# CS481/CS583: Bioinformatics Algorithms

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## GUIDE TREES AND EVOLUTIONARY TREES

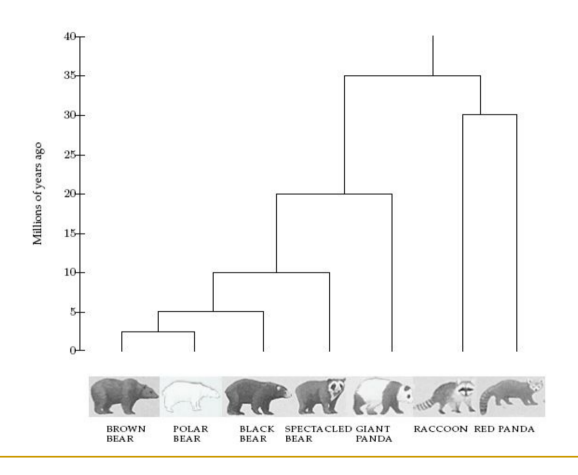
## Early Evolutionary Studies

- Anatomical features were the dominant criteria used to derive evolutionary relationships between species since Darwin till early 1960s
- The evolutionary relationships derived from these relatively subjective observations were often inconclusive. Some of them were later proved incorrect

# Evolution and DNA Analysis: the Giant Panda Riddle

- For roughly 100 years scientists were unable to figure out which family the giant panda belongs to
- Giant pandas look like bears but have features that are unusual for bears and typical for raccoons, e.g., they do not hibernate
- In 1985, Steven O'Brien and colleagues solved the giant panda classification problem using DNA sequences and algorithms

#### Evolutionary Tree of Bears and Raccoons



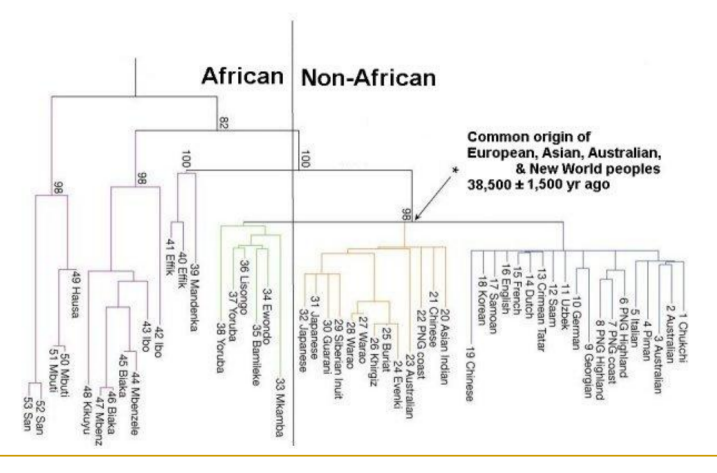
#### Evolutionary Trees: DNA-based Approach

- 40 years ago: Emile Zuckerkandl and Linus Pauling brought reconstructing evolutionary relationships with DNA into the spotlight
- In the first few years after Zuckerkandl and Pauling proposed using DNA for evolutionary studies, the possibility of reconstructing evolutionary trees by DNA analysis
- Now it is a dominant approach to study evolution.

## Out of Africa Hypothesis

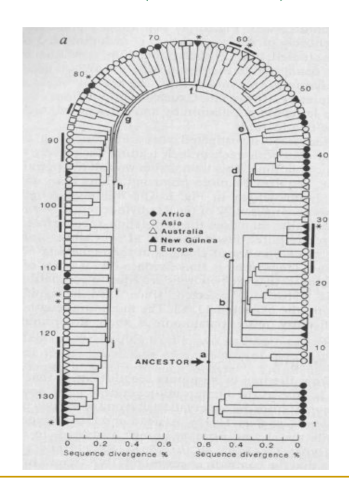
 Around the time the giant panda riddle was solved, a DNA-based reconstruction of the human evolutionary tree led to the Out of Africa Hypothesis that claims our common ancestor lived in Africa roughly 200,000 years ago

## Human Evolutionary Tree (cont'd)



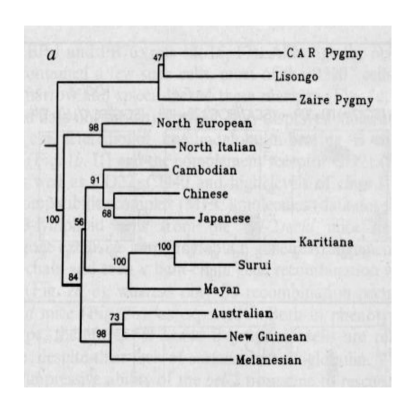
### Evolutionary Tree of Humans (mtDNA)

The evolutionary tree separates one group of Africans from a group containing all five populations.



#### Evolutionary Tree of Humans: (microsatellites)

 Neighbor joining tree for 14 human populations genotyped with 30 microsatellite loci.



## Evolutionary Trees

How are these trees built from DNA sequences?

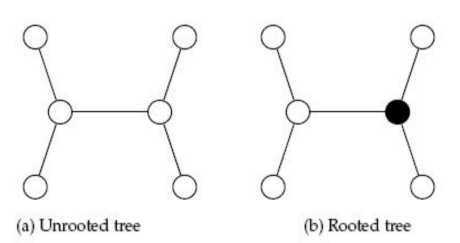
## Evolutionary Trees

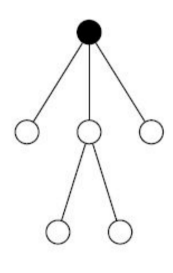
How are these trees built from DNA sequences?

- leaves represent existing species
- internal vertices represent ancestors
- root represents the oldest evolutionary ancestor

#### Rooted and Unrooted Trees

In the unrooted tree the position of the root ("oldest ancestor") is unknown. Otherwise, they are like rooted trees





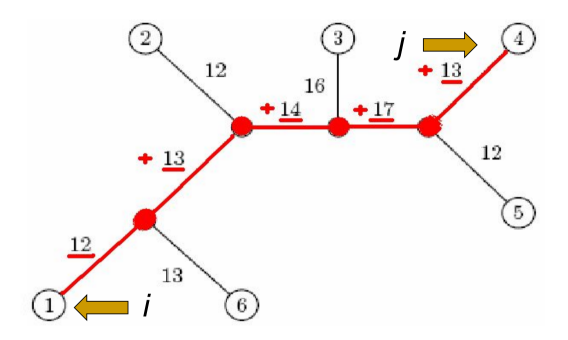
(c) The same rooted tree

#### Distances in Trees

- Edges may have weights reflecting:
  - Number of mutations on evolutionary path from one species to another
  - Time estimate for evolution of one species into another
- In a tree T, we often compute d<sub>ii</sub>(T) - the length of a path between leaves i and j

d<sub>ij</sub>(T) – tree distance between i and j

## Distance in Trees: an Example



$$d_{1,4} = 12 + 13 + 14 + 17 + 12 = 68$$

#### Distance Matrix

- Given n species, we can compute the n x n distance matrix D<sub>ii</sub>
- D<sub>ij</sub> may be defined as the edit distance between a gene in species i and species j, where the gene of interest is sequenced for all n species.

 $D_{ii}$  – edit distance between i and j

#### Edit Distance vs. Tree Distance

- Given n species, we can compute the n x n distance matrix D<sub>ii</sub>
- D<sub>ij</sub> may be defined as the edit distance between a gene in species i and species j, where the gene of interest is sequenced for all n species.
  - $D_{ii}$  edit distance between i and j
- Note the difference with
  - $d_{ij}(T)$  tree distance between i and j

## Fitting Distance Matrix

- Given n species, we can compute the n x n distance matrix D<sub>ij</sub>
- Evolution of these genes is described by a tree that we don't know.
- We need an algorithm to construct a tree that best *fits* the distance matrix D<sub>ii</sub>

## Fitting Distance Matrix

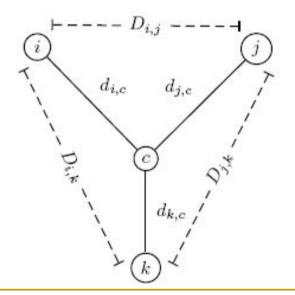
Lengths of path in an (*unknown*) tree T

• Fitting means  $D_{ij} = d_{ij}(T)$ 

Edit distance between species (*known*)

## Reconstructing a 3 Leaved Tree

- Tree reconstruction for any 3x3 matrix is straightforward
- We have 3 leaves i, j, k and a center vertex c



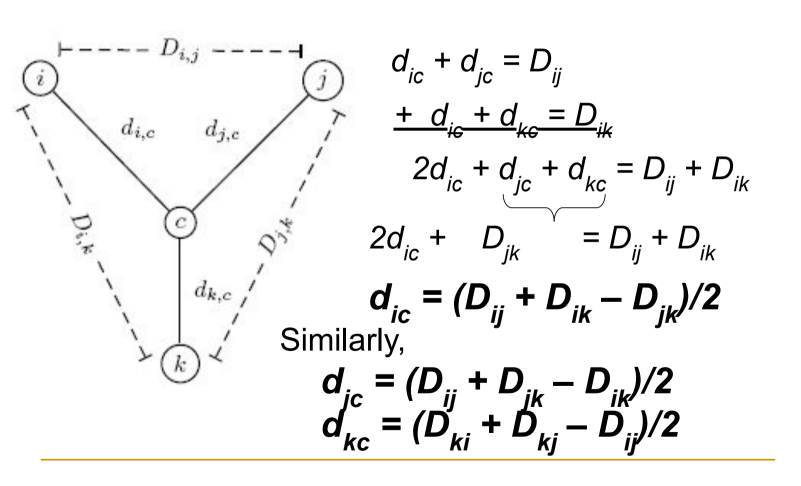
#### Observe:

$$d_{ic} + d_{jc} = D_{ij}$$

$$d_{ic} + d_{kc} = D_{ik}$$

$$d_{jc} + d_{kc} = D_{jk}$$

### Reconstructing a 3 Leaved Tree (cont'd)



## Trees with > 3 Leaves

A tree with n leaves has 2n-3 edges

This means fitting a given tree to a distance matrix D requires solving a system of "n choose 2" equations with 2n-3 variables

This is not always possible to solve optimally for n > 3

## Distance Based Phylogeny Problem

- Goal: Reconstruct an evolutionary tree from a distance matrix
- Input: n x n distance matrix D<sub>ij</sub>
- Output: weighted tree T with n leaves fitting D

If D is additive, this problem has a solution and there is a simple algorithm to solve it

### **UPGMA**

# UPGMA: Unweighted Pair Group Method with Arithmetic Mean

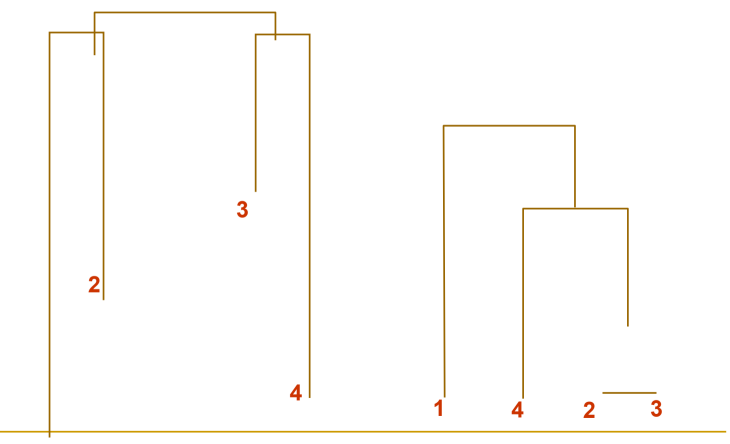
- UPGMA is a clustering algorithm that:
  - computes the distance between clusters using average pairwise distance
  - assigns a *height* to every vertex in the tree, effectively assuming the presence of a molecular clock and dating every vertex

#### UPGMA's Weakness

- The algorithm produces an ultrametric tree: the distance from the root to any leaf is the same
  - UPGMA assumes a constant molecular clock: all species represented by the leaves in the tree are assumed to accumulate mutations (and thus evolve) at the same rate. This is a major pitfalls of UPGMA.



## UPGMA's Weakness: Example



## Clustering in UPGMA

Given two disjoint clusters  $C_{i}$ ,  $C_{j}$  of sequences,

$$d_{ij} = \frac{\sum_{\{p \in Ci, q \in Cj\}} d_{pq}}{|C_i| \times |C_j|}$$
# of elements in i character between rands

Note that if  $C_k = C_j \cup C_j$ , then distance to another cluster  $C_i$  is:

$$d_{kl} = \frac{d_{il} |C_i| + d_{jl} |C_j|}{|C_i| + |C_i|}$$

# UPGMA Algorithm

#### **Initialization:**

Assign each  $x_i$  to its own cluster  $C_i$ 

Define one leaf per sequence, each at height 0

#### **Iteration:**

Find two clusters  $C_i$  and  $C_i$  such that  $d_{ii}$  is min

Let  $C_k = C_i \cup C_j$ 

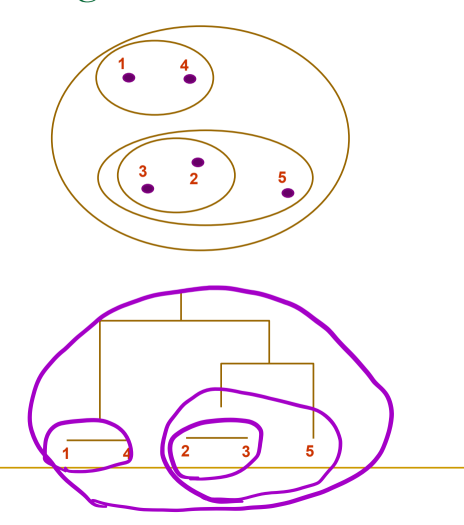
Add a vertex connecting  $C_{i'}$ ,  $C_{j}$  and place it at height  $d_{ij}/2$ 

Delete  $C_i$  and  $C_j$ 

#### **Termination:**

When a single cluster remains

# UPGMA Algorithm (cont'd)



Constructing the phylogenetic tree construction using UPGMA for the following score matrix:

	Α	В	С	D	Е	F
Α						
В	19					
С	27	31				
D	8	18	26			
E	33	36	41	31		
F	18	1	32	17	35	
	(A)	D) (BF				

	A	В	С	D	Ε	F
Α						
В	19					
С	27	31				
D	8	18	26			
Е	33	36	41	31		
F	18	$\left(\begin{array}{c}1\end{array}\right)$	32	17	35	

$$h_{BF} = \frac{1}{2} = 0.5$$

$$C = \{A,C, D, E, (B,F)\}$$

$$D_{BF, A} = \frac{1}{|\{B, F\}|^* |\{A\}|} D_{B, A} + D_{F, A}$$
$$= \frac{1}{2} (19 + 18) = 18.5$$

	Α	BF	С	D	Е
Α					
BF	18.5				
С	27	31.5			
D	8	17.5 <sub>c</sub>	26		
Е	33	35.5	41	31	

$$C = \{(A,D),C, E, (B,F)\}$$

	Α	BF	С	D	Е
Α					
BF	18.5				
С	27	31.5			
D	(8)	17.5	26		
Ε	33	35.5	41	31	

$$D_{BF, AD} = \frac{\left(D_{B, A} + D_{F, A} + D_{B, D} + D_{F, D}\right)}{|\{B, F\}| * |\{A, D\}|}$$
$$= \frac{\left(19 + 18 + 18 + 17\right)}{4} = 18$$

$h_{BF} = \frac{1}{2} = 0.5$
"BF /2 0.0
$h_{AD}^{-1} = 8/2 = 4$
···AD •/-
$h_{ADBF}^{AB} = 18/2 = 9$
··ADBF · · · ·

	AD	BF	С	Е
AD				
BF	(18)			
С	26.5	31.5		
Е	32	35.5	41	

$$C = \{((B,F),(A,D)),C,E\}$$

	AD	BF	С	Е
AD				
BF	18			
С	26.5	31.5		
Е	32	35.5	41	

$$D_{ADBF,C} = \frac{\left(D_{BF,C} \mid \{B,F\} \mid +D_{AD,C} \mid \{A,D\} \mid\right)}{\mid \{B,F\} \mid + \mid \{A,D\} \mid}$$
$$= \frac{\left(31.5 * 2 + 26.5 * 2\right)}{4} = 29$$

	ADBF	С	Е
ADBF			
С	29		
E	33.75	41	

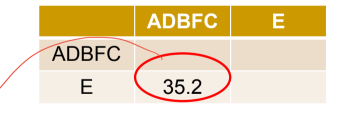
$$C = \{(((B,F),(A,D)),C), E\}$$

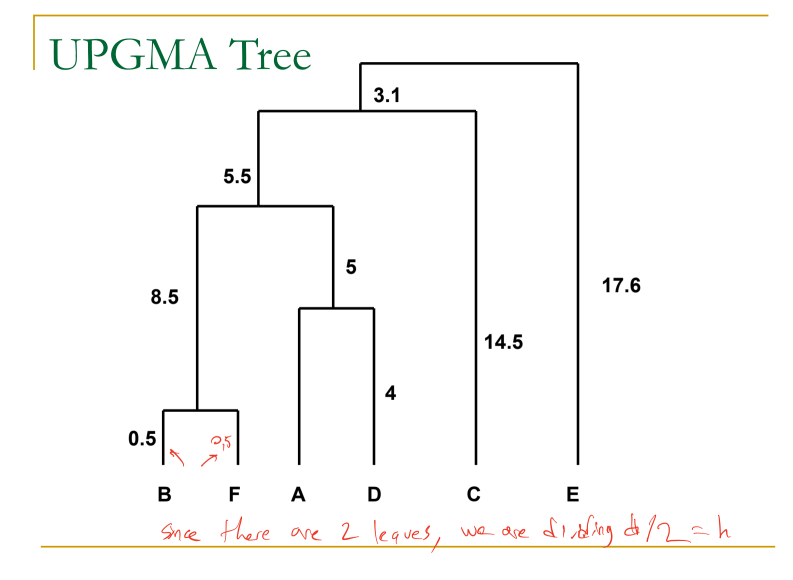
$$D_{ADBFC, E} = \frac{\left(D_{ADBF, E} \mid \{A, B, D, F\} \mid +D_{C, E} \mid \{C\} \mid\right)}{\mid \{A, B, D, F\} \mid + \mid \{C\} \mid}$$

	ADBF	С	E
ADBF			
С	(29)		
Е	33.75	41	

$$=\frac{(33.75*4+41*1)}{5}=35.2$$

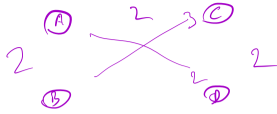
$h_{BF} = \frac{1}{2} = 0.5$
$h_{AB} = 8/2 = 4$
$h_{ADRE} = 18/2 = 9$
$h_{ADBFC}^{ADBFC} = 29/2 = 14.5$
$h_{ADBFCE}^{ADBFCE} = 35.2/2 = 17.0$





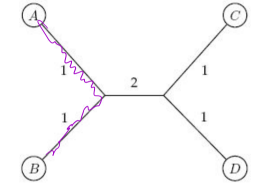
# **NEIGHBOR JOINING**

### Additive Distance Matrices



Matrix D is ADDITIVE if there exists a tree T with  $d_{ij}(T) = D_{ij}$ 

	A	В	C	D
Α	0	2	4	4
В	2	0	4	4
A B C D	4	4	0	2
D	4	4	2	0



NON-ADDITIVE otherwise

	A	В	C	D
A	0	2	2	2
В	2	0	3	2
C	2	3	0	2
D	2	2	2	0

? addithe

#### Using Neighboring Leaves to Construct the Tree

- Find neighboring leaves i and j with parent k
- Remove the rows and columns of i and j
- Add a new row and column corresponding to k, where the distance from k to any other leaf m can be computed as:

$$D_{km} = (D_{im} + D_{jm} - D_{ij})/2$$

Compress *i* and *j* into *k*, iterate algorithm for rest of tree

# Finding Neighboring Leaves

 To find neighboring leaves we simply select a pair of closest leaves.

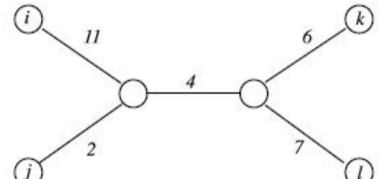
# Finding Neighboring Leaves

 To find neighboring leaves we simply select a pair of closest leaves.

### **WRONG**

# Finding Neighboring Leaves

- Closest leaves aren't necessarily neighbors
- *i* and *j* are neighbors, but  $(d_{ij} = 13) > (d_{jk} = 12)$



 Finding a pair of neighboring leaves is a nontrivial problem!

# Neighbor Joining Algorithm

- In 1987 Naruya Saitou and Masatoshi Nei developed a neighbor joining algorithm for phylogenetic tree reconstruction
- Finds a pair of leaves that are close to each other but far from other leaves: implicitly finds a pair of neighboring leaves
- Advantages: works well for additive and other non-additive matrices, it does not have the flawed molecular clock assumption

# Degenerate Triples

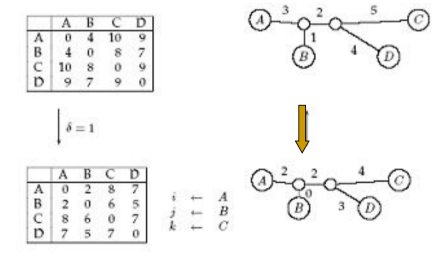
- A degenerate triple is a set of three distinct elements 1≤i,j,k≤n where D<sub>ij</sub> + D<sub>jk</sub> = D<sub>ik</sub>
- Element j in a degenerate triple i,j,k lies on the evolutionary path from i to k (or is attached to this path by an edge of length 0).

# Looking for Degenerate Triples

- If distance matrix D has a degenerate triple i,j,k then j can be "removed" from D thus reducing the size of the problem.
- If distance matrix D does not have a degenerate triple i,j,k, one can "create" a degenerate triple in D by shortening all hanging edges (in the tree).

# Shortening Hanging Edges to Produce Degenerate Triples

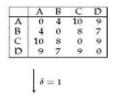
 Shorten all "hanging" edges (edges that connect leaves) until a degenerate triple is found



# Finding Degenerate Triples

- If there is no degenerate triple, all hanging edges are reduced by the same amount δ, so that all pair-wise distances in the matrix are reduced by 2δ.
- Eventually this process collapses one of the leaves (when δ = length of shortest hanging edge), forming a degenerate triple i,j,k and reducing the size of the distance matrix D.
- The attachment point for j can be recovered in the reverse transformations by saving D<sub>ij</sub> for each collapsed leaf.

#### Reconstructing Trees for Additive Distance Matrices



# AdditivePhylogeny Algorithm

```
AdditivePhylogeny(D)
       if D is a 2 x 2 matrix
          T = tree of a single edge of length D_{1,2}
3.
          return T
4
       if D is non-degenerate
          \delta = trimming parameter of matrix D
6.
          for all 1 \le i \ne j \le n
            D_{ii} = D_{ii} - 2\delta
8.
       else
9
          \delta = 0
10.
```

# AdditivePhylogeny (cont'd)

```
Find a triple i, j, k in D such that D_{ii} + D_{ik} = D_{ik}
 1.
       X = D_{ii}
       Remove j^{th} row and j^{th} column from D
3.
       T = AdditivePhylogeny(D)
 4
       Add a new vertex v to T at distance x from i to k
5
       Add j back to T by creating an edge (v,j) of length 0
6
       for every leaf l in T
7
          if distance from I to v in the tree \neq D_{I,i}
8
            output "matrix is not additive"
9.
            return
10.
       Extend all "hanging" edges by length \delta
11
       return T
12.
```

#### The Four Point Condition

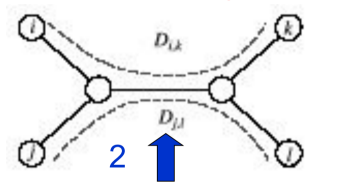
 AdditivePhylogeny provides a way to check if distance matrix D is additive

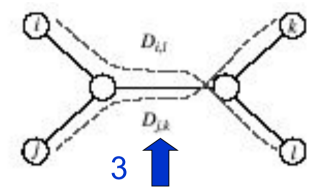
An even more efficient additivity check is the "four-point condition"

Let 1 ≤ i,j,k,l ≤ n be four distinct leaves in a tree

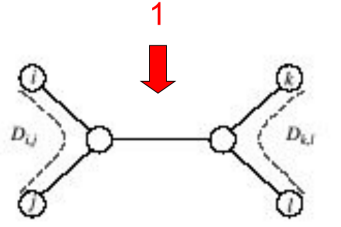
### The Four Point Condition (cont'd)

Compute: 1.  $D_{ij} + D_{kl}$ , 2.  $D_{ik} + D_{jl}$ , 3.  $D_{il} + D_{jk}$ 





2 and 3 represent the same number: the length of all edges + the middle edge (it is counted twice)



1 represents a smaller number: the length of all edges – the middle edge

#### The Four Point Condition: Theorem

The four point condition for the quartet i,j,k,l is satisfied if two of these sums are the same, with the third sum smaller than these first two

Theorem: An n x n matrix D is additive if and only if the four point condition holds for every quartet 1 ≤ i,j,k,l ≤ n

## Least Squares Distance Phylogeny Problem

If the distance matrix D is NOT additive, then we look for a tree T that approximates D the best:

**Squared Error**: 
$$\sum_{i,j} (d_{ij}(T) - D_{ij})^2$$

- Squared Error is a measure of the quality of the fit between distance matrix and the tree: we want to minimize it.
- Least Squares Distance Phylogeny Problem: finding the best approximation tree T for a non-additive matrix D (NP-hard).

### Alignment Matrix vs. Distance Matrix

Sequence a gene of length *m* nucleotides in *n* species to generate an...

n x m alignment matrix

CANNOT be transformed back into alignment matrix because information was lost on the forward transformation

Transform into...

n x n distance matrix

## MAX PARSIMONY

#### Character-Based Tree Reconstruction

#### Better technique:

- Character-based reconstruction algorithms use the n x m alignment matrix
   (n = # species, m = #characters)
   directly instead of using distance matrix.
- GOAL: determine what character strings at internal nodes would best explain the character strings for the n observed species

#### Character-Based Tree Reconstruction (cont'd)

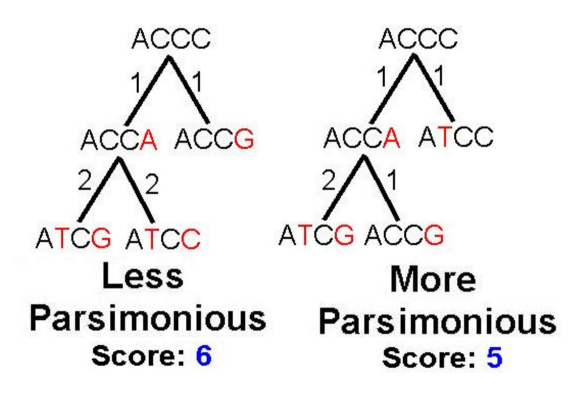
Characters may be nucleotides, where A, G,
 C, T are states of this character.

By setting the length of an edge in the tree to the Hamming distance, we may define the parsimony score of the tree as the sum of the lengths (weights) of the edges

# Parsimony Approach to Evolutionary Tree Reconstruction

- Applies Occam's razor principle to identify the simplest explanation for the data
- Assumes observed character differences resulted from the fewest possible mutations
- Seeks the tree that yields lowest possible parsimony score - sum of cost of all mutations found in the tree

#### Parsimony and Tree Reconstruction



## Small Parsimony Problem

- Input: Tree T with each leaf labeled by an m-character string.
- Output: Labeling of internal vertices of the tree T minimizing the parsimony score.
- Because the characters in the string are independent, the Small Parsimony problem can be solved independently for each character. Therefore, to devise an algorithm, we can assume that every leaf is labeled by a single character rather than by a string of m characters.

### Weighted Small Parsimony Problem

- A more general version of Small Parsimony Problem
- Input includes a k \* k scoring matrix describing the cost of transformation of each of k states into another one
- For Small Parsimony problem, the scoring matrix is based on Hamming distance

$$d_H(v, w) = 0$$
 if  $v=w$   
 $d_H(v, w) = 1$  otherwise

# Scoring Matrices

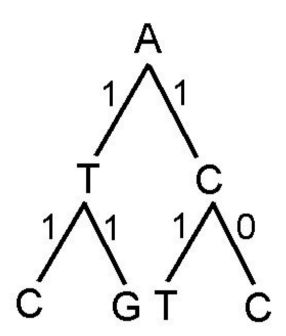
**Small Parsimony Problem** 

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Watashtad	Parsimony	Droblom
vvcivincu	Faisimonv	FIODICHI
1101511000	i wibilitoily	

	Α	Т	G	С
Α	0	1	1	1
Т	1	0	1	1
G	1	1	0	1
С	1	1	1	0

	Α	T	G	С
Α	0	3	4	9
Т	3	0	2	4
G	4	2	0	4
С	9	4	4	0

# Unweighted vs. Weighted

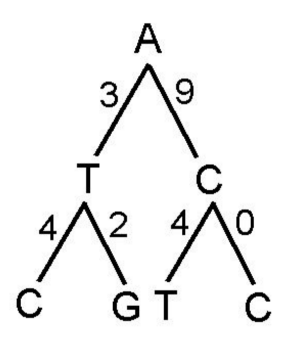


Small Parsimony Scoring Matrix:

	Α	Т	G	С
Α	0	1	1	1
Т	1	0	1	1
G	1	1	0	1
С	1	1	1	0

Small Parsimony Score: 5

# Unweighted vs. Weighted



Weighted Parsimony Scoring Matrix:

	Α	T	G	C
Α	0	3	4	9
Т	3	0	2	4
G	4	2	0	4
С	9	4	4	0

Weighted Parsimony Score: 22

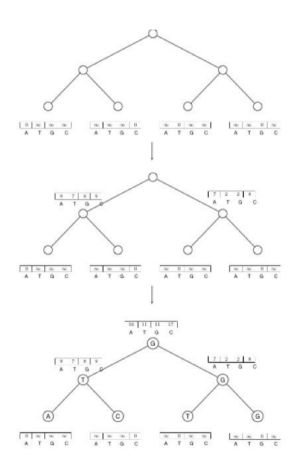
# Weighted Small Parsimony Problem: Formulation

- Input: Tree T with each leaf labeled by elements of a k-letter alphabet and a k x k scoring matrix (δ<sub>ii</sub>)
- Output: Labeling of internal vertices of the tree T minimizing the weighted parsimony score

## Sankoff's Algorithm

- Check children's every vertex and determine the minimum between them
- An example

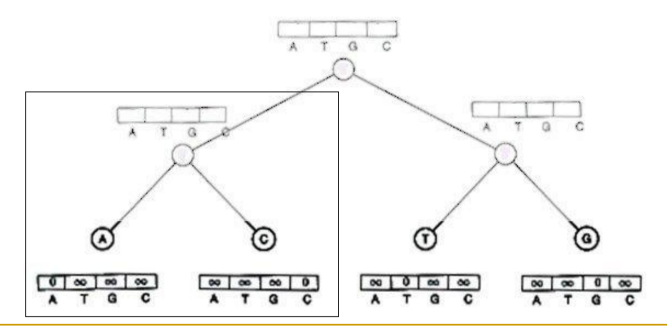
5	A	T	G	C
A	0	3	4	9
T	3	0	2	4
G	4	2	0	4
C	9	4	4	0

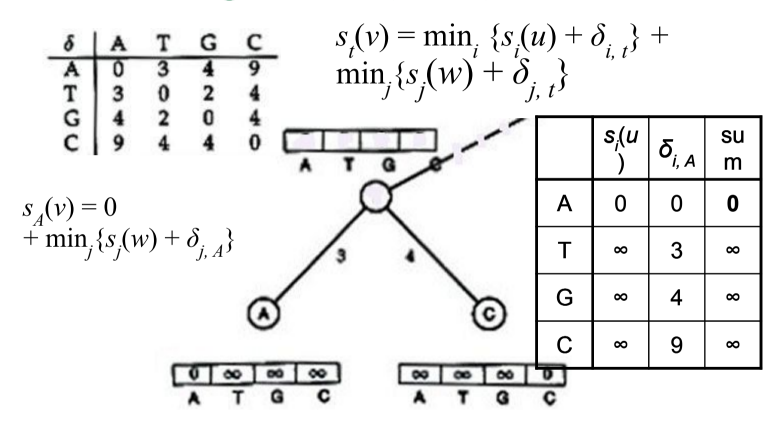


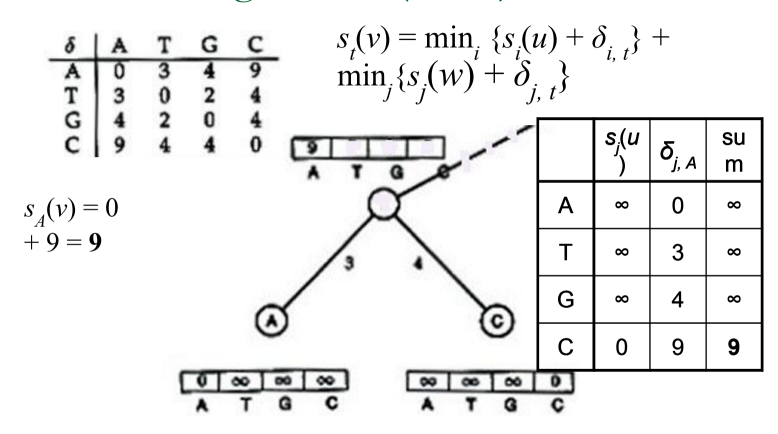
# Sankoff Algorithm: Dynamic Programming

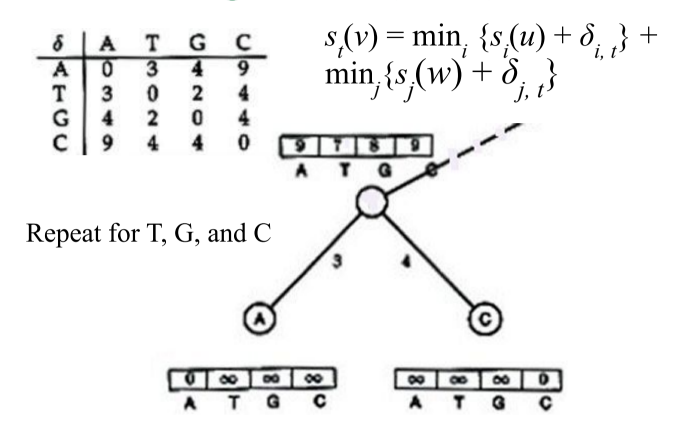
- Calculate and keep track of a score for every possible label at each vertex
  - $s_t(v)$  = minimum parsimony score of the **subtree** rooted at vertex v if v has character t
- The score at each vertex is based on scores of its children:
  - $s_{t}(parent) = \min_{i} \{s_{i}(left child) + \delta_{i,t}\} + \min_{j} \{s_{j}(left child) + \delta_{j,t}\}$

- Begin at leaves:
  - If leaf has the character in question, score is 0
  - □ Else, score is ∞

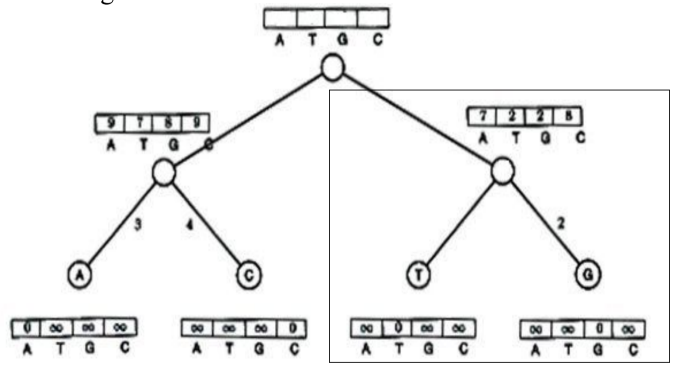




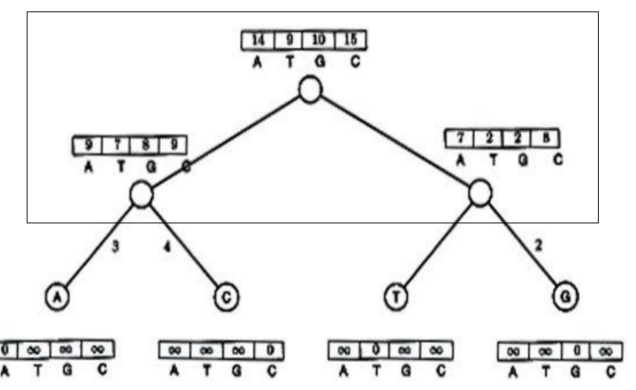




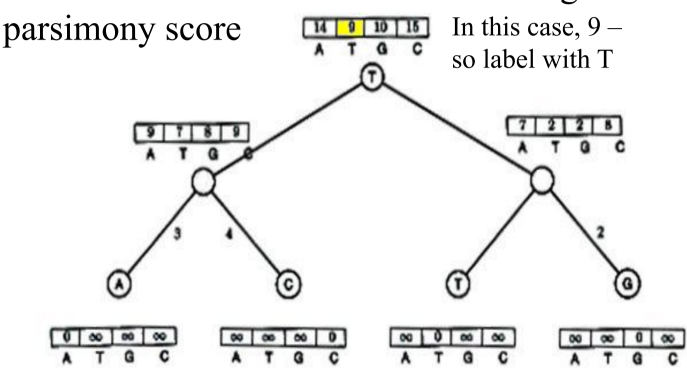
Repeat for right subtree



#### Repeat for root



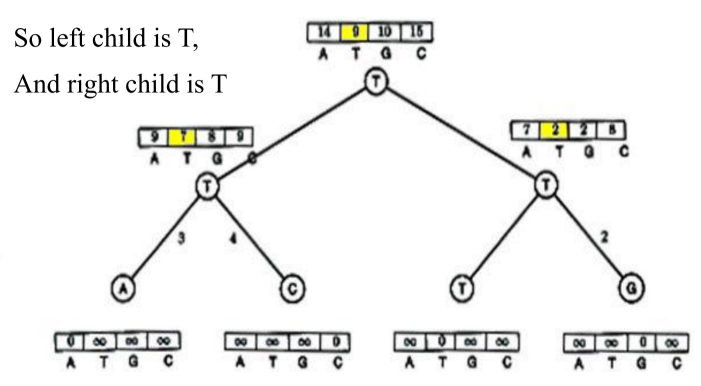
Smallest score at root is minimum weighted



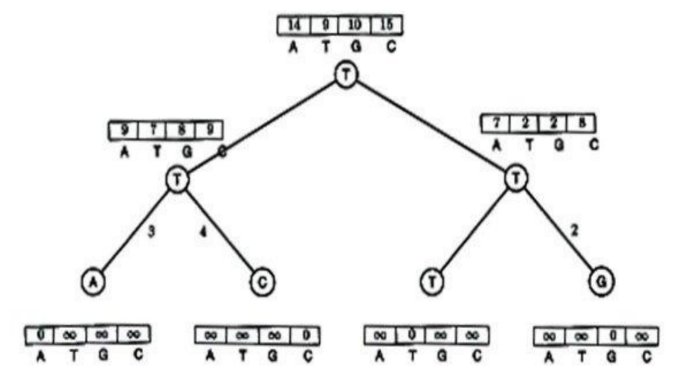
# Sankoff Algorithm: Traveling down the Tree

- The scores at the root vertex have been computed by going up the tree
- After the scores at root vertex are computed the Sankoff algorithm moves down the tree and assign each vertex with optimal character.

9 is derived from 7 + 2



And the tree is thus labeled...

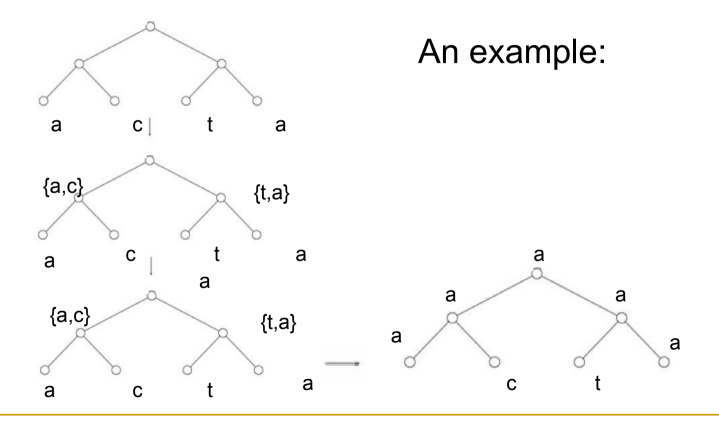


#### FITCH'S ALGORITHM

#### Fitch's Algorithm

- Solves Small Parsimony problem
- Dynamic programming in essence
- Assigns a set of letters to every vertex in the tree.
- If the two children's sets of characters overlap, it's the common set of them
- If not, it's the combined set of them.

## Fitch's Algorithm (cont'd)



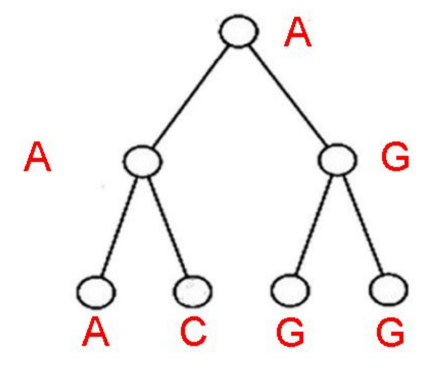
#### Fitch Algorithm

- 1) Assign a **set of possible letters** to every vertex, traversing the tree from leaves to root
- Each node's set is the combination of its children's sets (leaves contain their label)
  - E.g. if the node we are looking at has a left child labeled {A, C} and a right child labeled {A, T}, the node will be given the set {A, C, T}

## Fitch Algorithm (cont.)

- 2) Assign **labels** to each vertex, traversing the tree from root to leaves
- Assign root arbitrarily from its set of letters
- For all other vertices, if its parent's label is in its set of letters, assign it its parent's label
- Else, choose an arbitrary letter from its set as its label

## Fitch Algorithm (cont.)



#### Fitch vs. Sankoff

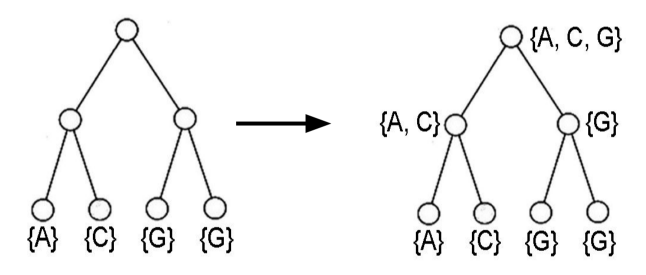
Both have an O(nk) runtime

Are they actually different?

Let's compare ...

#### Fitch

As seen previously:



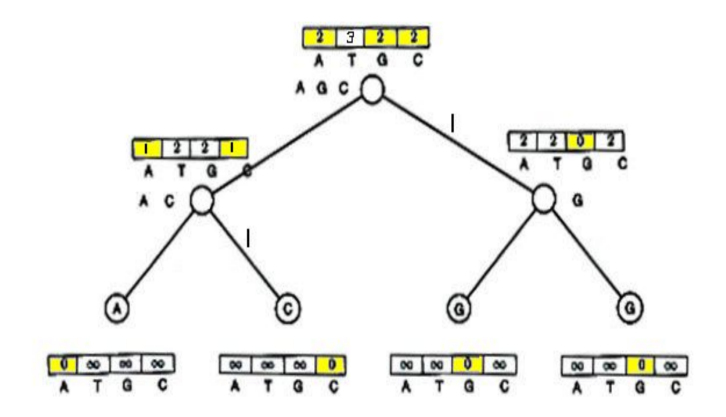
## Comparison of Fitch and Sankoff

As seen earlier, the scoring matrix for the Fitch algorithm is merely:

	Α	Т	G	С
Α	0	1	1	1
Т	1	0	1	1
G	1	1	0	1
С	1	1	1	0

 So let's do the same problem using Sankoff algorithm and this scoring matrix

#### Sankoff



#### Sankoff vs. Fitch

- The Sankoff algorithm gives the same set of optimal labels as the Fitch algorithm
- For Sankoff algorithm, character t is optimal for vertex v if s<sub>t</sub>(v) = min<sub>1≤i≤k</sub>s<sub>i</sub>(v)
  - $\Box$  Denote the set of optimal letters at vertex v as S(v)
    - If S(left child) and S(right child) overlap, S(parent) is the intersection
    - Else it's the union of S(left child) and S(right child)
    - This is also the Fitch recurrence
- The two algorithms are identical

#### Large Parsimony Problem

- Input: An n x m matrix M describing n species, each represented by an m-character string
- Output: A tree T with n leaves labeled by the n rows of matrix M, and a labeling of the internal vertices such that the parsimony score is minimized over all possible trees and all possible labelings of internal vertices

#### Large Parsimony Problem (cont.)

- Possible search space is huge, especially as n increases
  - $\square$  (2*n* 3)! possible rooted trees
  - $\square$  (2*n* 5)! possible unrooted trees
- Problem is NP-complete
  - $\Box$  Exhaustive search only possible w/ small n(< 10)
- Hence, branch and bound or heuristics used