Project 3 Markov chains and hidden Markov models

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Problem 6: Hidden Markov Models

According to slides 18, the HMM is parameterized by:

Initial state probabilities: $I_k = P(Z_1 = k)$ Transition probabilities: $T_{kl} = P(Z_n = l | Z_{n-1} = k)$ Emission probabilities: $E_{kx} = P(X_n = x | Z_n = k)$

(a)

 I_k should sum up to $1 \longrightarrow \text{degree}$ of freedom = K-1

In the $K \times K$ matrix T, each row needs to sum up to $1 \longrightarrow \text{degree}$ of freedom = K-1 for each row Similarly, in the $K \times M$ matrix X, each row needs to sum up to $1 \longrightarrow \text{degree}$ of freedom = M-1 for each row

Hence, the maximum number of free parameters is $(K-1)+K\times (K-1)+K\times (M-1)$

(b)

Computing the stationary distribution is equivalent to solving $\pi^t = \pi^t T$.

Let $\pi^t = \begin{bmatrix} \pi_1 & \pi_2 \end{bmatrix}$), we need to solve:

$$\begin{aligned} \left[\pi_1 \quad \pi_2\right] &= \left[\pi_1 \quad \pi_2\right] \begin{bmatrix} 0.3 & 0.7 \\ 0.2 & 0.8 \end{bmatrix} \\ &= \left[0.3\pi_1 + 0.2\pi_2 & 0.7\pi_1 + 0.8\pi_2\right] \end{aligned}$$

Given that $\pi_1 = 1 - \pi_2 e$ can solve the equations:

$$\begin{cases} \pi_1 = 0.3\pi_1 + 0.2\pi_2 \\ \pi_2 = 0.7\pi_1 + 0.8\pi_2 \Rightarrow \begin{cases} \pi_1 = \frac{2}{9} \\ \pi_1 = 1 - \pi_2 \end{cases} \Rightarrow \begin{cases} \pi_2 = \frac{7}{9} \end{cases}$$

Alternatively we can solve by solving eigens (which is porbably the more proper way to do it because solving the equations above by hand for high dimensional transition matrices is not a very smart idea). Since $\pi^t = \pi^t T$, we can re-write it into $\pi = T^t \pi$, which means π is just the corresponding eigenvector of eigenvalue $\lambda = 1$.

Note that the eigenvector should be normalized to satisfy the constraint that $\pi_1 + \pi_2 = 1$.

```
# initialize the transition matrix
Tr <- matrix(c(0.3, 0.7, 0.2, 0.8), 2, 2, byrow=TRUE)

# solve for the eigenvalues and eigenvectors for transpose of T
eigens <- eigen(t(Tr))

# get the index of lambda = 1
index <- which(eigens$values %>% near(1.0))[1]
ev <- eigens$vectors[, index]

# normalize so that the probs sum to 1
pi <- ev / sum(ev)
pi

## [1] 0.2222222 0.7777778

# check that the two computation confirms
near(pi[1], 2/9)

## [1] TRUE

near(pi[2], 7/9)</pre>
```

Problem 7: Predictinig protein secondary structure using HMMs

(a)

Read proteins train.tsv, proteins test.tsv and proteins new.tsv into the memory and store each in a data.frame

```
## identifier

## 1 >101M:A

## 2 >102L:A

## 3 >102M:A

## 4 >103L:A

## 5 >103M:A

## 6 >104L:B
```

1 MVLSEGEWQLVLHVWAKVEADVAGHGQDILIRLFKSHPETLEKFDRVKHLKTEAEMKASEDLKKHGVTVLTALGAILKKKGHHEA
2 MNIFEMLRIDEGLRLKIYKDTEGYYTIGIGHLLTKSPSLNAAAKSELDKAIGRNTNGVITKDEAEKLFNQDVDAAVRGILRNAKLKPVYDSLDAVR
3 MVLSEGEWQLVLHVWAKVEADVAGHGQDILIRLFKSHPETLEKFDRFKHLKTEAEMKASEDLKKAGVTVLTALGAILKKKGHHEA

(b)

Estimate the vector of initial state probabilities I, the matrix of transition probabilities T and the matrix for emission probabilities E by maximum likelihood.

```
First get all the possible amino acid and secondary structure states.
# get all the possible amino acid states
aa_states <- train$sequence %>%
  strsplit("") %>%
  unlist() %>%
  unique() %>%
  sort()
aa states
## [1] "A" "C" "D" "E" "F" "G" "H" "I" "K" "L" "M" "N" "P" "Q" "R" "S" "T" "U" "V"
## [20] "W" "X" "Y"
# get all the possible secondary structure states, and sort them
ss_states <- train$structure %>%
  strsplit("") %>%
  unlist() %>%
 unique() %>%
  sort()
ss_states
## [1] "B" "C" "E" "G" "H" "I" "S" "T"
MLE <- function(data, aa states, ss states) {</pre>
  k <- length(ss_states) # num of latent state</pre>
  m <- length(aa_states) # num of observed state
  # initialize the parameters
  I \leftarrow rep(0.0, k)
  names(I) <- ss_states</pre>
  E <- matrix(0.0, nrow=k, ncol=m)</pre>
  dimnames(E) <- list(ss_states, aa_states)</pre>
  Tr <- matrix(0.0, nrow=k, ncol=k)</pre>
  dimnames(Tr) <- list(ss_states, ss_states)</pre>
  N <- nrow(data) # num of data points
  # iterate over each row of the data
  # TODO: maybe rewrite the for loops
```

```
foreach (i=1:N) %do% {
    seq <- data$sequence[i]</pre>
    struct <- data$structure[i]</pre>
   ss_1st <- struct %>% substr(1, 1)
   I[ss_1st] \leftarrow I[ss_1st] + 1.0
   for (j in 1:nchar(seq)) {
      aa <- seq %>% substr(j, j)
      ss <- struct %>% substr(j, j)
     E[ss, aa] \leftarrow E[ss, aa] + 1.0
      if (j < nchar(seq)) {</pre>
       ss_next <- struct %>% substr(j+1, j+1)
       Tr[ss, ss_next] <- Tr[ss, ss_next] + 1.0</pre>
   }
  }
  # convert thee counts to log probs
  I <- I / sum(I)</pre>
 E <- E / rowSums(E)
  Tr <- Tr / rowSums(Tr)
 return(list(I=I, E=E, Tr=Tr))
}
res <- MLE(train, aa states, ss states)
res$I
## B C E G H I S T
## 0 1 0 0 0 0 0 0
res$E
## B 0.04524422 0.03239075 0.06580977 0.02467866 0.04215938 0.05192802 0.03239075
## C 0.06450873 0.01585481 0.07227845 0.04888328 0.03388859 0.08271453 0.02583216
## E 0.05804938 0.02348868 0.02798043 0.04002518 0.05247046 0.04809315 0.02752267
## G 0.10814116 0.01677149 0.09696017 0.08333333 0.03511530 0.06289308 0.03022362
## H 0.12500000 0.01249647 0.05261579 0.08629271 0.03646569 0.03503601 0.02188647
## S 0.05445111 0.01536492 0.08605701 0.05215985 0.02331694 0.14219287 0.02304738
## T 0.06342143 0.01168713 0.07736021 0.06015118 0.02348148 0.19911006 0.02122983
                        K
                                   T.
                                              М
                                                         N
                                                                    Ρ
## B 0.06221080 0.04575835 0.07506427 0.01645244 0.05449871 0.04215938 0.02982005
## C 0.04323518 0.06118295 0.07018550 0.02319447 0.05708306 0.08277187 0.03145159
## E 0.09306783 0.05767745 0.10511258 0.02374617 0.02400366 0.01782393 0.03135639
## G 0.03162124 0.05957372 0.05835080 0.01502446 0.05241090 0.06446541 0.03546471
## H 0.05491034 0.07296668 0.11285654 0.02974089 0.03641274 0.01929187 0.04131954
## I 0.05882353 0.00000000 0.23529412 0.05882353 0.00000000 0.00000000 0.00000000
## S 0.02749511 0.07244423 0.05067727 0.01489319 0.05492284 0.05761844 0.03093200
```

```
## T 0.02466091 0.07918297 0.04975071 0.01302740 0.06733501 0.06792473 0.03683054
##
             R.
                        S
                                  Τ
                                              IJ
                                                         V
                                                                     W
## B 0.04627249 0.04627249 0.08020566 0.000000000 0.13110540 0.008740360
## C 0.04733507 0.07981880 0.07184839 0.000000000 0.05106224 0.008171106
## E 0.03833720 0.05198409 0.07779018 0.000000000 0.12651275 0.019740795
## G 0.03913347 0.06970650 0.04297694 0.000174703 0.02428372 0.024633124
## H 0.06412384 0.04048997 0.04356114 0.000000000 0.07183705 0.013061282
## I 0.11764706 0.00000000 0.00000000 0.000000000 0.05882353 0.058823529
## S 0.05364243 0.08807871 0.07224206 0.000000000 0.03935575 0.016106207
## T 0.04122661 0.06438643 0.04444325 0.000000000 0.02021123 0.010132418
##
               Х
                          Y
## B 0.0010282776 0.06580977
## C 0.0009461280 0.02775309
## E 0.000000000 0.05521701
## G 0.000000000 0.04874214
## H 0.000000000 0.02963499
## I 0.00000000 0.00000000
## S 0.0002695599 0.02473212
## T 0.0001072214 0.02433925
res$Tr
                          C
                                      Ε
                                                  G
                                                                         Ι
##
               В
                                                             Η
## B 0.0185089974 0.60874036 0.0303341902 0.020565553 0.02365039 0.000000e+00
## C 0.0288740867 0.51503859 0.1113882450 0.025176795 0.08471493 0.000000e+00
## E 0.0039195491 0.10814522 0.8126054988 0.004520356 0.00557892 0.000000e+00
## G 0.0078616352 0.11198463 0.0186932215 0.697938505 0.03336827 0.000000e+00
## H 0.0004412595 0.01782689 0.0003000565 0.002859362 0.91049492 1.765038e-05
## $ 0.0258777546 0.38250556 0.0857874520 0.018262686 0.06738999 6.738999e-05
## T 0.0179059669 0.22709484 0.0695866617 0.012866563 0.04026162 5.361068e-05
##
             S
## B 0.15372751 0.14447301
## C 0.13873647 0.09607089
## E 0.02932509 0.03590536
## G 0.05765199 0.07250175
## H 0.01623835 0.05182152
## I 0.0000000 0.11764706
## S 0.35656042 0.06354876
## T 0.12040959 0.51182115
(c)
```

Estimate the stationary distribution π of the Markov chain by solving the eigenvalue problem and by using a brute-force approach.

Eigenvalue method

```
eigens <- eigen(t(res$Tr))
index <- which(eigens$values %>% near(1.0))[1]
ev <- eigens$vectors[, index]</pre>
```

```
pi_eigen <- ev / sum(ev)
pi_eigen

## [1] 0.0116560594 0.2042297412 0.2094674769 0.0343029736 0.3395299222
## [6] 0.0001018782 0.0889276425 0.1117843060</pre>
```

Brute-force

```
# compute Tr %*% Tr until there's no huge difference between the two
# matrices
Tr <- res$Tr
pi_brute <- rep(0.0, length(ss_states))</pre>
pi_brute[1] <- 1.0
while (TRUE) {
  Tr <- Tr %*% res$Tr
 pi_new <- Tr[1, ]</pre>
  if (all(near(pi_new, pi_brute))) {
    break
  }
 pi_brute <- pi_new</pre>
pi_brute
##
                            C
                                         Ε
## 0.0116560609 0.2042297695 0.2094675407 0.0343029787 0.3395298132 0.0001018781
## 0.0889276516 0.1117843074
near(pi_eigen, pi_brute, tol=1e-5)
##
                           Η
                                           Τ
## TRUE TRUE TRUE TRUE TRUE TRUE TRUE
```

(d)

Having estimated the parameters, i.e., the emission and transition matrices E,T and the vector of initial state probabilities I, you can predict the latent state sequence Z of a protein's amino acid sequence X using the Viterbi algorithm. Use the Viterbi algorithm provided in viterbi.r (carefully read the parameter description!) and iterate over each data.frame of proteins_test.tsv and proteins_new.tsv row by row and use the amino acid sequence to predict its secondary structure, which you add to the data.frame as a new column. Save the extended data.frame of proteins_new.tsv including the predicted secondary structure as a tsv file and hand it in together with your pdf.

```
predict_test <- data.frame(test[,2])
colnames(predict_test) <- c("AminoAcids")
predict_test <- viterbi(log(res$E), log(res$Tr), log(res$I), predict_test)

predict_new <- data.frame(new[,2])
colnames(predict_new) <- c("AminoAcids")
predict_new <- viterbi(log(res$E), log(res$Tr), log(res$I), predict_new)</pre>
```

```
# save predict new to a tsv file
write.table(predict_new, file="proteins_new.tsv", sep="\t", quote=FALSE, row.names=FALSE)
```

(e)

Estimate confidence intervals for each parameter in I, E and T with bootstrapping. In a single bootstrap run i estimate the probabilities for I_i , E_i and T_i the same as before, but not on the original data set proteins train.tsv, but on the resampled data set. i.e., sample with replacement as many rows from proteins train.tsv as the original data set has. Run a thousand bootstraps and compute the empirical 95% confidence intervals for each single parameter in $\{I_i\}_i$, $\{E_i\}_i$ and $\{T_i\}_i$.

```
and compute the empirical 95% confidence intervals for each single parameter in \{I_i\}_i, \{E_i\}_i
     and \{T_i\}_i.
num_cores <- detectCores()</pre>
registerDoParallel(num_cores)
# 1000 rounds of bootstrap
start_time <- Sys.time()</pre>
df <- foreach(i=1:1000, .combine=rbind, .packages=c("foreach", "tidyverse")) %dopar% {</pre>
  # allow duplicates
  picked <- sample(seq_len(nrow(train)), size=nrow(train), replace = TRUE)</pre>
  bootstrap <- train[picked, ]</pre>
  params <- MLE(bootstrap, aa_states = aa_states, ss_states = ss_states)</pre>
  return(params)
end_time <- Sys.time()</pre>
end_time - start_time
## Time difference of 3.843328 mins
stopImplicitCluster()
registerDoSEQ()
  # iterate over each matrix
  mat <- as.data.frame(list_of_matrix[[1]])</pre>
  dims <- dim(mat)</pre>
```

```
compute_ci <- function(list_of_matrix) {
    # iterate over each matrix
    mat <- as.data.frame(list_of_matrix[[1]])
    dims <- dim(mat)
    df <- data.frame(matrix(0, nrow=length(list_of_matrix), ncol=prod(dims)))
    colnames(df) <- sapply(1:dims[2], function(i) {
        sapply(1:dims[1], function(j) {
            paste0(rownames(mat)[j], colnames(mat)[i])
        })
    }) %>% unlist()

foreach (i=1:length(list_of_matrix)) %do% {
        v <- as.vector(list_of_matrix[[i]])
        foreach (j=1:length(v)) %do% {
            df[i, j] <- v[j]
        }
    }
    df <- apply(df, 2, quantile, probs=c(.025, .975), na.rm=TRUE)
    return(df)
}</pre>
```

```
I_emp <- as.data.frame(df[, "I"])</pre>
CI_I <- apply(I_emp, 1, quantile, probs=c(.025, .975))</pre>
##
         BCEGHIST
## 2.5% 0 1 0 0 0 0 0 0
## 97.5% 0 1 0 0 0 0 0
CI_E <- compute_ci(df[, "E"])</pre>
CI_E
                            CA
                                       EΑ
                                                  GA
                                                            HA IA
## 2.5% 0.03733706 0.06112611 0.05524126 0.09992364 0.1215970 0.0 0.05092880
## 97.5% 0.05353647 0.06785321 0.06055642 0.11702478 0.1284658 0.6 0.05804788
                 TA
                            BC
                                       CC
                                                  EC
                                                             GC
## 2.5% 0.05999845 0.02481786 0.01423457 0.02166365 0.01342027 0.01096700 0
## 97.5% 0.06715470 0.03981281 0.01747747 0.02534800 0.02055510 0.01410316
                 SC
                           TC
                                      BD
                                                 CD
                                                             ED
## 2.5% 0.01306597 0.01016416 0.05480235 0.06952154 0.02610076 0.08968065
## 97.5% 0.01800880 0.01325272 0.07620683 0.07530680 0.03000679 0.10440100
                 HD ID
                                SD
                                           TD
##
                                                      BF.
                                                                 CF.
                                                                            F.F.
## 2.5% 0.05105591 0.0 0.08120528 0.07314596 0.01783826 0.04577216 0.03800390
## 97.5% 0.05429762 0.5 0.09071893 0.08125889 0.03217520 0.05188908 0.04234227
                            HE IE
                                          SE
                                                     ΤE
## 2.5% 0.07610046 0.08369242 0 0.04820251 0.05630231 0.03342719 0.03220069
## 97.5% 0.09053595 0.08899312 0 0.05616438 0.06387202 0.05135091 0.03569510
                                       HF IF
                 EF
                            GF
                                                     SF
                                                               TF
## 2.5% 0.04999996 0.03075498 0.03479685 0 0.02080610 0.0213674 0.04159314
## 97.5% 0.05518106 0.03977019 0.03807638 0 0.02583164 0.0256363 0.06329178
                 CG
                            EG
                                       GG
                                                  HG IG
                                                               SG
## 2.5% 0.07939679 0.04553001 0.05627687 0.03335435 0 0.1362343 0.1937047
## 97.5% 0.08596570 0.05073376 0.06911339 0.03668840 0 0.1484953 0.2047660
                 BH
                            CH
                                       EH
                                                  GH
                                                             HH IH
## 2.5% 0.02520050 0.02377693 0.02565301 0.02596137 0.02009471 0 0.02043839
## 97.5% 0.04041123 0.02802430 0.02963243 0.03453233 0.02370681 0 0.02587384
                 TH
                            BI
                                       CI
                                                  ΕI
                                                             GI
## 2.5% 0.01890120 0.05198883 0.04123898 0.08847165 0.02693896 0.05288477 0.0
## 97.5% 0.02361752 0.07239259 0.04520458 0.09764383 0.03554891 0.05689105 0.2
                 SI
                           ΤI
                                       BK
                                                  CK
## 2.5% 0.02478721 0.02245736 0.03701670 0.05752782 0.05522655 0.05405877
## 97.5% 0.03036456 0.02690024 0.05596181 0.06479144 0.05996290 0.06557216
                                                     BL
                 HK IK
                               SK
                                                                CL
##
                                          ΤK
## 2.5% 0.07043873 0 0.06774436 0.07406220 0.06350628 0.06707379 0.1018696
## 97.5% 0.07560704 0 0.07687417 0.08462968 0.08834292 0.07362631 0.1086728
                 GL
                           HL
                                     IL
                                                SL
## 2.5% 0.05229527 0.1093864 0.0000000 0.04720163 0.04651404 0.01018273
## 97.5% 0.06459985 0.1162665 0.3333333 0.05406354 0.05293468 0.02321071
                            EM
                                       GM
                                                  MI MH
                 CM
## 2.5% 0.02175342 0.02206482 0.01179750 0.02841927 0.0 0.01278327 0.01144065
## 97.5% 0.02485315 0.02539844 0.01821219 0.03103604 0.2 0.01710857 0.01453235
                 BN
                            CN
                                                  GN
                                                                           SN
                                       EN
## 2.5% 0.04425958 0.05415652 0.02232413 0.04678921 0.03467575 0 0.05094471
## 97.5% 0.06477399 0.06008876 0.02590115 0.05838007 0.03802648 0 0.05921721
```

EΡ

GP

CP

##

TN

BP

```
## 2.5% 0.06395880 0.03338276 0.07937671 0.01638889 0.05893941 0.01805794
## 97.5% 0.07088224 0.05016907 0.08639285 0.01935019 0.06985875 0.02051871
                SP
                           ΤP
                                      BQ
                                                 CQ
## 2.5% 0.05433799 0.06430807 0.02239806 0.02956940 0.02940178 0.03109497
## 97.5% 0.06115424 0.07178236 0.03804065 0.03350015 0.03330924 0.03985298
                HQ IQ
                              SQ
                                         ΤQ
                                                    BR.
                                                                          F.R.
## 2.5% 0.03964397 0 0.02829652 0.03402817 0.03750481 0.04494813 0.03622575
## 97.5% 0.04301873 0 0.03386126 0.03940188 0.05558400 0.04961411 0.04061903
                GR
                           HR
                                     IR
                                                SR
                                                           TR
## 2.5% 0.03468293 0.06174005 0.0000000 0.05017069 0.03847974 0.03693761
## 97.5% 0.04380980 0.06649552 0.3333333 0.05710966 0.04403945 0.05552892
                                                 HS IS
                CS
                           ES
                                                                          TS
                                      GS
## 2.5% 0.07701044 0.04864705 0.06260879 0.03865092 0 0.08330459 0.05972254
## 97.5% 0.08270975 0.05551044 0.07707877 0.04247184 0 0.09284885 0.06895278
                           CT
                                      ΕT
                                                 GT
                ВT
                                                            HT IT
## 2.5% 0.06705311 0.06891383 0.07454346 0.03749383 0.04208555 0 0.06826523
## 97.5% 0.09538258 0.07469844 0.08113913 0.04884414 0.04514029 0 0.07642650
                TT BU CU EU
                                      GU HU IU SU TU
                                                            BV
## 2.5% 0.04131775 0 0 0.0000000000 0 0 0 0.1149178 0.04876068
## 97.5% 0.04764019 0 0 0.0005360972 0
                                             0 0 0 0.1467529 0.05349064
               EV
                          GV
                                     HV
                                               IV
                                                          SV
## 2.5% 0.1216927 0.02050779 0.06992141 0.0000000 0.03599240 0.01819445
## 97.5% 0.1314287 0.02826801 0.07368250 0.1666667 0.04278268 0.02253544
                             CW
                                        EW
                                                   GW
## 2.5% 0.004786727 0.007170102 0.01815913 0.02126852 0.01218079 0.0000000
## 97.5% 0.013314274 0.009219897 0.02118848 0.02814080 0.01413393 0.1666667
                SW
                           TW
                                      BX
                                                    CX EX GX HX IX
## 2.5% 0.01431749 0.00880367 0.000000000 0.0004653034 0 0 0 0.0000000000
## 97.5% 0.01788480 0.01157362 0.003212078 0.0015398320 0 0 0 0.0008245824
                  TX
                             BY
                                        CY
                                                   ΕY
                                                              GY
## 2.5% 0.0000000000 0.05427077 0.02606874 0.05277012 0.04399917 0.02796001 0
## 97.5% 0.0003309154 0.07816867 0.02945590 0.05788738 0.05419491 0.03137121 0
                SY
                           TY
## 2.5% 0.02228816 0.02208981
## 97.5% 0.02715243 0.02646527
CI_Tr <- compute_ci(df[, "Tr"])</pre>
CI_Tr
                                                   GB
                                                                HB IB
                                                                              SB
                           CB
## 2.5% 0.01307514 0.02687477 0.003366287 0.005717791 0.0002712940 0 0.02337039
## 97.5% 0.02385615 0.03098316 0.004500793 0.010183726 0.0006268562 0 0.02851486
                TB
                          BC
                                    CC
                                              EC
                                                        GC
                                                                   HC IC
## 2.5% 0.01620519 0.5895931 0.5033462 0.1045115 0.1059212 0.01680466 0
## 97.5% 0.01984288 0.6278534 0.5264039 0.1118928 0.1188862 0.01883271
                         TC
                                    ΒE
## 2.5% 0.3750835 0.2204611 0.02297971 0.1064347 0.8079827 0.01557574
## 97.5% 0.3899632 0.2330701 0.03813599 0.1164493 0.8170412 0.02187402
                  HE IE
                                SE
                                           ΤE
                                                     BG
## 2.5% 0.0001441769 0 0.08076129 0.06565904 0.01447739 0.02352102 0.003759800
## 97.5% 0.0004647994 0 0.09078146 0.07363596 0.02799887 0.02695228 0.005353479
##
               GG
                           HG IG
                                         SG
                                                    TG
## 2.5% 0.6951336 0.002444814 0 0.01614956 0.01144745 0.01726699 0.07978652
## 97.5% 0.7008745 0.003284930 0 0.02046556 0.01441291 0.03041572 0.08995787
                 EΗ
                            GH
                                      HH
                                                IH
                                                           SH
```

```
## 2.5% 0.004625707 0.02894329 0.9090937 0.0000000 0.06229492 0.03740477
## 97.5% 0.006541229 0.03779066 0.9118856 0.1666667 0.07252884 0.04338309
                                                                             0 0
         0 0.000000e+00 0.8000000 0.0000000000 0.000000000 0.1402201 0.1331952
## 2.5%
## 97.5% 0 5.525887e-05 0.8333333 0.0002077487 0.0001649936 0.1683101 0.1445062
                           GS
                                      HS IS
## 2.5% 0.02726135 0.05192919 0.01517788 0 0.3483722 0.1158081 0.1283118
## 97.5% 0.03122991 0.06381284 0.01735602 0 0.3647437 0.1250890 0.1589700
                CT
                          ET
                                     GT
                                                HT IT
                                                               ST
## 2.5% 0.0919086 0.03409808 0.06587820 0.05021728 0.0 0.05939931 0.5075686
## 97.5% 0.1004361 0.03774051 0.07900966 0.05354480 0.2 0.06787791 0.5162021
```

(f)

Use the following measure to compute the accuracy of the predicted secondary structure $P = (p_i)$ for the data.frame of proteins_test.tsv given the real secondary structure $S = (s_i)$:

$$a(P,S) = \frac{1}{L} \sum_{i} \begin{cases} 1 & \text{if } p_i = s_i \\ 0 & \text{if } p_i \neq s_i \end{cases}$$

with sequence length L. Compute the accuracy for every protein in your data.frame and store the accuracies in a vector. What is the accuracy of the Viterbi algorithm over all sequences (i.e. call summary on the vector of accuracies)?

```
registerDoParallel(num_cores)
accuracy_viterbi <- foreach (i=1:nrow(test), .combine = c, .packages=c("foreach", "tidyverse")) %dopar%
predicted <- predict_test$PredictedStructure[i] %>%
    strsplit("") %>%
    unlist()
    truth <- test$structure[i] %>%
        strsplit("") %>%
        unlist()
    acc <- sum(truth == predicted) / length(predicted)
    return(acc)
}
stopImplicitCluster()
registerDoSEQ()
summary(accuracy_viterbi)</pre>
```

Min. 1st Qu. Median Mean 3rd Qu. Max. ## 0.007752 0.226253 0.322917 0.319801 0.407240 0.857143

(g)

Instead of using the Viterbi algorithm, now randomly guess secondary structures for all sequences. Compare the global accuracies of the Viterbi and the random approach and plot all accuracy distributions using boxplots.

```
set.seed(42)
accuracy_random <- foreach (i=1:nrow(test), .combine = c) %dopar% {
  truth <- test$structure[i] %>%
    strsplit("") %>%
    unlist()
  predicted <- sample(ss_states, size=length(truth), replace = TRUE)
  acc <- sum(truth == predicted) / length(truth)</pre>
```

```
return(acc)
}
summary(accuracy_random)
      Min. 1st Qu. Median
##
                              Mean 3rd Qu.
                                               Max.
## 0.0000 0.1051 0.1213 0.1225 0.1398 0.2857
boxplot(accuracy_random, accuracy_viterbi,
        ylab = 'Accuracy', names = c('random', 'viterbi'))
                                                                0
     0.8
     9.0
Accuracy
     0.4
                              0
     0.2
     0.0
                              0
                                                             viterbi
                          random
```