Abstract—A pharmacological effect of a drug on cells, organs and systems refers to the specific biochemical interaction produced by a drug substance, which is called its mechanism of action. Drug repositioning (or drug repurposing) is a fundamental problem for the identification of new opportunities for the use of already approved or failed drugs. In this paper, we present a method based on a multi-relation unsupervised graph embedding model that learns latent representations for drugs and diseases so that the distance between these representations reveals repositioning opportunities. Once representations for drugs and diseases are obtained, we learn the likelihood of new links (that is, new indications) between drugs and diseases. Known drug indications are used for learning a model that predicts potential indications. Compared with existing unsupervised graph embedding methods, our method shows superior prediction performance in terms of area under the ROC curve, and we present examples of repositioning opportunities found on recent biomedical literature that were also predicted by our method.

Index Terms—Drug Repositioning, Graph Embedding

I. INTRODUCTION

Drug repositioning (aka repurposing) can be defined as renewing failed drugs and expanding successful ones by developing new therapeutic uses that are beyond their original uses or initial approved indications. Repositioned drugs account for approximately 30% of the US Food and Drug Administration (FDA) approved drugs in recent years [1]. A repositioned drug uses de-risked compounds, going directly to preclinical testing and clinical trials, thus providing inexpensive alternatives to the costly pipeline associated with the development of new drugs. One of the well-known examples is sildenafil citrate (brand name: Viagra), which was repositioned from a common hypertension drug to a therapy for erectile dysfunction [2].

Figure 1 illustrates the biochemical interaction that gives rise to the pharmacological effect of a drug. This paper is motivated by the problem of finding drug repositioning opportunities by modeling the mechanisms of action of drugs [3]. For instance, different biological solutions might be considered in order to chemically decrease the blood pressure such as removing the excess of salt from the body, thereby decreasing the tension in the vessels, or inhibiting the vasoconstrictive signalling of a hormone, or acting directly on the cells physically narrowing the vessels and preventing their unwanted action this way [4]. Each of the aforementioned solutions requires a different mechanism of action. The same drug can have several mechanisms of action and therefore it can potentially play a multitude of roles by perturbing proteins involved in various biological processes, which are accountable for the drug polypharmacology [5]. Thus, drug repositioning is a direct application of drug polypharmacology [6].

Our main goal is to discover new relations between current drugs and diseases by utilising existing public drug-disease-protein interactions. The main three steps of our proposed method are as follows. In the first step, we built a large and heterogeneous graph comprising drug, disease, and protein entities that are linked according to information collected from the biomedical literature, as shown in Figure 2. Specifically, we formulate the drug repositioning problem as a three layer multi-relation directed graph $G = (V, R, E)$, where $V$ is the set of entities (i.e., drugs, diseases and proteins), $R$ is a set of relations (i.e., drug-protein, drug-disease and protein-protein), and $E$ is a set of edges connecting different entities in $V$. In the graph, mechanisms of action are represented by relations involving drugs and proteins and repositioning opportunities are represented by (hidden) relations involving drugs and diseases. The graph also contains protein-protein interactions in order to increase connectivity and information propagation while learning node representations. The datasets used to build the graph are described in Section III.

In the second step, our goal is to find a low-dimensional latent representation for drugs and diseases, so that the latent representation embeds the relationship between mechanisms of action and drug indications. Drug-protein and drug-disease

![Fig. 1: A chemical is assigned as a treatment to a disease because it exhibits a particular mechanism of action that affects biological processes associated with the disease. Repositioning opportunities exist because the same drug perturbs multiple proteins themselves involved in multiple biological processes.](image-url)
interaction graphs usually exhibit a particular structure with many isolated sub-graphs, and often a protein is linked to drugs residing in different parts of a graph. We employ a SkipGram based algorithm [7] to learn node representations in an unsupervised way, but instead of performing deep random walks to produce contexts [8], [9], we employ a restricted number of permutations over the immediate neighborhood of a node as context to generate its representation [10]. This choice is motivated by the particularly sparse structure of drug-disease and drug-protein interaction graphs. Further, we exploit the multi-relation nature of the graph by employing two types of contexts while learning node representations: contexts composed of drugs and proteins (i.e., mechanisms of action) and contexts composed of drugs and diseases (i.e., drug indications). This results in an embedding for each drug and for each disease, so that adjacent entities are placed close to each other in the vector space, while unconnected entities are pushed apart. As a result, drugs and diseases that have a similar distribution of neighbors will end up being nearby in the vector space. Details for obtaining node embeddings can be seen in Section IV.

In the third step, we learn the likelihood of new links between drugs and diseases, as representations for drugs and diseases were obtained in previous step. Known drug indications are used for learning a parametric model which predicts other likely indications. Our evaluation follows the typical cross-validation framework, in which a subset of the known drug uses are hidden. Details of the experimental setup can be seen in Section V and the results obtained by different embedding algorithms we used can be seen in Section VI.

In the following we briefly summarize our contributions:

- We employ interaction graphs involving drugs, diseases and proteins in order to learn suitable vector representations for drugs and diseases. The input graph presents particular characteristics, such as high sparsity and low connectivity, so that contextual information based on the immediate neighborhood is likely to produce better representations than typical random walk approaches.
- Given the vector representations for drugs and diseases, we build a parametric model learned to identify if a specific drug is indicated to a specific disease.
- We evaluate the effectiveness of our model on predicting repositioning opportunities under a cross-validation framework. Our model reaches an area under the curve of +0.98, being significantly superior than predictive models built using contextual information produced by deep random walks.
- Finally, we compared our specific findings with reposition opportunities reported in the recent biomedical literature. We present some interesting cases that were predicted by our model, including the use of Amitriptyline for relief of Fibromyalgia in adults. Amitriptyline is an antidepressant that is now being reported in the literature to treat Fibromyalgia if used at doses below those at which the drugs act as antidepressants.

II. RELATED WORK

Several strategies have been proposed for drug repositioning: (i) structure-based, (ii) repositioning based on transcriptional signatures, (iii) ligand-based, and (iv) network-based. Structured-based methods follow the idea that similar proteins have similar functionality. Accordingly, this similarity comparison can be used to find secondary targets of already existing drugs [11]. Molecular transcriptional signatures can be compared to create relations between drugs and new disease indications. These relations provide useful information for finding new uses of known drugs [12]. Ligand-based approaches are based upon the concept that similar compounds tend to have similar biological properties. In drug repositioning, this method has been widely used to analyze and identify the activity of ligands for new disease indications [13].

In recent years, machine learning has been vastly employed in drug repositioning. Authors in [14] used a fully connected deep neural network for training the model using transcriptional data at gene level to predict drug therapeutics and to use them in drug repositioning. They analyzed the confusion matrix and found out that the misclassified cases can indeed be considered as an indication of their potential in novel uses. Donner et al. [15] proposed ligand-based approach based on the learning of embeddings of gene expression profiles using deep neural networks and considered it as a measure of compound functional similarity for drug repositioning. Hu and Agarwal [16] created a drug-disease network using publicly available gene expression.

Graphs are the typical structures used to model the relationships between drugs and diseases. The major challenge is to find a way to incorporate complex structures like graphs into the existing machine learning algorithms. Thereby, several models were proposed for learning a low dimensional representation of graphs known as graph embedding [8], [17]–[19]. In order to model the polypharmacy side-effect, Zitnik et al. [20] trained graph convolutional neural networks, with proteins and drugs as nodes and drug-protein and drug-drug interaction as edges. Deepika and Geetha [21] used node2vec [8] representations along with bagging Support Vector Machine (SVM) to predict drug-drug interactions. Gao et al. [22]
TABLE I: Basic statistics of the data.

<table>
<thead>
<tr>
<th></th>
<th>drug-protein</th>
<th>drug-disease</th>
<th>known indications</th>
</tr>
</thead>
<tbody>
<tr>
<td># of drugs</td>
<td>584</td>
<td># of drugs</td>
<td>600</td>
</tr>
<tr>
<td># of proteins</td>
<td>16,546</td>
<td># of diseases</td>
<td>508</td>
</tr>
<tr>
<td># of interactions</td>
<td>1,824,204</td>
<td># of interactions</td>
<td>2,836</td>
</tr>
</tbody>
</table>

Table I presents statistics about the data. We focused on 600 drugs that were indicated to 508 diseases, resulting in a total of 2,836 drug-disease links.

IV. UNSUPERVISED NODE EMBEDDING

In this section we aim to learn representations for drugs and diseases that best preserve the original graph structure, generalizing mechanisms of action in order to find novel uses and repositioning opportunities. Graph embedding consists in finding a continuous vector space representation for entities in the set of nodes \( V \). The task is to learn a dictionary \( Z \in \mathbb{R}^{|V| \times d} \), with one \( d \)-dimensional embedding for each node in \( V \). In other words, graph embeddings are the transformation of a graph to a set of vectors, by capturing the graph structure as well as node-to-node relationships. Unsupervised learning of graph embeddings has benefited from the information contained in contexts [10], and thus embedding methods usually work by simulating contexts and operate in two steps:

1) They sample pair-wise relationships from the graph through random walks. Each random walk generates a sequence of nodes, simulating a context.
2) They train an embedding model, e.g. using Skipgram algorithm [7], to learn representations that encode pairwise node similarities.

Embedding methods differentiate mainly on the first step, as there are many possible ways to extract context from a graph. The best strategy for producing context depends on specific characteristics of the graph. In this work the contexts are based solely on the first order neighborhoods of nodes, defined here as the nodes that are directly connected. Consequently, nodes’ representations will be mainly defined by their first order neighborhoods and nodes with similar neighborhoods (contexts) will be associated with similar representations. This results in embeddings focused mainly on the first-order proximity. More specifically, we first separate a node neighborhood in small groups and then we maximize the log likelihood of predicting a node given another in such a group [33].

A. Generating Contextual Groups

The first step is to group nodes based on their neighborhoods, so that context can be exploited. There are two main challenges in forming groups from neighborhoods, as follows:

- Nodes have different degrees, so groups containing all the neighbors from a node are difficult to treat.
- There is no explicit order in the nodes in a neighborhood. So there is no clear way to choose the order in which they would appear in a group.

To deal with these challenges, we create small groups with only \( k \) neighbors in each, using random permutations of their neighborhoods [10]. The number of permutations \( n \) is specified and controls the trade-off between training time and increasing the training dataset. Selecting a higher value for \( n \) creates a more uniform distribution on possible neighborhood groups, but also increases training time.
B. Learning Representations

The first step results in a set of groups \( S \), where each member of \( S \) is a subset of nodes in the graph. Then, we learn vector representations of nodes by maximizing the log likelihood of predicting a node given another node in a group and given a set of representations \( r \), making each node in a group predict all the others. The log likelihood to maximize is given by:

\[
\max_r \frac{1}{|S|} \sum_{s \in S} \left( \log (p(s|r)) \right)
\]

where \( p(s|r) \) is the probability of each group, given as:

\[
\log (p(s|r)) = \frac{1}{|s|} \sum v_i \in s \left( \sum v_j \in s, v_j \neq v_i \log (p(v_j|v_i, r)) \right)
\]

where \( v_i \) is a node in the graph and \( v_j \) are the other nodes in the same group. The probabilities in this model are learned using the feature vectors \( r_{v_i} \), which are then used as the node representations. The probability \( p(v_j|v_i, r) \) is given by:

\[
p(v_j|v_i, r) = \frac{\exp (r_{v_i}^T \times r_{v_j})}{\sum_{v \in V} \exp (r_{v_i}^T \times r_v)}
\]

where \( r_{v_i}^T \) is the transposed output feature vector of node \( j \), used to make predictions. The representations \( r_e \) and \( r_v \) are learned simultaneously by optimizing Equation 1. Essentially, by optimizing this log probability the algorithm maximizes the likelihood of predicting a neighbor given a node, creating node embeddings so that nodes with similar neighborhoods have similar representations [34]. Since there is more than one neighbor in each group, this model also makes connected nodes having similar representations, because they will both predict each other neighbors, resulting in representations also with first order similarities. A trade-off between first and second order proximities can be achieved by changing the parameter \( k \), which controls the number of nodes within each group.

V. Experimental Setup

Our data is a multi-relation graph compose of drug-protein and drug-disease interactions. Thus, in order to learn our models, it is necessary to select an efficient embedding space in order to better exploit the information within the graph. We discuss the choice for an appropriate embedding space which includes evaluating different graph-embedding algorithms and their corresponding hyper-parameters.

A. Learning the Embedding Space

We first find an efficient embedding space for the different node embedding algorithms that will be compared in our experiments. This involves 25 hyperparameter combinations that were randomly selected for each algorithm and embedding models are then learned in an unsupervised way. We consid- ered three graph-embedding algorithms in our experiments: (i) DeepWalk [9], where the best hyperparameters are: window size of 12, number and length of walks were set to 7 and 25, respectively; (ii) Node2Vec [8], with window size, number and length of walks equal to 5, 57 and 73, respectively; (iii) NBNE [10], with hyperparameters: window size of 6 and number of permutations set to 30.

B. Model Evaluation

We used the Multilayer Perceptron (MLP) as a binary classifier, which predicts possible links between drugs and diseases using the embedding space. Specifically, the vector of a drug and the vector of a possible indication (i.e., a disease) to the drug are concatenated, and the MLP model takes the final vector as input and makes a prediction (in this case, the output is the probability of a link existing between the drug and the disease). As shown in Table I, the known indications that form the drug-disease interaction graph contains 2,836 links. We used 5-fold cross-validation to assess the embedding’s quality. Thus, we divided 2,836 into five folds, each time one of them is used for the validation and the rest for the training. As the known indications data contains only positive examples, we have generated 30,196 negative examples using the complementary graph of the known indications in order to learn the MLP model [36], as shown in Table II. It is worth mentioning that there are more negative occurrences than positive ones in the real-world scenario, because drugs are produced for a small group of diseases or health issues, being ineffective for others. As can be seen in the table, the data is highly imbalanced, thus making our experiment close to a real-world scenario. Finally, we used area under the curve (AUROC score) as the basic measure for assessing the performance of the algorithms [35].

VI. Results

In this section we report results obtained by the three embedding algorithms DeepWalk, Node2Vec and NBNE. We also discuss examples of drug repositioning opportunities endorsed by recent biomedical literature.

**Prediction Performance:** As shown in Figure 3, NBNE has obtained the best result. Specifically, NBNE achieved numbers as high as 0.98 in terms of AUROC, while Node2vec and Deepwalk achieved 0.75 and 0.77, respectively. The improvement provided by NBNE compared to Deepwalk and Node2Vec is significant – 27% and 28% of improvement respectively. The main difference of NBNE from the other two algorithms is the context generation approach, as NBNE is based on the neighborhood while the other two algorithms are on random walks, as discussed in Section IV-A. It seems
that the neighborhood based approach generates more accurate representation in the drug repositioning scenario.

Related Diseases in the Embedding Space: We employ t-SNE in order to visualize the embedding space of drugs and diseases. T-SNE is a technique used to visualize high-dimensional data by giving each data point a location in a two-dimensional map [37]. The visualization suggests some insights about the reasons that lead to the good performance of our method. In order to have a clear visualization, the drugs are represented by triangles (blue) and the diseases by rectangles (green). Further, some points are also highlighted in the figures to demonstrate interesting properties of the embedding space. Figure 4 shows that similar diseases have close vector representations. Figure 4 (Left) shows a cluster of diseases (red points) representing pain related diseases, while Figure 4 (Right) shows a cluster of muscle related diseases. These visualizations suggest that our method generates meaningful representations as related diseases are located close to each other in the embedding space.

Diseases and their Corresponding Drugs in the Embedding Space: We have also analysed the spatial relation of diseases and their corresponding drugs. Figure 5 (Left) highlights drugs which are used to treat Breathing Difficulty. In this case, most of the indicated drugs are concentrated next to the disease. The same trend is observed in Figure 5 (Right), where we highlighted the disease HIV — again, most of the indicated drugs are placed next to the disease. These visualization give us a great insight of the embedding space generated by NBNE.

Model Explainability: Features within our embeddings are not meaningful as we do not know what each feature is actually representing. Fortunately, in order to better understand model predictions, we can still group the predictions based on the features used by the model while performing the prediction [38]. Intuitively, positive predictions involving the same disease should be located next to each other, as the corresponding drugs may share the same mechanism-of-action. We employ SHAP (SHapley Additive exPlanations [39]) in order to calculate the importance of each feature for each prediction.

Table III presents examples of repositioning opportunities. Our prediction model suggests Gabapentin as a candidate for bipolar II disorder, which has been confirmed by [43] and in several other studies. While Naproxen is used in treating balance problems, it can also be used for treating Myofascial Pain [44]. Recent research carried out by [45] shows that Amitriptyline, which has been used in the treatment of depression and Parkinson’s disease, is associated with the same mechanisms as those of other drugs used to treat Parkinson’s disease. In another example, Multiple Sclerosis (MS) and Toxoplasmosis were placed into the same cluster. According to a study carried out by [46], Toxoplasmosis should be considered as differential diagnosis of tumefactive MS. Another interesting result is the association between MS and other neurological conditions such as Amyotrophic Lateral Sclerosis (ALS). Our prediction model suggests Gabapentin as a candidate for treating ALS, which is confirmed by our model as it places these diseases in the same group. Recent studies show that fibromyalgia is associated with muscle tension and depression [45]. Recent research carried out by [47] confirms that both Amlodipine and Atorvastatin caused significant improvement in patients with high blood pressure which is in accordance with our results.
Drugs
Diseases
Pain Related Diseases

Drugs
Diseases
Muscle Related Diseases

Fig. 4: Proximity of related diseases. Left — Pain related diseases. Right — Muscle related diseases.

Drugs
Diseases
Drugs for Breathing Difficulty
Breathing Difficulty

Drugs
Diseases
Drugs for HIV
HIV

Fig. 5: Proximity of diseases and their corresponding medications. Left — Breathing Difficulty. Right — HIV. In general, drug indications and the corresponding health problems are located closely.

Fig. 6: The x-axis represents the instances and the y-axis is the corresponding SHAP explanation. Predictions that have similar explanation are placed next to each other in the x-axis.

VII. CONCLUSION

We utilize the existing drug-disease-protein interactions in form of graph structures for finding likely drug-disease interactions. Nonetheless, when using complex drug-disease-protein graph structures for drug repositioning, our main goal is the discovery of the hidden and unknown relations between the components’ interactions and finally using these unknown correlations to facilitate the complex and time-consuming process of drug discovery.

REFERENCES

