CS 466 Introduction to Bioinformatics Lecture 15

Mohammed El-Kebir October 14, 2020



Course Announcements

HW 3 will be released Oct 23 – due Oct 31 by 11:59pm

Project proposal due on Nov 5 by 11:59pm (Motivation, Datasets/papers, Planned method/experiments, Timeline)

Project report due on Dec 20

Outline

- Recap: RNA Secondary Structure Prediction
- Phylogenetics introduction
- Hierarchical clustering
- Additive distance phylogeny
- Four point condition
- Neighbor joining

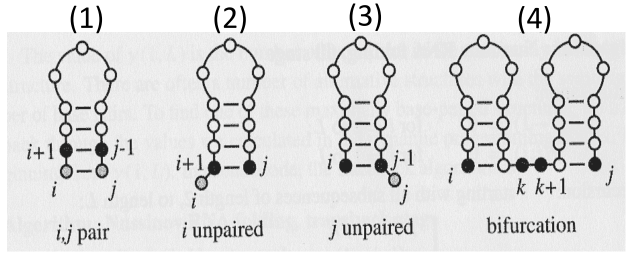
Reading:

• Chapter 10.2 and 10.5-10.8 in Jones and Pevzner

Nussinov Algorithm – Dynamic Programming

Problem: Given RNA sequence $\mathbf{v} \in \{A, U, C, G\}^n$, find a pseudoknot-free secondary structure with the maximum number of complementary base pairings

Let s[i, j] denote the maximum number of pseudoknot-free complementary base pairings in subsequence v_i, \dots, v_i

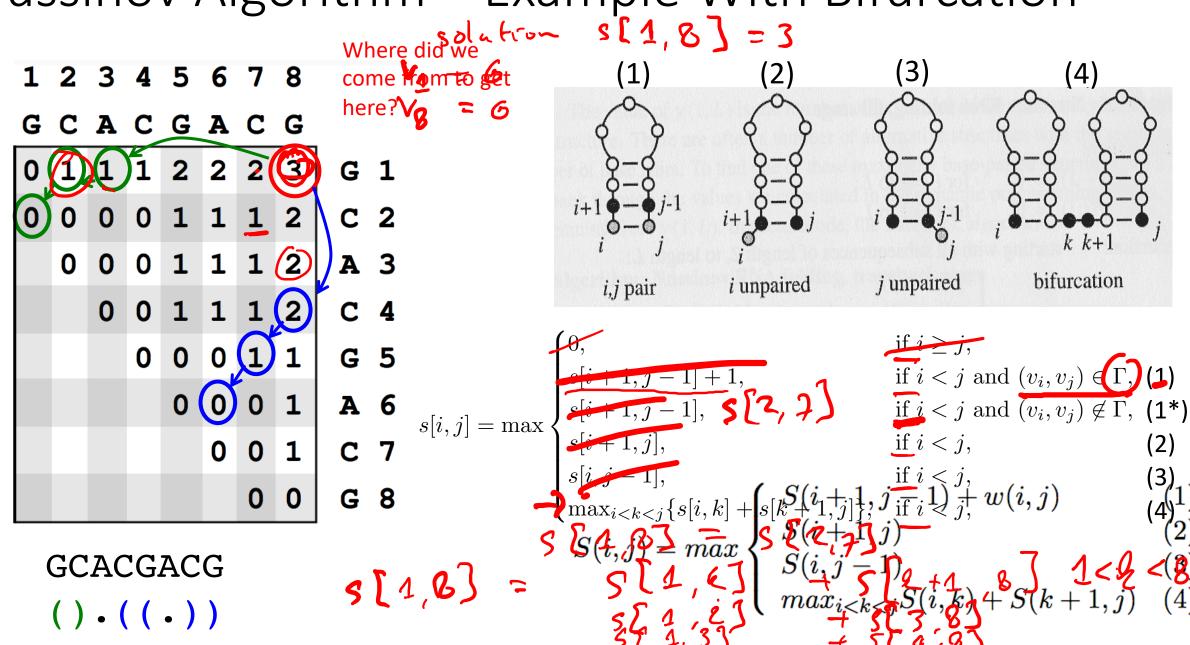


$$s[i,j] = \max egin{cases} 0, & ext{if } i \geq j, \ s[i+1,j-1]+1, & ext{if } i < j ext{ and } (v_i,v_j) \in \Gamma, \ s[i+1,j-1], & ext{if } i < j ext{ and } (v_i,v_j)
otin \Gamma, \ s[i+1,j], & ext{if } i < j, \ s[i,j-1], & ext{if } i < j, \ max_{i < k < j} \{s[i,k]+s[k+1,j]\}, & ext{if } i < j, \ S(i,j) = max \ \end{cases} egin{cases} 3 (i+1,j) \ (4)(i+1,j) \ S(i,j-1) \end{cases}$$

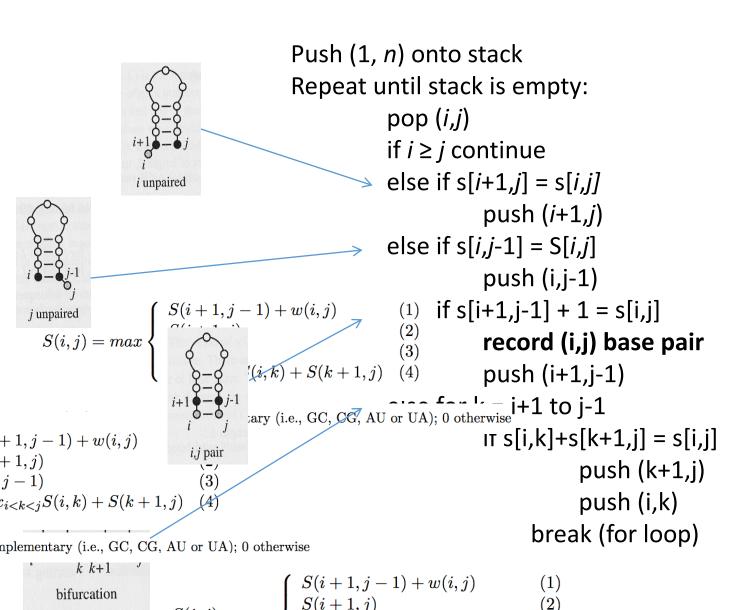
Question:

Which case is redundant?

Nussinov Algorithm – Example With Bifurcation

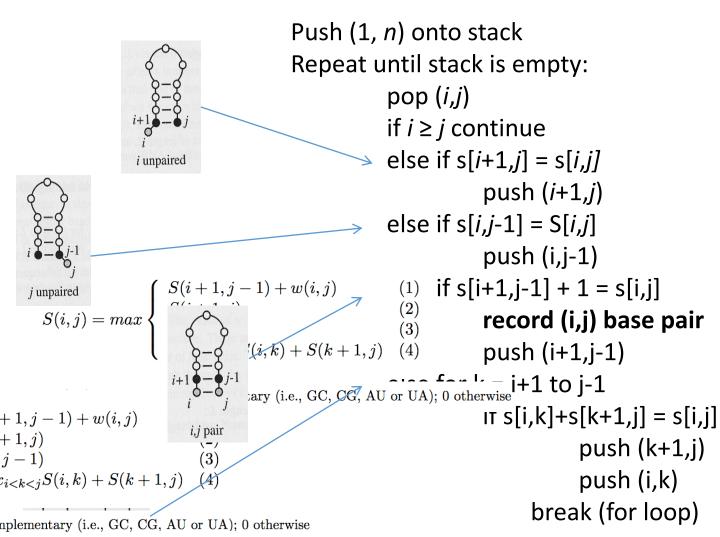


Nussinov Algorithm – Traceback Step



Nussinov Algorithm – Traceback Step

(1)(2)



```
Bach Track (1, n)
```

```
BackTrack(i, j)
if i < j
          if s[i+1, j] = s[i, j]
                     BackTrack(i+1, j)
          else if s[i, j-1] = S[i, j]
                    BackTrack(i, j-1)
          else if s[i+1,j-1] + 1 = s[i, j]
                     Output (i, j)
                     BackTrack(i+1, j-1)
          else for k = i+1 to j-1
                    if s[i, k] + s[k+1, j] = s[i, j]
                               BackTrack(k+1, j)
                               BackTrack(i, k)
                          break (for loop)
```

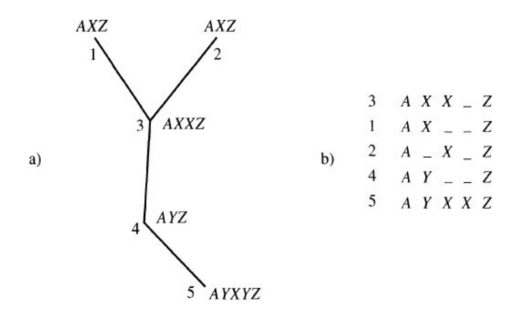
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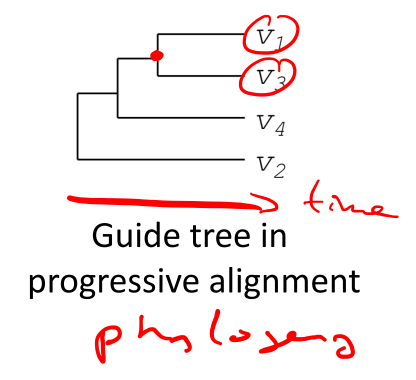
Reading:

Chapter 10.2 and 10.5-10.8 in Jones and Pevzner

Alignments and Trees



Tree / star alignment

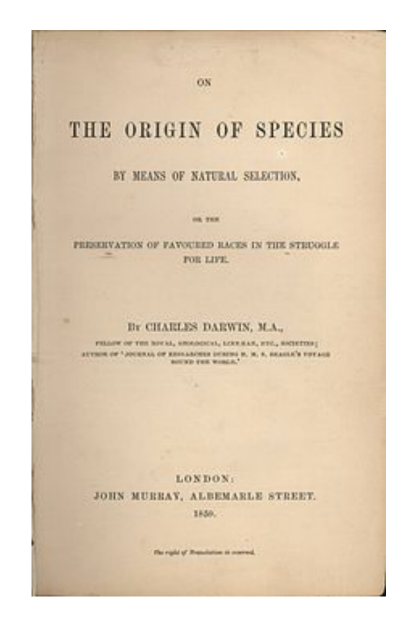


Tree topology represents similarity/distance between sequences

Biological sequences typically come from the present

Evolutionary Studies and Phylogenies

- Since Darwin's book (1859) until 1960s: Phylogeny reconstruction from anatomical features
- Subjective observations led to inconclusive/incorrect phylogenies

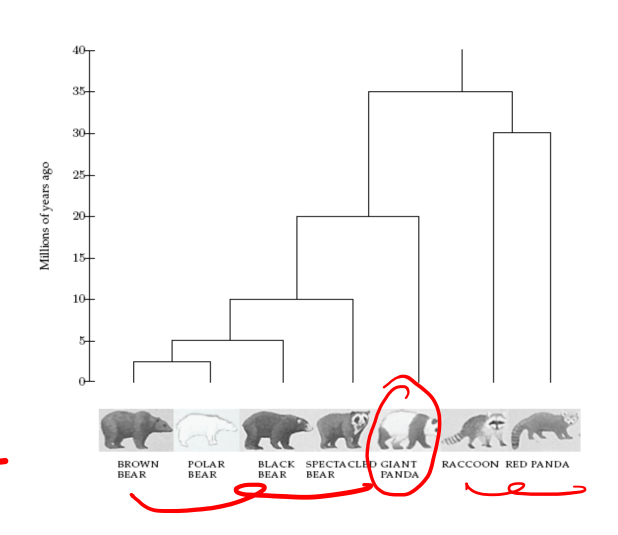


Evolutionary Studies and Phylogenies

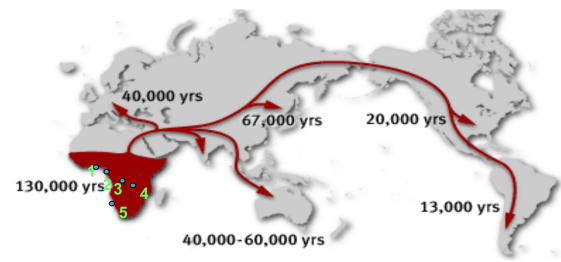
 Subjective observations led to inconclusive/incorrect phylogenies

Example

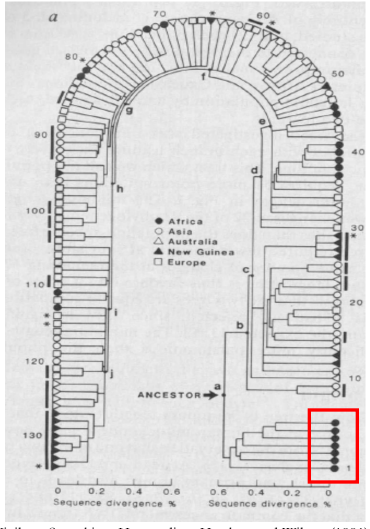
- Giant pandas look like bears but have features that are unusual for bears and typical for racoons
- In 1985, Steven O'Brien and colleagues solved the giant panda classification problem using DNA sequences and algorithms



Out of Africa Hypothesis



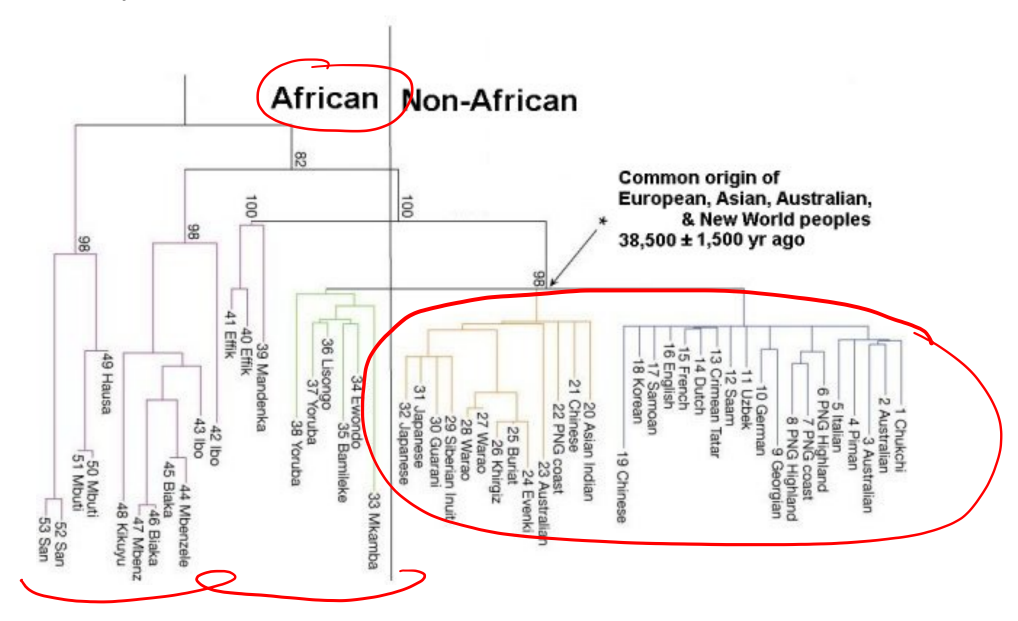
http://www.becominghuman.org



Vigilant, Stoneking, Harpending, Hawkes, and Wilson (1991)

Out of Africa Hypothesis claims that our most ancient ancestor lived in Africa roughly 200,000 years ago

Evolutionary Tree of Humans



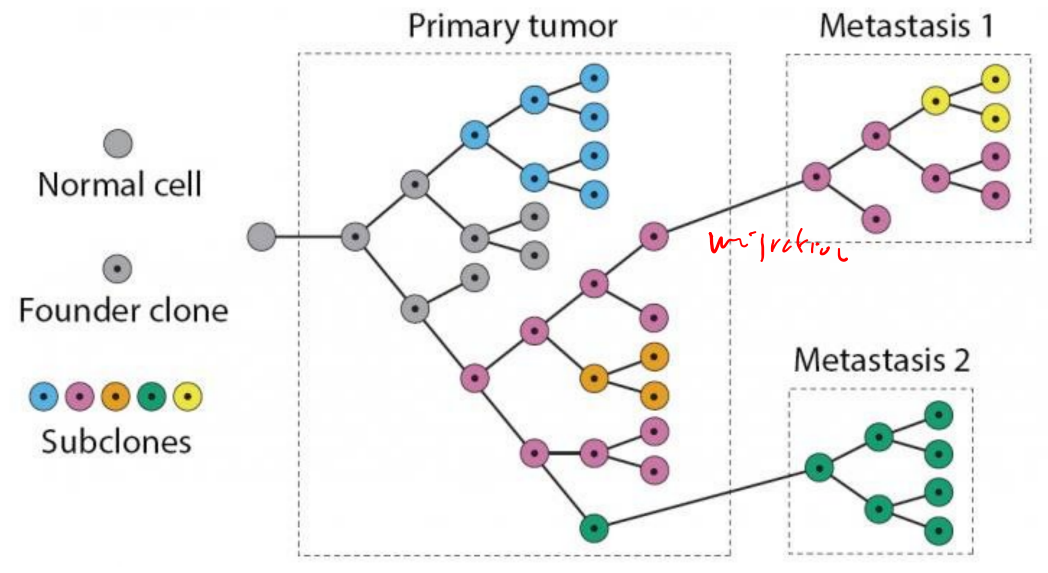
Evolutionary Tree of Species Norwegian Fores Sphynx Maine Siberian Germany Abyssinian American SH British SH Chartreux Millions of years ago F.s. silvestris . F.s. tristami Sokoke (China) Russ. Blue Egypt Turk. Van J. Bobtail Brown Singapore Turkey Tunisia Egypt. Korat RACCOON RED PANDA BEAR BEAR BEAR PANDA

http://bix.ucsd.edu/bioalgorithms/

[Lipinski *et al.*, 2008]

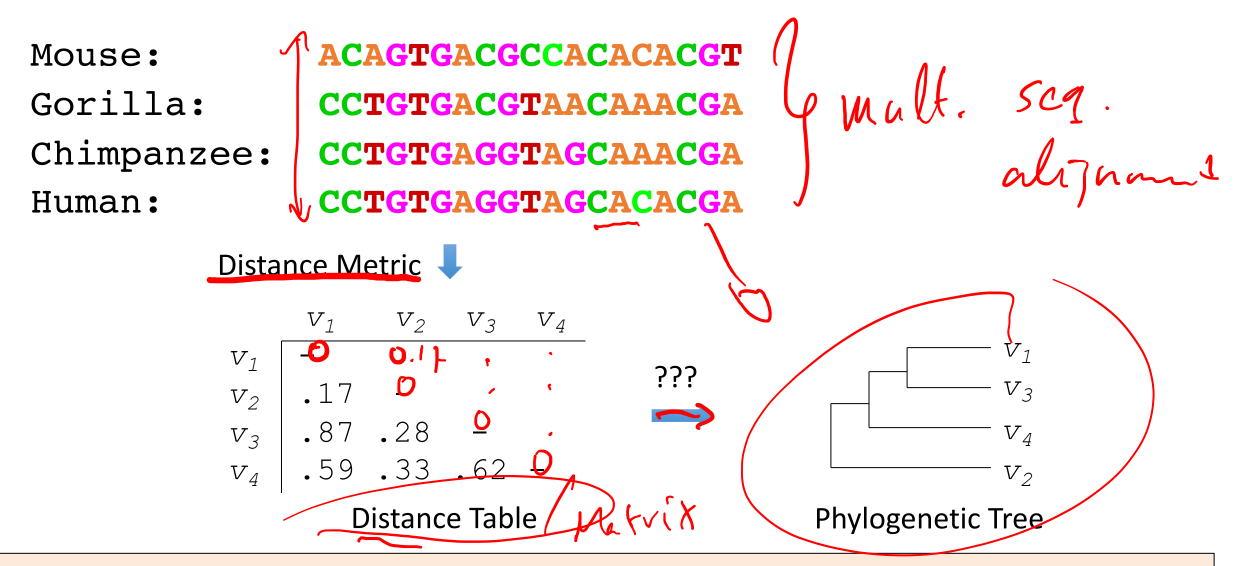
Question: What are the evolutionary relationships between species?

Evolutionary Tree of a Tumor



https://www.sciencedaily.com/releases/2016/09/160909223504.htm

Phylogenetic Tree Reconstruction



Question: Given sequence data, how to reconstruct tree?

Outline

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Reading:

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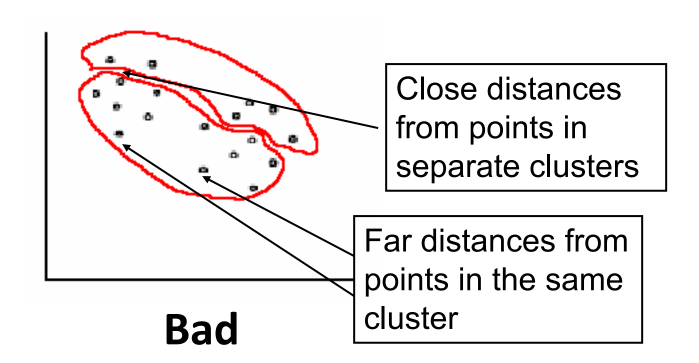
Clustering

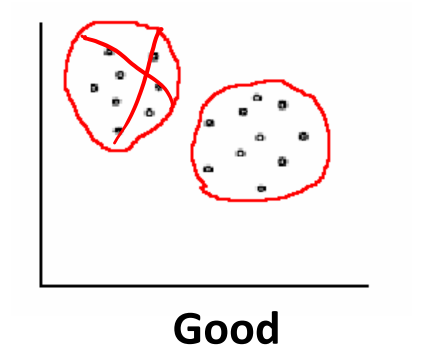
Given:

(1) $n \times n$ matrix $D = [d_{i,j}]$

Want:

- (1) Homogeneity within clusters
- (2) Separation between clusters

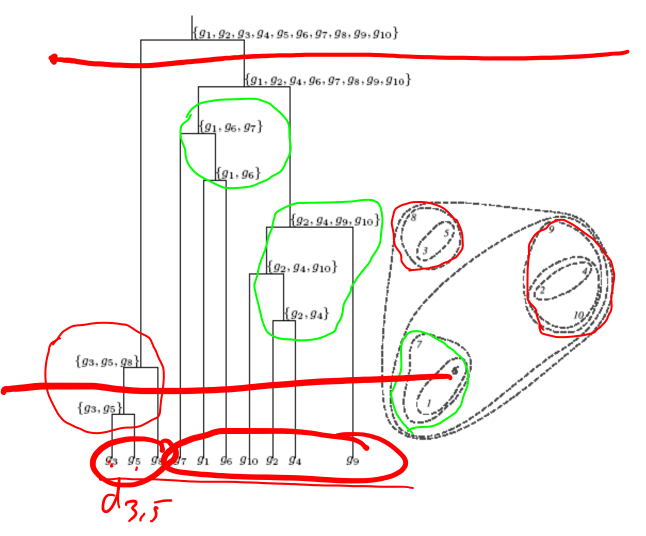




D=[dij]

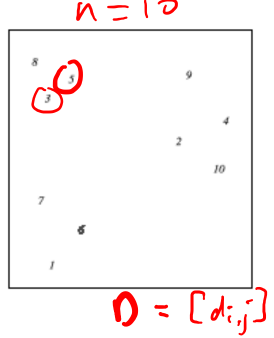
Organize elements into a tree such that:

- Leaves are elements
- Paths between leaves (i, j) represent pairwise element distance
- Similar elements lie within same subtrees



- 1. Hierarchical Clustering (**D**, n)
- 2. Form *n* clusters each with one element
- 3. Construct a graph **T** by assigning one vertex to each cluster
- **4. while** there is more than one cluster
- 5. Find the two closest clusters C_1 and C_2
- 6. Merge C_1 and C_2 into new cluster C with $|C_1| + |C_2|$ elements
- 7. Compute distance from *C* to all other clusters
- 8. Add a new vertex \boldsymbol{C} to \boldsymbol{T} and connect to vertices C_1 and C_2
- 9. Remove rows and columns of D corresponding to C_1 and C_2
- 10. Add a row and column to **D** corresponding to the new cluster **C**
- **11.** return *T*

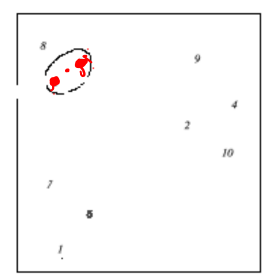




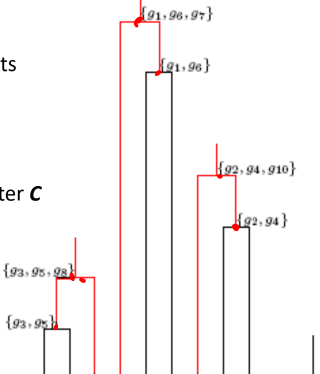
- Hierarchical Clustering (**D**, n)
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- Construct a graph **T** by assigning one vertex to each cluster
- while there is more than one cluster

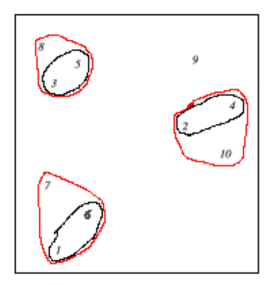
- Find the two closest clusters C_1 and C_2 Merge C_1 and C_2 into new cluster C with $|C_1| + |C_2|$ elements $C_1 = 93$ $C_2 = 95$ $C_3 = 95$
- Compute distance from *C* to all other clusters
- Add a new vertex \boldsymbol{C} to \boldsymbol{T} and connect to vertices C_1 and C_2
- 9. Remove rows and columns of **D** corresponding to C_1 and C_2
- Add a row and column to **D** corresponding to the new cluster **C** 10.
- **11.** return *T*





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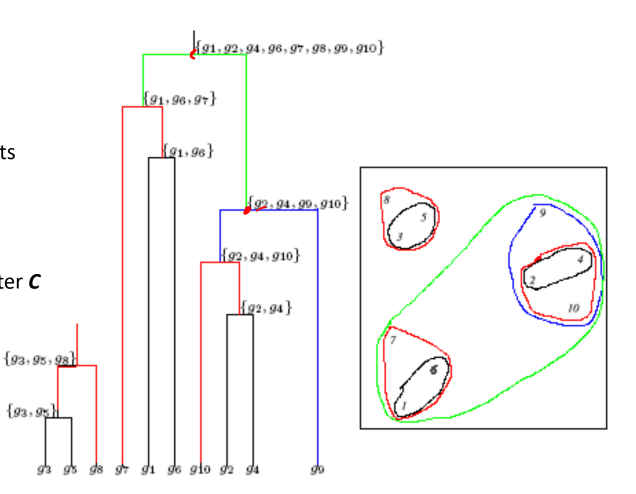




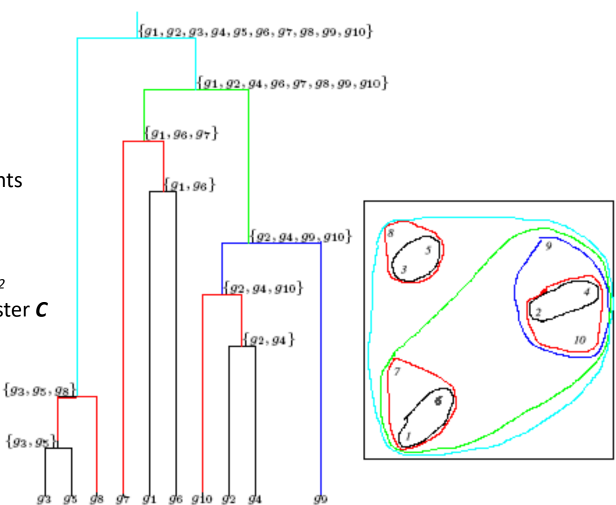
- Hierarchical Clustering (**D**, n)
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- **while** there is more than one cluster 4.
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- Merge C_1 and C_2 into new cluster C with $|C_1| + |C_2|$ elements 6.
- 7. Compute distance from *C* to all other clusters
- Add a new vertex \boldsymbol{C} to \boldsymbol{T} and connect to vertices C_1 and C_2 8.
- 9. Remove rows and columns of **D** corresponding to C_1 and C_2
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 $\{g_3, g_5\}$

11. return *T*

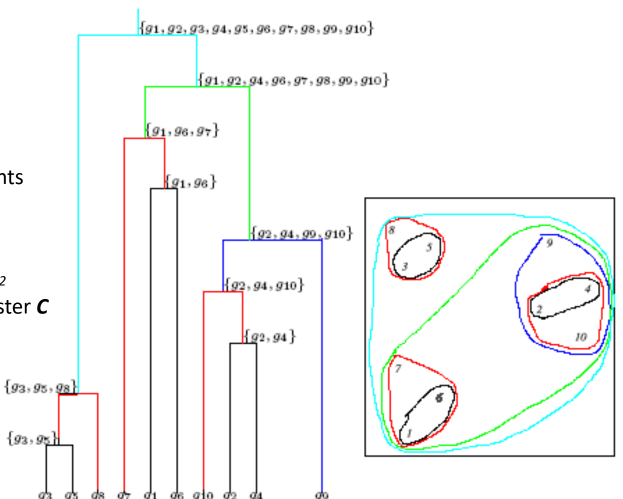


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Definition of distance between clusters affects clustering!



Hierarchical Clustering – Linkage Criteria

	Names	Formula			
	Maximum or complete-linkage clustering	$\max\{d(a,b):a\in A,b\in B\}.$			
	Minimum or single-inkage clustering	$\min\{d(a,b):a\in A,b\in B\}.$			
-> (Mean or average linkage clustering, or UPGMA	$\frac{1}{ A . B }\sum_{a\in A}\sum_{b\in B}d(a,b).$			
	Centroid linkage clustering, or UPGMC	$\ c_s-c_t\ $ where c_s and c_t are the centroids of clusters s and t , respectively.			
	Minimum energy clustering	$\frac{2}{nm} \sum_{i,j=1}^{n,m} \ a_i - b_j\ _2 - \frac{1}{n^2} \sum_{i,j=1}^n \ a_i - a_j\ _2 - \frac{1}{m^2} \sum_{i,j=1}^m \ b_i - b_j\ _2$			

https://en.wikipedia.org/wiki/Hierarchical_clustering#Linkage_criteria

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Phylogenetic Tree Reconstruction

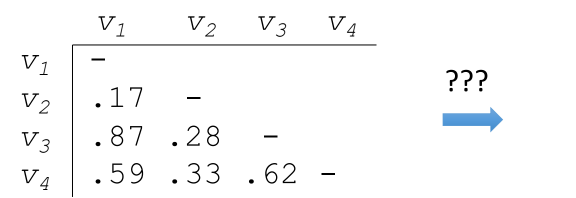
Mouse: ACAGTGACGCCACACGT

Gorilla: CCTGTGACGTAACAACGA

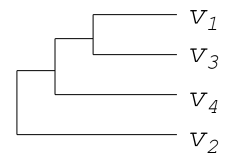
Chimpanzee: CCTGTGAGGTAGCAAACGA

Human: CCTGTGAGGTAGCACGA

Distance Metric -



Distance Table



Phylogenetic Tree

Question: Given sequence data, how to reconstruct tree?

Distance

Examples:

- $X = \mathbb{R}$ and $d(x, y) \in |x y|$
- $X = \Sigma^*$ and d is Hamming distance
- $X = \Sigma^*$ and d is edit distance

Alignment vs. Distance Matrices

 $A \rightarrow T$

T -> A

(-> G

Mouse:

ACAGTGACGCCACACGT

Gorilla:

CCTGTGACGTAACAAACGA

Chimpanzee:

CCTGTGAGGTAGCAAACGA

CCTGTGAGGTAGCACACGA

General of m in n species

Human:

Easy: use (weighted) edit distance

Reverse transformation not possible due to loss of information

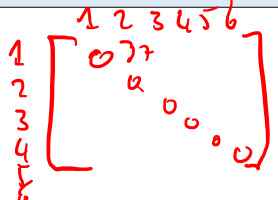
	v_1	V_2	$V_{\mathcal{J}}$	V_4
v_1	0	7	11	10
$egin{array}{c} V_1 \ V_2 \end{array}$	7	0	4	6
V_3	11	4	0	2
V_4	10	6	2	0

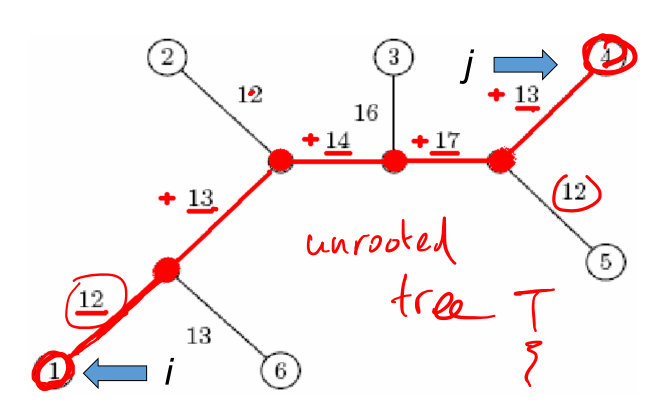
 $n \times n$ distance matrix

Distances in Trees

N=6 le aves /se15.

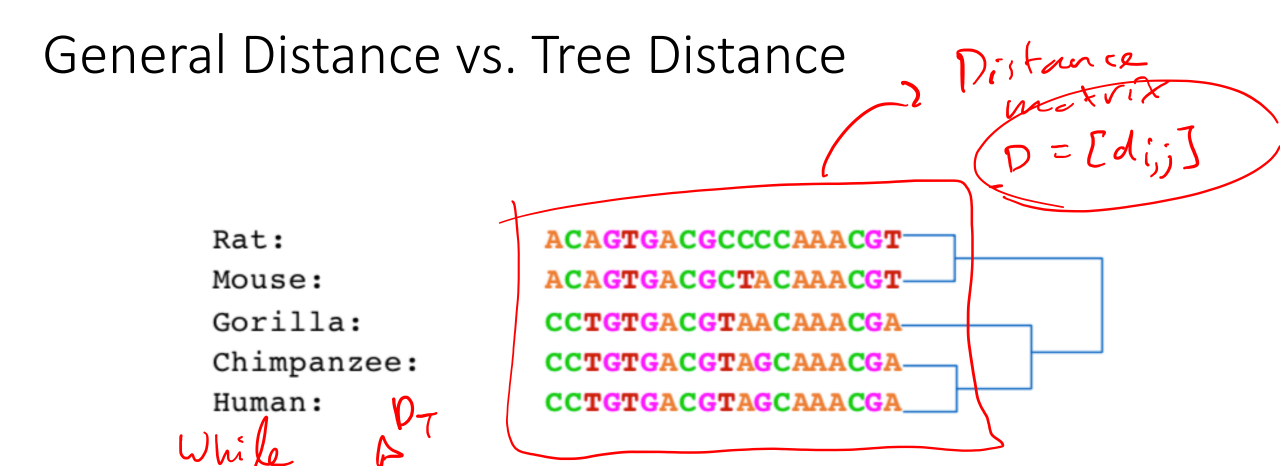
Given a tree T with positive edge weights w(e), tree distance $d_T(i,j)$ between two leaves *i* and *j* is the sum of weights of edges on the unique path from *i* to *j*





$$d_{\tau}(1,4) = 12 + 13 + 14 + 17 + 13 = 69$$

 $d_{\tau}(1,3) = 68$



Tree distance $d_T(i,j)$ not necessarily equal to $d_{i,j}$ as given by distance matrix obtained from alignment











Fitting a Tree to a Given Distance Matrix

- Given n species, we can compute $n \times n$ distance matrix $D = [d_{i,j}]$
- Evolution of these \underline{n} species is described by an unknown tree
- We need an algorithm to construct tree T that best fits D

Fitting a Tree to a Given Distance Matrix

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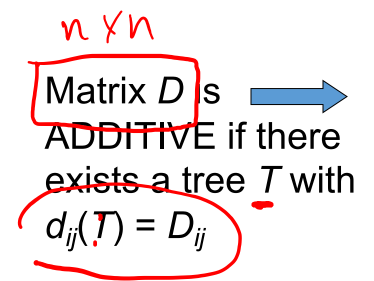
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Distance-Based Phylogeny: Given n \times n distance matrix D = [d_{i,j}], find edge-weighted tree T with n leaves that best fits D

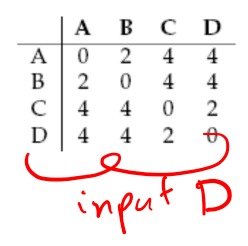
Question: How to define 'best fit'?

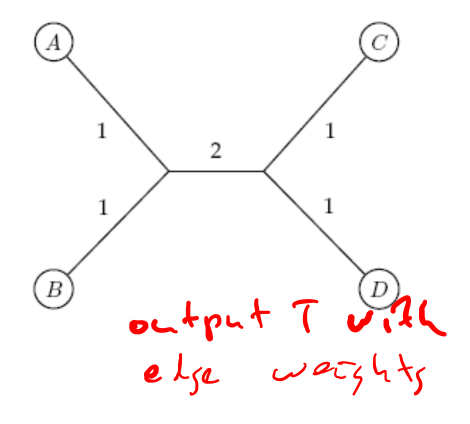
C = [d_{i,j}]

C = [d_{i,j}]
```

Additive Distance Matrices





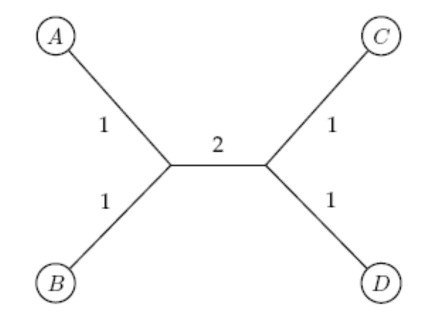


Additive Distance Matrices

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
C	4	4	0	2
A B C D	4	4	2	0



NON-ADDITIVE otherwise

?

This is a constructive definition

Small Additive Distance Phylogeny Problem: Given $n \times n$ distance matrix $D = [d_{i,j}]$ and unweighted tree T with

n leaves, determine edge weights such that $d_T(i,j) = d_{i,j}$

37

Small Additive Distance Phylogeny Problem:

Given $n \times n$ distance matrix $D = [d_{i,j}]$ and unweighted tree T with n leaves, determine edge weights such that $d_T(i,j) = d_{i,j}$

Large Additive Distance Phylogeny Problem:

Given $n \times n$ distance matrix $D = [d_{i,j}]$, find tree T with n leaves **and** edge weights such that $d_T(i,j) = d_{i,j}$

Small Additive Distance Phylogeny Problem:

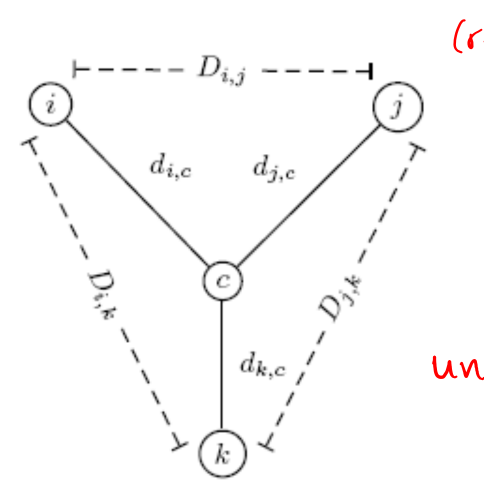
Given $n \times n$ distance matrix $D = [d_{i,j}]$ and unweighted tree T with n leaves, determine edge weights such that $d_T(i,j) = d_{f,j}$

Large Additive Distance Phylogeny Problem:

Given $n \times n$ distance matrix $D = [d_{i,i}]$, find tree T with n leaves and edge weights such that $d_T(i,j) = d_{i,i}$

Both problems can be solved in polynomial time .

Additive Distance Problem with n=3 Sequences



trees (rooted/unrooted) h vertices h-1 edges

rooted barry
trees

N leaves

N-1 intendrating

PM-1 vertices

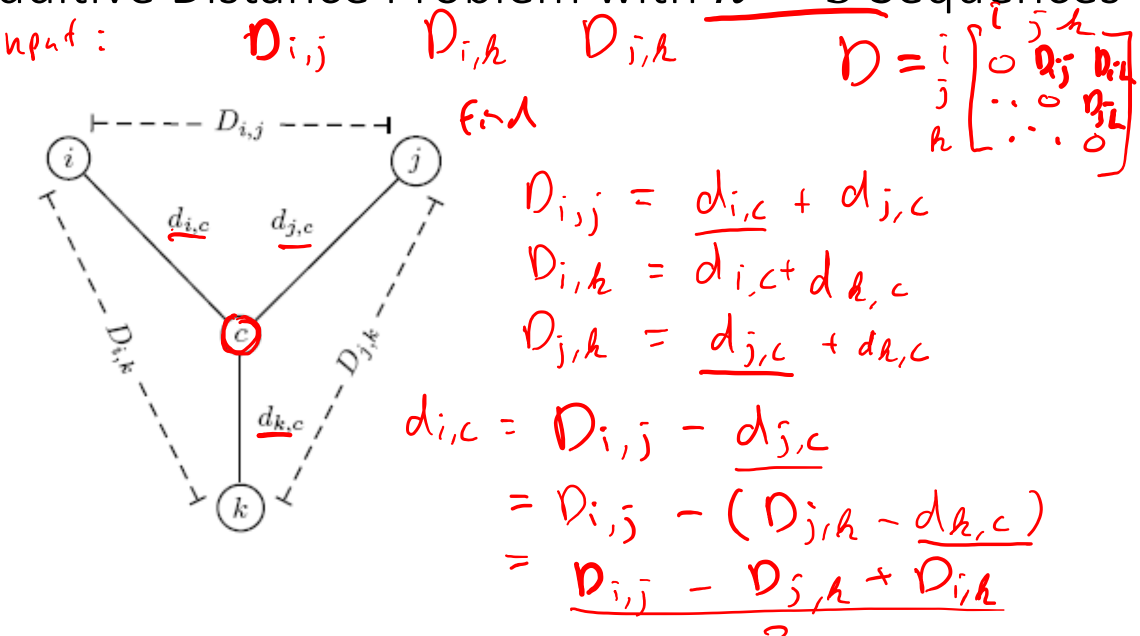
O2n-2 edges

N-2 leaves

unrooted sinary
trees

n learns
n-2 internal vertices
2n-2 vertices
2n-3 edges

Additive Distance Problem with n=3 Sequences



Additive Distance Problem with $n \geqslant 3$ Sequences

Unrooted binary tree with n leaves has 2n-3 edges and

- $\binom{n}{2}$ pairwise distances: 2n 3 variables $\binom{n}{2}$ equations $\binom{n}{2}$



	177
2n-3=	(7)
N=1"	

	A	В	C	D
Α	0	2	2	2
B C	2	0	3	2
C	2	3	0	2
D	2	2	2	0

Solution not always possible for n > 3

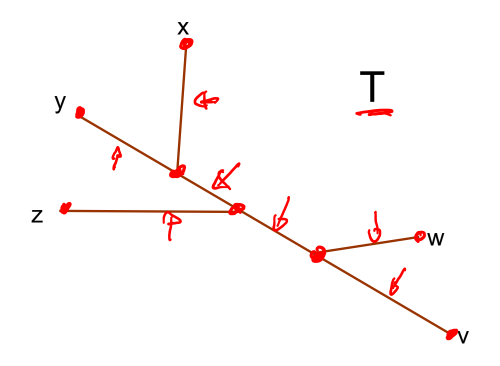
Small Additive Distance Phylogeny Problem:

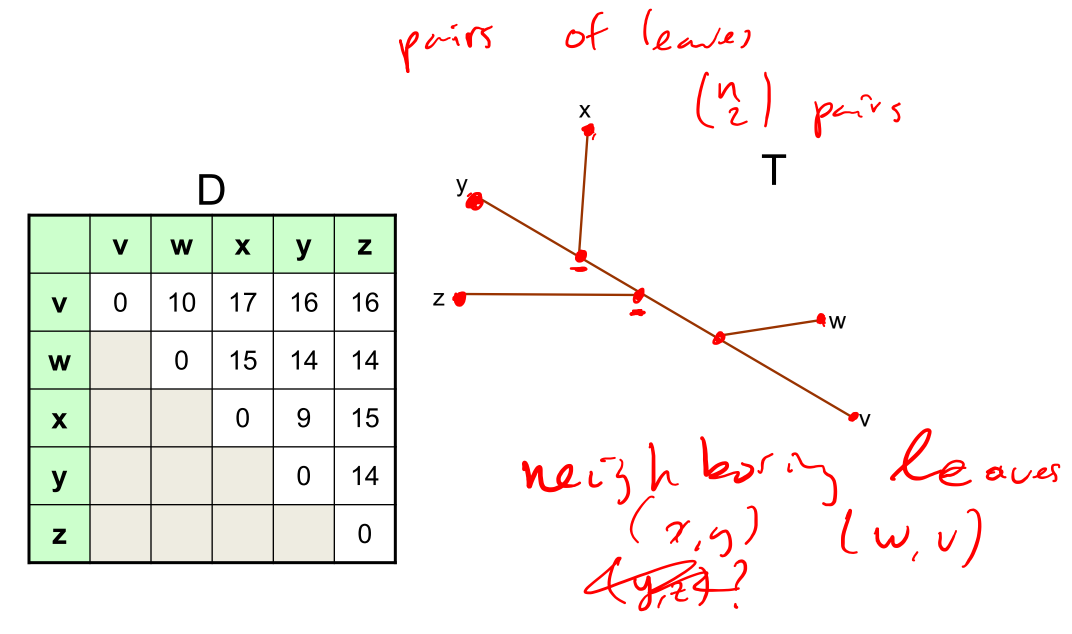
Given $n \times n$ distance matrix $D = [d_{i,j}]$ and unweighted tree T with n leaves, determine edge weights such that $d_T(i,j) = d_{i,j}$



	D
1 n	ynd

	V	W	X	у	Z
V	0	10	17	16	16
w		0	15	14	14
x			0	9	15
У				0	14
Z					0

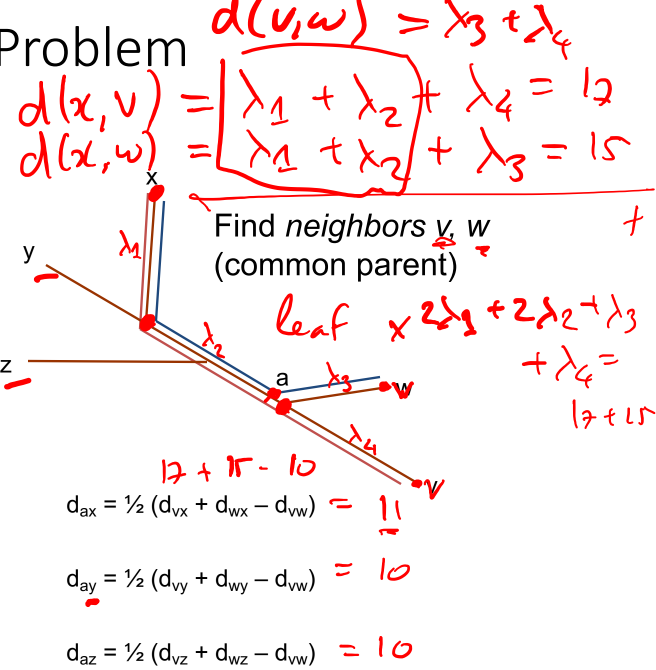


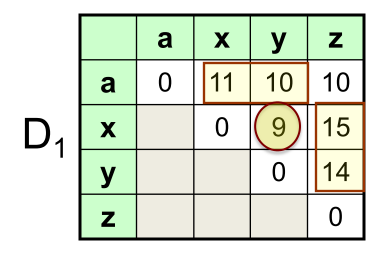


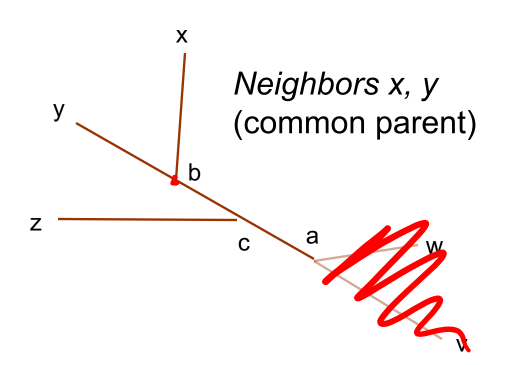
Small Additive Distance Problem $\frac{d(x,v)+d(y,w)}{d(x,v)}$

								_
		V	V	/	X	у	Z	
	V	C	(1	0)	17	16	16	
D	VV)	15	14	14	
	X				0	9	15	
	у					0	14	
	Z						0	

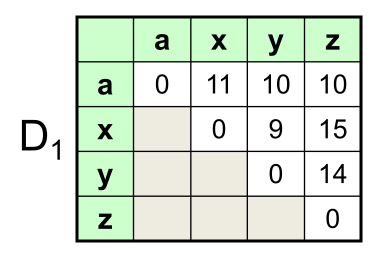
		a	X	У	Z
)	а	0	11	10	10
D_1	X		0	9	15
	у			0	14
	Z				0

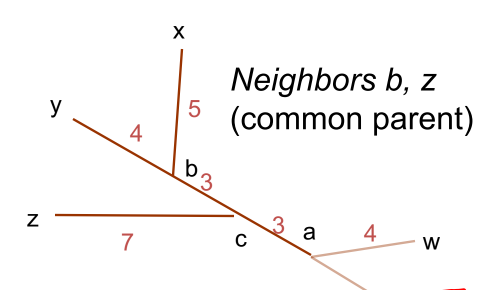




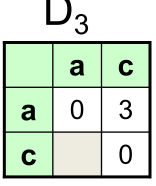


		а	b	Z
ח	а	0	6	10
D_2	b		0	10
	Z			0



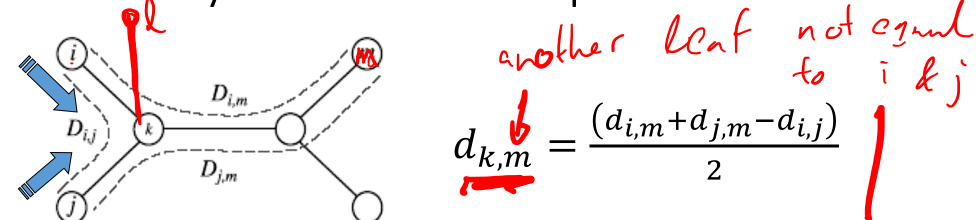


		а	b	Z
D_2	а	0	6	10
	b		0	10
	Z			0



d(a, c) = 3 d(b, c) = d(a, b) - d(a, c) = 3 d(c, z) = d(a, z) - d(a, c) = 7 d(b, x) = d(a, x) - d(a, b) = 5 d(b, y) = d(a, y) - d(a, b) = 4 d(a, w) = d(z, w) - d(a, z) = 4 d(a, v) = d(z, v) - d(a, z) = 6Correct!!!

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



4. Repeat steps 1-3 until tree has only two vertices

Small Additive Distance Phylogeny Problem:

Given $n \times n$ distance matrix $D = [d_{i,j}]$ and unweighted tree T with n leaves, determine edge weights such that $d_T(i,j) = d_{i,j}$

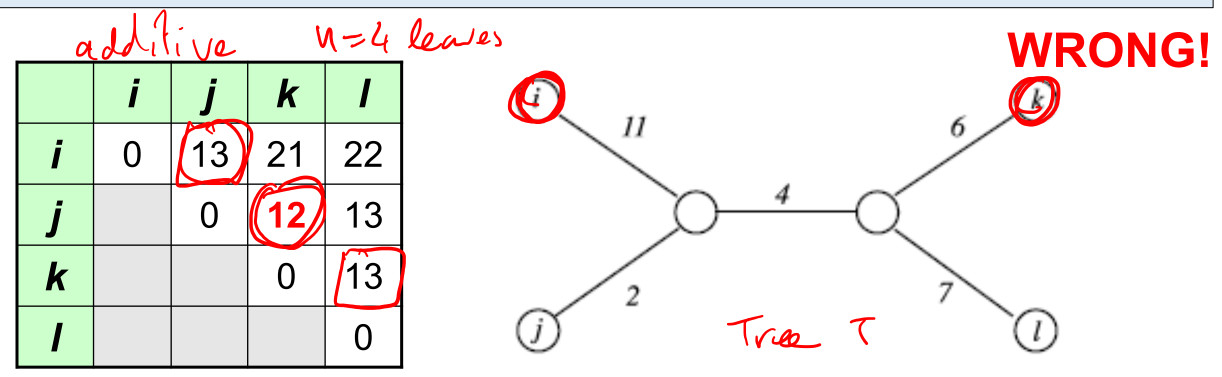
Large Additive Distance Phylogeny Problem:

Given $n \times n$ distance matrix $D = [d_{i,j}]$, find tree T with n leaves **and** edge weights such that $d_T(i,j) = d_{i,j}$

Both problems can be solved in polynomial time

Large Additive Distance Phylogeny Problem

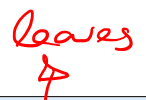
Idea: find neighboring leaves by simply selecting pair of closest leaves



i and *j* are neighbors, but $(d_{ij} = 13) > (d_{jk} = 12)$.

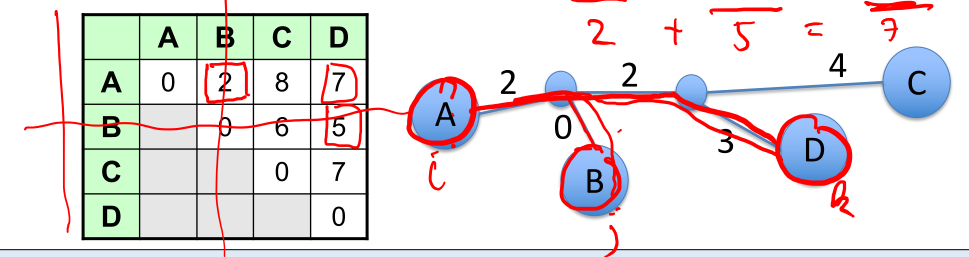
Finding a pair of neighboring leaves is a nontrivial problem!

Degenerate Triples



A degenerate triple is a set of three distinct elements

$$i,j,k \in [n]$$
 such that $d_{i,j}+d_{j,k}=d_{i,k}$



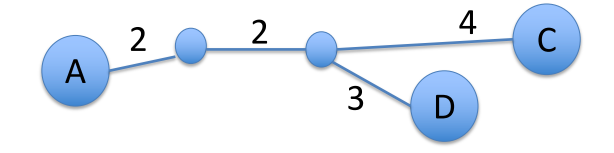
Element j in a degenerate triple (i, j, k) lies* on the evolutionary path from \underline{i} to \underline{k}

*or is attached to this path by an edge of length 0

Degenerate Triples can be Removed

A degenerate triple is a set of three distinct elements $i, j, k \in [n]$ such that $d_{i,j} + d_{j,k} = d_{i,k}$

	A	O	D
Α	0	8	7
С		0	7
D			0

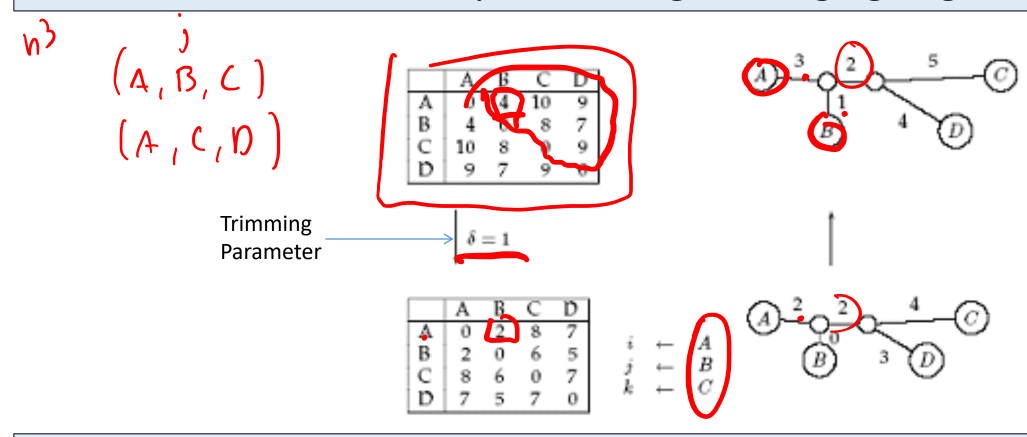


Element j in a degenerate triple (i, j, k) lies* on the evolutionary path from i to k

*or is attached to this path by an edge of length 0

Looking for Degenerate Triples

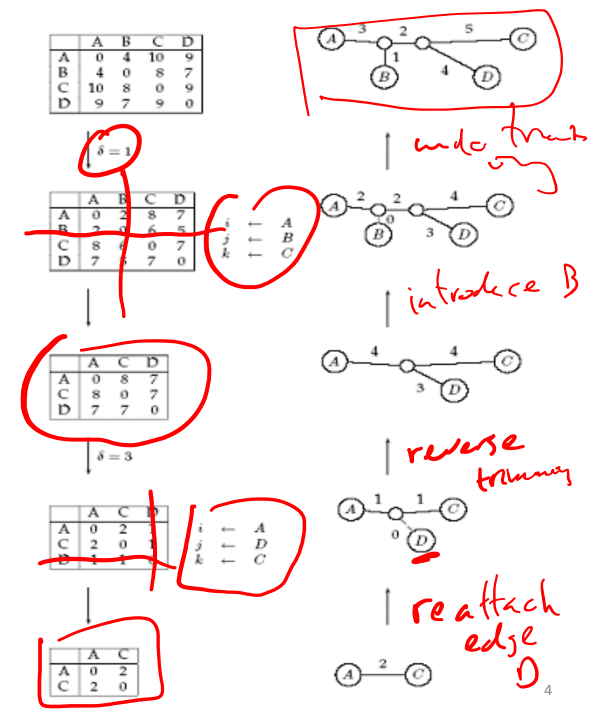
If distance matrix D does not have a degenerate triple, one can create one by shortening all hanging edges



Decrease entries in matrix D by 2δ

Additive Phylogeny

- If there is no degenerative triple:
 - Reduce all hanging edges by the same amount δ , so that all pairwise distances in the matrix are reduced by 2δ .
- This process will eventually collapse one of the leaves (when δ) equals the length of the 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 $d_{i,j}$ for each collapsed leaf.



Additive Phylogeny

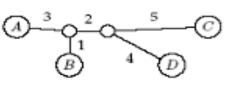
AdditivePhylogeny(D) if D is a 2 x 2 matrix T = tree of a single edge of length $D_{1,2}$ return T **if** *D* is non-degenerate Compute trimming parameter δ Trim(D, δ) Find a triple i, j, k in D such that $D_{ij} + D_{jk} = D_{ik}$ $x = D_{ii}$ Remove j^{th} row and j^{th} column from DT = AdditivePhylogeny(D). Add a new vertex v to T at distance x from i to k Add j back to T by creating an edge (v,j) of length 0 **for** every leaf *I* in *T* if distance from I to v in the tree $\neq D_{l,i}$ output "matrix is not additive" return Extend all "hanging" edges by length δ return T

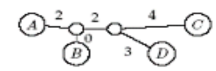
	Α	В	С	D
Α	0	4	10	9
В	4	0	8	7
C	10	8	0	9
D	9	7	9	0

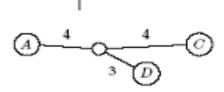


	Α	В	С	D
A	0	2	8	7
B	2	0	6	5
C	8	6	0	7
D	7	5	7	0









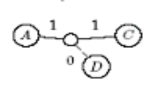
	А	_	ν	ı
Α	-0	8	7	
C	8	0	7	l
D	7	7	0	

$$\delta = 3$$

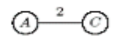
	Α	C	D			
Α	0	2	1	i	-	Α
C	2	0	1	j	\leftarrow	L
A C D	1	1	0	k:	\leftarrow	ϵ



	A	С
A	0	2
C	2	0







Outline

- Recap: RNA Secondary Structure Prediction
- Phylogenetics introduction
- Additive distance phylogeny
- Four point condition
- Neighbor joining

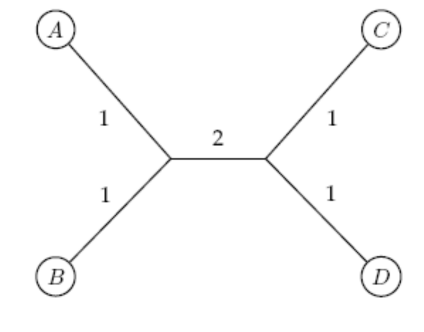
Reading:

Chapter 10.2 and 10.5-10.8 in Jones and Pevzner

Additive Distance Matrices

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

	Α	В	C	D
Α	0	2	4	4
В	2	0	4	4
C	4	4	0	2
D	0 2 4 4	4	2	0



NON-ADDITIVE otherwise

_	_				
Ī		A	В	С	D
 	Α	0	2	2	2
	В	2	0	3	2
	C D	2	3	0	2
1	D	2	2	2	Ø

?

This is a constructive definition

Question: Can we characterize set of additive matrices?

Four Point Condition (Zaretskii 1965, Buneman 1971)

Four point condition of matrix $D = [d_{i,i}]$:

Every four leaves (quartet) can be labeled as (i, j, k, l) such that

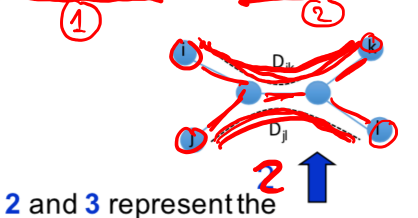
$$d_{i,j} + d_{k,l} \le d_{i,k} + d_{j,l} = d_{i,l} + d_{j,k}$$

Three sums:

1.
$$d_{i,j} + d_{k,l}$$

2.
$$d_{i,k} + d_{j,l}$$

3. $d_{i,l} + d_{i,k}$

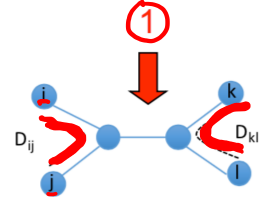


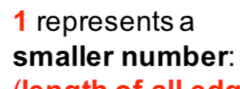
same number:

(length of all edges)

+ 2 * (length middle







(length of all edges) - (length middle edge)







Four Point Condition

Four point condition of matrix $D = [d_{i,j}]$:

Every four leaves (quartet) can be labeled as (i, j, k, l) such that

$$d_{i,j} + d_{k,l} \le d_{i,k} + d_{j,l} = d_{i,l} + d_{j,k}$$

$$d_{i,j} + d_{k,l} \leq d_{i,k} + d_{j,l}$$

$$d_{i,j} + d_{k,l} \leq d_{i,k} + d_{i,l}$$

$$d_{i,l} + d_{k,l} \leq d_{i,k} + d_{i,l}$$

$$d_{i,l} + d_{k,l} \leq d_{i,k} + d_{i,l}$$

If two leaves are the same, four point condition is triangle inequality (e.g. set (i = j))

Four point condition generalizes triangle inequality and defines a subset of distances, namely additive distances

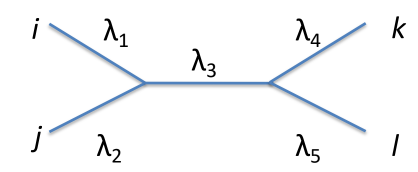
Every four leaves (quartet) can be labeled as (i, j, k, l) such that $d_{i,j} + d_{k,l} \le d_{i,k} + d_{j,l} = d_{i,l} + d_{j,k}$

Theorem: An $n \times n$ matrix D is additive if and only if the for point condition holds for every quartet $(i, j, k, l) \in [n]^4$

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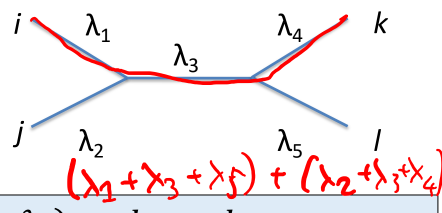
Proof: (=>) Since D is additive, there is a tree T such that $d_{i,j} = d_T(i,j)$ for all $(i,j) \in n^2$. Let (i,j,k,l) be a quartet. Assume w.l.o.g. that i,j and k,l are neighbors. Define λ_m as illustrated.



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$$d_{i,k} + d_{j,l} = (\lambda_1 + \lambda_3 + \lambda_4) + (\lambda_2 + \lambda_3 + \lambda_5) = d_{i,l} + d_{j,k}$$

$$\geq (\lambda_1 + \lambda_2) + (\lambda_4 + \lambda_5) = d_{i,j} + d_{k,l}$$

Every four leaves (quartet) can be labeled as (i, j, k, l) such that $d_{i,j} + d_{k,l} \le d_{i,k} + d_{j,l} = d_{i,l} + d_{j,k}$

Theorem: An $n \times n$ matrix D is additive if and only if the for point condition holds for every quartet $(i, j, k, l) \in [n]^4$

Proof: (<=) Assume four point condition holds. Need an algorithm to construct T. AdditivePhylogeny(T) is one such algorithm*. Neighbor joining is another algorithm.

^{*}we have not proved correctness nor shown how to correct δ

Additive Distance Matrix

Four point condition of matrix $D = [d_{i,j}]$:

Every four leaves (quartet) can be labeled as (i, j, k, l) such that $d_{i,j} + d_{k,l} \le d_{i,k} + d_{j,l} = d_{i,l} + d_{j,k}$

Theorem: Let D be an $n \times n$ matrix. The following statements are equivalent.

- 1. Matrix *D* is additive.
- 2. There exists a unique tree T (modulo isomorphism) s.t. $d_{i,j} = d_T(i,j)$ for all $(i,j) \in n^2$.
- 3. Four point condition holds for every quartet $(i, j, k, l) \in [n]^4$.

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Distance Based Phylogeny Problem

Large Additive Distance Phylogeny Problem:

Given $n \times n$ matrix $D = [d_{i,j}]$, find tree T with n leaves **and** edge weights such that $\max_{(i,j)\in[n]^2} |d_T(i,j) - d_{i,j}|$ is minimum.





Neighbor Joining Algorithm (Saitou and Nei 1987)

- Constructs binary unrooted trees.
- Recall: leaves a and b are neighbors γ if they have a common parent
- Recall: closest leaves are not necessarily neighbors
- NJ: Find pair of leaves that are "close" to each other but "far" from other leaves

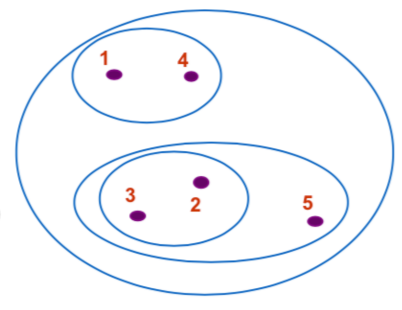
- Constructs binary unrooted trees.
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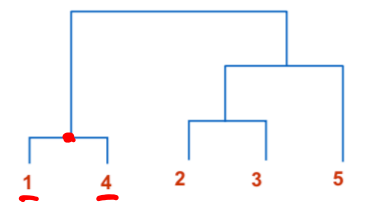
Two advantages: (1) reproduces correct tree for additive matrix, and (2) otherwise gives good approximation of correct tree

Distance Trees as Hierarchical Clustering

Leaves = Data points.

Data points clustered/grouped into hierarchy according to some distance criterion.



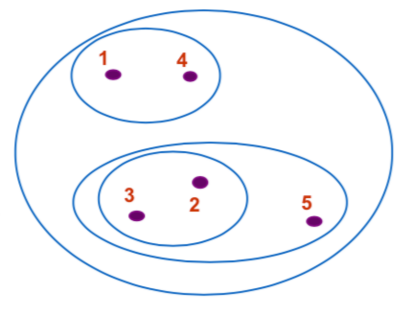


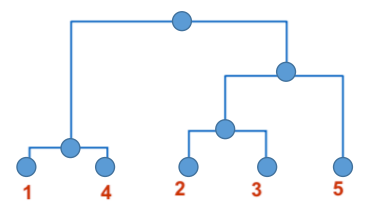
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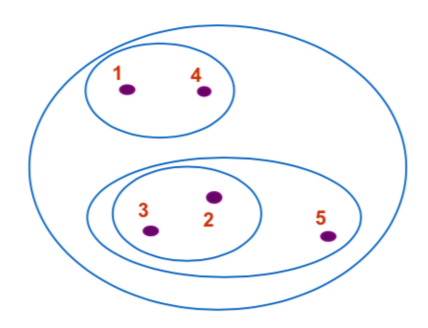


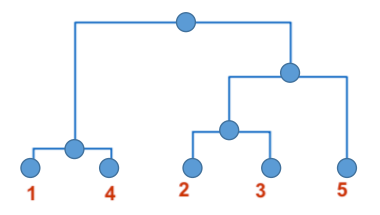


Distance Trees as Hierarchical Clustering

- 1. Hierarchical Clustering (**D**, n)
- Form n clusters each with one element
- 3. Construct a graph **T** by assigning one vertex to each cluster
- **4. while** there is more than one cluster
- Find the two closest clusters C_1 and C_2
- 6. Merge C_1 and C_2 into new cluster C with $|C_1| + |C_2|$ elements
- Compute distance from C to all other clusters
- 8. Add a new vertex \boldsymbol{C} to \boldsymbol{T} and connect to vertices C_1 and C_2
- 9. Remove rows and columns of **D** corresponding to C_1 and C_2
- 10. Add a row and column to **D** corresponding to the new cluster **C**
- **11.** return *T*

Selection criterion: distance between clusters affects clustering!





Neighbor Joining: Selection Criterion Input: Matrix D



Let $C = \{1, ..., n\}$ be current clusters/leaves.

Define: $u_i = \sum_k D(i, k)$.

Intuitively, u_i measures separation of i from other leaves.

Goal: Minimize D(i, j) and maximize $u_i + u_{j.}$

Solution: Find pair (i, j) that minimizes:

$$S_D(i, j) = (n - 2) D(i, j) - u_i - u_j$$
 $Claim$: Given additive matrix D.

 $S_D(x, y) = \min S_D(i, j)$ if and only if $x = 0$

 $S_D(x, y) = \min S_D(i, j)$ if and only if x and y are neighbors in tree T with $d_T = D$.

Neighboring Joining: Algorithm

Initialization:

Form n clusters C_1 , C_2 , ..., C_n , one for each leaf node.

Define tree T to be the set of leaf nodes, one per sequence.

Iteration: (D is $m \times m$)

Pick i, j such that $S_D(i, j) = (m - 2) D(i, j) - u_i - u_i$ is minimal.

Merge i and j into new node [ij] in T.

Assign length $\frac{1}{2} (D(i, j) + \frac{1}{(m-2)} (u_i - u_j))$ to edge (i, [ij])Assign length $\frac{1}{2} (D(i, j) + \frac{1}{(m-2)} (u_j - u_i))$ to edge (j, [ij])

Remove rows and columns from D corresponding to i and j.

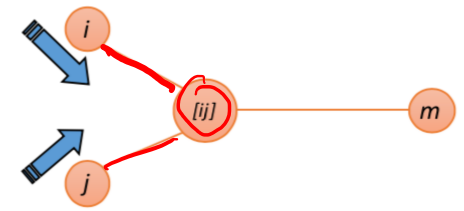
Add row and column to D for new vertex [ij]

Set D([ij], m) = $\frac{1}{2}$ [D(i, m) + D(j, m) - D(i,j)]

Termination:

When only one cluster

Question: Does this create rooted or unrooted trees?



Advantages of Neighbor Joining

distruce

Theorem: Let D be an $n \times n$ matrix. If matrix D is additive then neighbor joining produces the unique phylogenetic tree T (modulo isomorphism) such that $d_{i,j} = d_T(i,j)$ for all $(i,j) \in n^2$.

Theorem: Let D be an $n \times n$ matrix. If there exists an additive matrix D' such that $|D - D'|_{\infty} \leq 0.5$ then neighbor joining applied to D reconstructs the unique tree T (modulo isomorphism) such that $d'_{i,i} = d_T(i,j)$ for all $(i,j) \in n^2$.

Neighbor Joining in Practice F3 3,000 3,000 2,000 2,000 1,000 3,000 3,000 2,000 1,000 1,000 4,000 3,000 3,000 2,000 2,000 1,000 1,000 Euclidean distance Tumour subpopulations Diploids Pseudodiploids Hypodiploids Aneuploid A Cell number

Neighbor Joining tree relating copy number profiles from single cells in a tumor.

[Navin et al, Nature 2011]

Summary

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