GEFA: EARLY FUSION APPROACH IN DRUG-TARGET AFFINITY PREDICTION

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ABSTRACT

Predicting the interaction between a compound and a target is crucial for rapid drug repurposing. Deep learning has been successfully applied in drug-target affinity (DTA) problem. However, previous deep learning-based methods ignore modeling the direct interactions between drug and protein residues. This would lead to inaccurate learning of target representation which may change due to the drug binding effects. In addition, previous DTA methods learn protein representation solely based on a small number of protein sequences in DTA datasets while neglecting the use of proteins outside of the DTA datasets. We propose GEFA (Graph Early Fusion Affinity), a novel graph-in-graph neural network with attention mechanism to address the changes in target representation because of the binding effects. Specifically, a drug is modeled as a graph of atoms, which then serves as a node in a larger graph of residues-drug complex. The resulting model is an expressive deep nested graph neural network. We also use pre-trained protein representation powered by the recent effort of learning contextualized protein representation. The experiments are conducted under different settings to evaluate scenarios such as novel drugs or targets. The results demonstrate the effectiveness of the pre-trained protein embedding and the advantages our GEFA in modeling the nested graph for drug-target interaction.

Machine learning is an important component of the growing field of data science. Through the use of statistical methods, different type of algorithms is trained to make classifications or predictions, and to uncover key insights in this project. These insights subsequently drive decision making within applications and businesses, ideally impacting key growth metrics. Machine learning algorithms build a model based on this project data, known as training data, in order to make predictions or decisions without being explicitly programmed to do so. Machine learning algorithms
are used in a wide variety of datasets, where it is difficult or unfeasible to develop
c conventional algorithms to perform the needed tasks.

I. INTRODUCTION

The accurate prediction of drug-target
affinity (DTA) is a crucial aspect of drug
discovery, enabling the identification of
promising drug candidates that can
effectively interact with specific
biological targets. Traditional
experimental techniques for determining
DTA, such as high-throughput screening
and binding assays, are both time-
consuming and resource-intensive,
underscoring the need for efficient
computational approaches that can
accelerate the drug development process
【1】. In recent years, machine learning
(ML) and deep learning (DL) models
have gained traction in this field,
offering substantial improvements in the
speed and accuracy of DTA predictions
【2】【3】.

One of the innovative strategies in
advancing DTA prediction is the early
fusion approach. Early fusion involves
integrating multiple data sources or
feature representations at an initial stage,
allowing the model to learn complex
interactions between drug molecules and
their target proteins from the outset
【4】. This contrasts with late fusion
methods, where separate models are
trained on individual data modalities,
and their outputs are later combined. By
leveraging early fusion, models can
capture intricate relationships between
drugs and targets, leading to enhanced
predictive performance【5】.

The proposed GEFA (Generalized Early
Fusion Approach) builds on this concept
by developing a robust framework for
early fusion in DTA prediction. GEFA
aims to combine various molecular
representations of drugs, such as
SMILES strings and molecular
fingerprints, with protein sequences or
structural data, ensuring that the model
fully exploits the complementary
information from these diverse inputs.
This approach not only improves the
accuracy of DTA predictions but also
provides a scalable solution for large-
scale drug discovery projects【6】
【7】.

II.EXISTING SYSTEM

Drug re-purposing [18] is the process of
identifying well established medications
for the novel target disease. The
advantages of this drug re-purposing over developing a completely novel drug are lower risk and fast-track development [19]. The process of drug re-purposing consists of three key steps: identifying the candidate molecules given the target disease, drug effect assessment in the preclinical trial, and effectiveness assessment in clinical trial [20]. The first step, hypothesis generation, is critical as it decides the success of the whole process. Advanced computational approaches are used for hypothesis generation. Computational approaches in drug re-purposing can be categorized into six groups [20]: genetic association [21], [22], pathway pathing, retrospective clinical analysis, novel data sources, signature matching [29]– [31], molecular docking [32]–[34].

Drug-target binding affinity indicates the strength of the binding force between the target protein and its ligand (drug or inhibitor) [35]. The drug-target binding affinity prediction problem is a regression task predicting the value of the binding force. The binding strength is measured by the equilibrium dissociation constant (KD). A smaller KD value indicates a stronger binding affinity between protein and ligand [35]. There are two main approaches: structural approach and non-structural approach [1]. Structural methods utilize the 3D structure of protein and ligands to run the interaction simulation between protein and ligand. On the other hand, the non-structural approach relies on ligand and protein features such as sequence, hydrophobic, similarity or other alternative structural information.

The structure-based approach involves molecular docking, predicting the three-dimensional structure of the target-ligand complex. In molecular docking, there are a large number of target-ligand complex conformations. The conformations are evaluated by the scoring function. Based on the scoring function types, the structural approach can be categorized into three groups [1]: classical scoring function method [36]–[39], machine learning scoring function method [40], and deep learning scoring function method [41], [42]. In classical scoring approaches, Elanie et al. [36] uses DelPhi-calculated potential at each ligand atom for the contact scoring function. In machine learning approaches, Kundu et al. [40] extracts ligand features (e.g. atom count, physicochemical properties) and protein features (e.g. accessible surface, number of chains) from 3D structure data then applies machine learning to learn the scoring function. In deep learning
approaches, Marta et al. [41] uses 3D convolution with protein-ligand 3D structure to predict the binding affinity.

**Disadvantages**
- The system is not implemented compare the drug representation which extracted from the drug-protein fusion graph and drug representation extracted from the drug graph.
- The system is not implemented Graph Early Fusion for binding Affinity prediction (GEFA).

**III. PROPOSED SYSTEM**

- In summary, the contribution of our work is two-fold. First, we combine the protein sequence embedding feature and protein contact map to build the graph representation of a target protein. Second, in order to reflect the target representation change during the binding process, we propose a so-called Graph Early Fusion for binding Affinity prediction (GEFA) for more accurate biological modeling. We demonstrate the effects of the GEFA on Davis dataset [17] where it has shown superior performance against previous studies on different settings.

To address target protein representation change, the system proposes an early-fusion-based approach. Initially, we extract representation feature for a given drug molecule from its drug graph structure. Then, the drug representation is integrated into the protein graph structure before the protein representation learning phrase. This is basically a graph structure nested inside another graph structure. This graph-in-graph neural network design allows the model to learn changes in protein representation caused by the binding process with the drug molecule.

**Advantages**
- The proposed system refines the Graph-Graph Integration with Early Fusion and Graph Early Fusion for binding Affinity prediction (GEFA).
- The proposed system implemented the usage of attention mask as the graph edge. Instead of using attention as drug-residue edge weight, drug-residue edges are weighted the same as the residue-residue edges in the target graph.

**IV. MODULES**

**Service Provider**

In this module, the Service Provider has to login by using valid user name and
password.

After login successful he can do some operations such as Login, Train & Test Data Sets, View Trained Accuracy in Bar Chart,

View Trained Accuracy Results,

View Type, Find Type Ratio, Download Predicted Datasets, View Type Ratio Results, View All Remote Users.

**View and Authorize Users**

In this module, the admin can view the user’s details such as, user name, email, address and admin authorizes the users.

**Remote User**

In this module, there are n numbers of users are present. User should register before doing any operations. Once user registers, their details will be stored to the database. After registration successful, he has to login by using authorized user name and password. Once Login is successful user will do some operations like register and login, predict type, view your profile.

**V. CONCLUSION**

In this project, the GEFA (Generalized Early Fusion Approach) was introduced as an innovative framework to enhance drug-target affinity (DTA) prediction using machine learning techniques. By leveraging early fusion, the model is able to integrate multiple data sources such as molecular representations of drugs and protein sequences—at an initial stage, allowing for a more comprehensive understanding of the relationships between drugs and their biological targets. The integration of these diverse modalities improves the model’s ability to capture complex interactions, resulting in more accurate predictions.
and reliable predictions compared to traditional approaches.

GEFA not only demonstrates the potential to advance the field of computational drug discovery but also addresses critical challenges such as scalability and efficiency. By refining how input data is fused and processed, this method significantly reduces the time and cost of identifying potential drug candidates. As drug discovery continues to evolve with increasing reliance on computational techniques, GEFA provides a scalable and adaptable approach that can be extended to various types of biological targets and drug molecules. Future research could explore further optimizations in feature selection, model architectures, and real-world validations to fully unlock the potential of this early fusion approach.

VI. REFERENCES
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