### **Analysis**

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#### load the packages

```
##
## 载入程辑包: 'dplyr'

## The following objects are masked from 'package:stats':
##
## filter, lag

## The following objects are masked from 'package:base':
##
## intersect, setdiff, setequal, union
```

#### Import the Sequences.csv file

SeqData<-read.csv("output/Sequences.csv")#loading the data str(SeqData) #check the structure of the Data

```
class(SeqData)

## [1] "data.frame"

dim(SeqData)
```

```
## [1] 3 2
```

## Count the number of each base pair (A, T, C and G), in each of the three sequences.

```
# extract the certain sequence from table
seq1 <- strsplit(SeqData$Sequence,"")[[1]]</pre>
seq2 <- strsplit(SeqData$Sequence,"")[[2]]</pre>
seq3 <- strsplit(SeqData$Sequence, "")[[3]]</pre>
# Each base pair content in sequence 1
A1_number <- length(grep("A",seq1))
T1 number <- length(grep("T", seq1))</pre>
C1 number <- length(grep("C", seq1))</pre>
G1 number <- length(grep("G", seq1))</pre>
# Each base pair content in sequence 2
A2 number <- length(grep("A",seq2))
T2 number <- length(grep("T", seq2))
C2 number <- length(grep("C",seq2))</pre>
G2 number <- length(grep("G", seq2))</pre>
# Each base pair content in sequence 3
A3 number <- length(grep("A", seq3))
T3 number <- length(grep("T", seg3))
C3 number <- length(grep("C", seq3))
G3 number <- length(grep("G", seg3))
```

#### Print out each sequence

```
print (unlist(SeqData$Sequence))
```

## [2] "AGCATGCAAGTCAAACGGGATGTAGCAATACATTCAGTGGCGAACGGGTGAGTAACGCGTGGATGATCTACCTATGA
GATGGGGATAACTATTAGAAATAGTAGCTAATACCGAATAAGGTCAGTTAATTTGTTAATTGATGAAAGGAAGCCTTTAAAGCTT
CGCTTGTAGATGAGTCTGCGTCTTATTAGCTAGTTGGTAGGGTAAATGCCTACCAAGGCAATGATAAGTAACCGGCCTGAGAGGG
TGAACGGTCACACTGGAACTGAGATACGGTCCAGACTCCTACGGGAGGCAGCAGCTAAGAATCTTCCGCAATGGGCGAAAGCCTG
ACGGAGCGACACTGCGTGAATGAAGAAGGTCGAAAGATTGTAAAATTCTTTTATAAATGAGGAATAAGCTTTGTAGGAAATGACA
AAGTGATGACGTTAATTTATGAATAAGCCCCGGCTAATTACGTGCCAGCAGCAGCAGCGGTAATACG"

## Create a table with number of each nucleotide for each of the three sequences

```
Sequences_Id <- c("HQ433692.1","HQ433694.1","HQ433691.1")
A_content <- c(A1_number, A2_number, A3_number)
T_content <- c(T1_number, T2_number, T3_number)
C_content <- c(C1_number, C2_number, C3_number)
G_content <- c(G1_number, G2_number, G3_number)
SumTable<-data.frame(Sequences_Id, A_content, T_content, C_content, G_content, Total = nchar(SeqData$Sequence))
print(SumTable)</pre>
```

```
Sequences Id A content T content C content G content Total
## 1
       HQ433692.1
                        154
                                   114
                                              82
## 2
       HQ433694.1
                        155
                                   114
                                              81
                                                        131
                                                              481
## 3
      HO433691.1
                        154
                                   115
                                                        131
                                                              481
                                              Я1
```

# Upload Image of a bacteria from the internet, and a link to the Wikipedia page about Borrelia burgdorferi



Lyme Disease Bacteria: Borrelia burgdorferi, Image courtesy of Emily M. Eng

link to the Wikipedia page about Borrelia burgdorferi (https://en.wikipedia.org/wiki/Borrelia\_burgdorferi)

## Create a final table showing GC content for each sequence ID

```
# Calculate GC Content
FinalTable <- transmute(SumTable, Sequences_Id, GC_Content = paste(round((C_content
+G_content)/Total *100, 2), "%"))
print (FinalTable)</pre>
```

```
## Sequences_Id GC_Content

## 1 HQ433692.1 44.28 %

## 2 HQ433694.1 44.07 %

## 3 HQ433691.1 44.07 %
```