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>	> library(Rsamtools)						

Introduction

The Rsamtools package provides an interface to BAM files. BAM files are produced by samtools and other software, and represent a flexible format for storing 'short' reads aligned to reference genomes. BAM files typically contain sequence and base qualities, and alignment coordinates and quality measures. BAM files are appealing for several reasons. The format is flexible enough to represent reads generated and aligned using diverse technologies. The files are binary so that file access is relatively efficient. BAM files can be indexed, allowing ready access to localized chromosomal regions. BAM files can be accessed remotely, provided the remote hosting site supports such access and a local index is available. This means that specific regions of remote files can be accessed without retrieving the entire (large!) file. A full description is available in the BAM format specification (http://samtools.sourceforge.net/SAM1.pdf)

The main purpose of the Rsamtools package is to import BAM files into R. Rsamtools also provides some facility for file access such as record counting, index file creation, and filtering to create new files containing subsets of the original. An important use case for Rsamtools is as a starting point for creating R objects suitable for a diversity of work flows, e.g., AlignedRead objects in the ShortRead package (for quality assessment and read manipulation), or GAlignments objects in GenomicRanges package (for RNA-seq and other applications). Those desiring more functionality are encouraged to explore samtools and related software efforts.

Input

The essential capability provided by Rsamtools is BAM input. This is accomplished with the scanBam function. scanBam takes as input the name of the BAM file to be parsed. In addition, the param argument determines which genomic coordinates of the BAM file, and what components of each record, will be input. Rparam is an instance of the ScanBamParam

class. To create a param object, call ScanBamParam. Here we create a param object to extract reads aligned to three distinct ranges (one on seq1, two on seq2). From each of read in those ranges, we specify that we would like to extract the reference name (rname, e.g., seq1), strand, alignment position, query (i.e., read) width, and query sequence:

```
> which <- RangesList(seq1=IRanges(1000, 2000),</pre>
                       seq2=IRanges(c(100, 1000), c(1000, 2000)))
> what <- c("rname", "strand", "pos", "qwidth", "seq")</pre>
> param <- ScanBamParam(which=which, what=what)
```

Additional information can be found on the help page for ScanBamParam. Reading the relevant records from the BAM file is accomplished with

```
> bamFile <-
      system.file("extdata", "ex1.bam", package="Rsamtools")
> bam <- scanBam(bamFile, param=param)</pre>
```

Like scan, scanBam returns a list of values. Each element of the list corresponds to a range specified by the which argument to ScanBamParam.

```
> class(bam)
[1] "list"
> names(bam)
[1] "seq1:1000-2000" "seq2:100-1000" "seq2:1000-2000"
Each element is itself a list, containing the elements specified by the what and tag arguments to ScanBamParam.
> class(bam[[1]])
[1] "list"
> names(bam[[1]])
[1] "rname" "strand" "pos"
                                   "qwidth" "seq"
The elements are either basic R or IRanges data types
> sapply(bam[[1]], class)
          rname
                                                           qwidth
                         strand
                                             pos
                       "factor"
      "factor"
                                       "integer"
                                                        "integer"
            seq
"DNAStringSet"
A paradigm for collapsing the list-of-lists into a single list is
```

```
> .unlist <- function (x)</pre>
+ {
      ## do.call(c, ...) coerces factor to integer, which is undesired
      x1 <- x[[1L]]
      if (is.factor(x1)) {
          structure(unlist(x), class = "factor", levels = levels(x1))
          do.call(c, x)
+ }
> bam <- unname(bam) # names not useful in unlisted result
> elts <- setNames(bamWhat(param), bamWhat(param))</pre>
> lst <- lapply(elts, function(elt) .unlist(lapply(bam, "[[", elt)))
```

This might be further transformed, e.g., to a DataFrame, with

> head(do.call("DataFrame", lst))

DataFrame with 6 rows and 5 columns

	rname	strand	pos	qwidth
	< factor >	<factor></factor>	<pre><integer></integer></pre>	<integer></integer>
1	seq1	+	970	35
2	seq1	+	971	35
3	seq1	+	972	35
4	seq1	+	973	35
5	seq1	+	974	35
6	seq1	-	975	35
				sea

<DNAStringSet>

- 1 TATTAGGAAATGCTTTACTGTCATAACTATGAAGA
- 2 ATTAGGAAATGCTTTACTGTCATAACTATGAAGAG
- 3 TTAGGAAATGCTTTACTGTCATAACTATGAAGAGA
- 4 TAGGAAATGCTTTACTGTCATAACTATGAAGAGAC
- 5 AGGAAATGCTTTACTGTCATAACTATGAAGAGACT
- 6 GGAAATGCTTTACTGTCATAACTATGAAGAGACTA

Often, an alternative is to use a *ScanBamParam* object with desired fields specified in what as an argument to GenomicRanges::readGalignments; the specified fields are added as columns to the returned *Galignments*.

The BAM file in the previous example includes an index, represented by a separate file with extension .bai:

> list.files(dirname(bamFile), pattern="ex1.bam(.bai)?")

```
[1] "ex1.bam" "ex1.bam.bai"
```

Indexing provides two significant benefits. First, an index allows a BAM file to be efficiently accessed by range. A corollary is that providing a which argument to ScanBamParam requires an index. Second, coordinates for extracting information from a BAM file can be derived from the index, so a portion of a remote BAM file can be retrieved with local access only to the index. For instance, provided an index file exists on the local computer, it is possible to retrieve a small portion of a BAM file residing on the 1000 genomes HTTP server. The url ftp://ftp-trace.ncbi.nih.gov/1000genomes/ftp/pilot_data/NA19240/alignment/NA19240.chrom6.SLX.maq.SRP000032.2009_07.bam points to the BAM file corresponding to individual NA19240 chromosome 6 Solexa (Illumina) sequences aligned using MAQ. The remote file is very large (about 10 GB), but the corresponding index file is small (about 500 KB). With na19240url set to the above address, the following retrieves just those reads in the specified range

```
> which <- RangesList("6"=IRanges(100000L, 110000L))
> param <- ScanBamParam(which=which, what=scanBamWhat())
> na19240bam <- scanBam(na19240url, param=param)</pre>
```

Invoking scanBam without an index file, as above, first retrieves the index file from the remote location, and then queries the remote file using the index; for repeated queries, it is more efficient to retrieve the index file first (e.g., with download.file) and then use the local index as an argument to scanBam. Many BAM files were created in a way that causes scanBam to report that the "EOF marker is absent"; this message can safely be ignored.

BAM files may be read by functions in packages other than Rsamtools. In ShortRead, readAligned invoked with type="bam" will read BAM files in to an AlignedRead object. This function takes a param argument, just as scanBam, so the user can control which portions of the file are input. Similar input facilities are available through the readGAlignments function in GenomicRanges.

Additional ways of interacting with BAM files include scanBamHeader (to extract header information) and countBam (to count records matching param). filterBam filters reads from the source file according to the criteria of the Scan-BamParam parameter, writing reads passing the filter to a new file. The function sortBam sorts a previously unsorted BAM, while The function indexBam creates an index file from a sorted BAM file.

readPileup reads a pileup file created by samtools, importing SNP, indel, or all variants into a GRanges object.

2.1 Large bam files

BAM files can be large, containing more information on more genomic regions than are of immediate interest or than can fit in memory. The first strategy for dealing with this is to select, using the what and which arguments to ScanBamParam, just those portions of the BAM file that are essential to the current analysis, e.g., specifying what=c('rname', 'qname', 'pos') when wishing to calculate coverage of ungapped reads.

When selective input of BAM files is still too memory-intensive, the file can be processed in chunks, with each chunk distilled to the derived information of interest. Chromosomes will often be the natural chunk to process. For instance, here we write a summary function that takes a single sequence name (chromosome) as input, reads in specific information from the BAM file, and calculates coverage over that sequence.

This might be used as follows; it is an ideal candidate for evaluation in parallel, e.g., using the *parallel* package and srapply function in *ShortRead*.

```
> seqnames <- paste("seq", 1:2, sep="")</pre>
> cvg <- lapply(seqnames, summaryFunction, bamFile)
> names(cvg) <- seqnames
> cvg
$seq1
integer-Rle of length 1569 with 1054 runs
 Lengths: 2 2
                1 3 4 2 3 4 ...
                                        1
                                           1
 Values: 1 2 3 4 5 7 8 9 ... 9 7
$seq2
integer-Rle of length 1567 with 1092 runs
 Lengths: 1 3 1 1 1 3 1 4 ... 1
                                           2
                                       1
                                             1
 Values: 3 4 5 8 12 14 15 16 ... 10 8
```

The result of the function (a coverage vector, in this case) will often be much smaller than the input.

3 Views

The functions described in the previous section import data in to R. However, sequence data can be very large, and it does not always make sense to read the data in immediately. Instead, it can be useful to marshal *references* to the data into a container and then act on components of the container. The *BamViews* class provides a mechanism for creating 'views' into a set of BAM files. The view itself is light-weight, containing references to the relevant BAM files and metadata about the view (e.g., the phenotypic samples corresponding to each BAM file).

One way of understanding a *BamViews* instance is as a rectangular data structure. The columns represent BAM files (e.g., distinct samples). The rows represent ranges (i.e., genomic coordinates). For instance, a ChIP-seq experiment might identify a number of peaks of high read counts.

3.1 Assembling a BamViews instance

To illustrate, suppose we have an interest in caffeine metabolism in humans. The 'rows' contain coordinates of genomic regions associated with genes in a KEGG caffeine metabolism pathway. The 'columns' represent individuals in the 1000 genomes project.

To create the 'rows', we identify possible genes that KEGG associates with caffeine metabolism:

```
> library(KEGG.db)
> kid <- revmap(KEGGPATHID2NAME)[["Caffeine metabolism"]]
> egid <- KEGGPATHID2EXTID[[sprintf("hsa%s", kid)]]</pre>
```

Then we use the appropriate *TranscriptDb* package to translate Entrez identifiers to obtain ranges of interest (one could also use *biomaRt* to retrieve coordinates for non-model organisms, perhaps making a TranscriptDb object as outlined in the *GenomicFeatures* vignette). We'll find that the names used for chromosomes in the alignments differ from those used at Ensembl, so seqlevels<- is used to map between naming schemes and to drop unused levels.

Note that bamRanges 'knows' the genome for which the ranges are defined

```
> unique(genome(bamRanges))
```

```
[1] "hg18"
```

The details of creating the 'columns' of BAM files. Here we retrieve a vector of BAM file URLs (slxMaq09) from the package.

We now assemble the BamViews instance from these objects; we also provide information to annotated the BAM files (with the bamSamples function argument, which is a DataFrame instance with each row corresponding to a BAM file) and the instance as a whole (with bamExperiment, a simple named list containing information structured as the user sees fit).

3.2 Using BamViews instances

The BamViews object can be queried for its component parts, e.g.,

```
> bamExperiment(bv)
```

```
$description
```

[1] "Caffeine metabolism views on 1000 genomes samples"

\$created

```
[1] "Sat Nov 23 00:25:06 2013"
```

More usefully, Methods in *Rsamtools* are designed to work with *BamViews* objects, retrieving data from all files in the view. These operations can take substantial time and require reliable network access.

To illustrate, the following code (not evaluated when this vignette was created) downloads the index files associated with the by object

and then queries the 1000 genomes project for reads overlapping our transcripts; by loading the *parallel* package, we tell *Rsamtools* to use as many cores as are available on our machine (the *parallel* package does not provide parallel functionality on Windows computers)

```
> library(parallel)
> options(srapply_fapply="parallel", mc.cores=detectCores())
> olaps <- readGAlignmentsFromBam(bv)</pre>
```

The resulting object is about 11 MB in size. To avoid having to download this data each time the vignette is run, we instead load it from disk

```
> load(system.file("extdata", "olaps.Rda", package="Rsamtools"))
> olaps
List of length 24
names(24): NA06986.SLX.maq.SRP000031.2009_08.bam ...
```

> head(olaps[[1]])

GAlignments with 6 alignments and 0 metadata columns:

```
seqnames strand
                         cigar
                                   qwidth
      <Rle> <Rle> <character> <integer> <integer>
[1]
                                       51 31410650
          2
                  +
                            51M
          2
[2]
                  +
                            51M
                                       51 31410658
          2
[3]
                            51M
                                       51 31410663
[4]
           2
                            51M
                                       51 31410666
[5]
           2
                            51M
                                       51
                                           31410676
[6]
          2
                            51M
                                       51 31410676
                  +
                  width
          end
                             ngap
   <integer> <integer> <integer>
[1] 31410700
                     51
[2] 31410708
                     51
                                0
[3] 31410713
                     51
                                0
[4] 31410716
                                0
                     51
    31410726
                     51
                                0
[6]
    31410726
                     51
                                0
seqlengths:
                             3 ... NT_113898 NC_007605
                   2
247249719 242951149 199501827 ...
                                     1305230
                                                 171823
```

There are 34616 reads in NA06986.SLX.maq.SRP000031.2009_08.bam overlapping at least one of our transcripts. It is easy to explore this object, for instance discovering the range of read widths in each individual.

```
> head(t(sapply(olaps, function(elt) range(qwidth(elt)))))
```

```
[,1] [,2]
NA06986.SLX.maq.SRP000031.2009_08.bam
                                         51
                                              51
NA06994.SLX.maq.SRP000031.2009_08.bam
                                         36
                                              51
NA07051.SLX.maq.SRP000031.2009_08.bam
                                              51
                                              76
NA07346.SLX.maq.SRP000031.2009_08.bam
                                         48
NA07347.SLX.maq.SRP000031.2009_08.bam
                                         51
                                              51
NA10847.SLX.maq.SRP000031.2009_08.bam
                                         36
                                              51
```

Suppose we were particularly interested in the first transcript, which has a transcript id uc002ayr.1. Here we extract reads overlapping this transcript from each of our samples. As a consequence of the protocol used, reads aligning to either strand could be derived from this transcript. For this reason, we set the strand of our range of interest to *. We use the endoapply function, which is like lapply but returns an object of the same class (in this case, SimpleList) as its first argument.

```
> rng <- bamRanges(bv)[1]</pre>
> strand(rng) <- "*"
> olap1 <- endoapply(olaps, subsetByOverlaps, rng)</pre>
> olap1 <- lapply(olap1, "seqlevels<-", value=as.character(seqnames(rng)))</pre>
> head(olap1[[24]])
GAlignments with 6 alignments and 0 metadata columns:
      segnames strand
                                      qwidth
                            cigar
                                                  start
         <Rle> <Rle> <character> <integer> <integer>
  [1]
            15
                               36M
                                          36 72828212
  [2]
            15
                               36M
                                          36 72828226
  [3]
            15
                               36M
                                          36 72828237
  [4]
            15
                               36M
                                          36 72828272
  [5]
            15
                               36M
                                          36 72828291
  [6]
            15
                               36M
                                          36 72828297
            end
                    width
      <integer> <integer> <integer>
  [1] 72828247
                        36
                                   0
  [2] 72828261
                        36
                                   0
  [3] 72828272
                       36
                                   0
  [4] 72828307
                       36
                                   0
  [5] 72828326
                        36
                                   0
  [6] 72828332
                        36
  seqlengths:
          15
   100338915
```

To carry the example a little further, we calculate coverage of each sample:

Since the example includes a single region of uniform width across all samples, we can easily create a coverage matrix with rows representing nucleotide and columns sample and, e.g., document variability between samples and nucleotides

```
> m <- matrix(unlist(lapply(cvg, lapply, as.vector)),</pre>
             ncol=length(cvg))
> summary(rowSums(m))
  Min. 1st Qu. Median
                          Mean 3rd Qu.
                                           Max.
  38.00 69.00 78.00
                         77.86
                                 86.00 122.00
> summary(colSums(m))
  Min. 1st Qu. Median
                           Mean 3rd Qu.
                                          Max.
  12070
        15930
                 22560
                          25170
                                 33080
                                          51200
```

4 Directions

This vignette has summarized facilities in the *Rsamtools* package. Important additional packages include *GenomicRanges* (for representing and manipulating gapped alignments), *ShortRead* for I/O and quality assessment of ungapped short read alignments, *Biostrings* and *BSgenome* for DNA sequence and whole-genome manipulation, *IRanges* for range-based manipulation, and *rtracklayer* for I/O related to the UCSC genome browser. Users might also find the interface to the integrative genome browser (IGV) in *SRAdb* useful for visualizing BAM files.

```
> packageDescription("Rsamtools")
Package: Rsamtools
Type: Package
Title: Binary alignment (BAM), variant call (BCF), or
      tabix file import
Version: 1.14.2
Author: Martin Morgan, Herv\'e Pag\`es, Valerie
      Obenchain
Maintainer: Bioconductor Package Maintainer
      <maintainer@bioconductor.org>
Description: This package provides an interface to
      the 'samtools', 'bcftools', and 'tabix'
      utilities (see 'LICENCE') for manipulating SAM
      (Sequence Alignment / Map), binary variant call
      (BCF) and compressed indexed tab-delimited
      (tabix) files.
URL:
      http://bioconductor.org/packages/release/bioc/html/Rsamtools.html
License: Artistic-2.0 | file LICENSE
LazyLoad: yes
Depends: methods, IRanges (>= 1.19.11), GenomicRanges
      (>= 1.13.35), XVector, Biostrings (>= 2.29.7)
Imports: utils, BiocGenerics (>= 0.1.3), zlibbioc,
      bitops
Suggests: ShortRead (>= 1.19.10), GenomicFeatures,
      TxDb.Dmelanogaster.UCSC.dm3.ensGene, KEGG.db,
      TxDb.Hsapiens.UCSC.hg18.knownGene,
      RNAseqData.HNRNPC.bam.chr14,
      BSgenome. Hsapiens. UCSC. hg19, pasillaBamSubset,
      RUnit, BiocStyle
LinkingTo: IRanges, XVector, Biostrings
```

```
biocViews: DataImport, Sequencing,
      HighThroughputSequencing
Built: R 3.0.2; i386-w64-mingw32; 2013-11-23 08:23:40
      UTC; windows
-- File: D:/biocbld/bbs-2.13-bioc/tmpdir/RtmpcfLkCV/Rinst1ddc71711b00/Rsamtools/Meta/package.rds
> sessionInfo()
R version 3.0.2 Patched (2013-10-28 r64119)
Platform: i386-w64-mingw32/i386 (32-bit)
locale:
[1] LC_COLLATE=C
[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] parallel stats
                        graphics grDevices utils
[6] datasets methods
other attached packages:
 [1] TxDb.Hsapiens.UCSC.hg18.knownGene_2.10.1
 [2] GenomicFeatures_1.14.2
 [3] KEGG.db_2.10.1
 [4] RSQLite_0.11.4
 [5] DBI_0.2-7
 [6] AnnotationDbi_1.24.0
 [7] Biobase_2.22.0
 [8] Rsamtools_1.14.2
 [9] Biostrings_2.30.1
[10] GenomicRanges_1.14.3
[11] XVector_0.2.0
[12] IRanges_1.20.6
[13] BiocGenerics_0.8.0
loaded via a namespace (and not attached):
 [1] BSgenome_1.30.0 BiocStyle_1.0.0
 [3] RCurl_1.95-4.1
                        XML_3.98-1.1
 [5] biomaRt_2.18.0 bitops_1.0-6
 [7] rtracklayer_1.22.0 stats4_3.0.2
 [9] tools_3.0.2
                        zlibbioc_1.8.0
```

A Assembling a BamViews instance

A.1 Genomic ranges of interest

A.2 BAM files

Note: The following operations were performed at the time the vignette was written; location of on-line resources, in particular the organization of the 1000 genomes BAM files, may have changed.

We are interested in collecting the URLs of a number of BAM files from the 1000 genomes project. Our first goal is to identify files that might make for an interesting comparison. First, let's visit the 1000 genomes FTP site and discover available files. We'll use the *RCurl* package to retrieve the names of all files in an appropriate directory

```
> library(RCurl)
> ftpBase <-
+    "ftp://ftp-trace.ncbi.nih.gov/1000genomes/ftp/pilot_data/data/"
> indivDirs <-
+    strsplit(getURL(ftpBase, ftplistonly=TRUE), "\n")[[1]]
> alnDirs <-
+    paste(ftpBase, indivDirs, "/alignment/", sep="")
> urls0 <-
+    strsplit(getURL(alnDirs, dirlistonly=TRUE), "\n")</pre>
```

From these, we exclude directories without any files in them, select only the BAM index (extension .bai) files, and choose those files that exactly six '.' characters in their name.

```
> urls <- urls[sapply(urls0, length) != 0]
> fls0 <- unlist(unname(urls0))
> fls1 <- fls0[grepl("bai$", fls0)]
> fls <- fls1[sapply(strsplit(fls1, "\\."), length)==7]</pre>
```

After a little exploration, we focus on those files obtained form Solexa sequencing, aligned using MAQ, and archived in August, 2009; we remove the .bai extension, so that the URL refers to the underlying file rather than index. There are 24 files.

```
> urls1 <-
+ Filter(function(x) length(x) != 0,
+ lapply(urls,
+ function(x) x[grep1("SLX.maq.*2009_08.*bai$", x)]))
> slxMaq09.bai <-
+ mapply(paste, names(urls1), urls1, sep="", USE.NAMES=FALSE)
> slxMaq09 <- sub(".bai$", "", slxMaq09.bai) #$</pre>
```

As a final step to prepare for using a *BamViews* file, we create local copies of the *index* files. The index files are relatively small (about 190 Mb total).

```
> bamIndexDir <- tempfile()
> dir.create(bamIndexDir)
> idxFiles <- mapply(download.file, slxMaq09.bai,
+ file.path(bamIndexDir, basename(slxMaq09.bai)) ,
+ MoreArgs=list(method="curl"))</pre>
```