Genome Sequence Analysis: Assignment 1

University of Cambridge

Henrik Åhl

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Preface

This is an assignment report in connection to the *Genome Sequence Analysis* module in the Computational Biology course at the University of Cambridge, Michaelmas term 2016. All related code is as of January 27, 2017 available per request by contacting hpa22@cam.ac.uk. The approach is done after the guidelines specified by Rabiner [1], and as summarized by Shen [2]. All code is implemented using Python, and in particular by using the numpy package.

Exercises

```
# Get index for letter (symbol) occurring.
  def get_ind(probs):
      rand = random.random()
      for ii in xrange(len(probs)):
          if rand < probs[ii]:
              return ii
  # Read in transition matrix and initial distribution from files.
  # If sumrows == TRUE, sum the rows
  def read_and_output(mat_file, idist_file, seq_length, sumrows, transpose):
      # Read in data
      trans_mat = np.loadtxt(open(mat_file, "r"))
      init_dist = np.loadtxt(open(idist_file, "r"))
      # Act on the optional things
      if transpose:
          trans_mat = np.transpose(trans_mat)
      if sumrows:
19
          trans_mat = np.cumsum(trans_mat, 1)
21
      # Fill out the output probabilistically
      output = seq_length * [None]
      output[0] = np.random.choice(np.arange(0, len(trans_mat)), p=list(init_dist))
      for ii in xrange(1, seq_length):
          output[ii] = get_ind(trans_mat[output[ii - 1], :])
      "".join(str(s) for s in output)
      return output
  mat_file = "/home/henrik/compbio/src/assignments/gsa1/report/mat_file.dat"
  idist_file = "/home/henrik/compbio/src/assignments/gsa1/report/idist_file.dat"
  output = read_and_output(mat_file, idist_file, 115, True, False)
```

Evoking the print command we get the following:

Our program is thus able to produce an output corresponding to that of a Markov chain.

2

For this exercise, as we are only to read a single Markov chain, the initial distribution will correspond to a 100 % probability of ending up in the first state in the sequence.

```
# Get transtion matrix and initial distribution from one (!) sequence
 def infer_mat_and_init (sequence):
     # Get some data and create the matrices we need.
     elements = np.unique(list(sequence))
     trans\_mat = np.zeros((len(elements), len(elements)))
     init\_dist = np.zeros(len(elements)) # This is specific to this assignment
6
     init_dist[int(sequence[0])] = 1 # Count the first one
     # Count the occurrences of all events
     for ii in xrange(1, len(sequence))
         trans_mat[int(sequence[ii - 1]), int(sequence[ii])] += 1
     # What are the probabilities?
     for ss in xrange(len(trans_mat)):
        trans_mat[ss,] /= np.sum(trans_mat[ss,])
     return trans_mat, init_dist
 trans_mat, init_dist = infer_mat_and_init("
```

Again, looking at the inferred transfer matrix and initial distribution tells us that our program appears to give an output corresponding to our expectations:

$$A = \begin{pmatrix} 0.15 & 0.85 \\ 0.85 & 0.15 \end{pmatrix}, \quad \ \mu^0 = \begin{pmatrix} 1 \\ 0 \end{pmatrix}.$$

```
# Parameters
  S = \text{np.array}([0, 1])

V = \text{np.array}([0, 1, 2, 3, 4])
  A = np.matrix("0.8, 0.2;"
    "0.1, 0.9")
= np.matrix("0.2, 0.5, 0.2, 0.1, 0.0;"
                     "0.0, 0.1, 0.4, 0.4, 0.1")
8 \text{ mu0} = \text{np.array}([.5, .5])
   # \it Emit a sequence given some transition and emission matrices, as well as
    an initial distribution.
   def emit_sequence(seq_length, trans_matrix, emiss_matrix, init_dist):
        c_trans_mat = np.cumsum(trans_matrix, 1)
        c_emiss_mat = np.cumsum(emiss_matrix,
        emit = np.zeros(seq_length)
        hidden = np.zeros(seq_length)
18
        hidden[0] = np.random.choice(np.arange(0, len(trans_matrix)), p=list(init_dist))
        \operatorname{emit}[0] = \operatorname{get\_ind}((\operatorname{c\_emiss\_mat}[\operatorname{int}(\operatorname{hidden}[0]), :]) . \operatorname{tolist}()[0])
```

```
# Parse sequence
for ii in xrange(1, seq_length):
    hidden[ii] = get_ind((c_trans_mat[int(hidden[ii - 1]), :]).tolist()[0])
    emit[ii] = get_ind((c_emiss_mat[int(hidden[ii]), :]).tolist()[0])

emit = "".join(str(int(x)) for x in emit)
    hidden = "".join(str(int(x)) for x in hidden)
    return emit, hidden

emit, hidden = emit_sequence(115, A, B, mu0)
```

The result of our emitted sequence, as well as the underlying state, can be seen in fig. 3, and appears to correspond well to the expected behaviour.

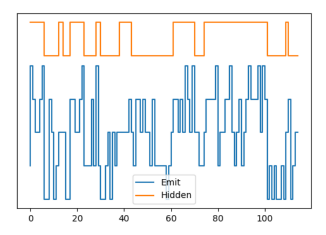


Figure 1: Output from the program emitting a sequence according to the designated parameters. 0-valued states tend to produce the lowest emission values on average, whereas the 1-valued states behave in the opposite manner.

4

We define a method for reading in a sequence from a file (which has been produced with our previously defined emission function).

```
# Read in sequence from file.
  def read_sequence(file, skipFirst):
      seq = "
      with open(file, 'r') as in_file:
          if skipFirst:
              in_file.readline()
          seq = in_file.read().replace('\n', '')
      return seq
  # Use the forward algorithm to determine (log) likelihood of sequence
  def forward(sequence, trans_matrix, emiss_matrix, hidden_states, init_dist):
12
      # Function for getting normalisation constant
      def get_constant(a):
14
          sum = np.sum(a)
          return 1.0 / len(trans_matrix) if sum == 0 else 1.0 / sum
      # Pre-allocate
18
      fw = np.zeros((len(hidden_states), len(sequence)))
      constants = []
20
```

```
# Initialise
      for ss in xrange(len(hidden_states)):
          fw[ss, 0] = init_dist[ss] * emiss_matrix[ss, int(sequence[0])]
      const = get\_constant(fw[0, :])
      constants.append(const)
      # Scale accordingly
      for ss in xrange(len(hidden_states)):
          fw[ss, 0] *= const
      # Now go through the rest of the sequence
      probs = np.zeros(len(hidden_states))
      for ii in xrange(1, len(sequence)):
          for ss in xrange(len(hidden_states)):
              probs[ss] = np.sum(
36
                  (fw[substate, ii - 1] * trans_matrix[substate, ss] * emiss_matrix[ss, int(
                      sequence[ii])]) for
                  substate in xrange(len(hidden_states)))
38
          # Scale again
          const = get_constant(probs)
          constants.append(const)
          for ss in xrange(len(hidden_states)):
              fw[ss, ii] = const * probs[ss]
      # Sum up the probabilities; disregard if O
46
      ln_prob = -np.sum([math.log(const) if const != 0 else 0 for const in constants])
      return ln_prob, constants, fw
  out_sequence, __ = emit_sequence(115, A, B, mu0)
  seq_file = '/home/henrik/compbio/src/assignments/gsa1/random_output_sequence.dat'
  with open(seq_file, "w") as f_out:
      f_out.write(out_sequence)
      f_out.close()
  that_same_sequence = read_sequence(seq_file, False)
prob, consts, fw = forward(that_same_sequence, A, B, S, mu0)
```

5

We use the *Saccharomyces cerevisiae* chromosome III genome from Ensembl [3]. The genome is binned by taking intervals of size 100, and subsequently identifying how many G or C bases are contained. In our implementation, bins are sequential and do not overlap.

Trying to approximate the correct distribution of categories, we label bins after percentage-wise GC content. Doing this rough fitting, the best fit appears to be found around 0-28.5%, 28.5-34.2%, 34.2-40.0%, 40.0-50.0% and a 50-100% split for label 1-5 respectively, with upper boundaries exclusive. As our genome is not evenly divisble by the bin size, the last bin will be misrepresentative. However, as our genome is large, and we thus will also have a large set of bins, the effects of this will be marginal. Thence we choose to disregard this effect.

```
G = bin.count("G")
          C = bin.count("C")
          gc_cont.append((G + C) / bin_size)
      return gc_cont
  # Read genome
  sc_file = '/home/henrik/compbio/src/assignments/gsa1/sc_gen.fa'
  sc_gen = read_sequence(sc_file , True)
     = calc_gc(sc_gen, 100)
  # Relabel according to predefined cuts
24
  def relabel (seq, cuts):
      relab = []
      for ii in seq:
          for cut in xrange(len(cuts)):
               if ii \le cuts[cut]:
                  relab.append(cut)
30
                   break
      relab = "".join(str(x) for x in relab)
      return relab
  # Compare the two strings
  relab = relabel(gc, [.285, .342, .40, .5, 1.]) # Split by percentages
  model3_seq , model3_hidden = emit_sequence(len(relab), A, B, mu0)
  log_p, const, fw_seq = forward(relab, A, B, S, mu0)
```

The final classification can be seen in fig. 2, where the two distributions are shown.

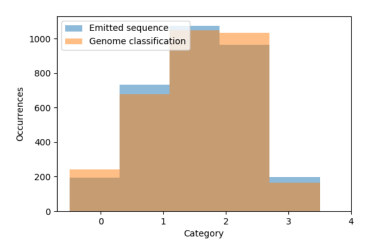


Figure 2: Distributions of emission values according to our given model and of the *Saccharomyces cerevisiae* genome after percentage-wise labelling

```
# We only use this for Baum-Welch, so we use the same constants as in our forward case
def backward(sequence, trans_matrix, emiss_matrix, hidden_states, init_dist, constants):
# Preallocate for our backwards sequence
bw = np.zeros((len(hidden_states), len(sequence)))
# Initialise
# Initialise
```

```
for ss in xrange(len(hidden_states)):
          bw[ss, len(sequence) - 1] = 1.
          bw[ss, len(sequence) - 1] = constants[len(sequence) - 1]
9
      # Go through the rest of the sequence, continue from the back end
11
      for ii in xrange(len(sequence) - 2, -1, -1):
          b = np.zeros(len(hidden_states))
          for ss in xrange(len(hidden_states)):
              b\,[\,ss\,]\,=\,np.sum\,(\,[\,(\,bw[\,substate\,,\ ii\,\,+\,\,1\,]\,\,*\,\,trans\_matrix\,[\,ss\,,\ substate\,]\,\,*
                                emiss_matrix[substate, int(sequence[ii + 1])]) for substate in
                                   xrange(len(hidden_states))])
          # Again, scale
          for ss in xrange(len(hidden_states)):
19
              bw[ss, ii] = constants[ii] * b[ss]
21
      ln_prob = -np.sum([math.log(c) if c != 0 else 0 for c in constants])
      return ln_prob, bw
  # Now we can go onto estimating our matrices
  {\tt def\ baum\_welch}({\tt sequence}\ ,\ {\tt trans\_matrix}\ ,\ {\tt emiss\_matrix}\ ,\ {\tt hidden\_states}\ ,\ {\tt init\_dist}):
27
      # Get those lists we need
      pl, constants, fw = forward(sequence, trans_matrix, emiss_matrix, hidden_states,
29
          init_dist)
      p1, bw = backward(sequence, trans_matrix, emiss_matrix, hidden_states, init_dist,
          constants)
31
      # Initial distribution
      for ss in xrange(len(hidden_states)):
          init_dist[ss] = fw[ss, 0] * constants[0] * bw[ss, 0]
      init_dist /= np.sum(init_dist)
      # New state matrix
      for ss in xrange(len(hidden_states)):
          for substate in xrange(len(hidden_states)):
              # Sum up the estimate, normalise and assign
              new_est = np.sum([(fw[ss, ii] * bw[substate, ii + 1] * trans_matrix[ss, substate]
41
                    * emiss_matrix[
                  substate, int(sequence[ii + 1])]) for ii in xrange(len(sequence) - 1)])
              sequence) - 1))
               trans_matrix[ss, substate] = new_est / norm if norm != 0 else 1. / len(
                  hidden_states)
      # New emission matrix (structure same as above)
      for ss in xrange(len(hidden_states)):
          for jj in xrange(emiss_matrix.shape[1]):
              new\_est = np.sum([(fw[ss , ii] * bw[ss , ii] / constants[ii] if int(sequence[ii]))
49
                  == jj else 0) for ii in
                                 xrange(len(sequence))])
              norm = np.sum([(fw[ss, ii] * bw[ss, ii] / constants[ii]) for ii in xrange(len(
                  sequence))])
              emiss_matrix[ss, jj] = new_est / norm if norm != 0 else 1. / emiss_matrix.shape
                  [1]
      return p1
  prev = change
  while change > 0.000001:
      prob = baum_welch(relab, A, B, S, mu0)
      change = prev - prob
59
      prev = prob
```

We here achieve a steady increase in the log likelihood for the given sequence, showing that our model adapts

to the data given. Ultimately, our matrices take the shapes of

$$A = \begin{pmatrix} 0.91 & 0.09 \\ 0.05 & 0.95 \end{pmatrix}, \qquad B = \begin{pmatrix} 0.01 & 0.05 & 0.21 & 0.59 & 0.14 \\ 0.11 & 0.31 & 0.40 & 0.18 & 0.00 \end{pmatrix}.$$

Our final log likelihood for the same situation evaluates as:

```
-4251.17034431
```

```
1 # Retrieve the most likely input hidden states given and output sequence and
    the corresponding matrices.
  def viterbi(sequence, trans_matrix, emiss_matrix, hidden_states, init_dist):
       # Define some things we're gonna use.
       constants = np.zeros(len(sequence))
       solution = np.zeros(len(sequence))
       dyn_mat = np.zeros((len(hidden_states), len(sequence)))
       store_path = np.zeros((len(hidden_states), len(sequence)))
9
       # Initialise rows
       \label{eq:dyn_mat} \texttt{dyn\_mat}\left[:\,,\ 0\right] \;=\; \texttt{init\_dist}\;\; *\;\; \texttt{emiss\_matrix}\left[:\,,\;\; \texttt{int}\left(\texttt{sequence}\left[0\right]\right)\right]
       constants [0] = 1.0 / \text{np.sum}(\text{dyn\_mat}[:, 0])
       dyn_mat[:, 0] *= constants[0]
13
       # Fill out the table
       for ii in xrange(1, len(sequence)):
            for ss in xrange(len(hidden_states)):
17
                 probabilities = pp.diag(dyn\_mat[:, ii - 1]) * trans\_matrix[:, ss] * \textit{Probability}
                     of coming from either state
                 store_path[ss, ii], dyn_mat[ss, ii] = max(enumerate(probabilities), key=
                     itemgetter(1))
                 dyn_mat[ss, ii] *= emiss_matrix[ss, int(sequence[ii])]
            # Now we scale
            constants [\,ii\,] \,=\, 1.0 \,\,/\,\, np.sum \left( [\,dyn\_mat\,[\,ss\,,\,\,ii\,] \,\,for\,\,ss\,\,in\,\,xrange \left( \,len \left( \,hidden\_states \,\right) \,\right) \,\right] \right)
            dyn_mat[:, ii] *= constants[ii]
25
       # Backtrack
       solution [len(sequence) - 1] = dyn_mat[:, len(sequence) - 1].argmax() # last state
       for ii in xrange(len(sequence) - 1, 0, -1):
            solution [ii - 1] = store_path [int(solution [ii]), ii]
       return solution
31
   model7\_seq = sc\_gen
gc = calc_gc(sc_gen, 100)
37 intab = "ATCG"
   outtab = "0123"
  trantab = maketrans(intab, outtab)
   model7_seq = model7_seq.translate(trantab)
41
   43 prev = change
   while change > 0.000001:
       prob = baum_welch(relab, A, B, S, mu0)
       change = prev - prob
47
       prev = prob
solution = viterbi(relab, A, B, S, mu0)
```

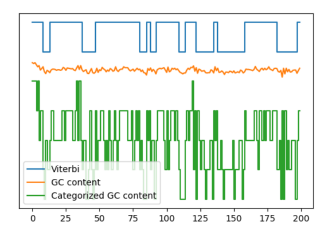


Figure 3: Inferred underlying hidden state sequence using the Viterbi algorithm, along with the corresponding categorized and non-categorized GC content under the previously determined binning scheme. Notably, the 1-valued states correspond well with the highest gc content, as expected from our matrices. Similarly, the lower state more consistently is affiliated with lower emission values.

8

Categorise according to a gaussian distribution with different means?