

Lecture 20: Clustering and Evolution

Study Chapter 10.4 – 10.8

Fall 2011

Clique Graphs

- A **clique** is a graph where every vertex is connected via an edge to every other vertex
- A **clique graph** is a graph where each connected component is a clique
- The concept of clustering is closely related to clique graphs. Every partition of *n* elements into *k* clusters can be represented as a clique graph on *n* vertices with *k* cliques.

Transforming an Arbitrary Graph into a Clique Graph

- Clusters are maximal cliques (cliques not contained in any other complete subgraph)
 - 1,6,7 is a non-maximal clique.
- An arbitrary graph can be transformed into a clique graph by adding or removing edges



Corrupted Cliques Problem

Determine the smallest number of edges that need be added or removed to transform a graph to a clique graph

Input: A graph *G*

Output: The smallest number of edge additions and/or removals that transforms *G* into a clique graph



Distance Graphs

- One can turn a distance matrix into a distance graph
 - Genes are represented as vertices in the graph
 - Choose a distance threshold θ
 - If the distance between two vertices is below θ , draw an edge between them
 - The resulting graph may contain cliques
 - These cliques represent clusters of closely located data points!



Transforming Distance Graph into Clique Graph

The distance graph (threshold θ =7) is transformed into a clique graph after removing the two highlighted edges



⁽a) Distance matrix, d (distances shorter than 7 are shown in bold).

After transforming the distance graph into the clique graph, the dataset is partitioned into three clusters



Figure 10.6 The distance graph (b) for $\theta = 7$ is not quite a clique graph. However, it can be transformed into a clique graph (c) by removing edges (g_1, g_{10}) and (g_1, g_9) .

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Heuristics for Corrupted Clique Problem

- Corrupted Cliques problem is NP-Hard, some heuristics exist to approximately solve it:
- **CAST** (Cluster Affinity Search Technique): a practical and fast algorithm:
 - CAST is based on the notion of genes *close* to cluster *C* or *distant* from cluster *C*
 - Distance between gene *i* and cluster *C*:

d(i,C) = average distance between gene *i* and *all* genes in *C*

Gene *i* is *close* to cluster *C* if $d(i,C) < \theta$ and *distant* otherwise



CAST Algorithm

- 1. <u>CAST(*S*, *G*, *θ*)</u>
- $2. \quad P \leftarrow \emptyset$
- 3. while $S \neq \emptyset$
- 4. $v \leftarrow$ vertex of maximal degree in the distance graph G
- 5. $C \leftarrow \{v\}$
- 6. while a close gene *i* not in *C* or distant gene *i* in *C* exists
- 7. Find the nearest close gene *i* not in *C* and add it to *C*
- 8. Remove the farthest distant gene *i* in *C*
- 9. Add cluster *C* to partition *P*
- 10. $S \leftarrow S \setminus C$
- 11. Remove vertices of cluster *C* from the distance graph *G*
- 12. return *P*

S – set of elements, G – distance graph, θ – distance threshold



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 was hotly debated
- Now it is a dominant approach to study evolution.



Out of Africa Hypothesis

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- 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 most ancient ancestor lived in Africa roughly 200,000 years ago
- Largely based on mitochondrial DNA



Human Evolutionary Tree (cont'd)

http://www.mun.ca/biology/scarr/Out_of_Africa2.htm



The Origin of Humans: "Out of Africa" vs Multiregional Hypothesis

Out of Africa:

- Humans evolved in Africa ~150,000 years ago
- Humans migrated out of Africa, replacing other shumanoids around the globe
- There is no direct descendence from Neanderthals

Multiregional:

- Humans evolved in the last two million years as a single species.
 Independent appearance of modern traits in different areas
- Humans migrated out of Africa mixing with other humanoids on the way
- There is a genetic continuity from Neanderthals to humans



mtDNA analysis supports "Out of Africa" Hypothesis

- African origin of humans inferred from:
 - African population was the most diverse (sub-populations had more time to diverge)
 - The evolutionary tree separated one group of Africans from a group containing all five populations.
 - Tree was rooted on branch between groups of greatest difference.



Evolutionary Tree of Humans (mtDNA)

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



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

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Evolutionary Tree of Humans: (microsatellites)

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



Human Migration Out of Africa



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







(b) Rooted tree

(c) The 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

$d_{ij}(T)$ – tree distance between i and j



Distance in Trees



$d_{1,4} = 12 + 13 + 14 + 17 + 13 = 69$



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Distance Matrix

- Given *n* species, we can compute the *n* x *n distance matrix D*_{*ij*}
- D_{ij} 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_{ij} – edit distance between i and j



Edit Distance vs. Tree Distance

- Given *n* species, we can compute the *n* x *n distance matrix D*_{*ij*}
- D_{ij} 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_{ij} – 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_{ij}
- 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*_{*ij*}



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*



<u>Observe:</u>

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

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

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



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Reconstructing a 3 Leaved Tree (cont'd)



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

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

$$2d_{ic} + d_{jc} + d_{kc} = D_{ij} + D_{ik}$$

$$2d_{ic} + D_{jk} = D_{ij} + D_{ik}$$

$$d_{ic} = (D_{ij} + D_{ik} - D_{jk})/2$$
nilarly,
$$d_{jc} = (D_{ij} + D_{jk} - D_{ik})/2$$

$$d_{kc} = (D_{ki} + D_{kj} - D_{ij})/2$$

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Trees with > 3 Leaves

- An unrooted tree with *n* leaves has 2*n*-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 for n > 3 given arbitrary/noisy distances



Additive Distance Matrices

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





NON-ADDITIVE otherwise

	Α	В	С	D
Α	0	2	2	2
В	2	0	3	2
С	2	3	0	2
D	2	2	2	0



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Distance Based Phylogeny Problem

- <u>Goal</u>: Reconstruct an evolutionary tree from a distance matrix
- Input: $n \ge n$ distance matrix D_{ij}
- <u>Output</u>: 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



Using Neighboring Leaves to Construct the Tree

- Find *neighboring leaves i* and *j* with common 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:



Finding Neighboring Leaves

- Or solution assumes that we can easily find neighboring leaves given only distance values
- How might one approach this problem?
- It is not as easy as selecting a pair of closest leaves.



Finding Neighboring Leaves

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- 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! (we'll return to it later)

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 nonadditive matrices, it does not have the flawed molecular clock assumption



Degenerate Triples

- A degenerate triple is a set of three distinct elements $1 \le i, j, k \le n$ where $D_{ij} + D_{jk} = D_{ik}$
- Called *degenerate* because it implies i, j, and k are collinear.
- 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 degenerative triple in *D* by shortening all hanging edges (in the tree).



Shortening Hanging Edges

• 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*_{*ij*} for each collapsed leaf.



Reconstructing Trees for Additive Distance Matrices





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AdditivePhylogeny Algorithm

1. AdditivePhylogeny(D)

3. T = tree of a single edge of length $D_{1,2}$

4. return *T*

- 5. if *D* is non-degenerate
- 6. δ = trimming parameter of matrix *D*

7. **for** all
$$1 \le i \ne j \le n$$

8.
$$D_{ij} = D_{ij} - 2\delta$$

10.
$$\delta = 0$$



AdditivePhylogeny (cont'd)

1. Find a triple *i*, *j*, *k* in *D* such that $D_{ij} + D_{jk} = D_{ik}$

$$2. \qquad x = D_{ij}$$

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



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 $1 \le i, j, k, l \le n$ be four distinct leaves in a tree



The Four Point Condition (cont'd)

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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 $n \ge n$ matrix D is additive if and only if the four point condition holds for *every* quartet $1 \le i, j, k, l \le n$



Next Time

• How to create trees if the matrices are not additive

