## BMI 651

Joshua Burkhart February 29, 2016

## **HW5: APPENDIX**

1

**1A** 

Transition matrix for two-state model:

```
AT = c(.85,.20)

GC = c(.15,.80)

df = data.frame(AT,GC)

row.names(df) <- c("AT","GC")

kable(df,row.names=TRUE)
```

	AT	GC
AT	0.85	0.15
GC	0.20	0.80

1B

1Bi

Emission matrix for AT-rich regions:

```
P = c(.30,.10,.20,.40)
df = data.frame(P)
row.names(df) <- c("A","C","G","T")
kable(df,row.names=TRUE)</pre>
```

```
\begin{array}{c|c} & P \\ \hline A & 0.3 \\ C & 0.1 \\ G & 0.2 \\ T & 0.4 \\ \end{array}
```

1Bii

Emission matrix for GC-rich regions:

```
P = c(.05,.30,.55,.10)
df = data.frame(P)
row.names(df) <- c("A","C","G","T")
kable(df,row.names=TRUE)</pre>
```

 $\begin{array}{c|c} & P \\ \hline A & 0.05 \\ C & 0.30 \\ G & 0.55 \\ T & 0.10 \\ \end{array}$ 

1C

```
GenerateNucSeq <- function(num_nucs)</pre>
  #set initial state using a uniform random distribution
  state <- sample(x=c("AT", "GC"), size=1)</pre>
  nuc_seq <- character()</pre>
  for(i in 1:num_nucs)
    if(state == "AT"){
      state <- sample(x=c("AT","GC"),size=1,prob=TM[1,])</pre>
      nuc <- sample(x=c("A","C","G","T"),size=1,prob=AT_EM)</pre>
    }
    else if(state == "GC")
      state <- sample(x=c("AT", "GC"), size=1, prob=TM[2,])</pre>
      nuc <- sample(x=c("A","C","G","T"),size=1,prob=GC_EM)</pre>
    }
    nuc_seq <- c(nuc_seq,nuc)</pre>
  return(nuc_seq)
}
```

## GenerateNucSeq(100)

## 1D

```
AT_DICT <- list(A=.30,
                 C = .10,
                 G = .20.
                 T=.40)
GC_DICT <- list(A=.05,</pre>
                 C=.30,
                 G = .55.
                 T=.10)
GenerateStateSeq <- function(nuc_seq)</pre>
  at_state_prob <- 0.5 * AT_DICT[[nuc_seq[1]]] # P(nuc_seq[1]/AT)
  gc_state_prob <- 0.5 * GC_DICT[[nuc_seq[1]]] # P(nuc_seq[1]/GC)</pre>
  for(i in 2:length(nuc_seq))
    # AT
    at_state_prob <- c(at_state_prob,</pre>
                            \# P(AT) * P(AT/AT) * P(qi/AT)
                        max(at_state_prob[i-1] * TM[1,1] * AT_DICT[[nuc_seq[i]]],
                            \# P(GC) * P(AT/GC) * P(qi/AT)
                            gc_state_prob[i-1] * TM[2,1] * AT_DICT[[nuc_seq[i]]]))
    # GC
    gc_state_prob <- c(gc_state_prob,</pre>
                            \# P(GC) * P(GC/GC) * P(qi/GC)
                        max(gc_state_prob[i-1] * TM[2,2] * GC_DICT[[nuc_seq[i]]],
                            \# P(AT) * P(GC/AT) * P(qi/GC)
                            at_state_prob[i-1] * TM[1,2] * GC_DICT[[nuc_seq[i]]]))
  state_trace <- character()</pre>
  for(j in 1:length(nuc_seq))
    state_trace <- c(state_trace,ifelse(at_state_prob[j] > gc_state_prob[j],"AT","GC"))
 return(state_trace)
}
GenerateStateSeq(c("A", "A", "G", "C", "G", "T", "G", "G", "G", "G",
```

```
"C", "C", "C", "C", "G", "G", "C", "G", "A", "C",
"A", "T", "G", "G", "G", "T", "G", "T", "C"))
```

- [1] "AT" "AT" "AT" "GC" "GC" "AT" "GC" "GC" "GC" "GC" "GC" "GC" "GC"
- [15] "GC" "GC" "GC" "GC" "AT" "GC" "AT" "AT" "AT" "GC" "GC" "GC" "GC"
- [29] "AT" "GC"