R and Stats - PDCB topic Infraestructure HTS

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Overview

ShortRead

SAM format

Rsamtools

Exercises

Infra-I/O

- ➤ This is the beginning of the infrastructure and input/output section of the course.
- Our goals: to learn how to read into R HTS data files, manipulate them and manipulate the information they contain.
- Today is mostly about reading in files and filtering reads we don't want.

Today's packages

- ➤ You should have them installed already, but if you don't then please do so:
 - > source("http://bioconductor.org/biocLite.R")
 - > biocLite(c("ShortRead", "Rsamtools"))

- Which HTS data formats are you familiar with or have heard about?
- **>** /

```
NAGAGGCCAGGCCATCTACCACCTTTTGTTGGAAATTTTGCTCTTTCAAC
+HWUSI-EAS636_0001:1:1:0:114#0/1
DOVYUQUYWSTWYYYYYYYYYYYYYYYYYYTPLSYYWWWRJRYYYYYWVTV
CGGAAGAGCGGTTCAGCAGGAATGCCGAGATCGGAAGAGCGGTTCAGCAT
+HWUSI-EAS636_0001:1:1:1:552#0/1
aaaaaaK_Y^_b_ZaaaPIXa_VZWRNHZ^LHUHRRPUPVJIRQWQYXB
AGCGCATCTTGCGCTATGTGCAGCAGAGCGTGAGCCTTAACCTGATGCGC
```

▶ B

HWUSI-EAS636	1	4	45	849	1900	0	1
GACTTAGGTCACTAAATACTTTAACCAATATAGGCA							
abbabbbaabbab	babababa	aaaaaaa	abaa]`aa	ECK12.fa	sta		113
F 36	146						
HWUSI-EAS636	1	4	101	43	603	0	1
CTTAGGTCACTA	AATACTTTA	ACCAATA	TAGGCATA				
Z_b]J_^_Zaa]]^bbabba^_`aaabaa_^TT`_W				ECK12.fasta			115
F	36 146						
HWUSI-EAS636	1	4	109	1181	363	0	1
CTTAGGTCACTAAATACTTTAACCAATATAGGCATA							
aa`baa`aaaaba	`baa`aaaabaaabaaaaa``aaaaa``_aaa			ECK12.fasta			115
F 36 1	146						

► C

```
##gff-version 2
##date 2010-09-13
Ecoli
      rtracklayer
                                         16
                                                 16
                                                         3
                        sequence
1.4
Ecoli
       rtracklayer
                                         38
                                                 38
                                                         24
                        sequence
2.4
Ecoli
       rtracklayer
                                                 50
                                                         6
                        sequence
                                         50
3.4
```

▶ D

```
HWUSI-EAS636:8:120:1791:562#0/1 -
                                        gi|49175990|ref|NC_000913.2|
1753519 GTCGGACTGTAGAACTCT
                                ::::::868;>>>:>B
0:A>T.15:T>C.17:T>G
HWUSI-EAS636:8:120:1791:393#0/1 -
                                        gi|49175990|ref|NC_000913.2|
2399840 TCGGACTGTAGAACTCTG
                                9>7@1B:<8@AA8A8AAB
2:T>C.3:G>T.15:T>G
HWUSI-EAS636:8:120:1791:1802#0/1
                                                gi|49175990|ref|
NC 000913.21
                1132065 GTTCAGAGTTCTACAGTC
                                                B::>9>4::::=67:?
        4:G>A,6:T>A,16:C>T
0
```

```
HWUSI-EAS636:8:120:1791:1350#0/1 - gi|49175990|ref|
NC_000913.2| 1753520 TCGGACTGTAGAACTCTG <:A8866?;8@:6><>?B
0 0:A>G,1:A>T,16:T>C
```

► E

► F

```
track name="R Track" type=bedGraph
chr_gi|49175990|ref|NC_000913.2| 18 45 2
chr_gi|49175990|ref|NC_000913.2| 81 95 0
chr_gi|49175990|ref|NC_000913.2| 95 104 2
```

▶ G

► H

```
HWI-EAS88_4_1_6_505_934 ChrA 1 + 0 0 15 15 15 1 12 0 1 35

aaagttagagaagtttgacttttgtaggcaccatc ------)))))))###

HWI-EAS88_4_1_7_163_963 ChrA 1 + 0 0 22 22 22 0 0 1 0 35

aaagttagagaagtttgacttctgtaggcaccatc ------)))))))###
```

HWI-EAS88 3 2 1 451 945 CCAGAGCCCCCCGCTCACTCCTGAACCAGTCTCTC
YQMIMIMMLMMIGIGMFICMFFFIMMHIIHAAGAH NM N
HWI-EAS88 3 2 1 409 991 AGCCTCCCTCTTTCTGAATATACGGCAGAGCTGTT
ZXZUYXZQYYXUZXYZYYZZXXZZIMFHXQSUPPO NM Y
HWI-EAS88 3 2 1 451 939 ACCAAAAACACCACATACACGAGCAACACACGTAC
LGDHLILLLLLLIGFLAALDIFDILLHFTAECAE NM N

▶ .]

I'm a HTS data file from your imagination :)

Answers

- A fastq
- B sorted
- C gff version 2
- D bowtie single end
- E bowtie paired end
- ▶ F bed
- G SAM
- ► H mag
- ▶ I export
- ▶ J:0

The ShortRead package

- It's one of the first BioC packages related to HTS data
- Has been the basic input/output package for HTS data
- It can read solexa, fastq, bowtie, and maq files. It can also read in other types of alignments.
- With it we can explore the quality of our reads/alignments, create a report and filter out reads.
- Current model: read all the reads into RAM and then manipulate them.

- Lets get into ShortRead!
- SR was originally designed to read in files from the Solexa set of directories.
- ▶ Lets look at the example data. Where is it for you?
 - > library(ShortRead)
 - > exptPath <- system.file("extdata",
 - + package = "ShortRead")
- ► For SR to recognize the path, we need to use SolexaPath:
 - > sp <- SolexaPath(exptPath)
 - > sp

```
class: SolexaPath
experimentPath: C:/PROGRA~1/R/R-212~1.0/library/ShortRe
dataPath: Data
scanPath: NA
```

imageAnalysisPath: C1-36Firecrest

baseCallPath: Bustard
analysisPath: GERALD

- Next, we can use some functions to find the path for several important files:
 - > imageAnalysisPath(sp)
 - [1] "C:/PROGRA~1/R/R-212~1.0/library/ShortRead/extdata/
 - > analysisPath(sp)

[1] "C:/PROGRA~1/R/R-212~1.0/library/ShortRead/extdata/

However, that isn't that interesting for us. We want to read in data! For example, an export file.

```
> aln <- readAligned(sp, "s_2_export.txt")</pre>
> aln
class: AlignedRead
length: 1000 reads; width: 35 cycles
chromosome: NM NM ... chr5.fa 29:255:255
position: NA NA ... 71805980 NA
strand: NA NA ... + NA
alignQuality: NumericQuality
alignData varLabels: run lane ... filtering contig
> class(aln)
```

```
[1] "AlignedRead"
attr(,"package")
[1] "ShortRead"
```

- AlignedRead objects are the main type of objects in SR. Multiple functions to access parts of it exist.
- ► For example, how would you extract the positions for all reads?

AlignedRead

As the names imply, we can extract the positions with:

```
> summary(position(aln))
```

```
Min. 1st Qu. Median Mean
11940 34710000 73390000 74160000
3rd Qu. Max. NA's
108500000 195500000 594
```

- Why do we have NAs?
- Some other useful accesors are:
 - > table(strand(aln))

```
+ - >
```

AlignedRead

```
> unique(width(aln))
[1] 35
> alignQuality(aln)
class: NumericQuality
quality: 0 0 ... 55 0 (1000 total)
> summary(quality(alignQuality(aln)))
  Min. 1st Qu. Median Mean 3rd Qu.
       0.00 0.00 17.04
  0.00
                                 37.00
  Max.
  72.00
> length(aln)
[1] 1000
```

AlignedRead

```
> head(table(chromosome(aln)))
0:0:187 0:0:19 0:0:21 0:0:25 0:0:255
 0:0:85
> head(id(aln))
  A BStringSet instance of length 6
    width seq
[1]
[2]
[3]
[4]
```

${\sf AlignedRead}$

[5] 0 [6] 0

Quick exercise

- Lets assume that the 5' end of our reads corresponds to transcription start sites.
- Get the TSSs positions.
- ▶ What is the TSSs for read number 10 in our aln object?
- Remember:
 - > summary(position(aln))[7]

NA's

594

Solution

Lets take advantage of how R works by using vectors.

```
> idx <- which(is.na(position(aln)) ==
+    FALSE)
> neg <- which(strand(aln)[idx] ==
+    "-")
> tss <- position(aln)[idx]
> tss[neg] <- tss[neg] + width(aln)[idx][neg] -
+    1

For the second part:
> tenth <- head(position(aln), 10)
> tenth
```

Solution

```
[1]
             NΑ
                       NA
                                NΑ
                                          NΑ
   [5]
             NA
                       NA 69345321 54982866
   [9]
             NA 80537786
  > tenth <- length(which(is.na(tenth) ==
        FALSE))
  > tenth
  [1] 3
  > tss[tenth]
  [1] 80537820
Is the answer correct?
  > aln[10]
```

Solution

```
class: AlignedRead
length: 1 reads; width: 35 cycles
chromosome: chr12.fa
position: 80537786
strand: -
alignQuality: NumericQuality
alignData varLabels: run lane ... filtering contig
> tss[tenth] == 80537786 + 35 - 1
[1] TRUE
```

Reading fastq files

- Before we continue with alignment files, SR is also capable of reading fastq files.
- Lets read the example file:

```
> args(readFastq)
function (dirPath, pattern = character(0), ...)
NULL
> sread <- readFastq(analysisPath(sp),
+    pattern = "sequence.txt")
> class(sread)
[1] "ShortReadQ"
attr(,"package")
[1] "ShortRead"
```

What did analysisPath do for us?

► In addition to AlignedRead, ShortReadQ objects completes the faimily of main objects in SR.

> sread

class: ShortReadQ

length: 256 reads; width: 36 cycles

Similar to AlignedRead objects, we can access parts of the information:

> head(id(sread))

```
A BStringSet instance of length 6
width seq
[1] 24 HWI-EAS88_1_1_1_1001_499
[2] 23 HWI-EAS88_1_1_1_898_392
[3] 23 HWI-EAS88_1_1_1_922_465
[4] 23 HWI-EAS88_1_1_1_895_493
[5] 23 HWI-EAS88_1_1_1_953_493
[6] 23 HWI-EAS88_1_1_1_868_763
> head(quality(sread))
```

```
class: SFastqQuality
quality:
 A BStringSet instance of length 6
  width seq
[1]
    36 1111111111111111111111111111VCHVMPLAS
[2]
    [3]
    Γ41
    36 ]]]]]]]]]]]]]]...]]]RJRZTQLOA
[5]
    [6]
    > length(sread)
[1] 256
> table(width(sread))
```

```
36
256
> head(sread(sread))
  A DNAStringSet instance of length 6
    width seq
[1]
       36 GGACTTTGTAGGAT...TTCCTTCTCCTGT
[2]
       36 GATTTCTTACCTAT...AACAGCATCGGAC
[3]
       36 GCGGTGGTCTATAG...TATCAATTTGGGT
Γ41
       36 GTTACCATGATGTT...TTTGGAGGTAAAA
[5]
       36 GTATGTTTCTCCTG...TTCTTGAAGGCTT
[6]
       36 GTTCTCTAAAAACC...CCCCTTCGGGGCG
> narrow(sread, start = 1, end = 10)
```

class: ShortReadQ

length: 256 reads; width: 10 cycles

Exercise

- ▶ Which are different dinucleotides in our reads? Only base 1 and 2 of our reads.
- What are the frequencies of the different dinucleotides?
- Coercion functions such as as.character can be useful:) You might need to check the help of:
 - > `?`(BStringSet)

Solution I

Lets use the sread, narrow, as.character and table functions:

```
> first2 <- sread(narrow(sread, start = 1,</pre>
     width = 2)
> head(first2)
 A DNAStringSet instance of length 6
   width seq
[1]
       2 GG
[2] 2 GA
[3] 2 GC
[4] 2 GT
[5] 2 GT
[6]
       2 GT
```

Solution I

```
> first2 <- as.character(first2)
> table(first2)
first2
GA GC GG GT
61 42 51 102
```

Solution II

- While the above solution was fine, it did involve changing between types of objects.
- Lets use the dinucleotideFrequency function:

+ start = 1, width = 2)))

> dinuc[1,]

AA AC AG AT CA CC CG CT GA GC GG GT TA

TC TG TT

0 0 0

> dinuc <- colSums(dinuc)</pre>

> dinuc[dinuc > 0]

Solution II

GA GC GG GT 61 42 51 102

Alphabet Frequency

- Now, lets try get the alphabet frequency per every sequencing cycle.
- ▶ This information is VERY useful to pick up errors!
- ► Any ideas?

```
Apropos is quite useful!
  > apropos("alphabet")
   [1] ".__T_alphabet:Biostrings"
   [2] ".__T_alphabet:Biostrings"
   [3] ".__T_alphabetByCycle:ShortRead"
   [4] ".__T_alphabetFrequency:Biostrings"
   [5] ".__T_alphabetFrequency:Biostrings"
   [6] ".__T_alphabetScore:ShortRead"
   [7] "AA ALPHABET"
   [8] "alphabet"
   [9] "alphabetByCycle"
  [10] "alphabetFrequency"
  [11] "alphabetScore"
```

```
[12] "DNA_ALPHABET"
[13] "RNA_ALPHABET"
```

- Lets use the function alphabetByCycle
 - > alph <- alphabetByCycle(sread(sread))</pre>
 - > dim(alph)

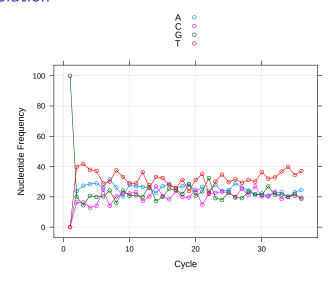
[1] 17 36

- Why did I use the sread accessor? Why does alph have 17 rows and 36 columns?
- Exercise: lets plot the alphabet by cycle relative frequency (only letters > 0) using lattice. Use only 1 panel and draw 1 line per alphabet letter present.
- Do you observe something unexpected?

```
vacio6
  > library(lattice)
  > alph2 <- as.data.frame(t(alph[rowSums(alph) >
       0.1)
  > head(alph2)
  1 0 0 256
 2 61 42 51 102
 3 70 42 37 107
 4 73 33 53
             97
 5 74 36 51 95
 6 67 63 52 74
```

```
> alph2 <- alph2/rowSums(alph2) *</pre>
      100
> head(alph2)
  0.00000 0.00000 100.00000
                               0.00000
2 23.82812 16.40625 19.92188 39.84375
3 27.34375 16.40625 14.45312 41.79688
4 28.51562 12.89062 20.70312 37.89062
5 28.90625 14.06250 19.92188 37.10938
6 26.17188 24.60938 20.31250 28.90625
```

```
> print(xyplot(A + C + G + T ~ 1:nrow(alph2),
+ data = alph2, type = c("o",
+ "g"), auto.key = TRUE,
+ xlab = "Cycle", ylab = "Nucleotide Frequency"))
```



qa report

- qa is a function that summarizes fastq files, export, etc and creates a series of summary plots.
- When working, it creates an html file.

```
> args(qa)
function (dirPath, ...)
NULL
> qa <- qa(sp)
> dir <- tempfile()
> report(qa, dest = dir)
> dir(paste(dir, "image", sep = "/"))
```

Yet, we can still access some of the data through R:

qa report

Which file did qa use by default?

▶ To end our cruise through SR, lets filter reads!

```
> apropos("filter")
 [1] ".__C_FilterRules"
 [2] ".__C__SRFilter"
 [3] ".__T_Filter:base"
 [4] ".__T_filterBam:Rsamtools"
 [5] ".__T_filterRules: IRanges"
 [6] ".__T__filterRules<-: IRanges"
 [7] ".__T_srFilter:ShortRead"
 [8] "alignDataFilter"
 [9] "alignQualityFilter"
[10] "chromosomeFilter"
[11] "dustyFilter"
```

```
[12] "filter"
[13] "Filter"
[14] "Filter"
[15] "filterBam"
[16] "filterRules"
[17] "FilterRules"
[18] "filterRules<-"
[19] "Filters"
[20] "idFilter"
[21] "nFilter"
[22] "occurrenceFilter"
[23] "polynFilter"
[24] "positionFilter"
[25] "srdistanceFilter"
```

```
[26] "srFilter"
[27] "strandFilter"
[28] "uniqueFilter"
```

► The main class is srFilter, though many types are already coded.

```
> nfilt <- nFilter()
> cfilt <- chromosomeFilter("chr5.fa")
> sfilt <- strandFilter("+")</pre>
```

- With the above filters we can now read in the reads from chromsome 5 in the plus strand.
- ► We can specify the filters when reading the file or to subset an AlignedRead object:

```
> chr5 <- readAligned(sp, "s_2_export.txt",
+ filter = cfilt)
> filt <- compose(cfilt, sfilt)
> chr5plus <- readAligned(sp, "s_2_export.txt",
+ filter = filt)
> length(chr5plus) == length(aln[filt(aln)])
[1] TRUE
```

Recap

- Universal format
- BAM is binary SAM
- ► SAM and BAM files can be ordered by the position of the reads (left to right on the genome)
- The model behind: read in pieces of files at a time.
- ▶ Definition: http://samtools.sourceforge.net/SAM1.pdf
- Related tools: http://samtools.sourceforge.net/
- SAM format is doomed, any clues why?

Overview

- Similar to ShortRead
- Can read in pieces of files at a time
- ▶ In the near future: will be able to handle gaps!

- ▶ After loading Rsamtools, next we need to construct a special object with ScanBamParam. Mainly this object specifies which parts of the chromsome / organism we want ot read in and the columns of information we want.
- ► To do so we need to use some of the IRanges functionality. Don't worry, we'll cover it next week :)
- ► Once we have the paramters, we can now read the BAM file using scanBam.
- scanBam can also read in files that are hosted on the web! :)

```
> library(Rsamtools)
> which <- RangesList(seq1 = IRanges(1000,</pre>
      2000), seq2 = IRanges(c(100),
      1000), c(1000, 2000)))
> which
SimpleRangesList of length 2
$seq1
IRanges of length 1
    start end width
[1] 1000 2000 1001
$seq2
IRanges of length 2
    start end width
```

```
[1] 100 1000 901
[2] 1000 2000 1001
> what <- c("rname", "strand", "pos",</pre>
+ "qwidth", "seq")
> what
[1] "rname" "strand" "pos" "qwidth"
[5] "seq"
> param <- ScanBamParam(which = which,</pre>
+ what = what)
> param
```

```
class: ScanBamParam
bamFlag: keep '0' bits: 2047; keep '1' bits: 2047
bamSimpleCigar: FALSE
bamReverseComplement: FALSE
bamTag:
bamWhich: 2 elements
bamWhat: rname, strand, pos,
 qwidth, seq
> bamFile <- system.file("extdata",</pre>
      "ex1.bam", package = "Rsamtools")
> bam <- scanBam(bamFile, param = param)</pre>
```

Exploring the output of scanBam

The output is a list with a second list inside. At the lowest level we can find an object for each of the what columns we specified

```
> class(bam)
[1] "list"
> names(bam)
[1] "seq1:1000-2000" "seq2:100-1000"
[3] "seq2:1000-2000"
> lapply(bam, class)
```

Exploring the output of scanBam

```
$`seq1:1000-2000`
[1] "list"
$`seq2:100-1000`
[1] "list"
$`seq2:1000-2000`
[1] "list"
> names(bam[[1]])
[1] "rname" "strand" "pos"
                                 "qwidth"
[5] "seq"
> sapply(bam[[1]], class)
```

Exploring the output of scanBam

rname strand

"factor" "factor"

pos qwidth

"integer" "integer"

seq

"DNAStringSet"

You might feel comfortable with such kind of data, though you might also like it in to view it in a tabular format such as a DataFrame:

```
> lst <- lapply(names(bam[[1]]),
+ function(elt) {
+ do.call(c, unname(lapply(bam,
+ "[[", elt)))
+ })
> names(lst) <- names(bam[[1]])
> head(do.call("DataFrame", lst))
```

	Da	ataFrame	with	6	rows	and	5	columns	
		rname		strand				pos	
		<integer< td=""><td>r> <i1< td=""><td colspan="3"><pre><integer></integer></pre></td><td colspan="3"><integer></integer></td></i1<></td></integer<>	r> <i1< td=""><td colspan="3"><pre><integer></integer></pre></td><td colspan="3"><integer></integer></td></i1<>	<pre><integer></integer></pre>			<integer></integer>		
	1		1		1			970	
	2		1		1			971	
	3		1		1			972	
	4		1		1			973	
	5		1		1			974	
	6		1		2			975	
		qwidt	th						
<integer></integer>			<u>r</u> >						
	1	3	35						
	2	3	35						
	3	3	35						

```
4 35
5 35
6 35
```

seq <DNAStringSet>

- 1 TATTAGGAAATGCTTTACTGTCATAACTATGAAGA
- 2 ATTAGGAAATGCTTTACTGTCATAACTATGAAGAG
- 3 TTAGGAAATGCTTTACTGTCATAACTATGAAGAGA
- 4 TAGGAAATGCTTTACTGTCATAACTATGAAGAGAC
- 5 AGGAAATGCTTTACTGTCATAACTATGAAGAGACT
- 6 GGAAATGCTTTACTGTCATAACTATGAAGAGACTA
- Note that it is a DataFrame and not a data.frame!

▶ We won't use this kind of object much since we can also transform it into a GRanges object (next session!).

From the web!

Just as an example, lets read in data from the web.

param = param)

- We'll get data only from chromosome 6 bases 100k to 110k from the 1000 genomes project.
- If we wanted to donwload all the data, well, that's around 10GB! The output with scanBam is only around 2Mb in memory.

```
> which <- RangesList(`6` = IRanges(100000L,
+ 110000L))
> param <- ScanBamParam(which = which)
> na19240url <- "ftp://ftp-trace.ncbi.nih.gov/1000genom
> na19240bam <- scanBam(na19240url,</pre>
```

From the web!

```
> print(object.size(na19240bam),
+ units = "Mb")
1.9 Mb
```

Rsamtools has much more to offer

- We only took a quick glimpse at Rsamtools. It still has other useful functions if you are working with BAM files such as RamViews:
 - use BamViews to reference a set of disk-based BAM files to be processed (e.g., queried using scanBam) as a single experiment.
- ▶ There is also a function to read in gapped alignments:
 - > aln1_file <- system.file("extdata",</pre>
 - + "ex1.bam", package = "Rsamtools")
 - > aln1 <- readBamGappedAlignments(aln1_file)</pre>
 - > head(aln1)

Rsamtools has much more to offer

```
GappedAlignments of length 6
   rname strand cigar qwidth start end
[1]
                36M
    seq1
            +
                       36
                                36
[2]
                35M
                       35
                             3 37
    seq1
            +
[3] seq1
            + 35M
                       35
                             5 39
[4] seq1
                36M
                       36
                               41
            +
[5]
                35M
                       35
                               43
    seq1
            +
[6]
                35M
                       35
                            13
                               47
    seq1
            +
   width ngap
Г17
      36
           0
[2]
      35
           0
[3]
      35
           0
[4]
      36
           0
```

Rsamtools has much more to offer

```
[5] 35 0
[6] 35 0
```

▶ And more to come as it's been actively developed :)

Some practice

- From the aln object, extract the dinucleotide frequency for the last 2 cycles.
 - Given the GC percentage of all cycles, did you expect the results you observe?
 - 2. Which is the read with NN at the end?
 - 3. Is there a significative difference vs the dinucleotide frequency of cycles 15 and 16?
- ► Load the na19240url object (note that fpt doesn't wort at IBt).
 - 1. Are all reads of the same length? If not, what is the distribution? Make a cumulative plot.
 - 2. Convert the PhredQuality instance to a quality matrix and make a plot of the median quality per cycle. Is there any trend in the quality?

Some practice

3. Make a third plot for the alphabet by cycle relative frequency (in percent). Do you observe anything unexpected?

Session Information

```
> sessionInfo()
R version 2.12.0 (2010-10-15)
Platform: i386-pc-mingw32/i386 (32-bit)
locale:
[1] LC_COLLATE=English_United States.1252
[2] LC_CTYPE=English_United States.1252
[3] LC_MONETARY=English_United States.1252
[4] LC NUMERIC=C
[5] LC_TIME=English_United States.1252
attached base packages:
[1] stats
             graphics grDevices
[4] utils
             datasets methods
[7] base
other attached packages:
[1] ShortRead_1.8.2
[2] Rsamtools 1.2.1
```

Session Information

[3] lattice_0.19-13

```
[4] Biostrings_2.18.0
[5] GenomicRanges_1.2.0
[6] IRanges_1.8.0
loaded via a namespace (and not attached):
[1] Biobase_2.10.0 grid_2.12.0
[3] hwriter_1.2
```