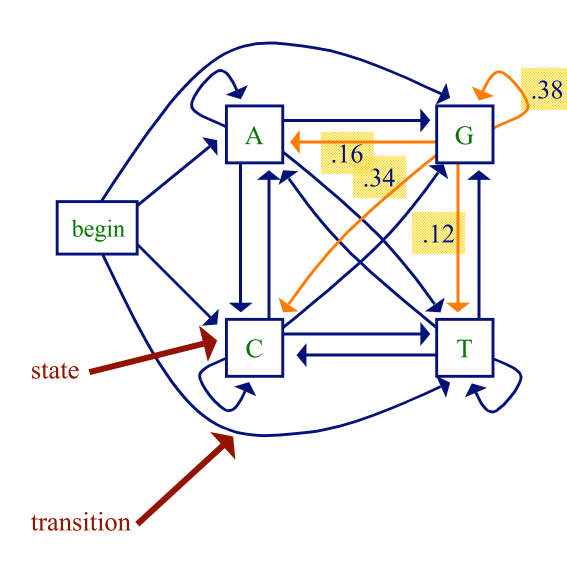
Markov Chain Models

(Slides courtesy of Dr. Mark Craven)

Motivation for Markov Models in Computational Biology

- there are many cases in which we would like to represent the statistical regularities of some class of sequences
 - genes
 - various regulatory sites in DNA (e.g. where RNA polymerase and transcription factors bind)
 - proteins in a given family
- Markov models are well suited to this type of task

A Markov Chain Model



transition probabilities

$$Pr(x_i = a \mid x_{i-1} = g) = 0.16$$

$$Pr(x_i = c \mid x_{i-1} = g) = 0.34$$

$$Pr(x_i = g \mid x_{i-1} = g) = 0.38$$

$$Pr(x_i = t \mid x_{i-1} = g) = 0.12$$

Markov Chain Models

- a Markov chain model is defined by
 - a set of states
 - some states *emit* symbols
 - other states (e.g. the *begin* state) are *silent*
 - a set of transitions with associated probabilities
 - the transitions emanating from a given state define a distribution over the possible next states

Markov Chain Models

- given some sequence x of length L, we can ask how probable the sequence is given our model
- for any probabilistic model of sequences, we can write this probability as

$$Pr(x) = Pr(x_L, x_{L-1}, ..., x_1)$$

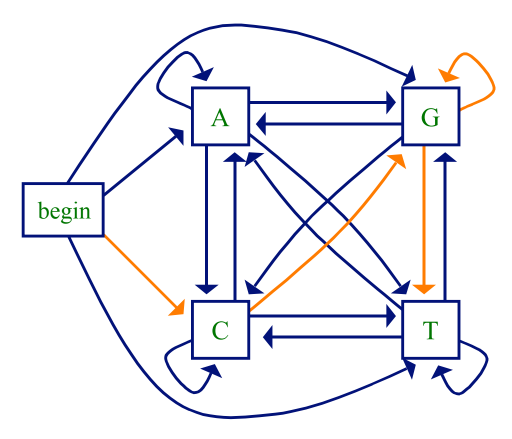
$$= Pr(x_L | x_{L-1}, ..., x_1) Pr(x_{L-1} | x_{L-2}, ..., x_1) ... Pr(x_1)$$

• key property of a (1st order) Markov chain: the probability of each X_i depends only on the value of X_{i-1}

$$Pr(x) = Pr(x_{L}|x_{L-1}) Pr(x_{L-1} | x_{L-2}) ... Pr(x_{2} | x_{1}) Pr(x_{1})$$

$$= Pr(x_{1}) \prod_{i=2}^{L} Pr(x_{i} | x_{i-1})$$

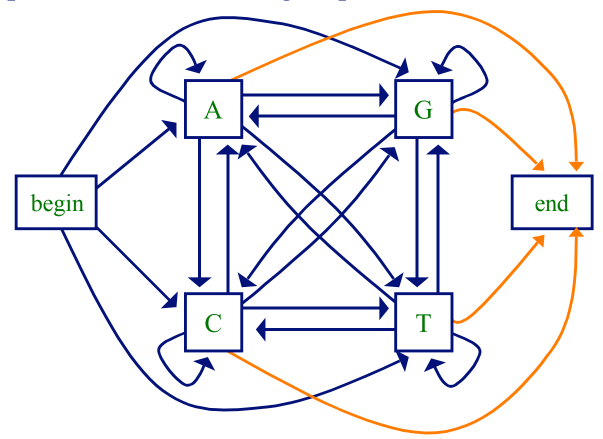
The Probability of a Sequence for a Given Markov Chain Model



Pr(cggt) = Pr(c) Pr(g | c) Pr(g | g) Pr(t|g)

Markov Chain Models

- can also have an *end* state; allows the model to represent
 - a distribution over sequences of different lengths
 - preferences for ending sequences with certain symbols



Markov Chain Notation

• the transition parameters can be denoted by $a_{x_{i-1}x_i}$ where

$$a_{x_{i-1}x_i} = \Pr(x_i \mid x_{i-1})$$

• similarly we can denote the probability of a sequence x as

$$a_{Bx_1} \prod_{i=2}^{L} a_{x_{i-1}x_i} = \Pr(x_1) \prod_{i=2}^{L} \Pr(x_i \mid x_{i-1})$$

where a_{Bx_1} represents the transition from the *begin* state

Example Application

- CpG islands
 - CG dinucleotides are rarer in eukaryotic genomes than expected given the marginal probabilities of C and G
 - but the regions upstream of genes are richer in CG dinucleotides than elsewhere – CpG islands
 - useful evidence for finding genes
- could predict CpG islands with Markov chains
 - one to represent CpG islands
 - one to represent the rest of the genome

Estimating the Model Parameters

- given some data (e.g. a set of sequences from CpG islands), how can we determine the probability parameters of our model?
- one approach: *maximum likelihood estimation*
 - given a set of data D
 - set the parameters θ to maximize

$$Pr(D | \theta)$$

– i.e. make the data D look likely under the model

Maximum Likelihood Estimation

- suppose we want to estimate the parameters Pr(a), Pr(c), Pr(g), Pr(t)
- and we're given the sequences

```
accgcgctta
```

gcttagtgac

tagccgttac

• then the maximum likelihood estimates are

$$Pr(a) = \frac{6}{30} = 0.2 Pr(g) = \frac{7}{30} = 0.233$$

$$Pr(c) = \frac{9}{30} = 0.3 Pr(t) = \frac{8}{30} = 0.267$$

Maximum Likelihood Estimation

• suppose instead we saw the following sequences

```
gccgcgcttg
gcttggtggc
tggccgttgc
```

• then the maximum likelihood estimates are

$$\Pr(a) = \frac{0}{30} = 0$$

$$\Pr(c) = \frac{9}{30} = 0.3$$

$$\Pr(t) = \frac{8}{30} = 0.267$$

do we really want to set this to 0?

A Bayesian Approach

- instead of estimating parameters strictly from the data, we could start with some prior belief for each
- for example, we could use *Laplace estimates*

$$Pr(a) = \frac{n_a + 1}{\sum_{i} (n_i + 1)}$$
 pseudocount

- where n_i represents the number of occurrences of character i
- using Laplace estimates with the sequences

gccgcgcttg
gcttggtggc
$$Pr(a) = \frac{0+1}{34}$$
tggccgttgc
$$Pr(c) = \frac{9+1}{34}$$

A Bayesian Approach

• a more general form: *m-estimates*

$$Pr(a) = \frac{n_a + p_a m}{\left(\sum_{i} n_i\right) + m}$$
 prior probability of a number of "virtual" instances

• with m=8 and uniform priors

gccgcgcttg

gcttggtggc
tggccgttgc

$$Pr(c) = \frac{9 + 0.25 \times 8}{30 + 8} = \frac{11}{38}$$

Estimation for 1st Order Probabilities

- to estimate a 1st order parameter, such as Pr(c|g), we count the number of times that g follows the history c in our given sequences
- using Laplace estimates with the sequences

gccgcgcttg gcttggtggc tggccgttgc

$$Pr(a \mid g) = \frac{0+1}{12+4} \quad Pr(a \mid c) = \frac{0+1}{7+4}$$

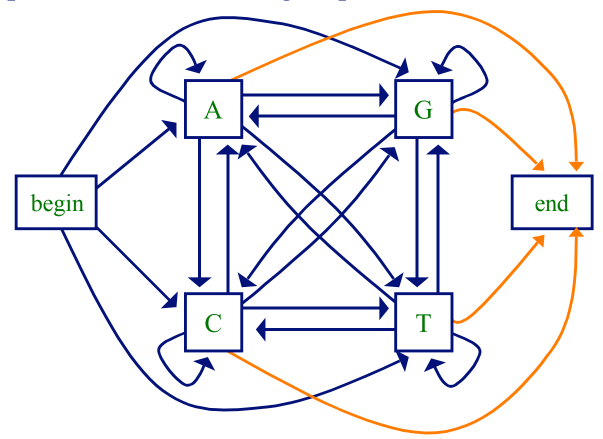
$$Pr(c \mid g) = \frac{7+1}{12+4} \quad M$$

$$Pr(g \mid g) = \frac{3+1}{12+4}$$

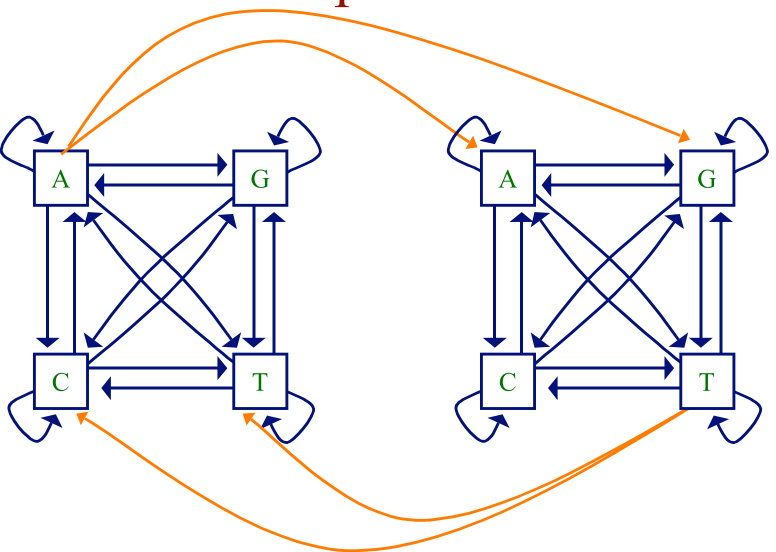
$$Pr(t \mid g) = \frac{2+1}{12+4}$$

Markov Chain Models

- can also have an *end* state; allows the model to represent
 - a distribution over sequences of different lengths
 - preferences for ending sequences with certain symbols



A Simple HMM



• given say a *T* in our input sequence, which state emitted it?

Hidden State

- we'll distinguish between the *observed* parts of a problem and the *hidden* parts
- in the Markov models we've considered previously, it is clear which state accounts for each part of the observed sequence
- in the model above, there are multiple states that could account for each part of the observed sequence
 - this is the hidden part of the problem

The Parameters of an HMM

• as in Markov chain models, we have transition probabilities

$$a_{kl} = \Pr(\pi_i = l \mid \pi_{i-1} = k)$$

probability of a transition from state k to l

 π represents a path (sequence of states) through the model

 since we've decoupled states and characters, we might also have emission probabilities

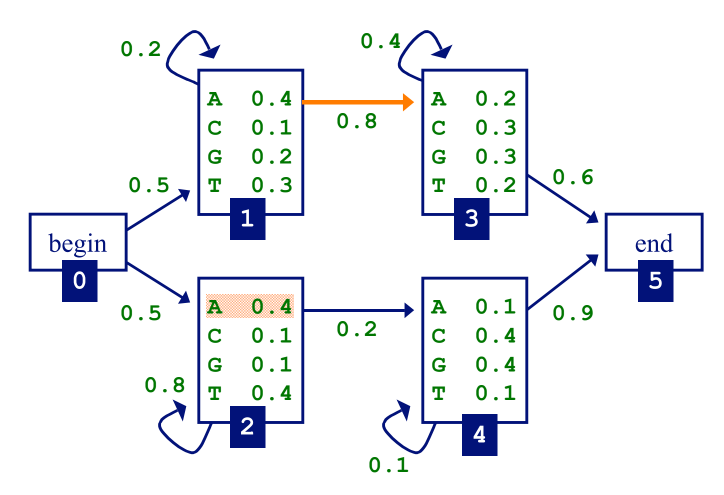
$$e_k(b) = \Pr(x_i = b \mid \pi_i = k)$$

probability of emitting character b in state k

A Simple HMM

 a_{13} probability of a transition from state 1 to state 3

 $e_2(A)$ probability of emitting character A in state 2



Three Important Questions

- How likely is a given sequence? the Forward algorithm
- What is the most probable "path" for generating a given sequence?
 - the Viterbi algorithm
- How can we learn the HMM parameters given a set of sequences?
 - the Forward-Backward (Baum-Welch) algorithm

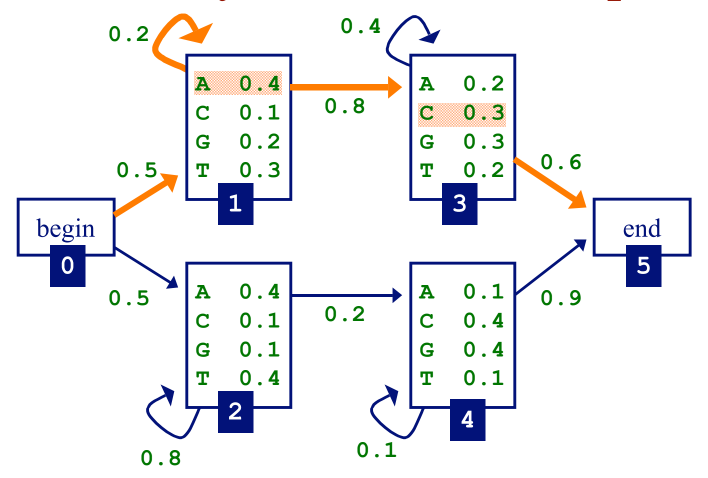
How Likely is a Given Sequence?

• the probability that the path is generated: $\chi_1 \dots \chi_I$

 π_0 is taken and the sequence

$$\Pr(x_1...x_L, \pi_0...\pi_N) = a_{0\pi_1} \prod_{i=1}^L e_{\pi_i}(x_i) a_{\pi_i\pi_{i+1}}$$
 (assuming begin/end are the only silent states on path)

How Likely Is A Given Sequence?



$$Pr(AAC,\pi) = a_{01} \times e_1(A) \times a_{11} \times e_1(A) \times a_{13} \times e_3(C) \times a_{35}$$
$$= 0.5 \times 0.4 \times 0.2 \times 0.4 \times 0.8 \times 0.3 \times 0.6$$

How Likely is a Given Sequence?

• the probability over *all* paths is:

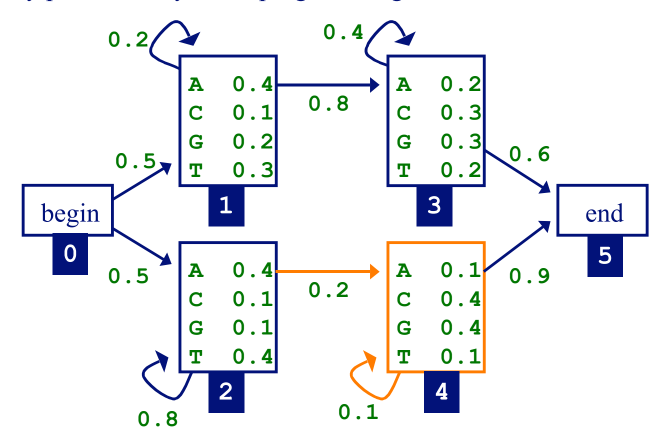
$$Pr(x_1...x_L) = \sum_{\pi} Pr(x_1...x_L, \pi_0...\pi_N)$$

- but the number of paths can be exponential in the length of the sequence...
- the Forward algorithm enables us to compute this efficiently

How Likely is a Given Sequence: The Forward Algorithm

- define $f_k(i)$ to be the probability of being in state k having observed the first i characters of x
- we want to compute $f_N(L)$, the probability of being in the end state having observed all of x
- can define this recursively

• because of the Markov property, don't have to explicitly enumerate every path – use dynamic programming instead



• e.g. compute $f_4(i)$ using $f_2(i-1)$, $f_4(i-1)$

• initialization:

$$f_0(0) = 1$$

probability that we're in start state and have observed 0 characters from the sequence

$$f_k(0) = 0$$
, for k that are not silent states

• recursion for emitting states (i=1...L):

$$f_l(i) = e_l(i) \sum_k f_k(i-1) a_{kl}$$

recursion for silent states:

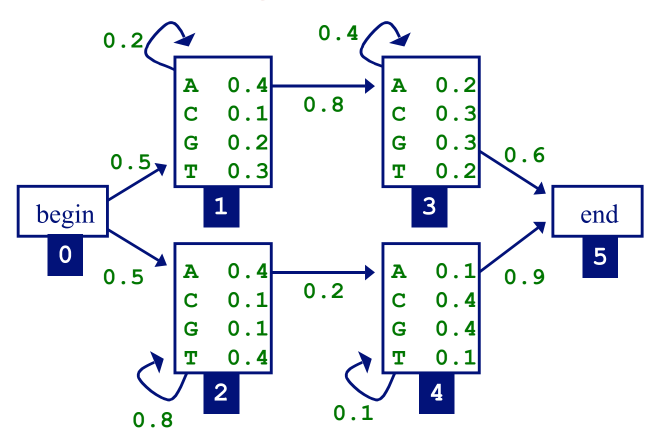
$$f_l(i) = \sum_k f_k(i) a_{kl}$$

• termination:

$$Pr(x) = Pr(x_1...x_L) = f_N(L) = \sum_k f_k(L)a_{kN}$$

probability that we're in the end state and have observed the entire sequence

Forward Algorithm Example



• given the sequence x = TAGA

Forward Algorithm Example

- given the sequence x = TAGA
- initialization

$$f_0(0) = 1$$
 $f_1(0) = 0$ K $f_5(0) = 0$

• computing other values

$$f_{1}(1) = e_{1}(T) \times (f_{0}(0) \times a_{01} + f_{1}(0)a_{11}) =$$

$$0.3 \times (1 \times 0.5 + 0 \times 0.2) = 0.15$$

$$f_{2}(1) = 0.4 \times (1 \times 0.5 + 0 \times 0.8)$$

$$f_{1}(2) = e_{1}(A) \times (f_{0}(1) \times a_{01} + f_{1}(1)a_{11}) =$$

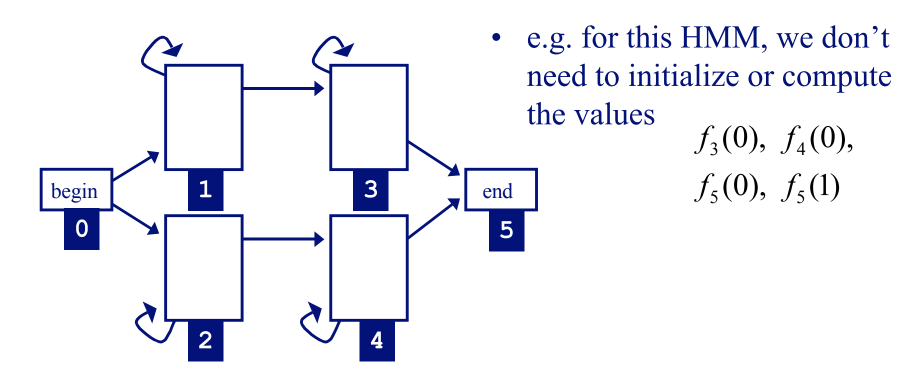
$$0.4 \times (0 \times 0.5 + 0.15 \times 0.2)$$

$$\bullet \bullet \bullet$$

$$Pr(TAGA) = f_{5}(4) = (f_{3}(4) \times a_{35} + f_{4}(4)a_{45})$$

Forward Algorithm Note

• in some cases, we can make the algorithm more efficient by taking into account the minimum number of steps that must be taken to reach a state



Three Important Questions

- How likely is a given sequence?
- What is the most probable "path" for generating a given sequence?
- How can we learn the HMM parameters given a set of sequences?

Finding the Most Probable Path: The Viterbi Algorithm

• define $v_k(i)$ to be the probability of the <u>most probable</u> <u>path</u> accounting for the first *i* characters of *x* and ending in state *k*

- we want to compute $v_N(L)$, the probability of the most probable path accounting for all of the sequence and ending in the end state
- can define recursively
- can use DP to find $v_N(L)$ efficiently

Finding the Most Probable Path: The Viterbi Algorithm

• initialization:

$$v_0(0) = 1$$

 $v_k(0) = 0$, for k that are not silent states

The Viterbi Algorithm

• recursion for emitting states (i=1...L):

$$v_{l}(i) = e_{l}(x_{i}) \max_{k} \left[v_{k}(i-1)a_{kl}\right]$$

$$ptr_{l}(i) = \arg\max_{k} \left[v_{k}(i-1)a_{kl}\right] \text{ keep track of most probable path}$$

recursion for silent states:

$$v_{l}(i) = \max_{k} \left[v_{k}(i) a_{kl} \right]$$
$$ptr_{l}(i) = \arg\max_{k} \left[v_{k}(i) a_{kl} \right]$$

The Viterbi Algorithm

• termination:

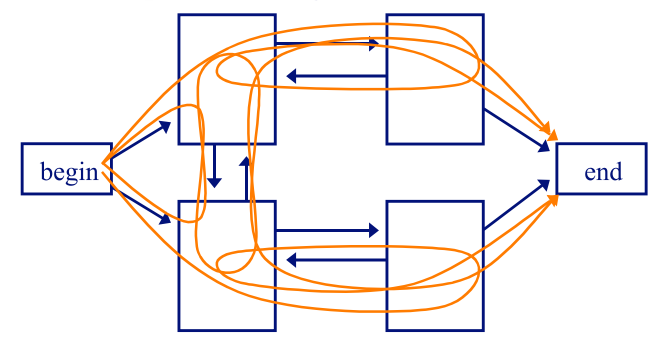
$$Pr(x,\pi) = \max_{k} (v_k(L)a_{kN})$$

$$\pi_L = \arg\max (v_k(L)a_{kN})$$

• traceback: follow pointers back starting at $\pi_{\rm L}$

Forward & Viterbi Algorithms

- Forward/Viterbi algorithms effectively consider all possible paths for a sequence
 - Forward to find probability of a sequence
 - Viterbi to find most probable path
- consider a sequence of length 4...



Three Important Questions

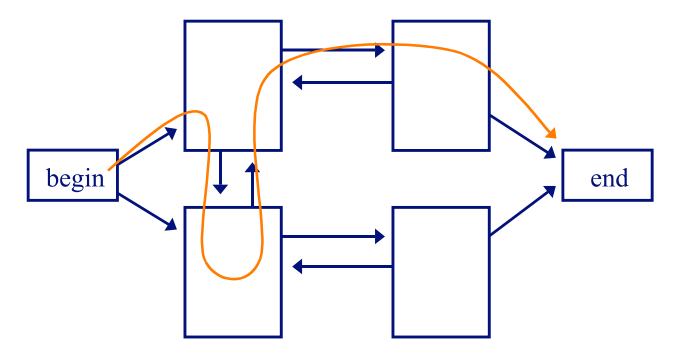
- How likely is a given sequence?
- What is the most probable "path" for generating a given sequence?
- How can we learn the HMM parameters given a set of sequences?

Learning Parameters

- if we know the state path for each training sequence, learning the model parameters is simple
 - no hidden state during training
 - count how often each parameter is used
 - normalize/smooth to get probabilities
 - process is just like it was for Markov chain models
- if we <u>don't</u> know the path for each training sequence, how can we determine the counts?
 - key insight: estimate the counts by considering every path weighted by its probability

Learning without Hidden State

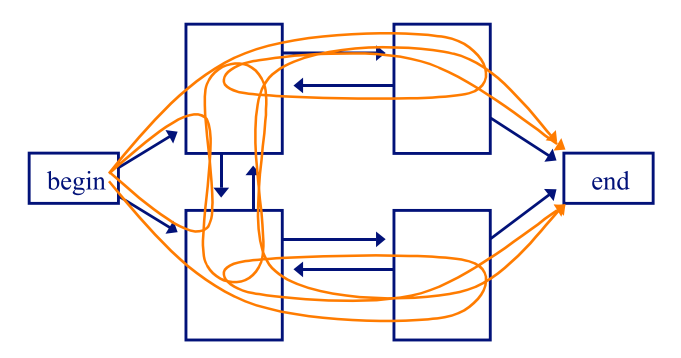
• learning is simple if we know the correct path for each sequence in our training set



• estimate parameters by counting the number of times each parameter is used across the training set

Learning with Hidden State

• if we don't know the correct path for each sequence in our training set, consider all possible paths for the sequence



• estimate parameters through a procedure that counts the expected number of times each parameter is used across the training set

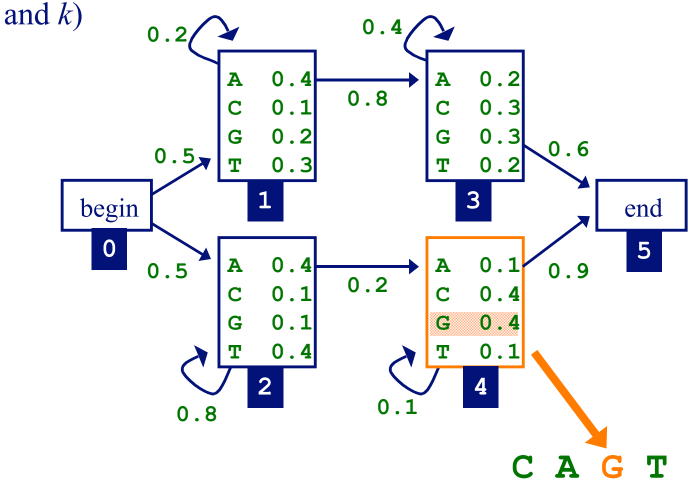
Learning Parameters: The Baum-Welch Algorithm

- *a.k.a* the Forward-Backward algorithm
- an Expectation Maximization (EM) algorithm
 - EM is a family of algorithms for learning probabilistic models in problems that involve hidden state
- in this context, the hidden state is the path that best explains each training sequence

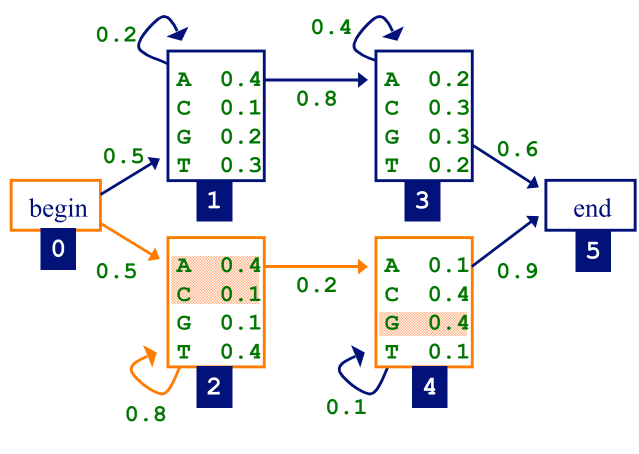
Learning Parameters: The Baum-Welch Algorithm

- algorithm sketch:
 - initialize parameters of model
 - iterate until convergence
 - calculate the *expected* number of times each transition or emission is used
 - adjust the parameters to *maximize* the likelihood of these expected values

• we want to know the probability of producing sequence *x* with the *i* th symbol being produced by state *k* (for all *x*, *i*

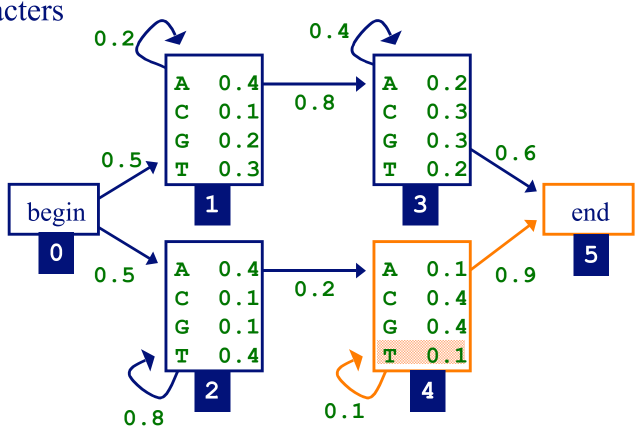


• the forward algorithm gives us $f_k(i)$, the probability of being in state k having observed the first i characters of x



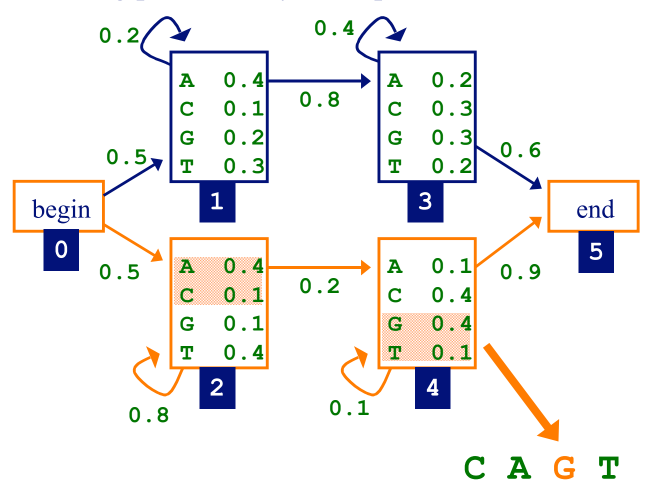
CAGT

• the backward algorithm gives us $b_k(i)$, the probability of observing the rest of x, given that we're in state k after i characters



CAGT

• putting forward and backward together, we can compute the probability of producing sequence *x* with the *i* th symbol being produced by state *q*



• first, we need to know the probability of the *i* th symbol being produced by state *k*, given sequence *x*

$$\Pr(\pi_i = k \mid x)$$

• given this we can compute our expected counts for state transitions, character emissions

• the probability of of producing *x* with the *i* th symbol being produced by state *k* is

$$Pr(\pi_i = k, x) = Pr(x_1...x_i, \pi_i = k) \times$$

$$Pr(x_{i+1}...x_L | \pi_i = k)$$

- the first term is $f_k(i)$, computed by the forward algorithm
- the second term is $b_k(i)$, computed by the backward algorithm

The Backward Algorithm

• initialization:

$$b_k(L) = a_{kN}$$

for states with a transition to end state

The Backward Algorithm

• recursion (i = L...1):

$$b_k(i) = \sum_{l} \begin{cases} a_{kl}b_l(i), & \text{if } l \text{ is silent state} \\ a_{kl}e_l(x_{i+1})b_l(i+1), & \text{otherwise} \end{cases}$$

The Backward Algorithm

termination:

$$Pr(x) = Pr(x_1...x_L) = \sum_{l} \begin{cases} a_{0l}b_l(0), & \text{if } l \text{ is silent state} \\ a_{0l}e_l(x_1)b_l(1), & \text{otherwise} \end{cases}$$

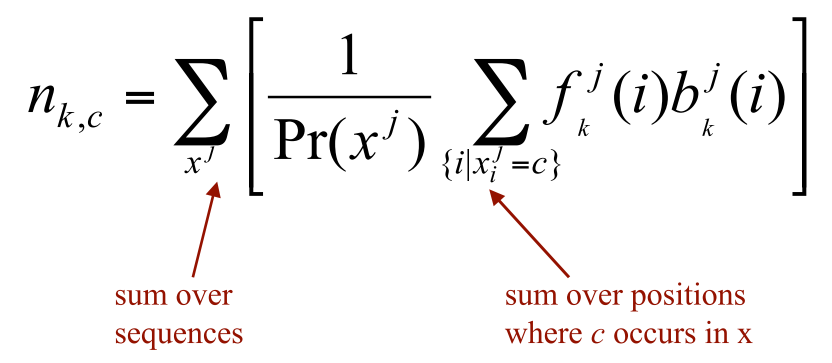
• now we can calculate the probability of the *i* th symbol being produced by state *k*, given *x*

$$Pr(\pi_i = k \mid x) = \frac{Pr(\pi_i = k, x)}{Pr(x)}$$

$$= \frac{f_k(i)b_k(i)}{Pr(x)}$$

$$= \frac{f_k(i)b_k(i)}{f_N(L)}$$

- now we can calculate the expected number of times letter *c* is emitted by state *k*
- here we've added the superscript *j* to refer to a specific sequence in the training set



• and we can calculate the expected number of times that the transition from *k* to *l* is used

$$n_{k\to l} = \sum_{x^{j}} \frac{\sum_{i} f_{k}^{j}(i) a_{kl} e_{l}(x_{i+1}^{j}) b_{l}^{j}(i+1)}{\Pr(x^{j})}$$

• or if *l* is a silent state

$$n_{k \to l} = \sum_{x^{j}} \frac{\sum_{i} f_{k}^{j}(i) a_{kl} b_{l}^{j}(i)}{\Pr(x^{j})}$$

The Maximization Step

- Let $n_{k,c}$ be the expected number of emissions of c from state k for the training set
- estimate new emission parameters by:

$$e_k(c) = \frac{n_{k,c}}{\sum_{c'} n_{k,c'}}$$

- just like in the simple case
- but typically we'll do some "smoothing" (e.g. add pseudocounts)

The Maximization Step

- let $n_{k\to l}$ be the expected number of transitions from state k to state l for the training set
- estimate new transition parameters by:

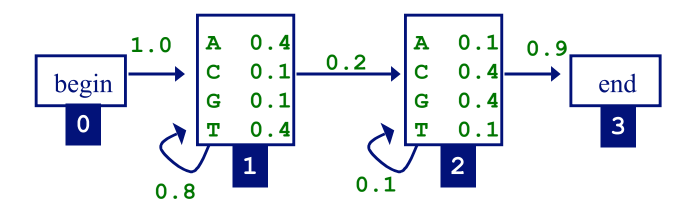
$$a_{kl} = \frac{n_{k \to l}}{\sum_{m} n_{k \to m}}$$

The Baum-Welch Algorithm

- initialize the parameters of the HMM
- iterate until convergence
 - initialize $n_{k,c}$, $n_{k\rightarrow l}$ with pseudocounts
 - **E-step**: for each training set sequence j = 1...n
 - calculate $f_k(i)$ values for sequence j
 - calculate $b_k(i)$ values for sequence j
 - add the contribution of sequence j to $n_{k,c}$, $n_{k\rightarrow l}$
 - **M-step**: update the HMM parameters using $n_{k,c}$, $n_{k\rightarrow l}$

Baum-Welch Algorithm Example

- given
 - the HMM with the parameters initialized as shown
 - the training sequences TAG, ACG



• we'll work through one iteration of Baum-Welch

determining the forward values for TAG

$$\begin{split} f_0(0) &= 1 \\ f_1(1) &= e_1(T) \times a_{01} \times f_0(0) = 0.4 \times 1 = 0.4 \\ f_1(2) &= e_1(A) \times a_{11} \times f_1(1) = 0.4 \times 0.8 \times 0.4 = 0.128 \\ f_2(2) &= e_2(A) \times a_{12} \times f_1(1) = 0.1 \times 0.2 \times 0.4 = 0.008 \\ f_2(3) &= e_2(G) \times \left(a_{12} \times f_1(2) + a_{22} \times f_2(2)\right) = \\ 0.4 \times (0.0008 + 0.0256) &= 0.01056 \end{split}$$

- here we compute just (he) values that depols the over to word a 5024 ro probability
- in a similar way, we also compute forward values for ACG

determining the backward values for TAG

$$b_3(3) = 1$$

$$b_2(3) = a_{23} \times b_3(3) = 0.9 \times 1 = 0.9$$

$$b_2(2) = a_{22} \times e_2(G) \times b_2(3) = 0.1 \times 0.4 \times 0.9 = 0.036$$

$$b_1(2) = a_{12} \times e_2(G) \times b_2(3) = 0.2 \times 0.4 \times 0.9 = 0.072$$

$$b_1(1) = a_{11} \times e_1(A) \times b_1(2) + a_{12} \times e_2(A) \times b_2(2) = 0.8 \times 0.4 \times 0.072 + 0.2 \times 0.1 \times 0.036 = 0.02376$$

- bo (There who compute Julies that is present and with the company of the probability
 - in a similar way, we also compute backward values for ACG

determining the expected emission counts for state 1

contribution of TAG contribution of ACG pseudocount
$$n_{1,A} = \frac{f_1(2)b_1(2)}{f_3(3)} + \frac{f_1(1)b_1(1)}{f_3(3)} + 1$$

$$n_{1,C} = \frac{f_1(2)b_1(2)}{f_3(3)} + 1$$

$$n_{1,C} = \frac{f_1(1)b_1(1)}{f_3(3)} + 1$$

$$n_{1,T} = \frac{f_1(1)b_1(1)}{f_3(3)} + 1$$

^{*}note that the forward/backward values in these two columns differ; in each column they are computed for the sequence associated with the column

• determining the expected transition counts for state 1 (not using pseudocounts)

contribution
of TAG

of ACG

$$n_{1\rightarrow 1} = \frac{f_1(1)a_{11}e_1(A)b_1(2)}{f_3(3)} + \frac{f_1(1)a_{11}e_1(C)b_1(2)}{f_3(3)}$$

$$n_{1\to 2} = \frac{f_1(1)a_{12}e_2(A)b_2(2) + f_1(2)a_{12}e_2(G)b_2(3)}{\text{in a similar way } f_1(2)a_{12}e_2(G)b_2(3)} + \frac{f_1(1)a_{12}e_2(C)b_2(2) + f_1(2)a_{12}e_2(G)b_2(3)}{\text{counts for state 2}}$$

• determining probabilities for state 1

$$e_{1}(A) = \frac{n_{1,A}}{n_{1,A} + n_{1,C} + n_{1,G} + n_{1,T}}$$

$$e_{1}(C) = \frac{n_{1,C}}{n_{1,A} + n_{1,C} + n_{1,G} + n_{1,T}}$$

$$M$$

$$a_{11} = \frac{n_{1 \to 1}}{n_{1 \to 1} + n_{1 \to 2}}$$

$$a_{12} = \frac{n_{1 \to 2}}{n_{1 \to 1} + n_{1 \to 2}}$$

Markov Models Summary

- we considered models that varied in terms of order, in/homogeneity, hidden state
- three DP-based algorithms for HMMs: Forward, Backward and Viterbi
- we discussed three key tasks: learning, classification and segmentation
- the algorithms used for each task depend on whether there is hidden state (correct path known) in the problem or not

Comments on Markov Models

- there are many successful applications in computational biology
 - gene recognition and associated subtasks
 - protein family modeling
 - motif modeling
 - etc.
- there are many variants of the models/algorithms we considered here (some of these are covered in BMI/CS 776)
 - fixed length motif models
 - semi-markov models
 - stochastic context free grammars
 - Gibbs sampling for learning parameters