CAIM: Cerca i Anàlisi d'Informació Massiva FIB, Grau en Enginyeria Informàtica

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Fall 2018

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7. Introduction to Network Analysis

Network Analysis, Part I

Today's contents

- 1. Examples of real networks
- 2. What do real networks look like?
 - real networks exhibit small diameter
 - .. and so does the Erdös-Rényi or random model
 - real networks have high clustering coefficient
 - .. and so does the Watts-Strogatz model
 - real networks' degree distribution follows a power-law
 - .. and so does the Barabasi-Albert or preferential attachment model

Examples of real networks

- Social networks
- Information networks
- Technological networks
- Biological networks

Social networks

Links denote social "interactions"

▶ friendship, collaborations, e-mail, etc.



Information networks

Nodes store information, links associate information

citation networks, the web, p2p networks, etc.



Technological networks

Man-built for the distribution of a commodity

 telephone networks, power grids, transportation networks, etc.



Biological networks

Represent biological systems

 protein-protein interaction networks, gene regulation networks, metabolic pathways, etc.



Representing networks

- Network \equiv Graph
- Networks are just collections of "points" joined by "lines"



points	lines	
vertices	edges, arcs	math
nodes	links	computer science
sites	bonds	physics
actors	ties, relations	sociology

Types of networks

From [Newman, 2003]



- (a) unweighted, undirected
- (b) discrete vertex and edge types, undirected
- (c) varying vertex and edge weights, undirected
- (d) directed

Small-world phenomenon

- A friend of a friend is also frequently a friend
- Only 6 hops separate any two people in the world



Measuring the small-world phenomenon, I

- Let d_{ij} be the shortest-path distance between nodes i and j
- To check whether "any two nodes are within 6 hops", we use:
 - The diameter (longest shortest-path distance) as

$$d = \max_{i,j} d_{ij}$$

The average shortest-path length as

$$l = \frac{2}{n \ (n+1)} \sum_{i>j} d_{ij}$$

The harmonic mean shortest-path length as

$$l^{-1} = \frac{2}{n \ (n+1)} \sum_{i>j} d_{ij}^{-1}$$

From [Newman, 2003]

	network	type	n	m	2	l	α	$C^{(1)}$	$C^{(2)}$	r	Ref(s).
	film actors	undirected	449 913	25516482	113.43	3.48	2.3	0.20	0.78	0.208	20, 416
	company directors	undirected	7 673	55 392	14.44	4.60	-	0.59	0.88	0.276	105, 323
	math coauthorship	undirected	253 339	496 489	3.92	7.57	-	0.15	0.34	0.120	107, 182
	physics coauthorship	undirected	52 909	245300	9.27	6.19	-	0.45	0.56	0.363	311, 313
lai	biology coauthorship	undirected	1520251	11 803 064	15.53	4.92	-	0.088	0.60	0.127	311, 313
õ	telephone call graph	undirected	47 000 000	80 000 000	3.16		2.1				8, 9
	email messages	directed	59 912	86 300	1.44	4.95	1.5/2.0		0.16		136
	email address books	directed	16 881	57 029	3.38	5.22	-	0.17	0.13	0.092	321
	student relationships	undirected	573	477	1.66	16.01	-	0.005	0.001	-0.029	45
	sexual contacts	undirected	2810				3.2				265, 266
-	WWW nd.edu	directed	269 504	1 497 135	5.55	11.27	2.1/2.4	0.11	0.29	-0.067	14, 34
mation	WWW Altavista	directed	203 549 046	2130000000	10.46	16.18	2.1/2.7				74
	citation network	directed	783 339	6716198	8.57		3.0/-				351
ufor	Roget's Thesaurus	directed	1 0 2 2	5 103	4.99	4.87	-	0.13	0.15	0.157	244
	word co-occurrence	undirected	460 902	17 000 000	70.13		2.7		0.44		119, 157
	Internet	undirected	10 697	31 992	5.98	3.31	2.5	0.035	0.39	-0.189	86, 148
5	power grid	undirected	4 941	6 5 9 4	2.67	18.99	-	0.10	0.080	-0.003	416
gic	train routes	undirected	587	19 603	66.79	2.16	-		0.69	-0.033	366
loc	software packages	directed	1 439	1 723	1.20	2.42	1.6/1.4	0.070	0.082	-0.016	318
-th	software classes	directed	1 377	2 213	1.61	1.51	-	0.033	0.012	-0.119	395
ţ	electronic circuits	undirected	24 097	53 248	4.34	11.05	3.0	0.010	0.030	-0.154	155
	peer-to-peer network	undirected	880	1 296	1.47	4.28	2.1	0.012	0.011	-0.366	6, 354
	metabolic network	undirected	765	3 686	9.64	2.56	2.2	0.090	0.67	-0.240	214
B	protein interactions	undirected	2 1 1 5	2 240	2.12	6.80	2.4	0.072	0.071	-0.156	212
logi	marine food web	directed	135	598	4.43	2.05	-	0.16	0.23	-0.263	204
bio	freshwater food web	directed	92	997	10.84	1.90	-	0.20	0.087	-0.326	272
	neural network	directed	307	2 359	7.68	3.97	-	0.18	0.28	-0.226	416, 421

- Can we mimic this phenomenon in simulated networks ("models")?
- The answer is YES!

The (basic) random graph model a.k.a. ER model

Basic $G_{n,p}$ Erdös-Rényi random graph model:

- parameter n is the number of vertices
- parameter p is s.t. $0 \le p \le 1$
- Generate and edge (i, j) independently at random with probability p

Measuring the diameter in ER networks

Want to show that the diameter in ER networks is small



- ▶ Let the average degree be *z*
- At distance l, can reach z^l nodes
- At distance $\frac{\log n}{\log z}$, reach all *n* nodes
- So, diameter is (roughly) $O(\log n)$

ER networks have small diameter

As shown by the following simulation



Measuring the small-world phenomenon, II

- To check whether "the friend of a friend is also frequently a friend", we use:
 - The transitivity or clustering coefficient, which basically measures the probability that two of my friends are also friends



Global clustering coefficient

$$C = \frac{3 \times \text{number of triangles}}{\text{number of connected triples}}$$



$$C = \frac{3 \times 1}{8} = 0.375$$

Local clustering coefficient

- For each vertex i, let n_i be the number of neighbors of i
- Let C_i be the fraction of pairs of neighbors that are connected within each other

$$C_i = rac{ {
m nr. of connections between } i {
m s neighbors}}{rac{1}{2} n_i \; (n_i-1)}$$

Finally, average C_i over all nodes i in the network

$$C = \frac{1}{n} \sum_{i} C_i$$

Local clustering coefficient example



C₁ = C₂ = 1/1
C₃ = 1/6

$$\blacktriangleright C_4 = C_5 = 0$$

• $C = \frac{1}{5}(1+1+1/6) = 13/30 = 0.433$

From [Newman, 2003]

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ER networks do not show transitivity

- C = p, since edges are added independently
- Given a graph with n nodes and e edges, we can "estimate" p as

$$\hat{p} = \frac{e}{1/2 \ n \ (n-1)}$$

- We say that clustering is high if $C \gg \hat{p}$
 - ► Hence, ER networks do not have high clustering coefficient since for them $C \approx \hat{p}$

ER networks do not show transitivity

Network	n	z	C	C for
			measured	random graph
Internet [153]	6,374	3.8	0.24	0.00060
World Wide Web (sites) [2]	153,127	35.2	0.11	0.00023
power grid [192]	4,941	2.7	0.080	0.00054
biology collaborations [140]	1,520,251	15.5	0.081	0.000010
mathematics collaborations [141]	253,339	3.9	0.15	0.000015
film actor collaborations [149]	449,913	113.4	0.20	0.00025
company directors [149]	7,673	14.4	0.59	0.0019
word co-occurrence [90]	460,902	70.1	0.44	0.00015
neural network [192]	282	14.0	0.28	0.049
metabolic network [69]	315	28.3	0.59	0.090
food web [138]	134	8.7	0.22	0.065

Table 1: Clustering coefficients, C, for a number of different networks; n is the number of node, z is the mean degree. Taken from [146].

So ER networks do not have high clustering, but..

- Can we mimic this phenomenon in simulated networks ("models"), while keeping the diameter small?
- The answer is YES!

The Watts-Strogatz model, I

From [Watts and Strogatz, 1998]

Reconciling two observations from real networks:

- High clustering: my friend's friends are also my friends
- small diameter



The Watts-Strogatz model, II

- Start with all n vertices arranged on a ring
- Each vertex has initially 4 connections to their closest nodes
 - mimics local or geographical connectivity
- With probability p, rewire each local connection to a random vertex
 - p = 0 high clustering, high diameter
 - p = 1 low clustering, low diameter (ER model)
- What happens in between?
 - As we increase p from 0 to 1
 - Fast decrease of mean distance
 - Slow decrease in clustering



Increasingly random connectivity

The Watts-Strogatz model, III

For an appropriate value of $p \approx 0.01$ (1 %), we observe that the model achieves high clustering and small diameter



Degree distribution

Histogram of nr of nodes having a particular degree



 $f_k =$ fraction of nodes of degree k

Scale-free networks

The degree distribution of most real-world networks follows a power-law distribution

$$f_k = ck^{-\alpha}$$



- "heavy-tail" distribution, implies existence of hubs
- hubs are nodes with very high degree

Random networks are not scale-free!

For random networks, the degree distribution follows the binomial distribution (or Poisson if n is large)

$$f_k = \binom{n}{k} p^k (1-p)^{(n-k)} \approx \frac{z^k e^{-z}}{k!}$$

- Where z = p(n-1) is the mean degree
- Probability of nodes with very large degree becomes exponentially small
 - so no hubs

So ER networks are not scale-free, but..

- Can we obtained scale-free simulated networks?
- The answer is YES!

Preferential attachment

- "Rich get richer" dynamics
 - The more someone has, the more she is likely to have
- Examples
 - the more friends you have, the easier it is to make new ones
 - the more business a firm has, the easier it is to win more
 - the more people there are at a restaurant, the more who want to go

Barabási-Albert model

From [Barabási and Albert, 1999]

- "Growth" model
 - The model controls how a network grows over time
- Uses preferential attachment as a guide to grow the network
 - new nodes prefer to attach to well-connected nodes
- (Simplified) process:
 - the process starts with some initial subgraph
 - each new node comes in with m edges
 - probability of connecting to existing node *i* is proportional to *i*'s degree
 - \blacktriangleright results in a power-law degree distribution with exponent $\alpha=3$

ER vs. BA

Experiment with 1000 nodes, 999 edges ($m_0 = 1$ in BA model).



random



preferential attachment

In summary..

phenomenon	real networks	ER	WS	BA
small diameter	yes	yes	yes	yes
high clustering	yes	no	yes	yes ¹
scale-free	yes	no	no	yes

¹clustering coefficient is higher than in random networks, but not as high as for example in WS networks
Network Analysis, Part II

Today's contents

1. Centrality

- Degree centrality
- Closeness centrality
- Betweenness centrality
- 2. Community finding algorithms
 - Hierarchical clustering
 - Agglomerative
 - Girvan-Newman
 - Modularity maximization: Louvain method

Centrality in Networks

Centrality is a node's measure w.r.t. others

- A central node is important and/or powerful
- A central node has an influential position in the network
- A central node has an advantageous position in the network



Power through connections

$$degree_centrality(i) \stackrel{def}{=} k(i)$$



Power through connections

$$in_degree_centrality(i) \stackrel{def}{=} k_{in}(i)$$



Power through connections

$$out_degree_centrality(i) \stackrel{def}{=} k_{out}(i)$$



Power through connections

By the way, there is a *normalized* version which divides the centrality of each degree by the maximum centrality value possible, i.e. n - 1 (so values are all between 0 and 1).



But look at these examples, does degree centrality look OK to you?

Closeness centrality

Power through proximity to others

$$closeness_centrality(i) \stackrel{def}{=} \left(\frac{\sum_{j \neq i} d(i, j)}{n - 1}\right)^{-1} = \frac{n - 1}{\sum_{j \neq i} d(i, j)}$$



Here, what matters is to be close to everybody else, i.e., to be easily reachable or have the power to quickly reach others.

Betweenness centrality

Power through brokerage

A node is important if it lies in many shortest-paths

► so it is essential in passing information through the network



Betweenness centrality

Power through brokerage

$$betweenness_centrality(i) \stackrel{def}{=} \sum_{j < k} \frac{g_{jk}(i)}{g_{jk}}$$

Where

- g_{jk} is the number of shortest-paths between j and k, and
- $g_{jk}(i)$ is the number of shortest-paths through i

Oftentimes it is normalized:

 $norm_betweenness_centrality(i) \stackrel{def}{=} \frac{betweenness_centrality(i)}{\binom{n-1}{2}}$

Betweenness centrality

Examples (non-normalized)



What is community structure?



Why is community structure important?



.. but don't trust visual perception

it is best to use objective algorithms













Main idea

A community is dense in the inside but sparse w.r.t. the outside

No universal definition! But some ideas are:

- A community should be densely connected
- A community should be *well-separated* from the rest of the network
- Members of a community should be more similar among themselves than with the rest

Most common..

nr. of intra-cluster edges > nr. of inter-cluster edges

Some definitions

Let G = (V, E) be a network with |V| = n nodes and |E| = m edges. Let C be a subset of nodes in the network (a "cluster" or "community") of size $|C| = n_c$. Then

intra-cluster density:

$$\delta_{int}(C) = rac{\operatorname{nr. internal edges of } C}{n_c(n_c - 1)/2}$$

inter-cluster density:

$$\delta_{ext}(C) = \frac{\text{nr. inter-cluster edges of } C}{n_c(n - n_c)}$$

A community should have $\delta_{int}(C) > \delta(G)$, where $\delta(G)$ is the average edge density of the whole graph *G*, i.e.

$$\delta(G) = \frac{\text{nr. edges in } G}{n(n-1)/2}$$

Most algorithms search for tradeoffs between large $\delta_{int}(C)$ and small $\delta_{ext}(C)$

• e.g. optimizing $\sum_C \delta_{int}(C) - \delta_{ext}(C)$ over all communities C

Define further:

- $m_c = \text{nr. edges within cluster } C = |\{(u, v)|u, v \in C\}|$
- ▶ f_c = nr. edges in the frontier of $C = |\{(u, v) | u \in C, v \notin C\}|$



- $n_{c_1} = 4, m_{c_1} = 5, f_{c_1} = 2$
- $n_{c_2} = 3, m_{c_2} = 3, f_{c_2} = 2$

$$\bullet \ n_{c_3} = 5, m_{c_3} = 8, f_{c_3} = 2$$

Community quality criteria

- **conductance**: fraction of edges leaving the cluster $\frac{f_c}{2m_c+f_c}$
- expansion: nr of edges per node leaving the cluster $\frac{f_c}{n_a}$
- internal density: a.k.a. "intra-cluster density" $\frac{m_c}{n_c(n_c-1)/2}$
- cut ratio: a.k.a. "inter-cluster density" $\frac{f_c}{n_c(n-n_c)}$
- ▶ modularity: difference between nr. of edges in *C* and the expected nr. of edges *E*[*m_c*] of a random graph with the same degree distribution

$$\frac{1}{4m}(m_c - E[m_c])$$

Methods we will cover

- Hierarchical clustering
 - Agglomerative
 - Divisive (Girvan-Newman algorithm)
- Modularity maximization algorithms
 - Louvain method

Hierarchical clustering

From hairball to dendogram



Suitable if input network has hierarchical structure





Agglomerative hierarchical clustering [Newman, 2010]

Ingredients

- Similarity measure between nodes
- Similarity measure between sets of nodes

Pseudocode

- 1. Assign each node to its own cluster
- 2. Find the cluster pair with highest similarity and join them together into a cluster
- 3. Compute new similarities between new joined cluster and others
- 4. Go to step 2 until all nodes form a single cluster



D. Blei Clustering 02



iteration 001

V1

D. Blei Clustering 02



iteration 002

V1

D. Blei Clustering 02



iteration 003

D. Blei Clustering 02



iteration 004

V1

D. Blei Clustering 02



iteration 005

D. Blei Clustering 02



iteration 006

D. Blei Clustering 02



iteration 007

D. Blei Clustering 02



iteration 008

D. Blei Clustering 02



iteration 009

D. Blei Clustering 02



iteration 010

D. Blei Clustering 02



iteration 011

D. Blei Clu



iteration 012

D. Blei Clustering 02



iteration 013

D. Blei Clustering 02



iteration 014

D. Blei Clustering 02


iteration 015



iteration 016



iteration 017



D. Blei Clus





iteration 020

V1

D. Blei Clustering 02



iteration 021

V1



iteration 022



iteration 023

V1



V1

Similarity measures w_{ij} for nodes I

Let A be the adjacency matrix of the network, i.e. $A_{ij} = 1$ if $(i, j) \in E$ and 0 otherwise.

Jaccard index:

$$w_{ij} = \frac{|\Gamma(i) \cap \Gamma(j)|}{|\Gamma(i) \cup \Gamma(j)|}$$

where $\Gamma(i)$ is the set of neighbors of node i

Cosine similarity:²

$$w_{ij} = \frac{\sum_k A_{ik} A_{kj}}{\sqrt{\sum_k A_{ik}^2} \sqrt{\sum_k A_{jk}^2}} = \frac{n_{ij}}{\sqrt{k_i k_j}}$$

where:

• $n_{ij} = |\Gamma(i) \cap \Gamma(j)| = \sum_k A_{ik} A_{kj}$, and • $k_i = \sum_k A_{ik}$ is the degree of node i

Similarity measures w_{ij} for nodes II

 Euclidean distance: (or rather Hamming distance since A is binary)

$$d_{ij} = \sum_{k} (A_{ik} - A_{jk})^2$$

Normalized Euclidean distance:³

$$d_{ij} = \frac{\sum_{k} (A_{ik} - A_{jk})^2}{k_i + k_j} = 1 - 2\frac{n_{ij}}{k_i + k_j}$$

Pearson correlation coefficient

$$r_{ij} = \frac{cov(A_i, A_j)}{\sigma_i \sigma_j} = \frac{\sum_k (A_{ik} - \mu_i)(A_{jk} - \mu_j)}{n\sigma_i \sigma_j}$$

where
$$\mu_i = rac{1}{n}\sum_k A_{ik}$$
 and $\sigma_i = \sqrt{rac{1}{n}\sum_k (A_{ik}-\mu_i)^2}$

²From the equation $\mathbf{xy} = |\mathbf{x}||\mathbf{y}|\cos\theta$

³Uses the idea that the maximum value of d_{ij} is when there are no common neighbors and then $d_{ij} = k_i + k_j$

Similarity measures for sets of nodes

- ► Single linkage: $s_{XY} = \max_{x \in X, y \in Y} s_{xy}$
- Complete linkage: $s_{XY} = \min_{x \in X, y \in Y} s_{xy}$

• Average linkage:
$$s_{XY} = \frac{\sum_{x \in X, y \in Y} s_{xy}}{|X| \times |Y|}$$

Agglomerative hierarchical clustering on Zachary's network

Using average linkage



The Girvan-Newman algorithm

A divisive hierarchical algorithm [Girvan and Newman, 2002]

Edge betweenness

The betweenness of an edge is the nr. of shortest-paths in the network that pass through that edge

It uses the idea that "bridges" between communities must have high edge betweenness



The Girvan-Newman algorithm

Pseudocode

- 1. Compute betweenness for all edges in the network
- 2. Remove the edge with highest betweenness
- 3. Go to step 1 until no edges left

Result is a dendogram



Definition of modularity [Newman, 2010] Using a *null* model

Random graphs are not expected to have community structure, so we will use them as null models.

Q = (nr. of intra-cluster communities) - (expected nr of edges)

In particular:

$$Q = \frac{1}{2m} \sum_{ij} (A_{ij} - P_{ij}) \,\delta(C_i, C_j)$$

where P_{ij} is the expected number of edges between nodes iand j under the null model, C_i is the community of vertex i, and $\delta(C_i, C_j) = 1$ if $C_i = C_j$ and 0 otherwise.

How do we compute P_{ij} ?

Using the "configuration" null model

The "configuration" random graph model choses a graph with the same degree distribution as the original graph uniformly at random.

- Let us compute P_{ij}
- ► There are 2*m* stubs or half-edges available in the configuration model
- Let p_i be the probability of picking at random a stub incident with i

$$p_i = \frac{k_i}{2m}$$

• The probability of connecting *i* to *j* is then $p_i p_j = \frac{k_i k_j}{4m^2}$

• And so
$$P_{ij} = 2mp_ip_j = \frac{k_ik_j}{2m}$$

Properties of modularity

$$Q = \frac{1}{2m} \sum_{ij} \left(A_{ij} - \frac{k_i k_j}{2m} \right) \, \delta(C_i, C_j)$$

- Q depends on nodes in the same clusters only
- Larger modularity means better communities (better than random intra-cluster density)

$$Q \leq \frac{1}{2m} \sum_{ij} A_{ij} \, \delta(C_i, C_j) \leq \frac{1}{2m} \sum_{ij} A_{ij} \leq 1$$

- Q may take negative values
 - partitions with large negative Q implies existence of cluster with small internal edge density and large inter-community edges

The Louvain method [Blondel et al., 2008]

Considered state-of-the-art



Pseudocode

- 1. Repeat until local optimum reached
 - 1.1 Phase 1: partition network greedily using modularity
 - 1.2 Phase 2: agglomerate found clusters into new nodes

The Louvain method

Phase 1: optimizing modularity

Pseudocode for phase 1

- 1. Assign a different community to each node
- 2. For each node *i*
 - For each neighbor j of i, consider removing i from its community and placing it to j's community
 - Greedily chose to place *i* into community of neighbor that leads to highest modularity gain
- 3. Repeat until no improvement can be done

The Louvain method

Phase 2: agglomerating clusters to form new network

Pseudocode for phase 2

- 1. Let each community C_i form a new node i
- 2. Let the edges between new nodes i and j be the sum of edges between nodes in C_i and C_j in the previous graph (notice there are self-loops)

The Louvain method

Observations

- The output is also a hierarchy
- Works for weighted graphs, and so modularity has to be generalized to

$$Q^{w} = \frac{1}{2W} \sum_{ij} \left(W_{ij} - \frac{s_i s_j}{2W} \right) \, \delta(C_i, C_j)$$

where W_{ij} is the weight of undirected edge (i, j), $W = \sum_{ij} W_{ij}$ and $s_i = \sum_k W_{ik}$.

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