Integrative Network Analysis: Unveiling Symptom-Disease Interactions and Enhancing Predictive Models

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Github repository: https://github.com/AndreaAlberti07/enhancing-disease-prediction

Abstract

We will write it once we have the results.

Keywords— Graph theory • Features Engineering • Community detection • Null models • Random forest • MLP

Contents

1	Introduction		2
	1.1	Network Creation (Not Weighted - Bipartite)	2
	1.2	L1 and L2 measures	2
	1.3	Betweenness Centrality	2
	1.4	Communities Detection	2
	1.5	L1 and L2 Metrics	3
	1.6	Betweenness Centrality	3
	1.7	Communities	2
	1.8	Most Important Symptoms/Diseases (4	
		Classes)	

1 Introduction

n the dynamic landscape of healthcare, understanding the intricate interplay between symptoms and diseases is paramount for effective diagnosis and prediction. This report embarks on a comprehensive journey through the realms of network analysis, leveraging both theoretical foundations and empirical data to unravel the complexities of symptom-disease interactions. Our dual-fold objective is to provide a nuanced descriptive analysis of these interactions while identifying key features to bolster predictive models.

The foundation of this endeavor lies in an extensive review of existing literature, drawing insights from seminal works on network theory and disease prediction. By establishing a baseline through prior research, we pave the way for a deeper understanding of the subject matter and ensure the relevance of our findings in the broader context of scientific inquiry.

Guided by insights gleaned from the literature, our exploration extends to the realm of data, where we meticulously curate and analyze datasets of varying sizes. Through a systematic process of exploratory data analysis and cleaning, we prepare the groundwork for constructing meaningful networks that encapsulate the relationships between symptoms and diseases.

The heart of our analysis lies in the creation of intricate network structures, employing bipartite models and non-weighted links to distill meaningful patterns. We delve into a spectrum of network metrics, from fundamental measures like degree distribution and clustering coefficients to more nuanced assessments of node importance and betweenness centrality. Statistical significance is rigorously assessed through the lens of a null model, ensuring that our observations transcend mere chance.

Community detection algorithms further dissect the network, revealing hidden structures and relationships between diseases. This not only enriches our understanding but also lays the groundwork for subsequent analyses. As we traverse the terrain of network analysis, we introduce novel metrics inspired by the Hidalgo-Hausmann framework, stratifying symptoms and diseases based on their predictive importance. These metrics, coupled with traditional measures like betweenness centrality, contribute to the definition of features that fuel our predictive models.

With a robust foundation established, we transition to the realm of predictive modeling, where our feature-rich

approach promises to enhance the performance of established models. Logistic regression, random forest, and multi-layer perceptron models are trained, tested, and validated, with a keen eye on feature importance and model improvement strategies.

This report unfolds as a holistic exploration, weaving together theoretical frameworks, empirical analyses, and predictive modeling into a cohesive narrative. As we traverse the intricate web of symptom-disease interactions, our aim is not only to elucidate the underlying dynamics but also to pave the way for more accurate and insightful predictive models in the realm of healthcare.

Network Creation (Not Weighted - Bipartite)

1.2 L1 and L2 measures

1.3 Betweenness Centrality

The betweenness centrality of a node v, according to Brandes [2], is defined as the sum of the fraction of all-pairs shortest paths that pass through v:

$$c_B(v) = \sum_{s,t \in V} \frac{\sigma(s,t|v)}{\sigma(s,t)} \tag{1}$$

where:

- V: The set of nodes.
- σ(s, t): The number of shortest paths from node s to node t.
- $\sigma(s, t|v)$: The number of those shortest paths from node s to node t that pass through some node v other than s and t.
- If s = t, then $\sigma(s, t) = 1$.
- If $v \in \{s, t\}$, then $\sigma(s, t|v) = 0$.

To compute the betweenness centrality we used the NetworkX function nx.bipartite.betweenness_centrality which implements the algorithm proposed by Brandes [1] and uses a proper normalization for bipartite graphs.

1.4 Communities Detection

Prior to apply any community detection algorithm, we need to perform two steps:

 Graph Projections: We need to project the bipartite graph into two graphs, one for each set of nodes. In our case the two sets are represented by symptoms and diseases. At this scope is available the NetworkX function nx.bipartite.projected_graph which returns the projection of the bipartite graph onto the specified nodes.

 Compute Similarity: We need to compute the similarity between nodes. For our purposes, it is possible to create a co-occurrence matrix, for each set of nodes. Taking as example the co-occurrence matrix of symptoms, each entry s_{ii} represents the number of times the symptom iand the symptom *j* co-occur in the same disease.

Once we have the two graphs, whose links are weighted by the similarity between nodes, we can apply the community detection algorithm. We used the Clauset-Newman-Moore greedy modularity maximization algorithm [3], implemented in the NetworkX function $\verb|mx.algorithms.community.greedy_modularity_communit| \textbf{biss}_{n} \ dissecting \ the \ centrality \ values \ into \ symptoms$ This algorithm aims at finding the partition of the graph that maximizes the modularity, which is defined by Newman [6] as:

$$Q = \frac{1}{2m} \sum_{ij} \left[A_{ij} - \frac{k_i k_j}{2m} \right] \delta(c_i, c_j)$$
 (2)

where:

- Q: Modularity of the network.
- Aii: Element of the adjacency matrix representing the connection between nodes *i* and *j*.
- k_i and k_i : Degrees of nodes i and j, respectively.
- m: Total number of edges in the network.
- $\delta(c_i, c_i)$: Kronecker delta function, which is 1 if c_i is equal to c_i (i.e., nodes i and j belong to the same community) and 0 otherwise.
- The sum is taken over all pairs of nodes i and j.

1.5 L1 and L2 Metrics

1.6 **Betweenness Centrality**

The examination of betweenness centrality in our bipartite network, as depicted in Figure 1, reveals a Power Law Distribution, indicative of a scale-free structure. This implies the presence of a few central nodes that act as pivotal connectors, while the majority of nodes exhibit lower betweenness centrality.

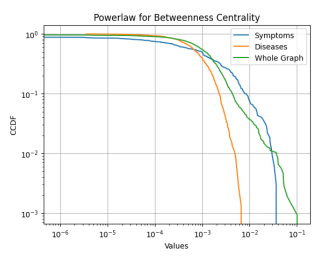


Figure 1. Betweenness Centrality CDFs

and diseases (see Figures 2 and 3), a notable observation emerges: symptoms tend to have higher betweenness centrality compared to diseases. To decipher the significance of this result, it's essential to delve into the interpretation of betweenness centrality.

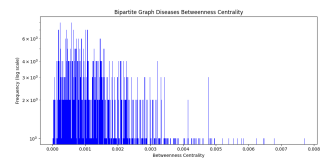


Figure 2. Betweenness Centrality of the diseases

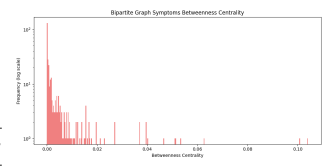


Figure 3. Betweenness Centrality of the symptoms

In general, a symptom exhibits high betweenness cen-

trality when it is linked to numerous diseases, and these diseases, in turn, are connected to a relatively limited set of symptoms. Conversely, a disease attains high betweenness centrality when it connects to numerous symptoms, and these symptoms are associated with relatively few diseases.

Analyzing our results (L1 and L2), it becomes evident that the higher betweenness centrality of symptoms is attributed to their connections with a multitude of diseases, while diseases, on the contrary, are linked to a relatively limited number of symptoms. From a predictive standpoint, this outcome presents a challenge as each symptom is not sufficiently specific, contributing to a broad array of disease classes.

Figure 4 highlights the top 10 nodes with the highest betweenness centrality, all of which are symptoms. As anticipated, these symptoms are more generic in nature, aligning with their central role in connecting various diseases.

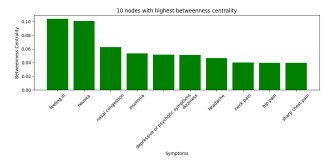


Figure 4. Top 10 nodes with the highest betweenness centrality

1.7 Communities

The identification of communities within the network serves a dual purpose – facilitating network interpretation and enhancing the capabilities of our ML prediction model.

From a network interpretation perspective, communities offer insights into disease-symptom relationships. A community of symptoms signifies a set of symptoms that frequently co-occur within the same diseases, while a community of diseases identifies a set of diseases often co-occurring within the same symptoms. The sizes of different communities are illustrated in Figure 5.

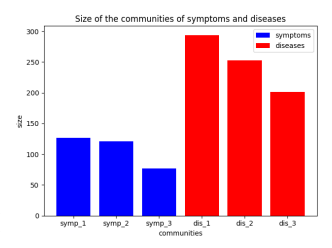


Figure 5. Sizes of the communities of symptoms and diseases

For clinical relevance, examining symptoms communities provides valuable information about diseases associated with these symptoms. This is exemplified in Figures 6, 8, and 9. As an illustration, in the community 1 of symptoms (Figure 6), 'herniated disk' is the third most pointed disease by the symptoms of the community, with each symptom pointing, on average, to three diseases.

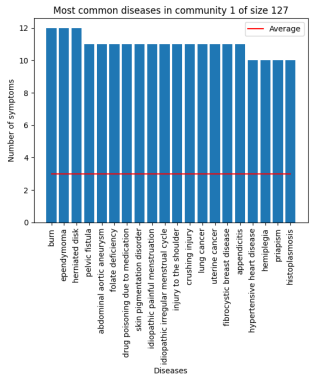


Figure 6. Community 1 of symptoms

A similar study can be conducted for communities of diseases, as depicted in Figures 7, 10, and 11. This information aids in profiling diseases and understanding the significance of each symptom. For instance, in community 1 of diseases (Figure 7), the symptom 'sharp abdominal pain' is present in almost half of the diseases in the community, indicating its generic nature and limited discriminatory value.

For example, a symptom associated with many diseases may be less informative and could potentially be removed from the symptom vector. However, we opted for a comprehensive approach using a combination of L1 and L2 measures to address this issue.

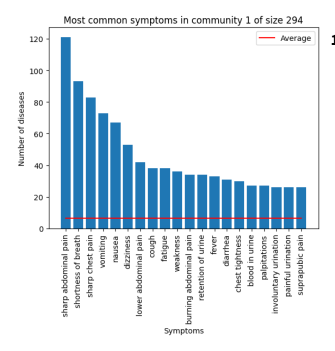


Figure 7. Community 1 of diseases

Transitioning to the creation of features for the ML model, two types of features were developed:

- Community Count: This feature counts how many symptoms of the symptom vector belong to each community. Each symptom community is characterized by different pointed diseases. The model can learn to prioritize diseases associated with the community with the highest count.
- Community Size: This feature replaces each symptom in the symptom vector with the size of the community to which the symptom belongs. It enables the model to distinguish between symptoms belonging to small and large communities, injecting community information into the model beyond basic one-hot encoding of symptoms.

It is noteworthy that communities can also contribute to improving the computational efficiency of the model.

1.8 Most Important Symptoms/Diseases (4 Classes)

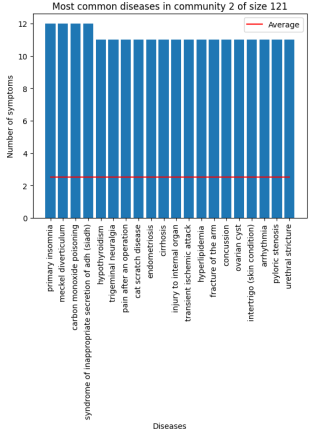
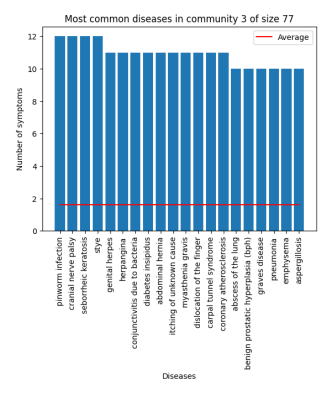


Figure 8. Community 2 of symptoms

Integrative Network Analysis



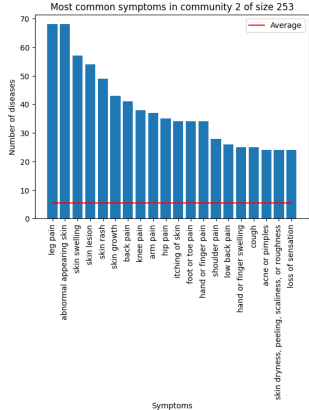


Figure 9. Community 3 of symptoms

Figure 10. Community 2 of diseases

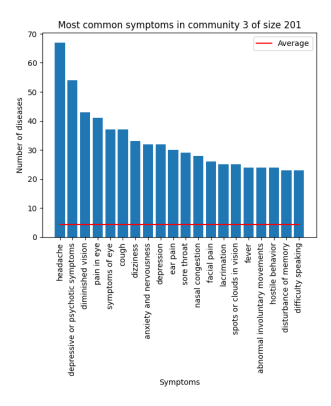


Figure 11. Community 3 of diseases

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