Bioinformatics - Computer Lab 2

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Question 1

At first, the dataset of the RAG1 gene sequences from 33 lizard species were downloaded from GenBank and saved in a fasta file using the provided R script 732A51 BioinformaticsHT2018 Lab02 GenBankGetCode.R.. The code can be found in Appendix 1 (Data Import of original dataset).

Question 1.1

The saved fasta-file has to be read in R so that we can work with that. The R code for the reading process can be found in Appendix 1.1 (Reading original data).

After that, the artificial dataset is built by considering that it contains 33 sequences (each length of the sequences is the same as in the lizard dataset) so that for each real sequence an artificial one is created. As mentioned, the simulation of the artificial sequences is based on the distribution given by the base composition of the original dataset.

The artificial dataset is submitted as the fasta file artificial_dataset_1_1.fasta. The written function for all these processes automatically prints the base composition in the simulated data compared to the base composition in the original data. An extract from the output can be seen here:

```
get_artificial_sequence_dataset(lizards_sequences)
```

```
## [1] "comparison of base compositions between original and artificial datasets (values rounded):"
     name_original name_artificial a_original a_artificial c_original
## 1 "JF806202"
                    "1"
                                      "0.29"
                                                 "0.32"
                                                               "0.2"
                    "2"
## 2 "HM161150"
                                     "0.31"
                                                 "0.31"
                                                               "0.21"
## 3 "FJ356743"
                    "3"
                                     "0.31"
                                                 "0.31"
                                                               "0.21"
                                                 "0.29"
## 4 "JF806205"
                    "4"
                                     "0.28"
                                                               "0.21"
                    "5"
                                                 "0.3"
                                                               "0.2"
## 5 "JQ073190"
                                     "0.31"
##
     c artificial
                   g_original g_artificial t_original t_artificial
                                             "0.26"
## 1 "0.2"
                               "0.23"
                                                         "0.26"
                   "0.24"
## 2 "0.21"
                   "0.23"
                               "0.23"
                                             "0.24"
## 3 "0.22"
                   "0.23"
                               "0.23"
                                             "0.24"
                                                         "0.23"
## 4 "0.21"
                   "0.24"
                               "0.23"
                                             "0.26"
                                                         "0.27"
## 5 "0.22"
                                                         "0.26"
                   "0.24"
                               "0.23"
                                             "0.26"
```

It becomes clear that the base compositions are very similar. The entire code for the function can be seen in Appendix 1.1 (Function code).

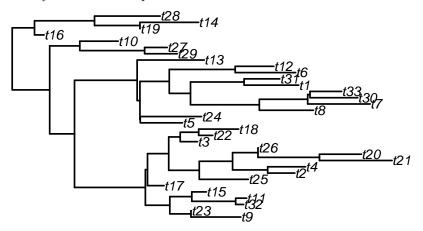
Question 1.2

In this part of the exercise do we use the prepared data from part 1, in Appendix 1 code can be found in (Data Import of original dataset).

We used the function rtree to create a tree object of the type phylo and the length of the original sequences.

```
tree <- rtree(n = length(lizards_sequences))</pre>
```

Here you can find the plot of the tree.



After the simulation of the phylogenetic tree, we had to simulate the sequence.

For this, we the had several things to do. 1. We simulated a transition rate matrix (Q-Matrix). In this case we choose one by yourself.

2. We had to choose the lengths of the sequences. We chose the average length of the original sequences and used this length for every artificial sequence.

```
# calculating average length of original sequences
avg_length = c()
for (seq in 1:33) {
   avg_length = c(avg_length, length(lizards_sequences[[seq]]))
}
avg_length = mean(avg_length)
```

Now we can simulate the sequences by using the function phangorn::simSeq().

```
sequences_artificial <- simSeq(tree, 1 = avg_length, Q=transition_matrix , type = "DNA")</pre>
```

Since in sequences are filled with integers from 1 to 4, do we have to replace the numbers by the letters a,b,c,d.

- 1 = a
- 2 = b
- 3 = c
- 4 = d

The code for this can be found in Appendix 1.2

The second simulate a artificial DNA sequence dataset do we save as "artificial_dataset_1_2.fasta".

```
ape::write.dna(sequences_artificial, file ="artificial_dataset_1_2.fasta", format = "fasta", colsep = "
```

Question 2

Question 2.1

```
lizards_sequences = read.fasta("lizard_seqs.fasta")
original_dataset <- lizards_sequences</pre>
artificial_sequences_1 <- read.fasta("artificial_dataset_1_1.fasta")</pre>
artificial_sequences_2 <- read.fasta("artificial_dataset_1_2.fasta")</pre>
original_base_compositions <- list()</pre>
artificial_1_base_compositions <- list()</pre>
artificial_2_base_compositions <- list()</pre>
for (i in 1:length(original_dataset)) {
    # getting base compositions for each original sequence
    original_base_compositions[[i]] =
      seqinr::count(original_dataset[[i]],1)
}
for (i in 1:length(artificial_sequences_1)) {
    # getting base compositions for each original sequence
    artificial_1_base_compositions[[i]] =
      seqinr::count(artificial_sequences_1[[i]],1)
}
for (i in 1:length(artificial_sequences_2)) {
    # getting base compositions for each original sequence
    artificial_2_base_compositions[[i]] =
      seqinr::count(artificial_sequences_2[[i]],1)
}
Reduce('+', original_base_compositions)
##
##
            С
                   g
## 20414 13422 15089 16474
sum(Reduce('+', original_base_compositions))
## [1] 65399
Reduce('+', original_base_compositions)/sum(Reduce('+', original_base_compositions))
##
##
                     С
## 0.3121454 0.2052325 0.2307222 0.2518999
Reduce('+', artificial_1_base_compositions)
##
##
       a
           С
                   g
## 20554 13530 15101 16250
sum(Reduce('+', artificial_1_base_compositions))
## [1] 65435
```

```
Reduce('+', artificial_1_base_compositions)/sum(Reduce('+', artificial_1_base_compositions))
##
##
                                  g
## 0.3141132 0.2067701 0.2307786 0.2483380
Reduce('+', artificial_2_base_compositions)
##
##
       a
              С
                     g
                           t
## 16344 16122 16588 16352
sum(Reduce('+', artificial_2_base_compositions))
## [1] 65406
Reduce('+', artificial_2_base_compositions)/sum(Reduce('+', artificial_2_base_compositions))
##
##
            a
                       С
                                  g
## 0.2498853 0.2464911 0.2536159 0.2500076
The original dataset and the first artificially created dataset are rather similar in their distributions for A, C,
T and G's. However, the second artificially created dataset has a slightly different distribution. This final
dataset has almost uniform distribution for A, C, T and G's, they all occur with an average frequency of
approximately 25%.
library(rDNAse)
original compositions <- list()</pre>
  for (i in 1:length(lizards_sequences)) {
  string1 <- paste(lizards_sequences[[i]], collapse = "")</pre>
  string1 <- toupper(string1)</pre>
  original_compositions[[i]] <- kmer(string1)</pre>
}
artificial_compositions_1 <- list()</pre>
  for (i in 1:length(artificial_sequences_1)) {
  string1 <- paste(artificial_sequences_1[[i]], collapse = "")</pre>
  string1 <- toupper(string1)</pre>
  artificial_compositions_1[[i]] <- kmer(string1)</pre>
}
artificial compositions 2 <- list()</pre>
  for (i in 1:length(artificial_sequences_2)) {
  string1 <- paste(artificial_sequences_2[[i]], collapse = "")</pre>
  string1 <- toupper(string1)</pre>
  artificial_compositions_2[[i]] <- kmer(string1)</pre>
}
Reduce('+', original_compositions)
        TG
                                                                                 TC
##
                  GA
                                             ΑT
                                                     CA
                                                              TT
                                                                       CT
                           AA
                                   AG
      6960
               4554
                                                   3909
                                                                     4296
                                                                              4947
##
                        5664
                                 4867
                                          5096
                                                            2277
                  GC
##
         CC
                           GT
                                   AC
                                            GG (Other)
                                                            NA's
##
      3207
               3076
                        3172
                                 2727
                                          3237
                                                   4728
                                                            2685
Reduce('+', artificial_compositions_1)
                                 CC
                                       CG
                                            CT
                                                  GA
                                                        GC
                                                             GG
                                                                                    TG
##
     AA
           AC
                AG
                      ΑT
                            CA
                                                                   GT
                                                                        TA
                                                                              TC
```

```
## 6410 4264 4821 5048 4284 2790 3087 3364 4747 3062 3495 3788 5102 3407 3690
##
    TT
## 4043
Reduce('+', artificial_compositions_2)
         AC
                AG
                     ΑT
                           CA
                                 CC CG
                                           CT
                                                 GA
                                                       GC
                                                            GG
                                                                  GT
                                                                       TA
                                                                                   TG
## 4082 4016 4206 4035 3946 4013 4162 3990 4147 4052 4210 4173 4165 4031 4002
     TT
##
## 4143
GC content is the largest for the second artificially created dataset. CG content is largest for the second
artificially created dataset. AT content is largest in the original dataset.
# Protein sequences
protein_original <- read.fasta("lizard_protein.fasta")</pre>
protein_artificial_1 <- read.fasta("artificial_1_protein.fasta")</pre>
protein_artificial_2 <- read.fasta("artificial_2_protein.fasta")</pre>
library(protr)
original_aac <- list()</pre>
for (i in 1:length(protein_original)) {
string1 <- paste(protein_original[[i]], collapse = "")</pre>
string1 <- toupper(string1)</pre>
string1 <- gsub(pattern = "[*]", replacement = "", x = string1)</pre>
string1 <- gsub(pattern = "B", replacement = "", x = string1)</pre>
string1 <- gsub(pattern = "J", replacement = "", x = string1)</pre>
string1 <- gsub(pattern = "0", replacement = "", x = string1)</pre>
string1 <- gsub(pattern = "U", replacement = "", x = string1)</pre>
string1 <- gsub(pattern = "X", replacement = "", x = string1)</pre>
string1 <- gsub(pattern = "Z", replacement = "", x = string1)</pre>
original_aac[[i]] <- extractAAC(string1)</pre>
}
artificial 1 aac <- list()
for (i in 1:length(protein_artificial_1)) {
string1 <- paste(protein_artificial_1[[i]], collapse = "")</pre>
string1 <- toupper(string1)</pre>
string1 <- gsub(pattern = "[*]", replacement = "", x = string1)</pre>
string1 <- gsub(pattern = "B", replacement = "", x = string1)</pre>
string1 <- gsub(pattern = "J", replacement = "", x = string1)</pre>
string1 <- gsub(pattern = "0", replacement = "", x = string1)</pre>
string1 <- gsub(pattern = "U", replacement = "", x = string1)</pre>
string1 <- gsub(pattern = "X", replacement = "", x = string1)</pre>
string1 <- gsub(pattern = "Z", replacement = "", x = string1)</pre>
artificial_1_aac[[i]] <- extractAAC(string1)</pre>
}
artificial 2 aac <- list()</pre>
for (i in 1:length(protein_artificial_2)) {
string1 <- paste(protein_artificial_2[[i]], collapse = "")</pre>
string1 <- toupper(string1)</pre>
string1 <- gsub(pattern = "[*]", replacement = "", x = string1)</pre>
string1 <- gsub(pattern = "B", replacement = "", x = string1)</pre>
string1 <- gsub(pattern = "J", replacement = "", x = string1)</pre>
string1 <- gsub(pattern = "0", replacement = "", x = string1)</pre>
string1 <- gsub(pattern = "U", replacement = "", x = string1)</pre>
```

```
string1 <- gsub(pattern = "X", replacement = "", x = string1)</pre>
string1 <- gsub(pattern = "Z", replacement = "", x = string1)</pre>
artificial_2_aac[[i]] <- extractAAC(string1)</pre>
Reduce('+', original_aac)/length(original_aac)
                        R
                                                          C
                                                                      Ε
            Α
## 0.04504567 0.06892454 0.03459927 0.03899492 0.04615282 0.06251367
##
                        G
                                   Η
                                               Ι
                                                          L
## 0.04665052 0.05212339 0.03881790 0.03970175 0.09512420 0.06888196
            М
                       F
                                   Ρ
                                               S
                                                          Т
## 0.02177691 0.04072228 0.06041408 0.09363222 0.05396491 0.02155234
            Y
##
## 0.02414733 0.04625931
Reduce('+', artificial_1_aac)/length(artificial_1_aac)
##
                        R
                                   N
                                               D
                                                          C
## 0.05057534 0.08796184 0.04476749 0.03495967 0.03066037 0.04049515
                        G
                                   Η
                                               Ι
                                                          L
## 0.04055862 0.05711907 0.02888776 0.06421406 0.09216594 0.05237905
                        F
##
                                   Ρ
                                               S
## 0.01695355 0.03323242 0.04400956 0.09865235 0.06662918 0.01463302
## 0.03807384 0.06307173
Reduce('+', artificial_2_aac)/length(artificial_2_aac)
##
            Α
                        R
                                   N
                                               D
                                                          C
                                                                      Ε
## 0.06681774 0.09424995 0.03355575 0.03320151 0.03318094 0.03266863
                        G
                                               Ι
                                   Η
## 0.03317184 0.06386547 0.03403190 0.04785987 0.09575508 0.03288863
            М
                        F
                                   P
                                               S
                                                          Т
## 0.01713940 0.03254250 0.06717763 0.10010313 0.06718395 0.01835640
            Y
## 0.03379739 0.06245228
```

After removing some unwanted letters and characters, the observed amino acids remain for the obtained protein sequences. Distribution of the amino acids among the three databases of obtained protein sequences is rather similar for all three protein databases.

```
library(seqinr)
library(stringr)

# reading original_dataset from fasta file
lizards_sequences = read.fasta("lizard_seqs.fasta")

# preparing data in fasta file (dna sequences include emtpy spaces which will be removed)
for (i in 1:length(lizards_sequences)) {
    lizards_sequences[[i]] = lizards_sequences[[i]][lizards_sequences[[i]] != " "]
}
taa_count <- c()
tag_count <- c()
tga_count <- c()</pre>
```

```
string <- lizards_sequences[[i]]</pre>
string <- paste(lizards_sequences[[i]], collapse = "")</pre>
taa_count[i] <-str_count(string, pattern = "taa")</pre>
tag_count[i] <- str_count(string, pattern = "tag")</pre>
tga_count[i] <- str_count(string, pattern = "tga")</pre>
names_sequences <- names(lizards_sequences)</pre>
df_original <- as.data.frame(cbind(names_sequences, taa_count, tag_count, tga_count,
                                      total_count_1 = taa_count + tag_count + tga_count))
artificial_sequences_1 <- read.fasta("artificial_dataset_1_1.fasta")</pre>
taa a1 <- c()
tag_a1 <- c()
tga_a1 <- c()
for (i in 1:33){
  string <- artificial_sequences_1[[i]]</pre>
  string <- paste(artificial_sequences_1[[i]], collapse = "")</pre>
  taa_a1[i] <-str_count(string, pattern = "taa")</pre>
  tag_a1[i] <- str_count(string, pattern = "tag")</pre>
  tga_a1[i] <- str_count(string, pattern = "tga")</pre>
names_a1 <- names(artificial_sequences_1)</pre>
df_a1 <- as.data.frame(cbind(names_a1, taa_a1, tag_a1, tga_a1, total_count_2 =
                                  taa_a1 + tag_a1 + tga_a1))
artificial_sequences_2 <- read.fasta("artificial_dataset_1_2.fasta")
taa_a2 <- c()
tag_a2 <- c()
tga_a2 <- c()
for (i in 1:33){
  string <- artificial_sequences_2[[i]]</pre>
  string <- paste(artificial_sequences_2[[i]], collapse = "")</pre>
  taa_a2[i] <-str_count(string, pattern = "taa")</pre>
  tag_a2[i] <- str_count(string, pattern = "tag")</pre>
  tga_a2[i] <- str_count(string, pattern = "tga")</pre>
names_a2 <- names(artificial_sequences_1)</pre>
df_a2 <- as.data.frame(cbind(names_a2, taa_a2, tag_a2, tga_a2, total_count_3 =</pre>
                                  taa_a2 + tag_a2 + tga_a2))
df_all <- as.data.frame(cbind(df_a1, df_a2))</pre>
df_all
##
      names_a1 taa_a1 tag_a1 tga_a1 total_count_2 names_a2 taa_a2 tag_a2
                    23
## 1
              1
                            26
                                    16
                                                              1
                                                                     40
                                                                             34
                                                   65
## 2
              2
                     64
                            54
                                    45
                                                  163
                                                              2
                                                                     33
                                                                             29
## 3
              3
                    62
                            39
                                    41
                                                  142
                                                              3
                                                                     25
                                                                             35
                                                                             27
## 4
              4
                    22
                            13
                                    23
                                                   58
                                                                     28
```

##	5	5	30	27	24	81	5	33	24
##		6	24	19	23	66	6	22	31
##		7	66	55	54	175	7	21	32
##		8	28	25	15	68	8	30	38
##		9	23	18	21	62	9	28	43
	10	10	87	49	44	180	10	23	35
	11	11	64	53	60	177	11	37	33
	12	12	77	61	50	188	12	33	27
	13	13	68	41	52	161	13	31	39
	14	14	75	59	53	187	14	44	18
##	15	15	71	55	48	174	15	41	27
##	16	16	18	21	10	49	16	28	35
##	17	17	21	14	24	59	17	33	30
##	18	18	86	50	43	179	18	23	29
##	19	19	62	58	46	166	19	32	31
##	20	20	24	17	17	58	20	43	30
##	21	21	38	29	26	93	21	39	36
##	22	22	67	45	54	166	22	26	32
##	23	23	33	27	20	80	23	34	39
##	24	24	18	21	14	53	24	29	30
##	25	25	73	59	54	186	25	29	36
	26	26	69	41	39	149	26	34	31
	27	27	27	18	15	60	27	44	39
	28	28	69	40	48	157	28	36	32
	29	29	79	56	59	194	29	23	37
	30	30	12	17	25	54	30	50	32
##	31	31	72	47	48	167	31	42	34
##	32	32	17	22	22	61	32	25	45
## ##	33	33	26	14	14	54	33	35	28
	1	tga_a2 tota 30	_	ւ_3 104					
	2	37	•	99					
##		31		91					
##		25		80					
##		33		90					
	6	42		95					
##	7	36		89					
##	8	31		99					
##	9	22		93					
##	10	34		92					
##	11	26		96					
##	12	37		97					
	13	36		106					
	14	34		96					
	15	35	:	103					
	16	32		95					
	17	34		97					
	18	28		80					
	19	30	93 07						
## ##		24		97 106					
##		31 29	106 87						
	23	29 21		94					
		26		9 4 85					
##	24								

```
## 25
           32
                          97
## 26
           35
                         100
## 27
           37
                         120
## 28
                          92
           24
## 29
           28
                          88
## 30
           25
                         107
## 31
           23
                          99
## 32
                         101
           31
## 33
           24
                          87
```

Interpreting stop codons as either "taa", "tag" or "tga" results in many stop codons for each sequence. In the original dataset this is highly unlikely, as a natural translation starts at a start codon and then continues until it reaches a stop codon. Or if it does not reach a stop codon at all.

```
reverse_complemented_lizards <- read.fasta("lizards_reverse_complement.fasta")
taa_reverse <- c()
tag_reverse <- c()

for (i in 1:33){
    string <- reverse_complemented_lizards[[i]]
    string <- paste(reverse_complemented_lizards[[i]], collapse = "")
    taa_reverse[i] <-str_count(string, pattern = "taa")
    tag_reverse[i] <- str_count(string, pattern = "tag")
    tga_reverse[i] <- str_count(string, pattern = "tga")
}

names_reverse <- names(reverse_complemented_lizards)
df_reverse <- as.data.frame(cbind(names_reverse, taa_reverse, tag_reverse, total_count_reverse)
    taa_reverse + tag_reverse + tag_reverse))
df_reverse</pre>
```

##		names_reverse	taa_reverse	tag_reverse	tga_reverse	total_count_reverse
##	1	JF806202	11	5	21	37
##	2	HM161150	28	22	55	105
##	3	FJ356743	29	25	57	111
##	4	JF806205	12	5	19	36
##	5	JQ073190	20	11	30	61
##	6	GU457971	10	7	20	37
##	7	FJ356741	26	27	59	112
##	8	JF806207	11	6	19	36
##	9	JF806210	11	7	20	38
##	10	AY662592	24	28	60	112
##	11	AY662591	26	25	55	106
##	12	FJ356748	29	26	54	109
##	13	JN112660	27	23	59	109
##	14	AY662594	27	24	55	106
##	15	JN112661	31	22	55	108
##	16	HQ876437	9	10	17	36
##	17	HQ876434	10	8	20	38
##	18	AY662590	35	29	50	114
##	19	FJ356740	30	26	56	112
##	20	JF806214	11	10	18	39
##	21	JQ073188	22	8	29	59
##	22	FJ356749	27	26	64	117

```
##
  25
            AY662598
                                29
                                             29
                                                          61
                                                                                119
                                             20
## 26
            JN112653
                                17
                                                          51
                                                                                 88
##
  27
            JF806204
                                10
                                              7
                                                          20
                                                                                 37
## 28
            FJ356747
                                31
                                             31
                                                          56
                                                                                118
## 29
            FJ356744
                                31
                                             28
                                                          61
                                                                                120
## 30
            HQ876440
                                11
                                              5
                                                          20
                                                                                 36
## 31
            JN112651
                                25
                                             23
                                                          57
                                                                                105
                                 8
                                              9
                                                          23
                                                                                 40
## 32
            JF806215
## 33
            JF806209
                                10
                                              6
                                                          18
                                                                                 34
stop_codons <- c(sum(as.numeric(df_all$total_count_2)), sum(as.numeric(df_all$total_count_3)), sum(as.n
names(stop_codons) <- c("a1", "a2", "reverse_complemented")</pre>
stop_codons
```

30

18

61

36

a1 a2 reverse_complemented ## 510 424 343

21

10

After reverse complementing the original dataset from the lizards, for this dataset we again determine the number of stopcodons observed. Compared to the two artificially created datasets, the numer of stop codons is smaller. We think this can be explained because creating artificial datasets of sequences completely randomly assigns letters in the sequence, whereas in the reverse complement, the sequence is based on the original sequences. The original sequences will never contain as much stop codons as the artifically created ones.

10

8

Question 2.2

23

24

JQ073189

JF806216

```
library(markovchain)
mcFitMle_original <- markovchainFit(lizards_sequences, method = "mle")</pre>
mcFitMle_original
## $estimate
## MLE Fit
   A 8 - dimensional discrete Markov Chain defined by the following states:
   a, c, g, m, r, s, t, y
##
   The transition matrix (by rows)
                                   is defined as follows:
##
                                                                     s
                                g
## a 0.3377604 0.1730948 0.27493261 4.900760e-05 0.0002450380 0.000000e+00
## c 0.3793901 0.2477071 0.05010812 0.000000e+00 0.0003728283 0.000000e+00
## g 0.3934372 0.2029168 0.19323832 6.629102e-05 0.0003314551 0.000000e+00
## m 0.0000000 0.0000000 0.66666667 0.000000e+00 0.0000000000 0.000000e+00
## r 0.4117647 0.1764706 0.11764706 0.000000e+00 0.0000000000 0.000000e+00
## t 0.1508047 0.2115396 0.35718190 6.073489e-05 0.0001214698 6.073489e-05
## y 0.3333333 0.2000000 0.13333333 0.000000e+00 0.0000000000 0.000000e+00
##
## a 0.2136731 0.0002450380
## c 0.3222728 0.0001491313
## g 0.2096122 0.0003977461
## m 0.3333333 0.0000000000
## r 0.2941176 0.0000000000
## s 0.0000000 0.000000000
## t 0.2801093 0.0001214698
```

```
## y 0.3333333 0.0000000000
##
##
## $standardError
##
            a
                      С
                                g
## a 0.004068516 0.002912552 0.003670666 4.900760e-05 1.095843e-04
## c 0.005318784 0.004297725 0.001932963 0.000000e+00 1.667339e-04
## g 0.005106990 0.003667637 0.003579101 6.629102e-05 1.482312e-04
## m 0.000000000 0.000000000 0.471404521 0.000000e+00 0.000000e+00
## r 0.155632430 0.101885342 0.083189033 0.000000e+00 0.000000e+00
## s 0.000000000 1.000000000 0.000000000 0.000000e+00 0.000000e+00
## t 0.003026402 0.003584388 0.004657618 6.073489e-05 8.589211e-05
## y 0.149071198 0.115470054 0.094280904 0.000000e+00 0.000000e+00
##
             s
                       t
## a 0.000000e+00 0.003235986 1.095843e-04
## c 0.000000e+00 0.004902089 1.054518e-04
## g 0.000000e+00 0.003727654 1.623792e-04
## m 0.000000e+00 0.33333333 0.000000e+00
## r 0.000000e+00 0.131533410 0.000000e+00
## s 0.000000e+00 0.00000000 0.000000e+00
## t 6.073489e-05 0.004124610 8.589211e-05
## y 0.000000e+00 0.149071198 0.000000e+00
##
## $confidenceLevel
## [1] 0.95
##
## $lowerEndpointMatrix
##
           а
                      С
                               g m
                                            r s
## a 0.33106824 0.168304107 0.26889491 0 6.478782e-05 0 0.20835040
## c 0.37064143 0.240637977 0.04692868 0 9.857546e-05 0 0.31420954
## g 0.38503694 0.196884079 0.18735122 0 8.763643e-05 0 0.20348075
## r 0.15577214 0.008884115 0.00000000 0 0.000000e+00 0 0.07776444
## t 0.14582675 0.205643836 0.34952080 0 0.000000e+00 0 0.27332494
## y 0.08813303 0.010068663 0.00000000 0 0.000000e+00 0 0.08813303
##
## a 6.478782e-05
## c 0.00000e+00
## g 1.306561e-04
## m 0.00000e+00
## r 0.00000e+00
## s 0.00000e+00
## t 0.00000e+00
## y 0.00000e+00
##
## $upperEndpointMatrix
          а
                            g
## a 0.3444525 0.1778856 0.28097032 0.0001296179 0.0004252881 0.0000000000
## c 0.3881387 0.2547762 0.05328756 0.0000000000 0.0006470811 0.0000000000
## g 0.4018374 0.2089495 0.19912541 0.0001753300 0.0005752738 0.0000000000
```

```
## t 0.1557827 0.2174354 0.36484300 0.0001606349 0.0002627497 0.0001606349
## a 0.2189958 0.0004252881
## c 0.3303360 0.0003225840
## g 0.2157436 0.0006648361
## m 0.8816179 0.0000000000
## r 0.5104709 0.0000000000
## s 0.0000000 0.000000000
## t 0.2868937 0.0002627497
## y 0.5785336 0.0000000000
mcFitMle_a1 <- markovchainFit(artificial_sequences_1, method = "mle")</pre>
mcFitMle_a1
## $estimate
## MLE Fit
## A 4 - dimensional discrete Markov Chain defined by the following states:
## a, c, g, t
## The transition matrix (by rows) is defined as follows:
            a
                     С
## a 0.3120284 0.2075646 0.2346785 0.2457285
## c 0.3167468 0.2062847 0.2282440 0.2487246
## g 0.3145375 0.2028889 0.2315796 0.2509939
## t 0.3141239 0.2097648 0.2271888 0.2489225
##
##
## $standardError
              а
                          C.
## a 0.003897312 0.003178665 0.003379907 0.003458564
## c 0.004839355 0.003905394 0.004108006 0.004288355
## g 0.004565232 0.003666535 0.003917209 0.004078103
## t 0.004397752 0.003593739 0.003740018 0.003914825
## $confidenceLevel
## [1] 0.95
##
## $lowerEndpointMatrix
##
            a
                      С
## a 0.3056179 0.2023362 0.2291190 0.2400396
## c 0.3087867 0.1998609 0.2214869 0.2416709
## g 0.3070284 0.1968580 0.2251364 0.2442860
## t 0.3068902 0.2038536 0.2210370 0.2424832
## $upperEndpointMatrix
            a
                      С
## a 0.3184389 0.2127931 0.2402379 0.2514173
## c 0.3247068 0.2127085 0.2350011 0.2557783
## g 0.3220466 0.2089199 0.2380229 0.2577018
## t 0.3213575 0.2156760 0.2333406 0.2553619
mcFitMle_a2 <- markovchainFit(artificial_sequences_2, method = "mle")</pre>
mcFitMle a2
```

\$estimate

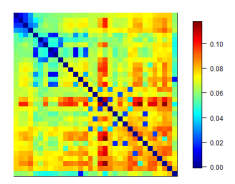
```
## MLE Fit
##
   A 4 - dimensional discrete Markov Chain defined by the following states:
   a, c, g, t
##
   The transition matrix (by rows)
                                       is defined as follows:
##
             a
                       С
                                  g
## a 0.2498317 0.2457923 0.2574209 0.2469551
## c 0.2449258 0.2490845 0.2583328 0.2476569
## g 0.2500905 0.2443614 0.2538898 0.2516584
## t 0.2548804 0.2466801 0.2449055 0.2535341
##
##
##
   $standardError
##
                                                    t
                           С
                                        g
## a 0.003910309 0.003878568 0.003969256 0.003887732
## c 0.003899025 0.003931987 0.004004318 0.003920703
## g 0.003883561 0.003838821 0.003912949 0.003895716
## t 0.003949380 0.003885329 0.003871327 0.003938935
##
## $confidenceLevel
##
   [1] 0.95
##
## $lowerEndpointMatrix
##
             а
                       С
                                  g
## a 0.2433998 0.2394126 0.2508920 0.2405604
  c 0.2385125 0.2426169 0.2517463 0.2412079
  g 0.2437026 0.2380471 0.2474535 0.2452505
  t 0.2483842 0.2402893 0.2385377 0.2470551
##
##
  $upperEndpointMatrix
##
             a
                       С
                                            t
## a 0.2562636 0.2521720 0.2639497 0.2533499
## c 0.2513392 0.2555520 0.2649193 0.2541059
## g 0.2564783 0.2506757 0.2603260 0.2580663
## t 0.2613765 0.2530709 0.2512732 0.2600130
```

We fitted a first order markov model on all sequences. Our assumption in our simulated datasets is that in the sequence the occurrence of a nucleotide does not depend on the rest of the sequence. This violates the limited horizon: which is that the probability of being in a state at time t depends only on the state at time t minus 1. We used sample {base} function, which obviously samples without taking into account past states.

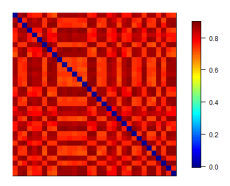
Question 2.3

To allign the sequences for each dataset (the original dataset $lizards_sequences$, the first artificial dataset $artificial_dataset_1_1$ and the second artificial dataset $artificial_dataset_1_2$), the plsgenomics package was used. The .fasta-files for the datasets were transformed to a DNAStringSet - class within R. The uncorrected distance matrices created represent the hamming distance between each of the sequences in each dataset. The results of these distance matrices are plotted as heatmaps (using plsgenomics package):

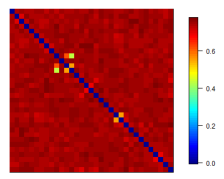
lizards sequences



$artificial_dataset_1_1$



 $artificial_dataset_1_2$



We see that for the original dataset, the allignment results are much better than for the artificial datasets. Based on the point that the artificial datasets were created by sampling randomly, the greater distances between the sequences compared to the distances within the original dataset make sense.

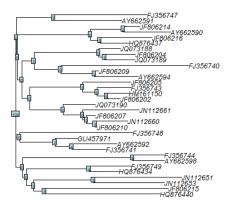
The R code for this Question 2.3 can be found in Appendix 2.3.

Question 3

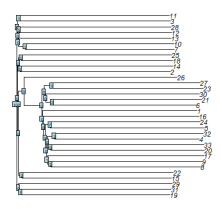
Question 3.1

Using the created distance matrix for each dataset (the original dataset *lizards_sequences*, the first artificial dataset *artificial_dataset_1_1* and the second artificial dataset *artificial_dataset_1_2*) with the aligned sequences, phylotrees were created. On top of that, a phylogenetic bootstrap analysis was performed. As a result, the bootstrap supports for the individual clades were integrated into the phylotrees.

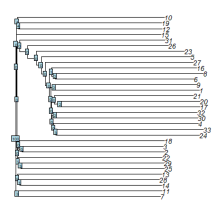
lizards_sequences



artificial_dataset_1_1



artificial dataset 1 2



The R code for the creation of the phylotrees and the bootstrap analysis can be found in Appendix 3.1.

Question 3.2

Different general characteristics can be comprared between phylogenetic trees, e.g.:

- number of tips
- different tips
- number of nodes

On top of that, different quantitative distances can be calculated, e.g.:

- symmetric difference
- branch score

The distances can be only calculated if the tips are named equally. Since the artificial datasets (artificial_dataset_1_1 and artificial_dataset_1_2) are not named as the original dataset (lizard_sequences), the distance measurements could be only processed for the comparison between the artificial datasets.

```
## => Comparing phylotree1 with phylotree2.
## Both trees have the same number of tips: 33.
## Tips in phylotree1 not in phylotree2 : JF806202, HM161150, FJ356743, JF806205, JQ073190, GU457971, F
## Tips in phylotree2 not in phylotree1 : 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18
## Both trees have the same number of nodes: 31.
## Both trees are unrooted.
## Both trees are not ultrametric.
## => Comparing phylotree1 with phylotree2.
## Both trees have the same number of tips: 33.
## Tips in phylotree1 not in phylotree2 : JF806202, HM161150, FJ356743, JF806205, JQ073190, GU457971, F
## Tips in phylotree2 not in phylotree1 : t6, t20, t3, t14, t21, t23, t15, t4, t7, t22, t18, t24, t5, t
## Both trees have the same number of nodes: 31.
## Both trees are unrooted.
## Both trees are not ultrametric.
## => Comparing phylotree1 with phylotree2.
## Both trees have the same number of tips: 33.
## Tips in phylotree1 not in phylotree2 : 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18
## Tips in phylotree2 not in phylotree1 : t6, t20, t3, t14, t21, t23, t15, t4, t7, t22, t18, t24, t5, t
## Both trees have the same number of nodes: 31.
## Both trees are unrooted.
## Both trees are not ultrametric.
```

Appendix 1

Data Import of original dataset

```
library(ape)
lizards_accession_numbers <- c("JF806202", "HM161150", "FJ356743", "JF806205",
                                "JQ073190", "GU457971", "FJ356741", "JF806207",
                                "JF806210", "AY662592", "AY662591", "FJ356748",
                                "JN112660", "AY662594", "JN112661", "HQ876437",
                                "HQ876434", "AY662590", "FJ356740", "JF806214",
                                "JQ073188", "FJ356749", "JQ073189", "JF806216",
                                "AY662598", "JN112653", "JF806204", "FJ356747",
                                "FJ356744", "HQ876440", "JN112651", "JF806215",
                                "JF806209")
lizards_sequences<-ape::read.GenBank(lizards_accession_numbers)</pre>
print(lizards_sequences)
ape::write.dna(lizards_sequences,
               file ="lizard_seqs.fasta",
               format = "fasta",
               append =FALSE,
               nbcol = 6,
               colsep = " "
               colw = 10)
```

Appendix 1.1

Reading original data

```
library(seqinr)
# reading original_dataset from fasta file
lizards_sequences = read.fasta("lizard_seqs.fasta")
```

Function code

```
library(seginr)
get_artificial_sequence_dataset = function(original_dataset) {
  # creating empty varibales which will be filled in following for-loop
  original_base_compositions = list()
  artificial_dataset = list()
  artificial base compositions = list()
  a_original = c(); c_original = c(); g_original = c(); t_original = c()
  a_artificial = c(); c_artificial = c(); g_artificial = c(); t_artificial = c()
  for (i in 1:length(original_dataset)) {
    # getting base compositions for each original sequence
   original base compositions[[i]] =
      seqinr::count(original dataset[[i]],1)/length(original dataset[[i]])
    # creating artificial sequences randomly drawn from the distribution
    # given by the base composition
    artificial_dataset[[as.character(i)]] = sample(x = c("a", "c", "g", "t"),
                                                   size = length(original_dataset[[i]]),
                                                   rep = TRUE,
                                                   prob = original_base_compositions[[i]])
    # creating dataframe to compare base compositions
    # between original and artificial sequences
   artificial_base_compositions[[i]] =
      seqinr::count(artificial_dataset[[i]],1)/length(artificial_dataset[[i]])
   a original = c(a original, round(original base compositions[[i]][1],2))
   a_artificial = c(a_artificial, round(artificial_base_compositions[[i]][1],2))
    c original = c(c original, round(original base compositions[[i]][2],2))
    c_artificial = c(c_artificial, round(artificial_base_compositions[[i]][2],2))
   g_original = c(g_original, round(original_base_compositions[[i]][3],2))
   g artificial = c(g artificial, round(artificial base compositions[[i]][3],2))
   t original = c(t original, round(original base compositions[[i]][4],2))
   t_artificial = c(t_artificial, round(artificial_base_compositions[[i]][4],2))
  comparison_base_compositions = cbind(
   name_original = names(original_dataset), name_artificial = names(artificial_dataset),
    a_original, a_artificial, c_original, c_artificial,
    g_original, g_artificial, t_original, t_artificial
  rownames(comparison_base_compositions) = 1:nrow(comparison_base_compositions)
  print("comparison of base compositions
       between original and artificial datasets (values rounded): ")
  print(comparison base compositions)
  # saving fasta file
  ape::write.dna(artificial dataset, file ="artificial dataset 1 1.fasta", format = "fasta",
                 colsep = "")
```

Appendix 1.2

Replace the integers by letters

```
for (k in 1:33){
  sequences_artificial[[k]] [sequences_artificial[[k]] == 1] = "a"
  sequences_artificial[[k]] [sequences_artificial[[k]] == "2"] = "c"
  sequences_artificial[[k]] [sequences_artificial[[k]] == "3"] = "g"
  sequences_artificial[[k]] [sequences_artificial[[k]] == "4"] = "t"
}
```

Appendix 2

Appendix 2.3

```
library(seqinr)
library(DECIPHER)
library(plsgenomics)
library(ape)
# getting all datasets in DNAStringSet format
  # original dataset
  lizards_sequences = readDNAStringSet("lizard_seqs.fasta")
  # artificial_dataset_1_1
  artificial_dataset_1_1 = readDNAStringSet("artificial_dataset_1_1.fasta")
  # artificial_dataset_1_2
  artificial_dataset_1_2 = readDNAStringSet("artificial_dataset_1_2.fasta")
# alliquing sequences for each dataset
sequence_alligning = function(dataset, name) {
  # alligning process
  sequences_alligned = AlignSeqs(dataset)
  # creating distance matrix
  dm_sequences_alligned = DistanceMatrix(sequences_alligned)
  saveRDS(dm sequences alligned, paste0("distanceMatrix ", name, ".RDS"))
  # creating matrix heatmap
  heatmap_dm_sequences_alligned = matrix.heatmap(dm_sequences_alligned)
  dev.copy(png,paste("heatmap_", name, ".png", sep=""))
  dev.off()
  return(sequences_alligned)
lizards_sequences_alligned = sequence_alligning(dataset = lizards_sequences, name = "lizards_sequences"
artificial_dataset_1_1_alligned = sequence_alligning(artificial_dataset_1_1, name = "artificial_dataset
artificial_dataset_1_2_alligned = sequence_alligning(artificial_dataset_1_2, name = "artificial_dataset
```

Appendix 3

Appendix 3.1

```
library(seqinr)
library(DECIPHER)
library(plsgenomics)
library(ape)
# creating phylotrees
create_phylotree = function(dataset_name) {
  distanceMatrix = readRDS(paste0("distanceMatrix_", dataset_name, ".RDS"))
  tree = nj(distanceMatrix)
  png(paste("phylotree_", dataset_name, ".png", sep=""))
 plot(tree)
 dev.off()
 return(tree)
tree_lizards_sequences = create_phylotree("lizards_sequences")
tree_artificial_dataset_1_1 = create_phylotree("artificial_dataset_1_1")
tree_artificial_dataset_1_2 = create_phylotree("artificial_dataset_1_2")
# performing bootstrap analysis
bootstrap_analysis = function(dataset_name, tree_object) {
  distanceMatrix = readRDS(paste0("distanceMatrix_", dataset_name, ".RDS"))
  bootstrap_result = boot.phylo(phy = tree_object,
                                x = distanceMatrix,
                                FUN = function(x) {
                                  nj(x)
                                })
  png(paste("bootstrap_phylotree_", dataset_name, ".png", sep=""))
  plot(tree object)
 nodelabels(bootstrap_result, cex=.6)
  dev.off()
}
bootstrap_analysis("lizards_sequences", tree_lizards_sequences)
bootstrap analysis ("artificial dataset 1 1", tree artificial dataset 1 1)
bootstrap_analysis("artificial_dataset_1_2", tree_artificial_dataset_1_2)
```

Appendix 3.2

```
library(phangorn)
compare_phylotrees = function(phylotree1, phylotree2) {
  if(all(phylotree1$tip.label == phylotree2$tip.label)) {
    comparePhylo(phylotree1, phylotree2)
    treedist(phylotree1, phylotree2)
  } else {
    comparePhylo(phylotree1, phylotree2)
  }
}
# Comparing tree_lizards_sequences & tree_artificial_dataset_1_1
```

```
compare_phylotrees(tree_lizards_sequences, tree_artificial_dataset_1_1)
# Comparing tree_lizards_sequences & tree_artificial_dataset_1_2
compare_phylotrees(tree_lizards_sequences, tree_artificial_dataset_1_2)
# Comparing tree_artificial_dataset_1_1 & tree_artificial_dataset_1_2
compare_phylotrees(tree_artificial_dataset_1_1, tree_artificial_dataset_1_2)
```