DT-ML: Drug-Target Metric Learning

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Abstract: The challenges of modern drug discovery motivate the use of machine learning-based methods, such as predicting drug-target interactions or novel indications for already approved drugs to speed up the early discovery or repositioning process. Publication bias has resulted in a shortage of known negative data points in large-scale repositioning datasets. However, training a good predictor requires both positive and negative samples. The problem of negative sampling has also recently been addressed in subfields of machine learning with utmost importance, namely in representation and metric learning. Although these novel negative sampling approaches proved to be efficient solutions for learning from imbalanced data sets, they have not yet been used in repositioning in such a way that the learned similarities give the predicted interactions. In this paper, we adapt representation learning-inspired methods in pairwise drug-target/drug-disease predictors and propose a modification to one of the loss functions to better manage the uncertainty of negative samples. We evaluate the methods using benchmark drug discovery and repositioning data sets. Results indicate that interaction prediction with metric learning is superior to previous approaches in highly imbalanced scenarios, such as drug repositioning.

1 INTRODUCTION

One of the main motivations for modern drug development is the discovery of new candidate compounds which can be used as medication. Developing a new drug molecule is a long and expensive process; bringing a new drug to market takes approximately 10–15 years and 1.5–2.0 billion USD (Wouters et al., 2020). One possible way to accelerate the development process is via drug repositioning. Repositioning or re-purposing refers to using a known drug in a new therapeutic application, which is a promising approach, considered less time-intensive, costly, and risky compared to de novo molecule design.

The different stages of drug development have been heavily influenced by the rise of artificial intelligence technologies in recent years. As a result of this, classical machine learning methods have become increasingly common among drug-target interaction (DTI) prediction approaches (Bagherian et al., 2021). We can reduce the cost and time required for measuring the interactions with their help. Besides, these models can later be used to estimate the interaction between an unknown protein and molecule, to search for candidates at the beginning of the development process that binds to a specific protein, or to reveal a new therapeutic application to a known drug, i.e., repositioning (Harrer et al., 2019). The simplest methods give estimates based only on the similarity of molecules (Lee et al., 2016), or treat the problem as a classification and apply neural networks (Arany et al., 2022). Utilizing matrix factorization (MF) is a common approach too (Bolgár and Antal, 2017). Still, most of the state-of-the-art (SOTA) solutions use a general version of MF, namely pairwise\textsuperscript{1} neural networks, such as the DeepDTA (Öztürk et al., 2018) or the AI-Bind (Chatterjee et al., 2021). While the AI-Bind method utilizes pre-trained representations, the DeepDTA model uses convolutional encoders to transform the SMILES representations from the molecular side and the amino acid sequences from the protein side, thus providing the latent embeddings. These are concatenated, and a multilayer perceptron (MLP) predicts the interactions. Most

\textsuperscript{1}Pairwise predictors have dual inputs, for instance, a molecule and a protein, hence their name.
of the aforementioned approaches were first applied in recommendation systems but are now considered SOTA in the field of DTI prediction too. In repositioning, the aim is not to estimate a specific interaction accurately but to establish a good disease or molecule ordering. Accordingly, several methods diverge from the traditional approach of treating interaction prediction as a binary classification and applying new loss functions better suited for ranking, such as the Bayesian Personalized Ranking (BPR) loss (Peska et al., 2017), also adopted from the field of recommendation systems.

Sufficient quality and quantity of data are necessary to apply a statistical learning approach. There are plenty of available data sets for DTI prediction tasks, but due to the high cost of interaction measurements, the sparsity of these sets is relatively high. Moreover, the number of known negative entries in drug-disease interactions is lower than expected due to publication bias, where negative results are often not published (Luo et al., 2021). Therefore, drug-disease matrices are not only sparse, but often only the positive entries are known. This is a common problem in repositioning tasks since SOTA predictors work with a loss function such as binary cross-entropy (BCE), which needs the negative samples too. One possible solution is negative sampling, but since the unknown entries can be either positive or negative, constructing a proper sampling method is challenging.

The problem of unknown negative samples has arisen in representation and (distance) metric learning too, especially in the field of contrastive learning (Le-Khac et al., 2020). The main motivation is to handle a large amount of available unlabeled data with machine learning. One way to do this is to learn representations in a self-supervised way. These embeddings can later be used in various supervised tasks if they correctly capture the underlying data distribution.

Contrastive representation learning is one of the first widely used solutions, both in the computer vision, natural language processing, and audio processing domains (Le-Khac et al., 2020). Architecturally, these methods can also be classified as pairwise, or rather, joint embedding methods, because in most cases, one input pair or triplet is compared at a time. The input embeddings are first processed by an encoder, thus creating the latent/metric representations, which are compared with a similarity function, and finally, a loss function is used to optimize the similarity between the pairs. The similarities of positive and negative pairs are maximized and minimized during optimization. Positive pairs can easily be produced with augmentation, but negative sampling is a challenging research problem. Unfortunately, using only positive pairs may lead to a collapse of the representation space since providing the same embedding for all entries can reduce the loss to zero. Therefore, negative sampling is necessary for contrastive methods.

Over the last few years, the development of different contrastive and, later, non-contrastive approaches has been an area of particular research interest. The first approaches were the energy-based contrastive loss functions, such as the Pair loss (Hadsell et al., 2006) and the Triplet loss (Collobert and Weston, 2008). They aim to associate low energy, i.e., low distance, to positive pairs and high energy to negative pairs.

A new, more effective family of methods is the probabilistic loss functions. Here, a likelihood is described by a SoftMax function with the similarity to the positive pair in the denominator and the similarity to all positive and all negative samples in the numerator. As opposed to the energy-based methods, it is not the quality but the quantity of the selected negative samples that matters since we want to approximate the denominator as accurately as possible. Therefore, we often take the samples not only from a single batch but keep their elements in a so-called memory bank over several batches. One way of sampling is using noise contrastive estimation (NCE), and a commonly used probabilistic loss function is the infoNCE or also known as normalized-temperature cross-entropy (NT-Xent) (Chen et al., 2020). A modified version of NT-Xent is called Supervised contrastive loss (SupCon) (Khosla et al., 2020). The authors of this paper performed supervised representation learning, where the labels resulted from a classification problem, and proposed SupCon, which can handle entities belonging to the same class. Another function is the circle loss (Sun et al., 2020). The novelty is that it does not increase the similarity of positive pairs and decrease the similarity of negative pairs equally but adaptively assigns different gradient weights. It does this by defining an optimum for the positive and negative similarities and then weighting each pair by the deviation from it.

Because of the efficiency problems associated with negative sampling, research in recent years has focused on non-contrastive approaches, which also aim to avoid latent collapse but do so without negative samples, for instance, the Variance-Invariance-Covariance Regularization (VICReg) (Bardes et al., 2021).

There are striking similarities between pairwise DTI prediction methods and joint embedding representation learning approaches. Namely, both utilize two inputs, from which two latent embeddings are
learned, and the output is given by comparing them. For example, the concatenation and MLP, mentioned by the DeepDTA and AI-Bind models, can be considered a special similarity function with trainable parameters.

This analogy is also true for the MF approaches, where the similarity function is a simple dot product. In the Metric Factorization (Zhang et al., 2018) collaborative filtering method, the idea appeared that during matrix factorization, we map users and products into a common space, and the similarity in this space is the prediction. But this was exploited only to the extent that instead of scalar multiplication, Euclidean distance was used. The first successful approach that combines the two is called Collaborative Metric Learning (CML) (Hsieh et al., 2017), which uses the Weighted Approximate-Rank Pairwise Loss (WARP) (Weston et al., 2010). The loss uses triplets of a user and a positive and negative item and gives weights to the triplets proportional to the approximated rank of the positive sample item in the row of the given user, i.e., those who are further back in the row are penalized more. This way, the WARP is better suited for ranking than the BCE loss, the authors have also tried the BPR ranking-based loss, but the WARP was found to be superior.

To the best of our knowledge, novel negative sampling solutions developed in the field of metric learning have not yet been applied to drug repositioning or DTI prediction this way, namely by treating the learned similarity as a predicted interaction. However, we have seen that ideas that have worked in the collaborative learning field are adopted sooner or later by interaction prediction methods. This would be particularly useful for drug repositioning because metric learning-based approaches provide better solutions to the problem of negative samples than the current repurposing methods, mainly using BCE loss with negative sampling. Some approaches use BPR, which is better suited for ranking, but the novel contrastive and non-contrastive loss functions have not yet been utilized to predict interactions. In current approaches, representation learning is only used in the pre-training phase, e.g., to learn node representations on a multimodal, heterogeneous knowledge graph. Later these embeddings are concatenated and used in interaction prediction tasks with a BCE loss function (Li et al., 2022), or the adjacency matrix is reconstructed from them (Chen et al., 2022).

To this end, we propose a drug-Target Metric Learning (DT-ML) approach that combines the two methods. In this paper, different metric learning-based methods are utilized and examined by their applicability to interaction prediction and drug repositioning. According to the results, among the various DT-ML approaches compared, the ones using probabilistic loss functions have proven superior, even better than the current SOTA. Additionally, we propose modifying one of the used loss functions, which could further improve the results.

2 METHODOLOGY

An overview of the DT-ML methodology is shown in Figure 1, detailing the data sets, architecture, similarity and loss functions, and metrics used in the evaluation.

2.1 Data and Representations

We utilized two widely used benchmark data sets to evaluate our models, namely KiBA (Tang et al., 2014) and ChEMBL (Gaulton et al., 2017). The former is a DTI data set, with interactions between molecules and proteins and known negative entries; the latter is used for repositioning, as it contains drugs with only positive indications of human conditions, i.e., associated diseases. In addition, a third data set was used to produce disease representations, namely the DisGeNET (Pihéro et al., 2016), which contains relationships between diseases and genes.

The KiBA set contains 467 kinase proteins, and their interactions with molecules are given with a dissociation constant (pK$_d$). After preprocessing the compounds, we retained only those for which the canonical SMILES descriptor is known, unique, and contains no more than 100 non-hydrogen atoms, yielding 50,418 molecules. We discretized the interaction data with a threshold of pK$_d$ = 3, as suggested by the authors of DeepDTA. This resulted in 72,944 positive and 162,681 negative entries; thus, the density of the interaction matrix is ~1%.

Among the many tried protein representations, the 512-dimensional CPCProt (Lu et al., 2020) proved to be the best. On the compound side, the pretrained, 300-dimensional Mol2vec embedding (Jaeger et al., 2018) gave the best results and was also the most efficient to work with, thus was chosen for all subsequent work.

Another data set we used is ChEMBL. It contains drug-like bioactive substances that are already FDA-approved or are in clinical trials, and associated indications as Medical Subject Headings (MeSH) (Lipson, 2000). MeSH is a controlled vocabulary of MeSH terms that is widely used in the literature, e.g., the AI-Bind model also uses it.

2Utilizing Mol2vec on the input is widespread in the literature, e.g., the AI-Bind model also uses it.
life science concepts and terms also used in the literature. The part of the data set we use has 21,042 known positive relationships between 4,755 drugs and 1,168 diseases, resulting in a density of 0.3789%.

To represent molecules, we used the Mol2vec. To obtain disease embeddings, we utilized the DisGeNET data set, which contains 1,134,924 positive entries between 30,170 diseases and 21,666 genes, giving a density of 0.1736%. Although there are no well-established methods for representing diseases as there are for proteins, a simple disease embedding can be easily obtained in a semi-supervised way based on the known disease-gene associations. We used truncated singular value decomposition (SVD) to convert the 21,666-dimensional sparse vectors into 64-dimensional dense, gene-based disease representations, later referred to as SVDDis. MeSH concepts map diseases from each column in the ChEMBL matrix to a row in DisGeNET, thus we can use SVDDis embeddings to represent diseases in the repositioning data set. Another possible option is to use one-hot representations, but this gave worse results, and without the embeddings, the model is no longer able to give predictions for new diseases.

2.2 Pairwise DTI Predictor

After preprocessing the data, we implemented a pairwise model using the PyTorch package.

We used the previously mentioned Mol2vec, CPCProt, and SVDDis embeddings as inputs. After scaling them on the training data, these embeddings are further transformed by two encoders, thus creating latent representations. We concluded that the method is less sensitive to the hyperparameters of the encoders. After trying several combinations, we finally chose two-layer MLP modules, with 512-dimensional hidden, and 256-dimensional output layers. Between layers, a Rectified Linear Unit (ReLU) activation and 20% dropout rate were used.

We defined fixed and trainable similarity functions to obtain a prediction of a given interaction from the metric embeddings and to measure the similarity between the entities. As a fixed similarity, we have tried Manhattan, Euclidean, mean squared, dot product, and cosine similarities, among them, the latter proved to be superior. Several trainable similarity functions were tested too. We found that instead of concatenation, it is better to take the Hadamard product and use an MLP module with a sigmoid activation to obtain the predictions. We used a multilayer perceptron with two, 256, and 128-dimensional hidden layers and a dropout with a rate of 10%. We refer to this later as the weighted dot product (WDP) similarity.

Figure 1 shows the architecture of the pairwise model and the used similarity functions. The model can be used with BCE loss function as a simple pairwise DTI predictor, or even with different loss functions according to the DT-ML approach.

2.3 Loss Functions

In our study, we tested several loss functions and negative sampling strategies.

As a ranking-based baseline, we implemented the WARP and used it with a margin of 0.1.

In the other baseline approaches, we have used BCE loss with sampling. The simplest approach is random sampling, in this case, we tried different ratios, but we found it best to sample twice as many negative samples as the number of known positives. This approach is later referred to as BCE random.
We tested the closed-world assumption, where all the unknown entries are assumed to be negative, in this case, the models worked with a fully completed matrix. We refer to this as BCE all.

This method is inefficient to use on the KiBA data set due to the number of possible interactions. On the other hand, in the KiBA data set, there are known negative interactions too, which can be utilized instead of sampling. In most cases, we discarded the negative entries of the KiBA set and used negative sampling just as with the ChEMBL data set so as not to compromise comparability, but we kept one case where we used the known negatives (BCE true).

We have also examined sampling during training, here, negative samples were given by unknown samples within a batch (BCE batch).

To improve the results, we weighted the positive and negative terms in the BCE loss. The weights are inversely proportional to the proportion of positive and negative samples in the data set. So even though there are more negative samples, they are taken into account with less weight, this way, we can express the uncertainty in the noisy negative sampling.

The above-listed baseline approaches represent the current SOTA, which we compared with several metric learning-inspired loss functions. Compared to the previous BCE and WARP functions, one of the main differences is that DT-ML methods are not only able to compare molecules with targets, but they also utilize molecule-molecule and target-target similarities in a semi-supervised way. This way, molecules, and targets are represented in a common latent space, and here the same similarity function is applied to compare the different types of modalities with themselves and with each other.

During the optimization of the DT-ML models, only the interacting molecule-target pairs within a batch are considered positive, negative pairs are sampled from the various possible molecule-molecule, molecule-target, and target-target combinations.

We have tried all loss functions implemented in the PyTorch Metric Learning framework (Musgrave et al., 2020). Of the several fixed and learnable similarity functions we tried, cosine proved to be the best for these approaches.

First, we examined energy-based loss functions, such as pair and triplet loss. We used them with a margin of 0.2, which we found to be optimal. The quality of the negative samples is a significant factor in using energy-based functions, hence it is important to select useful samples. With triplet loss, we only use triplets in which the positive pair has a greater similarity than the negative, but the difference between them is less than the predefined margin.

Among the tested probabilistic loss functions, NT-Xent, SupCon, and Circle losses were in the top three. For NT-Xent, a temperature hyperparameter of 0.01 was found to be optimal, and a memory bank capable of containing 512 interactions was used to further improve performance. SupCon can handle entities belonging to the same class better than previous loss functions. Indeed, when considering molecules as classes, we obtained better results. This means that targets binding to the same compound are forming positive pairs in the given batch. The temperature parameter was set to 0.01 for SupCon too. The best results were obtained with the Circle loss function. Besides the temperature hyperparameter, it has two optima and two margins for the positive and negative pairs, but for simplicity, the authors have used only one m hyperparameter to define them. Over the various investigated combinations, m = 0.4, γ = 10 proved to be optimal. Because of the uncertainty of the negative samples, we propose a modified version where the positive and negative samples have separate hyperparameters, m_p and n, respectively. With our Circle loss function, we gave negative samples a softer margin parameter of m_n = 0.6, and positive samples a harder margin of m_p = 0.3, this way, we were able to achieve further improvements.

Finally, we examined methods that do not require negative sampling at all, such as VICReg. This worked best when the weights of the variance, invariance, and covariance loss terms were equal.

2.4 Evaluation Methods

To evaluate the approaches mentioned above, we utilized a row-wise train-test split with 5-fold cross-validation. This way, the test data matrix contains only rows/molecules which were not included during training, but all the columns/targets used in the evaluation were seen in the training data too. We used five metrics in total to compare the various methods.

One of them is the area under the receiver operating characteristic curve (AUROC), which is frequently used to evaluate binary classification tasks.

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3One possible hypothesis is that these embeddings carry information about binding sites, individuals that share a common or related binding site will be close in the latent space.

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4The intuition behind this is the previously mentioned binding site analogy, as a protein or the proteins associated with one disease contain – on average – far more binding sites than the number of molecule substructures matching different sites. Thus, there is a high probability that proteins that share binding molecules will have a common binding site, so their representations should be similar indeed.
hence widespread in interaction prediction too. It only makes sense to use this metric with the KiBA data set, because the ChEMBL does not have any known negative entries. We calculated the AUROC values on the test columns, which had at least 50 positive and 50 negative entries in the whole KiBA data set, after that we took the column-wise average.

We also used four ranking-based metrics because, in repositioning, the order of the predicted interactions matters more than the actual predicted values of the interactions. To this end, we calculated the average precision@10 (later referred to as PREC) of rows in which there were at least ten entries among the test set, and the mean recall@50 (REC) value over rows in which there were at least 5 entries. We also used the Mean Reciprocal Rank (MRR) and the Mean Average Precision (MAP).

These values were calculated both at row and column levels. This was necessary, because, on one hand, most often, we are not looking for diseases for a known drug, but rather vice versa, and on the other hand, this way we can better detect overfitting.

3 RESULTS

We ran our models on a 32GB NVIDIA Tesla V100 GPU. Among the optimization algorithms tried, Adaptive Moment Estimation (Adam) proved to be the best, using an L2 weight decay with a weight of $10^{-5}$ and a learning rate of $5 \times 10^{-5}$. After a Xavier weight initialization, we trained the models over 24 epochs in the case of the KiBA, and over 128 epochs in the case of the ChEMBL data set, we used a batch size of 256 in both cases. Finally, we evaluated the aforementioned approaches according to the classification-based and the four ranking-based metrics.

On the KiBA data set, according to the AUROC metric, the SOTA BCE true approach outstandingly outperformed all the other methods, which is not surprising, since it is the only one using the known positive and negative entries as well. With the WDP and Cosine similarities, it managed to reach 0.7851 and 0.7391 AUROC respectively, which are the highest achieved values among all methods for both similarity functions.

Considering the ranking-based metrics, the results on the KiBA data set are shown in table 1 while the results on the ChEMBL set can be seen in table 2. Although BCE loss trained on the known negatives is still the most suitable for classification, some of the DT-ML approaches perform better for repositioning. Among them, the energy-based and the non-contrastive loss functions achieve poor scores, while the probabilistic methods perform particularly well, even better than the SOTA.

4 CONCLUSIONS

We have seen the challenges inherent in drug discovery and how deep learning-based interaction prediction and repositioning, can accelerate the development process. Most of the SOTA repositioning approaches utilize a DTI predictor, which needs both positive and negative entries to train. However, negative results are often not published, thus there is a shortage of negative samples among drug-disease interactions. We have also seen that in recent times negative sampling has been the main challenge in a subfield of machine learning with utmost importance, namely in metric learning too, and the attention invested in researching this area has led to a number of effective solutions.

There is a slight imbalance in the comparability of models. Much more iterations were performed during one epoch for the BCE all approach, and methods using the weighted similarity module have more trainable parameters. In these cases, baseline methods are better according to some row-based metrics. However, similar performance can also be achieved by using DT-ML methods with more parameters or more epochs.
Table 1: Row- and column-wise results on the KiBA data set, the best two methods are highlighted for each ranking-based metric. The first four rows contain the baseline, SOTA methods trained with the WDP similarity module, below them, there are the baseline and DT-ML approaches with our modified Circle loss at the bottom, with these methods, the cosine similarity was used.

<table>
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<tr>
<th>Loss function</th>
<th>BCE true</th>
<th>BCE random</th>
<th>BCE batch</th>
<th>WARP</th>
<th>Triplet</th>
<th>Circle</th>
<th>NT-Xent</th>
<th>SupCon</th>
<th>VICReg</th>
<th>our Circle</th>
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<tr>
<td>our Circle</td>
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<td>0.5401</td>
<td>0.6112</td>
<td>0.5669</td>
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The major contribution of this study is using these novel, metric learning-inspired approaches as pairwise DTI predictors in the domain of drug repositioning. We showed that DT-ML methods, which to the best of our knowledge have not yet been applied in this way, have performed particularly well according to the ranking metrics, not only at the row but also at the column level. And finally, we proposed a modification to the Circle loss to better manage the uncertainty of negative samples.

However, further research is needed, these methods need to be investigated more in-depth, and other modifications could be applied. One such possible improvement is to make better use of the intrinsically supervised nature of the approach. Molecules and targets can be augmented within a batch and compared to themselves, thus forming more positive pairs, and making the representations less sensitive to various augmentations. Another promising modification is to replace the cosine similarity with a trainable metric or even try a hyperbolic embedding space and similarities developed for non-Euclidean spaces.

**ACKNOWLEDGEMENT**

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**REFERENCES**