An Introduction to FastqCleaner

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Abstract

FastqCleaner is an interactive application for both quality control and pre-processing of FASTQ files. The interface, based on a Bioconductor back-end, provides diagnostic information for the input and output data and a series of filtering and trimming operations in an interactive framework.

Contents

1 Launching the application ........................................ 2
2 Description of the application .................................... 2
  2.1 First panel .................................................... 2
  2.2 Second panel ................................................ 5
  2.3 Third panel ................................................... 6
3 A worked example: FASTQ processing in a nutshell ........ 8
4 Advanced use of the package ..................................... 12
  4.1 Main functions ............................................... 12
  4.2 Auxiliary functions .......................................... 25
5 Contact information .............................................. 26
1 Launching the application

The interactive application can be launched in R with the following command:

```r
library('FastqCleaner')
launch_fqc()
```

As an alternative method, an RStudio addin (RStudio version 0.99.878 or higher required) installed with the package can be found in the Addins menu (Figure 1). This button allows the direct launch of the application with a single click.

![Figure 1: addin of the app in RStudio (RStudio version >= 0.99.878 required)](image)

2 Description of the application

The application contains three main panels, as described below.

2.1 First panel

The first panel includes two elements: a dashboard for selection of trimming and filtering operations, and a menu for selection of the input file/s (Fig. 2).
An Introduction to FastqCleaner

2.1.1 Selecting operations

The “operations menu” (Fig. 2, elements 1 to 8) shows the available operations for file processing:

1. **Remove by N(s):** removes sequences with a number of Ns (non-identified bases) above a selected threshold value.

2. **Remove low complexity sequences:** remove sequences with a value of complexity above a threshold value.

3. **Remove adapters:** removes adapters and partial adapters. Adapter sequences from both ends of single or paired read reads can be selected. Sequences can be reverse-complemented before processing. The program also allows to consider indels and/or anchored adapters. Two methods can be used for adapter remotion: “exact” or “error rate”. The methods are based on lower-level Biostrings infrastructure, working as the function trimLRpatterns, but optimized for both anchored and non-anchored adapters.

4. **Filter by average quality:** computes the average quality of sequences and removes those with a value below a given threshold.

5. **Trim low quality 3’ tails:** removes the 3’ tails of sequences that are below a given threshold.

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**Figure 2:** Panel 1, with each component indicated with a number. See explanation for each element below.
An Introduction to FastqCleaner

6. **Trim 3’ or 5’ by a fixed number**: removes a fixed number of bases from the 3’ and/or 5’ ends in the complete set of sequences

7. **Filter sequences by length**: removes all the sequences with a number of bases below a threshold value

8. **Remove duplicated sequences**: removes duplicated reads, conserving only one copy of each sequence present in the file

### 2.1.2 Loading files

The “file selection menu” (Fig. 2, elements 9 to 17) contains options to handle the input file (type of file, file selection), buttons to run, clear and reset the application, and the “advanced” submenu:

9. **Single-end reads / paired-end reads**: type of input files

10. “FILE” button: to select an input file

11. “RUN!” button: to run the program

12. Output format: to select whether the output file should be compressed (.gz) or not

13. “CLEAR” button: to clear the configuration of the operations menu that have been selected in the first panel, but keeping the input file(s)

14. “RESET” button: to restart the application, removing the input file(s) and the selected configurations

15. **Selection notificator**: information of the path of the selected file/s

16. **Encoding notificator**: information of the input file/s encoding

17. **Advanced options button**: to select a custom encoding and set the number of reads included in each chunk for processing, as described below

### 2.1.3 Advanced options

The “advanced options submenu” (Fig. 3) allows to customize some fine aspects of the trimming and filtering process:
An Introduction to FastqCleaner

Figure 3: Advanced options submenu

18. **Encoding menu**: in addition to the default approach used by the program (auto-detection of file encoding), users can select a standard encoding from a list.

19. **Chunk size**: the program takes this number of reads at random from the file (default: 1000000), for encoding detection.

### 2.2 Second panel

The second panel (“file operations” panel, Fig. 4) shows the operations that were successfully performed on the input file after running the program.
An Introduction to FastqCleaner

![FastqCleaner Interface](image)

**Figure 4**: File operations panel, with its elements

The panel contains the following elements:

20. **Files location**: location of input and output files

21. **Operations performed**: operations performed on the input file. Each individual display indicates the number of reads that passed the corresponding filter

### 2.3 Third panel

The third panel ( “live results” panel, Fig. 5) shows interactive diagnostics plots for both input and output files. The program takes a random sample of reads for construction of the plots (default: 10000 reads).
An Introduction to FastqCleaner

**Figure 5:** Live results panel

The panel includes the following options in the menu located on the left:

22. **Sample size:** the sample size used for construction of the plots. Default: 10000 reads

23. **Input / output:** show diagnostics plots for input or output files?

24. **Diagnostics plots:** the plot to be shown, that can be one of the following:

   - **Per cycle quality:** quality plots across reads for each cycle (i.e., sequence position)
   - **Per cycle mean quality:** average quality across reads per base, for each cycle (i.e., sequence position)
   - **Mean quality distribution:** Quality distribution, using for the construction of the histogram the mean quality of each read
   - **% reads with Phred scores > threshold:** % of reads with all the quality values > threshold
   - **Per cycle base proportion:** Proportion of each base (average across reads) in each cycle. It also shows the proportion of N’s
   - **CG content:** % CG and % AT (average across reads) for each cycle
   - **CG content distribution over all reads:** histogram for % reads with a given % CG
   - **Read length distribution:** % reads vs read length (bp)
   - **Read occurrence distribution:** % reads that occur at different frequencies values in the file. The plot also includes a table
   - **Relative k-mer diversity:** unique k-mers / all possible kmers for each cycle

25. **Select k-mer size:** k-mer size for the k-mers frequency plot

26. **Top sequences in duplication level analysis:** a list of duplicated sequences, ordered from high to low duplication level, can be desplegated from the “read occurrence distribution” plot. The number selected here indicates how many sequences should be shown. Note that the frequency of reads are relative to the sample size selected (i.e., fold-times in relation to those reads present only once in the sample)
3 A worked example: FASTQ processing in a nutshell

A sample FASTQ (gz-compressed) file ‘example.fastq.gz’ can be downloaded with the following command in R:

```r
download.file("https://goo.gl/hb4Kr9", "example_fastq.gz")
```

A direct download is provided in this link.

A typical FastqCleaner workflow starts with the input file/s upload (Fig. 6).

![File input menu](image)

**Figure 6:** File input menu. The example shows a single-end reads case (sample file 'example.fastq.gz'). For paired-end reads, the selection of the corresponding library type generates an additional button to upload the second file.

The file encoding is automatically detected by the program, but it can also be manually specified in the advanced submenu (Fig. 7). This menu also offers an option to customize the chunk size used for processing.
An Introduction to FastqCleaner

Next, the operations to be performed on the input file are selected from the operations menu (Fig. 8).

Figure 7: Advanced submanu
An Introduction to FastqCleaner

**Figure 8**: Selection of operations. A dialog box shows the input expected for the program. To use a filter, the “Use filter?” checkbox must be checked. A filter in use is indicated with a checkmark in the filter box.

The program then starts to run after pressing the “RUN!” button (Fig. 9).

![Select a file](image)

**Figure 9**: “RUN!” button action

Post-processing results are shown in the second panel (Fig. 10).
An Introduction to FastqCleaner

**Figure 10**: Second panel of the app, showing the operations performed and the paths of the input and output files

The type of plot to be displayed and the options for the construction of the plot are available in the third panel (Fig. 11). This panel also show the selected plot/s.
An Introduction to FastqCleaner

Figure 11: Third panel, showing as example a “CG” content plot. for the output file

To clean the operations, for example to run a different configuration, the “CLEAN” i(Fig. 11) must be pressed. The “RESET” button (Fig. 11) restarts the interface.

Additional help can be found in the “help” button located at the top-right of the app (Fig. 12).

Figure 12: help button. A webpage with information will be open

4 Advanced use of the package

FastqCleaner separates the interface from the implementation. In consequence, the processing functions of the package can be used as standard functions from the command line. Most of the functions make intensive use of Biostrings and ShortRead.Trimming and filtering is performed on ShortReadQ objects. A complete documentation for the functions is available in this link.

The functions included in the package are described in the following section.

4.1 Main functions

- adapter_filter

Based on the Biostrings isMatchingStartingAt and isMatchingEndingAt functions. It can remove adapters and partial adapters from the 3’ and 5’ sequence ends. Adapters can be anchored or not. Two methods are available: one based on the exact matching of the sequences and the adapter, and other in a mismatch rate. For this latter, when indels are allowed, the method is based on the “edit distance” of the sequences.

### Examples

```r
require("Biostrings")
require("ShortRead")
require("FastqCleaner")
```
# create sequences
```r
set.seed(10)
# nota that the use of set.seed before the call to the
# random generators allows reproducibility of the
# examples

input <- random_seq(6, 43)
input
```

## A DNAStringSet instance of length 6
## width seq
## [1] 43 TGGTCCGGTGTTCTGGCGGAATAGGTACAGTCCAGTAATTGCC
## [2] 43 TCCCGCAGACGCTGGGTCCGGAATGCCCTTTCTGAGCAGCTCC
## [3] 43 AGCCGTTTGACTTTCGCGGAAGTGAACTTAGATTCGGTCCTGA
## [4] 43 AACACGGTACTTCCACAGTCAACCGCCGACTTGGAGAATTGA
## [5] 43 TTAGCCGGGCGGTTATTCCCCTAGTAGACTTTACTAAGATTTGC
## [6] 43 AATACCTAAGCGAAGTGACAGATATGTTCGTCATTCATCCAGG

# create qualities of width 50
```r
set.seed(10)
input_q <- random_qual(c(30,40), slength = 6, swidth = 50, encod = "Sanger")
```

# create names
```r
input_names <- seq_names(length(input))
```

### FULL ADAPTER IN 3'
```r
adapter <- "ATCGACT"
```

# Create sequences with adapter
```r
my_seqs <- paste0(input, adapter)
my_seqs <- DNAStringSet(my_seqs)
```

## A DNAStringSet instance of length 6
## width seq
## [1] 50 TGGTCCGGTGTTCTGGCGGAATAGGTACAGTCCAGTAATTGCCATCGACT
## [2] 50 TCCCGCAGACGCTGGGTCCGGAATGCCCTTTCTGAGCAGCTCCATCGACT
## [3] 50 AGCCGTTTGACTTTCGCGGAAGTGAACTTAGATTCGGTCCTGATCGACT
## [4] 50 AACACGGTACTTCCACAGTCAACCGCCGACTTGGAGAATTGATCGACT
## [5] 50 TTAGCCGGGCGGTTATTCCCCTAGTAGACTTTACTAAGATTTGATCGACT
## [6] 50 AATACCTAAGCGAAGTGACAGATATGTTCGTCATTCATCCAGGATCGACT

# create ShortReadQ object
```r
my_read <- ShortReadQ(sread = my_seqs, quality = input_q, id = input_names)
```

# trim adapter
```r
filtered <- adapter_filter(my_read, Lpattern = adapter)
```

## A DNAStringSet instance of length 6
## width seq

---

13
An Introduction to FastqCleaner

### PARTIAL ADAPTER IN 5'

adapter <- "ATCGACT"
subadapter <- subseq(adapter, 1, 4)

# Create sequences with adapter
my_seqs <- paste0(input, subadapter)
my_seqs <- DNAStringSet(my_seqs)
my_seqs
## A DNAStringSet instance of length 6
## width seq
## [1] 47 TGGTCCGGTGTTCTGGCGGAATAGGTACAGTCCAGTAATTGCCATCG
## [2] 47 TCCCGCAGACGCTGGGTCCGGAATGCCCTTTCTGAGCAGCTCCATCG
## [3] 47 AGCCGTTTGACTTCGCGGAAAGTGAACTTAGATTCGGTCCTGAATCG
## [4] 47 AACACGGTACTTCCACAGTCAACCCGCCGACTTGGAGAATTTAATCG
## [5] 47 TTAGCCGGGCGGTTATTCCCCTAGTGATTTACTAAGATTGGCATCG
## [6] 47 AATACCTAAGCAGTGACAGATATGTTCGTCATTCATCCAGGATCG

# create ShortReadQ object
my_read <- ShortReadQ(sread = my_seqs, quality = subseq(input_q, 1, 47), id = input_names)

# trim adapter
filtered <- adapter_filter(my_read, Rpattern = adapter)
sread(filtered)
## A DNAStringSet instance of length 6
## width seq
## [1] 43 TGGTCCGGTGTTCTGGCGGAATAGGTACAGTCCAGTAATTGCC
## [2] 43 TCCCGCAGACGCTGGGTCCGGAATGCCCTTTCTGAGCAGCTCC
## [3] 43 AGCCGTTTGACTTCGCGGAAAGTGAACTTAGATTCGGTCCTGA
## [4] 43 AACACGGTACTTCCACAGTCAACCCGCCGACTTGGAGAATTTA
## [5] 43 TTAGCCGGGCGGTTATTCCCCTAGTGATTTACTAAGATTGG
## [6] 43 AATACCTAAGCAGTGACAGATATGTTCGTCATTCATCCAG

Documentation of the function

• **complex_filter**

Removes low complexity sequences, computing the entropy with the dinucleotide frequency:

\[ H_i = - \sum d_i \times \log_2(d_i) \]

where: \( d_i = \frac{D_i}{\sum_i^n D_i} \) represents the frequency of dinucleotides of the sequence \( i \) relative to the frequency in the whole pool of sequences.
An Introduction to FastqCleaner

The relation $H_i/H_r$ between $H_i$ and a reference entropy value $H_r$ is computed, and the obtained relations are compared with a given complexity threshold. By default the program uses a reference entropy of 3.908, that corresponds to the entropy of the human genome in bits, and a complexity threshold of 0.5.

```
# create sequences of different width
set.seed(10)
input <- lapply(c(0, 6, 10, 16, 20, 26, 30, 36, 40),
    function(x) random_seq(1, x))

# create repetitive "CG" sequences with length adequate
# for a total length input + CG = 40
CG <- lapply(c(20, 17, 15, 12, 10, 7, 5, 2, 0),
    function(x) paste(rep("CG", x), collapse = ""))

# concatenate input and CG
input <- mapply("paste", input, CG, sep = "")
input <- DNAStringSet(input)
input
## A DNAStringSet instance of length 9
## width  seq
## [1] 40   CGCGCGCGCGCGCGCGCGCGCGCGCGCGCGCGCGCGCGCG
## [2] 40   TGGTCCCGCGCGCGCGCGCGCGCGCGCGCGCGCGCGCGCG
## [3] 40   GGTGTTCTGGCGCGCGCGCGCGCGCGCGCGCGCGCGCGCG
## [4] 40   CGGAATAGGTACAGTCCGCGCGCGCGCGCGCGCGCGCGCG
## [5] 40   CAGTAATTGCCTCCCGCATGCCCGCGCGCGCGCGCGCGCG
## [6] 40   CGCTGGGTCCGGAATGCCCTTTCTGACGCGCGCGCGCGCG
## [7] 40   GCAGCTCCAGCGTTGGACTTCCGGAAGCGCGCGCGCG
## [8] 40   TGAACCTAGATTCGGTCCTGAAACACGGTACTTCCACGCG
## [9] 40   CAGTCAACCCCGCCAGTGGAGAATTTATTAGCCGGGCGG

# plot relative entropy (E, Shannon 1948)
H_plot <- function(x, H_max = 3.908135) {
    freq <- dinucleotideFrequency(x)
    freq <- freq / rowSums(freq)
    H <- -rowSums(freq * log2(freq), na.rm = TRUE)
    plot(H/H_max, type="l", xlab = "Sequence", ylab= "E")
    points(H/H_max, col = "#1a81c2", pch = 16, cex = 2)
}
H_plot(input)
```
An Introduction to FastqCleaner

**Figure 13:** Relative entropy plot for the sequences before the operation

```r
set.seed(10)
input_q <- random_qual(c(30,40), slength = 9, swidth = 40,
              encod = "Sanger")

input_names <- seq_names(9)

my_read <- ShortReadQ(sread = input, quality = input_q, id = input_names)

# apply the filter,
filtered <- complex_filter(my_read)

sread(filtered)

H_plot(sread(filtered))
```
An Introduction to FastqCleaner

Figure 14: Relative entropy plot for the sequences after the operation

Documentation of the function

- **fixed_filter**
  
  Removes the specified number of bases from 3’ or 5’.

```r
# create sequences, qualities and names of width 20
set.seed(10)
input <- random_seq(6, 20)
input
## A DNAStringSet instance of length 6
## width seq
##  [1] 20 TGGTCCGGTGTCTGGCGGA
##  [2] 20 ATAGGTACAGTCCAGTAATT
##  [3] 20 GCCTCCGCAGACGCTGGGT
##  [4] 20 CCGGAATGCCCTTTCTGAGC
##  [5] 20 AGCTCCAGCCGTTTGACTTC
##  [6] 20 GCGGAAAGTGAACCTTAGATT

set.seed(10)
input_q <- random_qual(c(30,40), slength = 6, swidth = 20, encod = "Sanger")

input_names <- seq_names(6)

# create ShortReadQ object
my_read <- ShortReadQ(sread = input, quality = input_q, id = input_names)

# apply the filter
filtered3 <- fixed_filter(my_read, trim5 = 5)
sread(filtered3)
## A DNAStringSet instance of length 6
## width seq
```
An Introduction to FastqCleaner

---

```r
## [1] 15 TGGTCCGGTGTTCTG
## [2] 15 ATAGGTACAGTCCAG
## [3] 15 GCCTCCCGCAGACGC
## [4] 15 CCGGAATGCCCTTTC
## [5] 15 AGCTCCAGCGGTGTT
## [6] 15 GCGGAAAGTGAACTT

filtered5 <- fixed_filter(my_read, trim3 = 5)
sread(filtered5)
## A DNAStringSet instance of length 6
## width seq
## [1] 15 CGGTGTTCTGGCGGA
## [2] 15 TACAGTCCAGTAATTGCCTCCCGCAGACGC
## [3] 15 CCGCAGACGCTGGGT
## [4] 15 ATGCCCTTTCCTGAGC
## [5] 15 CAGCCGTTTTGACTTC
## [6] 15 AAGTGAACTTAGATT

filtered3and5 <- fixed_filter(my_read, trim3 = 10, trim5 = 5)
sread(filtered3and5)
## A DNAStringSet instance of length 6
## width seq
## [1] 5 TTCTG
## [2] 5 TCCAG
## [3] 5 GACGC
## [4] 5 CTTTC
## [5] 5 GTTTG
## [6] 5 AACTT
```

Documentation of the function

- **length_filter**

Removes sequences with a length lower than minimum threshold value or/and higher than a maximum threshold value.

```r
# create ShortReadQ object width widths between 1 and 60
set.seed(10)
input <- random_length(10, widths = 1:60)
sread(input)
## A DNAStringSet instance of length 10
## width seq
## [1] 43 TCTGGCGGAATAGGTACAGTCCAGTAATTGCCTCCCGCAGACGC
## [2] 9 CTGGGTCCCG
## [3] 10 GAATGCCCTTTC
## [4] 48 TCTGAGCAGCTCCAGCGGTGTTACTTCGCGGAAAGTGAACTTAGATTCC
## [5] 12 GTCTCCGAAACA
## [6] 55 CGGACTTCCACAGTCAACCGCCCGACTTCGCGGAAATATTATCTAGCCCGGCGTTAT
## [7] 8 TCCCTTAG
## [8] 54 TGATCTTACTAAGATTTTGCATACCTAAGCGAAGTGACAGATATGTTCAT
```
## An Introduction to FastqCleaner

```r
# apply the filter, removing sequences with 5>length> 30
filtered <- length_filter(input, rm.min = 5, rm.max = 30)
ls(filtered)
```

```r
## A DNAStringSet instance of length 5
## width seq
## [1]  9 CTGGGTCCG
## [2] 10 GAATGCCCTT
## [3] 12 GGTCCTGAAACA
## [4]  8 TCCCCTAG
## [5] 19 AAATATGACTCGCGTATCG
```

### Documentation of the function

- **n_filter**

Wrapper of the `ShortRead nFilter` function. Removes all those sequences with a number of N's > a given threshold.

```r
# create 10 sequences of width 20
set.seed(10)
input <- random_seq(10, 20)
input
```

```r
## A DNAStringSet instance of length 10
## width seq
## [1] 20 TGGTCCGGTGTTCTGGCGGA
## [2] 20 ATAGGTACAGTCCAGTAATT
## [3] 20 GCCTCCCGCAGACGCTGGGT
## [4] 20 CCGGAATGCCCTTTCTGAGC
## [5] 20 AGCTCCAGCCGTTTGACTTC
## [6] 20 GCGGAAATGGAACCTTAGATT
## [7] 20 CGGTCCTGAACAGCGTGACT
## [8] 20 TCCACAGTCAACCGGCGAC
## [9] 20 TTGGAGAATTTATGCCCCG
## [10] 20 GCGGTATTCCTCCCTAGTGAT
```

# inject N's

```r
set.seed(10)
input <- inject_letter_random(input, how_many_seqs = 1:5,
how_many = 1:10)
input
```

```r
## A DNAStringSet instance of length 10
## width seq
## [1] 20 TGGTCCCGGTGTTCTGGCGGA
## [2] 20 ATAGGTACAGTCCAGTAATT
## [3] 20 GCCTCCCGCAGACGCTGGGT
## [4] 20 CCGGAATGCCCTTTCTGAGC
## [5] 20 AGCTCCAGCCGTTTGACTTC
## [6] 20 GCGGAAATGGAACCTTAGATT
## [7] 20 CGGTCCTGAACAGCGTGACT
## [8] 20 TCCACAGTCAACCGGCGAC
## [9] 20 TTGGAGAATTTATGCCCCG
## [10] 20 GCGGTATTCCTCCCTAGTGAT
```
An Introduction to FastqCleaner

```
## [7] 20 CNGTCNNNAANCNNNNTACN
## [8] 20 NCCANANTCAAANCCCNAC
## [9] 20 TTGGAGNATNTATTNGCNNN
## [10] 20 GCGGTTATTCCCCTAGTGAT

hist(letterFrequency(input, "N"), breaks = 0:10,
     main = "Ns Frequency", xlab = "# Ns",
     col = "#1a81c2")

hist(letterFrequency(sread(filtered), "N"),
     main = "Ns distribution", xlab = "",
     col = "#1a81c2")
```

**Figure 15:** N’s histogram for the sequences before the filtering operation

```r
# Create qualities, names and ShortReadQ object
set.seed(10)
input_q <- random_qual(10, 20)
input_names <- seq_names(10)
my_read <- ShortReadQ(sread = input, quality = input_q, id = input_names)

# Apply the filter
filtered <- n_filter(my_read, rm.N = 3)
sread(filtered)
## A DNAStringSet instance of length 7
## width seq
## [1] 20 TGGTCCGGGTTTCTGCGGA
## [2] 20 ATAGGTACAGTCCAGTAATT
## [3] 20 GCCTCCGAGACGGCTGGAT
## [4] 20 CCGGAATGCCTTCTGACTG
## [5] 20 AGCTCAGCCGTTTATGCTTC
## [6] 20 GCGGAAAGTGAATCTAGATT
## [7] 20 GCGGTTATTCCCCTAGTGAT
hist(letterFrequency(sread(filtered), "N"),
     main = "Ns distribution", xlab = "",
     col = "#1a81c2")
```
An Introduction to FastqCleaner

**Figure 16**: N’s histogram for the sequences after the filtering operation

**Documentation of the function**

- **qmean_filter**

Removes those sequences with quality < a given threshold.

```r
# create 30 sequences of width 20, 15 with low quality and 15 with high quality
set.seed(10)
input <- random_seq(30, 20)

set.seed(10)
my_qual_H <- random_qual(c(30,40), slength = 15, swidth = 20, encod = “Sanger”)

set.seed(10)
my_qual_L <- random_qual(c(5,30), slength = 15, swidth = 20, encod = “Sanger”)
input_q<- c(my_qual_H, my_qual_L)
input_names <- seq_names(30)
my_read <- ShortReadQ(sread = input, quality = input_q, id = input_names)

# Plot of average qualities
qual_plot <- function(x, cutoff) {
  q <- alphabetScore(x) / width(x)
  plot(q, type="l", xlab = "Sequence", ylab= "Average quality", ylim = c(0, 40))
  points(q, col = "#1a81c2", pch = 16, cex = 2)
  lines(seq_along(q), rep(cutoff, length(q)), type="l", col = "red", lty=2)
  text(length(q), cutoff+2, cutoff)
}

# Plot of average qualities before
qual_plot(my_read, cutoff = 30)
```
An Introduction to FastqCleaner

**Figure 17:** Average qualities before the filtering operation

```r
# Apply the filter
filtered <- qmean_filter(my_read, minq = 30)

# Average qualities after
qual_plot(filtered, cutoff = 30)
```

**Figure 18:** Average qualities after the filtering operation

Documentation of the function

- **seq_filter**

  Removes sequences that match those passed as argument.

```r
# Generate random sequences
set.seed(18)
input <- random_length(30, 3:7)

# Remove sequences that contain the following patterns:
```
An Introduction to FastqCleaner

```r
rm.seq = c("TGGTC", "CGGT", "GTTCT", "ATA")
match_before <- unlist(lapply(rm.seq, function(x) grep(x, as.character(sread(input)))))
match_before
## [1] 14 17 27

filtered <- seq_filter(input, rm.seq = rm.seq)
# Verify that matching sequences were removed
match_after <- unlist(lapply(rm.seq, function(x) {
  grep(x, as.character(sread(filtered))))))
match_after
## [1] 14 26
```

Documentation of the function

- **trim3q_filter**

Removes from the 3’ ends in-tandem nucleotides with a quality < a threshold value.

```r
# Create 6 sequences of width 20
set.seed(10)
input <- random_seq(6, 20)
input
## A DNAStringSet instance of length 6
## width seq
## [1] 20 TGGTCCGGTGTTCTGGCGGA
## [2] 20 ATAGGTACAGTCCAGTAATT
## [3] 20 GCCTCCCGCAGACGCTGGGT
## [4] 20 CCGGAATGCCCTTTCTGAGC
## [5] 20 AGCTCCAGCCGTTTGACTTC
## [6] 20 GCGGAAAGTGAACTTAGATT

# Create Phred+33 qualities of width 15 and paste to qualities of length # 5 used for the tails.
# for three of the sequences, put low qualities in tails
set.seed(10)
my_qual <- random_qual(c(30, 40), slength = 6, swidth = 15,
  encod = "Sanger")
set.seed(10)
tails <- random_qual(c(30, 40), slength = 6, swidth = 5,
  encod = "Sanger")

# Low quality tails in sequences 2, 3 & 4
set.seed(10)
tails[2:4] <- random_qual(c(3, 20), slength = 3, swidth = 5,
  encod = "Sanger")
my_qual <- paste0(my_qual, tails)
input_q <- BStringSet(my_qual)
input_q
```
An Introduction to FastqCleaner

### A BStringSet instance of length 6
### width seq
### [1] 20 EGFEDIBEH@C@DD?EAAID
### [2] 20 I?EDHIBEG?BHFG,%),4
### [3] 20 ACCFBBCFI?I@HBC402+,
### [4] 20 CGIAFG@?AIDF@I14)2+
### [5] 20 IB@CAAC7AGEDDH?BEB
### [6] 20 BH?GFFIHG?DABECFEDE

# Watch qualities before filtering
as.matrix(PhredQuality(input_q))

# Create names and ShortReadQ object
input_names <- seq_names(6)
my_read <- ShortReadQ(sread = input, quality = input_q, id = input_names)

# Apply the filter
filtered <- trim3q_filter(my_read, rm.3qual = 28)
sread(filtered)

Documentation of the function

- **unique_filter**

  Wrapper of the ShortRead occurrenceFilter function. that removes duplicated sequences.

  # Create duplicated sequences
  s <- random_seq(10, 10)
s <- sample(s, 30, replace = TRUE)

# Create a ShortReadQ object
q <- random_qual(30, 10)
n <- seq_names(30)
my_read <- ShortReadQ(sread = s, quality = q, id = n)

# Check presence of duplicates
isUnique(as.character(sread(my_read)))
## [1] FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE
## [13] FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE
## [25] FALSE FALSE FALSE FALSE FALSE TRUE

# Apply the filter
filtered <- unique_filter(my_read)
isUnique(as.character(sread(filtered)))
## [1] TRUE TRUE TRUE TRUE TRUE TRUE TRUE TRUE TRUE TRUE TRUE TRUE TRUE TRUE TRUE TRUE TRUE TRUE TRUE TRUE

4.2 Auxiliary functions

- **random_seq**
  Create a vector of random sequences, for a set of specified parameters.
  Documentation of the function

- **random_qual**
  Create a vector of random qualities for a given encoding and a set of specified parameters.
  Documentation of the function

- **seq_names**
  Create a vector of names for a set of sequences.
  Documentation of the function

- **random_length**
  Create a set of sequences with random lengths.
  Documentation of the function

- **inject_letter_random**
An Introduction to FastqCleaner

Inject a character (e.g., 'N') at random positions, given a set of parameters.

Documentation of the function

- **check_encoding**

  The function allows to check quality encoding. It detects encodings with the following formats:

  | Format      | Expected range |
  |-------------|----------------|
  | Sanger      | [0, 40]        |
  | Illumina 1.8| [0, 41]        |
  | Illumina 1.5| [0, 40]        |
  | Illumina 1.3| [3, 40]        |
  | Solexa      | [-5, 40]       |

Documentation of the function

5 Contact information

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