

Community detection and clustering

Social and Technological Networks

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Communities

- Groups of friends
- Colleagues/collaborators
- Web pages on similar topics
- Biological reaction groups
- Similar customers/users ...

Other applications

- A coarser representation of networks
- One or more meta-node for each community
- Identify bridges/weak-links
- Structural holes

Community detection

- Given a network
- What are the “communities”
 - Closely connected groups of nodes
 - Relatively few edges to outside the community
- Similar to clustering in data sets
 - Group together points that are more close or similar to each other than other points

Definitions of communities

- Varies. Depending on application
- General idea: **Dense subgraphs**: More links within community, few links outside
- Some types and considerations:
 - Partitions: Each node in exactly one community
 - Overlapping: Each node can be in multiple communities
 - We will usually consider partitions

Comment: Finding dense subgraphs is hard in general

- Finding largest clique
 - NP-hard
 - Computationally intractable
- Decision version: Does a clique of size k exist?
 - Also NP-complete
 - Computationally intractable
 - Polynomial time (efficient) algorithms unlikely to exist
- We will look for approximations

Dense subgraphs: Few preliminary definitions

- For S, T subgraphs of V
- $e(S, T)$: Set of edges from S to T
 - $e(S) = e(S, S)$: Edges within S
- $d_S(v)$: number of edges from v to S
- Edge density of S : $|e(S)|/|S|$
 - Largest for complete graphs or cliques

Dense subgraph Problem

- Find the subgraph with largest edge density
- There also exists a decision version:
 - Is there a subgraph with edge density $> \alpha$
- Can be solved using Max Flow algorithms
 - $O(n^2m)$: inefficient in large datasets
 - Finds the one densest subgraph
- Variant: Find densest S containing given subset X
- Other versions: Find subgraphs size k or less
- NP-hard

Efficient approximation for finding dense S containing X

Let $G_n \leftarrow G$.

for $k = n$ **downto** $|X| + 1$ **do**

 Let $v \notin X$ be the lowest degree node in $G_k \setminus X$.

 Let $G_{k-1} \leftarrow G_k \setminus \{v\}$.

Output the densest subgraph among $G_n, \dots, G_{|X|}$.

- Gives a $1/2$ approximation
- Edge density of output S set is at least half of optimal set S^*
- (Proof in Kempe 2018: <http://www-bcf.usc.edu/~dkempe/teaching/structure-dynamics.pdf>).

Betweenness & graph partitioning

- We want to split network into tightly knit groups (communities etc)
- Idea: Identify the edges connecting different communities and remove them
- These edges are “central” to the network
 - They lie on shortest paths
- Betweenness of edge (e) (can be considered for vertex (v)):
 - We send 1 unit of traffic between every pair of nodes in the network
 - measure what fraction passes through e , assuming the flow is split equally among all shortest paths.

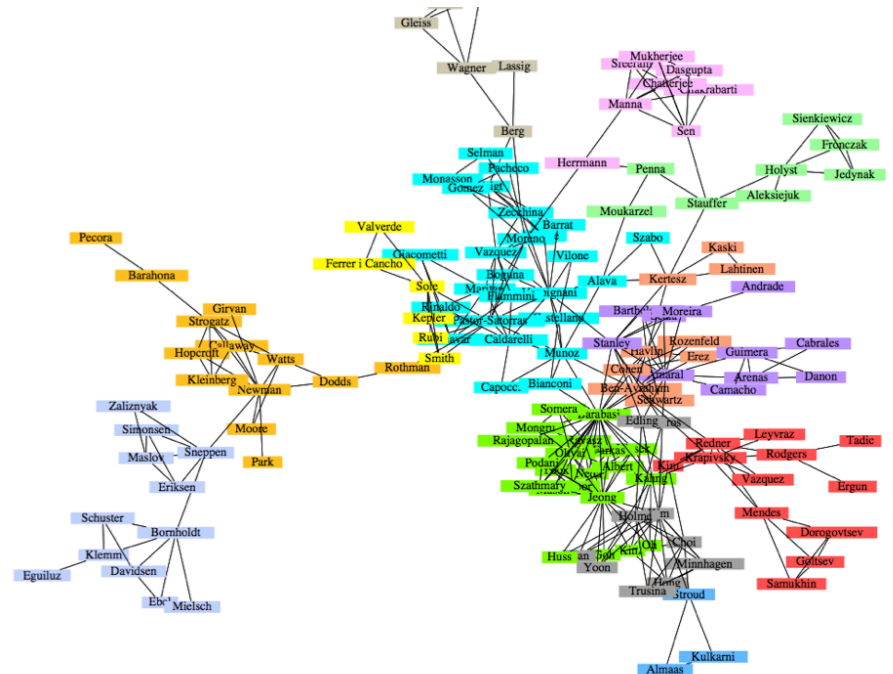
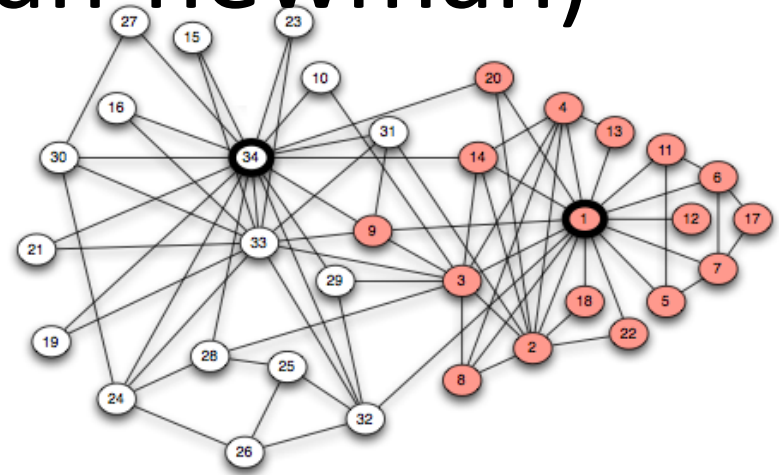
Computing betweenness

- Computing all shortest paths separately is inefficient
- A more efficient way:
- From each node:
 - Step 1: Compute BFS tree
 - Step 2: Find number of shortest paths to each node
 - Step 3: Find the flow through each edge
 - See Kleinberg-Easley for detailed algorithm

Partitioning (Girvan-newman)

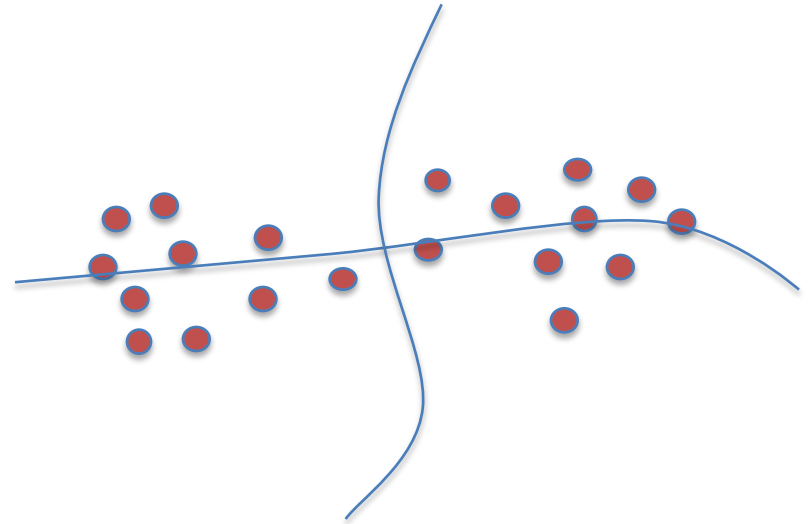
Repeat:

- Find edge e of highest betweenness
- Remove e
- Produces a hierarchic partitioning structure as the graph decomposes into smaller components
- Network version of hierarchic clustering



Modularity

- How do we evaluate the quality of detected communities?
 - How do you say the vertical cut is better than the horizontal cut?
- Idea: Maximize a quantity called *modularity*



Configuration model of Random graphs

- Suppose we want a graph that is random
- But has given degree for each vertex:

$$d_1, d_2, d_3, \dots, d_n$$

- At each vertex i we d_i *open-edges*
- Pair up the edges randomly
- Given a graph G , we can construct a random graph G' with same degrees but random edges

Modularity of subset S of V

- Given the graph G
- Consider a random G' graph with same node degrees
 - Number of edges in S in G : $|e(S)|_G$
 - Expected number of edges in S in G' : $E[|e(S)|_{G'}]$
 - Modularity of S : $|e(S)|_G - E[|e(S)|_{G'}]$
 - More coherent communities have more edges inside than would be expected in a random graph with same degrees
 - Note: modularity can be negative

Modularity of a clustering

- Take a partition (clustering) of V : $\mathcal{P} = \{S_1, \dots, S_k\}$
- Write $d(S_i)$ for sum of degrees of all nodes in S_i
- It can be shown that $E[|e(S)|_{G'}] \approx d(S_i)^2$
- Definition: Sum over the partition:

$$q(\mathcal{P}) = \frac{1}{m} \sum_i |e(S_i)|_G - \frac{1}{4m} d(S_i)^2$$

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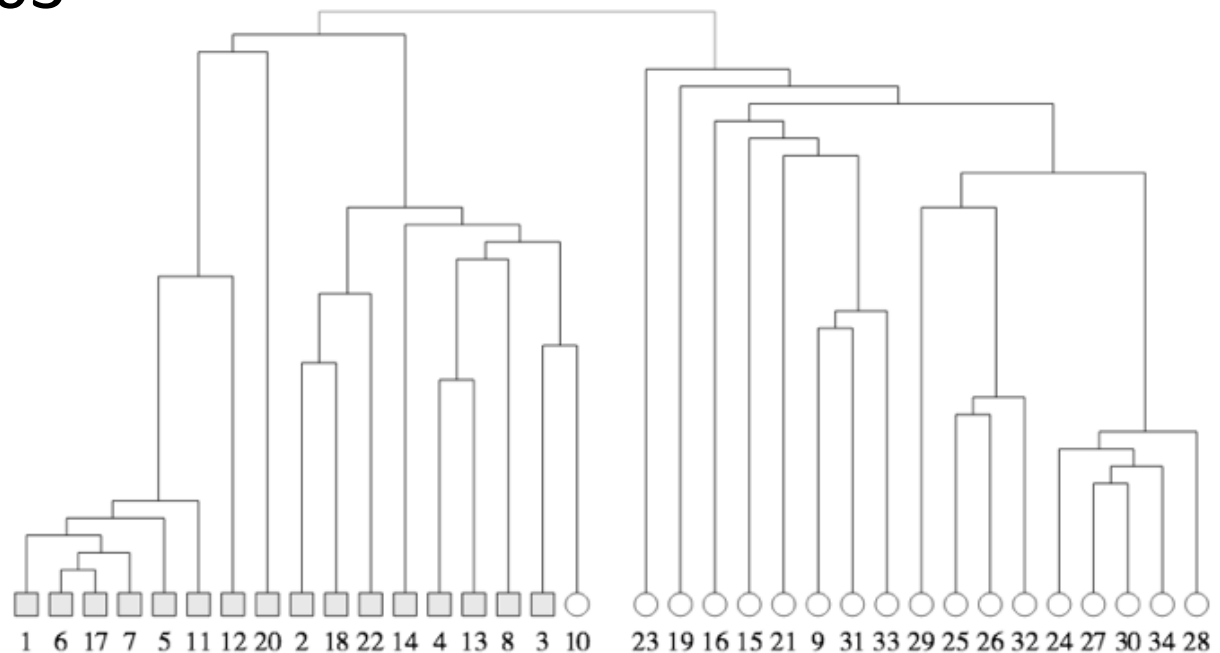
- Can be used as a stopping criterion (or finding right level of partitioning) in hierarchic methods
 - Eg. Girvan-newman

Modularity based clustering

- Modularity is meant for use more as a measure of quality, not so much as a clustering method
- Finding clustering with highest modularity is NP-hard
- Heuristic: Louvain method:
 - Place each node in its own community
 - For each community, consider merging with neighbor.
 - Make the greedy choice – make the merge that maximizes modularity
 - Or do not merge if none increases modularity
 - Repeat
- Note: Modularity is a relative measure for comparing community structure.
- Not entirely clear in which cases it may or may not give good results
- A threshold of 0.3 or more is sometimes considered to give good clustering

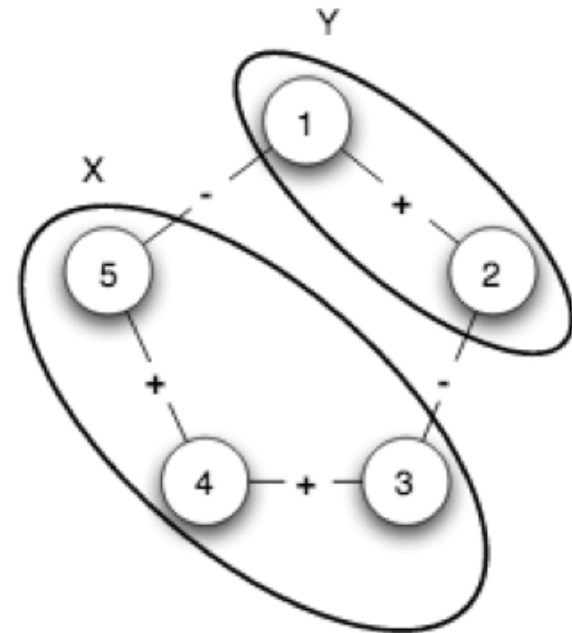
Karate club hierarchic clustering

- Shape of nodes gives actual split in the club due to internal conflicts
 - Newman 2003



Correlation clustering

- Some edges are known to be similar/friends/trusted
- marked “+”
- Some edges are known to be dissimilar/enemies/distrusted
- marked “-”
- Maximize the number of + edges inside clusters and
- Minimize the number of - edges between clusters



Applications

- Community detection based on similar people/users
- Document clustering based on known similarity or dissimilarity between documents
- Use of sentiments and/or other divisive attributes

Features

- Clustering without need to know number of clusters
 - k-means, medians, clusters etc need to know number of clusters or other parameters like threshold
 - Number of clusters depends on network structure
- Actually, does not need any parameter
- NP hard
- Note that graph may be complete or not complete
 - In some applications with unlabeled edges, it may be reasonable to change edges to “+” edges and non-edges to “-” edges

Approximation

- Naive 1/2 approximation:
 - If there are more + edges
 - Put them all in 1 cluster
 - If there are more - edges
 - Put nodes in n different clusters
- (not very useful)!

Better approximations

- 2 ways of looking at it:
 - Maximize agreement or Minimize disagreement
 - Similar idea, but we know different approximation algorithms
- Nikhil Bansal et al. develop PTAS (polynomial time approximation scheme) for maximizing agreement:
 - $(1-\epsilon)$ approximation, running time $O(n^2 e^{O(1/\epsilon)})$
- Min-disagree:
 - 4-approximation

Local detection of communities

- Suppose there is a huge graph, like www, or facebook network
- We often want to find the community that contains a particular node or group
 - E.g. to make recommendations: “your friends have watched this movie...”
 - To infer preferences and attributes
- Running a full scale community detection is computationally impractical
- We do not know the number of communities

Conductance: measure of edges inside community vs outside

- Given subsets S, T in V
- $e(S, T)$: set of edges between S and T
- Volume of edges: $vol(S) = \sum_{v \in S} d(v)$
- Conductance of S is defined as:

$$\Phi(S) := \frac{e(S, \bar{S})}{\min(vol(S), vol(\bar{S}))}$$

- Communities are likely to have low conductance

Personalised pagerank

- Given a seed set X
- Find the community S that contains X
- Pagerank style: Use random walks
- Algorithm
 - Set a limit k to number of steps in random walks
 - Repeat:
 - Select at random a start point from X
 - Take k random steps in the graph
 - Count how frequently each node occurs – pagerank
 - Nodes in the community have high pagerank

Personalised pagerank

- Alternative Algorithm
 - Set a probability to reset random walk
 - Repeat:
 - Select at random a start point from X
 - With probability $1 - \epsilon$ move to a random neighbor
 - With probability ϵ move to a random node in X
 - Count how frequently each node occurs – pagerank
 - Nodes in the community have high pagerank

Personalised pagerank

- Communities have low conductance
- Therefore, short random walks will leave the community only rarely
- Therefore, nodes in the community of X will have high pagerank compared to those outside
- It can be proved that if X is in a low conductance community, nodes outside this community will occur infrequently.
 - We will omit this proof

Community detection by clustering

- First, define a metric between nodes
 - Either compute intrinsic metrics like all pairs shortest paths [Floyd-Warshall algorithm $O(n^3)$]
 - Or embed the nodes in a Euclidean space, and use the metric there
 - We will later study embedding methods
- Apply a clustering algorithm with the metric

Clustering

- A core problem of machine learning:
 - Which items are in the same group?
- Identifies items that are similar relative to rest of data
- Simplifies information by grouping similar items
 - Helps in all types of other problems

Clustering

- Outline approach:
- Given a set of items
 - Define a distance between them
 - E.g. Euclidean distance between points in a plane; Euclidean distance between other attributes; non-euclidean distances; path lengths in a network; tie strengths in a network...
 - Determine a grouping (partitioning) that optimises some function (prefers 'close' items in same group).
- General references for clustering:
 - Charu Aggarwal: The Data Mining Textbook, Springer
 - Free on Springer site (from university network)
 - Blum et al. Foundations of Data Science (free online)

K-means clustering

- Find k-clusters $\mathcal{C} = \{C_1, \dots, C_k\}$
 - With centers $\mathbf{c}_1, \dots, \mathbf{c}_k,$
 - That minimize the sum of squared distances of nodes to their cluster centers (called the k-means cost)

$$\Phi_{kmeans}(\mathcal{C}) = \sum_{j=1}^k \sum_{\mathbf{a}_i \in C_j} d^2(\mathbf{a}_i, \mathbf{c}_j)$$

K-means clustering: Lloyd's algorithm

- There are n items
- Select k 'centers'
 - May be random k locations in space
 - May be location of k of the items selected randomly
 - May be chosen according to some method
- Iterate till convergence:
 - Assign each item to the cluster for its closest center
 - Recompute location of center as the mean location of all elements in the cluster (their centroid)
 - Repeat
- Warning: Lloyd's algorithm is a Heuristic. Does not guarantee that the k-means cost is minimised

K-means

- Visualisations
- <http://stanford.edu/class/ee103/visualizations/kmeans/kmeans.html>
- <http://shabal.in/visuals/kmeans/1.html>

K-means

- Ward's algorithm (also Heuristic)
 - Start with each node as its own cluster
 - At each round, find two clusters such that merging them will reduce the k-means cost the most
 - Merge these two clusters
 - Repeat until there are k-clusters

K means: discussion

- Tries to minimise squared sum of distances of items to cluster centers
 - NP-hard. Computationally intractable
 - Algorithm gives local optimum
- Depends on initialisation (starting set of centers)
 - Can give poor results
 - Submodular optimisation can help
- The right 'k' may be unknown
 - Possible strategy: try different possibilities and take the best
- Can be improved by heuristics like choosing centers carefully
 - E.g. choosing centers to be as far apart as possible: choose one, choose point farthest to it, choose point farthest to both (maximise min distance to existing set etc)...
 - Try multiple times and take best result..

K-medoids

- Similar, but now each center must be one of the given items
 - In each cluster, find the item that is the best ‘center’ and repeat
- Useful when there is no ambient space (extrinsic metric)
 - E.g. A distance between items can be computed between nodes, but they are not in any particular Euclidean space, so the ‘centroid’ in Lloyd’s algorithm is not a meaningful point

Other center based methods

- K-center: Minimise maximum distance to center:

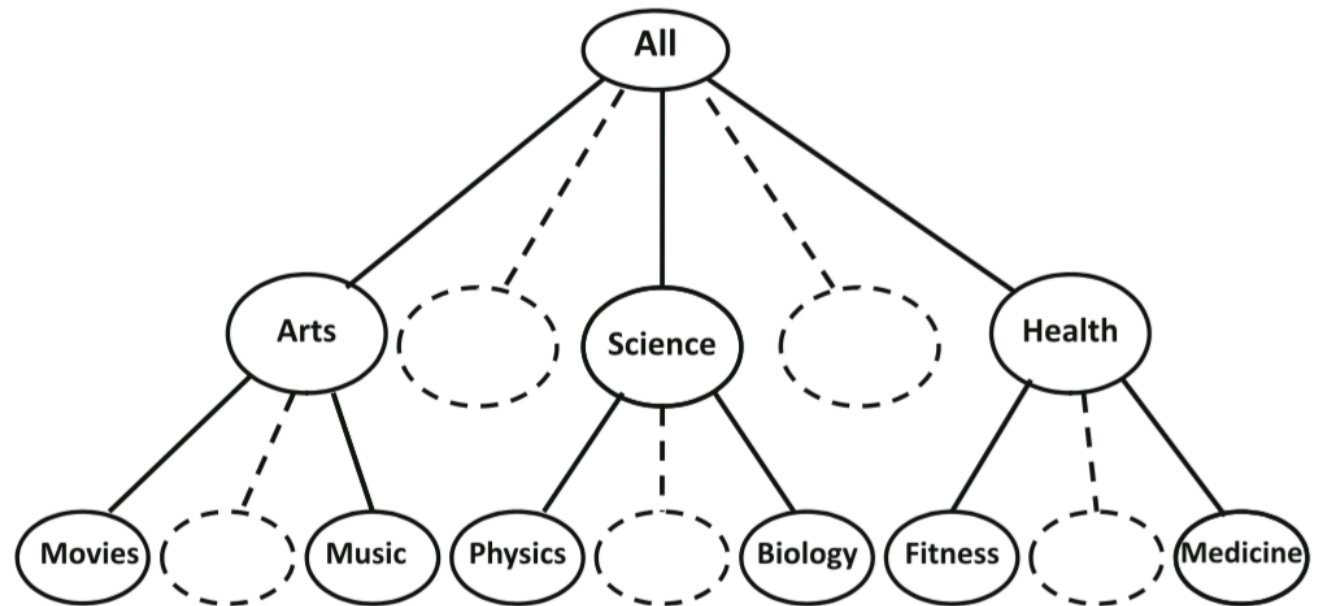
$$\Phi_{kcenter}(\mathcal{C}) = \max_{j=1}^k \max_{\mathbf{a}_i \in C_j} d(\mathbf{a}_i, \mathbf{c}_j)$$

- K-median: Minimise sum of distances:

$$\Phi_{kmedian}(\mathcal{C}) = \sum_{j=1}^k \sum_{\mathbf{a}_i \in C_j} d(\mathbf{a}_i, \mathbf{c}_j)$$

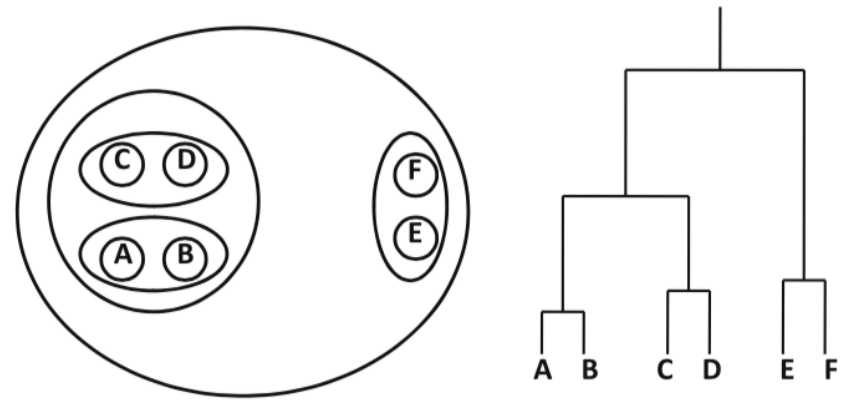
Hierarchical clustering

- Hierarchically group items
- Using some standard clustering method



Hierarchical clustering

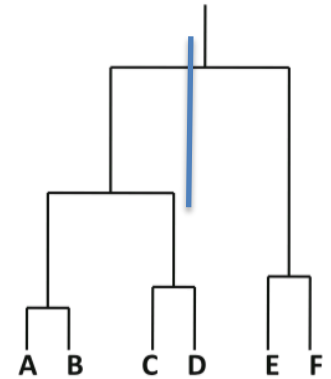
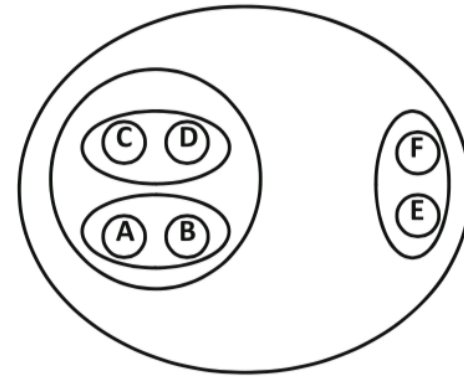
- Top down (divisive):
 - Start with everything in 1 cluster
 - Make the best division, and repeat in each subcluster
- Bottom up (agglomerative):
 - Start with n different clusters
 - Merge two at a time by finding pairs that give the best improvement



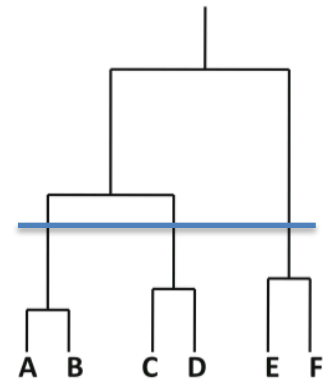
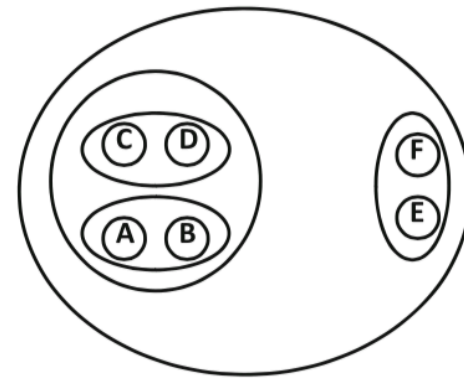
(a) Dendrogram

Hierarchical clustering

- Gives many options for a flat clustering
- Problem: what is a good 'cut' of the dendrogram?



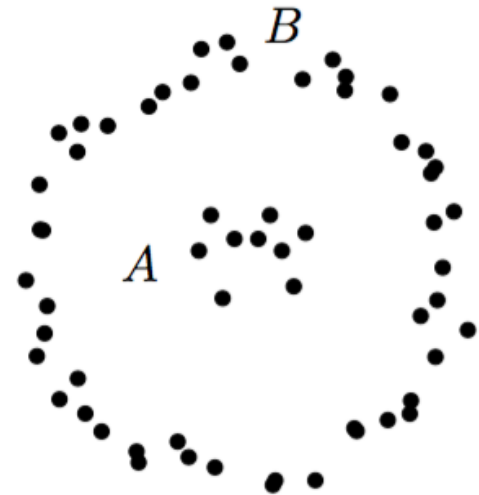
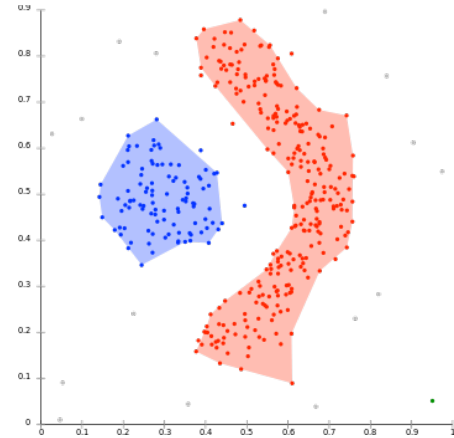
(a) Dendrogram



(a) Dendrogram

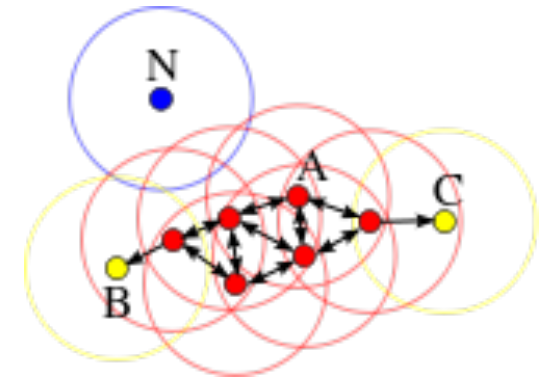
Density based clustering

- Group dense regions together
- Better at non-linear separations
- Works with unknown number of clusters



DBSCAN

- Density at a data point:
 - Number of data points within radius Eps
- A core point:
 - Point with density at least τ
- Border point
 - Density less than τ , but at least one core point within radius Eps
- Noise point
 - Neither core nor border. Far from dense regions



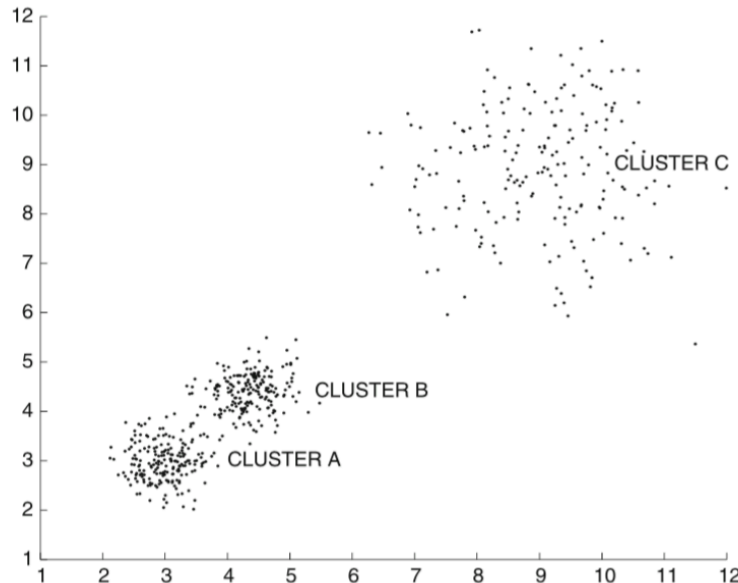
Algorithm

- Construct UDG of core points
- Connected components of the graph give the clusters
- Assign border points to suitable clusters (E.g. to the cluster to which it has most edges)

Algorithm *DBSCAN*(Data: \mathcal{D} , Radius: Eps , Density: τ)
begin
Determine core, border and noise points of \mathcal{D} at level (Eps, τ) ;
Create graph in which core points are connected
if they are within Eps of one another;
Determine connected components in graph;
Assign each border point to connected component
with which it is best connected;
return points in each connected component as a cluster;
end

DBSCAN: Discussions

- Requires knowledge of suitable radius and density parameters (Eps and τ)
- Does not allow for possibility that different clusters may have different densities



DBSCAN

- Useful in cases where it is clear which objects can be considered similar but number of clusters is not known
- Known to perform very well in real problems
- Worst case complexity: $O(n^2)$
- Current research: Making faster in special cases, approximations, distributed algorithms.

Other density based clustering

- Single linkage (same as Kruskal's MST algorithm)
 - Start with n clusters
 - Merge two clusters with the shortest bridging link
 - Repeat until k clusters
- Other, more robust methods exist

Community detection in networks

- A simple strategy:
 - Choose a suitable distance measure based on available data
 - E.g. Path lengths; distance based on inverse tie strengths; size of largest enclosing group or common attribute; distance in a spectral (eigenvector) embedding; etc..
 - Apply a standard clustering algorithm

Clustering is not always suitable in networks

- Small world networks have small diameter
 - And sometime integer distances
 - A distance based method does not have a lot of option to represent similarities/dissimilarities
- High degree nodes are common
 - Connect different communities
 - Hard to separate communities
- Edge densities vary across the network
 - Same threshold does not work well everywhere