AttentionSiteDTI: an interpretable graph-based model for drug-target interaction prediction using NLP sentence-level relation classification

Mehdi Yazdani-Jahromi†, Niloofar Yousefi†, Aida Tayebi, Elayaraja Kolanthai, Craig J. Neal, Sudipta Seal and Ozlem Ozmen Garibay

Corresponding author: Ozlem Ozmen Garibay, Industrial Engineering and Management Systems, University of Central Florida, Street, 32816, 4000 Central Florida Blvd. Orlando, USA; E-mail: ozlem@ucf.edu

†Mehdi Yazdani-Jahromi and Niloofar Yousefi contributed equally.

Abstract

In this study, we introduce an interpretable graph-based deep learning prediction model, AttentionSiteDTI, which utilizes protein binding sites along with a self-attention mechanism to address the problem of drug–target interaction prediction. Our proposed model is inspired by sentence classification models in the field of Natural Language Processing, where the drug–target complex is treated as a sentence with relational meaning between its biochemical entities a.k.a. protein pockets and drug molecule. AttentionSiteDTI enables interpretability by identifying the protein binding sites that contribute the most toward the drug–target interaction. Results on three benchmark datasets show improved performance compared with the current state-of-the-art models. More significantly, unlike previous studies, our model shows superior performance, when tested on new proteins (i.e. high generalizability). Through multidisciplinary collaboration, we further experimentally evaluate the practical potential of our proposed approach. To achieve this, we first computationally predict the binding interactions between some candidate compounds and a target protein, then experimentally validate the binding interactions for these pairs in the laboratory. The high agreement between the computationally predicted and experimentally observed (measured) drug–target interactions illustrates the potential of our method as an effective pre-screening tool in drug repurposing applications.

Keywords: Deep learning, Self-Attention, Binding Sites, Machine learning, drug–target interaction, SARS-CoV-2, DTI software, DTI database.

Introduction

Drug–target interaction (DTI) characterizes the binding between a drug and its target, critical to the discovery of novel drugs and repurposing of existing drugs. High-throughput screening remains the most reliable approach to examining the affinity of a drug toward its targets. However, the experimental characterization of every possible compound–protein pair quickly becomes impractical due to the immense space of chemical compounds, targets and mixtures. This motivates the use of computational approaches for DTI prediction. Molecular simulation and molecular docking are among the earlier
computational approaches, which typically require 3D structures of the target proteins to assess the DTI. The application of these structure-based methods is limited, as there are many proteins with unknown 3D structures, and they involve an expensive process. Machine Learning (ML) algorithms have emerged to overcome some of these challenges in the process of drug design and discovery. Traditional shallow ML-based models, such as KronRLS [1] and SimBoost [2], require hand-crafted features, which highly affect the performance of the models. Deep Learning has advanced these traditional models due to their ability to automatically capture useful latent features, leading to highly flexible models with extensive power in identifying, processing and extrapolating complex patterns in molecular data. Deep Learning models for DTI can be mainly categorized into two classes. One class is designed to work with sequence-based input data representation. Examples of this type include convolutional neural networks (CNNs) and recurrent neural networks (RNNs). These models are incapable of capturing structural information of the molecules, leading to the degraded predictive power. This limitation motivates the use of a more natural representation of molecules and the introduction of a second class of Deep Learning models, namely graph neural networks (GNNs). GNNs use graph descriptions of molecules, where atoms and chemical bonds correspond to nodes and edges, respectively. Graph convolutional neural network (GCNN) and graph attention network (GAT) [3] are the two widely used GNN-based models in computer-aided drug design, and discovery [4–6]. All these graph-based models use amino acid sequence representations for proteins, which cannot capture the 3D structural features that are key factors in the prediction of DTIs. Obtaining the high-resolution 3D structure of the proteins is a challenging task, besides the fact that proteins contain a large number of atoms requiring a large-scale 3D (sparse) matrix to capture the whole structure. To alleviate this issue, an alternative strategy has been adopted wherein the proteins are represented by a 2D contact (or distance) map that shows the interaction of proteins’ residue pairs in the form of a matrix [7, 8]. It is worth mentioning that a contact (or distance) map is typically the output of protein structure prediction, which is based on heuristics and provides only an approximation abstraction of the real structure of the protein; generally, different from the one determined experimentally via X-ray crystallography or by nuclear magnetic resonance spectroscopy (NMR) [9]. Taken all together, and considering the fact that the binding of a protein to many molecules occurs at different binding pockets rather than the whole protein, in this paper, we propose AttentionSiteDTI, a graph-based deep learning model that incorporates structural features of small molecules and proteins’ binding sites in the form of graphs into the pipeline of DTI prediction task. The structural features of small molecules and proteins, which are automatically learned by the model, will be more conducive to downstream tasks. Our approach is inspired by models developed for sentence classification in the field of Natural Language Processing (NLP) and is motivated by the fact that the structure of the drug–target complex can be very similar to the structure of a natural language sentence in that the structural and relational information of the entities is key to understanding the most important information of the sentence. In this regard, each protein pocket or drug is analogous to a word, and each drug–target pair is analogous to a sentence. Our AttentionSiteDTI, which utilizes a self-attention bidirectional Long Short-Term Memory (LSTM) mechanism, is highly explainable due to its self-attention. In essence, self-attention provides an understanding of which parts of the protein are most probable to bind with a given drug. Visualization of the proposed framework can be found in Figure 1. As the computational results show, our model outperforms many state-of-the-art models in terms of multiple evaluation metrics. The results of our ablation study show that the attention mechanism is very effective in capturing important factors governing DTI. Also, our in-lab experimental validation shows high agreement between computationally predicted and experimentally observed binding interaction between seven candidate compounds and spike (or ACE2) protein.

**Contribution**

The main contributions of our study are summarized as follows.

- We propose a novel formulation of DTI prediction problem inspired by recent developments in NLP that achieves state-of-the-art performance while exhibiting high generalizability and interpretability. Our approach is analogous to the sentence classification problem in NLP, where the drug–target complex is treated as a sentence with relational meaning between its biochemical entities a.k.a. protein pockets and drug molecule. We solve our sentence classification problem using an end-to-end GCNN-based model, which enables simultaneous learning of (1) context-sensitive embeddings from the graphs of molecules and protein pockets (that is, the embeddings are not fixed, but they change according to the context (i.e. sentence) in which they appear), and (2) a prediction model to capture the most important contextual semantic or relational information in a biochemical sequence (drug–target complex) for relation classification similar to sentence classification in NLP. Details of our GCNN-based model can be found in subsection Graph Attention Embedding Module.

- Our proposed model is highly generalizable due to our target protein input representation that uses protein pockets encoded as graphs. This helps the model concentrate on learning generic topological features from protein pockets, which can be generalized to new proteins that are not similar to the ones in the training data. Generalizability of our model is
AttentionSiteDTI specifically showcased by its superior performance on unseen proteins in BindingDB dataset (Figure 4).

Data Preparation and Graph Construction subsections discuss the details on how we prepare and use protein inputs for our model.

- Our proposed model is the first to be interpretable using the language of protein binding sites. This is important because it will help drug designers identify critical protein sites along with their functional properties. This is achieved by devising a self-attention mechanism, which makes the model learn which parts of the protein interact with the ligand, while achieving state-of-the-art prediction performance. Self-Attention subsection explains how the mechanism works by selectively focusing on the most relevant parts of the inputs.

- Finally, we validate the practical potential of our model in the prediction of binding interactions in real-world applications. We conduct in-lab experiments to measure the binding interaction between several compounds and Spike (or ACE2) protein, and compare the computationally predicted results against the ones experimentally observed in the laboratory. In-Lab Validation section provides the details of our in-lab experiments.

Related work

Deep learning-based approaches have been successfully deployed to address the problem of DTI prediction. The main difference between deep learning approaches is in their architecture as well as the representation of the input data. As previously mentioned, small molecules of the drugs can be easily and effectively represented in one-dimensional space, but proteins are much bigger molecules with complex interactions, and 1D representations can be insufficient. Although the datasets containing the 3D structure of the protein are limited, some recent deep learning-based literature has used them in their study. For example, AtomNet [10], is the first study that used the 3D structure of the protein as input to a 3D CNN to predict the binding of drug–target pairs using a binary classifier. Ragoza et al. [11] proposed a CNN scoring function that took the 3D representation of the protein–ligand complex and learned the features critical in binding prediction. This model outperformed the AutoDock Vina score in terms of discriminating and ranking the binding poses. Pafnucy [12] is a 3D CNNs that predicted the binding affinity values for the drug–target pairs. This study represented the input with a 3D grid and considered both proteins and ligands atoms similar. Using a regularization technique, their designed network focused on capturing the general properties of interactions between proteins and ligands.

There are limitations associated with all these studies. For example, it is a highly challenging task to experimentally obtain a high-quality 3D structure of proteins, which explains why the number of datasets with 3D structure information is very limited [8]. Most studies that use 3D structural information utilize CNNs, which are sensitive to different orientations of the 3D structure, besides the fact that these approaches are computationally expensive. To overcome these limitations, recent studies...
such as done in [13–15] have proposed graph convolutional network approaches, which take 3D structure of proteins as input for DTI prediction task. There are other studies that applied GCNN to the 3D structure of the protein–ligand complex. Among these studies, GraphBAR [6] is the first 3D graph CNN that used a regression approach to predicts drug–target binding affinities. They used graphs to represent the complex of protein–ligand instead of 3D voxelized grid cube. These graphs were in the form of multiple adjacency matrices in which the entries were calculated based on distance and feature matrices of molecular properties of the atoms. Also, they used a docking simulation method to augment additional data to their model. Lim et al. [5] proposed a graph convolution network model along with a distance-aware graph attention mechanism to extract features of the interactions binding pose directly from the 3D structure of drug–target complexes from docking software. Their model improved over docking and several deep learning-based models in terms of virtual screening and pose prediction tasks. However, their approach had limitations such as less explainability as well as additional docking errors added to the deep learning model. Pocket Feature is an unsupervised autoencoder model, which was proposed by Torng et al. [4], to learn representations from binding sites of the target proteins. The model used 3D graph representations for protein pockets along with 2D graph representations for drugs. They trained a GCNN model to extract features from the graphs of protein pockets and drugs’ SMILEs. Their model outperformed 3DCNN, as was done in [11] and docking simulation models such as AutoDock Vina [16], RF-Score [17] and NNScore [17]. Zheng et al. [8] pointed out the low efficiency of using direct input of three-dimensional structure and utilized a 2D distance map to represent the proteins. They further converted the problem of DTI prediction into a classical visual question and answering (VQA) problem, wherein, given a distance map of a protein, the question was whether or not a given drug interacts with the target protein. Although their model outperformed several state-of-the-art models, their VQA system is able to solve a classification task only, where it predicts if there is an interaction between drug–target pairs.

Materials and Methods

Overview

We introduce an end-to-end graph-based deep learning framework, AttentionSiteDTI, to address the problem of DTI prediction. We consider the DTI prediction task as a binary classification problem. The aim is to learn a predictive function \( \gamma() \), with graph inputs of drugs and proteins, to output binary predictions suggesting the occurrence or absence of interactions. The overall architecture of AttentionSiteDTI is shown in Figure 1. It consists of four modules: data preparation, graph embedding learning module, prediction module and interpretation module. In the data preparation, we find the binding sites of the proteins using the algorithm proposed in [18]. In the next step, the graphs of protein pockets and ligands are constructed and fed into a graph CNN to learn the embeddings from the corresponding graphs. Feature vectors of drug and protein are then integrated to obtain drug–target complex representation that incorporates their structural features. The concatenated representations are then fed into a binary classifier for predicting DTIs. The self-attention mechanism in the network uses concatenated embeddings of drug–target pairs as input to compute the attention output, which enables interpretability by making the model learn which parts of the protein interact with the ligand in a given drug–target pair.

Algorithm 1 AttentionSiteDTI Training Procedure

\[ P : \text{Protein graph convolution network} \]
\[ P_p : \text{Protein Pooling Layer} \]
\[ L : \text{Ligand graph convolution network} \]
\[ P_l : \text{Ligand pooling Layer} \]
\[ MLP : \text{Multi-layer perceptron classification layers} \]
\[ A : \text{Self-Attention Layer} \]
\[ LSTM : \text{Bi-LSTM Layer} \]
\[ N : \text{Number of epochs} \]
\[ G_i : \text{Graph of ligand in data instance} \]
\[ G_o : \text{List of graphs of protein binding sites in data instance} \]
\[ F_i : \text{List of feature vectors for protein binding sites in data instance} \]
\[ F_o : \text{Feature vector of ligand in data instance} \]
\[ S : \text{Sequence of protein binding sites features and ligand feature} \]
\[ y_i : \text{Label of the data instance} = 0, 1 \]
\[ p(y_i) : \text{Predicted label} \]

while Epoch < N do
  for \((G_i, G_o, label)\) in data do
    \[ F_p = [] \]
    for graph \(G_i, G_o\) do
      \[ F_p \leftarrow F_p \cup [P_p(P(graph))] \]
    end for
    \[ F_l \leftarrow P_l(L(G_o)) \]
    \[ S \leftarrow F_p \cup F_l \]
    \[ S \leftarrow LSTM(S) \]
    \[ S \leftarrow A(S) \]
    \[ p(y_i) \leftarrow MLP(Concat(S)) \]
    Update \(P, P_p, L, P_l, A, MLP\) by descending \(-\nabla_y \log(p(y_i)) + (1 - y_i) \log(1 - p(y_i))\)
  end for
end while

Data Preparation

We use the 3D structure of the proteins that are extracted from Protein Data Bank (PDB) files of proteins. PDB data are collections of submitted experimental values (e.g. from NMR, x-ray diffraction, cryo-electron microscopy) for proteins. In order to find the binding sites of the proteins, we use the algorithm proposed by Saberi Fathi et al. [18], which is one of the simplest methods to extract the binding sites of a protein from its 3D structure. This method is a simulation-based model, and it can be utilized prior to feeding the data into an end-to-end architecture, which helps reduce the model’s complexity.
As shown in [18], despite the simplicity of this model, its performance is comparable to other simulation-based methods with higher complexity.

This algorithm computes bounding box coordinates for each binding site of a protein. These coordinates are then used to reduce the complete protein structure into a subset of peptide fragments. Figure 2 provides a visualization of a protein’s binding sites extracted using the algorithm in [18].

### Graph Construction

#### Protein

Following the extraction of a protein’s binding sites, we then represent them as individual graphs wherein each atom is a node, and the connections between atoms are the edges in the graph. One-hot encoding of the atom type, atom degree, the total number of hydrogen atoms (attached) as well as the implicit valence of the atom are used to compute the feature vector of each atom. This approach yields a vector with a size of $1 \times 31$ for each node. These features are summarized in Table 1. Having multiple graphs, representing different binding pockets of a given protein, leads to the generation of multiple embeddings for that particular protein.

#### Ligand

A bidirectional graph is constructed for each ligand, which is represented in Simplified molecular-input line-entry system (SMILE) format in DTI datasets. Note that in this study, hydrogen atoms are not explicitly represented as the nodes in the graph. Similarly, a vector is constructed to represent the atom’s features in the graph of ligand. More specifically, the one-hot encoding of the atom type, the atom degree, the formal charge of the atom, number of radical electrons of the atom, the atom’s hybridization, atom’s aromaticity and the number of total hydrogens of the atom are used to construct the features of the atoms in a ligand. This approach yields a vector with a size of $1 \times 74$ for each node. Generated graphs for proteins and ligands are then fed into a GCNN to learn the corresponding embeddings.

### Graph Attention Embedding Module

#### Topology Adaptive Graph Convolutional Networks

We use a Topology Adaptive Graph CNN (TAGCN) [20], which is a variant of graph convolutional network [21], and it works by simultaneously sliding a set of fixed-size learnable filters on the input graph to produces a weighted sum of the filter’s outputs, representing both the strength correlation between graph vertices and the vertex features, themselves [20]. In other words, the output of a convolutional layer is the weighted sum of the feature maps resulting from filters with varying size $k$, for $k = 1, \ldots, K$. The graph convolutional layer for TAGCN is defined as

$$H^k = \sum_{k=1}^{K} \left( (D^{-1/2}AD^{-1/2})^k X \Theta_k + b_k \right).$$  

where $A$ denotes the adjacency matrix, $D_{ii} = \sum_j A_{ij}$ is its corresponding diagonal degree matrix, $X$ is the input feature matrix of the nodes, $\Theta_k$ is the vector of linear weights aggregating the results from all the adjacent vertices within a $k$-hop distance of a given node. Also, $b_k$ is the learnable bias, which is used in the summation after every hop.

#### Pooling Mechanism

Once the graphs of proteins and drugs are fed into a series of graph convolutional layers, we then extract embeddings from the corresponding graphs using the method proposed by Yujia Li et al. [22]. For the graph-level representation, they define a vector as

$$h_G = \tanh \left( \sum_{v \in V} \sigma \left(i(H_v^T, x_v)\right) \circ \tanh \left(i(H_v^T, x_v) \right) \right).$$

where, $\circ$ is element-wise multiplication, $i$ and $j$ are linear neural network layers, which are trainable to learn the

| Type                  | Encoding                          |
|-----------------------|-----------------------------------|
| Atom type             | C, N, O, S, F, P, Cl, Br, B, H (onehot encoding) |
| Degree of atom        | 0, 1, 2, 3, 4, 5, 6, 7, 8 (onehot encoding) |
| Number of hydrogen    | 0, 1, 2, 3, 4, 5 (onehot encoding)  |
| Implicit valence      | 0, 1, 2, 3, 4, 5 (onehot encoding)  |
| Is aromatic           | 0, 1 (onehot encoding)             |
best representation of the graph for classification. Also, concatenation of $H^{(T)}$ over the nodes is the output of the graph convolutional layer, and $x_ν$ is the feature vector of node $ν$. Note that the input and output are real-valued vectors, and $σ(i((H^{(T)}_ν, x_ν)))$ is a soft attention mechanism, which determines which node should contribute more toward the final graph embeddings, which allows learning a better representation of the graph [22]. This method is selected over the alternatives, i.e. max pooling and average pooling, as it enables learning higher order relations in the graphs by adding more parameters to the architecture.

**Self-Attention Module**

**Sequence Handling**

Following the extraction of embeddings, we then treat the problem as a text classification problem. We use the method proposed by Zhou et al. [23] to learn a classifier $γ : X → C$, where $X$ is the space of embeddings for protein pockets and ligands, and $C = \{0, 1\}$ is a set of classification labels, with 0 for non-active and 1 for active interactions in a drug-target pair. Given the protein-ligand complex $d ∈ X$ with label $c ∈ C$, we define $(d, c) ∈ X × C$ as

$$ (d, c) = \text{(sequence(protein pockets embeddings, ligand embedding), c)}. \quad (3) $$

**Self-Attention**

Self-Attention mechanism is a method for selectively concentrating on the most relevant parts of the input vector. It accomplishes this task by mapping a query and a set of key-value pairs to a weighted sum of the values, computed by the relationship of the query and the corresponding key [24]. Vaswani et al. [24] describe particular attention called ‘Scaled Dot-Product Attention’, where the input is composed of Queries (Q), Keys (K) and Values (V). They use a linear layer to project the input sequence to Q, K and V. Then, as described in [24], the matrix of the query vectors is multiplied by the transpose of the key vectors. Each element is then normalized by a factor of the square root of the keys’ dimension.

Here, we further apply a mask to prevent binding sites of a protein from paying attention to one another. In other words, we construct the mask in a way that each binding site only pays attention to itself and the interacting ligand. This can be achieved by creating a matrix with the same dimension as K and Q, where its diagonal elements are all set to one. Also, the last column of the matrix, corresponding to the ligand, is set to one. All other values in that matrix are all set to the small value of $9 \times 10^{-15}$ to prevent attention of the binding sites to one another. This mask will then be multiplied by the scaled results. Then row-wise Softmax will be applied to compute the attention map from the key K and the query Q. Finally, the computed matrix will be multiplied by the Values (V) to yield the final result. The sequence of these operations is illustrated in Figure 3, for better clarification.

$$ \text{Attention}(Q, K, V) = \text{Softmax} \left( \frac{QKT}{\sqrt{dk}} \right) V. \quad (4) $$

This self-attention mechanism uses a sequence of embeddings as input to extract Q, K and V. The attention output is then computed using Eq. 4.

**Bi-LSTM**

Long Short Term Memory (LSTM) is a variation of RNNs with memory cells, capable of learning long-term dependencies through the use of ‘additional gates’ incorporated in its architecture. Conventional LSTM only preserves information of the past, as the network receives the sequence inputs only in the original forward direction. Bidirectional LSTM networks add a second layer to LSTM networks to preserve the future and the past information by letting the sequence inputs flow in both forward and backward directions [23]. This feature of Bi-LSTM is especially useful in our particular application, as there is no meaningful order in which the protein binding sites and the ligand should interact. Thus, the future context in the sequence is as important as the past context. We use an element-wise sum to combine the outputs of the forward and backward passes.
Prediction Module

Classifier

The features previously computed are concatenated to a 1D vector \( I \), which is then passed to the classification layers. In this study, two fully connected layers are used to map the extracted features into a final classification output. We use multilayer perceptron with ReLU activation function. Also, dropout is used before each linear layer for better generalizability.

We use Logistic Sigmoid function in the final layer to predict the output in the form of a probability. Moreover, the average binary cross-entropy loss (Equation 5) is used to train the model by back-propagating the error to the network to update all parameters of the model in an end-to-end manner.

\[
L(y, \hat{y}) = -\frac{1}{N} \sum_{i=1}^{N} y_i \log(\hat{y}_i) + (1 - y_i) \log(1 - \hat{y}_i). \tag{5}
\]

Experiments

Datasets

We demonstrate the merit of the proposed model via a series of comparative experiments. We compare our AttentionSiteDTI with several state-of-the-art methods using three benchmark datasets, which provide 3D structure information of target proteins, required by our model. These datasets include the DUD-E dataset, the Human dataset and the customized BindingDB dataset. Table 2 summarizes the statistical information of these datasets. We use a simple docking-based model, proposed in [18], to find the binding sites of the target proteins. It is worth mentioning that we expect a boost in the performance of our model with the use of a more advanced prediction/computation approach that is more accurate in finding binding sites of the proteins.

DUD-E This dataset [25] consists of 102 targets from eight protein families. Each target has around 224 active compounds and more than 10 000 decoys, which were computationally generated in a way that their physical attributes are similar to active compounds but they are topologically dissimilar. Following the prior works by Ragoza et al. [11] and Zheng et al. [8], and in order to make a head-to-head comparison with the benchmark models, we performed 3-fold cross-validation in this dataset, and we reported the average performance on the evaluation metrics. Each fold was split based on the target in that similar targets were kept in the same fold. We used random undersampling on the decoys to make the training set balanced, and we used unbalanced test sets for evaluations.

Human This dataset was built by combining a set of highly credible and reliable negative drug–protein samples via a systematic in silico screening method with the known positive samples [26]. The dataset consists of 5423 interactions. For a head-to-head comparison, we used the same train, validation and test splits (80%,10%,10%) as were used in DrugVQA [8].

BindingDB This dataset [27] contains experimentally based assays of the interactions between small molecules and proteins. Following the work in DrugVQA [8], we used a small subset of the dataset, which consists of 39 747 positive and 31 218 negative samples. Further, in order to validate the generalization ability of our proposed model, the testing set was split into two groups of the proteins: those that are seen during the time of training versus those that are not being seen by the model.

Implementation and evaluation strategy

Experimentation strategies. We used Pytorch 1.8.2 (long-time support version) for our implementations. We trained the models with a batch size of 100 for 30 epochs, and we used Adam optimizer for training the network with a learning rate of 0.001. Also, for better generalization, a dropout with a probability of 0.3 was used before each fully connected layer. The GPU that we used for the experimentation was Nvidia RTX 3090 with 24 GB of memory. We used four as the number of hops in TAGCN for proteins and two for ligands. The size of the hidden state for the BiLSTM layer in our model was set to 31, which was the output of the graph convolution layer (TAGCN). We used padding of zero to reshape each matrix to the maximum number of binding pockets in the datasets. All hyperparameters were tuned to yield the best result for each dataset, which can be seen in Table 3. This hyperparameter tuning was performed by a simple grid search over a range of values, shown in Table 4.

Evaluation metrics. We evaluated our models in terms of several metrics, including the Area Under the receiver operating characteristic Curve (AUC). We additionally reported precision and recall for the human dataset, accuracy for the BindingDB dataset and ROC enrichment metric (RE) for the DUD-E dataset, defined as

\[
RE = \frac{TPR}{FPR} \text{ at a given FPR threshold.} \tag{6}
\]
The choice of different performance metrics as well as different benchmark models for different datasets is mainly for the head-to-head comparison with the results reported in the literature. Nevertheless, the reported metrics are the most widely used evaluation metrics in classification-based models in DTI literature [28]. Also, for comparison, we include a suite of different computational models including (a) traditional ML-based methods such as K-Nearest Neighbors (KNN), L2-logistic (L2), NN-score [29] and Random Forest-score (RF-score) [30]; (b) open-source molecular docking programs including AutoDock Vina [16] and Smina [31]; (c) string representation-based deep learning models such as AtomNet [10] and 3D-CNN [11]; and (d) graph representation-based deep learning models such as Graph CNNs (GCNs) [21], CPI-GNN [32], DrugVQA [8], TransformerCPI [33], PocketGCN [4], GNN [34] and GraphDTA [15] that have been developed to address the problem of DTI prediction.

**Ablation study** The ablation study illustrates the effectiveness of several text classification techniques in our proposed AttentionSiteDTI framework. We report the AUC and the ROC of all experiments, which is widely used in the literature. The results of this study can be found in **Table 5**, indicating that the attention mechanism is the most effective method in text classification, and it is particularly advantageous due to its power in explainability. The Bi-LSTM architecture cannot focus solely on interactions between ligand and binding sites; therefore, it has inferior results compared with other proposed architectures.

As the results show, the self-attention shows superior performance compared with Bi-LSTM+self-attention, except on the DUD-E dataset. This can be explained by the fact that DUD-E, in nature, is a more challenging dataset in the sense that DUD-E decoys (ligands with negative interactions) are matched to the physical chemistry of ligands with positive interactions (in terms of molecular weight, predicted logP, number of rotatable bonds and hydrogen bond donors and acceptors). Decoys should not truly bind to fulfill their duty as negative controls. Hence DUD-E employed 2D similarity fingerprints to reduce the topological similarity between decoys and ligands. In brief, DUD-E decoys were selected to physically mimic ligands with positive interactions (and hence to be difficult to dock), while they are also being topologically distinct to reduce the possibility of actual binding [25]. These decoys’ similarities are particularly challenging for approaches that utilizes ligands’ physical structure such as graph-based neural networks. To this end, a single attention module struggles to create features that are discriminating enough for the classifier head. Therefore, the additional use of a Bi-LSTM layer helps create richer features, which leads to the improved performance of the model on this challenging dataset. To be more specific, we achieved higher accuracy on the DUD-E dataset using an Bi-LSTM layer to capture the short-range relationships, followed by the self-attention mechanism to further improve the quality of the features. Since the other two datasets (Human and BindingDB) are not as challenging as the DUD-E dataset, adding a Bi-LSTM layer not only does not seem to be advantageous but it may also cause the model to overfit.

**Comparison on the DUD-E dataset** On DUD-E, we compared our proposed model with several state-of-the-art models that can be divided into four categories: (1) ML-based methods such as NN-score [29], and Random Forest-score (RF-score) [30]; (2) open-source molecular docking programs including AutoDock Vina [16] and Smina [31]; (3) deep learning-based models such as AtomNet [10] and 3D-CNN [11], which use neural networks to extract features from 3D structural information; and (4) graph-based models such as PocketGCN [4], GNN [34], DrugVQA [8], which are all based on graph representations. PocketGCN utilizes two Graph-CNNs that automatically extract features from the graph of protein pockets and ligands to capture protein–ligand binding interactions. CPI-GNN [32] is a prediction model that combines a graph neural network for ligands and a CNN for targets. DrugVQA utilizes a 2D distance map to represent proteins in a Visual Question Answering system, where the images are the distance maps of the proteins, the questions are the SMILES of the drugs and the answers are whether the drug–target pair will interact. Note that the scores of these models are collected from the work in [8]. Also, we employed the F1 score and ROC enrichment (RE) at thresholds 0.5%, 1%, 2% and 5%. As the results in **Table 6** indicate, our model achieves state-of-the-art performance in DTI prediction on all metrics with significant improvement at 0.5% RE. Also, we hypothesize that the poor performance of AtomNet and 3D-CNN may be due to the sparsity of 3D space, as they use the whole 3D structure of the proteins.

**Comparison on the human dataset** On the Human dataset, we compared our model against several traditional ML models such as K-Nearest Neighbors (KNN),

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**Table 3.** Dataset’s Training Hyperparameters (FC is representing number of fully connected layers, L-GCN and P-GCN are number of Graph convolutional layers for extracting ligands and protein binding sites embedding, respectively)

| Datasets  | FC  | L-GCN | P-GCN |
|-----------|-----|-------|-------|
| DUD-E     | 2   | 18    | 16    |
| Human     | 6   | 4     | 3     |
| Bindingdb | 6   | 4     | 3     |

**Table 4.** Grid search considered values for hyperparameter tuning

| TAGCN | FC       | L-GCN | P-GCN | LR   | Dropout | batch size |
|-------|----------|-------|-------|------|---------|------------|
| Hops  | [1–8]    | [1–8] | [4–20] | [4–20] | [10^{-2}, 10^{-3}] | [0.1–0.6] | [40, 60, 80, 100, 120] |

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Table 5. Ablation Study Results

| Ablation tests                  | BindingDB |            | Human |            | DUD-E |            |
|--------------------------------|-----------|------------|-------|------------|-------|------------|
|                                | AUC       | ROC        | AUC   | ROC        | AUC   | ROC        |
| Bi-LSTM                        | 0.863     | 0.889      | 0.976 | 0.979      | 0.954 | 0.562      |
| Self-Attention                 | 0.940     | 0.981      | 0.991 | 0.990      | 0.961 | 0.602      |
| Bi-LSTM + Self-Attention       | 0.925     | 0.973      | 0.984 | 0.982      | 0.971 | 0.623      |

Table 6. DUD-E Dataset Comparison

| Model            | AUC | 0.5% RE | 1.0% RE | 2.0% RE | 5.0% RE |
|------------------|-----|---------|---------|---------|---------|
| NN Score         | 0.584 | 4.166 | 2.980 | 2.460 | 1.891 |
| RF-score         | 0.622 | 5.628 | 4.274 | 3.499 | 2.678 |
| Vina             | 0.716 | 9.139 | 7.321 | 5.811 | 4.444 |
| Smina            | 0.696 | -      | -      | -      | -      |
| 3D-CNN           | 0.868 | 42.559 | 26.655 | 19.363 | 10.710 |
| AtomNet          | 0.895 | -      | -      | -      | -      |
| PocketGCN        | 0.886 | 44.406 | 29.748 | 19.408 | 10.735 |
| GNN              | 0.940 | -      | -      | -      | -      |
| DrugVQA          | 0.972 | 101.74 | 59.92 | 35.07 | 17.39 |
| AttentionSiteDTI | 0.971 | 101.74 | 59.92 | 35.07 | 16.74 |

Table 7. Human Dataset Comparison

| Model     | AUC   | Precision | Recall | F1 Score |
|-----------|-------|-----------|--------|----------|
| K-NN      | 0.86  | 0.798     | 0.927  | 0.858    |
| RF        | 0.940 | 0.861     | 0.897  | 0.879    |
| L2        | 0.911 | 0.861     | 0.913  | 0.902    |
| SVM       | 0.910 | 0.966     | 0.950  | 0.958    |
| GraphDTA  | 0.960 | 0.882     | 0.912  | 0.897    |
| GCN       | 0.956 | 0.862     | 0.928  | 0.894    |
| CPI-GNN   | 0.970 | 0.923     | 0.918  | 0.920    |
| E2E/GO    | 0.970 | 0.893     | 0.914  | 0.903    |
| DrugVQA   | 0.979 | 0.954     | 0.961  | 0.957    |
| BridgeDPI | 0.990 | 0.963     | 0.949  | 0.956    |
| AttentionSiteDTI | 0.991 | 0.951     | 0.975  | 0.963    |

Random Forest (RF), L2-logistic (L2) (these results were gathered from [17]); and some recently developed graph-based approaches including Graph CNNs [21], CPI-GNN [32], DrugVQA [8], TransformerCPI [33] as well as GraphDTA [15]. GraphDTA was originally designed for regression task, and later was tailored to binary classification task by [35]. For a head-to-head comparison with other models, we followed the same experimental setting as in [34, 36]. Also, we repeated our experiments with three different random seeds, similar to DrugVQA [8]. The performances of the aforementioned models were obtained from [35], as summarized in Table 7. It can be observed that the prediction performance of our proposed model is superior to all ML- and GNN-based models, and it achieves competitive performance with DrugVQA in terms of precision and recall. The relatively low performance of ML-based models is indeed in line with our expectations and is due to their use of low-quality features, unable of capturing complex nonlinear relationships in DTI. The deep learning models, on the other hand, are very powerful in extracting important features governing the complex interactions in a drug-target pair. On this basis, our model further improves on the accuracy, indicating that the quality of learned information in DTIs is guaranteed by the back propagation of the end-to-end learning of our AttentionSiteDTI.

Comparison on the BindingDB dataset On BindingDB dataset, we further compared our model against Tiresias [37], DBN [38], CPI-GNN [32], E2E [39], DrugVQA [8] and Bridge-DPI [35] as baselines. Tiresias uses similarity measures of drug and target pairs. DBN uses stacked restricted Boltzmann machines with inputs in the form of extended connectivity fingerprints. As mentioned earlier, CPI-GNN combines a graph neural network for compounds and a CNN for targets to capture DTIs. E2E is a GNN-based model that uses LSTM to learn drug-target pair information with Gene Ontology annotations. DrugVQA, as previously mentioned, is a Visual Question Answering system, where the images are the distance maps of the proteins, the questions are the SMILES of the drugs and the answers are whether the drug-target pair will interact. Finally, BridgeDPI uses CNNs to obtain embeddings for drugs and proteins, as well as a GNN to learn the associations between proteins/drugs using some hyper-nodes that are connections between proteins/drugs.

Note that the scores for all these models were derived from [35]. Also, following suggestions from previous works, we report the prediction results in terms of AUC and Accuracy (ACC) on the test set, which is divided into a set of unseen proteins (the proteins that are not observed in the training set) and a set of seen protein (the proteins...
that are observed in the training set). This, indeed, makes the customized BindingDB dataset suitable to assess models’ generalization ability to unknown proteins, which should be the focus in prediction problems (i.e., cold-start problem), as there are a large number of unknown proteins in nature.

As experimental results indicate in Figure 4, all models generally perform well on seen proteins with AUC above 0.9 and ACC exceeding 0.85. However, these models show different and much worse performance on unseen proteins, which reflects the complexity of this more realistic learning scenario. Tiresias is a similarity-based model that uses a set of expert-designed similarity measures as the features for proteins and drugs. The poor performance of Tiresias on the unseen proteins is perhaps due to the fact that these handcrafted features are not sufficient in capturing interactions between drug–target pairs, thus resulting in the accuracy of even less than 0.5 on unseen proteins. On the other hand, the good performance of deep learning-based models, including DBN, CPI-GNN, E2E, DrugVQA, BridgeDPI, as well as our AttentionSiteDTI, shows the effectiveness of these models in capturing relevant features that are critical in DTI prediction problem. As the results show, our model achieves the best performance with an AUC of 0.97 and 0.94 on seen and unseen proteins, respectively. Also, in terms of accuracy, our AttentionSiteDTI outperforms all other models, with accuracy reaching 0.89 in unseen proteins. This is an indication that our attention-based bidirectional LSTM network is, indeed, effective in relation classification of drug–target (protein pocket) pairs by learning the deeper interaction rules governing the relationship between proteins’ binding sites (pockets) and drugs. Also, the seemingly good performance of baselines on seen proteins can be an indication of over-fitting.

Discussion

We believe that the improved performance of our proposed model can be explained by several factors, including (1) the input representations, which, as argued in [40], significantly affect the prediction performance of the model. The use of more advanced input feature representations such as structured graphs can help capture the structural information of the molecules, (2) the prediction technique. For example, traditional ML-based techniques such as NNScore and FRscore often depend on the quality of hand-crafted features, which most often fail to learn complex nonlinear relationships in DTI [35]. In contrast, self-attention provides a powerful and automatic feature extraction mechanism to learn higher order nonlinear relationships. Also, deep learning approaches that use string representations as the input to their models are unable to capture the structural information of drugs and/or proteins [41]. On the other hand, graph-based neural networks (that use graph representations of drugs and proteins) can effectively capture topological relationships of drug molecules and target proteins, which enables further performance improvement [41], and most importantly (3) our context-sensitive embedding approach to drug–target complexes. In our approach, a drug–target complex is treated as a sentence with relational meaning between its biochemical entities a.k.a. protein pockets and drug molecule; This enables capturing the most important contextual semantic and relational information in a biochemical sequence (i.e. sentence) for relation classification similar to sentence classification in NLP.

In addition to improved performance, our model enables interpretability by exploring which binding sites of the protein interact with a given ligand. This is especially crucial in the design and development of new
pharmaceutically active molecules, where it is critical to know which parts of a molecule are important for its biological properties.

**Interpretation Module**

Ligands bind to certain parts (active sites) of proteins, either blocking the binding of other ligands or inducing a change in the protein structure, which produces a therapeutic effect. Binding at other sites that provide no therapeutic value is ‘non-active’, and generally does not cause a direct biological effect. Ligands binding to active sites and inducing a change in protein structure (conformation) are less likely in our system of study and are probably not as helpful for building models (usually, these ligands/therapeutic agents are employed/considered when a patient has an ailment, which causes natural biochemicals to be produced in insufficient quantities).

In this work, the attention mechanism enables the model to predict which protein binding sites are more probable to bind with a given ligand. These probabilities are contained in the attention matrix, computed by the model. The attention visualization can be found in Figure 5 as the heat map of the protein for the complex of SARS-CoV2 Spike protein and human, host cell-expressing ACE2 in the interaction with the drug named Darunavir. The projection of the heat map on the protein is depicted in this figure, as well.

**In-Lab Validation**

**SARS-CoV-2 Case study**

To further evaluate the practical potential of our proposed model, we experimentally tested and validated the binding interactions between spike (or ACE2) protein and seven candidate compounds (N-acetyl-neuraminic acid, 3α,6α-Mannopentaose, N-glycolyneuraminic acid, 2-Keto3-deoxyoctonate, N-acetyllactosamine, cytidine5-monophospho-N-acetylneuraminic acid sodium salt and Darunavir). Evaluation was performed using a binding inhibition assay kit. Here, candidate molecules were used as inhibitor compounds toward formation of the spike protein-ACE2 complex. Greater performance, of a given candidate molecule, in these assays translates directly to therapeutic value as formation of this complex has been identified as the first step toward host cell infection by the SARS-CoV2 virus. Interaction between these two proteins is facilitated by a glycans on the host cell. These glycan modifications of surface proteins are unique in composition, surface density and in linkage (i.e. ‘O’ and ‘N’ type); as well as often being cell type and tissue-specific [42]. The glycan modifications on ACE2 are common for cells of the respiratory system (oral cavity, lung epithelial cells), allowing recognition by certain upper respiratory tract infecting viruses (e.g. SARS-CoV2, porcine epidemic diarrhea virus and alphacoronavirus transmissible gastroenteritis virus) [43, 44]. It has been shown that SARS-CoV2 binds specifically to neuraminic
We proposed using graph representations of both drug and target complex as a natural language sentence. This enables interpreting and relational information contained in the sentence.

A graph-based deep learning model was developed to solve the DTI prediction problem in an end-to-end manner, where both the graph embeddings and the DTI prediction model were learned simultaneously. This enables more efficient and effective training of the network as a whole with minimizing over one loss function.

We proposed using graph representations of both drug and target as the inputs to the network. More specifically, we used graph representations of protein pockets as inputs for the target protein, which helps with better generalizability of the model.

We devised self-attention mechanism that enables interpretability through identification of most probable binding sites of a protein with a given ligand.

Our proposed model showed improved performance compared with many state-of-the-art models in terms of several performance metrics on three benchmark datasets.

We validated the practical potential of our computation model through in-lab validation, where we measured the binding interaction between several compounds and Spike (ACE2) protein, and compared the results with computational predictions.

We observed high agreement between computationally predicted and experimentally observed binding interactions, which illustrates the potential of our model in virtual screening tasks.

### Conclusion

In this work, we proposed an end-to-end GCNN-based model, built on a self-attention mechanism, to capture any relationship between binding sites of a given protein and the drug in a sequence analogous to a sentence with relational meaning between its biochemical entities a.k.a. protein pockets and drug molecule. Our proposed framework enables learning which binding sites of a protein interact with a given ligand, thus allowing interpretability and better generalizability, while outperforming state-of-the-art methods in the prediction of DTI. We experimentally validated the predicted binding interactions between seven candidate compounds and the Spike (or ACE2) protein. The results of our in-lab validation showed high agreement between the computationally predicted and experimentally observed binding interactions. Our model exhibits state-of-the-art performance, is highly generalizable, provides interpretable outputs and performs well when validated against in-lab experiments. As a result, we expect it to be an effective virtual screening tool in drug discovery applications.

### Data and Code availability

All datasets are publicly available. DUD-E dataset is available at [http://dude.docking.org](http://dude.docking.org). Human dataset is available at [https://github.com/IBMInterpretableDTIP](https://github.com/IBMInterpretableDTIP) and finally the customized BindingDB-IBM dataset.
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