## project 2

### March 10, 2023

In this project, you will be working with the protein interaction network (PIN) of Saccharomyces cerevisiae from the BioGRID Multi-Validated (MV) Datasets. The nodes and edges of this PIN denote proteins and their physical interactions, respectively. These interactions have been identified from large-scale protein interaction experiments such as yeast two-hybrid screenings and affinity purification with mass spectrometry. Additionally, they pass a specific set of criteria as defined here (BioGRID MV Datasets). See the following link to familiarize yourself with the contents of this BioGRID file. You will be analyzing the yeast PIN using k-core analysis, and further use the network as a scaffold for incorporating gene expression data, allowing you to perform s-core analysis using pairwise gene expression correlations as weights.

### 0.0.1 1.1 Yeast protein interaction network

Download the file containing all multi-validated protein-protein interactions from BioGRID found on Blackboard BioGRID\_PIN.txt. Create a network with NetworkX using the systematic names to set source and target nodes. These corresponds to the columns named Systematic Name Interactor A and Systematic Name Interactor B. Remove all self-edges from the network.

```
[]: # Imports
import pandas as pd
import numpy as np
import networkx as nx
from pyvis.network import Network
```

```
[]: import sys
import os
path_ = sys.path[0].split("/")
os.chdir("/".join(path_[:-1]))
os.getcwd()
from SysBio_fun import *
os.chdir("/".join(path_))
```

```
[]: # Read file:
df = pd.read_table("BioGRID_PIN.txt", sep="\t", low_memory=False)

# Define source and target column:
source = "Official Symbol Interactor A"
target = "Official Symbol Interactor B"
# Rename columns:
```

```
df.rename(columns={source:"source",
                   target:"target"},
                   inplace=True)
# Drop rows where column "source" value is identical to column "target" value.
# I.e., drop self loops:
# **NB!** This the number of rows left in the df may vary depending on which
# column-pairs were treated as source-target. E.g. source:"-" and target:"-"
# will be removed, even though they may have different Entrez ID.
df = df[df["source"] != df["target"]]
# Generate the graph:
# **NB!** I treat the graph as multi directed, i.e. each row in the df
# amounts to a single edge in the network!
PIN = nx.from_pandas_edgelist(df, edge_attr=True, create_using=nx.
 →MultiDiGraph())
# Make a copy
PIN_copy = PIN.copy()
```

(i) How large is the network (i.e. number of nodes and edges), and what is the average node degree?

The number of nodes are: 20550
The number of edges are: 245288
The average node degree is: 23.872311435523116

(ii) Look up *S. cerevisiae* S288C in the NCBI Taxonomy Database here and take note of the taxonomy ID. Using the Organism Interactor A and Organism Interactor B edge attributes in the network, use this taxonomy ID to filter out all non-*S. cerevisiae* S288C nodes from the network. Also, remove all self-loops and isolated nodes. How many nodes and edges remain? Create a plot of its degree distribution. How would you characterize this network and its degree distribution?

```
[]: PIN = PIN_copy.copy()
```

```
# The taxonomic ID of S. cerevisiae S288C is: 559292
# Generate a list of edges that does not fit:
r1 = [edge for edge,tax \
      in nx.get_edge_attributes(PIN, "Organism Interactor A").items() \
          if tax != 559292] # The edge information is a tuple and the key
r2 = [edge for edge,tax \
      in nx.get_edge_attributes(PIN, "Organism Interactor B").items() \
        if tax != 559292]
rem = set(r1+r2) # set() is used to remove potential duplicates
# Use the list to remove all edges
PIN.remove_edges_from(rem)
# In pandas:
# len(df[(df["Organism Interactor A"] == 559292)) & (df["Organism Interactor B"]_{\square}
 →== 559292)])
# Remove nodes with a node degree of O (i.e. isolated nodes):
PIN.remove nodes from( # Remove nodes from list ...
    [nodes for nodes,degree \
     in dict(PIN.degree()).items() if degree == 0] # nodes with degree 0
)
N = PIN.number_of_nodes() # New number of nodes
E = PIN.number_of_edges() # New number of edges
print(f"""
The number of nodes is now {N},
while number of edges is {E}
""")
```

The number of nodes is now 4104, while number of edges is 68704

(iii) Compare the yeast PIN with Barabási–Albert (BA) and Erdős–Rényi (ER) networks with approximately the same number of nodes and edges using a few selected network measurements and the degree distributions. Do these models do a good job in describing the yeast PIN? Discuss why/why not.

```
[0]
     ).copy())
     n = 1_comp.number_of_nodes()
     e = 1_comp.number_of_edges()
     ER = connected_ER(n, e)
     BA = connected_BA(n, round(((e*2)/n)))
[]: | #type(PIN.subgraph(nx.connected_components(PIN.to_undirected())[0]))
     type(
         nx.Graph(PIN.subgraph(
             sorted(
                 nx.connected_components(
                     nx.Graph(PIN.copy())
                 ), reverse=True)[0]
             ))
         )
     type(PIN)
     \# PIN2 = nx.Graph(PIN.copy())
     # print(type(PIN2))
     # print(PIN == PIN2)
     # PIN2 = PIN2.to undirected()
     # print(type(PIN2))
[]: networkx.classes.multidigraph.MultiDiGraph
[]: graph_info(l_comp)
     graph_info(ER)
     graph_info(BA)
    Number of nodes: 4022
    Number of edges: 16700
    Is connected:
                     True
    Number of nodes: 4022
    Number of edges: 16700
    Is connected:
                     True
    Number of nodes: 4022
    Number of edges: 32112
    Is connected:
                     True
[]: def degree_distribution(graph):
         Function that returns a scatter plot of the degree
         distribution of a graph.
         N = graph.number_of_nodes() # Total number of nodes
```

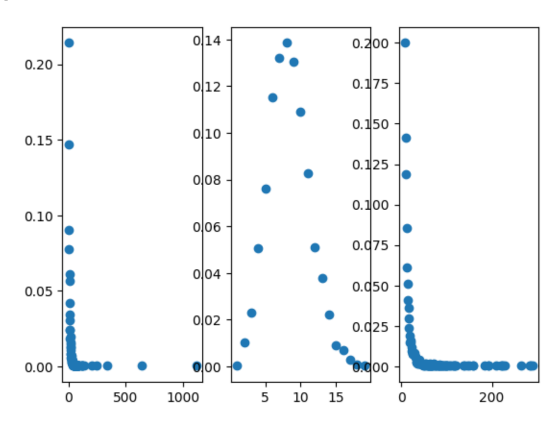
```
y = np.array(nx.degree_histogram(graph))/N # All occurrences of "n"_
degree, divided by total number of nodes
x = np.arange(len(y))[y != 0] # x values
y = y[y != 0] # Remove the values = 0 from the array

return x, y

x_l_comp, y_l_comp = degree_distribution(l_comp)
x_ER, y_ER = degree_distribution(ER)
x_BA, y_BA = degree_distribution(BA)

figure, axis = plt.subplots(1, 3)
axis[0].scatter(x_l_comp, y_l_comp)
axis[1].scatter(x_ER, y_ER)
axis[2].scatter(x_BA, y_BA)
```

## []: <matplotlib.collections.PathCollection at 0x7c30ebb3e700>



# []: #plt.scatter(nx.clustering(BA).keys(), nx.clustering(BA).values())

(iv) You will now start peeling away layers of the yeast PIN by k-core analysis. In your own words,

describe/define k-core analysis, and explain how it works on a network. What network does the 1-core correspond to?

k-core analysis is a method to

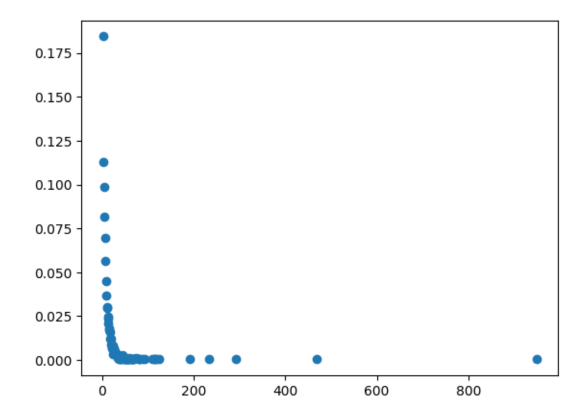
(v) Calculate the 2-core of the yeast PIN. How large is this network? Plot and characterize its degree distribution and compare it to the original yeast PIN.

```
[]: l_comp_2_core = nx.k_core(l_comp, k=2)
graph_info(l_comp_2_core)

x, y = degree_distribution(l_comp_2_core)
plt.scatter(x,y)
```

Number of nodes: 3102 Number of edges: 15780 Is connected: True

[]: <matplotlib.collections.PathCollection at 0x7c30ee367190>



(vi) Find the innermost k-core (the last k before the network is empty). How many edges and nodes are there in the innermost k-core? What is the absolute maximal amount of edges that there could be between this number of nodes, and how does the innermost core compare to that?

```
[]: current_k = 0
g = l_comp.copy()
while nx.number_of_nodes(g) > 0:
    current_k += 1
    g = nx.k_core(g, k=current_k)
innermost_k_core = current_k-1

print(f"The innermost k-core is: {innermost_k_core}")

graph_info(nx.k_core(l_comp.copy(), k=innermost_k_core))
```

The innermost k-core is: 18

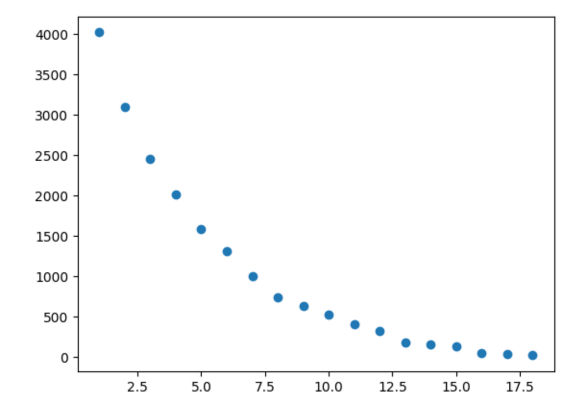
Number of nodes: 19 Number of edges: 171 Is connected: True

(vii) Create a plot of the number of nodes and the number of edges in a k-core against k. How would you characterize this plot? What does it tell you?

```
y = [nx.number_of_nodes(nx.k_core(l_comp.copy(), k=k_)) for k_ in range(1,
innermost_k_core+1)]
x = range(1, innermost_k_core+1)
```

```
[]: plt.scatter(x, y)
```

[]: <matplotlib.collections.PathCollection at 0x7c30e126ea60>



(viii) Generate the second innermost k-core and visualize it using pyvis. Describe the network.

```
[]: # def show_html(graph, name="nx", show=True, size="small"):
     #
     #
           Generate and display the graph through pyvis
     #
     #
           if size == "small":
     #
               nt = Network("500px", "500px")
     #
           else:
     #
               nt = Network("1080px", "1920px")
     #
           nt.from_nx(graph)
     #
           if show == True:
     #
               nt.show(f"{name}.html")
     s = nx.k_core(l_comp.copy(), k=innermost_k_core-1)
     show html(s, name=f"{innermost k core-1}-core network", show=True, size="large")
```

(ix) We will now load gene expression data onto our yeast PIN. Go to this url containing the gene expression data set from  $Transcription\ profiling\ by\ array\ of\ yeast\ to\ investigate\ expression\ of\ the\ beta-subunit\ of\ Snf1\ kinase.$  For those interested, you can find detailed information about the dataset in the original publication here. Download normalized\_expressions.tsv from Blackboard and import the list of gene names, as well as corresponding expression data from 24 experimental conditions. For all pairwise gene combinations, calculate the Pearson correlation coefficients and associated p-values for the N genes present in the yeast PIN (both as  $N \times N$  Numpy arrays). Out of the gene pairs with positive correlation values, how many of them have a significance value below 0.0001?

```
[]: norm_ex = pd.read_table("normalized_expressions.tsv", sep="\t")
norm_ex.head(10)
```

```
[]:
        Gene ID
                 GSM517188
                            GSM517189
                                        GSM517170
                                                   GSM517171
                                                               GSM517180
                                                                          GSM517186
     O YALOO1C
                 10.862551
                            10.787718
                                        10.609314
                                                   10.533807
                                                               10.683814
                                                                          10.884063
      YALOO2W
                 10.318229
                            10.376402
                                        10.307895
                                                   10.231124
                                                               10.217388
                                                                          10.289048
     1
     2
      YALOO3W
                                        14.038039
                                                   14.042464
                 14.116945
                            14.110254
                                                               14.116978
                                                                          14.100915
     3
       YAL005C
                 14.372108
                            14.419899
                                        14.327958
                                                   14.283262
                                                               14.436713
                                                                          14.418395
       YALO07C
                 11.717200
                             11.861313
                                        11.594489
                                                   11.750768
                                                               11.574141
     4
                                                                          11.824636
     5
      YAL008W
                 11.275044
                            11.248086
                                        11.370819
                                                   11.108830
                                                               11.510864
                                                                          11.330115
     6
       YALOO9W
                  9.971481
                             9.935119
                                         9.988509
                                                    9.993210
                                                                9.915578
                                                                           9.936548
     7
       YAL010C
                  9.788592
                              9.882031
                                         9.969160
                                                    9.935823
                                                                9.869899
                                                                           9.830950
     8
       YALO11W
                  9.434274
                              9.396159
                                         9.500599
                                                    9.525675
                                                                9.558400
                                                                           9.427803
        YAL012W
                             13.565094
                                        13.187785
                                                   13.172306
                 13.515144
                                                               13.216801
                                                                          13.380632
```

GSM517187 GSM517183 GSM517172 ... GSM517184 GSM517168 GSM517181 \

```
10.991316
               10.793569
                           10.641122
                                          10.776250
                                                      10.829294
                                                                  10.706164
0
1
   10.571356
               10.192751
                           10.307293
                                          10.166869
                                                      10.360606
                                                                  10.282979
2
   14.101845
               14.105961
                           13.999617
                                          14.085988
                                                      14.127210
                                                                  14.048146
3
   14.354286
               14.369767
                           14.285516
                                          14.243826
                                                      14.349704
                                                                  14.361417
   11.856330
4
               11.650583
                           11.548949
                                          11.612014
                                                      11.475502
                                                                  11.608367
                                                      11.554264
   11.461117
                           11.246455
                                          11.431879
5
               11.363152
                                                                  11.363908
6
   10.152794
                9.824603
                           10.018877
                                           9.809905
                                                       9.740123
                                                                   9.854044
7
   10.069362
                9.904191
                            9.780692
                                           9.966418
                                                       9.828681
                                                                   9.752460
    9.519610
                                           9.538738
8
                9.578430
                            9.450371
                                                       9.512650
                                                                   9.614558
9
   13.419438
               13.374047
                           13.050084
                                          13.449695
                                                      13.217452
                                                                  13.372705
   GSM517185
               GSM517167
                           GSM517169
                                       GSM517190
                                                   GSM517173
                                                               GSM517174
                                                                           GSM517175
0
   11.036464
               10.876691
                           10.778201
                                       10.860265
                                                   10.581020
                                                               10.698511
                                                                           10.561507
1
   10.015871
               10.210560
                           10.433530
                                       10.334844
                                                   10.246341
                                                               10.279370
                                                                           10.347147
2
   14.031750
               14.115299
                                       14.117709
                                                               14.010271
                                                                           13.993024
                           14.085665
                                                   13.981556
3
   14.238842
               14.298123
                           14.325895
                                       14.371525
                                                   14.310817
                                                               14.346674
                                                                           14.345590
4
   11.670285
               11.479741
                           11.490664
                                       11.975316
                                                   11.606178
                                                               11.605336
                                                                           11.535457
5
   11.351615
               11.426774
                           11.533108
                                       11.299099
                                                   11.358341
                                                               11.310492
                                                                           11.474645
6
   10.033609
                9.888328
                            9.792665
                                       10.048278
                                                    9.966829
                                                               10.066921
                                                                            9.871540
7
   10.116446
                            9.906184
                9.899894
                                        9.823420
                                                    9.742537
                                                                9.947832
                                                                            9.807137
8
    9.257099
                9.566624
                            9.492080
                                        9.458073
                                                    9.336601
                                                                9.489274
                                                                            9.422245
9
   13.160807
               13.213176
                           13.258713
                                       13.537055
                                                   13.238104
                                                               13.163476
                                                                           13.089239
```

#### [10 rows x 25 columns]

- (x) Download s\_core.py. This is similar to the NetworkX function k\_core, but uses node strength instead of node degree. Strength is defined as the sum of the weights of all the links attached to a node (i.e. Person correlations). With negative weights, the absolute values of the weights are used to calculate the strength. In this case, however, we are only looking at positively correlated nodes. Do you think it would make sense to include negative correlations in this particular case? Discuss.
- (xi) Calculate the s-core with s = 2.5, 3, 3.5, and 4, and characterize their degree distributions. Discuss how the distribution changes for increasing values of s.

#### [ ]: from s\_core import \*

(xii) Find an integer value for s that gives as good match as possible relative to the second innermost k-core in terms of number of distinct modules and number of nodes. Visualize this s-core with pyvis and describe the network.

### []:

- (xiii) Using the full yeast PIN network (network from (ii)), extract the nodes that are either part of the second innermost k-core, the s-core in (xii), or part of both cores. How many nodes and edges are there in this new network? Create a visualization of this subnetwork using pyvis, and color the nodes according to the following classifications:
  - Red if it was only found in the second innermost k-core.
  - Green if it was only found in the s-core.

• Blue if it was found in both.

Describe the network and discuss what you observe. Is it a single connected component? What do the differences between the s- and k-cores tell you?

```
[]: path_ = "/".join(sys.path[0].split("/")[:-1])
    #os.chdir("/".join(path_[:-1]))
    #file_path = '~/TBT4165/SysBio_fun.py'
    #print(path_+"/SysBio_fun.py")
    with open(path_+"/SysBio_fun.py") as file:
        print(file.read())
    import networkx as nx
    import matplotlib.pyplot as plt
    import numpy as np
    from pyvis.network import Network
    This is a module created to hold
       all custom functions used in multiple
                                             #
       project throughout the TBT4165 course
    def connected_ER(N, E):
       N : int, Number of nodes
       E : int, Number of edges
       g = nx.gnm_random_graph(N, E)
       while nx.is_connected(g) == False:
           g = nx.gnm_random_graph(N, E)
       return g
    def connected_BA(N, m):
       11 11 11
       N : int, Number of nodes
       m : int, Number of edges to attach from a new node to existing nodes
       g = nx.barabasi_albert_graph(n=N, m=m)
       while nx.is_connected(g) == False:
           g = nx.barabasi_albert_graph(n=N, m=m)
       return g
    def get_largest_component(g, is_undirected=True):
       if is_undirected:
           1_comp = g.subgraph(
```

```
sorted(
                nx.connected_components(
                    nx.to_undirected(g.copy())
                    ),
                reverse=True
            [0]
        ).copy()
    else:
        l_comp = g.subgraph(
            sorted(
                nx.connected_components(g),
                reverse=True
            [0]
        ).copy()
    return l_comp
def graph_info(g):
    print(f"\nNumber of nodes: {g.number_of_nodes()}")
    print(f"Number of edges: {g.number_of_edges()}")
                                 {nx.is_connected(g)}")
        print(f"Is connected:
    except:
        print("""\
The graph is not undirected. Therefore .is_connected() does not work.
def show_html(graph, name="nx", show=True, size="small"):
    Generate and display the graph through pyvis
    if size == "small":
       nt = Network("500px", "500px")
    else:
        nt = Network("1080px", "1920px")
   nt.from_nx(graph)
    if show == True:
        nt.show(f"{name}.html")
def nice_plot(title, xlab, ylab, show=True):
    Function to generate title and axis labels to a plot.
    The argument "show" is by default True.
```

```
11 11 11
   plt.title(title)
   plt.xlabel(xlab)
   plt.ylabel(ylab)
    if (show == True):
        plt.show()
def plot_degree_distribution(graph):
    Function that returns a scatter plot of the degree
    distribution of a graph.
    N = graph.number_of_nodes() # Total number of nodes
    y = np.array(nx.degree_histogram(graph))/N # All occurrences of "n" degree,
divided by total number of nodes
    x = np.arange(len(y))[y != 0] # x values
   y = y[y != 0] # Remove the values = 0 from the array
    return [x, y]
    #return plt.scatter(x,y)
```