



# A DEEP LEARNING APPROACH FOR DRUG SIDE EFFECT PREDICTION: INTRODUCING DEEP SIDE FRAMEWORK

<sup>1</sup>S. Vijay kumar, <sup>2</sup>K. Anil

<sup>1</sup>Assistant Professor, <sup>2</sup>MCA Student

Department Of MCA Student

Sree Chaitanya College of Engineering, Karimnagar

## ABSTRACT

Clinical trial drug failures brought on by unanticipated side effects put participants' health at risk and result in significant financial losses. Algorithms for predicting side effects might potentially direct the medication creation process. The LINCS L1000 dataset builds a knowledge foundation for context-specific aspects and offers a wealth of cell line gene expression data disturbed by various medications. Only the high-quality trials in LINCS L1000 are used in the state-of-the-art method, which discards a significant percentage of the experiments in order to use context-specific information. Our aim in this study is to fully utilise this data in order to improve the prediction performance. Five deep learning architectures are used in our experiments. When using drug chemical structure (CS) and the entire collection of drug altered gene expression profiles (GEX) as modalities, we find that a multi-modal design yields the greatest prediction performance among multi-layer perceptron-based architectures. In general, we find the CS to be more informative than the GEX. The best results are obtained by a convolutional neural network-based model that just employs the SMILES string representation of the

medications; this model outperforms the state-of-the-art by 13:0% in macro-AUC and 3:1% in micro-AUC. Additionally, we demonstrate that the model can predict drug-side effect couples that have been documented in the literature but were absent from the ground truth side effect dataset.

## 1. INTRODUCTION

Computational methods hold great promise for mitigating the health and financial risks of drug development by predicting possible side effects before entering into the clinical trials. Several learning based methods have been proposed for predicting the side effects of drugs based on various features such as: chemical structures of drugs drug-protein interactions, protein-protein interactions (PPI), activity in metabolic networks, pathways, phenotype information and gene annotations [8]. In parallel to the above mentioned approaches, recently, deep learning models have been employed to predict side effects: (i) [31] uses biological, chemical and semantic information on drugs in addition to clinical notes and case reports and (ii) [4] uses various chemical fingerprints extracted using deep



architectures to compare the side effect prediction performance.

While these methods have proven useful for predicting adverse drug reactions (ADRs – used interchangeably with drug side effects), the features they use are solely based on external knowledge about the drugs (i.e., drug-protein interactions, etc.) and are not cell or condition (i.e., dosage) specific. To address this issue, Wang et al. (2016) utilize the data from the LINCS L1000 project [32]. This project profiles gene expression changes in numerous human cell lines after treating them with a large number of drugs and small-molecule compounds. By using the gene expression profiles of the treated cells, [32] provides the first comprehensive, unbiased, and cost-effective prediction of ADRs. The paper formulates the problem as a multi-label classification task. Their results suggest that the gene expression profiles provide context-dependent information for the side-effect prediction task. While the LINCS dataset contains a total of 473,647 experiments for 20,338 compounds, their method utilizes only the highest quality experiment for each drug to minimize noise. This means that most of the expression data are left unused, suggesting a potential room for improvement in the prediction performance. Moreover, their framework performs feature engineering by transforming gene expression features to enrichment vectors of biological terms. In this work, we investigate whether the incorporation of gene expression data along with the drug structure data can be leveraged

Page | 432

better in a deep learning framework without the need for feature engineering.

In this study, we propose a deep learning framework, Deep Side, for ADR prediction. Deep Side uses only (i) in vitro gene expression profiling experiments (GEX) and their experimental meta data (i.e., cell line and dosage - META), and (ii) the chemical structure of the compounds (CS). Our models train on the full LINCS L1000 dataset and use the SIDER dataset as the ground truth for drug - ADR pair labels [13]. We experiment with five architectures: (i) a multi-layer perceptron (MLP), (ii) MLP with residual connections (Res MLP), (iii) multi-modal neural networks (MMNN. Concat and MMNN. Sum), (iv) multi-task neural network (MTNN), and finally, (v) SMILES convolutional neural network (SMILES Conv).

We present an extensive evaluation of the above-mentioned architectures and investigate the contribution of different features. Our experiments show that CS is a robust predictor of side effects. The base MLP model, which uses CS features as input, produces \_11% macro-AUC and \_2% micro-AUC improvement over the state-of-the-art results provided in [32], which uses both GEX (high quality) and CS features. The multi-modal neural network model, which uses CS, GEX and META features and uses summation in the fusion layer (MMNN. Sum) achieves 0:79 macro-AUC and 0:877 micro-AUC which is the best result among MLP based approaches. We also find out that when the



<https://doi.org/10.5281/zenodo.14066203>

chemical structure features are fully utilized in a complex model like ours, it overpowers the information that is obtained from the GEX dataset. The convolutional neural network that only uses the SMILES string representation of the drug structures achieves the best result among all the proposed architectures with provides 13:0% macro-AUC and 3:1% micro-AUC improvement over the state-of-the-art algorithm.

## 2. EXISTING SYSTEM

A drug-drug interaction (DDI) is defined as an association between two drugs where the pharmacological effects of a drug are influenced by another drug. Positive DDIs can usually improve the therapeutic effects of patients, but negative DDIs cause the major cause of adverse drug reactions and even result in the drug withdrawal from the market and the patient death. Therefore, identifying DDIs has become a key component of the drug development and disease treatment.

In this study, an existing system, develops a method to predict DDIs based on the integrated similarity and semi-supervised learning (DDI-IS-SL). DDI-IS-SL integrates the drug chemical, biological and phenotype data to calculate the feature similarity of drugs with the cosine similarity method. The Gaussian Interaction Profile kernel similarity of drugs is also calculated based on known DDIs. A semi-supervised learning method (the Regularized Least Squares classifier) is used to calculate the interaction possibility scores of drug-drug pairs. In terms of the 5-fold cross validation, 10-fold

Page | 433

cross validation and de novo drug validation, DDI-IS-SL can achieve the better prediction performance than other comparative methods. In addition, the average computation time of DDI-IS-SL is shorter than that of other comparative methods. Finally, case studies further demonstrate the performance of DDI-IS-SL in practical applications.

### Disadvantages

- The complexity of data: Most of the existing machine learning models must be able to accurately interpret large and complex datasets to detect an accurate Drug Side Effect.
- Data availability: Most machine learning models require large amounts of data to create accurate predictions. If data is unavailable in sufficient quantities, then model accuracy may suffer.
- Incorrect labeling: The existing machine learning models are only as accurate as the data trained using the input dataset. If the data has been incorrectly labeled, the model cannot make accurate predictions.

## 3. PROPOSED SYSTEM

Multi-layer perceptron (MLP) Our MLP [22] model takes the concatenation of all input vectors and applies a series of fully-connected (FC) layers. Each FC layer is followed by a batch normalization layer [10]. We use ReLU activation [16], and dropout regularization [27] with a drop probability of 0:2. The sigmoid activation function is applied to the final layer outputs, which yields the ADR prediction probabilities. The loss function is defined as the sum of negative log- probabilities over



ADR classes, i.e. the multi-label binary cross-entropy loss (BCE). An illustration of the architecture for CS and GEX features is given in this system.

**Residual multi-layer perceptron (ResMLP)**  
The residual multi-layer perceptron (ResMLP) architecture is very similar to MLP, except that it uses residual-connections across the fully- connected layers. More specifically, the input of each intermediate layer is element-wise added to its output, before getting processed by the next layer. Such residual connections have been shown to reduce the vanishing gradient problem to a large extent [7].

This effectively allows deeper architectures, therefore, potentially learning more complex and parameter-efficient feature extractors. **Multi-modal neural networks (MMNN)** The multi-modal neural network approach contains distinct MLP sub-networks where each one extract features from one data modality only. The outputs of these sub-networks are then fused and fed to the classification block. For feature fusion, we consider two strategies: concatenation and summation. While the former one concatenates the domain-specific feature vectors to a larger one, the latter one performs element-wise summation. By definition, for summation based fusion, the domain-specific feature extraction sub-networks have to be designed to produce vectors of equivalent sizes. We refer to the concatenation and summation based MMNN networks as MMNN.Concat and MMNN.Sum, respectively.

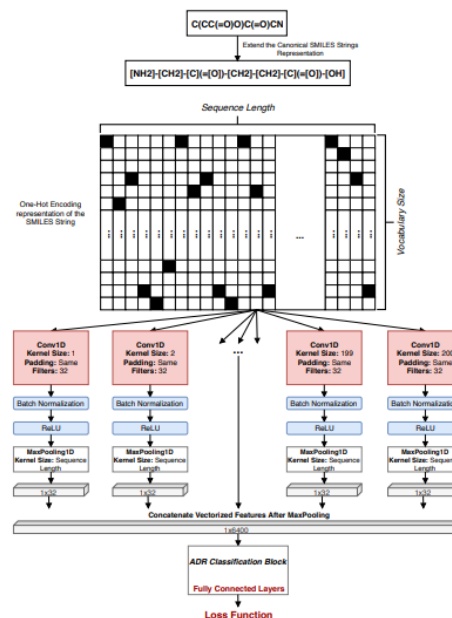
**Multi-task neural network (MTNN)** our multitask learning (MTL) based architecture

aims to take the side effect groups obtained from the taxonomy of ADReCS into account. For this purpose, the approach defines shared and task-specific MLP sub-network blocks. The shared block takes the concatenation of GEX and CS features as input and outputs a joint embedding. Each task-specific sub-network then converts the joint embedding into a vector of binary prediction scores for a set of inter-related side-effect classes.

### Advantages

- The proposed system implemented many ml classifiers for testing and training on datasets.
- The proposed system developed Convolutional neural networks (CNN) which are known to provide a powerful way of automatically learning complex features in vision tasks to find an accurate accuracy on the datasets.

## 4. SYSTEM ARCHITECTURE





## 5. 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 Train & Test Drug Data Sets, View Trained and Tested Drug Datasets Accuracy in Bar Chart, View Trained and Tested Drug Datasets Accuracy Results, View Drug Side Effect Prediction Type, Find Drug Side Effect Prediction Type Ratio, Download Predicted Data Sets, View Drug Side Effect Prediction Type Ratio Results, View All Remote Users.

### View and Authorize Users

In this module, the admin can view the list of users who all registered. In this, 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 DRUG SIDE EFFECT TYPE, VIEW YOUR PROFILE.

## 6. ALGORITHM

### Gradient boosting

**Gradient boosting** is a machine learning technique used in regression and classification tasks, among others. It gives a prediction model in the form of an ensemble of weak prediction

Page | 435

models, which are typically decision trees.<sup>[1][2]</sup> When a decision tree is the weak learner, the resulting algorithm is called gradient-boosted trees; it usually outperforms random forest. A gradient-boosted trees model is built in a stage-wise fashion as in other boosting methods, but it generalizes the other methods by allowing optimization of an arbitrary differentiable loss function.

### Logistic regression Classifiers

*Logistic regression analysis* studies the association between a categorical dependent variable and a set of independent (explanatory) variables. The name *logistic regression* is used when the dependent variable has only two values, such as 0 and 1 or Yes and No. The name *multinomial logistic regression* is usually reserved for the case when the dependent variable has three or more unique values, such as Married, Single, Divorced, or Widowed. Although the type of data used for the dependent variable is different from that of multiple regression, the practical use of the procedure is similar.

Logistic regression competes with discriminant analysis as a method for analyzing categorical-response variables. Many statisticians feel that logistic regression is more versatile and better suited for modeling most situations than is discriminant analysis. This is because logistic regression does not assume that the independent variables are normally distributed, as discriminant analysis does.





<https://doi.org/10.5281/zenodo.14066203>

This program computes binary logistic regression and multinomial logistic regression on both numeric and categorical independent variables. It reports on the regression equation as well as the goodness of fit, odds ratios, confidence limits, likelihood, and deviance. It performs a comprehensive residual analysis including diagnostic residual reports and plots. It can perform an independent variable subset selection search, looking for the best regression model with the fewest independent variables. It provides confidence intervals on predicted values and provides ROC curves to help determine the best cutoff point for classification. It allows you to validate your results by automatically classifying rows that are not used during the analysis.

### SVM

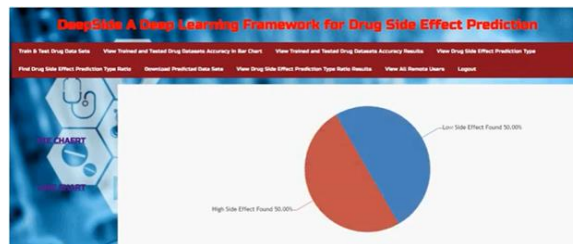
In classification tasks a discriminant machine learning technique aims at finding, based on an *independent and identically distributed* (iid) training dataset, a discriminant function that can correctly predict labels for newly acquired instances. Unlike generative machine learning approaches, which require computations of conditional probability distributions, a discriminant classification function takes a data point  $x$  and assigns it to one of the different classes that are a part of the classification task. Less powerful than generative approaches, which are mostly used when prediction involves outlier detection, discriminant approaches require fewer computational resources and less training data, especially for a multidimensional feature space and when

only posterior probabilities are needed. From a geometric perspective, learning a classifier is equivalent to finding the equation for a multidimensional surface that best separates the different classes in the feature space.

SVM is a discriminant technique, and, because it solves the convex optimization problem analytically, it always returns the same optimal hyperplane parameter—in contrast to *genetic algorithms* (GAs) or *perceptrons*, both of which are widely used for classification in machine learning. For perceptrons, solutions are highly dependent on the initialization and termination criteria. For a specific kernel that transforms the data from the input space to the feature space, training returns uniquely defined SVM model parameters for a given training set, whereas the perceptron and GA classifier models are different each time training is initialized. The aim of GAs and perceptrons is only to minimize error during training, which will translate into several hyperplanes' meeting this requirement.

## 7. SCREEN SHOTS





## 8. CONCLUSION

The process of developing pharmaceutical drugs is a drawn-out and difficult one. Unexpected adverse drug reactions (ADRs) during the drug development process have the potential to halt or restart the entire process. As a result, it is crucial to foresee the drug's adverse effects in advance during the design stage. To account for factors like dosage, time interval, and cell line, we forecast ADRs using our Deep Side framework, which combines chemical structure with context-related (gene expression) information. The suggested MMNN model outperforms models that just employ chemical structure (CS) fingerprints in terms of accuracy by utilising GEX and CS as integrated features. Considering that we are simply attempting to predict the condition-independent side effects, the stated accuracy is significant. Lastly, by using convolution on the SMILES representation of drug chemical structure, the SMILES Conv model performs better than any other method.

## REFERENCES

1. Atias, N., Sharan, R.: An algorithmic framework for predicting side effects of drugs. Journal of Computational Biology 18(3), 207{218 (2011)



<https://doi.org/10.5281/zenodo.14066203>

2. Bresso, E., Grisoni, R., Marchetti, G., Karaboga, A.S., Souchet, M., Devignes, M.D., Sma • \_l-Tabbone, M.: Integrative relational machine-learning for understanding drug side-effect profiles. BMC bioinformatics 14(1), 207 (2013)
3. Cai, M.C., Xu, Q., Pan, Y.J., Pan, W., Ji, N., Li, Y.B., Jin, H.J., Liu, K., Ji, Z.L.: Adres: an ontology database for aiding standardization and hierarchical classification of adverse drug reaction terms. Nucleic acids research 43(D1), D907{D913 (2014)
4. Dey, S., Luo, H., Fokoue, A., Hu, J., Zhang, P.: Predicting adverse drug reactions through interpretable deep learning framework. BMC Bioinformatics 19 (12 (2018). <https://doi.org/10.1186/s12859-018-2544-0>
5. Dimitri, G.M., Li\_o, P.: Drugclust: A machine learning approach for drugs side effects prediction. Computational Biology and Chemistry 68, 204 { 210 (2017). <https://doi.org/https://doi.org/10.1016/j.compbiochem.2017.03.008>
6. Groopman, J.E., Itri, L.M.: Chemotherapy-induced anemia in adults: incidence and treatment. Journal of the National Cancer Institute 91(19), 1616{1634 (1999)
7. He, K., Zhang, X., Ren, S., Sun, J.: Deep residual learning for image recognition. In: The IEEE Conference on Computer Vision and Pattern Recognition (CVPR) (June 2016)
8. Huang, L.C., Wu, X., Chen, J.Y.: Predicting adverse side effects of drugs. BMC genomics 12(5), S11 (2011)
9. Huang, L.C., Wu, X., Chen, J.Y.: Predicting adverse drug reaction profiles by integrating protein interaction networks with drug structures. Proteomics 13(2), 313{324 (2013)
10. Io\_e, S., Szegedy, C.: Batch normalization: Accelerating deep network training by reducing internal covariate shift. arXiv preprint arXiv:1502.03167 (2015)
11. Kim, S., Thiessen, P.A., Bolton, E.E., Chen, J., Fu, G., Gindulyte, A., Han, L., He, J., He, S., Shoemaker, B.A., et al.: Pubchem substance and compound databases. Nucleic acids research 44(D1), D1202{D1213 (2015)
12. Krizhevsky, A., Sutskever, I., Hinton, G.E.: Imagenet classification with deep convolutional neural networks. In: Advances in neural information processing systems. pp. 1097{1105 (2012)
13. Kuhn, M., Letunic, I., Jensen, L.J., Bork, P.: The sider database of drugs and side effects. Nucleic acids research 44(D1), D1075{D1079 (2015)
14. Landrum, G., et al.: Rdkit: Open-source cheminformatics (2006)
15. Lee, W., Huang, J., Chang, H., Lee, K., Lai, C.: Predicting drug side effects using data analytics and the integration of multiple data sources. IEEE Access 5, 20449{20462 (2017). <https://doi.org/10.1109/ACCESS.2017.2755045>