FEAT: A Fast, Effective, and Feasible Model for Molecular Property Prediction Based on Graph Neural Network

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Abstract: Artificial Intelligence based methods and algorithms are being increasingly used by chemists to perform various tasks that would be rather difficult to perform using conventional methods. Whenever scientists design a new set of molecules for certain application, they need to experimentally validate if it possesses the desirable properties. Such (iterative) methods are often expensive and time-consuming. In the realm of Artificial Intelligence and Machine Learning, the molecules can themselves be viewed as graphs present in nature with bonds as edges and nodes as atoms. Therefore, it is worthwhile to exploit Graph Neural Networks for extracting the structural properties of these atoms and bonds, so as to further leverage these to predict the properties of these molecules (represented as graphs) as a whole. We propose a Graph Neural Network based model, FEAT, for this purpose. FEAT’s performance has been evaluated on multiple publicly available datasets and the results obtained are promising.

1 INTRODUCTION

In recent times various Artificial Intelligence (AI) methods and algorithms are being used more and more by chemists to perform various tasks that would be rather difficult to perform using traditional methods. Latest applications of AI in chemistry are targeted towards designing new molecules for drug discovery, attaining knowledge of universe etc. Whenever researchers design a new molecule, its properties have to be experimentally identified and analysis is to be performed whether they are the desired ones or not. Many times the process of experimentally identifying the properties becomes rather expensive and time-consuming as we may have hundreds or even thousands of new molecules designed and waiting to test for properties. These experiments can be iteratively done with the help of artificial intelligence.

1.1 Quantitative Structure, Activity Relationship

Quantitative Structure-Activity Relationship (QSAR) models have been used in the past to predict molecular properties using physical laws or empirical relationships relating the structure of the molecule to their properties.

The physiological activity $\Phi$ was expressed as a function of chemical structure $C$.

$$\Phi = f(C)$$  \hspace{1cm} (1)

One of the first historical QSAR applications was to predict boiling points for different molecules (Dearden, 2003). A general trend was observed between number of carbons in alkanes and their boiling points. The boiling points for alkanes increases as the number of carbons increased in the molecules (Dearden, 2003).

But generally, the patterns within structure and properties are not usually very straightforward and are very complex in nature. Hence, we try to implement the same idea using Artificial Intelligence and Machine Learning so that complex structural-property relations could be extracted from molecules to predict some property (e.g. solubility) accurately.

1.2 Graph Neural Networks

Graphs are a type of data structures having two
components: Vertices (or Nodes) and edges. An edge connects two nodes as shown in Figure 1. A graph can be represented by an adjacency matrix.

Figure 1: Graph Representation: Nodes and edges both can have a set of features that are unique for that node or edge.

Graph neural networks (GNN) refer to the neural network architectures that operate on graph data. Each node has some set of features that defines it. For example, in case of a social network these features could be gender, age, country, friends etc. Similarly, edges may also contain some set of features e.g. friendship between two people, distance between two persons’ locations etc.

The GNNs, as shown in Figure 2, work by gathering and passing information between nodes and its neighbours. Nodes receive information from neighbours via edges. Data from these edges are aggregated (using many different techniques like max pooling, averaging etc.) and are passed to an activation function to get new set of embeddings for the node.

Figure 2: Working of GNN, current node embeddings $h_v$ are calculated with the help if neighbouring node embeddings, neighbouring edge features and current node features.

Every node in the initial state has features $X_v$. The embeddings for a node after message passing can be defined as:

$$h_v = f(X_v, X_{neighbours}, h_{neighbours}, X_{neighbours})$$  \hspace{1cm} (2)

Where,

$f$ is a local transition function,

$X_{neighbours}$ denotes the features of the neighbours of $v$,

$X_{neighbours}$ denotes the edge features connected to $v$,

$h_{neighbours}$ denotes the embeddings of the neighbours of $v$.

In the Figure 2 above, $h_v^{(1)}$ is the initial embedding of the node, $h_{neighbours}^{(1)}$ is the aggregated embeddings of its neighbours. Combining these and passing to the node’s activation function will provide the new embeddings for the node $A$. In different steps, as shown in Figure 3, each node gets a new set of embeddings for itself. After various iterations or K layers of message passing, a node learns more and more about its neighbourhood and its distant neighbours as well.

Figure 3: GNN-produced node level, edge level and graph level predictions.

QSAR relations can be utilised and built upon to extract information from structural composition of a molecule. We could input a molecule to a simple feed-forward recurrent neural network; the predicted molecular properties would be as a result of the model learning the sequential grammar of molecular structure (SMILES strings) rather than extracting information from structural properties of it.

A molecule itself could be viewed as a graph as depicted in Figure 4. The atoms correspond to the vertices and edges corresponds to the bonds between those vertices / atoms. Vertices can contain features as electronic configuration, formal charge, etc. Edges may contain features like bond type, bond distance, etc.

Figure 4: Molecules represented as graphs with individual atoms representing the nodes of the graph and bonds representing the edges of the graph.

2 DATASETS AND ARCHITECTURE

2.1 Datasets and Pre-Processing

Three MoleculeNet datasets have been used to train and test the model (Wu et al., 2018).
a) **ESOL:** (1128 molecules) Water solubility for common organic molecules (Wu et al., 2018).

b) **Free Solv:** (642 molecules) Experimental and calculated hydration free energy of molecules in water (Wu et al., 2018).

c) **Lipophilicity:** (4200 molecules) Octanol/water distribution coefficient (Wu et al., 2018).

Using rdkit library molecules are pre-processed to attach the following atomic properties to the individual atoms in the molecule: Atomic Number, Chirality, Degree, Formal Charge, Number of Hydrogens, Number of Radical Electrons, Hybridization, Aromatic or not, Atom inside the ring or not. Bond type, stereo configuration and conjugation were the three properties chosen as the bond properties that the edges of the graph will hold.

### 2.2 Model Architecture

The model, which we call as FEAT, consists of five message passing layers as listed in Table 1. The number of layers itself is a hyper parameter hence can be changed accordingly whether the data under fit or over fit the model. The first message passing layer has an input dimension of which refers to the 9 atomic properties that we shortlisted for the data. These message passing layers constitute of Graph Convolution Network layer (Kipf and Welling, 2017) with tanh activation function (Wang et al., 2020). The dimensions of hidden message passing layers are hyper-parameters too, here 64 has been chosen as the dimension of hidden message passing layers.

A concatenation of global mean and max pooling layer was added to aggregate the features together. The final layer consists of a linear layer which takes in the output of global pooling layer as input and outputs the value of the predicted property.

The tanh activation function was chosen over ReLu or Sigmoid due to its higher performance for convolutional systems (Wang et al., 2020), (Wieder et al., 2020).

### 3 RESULTS AND DISCUSSION

The model FEAT was run on Dataset-1 (ESOL) and the plot shown in Figure 5 is obtained, it contains a graph that has been plotted between Mean Square Error and Number of Epochs for the ESOL dataset, we can see a steep loss in the mean squared error value which indicated the training going in the right direction with a good learning rate.

To evaluate the predictions, 128 data points (1 batch) from the test dataset (20% of the whole dataset) were plotted against the actual experimentally determined values. The Figure 6 presents the graph between the experimentally determined solubility values vs the solubility values predicted by the model. The line $y = x$ has been added to visualize the accuracy of the predicted property values by FEAT as compared to experimentally determined values.

![Figure 5: MSE vs Epoch for ESOL dataset, good descent for loss observed for the MSE function as epochs increase.](image-url)
Similar graphs were obtained for model run on Dataset-2 (Free Solv) and Dataset-3 (Lipophilicity). In both the cases, the model FEAT seems to underfit the data. Nevertheless, various hyper-parameters can be tweaked and experimented with to generate more accurate results.

4 CONCLUSIONS AND SCOPE FOR FUTURE RESEARCH

It is known that physiological properties of the chemicals can be related to structural properties of the chemicals by some functions as we call these relations as QSAR relations. QSAR relations are not usually linear in nature, they might be very complex in higher order polynomials, hence we try to take help of artificial neural networks in achieving this task. As molecules can be views as graphs present in nature, the bonds correspond to the edges and atoms correspond to the nodes in the graph. It makes sense to use Graph Neural Network to extract structural information from the molecular graphs and use this information to predict various properties from it.

The graph neural network based model FEAT was designed by having five linear layers of Graph Convolutional Network Layers with a concatenated Global Mean and Max pooling layer with a final linear layer to predict the property. The presented model FEAT gave nearly accurate results. In addition, it promises to be a fast and feasible means to predict molecular properties in comparison to conventional methods based off manual experimentation by chemists. Considering the present state of this research, it definitely cannot be taken as an alternative to conventional experimentation; however, it could potentially serve as a means for the required predictions in environments or situations that are resource- and time-constrained. As a scope for future research in this relatively less researched domain, the model can be further improved by adding more message passing layers, experimenting with different learning rates, changing atomic properties chosen for atoms, etc.

REFERENCES


