Outline: Feb 19

- Notes on Lab 2
- Phylogenetic Trees
- UPGMA algorithm
- Ultrametrics

Notes:
- Office hours today: 3-5pm
- No office hours on Wednesday, moved to Tuesday 1-3pm
- Lab this week: practice midterm
- No class on Friday
- Reading posted (Durbin Chap. 7)
Lab 2 Notes
Additional test case

Welcome to the de Bruijn graph assembler

Creating de Bruijn graph with k=3...DONE

Display text version of graph (y/n): y

Nodes: AZ, ZB, BC, CD, DF, FZ, ZR, RT, TT, TB, CF, FR, TA, ZC, RZ, ZT, TX

Edges:
AZ: ['ZB', 'ZC']
ZB: ['BC']
BC: ['CD', 'CF']
CD: ['DF', 'DF']
DF: ['FZ', 'FR']
FZ: ['ZR']
ZR: ['RT']
RT: ['TT', 'TA']
TT: ['TB', 'TX']
TB: ['BC']
CF: ['FR']
FR: ['RT', 'RZ']
TA: ['AZ']
ZC: ['CD']
RZ: ['ZT']
ZT: ['TT']

Write graph visualization to file (y/n): y
Enter filename prefix: test4

Determine whether graph is Eulerian (y/n): y
Graph Eulerian? True

Traverse graph and find contigs (y/n): y
Write contigs to file (y/n): test4.fasta

Starting path 1 / 1 from AZ:
>contig_0
AZCDRFTTBFCRTHZABCDFZRTTX
Genome X examples

k=12

k=15
Lab 2: genome X, k=15

- Balance between low and high k
- If k is too small, will end up with overlaps that are not truly in the reads
- Graph becomes too connected and coverage differences create many non-balanced nodes
- If k is too large, graph becomes very disconnected and we end up building back only the reads
Runtime plot: Daniel and Sayed
Runtime plot: Nathan and Tyler

de Bruijn Graph Assembly Runtime Plot (k = 13)
Phylogenetic Trees: UPGMA
(Unweighted Pair Group Method with Arithmetic mean)
Phylogenetic trees

- **Definition**: diagram of inferred evolutionary relationships between samples (species, genes, individuals, etc)

- **Input**: usually genetic data, although it could be from the fossil record. Preprocessing usually involves alignment (either pairwise or multiple sequence). Then process the alignments to obtain the number of pairwise differences or another form of “dissimilarity”

- **Output**: tree structure PLUS branch lengths which represent time

- **We can learn**: evolutionary history! Sequence of speciation events, function and evolution of common traits and genes, biology of common ancestors, tempo and mode of mutation, natural selection, recombination, migration, population size changes
Great Panda Mystery
Phylogenetic tree of bears and raccoons

Credit: Ameet Soni
1) In phylogenetic trees, observed sequences usually exist at the: (a) root of the tree, (b) internal nodes of the tree, (c) leaves of the tree, (d) all of the above

2) Why do we make the assumption that alleles at each site can be encoded as 0’s and 1’s?

3) Is the reference sequence always the ancestral sequence?
Recap + extensions (discuss with a partner)

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   Answer: No! usually not. The reference happened to be sequenced first. Most of the time we don’t know the ancestral sequence, but phylogenetic trees can help us reconstruct it.
Rooted vs. unrooted trees

Credit: Ameet Soni
Example from last time

- Record pairwise differences (which could be obtained from a pairwise sequence alignment)
- We will use this dissimilarity map as input to our first phylogenetic tree algorithm

<table>
<thead>
<tr>
<th></th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>0</td>
<td>1</td>
<td>3</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>B</td>
<td></td>
<td>0</td>
<td>2</td>
<td>5</td>
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<td>C</td>
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<td>D</td>
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<td>E</td>
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<td>0</td>
</tr>
</tbody>
</table>
find min + merge \{A\} + \{B\}
"Guess" tree, example of the type of output we want

1. $s(x,x) = 0$
2. $s(x,y) = s(y,x)$
(no triangle inequality)

$\begin{array}{cccc}
A & B & C & D \\
A & 0 & 1 & 3 & 6 & 6 \\
B & 0 & 3 & 0 & 6 & 6 \\
C & 0 & 6 & 6 & 0 & \\
D & 0 & 6 & 6 & 0 & \\
E & 0 & 6 & 6 & 0 & \\
\end{array}$

"induced map"

$X = \{A, B, C, D, E\}$

A - C - B
goal: minimize the difference between $S$ and $S'$

(Original) (induced tree metric)

$J(S, S') = \sum_{(i, j) \in X} [\delta(i, j) - \delta'(i, j)]^2$

$(NP \text{- complete})$

UPGMA

**Initialization**

1. Each sample $x \in \mathcal{X}$ starts in its own cluster $C_x$

2. $\Delta$ cluster distance matrix

**Update**

1. Find minimum in $\Delta$ (non-zero) & merge $C_i$ & $C_j$
\[ \Delta(c_1c_2c_3, c_4c_5) = \frac{|c_1|}{|c_1| + |c_3|} \Delta(c_1, c_3) + \frac{|c_3|}{|c_1| + |c_3|} \Delta(c_3, c_5) \]

\[ \Delta(c_1, c_2, c_3) = \frac{1}{2} \cdot 3 + \frac{1}{2} \cdot 2 = 2.5 \]