---

output: html\_document

editor\_options:

chunk\_output\_type: console

---

# DNA descriptive statics - Part 1

\*\*By\*\*: Avril Coghlan

\*\*Adapted, edited and expanded\*\*: Nathan Brouwer (brouwern@gmail.com) under the Creative Commons 3.0 Attribution License [(CC BY 3.0)](https://creativecommons.org/licenses/by/3.0/).

## Preface

This is a modification of ["DNA Sequence Statistics (1)"](https://a-little-book-of-r-for-bioinformatics.readthedocs.io/en/latest/src/chapter1.html) from Avril Coghlan's [\*A little book of R for bioinformatics.\*](https://a-little-book-of-r-for-bioinformatics.readthedocs.io/en/latest/index.html). The text and code were originally written by Dr. Coghlan and distributed under the [Creative Commons 3.0](https://creativecommons.org/licenses/by/3.0/us/) license.

## Writing TODO:

\* Add biology introduction

\* Work on flow

\* organize intial sections (intro, vocab, preliminaries)

## Introduction

<!-- Check the current status of the biology of this section -->

<!-- ### Note on the biology in this section -->

<!-- Some of the biology in this tutorial appears to be out of date. Specifically, research on the biological basis of unusual frequencies of DNA "words" does not appear to be much current interest in bioinformatics and genomics. (In contrast, methylation of bases within certain DNA words is being worked on.) The examples are still good for practicing R skills. -->

## Vocabulary

\* GC content

\* DNA words

\* scatterplots, histograms, piecharts, and boxplots

## Functions

\* `seqinr::GC()`

\* `seqinr::count()`

## Preliminaries

```{r}

library(compbio4all)

library(seqinr)

```

## Converting DNA from FASTA format

In a previous exercise we downloaded and examined DNA sequence in the FASTA format. The sequence we worked with is also stored as a data file within the `compbio4all` pa package and can be brought into memory using the `data()` command.

```{r}

data("dengueseq\_fasta")

```

We can look at this data object with the `str()` command

```{r}

str(dengueseq\_fasta)

```

This isn't in a format we can work with directly so we'll use the function fasta\_cleaner() to set it up.

```{r}

header. <- ">NC\_001477.1 Dengue virus 1, complete genome"

dengueseq\_vector <- compbio4all::fasta\_cleaner(dengueseq\_fasta)

```

Now check it out.

```{r}

str(dengueseq\_vector)

```

What we have here is each base of the sequence in a seperate slot of our vector.

The first four bases are "AGTT"

We can see the first one like this

```{r}

dengueseq\_vector[1]

```

The second one like this

```{r}

dengueseq\_vector[2]

```

The first and second like this

```{r}

dengueseq\_vector[1:2]

```

and all four like this

```{r}

dengueseq\_vector[1:4]

```

## Length of a DNA sequence

Once you have retrieved a DNA sequence, we can obtain some simple statistics to describe that sequence, such as the sequence’s total length in nucleotides. In the above example, we retrieved the DEN-1 Dengue virus genome sequence, and stored it in the vector variable dengueseq\_vector To obtain the length of the genome sequence, we would use the length() function, typing:

```{r}

length(dengueseq\_vector)

```

The length() function will give you back the length of the sequence stored in variable dengueseq\_vector, in nucleotides. The length() function actually gives the number of \*\*elements\*\* (slots) in the input vector that you passed to it, which in this case in the number of elements in the vector dengueseq\_vector. Since each element of the vector dengueseq\_vector contains one nucleotide of the DEN-1 Dengue virus sequence, the result for the DEN-1 Dengue virus genome tells us the length of its genome sequence (ie. 10735 nucleotides long).

### Base composition of a DNA sequence

An obvious first analysis of any DNA sequence is to count the number of occurrences of the four different nucleotides (“A”, “C”, “G”, and “T”) in the sequence. This can be done using the the table() function. For example, to find the number of As, Cs, Gs, and Ts in the DEN-1 Dengue virus sequence (which you have put into vector variable dengueseq\_vector, using the commands above), you would type:

```{r}

table(dengueseq\_vector)

```

This means that the DEN-1 Dengue virus genome sequence has 3426 As occurring throughout the genome, 2240 Cs, and so forth.

### GC Content of DNA

One of the most fundamental properties of a genome sequence is its \*\*GC content\*\*, the fraction of the sequence that consists of Gs and Cs, ie. the %(G+C).

The GC content can be calculated as the percentage of the bases in the genome that are Gs or Cs. That is, GC content = (number of Gs + number of Cs)\*100/(genome length). For example, if the genome is 100 bp, and 20 bases are Gs and 21 bases are Cs, then the GC content is (20 + 21)\*100/100 = 41%.

You can easily calculate the GC content based on the number of As, Gs, Cs, and Ts in the genome sequence. For example, for the DEN-1 Dengue virus genome sequence, we know from using the table() function above that the genome contains 3426 As, 2240 Cs, 2770 Gs and 2299 Ts. Therefore, we can calculate the GC content using the command:

```{r}

(2240+2770)\*100/(3426+2240+2770+2299)

```

Alternatively, if you are feeling lazy, you can use the GC() function in the SeqinR package, which gives the fraction of bases in the sequence that are Gs or Cs.

```{r}

seqinr::GC(dengueseq\_vector)

```

The result above means that the fraction of bases in the DEN-1 Dengue virus genome that are Gs or Cs is 0.4666977. To convert the fraction to a percentage, we have to multiply by 100, so the GC content as a percentage is 46.66977%.

### DNA words

As well as the frequency of each of the individual nucleotides (“A”, “G”, “T”, “C”) in a DNA sequence, it is also interesting to know the frequency of longer \*\*DNA words\*\*, also referred to as \*\*genomic words\*\*. The individual nucleotides are DNA words that are 1 nucleotide long, but we may also want to find out the frequency of DNA words that are 2 nucleotides long (ie. “AA”, “AG”, “AC”, “AT”, “CA”, “CG”, “CC”, “CT”, “GA”, “GG”, “GC”, “GT”, “TA”, “TG”, “TC”, and “TT”), 3 nucleotides long (eg. “AAA”, “AAT”, “ACG”, etc.), 4 nucleotides long, etc.

To find the number of occurrences of DNA words of a particular length, we can use the count() function from the R SeqinR package.

The count() function only works with lower-case letters, so first we have to use the tolower() function to convert our upper class genome to lower case

```{r}

dengueseq\_vector <-tolower(dengueseq\_vector)

```

Now we can look for words. For example, to find the number of occurrences of DNA words that are 1 nucleotide long in the sequence dengueseq\_vector, we type:

```{r}

seqinr::count(dengueseq\_vector, 1)

```

As expected, this gives us the number of occurrences of the individual nucleotides. To find the number of occurrences of DNA words that are 2 nucleotides long, we type:

```{r}

seqinr::count(dengueseq\_vector, 2)

```

Note that by default the count() function includes all overlapping DNA words in a sequence. Therefore, for example, the sequence “ATG” is considered to contain two words that are two nucleotides long: “AT” and “TG”.

If you type help(‘count’), you will see that the result (output) of the function count() is a table object. This means that you can use double square brackets to extract the values of elements from the table. For example, to extract the value of the third element (the number of Gs in the DEN-1 Dengue virus sequence), you can type:

```{r}

denguetable\_2 <- seqinr::count(dengueseq\_vector,2)

denguetable\_2[[3]]

```

The command above extracts the third element of the table produced by count(dengueseq\_vector,1), which we have stored in the table variable denguetable.

Alternatively, you can find the value of the element of the table in column “g” by typing:

```{r}

denguetable\_2[["aa"]]

```

Once you have table you can make a basic plot

```{r}

barplot(denguetable\_2)

```

We can sort by the number of words using the sort() command

```{r}

sort(denguetable\_2)

```

Let's save over the original object

```{r}

denguetable\_2 <- sort(denguetable\_2)

```

```{r}

barplot(denguetable\_2)

```

R will automatically try to optimize the appearance of the labels on the graph so you may not see all of them; no worries.

R can also make pie charts. Piecharts only really work when there are a few items being plots, like the four bases.

```{r}

denguetable\_1 <- seqinr::count(dengueseq\_vector,1)

```

Make a piechart with pie()

```{r}

pie(denguetable\_1)

```

### Summary

In this practical, have learned to use the following R functions:

length() for finding the length of a vector or list

table() for printing out a table of the number of occurrences of each type of item in a vector or list.

These functions belong to the standard installation of R.

You have also learnt the following R functions that belong to the SeqinR package:

GC() for calculating the GC content for a DNA sequence

count() for calculating the number of occurrences of DNA words of a particular length in a DNA sequence

## Acknowledgements

This is a modification of ["DNA Sequence Statistics (1)"](https://a-little-book-of-r-for-bioinformatics.readthedocs.io/en/latest/src/chapter1.html) from Avril Coghlan's [\*A little book of R for bioinformatics.\*](https://a-little-book-of-r-for-bioinformatics.readthedocs.io/en/latest/index.html). Almost all of text and code was originally written by Dr. Coghlan and distributed under the [Creative Commons 3.0](https://creativecommons.org/licenses/by/3.0/us/) license.

In "A little book..." Coghlan noted: "Many of the ideas for the examples and exercises for this chapter were inspired by the Matlab case studies on Haemophilus influenzae (www.computational-genomics.net/case\_studies/haemophilus\_demo.html) and Bacteriophage lambda (http://www.computational-genomics.net/case\_studies/lambdaphage\_demo.html) from the website that accompanies the book Introduction to Computational Genomics: a case studies approach by Cristianini and Hahn (Cambridge University Press; www.computational-genomics.net/book/)."

### License

The content in this book is licensed under a Creative Commons Attribution 3.0 License.

https://creativecommons.org/licenses/by/3.0/us/

### Exercises

Answer the following questions, using the R package. For each question, please record your answer, and what you typed into R to get this answer.

Model answers to the exercises are given in Answers to the exercises on DNA Sequence Statistics (1).

1. What are the last twenty nucleotides of the Dengue virus genome sequence?

1. What is the length in nucleotides of the genome sequence for the bacterium Mycobacterium leprae strain TN (accession NC\_002677)?

Note: Mycobacterium leprae is a bacterium that is responsible for causing leprosy, which is classified by the WHO as a neglected tropical disease. As the genome sequence is a DNA sequence, if you are retrieving its sequence via the NCBI website, you will need to look for it in the NCBI Nucleotide database.

1. How many of each of the four nucleotides A, C, T and G, and any other symbols, are there in the Mycobacterium leprae TN genome sequence?

Note: other symbols apart from the four nucleotides A/C/T/G may appear in a sequence. They correspond to positions in the sequence that are are not clearly one base or another and they are due, for example, to sequencing uncertainties. or example, the symbol ‘N’ means ‘aNy base’, while ‘R’ means ‘A or G’ (puRine). There is a table of symbols at www.bioinformatics.org/sms/iupac.html.

1. What is the GC content of the \*Mycobacterium leprae TN\* genome sequence, when (i) all non-A/C/T/G nucleotides are included, (ii) non-A/C/T/G nucleotides are discarded?

Hint: look at the help page for the GC() function to find out how it deals with non-A/C/T/G nucleotides.

1. How many of each of the four nucleotides A, C, T and G are there in the complement of the Mycobacterium leprae TN genome sequence? \*Hint\*: you will first need to search for a function to calculate the complement of a sequence. Once you have found out what function to use, remember to use the help() function to find out what are the arguments (inputs) and results (outputs) of that function. How does the function deal with symbols other than the four nucleotides A, C, T and G? Are the numbers of As, Cs, Ts, and Gs in the complementary sequence what you would expect?

1. How many occurrences of the DNA words CC, CG and GC occur in the Mycobacterium leprae TN genome sequence?

1. How many occurrences of the DNA words CC, CG and GC occur in the (i) first 1000 and (ii) last 1000 nucleotides of the Mycobacterium leprae TN genome sequence?

1.How can you check that the subsequence that you have looked at is 1000 nucleotides long?

```{r}

```