Comp 555 - BioAlgorithms - Spring 2019



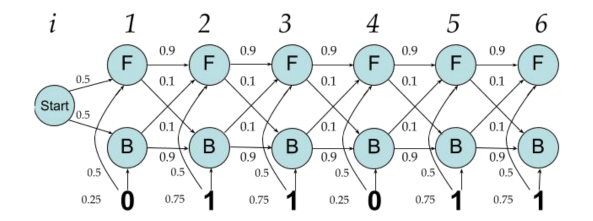
• PAY ATTENTION, PROBLEM SET #5, IS BASED ON THIS MATERIAL

Inferring Ancestory using HMMs

Decoding Problem Solution



• The *Decoding Problem* is equivalent to finding a longest path in the directed acyclic graph (DAG), where "longest" is defined as the maximum product of the probabilities along the path.



Viterbi Decoding Algorithm



- Since the *longest path* is a product of edge weights, if we use the **log** of the weights we can make it a sum again!
- The value of the product can become extremely small, which leads to underflow.
- Many common probability distributions have an exponential form. Taking their log simplifies these distributions.
- Improves numerical accurracy and stability.

$$s_{k,i+1} = \log(e_l(x_{i+1})) + \max_{k \in Q} \{s_{k,i} + \log(a_{kl})\}$$

Viterbi Decoding Algorithm (cont)



- Every path in the graph has the probability $P(x|\pi)$.
- The Viterbi decoding algorithm finds the path that maximizes $P(x|\pi)$ among all possible paths.
- The Viterbi decoding algorithm runs in *O*(*n*/*Q*/²) time (length of sequence times number of states squared).
- The Viterbi decoding algorithm can be efficiently implemented as a dynamic program

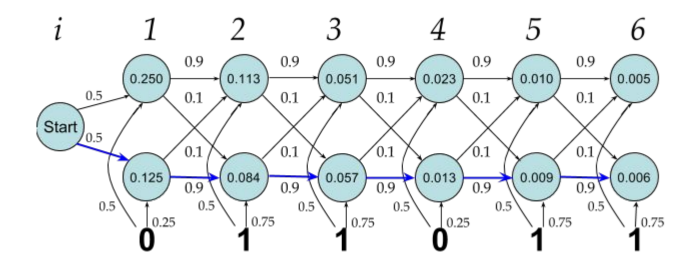
Dynamic Program's Recursion:

$$s_{l,i+1} = \max_{k \in Q} \{s_{k,i} \cdot \text{weight of edge between } (k,i) \text{ and } (l,i+1)\}$$
$$= \max_{k \in Q} \{s_{k,i} \cdot a_{kl} \cdot e_l(x_{i+1})\}$$
$$= e_l(x_{i+1}) \cdot \max_{k \in Q} \{s_{k,i} \cdot a_{kl}\}$$

Viterbi Example



- Solves all subproblems implied by observed sequence
- How likely is this path? 0.006
- What is it? **BBBBBB**



How likely is "most likely?



- The "most likely path" may not be a lot more likely than a 2nd or 3rd most likely paths (even more so in more realistic cases than this one).
- Actual probability of the "most likely path" is not that high.

Р	π	Р	π	Р	π	Р	π	
0.0058	BBBBBB	0.0001	BBBFFB	0.0000	FFFBFF	0.0000	FBBFBF	
0.0046	FFFFF	0.0001	FFFFBF	0.0000	FFBFBB	0.0000	BFBBFF ("
0.0013	FBBBBB	0.0001	FFBFFF	0.0000	FBFFBB	0.0000	BFFBBF	" FFFFFF " is
0.0012	FFFFBB	0.0001	FBFFFF	0.0000	FBBFFB	0.0000	BBFBFF	nearly as good
0.0009	FFBBBB	0.0001	FFBBBF	0.0000	FFBFFB	0.0000	FFBFBF	as " BBBBBB "
0.0008	FFFFB	0.0001	BFFFBB	0.0000	FBFFFB	0.0000	FBFFBF	
0.0006	FFFBBB	0.0001	FBBBFF	0.0000	FBFBBB	0.0000	BFFBFF	
0.0006	BBBFFF	0.0001	BBFFFB	0.0000	FBBBFB	0.0000	BFBFBB	
0.0004	BBBBBF	0.0000	BFBBBB	0.0000	BBBFBF	0.0000	FBFBBF	
0.0004	BBFFFF	0.0000	BBBBFB	0.0000	FFBBFB	0.0000	BFBFFB	
0.0003	BBBBFF	0.0000	BBFBBB	0.0000	BBFFBF	0.0000	FBFBFF	
0.0003	BFFFFF	0.0000	BFFFFB	0.0000	BFFFBF	0.0000	BFBBFB	Π
0.0001	BBBFBB	0.0000	FFFBBF	0.0000	BFBFFF	0.0000	BBFBFB	11
0.0001	FBBFFF	0.0000	FFBBFF	0.0000	FFFBFB	0.0000	BFFBFB	
0.0001	FBBBBF	0.0000	FBBFBB	0.0000	BFBBBF	0.0000	FBFBFB	-
0.0001	BBFFBB	0.0000	BFFBBB	0.0000	BBFBBF	0.0000	BFBFBF	

HMMs in Biology



- Inferring ancestral contributions of a descendant
- Collaborative Cross project
- Maintained at UNC since 2006

Objective:

Create new reproducible mouse strains by randomly combining the genomes of eight diverse mice strains

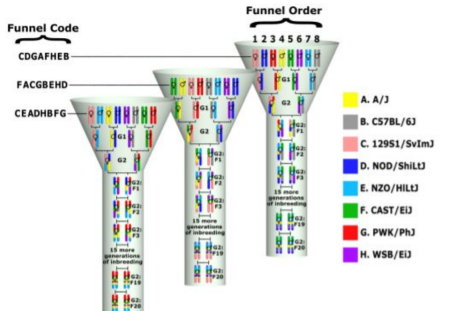
Problem:

Given an extant strain, which parts of its genome came from which founder?



Mixing Genome

- A randomized breeding scheme was used to
 - Mix the genomes by recombination
 - Fix the genomes by inbreeding
- A breeding funnel 8 genomes go in a mosaic comes out
- Genotyping was used to track founder contributions

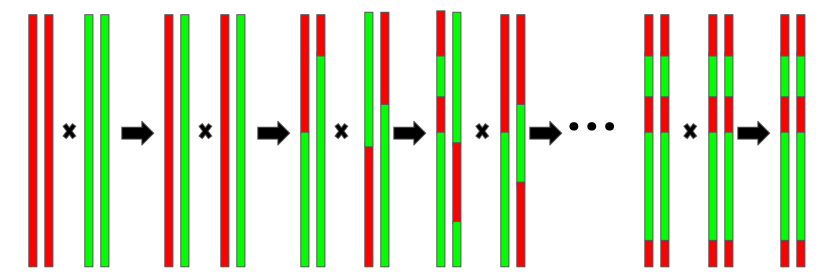




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Instead of "Birds and Bees," Mouse and Flies

- Recombination mixes the genomes of the two chromosomes
- Sib-mating causes the genomes to fix

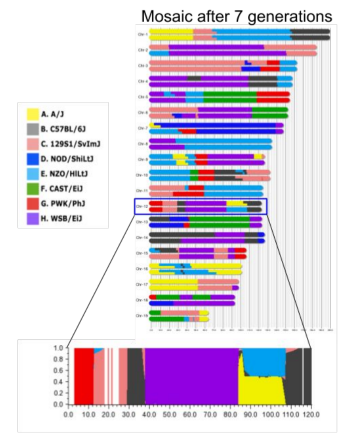




A Genome Mosaic



- A Hidden Markov Model is used to infer the "hidden" state of which of the 8 founders contributed to what parts of the genome
- A Viterbi Solution finds the most likely mosaic given a set of "genotypes"



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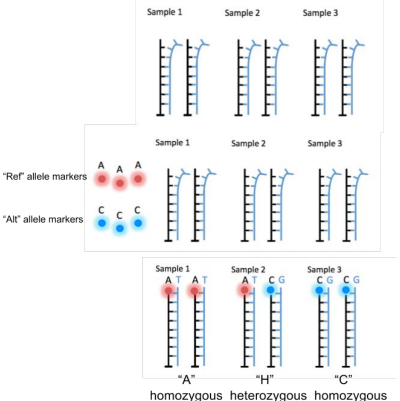
Genotyping Microarrays

 DNA probes to query the state of specific "known" and "informative" Single Nucleotide Polymorphisms (SNPs)

Bases in the genome that vary within a population

 Each probe distinguishes 4 cases ("Ref", "Alt", "H", "N")

• From these observations we infer the founder at every marker







Example Genotypes

- Genotypes for a chromosome
- 1000s of probes with positions of variant
- Alleles are indicated by the nucleotide
- Rarely can a single maker resolve the founder
- Which strain would you guess?

Probe info			Fo	und	er (Gen	otyp	bes	Т	arg
	<u> </u>	Г								¥
chromosome	positionB38	C/A	C57BL/6J	129S1/SvImJ	NOD/ShiLtJ	NZO/H1LtJ	CAST/EiJ	PWK/PhJ	WSB/EiJ	OR3199m266
2	3176721	G	Т	G	Т	G	G	Т	Т	Т
2	3180256	G	G	G	G	G	A	A	G	G
2	3182308	А	G	А	G	А	G	G	А	Α
2	3183784	Т	G	Т	G	Т	G	G	Т	Т
2	3233750	G	G	G	G	G	A	G	G	G
2	3350920	А	A	Α	A	А	G	G	A	А
2	3353380	Т	Т	C	Т	С	С	С	С	С
2	3362696	Т	Т	Т	Т	Т	Т	C	Т	Т
2	3420272	С	С	Т	С	Т	Т	Т	С	С
2	3433708	G	G	G	G	G	A	A	G	G
2	3438642	С	С	Т	С	т	С	Т	С	С
2	3456515	С	С	С	С	С	Т	С	С	С
2	3503822	Т	Т	Т	Т	С	Т	С	т	Т
2	3557793	Α	A	Α	A	А	G	G	Α	A
2	3595443	Т	Т	G	Т	G	G	G	Т	Т
2	3613854	Α	A	A	A	G	G	G	A	A
2	3663247	Т	Т	Т	т	Т	C	C	т	Т
2	3666094	G	G	G	G	G	G	т	т	Т
2	3681891	G	G	G	G	G	A	G	G	G
2	3715097	G	G	G	G	G	т	т	G	G

Genotype Noise



- One last issue, between 1% and 5% of genotypes are simply wrong
- Source of errors
 - A probe didn't glow bright enough
 - A section of the array was damaged (fingerprints, cracks, hair, etc.)
 - Mess ups when fabricating a probe's sequence
 - DNA itself was contaminated
- Error types:
 - "No" calls (observation is uninformative)
 - A possible, but incorrect call

Reading Genotypes



```
In [1]: fp = open("CCGenotypes.csv", 'r')
                              # break file into lines
     data = fp.read().split('\n')
      fp.close()
      header = data.pop(0).split(',')  # First line is header
     while (len(data[-1].strip()) < 1): # remove extra lines</pre>
        data.pop()
      for i, line in enumerate(data):
                              # make a list from each row
        field = line.split(',')
        field[1] = int(field[1])
                             # convert position to integer
        data[i] = field
      fp.close()
      print(header)
      print("Number of probes", len(data))
     for i in range(100,110):
        print("data[%d] = %s" % (i, data[i]))
      ['Chromosome', 'Position', 'A/J', 'C57BL/6J', '129S1/SvImJ', 'NOD/ShiLtJ', 'NZO/HlLtJ', 'CAST/EiJ', 'PWK/PhJ', 'WSB/
     EiJ', 'CC004/TauUnc']
     Number of probes 419
     data[101] = ['1', 59995627, 'A', 'A', 'A', 'C',
                                       'C', 'C', 'C', 'C',
                                                     'C']
                                   'G',
                                       'G',
     data[102] = ['1', 60400655, 'A', 'A', 'A',
                                              'A',
                                                 'G',
                                          'G',
     data[103] = ['1', 60761817, 'G', 'G', 'G',
                                   'A',
                                       'A',
                                          'A',
                                              'G',
                                                 'G',
                                                     'G']
                                   'C', 'C', 'C',
                                              'T', 'C', 'C']
     data[104] = ['1', 61312969, 'C', 'C', 'C',
```

Emission Probabilites based on Genotypes



Each probe has its own emission probabilites

```
In [2]: i = int(input("Enter locus [0, %d] to see its Emission probability:" % len(data)))
       print(data[i])
       Nstates = 8
       ErrorRate = 0.05
       # Count expected genotypes
       count = dict([(call, data[i][2:2+Nstates].count(call)) for call in "ACGTHN"])
                        ", ', '.join(["%8s" % v[0:8] for v in header[2:2+Nstates]]))
       print("
       for base in count.keys():
           # Compute emission probability, assuming 5% error rate
           if (count[base] == 0):
               emission = [1.0/Nstates for j in range(2,2+Nstates)] # unexpected
           else:
               emission = [(1.0 - ErrorRate)/count[base] if data[i][j] == base else ErrorRate/(Nstates - count[base])
                          for j in range(2,2+Nstates)]
           emission = ["%6.4f" % v for v in emission]
           print(" %s: %2d %s" % (base, count[base], emission))
       Enter locus [0, 419] to see its Emission probability:103
       A/J, C57BL/6J, 129S1/Sv, NOD/ShiL, NZO/HlLt, CAST/EiJ, PWK/PhJ, WSB/EiJ
           A: 3 ['0.0100', '0.0100', '0.0100', '0.3167', '0.3167', '0.3167', '0.0100', '0.0100']
           C: 0 ['0.1250', '0.1250', '0.1250', '0.1250', '0.1250', '0.1250', '0.1250', '0.1250']
           G: 5 ['0.1900', '0.1900', '0.1900', '0.0167', '0.0167', '0.0167', '0.1900', '0.1900']
```

T: 0 ['0.1250', '0.1250', '0.1250', '0.1250', '0.1250', '0.1250', '0.1250', '0.1250']

H: 0 ['0.1250', '0.1250', '0.1250', '0.1250', '0.1250', '0.1250', '0.1250', '0.1250'] N: 0 ['0.1250', '0.1250', '0.1250', '0.1250', '0.1250', '0.1250', '0.1250'] Comp 555 - Fall 2019

Transition probabilities

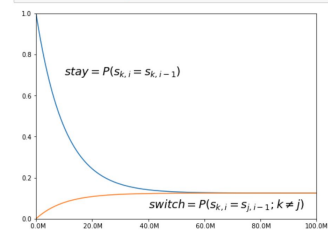
In [8]: %matplotlib inline
 import numpy

import matplotlib.pyplot as plot



- Recombination likelihood is modeled using an exponential distribution
- Recombinations between nearby probes are unlikely
- Distant probes are more likely to be from different founders

```
fig = plot.figure(figsize = (8,6))
axes = fig.add_subplot(111)
Nstates = 8
scale = 10000000.0
x = numpy.arange(0,100000000.0,200000.0)
stay = ((Nstates - 1.0) * numpy.exp(-x/scale) + 1.0) / Nstates
switch = (1.0 - stay) / (Nstates - 1.0)
plot.plot(x, stay, x, switch)
plot.text(10000000, 0.7, r'$stay = P(s_{k,i} = s_{k,i-1})$', size="18")
plot.text(4000000, 0.05, r'$switch = P(s_{k,i} = s_{j,i-1}; k \neq j)$', size="18")
plot.xlim((0,10000000.0))
plot.ylim((0,1.0))
pos, labels = plot.xticks()
result = plot.xticks(pos, ["%5.1fM" % (p/1000000) for p in pos])
```



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Viterbi Algorithm as a Dynamic Program



```
In [18]: from math import exp, log10
         Nstates = 8
         prevpos = 1
         state = [[(float(len(data)),i) for i in range(Nstates)]] # (log(p), PathToHere)
         for i in range(len(data)):
             # Count expected genotypes
             count = dict([(call, data[i][2:2+Nstates].count(call)) for call in "ACGTHN"])
             # Get the target genotype at this probe
             observed = data[i][-1]
             # Compute emission probability, assuming 5% error rate
             if (count[observed] == 0):
                 emission = [1.0/Nstates for j in xrange(2,2+Nstates)] # unexpected
             else:
                 emission = [0.95/count[data[i][j]] if data[i][j] == observed else 0.05/(Nstates - count[data[i][j]])
                             for j in range(2,2+Nstates)]
             # compute transition probability
             position = data[i][1]
             delta = position - prevpos
             prevpos = position
             stay = ((Nstates - 1.0)*exp(-delta/10000000.0) + 1.0)/Nstates
             switch = (1.0 - stay)/(Nstates - 1.0)
             # update state probailities for all paths leading to the ith state
             path = [1]
             for j in range(Nstates):
                 choices = [(log10(emission[j])+(log10(stay) if (k==j) else log10(switch))+state[-1][k][0],k)
                            for k in range(Nstates)]
                 path.append(max(choices)) # choices is a list of tuples of (score[i], from_whence_I_arrived[i])
             state.append(path)
         print("Length of paths:", len(state))
```

Length of paths: 418

Backtrack to find solution

```
In [24]: # backtrack
         path = state[-1]
         maxi = 0
         maxp = path[0][0]
         for i in range(1, Nstates):
             if (path[i][0] > maxp):
                 maxp = path[i][0]
                  maxi = i
         print(maxi, path[maxi], header[2+maxi])
         for j in range(len(state)-2,-1,-1):
             data[j].append(header[2+maxi])
             maxi = state[j+1][maxi][1]
         header.append("Founder")
         fp = open("result.csv", 'w')
         fp.write(','.join(header)+'\n')
         prev = ''
          for row in data:
             line = ','.join([str(v) for v in row])
             fp.write(line+' \ n')
             if (row[-1] != prev):
                  print(line)
                 prev = row[-1]
         print(line)
         fp.close()
```

5 (129.58171061177885, 5) CAST/EiJ 1,3409090, C, C, A, A, C, A, A, A, A, PWK/PhJ, PWK/PhJ 1,14334166, A, G, A, A, A, G, G, G, G, AJS1/SVImJ, 129S1/SVImJ 1,41477940, G, A, A, A, G, G, G, G, A, /J, A/J 1,52869070, G, G, G, A, A, G, G, G, A, A, WSB/EiJ, WSB/EiJ 1,67749123, A, G, A, A, G, G, G, A, A, WSB/EiJ, WSB/EiJ 1,12786434, C, C, C, C, C, T, C, C, C, C, C57BL/6J, C57BL/6J 1,172685919, A, G, A, A, G, G, G, A, A/J, A/J 1,176674355, A, G, G, G, A, G, CAST/EiJ, CAST/EiJ 1,194886567, G, G, T, G, T, T, C, T, T, CAST/EiJ, CAST/EiJ



A peek at the result

There are a second

In [23]: !head result.csv; echo '...'; tail result.csv

Chromosome, Position, A/J, C57BL/6J, 129S1/SvImJ, NOD/ShiLtJ, NZO/HlLtJ, CAST/EiJ, PWK/PhJ, WSB/EiJ, CC004/TauUnc, Founder

1, 3409090, C, C, A, A, C, A, A, A, A, PWK/PhJ 1, 3427467, A, A, A, A, A, G, G, A, G, PWK/PhJ 1, 3439034, C, C, T, T, C, C, C, T, C, PWK/PhJ 1, 3668628, A, G, G, G, G, A, A, G, A, PWK/PhJ 1, 4504223, G, G, G, G, G, A, G, A, G, PWK/PhJ 1, 4744395, T, T, T, T, T, T, G, T, G, T, PWK/PhJ 1, 5069641, A, A, A, A, A, G, A, A, A, PWK/PhJ 1, 5149169, T, G, T, G, T, G, T, T, T, PWK/PhJ 1, 7698048, A, G, A, A, A, G, G, G, G, PWK/PhJ

1 10'

1,193654902,G,A,A,G,A,G,G,G,G,CAST/EiJ 1,193673297,G,A,A,G,A,G,G,G,G,G,CAST/EiJ 1,193688845,A,C,C,A,C,C,A,A,C,CAST/EiJ 1,193709621,G,A,A,A,A,A,G,G,A,CAST/EiJ 1,193732571,T,C,C,C,C,C,T,C,C,CAST/EiJ 1,193928056,A,G,G,A,G,A,A,A,A,CAST/EiJ 1,194000258,C,C,C,T,C,C,C,T,C,CAST/EiJ 1,194149219,G,A,A,G,G,G,G,G,G,G,CAST/EiJ 1,194886567,G,G,T,G,T,T,G,T,T,CAST/EiJ





- The inferred Mosaic
- Repeat for every chromosome
- Most likely, but how likely?
- Other approaches

Back to the Casino with new questions



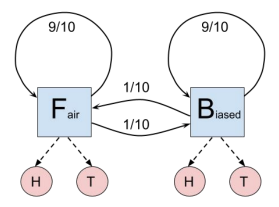
- Are there common aspects of the most likely solutions?
- Which coin was I most likely using on the 4th roll

Р	π	Р	π	Р	π	Р	π
0.0058	BBBBBB	0.0001	BBBFFB	0.0000	FFFBFF	0.0000	FBBFBF
0.0046	FFFFF	0.0001	FFFFBF	0.0000	FFBFBB	0.0000	BFBBFF
0.0013	FBBBBB	0.0001	FFBFFF	0.0000	FBFFBB	0.0000	BFFBBF
0.0012	FFFFBB	0.0001	FBFFFF	0.0000	FBBFFB	0.0000	BBFBFF
0.0009	FFBBBB	0.0001	FFBBBF	0.0000	FFBFFB	0.0000	FFBFBF
0.0008	FFFFB	0.0001	BFFFBB	0.0000	FBFFFB	0.0000	FBFFBF
0.0006	FFFBBB	0.0001	FBBBFF	0.0000	FBFBBB	0.0000	BFFBFF
0.0006	BBBFFF	0.0001	BBFFFB	0.0000	FBBBFB	0.0000	BFBFBB
0.0004	BBBBBF	0.0000	BFBBBB	0.0000	BBBFBF	0.0000	FBFBBF
0.0004	BBFFFF	0.0000	BBBBFB	0.0000	FFBBFB	0.0000	BFBFFB
0.0003	BBBBFF	0.0000	BBFBBB	0.0000	BBFFBF	0.0000	FBFBFF
0.0003	BFFFFF	0.0000	BFFFFB	0.0000	BFFFBF	0.0000	BFBBFB
0.0001	BBBFBB	0.0000	FFFBBF	0.0000	BFBFFF	0.0000	BBFBFB
0.0001	FBBFFF	0.0000	FFBBFF	0.0000	FFFBFB	0.0000	BFFBFB
0.0001	FBBBBF	0.0000	FBBFBB	0.0000	BFBBBF	0.0000	FBFBFB
0.0001	BBFFBB	0.0000	BFFBBB	0.0000	BBFBBF	0.0000	BFBFBF

Forward-Backward Problem



Given: A sequence of coin tosses generated by an HMM.



Goal: Find the most probable coin that was in use at a particular flip.

$$P(\pi_i = k | x) = \frac{P(x, \pi_i = k)}{P(x)}$$

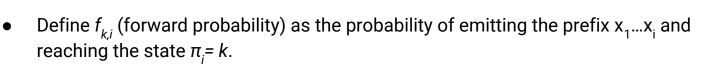
Where $P(x, \pi_i = k)$ is the probabilites of all paths in state *k* at *i*, and P(x) is the probability of sequence *x*. Comp 555 - Fall 2019



Illustrating the difference using 4 flips

Not a lot worse than the best solution $X = (T, H, H, H, H)$ X = (T, H, H, H, H) Viterbi solution, the most likely sequence states.	<pre>FFFF (0.0228) BFFF (0.0013) FBFF (0.0004) BBFF (0.0004) BFBF (0.0004) BFBF (0.0000) FBBF (0.0006) BBBF (0.0028) FFFB (0.0028) FFFB (0.0002) FBFB (0.0002) FBFB (0.0001) BBFB (0.0003) FFBB (0.0003) FFBB (0.0003) FBBB (0.0085) BBBBB (0.0384) P(x) = 0.0877 High probability output (>0.0625)</pre>	x = THHH p FFFF (0.0228) FFBF (0.0004) FFBB (0.0038) FFBB (0.0057) BFFF (0.0013) BFBF (0.0000) BFBB (0.0002) BFBB (0.0003) P(π_2 =F x) = 0.0345/0.0877 = 0.3936 FBFF (0.0004) FBFF (0.0004) FBFF (0.0006) FBFB (0.0006) FBFB (0.0006) BBFF (0.0001) FBBB (0.0085) BFFF (0.0019) BBFF (0.0019) BBFF (0.0003) BBFB (0.0003) BBFB (0.00384) P(π_2 =B x) = 0.0532/0.0877 = 0.6064
Comp 555 - Fall 2019		· (₂ D ₁ X) 0.0002/0.0077 0.0004

Forward Algorithm



• The recurrence for the forward algorithm is:

$$f_{k,i} = e_k(x_i) \cdot \sum_{l \in Q} f_{l,i-1} \cdot A_{l,k}$$

• Similar to Viterbi solution to *i*, except all paths are multipled together rather than taking the Max





However, forward probability is not the only factor effecting $P(\pi_i = k | x)$.

- The sequence of transitions and emissions that the HMM undergoes between π_i and π_{i+1} also affect $P(\pi_i = k | x)$.
- Backward probability $b_{ki} \equiv$ the probability of being in state $\pi_i = k$ and emitting the suffix $x_{i+1} \dots x_n$.
- The backward algorithm's recurrence:

$$b_{k,i} = \sum_{l \in Q} e_l(x_{i+1}) \cdot b_{l,i+1} \cdot A_{k,l}$$

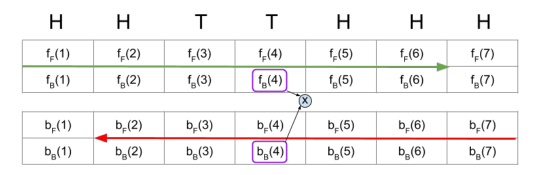
Forward-Backward Algorithm



• The probability that the dealer used a biased coin at any moment *i* is as follows:

$$P(\pi_{i} = k | x) = \frac{P(x, \pi_{i} = k)}{P(x)} = \frac{f_{k}(i) \cdot b_{k}(i)}{P(x)}$$

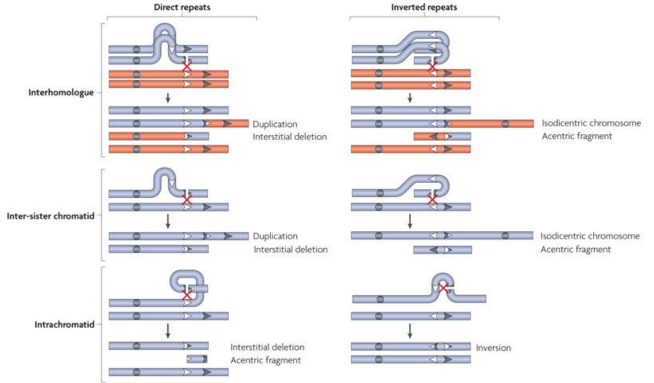
- So, to find $P(\pi_i = k | x)$ for all *i*, we solve two dynamic programs
 - One from beginning to end
 - One from the end to the beginning
 - Combine the corresponding states



Next Time



Genome Rearrangements



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