# An Introduction to ShortRead

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```
> library("ShortRead")
```

The ShortRead package provides functionality for working with FASTQ files from high throughput sequence analysis. The package also contains functions for legacy (single-end, ungapped) aligned reads; for working with BAM files, please see the Rsamtools, GenomicRanges, GenomicAlignments and related packages.

## 1 Sample data

Sample FASTQ data are derived from ArrayExpress record E-MTAB-1147. Paired-end FASTQ files were retrieved and then sampled to 20,000 records with

```
> sampler <- FastqSampler('E-MTAB-1147/fastq/ERR127302_1.fastq.gz', 20000)
> set.seed(123); ERR127302_1 <- yield(sampler)
> sampler <- FastqSampler('E-MTAB-1147/fastq/ERR127302_2.fastq.gz', 20000)
> set.seed(123); ERR127302_2 <- yield(sampler)</pre>
```

## 2 Functionality

Functionality is summarized in Table 1.

Input FASTQ files are large so processing involves iteration in 'chunks' using FastqStreamer

```
> strm <- FastqStreamer("a.fastq.gz")
> repeat {
+    fq <- yield(strm)
+    if (length(fq) == 0)
+        break
+    ## process chunk
+ }</pre>
```

or drawing a random sample from the file

```
> sampler <- FastqSampler("a.fastq.gz")
> fq <- yield(sampler)</pre>
```

The default size for both streams and samples is 1M records; this volume of data fits easily into memory. Use <a href="countFastq">countFastq</a> to get a quick and memory-efficient count of the number of records and nucleotides in a file

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Input	
FastqStreamer	Iterate through FASTQ files in chunks
FastqSampler	Draw random samples from FASTQ files
readFastq	Read an entire FASTQ file into memory
writeFastq	Write FASTQ objects to a connection (file)
countFastq	Quickly count FASTQ records in files
Sequence and quality su	mmary
alphabetFrequency	Nucleotide or quality score use per read
alphabetByCycle	Nucleotide or quality score use by cycle
alphabetScore	Whole-read quality summary
encoding	Character / 'phred' score mapping
Quality assessment	
qa	Visit FASTQ files to collect QA statistics
report	Generate a quality assessment report
Filtering and trimming	
srFilter	Pre-defined and bespoke filters
trimTails, etc.	Trim low-quality nucleotides
narrow	Remove leading / trailing nucleotides
tables	Summarize read occurrence
srduplicated, etc.	Identify duplicate reads
filterFastq	Filter reads from one file to another

Table 1: Key functions for working with FASTQ files

Small FASTQ files can be read in to memory in their entirety using readFastq; we do this
for our sample data

```
> fq <- readFastq(fl)
```

The result of data input is an instance of class ShortReadQ (Table 2). This class contains

```
DNAStringSet(Biostrings)Short read sequencesFastqQuality, etc.Quality encodingsShortReadQReads, quality scores, and ids
```

Table 2: Primary data types in the ShortRead package

reads, their quality scores, and optionally the id of the read.

```
> fq
class: ShortReadQ
length: 20000 reads; width: 72 cycles
> fq[1:5]
```

```
class: ShortReadQ
length: 5 reads; width: 72 cycles
> head(sread(fq), 3)
DNAStringSet object of length 3:
   width seq
[1]
      72 GTCTGCTGTATCTGTGTCGGCTGTCTCGCGGGAC...GTCAATGAAGGCCTGGAATGTCACTACCCCCAG
      72 CTAGGGCAATCTTTGCAGCAATGAATGCCAATGG...CAGTGGCTTTTGAGGCCAGAGCAGACCTTCGGG
[2]
[3]
      72 TGGGCTGTTCCTCACTGTGGCCTGACTAAAA...TGGCATTAAGAAAGAGTCACGTTTCCCAAGTCT
> head(quality(fq), 3)
class: FastqQuality
quality:
BStringSet object of length 3:
   width seq
      72 HHHHHHHHHHHHHHHHHHHHHBBBB?B:BBGG<D...ABFEFBDBD@DDECEE3>:?;@@@>?=BAB?##
[1]
[2]
      72 IIIIHIIIGIIIIIIHIIIIEGBGHIIIIHGII...IIIHIIIHIIIIIGIIIEGIIGBGE@DDGGGIG
      [3]
```

The reads are represented as *DNAStringSet* instances, and can be manipulated with the rich tools defined in the *Biostrings* package. The quality scores are represented by a class that represents the quality encoding inferred from the file; the encoding in use can be discovered with

The primary source of documentation for these classes is ?ShortReadQ and ?QualityScore.

## 3 Common work flows

## 3.1 Quality assessment

FASTQ files are often used for basic quality assessment, often to augment the purely technical QA that might be provided by the sequencing center with QA relevant to overall experimental design. A QA report is generated by creating a vector of paths to FASTQ files

```
> fls <- dir("/path/to", "*fastq$", full=TRUE)
```

collecting statistics over the files

```
> qaSummary <- qa(fls, type="fastq")</pre>
```

and creating and viewing a report

```
> browseURL(report(qaSummary))
```

By default, the report is based on a sample of 1M reads.

These QA facilities are easily augmented by writing custom functions for reads sampled from files, or by explorting the elements of the object returned from qa(), e.g., for an analysis of ArrayExpress experiment E-MTAB-1147:

```
> qaSummary
class: FastqQA(10)
QA elements (access with qa[["elt"]]):
  readCounts: data.frame(16 3)
  baseCalls: data.frame(16 5)
  readQualityScore: data.frame(8192 4)
  baseQuality: data.frame(1504 3)
  alignQuality: data.frame(16 3)
  frequentSequences: data.frame(800 4)
  sequenceDistribution: data.frame(1953 4)
  perCycle: list(2)
    baseCall: data.frame(5681 4)
    quality: data.frame(44246 5)
  perTile: list(2)
    readCounts: data.frame(0 4)
    medianReadQualityScore: data.frame(0 4)
  adapterContamination: data.frame(16 1)
```

For instance, the count of reads in each lane is summarized in the readCounts element, and can be displayed as

```
> head(qaSummary[["readCounts"]])
                         read filter aligned
ERR127302_1.fastq.gz 29741852
                                  NA
                                           NA
ERR127302_2.fastq.gz 29741852
                                  NA
                                           NA
ERR127303_1.fastq.gz 32665567
                                  NA
                                           NA
ERR127303_2.fastq.gz 32665567
                                           NA
                                  NA
ERR127304_1.fastq.gz 31876181
                                  NA
                                           NA
ERR127304_2.fastq.gz 31876181
                                  NA
                                           NA
> head(qaSummary[["baseCalls"]])
                                     C
                                              G
ERR127302_1.fastq.gz 16439860 19641395 19547421 16335620 35704
ERR127302_2.fastq.gz 16238041 20020655 19608896 16060661 71747
ERR127303_1.fastq.qz 16826258 19204659 19448727 16507994 12362
ERR127303_2.fastq.gz 16426991 19822132 19374419 16324978 51480
ERR127304_1.fastq.qz 16279217 19740457 19879137 16089405 11784
ERR127304_2.fastq.gz 15984998 20297064 19812474 15853510 51954
```

The readCounts element contains a data frame with 1 row and 3 columns (these dimensions are indicated in the parenthetical annotation of readCounts in the output of qaSummary). The rows represent different lanes. The columns indicated the number of reads, the number of reads surviving the Solexa filtering criteria, and the number of reads aligned to the reference genome for the lane. The baseCalls element summarizes base calls in the unfiltered reads.

The functions that produce the report tables and graphics are internal to the package. They can be accessed through calling ShortRead:::functionName where functionName is one of the functions listed below, organized by report section.

Run Summary: .ppnCount, .df2a, .laneLbl, .plotReadQuality Read Distribution: .plotReadOccurrences, .freqSequences Cycle Specific: .plotCycleBaseCall, .plotCycleQuality

 $\label{thm:polynomial} Tile\ Performance: .atQuantile, .colorkeyNames, .plotTileLocalCoords, .tileGeometry, .plotTileCounts, .plotTileQualityScore$ 

Alignment : .plotAlignQuality

 $Multiple\ Alignment:\ .plotMultipleAlignmentCount$ 

Depth of Coverage : .plotDepthOfCoverage

Adapter Contamination: .ppnCount

### 3.2 Filtering and trimming

It is straight-forward to create filters to eliminate reads or to trim reads based on diverse characteristics. The basic structure is to open a FASTQ file, iterate through chunks of the file performing filtering or trimming steps, and appending the filtered data to a new file.

```
> myFilterAndTrim <-
+
      function(fl, destination=sprintf("%s_subset", fl))
+ {
      ## open input stream
      stream <- open(FastqStreamer(fl))</pre>
      on.exit(close(stream))
      repeat {
          ## input chunk
          fq <- yield(stream)</pre>
          if (length(fq) == 0)
               break
          ## trim and filter, e.g., reads cannot contain 'N'...
          fq <- fq[nFilter()(fq)] # see ?srFilter for pre-defined filters</pre>
          ## trim as soon as 2 of 5 nucleotides has quality encoding less
          ## than "4" (phred score 20)
          fq <- trimTailw(fq, 2, "4", 2)
          ## drop reads that are less than 36nt
          fq \leftarrow fq[width(fq) >= 36]
          ## append to destination
          writeFastq(fq, destination, "a")
```

```
+ }
+ }
```

This is memory efficient and flexible. Care must be taken to coordinate pairs of FASTQ files representing paired-end reads, to preserve order.

# 4 Using ShortRead for data exploration

### 4.1 Data I/O

ShortRead provides a variety of methods to read data into R, in addition to readAligned.

### 4.1.1 readXStringColumns

readXStringColumns reads a column of DNA or other sequence-like data. For instance, the Solexa files s\_N\_export.txt contain lines with the following information:

```
> ## location of file
> exptPath <- system.file("extdata", package="ShortRead")</pre>
> sp <- SolexaPath(exptPath)</pre>
> pattern <- "s_2_export.txt"
> fl <- file.path(analysisPath(sp), pattern)</pre>
> strsplit(readLines(fl, n=1), "\t")
[[1]]
 [1] "HWI-EAS88"
                                               "3"
 [3] "2"
                                               "1"
 [5] "451"
                                               "945"
 [7] ""
 [9] "CCAGAGCCCCCGCTCACTCCTGAACCAGTCTCTC"
                                               "YQMIMIMMLMMIGIGMFICMFFFIMMHIIHAAGAH"
[11] "NM"
[13] ""
[15] ""
[17] ""
                                               .. ..
                                               ....
[19] ""
[21] ""
                                               "N"
> length(readLines(fl))
[1] 1000
```

Column 9 is the read, and column 10 the ASCII-encoded Solexa Fastq quality score; there are 1000 lines (i.e., 1000 reads) in this sample file.

Suppose the task is to read column 9 as a *DNAStringSet* and column 10 as a *BStringSet*. *DNAStringSet* is a class that contains IUPAC-encoded DNA strings (IUPAC code allows for nucleotide ambiguity); *BStringSet* is a class that contains any character with ASCII code 0 through 255. Both of these classes are defined in the *Biostrings* package. readXString Columns allows us to read in columns of text as these classes.

Important arguments for readXStringColumns are the dirPath in which to look for files, the pattern of files to parse, and the colClasses of the columns to be parsed. The dirPath and pattern arguments are like list.files. colClasses is like the corresponding argument to

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**read.table**: it is a *list* specifying the class of each column to be read, or NULL if the column is to be ignored. In our case there are 21 columns, and we would like to read in columns 9 and 10. Hence

```
> colClasses <- rep(list(NULL), 21)
> colClasses[9:10] <- c("DNAString", "BString")
> names(colClasses)[9:10] <- c("read", "quality")</pre>
```

We use the class of the type of sequence (e.g., *DNAString* or *BString*), rather than the class of the set that we will create (e.g., *DNAStringSet* or *BStringSet*). Applying names to colClasses is not required, but makes subsequent manipulation easier. We are now ready to read our file

```
> cols <- readXStringColumns(analysisPath(sp), pattern, colClasses)</pre>
> cols
$read
DNAStringSet object of length 1000:
       width seq
   [1]
          35 CCAGAGCCCCCGCTCACTCCTGAACCAGTCTCTC
   [2]
          35 AGCCTCCCTCTTTCTGAATATACGGCAGAGCTGTT
          35 ACCAAAAACACCACATACACGAGCAACACACGTAC
   [3]
   [4]
          35 AATCGGAAGAGCTCGTATGCCGGCTTCTGCTTGGA
   [5]
          35 AAAGATAAACTCTAGGCCACCTCCTCCTTCTTA
         35 GTGGCAGCGGTGAGGCGGCGGGGGGGGGTTGTTTG
 [996]
          35 GTCGGAGGTCAGCAAGCTGTAGTCGGTGTAAAGCT
 [997]
          35 GTCATAAATTGGACAGTGTGGCTCCAGTATTCTCA
 [998]
 [999]
          35 ATCTACATTAAGGTCAATTACAATGATAAATAAAA
[1000]
          35 TTCTCAGCCATTCAGTATTCCTCAGGTGAAAATTC
$quality
BStringSet object of length 1000:
       width seq
   [1]
          35 YQMIMIMMLMMIGIGMFICMFFFIMMHIIHAAGAH
   [2]
          35 ZXZUYXZQYYXUZXYZYYZZXXZZIMFHXQSUPPO
          35 LGDHLILLLLLIGFLLALDIFDILLHFIAECAE
   [3]
   [4]
          35 JJYYIYVSYYYYYYYSDYYWVUYYNNVSVQQELQ
   [5]
          35 LLLILIIIDLLHLLLLLLLLLLLLLLLHLLLEL
          35 ZZZZZZZYZZYUYZYUYZKYUDUZIYYODJGUGAA
 [996]
 [997]
          35 ZZZZZZZZZZZZZZZZZZZZYZZYXXZYSSXXUUHHQ
          35 ZZZZZZZZZZZZZZZZYZZZYZZZYZZXZUUUUS
 [998]
          35 ZZZZZZZZZZXXZYZYZYZZYZZXXZSYXUUNUN
 [999]
          35 ZZZZZZZZZZZZZZZZZZZZZZZZYYSYSZXUUUUU
[1000]
```

The file has been parsed, and appropriate data objects were created.

A feature of readXStringColumns and other input functions in the ShortRead package is that all files matching pattern in the specified dirPath will be read into a single object. This provides a convenient way to, for instance, parse all tiles in a Solexa lane into a single DNAStringSet object.

There are several advantages to reading columns as *XStringSet* objects. These are more compact than the corresponding character representation:

```
> object.size(cols$read)
51032 bytes
> object.size(as.character(cols$read))
102128 bytes
```

They are also created much more quickly. And the *DNAStringSet* and related classes are used extensively in *ShortRead*, *Biostrings*, *BSgenome* and other packages relevant to short read technology.

### 4.2 Sorting

Short reads can be sorted using **srsort**, or the permutation required to bring the short read into lexicographic order can be determined using **srorder**. These functions are different from **sort** and **order** because the result is independent of the locale, and they operate quickly on *DNAStringSet* and *BStringSet* objects.

The function srduplicated identifies duplicate reads. This function returns a logical vector, similar to duplicated. The negation of the result from srduplicated is useful to create a collection of unique reads. An experimental scenario where this might be useful is when the sample preparation involved PCR. In this case, replicate reads may be due to artifacts of sample preparation, rather than differential representation of sequence in the sample prior to PCR.

## 4.3 Summarizing read occurrence

The tables function summarizes read occurrences, for instance,

```
> tbls <- tables(fq)
> names(tbls)
[1] "top"
                "distribution"
> tbls$top[1:5]
CTATTCTCTACAAACCACAAAGACATTGGAACACTATACCTATTATTCGGCGCATGAGCTGGAGTCCTAGGC
CGATAACGTTGTAGATGTCGTTACCTAGAAGGTTGCCTGGCTCGCCCAGCTCGGCTCGAATAAGGAGGC
CTAGCATTTACCATCTCACTTCTAGGAATACTAGTATATCGCTCACACCTCATATCCTCCCTACTATGCCTA
CACGAGCATATTTCACCTCCGCTACCATAATCATCGCTATCCCCACCGGCGTCAAAGTATTTAGCTGACTCG
> head(tbls$distribution)
 nOccurrences nReads
1
           1 19291
2
           2
               247
```

3	3	34
4	4	18
5	5	3
6	6	2

The top component returned by tables is a list tallying the most commonly occurring sequences in the short reads. Knowledgeable readers will recognize the top-occurring read as a close match to one of the manufacturer adapters.

The distribution component returned by tables is a data frame that summarizes how many reads (e.g., 19291) are represented exactly 1 times.

### 4.4 Finding near matches to short sequences

Facilities exist for finding reads that are near matches to specific sequences, e.g., manufacturer adapter or primer sequences. <a href="srdistance">srdistance</a> reports the edit distance between each read and a reference sequence. <a href="srdistance">srdistance</a> is implemented to work efficiently for reference sequences whose length is of the same order as the reads themselves (10's to 100's of bases). To find reads close to the most common read in the example above, one might say

```
> dist <- srdistance(sread(fq), names(tbls$top)[1])[[1]]
> table(dist)[1:10]

dist
    0     4     6     10     14     18     20     21     31     32
    7     1     3     1     4     1     3     11
```

'Near' matches can be filtered, e.g.,

```
> fqSubset <- fq[dist>4]
```

A different strategy can be used to tally or eliminate reads that consist predominantly of a single nucleotide. alphabetFrequency calculates the frequency of each nucleotide (in DNA strings) or letter (for other string sets) in each read. Thus one could identify and eliminate reads with more than 30 adenine nucleotides with

```
> countA <- alphabetFrequency(sread(fq))[,"A"]
> fqNoPolyA <- fq[countA < 30]</pre>
```

alphabetFrequency, which simply counts nucleotides, is much faster than srdistance, which performs full pairwise alignment of each read to the subject.

Users wanting to use R for whole-genome alignments or more flexible pairwise alignment are encouraged to investigate the *Biostrings* package, especially the *PDict* class and matchPDict and pairwiseAlignment functions.

# 5 Legacy support for early file formats

The *ShortRead* package contains functions and classes to support early file formats and ungapped alignments. Help pages are flagged as 'legacy'; versions of *ShortRead* prior to 1.21 (*Bioconductor* version 2.13) contain a vignette illustrating common work flows with these file formats.

### 6 sessionInfo

- > toLatex(sessionInfo())
  - R version 4.3.0 RC (2023-04-18 r84287), x86\_64-pc-linux-gnu
  - Locale: LC\_CTYPE=en\_US.UTF-8, LC\_NUMERIC=C, LC\_TIME=en\_GB, LC\_COLLATE=C, LC\_MONETARY=en\_US.UTF-8, LC\_MESSAGES=en\_US.UTF-8, LC\_PAPER=en\_US.UTF-8, LC\_NAME=C, LC\_ADDRESS=C, LC\_TELEPHONE=C, LC\_MEASUREMENT=en\_US.UTF-8, LC\_IDENTIFICATION=C
  - Time zone: America/New\_York
  - TZcode source: system (glibc)
  - Running under: Ubuntu 22.04.2 LTS
  - Matrix products: default
  - BLAS: /home/biocbuild/bbs-3.18-bioc/R/lib/libRblas.so
  - LAPACK: /usr/lib/x86\_64-linux-gnu/lapack/liblapack.so.3.10.0
  - Base packages: base, datasets, grDevices, graphics, methods, stats, stats4, utils
  - Other packages: Biobase 2.61.0, BiocGenerics 0.47.0, BiocParallel 1.35.0, Biostrings 2.69.0, GenomeInfoDb 1.37.0, GenomicAlignments 1.37.0, GenomicRanges 1.53.0, IRanges 2.35.0, MatrixGenerics 1.13.0, Rsamtools 2.17.0, S4Vectors 0.39.0, ShortRead 1.59.0, SummarizedExperiment 1.31.0, XVector 0.41.0, matrixStats 0.63.0
  - Loaded via a namespace (and not attached): BiocManager 1.30.20, BiocStyle 2.29.0, DelayedArray 0.27.0, GenomeInfoDbData 1.2.10, Matrix 1.5-4, RColorBrewer 1.1-3, RCurl 1.98-1.12, Rcpp 1.0.10, bitops 1.0-7, cli 3.6.1, codetools 0.2-19, compiler 4.3.0, crayon 1.5.2, deldir 1.0-6, digest 0.6.31, evaluate 0.20, fastmap 1.1.1, grid 4.3.0, htmltools 0.5.5, hwriter 1.3.2.1, interp 1.1-4, jpeg 0.1-10, knitr 1.42, lattice 0.21-8, latticeExtra 0.6-30, parallel 4.3.0, png 0.1-8, rlang 1.1.0, rmarkdown 2.21, tools 4.3.0, xfun 0.39, yaml 2.3.7, zlibbioc 1.47.0