Module IV - Basic Analysis

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Loading data from multiple sources The Python dict Running basic centralities Getting things out of NetworkX Basic visualization

Agenda for Module IV

Loading data from multiple sources

- Local network data files
- Building directly from the Internet

Loading data from multiple sources
The Python dict
Running basic centralities
Getting things out of NetworkX
Basic visualization

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Brief review of Python dictionaries

- Why is the dict so useful?
- ► How NetworkX utilizes it?

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- Degree, Closeness, Betweeness Eigenvector
- Calculating degree distribution
- Plotting statistics using matplotlib
- Calculating cliques, clustering and transitivity

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- Writing network data
- Saving network analysis statistics

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Basic visualization

- Review of NetworkX's plotting algorithms
- Adding analysis to visualization

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For those that are unfamiliar with working at the command-line, however, the process can be confusing



Let's try!

- We will load the edge list of Hartford drug users network
- Specify that the network be a directed graph, and the nodes be integers
- Use nx.info() to check that data has been loaded correctly

Local network data Building directly from the Interne

Loading the Hartford drug users network

Starting NetworkX and loading data

>>> hartford=nx.read_edgelist("../../data/hartford_drug.txt",create_using=nx.DiGraph(),nodetyp
>>> nx.info(hartford)

Name:

Type: DiGraph
Number of nodes: 212
Number of edges: 337
Average in degree: 1.5896
Average out degree: 1.5896

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What did we just do?

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- Specified path to Hartford drug users file

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- Used the create_using option to force NX to create as a directed graph

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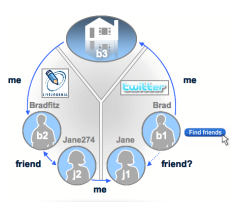
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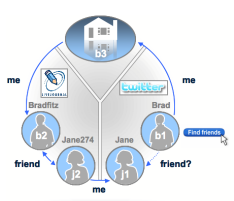
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- Used the read_edgelist function to load EL file
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- Used the info function to check that it all worked

Some formats may have more or less options, always check the documentations!

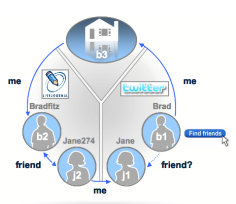


Building the social network among LiveJournal users

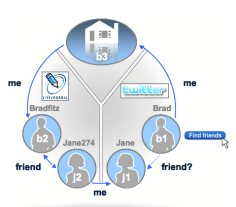


Perhaps the most powerful aspect of NetworkX is its ability to work in Python to generate networks from live-streaming data

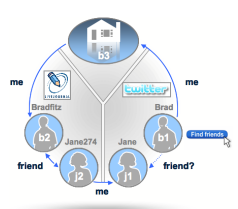
In Python, use NetworkX, cjson and a other standard scientific libraries to parse Google's SocialGraph data



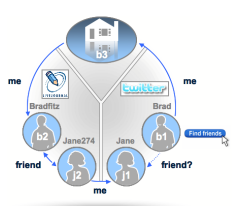
- ► In Python, use NetworkX, cjson and a other standard scientific libraries to parse Google's SocialGraph data
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- ► Through a process called "k-snowball searching" seed → friend → · · · → friend_k



- In Python, use NetworkX, cjson and a other standard scientific libraries to parse Google's SocialGraph data
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 - Seed: imichaeldotorg.livejournal.com
 - k = 3



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- Using a "seed" user, we will build out a network
- Through a process called "k-snowball searching" seed → friend → · · · → friend_k
 - Seed: imichaeldotorg.livejournal.comk = 3
- Note the low value of k

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from cjson import *
from urllib import *
from time import *
from scipy import array,unique
...
if __name__ == "__main__":
seed="imichaeldotorg"
seed_url="http://"+seed+".livejournal.com"
# 3.1 Scrape, parse and build seed's ego net
sg=get_sg(seed_url)
net,newnodes=create_egonet(sg)
nx.write_pajek(net,"../../data/"+seed+"_ego.net")
nx.info(net)
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def get_sg(seed_url):
    sgapi_url="http://socialgraph.apis.google.com/lookup?q="+seed_url+"&edo=1&edi=1&fme=1&pretty=0"
    try:
        furl=urlopen(sgapi_url)
        fr=furl.read()
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        return fr
except IOError:
        print "Could not connect to website"
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Loading the libraries and scraping egonet

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Local network data Building directly from the Internet

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Average in degree:

Average out degree:

The code, part 1

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```
Name: ['http://imichaeldotorg.livejourna
Type: DiGraph
Number of edges: 5
```

1.0

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Build egonet and snowball

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Creating the egonet
def create_egonet(s):
    trv:
        raw=decode(s)
        G=nx.DiGraph()
        pendants=[]
        n=raw['nodes']
        nk=n.kevs()
        G.name=str(nk)
        pendants=[]
        for a in range(0.len(nk)):
            for b in range(0,len(nk)):
                if al=h:
                    G.add_edge(nk[a].nk[b])
        for k in nk:
            eao=n[k]
            ego_out=ego['nodes_referenced']
            for o in ego_out:
                G.add_edge(k.o)
                pendants.append(o)
            ego_in=ego['nodes_referenced_by']
            for i in eqo_in:
                G.add_edge(i,k)
                pendants.append(i)
        pendants=array(pendants,dtype=str)
        pendants.flatten()
        pendants=unique(pendants)
        return G,pendants
   except DecodeError:
   except KeyError:
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Rolling the snowball

```
def snowball_round(G.seeds.mvspace=False):
    t0=time()
    if myspace:
        seeds=get_myspace_url(seeds)
    sb data=[]
    for s in range(0,len(seeds)):
        s_sq=qet_sq(seeds[s])
        new_eqo,pen=create_eqonet(s_sq)
        for p in pen:
                sb_data.append(p)
        if s<1:
            sb_net=nx.compose(G,new_eqo)
        else:
            sb_net=nx.compose(new_eqo,sb_net)
        del new_ego
        if s==round(len(seeds)*0.2):
            sb_net.name='20% complete'
            nx.info(sb net)
            print 'AT: '+strftime('%m/%d/%Y, %H:%M:%S', gmtime())
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    # More time keeping, probably a MUCH better way to do this
    sb_data=arrav(sb_data)
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Getting things out of NetworkX
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Running basic centralities
Getting things out of NetworkX
Basic visualization

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Basic visualization

Build the whole network

Step	Nodes	Edges	Mean Degree	Density
Seed	5	5	2.0	0.25
k = 2	75	115	3.0	0.02
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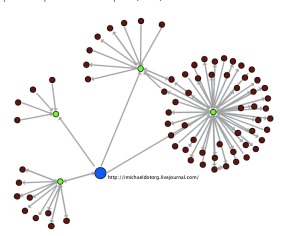
Our seed is abnormally isolated, with only four neighbors

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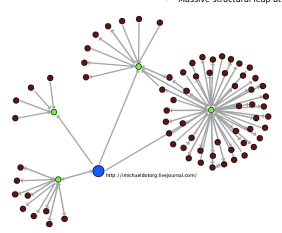
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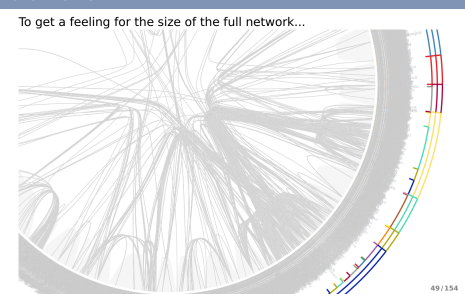
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The full network



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>> fruit_dict={"apple":1,"orange":[0.23,0.11],"banana":True }
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The Python dictionary is an extremely flexible and useful data structure, making it one of the primary advantages of Python over other languages

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Using dictionaries for network analysis

From the documentation...

networkx.closeness_centrality

 $closeness_centrality(G, v=None, weighted_edges=False)$

Compute closeness centrality for nodes.

Closeness centrality at a node is 1/average distance to all other

nodes.

Parameters: G: graph

A networkx graph

v : node, optional

Return only the value for node v.

weighted_edges : bool, optional

Consider the edge weights in determining the shortest paths. If False, all edge weights are considered equal.

Returns: nodes : dictionary

Dictionary of nodes with closeness centrality as the value.

NetworkX's metric's make extensive use of the dict type

In this case the key→value mapping is of the form: {node_label: metric}

Let's look at an example:

```
>> in_cen=nx.in_degree_centrality(hartford)
>> in_cen
{1: 0.014218009478672987, 2: 0.018957345971563982,...
90: 0.0047393364928909956, 293: 0.0}
```

We can see that node #90 has in-degree centrality 0.0047

But we can do so much more!

For our first analysis in NetworkX, we will do some basic network manipulation, then run multiple measures to find highest centrality nodes

 First, we will need to convert to an undirected network, and extract the main component

```
# Many of the centrality metrics require undirected graphs, so we will symmetriz
>> hartford_ud=hartford.to_undirected()
# The network also has many small components, but for
# this analysis we are interested in the largest
>> hartford_mc=hartford_main=nx.connected_component_subgraphs(hartford_ud)[0]
```

```
# Betweenness centrality
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To find the most central actors we will use Python's list comprehension technique to do basic data manipulation on our centrality dictionaries

```
def highest_centrality(cent_dict):
    """Returns node key with largest value from
    NX centrality dict"""
    # Create ordered tuple of centrality data
    cent_items=cent_dict.items()
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Now, just ask for the answer

Finding Most central actors

»> print("Actor "+str(highest_centrality(bet_cen))+" has the highest Betweenness centrality"
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def highest_centrality(cent_dict):
    """Returns node key with largest value from
NX centrality dict"""
    # Create ordered tuple of centrality data
    cent_items=cent_dict.items()
    # List comprehension!
    cent_items=[(b,a) for (a,b) in cent_items]
    # Sort in descending order
    cent_items.sort()
    cent_items.reverse()
    return cent_items[0][1]
```

```
List comprehension
```

```
Given a dict: d={1: 0.15, 2: 0.67}

d.items() → [(1,0.15),(2,0.67)]

d=[(b,a) for (a,b in d)] → [(0.15,1),(0.67,2)]
```

Now, just ask for the answer

Finding Most central actors

```
»> print("Actor "+str(highest_centrality(bet_cen))+" has the highest Betweenness centrality"
Actor 82 has the highest Betweenness centrality
```

To find the most central actors we will use Python's list comprehension technique to do basic data manipulation on our centrality dictionaries

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    # Create ordered tuple of centrality data
    cent_items=cent_dict.items()
                                                         d=[(b.a) \text{ for } (a.b \text{ in } d)] \rightarrow
    # List comprehension!
                                                         [(0.15,1),(0.67,2)]
    cent_items=[(b,a) for (a,b) in cent_items] Here, we use list
    # Sort in descending order
                                                    comprehension in order to
    cent_items.sort()
    cent items.reverse()
                                                    use Python's built-in sort
    return cent_items[0][1]
                                                    and reverse list functions
```

Now, just ask for the answer

Finding Most central actors

»> print("Actor "+str(highest_centrality(bet_cen))+" has the highest Betweenness centrality"
Actor 82 has the highest Betweenness centrality

To find the most central actors we will use Python's list comprehension technique to do basic data manipulation on our centrality dictionaries

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Calculating degree distribution

One of the most popular network level statistical description of a network is its degree distribution

► In NetworkX this is a simply one-line operation

Get list of degree rank frequency

- # Create a Barabasi-Albert network
- »> ba_net=barabasi_albert_graph(1000,2)
 # 6.1 Built-in function for degree distribution
- »> dh=degree_histogram(ba_net)

Calculating degree distribution

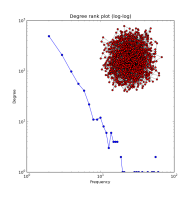
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>>> ba_net=barabasi_albert_graph(1000,2)
6.1 Built-in function for degree distribution
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- As we will see next, we can use matplotlib to take this data and create publication ready plots
- Ex. from http: //networkx.lanl.gov/examples/ drawing/degree_histogram.html



Often in network analysis we are interested in estimating the cohesiveness of a network, or the communities that exists within the structure

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Cliques

Maximal cliques are the largest complete subgraph containing a given point. There are several algorithms for finding cliques, including Bron & Kerbosch (1973), Tomita, Tanaka and Takahashi (2006), Cazals and Karande (2008)

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Clustering

For each node find the fraction of possible triangles that exist, $c_V = \frac{2T(v)}{deg(v)(deg(v)-1)}$, where T(v) is the number of triangles through node v.

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The fraction of all possible triangles which are in fact triangles. Or, $Trans = 3\left(\frac{T}{t}\right)$, where T = # of possible triangles and t = # of actual triads

Calculating basic community structure

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We will use clustering coefficients to identify community structure

```
# Calculate clustering coefficients of each node (return as dict)
clus=clustering(hartford_mc,with_labels=True)
# Get counts of nodes membership for each clustering coefficient, and clean up
unique_clus=list(unique(clus.values()))
clus_counts=zip(map(lambda c: clus.values().count(c),unique_clus),unique_clus)
clus_counts.sort()
clus_counts.reverse()
# Create a subgraph from nodes with most frequent clustering coefficient
mode clus sg=subgraph(hartford mc.[(a) for (a.b) in clus.items() if b==clus counts[0][1]])
```

Calculating clustering coefficients

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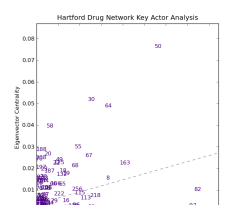
Later, we'll learn how to create a network visualization like the one 79/154

Recall Python's scientific computing trinity: NumPy, SciPy and matplotlib

While NumPy and SciPy do most of the behind the scenes work, you will interact with matplotlib frequently for when doing network analysis

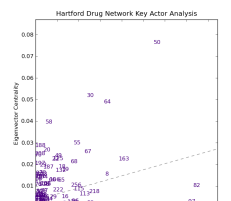
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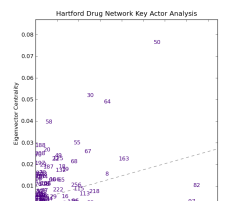
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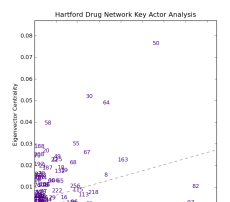


We will need to create a function that takes two centrality dict and generates this plot

Create a matplotlib figure

Recall Python's scientific computing trinity: NumPy, SciPy and matplotlib

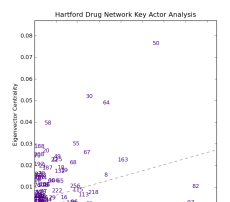
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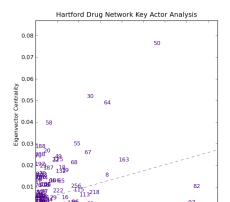
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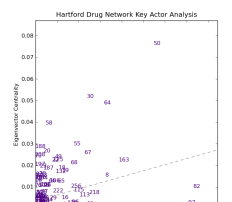
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- 4. Add axis and title labels

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- Create a matplotlib figure
- Plot each node label as a point
- 3. Add a "best fit" line
- 4. Add axis and title labels
- 5. Save figure as a PNG file 87/154

Basic visualization

Creating a key actor plot in matplotlib

The centrality $_{-}$ scatter function, part one

```
def centrality_scatter(met_dict1,met_dict2,path="",ylab="",xlab="",title="",reg=False):
    # Create figure and drawing axis
    fig=P.figure(figsize=(7,7))
    axl=fig.add_subplot(111)
    # Create items so actors can be sorted properly
    met_itemsl=met_dict1.items()
    met_items2=met_dict2.items()
    met_items2.sort()
    # Grab data
    xdata=[(b) for (a,b) in met_items1]
    ydata=[(b) for (a,b) in met_items2]
    # Add each actor to the plot by ID
    for p in xrange(len(met_items1)):
        axl.text(x=xdata[p],y=ydata[p],s=str(met_items1[p][0]),color="indigo")
```

```
The centrality_scatter function, part one
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   met_items1.sort()
   met items2.sort()
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   vdata=[(b) for (a,b) in met_items2]
   # Add each actor to the plot by ID
    for p in xrange(len(met_items1)):
        ax1.text(x=xdata[p].v=vdata[p].s=str(met items1[p][0]).color="indigo")
```

Create a canvas to draw on

```
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def centrality_scatter(met_dict1,met_dict2,path="",ylab="",xlab="",title="",reg=False):
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- Create a canvas to draw on
- manipulate and store centrality data

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```

- Create a canvas to draw on
- manipulate and store centrality data
- Add points to plot as node labels

The centrality_scatter function, part one

```
def centrality_scatter(met_dict1,met_dict2,path="",ylab="",xlab="",title="",reg=False):
    # If adding a best fit line, we will use NumPy to calculate the points.
    if rea:
        #Function returns v-intercept and slope. So, we create a function to
        # draw LOBF from this data
        slope, yint=polyfit(xdata, ydata, 1)
        xline=P.xticks()[0]
        yline=map(lambda x: slope*x+yint,xline)
        # Add line
        ax1.plot(xline,yline,ls='-',color='grey')
    # Set new x- and y-axis limits to data
    P.xlim((0.0,max(xdata)+(.15*max(xdata))))
                                                # Give a little buffer
    P.vlim((0.0.max(vdata)+(.15*max(vdata))))
    # Add labels
    ax1.set_title(title)
    ax1.set_xlabel(xlab)
    ax1.set_vlabel(vlab)
    # Save figure
    P.savefig(path,dpi=100)
```

```
The centrality_scatter function, part one
def centrality_scatter(met_dict1,met_dict2,path="",ylab="",xlab="",title="",reg=False):
   # If adding a best fit line, we will use NumPy to calculate the points.
   if rea:
        # Function returns y-intercept and slope. So, we create a function to
        # draw LOBF from this data
        slope, yint=polyfit(xdata, ydata, 1)
        xline=P.xticks()[0]
       vline=map(lambda x: slope*x+yint,xline)
        # Add line
        ax1.plot(xline.vline.ls='-'.color='grev')
   # Set new x- and y-axis limits to data
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Add a best fit line

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   # Save figure
```

- Add a best fit line
- Resize figure to fit data

P.savefig(path,dpi=100)

The centrality_scatter function, part one

Creating a key actor plot in matplotlib

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    # Set new x- and y-axis limits to data
    P.xlim((0.0,max(xdata)+(.15*max(xdata))))
                                                # Give a little buffer
    P.vlim((0.0.max(vdata)+(.15*max(vdata))))
    # Add labels
    ax1.set title(title)
```

- Add a best fit line
- Resize figure to fit data

ax1.set_xlabel(xlab)
ax1.set_ylabel(ylab)
Save figure
P.savefig(path,dpi=100)

Add labels, and save the figure as a PNG file

As powerful as NetworkX and the complementing scientific computing packages in Python are, it may often be useful or necessary to output your data for additional analysis

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Suite of tools lacks your specific need

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- Suite of tools lacks your specific need
- Require alternate visualization

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In most cases this will entail either exporting the raw network data, or metrics from some network analysis

1. NetworkX can write out network data in as many formats as it can read them, and the process is equally straightforward

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- When you want to export metrics we can use Python's built-in XML and CSV libraries

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- NetworkX can write out network data in as many formats as it can read them, and the process is equally straightforward
- When you want to export metrics we can use Python's built-in XML and CSV libraries
- 3. Depending on your needs you may prefer one, the other or both Next, we will review how to save data in different formats and export metrics to a CSV file using the Hartford drug net data

```
NX syntax for writing a network file

>>> nx.write_format(G, "path/to/file.txt", ...options...)

↑ ↑ ↑ ↑

NX function, net variable File to be written Nodes/edge data, etc.
```

```
NX syntax for writing a network file

>>> nx.write_format(G, "path/to/file.txt", ....options...)

↑ ↑ ↑ ↑

NX function, net variable File to be written Nodes/edge data, etc.
```

Saving network data in different formats

The syntax for exporting network data follows exactly the syntax for loading it



Let's try!

- Output the Hartford drug net data as an adjacency list
- Add metric data to each node of the network
- Output new network in Pajek format with node attributes

Saving network data and adding node attributes

As shown, this is a simple one line operation

Output Hartford drug net data as an adjacency list

```
\verb|nx.write_adjlist(hartford_mc,".../.../data/hartford_mc_adj.txt")| \\
```

Next, we will add the Eigenvector centrality of each node to the graph object

Adding node attributes

```
def add_metric(G,met_dict):
    """Adds metric data to G from a dictionary keyed by node labels"""
    if(G.nodes().sort()==met_dict.keys().sort()):
        for i in met_dict.keys():
            G.add_node(i,metric=met_dict[i])
        return G
    else:
        raise ValueError("Node labels do not match")
```

Saving network data and adding node attributes

As shown, this is a simple one line operation

Output Hartford drug net data as an adjacency list

```
nx.write_adjlist(hartford_mc,"../../data/hartford_mc_adj.txt")
```

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            G.add_node(i,metric=met_dict[i])
        return G
    else:
        raise ValueError("Node labels do not match")
```

Quick error checking

Saving network data and adding node attributes

As shown, this is a simple one line operation

Output Hartford drug net data as an adjacency list

```
nx.write_adjlist(hartford_mc,"../../data/hartford_mc_adj.txt")
```

Next, we will add the Eigenvector centrality of each node to the graph object

Adding node attributes

```
def add_metric(G,met_dict):
    """Adds metric data to G from a dictionary keyed by node labels"""
    if(G.nodes().sort()==met_dict.keys().sort()):
        for i in met_dict.keys():
             G.add_node(i,metric=met_dict[i])
        return G
    else:
        raise ValueError("Node labels do not match")
```

- Quick error checking
- Add node attribute as "metric"

Python has powerful built-in tools for reading and writing standard data formats

```
import csv
def csv exporter(data dict.path):
    """Takes a dict of centralities keyed by column headers and exports
    data as a CSV file"""
    # Create column header list
    col_headers=["Actor"]
    col_headers.extend(data_dict.kevs())
    # Create CSV writer and write column headers
    writer=csv.DictWriter(open(path, "w"), fieldnames=col_headers)
    writer.writerow(dict((h,h) for h in col_headers))
    # Write each row of data
    for j in data_dict[col_headers[1]].keys():
        # Create a new dict for each row
        row=dict.fromkeys(col_headers)
        row["Actor"]=i
        for k in data_dict.keys():
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The results of CSV export

We can now open the CSV file in our favorite spreadsheet program

- Perform traditional data exploration
- Load into other analytics platforms for additional analysis (e.g., R)
- Store for latter use

~	A	В	С	D
1	Actor	Closeness	Betweeness	Eigenvector
2	1	0.12467532	0.0072576	0.00025176
3	2	0.12475634	0.01767427	0.00025964
4	3	0.12565445	0.05687441	0.00023185
5	4	0.10223642	0.03108639	1.44E-05
6	5	0.1443609	0	0.00313152
7	6	0.09943035	0.01041667	1.49E-07
8	7	0.11340815	0.04362093	6.78E-05
9	8	0.20512821	0.16354003	0.01471888
10	9	0.11267606	0.00741624	0.0001101
11	10	0.13983977	0.05258239	0.00095456
12	11	0.1703638	0.01250999	0.0032333
13	13	0.13892909	0	1.79E-05
14	14	0.17219731	0.11848775	0.00029737
15	15	0.13521127	0.00079897	2.11E-05
16	16	0.15907208	0.06203647	0.00432838

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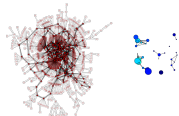
NetworkX was designed as a data manipulation and analysis tool, and therefore is not meant as a visualization platform

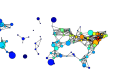
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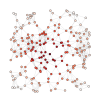
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NetworkX was designed as a data manipulation and analysis tool, and therefore is not meant as a visualization platform

▶ It is, however, still capable of making very nice visualization







- ► The random layout places nodes in...random positions
- The circular layout places nodes in...a circle





```
# Use subplots to draw random and circular layouts
# of drug net side-by-side
fig1=P.figure(figsize=(9,4))
fig1.add_subplot(121)
nx.draw_random(hartford_mc,with_labels=False,node_size=60)
fig1.add_subplot(122)
nx.draw_circular(hartford_mc,with_labels=False,node_size=60)
P.savefig("../../images/networks/rand_circ.png")
```

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Visualization algorithms in NetworkX - Spring & Spectral

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Visualization algorithms in NetworkX - Spring & Spectral

More commonly used visualization techniques include the spring and spectral layouts

- The spring layout is a version of the Fruchterman-Reingold force-directed algorithm, which attempts to minimize overlapping edges
- The spectral layout finds node position using the eigenvectors of the graph Laplacian, which is useful for quickly visualizing structural clustering



The shell layout draws nodes as concentric circles

- Two dimensional extension of the circle layout
- We may have some reason to isolate certain nodes

```
P.figure(figsize=(8,8))
# Find actors in 25th percentile
max_eig=max([(b) for (a,b) in eig_cen.items()])
s1=[(a) for (a,b) in eig_cen.items() if b>=.25*max_eig]
s2=hartford_mc.nodes()
# setdiffId is a very useful NumPy function!
s2=list(setdiffId(s2,s1))
shells=[s1,s2]
# Calculate position and draw
shell_pos=shell_layout(hartford_mc,shells)
draw_networkx(hartford_mc,shell_pos,with_labels=False,node_size=60)
P.savefig("./../images/networks/shell.png")
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s2=hartford_mc.nodes()
# setdiff1d is a very useful NumPy function!
s2=list(setdiff1d(s2,s1))
shells=[s1,s2]
# Calculate position and draw
shell_pos=shell_layout(hartford_mc,shells)
draw_networkx(hartford_mc,shell_pos,with_labels=False,node_size=6
P.savefig("../../images/networks/shell.png")
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25th percentile Eigenvector centrality actors

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Beyond layout, we may also want to add analytical data to our visualization

Changing node and edge size and colors

NetworkX allows you to alter the size, color and shape of the nodes and edges in any visualization

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In our final exercise, we will add the following analysis to the Hartford drug network

- Node size by Eigenvector centrality
- Intensity of node color by betweenness centrality
- Edge thickness by edge betweenness

```
# Adding analysis to visualization
P.figure(figsize=(15,15))
P.subplot(111,axisbg="lightgrey")
spring_pos=nx.spring_layout(hartford_mc,iterations=1000)
# Use betweeneess centrality for node color intensity
bet_color=bet_cen.items()
bet_color.sort()
bet_color=[(b) for (a.b) in bet_color]
# Use Eigenvector centrality to set node size
eig_size=eig_cen.items()
eia_size.sort()
eig\_size=[((b)*2000)+20 \text{ for } (a,b) \text{ in } eig\_size]
# Use matplotlib's colormap for node intensity
draw_networkx(hartford_mc,spring_pos,node_color=bet_color,...
    ...cmap=P.cm.Greens,node_size=eig_size,with_labels=False)
P.savefig("../../images/networks/analysis.png")
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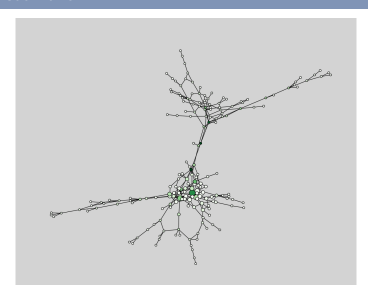
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Final visualization



Basic Analysis

How to load local data, and an example of building networks from data streamed directly from the Internet

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- ▶ A brief review of the Python dict data type

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Questions?