## Complex Networks network algorithms (1)

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## contents of this chapter

- algorithms for calculating centrality indices, finding components, calculating shortest paths and maximum flow


# software implementing common network algorithms 

| Name | Availability | Platform | Description |
| :--- | :--- | :--- | :--- |
| Pajek | Free | W | Interactive social network analysis and visualization |
| Net Workbench | Free | WML | Interactive network analysis and visualization |
| Netminer | Commercial | W | Interactive social network analysis and visualization |
| InFlow | Commercial | W | Interactive social network analysis and visualization |
| UCINET | Commercial | W | Interactive social network analysis |
| yEd | Free | WML | Interactive visualization |
| Visone | Free | WL | Interactive visualization |
| Graphviz | Free | L | Visualization |
| NetworkX | Free | WML | Interactive network analysis and Python library |
| JUNG | Free | WML | JAVA library for network analysis and visualization |
| igraph | Free | WML | C/R/Python libraries for network analysis |
| GTL | Free | WML | C++ library for network analysis |
| LEDA/AGD | Commercial | WL | C++ library for network analysis |

Table 9.1: A selection of software implementing common network algorithms. Platforms are Microsoft Windows (W), Apple Macintosh (M), and Linux (L). Most Linux programs also run under Unix and Unix-like systems such as BSD, and many Windows programs can run on Macs and Linux systems using emulation software.

Reasons for studying algorithms (even though ready-made software packages are available)

- much time can be wasted when people misunderstand the answers the software can give them
- you will sooner or later find that you need to do something that cannot be done with standard software and you'll have to write some programs of your own
- by relying on ready-made packages, researchers have become limited in what types of analysis they can perform


## running time and computational

## complexity

- if the calculation you have started will take a thousand years to finish, it is basically useless
- concept of complexity is useful because it helps us to avoid wasting our energies on programs that will not finish running in any reasonable amount of time


## computational complexity

- a measure of the running time of a computer algorithm
- example: to find the largest number in a list of n numbers
- simply go through the list to the end, keeping a record of the largest number we have seen
- worst case: every number is bigger than all the ones before it
- its running time is $n \tau$
- time complexity of this algorithm is order $\mathrm{n}(\mathrm{O}(\mathrm{n})$ )


## algorithms for degrees and degree

 distributions- degree of a vertex
- if network is stored in adjacency list: $\mathrm{O}(1)$
- if network is stored in adjacency matrix: O(n)
- degree distribution $\mathrm{p}_{\mathrm{k}}: \mathrm{O}(\mathrm{n})$

- cumulative degree distribution $\mathrm{P}_{\mathrm{k}}$
- form a histogram of the degrees and calculate : $\mathrm{O}(\mathrm{n})$

$$
P_{k}=\sum_{k^{\prime}=k}^{\infty} p_{k^{\prime}}=-p_{k-1}+\sum_{k=k_{k}-1}^{\infty} p_{k^{\prime}}=P_{k-1}-p_{k-1} \quad P_{0}=\sum_{k^{\prime}=0}^{\infty} p_{k^{\prime}}=1
$$

- sorting degrees in descending order and rank from 1 to $\mathrm{n}: \mathrm{O}(\mathrm{nlogn})$


## cf. assortative mix, homophily

- high-degree vertices tend to connect high-
degree ones correlation coefficient $r=\frac{\sum_{i j}\left(A_{i j}-k_{i} k_{j} / 2 m\right) k_{i} k_{j}}{\sum_{i j}\left(k_{i} \delta_{i j}-k_{i} k_{j} / 2 m\right) k_{i} k_{j}}$
- faster computation of $r$

$$
r=\frac{S_{1} S_{e}-S_{2}^{2}}{S_{1} S_{3}-S_{2}^{2}} \quad \begin{aligned}
& S_{e}=\sum_{i j} A_{i j} k_{i} k_{j}=2 \sum_{i} \sum_{i} k_{i} k_{j} \\
& S_{1}, S_{2}=\sum_{i}^{\operatorname{cedges}(i, j)} k_{i}^{2}, S_{3}=\sum_{i} k_{i}^{3}
\end{aligned}
$$

- social networks: positive $r$
- other networks: negative r


## correlation coefficient

- computing $\mathrm{S}_{\mathrm{e}}$ takes time $\mathrm{O}(\mathrm{m})$
- computing $S_{1}, S_{2}$, and $S_{3}$ take time $O(n)$
- total time is $\mathrm{O}(\mathrm{m}+\mathrm{n}) \quad$| mean degree |
| :---: |
| $=2 \mathrm{~m} / \mathrm{m}$ is constant |
- sparse networks ( $m \propto n$ ): $O(m+n) \equiv O(n)$
- dense networks ( $\mathrm{m} \propto \mathrm{n}^{2}$ ): $\mathrm{O}(\mathrm{m}) \equiv \mathrm{O}\left(\mathrm{n}^{2}\right)$


## cf. transitivity

- $a \bullet b$ and $b \cdot c->a \bullet c$
- $u \& v$ are friends and $v \& w$ are friends
- clustering coefficient: $C=\frac{(\# \text { of closed paths of length two) }}{(\# \text { of paths of length two) }}$
- $\mathrm{C}=1$ :clique
- $\mathrm{C}=0$ :tree, square lattice
- $\mathrm{C}=\frac{\text { (\# of triangles) } \times 6}{\text { (\# of paths of length two) }}=\frac{\text { (\# of triangles) } \times 3}{\text { (\# of connected triples) }}$
- social networks tend to have high values


## cf. local clustering coefficient

- $\mathrm{C}_{\mathrm{i}}=\frac{\text { (\# of pairs of neighbors of } \mathrm{i} \text { that are connected) }}{\text { (\# of paths of neighbors of } \mathrm{i} \text { ) }}$
- vertices with higher degree have lower local clustering coefficient on average
- structural holes
- bad for info spread or traffic
- good for the central vertex
- it can control the flow of information
- similar to betweenness centrality


## clustering coefficients

<naive method>

- local clustering coefficient
$C_{i}=\frac{\text { (\# of pairs of neighbors of } i \text { that are connected) }}{(\# \text { of paths of neighbors of } i)}$
- go through every pair of neighbors of $i$, and count how many are connected
- overall clustering coefficient
$C=\frac{\text { (\# of triangles) } \times 3}{\text { (\# of connected triples) }}$
- we consider for every vertex each pair of neighbors and find whether they are connected


## complexity of naive method

- vertex i with degree $\mathrm{k}_{\mathrm{i}}$ has $\frac{1}{2} k_{i}\left(k_{i}-1\right)$ pairs of neighbors
- total number of checks:

$$
\sum_{i} \frac{1}{2} k_{i}\left(k_{i}-1\right)=\frac{1}{2} n\left(\left\langle k^{2}\right\rangle-\langle k\rangle\right) \quad\langle k\rangle=\frac{1}{n} \sum_{i} k_{i}=\frac{2 m}{n}\left\langle k^{2}\right\rangle=\frac{1}{n} \sum_{i} k_{i}^{2}
$$

- depends on degree distribution
- if a network follows a power law $p_{k} \approx k^{-\alpha}$, the second moment $\left\langle k^{2}\right\rangle$ formally diverges if $\alpha<3$
- it will take an infinite amount of time!


## more careful calculation

- a network is stored in adjacency list
- for a single vertex i
- \# of pairs of neighbors ( j and I): $: \frac{1}{2} k_{i}\left(k_{i}-1\right)$
- for each pair, check whether an edge exist
 between them : time proportional to $\mathrm{k}_{\mathrm{j}}$ or $\mathrm{k}_{\mathrm{l}}$
- if chosen at random : time proportional to $k_{j}+k_{l}$
$-\Gamma_{i}$ : set of neighbors of vertex $i$
- total time to check for edges between all pairs:
$\sum_{j, l \in \Gamma_{i}^{j} ;<1}\left(k_{j}+k_{l}\right)=\frac{1}{2} \sum_{j, l \in \Gamma_{i}, j \neq l}\left(k_{j}+k_{l}\right)=\sum_{j, l \Gamma_{i}, j \neq l} k_{j}=\left(k_{i}-1\right) \sum_{j \in \Gamma_{i}} k_{j}$


## more careful calculation (2)

- sum of the quantity over all vertices i :

$$
\sum_{i}\left(k_{i}-1\right) \sum_{j \in \Gamma_{j}} k_{j}=\sum_{i i} A_{i j}\left(k_{i}-1\right) k_{j}=\sum_{i j} A_{i j} k_{i} k_{j}-\sum_{j} k_{j}^{2} \quad \sum_{i} A_{i j}=k_{j}
$$

- compare ${ }_{j \in \Gamma_{i}}$ with the expression ${ }_{i j}^{i j}$ of assortativity

$$
r=\frac{\sum_{i j}\left(A_{j}-k_{i} k_{j} / 2 m\right) k_{i} k_{j}}{\sum_{j}\left(k_{j} \delta_{j}-k_{j} k_{j} / 2 m\right) k_{i} k_{j}} \text { same }
$$

- time tó calculate clustering coefficient depends on whether the degrees of vertices are correlated or not
- if no correlation, $\mathrm{r}=0 \sum_{i j} A_{i j} k_{i} k_{j}=\frac{1}{2 m} \sum_{i j} k_{i}^{2} k_{j}^{2}=\frac{1}{2 m}\left[\sum_{i} k_{i}^{2}\right]^{2}$
- running time for clustering coefficient:

$$
\frac{1}{2 m}\left[\sum_{i} k_{i}^{2}\right]^{2}-\sum_{j} k_{j}^{2}=n\left\langle k^{2}\right\rangle\left[\frac{\left\langle k^{2}\right\rangle}{\langle k\rangle}-1\right] \frac{2 m=\sum_{i} k_{i}=n\langle k\rangle}{\begin{array}{c}
\text { depends on }\left\langle k^{2}\right\rangle \text { : it can become large for } \\
\text { highly skewed degree distributions }
\end{array}}
$$

## cf. assortative mixing by degree

- assortative: high-degree vertices connect to other high-degree vertices
- core/periphery structure :

common feature of social network;-
- covariance

$$
\operatorname{cov}\left(k_{i}, k_{j}\right)=\frac{1}{2 m} \sum_{v j}\left(A_{j}-\frac{k, k_{j}}{2 m}\right)_{k, k_{j}}
$$


$\odot$

- correlation coefficient (assortativity coefficient)
$r=\frac{\sum_{i j}\left(A_{i j}-k_{i} k_{j} / 2 m\right) k_{i} k_{j}}{\sum_{i j}\left(k_{i} \delta_{i j}-k_{i} k_{j} / 2 m\right) k_{i} k_{j}}$


## more careful calculation (3)

- if a network is simple with no multiedges, $\rightarrow$ maximum degree is $\mathrm{k}=\mathrm{n}$, and the degree distribution is cut off $\rightarrow\left\langle\mathrm{k}^{2}\right\rangle$ scales at worst as $\mathrm{n}^{3-\alpha}$
- running time of clustering coefficient would go as $n \times n^{3-\alpha} \times n^{3-\alpha}=n^{7-2 \alpha}$
- $\alpha=3: O(n)$
- $\alpha=2: O\left(n^{3}\right)$


## shortest paths and breadth-first search

- the study of each of these algorithms has three parts
- description of the algorithm
- analysis of its running time
- proof that the algorithm performs the calculation it claims to


## breadth-first search (1)

- finds the shortest distance from a given starting vertex s to every other vertex in the same component as s
- s have distance 0 from itself
- find all the neighbors of $s$ (distance 1)
- find all the neighbors of those vertices (distance 2 )


## breadth-first search (2)

- every vertex whose shortest distance from s is d has a network neighbor whose shortest distance from $s$ is $d-1$
- breadth-first search finds the component to which vertex s belongs



## naive implementation



- array of $n$ elements to store the distance of
- distance variable d

1. find all vertices that are distance $d$ from $s$ vertices of distance $d$ : $\mathrm{n} \times \mathrm{O}(\mathrm{m} / \mathrm{n})=0(\mathrm{~m})$
2. find all the neighbors of those vertices and check to see if distance from s is unknown
3. if \# of neighbors with unknown distance is zero, the algorithm is over. otherwise, set the distance of each of those neighbors to $d+1$
4. increase the value of $d$ by 1
5. repeat from step 1
total running time:

n (in the worst case) $->0\left(m+\mathrm{n}^{2}\right)$ $\log \mathrm{n}$ (in most cases) $->0(\mathrm{~m}+\mathrm{nlog} \mathrm{n})$

## better implementation (1)

- time consuming part is step 1 (find vertices that are distance $d$ from the starting vertex s)
- use distance array and queue

| vertex | distance |
| :--- | :--- |
| 1 | -1 |
| 2 | -1 |
| 5 | 0 |
| 4 | -1 |
| 5 | -1 |



## better implementation (2)

1. place $s$ in the queue, set 0 in distance array ${ }^{5}$
2. if read and write pointers are pointing the same element, the algorithm is finished. otherwise, read the vertex label pointed by read pointer
3. find the distance $d$ for that vertex
4. go through each neighboring vertex in turn
5. if it has a known distance, leave it alone
6. if it has unknown distance, assign it distance $d+1$, store its label in the queue array


## better implementation (3)

- running time
$-\mathrm{O}(\mathrm{n})$ to set up the distance array
- for each element in the queue, we run through its neighbors ( $\mathrm{O}(\mathrm{m} / \mathrm{n}$ ) on average), calculate their distance and add them to the queue or do nothing (O(1))
- we require overall at most at time $\mathrm{n} \times \mathrm{O}(\mathrm{m} / \mathrm{n})=\mathrm{O}(\mathrm{m})$
- the whole algorithm takes time $\mathrm{O}(\mathrm{m}+\mathrm{n})$
- better than O(m+nlogn)
- for sparse network with $m \approx n, O(n)$


## finding shortest paths (1)

- the previous breadth-first search does not tell us the particular path(s) by which that shortest distance is achieved
- a small modification of the bread-first search algorithm
- if a vertex $j$ (neighbor of i) has unknown distanfice
- assign j a distance and store it in a queug
- add a directed edge from $j$ to $i$
- its running time is still $O(m+n)$


## finding shortest paths (2)

- for multiple shortest paths
- if a vertex $j$ (neighbor of i) has unknown distance
- assign j a distance and store it in a queue
- add a directed edge from $j$ to $i$
- if a vertex j has distance $\mathrm{d}+1$
- add an extra directed edge from $j$ to $i$


## betweenness centrality

- betweenness of v:\# of geodesic paths between pairs of vertices that pass through $v$
- simplest way: (1) use breadth-first search to find the shortest path between $s$ and $t$, and see if the vertex $v$ lies along the path (2) repeat this process for every distinct pair $s, t$.
- this is correct, but slow
$-\mathrm{O}(\mathrm{m}+\mathrm{n})$ to find a shortest path between two vertices
$-n(n-1) / 2$ distinct pairs of vertices
- calculating betweenness for $v$ will take $O\left(n^{2}(m+n)\right.$ ) (or $\mathrm{O}\left(\mathrm{n}^{3}\right)$ when $\mathrm{m} \approx \mathrm{n}$ (sparse network))


## calculation of betweenness centrality

(1)

- $\mathrm{O}(\mathrm{m}+\mathrm{n})$ for finding paths between s and all other vertices (a shortest path tree)
- trace the paths from each vertex back to $s$ and count the number of paths that go through $v$ : O(nlogn)


## $n$ paths of length logn

- O(m+nlogn) for each s
- repeat this calculation for all s -> O(n(m+nlogn)) (or O(n²logn))
- much better than $O\left(n^{3}\right)$
- betweenness of all vertices will be calculated at the same time


## we can still do better

- many of the shortest paths share many of the same edges
- score of each vertex (betweenness count for paths that end at vertex s)
- leaves:1
- others:1+ $\sum$ (score of its immediate children)
- the queue array created by BFS have a vertex $\rightarrow 0(\mathrm{~m}+\mathrm{n})$ list of vertices in order of decreasing
distance from $s->O(m+n)$ for each $s$
- $\mathrm{O}(\mathrm{n}(\mathrm{m}+\mathrm{n}))$ for all vertices $\mathrm{BFS}: \mathrm{O}(\mathrm{m}+\mathrm{n})$
worst case: checking every neighbor of every



## more than one shortest path (1)

counting the number of shortest paths from each vertex

1. assign vertex $s$ distance zero, set $d=0$, and assign $s$ a weight $\mathrm{w}_{\mathrm{s}}=1$
2. for each vertex i whose distance is d, follow each attached edge to the vertex $j$ at its other end, and then do one of the following three things:

- if j has not yet been assigned a distance, assign it distance $\mathrm{d}+1$ and weight $w_{j}=w_{i}$
- if $j$ has already been assigned distance $d+1$, then $w_{j}<-w_{j}+w_{i}$
- if $j$ has already been assigned a distance less than $d+1$, do nothing

3. increase $d$ by 1
4. repeat from 2 until there are no vertices that have ${ }^{1}$ distance d


## more than one shortest path (2)

- the fraction of the paths to $s$ that pass through $j$ and that also pass through $i$ is $w_{i} / w_{j}$

1. find every leaf vertex $t$ and assign it a score of $\mathrm{x}_{\mathrm{i}}=1$
2. from the bottom of the tree, assign to each vertex i a score $x_{i}=1+\sum_{j} x_{j} w_{i} / w_{j}$ ( j is a neighbor immediately below vertex i)
3. repeat 2 until vertex $s$ is reached

## shortest paths in networks with varying edge lengths

- the shortest path may traverse many edges
- Dijkstra's algorithm
- finds the shortest distance from a given source vertex s to every other vertex,
- it takes the lengths of edges into account
- it keeps a record of the shortest distance so far and update that record whenever a shorter one is found



##  <br> - two arrays of n elements certainty of the

- current estimates of the distance from
- certainty of the estimates
- find vertex $j$ that has the smallest estimated distance (smallest and not yet certain)
- mark this distance as being certain
- calculate the distances from $s$ via $v$ to each of the neighbors of $v$. if the resulting distance is smaller, replace the estimate
- repeat from 1 until the distances to all vertices are certain


## the smallest estimated distance is certain

- if $v$ is the vertex with the smallest estimated distance from s , then that estimated distance must be the true shortest distance to $v$.
- if there were a shorter path $\mathrm{s}, \ldots, \mathrm{x}, \mathrm{y}, \ldots, \mathrm{v}$,
- all points along the path must have shorter distances from $s$ than v's estimated distance
- it means that y has a smaller estimated distance than v , which is impossible.


## running time of Dijkstra's algorithm

- simplest implementation
- find the smallest estimated distance : $O(n) \longrightarrow O(m / n+n)$
- calculate new estimates of its neighbors: $O(\mathrm{~m} / \mathrm{n})$
- repeat above for $n$ rounds: $O\left(m+n^{2}\right)$
- better implementation
- find the smallest estimate from binary heap: $\mathrm{O}(\operatorname{logn})$
- replacing an estimated distance a new one : O(logn) (x $\mathrm{O}(\mathrm{m} / \mathrm{n})$ times in the worst case)
- repeat $n$ rounds: $O((m+n) \operatorname{logn})$ (or $O(n \operatorname{logn})$ for a sparse network)


## maximum flows and minimum cuts

- augmenting path algorithm (Ford-Fulkerson) : $\mathrm{O}((\mathrm{m}+\mathrm{n}) \mathrm{m} / \mathrm{n})$
- basic idea:
- find a path from s to $t$ using BFS
- find another using only the edges not used in the path



## augmenting path algorithm

- the procedure will not always find the maximum flow but here are ww edge
- if we remove the edges of the shortest path, there is no more paths
- if we allow flows in both directions
along an edge, we can find another
- the maximum possible flow is 2 s


## residual graph (1)

- a directed network in which the edges connect the pairs of vertices on the original network between which we still have capacity available to carry one or more units of flow in the given direction
initial



## residual graph (2)



- the largest number of edges we update: $\mathrm{O}(\mathrm{m})$
- BFS: $\mathrm{O}(m+n)$
- the number of independent paths $\leq \min \left(k_{s}, k_{t}\right)$
- time complexity of this algorithm: $\mathrm{O}\left(\min \left(\mathrm{k}_{s}, \mathrm{k}_{\mathrm{t}}\right)(\mathrm{m}+\mathrm{n})\right)$
- average running time: $\mathrm{O}((\mathrm{m}+\mathrm{n}) \mathrm{m} / \mathrm{n}) \because\left\langle\min \left(k_{s}, k_{t}\right)\right\rangle \leq\langle k\rangle=2 m / n$


## why augmenting path algorithm correctly finds maximum flows

- if at some point in our algorithm the flow from $s$ to $t$ is less than the maximum possible flow, then there must exist at least one augmenting path on the current residual graph
- f: flows on the network
- $\mathrm{f}_{\text {max }}$ : maximum possible flow
- $\Delta f=f_{\text {max }}-f$


## correctness of the augmenting path algorithm



## finding independent paths

- augmenting paths $\neq$ independent paths

reconstructing the independent paths from the residual graph
- deleting every pair of edges that join the same two vertices in opposite directions -> graph consisting of the independent paths only



## finding a minimum cut set

- maximum flows residual graph



## finding vertex-independent paths

- almost the same as augmenting path
algorithm
- but no two paths may pass through the same vertex
- mapping from the vertex-independent path problem to the edge-independent path problem
- replace each vertex (except for $s$ and $t$ ) with a pair of vertices with a directed edge between them
vertex transformation for the vertexindependent path algorithm
- each vertex is replaced by a pair of vertices joined by a single directed edge


