Pairwise Sequence Alignments

Patrick Aboyoun Gentleman Lab Fred Hutchinson Cancer Research Center Seattle, WA

October 14, 2013

Contents

1	Introduction 2										
2	Pairwise Sequence Alignment Problems	2									
3	Main Pairwise Sequence Alignment Function 3.1 Exercise 1	3 5									
4	Pairwise Sequence Alignment Classes 4.1 Exercise 2	5									
5	Pairwise Sequence Alignment Helper Functions 5.1 Exercise 3	6									
6	Edit Distances 6.1 Exercise 4	11 12									
7	Application: Using Evolutionary Models in Protein Alignments 7.1 Exercise 5	12 12									
8	Application: Removing Adapters from Sequence Reads 8.1 Exercise 6	13 17									
9	Application: Quality Assurance in Sequencing Experiments 9.1 Exercise 7	17 20									
10	Computation Profiling 10.1 Exercise 8	20 23									
11	Computing alignment consensus matrices	23									
12	Exercise Answers 12.1 Exercise 1										
	12.2 Exercise 2	25									
	12.5 Exercise 5										

13 Sess	sion Inform	natio	n																33
12.8	Exercise 8			 			 	•	 •					 •					31
12.7	Exercise 7			 			 												30
12.6	Exercise 6			 			 												26

1 Introduction

In this document we illustrate how to perform pairwise sequence alignments using the Biostrings package through the use of the pairwiseAlignment function. This function aligns a set of pattern strings to a subject string in a global, local, or overlap (ends-free) fashion with or without affine gaps using either a fixed or quality-based substitution scoring scheme. This function's computation time is proportional to the product of the two string lengths being aligned.

2 Pairwise Sequence Alignment Problems

The (Needleman-Wunsch) global, the (Smith-Waterman) local, and (ends-free) overlap pairwise sequence alignment problems are described as follows. Let string S_i have n_i characters $c_{(i,j)}$ with $j \in \{1, \ldots, n_i\}$. A pairwise sequence alignment is a mapping of strings S_1 and S_2 to gapped substrings S'_1 and S'_2 that are defined by

$$S'_{1} = g_{(1,a_{1})}c_{(1,a_{1})} \cdots g_{(1,b_{1})}c_{(1,b_{1})}g_{(1,b_{1}+1)}$$

$$S'_{2} = g_{(2,a_{2})}c_{(2,a_{2})} \cdots g_{(2,b_{2})}c_{(2,b_{2})}g_{(2,b_{2}+1)}$$

where

```
a_i, b_i \in \{1, \dots, n_i\} with a_i \leq b_i
g_{(i,j)} = 0 or more gaps at the specified position j for aligned string i length(S'_1) = length(S'_2)
```

Each of these pairwise sequence alignment problems is solved by maximizing the alignment score. An alignment score is determined by the type of pairwise sequence alignment (global, local, overlap), which sets the $[a_i, b_i]$ ranges for the substrings; the substitution scoring scheme, which sets the distance between aligned characters; and the gap penalties, which is divided into opening and extension components. The optimal pairwise sequence alignment is the pairwise sequence alignment with the largest score for the specified alignment type, substitution scoring scheme, and gap penalties. The pairwise sequence alignment types, substitution scoring schemes, and gap penalties influence alignment scores in the following manner:

Pairwise Sequence Alignment Types: The type of pairwise sequence alignment determines the substring ranges to apply the substitution scoring and gap penalty schemes. For the three primary (global, local, overlap) and two derivative (subject overlap, pattern overlap) pairwise sequence alignment types, the resulting substring ranges are as follows:

```
Global - [a_1,b_1]=[1,n_1] and [a_2,b_2]=[1,n_2]

Local - [a_1,b_1] and [a_2,b_2]

Overlap - \{[a_1,b_1]=[a_1,n_1],[a_2,b_2]=[1,b_2]\} or \{[a_1,b_1]=[1,b_1],[a_2,b_2]=[a_2,n_2]\}

Subject Overlap - [a_1,b_1]=[1,n_1] and [a_2,b_2]

Pattern Overlap - [a_1,b_1] and [a_2,b_2]=[1,n_2]
```

Substitution Scoring Schemes: The substitution scoring scheme sets the values for the aligned character pairings within the substring ranges determined by the type of pairwise sequence alignment. This scoring scheme can be fixed for character pairings or quality-dependent for character pairings. (Characters that align with a gap are penalized according to the "Gap Penalty" framework.)

Fixed substitution scoring - Fixed substitution scoring schemes associate each aligned character pairing with a value. These schemes are very common and include awarding one value for a match and another for a mismatch, Point Accepted Mutation (PAM) matrices, and Block Substitution Matrix (BLOSUM) matrices.

Quality-based substitution scoring - Quality-based substitution scoring schemes derive the value for the aligned character pairing based on the probabilities of character recording errors [3]. Let ϵ_i be the probability of a character recording error. Assuming independence within and between recordings and a uniform background frequency of the different characters, the combined error probability of a mismatch when the underlying characters do match is $\epsilon_c = \epsilon_1 + \epsilon_2 - (n/(n-1)) * \epsilon_1 * \epsilon_2$, where n is the number of characters in the underlying alphabet (e.g. in DNA and RNA, n = 4). Using ϵ_c , the substitution score is given by $b*\log_2(\gamma_{(x,y)}*(1-\epsilon_c)*n+(1-\gamma_{(x,y)})*\epsilon_c*(n/(n-1))$), where b is the bit-scaling for the scoring and $\gamma_{(x,y)}$ is the probability that characters x and y represents the same underlying letters (e.g. using IUPAC, $\gamma_{(A,A)} = 1$ and $\gamma_{(A,N)} = 1/4$).

Gap Penalties: Gap penalties are the values associated with the gaps within the substring ranges determined by the type of pairwise sequence alignment. These penalties are divided into gap opening and gap extension components, where the gap opening penalty is the cost for adding a new gap and the gap extension penalty is the incremental cost incurred along the length of the gap. A constant gap penalty occurs when there is a cost associated with opening a gap, but no cost for the length of a gap (i.e. gap extension is zero). A linear gap penalty occurs when there is no cost associated for opening a gap (i.e. gap opening is zero), but there is a cost for the length of the gap. An affine gap penalty occurs when both the gap opening and gap extension have a non-zero associated cost.

3 Main Pairwise Sequence Alignment Function

The pairwiseAlignment function solves the pairwise sequence alignment problems mentioned above. It aligns one or more strings specified in the *pattern* argument with a single string specified in the *subject* argument.

The gap penalties are regulated by the gap Opening and gap Extension arguments.

The substitution scoring scheme is set using three arguments, two of which are quality-based related (patternQuality, subjectQuality) and one is fixed substitution related (substitutionMatrix). When the substitution scores are fixed by character pairing, the substitutionMatrix argument takes a matrix with the appropriate alphabets as dimension names. The nucleotideSubstitutionMatrix function translates simple match and mismatch scores to the full spectrum of IUPAC nucleotide codes.

```
> submat <-
+ matrix(-1, nrow = 26, ncol = 26, dimnames = list(letters, letters))
> diag(submat) <- 0
> pairwiseAlignment(pattern = c("succeed", "precede"), subject = "supersede",
+ substitutionMatrix = submat,
+ gapOpening = 0, gapExtension = -1)

Global PairwiseAlignmentsSingleSubject (1 of 2)
pattern: [1] succe-ed
subject: [1] supersed
score: -5
```

When the substitution scores are quality-based, the patternQuality and subjectQuality arguments represent the equivalent of [x-99] numeric quality values for the respective strings, and the optional fuzzyMatrix argument represents how the closely two characters match on a [0,1] scale. The patternQuality and subjectQuality arguments accept quality measures in either a PhredQuality, SolexaQuality, or IlluminaQuality scaling. For PhredQuality and IlluminaQuality measures $Q \in [0,99]$, the probability of an error in the base read is given by $10^{-Q/10}$ and for SolexaQuality measures $Q \in [-5,99]$, they are given by $1-1/(1+10^{-Q/10})$. The qualitySubstitutionMatrices function maps the patternQuality and subjectQuality scores to match and mismatch penalties. These three arguments will be demonstrated in later sections.

The final argument, scoreOnly, to the pairwiseAlignment function accepts a logical value to specify whether or not to return just the pairwise sequence alignment score. If scoreOnly is FALSE, the pairwise alignment with the maximum alignment score is returned. If more than one pairwise alignment has the maximum alignment score exists, the first alignment along the subject is returned. If there are multiple pairwise alignments with the maximum alignment score at the chosen subject location, then at each location along the alignment mismatches are given preference to insertions/deletions. For example, pattern: [1] ATTA; subject: [1] ATTA is chosen above pattern: [1] ATTA; subject: [1] A-TA if they both have the maximum alignment score.

```
> submat <-
+ matrix(-1, nrow = 26, ncol = 26, dimnames = list(letters, letters))
> diag(submat) <- 0
> pairwiseAlignment(pattern = c("succeed", "precede"), subject = "supersede",
+ substitutionMatrix = submat,
+ gapOpening = 0, gapExtension = -1, scoreOnly = TRUE)

[1] -5 -5
```

- 1. Using pairwiseAlignment, fit the global, local, and overlap pairwise sequence alignment of the strings "syzygy" and "zyzzyx" using the default settings.
- 2. Do any of the alignments change if the gapExtension argument is set to -Inf?

[Answers provided in section 12.1.]

4 Pairwise Sequence Alignment Classes

Following the design principles of Bioconductor and R, the pairwise sequence alignment functionality in the Biostrings package keeps the end-user close to their data through the use of five specialty classes: PairwiseAlignments, PairwiseAlignmentsSingleSubject, PairwiseAlignmentsSingleSubjectSummary, AlignedXStringSet, and QualityAlignedXStringSet. The PairwiseAlignmentsSingleSubject class inherits from the PairwiseAlignments class and they both hold the results of a fit from the pairwiseAlignment function, with the former class being used to represent all patterns aligning to a single subject and the latter being used to represent elementwise alignments between a set of patterns and a set of subjects.

```
> psa1 <- pairwiseAlignment(pattern = c("succeed", "precede"), subject = "supersede")
> class(psa1)

[1] "PairwiseAlignmentsSingleSubject"
attr(,"package")
[1] "Biostrings"
```

and the pairwiseAlignmentSummary function holds the results of a summarized pairwise sequence alignment.

```
> summary(psa1)
Global Single Subject Pairwise Alignments
Number of Alignments: 2
```

Scores:

```
Min. 1st Qu. Median Mean 3rd Qu. Max. -34.00 -31.78 -29.56 -29.56 -27.34 -25.12
```

Number of matches:

Min.	1st Qu.	Median	Mean 3	rd Qu.	${\tt Max.}$
3.00	3.25	3.50	3.50	3.75	4.00

Top 7 Mismatch Counts:

	SubjectPosition	Subject	Pattern	Count	Probability
1	3	р	С	1	0.5
2	4	e	С	1	0.5
3	4	е	r	1	0.5
4	5	r	е	1	0.5
5	6	s	С	1	0.5
6	8	d	е	1	0.5
7	9	е	d	1	0.5

> class(summary(psa1))

```
[1] "PairwiseAlignmentsSingleSubjectSummary" attr(,"package")[1] "Biostrings"
```

The AlignedXStringSet and QualityAlignedXStringSet classes hold the "gapped" S'_i substrings with the former class holding the results when the pairwise sequence alignment is performed with a fixed substitution scoring scheme and the latter class a quality-based scoring scheme.

4.1 Exercise 2

1. What is the primary benefit of formal summary classes like *PairwiseAlignmentsSingleSubjectSummary* and *summary.lm* to end-users?

[Answer provided in section 12.2.]

5 Pairwise Sequence Alignment Helper Functions

Tables 1, 1 and 3 show functions that interact with objects of class PairwiseAlignments, PairwiseAlignmentsSingleSubject, and AlignedXStringSet. These functions should be used in preference to direct slot extraction from the alignment objects.

The score, nedit, nmatch, nmismatch, and nchar functions return numeric vectors containing information on the pairwise sequence alignment score, number of matches, number of mismatches, and number of aligned characters respectively.

```
> submat <-
+ matrix(-1, nrow = 26, ncol = 26, dimnames = list(letters, letters))
> diag(submat) <- 0
> psa2 <-
+ pairwiseAlignment(pattern = c("succeed", "precede"), subject = "supersede",
+ substitutionMatrix = submat,
+ gapOpening = 0, gapExtension = -1)
> score(psa2)
[1] -5 -5
```

Function	Description
[Extracts the specified elements of the alignment object
alphabet	Extracts the allowable characters in the original strings
compareStrings	Creates character string mashups of the alignments
deletion	Extracts the locations of the gaps inserted into the pattern for the alignments
length	Extracts the number of patterns aligned
mismatchTable	Creates a table for the mismatching positions
nchar	Computes the length of "gapped" substrings
nedit	Computes the Levenshtein edit distance of the alignments
indel	Extracts the locations of the insertion & deletion gaps in the alignments
insertion	Extracts the locations of the gaps inserted into the subject for the alignments
nindel	Computes the number of insertions & deletions in the alignments
nmatch	Computes the number of matching characters in the alignments
nmismatch	Computes the number of mismatching characters in the alignments
pattern, subject	Extracts the aligned pattern/subject
pid	Computes the percent sequence identity
rep	Replicates the elements of the alignment object
score	Extracts the pairwise sequence alignment scores
type	Extracts the type of pairwise sequence alignment

 ${\bf Table~1:~Functions~for~\it Pairwise Alignments~and~\it Pairwise Alignments Single Subject~objects.}$

```
> nedit(psa2)
[1] 4 5
> nmatch(psa2)
[1] 4 4
> nmismatch(psa2)
[1] 3 3
> nchar(psa2)
[1] 8 9
> aligned(psa2)
 A BStringSet instance of length 2
   width seq
[1]
       9 succe-ed-
[2]
       9 pr-ec-ede
> as.character(psa2)
[1] "succe-ed-" "pr-ec-ede"
> as.matrix(psa2)
     [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9]
[1,] "s" "u" "c" "c" "e" "-" "e" "d" "-"
              "-" "e" "c" "-" "e" "d" "e"
[2,] "p" "r"
```

Function	Description
aligned	Creates an XStringSet containing either "filled-with-gaps" or degapped aligned strings
as.character	Creates a character vector version of aligned
as.matrix	Creates an "exploded" character matrix version of aligned
consensusMatrix	Computes a consensus matrix for the alignments
consensusString	Creates the string based on a $50\% + 1$ vote from the consensus matrix
coverage	Computes the alignment coverage along the subject
mismatchSummary	Summarizes the information of the mismatchTable
summary	Summarizes a pairwise sequence alignment
toString	Creates a concatenated string version of aligned
Views	Creates an XStringViews representing the aligned region along the subject

 ${\bf Table~2:~Additional~functions~for~\it Pairwise Alignments Single Subject~objects.}$

> consensusMatrix(psa2)

	[,1]	[,2]	[,3]	[,4]	[,5]	[,6]	[,7]	[,8]	[,9]
-	0	0	1	0	0	2	0	0	1
С	0	0	1	1	1	0	0	0	0
d	0	0	0	0	0	0	0	2	0
е	0	0	0	1	1	0	2	0	1
р	1	0	0	0	0	0	0	0	0
r	0	1	0	0	0	0	0	0	0
s	1	0	0	0	0	0	0	0	0
u	0	1	0	0	0	0	0	0	0

The summary, mismatchTable, and mismatchSummary functions return various summaries of the pairwise sequence alignments.

> summary(psa2)

Global Single Subject Pairwise Alignments Number of Alignments: 2

Scores:

Min. 1st Qu. Median Mean 3rd Qu. Max. -5 -5 -5 -5 -5 -5

Number of matches:

Top 6 Mismatch Counts:

	SubjectPosition	Subject	Pattern	Count	Probability
1	1	s	р	1	0.5
2	2	u	r	1	0.5
3	3	p	С	1	0.5
4	4	е	С	1	0.5
5	5	r	С	1	0.5
6	5	r	е	1	0.5

> mismatchTable(psa2)

PatternId PatternStart PatternEnd PatternSubstring SubjectStart 1 3 3 6 6 3

1	1	3	3	С	3
2	1	4	4	С	4
3	1	5	5	е	5
4	2	1	1	p	1
5	2	2	2	r	2
6	2	4	4	С	5

SubjectEnd SubjectSubstring

1	3	р
2	4	е
3	5	r
4	1	s
5	2	u
6	5	r

> mismatchSummary(psa2)

\$pattern

\$pattern\$position

	Position	Count	Probability
1	1	1	0.5
2	2	1	0.5
3	3	1	0.5
4	4	2	1.0
5	5	1	0.5
6	6	0	0.0
7	7	0	0.0

\$subject

	SubjectPosition	Subject	${\tt Pattern}$	${\tt Count}$	Probability
	1	s	р	1	0.5
:	2 2	u	r	1	0.5
;	3	p	С	1	0.5
4	1 4	е	С	1	0.5
į	5 5	r	С	1	0.5
(5 5	r	е	1	0.5

The pattern and subject functions extract the aligned pattern and subject objects for further analysis. Most of the actions that can be performed on PairwiseAlignments objects can also be performed on AlignedXStringSet and QualityAlignedXStringSet objects as well as operations including start, end, and width that extracts the start, end, and width of the alignment ranges.

> class(pattern(psa2))

- [1] "AlignedXStringSet"
 attr(,"package")
 [1] "Biostrings"
- > aligned(pattern(psa2))
 - A BStringSet instance of length 2 width seq

Function	Description
]	Extracts the specified elements of the alignment object
aligned, unaligned	Extracts the aligned/unaligned strings
alphabet	Extracts the allowable characters in the original strings
as.character, toString	Converts the alignments to character strings
coverage	Computes the alignment coverage
end	Extracts the ending index of the aligned range
indel	Extracts the insertion/deletion locations
length	Extracts the number of patterns aligned
mismatch	Extracts the position of the mismatches
mismatchSummary	Summarizes the information of the mismatchTable
mismatchTable	Creates a table for the mismatching positions
nchar	Computes the length of "gapped" substrings
nindel	Computes the number of insertions/deletions in the alignments
nmismatch	Computes the number of mismatching characters in the alignments
rep	Replicates the elements of the alignment object
start	Extracts the starting index of the aligned range
toString	Creates a concatenated string containing the alignments
width	Extracts the width of the aligned range

Table 3: Functions for AlignedXString and QualityAlignedXString objects.

```
[1] 8 succe-ed
[2] 9 pr-ec-ede

> nindel(pattern(psa2))
    Length WidthSum
[1,] 1 1
[2,] 2 2

> start(subject(psa2))
[1] 1 1

> end(subject(psa2))
[1] 8 9
```

For the overlap pairwise sequence alignment of the strings "syzygy" and "zyzzyx" with the pairwiseAlignment default settings, perform the following operations:

- 1. Use nmatch and nmismath to extract the number of matches and mismatches respectively.
- 2. Use the compareStrings function to get the symbolic representation of the alignment.
- 3. Use the as.character function to the get the character string versions of the alignments.
- 4. Use the pattern function to extract the aligned pattern and apply the mismatch function to it to find the locations of the mismatches.
- 5. Use the subject function to extract the aligned subject and apply the aligned function to it to get the aligned strings.

[Answers provided in section 12.3.]

6 Edit Distances

One of the earliest uses of pairwise sequence alignment is in the area of text analysis. In 1965 Vladimir Levenshtein considered a metric, now called the *Levenshtein edit distance*, that measures the similarity between two strings. This distance metric is equivalent to the negative of the score of a pairwise sequence alignment with a match cost of 0, a mismatch cost of -1, a gap opening penalty of 0, and a gap extension cost of -1.

The stringDist uses the internals of the pairwiseAlignment function to calculate the Levenshtein edit distance matrix for a set of strings.

There is also an implementation of approximate string matching using Levenshtein edit distance in the agrep (approximate grep) function of the base R package. As the following example shows, it is possible to replicate the agrep function using the pairwiseAlignment function. Since the agrep function is vectorized in x rather than pattern, these arguments are flipped in the call to pairwiseAlignment.

```
> agrepBioC <-
+ function(pattern, x, ignore.case = FALSE, value = FALSE, max.distance = 0.1)
    if (!is.character(pattern)) pattern <- as.character(pattern)</pre>
    if (!is.character(x)) x <- as.character(x)</pre>
    if (max.distance < 1)
      max.distance <- ceiling(max.distance / nchar(pattern))</pre>
    characters <- unique(unlist(strsplit(c(pattern, x), "", fixed = TRUE)))</pre>
    if (ignore.case)
      substitutionMatrix <-
+
        outer(tolower(characters), tolower(characters), function(x,y) -as.numeric(x!=y))
      substitutionMatrix <-
        outer(characters, characters, function(x,y) -as.numeric(x!=y))
    dimnames(substitutionMatrix) <- list(characters, characters)</pre>
+
    distance <-
+
      - pairwiseAlignment(pattern = x, subject = pattern,
                           substitutionMatrix = substitutionMatrix,
                           type = "local-global",
                           gapOpening = 0, gapExtension = -1,
                           scoreOnly = TRUE)
+
    whichClose <- which(distance <= max.distance)</pre>
    if (value)
      whichClose <- x[whichClose]</pre>
+
    whichClose
+ }
> cbind(base = agrep("laysy", c("1 lazy", "1", "1 LAZY"), max = 2, value = TRUE),
        bioc = agrepBioC("laysy", c("1 lazy", "1", "1 LAZY"), max = 2, value = TRUE))
     base
[1,] "1 lazy" "1 lazy"
> cbind(base = agrep("laysy", c("1 lazy", "1", "1 LAZY"), max = 2, ignore.case = TRUE),
        bioc = agrepBioC("laysy", c("1 lazy", "1", "1 LAZY"), max = 2, ignore.case = TRUE))
     base bioc
[1,]
        1
[2,]
             3
        3
```

- 1. Use the pairwiseAlignment function to find the Levenshtein edit distance between "syzygy" and "zyzzyx".
- 2. Use the stringDist function to find the Levenshtein edit distance for the vector c("zyzzyx", "syzygy", "succeed", "precede", "supersede").

[Answers provided in section 12.4.]

7 Application: Using Evolutionary Models in Protein Alignments

When proteins are believed to descend from a common ancestor, evolutionary models can be used as a guide in pairwise sequence alignments. The two most common families evolutionary models of proteins used in pairwise sequence alignments are Point Accepted Mutation (PAM) matrices, which are based on explicit evolutionary models, and Block Substitution Matrix (BLOSUM) matrices, which are based on data-derived evolution models. The Biostrings package contains 5 PAM and 5 BLOSUM matrices (PAM30 PAM40, PAM70, PAM120, PAM250, BLOSUM45, BLOSUM50, BLOSUM62, BLOSUM80, and BLOSUM100) that can be used in the substitutionMatrix argument to the pairwiseAlignment function.

Here is an example pairwise sequence alignment of amino acids from Durbin, Eddy et al being fit by the pairwiseAlignment function using the BLOSUM50 matrix:

```
> data(BLOSUM50)
> BLOSUM50[1:4,1:4]
   A R N D
  5 -2 -1 -2
R -2 7 -1 -2
N - 1 - 1 7
D -2 -2 2 8
> nwdemo <-
    pairwiseAlignment(AAString("PAWHEAE"), AAString("HEAGAWGHEE"), substitutionMatrix = BLOSUM50,
                      gapOpening = 0, gapExtension = -8)
> nwdemo
Global PairwiseAlignmentsSingleSubject (1 of 1)
pattern: [1] PA--W-HEAE
subject: [2] EAGAWGHE-E
score: 1
> compareStrings(nwdemo)
[1] "?A--W-HE+E"
> pid(nwdemo)
[1] 50
```

7.1 Exercise 5

- 1. Repeat the alignment exercise above using BLOSUM62, a gap opening penalty of -12, and a gap extension penalty of -4.
- 2. Explore to find out what caused the alignment to change.

[Answers provided in section 12.5.]

8 Application: Removing Adapters from Sequence Reads

Finding and removing uninteresting experiment process-related fragments like adapters is a common problem in genetic sequencing, and pairwise sequence alignment is well-suited to address this issue. When adapters are used to anchor or extend a sequence during the experiment process, they either intentionally or unintentionally become sequenced during the read process. The following code simulates what sequences with adapter fragments at either end could look like during an experiment.

```
> simulateReads <-
+ function(N, adapter, experiment, substitutionRate = 0.01, gapRate = 0.001) {
    chars <- strsplit(as.character(adapter), "")[[1]]</pre>
    sapply(seq_len(N), function(i, experiment, substitutionRate, gapRate) {
      width <- experiment[["width"]][i]</pre>
      side <- experiment[["side"]][i]</pre>
      randomLetters <-
        function(n) sample(DNA_ALPHABET[1:4], n, replace = TRUE)
      randomLettersWithEmpty <-
        function(n)
        sample(c("", DNA_ALPHABET[1:4]), n, replace = TRUE,
               prob = c(1 - gapRate, rep(gapRate/4, 4)))
      nChars <- length(chars)</pre>
      value <-
        paste(ifelse(rbinom(nChars,1,substitutionRate), randomLetters(nChars), chars),
              randomLettersWithEmpty(nChars),
              sep = "", collapse = "")
+
      if (side)
        value <-
+
          paste(c(randomLetters(36 - width), substring(value, 1, width)),
                  sep = "", collapse = "")
+
      else
        value <-
          paste(c(substring(value, 37 - width, 36), randomLetters(36 - width)),
                  sep = "", collapse = "")
      value
    }, experiment = experiment, substitutionRate = substitutionRate, gapRate = gapRate)
+ }
> adapter <- DNAString("GATCGGAAGAGCTCGTATGCCGTCTTCTGCTTGAAA")</pre>
> set.seed(123)
> N <- 1000
> experiment <-
   list(side = rbinom(N, 1, 0.5), width = sample(0:36, N, replace = TRUE))
> table(experiment[["side"]], experiment[["width"]])
     0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21
 0 13 10 8 7 11 18 9 15 15 18 16 10 11 9 13 13 18 18 14 9 19 13
  1 15 21 21 12 17 14 8 11 12 10 14 16 7 14 19 14 14 16 14 16 16 16
    22 23 24 25 26 27 28 29 30 31 32 33 34 35 36
  0 19 19 12 6 15 18 12 15 16 17 19 6 13 18 15
  1 13 11 9 11 15 10 10 16 15 15 11 7 12 7 14
> adapterStrings <-
   simulateReads(N, adapter, experiment, substitutionRate = 0.01, gapRate = 0.001)
```

> adapterStrings <- DNAStringSet(adapterStrings)</pre>

These simulated strings above have 0 to 36 characters from the adapters attached to either end. We can use completely random strings as a baseline for any pairwise sequence alignment methodology we develop to remove the adapter characters.

```
> M <- 5000
> randomStrings <-
+ apply(matrix(sample(DNA_ALPHABET[1:4], 36 * M, replace = TRUE),
+ nrow = M), 1, paste, collapse = "")
> randomStrings <- DNAStringSet(randomStrings)</pre>
```

Since edit distances are easy to explain, it serves as a good place to start for developing a adapter removal methodology. Unfortunately given that it is based on a global alignment, it only is useful for filtering out sequences that are derived primarily from the adapter.

```
> ## Method 1: Use edit distance with an FDR of 1e-03
> submat1 <- nucleotideSubstitutionMatrix(match = 0, mismatch = -1, baseOnly = TRUE)
> randomScores1 <-
   pairwiseAlignment(randomStrings, adapter, substitutionMatrix = submat1,
                     gapOpening = 0, gapExtension = -1, scoreOnly = TRUE)
> quantile(randomScores1, seq(0.99, 1, by = 0.001))
         99.1%
                 99.2%
                        99.3%
                                99.4%
                                        99.5%
                                                99.6%
                                                       99.7%
   99%
                                                               99.8%
-16.990 -16.000 -16.000 -16.000 -16.000 -16.000 -16.000 -16.000 -15.998
 99.9%
          100%
-15.000 -14.000
> adapterAligns1 <-
   pairwiseAlignment(adapterStrings, adapter, substitutionMatrix = submat1,
                     gapOpening = 0, gapExtension = -1)
> table(score(adapterAligns1) > quantile(randomScores1, 0.999), experiment[["width"]])
        0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20
 FALSE 28 31 29 19 28 32 17 26 27 28 30 26 18 23 32 27 32 34 28 25 35
        0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
       21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36
 FALSE 29 32 30 21 17 30 28 22 7 0 0 0 0 0 0
 TRUE
        0 0 0 0 0 0 0 24 31 32 30 13 25 25 29
```

One improvement to removing adapters is to look at consecutive matches anywhere within the sequence. This is more versatile than the edit distance method, but it requires a relatively large number of consecutive matches and is susceptible to issues related to error related substitutions and insertions/deletions.

```
> ## Method 2: Use consecutive matches anywhere in string with an FDR of 1e-03
> submat2 <- nucleotideSubstitutionMatrix(match = 1, mismatch = -Inf, baseOnly = TRUE)
> randomScores2 <-
+ pairwiseAlignment(randomStrings, adapter, substitutionMatrix = submat2,
+ type = "local", gapOpening = 0, gapExtension = -Inf,
+ scoreOnly = TRUE)
> quantile(randomScores2, seq(0.99, 1, by = 0.001))
```

```
99% 99.1% 99.2% 99.3% 99.4% 99.5% 99.6% 99.7% 99.8% 99.9% 100%
   7
        8
               8
                   8
                          8
                                8
                                      8
                                          8
                                                  9
                                                            11
> adapterAligns2 <-
   pairwiseAlignment(adapterStrings, adapter, substitutionMatrix = submat2,
                     type = "local", gapOpening = 0, gapExtension = -Inf)
> table(score(adapterAligns2) > quantile(randomScores2, 0.999), experiment[["width"]])
        0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20
 FALSE 28 31 29 19 28 32 17 26 27 28 5 1 3 1 1 1 1 2 1 1 0
 TRUE 0 0 0 0 0 0 0 0 0 0 25 25 15 22 31 26 31 32 27 24 35
       21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36
 FALSE 0 0 0 0 0 0 0 0 0 0 0 0 0 0
 TRUE 29 32 30 21 17 30 28 22 31 31 32 30 13 25 25 29
> # Determine if the correct end was chosen
> table(start(pattern(adapterAligns2)) > 37 - end(pattern(adapterAligns2)),
       experiment[["side"]])
         0
             1
 FALSE 469 53
 TRUE
        38 440
  Limiting consecutive matches to the ends provides better results, but it doesn't resolve the issues related
to substitutions and insertions/deletions errors.
> ## Method 3: Use consecutive matches on the ends with an FDR of 1e-03
> submat3 <- nucleotideSubstitutionMatrix(match = 1, mismatch = -Inf, baseOnly = TRUE)
> randomScores3 <-
   pairwiseAlignment(randomStrings, adapter, substitutionMatrix = submat3,
                     type = "overlap", gapOpening = 0, gapExtension = -Inf,
                     scoreOnly = TRUE)
> quantile(randomScores3, seq(0.99, 1, by = 0.001))
  99% 99.1% 99.2% 99.3% 99.4% 99.5% 99.6% 99.7% 99.8% 99.9% 100%
                          4
                                4
                                      4
                                            4
                                                  5 6
> adapterAligns3 <-
   pairwiseAlignment(adapterStrings, adapter, substitutionMatrix = submat3,
                     type = "overlap", gapOpening = 0, gapExtension = -Inf)
> table(score(adapterAligns3) > quantile(randomScores3, 0.999), experiment[["width"]])
        0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20
 FALSE 28 31 29 19 28 32 17 1 2 5 6 1 4 2 2 2 5 7 4 3 3
 TRUE 0 0 0 0 0 0 0 25 25 23 24 25 14 21 30 25 27 27 24 22 32
       21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36
 FALSE 6 2 4 6 2 8 6 8 7 12 7 8 2 7 7 5
 TRUE 23 30 26 15 15 22 22 14 24 19 25 22 11 18 18 24
> # Determine if the correct end was chosen
```

> table(end(pattern(adapterAligns3)) == 36, experiment[["side"]])

```
0 1
FALSE 466 71
TRUE 41 422
```

Allowing for substitutions and insertions/deletions errors in the pairwise sequence alignments provides much better results for finding adapter fragments.

```
> ## Method 4: Allow mismatches and indels on the ends with an FDR of 1e-03
> randomScores4 <-
   pairwiseAlignment(randomStrings, adapter, type = "overlap", scoreOnly = TRUE)
> quantile(randomScores4, seq(0.99, 1, by = 0.001))
             99.1%
                      99.2%
                                99.3%
                                         99.4%
                                                   99.5%
7.927024 7.927024 7.927024 7.927024 7.927024 7.927024 7.927024
   99.7%
             99.8%
                      99.9%
                                 100%
9.908780 9.908872 11.890536 13.872293
> adapterAligns4 <-
   pairwiseAlignment(adapterStrings, adapter, type = "overlap")
> table(score(adapterAligns4) > quantile(randomScores4, 0.999), experiment[["width"]])
        0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20
 FALSE 28 31 29 19 28 32 17 1 2 5 1 0 1 0 1 0 1 0 0 0
        0 0 0 0 0 0 25 25 23 29 26 17 23 31 27 31 34 28 25 35
       21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36
 FALSE 0 0 0 0 0 0 0 0 0 0 0 0 0 0
 TRUE 29 32 30 21 17 30 28 22 31 31 32 30 13 25 25 29
> # Determine if the correct end was chosen
> table(end(pattern(adapterAligns4)) == 36, experiment[["side"]])
         0
             1
 FALSE 484 17
 TRUE.
        23 476
```

Using the results that allow for substitutions and insertions/deletions errors, the cleaned sequence fragments can be generated as follows:

```
> ## Method 4 continued: Remove adapter fragments
> fragmentFound <-
    score(adapterAligns4) > quantile(randomScores4, 0.999)
> fragmentFoundAt1 <-
    fragmentFound & (start(pattern(adapterAligns4)) == 1)
> fragmentFoundAt36 <-
    fragmentFound & (end(pattern(adapterAligns4)) == 36)
> cleanedStrings <- as.character(adapterStrings)</pre>
> cleanedStrings[fragmentFoundAt1] <-</pre>
    as.character(narrow(adapterStrings[fragmentFoundAt1], end = 36,
           width = 36 - end(pattern(adapterAligns4[fragmentFoundAt1]))))
> cleanedStrings[fragmentFoundAt36] <-
    as.character(narrow(adapterStrings[fragmentFoundAt36], start = 1,
           width = start(pattern(adapterAligns4[fragmentFoundAt36])) - 1))
> cleanedStrings <- DNAStringSet(cleanedStrings)</pre>
> cleanedStrings
```

```
A DNAStringSet instance of length 1000
       width seq
   Γ17
          26 TTGCACGATAGTTGCATATGCTACAA
   [2]
          15 ATTTCTCCTTCTCAG
   [3]
          36 TGAAACCGGGTCACCATCATCATCTAGTAGGACCTT
   [4]
           5 GCGTT
   [5]
           5 GCTTA
 [996]
          3 TAA
 [997]
          25 TTGTTTGGGAAGGCTTCAACCAAAC
          9 CAGCTAATA
 [998]
[999]
          22 CGGATGGGAGCCGAGCATGACT
[1000]
          20 TCGCTATCTGAAGTCTGTTT
```

- 1. Rerun the simulation time using the simulateReads function with a *substitutionRate* of 0.005 and *gapRate* of 0.0005. How do the different pairwise sequence alignment methods compare?
- 2. (Advanced) Modify the simulateReads function to accept different equal length adapters on either side (left & right) of the reads. How would the methods for trimming the reads change?

[Answers provided in section 12.6.]

9 Application: Quality Assurance in Sequencing Experiments

Due to its flexibility, the pairwiseAlignment function is able to diagnose sequence matching-related issues that arise when matchPDict and its related functions don't find a match. This section contains an example involving a short read Solexa sequencing experiment of bacteriophage ϕ X174 DNA produced by New England BioLabs (NEB). This experiment contains slightly less than 5000 unique short reads in srPhiX174, with quality measures in quPhiX174, and frequency for those short reads in wtPhiX174.

In order to demonstrate how to find sequence differences in the target, these short reads will be compared against the bacteriophage ϕ X174 genome NC_001422 from the GenBank database.

```
> data(phiX174Phage)
> genBankPhage <- phiX174Phage[[1]]</pre>
> nchar(genBankPhage)
[1] 5386
> data(srPhiX174)
> srPhiX174
  A DNAStringSet instance of length 1113
       width seq
          35 GTTATTATACCGTCAAGGACTGTGTGACTATTGAC
   [1]
   [2]
          35 GGTGGTTATTATACCGTCAAGGACTGTGTGACTAT
   [3]
          35 TACCGTCAAGGACTGTGTGACTATTGACGTCCTTC
   ۲4٦
          35 GTACGCCGGGCAATAATGTTTATGTTGGTTTCATG
   [5]
          35 GGTTTCATGGTTTGGTCTAACTTTACCGCTACTAA
[1109]
          35 ATAATGTTTATGTTGGTTTCATGGTTTGTTCTATC
```

```
[1111]
         35 CAATAATGTTTATGTTGGTTTCATGGTTTGTTTTA
[1112]
         35 GACGTCCTTCCTCGTACGCCGGGCAATGATGTTTA
Γ11137
         35 ACGCCGGCAATAATGTTTATGTTGTTTTCATTGT
> quPhiX174
 A BStringSet instance of length 1113
      width seq
  [1]
         35 ZYZZZZZZZZZYYZZYYYYYYYYYYYYYYYYYYYYY
  [2]
         [3]
         35 ZZZYZYYZYYZYYZYYYYYYYYYYYYYYYYYYYYY
  [4]
         35 ZZZZZYZYYZZZYYYYYYYYYYYYYYYYYYYYY
  [5]
  . . .
[1109]
         35 ZZZZZYZZZYZYZYZYYYYYYYYYYYYQYYYQCYQYQCT
[1110]
         35 YYYYTYYYYYYYYYYYTJTTYOAYIIYYYGAY
         35 ZZYZZZZZZZZZZZYZYYVYYYYYYVQYYYIQYAYW
[1111]
         35 YZYZZYYYZYYYYYYVYYVYYYWWVYYYYYWYYV
[1112]
Γ11137
         35 ZZYYZYYYYYYZYVZYYYYYYYYYYJAYYYIGYCJY
> summary(wtPhiX174)
  Min. 1st Qu. Median
                       Mean 3rd Qu.
                                       Max.
  2.00
         2.00 3.00 48.34 6.00 965.00
> fullShortReads <- rep(srPhiX174, wtPhiX174)</pre>
> srPDict <- PDict(fullShortReads)
> table(countPDict(srPDict, genBankPhage))
37018 16784
```

35 GGGCAATAATGTTTATGTTGGTTTCATTTTTTT

Γ11107

For these short reads, the pairwiseAlignment function finds that the small number of perfect matches is due to two locations on the bacteriophage $\phi X174$ genome.

Unlike the countPDict function, the pairwiseAlignment function works off of the original strings, rather than PDict processed strings, and to be computationally efficient it is recommended that the unique sequences are supplied to the pairwiseAlignment function, and the frequencies of those sequences are supplied to the weight argument of functions like summary, mismatchSummary, and coverage. For the purposes of this exercise, a substring of the GenBank bacteriophage ϕ X174 genome is supplied to the subject argument of the pairwiseAlignment function to reduce the computation time.

```
> genBankSubstring <- substring(genBankPhage, 2793-34, 2811+34)
> genBankAlign <-
+ pairwiseAlignment(srPhiX174, genBankSubstring,
+ patternQuality = SolexaQuality(quPhiX174),
+ subjectQuality = SolexaQuality(99L),
+ type = "global-local")
> summary(genBankAlign, weight = wtPhiX174)

Global-Local Single Subject Pairwise Alignments
Number of Alignments: 53802
```

```
-45.08
          35.81 50.07
                           41.24
                                   59.50
                                           67.35
Number of matches:
  Min. 1st Qu. Median
                           Mean 3rd Qu.
                                            Max.
  21.00
          31.00
                  33.00
                           31.46
                                   34.00
                                           35.00
Top 10 Mismatch Counts:
   SubjectPosition Subject Pattern Count Probability
                         С
                53
                                  T 22965 0.95536234
1
2
                35
                          С
                                  T 22849
                                          0.99969373
3
                76
                         G
                                  T 1985 0.10062351
4
                69
                         Α
                                  Т
                                     1296 0.05654697
                                  T 1289 0.07289899
5
                79
                         С
6
                                 C 1153 0.04783637
                58
                         Α
7
                72
                         G
                                 A 1130 0.05248978
8
                63
                         G
                                  A 1130 0.04767731
9
                         Т
                67
                                  G 1130 0.04721514
10
                81
                         Α
                                  G 1103 0.06672313
> revisedPhage <-
    replaceLetterAt(genBankPhage, c(2793, 2811), "TT")
> table(countPDict(srPDict, revisedPhage))
    0
          1
 6768 47034
   The following plot shows the coverage of the aligned short reads along the substring of the bacteriophage
\phi X174 genome. Applying the slice function to the coverage shows the entire substring is covered by aligned
short reads.
> genBankCoverage <- coverage(genBankAlign, weight = wtPhiX174)
> plot((2793-34):(2811+34), as.integer(genBankCoverage), xlab = "Position", ylab = "Coverage",
       type = "1")
> nchar(genBankSubstring)
[1] 87
```

Max.

> slice(genBankCoverage, lower = 1)

Views on a 87-length Rle subject

start end width 1 87 87

views:

Mean 3rd Qu.

Scores:

Min. 1st Qu. Median

87 [8899 9698 10484 11228 11951 12995 13547 ...]



- 1. Rerun the global-local alignment of the short reads against the entire genome. (This may take a few minutes.)
- 2. Plot the coverage of these alignments and use the slice function to find the ranges of alignment. Are there any alignments outside of the substring region that was used above?
- 3. Use the reverseComplement function on the bacteriophage ϕ X174 genome. Do any short reads have a higher alignment score on this new sequence than on the original sequence?

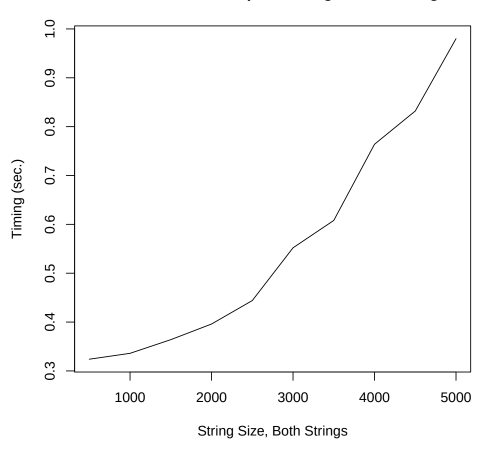
[Answers provided in section 12.7.]

10 Computation Profiling

The pairwiseAlignment function uses a dynamic programming algorithm based on the Needleman-Wunsch and Smith-Waterman algorithms for global and local pairwise sequence alignments respectively. The algorithm consumes memory and computation time proportional to the product of the length of the two strings being aligned.

```
> N <- as.integer(seq(500, 5000, by = 500))
> timings <- rep(0, length(N))</pre>
> names(timings) <- as.character(N)</pre>
> for (i in seq_len(length(N))) {
+ string1 <- DNAString(paste(sample(DNA_ALPHABET[1:4], N[i], replace = TRUE), collapse = ""))
+ string2 <- DNAString(paste(sample(DNA_ALPHABET[1:4], N[i], replace = TRUE), collapse = ""))
  timings[i] <- system.time(pairwiseAlignment(string1, string2, type = "global"))[["user.self"]]</pre>
+ }
> timings
  500 1000 1500 2000 2500 3000 3500 4000 4500 5000
0.324 0.336 0.364 0.396 0.444 0.552 0.608 0.764 0.832 0.980
> coef(summary(lm(timings ~ poly(N, 2))))
             Estimate Std. Error t value
(Intercept) 0.5600000 0.005939814 94.279053 3.979285e-12
poly(N, 2)1 0.6671840 0.018783340 35.519990 3.639060e-09
poly(N, 2)2 0.1772111 0.018783340 9.434481 3.136055e-05
> plot(N, timings, xlab = "String Size, Both Strings", ylab = "Timing (sec.)", type = "1",
      main = "Global Pairwise Sequence Alignment Timings")
```

Global Pairwise Sequence Alignment Timings



When a problem only requires the pairwise sequence alignment score, setting the *scoreOnly* argument to TRUE will more than halve the computation time.

4500

0.01 0.00 0.09

4000

500 1000 1500 2000 2500 3000 3500

0.01 -0.01 -0.01 -0.09 -0.12 -0.02 -0.01

- 1. Rerun the first set of profiling code, but this time fix the number of characters in string1 to 35 and have the number of characters in string2 range from 5000, 50000, by increments of 5000. What is the computational order of this simulation exercise?
- 2. Rerun the second set of profiling code using the simulations from the previous exercise with *scoreOnly* argument set to TRUE. Is is still twice as fast?

[Answers provided in section 12.8.]

11 Computing alignment consensus matrices

The consensusMatrix function is provided for computing a consensus matrix for a set of equal-length strings assumed to be aligned. To illustrate, the following application assumes the ORF data to be aligned for the first 10 positions (patently false):

```
> file <- system.file("extdata", "someORF.fa", package="Biostrings")</pre>
> orf <- readDNAStringSet(file)</pre>
> orf
 A DNAStringSet instance of length 7
                                                 names
   5573 ACTTGTAAATATCTTTT...TCGACCTTATTGTTGATAT YALOO1C TFC3 SGDI...
[1]
    5825 TTCCAAGGCCGATGAATTC...AATTTTTTTCTATTCTCTT YALOO2W VPS8 SGDI...
    2987 CTTCATGTCAGCCTGCACT...ACTCATGTAGCTGCCTCAT YALOO3W EFB1 SGDI...
    3929 CACTCATATCGGGGGTCTT...CCGAAACACGAAAAAGTAC YALOO5C SSA1 SGDI...
    2648 AGAGAAAGAGTTTCACTTC...AATTTATGTGTGAACATAG YALOO7C ERP2 SGDI...
[5]
    2597 GTGTCCGGGCCTCGCAGGC...TTTGGCAGAATGTACTTTT YALOO8W FUN14 SGD...
    2780 CAAGATAATGTCAAAGTTA...AGGAAGAAAAAAAAATCAC YALOO9W SPO7 SGDI...
> orf10 <- DNAStringSet(orf, end=10)</pre>
> consensusMatrix(orf10, as.prob=TRUE, baseOnly=TRUE)
          [,1]
                    [,2]
                             [,3]
                                       [,4]
                                                [,5]
                                                          [,6]
     0.2857143 0.2857143 0.2857143 0.0000000 0.5714286 0.4285714
Α
C
     0.4285714 0.1428571 0.2857143 0.2857143 0.2857143 0.1428571
G
     0.1428571 0.1428571 0.1428571 0.2857143 0.1428571 0.0000000
Т
     0.1428571 0.4285714 0.2857143 0.4285714 0.0000000 0.4285714
[,7]
                    [,8]
                             [,9]
     0.4285714 0.4285714 0.2857143 0.1428571
Α
C
     0.0000000 0.0000000 0.2857143 0.4285714
     0.4285714 0.4285714 0.1428571 0.2857143
G
     0.1428571 0.1428571 0.2857143 0.1428571
```

The information content as defined by Hertz and Stormo 1995 is computed as follows:

```
> informationContent <- function(Lmers) {
+ zlog <- function(x) ifelse(x==0,0,log(x))
+ co <- consensusMatrix(Lmers, as.prob=TRUE)
+ lets <- rownames(co)</pre>
```

```
+ fr <- alphabetFrequency(Lmers, collapse=TRUE)[lets]
+ fr <- fr / sum(fr)
+ sum(co*zlog(co/fr), na.rm=TRUE)
+ }
> informationContent(orf10)
[1] 2.167186
```

12 Exercise Answers

12.1 Exercise 1

1. Using pairwiseAlignment, fit the global, local, and overlap pairwise sequence alignment of the strings "syzygy" and "zyzzyx" using the default settings.

```
> pairwiseAlignment("zyzzyx", "syzygy")
  Global PairwiseAlignmentsSingleSubject (1 of 1)
  pattern: [1] zyzzyx
  subject: [1] syzygy
  score: -19.3607
  > pairwiseAlignment("zyzzyx", "syzygy", type = "local")
  Local PairwiseAlignmentsSingleSubject (1 of 1)
  pattern: [2] yz
  subject: [2] yz
  score: 4.607359
  > pairwiseAlignment("zyzzyx", "syzygy", type = "overlap")
  Overlap PairwiseAlignmentsSingleSubject (1 of 1)
  pattern: [1] ""
  subject: [1] ""
  score: 0
2. Do any of the alignments change if the gapExtension argument is set to -Inf? Yes, the overlap pairwise
  sequence alignment changes.
  > pairwiseAlignment("zyzzyx", "syzygy", type = "overlap", gapExtension = -Inf)
  Overlap PairwiseAlignmentsSingleSubject (1 of 1)
  pattern: [1] ""
```

12.2 Exercise 2

score: 0

subject: [1] ""

1. What is the primary benefit of formal summary classes like PairwiseAlignmentsSingleSubjectSummary and summary.lm to end-users? These classes allow the end-user to extract the summary output for further operations.

```
> ex2 <- summary(pairwiseAlignment("zyzzyx", "syzygy"))
> nmatch(ex2) / nmismatch(ex2)
[1] 0.5
```

For the overlap pairwise sequence alignment of the strings "syzygy" and "zyzzyx" with the pairwiseAlignment default settings, perform the following operations:

```
> ex3 <- pairwiseAlignment("zyzzyx", "syzygy", type = "overlap")</pre>
```

1. Use nmatch and nmismath to extract the number of matches and mismatches respectively.

```
> nmatch(ex3)
[1] 0
> nmismatch(ex3)
```

[1] 0

```
2. Use the compareStrings function to get the symbolic representation of the alignment.> compareStrings(ex3)
```

```
[1] ""
```

3. Use the as.character function to the get the character string versions of the alignments.

```
> as.character(ex3)
```

4. Use the pattern function to extract the aligned pattern and apply the mismatch function to it to find the locations of the mismatches.

```
> mismatch(pattern(ex3))
IntegerList of length 1
[[1]] integer(0)
```

5. Use the subject function to extract the aligned subject and apply the aligned function to it to get the aligned strings.

```
> aligned(subject(ex3))

A BStringSet instance of length 1
   width seq
[1] 0
```

12.4 Exercise 4

1. Use the pairwiseAlignment function to find the Levenshtein edit distance between "syzygy" and "zyzzyx".

```
2. Use the stringDist function to find the Levenshtein edit distance for the vector c("zyzzyx", "syzygy",
    "succeed", "precede", "supersede").

> stringDist(c("zyzzyx", "syzygy", "succeed", "precede", "supersede"))

1 2 3 4
2 4
3 7 6
4 7 7 5
5 9 8 5 5
```

1. Repeat the alignment exercise above using BLOSUM62, a gap opening penalty of -12, and a gap extension penalty of -4.

2. Explore to find out what caused the alignment to change. The sift in gap penalties favored infrequent long gaps to frequent short ones.

12.6 Exercise 6

1. Rerun the simulation time using the simulateReads function with a substitutionRate of 0.005 and gapRate of 0.0005. How do the different pairwise sequence alignment methods compare? The different methods are much more comprobable when the error rates are lower.

```
> adapter <- DNAString("GATCGGAAGAGCTCGTATGCCGTCTTCTGCTTGAAA")</pre>
> set.seed(123)
> N <- 1000
> experiment <-
   list(side = rbinom(N, 1, 0.5), width = sample(0.36, N, replace = TRUE))
> table(experiment[["side"]], experiment[["width"]])
     0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21
  0 13 10 8 7 11 18 9 15 15 18 16 10 11 9 13 13 18 18 14 9 19 13
  1 15 21 21 12 17 14 8 11 12 10 14 16 7 14 19 14 14 16 14 16 16 16
    22 23 24 25 26 27 28 29 30 31 32 33 34 35 36
  0 19 19 12 6 15 18 12 15 16 17 19 6 13 18 15
  1 13 11 9 11 15 10 10 16 15 15 11 7 12 7 14
> ex6Strings <-
+ simulateReads(N, adapter, experiment, substitutionRate = 0.005, gapRate = 0.0005)
> ex6Strings <- DNAStringSet(ex6Strings)</pre>
> ex6Strings
```

```
A DNAStringSet instance of length 1000
      width seq
   [1]
         36 CTGCTTGAAATTGCACGATAGTTGCATATGCTACAA
   [2]
         36 ATTTCTCCTTCTCAGGATCGGAAGAGCTCGTATGCC
   [3]
         36 TGAAACCGGGTCACCATCATCATCTAGTAGGACCTT
   [4]
         36 GCGTTGATCGGAAGAGCTCGTATGCCGTCTTCTGCT
   [5]
         36 GCTTAGATCGGAAGAGCTCGTATGCCGTCTTCTGCT
   . . .
        . . . . . . .
 [996]
         36 TAAGATCGGAAGAGCTCGTATGCCGTCTTCTGCTTG
 [997]
         36 TTGTTTGGGAAGGCTTCAACCAAACGATCGGAAGAG
 [998]
         36 AGCTCGTATGCCGTCTTCTGCTTGAAACAGCTAATA
 [999]
         36 CGGATGGGAGCCGAGCATGACTGATCGGAAGAGCTC
[1000]
         36 CGTCTTCTGCTTGAAATCGCTATCTGAAGTCTGTTT
> ## Method 1: Use edit distance with an FDR of 1e-03
> submat1 <- nucleotideSubstitutionMatrix(match = 0, mismatch = -1, baseOnly = TRUE)
> quantile(randomScores1, seq(0.99, 1, by = 0.001))
         99.1%
                 99.2%
                         99.3%
                                99.4%
                                        99.5%
                                               99.6%
                                                       99.7%
-16.990 -16.000 -16.000 -16.000 -16.000 -16.000 -16.000 -16.000 -15.998
 99.9%
          100%
-15.000 -14.000
> ex6Aligns1 <-
+ pairwiseAlignment(ex6Strings, adapter, substitutionMatrix = submat1,
                     gapOpening = 0, gapExtension = -1)
> table(score(ex6Aligns1) > quantile(randomScores1, 0.999), experiment[["width"]])
        0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20
 FALSE 28 31 29 19 28 32 17 26 27 28 30 26 18 23 32 27 32 34 28 25 35
       0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
       21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36
 FALSE 29 32 30 21 17 30 28 22 3 0 0 0 0 0 0
        0 0 0 0 0 0 0 28 31 32 30 13 25 25 29
> ## Method 2: Use consecutive matches anywhere in string with an FDR of 1e-03 \,
> submat2 <- nucleotideSubstitutionMatrix(match = 1, mismatch = -Inf, baseOnly = TRUE)
> quantile(randomScores2, seq(0.99, 1, by = 0.001))
  99% 99.1% 99.2% 99.3% 99.4% 99.5% 99.6% 99.7% 99.8% 99.9% 100%
   7
             8
                    8
                        8
                             8
                                   8
                                         8
                                               9
                                                            11
> ex6Aligns2 <-
  pairwiseAlignment(ex6Strings, adapter, substitutionMatrix = submat2,
                     type = "local", gapOpening = 0, gapExtension = -Inf)
> table(score(ex6Aligns2) > quantile(randomScores2, 0.999), experiment[["width"]])
        0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20
 FALSE 28 31 29 19 28 32 17 26 27 28 4 1 3 0 1 1 0 0 1 0 0
  TRUE 0 0 0 0 0 0 0 0 0 0 26 25 15 23 31 26 32 34 27 25 35
       21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36
```

```
FALSE 0 0 0 0 0 0 0 0 0 0 0 0 0 0
 TRUE 29 32 30 21 17 30 28 22 31 31 32 30 13 25 25 29
> # Determine if the correct end was chosen
> table(start(pattern(ex6Aligns2)) > 37 - end(pattern(ex6Aligns2)),
       experiment[["side"]])
         0
            1
 FALSE 473 48
 TRUE
        34 445
> ## Method 3: Use consecutive matches on the ends with an FDR of 1e-03
> submat3 <- nucleotideSubstitutionMatrix(match = 1, mismatch = -Inf, baseOnly = TRUE)
> ex6Aligns3 <-
  pairwiseAlignment(ex6Strings, adapter, substitutionMatrix = submat3,
                     type = "overlap", gapOpening = 0, gapExtension = -Inf)
> table(score(ex6Aligns3) > quantile(randomScores3, 0.999), experiment[["width"]])
        0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20
 FALSE 28 31 29 19 28 32 17 0 1 2 4 1 3 1 1 2 2 3 2 2 3
       0 0 0 0 0 0 26 26 26 26 25 15 22 31 25 30 31 26 23 32
       21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36
 FALSE 5 2 3 2 1 3 5 3 3 3 2 3 2 4 4 3
 TRUE 24 30 27 19 16 27 23 19 28 28 30 27 11 21 21 26
> # Determine if the correct end was chosen
> table(end(pattern(ex6Aligns3)) == 36, experiment[["side"]])
         0 1
 FALSE 473 42
 TRUE
       34 451
> ## Method 4: Allow mismatches and indels on the ends with an FDR of 1e-03
> quantile(randomScores4, seq(0.99, 1, by = 0.001))
     99%
             99.1%
                      99.2%
                                99.3%
                                         99.4%
                                                   99.5%
                                                            99.6%
7.927024 7.927024 7.927024 7.927024 7.927024 7.927024 7.927024 7.927024
   99.7%
            99.8%
                      99.9%
                                 100%
9.908780 9.908872 11.890536 13.872293
> ex6Aligns4 <- pairwiseAlignment(ex6Strings, adapter, type = "overlap")</pre>
> table(score(ex6Aligns4) > quantile(randomScores4, 0.999), experiment[["width"]])
        0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20
 FALSE 28 31 29 19 28 32 17 0 1 2 1 0 1 0 1 0 0 0 0 0
       0 0 0 0 0 0 26 26 26 29 26 17 23 31 27 32 34 28 25 35
       21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36
 FALSE 0 0 0 0 0 0 0 0 0 0 0 0 0 0
 TRUE 29 32 30 21 17 30 28 22 31 31 32 30 13 25 25 29
> # Determine if the correct end was chosen
> table(end(pattern(ex6Aligns4)) == 36, experiment[["side"]])
```

```
0 1
FALSE 485 17
TRUE 22 476
```

2. (Advanced) Modify the simulateReads function to accept different equal length adapters on either side (left & right) of the reads. How would the methods for trimming the reads change?

```
> simulateReads <-
+ function(N, left, right = left, experiment, substitutionRate = 0.01, gapRate = 0.001) {
    leftChars <- strsplit(as.character(left), "")[[1]]</pre>
    rightChars <- strsplit(as.character(right), "")[[1]]
    if (length(leftChars) != length(rightChars))
      stop("left and right adapters must have the same number of characters")
    nChars <- length(leftChars)</pre>
    sapply(seq_len(N), function(i) {
      width <- experiment[["width"]][i]</pre>
      side <- experiment[["side"]][i]</pre>
      randomLetters <-
        function(n) sample(DNA_ALPHABET[1:4], n, replace = TRUE)
+
      randomLettersWithEmpty <-
        function(n)
        sample(c("", DNA_ALPHABET[1:4]), n, replace = TRUE,
+
               prob = c(1 - gapRate, rep(gapRate/4, 4)))
      if (side) {
+
        value <-
          paste(ifelse(rbinom(nChars,1,substitutionRate), randomLetters(nChars), rightChars),
                randomLettersWithEmpty(nChars),
                sep = "", collapse = "")
        value <-
          paste(c(randomLetters(36 - width), substring(value, 1, width)),
+
                  sep = "", collapse = "")
      } else {
        value <-
          paste(ifelse(rbinom(nChars,1,substitutionRate), randomLetters(nChars), leftChars),
                randomLettersWithEmpty(nChars),
                sep = "", collapse = "")
+
        value <-
          paste(c(substring(value, 37 - width, 36), randomLetters(36 - width)),
                  sep = "", collapse = "")
      }
      value
    })
+ }
> leftAdapter <- adapter
> rightAdapter <- reverseComplement(adapter)
> ex6LeftRightStrings <- simulateReads(N, leftAdapter, rightAdapter, experiment)
> ex6LeftAligns4 <-
   pairwiseAlignment(ex6LeftRightStrings, leftAdapter, type = "overlap")
> ex6RightAligns4 <-
    pairwiseAlignment(ex6LeftRightStrings, rightAdapter, type = "overlap")
> scoreCutoff <- quantile(randomScores4, 0.999)</pre>
> leftAligned <-
```

```
+ start(pattern(ex6LeftAligns4)) == 1 & score(ex6LeftAligns4) > pmax(scoreCutoff, score(ex6RightAligns4))
    > rightAligned <-
        end(pattern(ex6RightAligns4)) == 36 & score(ex6RightAligns4) > pmax(scoreCutoff, score(ex6LeftAl
    > table(leftAligned, rightAligned)
               rightAligned
    leftAligned FALSE TRUE
          FALSE
                  195 382
          TRUE
                  423
                         0
    > table(leftAligned | rightAligned, experiment[["width"]])
             0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20
      FALSE 28 31 29 19 28 32 17 3 3 2 0 0 1 1 0 0 0 0 0 1
             0 0 0 0 0 0 23 24 26 30 26 17 22 32 27 32 34 28 25 34
            21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36
      FALSE 0 0 0 0 0 0 0 0 0 0 0 0 0 0
      TRUE 29 32 30 21 17 30 28 22 31 31 32 30 13 25 25 29
12.7
      Exercise 7
  1. Rerun the global-local alignment of the short reads against the entire genome. (This may take a few
    minutes.)
    > genBankFullAlign <-
        pairwiseAlignment(srPhiX174, genBankPhage,
                          patternQuality = SolexaQuality(quPhiX174),
                          subjectQuality = SolexaQuality(99L),
                          type = "global-local")
    > summary(genBankFullAlign, weight = wtPhiX174)
    Global-Local Single Subject Pairwise Alignments
    Number of Alignments: 53802
    Scores:
       Min. 1st Qu. Median
                              Mean 3rd Qu.
                                              Max.
     -45.08 56.72
                    59.89
                             60.59
                                     69.56
                                             69.85
    Number of matches:
       Min. 1st Qu. Median
                              Mean 3rd Qu.
                                              Max.
      24.00
              33.00
                    34.00
                             34.01 35.00
                                             35.00
    Top 10 Mismatch Counts:
       SubjectPosition Subject Pattern Count Probability
```

T 22965 0.999912919

T 22845 0.999693681

T 1985 0.106800818

T 287 0.018648473

605 0.033570081

489 0.023314580

325 0.013882363

2811

2793

2834

2835

2829

2782

2839

2

3

4

5

6

7

C

С

G

G

G

G

Т

T

T

```
8 2807 A C 169 0.007657801
9 2827 A T 168 0.007714207
10 2837 C T 159 0.009612478
```

2. Plot the coverage of these alignments and use the slice function to find the ranges of alignment. Are there any alignments outside of the substring region that was used above? Yes, there are some alignments outside of the specified substring region.

```
> genBankFullCoverage <- coverage(genBankFullAlign, weight = wtPhiX174)
> plot(as.integer(genBankFullCoverage), xlab = "Position", ylab = "Coverage", type = "1")
> slice(genBankFullCoverage, lower = 1)
Views on a 5386-length Rle subject
views:
  start end width
[1] 1195 1230
             2514 2548
[3]
   2745 2859
            115 [ 416
                     946 1536 2135 2797 3374 4011 ...]
[4]
   3209 3247
             39 [
                 32
                     54 440 1069 1130 1130 1130 1130 ...]
             [5]
   3964 3998
```

3. Use the reverseComplement function on the bacteriophage ϕ X174 genome. Do any short reads have a higher alignment score on this new sequence than on the original sequence? Yes, there are some strings with a higher score on the new sequence.

```
> genBankFullAlignRevComp <-
+ pairwiseAlignment(srPhiX174, reverseComplement(genBankPhage),
+ patternQuality = SolexaQuality(quPhiX174),
+ subjectQuality = SolexaQuality(99L),
+ type = "global-local")
> table(score(genBankFullAlignRevComp) > score(genBankFullAlign))

FALSE TRUE
1112 1
```

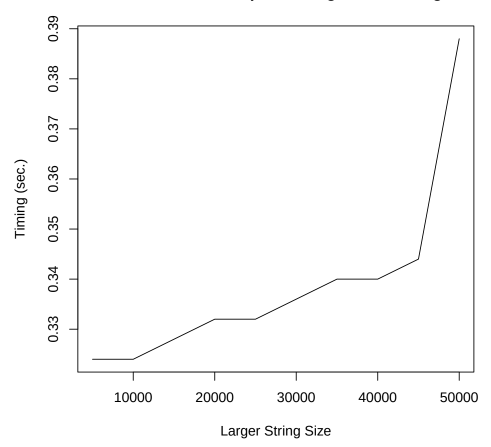
12.8 Exercise 8

1. Rerun the first set of profiling code, but this time fix the number of characters in **string1** to 35 and have the number of characters in **string2** range from 5000, 50000, by increments of 5000. What is the computational order of this simulation exercise? As expected, the growth in time is now linear.

```
> N <- as.integer(seq(5000, 50000, by = 5000))
> newTimings <- rep(0, length(N))
> names(newTimings) <- as.character(N)
> for (i in seq_len(length(N))) {
+ string1 <- DNAString(paste(sample(DNA_ALPHABET[1:4], 35, replace = TRUE), collapse = ""))
+ string2 <- DNAString(paste(sample(DNA_ALPHABET[1:4], N[i], replace = TRUE), collapse = ""))
+ newTimings[i] <- system.time(pairwiseAlignment(string1, string2, type = "global"))[["user.self"]
+ }
> newTimings
5000 10000 15000 20000 25000 30000 35000 40000 45000 50000
```

0.324 0.324 0.328 0.332 0.332 0.336 0.340 0.340 0.344 0.388

Global Pairwise Sequence Alignment Timings



2. Rerun the second set of profiling code using the simulations from the previous exercise with *scoreOnly* argument set to TRUE. Is is still twice as fast? Yes, it is still over twice as fast.

```
> newScoreOnlyTimings <- rep(0, length(N))
> names(newScoreOnlyTimings) <- as.character(N)
> for (i in seq_len(length(N))) {
+    string1 <- DNAString(paste(sample(DNA_ALPHABET[1:4], 35, replace = TRUE), collapse = ""))
+    string2 <- DNAString(paste(sample(DNA_ALPHABET[1:4], N[i], replace = TRUE), collapse = ""))
+    newScoreOnlyTimings[i] <- system.time(pairwiseAlignment(string1, string2, type = "global", score</pre>
```

```
+ }
> newScoreOnlyTimings

5000 10000 15000 20000 25000 30000 35000 40000 45000 50000
0.356 0.332 0.328 0.329 0.332 0.332 0.336 0.336 0.340 0.348
> round((newTimings - newScoreOnlyTimings) / newTimings, 2)

5000 10000 15000 20000 25000 30000 35000 40000 45000 50000
-0.10 -0.02 0.00 0.01 0.00 0.01 0.01 0.01 0.01
```

13 Session Information

All of the output in this vignette was produced under the following conditions:

```
> toLatex(sessionInfo())
```

```
\begin{itemize}\raggedright
  \item R version 3.0.2 (2013-09-25), \verb|x86_64-unknown-linux-gnu|
  \item Locale: \verb|LC_CTYPE=en_US.UTF-8|, \verb|LC_NUMERIC=C|, \verb|LC_TIME=en_US.UTF-8|, \verb|LC_CCO
  \item Base packages: base, datasets, grDevices, graphics,
    methods, parallel, stats, utils
  \item Other packages: BiocGenerics~0.8.0, Biostrings~2.30.0,
    IRanges~1.20.0, XVector~0.2.0
  \item Loaded via a namespace (and not attached):
    stats4~3.0.2, tools~3.0.2
  \end{itemize}
```

References

- [1] Durbin, R., Eddy, S., Krogh, A., and Mitchison G. *Biological Sequence Analysis*. Cambridge UP 1998, sec 2.3.
- [2] Haubold, B. and Wiehe, T. Introduction to Computational Biology. Birkhauser Verlag 2006, Chapter 2.
- [3] Malde, K. The effect of sequence quality on sequence alignment. Bioinformatics, 24(7):897-900, 2008.
- [4] Needleman, S. and Wunsch, C. A general method applicable to the search for similarities in the amino acid sequence of two proteins. *Journal of Molecular Biology*, 48, 443-453, 1970.
- [5] Smith, H.; Hutchison, C.; Pfannkoch, C.; and Venter, C. Generating a synthetic genome by whole genome assembly: {phi}X174 bacteriophage from synthetic oligonucleotides. *Proceedings of the National Academy of Sciences*, 100(26): 15440-15445, 2003.
- [6] Smith, T.F. and Waterman, M.S. Identification of common molecular subsequences. *Journal of Molecular Biology*, 147, 195-197, 1981.