

# **Network Analysis**

Héctor Corrada Bravo

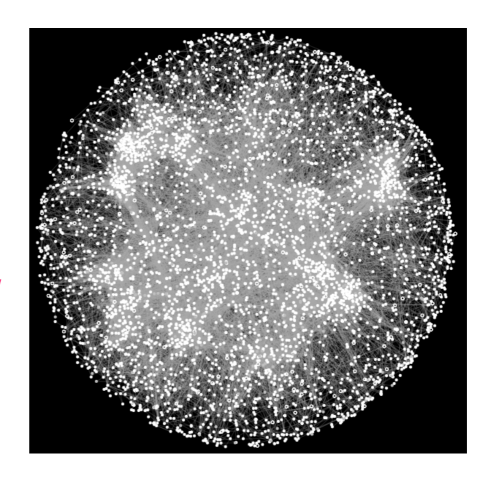
University of Maryland, College Park, USA CMSC644 2019-05-01



#### Genetic Interaction Network

- Yeast high-throuput doubleknockdown assay
- ~5000 genes
- ~800k interactions

http://www.geneticinteractions.org/



Costanzo et al. (2016) Science. DOI: 10.1126/science.aaf1420

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#### Genetic Interaction Network

• Number of vertices: 2803

• Number of edges: 67,268

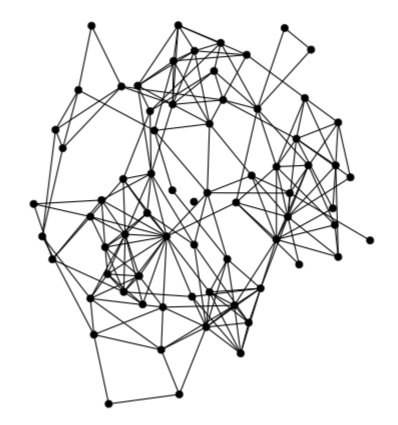
#### Preliminaries

Network: abstraction of entities and their interactions
Graph: mathematical representation

vertices: nodes

edges: links

#### Undirected graph



#### **Preliminaries**

Network: abstraction of entities and their interactions
Graph: mathematical representation

vertices: nodes

edges: links

# Directed graph

Number of vertices: n

In our example: *number of genes* 

Number of vertices: n

In our example: *number of genes* 

Number of edges: m

In our example: *number of genetic interactions* 

Number of vertices: *n* 

In our example: *number of genes* 

Number of edges: m

In our example: *number of genetic interactions* 

Degree of vertex i:  $k_i$ 

Number of genetic interactions for gene i

#### On the board:

- ullet Calculate number of edges m using degrees  $k_i$  (for both directed and undirected networks)
- Calculate *average degree c*
- Calculate *density*  $\rho$

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- Calculate number of edges m using degrees  $k_i$  (for both directed and undirected networks)
- Calculate *average degree c*
- Calculate density  $\rho$

#### In our example:

Average degree: 47.9971459

Density: 0.0171296

#### (On the board)

Number of edges using degrees (undirected)

$$m=rac{1}{2}\sum_{i=1}^n k_i$$

Number of edges using degrees (directed)

$$m = \sum_{i=1}^n k_i^{ ext{in}} = \sum_{i=1}^n k_i^{ ext{out}}$$

## (On the board)

Average degree

$$c = rac{1}{n} \sum_{i=1}^n k_i$$

Density

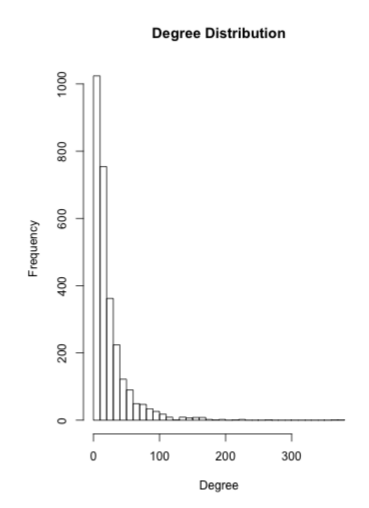
$$ho=rac{m}{inom{n}{2}}=rac{2m}{n(n-1)}=rac{c}{n-1}pproxrac{c}{n}$$

## Degree distribution

Fundamental analytical tool to characterize networks

 $p_k$ : probability randomly chosen vertex has degree k

On the board: how to calculate  $p_k$  and how to calculate average degree c using degree distribution.



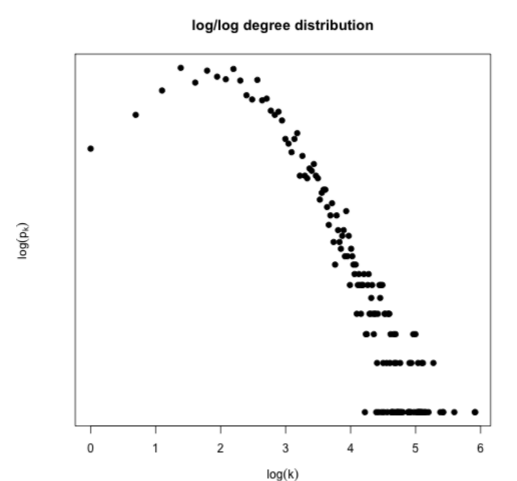
## (On the board)

Degree distribution

$$p_k=rac{n_k}{n}$$

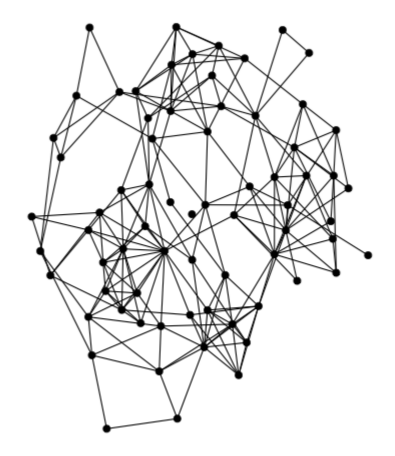
 $n_k$ : number of nodes in graph with degree k

## Degree Distribution



#### Paths and Distances

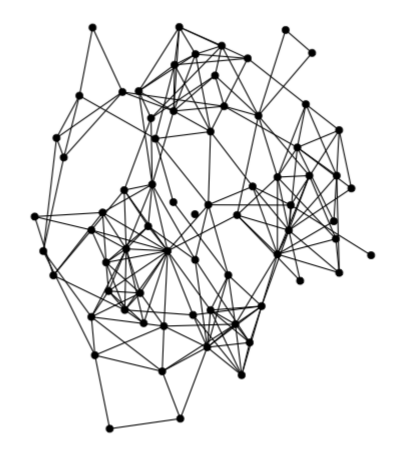
Distance  $d_{ij}$ : length of shortest path between vertices i and j.



#### Paths and Distances

Distance  $d_{ij}$ : length of shortest path between vertices i and j.

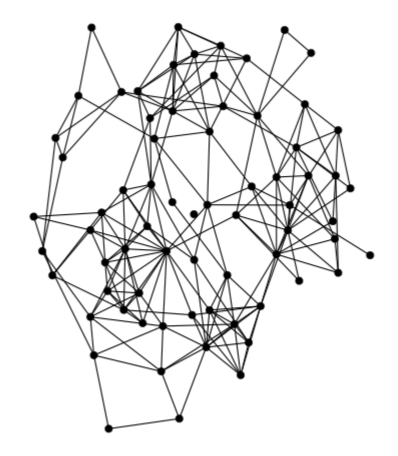
Diameter. longest shortest path  $\max_{ij} d_{ij}$ 



#### Paths and Distances

Distance  $d_{ij}$ : length of shortest path between vertices i and j.

On the board: average path length

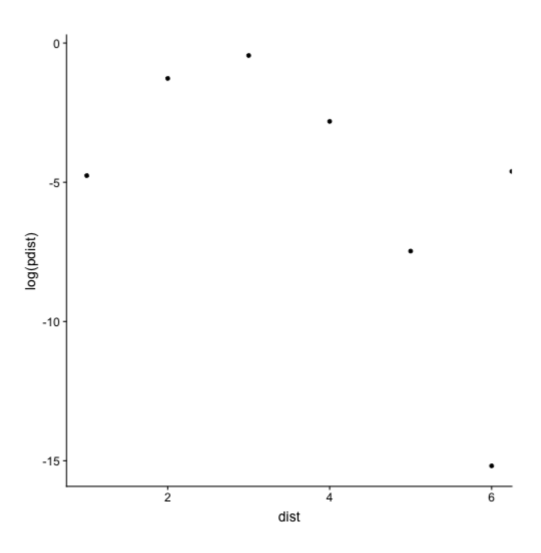


## (On the board)

Average path length

$$\overline{d} = rac{1}{n(n-1)} \sum_{i,j;i 
eq j} d_{ij}$$

## Distance Distribution



By convention: if there is no path between vertices i and j then  $d_{ij}=\infty$ 

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*Vertices* i and j are connected if  $d_{ij} < \infty$ 

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*Graph* is connected if  $d_{ij} < \infty$  for all i,j

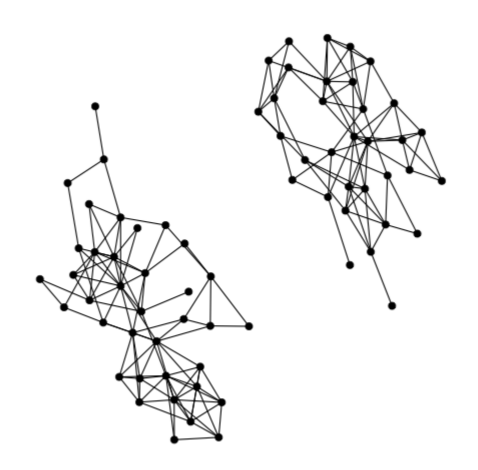
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*Vertices* i and j are connected if  $d_{ij} < \infty$ 

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Components maximal subset of connected components

# Components



# Clustering Coefficient

Another quantity of interest: how dense is the neighborhood around vertex i?

Do the genes that interact with me also interact with each other?

Related to the *locality* property.

Definition on the board

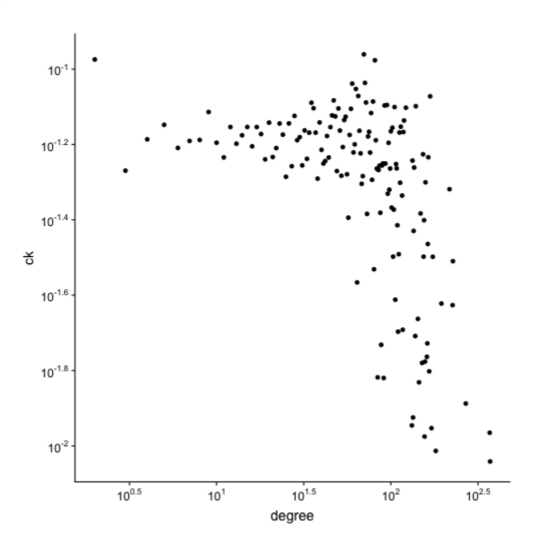
#### (On the board)

Clustering coefficient

$$c_i = rac{2m_i}{k_i(k_i-1)}$$

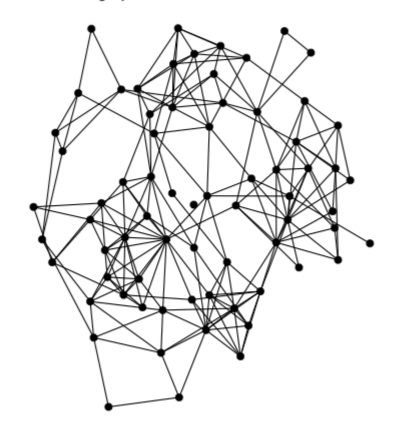
 $m_i$ : number of edges between neighbors of vertex i

# Clustering coefficient



# Adjacency Matrix

#### Undirected graph





## Adjacency Matrix

#### On the board:

- Definition
- Computing degree with adj.
   matrix
- ullet Computing num. edges m with adj. matrix
- Computing paths with adj. matrix



# Adjacency Matrix

#### Directed graph





## Weighted networks

Edges are assigned a weight indicating quantitative property of interaction

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Edges are assigned a weight indicating quantitative property of interaction

- Strength of genetic interaction (evidence from experiment)
- Rates in a metabolic network
- Spatial distance in an ecological network

Adjacency matrix contains weights instead of 0/1 entries

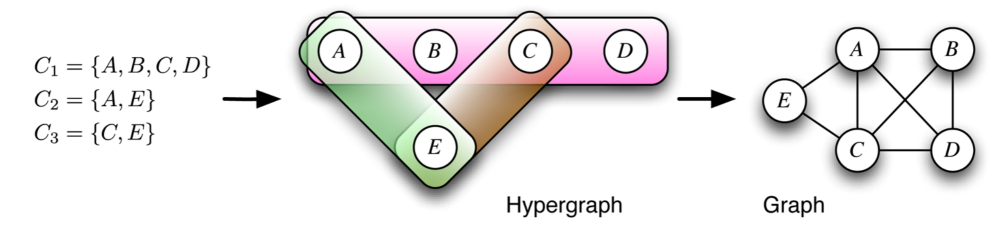
Adjacency matrix contains weights instead of 0/1 entries

Path lengths are the sum of edge weights in a path

# Hypergraphs

#### Edges connect more than two vertices

A Protein-protein interaction network

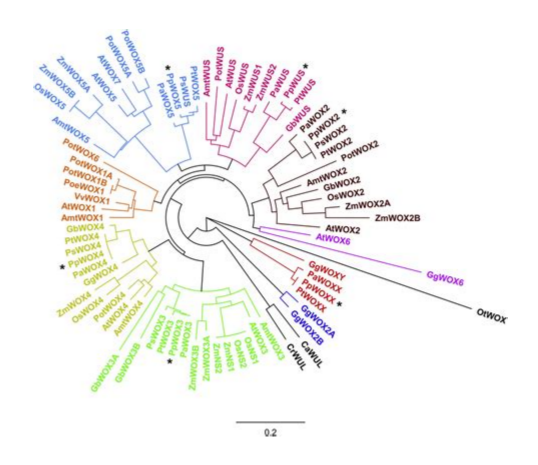


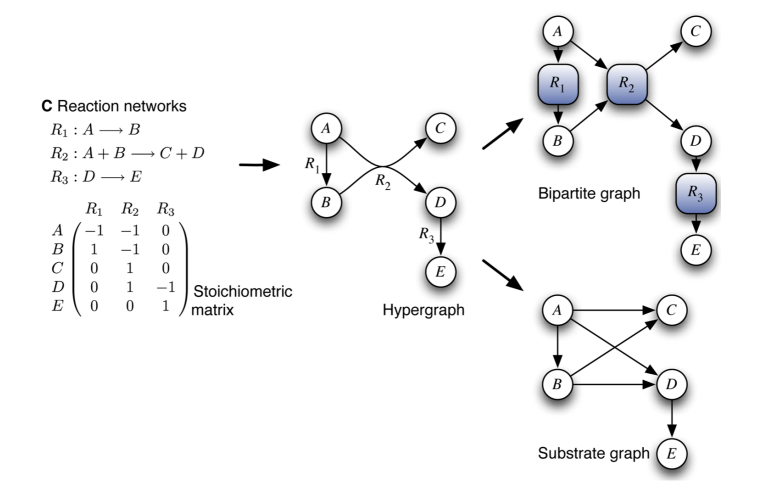
https://journals.plos.org/ploscompbiol/article?id=10.1371/journal.pcbi.1000385

#### Trees

Acyclic graphs

Single path between any pair of vertices





We use an *Incidence Matrix* B instead of *Adjacency Matrix* 

(On the board): definition

**Projections** 

 $\mathit{vertex\ projection}$ :  $P_{ij}$ , num. of groups in which vertices i and j co-occur

group projection:  $P_{ij}^{\prime}$ , num. of members groups i and j share

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(On the board)

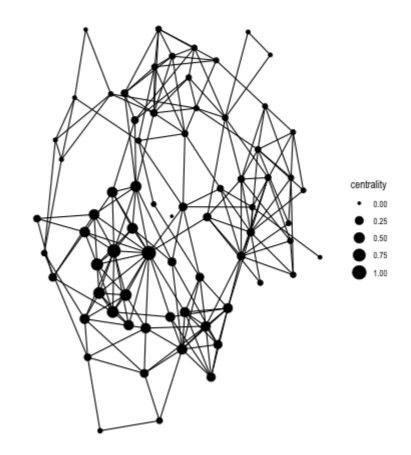
$$P = B^T B$$

$$P' = BB^T$$

# Centrality

What are the *important* nodes in the network?

What are *central* nodes in the network?



# Centrality

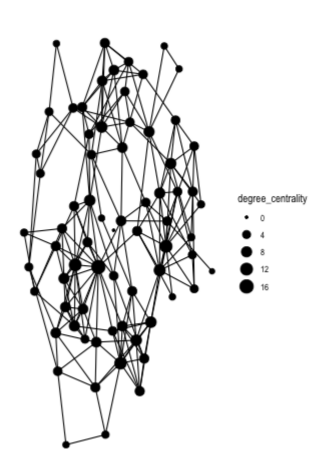
**Undirected Graphs** 

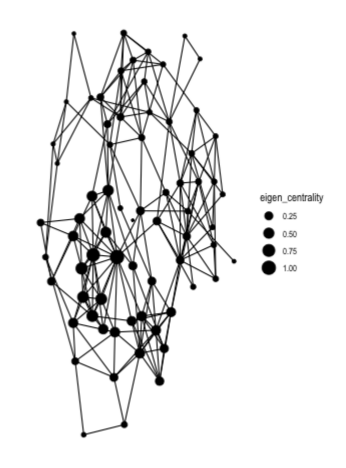
• Eigenvalue Centrality

Directed Graphs

- Katz Centrality
- Pagerank

# Centrality

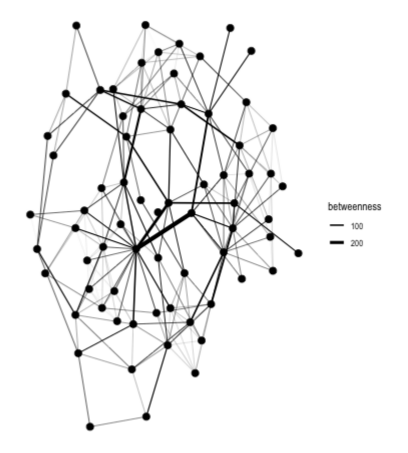




#### Betweenness

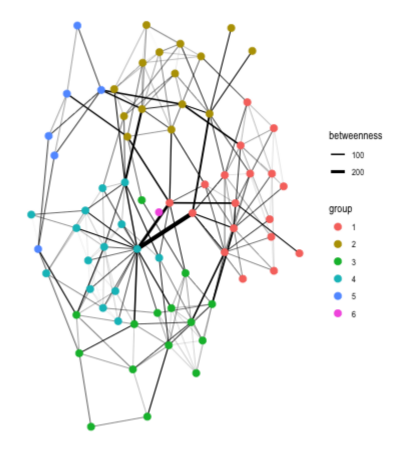
What are the *important* edges in the network?

What are edges that may connect clusters of nodes in the network?



#### Betweenness

Girvan-Newman Algorithm hierarchical method to
partition nodes into
communities using edge
betweenness



### Girvan-Newman Algorithm

Two phases:

Phase One: Compute betweenness for every edge

Phase Two: Discover communities by removing *high* betweenness

edges (similar to hierarchical clustering)

### Girvan-Newman Algorithm

Calculating Betweenness

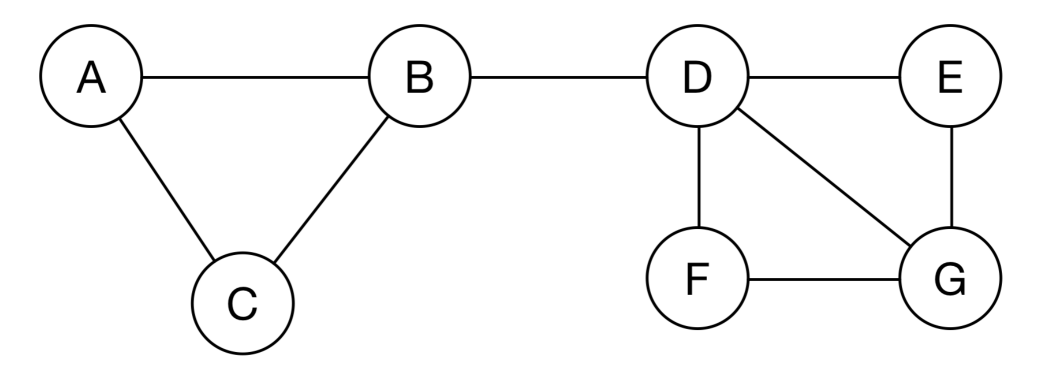
Formally,  $\operatorname{betweenness}(e)$ : fraction of node pairs (x,y) where shortest path crossess edge e

For each node x, use breadth-first-search to count number of shortest paths through each edge in graph

Sum result across nodes, and divide by two

# Girvan-Newman Algorithm

### Example



### Resources

Cross-language

igraph: http://igraph.org/

#### Resources

R

#### Workhorses:

- igraph
- Rgraphviz

#### Tidyverse (https://tidyverse.org):

- tidygraph
- ggraph

### Resources

### Python

- igraph
- networkx