RESEARCH ARTICLE

A Graph Sequence Neural Architecture for Code Completion with Semantic Structure Features

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Summary

Code completion plays an important role in intelligent software development for accelerating coding efficiency. Recently, the prediction model based on deep learning has achieved good performance in code completion task. However, the existing models cannot avoid three drawbacks:(i) In the existing models, the code representation loses the information (parent-child information between nodes) and lacks many effective features (orientation between nodes). (ii) The known code structure information is not fully utilized, which will cause the model to generate completely irrelevant results. (iii) Simple sequence modeling ignores repeated patterns and structural information, and cannot capture the characteristics of correlation and directionality between nodes. In this paper, we propose a Code Completion approach named CC-GGNN, which is graph model based on Gated Graph Neural Networks(GGNN) to address the problems. We propose a new architecture to obtain the effective code features from code representation. In order to utilize the known information, we propose Classification Mechanism, which classifies the representation of the node using the known parent node and trains training graph in the model. The experimental results show that our model outperforms the state-of-the-art methods MRR@5 at most 9.2% and ACC at most 11.4% in datasets.

KEYWORDS:

Code Completion; Deep Learning; Program Comprehension.

1 | INTRODUCTION

As the scale of software continues to increase, developers benefit from code completion to accelerate software development intelligently. It makes use of the existing code in the context to recommend the next code token, such as method calls or object fields. Since the source code has obvious repeatability and predictability¹, probabilistic statistical model can obtain this information well. N-gram^{1,2,3} is the most widely used probabilistic statistical models in code completion. However, the input of the probabilistic statistical model is simple code tokens, which lacks the semantic structure information of the source code. Besides, the source code files are written by different programmers and have no fixed structure. For example, Python has a more casual programming style than Java. These reasons lead to the low accuracy of the prediction model, and we may even get the opposite result. To address these problems, the higher-order probabilistic model of probabilistic high-order grammar model (PHOG)⁴, which uses context-free grammar to transform the source code into an Abstract Syntactic Tree (AST) as model input. AST not

only reflects the grammatical structure of the code, but also contains important lexical information. The AST nodes may represent identifiers and strings in the source code. With the development of deep learning in natural language processing tasks, many researchers believe that code is also a natural language. Recurrent Neural Network models (RNN)^{5,6,7} are the most common neural network technique for next code prediction. Source code is represented as a node sequence or many node paths in these models. The model computes the probability of the candidate node and recommends the AST node with the highest probability.

However, how to effectively combine the syntactic information and structural features of the code to better complete the code is still a problem in the above model. There exist three fundamental problems not well solved in this regard.

The first one is the representation of code. The previous works are mainly divided into three categories for code representation: code tokens, AST sequence, and AST node's path. The code token models^{8,9} just uses statistical influence of the previous n tokens on the predicted value. The AST sequence model⁶ obtains the sequence of the AST through the depth-first traversal method. Although the extracted sequence information is rich, the hierarchical information of the node is not represented in the sequence. The same problem also exists in the AST path model^{10,11,12}, which only get the path between the terminal nodes. To address this limitation, Liu et al.¹³ uses the root path and AST sequence to get the hierarchical structural information. However, the sequence models lost many information. The more forks of the AST, the more information lost in the sequence code representation. Therefore, we directly convert the AST of the node into a directed graph. AST Graph will completely retain the structural information between nodes, including directions.

The second problem is how to utilize the known structural information in node representation. In previous works^{6,11}, researches directly brings the node's sequence into the model for calculation. In order to make full use of the existing information, we classify these ASTs by their parent nodes. The prediction node with different parent nodes should consider different candidate values, such as *number* and *string*. This problem was not considered in before works. Experimental results show that this operation can improve the effect of the prediction, and the prediction for some parent nodes in the type prediction can reach 100% Accuracy(Section 6.3).

Finally, the structural information of the AST provides a strong indication of the dependencies between link nodes that should be considered. However, simple sequential modeling ignores repetitive patterns and structural information, while undirected graph modeling ignores sequential and repetitive patterns. In this paper, we combine the correlation and directionality between AST nodes to model together.

To address the above problems, we propose CC-GGNN model for code completion with GGNN by integrating AST into a comprehensive code completion model that can leverage both semantic and syntactical information of code. CC-GGNN model use classification mechanism to classify ASTs by known parent node and transform them into different training graph. Finally, our model use the AST structure can effectively obtain the format of code repetitive pattern and obvious structural information. The GRU structure in GGNN can help us solve the problem of long-distance dependence.

To evaluate the performance of our proposed model, we conduct experiments on two real-world datasets, including Python and JavaScript. The data that support the findings of this study are openly available in previous work^{4,6}. We compared CC-GGNN model with the state-of-the-art models: For the node's type prediction, our model achieves the best accuracy of 84.7% and 90.2% on Python and JavaScript datasets respectively, which improves the state-of-the-art methods from 2.3% to 11.4% in PY50k dataset and improves the state-of-the-art methods from 3.2% to 10.1% in JS50k dataset. For the node's value prediction, our model achieves the accuracy of 73.5% and 78.8% on Python and JavaScript datasets respectively, which improves the state-of-the-art methods from 1.6% to 9.0% in JS50k dataset and improves the state-of-the-art methods from 1.8% to 9.0% in JS50k dataset. Statistical testing shows that the improvements over the baseline methods are statistically significant.

To sum up, the contributions of this work are as follows:

- We extract the AST representation from the predicted node, which contains the structure of the node in the source code. Construct a directed training graph that contains rich structural information. The graph representation can effectively obtain the format of code repetitive pattern and obvious structural information.
- We use the known structure information to divide the training data into different training graphs. The prediction nodes with different parent nodes are considered separately. This classification mechanism can narrow the range of candidate values and improve the accuracy of prediction.
- We evaluate our proposed model on two real-world datasets. Experimental results show that CC-GGNN achieves the best performance compared with the state-of-the-art models.

The paper is organized as follows. We give a motivating example in Section 2. Sections 3 introduces the relevant background. The proposed apporach is explained in Section 4. In Section 5 describes about dataset, data processing and baselines, respectively.

The evaluation and discussion are introduced in Section 6. Section 7 discuss the threats of experiments. Related works are addressed in Section 8. Finally, conclusions and future work are presented in Section 9.

2 | MOTIVATING EXAMPLE

As shown in Figure 1, it shows a snippet of Python code and the corresponding AST. Each AST consists of terminal nodes and non-terminal nodes. The terminal node format is *type: value*, which appears in the AST leaf node. Meanwhile, the non-terminal node format is *type: EMPTY*, which is the intermediate node of the AST. The setting of *EMPTY* is the same as the previous work⁶(not shown in Figure 1). The traditional sequence model obtains the code representation by depth-first traversal of the AST, but this code representation method will cause information loss due to the AST's node bifurcation. For example, the node *for* of the second for loop in Figure 1, its parent node should be *Module*, but the parent node becomes *break* in the sequence model. The more branches of the AST, the more information is lost. In this short example, there are 29 node branches. In order to avoid information loss, we input AST directly into the model as graph structure data. Such code representation can make full use of the structural information of the AST, and can effectively avoid the loss of node information.



FIGURE 1 An example of Python program and their corresponding AST.

Secondly, in order to make full use of the known information, we process the parent node of the predicted node. Through the parent node classification mechanism, we can construct the most relevant training data and candidate values. For example, the node *30* in code snippet, we know the type of the predicted terminal node: *num*. Then, more attention should be paid to the number in the next prediction, and the candidate value of the type string should not be considered. This known information can help us complete the code completion task through the classification mechanism.

Repetitive codes have obvious repetitive structures in the tree structure of AST, and these repetitive structural features are difficult to capture in the sequence model as a sequence code representation. Using AST to construct training graphs with tree structure data as input can clearly obtain repeated structures. Combined with the GGNN model, it has strong fitting performance for nonlinear structure data, and the model is more sensitive to repeated structures. For example, the two *for* loop codes in the example can easily predict the predicted *NameLoad: range* of the second *for* loop by using the structure information of the repeated AST.

3

3 | BACKGROUND

3.1 | GGNN

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The Gated Graph Neural Network (GGNN) is a classical spatial domain message passing model based on Gate Recurrent Unit (GRU). In order to extend the neural network method for processing graph structured data, Scarselli et al.¹⁴ proposed the vanilla graph neural network. Li et al.¹⁵ further introduced gated recurrent units and propose Gated Graph Neural Network in 2015. And Daniel et al.¹⁶ proposed a new variant of the graph model. Formally, the GGNN model update functions are given as follows:

$$h_v^{(1)} = [x_v^T, 0]^T \tag{1}$$

$$a_{v}^{(t)} = A_{v:}^{T} [h_{1}^{(t-1)T}, ..., h_{|v|}^{(t-1)T}]^{T} + b$$
⁽²⁾

$$z_{v}^{t} = \sigma(W^{z}a_{v}^{(t)} + U^{z}h_{v}^{(t-1)})$$
(3)

$$r_{v}^{t} = \sigma(W^{r}a_{v}^{(t)} + U^{r}h_{v}^{(t-1)})$$
(4)

$$\widetilde{h_v^{(t)}} = tanh(Wa_v^{(t)} + U(r_v^t \odot h_v^{(t-1)}))$$
(5)

$$h_{v}^{(t)} = (1 - z_{v}^{t}) \odot h_{v}^{(t-1)} + z_{v}^{t} \odot h_{v}^{(t)}$$
(6)

 x_v^T is graph node annotations, which bring into hidden state to calculate the hidden layer vector of the first layer, and pads the rest with zeros in Equation (1). Each node in the graph accepts hidden state information from neighboring nodes and transmits similar information to neighboring nodes too. The model passes message between different nodes of the graph via incoming and outgoing edges, which dependents on the edge type and node direction. The incoming and outgoing connection matrix combined final connection matrix $A_{v:}^T$ corresponding to node v in Equation (2). z_v^t is the update gates, which determines the next step to update the hidden layer information. Equation (4) is reset gates and r_v^t controls reset node's hidden layer information $a_v^{(t)} \in \mathbb{R}^{2D}$ contains activations from edges in both directions. The remaining are GRU-like updates that incorporate information from the other nodes and from the previous timestep to update each node's hidden state. Equation (3) and Equation (4) are the update gates z and reset gates r, which σ is the sigmoid function $\sigma(x) = 1/(1 + e^{-x})$, and \odot is element-wise multiplication in Equation (5). Finally, we get the final updated node status $h_v^{(t)}$ by Equation (6). Li et al.¹⁵ found this GRU-like propagation step is more effective then vanilla recurrent neural network-style in experiments.

The GGNN model can not only obtain the structural features of the abstract syntax tree, but also use the GRU-like propagation step to increase the long-distance dependency of the node's information. So it is suitable for solving related problems of code semantic structure.

3.2 | AST Graph

To learn from sources programs, we need to find a suitable representation that captures semantics features of ASTs. In previous work, one way to flatten each AST as a sequence of nodes in the in-order depth-first traversal^{6,16}. The sequence code contains all the information of the source file, the most relevant node information of the predicted node, such as the father and brother nodes. However, the code of sequence representation only keep the order of nodes, it lacks the hierarchical information of the node and lost many nodes parent information. Another representation is to decompose the path between nodes in the AST. An AST path is a path between nodes in the AST, starting from one node, ending in another node, and passing through an intermediate non-terminal in the path which is a common ancestor of both terminals. However, these methods of code representation have some shortcomings, such as the loss of structure information, insensitivity to repeated structure, and insufficient utilization of known information. So, we directly use the node's AST to construct the training data graph. Each node can be represented by the AST which constructed by the node before it. We use these nodes' AST to construct the AST graph. Pairs of connected nodes are parent-child relationships that can retain structural characteristics in the AST. Besides, the AST Graph will keep the parent node information between nodes. More formally:

Definition 1 (AST Graph): An AST is a tree structure consisting of k nodes: $[node_1, node_2...node_{k+1}]$. The node $node_{k+1}$'s AST can be represented by the partial AST constructed by the previous k nodes. Through the classification of the $node_k$, all

nodes' ASTs are divided into different sets: $[set_1, ..., set_i]$, for $i \in [2...t]$, each set_i of nodes' ASTs builds a training graph together. Every two adjacent nodes are parent-child or child-parent relationship in each AST. Duplicated nodes in the ASTs are merged into one node in the AST Graph.



(a) Nodes' AST with parent node: n₁ (b) AST Graph

FIGURE 2 An example of AST Graph

In this paper, we have the similar points as the code representation in the previous related work²¹: 1. Keep the parent-child relationship of each node, and de-duplicate the nodes to build a training graph. 2. The weight of node's edge is the frequency of the occurrence of the edge in the corresponding AST.

We also have unique characteristics of code representation: 1. We consider the directions between nodes, and the weights of edges between directions are also different. The direction of the node can effectively maintain the structural characteristics and hierarchical information. For example, an undirected graph can only determine the parent-child relationship between two nodes, but cannot determine who is the parent node. 2. Classification mechanism, we combine the characteristics of predicting nodes with different parent nodes, and divide the data into different training graphs. At the same time, we will also filter the candidate values of these graphs. For example, the candidate values of graphs whose parent nodes are *Num* and *Str* are different. We will introduce it in the Section 4.

Figure 2 shows the process of converting nodes' AST to AST graphs. Figure 2(a) is the AST set with the parent node n_1 . Through the connection between nodes and removing the duplicate nodes, we can obtain the training graph (Figure 2(b)) with the parent node of the predicted node n_1 .

4 | PROPOSED MODEL

In this section, we introduce CC-GGNN model in detail. We formulate the problem at first, then explain how to construct the graph from node's AST. Meanwhile, we describe our approaches to representing learning node embedding on AST graphs, show how they are integrated into a embedding vector. As illustrated in Figure 3, it includes two steps: 1. Data processing, 2. Graph representation and model prediction.

4.1 | Problem Definition

Code completion is to predict the next node in source code. We get each node's AST to construct the train graph G. For prediction AST P_ast , we output probabilities \hat{y} for all possible nodes, where an element value of vector \hat{y} is the recommendation score of the corresponding node. The node with *Top-K* types T_i and values V_j in source code will be the candidate nodes for prediction⁶. Next node's type and value prediction process are transformed into the math problem of finding the maximum \hat{y} in candidate nodes, as shown in the following Equation (7), (8).

$$\exists i \in \{1, 2, 3, ...s\} : \arg\max_{T_i} \hat{y} (G, T_i, P_ast)$$
(7)



FIGURE 3 The overall architecture of the model

$$\exists j \in \{1, 2, 3, ...k\} : \arg\max_{V_j} \hat{y}(G, V_j, P_ast)$$
(8)

4.2 | Constructing AST Graphs

Each node will get a corresponding node's AST *p* in source code(Section 3.2). In the AST *p*, each node represents an code node $v_i \in V$ and these nodes are transformed into connected edges in the graph, that is, adjacent node (v_{i-1}, v_i) means edge $(v_{i-1}, v_i) \in E$ in graph. And these AST information will be jointly converted into a training graph *G*.

In Figure 3, according to the parent node n_1 of the test AST, we filtered (data processing) the training data's AST to construct a training graph. The test data does not participate in the construction of the training graph. The red nodes in the graph are just the reflection of the test data in the training graph. The operation of converting ASTs classification into multiple graphs can avoid the excessive number of nodes in one train AST graph, and can reduce the impact of different AST on the accuracy of the predicted node. After we get the training graph, each node can be embed as a vector of fixed dimensions by GGNN model: $v_i \in \mathbb{R}^d$, where d is the dimensionality. Each p can be represented a vector based on those node vectors, we will introduce it in detail in next section.

4.3 | Learning Node Embedding

In this section, we will learn each node vector in graph. For example, the graph *G* and the matrix *A* are shown in Figure 4. In directed graph *G*, the *A* is defined as the concatenation of two adjacency matrices $A^{(out)}$ and $A^{(in)}$, which represents weighted connections of outgoing and incoming edges in the graph respectively. The sparsity structure matrices corresponds to the edges of the graph and these matrices help us calculate the vector of nodes.

For each graph G, the GGNN model gets node's vector. According to the graph, we rewrite Equation (2):

$$a_i^t = A_{i:}[v_1^{t-1}, ..., v_n^{t-1}]^T H + b$$
(9)



FIGURE 4 An example graph's connection matrix.

The $[v_1^{t-1}, ..., v_n^{t-1}]$ is the list of node vectors in p, where $H \in \mathbb{R}^{d \times 2d}$. In GGNN model, Equation (2) can propagation the information between different nodes by matrix A. Each node in the graph accepts hidden state information from neighboring nodes and transmits similar information to neighboring nodes too. z_v^t is the update gates, which determine hidden layer information update in next step. Equation (4) is reset gates and r_v^t controls reset node's hidden layer information. Equation (3) and Equation (4) will determine which node's information will be passed on. Using the GRU's update gate, the Equation (6) is the combination of the previous hidden state and the candidate state. After updating all nodes in AST graphs until convergence, we can obtain the final node vectors by Equation (10).

There are several types of outputs that GGNN model produce in different situations. In this task, we get each node's embedding vector by Equation (10).

$$o_v = g(h_v^T, x_v) \tag{10}$$

For each node $v \in V$ output node scores and applying a softmax over node scores. In Equation (10), g is a specific function, using the final state h_v^T and initial state x_v of each node to calculate its output. The final state h_v^T is derived from Equation (6).

4.4 | AST Embedding and Prediction

In our model, the predicted node is represented by its related nodes. It similar to the previous path and sequence-based method, after we transform each node into a different vector, we convert the AST composed of nodes into a vector of fixed dimensions. However, the previous work is only a simple vector addition calculation, and it is hard to obtain effective features. So, we plan to apply a new strategy to combine long-term preference and node information.

We obtain the node of vectors by CC-GGNN model, then generate global vector together. The representation of each graph is an embedding vector $P_s \in \mathbb{R}^d$, which can divide in two parts: 1.local embedding P_1 . 2. global embedding P_g .

In AST, the parent node contains the most abundant information of the predicted node, such as parent-child characteristics. So, the local embedding defined as the parent node's vector. For example, in the test AST in Figure 3(red AST structure), the parent node is n_1 , $P_1 = n_1$.

Then, we aggregate all node vectors to get global embedding p_g in AST. The model obtain the priority of node information by adding an attention mechanism, which shows in Equation (11)

$$\alpha_i = W_1^T \sigma(W_2 n_1 + W_3 v_i) \tag{11}$$

The parameters $W_1 \in \mathbb{R}^d$ and $W_2, W_3 \in \mathbb{R}^{d \cdot d}$ control the weights of item embedding vectors. And, the global embedding vector is calculated by parameter α_i .

$$p_g = \sum_{i=1}^n \alpha_i v_i \tag{12}$$

Finally, we compute the hybrid embedding p_h by taking vector transformation over the vector splicing of the local vector P_1 and global embedding vector $\alpha_i v_i$:

$$P_1 = n_1 \tag{13}$$

$$P_h = W_4[n_1; \sum_{i=1}^n \alpha_i v_i] = W_4[P_1; P_g]$$
(14)

where matrix $W_4 \in \mathbb{R}^{d \cdot 2d}$ control combined vectors into the latent space \mathbb{R}^d . Finally, we get the embedding of each global representation p_h from the model. Then, we compute the score \hat{y}_i for each candidate node $v \in V$ by multiplying its embedding v_i with P_h for prediction:

$$\hat{y}_i = softmax(P_h^T v_i) \tag{15}$$

where $\hat{y} \in \mathbb{R}^m$ denotes the probabilities of nodes appearing to be the next node in AST. And, We get the output vector of the model by softmax function, which is Normalization function in output layer of the neural network. For each graph, the loss function is defined as the cross-entropy of the prediction and the ground truth. It can be written as follows:

$$Loss(\hat{y}) = -\sum_{i=1}^{m} y_i log(\hat{y}_i) + (1 - y_i) log(1 - \hat{y}_i)$$
(16)

In Equation (16), the y_i denotes the one-hot encoding vector of the ground truth node. And, we use the Back-Propagation Through Time (BPTT) algorithm to train the proposed graph model. In the prediction process, the graph model built by classification can improve the screening of more relevant candidate values.

5 | EXPERIMENT SET UP

5.1 | Data Availability Statement

The data that support the findings of this study are openly available in Machine Learning for Programming at http://plml.ethz.ch. The data¹⁷ consists of two parts: 150k Python Dataset and 150k JavaScript Dataset.

5.2 | Dataset and Preprocessing

We evaluate different approaches on benchmarked datasets: Python(PY150) and JavaScript(JS150). PY150 is summerized in Table 1 and JS150 is summerized in Table 2.

As shown in Table 1 and Table 2, it is overall dataset statistics. In the type prediction task, there are only 181 types in training data and test data in Table 3. Such as *NameLoad*, *alias*, *NameParam*, etc. These types are determined by the Python programming language and cannot be defined by the programmer, which results in fewer candidate values. In the value prediction task, the source file has $3.4*10^6$ different node values in Table 4. There are arbitrary possibilities for encoding the program text. The value can be any program identifier (such as *None*, *format*), literals (such as 0.035, 1075), program operators (such as /, -, *), etc. It is impossible to use all of them for calculation, especially some of these values only appear once, so we need to filter the vocabulary. Compared with the PY150 dataset, the structure of the JS150 dataset is simpler. The JS150 only contains 44 types of JavaScript language customization, and $2.6*10^6$ different value candidates.

The number of unique node values in dataset is too large to directly apply neural networks models, thus we only choose K most frequent values in training set to build the global vocabulary, where K is a free parameter. In order to effectively compare with the baseline experiment, we set the value of K to 1k, 10k, and 50k. We further add two special values: *EMPTY* being the value of non-terminal AST nodes and mark all out-of-vocabulary node values in training set and test set as *UNK*. The number of nodes in the training graph is limited by the parameter K. When K=1k, the number of nodes in the AST graph is the least, and there are only 1000 nodes at most. When K=50k, the number of graph nodes is the largest, and the largest graph has more than 10,000 nodes.

Classification mechanism: This mechanism is a method of data processing. We obtain the known parent node of the predicted node as a classification node, and divide the AST of each node into different sets. We convert each set into a training graph and combine the attention mechanism (Section 4.4) to predict the completion result. For example, if we predict the node $(n_1 :?)$ in Figure 2, the known information is that the parent node is n_1 , and obtain the training graph constructed by n_1 to predict the type.

TABLE 1 Dataset statistics for PY

	Category	Size
1	Training files	1.0*10 ⁵
2	Test files	5.0*104
3	leaf nodes	1.6*10 ⁷
4	Non-terminal nodes	1.4*10 ⁷
5	Training Queries	6.2*10 ⁷
6	Test Queries	3.0*107
7	Type Vocabulary	181
8	Value Vocabulary	3.4*10 ⁶

TABLE 3 TYPE Nodes type for PY

TABLE 2 Dataset statistics for JS

	Category	Size
1	Training files	$1.0*10^5$
2	Test files	5.0*10 ⁴
3	leaf nodes	1.6*10 ⁷
4	Non-terminal nodes	$1.4^{*}10^{7}$
5	Training Queries	$10.7*10^7$
6	Test Queries	5.3*10 ⁷
7	Type Vocabulary	44
8	Value Vocabulary	$2.6*10^{6}$

TABLE 4 VALUE Nodes type for PY

	Types	Size
1	NameLoad	1.8*10 ⁷
2	attr	9.2*10 ⁶
3	AttributeLoad	8.4*10 ⁶
4	Str	$7.2^{*10^{6}}$
5	Call	6.9*10 ⁶
6	Assign	3.9*10 ⁶
181	CompareLtELtELtE	1

	Types	Size
1	self	$4.1*10^{6}$
2	0	6.3*10 ⁵
3	None	6.1*10 ⁵
4	1	5.5*10 ⁵
5	True	4.6*10 ⁵
6	name	3.5*10 ⁵
$3.4*10^{6}$	http://github.com/lepture/flask-wtf	1

We can transform the most relevant AST into the one graph through the classification mechanism. And, we can narrow down the candidate values of each graph by classification mechanism. For example, the candidate value ranges of Num and Str as the parent node are different. We traverse the training data to find the candidate value of each parent node, and intersect the K candidate values, the remaining values are the new candidates. We can avoid meaningless calculations by this process.

Through experiments, we found that the types of terminal nodes and non-terminal nodes in the PY150 are divided into 153 graphs, of which there are 141 types of terminal nodes (Section 6.3), and value can be divided into 15 types. The types of terminal nodes and non-terminal nodes in the JS150 are divided into 44 graphs, of which there are 33 types of terminal nodes (Section 6.3), and value can be divided into 16 types.

5.3 | Baselines

As there are already prior investigations conducting code completion on the benchmarked dataset, to validate the effectiveness of our proposed approaches, we need to compare them against the state-of-the-art. We compare CC-GGNN model with five code completion models from previous work and these baselines can be divided into two categories: 1.Probabilistic Model. 2.Deep learning Network.

Probabilistic Model DEEP3¹⁸: The approach is general and can be used to learn a probabilistic model of any programming language. Model implemented approach in a system called DEEP3 and evaluated it for the challenging task of learning probabilistic models of JavaScript and Python. DEEP3 model achieves the highest prediction accuracy of the probabilistic models in code completion task.

VanillaLSTM^{6,19}: The model uses LSTM to extract AST node features from the flattened AST sequence. The input of each LSTM unit is the concatenation of the previous hidden state and the type and value embedding of the current AST node. It is a standard LSTM network without any attention or pointer mechanisms.

Pointer Mixture Network⁶: The model is an an attention and pointer generator network-based code completion method. Model inspired by the prevalence of locally repeated terms in program source code, and the recently proposed pointer copy mechanism, they propose a pointer mixture network for better predicting Out-of-Vocabulary(OoV) words in code completion. Based on the

context, the pointer mixture network learns to either generate a within-vocabulary word through an RNN component(Attentional LSTM model), or regenerate an OoV word from local context through a pointer component.

Transformer-XL¹³: The model is an improved Transformer architecture²⁰ with the flattened AST sequence. In order to make effective use of the code structure, the model adds a path from the predicted node to the root. It is biggest difference with Pointer Mixture Network model in data process part. The naive multi-task learning is used, and task weights are fixed to be equal.

CCAG²¹: CCAG models the flattened sequence of AST as an AST graph. CCAG uses proposed AST Graph Attention Block to capture different dependencies in the AST graph for representation learning in code completion. The sub-tasks of code completion are optimized via multi-task learning in CCAG, and the task balance is automatically achieved using uncertainty without the need to tune task weights.

For a fair comparison, we set batch size as 128, hidden unit size of 1500 and train all model for 8 epochs. The initial learning rate set as 0.001 and decay it by multiplying 0.6 after every epoch in all methods. These configurations are same as Li et al.⁶, because this paper is widely cited. For VanillaLSTM and PointerMixtureNet, the gradient norm is clipped to 5, and the size of context windows is set to 50. For Transformer based methods, we employ h = 6 parallel heads, and the dimension of each head d_{head} is set to 64. We set the segment length to 50 and use a 6-layer layer number Transformer-XL network. We reproduce the baseline: DEEP3, VanillaLSTM, PointerMixtureNet, Transformer-XL. The experimental results are shown in the Table 5. For the CCAG model, our model uses the same parameter configuration for comparison experiments. Embedding size, hidden size and batch size are all set to 128. The experimental results are shown in the Table 6.

5.4 | Evaluation Mertics

We evaluate the quality of code completion using two automatic metrics: Accuracy and MRR@5. In the code completion task, the previous model provides an *Top-1* list of suggestion for each node's type or value in the source code file. Those models use Accuracy to evaluate the performance of our model. The calculation of Accuracy is shown in Equation 17.

$$Acc = \frac{1}{n} \sum_{i=1}^{n} acc_i \tag{17}$$

For the *ith* predicted node, if *Top-1* prediction is ground truth $acc_i = 1$, otherwise $acc_i = 0$. However, in actual code completion tools, models often output more than one reference result. To measure performance on these tasks, we also use mean reciprocal rank (MRR@5). Comparing to the metric (Acc@1), this is closer to the realistic scenario when completion suggestions are presented to developers. The rank is defined as

$$MRR = \frac{1}{n} \sum_{i=1}^{n} \frac{1}{rank_i}$$
(18)

where *n* is the number of predicting locations and $rank_i$ is the rank of the correct label given by the model for the *ith* data point. We present MRR@5 as a percentage, in keeping with prior work²².

6 | EVALUATION AND DISCUSSION

6.1 | Research Questions

To evaluate our proposed approach, we conduct experiments to investigate the following research questions:

RQ1: How does the overall performance of our proposed method in terms of code completion compare with the stateof-the-art model?

We compared the state-of-the-art method in the probabilistic model and the results of the Deep neural network models used sequence and graph as input in the same dataset.

RQ2: For terminal nodes that are more complex and difficult to predict, how effective is our proposed model?

In previous work⁶ for the completeness of prediction, the model artificially added value: *EMPTY* to non-terminal nodes in the process of data processing (Section 5.2). And the non-terminal node has $1.4*10^7$ (about 43%) in the dataset. In other words, about half of the value prediction tasks are *EMPTY*, so in order to further show the prediction effect of the model, we separately increase the value prediction experiment.

RQ3: What is the effectiveness of code representation and classification mechanism for our proposed model?

In this part, we calculate the impact of each component for our proposed model. We experimentally study the impact of the semantic structure information of AST on the model.

RQ4: What is the effectiveness of the hidden size parameter and OoV problem on the experiment?

We analyze the experimental results of the hidden size values of 128, 256, 512, 1024, 1500. And, we discuss the influence of the OoV problem in our proposed method.

6.2 | Overall Performance

For **RQ1**, Table 5 and Table 6 illustrate the overall performance of our combined model compared to baselines. Our combined model obviously outperforms all baselines on Accuracy in six datasets. Comparing to the purely AST sequence based baselines, that is, Pointer Mixture Network and VanillaLSTM, our model achieves obvious improvements, indicating that it is much more capable for modeling ASTs. CC-GGNN compared with the Transformer-XL model, the improvement is more bigger in type prediction task. Our model is also better than CCAG model, which also takes graph data as input.

TABLE 5 Accuracy of various types and values for six datasets.

	JS	1k	JS	10k	JS	50k	РҮ	′1k	PY	10k	PY	50k
	value	type	value	type	value	type	value	type	value	type	value	type
VanillaLSTM	62.6%	74.7%	66.3%	78.4%	69.8%	80.3%	59.2%	68.6%	61.3%	72.9%	63.4%	77.5%
PointerMixtureNet	69.8%	78.5%	73.4%	83.2%	75.3%	85.9%	64.7%	76.4%	68.3%	77.4%	69.2%	78.1%
Transformer-XL	73.9%	84.1%	75.3%	85.7%	77.2%	87.0%	68.1%	78.9 %	70.6%	80.7%	71.9 %	82.4%
CC-GGNN	75.3%	87.1%	77.1%	88.4%	78.8 %	90.2%	69.9 %	80.3%	71.8%	83.2%	73.5%	84.7 %

TABLE 6 Accuracy of various types and values for six datasets with CCAG model.

	JS	1k	JS	10k	JS	50k	РУ	/1k	PY	10k	PY	50k
	value	type	value	type	value	type	value	type	value	type	value	type
CCAG	62.8%	75.7%	66.7%	78.6%	68.2%	80.1%	61.9%	76.7%	63.2%	80.9%	64.2%	75.3%
CC-GGNN	64.1%	78.8 %	67.9%	81.1%	70.5%	83.3%	63.3%	78.4 %	65.7%	81.3%	66.9%	80.1%

Experiments first discuss the application of our model for predicting types of AST nodes, it is essentially a prediction of the code structure. As can be seen from the Table 5, for the node's type prediction, our model achieves the accuracy of 84.7% on the PY50k datasets respectively, which improves probabilistic model VanillaLSTM by 7.2%, improves Pointer Mixture Network by 6.6% and improves Transformer-XL 2.3%. Transformer-XL model can obtain the hierarchical structure information of sequence data, the experimental results are better than the Pointer Mixture Network model.

Compared with the CCAG model in type task, our model narrows the range of candidate values and considers the directionality of the nodes. Therefore, the CC-GGNN model can obtains more AST structure information, and the maximum effect of the model is improved by 3.6% in the JS50K dataset.

For the node's value prediction, our model achieves the accuracy of 73.5% in PY50k dataset. Compared with the Pointer Mixture Network, CCAG and Transformer-XL models, our model has increased by 4.3%, 2.7%, and 1.6% respectively. Through the classification mechanism of the model, we can effectively filter out the most relevant candidate values, which can effectively help us improve the accuracy of prediction. In the value prediction, the predictions of the *UNK* targets are treated as wrong predictions. When the parameter K = 50K, the *UNK* rates for Python is 11%.

6.3 | Terminal Node Prediction

In the Python dataset, there are $1.4*10^7$ non-terminal nodes form the intermediate structure of ASTs and $1.6*10^7$ terminal node form the leaf node. Non-terminal nodes have no value type in original AST, so the most predicted value in the overall experiment is *EMPTY*, which is artificial definition. For **RQ2**, we add experiments to predict 16,003,628 leaf nodes in PY50k and predict 83,118,887 leaf nodes in JS50k. The difficulty of leaf nodes prediction mainly lies in the leaf's value prediction, and the difficulty can be divided into several points:

- The leaf node has a wide range of value candidates. In our training data, the type of value is as much as $3.4*10^6$ in Python dataset, which is much larger than the range of type 181. But, most of them are values that have appeared several times, so we limit the candidates to calculate by the parameter *K* value. Values that are not within the *K* value range are marked as *UNK*.
- The value of the leaf node appears random. Even if the range of candidates is selected by the parameter *K* value, the value of the leaf node appears random. These leaf node values are randomly defined by programmers. If the relevant dataset is small, the prediction results are very poor.
- The value of the leaf node has little correlation with the structure of the AST. The same parent node *Nameload* has tens of thousands of candidate values, the AST structure has little influence on prediction. The prediction of value requires an overall understanding of the entire snippet code, and the dependent nodes may have a long distance.

TABLE 7 MRR@5 of various types and values of leaf token predictions for PY50k and JS50k

	PY:	50k	JS	50k
	value	type	value	type
DEEP3	42.1%	82.6%	49.7	86.5%
PointerMixtureNet	44.8%	85.2%	56.3	89.4%
Transformer-XL	46.4%	87.3%	58.6	90.3%
CC-GGNN	49.3%	91.8%	59.1	93.6%

Overall, Table 6 shows that our model achieved better scores than the probability model DEEP3 in PY50k and JS50k. Besides, our model achieves better scores than deep neural network models: PointerMixtureNet and Transformer-XL on both datasets. Our best performing model is 4.5% higher than the MRR@5 of PointerMixtureNet (from 44.8% to 49.3%), and is 2.9% higher than the MRR@5 indicator of the Transformer-XL model (from 46.4% to 49.3%) in leaf's value task.

TYPE	ACC	MRR@5	TYPE	ACC	MRR@5	TYPE	ACC	MRR@5
Module	81.04%	90.22%	CompareLtE	69.41%	83.21%	CompareLtGt	33.33%	66.67%
ImportFrom	100.00%	100.00%	CompareIn	80.06%	89.20%	AugAssignMod	54.22%	63.72%
alias	100.00%	100.00%	name	100.00%	100.00%	CompareGtGt	39.65%	60.13%
bases	%66.66	%66.66	CompareNotIn	80.31%	89.79%	CompareEqEqEq	54.15%	71.55%
AttributeLoad	99.36%	99.68%	CompareIsNot	98.14%	99.11%	CompareIsIs	90.12%	91.33%
body	94.71%	96.82%	finalbody	100.00%	100.00%	CompareLtLtEEqGtEGtNotEq	100.00%	100.00%
Assign	90.19%	95.30%	GeneratorExp	92.38%	94.54%	CompareLtLtEEqLtLtLt	100.00%	100.00%
Call	68.45%	73.31%	comprehension	99.88%	99.94%	CompareLtELtLtE	75.00%	87.50%
arguments	80.01%	87.62%	BinOpDiv	61.32%	72.65%	CompareEqIs	70.11%	73.49%
args	100.00%	100.00%	UnaryOpUSub	99.87%	99.87%	AugAssignBitXor	50.97%	67.33%
defaults	71.31%	79.05%	IfExp	72.26%	81.33%	CompareLtGtEqGtELtENotEqInNotInIsIsNot	0.00%	0.00%
Return	81.13%	86.62%	BinOpPow	72.62%	84.55%	CompareGtGtGt	100.00%	100.00%
decorator_list	100.00%	100.00%	ListComp	96.21%	97.41%	CompareGtGtE	58.31%	66.75%
TupleLoad	85.36%	93.18%	Print	74.16%	84.24%	AugAssignPow	33.33%	67.63%
Expr	%06.66	99.95%	Lambda	80.15%	89.73%	CompareGtEGtE	50.00%	75.00%
Import	100.00%	100.00%	Assert	61.21%	78.43%	AugAssignRShift	72.65%	84.33%
If	99.75%	99.87%	Set	77.32%	83.22%	CompareGtELt	0.00%	0.00%
SubscriptLoad	%96.66	99.98%	CompareLtELtE	63.18%	64.68%	AugAssignLShift	54.31%	72.13%
Slice	77.36%	83.15%	AugAssignMult	54.21%	67.14%	CompareLtEGtE	100.00%	100.00%
keyword	53.24%	71.45%	AugAssignBitAnd	71.91%	79.94%	CompareLtELtE	50.18%	63.54%
With	99.25%	99.62%	SetComp	100.00%	100.00%	CompareEqGt	50.00%	75.00%
Dict	89.25%	93.09%	AugAssignBitOr	72.08%	81.88%	CompareLtEq	100.00%	100.00%
Index	64.27%	78.88%	DictComp	93.21%	95.36%	CompareIsIsIs	100.00%	100.00%
For	99.93%	96. <u>9</u> 6%	Yield	79.10%	86.25%	CompareNotEqEq	100.00%	100.00%
TupleStore	100.00%	100.00%	Delete	100.00%	100.00%	CompareEqNotEqEq	80.00%	90.00%
UnaryOpNot	92.13%	94.90%	SubscriptDel	100.00%	100.00%	CompareGtEGt	64.37%	76.19%
Raise	89.35%	96.44%	CompareEqEq	43.82%	65.16%	CompareEqNotEq	33.33%	50.00%
compareEq	74.55%	85.85%	Global	100.00%	100.00%	TupleDel	100.00%	100.00%
AttributeStore	%66.66	%66.66	BinOpBitAnd	69.14%	78.31%	CompareEqEqEqEq	50.00%	75.00%
SubscriptStore	100.00%	100.00%	BinOpBitOr	84.31%	92.62%	CompareGtLt	52.13%	73.81%
orelse	50.42%	68.67%	ListStore	100.00%	100.00%	CompareGtEqGt	0.00%	0.00%
BoolOpAnd	90.22%	95.12%	Exec	70.34%	83.33%	CompareGtELtE	0.00%	0.00%
BinOpAdd	68.47%	80.62%	CompareLtLt	59.53%	73.36%	CompareLtEGt	25.00%	25.00%
BinOpMult	57.12%	78.31%	UnaryOpInvert	86.42%	89.34%	CompareLtEGtGt	0.00%	0.00%

TABLE 8 The *type* prediction result of the leaf node

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0.00%	0.00%	0.00%	100.00	100.009	100.009	0.00%	50.00%	100.009	0.00%	0.00%	0.00%	0.00%
0.00%	0.00%	0.00%	100.00%	100.00%	100.00%	0.00%	0.00%	100.00%	0.00%	0.00%	0.00%	0.00%
CompareInIn	CompareInIs	CompareLtGtE	CompareNotEqNotEq	CompareNotEqNotEqNotEq	CompareIsIsIsIsIs	CompareLtLtLtLt	CompareNotEqNotEq	CompareEqGtEq	CompareGtGtGtGt	CompareLtLtLt	CompareIsNotIsNot	CompareInNotIn
82.17%	80.10%	70.43%	73.56%	100.00%	81.49%	70.11%	90.13%	76.98%	87.66%	50.00%	85.55%	58.73%
64.38%	65.95%	60.58%	66.42%	100.00%	69.63%	48.14%	89.31%	64.52%	80.31%	50.00%	75.00%	39.67%
BinOpFloorDiv	AugAssignDiv	BinOpRShift	UnaryOpUAdd	Repr	BinOpLShift	AugAssignFloorDiv	AttributeDel	CompareLtLtE	BinOpBitXor	CompareEqEqEqEqEqEq	CompareEqEqEq	CompareLtELt
87.43%	95.43%	89.41%	100.00%	84.42%	76.19%	90.74%	83.46%	78.55%	87.35%	76.95%	83.48%	99.67%
85.00%	91.07%	81.77%	100.00%	82.55%	63.02%	82.45%	79.35%	70.34%	76.12%	73.01%	70.49%	99.39%
ListLoad	BinOpMod	CompareGt	type	While	CompareNotEq	AugAssignAdd	BoolOpOr	BinOpSub	CompareGtE	CompareLt	AugAssignSub	CompareIs

TABLE 8 The *type* prediction result of the leaf node(Continued))

Our model improves the type prediction more obviously. For PY50k dataset, the experimental result of MRR@5 is 91.8%. For JS50k dataset, the experimental result of MRR@5 is 93.6%. We will further show the experimental results in the Tables 8 and Tables 9.

For the type prediction of leaf nodes, our model divides all ASTs into 141 different training graphs by the known parent nodes of the leaf nodes. Each training graph contains different number of AST. The training graph constructed with *AttributeLoad* has 5003371 nodes' AST, which contains the largest number of nodes in all training graphs. Our approach has achieve high accuracy and MRR@5 scores. There are also training graphs with only one AST, such as *CompareNotEqNotEqNotEq* and *CompareIsIsIsIsIs*. Because the number of training data is too small, the prediction Accuracy and MRR@5 value of the model decrease.

The detail of experimental results of the model are shown in Table 8 and Table 8. In PY50K dataset, the experimental results show that the model can achieve 100% prediction accuracy for 28 training graphs such as *ImportForm*. These nodes account for approximately 8.8% of the total and mark as gray in the table. But there are also 13 training graphs constructed by parent nodes whose prediction accuracy is zero. We found that these training data are less than a hundred ASTs through analysis. Most of them only appeared once. In other words, the training data rarely leads to low prediction accuracy.

TYPE	ACC	MRR@5	SIZE	TYPE	ACC	MRR@5	SIZE
VariableDeclarator	72.85%	85.74%	9.14*10 ⁵	CallExpression	85.64%	91.85%	$1.21*10^{7}$
FunctionDeclaration	99.99%	99.99%	$1.10^{*}10^{6}$	AssignmentExpression	89.07%	93.01%	$4.56*10^{6}$
ArrayExpression	93.48%	95.73%	$4.64*10^{6}$	ExpressionStatement	95.16%	96.86%	$6.28*10^4$
MemberExpression	94.35%	97.34%	3.53*10 ⁷	FunctionExpression	98.36%	99.65%	$3.06*10^{6}$
ArrayAccess	82.36%	87.64%	$3.96*10^{6}$	UnaryExpression	92.28%	93.62%	$8.24*10^5$
NewExpression	89.72%	92.45%	$7.52*10^5$	BinaryExpression	77.82%	82.43%	$7.04*10^{6}$
Property	76.55%	79.40%	$3.58*10^{6}$	IfStatement	86.16%	88.47%	$4.30*10^5$
ConditionalExpression	81.01%	86.18%	$7.09*10^5$	LogicalExpression	67.87%	72.73%	$7.74*10^5$
ReturnStatement	78.65%	83.84%	$7.80*10^5$	VariableDeclaration	100.00%	100.00%	$1.01*10^{6}$
BlockStatement	82.37%	87.55%	3.94*10 ⁵	UpdateExpression	100.00%	100.00%	$4.08*10^5$
ForInStatement	99.78%	99.81%	$1.18*10^{5}$	ForStatement	91.14%	93.32%	7.46*10 ⁴
WhileStatement	76.25%	81.41%	$1.56*10^4$	CatchClause	95.89%	97.10%	$7.87*10^4$
TryStatement	99.39%	99.60%	$6.09*10^4$	SwitchStatement	98.64%	99.17%	$2.00*10^4$
SwitchCase	84.89%	88.10%	3.59*10 ⁵	ThrowStatement	95.02%	96.68%	$1.98*10^4$
DoWhileStatement	65.64%	69.41%	2914	Program	99.52%	99.56%	7784
SequenceExpression	89.02%	92.47%	$3.44*10^4$	LabeledStatement	50.00%	75.00%	8
AssignmentPattern	44.44%	74.07%	9				

TABLE 9 The type prediction result of the leaf node

Compared with the PY50k dataset, JS50K is relatively simple. Although JS50K has more leaf nodes, there are fewer types and value categories. There are only 33 types of parent nodes of leaf nodes, which greatly reduces the difficulty of prediction. It can be seen from Table 8 that the *VariableDeclaration* and *UpdateExpression* training graphs of the parent node marked in gray can achieve a prediction accuracy of 100%, accounting for 1.71% of all leaf nodes.

Case Study: Figure 5. is a case in CCAG, it is contained in the test set of Python. The code segment is used to illustrate how the incoming email is handled in Google Cloud. NameStore:body highlighted with green is the next node to predict.

First, it is easily find that the parent node of the prediction node is *TupleStore*, after we convert the code file into AST. Combining our classification mechanism and traversing the training dataset, we found that there are only five candidates whose parent node is *TupleStore*. Candidates:[*NameStore*, *AttributeStore*, *TupleStore*, *ListStore*, *SubscriptStore*].

However, these model results in Figure 5. all contain values that should not be considered, we marked by the red box. $CCAG_g$ and PointerMixtureNet models even predict that the maximum probability value is a non-candidate value. Therefore, a large and invalid candidate value table not only wastes computing power and time, but also reduces the accuracy of the model. Transformer-XL mainly gives high probability to the node type on the path from the predicted node to the root in the AST.



FIGURE 5 A code completion example.

Similarly, our model successfully captures the repetitive pattern (the repeated local ASTs) and has a higher probability than the CCAG model. We think there are two reasons: 1. The Gated Graph Neural Network model has a strong nonlinear fitting ability to graph structure data. Besides CC-GGNN model can learn the inherent characteristics of the directions and edges between nodes in the graph structure, which is helpful for accurate prediction of repeated structure data. 2.In the GGNN model, the way of information transmission between nodes is that each node spreads to surrounding nodes, and the information transmitted by the same graph structure data is the same. These reasons led to the success of CC-GGNN captures the repetitive pattern. Another point is that for the MRR@5 metric, our model can always display the predicted value correctly.

6.4 | Each Component Effect on the Model

For **RQ3**, we conducted an ablation study to examine the impact of the two proposed components used in our model: classification mechanism and different AST representations. We conduct experiments without classification mechanism and use the AST representation of the CCAG and Transformer-XL models respectively.

For type prediction task, the results are shown in Figure 6. The blue line shows the results of our full model. The yellow line shows the code representation of our model combined with Transformer-XL model's code representation. The green line shows the our model with CCAG model's undirected graph representation settings. And the red line removes the classification mechanism from the full model. It is obvious that full model is better than other three models. It can be seen that the effect of using the graph model in combination with other AST representations is weaker than the full model. Besides, the model effect of removing the classification mechanism decreases more, which demonstrates that both the classification mechanism and our AST representations are necessary to improve the performance, and classification mechanism contributes more to the improvements.

For value prediction task, the results are shown in Figure 7. The blue line shows the results of our full model. The yellow and green lines present the results of our model with other AST representation, and the red line removes the classification mechanism from the full model. Similar to the experimental results of type prediction, these models have worse results with no classification mechanism and other AST representation.

For **RQ3**, we draw conclusions from the above experimental results. Both classification mechanism and structure characteristics of AST graph representation can effectively improve the predictive ability of the model. The classification mechanism has the most obvious effect on the overall model, and it is effective to increase the attention mechanism to the parent node of the predicted node.

6.5 | Hidden Size and OoV Problem

For **RQ4**, we take the PY50k data set as an example. First, we analyze impact on the model by adjusting the range of hidden size. Then, we discussed the impact of the OoV problem on the value prediction task.

As shown in Figure 8, with the increase of the hidden size parameter, the accuracy of the value and type prediction tasks are improved. The orange line shows that the change of hidden size has no obvious effect on the prediction accuracy of type.



FIGURE 6 Effectiveness of each component in type prediction task

However, the trend of value prediction task changes is more significant from line blue. The reason is that value has more nodes than type in dataset, so it is more sensitive to hidden size.

The influence of the OoV problem: 1. The OoV problem will affect the upper limit of the model's prediction accuracy. For the six data sets we use, the OoV rates of JS1k, JS10k, JS50k, PY1k, PY10k, and PY50k are 20%, 11%, 7%, and 24%, respectively. 16%, 11%. This leads to the upper limit of the prediction accuracy of our CC-GGNN model of 80%, 89%, 93%, 76%, 84%, 89%. 2. Because we define the OoV word to participate in the training of the model by *UNK*, and we successfully predict the OoV word as *UNK*, we think it is wrong. Therefore, some nodes are predicted to be *UNK*, which will reduce the accuracy of model prediction.

6.6 | Training Cost Analysis

In order to compare the training time of the model, we selected the pointer network and the Transformer-XL model. With help of the Transformer-XL model, the representation of each input of each segment depends on the self-attention layer calculation, and repetition only occurs between segments. Therefore, it allows more parallelization and requires less time to train. In Pointer Mixture Network, they adopt LSTM as the language model, where most of the recurrent computations are performed during the hidden states' updating process. Similar to the pointer network, our model also spend a lot of time in the vector transfer of the hidden layer, so the training time is longer than the Transformer-XL model, but our classification mechanism reduces the range of candidate values and reduces the training time. The experimental results are shown in Table 9, Transformer-XL model spend 78% of the time compared to our model. Compared with the pointer network model, our training time is shortened by about 10%.

Model	# of Parameters	Training Time(h/Epoch)
PointerMixtureNet	162.6M	9.4
Transformer-XL	98.9M	6.7
CC-GGNN	147.5M	8.5

TABLE 10 Training cost analysis in the Python dataset



FIGURE 7 Effectiveness of each component in *value* prediction task

7 | THREATS TO VALIDITY

Threats to external validity relate to the quality of the datasets we use. Take the Python dataset as an example. All the data comes from more than 3000 projects. There are 150,000 training data and test data, so there are multiple data sources for one project. Then the code habits of the programmers of this project have a greater impact on the training of the model. Python and JavaScript are two benchmark datasets that have been used in previous code completion work^{6,13}. All programs in the dataset are from GitHub repositories. Therefore, further research is needed to verify our findings and extend them to other programming languages. For example, we will consider the Java dataset¹³.

Threats to internal validity includes the influence of classification mechanism on the number of training graph. The performance of our model would be affected by the different graph. For example, in section 5.3, the node with 0% prediction accuracy. We divide the training graph by the parent node, resulting in less training data for this node and poor prediction results.

Threats to construct validity relate to the suitability of our evaluation measure. We use accuracy and MRR@5 as the metric which evaluates the proportion of correctly predicted next token. It is a classical evaluation measure for code completion and is used in almost all the previous code completion work. Recently, the MRR metric begin to use in code completion work^{22 23}.

8 | RELATED WORK

8.1 | Code Completion

The n-gram language models 24,25,26,27 are the early Machine Learning models for code completion. These probability models simple and effective for predict the next token. An n-gram language model computes the probability of the next token given previous *n* code tokens as model input. These models only use the most primitive tokens representation of the code and do not consider effective semantic structure features. Therefore, in subsequent research, the researchers used the abstract syntax trees representation code and bring its into the probability model to improve the accuracy of the prediction. These models include probabilistic context-free grammars 28 and probabilistic higher-order grammars 4 . This class of model uses context-free grammar to convert the source code into an AST, and predicts next token based on the information selectively collected across paths in



FIGURE 8 The impact of the hidden sizes in our model

AST. The AST path contains rich structural and semantic features, which is the key to improve the prediction accuracy of the next node in probabilistic grammar models. Bruch et al.²⁹ used the KNN algorithm to find the most similar completed fragments in the existing code base and provided candidates for method calls. It also points out that many machine learning algorithms can be introduced into the research of code completion. Subsequently, Raychev et al.¹¹ used the classic model of decision tree model combined with the path information of AST to achieve the highest prediction accuracy of this line of works. Their model learns a decision tree model that uses this information essentially as features. Our work is significantly better than DEEP3.

With the development of deep learning in natural language processing tasks, many researchers believe that code is also a natural language. Deep learning is able to capture longer dependency within a sequence or node's path. Recurrent Neural Network (RNN) is the most common neural technique for code prediction. And, their variants feed input code as a linear AST sequence. In this way, the RNN essentially learns a language model over the training corpus, and it is generally believed to do so better than n-grams²² model. Indeed, several papers in the literature, including work done in industrial contexts¹⁹ has used RNNs. Liu et al.³⁰ attempt has been made on feeding RNN with serialized ASTs. The further research accuracy is improved by using more AST guided architectures⁶. In addition to the above models, there are other different types of code completion models. For example, use the data after the prediction node to make predictions^{31,32,33,34}.

8.2 | Graph Embedding

Graph models are widely used in graph structure node quantization processing. The graph embedding obtains the link conversion sequence of the nodes and edges in the structure, and then embeds the sequence into a fixed-dimensional vector through the graph model. Bryan Perozzi et al.³⁵ proposed DeepWalk model to process graph problem, which is uses a random walk algorithm. However, the random walk algorithm extract path sequence is random, and no attention is paid to the depth-first search (DFS) and breadth-first search (BFS) paths between fusion nodes. To solve this problem, Aditya Grover et al.³⁶ proposed Node2vec model, which is a biased random walk algorithm based on the edge weight of graph nodes. The Node2vec algorithm can fuse the structural information of the node DFS and BFS. Node2vec can effectively integrate the information around the node, so the extracted features are more effective. The powerful method of in modeling the dependency relationship between graph nodes has made the research field related to graph analysis achieve good results. Graph models can also be combined with depth models to predict tasks. Structural deep network embedding (SDNE)³⁷ recommends using a deep autoencoder to maintain the

proximity of the first-order and second-order networks. It achieves this by jointly optimizing these two approximations. The graph convolutional network $(\text{GCN})^{38,39,40}$ model aggregates the neighborhood embedding representations of nodes through iteration, and and uses the previous iteration embedding combined with the embedding function to obtain a new embedding. Only the aggregation embedding of the local neighborhood makes it scalable, and iteratively learns to embed a node to describe the global neighborhood. GGNN^{14,15} is a classical spatial domain message passing model based on GRU^{41,42}, which can solve the problem of gradient in sequence long-term memory and back propagation.

Allamanis et al.³¹ use graph model to represent the structure and semantic structure information of the source code, and found that graph neural networks have better performance than convolutional neural networks in variable completion and variable misuse tasks. The research of Rahman et al.⁴³ pointed out that the graph model has a better predictive effect than the N-gram model. However, there is a lack of comparison with the deep neural network model, and it is recommended to further study the statistical code graph model to accurately capture more complex coding models.

9 | CONCLUSION AND FUTURE WORKS

In this work, we propose CC-GGNN model to solve the problems of the code completion. To better represent AST nodes, we classify node's ASTs by known parent node and transform them into training directed graph. Finally, we combine powerful GGNN model with the ability to integrate structural information to predict the next node. Compared with previous work, the experimental results show that our model can effectively improve the results of code completion. Especially for the *type* prediction of leaf nodes, our model has a significant improvement effect.

In future work, we will apply the CC-GGNN model to more programming languages, such as Java. Since there are many comparison methods and the Python grammar has more casual style. This style leads to the Python dataset being one of the most difficult to predict.

Secondly, our model cannot predict OoV problem. It is a great challenge for predicting the rare words artificially defined by programmers in source code. In previous work, researchers used Pointer Mixture Network and copying mechanisms⁴⁴ to predict Out-of-Vocabulary(OoV). We will try these strategies in the next experiments to improve our model.

Finally, we will try to bring more code information into the model, such as using Control Flow Graph(CFG)^{45,46}. CFG contains a lot of call relationships between nodes, which may help us improve the prediction accuracy of value prediction task.

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