

DISEASE PREDICTION VIA GRAPH-BASED MODEL

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VIETTEL DIGITAL TALENT PROGRAM 2023

ABSTRACT

Disease prediction is a well-known classification problem in medical applications. Graph Convolutional Networks (GCNs) provide a powerful tool for analyzing the patients' features relative to each other. This can be achieved by modeling the problem as a graph node classification task, where each node is a patient or disease. Graph-based models overcome the limitations of traditional machine learning models by not considering individual patient data records as discrete entities but rather establishing connections between symptoms, similar diseases, and generating different disease groups. In this report, the author proposes a Graph Neural Network model with various information aggregation mechanisms such as Gated Recurrent Unit (GRU), ... the mechanism updates information from neighboring nodes through Attention mechanism, building node features for the disease graph based on symptoms using the TF-IDF formula and achieves a 100% F1-Macro evaluation result for disease prediction based on symptoms dataset from India.

KEYWORDS

Graph-based model, Graph Attention, Create node feature by TF-IDF

1 INTRODUCTION

EMRs, short for electronic medical records, are a commonly utilized data management scheme employed in hospitals to store comprehensive clinical information obtained from patients' visits. In recent times, advancements in information technology and machine learning have led to a more manageable volume of EMRs. Consequently, the analysis of EMRs using machine learning and data mining techniques has emerged as a prominent research direction, aiming to enhance healthcare services [5]. One notable application of machine learning in the healthcare domain involves disease prediction, which focuses on determining whether a patient is afflicted with a specific ailment. This task typically involves training a classifier to make predictions based on the information derived from EMRs [17], [11]. For instance, Palaniappan and Awang employed various data mining techniques, such as Decision Trees [14], and Neural Networks [8], to construct a predictive system for heart diseases [12]. Leveraging the capabilities of Convolutional Neural Networks (CNNs), Suo et al. [17] initially identified similarities among patients based on their EMRs and subsequently conducted personalized disease predictions.

Existing deep neural networks lack the incorporation of patient interactions and associations within their architecture. However, considering these relationships can prove advantageous as it facilitates the analysis and study of similar patient cohorts. Graphs offer a natural means of representing the interactions among a

population by treating patients as nodes and their associations as edges. Constructing a graph between patients based on a subset of their features allows for the summarization of these features within the graph edges, reducing feature dimensionality and mitigating overfitting caused by a large number of features [2], [18].

Related work

In recent years, GCNs have been adopted in different applications, especially in medical domains e.g., brain analysis [9], mammogram analysis [4],.... The disease prediction problem has been also widely explored by GCN-based methods [1], [10]. Following the promising results of GCNs in the medical domain, [13] exploit GCNs in the disease prediction problem with multi-modal datasets. They chose ChebyNet with constant filter size as the classifier and investigated the affinity graph construction by computing the distance between the subjects for brain analysis. InceptionGCN [7] extends ChebyNet and designs filters with different kernel sizes, and chooses the optimal one for the final disease prediction. A branch of methods focuses on improving the graph structure. [6] and [16] start with a pre-constructed graph and update it during the training, but [3] learn the whole graph from features directly in an end-to-end manner.

Main contribution

In this report, my main contributions are as follows:

- Combining GCN layers, Multi Layer Perceptron (MLP), and mechanisms for information exchange and update, such as using Gated Recurrent Neural Network (GRU) or Attention mechanism, to build a disease prediction model with a dataset from India.
- Proposing methods for constructing different graphs: disease graph and patient graph.
- Introducing a training method that combines Multi Head Attention layer and reduces the size of the input graph while still achieving good prediction results with various metrics like Recall, Precision, and F1.
- Proposing a method for building a feature set for the disease graph based on the TF-IDF formula.

2 DATA PREPROCESSING

In this report, I utilize two datasets for disease prediction based on relevant symptoms to construct multi-class classification and binary classification problems.

2.1 Disease Symptom Prediction

This dataset ¹ consists of 4920 records with 18 different fields representing disease symptoms. There are 41 distinct diseases, with each disease having 120 records. After conducting the Exploratory Data Analysis (EDA), I observed that each disease exhibits 2-3 distinctive symptoms. There are also common symptoms shared among different diseases, such as headaches and fatigue. Additionally, patients infected with the same disease tend to have 1-2 similar

Supervised by LE HOANG NGAN.

Mini Project in VDT, gen 3, (2023)
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¹<https://www.kaggle.com/datasets/itachi9604/disease-symptom-description-dataset>

symptoms. These findings inspire the idea of constructing graphs for different disease types and patient graphs based on shared symptoms. These graphs serve to generalize the information of the patients, and by utilizing Graph Neural Networks (GNN), we can extract latent features from the data.

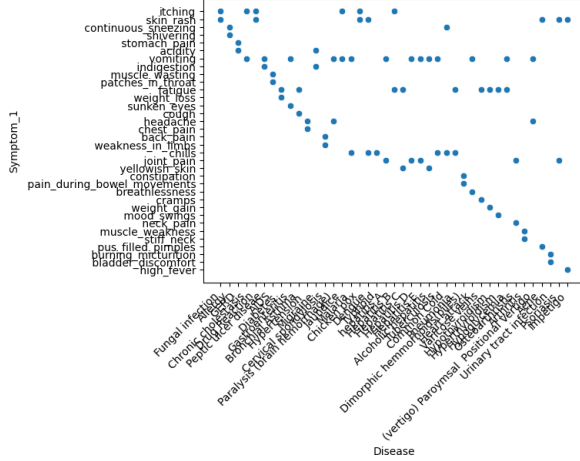


Figure 1: Relation between Symptom and Disease

2.2 Pima Indian Diabetes (Diabetes) [15]

The dataset is produced by the "National Institute of Diabetes and Digestive and Kidney Diseases". The goal of this dataset is to recognize the diabetic status of patients (binary classification). Every patient has 7 numeric features which show the diagnostic measurements of diabetes including the number of pregnancies, plasma glucose, blood pressure, skin thickness, Body Mass Index (BMI), insulin level, diabetes pedigree function, and age.

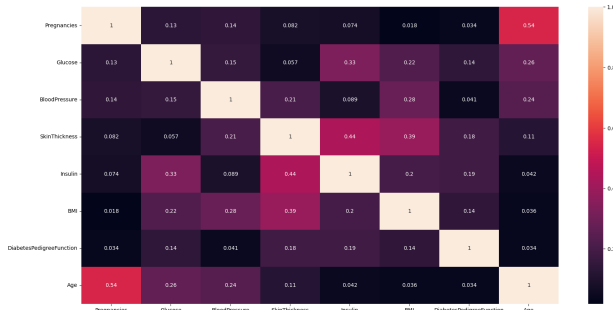


Figure 2: Correlation between feature columns

Based on the data, we can observe a relatively high correlation between the age field and other fields in the dataset. Therefore, I perform binning on the age field to create different age groups and utilize them to construct graph connections between these age groups.

3 THE PROPOSED METHOD

In this report, I will build various graph-based models, including both single-graph and multi-graph inputs. Additionally, I will utilize different mechanisms to update and aggregate messages between nodes.

3.1 Definitions

We consider a graph G with N nodes, denoted by $G(V, E, X)$, where V represents the set of nodes ($|V| = N$), E is the set of edges, and $X \in \mathbb{R}^{N \times F}$ denotes the node feature matrix. The adjacency matrix $A \in \mathbb{R}^{N \times N}$ is unweighted and undirected, reflecting the connections between nodes. Each node v_i has an associated feature vector x_i , a one-hot label vector y_i , and a true class label $c_i \in C$, where C represents the set of classes. The label information is available only for a subset of nodes, and the objective is to learn a parametric function $f_\theta(X, A)$ that takes the adjacency matrix and node features as input and predicts the true labels for the unlabeled nodes. It is important to note that a probabilistic classifier generates a probability distribution over the $|C|$ classes, and the label with the highest probability is selected. In our proposed method, q_i represents the output probability distribution of the classifier, defined over the $|C|$ classes, for the sample x_i , where the c -th element represents the classifier's confidence in assigning the label c to x_i . Thus, the problem can be formulated as follows:

$$Q = f_\theta(X, A),$$

where $Q \in \mathbb{R}^{N \times |C|}$ is the prediction matrix of the classifier for all nodes, including the unlabeled ones.

3.2 Neural Graph Encoder

To start, in the context of a graph G , we adopt a uniform representation for disease, symptom, or patient nodes as $v \in G$ for brevity. At the l -th layer of information propagation, the embedding \mathbf{h}_v^l of node v is computed as follows:

$$\mathbf{h}_{\mathcal{N}(v)}^l = \text{AGGREGATE} \left(\left\{ \mathbf{h}_{v'}^{l-1}, \forall v' \in \mathcal{N}(v) \right\} \right)$$

$$\mathbf{h}_v^l = \sigma \left(\mathbf{W}^l \cdot \left[\mathbf{h}_v^{l-1}; \mathbf{h}_{\mathcal{N}(v)}^l \right] \right)$$

Here, \mathbf{W}^l denotes the weight matrix to be learned at the l -th layer, \mathbf{h}_v^{l-1} represents the embedding of node v at the previous layer, and L denotes the total number of layers. The concatenation of two vectors is denoted by $[\cdot; \cdot]$, and $\mathcal{N}(v)$ represents the set of neighbor nodes of v sampled uniformly. It is important to note that for $l = 0$, the node embedding $\mathbf{h}_v^0 \in \mathbb{R}^d$ is initialized using either random values or side information from the data, depending on its availability. For example, in the case of a patient node, if patient demographics and medical profiles are available in the electronic medical record (EMR) data, \mathbf{h}_v^0 will be initialized as a dense, real-valued feature vector. Each element in \mathbf{h}_v^0 would then represent an observed value of a specific feature dimension (e.g., age). $\mathbf{h}_{\mathcal{N}(v)}^l$ represents the synergistic representation obtained through the aggregation function, which is designed to combine the embeddings of node v 's neighbors from the $(l-1)$ -th layer. σ denotes a non-linear activation function such as the hyperbolic tangent (tanh), and the aggregator can be selected from options like mean, max pooling, recurrent neural networks (RNNs), and so on.

Then, we take a normalization step before reaching the final embedding for all nodes at the last layer L :

$$\mathbf{h}_v = \frac{\mathbf{h}_v^L}{\|\mathbf{h}_v^L\|_2}, \forall v \in G$$

After passing through the GCN layers, we will feed \mathbf{h}_v through an MLP layer to obtain the final embedding representation.

$$\mathbf{z}_v = \sigma(\mathbf{W}\mathbf{h}_v), \forall v \in G$$

where \mathbf{W} is the learnable weight, and \mathbf{z}_v is the final embedding for node v .

Attention mechanism

We can apply attention mechanisms to selectively encode the information from neighbors according to their importance to the target node v . This is achieved by taking a weighted sum of the representations of all v 's neighbor nodes:

$$\mathbf{h}_v^l = \sum_{v' \in \mathcal{N}(v)} \alpha_{v'v} \mathbf{M} \mathbf{h}_{v'}^{l-1}$$

where \mathbf{M} is the transformation weight matrix, and $\alpha_{v'v}$ is the attentive weights indicating the importance of neighbor node $v' \in \mathcal{N}(v)$ when calculating \mathbf{h}_v^l . Each $\alpha_{v'v}$ is computed via the following attention network:

$$\alpha_{v'v} = \frac{\exp\left(\text{LeakyReLU}\left(\mathbf{a}^T \left[\mathbf{N} \mathbf{h}_v^{l-1} \parallel \mathbf{N} \mathbf{h}_{v'}^{l-1}\right]\right)\right)}{\sum_{k \in \mathcal{N}(v)} \exp\left(\text{LeakyReLU}\left(\mathbf{a}^T \left[\mathbf{N} \mathbf{h}_v^{l-1} \parallel \mathbf{N} \mathbf{h}_k^{l-1}\right]\right)\right)}$$

with a projection vector \mathbf{a} and the weight matrix \mathbf{N} . Essentially, the learned attentive weights allows the aggregator to lay more emphasis on neighbor nodes having more contributions to the message passing process, thus being able to generate highly expressive node embeddings.

Disease Prediction for Patients

The graph decoder in our model plays a crucial role in translating the information captured by symptom, disease, and patient node embeddings into predictions of potential diseases associated with a given patient. To elaborate, when provided with the embedding \mathbf{z}_p of a patient p , the graph decoder transforms it into a vectorized output $\hat{\mathbf{c}}_p \in [0, 1]^{|C|}$ that serves as an approximation of the patient's multi-hot disease label $\mathbf{c}_p \in [0, 1]^{|C|}$. Specifically, during the decoding process, each element of $\hat{\mathbf{c}}_p$ is computed as follows:

$$\hat{c}_{p,n} = \text{sigmoid}\left(\mathbf{z}_p^T \mathbf{Q}_n\right), \forall m_n \in \mathcal{V}_M$$

where $\hat{c}_{p,n}$ is the n -th element in $\hat{\mathbf{c}}_p$, while $n \leq |\mathcal{V}_M|$ is used for indexing all the diseases m . The closer $\hat{c}_{p,n} \in \hat{\mathbf{c}}_p$ is to 1, the more likely patient p is diagnosed with disease m_n . $\mathbf{Q} \in \mathbb{R}^{|\mathcal{V}_M| \times d}$ carries the corresponding regression weights for all diseases, and \mathbf{Q}_n is the n -th column of it. To train our model, we quantify the prediction error via the following negative log likelihood loss function:

$$\mathcal{L} = - \sum_{n=1}^{|\mathcal{V}_M|} c_{p,n} \log(\hat{c}_{p,n})$$

3.3 Proposed Model

I will build two different types of Graph-based models, using either a single graph or multiple graphs as inputs. To avoid the over-smoothing problem, I will only use two layers of Graph Convolution Network (GCN) and incorporate skip connections to preserve information. To enhance the model's representational capacity, I will combine MLP layers, including Fully Connected and Batch Normalization layers, to extract information effectively. Moreover, during the training process, I will employ various mechanisms to aggregate information, such as SUM, MAX, MEAN, and Attention, to effectively combine the information from neighboring nodes. For updating node representation, I will utilize MLP layers to learn isomorphic or non-isomorphic sub-graphs and employ a GRU network to learn sequential representations of neighboring nodes.

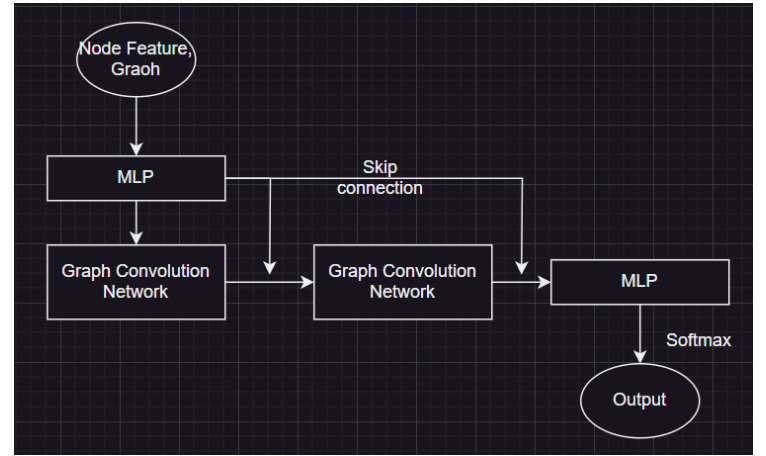


Figure 3: Proposed model with 1 input graph

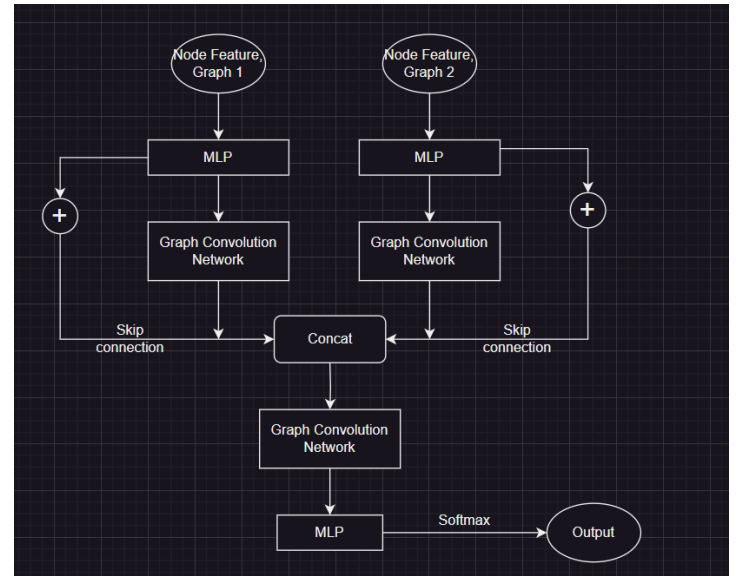


Figure 4: Proposed model with 2 input graph

Training method

When training model, I have this Hypothesis: Reducing the size of the input graph by sampling a small percentage of connections (5%, 10%, 20%, 30%, etc.) does not decrease the model's performance if information is combined using the Attention mechanism. Because, based on the data, each individual is only grouped with 120 other individuals (1/41) with the same disease, so only a small representative group is needed for each disease group. The Attention mechanism helps the model learn important connections within the graph. A node learns which connections are significant with its neighboring nodes. Therefore, removing edges with low probability does not affect the performance. The experimental results in the following section demonstrate the effectiveness of this training approach.

Create feature for Disease Graph

I propose to construct features for the disease graph based on the TF-IDF formula by these steps:

- (1) Calculate the top 3 most common symptoms for each disease.
- (2) TF (Term Frequency): Count the number of individuals with each disease who exhibit the corresponding common symptom.
- (3) IDF (Inverse Document Frequency): Count the occurrence of each common symptom across the entire dataset.
- (4) Calculate the weight by multiplying TF with 1/IDF.

So we have a feature matrix for graph disease.

4 RESULTS & CONCLUSION

4.1 Result

In this report, to evaluate the classification performance of the model, I optimize the parameters based on the F1-macro, Recall-macro, and Precision-macro metrics. This approach is chosen because in reality, there are numerous rare diseases with very few data records. Optimizing parameters using the macro strategy helps to address the issue of data imbalance to some extent.

Graph Construction

Firstly, we need to define how to construct graph. I proposed some strategy to construct 2 type of graph: Disease Graph, Patient Graph.

For establishing connections between two patients, we can:

- Strategy 1: Create an edge between two patients if they have the same type of disease (To avoid data leakage, only use the training set to construct the graph).
- Strategy 2: Create an edge between two patients if they share two similar symptoms. (This approach can be combined with Strategy 1 to create a new graph).
- Strategy 3: Create an edge between two patients if their cosine similarity, measured by a threshold γ , exceeds a certain threshold.

$$a_{ij} = \begin{cases} 1, & \text{if cosine-similarity}(x_i, x_j) \geq \gamma \\ 0, & \text{otherwise} \end{cases}$$

where a_{ij} is indicator for representing the existence of an edge connecting patients i -th and j -th and x_i is sample of patient i -th.

For establishing connections between two diseases, we can create a connection between two diseases if they share the top two prominent symptoms.

We divided the dataset into train/validation/test sets with a ratio of 70/10/20, respectively. The results on 2 data sets will be represented below.

4.1.1 Multi-class Classification. After conducting experiments on the Kaggle dataset, we obtained the following results for each graph construction strategy and methods of aggregating information from neighboring nodes and updating node representations:

Model	Accuracy	F1-macro	Recall-macro	Precision-macro
Baseline Random Forest	0.91 ± 0.01	0.90 ± 0.01	0.90 ± 0.01	0.95 ± 0.01
Strategy 1 (Mean + GRU)	0.99 ± 0.01	0.99 ± 0.01	0.99 ± 0.01	0.99 ± 0.01
Strategy 2 (Sum + Concat)	0.98 ± 0.01	0.98 ± 0.01	0.99 ± 0.01	0.99 ± 0.01
Strategy 3 (Attention + GRU + sample 20%)	0.95 ± 0.01	0.94 ± 0.01	0.95 ± 0.01	0.95 ± 0.01
Strategy 3 (MultiHeadAttention + concat + sample 5%)	0.99 ± 0.01	0.99 ± 0.01	1 ± 0.01	0.99 ± 0.01
Multi Graph Input (Mean + GRU)	1 ± 0.01	1 ± 0.01	1 ± 0.01	1 ± 0.01

We can observe that the Graph-based models all outperform the baseline model, Random Forest, demonstrating their effectiveness on this dataset. Furthermore, the training method that reduces the size of the input graph and utilizes the Multi Head Attention layer helps the model focus on important connections within the graph, resulting in excellent classification results.

4.1.2 Binary Classification. We proceeded to build a binary classification model using the Diabetes dataset. To construct the graph according to Strategy 2, I performed age binning to form age groups and established connections between patient age groups. We can see that the Graph-based models are still very effective; however, constructing the graph according to Strategy 2 resulted in less effective learning for the model.

Model	Accuracy	F1-macro	Recall-macro	Precision-macro
Baseline KNN	0.73 ± 0.01	0.73 ± 0.01	0.73 ± 0.01	0.74 ± 0.01
Strategy 1 (Mean + GRU)	0.77 ± 0.01	0.76 ± 0.01	0.76 ± 0.01	0.79 ± 0.01
Strategy 2 (Sum + Concat)	0.76 ± 0.01	0.75 ± 0.01	0.75 ± 0.01	0.78 ± 0.01
Strategy 3 (Attention + GRU + sample 20%)	0.77 ± 0.01	0.77 ± 0.01	0.77 ± 0.01	0.79 ± 0.01

4.2 Conclusion

Through this project, I have achieved the following results:

- Understanding and constructing Graph-based models such as GraphSage, Graph Attention, etc., with one or multiple graphs as inputs.

- Proposing different approaches to construct various graphs (e.g., disease graph, symptom graph) and building a feature set for diseases based on applying TF-IDF measure to prominent symptoms.
- Introducing the Graph-based MultiHeadAttention model with a mechanism to sample edges, which helps reduce the size of the graph while ensuring classification performance.

In the future, I will supplement the data by crawling additional data to evaluate the effectiveness of the model with a larger dataset. Also, Constructing various weighted graphs with different types of nodes (including graphs consisting of patient nodes, disease nodes, and interconnected symptom nodes).

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