# Evolutionary Time and Protein-Protein Interaction Networks

Preliminary Analysis

STA 596: Practical Data Science
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## I. Introduction

Proteins control all biological systems in a cell and, through various interactions with each other, enable cells to complete tasks such as: enzyme activation, gene regulation, and intercellular communication. Protein interactions can be modeled by an undirected network of protein-protein interactions (a PPI network, or PPIN), with nodes representing proteins and edges representing interactions (see Figure 1). The complete set of such interactions for a species is called the protein interactome. When interaction relationships between proteins break, possibly due to environmental factors or random mutations, such breakage can cause disease and death, of the cell and of the organism. We hypothesize that the evolutionary time of a species, which is defined as the total branch length from the root to the leaf representing that species in the tree of life, is directly related to network statistics which describe the topological stability of the species' protein interactome.

#### II. Background

Advances in proteomics allow researchers to study the protein interactome, but limitations of experimental methods in practice prevent PPI networks from being comprehensive and free of noise. We regard extant PPIN data, including the data used in this project, as a noisy sample from the true protein interactome. For example, the yeast-two-hybrid method for mapping protein interactomes was first developed in 1989 by Fields and Song using *Saccharomyces cerevisiae* as a biological model. The accuracy of this experimental method is estimated to be less than 10 percent. Consequently, the population being studied is the true protein interactome for each species and the variables of interest are the network statistics.

Understanding how protein interactomes evolve and developing methods for analyzing PPI networks is a central goal of evolutionary systems biology (Maddamsetti (2021)). In a paper by Rohan Maddamsetti they provided evidence that protein interactomes in E-Coli appear to show a generational increase in network resilience. Marinka Zitnik (Zitnik et al. (2019)) defined network resilience as the measure of how quickly a network breaks down as edges between nodes are randomly removed. A resilience rating of 1 implies that the network is most resilient while a rating of 0 implies a complete loss of connectivity in the PPI network. The current research identified a positive linear relationship between the resilience of an interactome and evolutionary time of the species.

#### III. Methods

## III.I. Acquisition of Data

Our dataset comes from the Stanford Network Analysis Platform (or SNAP). This data was collected using the Search Tool for the Retrieval of Interacting Genes/Proteins (or STRING), from the European Molecular Biology Laboratory, EMBL and is organized in multiple text files, joined by species ID. It comprises taxonomy information, an edge set for each interactome, and a numerical variable for evolutionary time of the species.

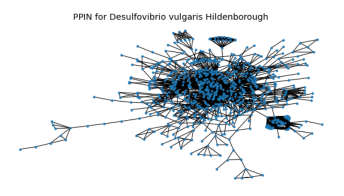


Figure 1: An example PPIN

In order to apply common network analysis techniques, we restructured the PPIN data as a series of adjacency matrices. To reduce the computational burden during initial algorithm development, we selected an arbitrary subset of 100 species of Proteobacteria, a major phylum which includes a wide variety of pathogenic genera such as Salmonella. For network statistics that only pertain to connected graphs, we use the largest connected subgraph (or LCSG).

To extract pertinent information about net-

work stability from each network we use the concept of Exponential Random Graph Models. The basic assumption of these models is that the structure in an observed network can be explained by a vector of sufficient statistics which are a function of the observed network. To construct our models, we first need to find the vector of sufficient statistics for each network.

Using functions of NetworkX, a Python library for network analysis, we calculated statistics for each PPIN, including (but not limited to): average degree centrality, density, number of triangles, modularity, and maximal clique statistics for the complete network and for the LCSG. Where network metrics are derived from the LCSG, the metric name is prefixed with "LCSG".

Average centrality for a network describes the average number of edges for all nodes. In other words, this statistic describes the average number of connections for all nodes. If a network has a high average degree centrality then we interpret this network as being dense with respect to the number of nodes in the network.

Network **Density** describes the portion of the potential connections in a network that are actual connections. Suppose there is a PPIN which has high density and another which has low density in terms of the interactions among the proteins. We could posit that one issue might be that information doesn't transmit very efficiently across the low density network because it has to go from protein to protein, rather than diffusing from one protein rapidly to all the others. Moreover, if proteins are taken out of a network with low density, the network may suffer due to the structural holes caused by the deletion of these nodes.

The **Number of Triangles** statistic counts the number of triangles in the network—we sum up the number of triangles each node is a part of then divide this number by three. A triangle is a set of three nodes where each node has a relationship to the other two—this is sometimes referred to as a 3-clique. Networks that have a large number of triangles tend to be highly interconnected. However, networks that have a low number of triangles turn out to be poorly connected and may suffer from instability.

**Modularity** is a measure of the structure of networks or graphs which measures the strength of division of a network into modules. Networks with high modularity have dense connections between the nodes within modules but sparse connections between nodes in different modules.

Cliques are fully connected subgraphs, meaning each node in a clique is directly connected to every other node in the clique. Therefore any clique of size n > 1 necessarily includes  $\binom{n}{n-1}$  subcliques. For our network statistics (e.g. **Clique-Size Mean**) we only measure maximal cliques, i.e. those cliques which are not sub-cliques of a larger clique.

**GiantProportion** is a simple metric that seems to bring unique information into the model. It is the ratio of nodes in the complete graph that are also members of the LCSG, this is *LCSG Node Count* divided by *Total Node Count*.

Next, we wrote a function which would find the **number of k-stars** in each PPIN from one to the maximum size star in the network. For example, a 1-star is a node in the network with one degree, a 2-star is a node in the network with two degrees, a 3-star is a node with three degrees, and so on.

Above we discussed how random mutations or environmental factors can cause failure of protein interaction networks. Through random and iterative removing of nodes, we simulated systemic damage. We observed changes in the network statistics over progressive simulated damage to the PPIN. The **critical threshold** is the point where node removal would cause less than 10% of the total graph to be included in the Giant Component. In the appendix of the paper from SNAP,

researchers specify that "when a network is so fragmented by the removal of nodes that the largest connected part of the network is sufficiently small (only 10% of the size of the original network), then any sensible dynamical process" of the PPIN will be unable to function. **Processing time?** 

We ran 10 simulations for each PPIN and took the mean value for critical threshold as a network statistic to feed into our models.

Finally, we explored the main core of each graph. Any complete graph is a core and every graph has a core. The main core is the core with the largest degree. That is, the main core is the largest maximal clique within each network. As a network statistic we considered how many edges each main core has. We expect that the larger the main core of a network, the more resilient it will be to our simulated network failure algorithm. Please see figure 2. Here you can see that the main core of the network shown

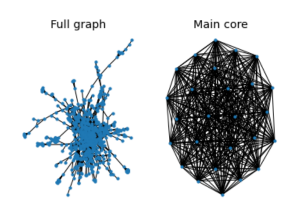


Figure 2: Main core

in the beginning of this paper is a 26-clique with 325 edges. Note that a complete graph of n vertices has  $\frac{n(n-1)}{2}$  edges.

#### III.II. Models

Our research question aims to find network statistics, which measure topological stability of the species' protein interactome, that are significant predictors of the evolutionary time of a species. That is, we suspect that there is a relationship between some of these network statistics and the response–evolutionary time. The question is, which of the network statistics have the most influence on the response variable? The nature of our data is that there are more variables than data points. Moreover, our network statistics all describe some aspect of topological stability for the network—meaning that they are all highly correlated with each other. For this reason we need models that work well with such data—we need models which perform feature selection, are interpretable, and highly accurate. For this we chose two supervised learning models: Ridge regression and Random Forest regression.

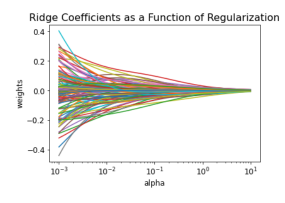


Figure 3: Ridge Tuning

The Ridge method regularizes model parameters by shrinking the regression coefficients, using the L2 norm, i.e. adds penalty equivalent to square the magnitude of coefficients. This regularization minimizes the sum of squared coefficients to reduce the impact of correlated predictors. This property allows Ridge regression to handle the multicollinearity of our network statistics while producing interpretable results. The feature selection phase occurs after the shrinkage, where variables with coefficients of relatively small magnitude are interpreted as having minor contribution to the response. Hence, important network statistics

for this model where taken to be those which have the coefficients of greatest magnitude. Furthermore, we tuned the penalty term of the Ridge regression using RidgeCV. Please see figure 3 for a plot of the Ridge coefficients as a function of the regularization penalty. We found that the optimal regularization penalty was alpha = 5.

Random forests build a collection of de-correlated decision trees, by randomly choosing only m predictors from the full set of predictors when performing a split, the split is only allowed to use one of those m predictors. Finally, the average of the resulting trees is taken. In random forest regression, features are selected that improve the variance reduction. That is, correlation between trees is reduced without increasing the variance too much. To tune the hyperparameters for this model we performed an exhaustive search using GridSearchCV from Scikit-learn. We tuned the number of estimators, max depth, and max number of features.

#### IV. Results

In figure 4, see that the network statistics with the most influence on the evolutionary time of each species for the Ridge model were average degree centrality, total network density, LCSG density, LCSG algebraic connectivity, modularity, and a handful of the larger number of k-stars variables. The sign of coefficients tells you if the network statistic is positively or negatively related to the response, evolutionary time. See that all of the listed significant statistics are positively related to evolutionary time.

We can expect that as the total network density and LCSG density increase, the number of actual connections in a network approaches the number of potential connections. That is, stability increases—making the network more resilient to failure or breakage. Recall that network resilience is the measure of how quickly a network breaks down as edges between nodes are randomly removed and it has been shown that networks with low network resilience also have a low evolutionary time. When a network is highly connected, though not necessarily fully connected, it has higher resilience. In this same way, we can expect average degree centrality and modularity to be positively related to network

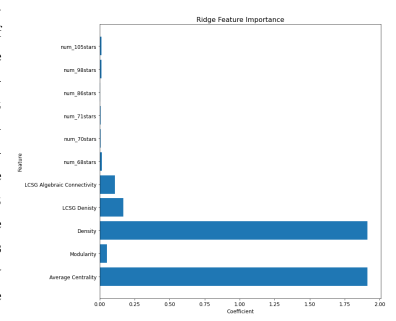


Figure 4: Ridge Feature Selection

resilience and evolutionary time. The numerically second smallest eigenvalue of the Laplacian matrix of a graph is known as the algebraic connectivity of the graph. This eigenvalue is greater than 0 only if the graph is connected. For this reason, we only find algebraic connectivity of the LCSG. The magnitude of this value reflects how well connected the overall graph is and it is always bounded above by node connectivity. The mean squared training error for the Ridge model was 0.01 and the mean squared test error was 0.126.

The random forest regression model was consistent with the results from above, please see the important variables for this model (See Figure 5). Arguably, the network statistics with the most influence on the evolutionary time of each species for this model were modularity, clique count, LCSG modularity, average centrality, clique-size mean, density, and number of other k-stars statistics.

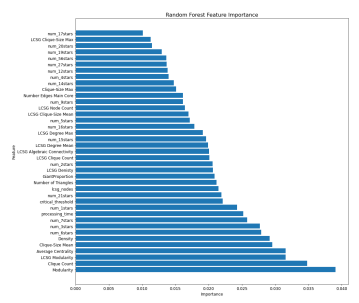


Figure 5: Random Forest Feature Selection

We explained before how modularity, density, and average degree centrality influence the topological stability of a network. The LCSG clique-size mean denotes the average size of a maximal clique within the LCSG. If this number is large, then we can expect the stability of this network to be strong and this translates to a higher network resilience. The number of cliques in the network, of clique count has a similar interpretation. In the random forest regression, the model did note the critical threshold, processing time, and number of edges in the main core as significant predictors. The mean squared training

error for the random forest regression model was 0.005 and the mean squared rest error was 0.028.

#### V. Conclusion

There are many possible confounders which should be considered along with our preliminary results: investigative biases towards modeling common or popular organisms, network size, and genome size. It seems that the network statistics we have found so far and entered into the models do a fair job of describing topological stability. Moreover, many of these statistics do have a significant relationship to evolutionary time of a species.

# VI. Appendix

# VI.I. Exhaustive Enumeration of Cliques is Memory-bound

We tried implementing exhaustive enumeration of all cliques (as opposed to maximal cliques) but found this to be impractical, as the supporting NetworkX algorithm (enumerate\_all\_cliques) is memory-bound. We tested this limitation via execution with 64GB of RAM and 3 Worker processes; execution failed with a MemoryError after  $\sim$  8 hours of processing, when one of a workers attempted to allocate additional memory beyond available capacity (see Figure 6).

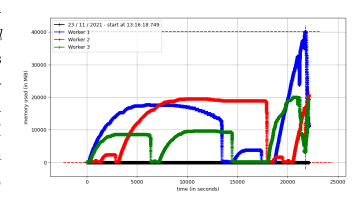


Figure 6: Worker memory usage

# VI.II. K-Stars Algorithm

Below is pseudocode for how we constructed the data frame of stars counts for each network.

# Algorithm 1 Get Stars Algorithm

 $LCSG \leftarrow giant component for undirected network$ 

 $A \leftarrow \text{convert LCSG to adjacency matrix}$ 

 $d \leftarrow sum row elements of A$ 

values, counts  $\leftarrow$  find unique elements and counts for each

stars  $\leftarrow$  pandas DataFrame of counts with index names being values

Table 1: Count K-Stars Data Frame

Species_ID	882	883	36870	52598	56780	•••
num_1stars	109.0	39.0	22.0	17.0	26.0	
$num\_2stars$	105.0	57.0	54.0	30.0	70.0	
$num\_3stars$	65.0	63.0	31.0	20.0	53.0	
:	:	:	:	:	:	٠

# VI.III. Predictors Using NetworkX

Table 2: Predictors Using NetworkX Data Frame

Species_ID	882	883	36870	52598	56780	
Average Centrality	0.007	0.011	0.017	0.008	0.011	
Number of Triangles	12883.000	12192.000	695.000	339.000	11755.000	
Modularity	0.710	0.730	0.695	0.796	0.640	
Density	0.007	0.011	0.017	0.008	0.011	
Clique Count	1060.000	3580.000	209.000	823.000	547.000	
Clique-Size Max	26.000	19.000	10.000	6.000	27.000	
Clique-Size Mode	2.000	5.000	2.000	2.000	2.000	
Clique-Size Mean	4.726	5.004	2.923	2.335	4.075	
LCSG Clique Count	916.000	371.000	180.000	249.000	451.000	
LCSG Clique-Size Max	26.000	19.000	10.000	5.000	27.000	
LCSG Clique-Size Mode	2.000	2.000	2.000	2.000	2.000	
LCSG Clique-Size Mean	5.102	4.647	3.039	2.116	4.237	
LCSG Node Count	736.000	502.000	217.000	139.000	536.000	
LCSG Degree Max	60.000	58.000	27.000	11.000	66.000	
LCSG Degree Mode	1.000	3.000	2.000	2.000	2.000	
LCSG Degree Mean	9.465	9.661	5.060	4.144	10.590	
LCSG Denisty	0.013	0.019	0.023	0.030	0.020	
LCSG Algebraic Connectivity	0.016	0.011	0.008	0.017	0.018	
LCSG Modularity	0.679	0.559	0.674	0.741	0.556	
Number Edges Main Core	325.000	171.000	45.000	10.000	351.000	
Critical Threshold	876.500	723.900	215.500	200.500	675.100	
Processing Time	102.449	61.957	6.729	14.796	54.736	

# VII. References

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