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Voodoo: Combining Bottom-up and top-down approaches through graph learning over interaction networks for drug-target-interaction prediction

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Abstract

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1 Introduction

(Wang and Kurgan, 2018)

In history, traditional remedies, that were known for their medicinal properties lead to drugs by extraction of the functional ingredients. Alternatively, characteristics and features of potential drugs were detected by accident like in the case of penicillin. More recently, biological drug targets can be found *in silico* through discovery of suitable computational predictors.

The challenge of accurately predicting drug-target-interactions (DTI) has shown its importance in the fields of drug repurposing and repositioning, and in the exploration of novel drugs and their interaction partners. Knowledge about those links between compounds and their target proteins help in an array of medical and pharmaceutical studies. Additionally, those associations can be utilized to identify disease specific targets, leading to desirable therapeutic effects.

With the rapidly growing field of machine learning approaches and their application to bioscientifical problems in the realm of bioinformatics, different kinds of data, such as long DNA sequences could be utilized for feature generation, while rapid advances were made. Almost all state of the

art models for drug-target-interaction prediction were based on the usage of neural networks with increasing size.

Only recently, the technique of graph learning was introduced by Kipf and Welling (2016) through graph convolution algorithms, and improved and altered under usage of different kernels (Defferrard et al., 2016; Bianchi et al., 2019), attention mechanisms (Veličković et al., 2017), random walks (Klicpera et al., 2018), and mixtures of both (Hamilton et al., 2017). While based on diverse systems, they can be relevant for testing distinct hypothesis for given graphs. While convolutional filters are suitable for finding patterns among the the given graph, attention mechanisms are more relevant for discovery of important regions within. Lately, graph learning approaches found application for computing compound representations for DTI prediction.

Approaches on this rather sophisticated problem can divided into topdown or network approaches (CITATION), and bottom-up or molecular approaches (CITATION). Top-down approaches take advantage of other data such as diseases (CITATION), side effects, knowledge graphs or ontologies, in order to learn representations for both compound and protein. Γry to

Need to state the problem clearly here or at end of prev paragraph

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We call these approaches "top-down" because they start with the observable characteristics induced by a drug and infer the targets based on the likely molecular mechanisms that result in these phenotypes.

On the other hand, bottom-up approaches attempt to learn from chemical properties of proteins or drugs to infer candidate drug-target interactions. For drugs, molecular structure (CITATION GraphDTA), molecular fingerprints, similarity to other drugs (See Bioinf Survey), and other molecular features may be used. On the protein side, secondary structure prediction (CITATION), contact prediction (CITATION), or simply convolution over the amino acid sequences can be used to obtain a feature representation for a given proteins. However, both bottom-up and top-down approaches to drug-target interaction prediction

replace: "contain and share some problems" with something like "have some limitations"

that are not solvable within themselves.

Following is not sufficiently precise; here, you need to clearly state the challenges faced by both approaches, ideally with references.

Thus, bottom-up approaches share the lack of ability to generalize, which we will show in later sections, and usually focus on engineering sophisticated features for the drugs, while neglecting to formulate meaningful features on the protein side. Top-down approaches lack the ability to spot small differences to cope with small differences within the drug structure and rely heavily on given data for the considered drug-target pair. The latter is not suitable for predictions on novel or unseen compounds, as e.g., data on side effects or its impact on diseases is seldom given for novel drugs.

In order to design such a feature for proteins and drugs, respectively, we make use of the interaction networks for both proteins and compounds. Drug-drug interaction networks were introduced and standardized by Ayvaz *et al.* (2015) and have been used for clinical decision support (Scheife *et al.*, 2015). Drug-drug interaction networks may give a hint on common targeted pathways. As an additional compound feature we will use semantic side effect similarity, which we will discuss later on.

Generally try to avoid pointers to "later".

Don't

Protein-protein interaction networks have shown great results in ... ((Vazquez et al., 2003), (Ackerman et al., 2019)) in granting context for molecular system biology. However, these contexts were never applied to the problem of drug-target-interaction prediction. Thus we formalized our hypotheses over these interaction graphs and will test them in the following chapters.

2 Methods

2.1 Problem Description

The issue of predicting drug-target interactions can be described quite briefly: For a given drug and a given protein we want to determine whether those interact or not. We do not differentiate between types of interaction such as activation and inhibition, and do not predict the strength of the interaction. If we additionally make the closed world assumption, i.e., assume that our knowledge is complete and all drug-protein pairs without a known interaction do not interact, we can formulate the problem as a binary classification task.

2.2 Datasets

precise number The data for the different parts of this model were obtained from various sources. Starting with the protein-protein interactions, we fetched ≈ 11000 human proteins with over 170.000 links from STRING (Szklarczyk et al., 2014). For the drug-target interactions themselves, we fetched 137.000 links from STITCH database (Szklarczyk et al., 2015). As both STRING and STITCH provide probability scores for each association, we

filtered them as advised by a threshold of 700, thus only obtaining likely interactions.

For the ontology segment we utilized PhenomeNET (Hoehndorf et al., 2011), a collection of various ontologies such as Human Phenotype Ontology (Köhler et al., 2018), Gene Ontology (Ashburner et al. (2000) and Seth Carbon et al. (2020)), Mammalian Phenotype Ontology (Smith and Eppig, 2009) and numerous others. Side effects and their links to drugs were obtained Side Effect Resource (SIDER)(Kuhn et al., 2015) and structured according MedDRA database (Mozzicato, 2009). They were mapped to PhenomeNET with aid of Phenomebrowser.net, which provides a SPARQL query endpoint for the mentioned resources.

which ontologies to cite

For comparative evaluation we used the gold standard dataset introduced by ?, which includes both drug-target interaction pairs and side effects

Eventually, we only considered proteins that had at least one link in either STITCH or STRING, and drugs with at least one side effect and one existing target. Thus, the intersection between these resources yielded 1160 drugs and 6680 human proteins for the training phase. We provide links to and methods for the necessary data in the provided Github repository.

2.3 Evaluation and metrics

As the number of drug-targets are sparsely given w.r.t. to the number of both drugs and proteins considered, the resulting training, validation and testing datasets are highly imbalanced. As there are only 22.000 links in the considered subset the ratio

$$\frac{\#dti_links}{\#drugs \cdot \#proteins} \approx 360,$$

consequently needing compensation in the computed loss function and appropriate metrics for the evaluation.

On that account, we weighted all positive drug-protein pair samples with this ratio by introducing the following loss function w.r.t. to binary cross-entropy:

$$l(x,y) = -w [y \cdot \log x + (1-y) \cdot \log(1-x)]$$
 (1)

for given prediction x and target y. We average this loss among all drugprotein pairs in the training set, leading to a stable environment for the used optimizing scheme Adam (Kingma and Ba, 2014). We implemented a 5-fold cross validation among the proteins as justified in the results section. Furthermore, we used early stopping to detect plateaus in the training process.

To assess each model, we computed the area under receiver operating characteristic-score (AUROC), F1-score (F1) and Matthews correlation coefficient (MCC) for each considered hyperparameter configuration setting.

Avoiding validation overfitting, we eventually tested the best performing model on the Yamanishi dataset as presented and discussed. Furthermore, we compare our results on this very dataset with various other cutting edge approaches to DTI-prediction.

2.4 Model

In order to build a method that incorporates both top-down and bottom-up features, we first created a model for each individually. Hence, we assemble

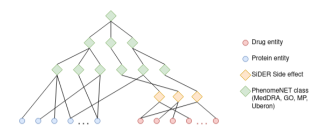


Figure 1. Drugs and proteins with annotations to SiDER and PhenomeNET

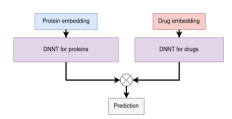


Figure 2. Siamese network applied to molecular and DL2vec features, utilizing deep neural network transformations (DNNT). The similarity function \otimes yields the similarity between both transformed embeddings e.g. by computing the cosine similarity.

rich molecular structure based features for drugs from *SmilesTransformer*(Honda *et al.*, 2019) and proteins from *DeepGOPlus*(Kulmanov and Hoehndorf, 2019). *SmilesTransformer* introduces an autoencoder, learning over the SMILES strings and thus the molecular organization of each drug in an unsupervised manner. On the other hand, we take advantage of the pretrained models of *DeepGOPlus*, obtaining features from the proteins amino acid sequence, showing significant performance in the field of protein function prediction. Thus, both embeddings seem to suitably supplement the following ontology based representations.

In the top-down section, we used *DL2vec* (Chen *et al.*, 2020) to obtain ontology based representations. Hereby, DL2vec constructs a graph by introducing vertices and edges for each ontology class and axiom, respectively, followed by random walks starting from each entity. These walks are eventually learned on using a Word2vec (Mikolov *et al.*, 2013) model. Thus, we pick up rich, neighbourhood focused representations for each entity, which has shown great results for representing protein function and phenotypes. The overall structure of the ontology can be seen in Figure 1.

As we wanted to learn from the similarity of drug side effects and protein phenotypes we opted for a deep siamese network approach, hence learning a high-dimensional embedding emphasizing this identity by forcing a maximal cosine similarity between these embeddings. On the other hand we built a deep neural network for the molecular structure based features, also benefiting from the siamese network architecture. Therefore, the precomputed embeddings are run through a deep neural network transformation (DNNT) which also reduces the eventual representation size for drugs and proteins separately. An example structure for both types of features can be found in Figure 2.

2.4.1 Graph convolutional layers

These molecular and ontology based sub-models were added to a larger graph convolutional model, based on the protein-protein interaction (PPI)

graph. The PPI dataset is represented by a graph G=(V,E), where each protein is represented by a vertex $v\in V$, and each edge $e\in E\subseteq V\times V$ symbolizes an interaction between two proteins. Additionally, we introduce mapping $x:V\to \mathbb{R}^d$ projecting each vertex v to its node feature $x_v=x(v)$, where d denotes the dimensionality of the node features.

As described before, graph convolution has shown significant performance increase in a variety of tasks. While there are various methods out there we will only introduce the most basic one here. A graph convolutional layer w.r.t. Kipf and Welling (2016) hereby consists of a learnable weight matrix followed by an aggregation step, formalized by

$$\mathbf{X}' = \hat{\mathbf{D}}^{-1/2} \hat{\mathbf{A}} \hat{\mathbf{D}}^{-1/2} \mathbf{X} \mathbf{\Theta}$$
 (2)

where for a given graph G=(V,E), $\hat{A}=A+I$ denotes the adjacency matrix with added self-loops for each vertex, D is described by $\hat{D}_{ii}=\sum_{j=0}\hat{A}_{ij}$, a diagonal matrix displaying the degree of each node, and Θ denotes the learnable weight matrix. Added self-loops enforce that each node representation is directly dependent on its own preceding one. Notably, the number of graph convolutional layers stacked equals the radius of relevant nodes for each vertex within the graph.

The update rule for each individual node is denoted by

$$\mathbf{x}_{i}' = \boldsymbol{\Theta} \sum_{j}^{N} \frac{1}{\sqrt{\hat{d}_{j} \hat{d}_{i}}} \mathbf{x}_{j}$$
 (3)

where both \hat{d}_i,\hat{d}_j are dependent on the edge weights e_{ij} of the graph. With simple, single valued edge weights such as $e_{ij}=1\ \forall (i,j)\in E$, all \hat{d}_i reduce to d_i , i.e. the degree of each vertex i. We will denote this type of graph convolutional neural layers with GCNConv.

While in this initial formulation the node-wise update step is defined by the sum over all neighbouring node representations, we are able to alter this formulation for a more sophisticated message passing scheme. As described we are able to rearrange the order of activation function σ , aggregation AGG and linear neural layer MLP with this formulation as proposed by Li *et al.* (2020a):

$$\mathbf{x}'_{i} = \text{MLP}\left(\mathbf{x}_{i} + \text{AGG}\left(\left\{\sigma\left(\mathbf{x}_{j} + \mathbf{e}_{ji}\right) + \epsilon : j \in \mathcal{N}(i)\right\}\right)\right)$$
 (4)

where we will generally only consider $\sigma \in \{ ReLU, LeakyReLU \}$. We will denote this generalized layer type as GENConv, following the notation of Fey and Lenssen (2019). While the reordering is merely import for numerical stability, the authors claim that this alteration of the original formulation eases the vanishing gradient problem for deeper convolutional networks. Additionally, the authors revise the aggregation function by generalizing it, allowing more sophisticated weighting functions such as learnable SoftMax or Power for the incoming signals for each vertex, substituting the previously mentioned averaging step. Hence, while GC-NConv suffers from both vanishing gradients and signal fading for large scale, highly connected graphs, each propagation step in GENConv emphasizes strong signals with values close to 0 and 1. Note particularly, that the same convolutional filter and weight matrix are applied to and learned for all nodes simultaneously. Note additionally, that the resulting information hold no information on their own connectivity.

We employ another mechanism to avoid redundancy and fading signals in stacked graph convolutional networks, which was introduced by Li et al. (2019) and refined in Li et al. (2020b). The authors propose a residual network architecture and normalization scheme, solving these issues in a

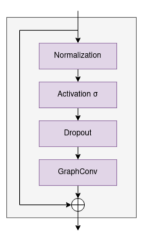


Figure 3. Residual architecture built by Li et al. (2019) and Li et al. (2020b) enabling deeper graph convolutional models

variety of graph predicting tasks. This structure is depicted in Figure 3 resulting in reusable residual blocks, which can be stacked multiple times, thereby not losing focus of each node neighbourhood.

2.4.2 Hyperparameter tuning

To find the best hyperparameter configuration for the proposed model we performed a grid search, to find the most expressive and non-redundant representation. Therefor, we pretrained the bottom-up and the top-down model separately and aimed at best performing models w.r.t. previously described metrics. Embedding sizes, depth of the neural network, optimizer, learning rate and layer types were found from an extensive, manual grid search.

3 Results

3.1 Choosing a cross-validation splitting scheme

As in machine learning inference is derived from the underlying data, models and data are naturally and intrinsically linked. Thus, the more we understand about the pitfalls and biases within the data, the more we can try to bypass these difficulties. We will hereby abbreviate, e.g., a cross validation splitting scheme as "split" determining the train, validation and test subset of a given dataset.

Especially within DTI prediction, one can observe elemental and basic skews (Pahikkala *et al.*, 2014). (1.) Frequently, novel drugs are designed by altering non-functional components of a drug, leading to two and more very similar drugs, designed to target the same proteins (CITATION?). This bias can lead to "hidden duplicates" that can distribute among the train-test split, skewing the ultimate predictive performance. (2.) Additionally, within the realm of DTI prediction, some proteins, which we will denote as *hub proteins*, have significantly more known interactions with drugs than others. In the considered subset of STITCH database, 5% of the proteins had 40% of the interactions, with similar biases for Yamanishi and Drugbank (Wishart *et al.* (2007) and Wishart *et al.* (2017)) datasets, making them easy and reliable predictors. (3.) The second distortion could be based on a publication bias for more "valuable" proteins due to their disease involvement, such as e.g., cancer, leading to more research and eventually more interacting drugs regarding those. Thus if we aim to find

a drug for a given protein, the challenge is to develop a training and evaluation scheme, that does not simply overfit to those biases.

In general, when concerning cross-validation splitting schemes for DTI prediction, there are the following three options:

- 1. Build split over drugs
- 2. Build split over drug-target pairs
- 3. Build split over proteins

where the first and third option concern splitting drugs and proteins, respectively, into train, validation and test sets, and arranging the corresponding drug-target interactions. Hereby, different training and prediction schemes lead to divergent expressiveness of the resulting model.

The most common scheme for DTI prediction is the split over drug-target pairs (Wang and Kurgan, 2018), where likely all drugs and targets of the validation and testing phase have already occurred in the training phase, as part of other drug-target pairs. Thus, the first and second splitting scheme are exposed to the first dataset bias and are hence more likely vulnerable to transductive inference by just predicting recently seen structures, rather than implementing inductive inference and generalizing over the drug representations. Second, these two strategies are more susceptible to the second and third bias, as only in these cases the model may overfit on the number of existing interactions, while in the third scheme the number of interactions of the test proteins is not known.

Considering the publications of the last

In general, drug-target interaction prediction is the task of accurately predicting, whether for a given drug and a given protein there is a biological interaction within the target organism. Hereby, different training and prediction schemes lead to divergent expressiveness of the resulting model. However, when building the train-test split over compound-protein pairs for building the actual model, there are the following three options:

- 1. Build split over drugs
- 2. Build split over drug-target pairs
- 3. Build split over proteins

In general, recent works do perform their split over the drugs or drug-target pairs ((Wang and Kurgan, 2018), CITATION). As there are hopefully many more drugs to discover, the drug split scheme both emphasizes the drug repurposing idea, by applying unseen compounds to existing targets, but also benefits from more complicated drug representations, leading to tremendous results. This performance gain is based on minor variations among large groups of pharmaceuticals, that are easy to acquire. The second scheme has knowledge on all drugs and all proteins, and is thus prone to overfitting and the same development bias. Eventually, as there only limited drug-targets (Overington *et al.*, 2006), predicting per protein is rather counter-intuitive. As it is hard to generalize over proteins representations, we aim at reaching similar performances for both drug and protein splitting schemes.

In general, recent works do perform their split over the drugs or drugtarget pairs ((Wang and Kurgan, 2018), CITATION). The first is more relevant for novel drugs, as it is much more likely to test a new compound than a innovative protein. However, it lies in the very nature of the used datasets, making the prediction for new drugs much easier. Thus, drugs are often built by minor variations of existing drugs, thus leading to no deviations in the functional group of that very compound (CITATION/EXAMPLE). When distributed over both train and test split, the models do not perform inductive inference and generalize, but rather implement transductive inference by just predicting the recently seen structures. Hence, when entirely new molecules are seen, the models perform much worse

The same applies to splits of drug-target pairs, as all drugs were already seen, and novelty cannot be coped with.

As mentioned in the introduction it is quite difficult to learn suitable features from proteins. In general, attempts search for motifs in the protein sequences under usage of convolutional neural networks and filters, which is more suitable for tasks like protein function prediction, than for for drug-target interaction prediction, and lack a more in-depth hypothesis on the protein side, while investing in refined drug features.

Thus, building splitting over proteins is the most challenging of the three options.

3.2 Baseline model

Our aim is to develop and evaluate a computational model to predict drug-target interactions. Specifically, given a protein, we are interested in identifying the drugs that may target the protein; this scenario is motivated by the need to identify drugs that can target a

3.3 Voodoo identifies drugs that target a protein

We developed Voodoo as a computational model to predict drug-target interactions. Specifically, given a protein, Voodoo will identify and rank drugs that likely target this protein. Voodoo combines two types of features: structural information for drugs and proteins that can be used to determine if they may physically interact, and

This scenario is motivated by the need to identify drugs that can target a specific protein that may be associated with a disease or abnormal phenotypes. Our aim is further to combine two different types of features: structural information based on protein structure and molecular drug composition, and their respective observable characteristics.

Before introducing the core of our eventual model, we propose a suitable naive baseline model in order to analyze and understand the existing approaches and this novel method for drug target interaction prediction. For given lists D,P of drugs and proteins, respectively, and a set of known interactions $Int := \{(d_i,p_i)\}$, we construct an interaction matrix

$$M_{int} \in \{0,1\}^{|D| \times |P|}$$
 with

$$M_{ij} = \begin{cases} 1 & \text{if } (d_i, p_j) \in Int \\ 0 & \text{otherwise} \end{cases}$$

describing for all drug–protein whether there is a known interaction or not. We now rank all proteins $p_j \in P$ descending by their number of drug interactors, characterized by

$$f:P o\mathbb{N}$$
 with $f:p_j\mapsto \sum_{i=0}^{|D|}M_{ij}$

by summing over the columns of M_{ij} and ranking these sums. We now finish our baseline predictor P_k by predicting all drugs to interact with the top k targets, denoted by TopK(P) w.r.t. M_{ij} from the previously introduced ranking, formalized by

$$P_k:D\times P\to \{0,1\} \text{ with } P_k:d_i,p_j\mapsto \begin{cases} 1 & \text{if } p_j\in TopK(P)\\ 0 & otherwise \end{cases}$$

with the only hyperparameter k. Note particularly, that the consequent prediction of P_k is not dependent on the considered drug d_i , and will thus predict all drugs similarly for a given protein p_j . Subsequently, this naive predictor is not yielding any valuable information on the individual interactions with drugs for a given protein. Also note the possibility for a similar predictor by calculating the top k interacting drugs, respectively.

To obtain a prediction for a given drug-protein pair, we first build DL2vec representations for both drugs and proteins over PhenomeNET, while also preparing structural, molecular representations. Hence, we pretrain the siamese networks for both molecular and ontology features, yielding the deep neural network transformers for both methods and both drugs and proteins, individually. These embeddings are now added to the graph neural network as node features, where we start an end-to-end learning and training process. Notably, like in the pretraining phase, we rely on siamese neural networks, enforcing similar representations for fitting drug-protein pairs. The overall architecture can be seen in Figure 4.

3.4 High level description

For a given protein, we want to find all suitable interacting drugs, **among** a set of known drugs. To obtain a prediction for a given drug-protein pair, we first build representations embodying both phenotypes and functions for both drugs and proteins, while also preparing structural, molecular representations, e.g., utilizing DL2vec over PhenomeNET and the prebuilt features from SMILES transformer and DeepGOPlus. Note that this approach can be easily generalized, profiting from other protein function and phenotype representation methods.

Hence, we pretrain the siamese networks for both molecular and ontology features, yielding the deep neural network transformers (DNNT) for both methods, and both drugs and proteins, individually. These embeddings are now added to the graph neural network as node features, where we start an end-to-end learning and training process. Notably, like in the pretraining phase, we rely on siamese neural networks, enforcing similar representations for fitting drug-protein pairs. The overall architecture is depicted in Figure 4.

Through the very nature of the graph convolutional neural network, we build the transformed representation for all proteins in every forwarding step of the model. Note particularly, that the same convolutional filter and weight matrix are applied to and learned for all nodes simultaneously. By construction, for a single drug we can compute and predict all its interactors in a single run of the model, leading to significantly less computing time.

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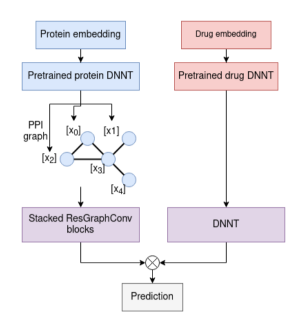


Figure 4. Full DTI prediction model based on the pretrained deep neural network transformers (DNNT) for either molecular structure or ontology based features. The transformed protein representations are added to each corresponded protein as node features for the graph convolutional steps

4 Findings

4.1 Deification of our method

- we built protein function and ontology based features based on DL2vec
- Ontology derived protein function focused features are highly predictive for dtis
- We built a versatile template for various features to test localization on the PPI graph
- normal GCNs don't work on PPI graph, as it is highly connected ightarrowneeds stronger more expressive aggregation function \rightarrow GENConv in residual blocks for better numerical stability
- protein functions localize on the PPI graph, while molecular features don't

Approach	Splitting	Original	Protein	MicroAUC
	scheme	AUROC	split	
		score	AUROC	
DeepDTI	Drugs	87.6	75.9	70.1
DeepDTA	DP pairs	87.6	76.7	69.4
DeepConv-DTI	DP pairs	88.3	76.6	73.0
MolTrans	DP pairs	89.5	77.0	74.0

• all AUROC in % AUROC score on STITCH

1	AUROC in % AUROC score on s	SIIICH	• drug indications are nightly predictive for downstream		
	DNNT model	Without graph model	With graph model	apability to differentiate highly related drugs/proteir	
	MolPred	69	698	tratified Cross validation is suitable for training.	
	PhenomeNETPred	88	92v	alidating and testing (Uselessly high AUPRC)	
	MolPred + PhenomeNETPred	89	9 3r	nicroAUC is a superior and more intuitive metric for d	

• microAUC for MolPred + PhenomeNETPred on graph is about 93+-

• and on yamanishi dataset

DNNT model	Without graph model	With graph model
PhenomeNETPred	83	84
MolPred + PhenomeNETPred	83	84.5

MicroAUC is about 83

4.2 How to insult other methods

- Only few other methods perform their split over proteins (Wang and Kurgan, 2018), DTI-CDF does it
- Running split over proteins is harder than, drug and drug protein pair split (see below table)
- this applies for both DTI prediction and drug target affinity prediction (and Saras gene-disease association)

•	using indications is like cheating, as not applicable for searching new
	drugs
•	drug indications are highly predictive for downstream tasks, but lack

- ng, but **NOT** for
 - drug repurposing

 $\xrightarrow{}$ why for each protein and not for each drug

Approach	Splitting	Original	Protein	MicroAUC
	scheme	AUROC	split	
		score	AUROC	
DTINet	DP pairs	91	84.1	67.2
DTIGEMS+	DP pairs	93	72.2	67.8
DTI-CDF	Proteins	83	83	79

- A naive predictor (ranking proteins) and predict each drug similarly achieves cutting edge performance (87.5 AUROC for whole dataset, $85.5~{\rm for}~5\text{-fold}~{\rm cross}~{\rm validation}~{\rm in}~{\rm drug\text{-}split}) \rightarrow {\rm No}~{\rm prot}~{\rm focused}$ microAUC possible. \rightarrow hub proteins
- Yamanishi Dataset is only partially suitable, if everybody just derives a suitable subset (DTIGEMS)
- This also applies to drug target affinity prediction. We were hereby able to roughly reproduce the results from MolTrans (Bioinformatics) on BioSnap

4.3 Tested hypotheses

In this work we are testing the following hypotheses:

- 1. Can we build a model that outperforms state of the art approaches, combining top-down and bottom-up approaches?
- 2. Are interaction networks sufficient to improve the performance of simple molecular predictors?

We will test the first hypothesis by building a model that takes both topdown and bottom-up features into account. Thus, we propose a novel approach to combine those mutual exclusive attempts, through the usage of interaction networks, similarity and molecular features. Additionally, we test the latter by building a simple molecular DTI predictor and enhance it under usage of the interaction networks.

For the bottom-up approach we build a model that only relies on molecular features, which we will discuss in more detail in the following methods chapter. For the combination of both approaches we now attach the predictions to the protein-protein interaction graph as node features for future graph learning steps. In this graph we tried to find both patterns and regions for each drug that could be of interest through application of different graph convolutional layers, which in return represent the feature for each protein. Representing the drug we take the drug-drug interaction graph and the semantic similarity over side effects which we will explain in the following paragraphs.

5 Methods

5.1 Models

The used model consists of two separate models, that help to fuse together the two methods:

- 1. The molecular predictor
- 2. The interaction network based predictor

We build the molecular predictor by using pretrained, molecular fingerprints models for both drugs and proteins. Regarding proteins, we used the pretrained feature generator from *DeepGoPlus* ((Kulmanov and Hoehndorf, 2019)) that was originally designed for protein function prediction and is regarded as state of the art for this purpose. For drugs we used a pretrained fingerprint model from *SMILES transformer* ((Honda *et al.*, 2019)), that provides a simple and fast method to compute fingerprints through autoencoder models. The encodings from these two models were funneled into a simple deep neural network (see Figure ??) with few fully connected.

The results of that prediction flow into the annotation of the protein-protein interaction (PPI) graph as depicted in (IMAGE). Hereby, the predictions of the molecular predictor are used as node features for the graph, with respect to the given drug. Thus, given a compound-target pair, the nodes of the PPI graph now hold bottom-up features, which can now be processed by the graph learning algorithms.

The PPI graph is processed by different graph convolutional layers, that may underline the importance of either patterns or regions within the graph, to obtain a feature vector for the wanted node. In contrast to learning over whole graphs we perform node classification within the graph. These layers are either graph convolutional layers, that learn a certain kernel over the graph, or attention based. Different layers of both and other types such as were tested.

The drug-drug interaction features are retrieved by choosing the corresponding row in the adjacency matrix of the graph, thus leading to quite simple features.

For the semantic similarity feature, that once again represents a top-down

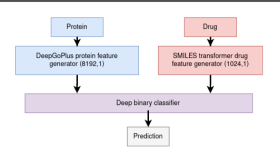


Figure 5. Molecular predictor based on the generated features from DeepGoPlus and SMILES transformer.

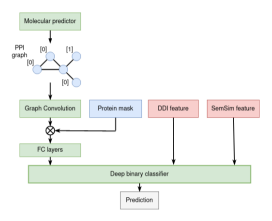


Figure 6. Deep neural network that predicts based on drug-drug interaction features and semantic similarity features over side effects for drugs, and graph convolution over protein-protein interaction networks for proteins. Protein and drug features are represented by blue and red, respectively.

attribute, we artificially link each drug to its corresponding side effects in the MedDRA hierarchy. Concerning this hierarchy, drug-drug similarity is computed by the Resnik similarity ((Resnik, 1995)). For the given compound we take the corresponding row of this symmetric similarity matrix.

Thereby, we concatenate these three features together and funnel them into another deep neural network as depicted in figure 6. This network finally yields our prediction. We hereby perform splits over both drugs and proteins, in order to test and show the discrepancy and increasing difficulty.

Implementation was done in PyTorch ((Paszke et al., 2019)) and is available on Github under github.com/thinnerichs/KAUST-dti-metabol. Graph learning methods were build with help of PyTorch-Geometric ((Fey and Lenssen, 2019)), a geometric deep learning extension library for PyTorch, that recently got a lot of attention in the machine learning community. This library gives the potential to use many state of the art graph learning mechanisms, such as plain but effective graph convolution (Kipf and Welling (2016)), Chebychev kernels ((Defferrard et al., 2016)), ARMA kernels ((Bianchi et al., 2019)), translation-invariant operators ((Verma et al., 2017)), attention mechanisms ((Veličković et al., 2017)), random walks ((Klicpera et al., 2018)) and mixtures of the latter two ((Hamilton et al., 2017)). The performance of these various layer types were tested for this particular problem, as discussed in the results section.

6 Discussion

7 Conclusion ACKNOWLEGGEMENTS

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