

CMSC423: Bioinformatic Algorithms, Databases and Tools

Lecture 10

Sequence alignment: inexact
alignment, multiple sequence
alignment

Iterative alignment

```
SC YFPHF1DLSHGSAQVKAHGKKVGDALTLAVGHLDDLPGAL
```

- Take sequences s_i in order:
 - align s_1 with s_c - results in gaps being inserted in both sequences

```
SC YFPHF1DLSHGSAQVKAHGKKVGDALTLAVGHLDDLPGAL
```

```
S1 YFPHF1DLSHG-AQVKG--KKVADALTNVAHVDDMPNAL
```

- align s_2 with s_c - if gaps must be inserted – insert in previously aligned sequences

```
SC YFPHF-DLS-----HGSAQVKAHGKKVGDALTLAVGHLDDLPGAL
```

```
S1 YFPHF-DLS-----HG-AQVKG-GKKVA-----DALTNVAHVDDMPNAL
```

```
S2 FFPKFKGLTTADQLKKSADVRWHAERII-----NAVNDAVASMDDEKMS
```

- and so on (note: if gaps coincide with previously introduced gaps no need to change previously aligned sequences)

```
SC YFPHF-DLS-----HGSAQVKAHGKKVGDALTLAVGHLDDLPGAL
```

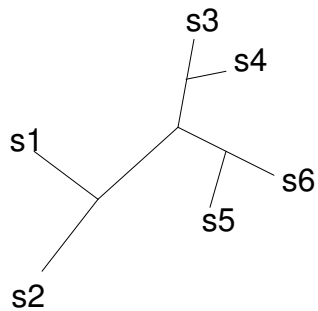
```
S1 YFPHF-DLS-----HG-AQVKG-GKKVA-----DALTNVAHVDDMPNAL
```

```
S2 FFPKFKGLTTADQLKKSADVRWHAERII-----NAVNDAVASMDDEKMS
```

```
S3 LFSFLKGTSEVP--QNNPELQAAGKVF2KLVEAAIQLVTVVVDATL
```

CLUSTALW

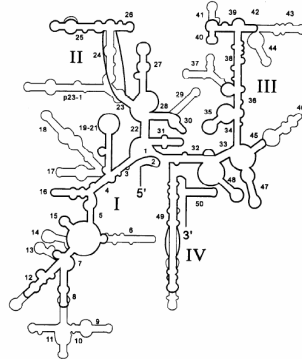
- Compute pairwise distances between strings
- Build phylogenetic tree
- Build iterative alignment by following tree edges



MUSCLE

- Just like ClustalW but different
- Build pairwise distances – uses fast heuristic (just count # of k-mers in common)
- Build phylogenetic tree
- Build multiple alignment based on tree
- Re-estimate distances based on tree
- Re-build tree
- Re-build multiple alignment
- etc. etc. etc.

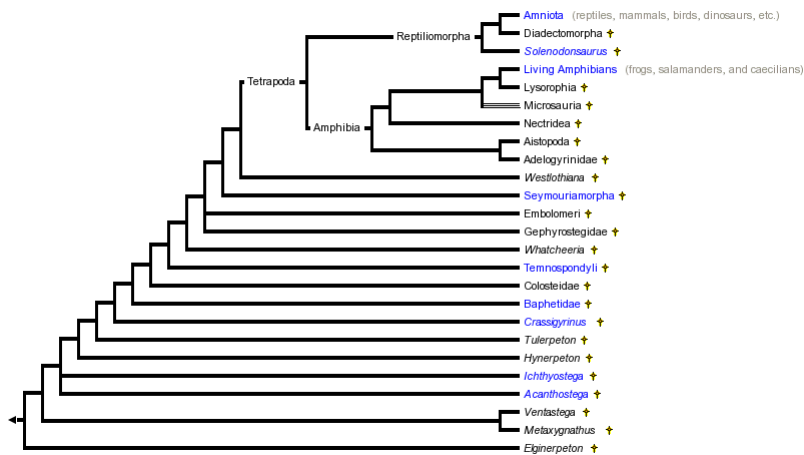
Biological relevance of multiple alignments



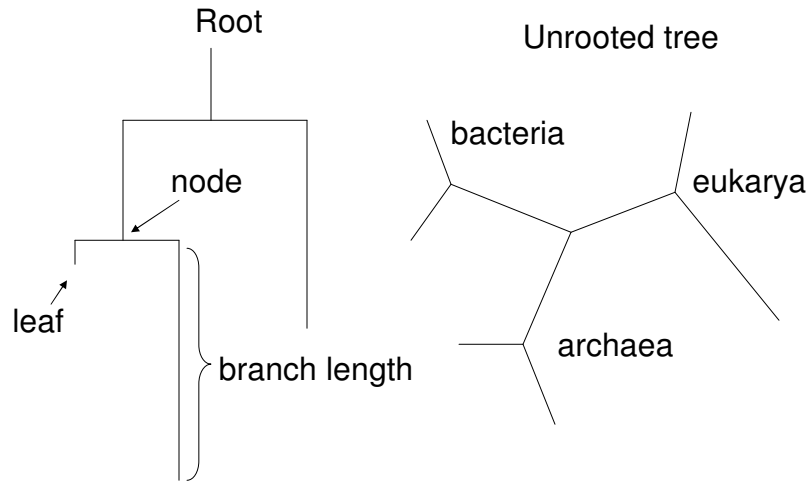
Species	3 2	3 3	3 4	3 5	3 5'
Cow	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - A A A				
Reticle blood	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - A C A				
Scindid lizard	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - C W A				
Xenopus	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - A A A				
Oppreid fish	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - A A A				
Sea Urchin	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - A A A				
Drosophila	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - G G A				
Hemaphys	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - A A U				
Loquat	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - U A A				
Onesidly	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - U A A				
Castipade	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - U A A				
Sorapim	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - U A A				
Byline thran	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - A C A				
Orychophoran	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - A A A				
Small	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - A A A				
Bathoon	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - U U U				
NCIFP	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - Y				
Meatcde	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - U - -				
Sea Anemone	U G G U G G U G G C U U U A A U C C U U C U A G U A G U G C C U G U C U G U A - A U C U G U A U A A - A C C C G G A U - U A A				

Phylogenetic trees – how evolution works

- <http://www.tolweb.org/tree/> - the tree of life



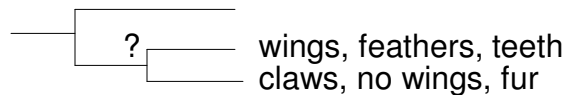
Anatomy of a tree



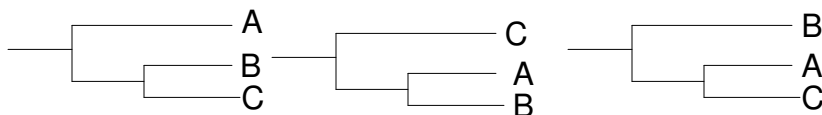
Phylogenetic trees are usually binary (though they don't have to)

Phylogeny questions

- Given several organisms & a set of features (usually sequence, but also morphological: wing shape/color...)
- A. Given a phylogenetic tree – figure out what the ancestors looked like (what are the features of internal nodes)



- B. Find the phylogenetic tree that best describes the common evolutionary heritage of the organisms



Phylogeny questions

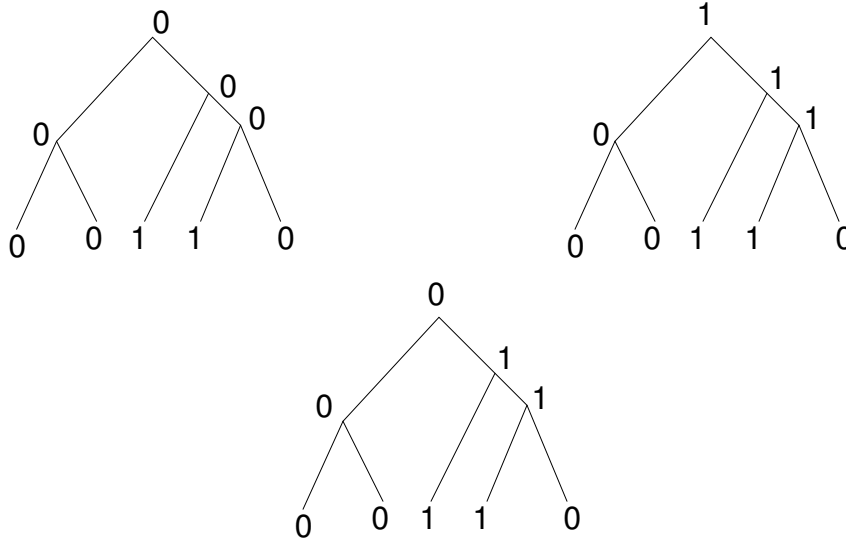
- A. Easy-ish – can be done with dynamic programming
- B. Hard – Many possible trees

$$\frac{(2n-3)!}{2^{n-2}(n-2)!} \quad \text{rooted trees with } n \text{ leaves}$$

Scoring a tree – Sankoff's algorithm

- Assumption – we try to minimize # of state changes from root to leaves – Parsimony approach
- Small parsimony
 - given a tree where leaves are labeled with m-character strings
 - find labels at internal nodes s.t. # of state transitions is minimized
- Weighted small parsimony
 - same as parsimony except that state transitions are assigned weights
 - minimize the overall weight of the tree

Example



Sankoff's algorithm

- At each node v in the tree store $s(v,t)$ – best parsimony score for subtree rooted at v if character stored at v is t
- Traverse the tree in post-order and update $s(v,t)$ as follows
 - assume node v has children u and w
 - $s(v,t) = \min_i \{s(u,i) + \text{score}(i,t)\} + \min_j \{s(w,j) + \text{score}(j,t)\}$
- Character at root will be the one that maximizes $s(\text{root}, t)$
- Note – this solves the weighted version. For unweighted set $\text{score}(i,i) = 0$, $\text{score}(i,j) = 1$ for any i, j