ENSE00002684279
##
     [3]
               17 38077929-38078002
                                         - 1
                                                549059 ENSE00002697088
                                                549060 ENSE00002718599
##
     [4]
               17 38078351-38078552
                                         - 1
##
     seqinfo: 722 sequences (1 circular) from an unspecified genome
##
# Get the sample-level data
colData(airway)
## DataFrame with 8 rows and 9 columns
##
              SampleName
                             cell
                                      dex
                                              albut
                                                           Run avgLength
##
                <factor> <factor> <factor> <factor>
                                                      <factor> <integer>
## SRR1039508 GSM1275862 N61311
                                     untrt
                                              untrt SRR1039508
                                                                      126
## SRR1039509 GSM1275863 N61311
                                              untrt SRR1039509
                                                                      126
                                     trt
## SRR1039512 GSM1275866 N052611
                                     untrt
                                              untrt SRR1039512
                                                                      126
## SRR1039513 GSM1275867 N052611
                                     trt
                                              untrt SRR1039513
                                                                      87
## SRR1039516 GSM1275870 N080611
                                     untrt
                                              untrt SRR1039516
                                                                      120
## SRR1039517 GSM1275871 N080611
                                    trt
                                              untrt SRR1039517
                                                                      126
## SRR1039520 GSM1275874 N061011
                                              untrt SRR1039520
                                                                      101
                                     untrt
## SRR1039521 GSM1275875 N061011
                                              untrt SRR1039521
                                                                      98
                                     trt
##
              Experiment
                                      BioSample
                            Sample
##
                <factor> <factor>
                                       <factor>
## SRR1039508 SRX384345 SRS508568 SAMN02422669
## SRR1039509 SRX384346 SRS508567 SAMN02422675
```

```
## SRR1039512 SRX384349 SRS508571 SAMN02422678
## SRR1039513 SRX384350 SRS508572 SAMN02422670
## SRR1039516 SRX384353 SRS508575 SAMN02422682
## SRR1039517 SRX384354 SRS508576 SAMN02422673
## SRR1039520
               SRX384357 SRS508579 SAMN02422683
## SRR1039521 SRX384358 SRS508580 SAMN02422677
# Check for the existence of overlapping regions
# in the exon coordinates of one specific gene
reduce(rowRanges(airway)$ENSG00000172057)
  GRanges object with 8 ranges and 0 metadata columns:
##
         seqnames
                             ranges strand
##
            <Rle>
                           <IRanges>
                                      <Rle>
##
     [1]
               17 38077294-38078938
     [2]
##
               17 38079365-38079516
##
     [3]
               17 38080283-38080478
##
     [4]
               17 38081008-38081058
##
     [5]
               17 38081422-38081624
##
     [6]
               17 38081876-38083099
               17 38083320-38083482
##
     [7]
##
     [8]
               17 38083737-38083854
##
##
     seqinfo: 722 sequences (1 circular) from an unspecified genome
# Check sample groups
table(airway$dex)
##
##
     trt untrt
##
       4
             4
```

5.5 Expression array archives

Expression array archives contain the ExpressionSet for a lot of various species and experiments. Expression array archive in the US:

- Gene Expression Omnibus (GEO)
- Package: GEOquery

Expression array archive in Europe:

- European Molecular Biology Laboratories (EMBL)
- Package: ArrayExpress

5.5.1 Obtaining the ExpressionSet for a GEO series

```
library(GEOquery)
# glioMA = getGEO("GSE78703")[[1]]
```

5.6 Obtaining the ExpressionSet for an EMBL series

```
library(ArrayExpress)
# glioMA = getAE("E-MTAB-5797")
```

5.7 Storing data

In this section, we will learn how to store data in a database and manipulate that data.

This dataset can quite easily be handled directly in R, but for larger datasets dimensionality can become a problem. The simplest workaround is to store the data in a SQL query and retrieve only the parts of the data that are needed at any given time.

In here, we will use two files:

- a genotypes file: contains all genotype calls for all SNP and all samples.
- a map file: holds mapping information of the SNP, e.g., chromosome, physical location, etc.

We want to create a database. There are two ways:

5.7.1 Using command line

For our database we need to create a schema (a schema is simply a text file that describes the database structure and is used to create the initial empty DB). A simple schema will consist of two tables, one for each of the files and one column for each source of information.

```
cat Data/snpDB.sql
```

We are ready to create the database, named "SNPsmall":

```
sqlite3 Data/SNPsmall < Data/snpDB.sql
```

5.7.2 Using R

This process does not need an schema.

```
# Load package
library(RSQLite)
# Create a new blank database called SNPsmall
con = dbConnect(dbDriver("SQLite"), dbname = "Output/SNPsmall")
# Upload files to the database
dbWriteTable(
  con,
  "snpmap",
  "Data/SNPmap.txt",
  header = TRUE,
  append = T,
  sep = "\t"
dbWriteTable(
  con,
  "SNP",
  "Data/SNPSample1.txt",
  append = T,
  header = TRUE,
  skip = 0,
 sep = "\t"
# Take a look at the tables and fields in the databse
```

```
con = dbConnect(dbDriver("SQLite"), dbname = "Output/SNPsmall")
dbListTables(con)
## [1] "SNP"
                "snpmap"
dbListFields(con, "snpmap")
## [1] "name"
                    "chromosome" "position"
dbListFields(con, "SNP")
                 "animal" "allele1" "allele2" "x"
## [1] "snp"
                                                                     "gcscore"
You can add indexes to the database to make it faster.
dbGetQuery(con, "CREATE INDEX chromosome_idx ON snpmap(chromosome)")
## data frame with 0 columns and 0 rows
dbGetQuery(con, "CREATE INDEX snp idx ON SNP(animal)")
## data frame with 0 columns and 0 rows
dbGetQuery(con, "CREATE INDEX ID_idx ON SNP(snp)")
## data frame with 0 columns and 0 rows
The function dbGetQuery is used to send an SQL query to the DB and return the data in a single step.
# Retrieve the number of records in a table
dbGetQuery(con, "select count (*) from snpmap")
##
     count (*)
         20000
## 1
# Retrieve sample ids
sampleids s1 = as.vector(dbGetQuery(con, "select distinct animal from SNP")) $animal
head(sampleids_s1)
## [1] "\"sample1\"" "\"sample10\"" "\"sample11\"" "\"sample12\"" "\"sample13\""
## [6] "\"sample14\""
# Retrieve all data associated with the first sample
hold_s1 = dbGetQuery(con,
                  paste("select * from SNP where animal='", sampleids_s1[1], "'", sep = ""))
head(hold_s1)
##
                                        animal allele1 allele2
                                                                          y gcscore
                                snp
## 1 "250506CS3900065000002_1238.1" "sample1"
                                                   "A"
                                                           "B" 0.833 0.707 0.8446
## 2 "250506CS3900140500001_312.1" "sample1"
                                                   "B"
                                                           "B" 0.018 0.679 0.9629
## 3 "250506CS3900176800001_906.1" "sample1"
                                                   "B"
                                                           "B" 0.008 1.022 0.9484
## 4 "250506CS3900211600001 1041.1" "sample1"
                                                   "B"
                                                           "B" 0.010 0.769 0.9398
## 5 "250506CS3900218700001_1294.1" "sample1"
                                                   "B"
                                                           "B" 0.000 0.808 0.9272
## 6 "250506CS3900283200001_442.1" "sample1"
                                                   "B"
                                                           "B" 0.019 0.583 0.9552
When we are finished with the database we should close the connection.
dbDisconnect(con)
```

5.8 Reference genomes

Let's check how to load some reference genomes and specially human reference genome:

```
# Import packages
library(BSgenome)
library(Biostrings)
# Check some of the available reference genomes
available.genomes()[1:5]
## [1] "BSgenome.Alyrata.JGI.v1"
## [2] "BSgenome.Amellifera.BeeBase.assembly4"
## [3] "BSgenome.Amellifera.NCBI.AmelHAv3.1"
## [4] "BSgenome.Amellifera.UCSC.apiMel2"
## [5] "BSgenome.Amellifera.UCSC.apiMel2.masked"
# Load Hsapiens. UCSC. hq19 reference seq
library(BSgenome.Hsapiens.UCSC.hg19)
hs = BSgenome.Hsapiens.UCSC.hg19
# Acquire a chromosome's sequence
hs$chr17
## 81195210-letter DNAString object
## seq: AAGCTTCTCACCCTGTTCCTGCATAGATAATTGCAT...GTGGGTGTGGGTGTGTGTGTGGGTGTGGGTGTGGT
```

5.9 Annotations

Let's see how to annotate the data now. We will try with human reference genome annotation.

5.9.1 UCSC annotation

```
# Load Hsapiens. UCSC. hg19 transcripts and genes
library(TxDb.Hsapiens.UCSC.hg19.knownGene)
hs_txdb = TxDb.Hsapiens.UCSC.hg19.knownGene
hs_txdb
## TxDb object:
## # Db type: TxDb
## # Supporting package: GenomicFeatures
## # Data source: UCSC
## # Genome: hg19
## # Organism: Homo sapiens
## # Taxonomy ID: 9606
## # UCSC Table: knownGene
## # Resource URL: http://genome.ucsc.edu/
## # Type of Gene ID: Entrez Gene ID
## # Full dataset: yes
## # miRBase build ID: GRCh37
## # transcript_nrow: 82960
## # exon_nrow: 289969
## # cds_nrow: 237533
## # Db created by: GenomicFeatures package from Bioconductor
## # Creation time: 2015-10-07 18:11:28 +0000 (Wed, 07 Oct 2015)
## # GenomicFeatures version at creation time: 1.21.30
## # RSQLite version at creation time: 1.0.0
## # DBSCHEMAVERSION: 1.1
```

Get the addresses of genes by Entrez gene IDs genes(hs_txdb) ## GRanges object with 23056 ranges and 1 metadata column: ## segnames ranges strand | gene id ## <Rle> <IRanges> <Rle> | <character> ## chr19 58858172-58874214 - | ## 10 chr8 18248755-18258723 10 ## 100 chr20 43248163-43280376 100 25530930-25757445 ## 1000 1000 chr18 ## 10000 chr1 243651535-244006886 10000 ## ## 9991 chr9 114979995-115095944 - | 9991 ## + | 9992 chr21 35736323-35743440 9992 9993 ## chr22 19023795-19109967 9993 ## 9994 chr6 90539619-90584155 9994 ## 9997 chr22 50961997-50964905 9997 _____ ## seqinfo: 93 sequences (1 circular) from hg19 genome # Get the addresses of exons exons(hs_txdb) ## GRanges object with 289969 ranges and 1 metadata column: ## seqnames ranges strand | exon_id ## <Rle> <IRanges> <Rle> | <integer> ## [1] chr1 11874-12227 + | ## + | [2] chr1 12595-12721 ## [3] chr1 12613-12721 + | 3 ## [4] chr1 12646-12697 + | 4 ## [5] 5 chr1 13221-14409 + | ## ## [289965] chrUn_gl000241 35706-35859 289965 ## [289966] chrUn_gl000241 36711-36875 - 1 289966 ## [289967] chrUn_gl000243 11501-11530 + | 289967 ## [289968] chrUn_gl000243 13608-13637 + | 289968 ## [289969] chrUn_gl000247 5787-5816 - | 289969 ## seqinfo: 93 sequences (1 circular) from hg19 genome # Get the addresses of transcripts transcripts(hs txdb) ## GRanges object with 82960 ranges and 2 metadata columns: ## seqnames ranges strand | tx_id tx_name ## <Rle> <IRanges> <Rle> | <integer> <character> ## [1] chr1 11874-14409 + | 1 uc001aaa.3 [2] ## chr1 11874-14409 + | 2 uc010nxq.1 ## [3] 11874-14409 3 uc010nxr.1 chr1 + | ## [4] 69091-70008 4 uc001aal.1 chr1 + | ## [5] chr1 321084-321115 + | 5 uc001aaq.2 ## ## [82956] chrUn_gl000237 1-2686 82956 uc011mgu.1 - 1

20433-36875

11501-11530

##

##

[82957] chrUn gl000241

[82958] chrUn_gl000243

- |

+ |

82957 uc011mgv.2

82958 uc011mgw.1

```
5787-5816
##
     [82960] chrUn_gl000247
                                                         82960 uc022brr.1
##
     seqinfo: 93 sequences (1 circular) from hg19 genome
##
# Filter all exons identified for two different genes (by their Entrez gene IDs)
exons(
 hs_txdb,
  columns = c("EXONID", "TXNAME", "GENEID"),
  filter = list(gene_id = c(100, 101))
## GRanges object with 39 ranges and 3 metadata columns:
##
          segnames
                                 ranges strand |
##
             <Rle>
                              <IRanges> <Rle> | <integer>
##
      [1]
             chr10 135075920-135076737
                                              - |
                                                     144421
##
      [2]
             chr10 135077192-135077269
                                              - 1
                                                     144422
##
      [3]
             chr10 135080856-135080921
                                              - 1
                                                     144423
                                              - |
##
      [4]
             chr10 135081433-135081570
                                                     144424
##
      [5]
             chr10 135081433-135081622
                                              - |
                                                     144425
##
      . . .
##
     [35]
             chr20
                      43254210-43254325
                                                     256371
                                              - |
##
     [36]
             chr20
                      43255097-43255240
                                              - |
                                                     256372
                                              - 1
##
     [37]
             chr20
                      43257688-43257810
                                                     256373
##
     [38]
             chr20
                      43264868-43264929
                                              - 1
                                                     256374
##
     [39]
             chr20
                      43280216-43280376
                                                     256375
##
                                                      GENEID
                                     TXNAME
##
                            <CharacterList> <CharacterList>
##
      [1] uc009ybi.3,uc010qva.2,uc021qbe.1
##
      [2]
                      uc009ybi.3,uc021qbe.1
                                                         101
##
      [3] uc009ybi.3,uc010qva.2,uc021qbe.1
                                                         101
##
      ۲4٦
                                                         101
                                 uc009ybi.3
##
      [5]
                      uc010qva.2,uc021qbe.1
                                                         101
##
      . . .
                                                          . . .
##
     [35]
                                 uc002xmj.3
                                                         100
##
     [36]
                                 uc002xmj.3
                                                         100
##
     [37]
                                 uc002xmj.3
                                                         100
##
     [38]
                                 uc002xmj.3
                                                         100
##
                                 uc002xmj.3
                                                         100
     [39]
##
##
     seqinfo: 93 sequences (1 circular) from hg19 genome
5.9.2 ENSEMBL annotation
# Load Hsapiens.v75 transcripts, genes, and proteins
library(ensembldb)
library(EnsDb.Hsapiens.v75)
hs_edb = EnsDb.Hsapiens.v75
hs edb
## EnsDb for Ensembl:
## |Backend: SQLite
## |Db type: EnsDb
## |Type of Gene ID: Ensembl Gene ID
## |Supporting package: ensembldb
```

##

[82959] chrUn_gl000243

13608-13637

+ |

82959 uc022brg.1

```
## |Db created by: ensembldb package from Bioconductor
## |script version: 0.3.0
## |Creation time: Thu May 18 09:15:45 2017
## |ensembl_version: 75
## |ensembl_host: localhost
## |Organism: homo sapiens
## |taxonomy id: 9606
## |genome build: GRCh37
## | DBSCHEMAVERSION: 2.0
## | No. of genes: 64102.
## | No. of transcripts: 215647.
## | Protein data available.
# List attributes
listTables(hs_edb)
## $gene
## [1] "gene_id"
                          "gene_name"
                                              "gene_biotype"
                                                                  "gene_seq_start"
## [5] "gene_seq_end"
                                              "seq_strand"
                          "seq_name"
                                                                  "seq_coord_system"
## [9] "symbol"
##
## $tx
## [1] "tx id"
                          "tx biotype"
                                              "tx_seq_start"
                                                                  "tx seq end"
## [5] "tx_cds_seq_start" "tx_cds_seq_end"
                                              "gene id"
                                                                  "tx name"
##
## $tx2exon
## [1] "tx_id"
                  "exon_id" "exon_idx"
##
## $exon
## [1] "exon_id"
                         "exon_seq_start" "exon_seq_end"
##
## $chromosome
                     "seq_length" "is_circular"
## [1] "seq_name"
##
## $protein
## [1] "tx_id"
                          "protein_id"
                                              "protein_sequence"
## $uniprot
                               "uniprot_id"
## [1] "protein_id"
                                                      "uniprot_db"
## [4] "uniprot_mapping_type"
## $protein_domain
## [1] "protein_id"
                                "protein_domain_id"
                                                        "protein_domain_source"
## [4] "interpro_accession"
                                "prot_dom_start"
                                                        "prot_dom_end"
##
## $entrezgene
## [1] "gene_id" "entrezid"
##
## $metadata
## [1] "name" "value"
# Return seq_name (i.e., chr name) for transcripts of a specific gene
transcripts(hs_edb,
            filter = GenenameFilter("ZBTB16"),
            columns = "seq_name")
```

```
## GRanges object with 9 ranges and 2 metadata columns:
##
                     segnames
                                           ranges strand |
                                                                      tx id
                                                                <character>
##
                        <Rle>
                                        <IRanges> <Rle> |
##
     ENST00000335953
                           11 113930315-114121398
                                                        + | ENST00000335953
##
     ENST00000541602
                           11 113930447-114060486
                                                        + | ENST00000541602
##
    ENST00000544220
                           11 113930459-113934368
                                                        + | ENST00000544220
     ENST00000535700
                           11 113930979-113934466
                                                        + | ENST00000535700
##
##
                           11 113931229-114121374
                                                        + | ENST00000392996
     ENST00000392996
##
     ENST00000539918
                           11 113935134-114118066
                                                       + | ENST00000539918
##
                                                        + | ENST00000545851
     ENST00000545851
                           11 114051488-114118018
##
     ENST00000535379
                           11 114107929-114121279
                                                        + | ENST00000535379
                                                       + | ENST00000535509
##
     ENST00000535509
                           11 114117512-114121198
##
                       gene_name
##
                     <character>
##
     ENST00000335953
                          ZBTB16
##
     ENST00000541602
                          ZBTB16
##
     ENST00000544220
                          ZBTB16
##
     ENST00000535700
                          ZBTB16
##
     ENST00000392996
                          ZBTB16
##
    ENST00000539918
                          ZBTB16
##
    ENST00000545851
                          ZBTB16
##
    ENST00000535379
                          ZBTB16
##
    ENST00000535509
                          ZBTB16
##
     seqinfo: 1 sequence from GRCh37 genome
##
5.9.3 OrgDb annotation
# Import packages
library(AnnotationDbi)
library(org.Hs.eg.db)
# View the first five columns' names
columns(org.Hs.eg.db)[1:5]
## [1] "ACCNUM"
                      "ALIAS"
                                     "ENSEMBL"
                                                     "ENSEMBLPROT" "ENSEMBLTRANS"
# View the first 5 keys (i.e., values) in a specific column (i.e., keytype)
keys(org.Hs.eg.db, keytype = "ENSEMBL")[1:5]
## [1] "ENSG00000121410" "ENSG00000175899" "ENSG00000291190" "ENSG00000171428"
## [5] "ENSG00000156006"
# Map the ID from a specific key (e.g., ENSG00000175899) from its
# corresponding keytype (e.g., ENSEMBL) to another keytype (e.g., ENTREZID)
mapIds(org.Hs.eg.db,
       keys = "ENSG00000175899",
       column = "ENTREZID",
       keytype = "ENSEMBL")
## ENSG0000175899
# A process similar to the mapIds function from the AnnotationDbi package,
# this time in the biomaRt package
library(biomaRt)
```

```
mart <-
  useMart(biomart = "ensembl",
          dataset = "hsapiens_gene_ensembl",
          host = "https://useast.ensembl.org")
getBM(
  mart = mart,
  attributes = "entrezgene_id",
 filters = "ensembl_gene_id",
  values = "ENSG00000175899"
)
##
     entrezgene id
## 1
# List the first 5 attributes for a specific mart object
listAttributes(mart)[1:5,]
##
                               name
                                                      description
## 1
                    ensembl_gene_id
                                                   Gene stable ID feature_page
## 2
           ensembl_gene_id_version
                                          Gene stable ID version feature_page
             ensembl_transcript_id
## 3
                                            Transcript stable ID feature_page
## 4 ensembl_transcript_id_version Transcript stable ID version feature_page
                                                Protein stable ID feature_page
                ensembl_peptide_id
# List the first 5 filters for a specific mart object
listFilters(mart)[1:5,]
                                   description
##
                name
## 1 chromosome name Chromosome/scaffold name
## 2
                                         Start
               start
## 3
                                           End
                 end
## 4
                                    Band Start
          band_start
## 5
            band_end
                                      Band End
5.9.4 Gene sets and pathways
5.9.4.1 Gene Ontology Gene Ontology (GO) organizes terms relevant to the roles of genes and gene
products in:

    biological processes

  • molecular functions
  • cellular components
# Import packages
library(GO.db)
library(AnnotationDbi)
# View the columns' names
columns(GO.db)
## [1] "DEFINITION" "GOID"
                                  "ONTOLOGY"
                                                "TERM"
# View the first 5 keys (i.e., values) in a specific column (i.e., keytype)
keys(GO.db, keytype = "ONTOLOGY")[1:5]
```

"universal" NA

[1] "BP"

"CC"

"MF"

```
# Import packages
library(KEGGREST)
# Get data for the gene with the Entrez ID 675 (i.e., BRCA2 gene)
brca2K = keggGet("hsa:675")
# List gene attributes
names(brca2K[[1]])
5.9.4.2 KEGG: Kyoto Encyclopedia of Genes and Genomes
## [1] "ENTRY"
                    "SYMBOL"
                                "NAME"
                                            "ORTHOLOGY" "ORGANISM"
                                                                    "PATHWAY"
## [7] "NETWORK"
                    "DISEASE"
                                "BRITE"
                                            "POSITION" "MOTIF"
                                                                    "DBLINKS"
## [13] "STRUCTURE" "AASEQ"
                                "NTSEQ"
# Get its seq data
brca2K[[1]] $NTSEQ
## DNAStringSet object of length 1:
      width seq
## [1] 10257 ATGCCTATTGGATCCAAAGAGAGGCCAACATTTT...CAGGACACAATTACAACTAAAAAATATATCTAA
# Get the list of genes making up a pathway model
brpat = keggGet("path:hsa05212")
# List pathway attributes
names(brpat[[1]])
                      "NAME"
## [1] "ENTRY"
                                    "DESCRIPTION" "CLASS"
                                                                "PATHWAY_MAP"
## [6] "NETWORK"
                     "DISEASE"
                                    "DRUG"
                                                                "GENE"
                                                  "ORGANISM"
## [11] "COMPOUND"
                     "REL_PATHWAY" "KO_PATHWAY"
                                                  "REFERENCE"
# Get the Entrez IDs for the first five genes
brpat[[1]] $GENE[seq(1, 10, 2)]
## [1] "3845" "5290" "5293" "5291" "5295"
# Get an illustration of the pathway
library(png)
library(grid)
brpng = keggGet("hsa05212", "image")
grid.raster(brpng)
```

