

A Novel Composite Graph Neural Network

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Abstract—Graph neural networks (GNNs) have achieved great success in many fields due to their powerful capabilities of processing graph-structured data. However, most GNNs can only be applied to scenarios where graphs are known, but real-world data are often noisy or even do not have available graph structures. Recently, graph learning has attracted increasing attention in dealing with these problems. In this article, we develop a novel approach to improving the robustness of the GNNs, called composite GNN. Different from existing methods, our method uses composite graphs (C-graphs) to characterize both sample and feature relations. The C-graph is a unified graph that unifies these two kinds of relations, where edges between samples represent sample similarities, and each sample has a tree-based feature graph to model feature importance and combination preference. By jointly learning multiaspect C-graphs and neural network parameters, our method improves the performance of semisupervised node classification and ensures robustness. We conduct a series of experiments to evaluate the performance of our method and the variants of our method that only learn sample relations or feature relations. Extensive experimental results on nine benchmark datasets demonstrate that our proposed method achieves the best performance on almost all the datasets and is robust to feature noises.

Index Terms—Composite graph (C-graph), graph neural networks (GNNs), sample graph, tree-based feature graph.

NOMENCLATURE

Symbol	Meaning.
\mathbb{R}	Set of real numbers.
N	Number of samples.
F	Number of features.
C	Number of classes.
R	Number of submodules.
$\mathbf{0}$	C -dimensional zero vector.

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$\mathbf{1}^\top$	N -dimensional all-one column vector.
$\mathbf{I} \in \mathbb{R}^{N \times N}$	Identity matrix.
$\mathbf{X} \in \mathbb{R}^{N \times F}$	Input dataset.
$\mathbf{Y} \in \mathbb{R}^{N \times C}$	One-hot labels.
$\mathring{\mathbf{Y}} \in \mathbb{R}^{N \times C}$	Pseudolabels.
$\mathbf{A}_S \in \mathbb{R}^{N \times N}$	Adjacency matrix of the sample graph.
$\mathbf{A}_F \in \mathbb{R}^{F \times F}$	Adjacency matrix of the tree-based feature graph.
\mathcal{G}	Composite graph (C-graph).
$\mathbf{A}_G = \{\mathbf{A}_S, \mathbf{A}_F\}$	Adjacency matrix set of \mathcal{G} .
$\boldsymbol{\alpha} \in \mathbb{R}^{F+C}, \tilde{\boldsymbol{\alpha}} \in \mathbb{R}^F$	Trainable vectors.
Θ_1 and Θ_2	Trainable weight matrices.
\odot	Elementwise product.
ℓ	Differentiable error function.
σ	Nonlinear activation function.

I. INTRODUCTION

IN RECENT years, graph neural networks (GNNs) have been successfully used to deal with graph-structured data [1], [2] and have been widely applied in many domains, such as computer vision systems [3], [4], natural language processing [5], [6], neural-symbolic computing [7], [8], and COVID-19 prevention and control [9], [10]. The good performance of GNNs depends on their capabilities of utilizing data graph structure [11]. Many GNNs are developed to aggregate node information along edges of given sample graphs (the graphs whose nodes are samples and the edges are the relations between samples), such as graph convolutional network (GCN) [12] and graph attention network (GAT) [13]. These information aggregation methods are also further improved in recently published works [14], [15]. However, most of these methods do not learn the graph structure of samples (i.e., the sample graph), and thus, can be hardly applied to real scenarios where graphs are noisy or even unavailable. Many link prediction methods proposed in the literature can learn sample graphs [16], [17], but these methods either are difficult to be directly incorporated into GNNs or cannot jointly learn neural network parameters and sample graphs.

Jointly learning the graph structure of samples and the parameters of a neural network has shown to be important to the robustness of GNNs [18], [19]. To this end, some GNN methods capable of simultaneously learning or refining the graph structure of samples and neural network parameters are developed [19], [20]. However, none of them consider learning feature graphs (the graphs whose nodes are features and edges are feature relationships), thereby incapable of capturing the structural information between features. Similar to samples, features can also exhibit multifarious structures, such as tree structures [21], [22] and graph structures [23], [24]. The structures of features have been proved to be very important in many real-world applications [25], [26].

Recently, some studies noticed that learning the unknown graph structure of features can help improve the performance of GNNs, such as Fi-GNN [27] and graph convolutional networks with categorical node features (CatGCN) [28]. These methods consider learning the feature graphs, but the sample graphs are required to be known, which hinders their applications in scenarios where the sample graphs are inaccessible. High-quality sample relations and feature relations depend on each other, and thus, should be jointly learned instead of being learned in two stages. To this end, we need to use a graph to characterize both sample relations and feature relations. Both feature and sample relations can be learned from the Euclidean distance or the cosine similarity of each pair of samples and features. However, pairwise similarities (distances) can neither select important features nor obtain important feature combinations, which have been shown to be important for feature relationship learning [28], [29]. For example, men aged 20–25 are more likely to be digital enthusiasts [28]. This gender–age feature combination is more discriminating than either gender or age alone for judging digital enthusiasts. Sample relations usually represent the pairwise similarity between samples, while feature relations need to show their combination preference. Hence, it is a natural way to model sample relations with a sample graph, and on each node, feature relations should be modeled with trees.

Thus, a natural question arises: *how to model a graph of samples and trees of features with a unified graph and jointly learn this unified graph and the neural network parameters?*

We propose a composite GNN (CGNN) to learn and utilize C-graphs for node embeddings. First, we propose a new type of feature graph called the tree-based feature graph. When using features to build a tree in CGNN, the important features are selected as nodes, and the combination of two connected features is preferred. The reasons are because: 1) a feature tree’s hierarchy corresponds to how informative each feature is, with more informative features closer to the tree’s root node [30] and 2) the features used to build a tree are more closely related to one another than the ones, not used [31]. Then, we propose the C-graph, which uses a sample graph to model the sample similarities, and a tree-based feature graph to model the feature relations of each sample. Finally, we propose a C-graph transformation operation to aggregate features using the C-graph. In our method, both sample relations and feature relations can be jointly updated by minimizing the same loss function, which enables us to utilize their dependencies and obtain a better C-graph.

To the best of our knowledge, no previous studies discuss the unified modeling of the tree structure of features and the graph structure of samples. We highlight the contributions and the advantages of our work as follows.

- 1) *Unifying the Tree-Based Feature Graph and the Sample Graph Into a Composite Graph:* The C-graph reveals both sample similarities and feature combination preferences.
- 2) *Proposing a C-Graph Transformation Operation to Aggregate Features Using the C-Graph:* This operation jointly uses sample and feature relations to update features.
- 3) *Proposing a Composite Graph Neural Network for Semisupervised Node Classification Problems:* We develop a C-graph-based neural network method, which can jointly learn C-graphs from different aspects and neural network parameters, and thus, can be directly

used to handle *semisupervised problems without knowing graphs a priori*.

In addition, extensive validations on different datasets demonstrate that CGNN achieves better performance than state-of-the-art methods in semisupervised classification accuracy, regardless of whether the sample graph is known or not. Moreover, additional experiments show that CGNN is robust to feature noises even when the sample graph is unknown.

The remainder of this article is organized as follows. Section II reviews the related work. In Section III, we propose our framework CGNN that uses C-graphs to deal with semisupervised problems and give the details of jointly learning the graph structure of samples and the tree structure of features to obtain C-graph. Comprehensive evaluation and experimental result analysis are conducted in Section IV. Finally, conclusion is given in Section V.

Notations: In this article, we use boldfaced characters to represent vectors and matrices. Suppose that \mathbf{M} is a matrix, then $\mathbf{M}[n, \cdot]$, $\mathbf{M}[\cdot, f]$, and $\mathbf{M}[n, f]$ denote its n th row, f th column, and (n, f) th element, respectively. A vector $\mathbf{x} = (x_1, x_2, \dots, x_N)$ is abbreviated as $(x[i])_{i=1}^N$ if the index set that i runs over is clear from the context, and the i th element of \mathbf{x} is $x[i]$. We summarize the important notations used throughout this article in the Nomenclature, where \mathbf{X} is a matrix of node feature vectors, and the feature vector of each node is called a sample. We use each row in \mathbf{X} to denote a sample, each column in \mathbf{X} to denote a feature, and α , $\tilde{\alpha}$, Θ_1 , and Θ_2 to denote the neural network parameters.

II. RELATED WORK

Many graph-based methods have been proposed for relational data with known graph structures, such as semisupervised embedding (SemiEmb) [32], DeepWalk [33], and large-scale information network embedding (LINE) [34]. Among these methods, graph embedding-based method receives considerable attention. In [35], a method named planetoid is proposed, which trains an embedding for each sample by jointly predicting the class label and capturing structural information in the given sample graph. However, planetoid cannot fully embed all the graph structure and label information limited by its sampling strategy [36]. Recently, GNNs closely related to graph embedding have attracted increasing interest in the field of machine learning and data mining [37]. In [12] and [13], the well-known GCN and GAT are proposed, which directly aggregate neighbor information using graph convolution and graph attention, respectively. Inspired by GCN and GAT, many GNNs with better performances are proposed, such as graph convolutional network via initial residual and identity mapping (GCNII) [14], adaboosting graph convolutional network (AdaGCN) [38], and bilinear graph neural network (BGAT-T) [39].

A foundational assumption of the previous methods is that the given graph structure of samples is of high quality, limiting their application in scenarios with unknown or noisy graph structures. Recently, graph learning has drawn significant attention in dealing with these problems [40], [41]. Some link prediction methods and GNN methods can extend a sample graph based on some prior known structural relationships between nodes [42], [43], but these methods cannot learn the sample graph and neural network parameters simultaneously. Recently, some methods that can jointly learn the sample graph and neural network parameters have been proposed. k -nearest neighbors (k NN)-learning discrete structures for graph

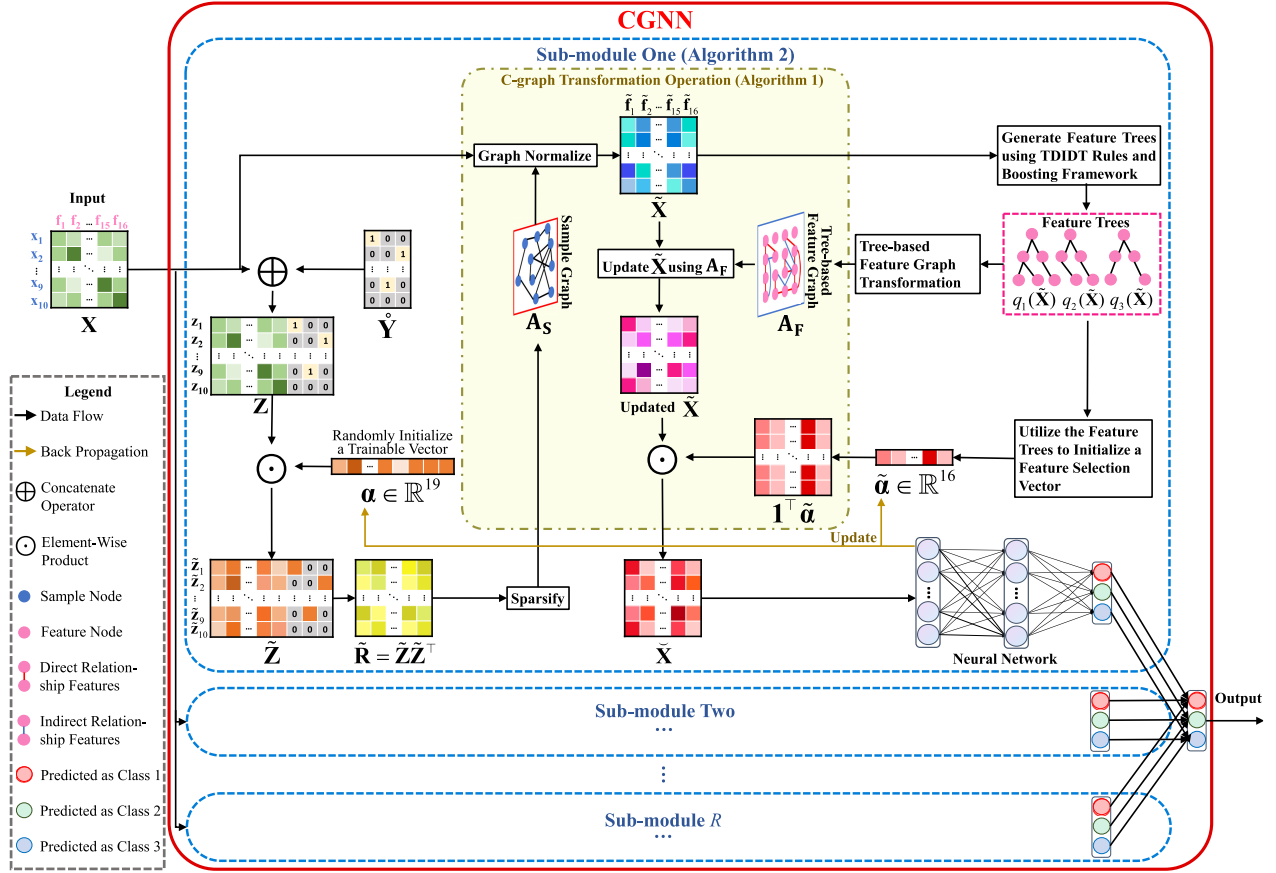


Fig. 1. Toy example of CGNN with R submodules for a three-class classification of input $X \in \mathbb{R}^{10 \times 16}$. Each blue dotted box denotes a submodule, and there are R blue dotted boxes in total. $\tilde{Y} \in \mathbb{R}^{10 \times 3}$ denotes the pseudolabels generated from the training labels. \tilde{X} denotes the node embeddings obtained by learning the C-graph defined in Definition 1. In our method, the C-graph $A_G = \{A_S, A_F\}$ and the parameters of the neural network are jointly learned. In the C-graph A_G , A_S and A_F depend on each other in the update process. The proposed C-graph transformation operation takes X , $A_G = \{A_S, A_F\}$ and $\tilde{\alpha}$ as input, and outputs \tilde{X} . Best viewed in color.

neural networks (LDS) gives a bilevel programming approach to jointly learn the sample graph and a neural network [19]. Deep iterative and adaptive learning for graph neural networks (DIAL-GNN) iteratively learns the parameters of a neural network and augments the given graph structure [44]. Graphite proposed in the literature [45] uses a deep latent variable generative model to learn a sample graph and the sample graph can be iteratively refined via a message-passing operation. Pro-GNN simultaneously learns the parameters of a neural network and a robust graph structure of samples by exploring graph properties of sparsity, low rank, and feature smoothness [18]. A GCN method proposed in the literature [46] learns the graph and graph representation from the low-dimensional space of the original data simultaneously by using a new multigraph fusion method. Peng et al. [47] proposed a GNN method that learns the graph structure from the given data points by using reverse graph learning and out-of-sample extension strategies.

Although the aforementioned methods jointly learn sample graph and neural network parameters, they do not consider the unknown underlying relations between features. However, in many applications, the hierarchical structure of features and the combination of related features are essential. In our method, we jointly learn the graph structure of samples and the tree structure of features, which further compose the C-graph. Hence, our proposed C-graph can characterize the similarity between samples, relative importance between features, and combination preference of features. However, it is still an unresolved problem in the literature to learn and utilize the

C-graph. Next, we will provide the details of our method in Section III.

III. COMPOSITE GRAPH NEURAL NETWORK

In this section, we propose the GNN method CGNN to address the challenging problem where data samples influence each other through edges of multiple unknown graphs and these edges are closely related to the features of samples. CGNN is a unified framework to learn a generalized graph, C-graph, capable of demonstrating both sample relationships and feature relationships. Specifically, CGNN is proposed to satisfy the following.

- 1) Learn a C-graph of features and samples, which characterizes the similarity between samples, the hierarchical structure of features, and the combination preference of related features.
- 2) Learn multiple C-graphs from different aspects using different submodules.
- 3) Jointly learn the C-graphs and neural network parameters for semisupervised tasks. The joint learning of sample relations and neural network parameters is shown to be essential to the robustness of GNNs.

A toy example for illustrating CGNN is shown in Fig. 1. The pseudocode of CGNN is given in Algorithm 2 in Section IV.

A. Main Definitions

Consider a dataset with N samples $X = [x_1, x_2, \dots, x_N]^T$. For any $n \in \{1, 2, \dots, N\}$, $x_n \in \mathbb{R}^F$ has F features. The

one-hot labels of these samples are $\mathbf{Y} = [\mathbf{y}_1, \mathbf{y}_2, \dots, \mathbf{y}_N]^\top$, where each element $\mathbf{y}_n \in \mathbb{R}^C$ with C being the number of classes. In this article, we focus on the semisupervised classification problem, where \mathbf{X} and a subset of \mathbf{Y} are used to learn a model to predict the remaining unknown subset of \mathbf{Y} . Incorporating the relationship between samples (i.e., the sample graph) has shown great potential in semisupervised classification problems [12]. Most GNNs can only deal with the case where graphs are known and some other GNNs capable of learning graphs only consider the sample relations. In this article, we consider a C-graph characterizing both the sample relations and feature relations.

Definition 1 (Composite Graph): In this article, the C-graph is defined as $\mathcal{G} \triangleq (\mathbf{S}, \mathbf{E}_\mathbf{S}, \mathbf{E}_\mathbf{F})$, where \mathbf{S} , $\mathbf{E}_\mathbf{S}$, and $\mathbf{E}_\mathbf{F}$ are the sample node set, the sample edge set, and the feature edge set, respectively. We define the adjacency matrix set of \mathcal{G} as $\mathbf{A}_\mathcal{G} \triangleq \{\mathbf{A}_\mathbf{S}, \mathbf{A}_\mathbf{F}\}$, where $\mathbf{A}_\mathbf{S}$ and $\mathbf{A}_\mathbf{F}$ are the adjacency matrices of the sample graph and the tree-based feature graph, respectively. We say $\mathbf{A}_\mathbf{S}$ and $\mathbf{A}_\mathbf{F}$ are unified into a C-graph since they are simultaneously used to update features in Algorithm 1, jointly learned by optimizing the same loss function $\ell(\mathbf{Y}, \hat{\mathbf{Y}})$, and depend on each other in the update process (see Section III-C3).

Given a dataset \mathbf{X} and a known subset of \mathbf{Y} , we aim at learning the C-graph $\mathbf{A}_\mathcal{G} = \{\mathbf{A}_\mathbf{S}, \mathbf{A}_\mathbf{F}\}$ and use the learned graph to obtain node embeddings and predicted labels. It is worth noting that $\mathbf{A}_\mathbf{S}$ and $\mathbf{A}_\mathbf{F}$ are closely related to each other and are jointly learned in a unified framework. In Sections III-B–III-D, $\mathbf{D}_\mathbf{S}$ denotes an $N \times N$ diagonal matrix with $\mathbf{D}_\mathbf{S}[n, n] \triangleq 1 + \sum_{p=1}^n \mathbf{A}_\mathbf{S}[n, p]$ for each $n \in \{1, 2, \dots, N\}$. $\sigma(\cdot)$ denotes an activation function. $\ell(\mathbf{Y}, \hat{\mathbf{Y}})$ denotes a differentiable function that measures the difference between the target \mathbf{Y} and the prediction $\hat{\mathbf{Y}}$, and we use the cross-entropy error function in our experiments.

B. General Framework

In this section, we present the general framework of CGNN. Our framework can jointly learn C-graphs from different aspects and neural network parameters. The learned C-graphs can reveal both the underline structure information of sample similarities and feature combination preferences for helping obtain better node embeddings and deal with semisupervised problems without knowing graphs a priori. Let R be the number of submodules in our method. In Fig. 1, each blue dotted box denotes a submodule, and there are R submodules in total. The following formulas (1)–(3) are repeated for each submodule $r \in \{1, 2, \dots, R\}$, and thus, in these formulas, we omit submodule index r . First, we use a C-graph transformation operation to embed \mathbf{X} with the C-graph, and the obtained node embeddings are further input to a fully connected hidden layer. This process is given by

$$\mathbf{X}^{(1)} = \sigma(\text{Cgto}(\mathbf{X}, \mathbf{A}_\mathcal{G}, \tilde{\alpha})\Theta_1) \quad (1)$$

where $\text{Cgto}(\cdot, \cdot, \cdot)$ is the C-graph transformation operation over $\mathbf{A}_\mathcal{G}$, Θ_1 is a trainable $F \times H$ matrix, where H denotes the number of features of the first hidden layer, and $\mathbf{X}^{(1)}$ is the output of the first hidden layer. The Cgto can not only enhance the feature weight by extracting information between neighbors but also can highlight the important features. The pseudocode of Cgto is shown in Algorithm 1. In line 6 of Algorithm 1, \mathbf{X} is normalized to $\tilde{\mathbf{X}}$ by $\mathbf{A}_\mathbf{S}$ and $\mathbf{D}_\mathbf{S}$. From line 7 to line 16, $\tilde{\mathbf{X}}$ is further updated using $\mathbf{A}_\mathbf{F}$. It is worth noting that in line 17, the important features will be assigned larger

Algorithm 1 C-Graph Transformation Operation

Input : $\mathbf{X}, \mathbf{A}_\mathcal{G} = \{\mathbf{A}_\mathbf{S}, \mathbf{A}_\mathbf{F}\}, \tilde{\alpha}$
Output: the transformed \mathbf{X}

- 1 Initialize the three hyperparameters w_C, w_D , and w_I ;
- 2 Initialize $\mathbf{D}_\mathbf{S} \in \mathbb{R}^{N \times N}$ as a zero matrix;
- 3 **for** each $n \in \{1, 2, \dots, N\}$ **do**
- 4 $\mathbf{D}_\mathbf{S}[n, n] = 1 + \sum_{p=1}^n \mathbf{A}_\mathbf{S}[n, p]$;
- 5 **end**
- 6 $\tilde{\mathbf{X}} = \mathbf{D}_\mathbf{S}^{-\frac{1}{2}}(\mathbf{A}_\mathbf{S} + \mathbf{I})\mathbf{D}_\mathbf{S}^{-\frac{1}{2}}\mathbf{X}$;
- 7 **for** each feature $\tilde{\mathbf{f}}_i$ in $\tilde{\mathbf{X}}$ **do**
- 8 $\tilde{\mathbf{f}}_i = w_C \mathbf{f}_i$;
- 9 **for** each $f \in \{1, 2, \dots, F\}$ **do**
- 10 **if** $\mathbf{A}_\mathbf{F}[i, f] = 2$ **then**
- 11 $\tilde{\mathbf{f}}_i = \tilde{\mathbf{f}}_i + w_D \mathbf{f}_f$;
- 12 **else if** $\mathbf{A}_\mathbf{F}[i, f] = 1$ **then**
- 13 $\tilde{\mathbf{f}}_i = \tilde{\mathbf{f}}_i + w_I \mathbf{f}_f$;
- 14 **end**
- 15 **end**
- 16 **end**
- 17 $\tilde{\mathbf{X}} = \mathbf{1}^\top \tilde{\alpha} \odot \tilde{\mathbf{X}}$;
- 18 **return** $\tilde{\mathbf{X}}$;

weights by $\tilde{\alpha}$ in the training process, which will be illustrated in Section III-C2 in detail.

Next, we use a generalized GCN layer to update $\mathbf{X}^{(1)}$ and obtain

$$\mathbf{X}^{(2)} = \sigma\left(\mathbf{D}_\mathbf{S}^{-\frac{1}{2}}(\mathbf{A}_\mathbf{S} + \mathbf{I})\mathbf{D}_\mathbf{S}^{-\frac{1}{2}}\mathbf{X}^{(1)}\Theta_2\right) \quad (2)$$

where Θ_2 is a trainable $H \times C$ matrix. Different from the GCN [12] that uses a fixed graph in the training process, in formula (2), the adjacency matrix of the samples $\mathbf{A}_\mathbf{S}$ are trained together with Θ_2 .

Finally, we use a softmax function to process $\mathbf{X}^{(2)}$ and obtain

$$\mathbf{X}^{(3)} = \text{Softmax}(\mathbf{X}^{(2)}). \quad (3)$$

In our method, (1)–(3) are jointly regarded as a submodule of CGNN. CGNN uses multiple submodules to learn multiple C-graphs from different aspects, and thus, can adapt to the case where the relations of samples and features are influenced by multiple factors.

We use R submodules and for each $r \in \{1, 2, \dots, R\}$, the loss function of submodule r is given by

$$L_r = \ell(\mathbf{Y}, \mathbf{X}^{(3)}). \quad (4)$$

Note that the submodule index r is omitted in $\mathbf{X}^{(3)}$.

Then, the loss function of CGNN is

$$\mathcal{L}(\theta) = \frac{1}{R} \sum_{r=1}^R L_r + \Omega(\theta) \quad (5)$$

where R is the number of submodules, Ω is a regularizer and θ denotes the set of trainable parameters.

C. Learning the C-Graph

In our model, C-graphs are used for node embeddings and the proposed C-graph framework considers both the relationship between samples and the relationship between features. In this section, we first introduce how to learn the sample

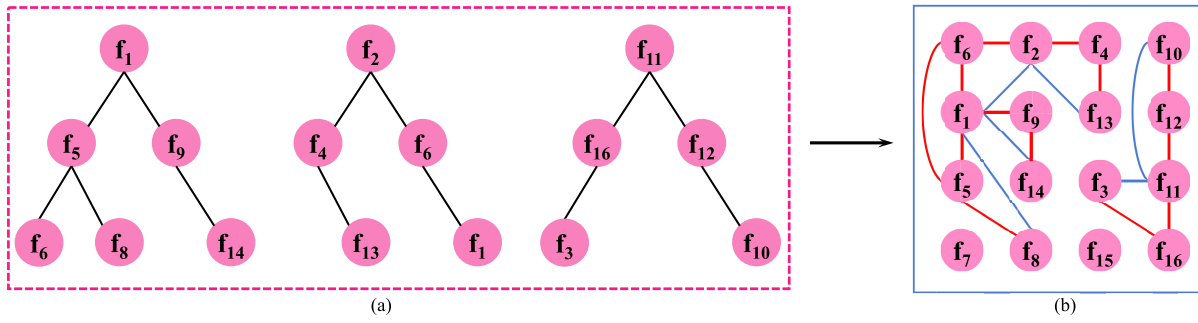


Fig. 2. Example of constructing a tree-based feature graph of 16 features from multiple binary trees. The red lines and the blue lines indicate the direct relationships and the indirect relationships, respectively. (a) Three binary trees. (b) Tree-based feature graph transformed by the three binary trees.

graph \mathbf{A}_S and the tree-based feature graph \mathbf{A}_F . Moreover, we introduce how \mathbf{A}_S and \mathbf{A}_F depend on each other in the learning process.

1) *Learning Sample Relations*: In this section, we propose the method to learn the adjacency matrix of the sample graph \mathbf{A}_S . We consider the known class label as an important feature to learn the sample graph. We first construct pseudolabels $\hat{\mathbf{Y}} = [\hat{y}_1, \hat{y}_2, \dots, \hat{y}_N]^T$ using the training labels by

$$\hat{y}_n = \begin{cases} y_n, & \text{if } y_n \text{ is a training label} \\ 0, & \text{otherwise} \end{cases} \quad (6)$$

where $n \in \{1, 2, \dots, N\}$, and $\mathbf{0}$ denotes the C -dimensional zero vector. Then, we concatenate \mathbf{x}_n and \hat{y}_n and obtain

$$\mathbf{z}_n = \text{Concatenate}(\mathbf{x}_n, \hat{y}_n) \quad (7)$$

where $\text{Concatenate}(\cdot, \cdot)$ denotes the concatenation operation. It should be noted that the test labels are all-zero vectors in the training and testing process. Besides, as pointed out in the literature [48], noisy features can hinder the model training process. Hence, a trainable vector α is utilized to diminish the noisy features, which is given by

$$\tilde{\mathbf{z}}_n = \mathbf{z}_n \odot \alpha. \quad (8)$$

We regard α as a feature improvement operator. The effectiveness of this simple operator is also shown in [49]. Different from most feature selection methods, α is trained mainly to assign different weights to different features, rather than reduce the feature dimension. In other words, at the end of the training process, the smaller values of α will indicate the noisy dimensions of features. Given that α is initialized randomly, different α can have different values and focus on different features in different submodules. From (8), a new matrix $\tilde{\mathbf{Z}} = [\tilde{\mathbf{z}}_1, \tilde{\mathbf{z}}_2, \dots, \tilde{\mathbf{z}}_N]^T$ can be obtained and then we compute the relationship matrix $\tilde{\mathbf{R}}$ using $\tilde{\mathbf{Z}}$:

$$\tilde{\mathbf{R}} = \tilde{\mathbf{Z}}\tilde{\mathbf{Z}}^T. \quad (9)$$

Next, we find the top η values in each row of $\tilde{\mathbf{R}}$ and set the others to 0 to sparsify $\tilde{\mathbf{R}}$. The sparsified $\tilde{\mathbf{R}}$ is regarded as the \mathbf{A}_S .

2) *Learning Feature Relations*: In this section, the method to learn the tree-based feature graph (i.e., \mathbf{A}_F) is presented. We decompose the learning of a tree-based feature graph into the learning of multiple binary decision trees. There are three main advantages of using binary decision trees: 1) features used to construct a tree are more closely related to each other than the unused features [31]; 2) the hierarchy of features in a tree reflects their informativeness and the

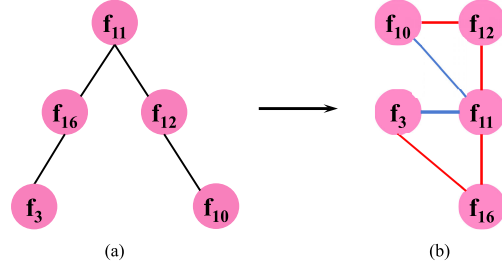


Fig. 3. Example of constructing a tree-based feature graph from a binary tree. The red lines and the blue lines indicate the direct relationships and the indirect relationships, respectively. (a) Binary tree. (b) Tree-based feature graph transformed by the binary tree.

features closer to the root are more important [30]; and 3) the binary decision tree structure is easy to implement and has high computational efficiency [50]. Take Fig. 2 as an example. We first learn three binary trees [see Fig. 2(a)] and then turn them into the tree-based feature graph [see Fig. 2(b)]. The specific tree-based feature graph transformation method is as follows. First, we define two kinds of relationships: the direct relationship and the indirect relationship. The direct relationship is between each pair of directly connected nodes [e.g., the relationship between f_{11} and f_{16} in Fig. 2(a)] and the indirect relationship is between each pair of nodes that belong to the same branch of the tree but are not directly connected [e.g., the relationship between f_{11} and f_{10} in Fig. 2(a)].

Second, we assume that direct relationships represent stronger influence than indirect relationships and use 0, 1, and 2 to denote no relationship, an indirect relationship, and a direct relationship between each pair of features, respectively. Then we connect all the related features by finding all the direct and indirect relationships in the tree [see the transformation from Fig. 3(a) and (b)]. Similarly, for multiple binary trees in Fig. 2, the same nodes and edges generated by each tree are merged. The adjacency matrix of the merged graph with values from $\{0, 1, 2\}$ is the tree-based feature graph \mathbf{A}_F that we want to learn.

In this article, multiple trees are employed to learn the tree-based feature graph since a very deep tree will be generated when the number of features is becoming larger. Thus, in our learning process, ensemble learning methods are considered [51] to generate multiple binary trees to express the relationship between many features with the purpose of constructing the tree-based feature graph. The main advantage of ensemble learning is that it can combine many weak models into strong models, so each generated tree can be relatively simple. We use the boosting framework in ensemble learning to generate multiple trees and these trees are all finally merged into one graph. Here, the boosting framework is used due to the following reason: in the boosting framework, the

generation of the latter tree depends on the information of the former tree [52], and thus, the trees generated in the framework of boosting have a stronger correlation with each other than other frameworks in ensemble learning.

Specifically, we build trees according to the top-down induction of decision trees (TDIDT) method [53], which splits the features of $\tilde{\mathbf{X}} = \mathbf{D}_S^{(1/2)}(\mathbf{A}_S + \mathbf{I})\mathbf{D}_S^{-(1/2)}\mathbf{X}$ recursively, creates the successor children by searching the greatest gain value to the predictions and stops when the gain value does not increase. For $\tilde{\mathbf{X}} = [\tilde{\mathbf{x}}_1, \tilde{\mathbf{x}}_2, \dots, \tilde{\mathbf{x}}_N]^\top$, our method builds trees by minimizing $\ell(\mathbf{Y}, \tilde{\mathbf{Y}})$, where $\tilde{\mathbf{Y}} = [\tilde{\mathbf{y}}_1, \tilde{\mathbf{y}}_2, \dots, \tilde{\mathbf{y}}_N]^\top$ is the prediction results of the feature trees and each element $\tilde{\mathbf{y}}_n \in \mathbb{R}^C$. We train the trees in an additive manner. Suppose that T is the total number of iterations for training trees; then, we build CT trees in total because C trees are generated in each iteration by minimizing $\ell(\mathbf{Y}, \tilde{\mathbf{Y}})$. Thus, minimizing $\ell(\mathbf{Y}, \tilde{\mathbf{Y}})$ is equivalent to minimize

$$\begin{aligned} \ell^{(T)}(\mathbf{Y}, \tilde{\mathbf{Y}}^{(T)}) &= \sum_{n=1}^N l(\mathbf{y}_n, \tilde{\mathbf{y}}_n^{(T)}) \\ &= \sum_{n=1}^N l\left(\mathbf{y}_n, (\tilde{\mathbf{y}}_n^{(T)}[c])_{c=1}^C\right) \end{aligned} \quad (10)$$

where $\ell^{(T)}(\mathbf{Y}, \tilde{\mathbf{Y}}^{(T)})$ and $\tilde{\mathbf{Y}}^{(T)} = [\tilde{\mathbf{y}}_1^{(T)}, \tilde{\mathbf{y}}_2^{(T)}, \dots, \tilde{\mathbf{y}}_N^{(T)}]^\top$ are the loss function value and the prediction results, respectively, and l denotes a differentiable function that measures the difference between the target \mathbf{y}_n and the prediction $\tilde{\mathbf{y}}_n^{(T)}$.

Let $B^{(T)}[c]$ be the c th tree ($c \in \{1, 2, \dots, C\}$) generated at the final iteration. Next, we take the process of generating $B^{(T)}[c]$ as an example to introduce our approach to building feature trees. At the T th iteration, for each $n \in \{1, 2, \dots, N\}$, $\tilde{\mathbf{y}}_n^{(T)}[c]$ is computed by

$$\begin{aligned} \tilde{\mathbf{y}}_n^{(T)}[c] &= \tilde{\mathbf{y}}_n^{(T-1)}[c] + B^{(T)}[c](\tilde{\mathbf{x}}_n) \\ &= \tilde{\mathbf{y}}_n^{(T-1)}[c] + \sum_{m=1}^{M^{(T)}[c]} b_m^{(T)}[c] \mathbb{1}(\tilde{\mathbf{x}}_n \in B_m^{(T)}[c]) \end{aligned} \quad (11)$$

where $B^{(T)}[c](\tilde{\mathbf{x}}_n)$ denotes the value of using $B^{(T)}[c]$ to predict $\tilde{\mathbf{x}}_n$ as the c th class, $M^{(T)}[c]$ denotes the total number of leaves in $B^{(T)}[c]$, $B_m^{(T)}[c]$ and $b_m^{(T)}[c]$ denote the m th leaf of $B^{(T)}[c]$ and the value of $B_m^{(T)}[c]$, respectively, and $\mathbb{1}(\cdot)$ denotes the indicator function. Notably, $\tilde{\mathbf{y}}_n^{(0)}[c] = 0$ for $n \in \{1, 2, \dots, N\}$ and $c \in \{1, 2, \dots, C\}$. We use second-order approximation [54] to optimize (11) and compute the optimal $b_m^{(T)}[c]$. Then, we further rewrite (10) as the following regularized objective:

$$\begin{aligned} \ell^{(T)}(\mathbf{Y}, \tilde{\mathbf{Y}}^{(T)}) &+ \sum_{c=1}^C \Omega(B^{(T)}[c]) \\ &= \sum_{n=1}^N l(\mathbf{y}_n, \tilde{\mathbf{y}}_n^{(T-1)} + \mathbf{B}^{(T)}(\tilde{\mathbf{x}}_n)) + \sum_{c=1}^C \Omega(B^{(T)}[c]) \\ &\simeq \sum_{n=1}^N \left[l(\mathbf{y}_n, \tilde{\mathbf{y}}_n^{(T-1)}) + \sum_{c=1}^C \frac{\partial l(\mathbf{y}_n, \tilde{\mathbf{y}}_n^{(T-1)})}{\partial \tilde{\mathbf{y}}_n^{(T-1)}[c]} B^{(T)}[c] \right. \\ &\quad \left. + \frac{1}{2} \sum_{c_1=1}^C \sum_{c_2=1}^C \frac{\partial^2 l(\mathbf{y}_n, \tilde{\mathbf{y}}_n^{(T-1)})}{\partial \tilde{\mathbf{y}}_n^{(T-1)}[c_1] \partial \tilde{\mathbf{y}}_n^{(T-1)}[c_2]} B^{(T)}[c_1] B^{(T)}[c_2] \right] \\ &\quad + \sum_{c=1}^C \Omega(B^{(T)}[c]) \end{aligned} \quad (12)$$

where $\Omega(B^{(T)}[c])$ denotes the regularizer on the leaves of $B^{(T)}[c]$ and $\mathbf{B}^{(T)}(\tilde{\mathbf{x}}_n)$ denotes a C -dimension vector consisting of $B^{(T)}[1](\tilde{\mathbf{x}}_n), B^{(T)}[2](\tilde{\mathbf{x}}_n), \dots, B^{(T)}[C](\tilde{\mathbf{x}}_n)$. Since $B^{(T)}[c](\tilde{\mathbf{x}}_n) = \sum_{m=1}^{M^{(T)}[c]} b_m^{(T)}[c] \mathbb{1}(\tilde{\mathbf{x}}_n \in B_m^{(T)}[c])$, we then substitute $\sum_{m=1}^{M^{(T)}[c]} b_m^{(T)}[c] \mathbb{1}(\tilde{\mathbf{x}}_n \in B_m^{(T)}[c])$ into (12) and transform it into an optimization problem with respect to $b_m^{(T)}[c]$. Therefore, we can train $B^{(T)}[c]$ by optimizing $b_m^{(T)}[c]$. As for how to set and split the nodes of $B^{(T)}[c]$, there are many good algorithms proposed in the past literature, such as greedy top-down approach [55], multiple classification rank method [56] and candidate node split finding approximate algorithm [57]. Since the main contribution of this article is not to improve the node splitting of decision trees, we choose loss reduction as gain value and use the candidate node split finding approximate algorithm to conduct node selection and splitting of $B^{(T)}[c]$ after considering a tradeoff between the computational efficiency and practical effect.

3) Dependencies Between \mathbf{A}_S and \mathbf{A}_F in the Learning Process: We have presented the methods to learn \mathbf{A}_S and \mathbf{A}_F . In the following, we will show how \mathbf{A}_S and \mathbf{A}_F depend on each other in the learning process. The feature enhancement in learning \mathbf{A}_S can help the learning of binary trees when learning \mathbf{A}_F . The operator α in \mathbf{A}_S can enhance the important features through continuous optimization. The enhanced features can further help \mathbf{A}_F construct the ensemble trees with nodes being features. Recall that in training \mathbf{A}_S , α is randomly initialized with different values to ensure learning different \mathbf{A}_S in different submodules, which can increase the diversity of trees in \mathbf{A}_F .

The sample graph and feature graph are jointly used to generate the node embeddings. Firstly, $\tilde{\mathbf{X}}$ is computed using \mathbf{A}_S (see Section III-C2). Then, for each $i \in \{1, 2, \dots, F\}$, we further update each feature (i.e., column vector) $\tilde{\mathbf{f}}_i$ in $\tilde{\mathbf{X}}$ using the learned \mathbf{A}_F , which is given by

$$\tilde{\mathbf{f}}_i = \sum_{j \in \mathbf{J}} w_D \tilde{\mathbf{f}}_j + \sum_{k \in \mathbf{K}} w_I \tilde{\mathbf{f}}_k + w_C \tilde{\mathbf{f}}_i \quad (13)$$

where \mathbf{J} and \mathbf{K} represent the index sets of features having indirect and direct relations with $\tilde{\mathbf{f}}_i$, respectively, and w_D, w_I, w_C are three hyperparameters denoting the weight of the direct relationship, the weight of the indirect relationship, and the weight of the self-relations, respectively. Next, we perform feature selection on the updated $\tilde{\mathbf{X}} = [\tilde{\mathbf{f}}_1, \tilde{\mathbf{f}}_2, \dots, \tilde{\mathbf{f}}_F]$ by using a new trainable row vector $\tilde{\alpha}$

$$\tilde{\mathbf{X}} = \mathbf{1}^\top \tilde{\alpha} \odot \tilde{\mathbf{X}} \quad (14)$$

where $\mathbf{1}^\top$ is the N -dimensional all-one column vector and $\tilde{\alpha}$ is initialized with feature importance values, namely, $\tilde{\alpha}[i] = \exp(v_i)$. Here, $\exp(\cdot)$ denotes the exponential function, and $v_i \in [0, 1]$ is the importance value of $\tilde{\mathbf{f}}_i$. Notably, $\{v_i\}_{i=1}^F$ are obtained in the process of building trees in learning \mathbf{A}_F .

The obtained $\tilde{\mathbf{X}}$ is further updated in (2) using \mathbf{A}_S .

D. Model Training and Pseudocode of CGNN

It is worth noting that the C-graph transformation operation in (1) is non-differentiable for \mathbf{A}_F . Thus, the three variables \mathbf{A}_F, θ , and \mathbf{A}_S share the optimization objective, but have different update methods. Specifically, we use ensemble tree model to update \mathbf{A}_F , as shown in (11) and (12). It can be seen that in formulas (1)–(5), the loss function is differentiable with

Algorithm 2 CGNN (One Submodule)

```

1 Input:  $\mathbf{X}, \mathbf{Y}$ ;
2 Output: Best learned C-graph  $\mathbf{A}_G^*$  and the values of
   parameters  $\alpha^*, \tilde{\alpha}^*$  and  $\theta^*$ ;
3 Initialize parameters:  $\alpha, \tilde{\alpha}, \theta = \{\Theta_1, \Theta_2\}$ ;
4 Initialize hyperparameters;
5 while Stopping condition is not met do
6   Substitute  $\mathbf{X}, \mathbf{Y}$  and  $\alpha$  into (6)-(9) to compute the
     sample graph  $\mathbf{A}_S$ ;
7   Use  $\mathbf{A}_S$  to obtain  $\mathbf{D}_S$ ;
8    $\tilde{\mathbf{X}} \leftarrow \mathbf{D}_S^{-\frac{1}{2}} (\mathbf{A}_S + \mathbf{I}) \mathbf{D}_S^{-\frac{1}{2}} \mathbf{X}$ ;
9   Use  $\tilde{\mathbf{X}}$  and (11)-(12) to compute the trees of
     features and obtain the tree-based feature graph
      $\mathbf{A}_F$ ;
10  Initialize  $\tilde{\alpha}$  using  $\mathbf{A}_F$ ;
11  Generate  $\mathbf{A}_G$  using  $\mathbf{A}_S$  and  $\mathbf{A}_F$ ;
12  while Inner objective decreases do
13    Substitute  $\mathbf{X}, \mathbf{A}_G, \tilde{\alpha}, \theta, \mathbf{Y}$  into (1)-(5) to
      compute  $\mathcal{L}(\theta)$ ;
14    Update  $\alpha, \tilde{\alpha}, \theta$  by using  $\mathcal{L}(\theta)$ ;
15  end
16 end
17 Denote the last iteration values of  $\alpha, \tilde{\alpha}$ , and  $\theta$  as  $\alpha^*,$ 
    $\tilde{\alpha}^*$  and  $\theta^*$ , respectively;
18 Substitute  $\mathbf{X}, \mathbf{Y}$  and  $\alpha^*$  into (6)-(9) to compute the
   best sample graph  $\mathbf{A}_S^*$ ;
19 Use  $\mathbf{A}_S^*$  to obtain  $\mathbf{D}_S^*$ ;
20  $\tilde{\mathbf{X}}^* \leftarrow \mathbf{D}_S^{*- \frac{1}{2}} (\mathbf{A}_S^* + \mathbf{I}) \mathbf{D}_S^{*- \frac{1}{2}} \mathbf{X}$ ;
21 Use  $\tilde{\mathbf{X}}^*$  and (11)-(12) to compute the trees of features
   and obtain the best tree-based feature graph  $\mathbf{A}_F^*$ ;
22 Generate  $\mathbf{A}_G^*$  using  $\mathbf{A}_S^*$  and  $\mathbf{A}_F^*$ ;
23 return  $\mathbf{A}_G^*, \alpha^*, \tilde{\alpha}^*$  and  $\theta^*$ .

```

respect to θ and \mathbf{A}_S , and thus, we use error backpropagation to update these two variables.

The pseudocode of CGNN with one submodule is given in Algorithm 2.

IV. EXPERIMENTS

In the experiments, we use 28 baseline methods and nine datasets to evaluate the performance of our model. We evaluate CGNN on node classification problems. Our experiments are conducted mainly to answer the following questions.

- Q_1 : Does CGNN have better performance than other baseline methods in semisupervised node classification problems when graphs are unknown?
- Q_2 : Will CGNN perform worse if the C-graph is replaced by a sample graph or a feature graph?
- Q_3 : Is the proposed tree-based feature graph better than the feature graph based on pairwise similarity?
- Q_4 : Does CGNN have stable performance for the three proposed hyperparameters W_C , W_D , and W_I ?
- Q_5 : CGNN can learn multiple C-graphs using multiple submodules, but do multiple C-graphs really improve performance?
- Q_6 : What is the relationship between the number of submodules and the time and space consumption of CGNN?

- Q_7 : Does jointly learning graph structures and parameters improve the classification performance of our proposed CGNN when the structure of the sample graph is known?
- Q_8 : Does CGNN have the scalability to be applied in large datasets?

A. Details of Experiments

1) *Datasets Settings*: In the experiments, we use nine benchmark datasets to evaluate the performance of CGNN. The datasets include three frequently utilized citation network datasets in GNNs, such as Citeseer [58], Cora [59], and Pubmed [60], a music genre classification dataset free music archive (FMA) [61], and four classic datasets from scikit-learn [62]: wine, cancer, digits, and 20news. These datasets are commonly used to evaluate graph learning methods [19]. In addition, ogbn-arxiv [63] is used in our experiments to evaluate the scalability and performance of CGNN in large graph applications. We use the same dataset split of previous work [19], [64]. Details of the above-mentioned datasets are given in Table I.

2) *Experimental Settings*: For the sake of fair comparisons, the hyperparameters of baselines are selected according to the original implementations of the authors. Specifically, we tune hyperparameters using validation datasets and select their values through the validation loss and accuracy. The values of main hyperparameters are given in Table II. In our method, the activation function σ is set as the ReLU function, the function ℓ is set as the cross-entropy error function, and the optimizer is set as AdaMax [65].

3) *Experimental Environment*: We use Python 3.7, TensorFlow 2.0, and XGBoost [57] to implement our method,¹ and we run all the codes on a Hewlett-Packard server with 4× NVIDIA Quadro RTX 6000 24-GB GPUs, 4× Intel Xeon Gold 5210 CPUs, 16 × 32-GB DDR4 RAM, and 5 × 2-TB hard disks.

B. Performance of CGNN Without Knowing Graphs in Advance (Q_1)

CGNN tries to solve the semisupervised node classification problem when graph structures are completely unknown. In this case, we compare CGNN with 14 baseline methods, including the following.

- 1) *Four Classical Machine Learning Algorithms*: Logistic regression (LogReg), feed-forward neural networks (FFNNs), and support vector machines (Linear SVM and RBF SVM).
- 2) *Two Ensemble Learning Methods*: Random forests (RFs) [66], and eXtreme Gradient Boosting (XGB) [57] which is the basis of constructing ensemble trees in our model.
- 3) *Three Popular Semisupervised Learning Methods*: Label propagation (LP) [67], manifold regularization (ManiReg) [68], and SemiEmb [32]. ManiReg and SemiEmb are provided with a k NN sample graph for graph Laplacian regularization.
- 4) *Seven GNN Methods*: Sparse-GCN, Dense-GCN, RBF-GCN, k NN-GCN, k NN-GAT, k NN-AdaGCN, and k NN-LDS. Sparse-GCN creates a sparse Erdős-Rényi random sample graph, dense-GCN creates a dense sample graph

¹Codes is available at <https://github.com/Peter7777777/CGNN>.

TABLE I
DETAILS OF THE EXPERIMENTAL DATASETS. “No” INDICATES THAT THE SAMPLE GRAPH IS UNKNOWN IN THIS DATASET

Dataset	# Samples	# Features	# Sample Graph Edges	# Classes	Train / Val / Test	Type
Wine	178	13	No	3	10 / 20 / 158	Chemical
Cancer	569	30	No	2	10 / 20 / 539	Medical
Digits	1,797	64	No	10	50 / 100 / 1,647	Image
20news	9,607	236	No	10	100 / 200 / 9,307	News
FMA	7,994	140	No	8	160 / 320 / 7,514	Music
Citeseer	3,327	3,703	4,732	6	120 / 500 / 1,000	Citation
Cora	2,708	1,433	5,429	7	140 / 500 / 1,000	Citation
Pubmed	19,717	500	44,338	3	60 / 500 / 1,000	Citation
ogbn-arxiv	169,343	128	1,166,243	40	90,941 / 29,799 / 48,603	Citation

TABLE II
VALUES OF MAIN HYPERPARAMETERS

Hyperparameters	Wine	Cancer	Digits	Citeseer	Cora	20news	FMA	Pubmed	ogbn-arxiv
Learning rate	0.01	0.01	0.1	0.01	0.01	0.01	0.01	0.1	0.01
Number of epochs	800	700	800	800	800	800	1000	700	800
η (in Section III-C1)	50	85	30	12	12	12	12	12	10
w_C (in Section III-C3)	1.2	1	1.2	1.3	1.3	1.3	1.2	1.5	1.5
w_D (in Section III-C3)	1.1	0.8	0.02	1.1	1.1	1.1	0.2	0.2	0.02
w_I (in Section III-C3)	1	0.5	0.01	0.9	0.9	0.9	0.1	0.1	0.01
Number of trees for each class	100								
Max depth of each tree	5								
Default number of submodules	3								
Features of the first hidden layer	64								
L_2 Regularizer parameter	5.00E-05								
Dropout rate	0.5								

TABLE III
PERCENTAGE OF THE AVERAGE ACCURACY (\pm STANDARD DEVIATION) WITHOUT GRAPH STRUCTURE AS INPUT, AND WE FUSE THREE SUBMODULES IN CGNN FOR REPORT ITS RESULTS. EACH RESULT OF CGNN IS REPORTED BASED ON 30 MONTE CARLO EXPERIMENTS. THE TOP THREE RESULTS ARE IN BOLD, AND CGNN ACHIEVES THE BEST RESULTS ON ALL THE DATASETS

Method	Wine	Cancer	Digits	Citeseer	Cora	20news	FMA
LogReg	92.1 (1.3)	93.3 (0.5)	85.5 (1.5)	62.2 (0.0)	60.8 (0.0)	42.7 (1.7)	37.3 (0.7)
Linear SVM	93.9 (1.6)	90.6 (4.5)	87.1 (1.8)	58.3 (0.0)	58.9 (0.0)	40.3 (1.4)	35.7 (1.5)
RBF SVM	94.1 (2.9)	91.7 (3.1)	86.9 (3.2)	60.2 (0.0)	59.7 (0.0)	41.0 (1.1)	38.3 (1.0)
FFNN	89.7 (1.9)	92.9 (1.2)	36.3 (10.3)	56.7 (1.7)	56.1 (1.6)	38.6 (1.4)	33.2 (1.3)
RF	93.7 (1.6)	92.1 (1.7)	83.1 (2.6)	60.7 (0.7)	58.7 (0.4)	40.0 (1.1)	37.9 (0.6)
XGB	85.1 (0.0)	85.7 (0.0)	64.8 (0.0)	56.6 (0.0)	57.6 (0.0)	38.4 (0.0)	25.9 (0.0)
LP	89.8 (3.7)	76.6 (0.5)	91.9 (3.1)	23.2 (6.7)	37.8 (0.2)	35.3 (0.9)	14.1 (2.1)
ManiReg	90.5 (0.1)	81.8 (0.1)	83.9 (0.1)	67.7 (1.6)	62.3 (0.9)	46.6 (1.5)	34.2 (1.1)
SemiEmb	91.9 (0.1)	89.7 (0.1)	90.9 (0.1)	68.1 (0.1)	63.1 (0.1)	46.9 (0.1)	34.1 (1.9)
Sparse-GCN	63.5 (6.6)	72.5 (2.9)	13.4 (1.5)	33.1 (0.9)	30.6 (2.1)	24.7 (1.2)	23.4 (1.4)
Dense-GCN	90.6 (2.8)	90.5 (2.7)	35.6 (21.8)	58.4 (1.1)	59.1 (0.6)	40.1 (1.5)	34.5 (0.9)
RBF-GCN	90.6 (2.3)	92.6 (2.2)	70.8 (5.5)	58.1 (1.2)	57.1 (1.9)	39.3 (1.4)	33.7 (1.4)
k NN-GCN	93.2 (3.1)	93.8 (1.4)	91.3 (0.5)	68.3 (1.3)	66.5 (0.4)	41.3 (0.6)	37.8 (0.9)
k NN-GAT	97.3 (0.1)	93.1 (0.1)	93.4 (0.1)	69.3 (1.1)	65.7 (0.4)	41.3 (0.7)	38.8 (0.2)
k NN-AdaGCN	97.8 (0.5)	94.1 (2.7)	91.9 (3.6)	59.0 (0.6)	58.1 (2.0)	48.5 (1.4)	38.6 (0.5)
k NN-LDS	97.3 (0.4)	94.4 (1.9)	92.5 (0.7)	71.5 (1.1)	71.5 (0.8)	46.4 (1.6)	39.7 (1.4)
SCRL	93.9 (2.1)	92.2 (0.3)	89.5 (1.3)	66.2 (2.7)	65.7 (2.2)	40.3 (1.9)	38.9 (1.7)
SPGRL	97.7 (0.3)	94.0 (0.2)	92.6 (0.4)	69.0 (1.3)	68.9 (0.9)	42.4 (0.6)	37.4 (0.4)
CGNN	98.0 (0.1)	94.5 (0.1)	92.7 (0.4)	73.6 (0.3)	73.4 (0.4)	50.2 (0.6)	40.5 (0.2)

with equal edge probabilities, and RBF-GCN creates a dense RBF kernel sample graph. k NN-GCN, k NN-GAT, k NN-AdaGCN, and k NN-LDS create a k NN sample graph.

To ensure a fair comparison with baseline methods, we adopt the same dataset preprocessing and splitting way as that in k NN-LDS [19]. Table III shows the comparative experiment results of CGNN and the baseline methods. In terms of classification accuracy, we can observe that CGNN achieves the highest node classification accuracies than other methods on all datasets except Digits. CGNN underperforms k NN-GAT

on the digits dataset because every neighbor in the C-graph has the same importance, while k NN-GAT considers the importance of each neighbor utilizing an attention mechanism to assign different weights to nodes, which helps k NN-GAT learn better embeddings than CGNN on Digits. Comparing CGNN with k NN-LDS, we can find that CGNN can achieve 2% higher accuracies than k NN-LDS on Citeseer and Cora, and about 4% higher than k NN-LDS on 20news. CGNN also outperforms k NN-AdaGCN by a significant margin on Citeseer and Cora. CGNN outperforms baseline methods since it captures and utilizes the relationships between features by

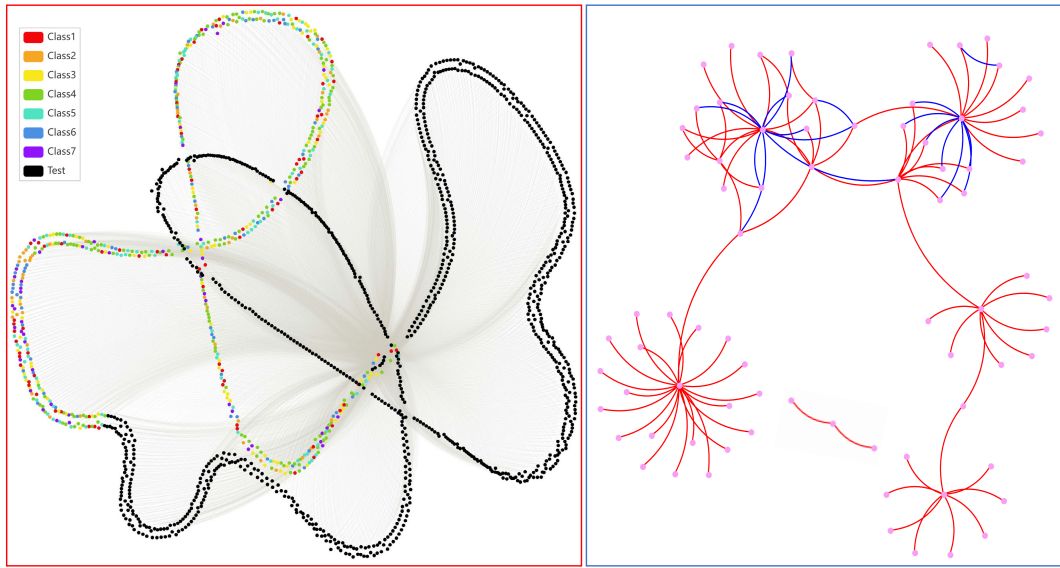


Fig. 4. C-graph learned on the Cora dataset when our method has one submodule. The graph in the red box is the sample graph. In the sample graph, the classes of the black nodes are unknown, the other nodes use different colors to represent their classes, and each gray line is an edge between two samples. The graph in the blue box is a tree-based feature graph. For the convenience of observation, we do not show the feature nodes without direct or indirect relations to other feature nodes. The pink nodes are features in the tree-based feature graph, and the red and blue edges are the direct and indirect relations between features, respectively.

learning C-graphs. The comparison between CGNN and the ensemble learning algorithms RF and XGB also shows that although CGNN uses the same scheme as XGB to implement the ensemble tree models, the performance of CGNN is much higher than that of XGB. This result further highlights the importance of learning C-graphs. As for the standard deviation, the experimental results of CGNN on all the datasets are also significantly lower than those of other GNN baseline methods. We also give a visualization result of the learned C-graph in Fig. 4 and visualization results of α and $\tilde{\alpha}$ in Fig. 5.

We conduct some experiments to show the robustness of CGNN and the results are shown in Table IV. We add random noises to the original dataset. The noises are randomly sampled from a uniform distribution over the interval $[0, 0.2]$, and the proportion of features that are added noises is given in the first column of Table IV. From Table IV, we can find that when the noise ratio reaches 10%, the performance of our algorithm is still better than that of k NN-LDS. When the ratio reaches 20%, the performance of our algorithm is still close to the performance of k NN-LDS on clean datasets. When the noise ratio is 50%, the performance of our algorithm decreases on Cora, Citeseer, and 20news, but, in general, is still comparable to the performance of other baselines on clean datasets. The average node classification results on several datasets (e.g., wine, cancer, and digits) do not change because these datasets are relatively simple, within a certain degree of noise ratio. After reducing the impact of noises through the proposed trainable vector α , the sample graph learned after adding noises is high-quality. Moreover, the features used to generate trees are first filtered by α , which helps find valuable features in noisy data. Furthermore, the trees are constructed by minimizing the classification loss, which ensures the features selected for data splitting are essential for the target classification task and high-quality feature trees. Consequently, our model gets high-quality C-graphs, equivalent to the C-graphs we are learning from these datasets when we do not add noises.

TABLE IV
AVERAGE NODE CLASSIFICATION ACCURACY (%) OF CGNN IN THE ROBUSTNESS EXPERIMENTS. THE FIRST COLUMN INDICATES THE PROPORTION OF FEATURES THAT ARE ADDED NOISES

Noise	Wine	Cancer	Digits	Citeseer	Cora	20news	FMA
1%	98.0	94.5	92.6	73.2	73.1	49.3	40.5
2%	98.0	94.5	92.6	72.9	73.1	49.2	40.5
5%	98.0	94.5	92.6	72.5	72.9	48.2	40.5
10%	98.0	94.5	92.6	72.5	72.5	46.5	40.4
20%	98.0	94.2	92.6	71.7	71.1	44.3	40.4
50%	98.0	94.1	92.4	70.6	70.1	38.7	40.3

C. Ablation Study (Q_2 and Q_3)

Our proposed CGNN has two key components to learning C-graphs: learning sample graphs and learning tree-based feature graphs. In this section, we validate their impacts on the performance via ablation experiments. We detail the variants of CGNN as follows.

- 1) *SGNN*: Sample graph neural network (SGNN) only learns the sample graphs and uses them to replace the C-graphs in CGNN. In other words, SGNN is equivalent to CGNN without learning the tree-based feature graphs.
- 2) *FGNN*: Tree-based feature graph neural network (FGNN) is the method that only learns the feature graphs. Notably, FGNN uses the tree-based feature graphs generated in the final iteration and the sample graphs generated in the first iteration to conduct node classification. FGNN requires sample graphs since learning feature graphs and solving node classification depend on sample relations in our approach. Using the sample graphs generated at the initial iteration facilitates us to compare the difference between learning C-graphs and only learning tree-based feature graphs without learning the sample graphs.

The results of the ablation experiments are given in Fig. 6. From the node classification accuracies in Fig. 6, we see that CGNN outperforms SGNN and FGNN. In general, the results imply that both learning sample graphs and tree-based feature

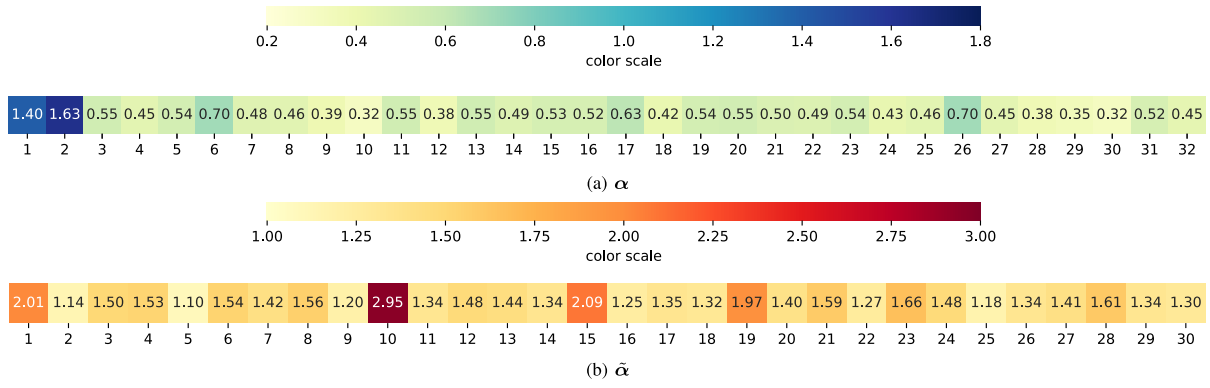


Fig. 5. Heatmap visualization results of α and $\tilde{\alpha}$ on the cancer dataset when our method uses one submodule. The scale value in the horizontal direction represents the feature number. Recall that $\alpha \in \mathbb{R}^{F+C}$ and $\tilde{\alpha} \in \mathbb{R}^F$, so α is longer than $\tilde{\alpha}$. (a) α . (b) $\tilde{\alpha}$.

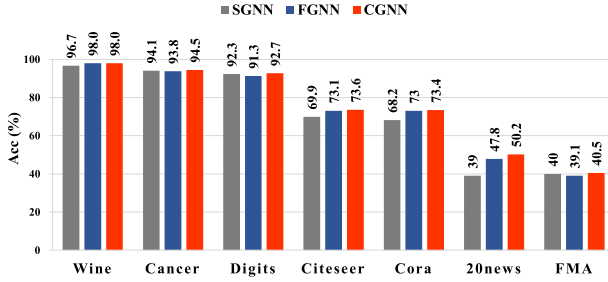


Fig. 6. Comparison of the percentage of the average node classification accuracy (Acc) of CGNN and its two variants, SGNN and FGNN. SGNN denotes only learning the sample graphs in CGNN. FGNN denotes only learning the feature graphs in CGNN.

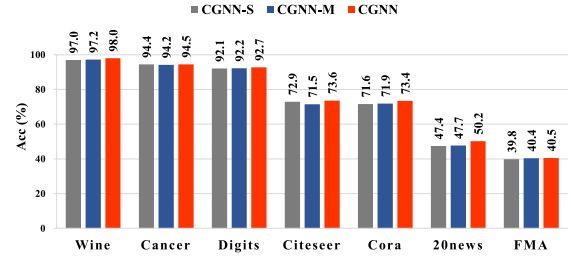


Fig. 7. Comparison of the percentage of the average node classification accuracy (Acc) of CGNN and its other two variants, CGNN-S and CGNN-M. Difference from CGNN, which uses two hidden layers to learn the embeddings for classification, CGNN-S uses a simple fully connected layer, and CGNN-M uses multiple hidden layers to learn the embeddings for classification. The hidden layers number of CGNN-M is set as five to report the results.

graphs are conducive to the model performance. It is worth noting that although the results of SGNN or FGNN are close to those of CGNN sometimes on some datasets, the performance of CGNN is much better than its two variants on 20news. The main reason is that only learning sample relations or feature relations are unable to get satisfactory embeddings for the downstream neural network to classify nodes nicely on the dataset. Therefore, both the sample graph and the tree-based feature graph are important in graph node embedding.

We also compare CGNN with its two other variants.

- 1) *CGNN-S*: CGNN with a simple (CGNN-S) fully connected layer only uses a simple fully connected layer to learn the embeddings for classification.
- 2) *CGNN-M*: CGNN with multiple (CGNN-M) layers uses multiple hidden layers to learn the embeddings for classification.

The main difference between CGNN, CGNN-S, and CGNN-M is the number of network layers. The neural network in CGNN has two hidden layers. Fig. 7 gives the comparison results between the three methods, and we set the hidden layer number as five to report the results of CGNN-M. From Fig. 7, we observe that CGNN has the best performance, which means two hidden layers are most appropriate for the current CGNN. However, since the results of CGNN-M on several datasets (e.g., Citeseer, Cora, and 20news) are inferior to that of CGNN, this indicates that CGNN also exists the limitation of oversmoothing, i.e., with the number of layers increasing, the representations of nodes in CGNN become indistinguishable. Thus, we think it is significant for future work to design an improved CGNN that not only jointly learns the C-graph and neural network parameters but also deals with the problem of oversmoothing.

To further compare the difference between our proposed tree-based feature graph and pairwise similarity-based feature graph, we propose tree-based feature graph embedding neural network (T-EmbNN). T-EmbNN is an extremely simplified variant method of CGNN, which only uses tree-based feature graphs to conduct node classification. Specifically, we use $\tilde{\mathbf{X}} = \mathbf{X}$ to replace $\tilde{\mathbf{X}} = \mathbf{D}_S^{-(1/2)}(\mathbf{A}_S + \mathbf{I})\mathbf{D}_S^{-(1/2)}\mathbf{X}$ in (11) and (12) and Algorithm 1, and we use $\mathbf{X}^{(2)} = \sigma(\mathbf{X}^{(1)}\mathbf{\Theta}_2)$ to replace $\mathbf{X}^{(2)} = \sigma(\mathbf{D}_S^{-(1/2)}(\mathbf{A}_S + \mathbf{I})\mathbf{D}_S^{-(1/2)}\mathbf{X}^{(1)}\mathbf{\Theta}_2)$ in (2). In order to control variables, the number of submodules of T-EmbNN is set to one. Then, we compare T-EmbNN with two methods that use the pairwise similarity of features to generate a feature graph. These two methods are denoted as E-EmbNN and C-EmbNN and use the popular Euclidean distance and cosine similarity to measure the similarity between paired features, respectively. Both E-EmbNN and C-EmbNN use a k NN graph, $k \in \{2, 3, \dots, 20\}$, to construct the sparsified feature graph. In addition to E-EmbNN and C-EmbNN, we also compare T-EmbNN with a neural network approach that does not use any graph structures, denoted as NN. NN takes the simple form: $\text{Softmax}(\sigma(\sigma(\mathbf{X}\mathbf{\Theta}_1)\mathbf{\Theta}_2))$. In other words, the model of NN only has an input layer, two hidden fully connected layers, and an output layer. The results of the comparative experiments are given in Fig. 8. From Fig. 8, we can observe that T-EmbNN achieves the best performance in most cases. It is worth noting that T-EmbNN is significantly superior to other methods on Citeseer, Cora, and 20news. It can also be observed that E-EmbNN is not always better than C-EmbNN, and vice versa. This implies that it is difficult to find a pairwise feature similarity metric that is appropriate to all these datasets. Moreover, on some datasets, both the performance

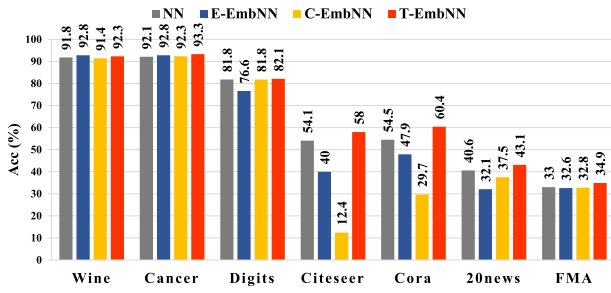


Fig. 8. Comparative experiments on the average node classification accuracy (Acc) of learning tree-based feature graphs. T-EmbNN only learns a tree-based feature graph, which is an extremely simplified variant method of CGNN. Both E-EmbNN and C-EmbNN learn a k NN feature graph, and they use the Euclidean distance and cosine similarity to measure the pairwise feature similarity, respectively. NN denotes a neural network approach that does not use any graph structures.

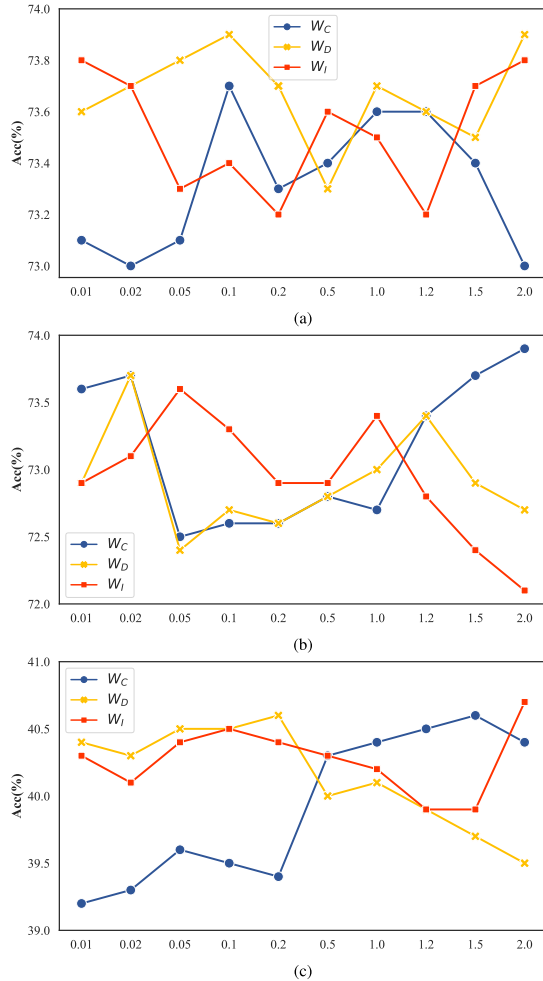


Fig. 9. Influence of the hyperparameters, W_C , W_D , and W_I , on Citeseer, Cora, and FMA. (a) Citeseer. (b) Cora. (c) FMA.

of E-EmbNN and C-EmbNN are even inferior to NN. In this experiment, T-EmbNN achieves better performance on almost all the datasets, which demonstrates the importance of learning feature importance and proper feature combination.

D. Sensitivity Analysis of Hyperparameters (Q_4)

In this section, we analyze the three proposed hyperparameters, W_C , W_D , and W_I . First, we test them by setting each of their values via a grid search in $\{0.01, 0.02, 0.05, 0.1, 0.2, 0.5, 1.0, 1.2, 1.5, 2.0\}$, and fix others values as

TABLE V

AVERAGE NODE CLASSIFICATION ACCURACY (%) OF CGNN WITH VARYING NUMBERS OF SUBMODULES WHEN NO GRAPH STRUCTURE IS PROVIDED AS INPUT. HERE, R DENOTES THE NUMBER OF SUBMODULES

R	Wine	Cancer	Digits	Citeseer	Cora	20news	FMA
1	98.0	94.4	92.5	73.0	73.1	48.5	40.4
2	98.0	94.5	92.6	73.1	73.2	49.9	40.4
3	98.0	94.5	92.7	73.6	73.4	50.2	40.5
4	98.0	94.5	92.8	73.1	73.7	50.3	40.2
5	98.0	94.4	92.6	73.6	73.8	50.7	40.4

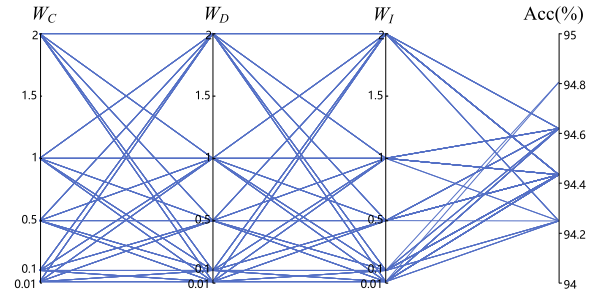


Fig. 10. Influence of the hyperparameters, W_C , W_D , and W_I on Cancer.

given in Table II. We report the average classification accuracy of ten runs, and the results are shown in Fig. 9. We find that our model mostly performs better than the baselines in Table III. Even if we only look at the worst results on each dataset, the performance of CGNN still stays comparable. Furthermore, in order to analyze the sensitivity of the three hyperparameters more comprehensively, we conduct an experiment by setting all of their values in $\{0.01, 0.1, 0.5, 1.0, 2.0\}$ through grid search. Since there are a total of 125 combinations and three variables are involved, we choose parallel coordinates to show the experimental results. We report the average classification accuracy of five runs on Cancer, and the results are shown in Fig. 10. We observe that our model has stable performance as the results mostly reach the fine-tuned accuracy in Table III.

E. Space and Time Consumption (Q_5 and Q_6)

We consider that CGNN can improve the classification accuracy by fusing different C-graphs learned in multiple submodules. We perform experiments to study the impact of the number of submodules on the classification accuracy of CGNN. Table V shows the classification accuracy of CGNN with varying numbers of submodules when graphs are unknown.

In Table V, the number in the first column of each row indicates the number of submodules in CGNN. From Table V, we see that properly increasing the number of submodules, especially from 1 to 3, can improve the performance of our method. Thus, after considering the tradeoff between computation complexity and performance, we choose $R = 3$ as the default number of submodules in our method. Figs. 11 and 12 give the results of average running time (second) and memory (MB) of CGNN under different submodules, respectively. From Figs. 11 and 12, we observe that the time and space consumption of CGNN increase linearly with the number of submodules in general.

Complexity: The time complexity of the learning sample graph and tree-based feature graph are $O(N^2)$ and $O(NFTD)$, respectively. N and F are the number of samples and features of the input dataset, respectively; T and D

TABLE VI

BRIEF DESCRIPTIONS OF THE BASELINE METHODS FOR COMPARING WITH CGNN WHEN THE SAMPLE GRAPH IS KNOWN A PRIORI

Algorithm	Description
GIN [69]	A GNN method under the neighborhood aggregation framework.
GCN [12]	A method aggregates neighbor information with spectral convolutions on graphs.
GAT [13]	A method aggregates neighbor information by attention mechanism.
Graphite [45]	A latent variable generative model based on variational autoencoding.
BGCN-T [39]	A method improves GCN with bilinear interactions between neighbor nodes.
BGAT-T [39]	A method improves GAT with bilinear interactions between neighbor nodes.
SGC [70]	A simple and efficient GCN-based method.
Shoestring [71]	A graph-based method with severely limited labeled data.
APNP [72]	A method utilizes the relationship between GCN and personalized PageRank.
DAGNN [15]	A deep adaptive GNN method can capture sample relations from large and adaptive receptive fields.
SelfSAGCN [73]	A GCN-based simple and effective self-supervised semantic alignment method.
k NN-LDS [19]	A method jointly learns the graph structure of samples and parameters of a neural network.
GCNII [14]	A simple and deep GCN-based method via initial residual and identity mapping.
(A+ k NN)-GCN [64]	A GCN-based method uses the given sample graph and a constructed k NN feature graph.
SimP-GCN [64]	A node similarity preserving GCN-based method.
SCRL [74]	A self-supervised consensus graph representation learning method.
SPGRL [75]	A structure-preserving graph representation learning method.
Ortho-GCN [76]	A method improves GCN with orthogonal graph convolutions.
Ortho-GCNII [76]	A method improves GCNII with orthogonal graph convolutions.

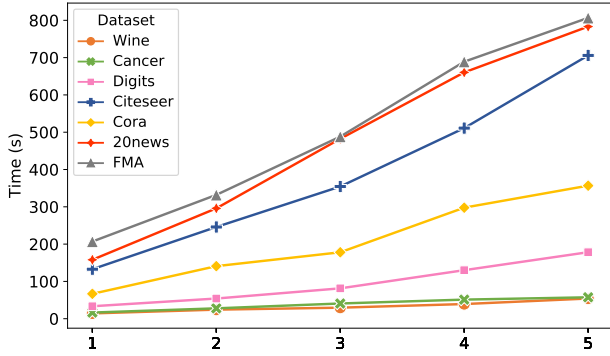


Fig. 11. Average training time (second) of CGNN when using different submodules. The horizontal coordinate indicates the number of submodules.

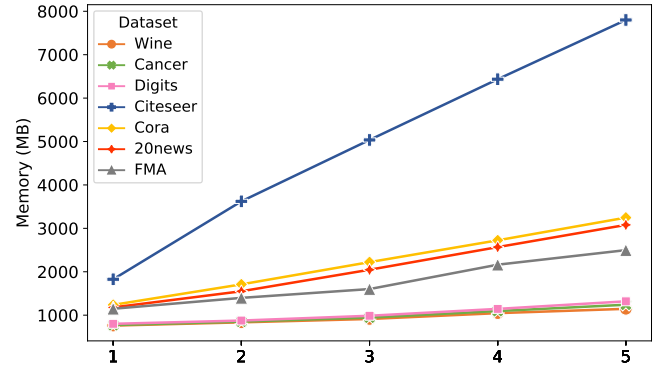


Fig. 12. Average memory (MB) of CGNN when using different submodules. The horizontal coordinate indicates the number of submodules.

denote the number of trees and the max depth of each tree, respectively. Thus, the time complexity of our method is $O(N^2 + NFTD + |\mathcal{E}_S||\mathcal{E}_F|U)$, where U is the number of hidden units in the network; \mathcal{E}_S and \mathcal{E}_F denote the edge number of the learned sample graph and tree-based feature graph, respectively. The time complexity of the baseline method k NN-GCN is $O(N^2 + |\mathcal{E}_S|U)$, and other GCN-based baselines have similar time complexity with k NN-GCN. Our method has higher time complexity than these methods since it learns and captures feature relations while others do not.

F. Performance of CGNN When the Sample Graph Is Known (Q_7)

Our method is proposed mainly for scenarios where graph structures are completely inaccessible and noisy. In this experiment, we will show our method is better than baselines when graph structures are known a priori. To this end, we compare our method with many GNNs on datasets with known graphs. Specifically, we learn the tree-based feature graph using the known sample graph and then obtain the C-graph. We select some recently proposed and state-of-the-art algorithms for comparative experiments. Brief descriptions of the baseline methods are given in Table VI.

Table VII shows the results of CGNN and the baseline methods. We provide a comparison on each dataset by following the same dataset splitting setup as that in previous works [39], [64], and some reported results of the baseline methods also come from these publications. From Table VII, we observe CGNN achieves the best accuracy on all three datasets. The main reason why CGNN performs better than other baselines is that CGNN learns the tree-based feature graphs and thus captures and utilizes the beneficial feature relationships.

G. Scalability of CGNN in Large Dataset Applications (Q_8)

We evaluate the performance of our method on eight datasets with less than 20 000 nodes in both scenarios, where the graph structure is unknown or known. In this section, we conduct experiments on ogbn-arxiv with 169 343 nodes to further evaluate the scalability and performance of CGNN on large datasets. We compare CGNN with baseline methods on ogbn-arxiv under the same dataset splitting setup as that in previous works [63], [76]. In Table VIII, we give the results of CGNN and baselines, and the number of hidden layers is set to two for a fair comparison. From Table VIII, we know that CGNN is applicable to large datasets in scenarios

TABLE VII

PERCENTAGE OF THE AVERAGE ACCURACY (\pm STANDARD DEVIATION) WHEN GRAPH IS PROVIDED AS INPUT. EACH RESULT OF CGNN IS REPORTED BASED ON 30 MONTE CARLO EXPERIMENTS. THE TOP THREE RESULTS ARE IN BOLD, AND CGNN ACHIEVES THE BEST RESULTS ON ALL THE DATASETS

Method	Cora	Citeseer	Pubmed
GIN	79.7 (0.8)	69.4 (0.6)	78.5 (0.2)
GCN	81.2 (0.4)	71.1 (0.7)	78.5 (1.0)
GAT	83.1 (0.7)	72.5 (0.7)	79.0 (0.3)
Graphite	82.1 (0.1)	71.0 (0.1)	79.3 (0.1)
BGCN-T	82.0 (0.1)	71.9 (0.1)	79.4 (0.1)
BGAT-T	84.2 (0.4)	74.0 (0.3)	79.8 (0.3)
SGC	81.7 (0.6)	71.3 (1.1)	78.9 (1.3)
Shoestring	81.9 (2.1)	69.5 (2.4)	79.7 (4.5)
APNP	83.3 (0.4)	71.8 (0.9)	80.1 (1.3)
DAGNN	84.4 (0.4)	73.3 (0.7)	80.5 (0.9)
SelfSAGCN	83.8 (0.5)	73.5 (1.2)	80.7 (1.5)
kNN-LDS	84.1 (0.4)	75.0 (0.4)	80.0 (0.5)
GCNII	85.5 (0.5)	73.4 (0.6)	80.2 (0.4)
(A+kNN)-GCN	79.1 (0.7)	71.1 (0.7)	80.8 (0.9)
SimP-GCN	82.8 (0.5)	72.6 (0.7)	81.1 (0.6)
SCRL	72.5 (1.0)	73.6 (1.4)	79.6 (1.1)
SPGRL	81.3 (0.4)	75.1 (1.1)	77.6 (1.0)
Ortho-GCN	82.8 (1.0)	72.3 (1.2)	80.7 (0.5)
Ortho-GCNII	85.6 (0.7)	74.1 (1.1)	81.2 (0.6)
CGNN	86.3 (0.5)	75.7 (0.7)	81.6 (0.7)

TABLE VIII

PERCENTAGE OF THE AVERAGE ACCURACY (\pm STANDARD DEVIATION) IN THE LARGE GRAPH APPLICATION. EACH RESULT OF CGNN IS REPORTED BASED ON 30 MONTE CARLO EXPERIMENTS

Dataset	Scenario	Method	Result
ogbn-arxiv	graph unknown a priori	kNN-GCN	53.5 (0.3)
		kNN-GAT	54.2 (0.2)
		CGNN	54.9 (0.3)
	graph known a priori	GCN	71.3 (0.3)
		GAT	71.3 (0.2)
		GCNII	71.2 (0.2)
		Ortho-GCN	71.3 (0.3)
		Ortho-GCNII	71.4 (0.2)
		CGNN	72.2 (0.3)

where graph structure is unknown or known a priori, and CGNN achieves the best performance in both scenarios. The results demonstrate that learning the C-graph is vital for graph learning, especially when graph structure is unknown a priori.

V. CONCLUSION

In this work, we have presented a new GNN method named CGNN to solve semisupervised classification problems. Unlike other GNN methods, our C-graphs not only characterize sample similarities but also imply feature importance and combination preference. Furthermore, CGNN can learn multiple C-graphs from different aspects. Each C-graph is obtained by simultaneously optimizing the graph structure of samples and the tree structure of features, which ensures robustness. Experiments on multiple datasets show our model achieves the best performance on almost all the datasets and is robust to feature noises.

In terms of limitations, just like many GNN methods that face the problem of over-smoothing, there is also an over-smoothing phenomenon in CGNN. Although the

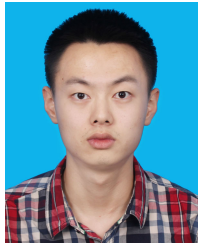
oversmoothing problem has aroused many concerns recently, the existing methods are not good enough to help CGNN. The key reason is that existing approaches do not have optimization schemes for the C-graph. Thus, employing existing techniques in our framework cannot ensure a high-quality C-graph is obtained by jointly learning the C-graph and neural network parameters. How to improve CGNN to avoid oversmoothing deserves further study. Another limitation is that our method has more time complexity than most baselines since it learns and captures feature relations while others do not. Two solutions may be investigated in the future to improve the computation efficiency: 1) our method learns the sample graph of the C-graph using all the samples, a node cluster algorithm proposed in [77] may help our model use a part of samples to infer the sample graph of all samples quickly and 2) since our model includes many submodules to learn C-graphs, an approach that can split the training process into optimizing many subtasks [78] may be involved in our model to speed up the training process.

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