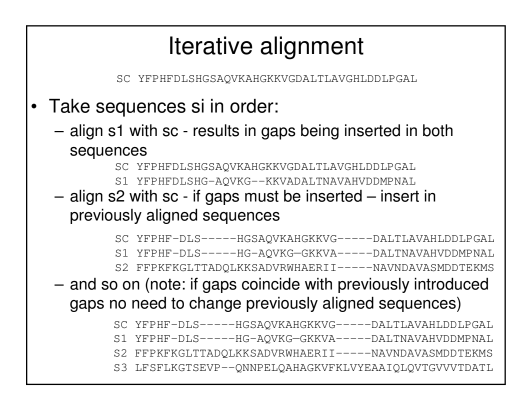
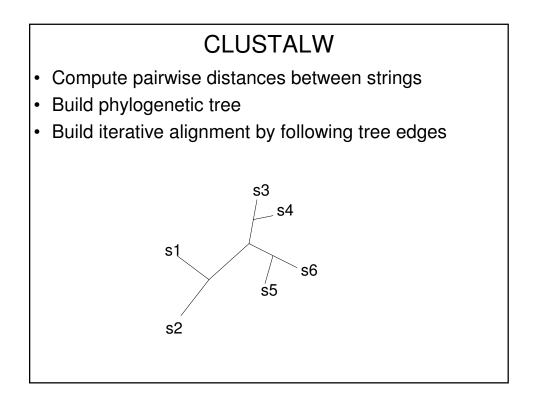
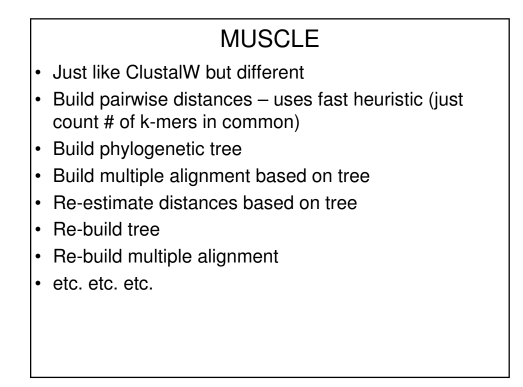
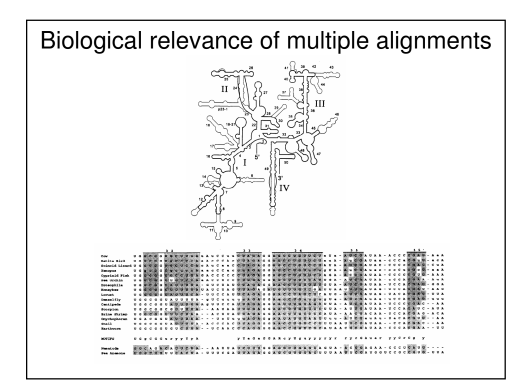
## CMSC423: Bioinformatic Algorithms, Databases and Tools Lecture 10

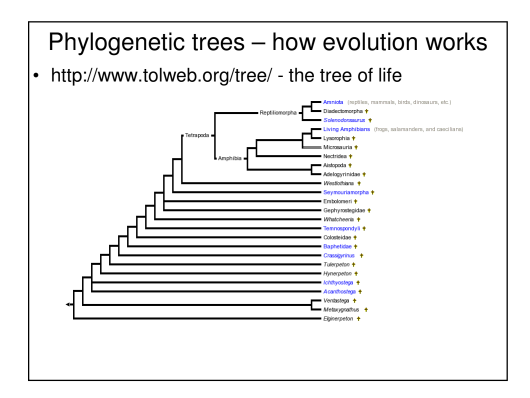
Sequence alignment: inexact alignment, multiple sequence alignment

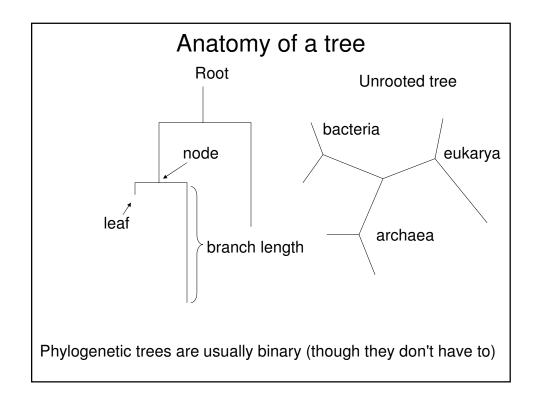


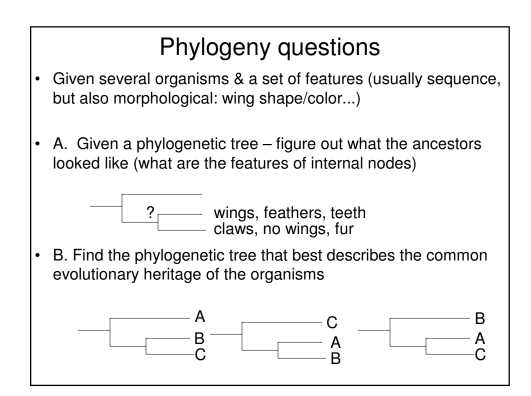


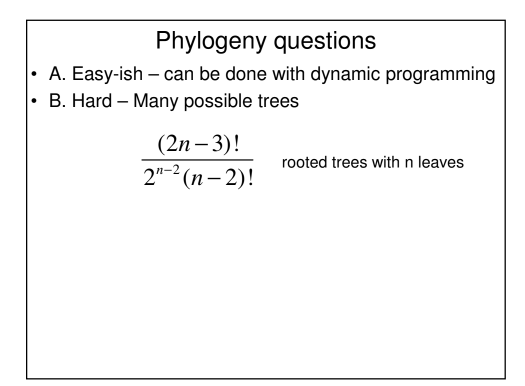


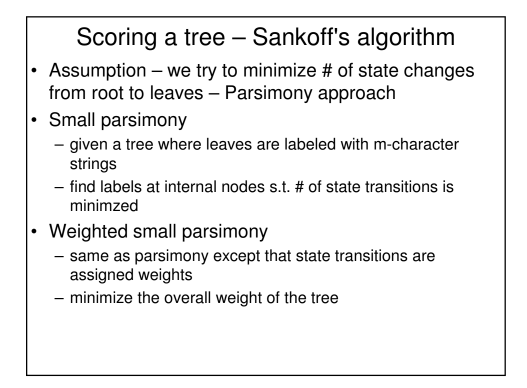


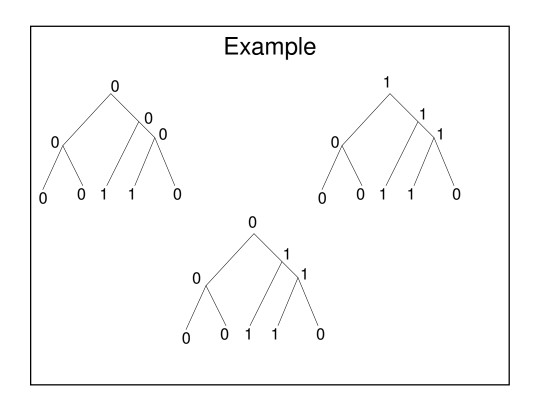












## Sankoff's algorithm

- 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
  - $s(v,t) = min_i \{s(u,i) + score(i,t)\} + min_j \{s(w,j) + score(j,t)\}$
- Character at root will be the one that maximizes s(root, t)
- Note this solves the weighted version. For unweighted set score (i,i) = 0, score(i,j) = 1 for any i,j