Disease gene prediction with privileged information and heteroscedastic dropout

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Abstract

Motivation: Recently, machine learning models have achieved tremendous success in prioritizing candidate genes for genetic diseases. These models are able to accurately quantify the similarity among disease and genes based on the intuition that similar genes are more likely to be associated with similar diseases. However, the genetic features these methods rely on are often hard to collect due to high experimental cost and various other technical limitations. Existing solutions of this problem significantly increase the risk of overfitting and decrease the generalizability of the models.

Results: In this work, we propose a graph neural network (GNN) version of the Learning under Privileged Information paradigm to predict new disease gene associations. Unlike previous gene prioritization approaches, our model does not require the genetic features to be the same at training and test stages. If a genetic feature is hard to measure and therefore missing at the test stage, our model could still efficiently incorporate its information during the training process. To implement this, we develop a Heteroscedastic Gaussian Dropout algorithm, where the dropout probability of the GNN model is determined by another GNN model with a mirrored GNN architecture. To evaluate our method, we compared our method with four state-of-the-art methods on the Online Mendelian Inheritance in Man dataset to prioritize candidate disease genes. Extensive evaluations show that our model could improve the prediction accuracy when all the features are available compared to other methods. More importantly, our model could make very accurate predictions when >90% of the features are missing at the test stage.

Availability and implementation: Our method is realized with Python 3.7 and Pytorch 1.5.0 and method and data are freely available at: https://github.com/juanshu30/Disease-Gene-Prioritization-with-Privileged-Information-and-Heteroscedastic-Dropout.

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1 Introduction

Identifying disease genes is the most important step for understanding parthenogenesis and for searching therapeutic targets. Experimentally verifying a causal link between gene and disease is time-consuming and expensive. The past decades have witnessed the success of a number of computational models in prioritizing new genes based on known disease and gene associations. Network fusion algorithms (Aerts et al., 2006; Britto et al., 2012; Chen et al., 2007, 2009; De Bie et al., 2007; Gefen et al., 2010; Gerstein et al., 2012; Guney and Oliva, 2012; Kim et al., 2015; Kumar et al., 2018; Magger et al., 2012; Robinson et al., 2014; Tranchevent et al., 2008; Yang et al., 2015; Zakeri et al., 2018; Zitnik et al., 2015) were first proposed to combine different sources of information on both diseases and genes and provide an universal ranking of associations for any disease gene pairs. Another type of approaches implementing the same intuition try to model this problem as a recommender system, in which diseases and genes represent customers and products, respectively (Natarajan and Dhillon, 2014; Zakeri et al., 2018). Disease gene prioritization problem can also be modeled as a link prediction problem based on the node and edge features on graphs. The early graph-based approaches mainly relies on standard random walk (Chen et al., 2009; Köhler et al., 2008; Lee et al., 2011) and its alternative forms (Adie et al., 2005; Chen et al., 2009; Erten et al., 2011; Franke et al., 2006; Köhler et al., 2008; Linghu et al., 2009; Martinez et al., 2015) to smooth the signal on the graphs.

The main intuition behind these computational models is that similar genes are more likely to be associated with a similar set of diseases and similar diseases tend to share similar disease genes (Latif et al., 1993; Tanzi et al., 1993). Therefore, an important component of these models is the features adopted to quantify the
similarity among diseases and genes. For genes to be prioritized, their similarity should reflect their functional similarities, which could be characterized by various types of information such as their involvement in the same biological processes, their physical or genetic interactions, their co-expression patterns in certain cell lines, co-occurrence evidence from the literature and their phenotypic similarity after gene deletions. Disease similarity could be calculated based on the overlap of their symptoms or their semantic distance on the disease ontology.

However, in practice such features are often hard to collect on a large scale due to technical limitations. For instance, the widely adopted gene feature dataset (Lamb et al., 2006; Subramanian et al., 2017) only measures the knocking out effects on 500 genes in humans. Liu et al. (Liu et al., 2016) constructed a sample-specific network to describe an individual’s disease state by integrating the co-expression correlations and PI interactions within a tissue. However, those tissue-specific data are only available for a limited number of cell lines and tissues. A third example is Shim et al. (Shim et al., 2019) which relies on the pathway-specific protein domains to predict disease genes. However, they found that only 27% of InterPro (Mitchell et al., 2015) domains had function/pathway annotation from InterPro2GO. In addition, many other gene features such as the associated drug information and evolutionary history are also hard to collect for all the genes due to experimental limitations (Wang et al., 2019a).

A trivial solution adopted by most of these algorithms is to take the union of all the features and create a very large and sparse feature matrix by filling the missing values as zeros. In this so-called gene prediction problem, the number of positive training samples corresponding to the known disease–gene associations is very small. Therefore, merging all features together increases the risk of overfitting the training data by significantly enlarging the feature dimension. Second, this trivial solution significantly limits the generalizability of the machine learning models. Consider the gene knocking out features in the CMAP project, if only 500 genes have such features, then the model has to rely on many zeros to make predictions for the rest of the genes and lead to less accurate results. Third, such a high feature dimension requires a more complicated model, which poses technical challenges for certain types of machine learning models. For instance, graph neural networks (GNN) were recently adopted to predict disease genes (Liu et al., 2020; Li et al., 2019b; Testolin and Zorzi, 2016; Wang et al., 2019b) and it is widely known that GNN models suffer from an ‘over-smoothing’ problem when their architectures become very deep (Li et al., 2018). In addition, when a data observation has a missing feature, we usually impute this missing value by taking average of this feature values among all the other observations that have this feature. However, sometimes there are many observations missing this feature, then impute this missing value by taking average is not a good choice. In this case, we choose to delete this observation. Therefore, it is typical for the above methods to remove all the genes without any of the above features from both training and test data, which further limits their usage and generalizability (Li et al., 2019b). However, if we still include these sparse feature genes in the model and fill the missing features with zero, then the learnt embedding of these genes will have a high variance. This side effect might become even larger when we train the model in a batch fashion based on neighborhood sampler or reconstructed subgraphs, which will make the model very unstable and hard to generalize.

To address these problems, we developed a new computational framework to predict disease gene associations, which belongs to the family of Learning under Privileged Information (LUI) (Vapnik and Vashist, 2009). We implemented it using a relational graph neural network (RGCN) (Schlichtkrull et al., 2020) with the Heteroscedastic Gaussian Dropout. One of the intriguing properties of our model is that the training and test samples could take different sets of features. In particular, we introduce a concept called ‘privileged features’ to indicate those features that are only available for the training samples. The main computational challenge we addressed here is how to leverage the information provided by these privileged features to improve the prediction accuracy on the test samples. The backbone of our model is two connected RGCN models with the same architecture running on top of a heterogeneous network of diseases and genes. One RGCN model is used to map the features of diseases and genes to low dimensional embedding by using a non-linear aggregation of the neighborhood information. The second RGCN takes the privileged features which might be missing at the test stage to control the dropout probability of the first RGCN model. Since the dropout operation is not needed at the test time, the second RGCN and the privileged features can be excluded when making predictions. The second RGCN model injects the information provided by the privileged features into the first RGCN by determining the importance and scales of noise of each hidden neuron at each training epoch. As a result, we can include the information of those genes with sparse features and control the variance of the learnt embedding. This learnable and informative dropout model is very useful because usually not all the nodes in a graph have the same set of features, this is especially true in the biology domain where features are difficult to get due to high experiment cost. This framework is not only suitable to the graph convolutional networks, but also can be extended to various of GNNs, such as GraphSage (Hamilton et al., 2017) and GAT (Velickovic et al., 2017), which has been more and more applied to biology domain (Kwak et al., 2020; Zitnik et al., 2018).

To evaluate our method, we compared our method with four state-of-the-art disease gene prediction algorithms, including IMC (Natarajan and Dhillon, 2014), GeneHound (Zakeri et al., 2018), Catapult (Singh-Blom et al., 2013), Katz (Singh-Blom et al., 2013) and a basic RGCN model on the Online Mendelian Inheritance in Man (OMIM) dataset (Hamosh, 2002). We evaluated the performance of these models on various graph learning settings including general link prediction, link prediction on unseen nodes and on sparse features. Extensive experiments show that our method significantly outperforms other methods on these machine learning tasks related to disease gene prediction. More importantly, our model can accurately make link predictions for those genes with sparse features or even without any genetic features by successfully incorporating privileged features during the training process.

2 Materials and methods

In this work, we model the disease gene prioritization problem as a link prediction task. We integrate disease similarity, gene similarity and disease–gene associations into a multi-relational network. As shown in Figure 1, two RGCN models are constructed to transform two separate feature sets into one low dimensional embedding which represents either a disease or a gene. Another edge decoding layer then transforms the embedding’s of a disease–gene pair to their strength of associations. Note that the parameters in both RGCNs and the decoding layer are trained in an end-to-end manner so that they could act as regularizers for each other.

2.1 Relational graph convolutional networks

To capture the non-linear relationship between features, labels and network topologies, we adopt the relational graph convolutional network (RGCN) (Schlichtkrull et al., 2018) framework to jointly model different types of edges in the disease–gene graph. The convolution operation in RGCN is the same as the one defined in the graph convolutional neural networks (GCN) (Bruna et al., 2013) in which the parameters are shared over all the locations in the graph. The central idea of GCN is to generate a node representation by aggregating its own features and neighbors’ features. This aggregation process can be viewed as a message passing operation over the entire graph. The propagation rule on a regular graph for the lth layer is defined as follows,

\[
H^{l+1} = \sigma \left( D^{-1/2} \cdot AD \cdot H^l W^l \right)
\]  

(1)

where \(H^l\) is the node embedding in the lth layer and \(H^0\) is the raw feature matrix. \(A\) is the adjacency matrix of the regular graph and \(D\)
is the degree matrix in which dropout of the main net whose outputs are used as the variance of a Gaussian distribution in the Gaussian neighborhood information. The dropout net (right) is another RGCN model whose outputs are used as the variance of a Gaussian distribution in the Gaussian neighborhood information.

Our framework is also built on the RGCN model, but we incorporate the privileged features during training, we introduce two multi-layer RGCNs with the same architecture, i.e. the numbers of layers, hidden neurons and their connectivities in each layer are exactly the same. As shown in Figure 1, the first RGCN (left), named the main net, is used to transform the public features $x$ to node representations and later to the final predictions. The second RGCN (right), named the dropout net, takes the privileged features $x$ and transforms it to the variance of a Gaussian distribution which controls the confidence of each hidden neuron in the main net. At each training epoch, the two RGCN models perform message passing separately on the same graph using the updating rule defined at Equation (2). Let $m(x)$ and $d(x')$ denote the node embedding of the main net and dropout net, respectively, then the value of the $i$th dimension $v_i$ of the joint representation of both $x$ and $x'$ is defined as the follows,

$$v_i = m_i(x) + d_i(x')$$

For conventional dropout (Srivastava et al., 2014), the second term of the right-hand side of Equation (3) is a Bernoulli distribution with a fixed probability. For Gaussian dropout, it simply replaced the Bernoulli distribution with a Gaussian distribution (Lambert et al., 2018). The mean of the Gaussian distribution is 1 and the variance is derived from the dropout net. To calculate Equation (3) at each training epoch, we need to sample a weight from the Gaussian distribution and then multiply this weight with the embedding’s from the main net. This technique is called Heteroscedastic Gaussian Dropout (Lambert et al., 2018) which is a natural extension of the Gaussian dropout in which the variance of Gaussian distribution is a constant. The intuition is that for an important hidden neuron, which captures the feature patterns within a local network region, tends to get a Gaussian distribution with a smaller standard deviation because its exact value is necessary for the final prediction. For a less important hidden neuron, the RGCN allows its value to be flexible within a certain range based on their privileged features.

To the best of our knowledge, our method is the first to adopt this technique on GNNs. A naive implementation of Equation (3) is to generate different variance values for different dimensions of the node embedding’s. However, such implementation significantly increases the risk of overfitting by introducing extra model complexity. To address this problem, we consider using a shared variance across all the dimensions for each node, then Equation (3) becomes,

$$v_i = m_i(x) + N(1, \sigma^2)$$

Where $g(\cdot)$ is an aggregation function that can output a summary statistic, such as sum, mean or median.

2.3 Training via variational graph auto-encoders

Equation (3) provides a way to calculate the value of each hidden neuron. However, it is hard to calculate its gradient and propagate it back to train the parameters in the neural networks. Therefore, we formalize the entire model as a Graph Variational Auto-Encoder (VAE) (Simonyov and Komodakis, 2018). Let $X$ denote the feature matrix by collecting all the disease and gene nodes and $A$ the adjacent matrix of the disease–gene association graph. We optimize the variational lower bound of the log likelihood with respect to the parameters was follows,
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\[ L = E_{q(Z|X,A)}[\log p(A|Z)] - KL[q(Z|X,A)||p(Z)] \]  

(5)

Here, \( q(Z|X,A) \) is the encoder function transforming the input feature \( X \) into a latent variable \( Z \) on the low dimensional space, which is defined as a Gaussian distribution as follows,

\[ q(Z|X,A) = \prod_i q(z_i|X,A) = \prod_i N(z_i|x_i, \text{diag}(\sigma^2_i)) \]  

(6)

The mean \( \mu_i \) and variance \( \sigma^2_i \) of the Gaussian functions are the main net and dropout net RGCN models with the form defined in Equation (2). That is, the hidden variable \( Z \) is sampled from a multi-variate Gaussian distribution in which the mean and variance are derived from two RGCN models. \( p(A|Z) \) is the decoder function which transformed the hidden variable \( Z \) sampled from the encoder function back to the adjacent matrix \( A \) of the disease gene graph. \( p(Z) \) is the prior distribution of \( Z \) following the Gaussian distribution defined as follows,

\[ p(Z) = \prod_i N(z_i|0, I) \]  

(7)

Note that the hidden variable \( Z \) in the first term of Equation (5) is the same as what we defined in Equation (3). The second KL divergence of Equation (5) restricts the posterior distribution to be close to the prior distribution we set beforehand, which acts as a regularization term to prohibit overfitting. Different from conventional VAE, the mean and variance are two RGCN models instead of one CNN or Multilayer perceptron. In addition, the features used to calculate the mean and variance are different. The mean is calculated by the main net using the public features and the variance is calculated by the dropout net using the privileged features. To train the model, we adopt the reparameterization trick (Kingma and Welling, 2013) to help the gradient back-propagate properly, which is the same as used in VAE. Please refer to (Xu et al., 2018) for more technical details for training the VAE. To optimize \( L \) defined in Equation (5), we adopted the ADAM algorithm with learning rate \( 1 \times 10^{-4} \).

2.4 Link prediction

After the model is trained, we only need to calculate the hidden variable \( Z \) from using the main net and no dropout net or privileged features are involved at this step. The probability for disease \( j \) to be associated with gene \( i \) is defined as:

\[ s(x_i, x_j) = \text{sigmoid}(<m(x_i), m(x_j)>) \]  

(8)

where \( m(x_i) \) and \( m(x_j) \) are the node embeddings of gene node \( i \) and disease node \( j \), respectively. Link prediction is a binary classification problem given two nodes, if there is an edge between them, we denote the left variable as 1 and 0 otherwise. Therefore, we view a disease–gene association as a positive sample and no association between a disease and gene as a negative sample. This disease gene prediction task is a highly imbalanced problem, in which there are overwhelmingly more negative samples compared to positive samples. In practice, the disease gene associations only occupy 0.5% of all the possible links. To address this problem, we did negative sampling, and each positive sample has one negative sample from the entire graph that will be included in our objective functions.

2.5 Heterogeneous network

The heterogeneous disease gene network contains three types of edges: gene similarity, disease similarity and disease–gene associations.

Gene similarity. Gene similarity network is extracted from the HumanNet database (Lee et al., 2011), which characterizes the function similarity between two genes using multiple resources, including mRNA co-expression, protein–protein physical interaction and evolution information based on comparative genomics.

Disease similarity. The disease similarity network is constructed based on their phenotype similarity extracted from MimMiner (Van Driel et al., 2006) using text mining techniques. There is an edge connecting two diseases if their phenotypic similarity score exceeds 0.2.

Gene–disease association. Gene–disease association is the most important association we studied in this paper. It is constructed from the OMIM database (Hamosh et al., 2005), in which data were collected and edited at Johns Hopkins University with input from scientists and physicians around the world. In total, we include 3215 diseases, 12 331 genes, 321 375 disease–disease edges, 366 918 gene–gene edges and 3988 disease–gene edges. All the edges are undirected edges.

2.6 Features

We adopted the same features and data pre-processing pipeline as described in (Natarajan and Dhillon, 2014) for both genes and diseases. In particular, we collected two types of features for genes:

Gene expression. We extracted the gene expression measured by the microarray experiments in different samples from the BioGPS database (www.biogps.org) and the Connectivity Map (www.broadinstitute/cmap) (Natarajan and Dhillon, 2014). In particular, each feature represents the value of the mRNA expression in a particular sample in a given cell type. Two genes with similar feature vectors indicates that their expressions are similar across different individuals in multiple tissues and cell types. To remove the redundancy among highly correlated individual samples, we used principal component analysis on the feature matrix and used the first 100 eigenvectors as the feature representations.

Function associations from different species. Natarajan and Dhillon collected (Natarajan and Dhillon, 2014) the gene–phenotype association’s studies in eight species including plant, worm, fruit fly, yeast, E.coli, mouse, zebrafish and chicken. If two genes have similar phenotype features, it indicates two genes are associated with a similar set of phenotypes across different species. In total, we considered 17 480 gene features in the model.

For each disease, we also consider two sets of features: the ontology annotations from the Human Disease Ontology (https://www.ebi.ac.uk/ols/ontologies/doid) and the clinical features drawn from the OMIM webpage (https://bioportal.bioontology.org/ontologies/OMIM).

3 Results

In this section, we present the results of the performance evaluation for different models to recover the gene disease edges. We consider the other three scenarios: First, we evaluated the predicted associations for new diseases, for which all the associations with genes were removed during test and validation sets (Li et al., 2019b). In the second experiment, we tested the performance of different methods on recovering new associations between a gene and a disease, both of which have no gene–disease association in the training and validation dataset (Li et al., 2019b). In addition, we also focused on a special group of genes, named ‘singleton genes’ (Li et al., 2019b; Singh-Blom et al., 2013), which only has one association with one of the many diseases. This association will be removed from the training data and the task is to recover this association. For all these conditions, we considered two scenarios: (i) Prediction performance using all the features and (ii) prediction performance when the genomics features on the test sample are sparse. Besides, the missing features in the training, validation and test sets will be filled with zero. The purpose of our framework is not to impute the missing features of each node. Instead, it tries to alleviate the effect of missing features by applying a learnable dropout to each node so that each node can learn its representation from additional information. In addition, the gene disease network is an undirected graph and that is to say edges ‘\( i \rightarrow j \)’ and ‘\( j \rightarrow i \)’ between nodes \( i \) and \( j \) are the same edges. Therefore, we need to avoid the data leakage problem, which means that if ‘\( i \rightarrow j \)’ has been moved to test set, then ‘\( j \rightarrow i \)’ should not appear in the training set. This has been violated in (Li et al., 2019b), where they put the edge ‘\( i \rightarrow j \)’ in the test set but fail to remove ‘\( j \rightarrow i \)’ from the training set. We believe the results they reported in their paper are not reliable. Therefore, we retrained the
PGCN model (we called RGCN model in our experiment) using the same train/val/test dataset as ours.

3.1 Overall performance

3.1.1 Performance on all the genes

We used the 10-fold cross validation strategy to test each computational model. In each fold, we use 10% of the data as validation set, 10% as test set and the remaining 80% as the training set. AUROC (Provost Foster et al., 1998) and AUPRC (Manning and Schutze, 1999; Raghavan et al., 1989) were adapted to evaluate the performance of each model. Both AUROC and AUPRC are important metrics to evaluate the model performance when dealing with imbalanced prediction problems. Hence, we used AUPRC and AUROC to measure the model performance in the following sections. For the four competing methods, Katz is a similarity score-based method, where the similarity is measured through the number of walks connecting the two nodes with different lengths. Katz makes prediction that merely depend on the network structure but cannot incorporate node features into the modeling process. Hence, it is hard to make a good prediction for both genes and diseases. Catapult improves the results through supervised learning and the features used are merely network topological features. GeneHound and IMC prioritize the disease genes by modeling this problem as a recommendation system through matrix factorization. In this case, they can take full advantage of the gene- and disease-specific information (Li et al., 2019b). In this experiment, the public features and the privileged features of our method are the same and our model is equivalent to the variational dropout framework (Kingma et al., 2015; Molchanov et al., 2017). To evaluate the contribution of Heteroscedastic Gaussian Dropout, we also implemented a relation-GCN model with the exact same architecture but without using the LUPI paradigm. To further evaluate the contribution of network topology and genetic features, we also compared our methods with a RGCN model without using any features on diseases and genes.

As demonstrated in Figure 2, LUPI-based RGCN models outperform other models by at least 8.8% in AUROC and 7.0% in AUPRC. In comparison to the simple RGCN model, LUPI-based RGCN model could achieve 11.7% of improvement in AUROC and 12.4% in AUPRC. This demonstrates that even though we could collect all the features easily for all the genes, our framework is stable predictions (Kingma et al., 2019). finding that variational dropout could provide more accurate and more robust compared to other models by introducing a feature-de-

3.1.2 Performance of genes with sparse features

Due to technical limitations and high experiment cost, a significant amount of genes contains sparse features. As shown in Figure 3, all the genes in our dataset contain less than 30% of non-zero features and 27.7% of the genes only have less than 5% of non-zero features. Note that sparse features is a very common phenomenon in biological applications especially in system biology (Kuenzi et al., 2020; Ma et al., 2018; Severson et al., 2017; Yu et al., 2016). In this section, we specifically examine the performance on this subset of genes with extreme sparse features. We use all the gene features as the public features as well as the privileged ones to train the LUPI-based RGCN model.

Overall, the results show that most of the model performance decreases compared to the performance on all the genes. For LUPI-based RGCN method, the feature-dependent Gaussian dropout, to some extent, alleviates this sparse-feature problem (Figure 4). Our LUPI-based model outperforms the second-best model by 10% in AUROC and 8.7% in AUPRC, respectively. This improvement is larger than the one achieved when compared on all the genes. This result suggests that a better place to deploy our model is when the feature is sparse and Heteroscedastic Gaussian Dropout could more efficiently select the important features by reassigning the variance of each feature dimension.

3.1.3 Performance for different numbers of features

We further analysed how the performance changes with respect to different feature sparsity. We randomly sampled k% of gene features in the test dataset and used all the features in the training dataset. Note that only our model allows the features for training data and test data are different and RGCN-based models could take genetic features. Therefore, we compared with the RGCN-based model and filled the missing feature as 0 in the test data. For each k, we repeated this sampling process ten times. In this experiment, we applied the F1-score to measure the performance, which is an integration of both recall and precision:
3.2 Performance for singleton genes

From the previous studies, we know that the main signal of the prediction comes from the topological structure of the disease gene network. Next, we evaluate the prediction performance on another set of genes on which genetic features play more important roles in comparison to network structures. We evaluated the prediction performance on recovering gene and disease associations for singleton genes. Singleton genes are those genes which are only associated with one disease, so their network topology information is weak. In the dataset, we found 577 singleton genes. The evaluation metric is called recall@K, which is also used in (Li et al., 2019b). Recall@K suggests the probability of an actual association being retrieved when checking the top-K predictions and is frequently used in the context of recommendation systems (Herlocker et al., 2004), whose target is to recommend top-N items to the user. In disease gene prioritization, recall@K is also an important indicator because the top-ranked genes are candidates for further investigation. As shown in Figure 6A, the results show that the LUPI-based RGCN model outperforms the other baseline models. When K is small, the performance of Katz, RGCN and LUPI-based RGCN model are almost the same. When K increases, our LUPI-based method starts to significantly outperform other baseline methods. In particular, our method outperforms the RGCN model without genetic features 17.3% when considered the top 10 candidate genes, suggesting our model could better combine the information provided by genetic features when the network topology signal is weak. We further investigate the performance of different models for different number of features in test data. That is, the training and test features are different only for these singleton genes. As demonstrated in Figure 6B, our method is constantly better than the RGCN. When using more features in the test data, the performance can be improved in both LUPI_RGCN and RGCN model, suggesting the advantage of using Heteroscedastic Gaussian Dropout.

3.4 Performance for new associations

The prediction for new association is the most difficult task compared with the previous two machine learning tasks, because neither the disease nor the gene of the association has been seen in the training set. As a result, it requires the model to learn a representation that understands well why there is a link between certain genes and diseases. As shown in Figure 8A, there is a clear performance drop in the recall value compared to the previous two experiments. Katz and Catapult cannot recover any associations within the top 10 predictions. The reason is that Katz and Catapult merely rely on the network topology information. However, in this setting, the missing associations significantly affect the topological structure of the graph. In this case, our method can better use the node features to help to learn an expressive embedding and improve the performance in recovering novel associations. As LUPI-based RGCN models...
4 Discussion

In this work, we formalized the gene disease prioritization task as a link prediction problem, which considers both the similarity of network topology and genetic features. We designed a RGCN version of LUPI framework which could integrate these two information sources in a more efficient way. By introducing an extra neural network model to control the dropout variance, we showed that our model could better leverage these genetic features in various conditions such as sparse features, singleton genes, new diseases and novel associations. In particular, our model could achieve stable performance improvements even when the public features and privileged features are the same.

In this work, we mainly focused on the disease gene prediction, we view our computational framework as a very general learning paradigm which could be easily adapted to other biology applications where collecting certain genetic features is hard. For example, in cancer study, it is very common to train supervised learning models to understand the relationship between rich genomic and genetic features and tumors’ phenotypes (Ing et al., 2017; Lu et al., 2019; Nevins and Potti, 2007; Warnat et al., 2005). However, such models are very hard to deploy in clinics as it is typically hard to measure the same set of features for real patients. Our LUPI framework could be easily applied here to bridge this gap.

Nowadays, there are a lot of interests in studying the interpretation of GNNs. Under the LUPI framework, the interpretation task is not only to understand the behavior of the main GCN but also to understand the behavior of the dropout GCN and their interactions. This requires non-trivial extensions of current GCN interpretation frameworks such as GraphLIME (Huang et al., 2020), GNNeXplainer (Ying et al., 2019) and Xggn (Yuan et al., 2020). In the future work, we will focus on generalizing such interpretations frameworks to interpret our LUPI predictions.

Financial Support: none declared.

Conflict of Interest: none declared.

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