

# Large-Scale Learnable Graph Convolutional Networks

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## ABSTRACT

Convolutional neural networks (CNNs) have achieved great success on grid-like data such as images, but face tremendous challenges in learning from more generic data such as graphs. In CNNs, the trainable local filters enable the automatic extraction of high-level features. The computation with filters requires a fixed number of ordered units in the receptive fields. However, the number of neighboring units is neither fixed nor are they ordered in generic graphs, thereby hindering the applications of convolutional operations. Here, we address these challenges by proposing the learnable graph convolutional layer (LGCL). LGCL automatically selects a fixed number of neighboring nodes for each feature based on value ranking in order to transform graph data into grid-like structures in 1-D format, thereby enabling the use of regular convolutional operations on generic graphs. To enable model training on large-scale graphs, we propose a sub-graph training method to reduce the excessive memory and computational resource requirements suffered by prior methods on graph convolutions. Our experimental results on node classification tasks in both transductive and inductive learning settings demonstrate that our methods can achieve consistently better performance on the Cora, Citeseer, Pubmed citation network, and protein-protein interaction network datasets. Our results also indicate that the proposed methods using sub-graph training strategy are more efficient as compared to prior approaches.

## CCS CONCEPTS

• **Computing methodologies** → **Neural networks**; *Structured outputs*; Artificial intelligence;

## KEYWORDS

Deep learning, graph convolutional networks, graph mining, large-scale learning

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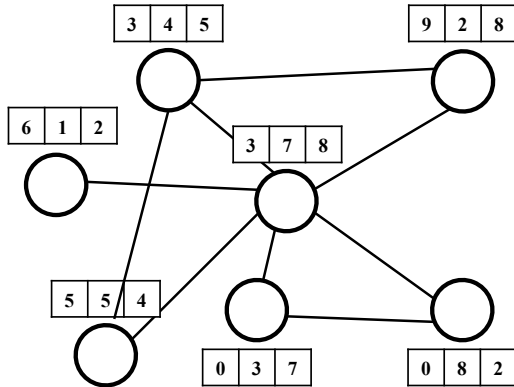
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## 1 INTRODUCTION

Deep learning methods are becoming increasingly powerful in solving various challenging artificial intelligence tasks. Among these deep learning methods, convolutional neural networks (CNNs) [18] have demonstrated promising performance in many image-related applications, such as image classification [5], semantic segmentation [2], and object detection [10, 22]. A variety of CNN models have been proposed to continuously set the performance records [11, 17, 24, 26]. In addition to images, CNNs have also been successfully applied to natural language processing tasks such as neural machine translation [3, 6, 19]. One common characteristic behind these tasks is that the data can be represented by grid-like structures. This enables the use of convolutional operations in the form of the same local filters scanning every position on the input. Unlike traditional hand-crafted filters, the local filters used in convolutional layers are trainable. The networks can automatically decide what kind of features to extract by learning the weights in these trainable filters, thereby avoiding hand-crafted feature extraction [29].

In many real-world applications, the data can be naturally represented as graphs, such as social, citation, and biological networks. Figure 1 provides an illustration of graph data. Many interesting discoveries can be made by analyzing these graph data, such as social network analysis [8]. An important task on graph data is node classification [15, 28], in which models make predictions for every node in a graph based on node features and graph topology. As mentioned above, CNNs, with the power of automatic feature extraction, have achieved great success on tasks with grid-like data, which can be considered as special cases of graph data. Therefore, applying deep learning models, especially CNNs, on graph tasks is appealing. However, using regular convolutional operations on generic graphs faces two main challenges. These challenges are resulted from the fact that regular convolutions require the number of neighboring nodes for each node remains the same, and these neighboring nodes are ordered. In generic graphs, the numbers of neighboring nodes usually differ for different nodes in a graph. In addition, among the neighboring nodes of a node, there is no ranking information based on which we can order them to ensure the output is deterministic. In this work, we analyze the necessity of having a fixed number of ordered neighboring nodes in regular convolutional operations and propose elegant solutions to address these challenges.

Several recent studies tried to apply convolutional operations on generic graphs. Graph convolutional networks (GCNs) [15] proposed to use a convolution-like operation to aggregate features of all adjacent nodes for each node, followed by a linear transformation to generate new a feature representation for a given node. Specifically, all feature vectors in the neighborhood, including the feature vector of the central node itself, are summed up, weighted by non-trainable weights depending on the number of



**Figure 1: An illustration of graph data. There are 7 nodes in this graph and each node has 3 features. Each node in this graph may have a different number of neighboring nodes, and there is no relative order among them.**

neighbors. This can be thought of as a convolution-like operation which, however, is intrinsically different from the regular convolutional operation in two aspects. First, it does not use the same local filter to scan every node; that is, nodes that have different numbers of adjacent nodes have filters of different sizes and weights. Second, the weights in the filters are the same for all neighboring nodes in the receptive field as they are determined by the number of neighbors. Consequently, the weights are not learned. Graph attention networks (GATs) [28] employed the attention mechanism [1] to obtain different and trainable weights for adjacent nodes by measuring the correlation between their feature vectors and that of the central node. Yet graph attention operation still differs from the regular convolution which learns weights in local filters directly. Moreover, the attention mechanism requires extra computation in terms of pairs of feature vectors, resulting in excessive memory and computational resource requirements in practice.

In this work, we make two major contributions to applying CNNs on generic graph data. First, we propose the learnable graph convolutional layer (LGCL) to enable the use of regular convolutional operations on graphs. Note that prior studies modified the original convolutional operations to fit them for graph data. In contrast, our LGCL transforms the graphs to enable the use of regular convolutions. Our models based on LGCL achieve better performance on both transductive learning and inductive node classification tasks, as demonstrated by our experimental results. Second, we observe another limitation of prior methods; that is, their training process takes the adjacency matrix of the whole graph as an input. This requires excessive memory and computational resources when the graph has a large amount of nodes, which is usually the case in real-world tasks. In order to overcome this limitation, we develop a sub-graph training method, which is a simple yet effective approach to allow the training of deep learning methods on large-scale graph data. The sub-graph training method can significantly reduce the amount of required memory and computational resources, with negligible loss in terms of model performance.

## 2 RELATED WORK

A few recent studies have tried to apply convolutional operations on graph data. Graph convolutional networks (GCNs) were introduced in [15] and achieved the state-of-art performance on several node classification tasks. The authors defined and used a convolution-like operation termed the spectral graph convolution. This enables CNNs to directly operate on graphs. Basically, each layer in GCNs updates the feature vector representation of each node in the graph by considering the features of neighboring nodes. To be specific, the layer-wise forward-propagation operation of GCNs can be expressed as

$$X_{l+1} = \sigma(\hat{D}^{-\frac{1}{2}} \hat{A} \hat{D}^{-\frac{1}{2}} X_l W_l), \quad (1)$$

where  $X_l$  and  $X_{l+1}$  are the input and output matrices of layer  $l$ , respectively. For both matrices, the numbers of rows are the same, corresponding to the number of nodes in the graph, while the numbers of columns can be different, depending on the dimensions of the input and output feature space. In Eq (1),  $\hat{A} = A + I$  is used to aggregate feature vectors of adjacent nodes, where  $A$  is the adjacency matrix of the graph, and  $I$  is the identity matrix. Also,  $\hat{A}$  is used, instead of  $A$ , because the layers need to add self-loop connections to make sure that the old feature vector of the node itself is taken into consideration when updating the representation of a node.  $\hat{D}$  is the diagonal node degree matrix, which is used to normalize  $\hat{A}$  so that the scale of feature vectors after aggregation remains the same.  $W_l$  is a trainable weight matrix and represents a linear transformation that changes the dimension of feature space. Therefore, the dimension of  $W^l$  depends on how many features that each node in the input and output have, *i.e.*, the number of columns in  $X_l$  and  $X_{l+1}$ , respectively.  $\sigma(\cdot)$  denotes an activation function like ReLU.

We analyze the convolution-like operation, which is the feature aggregation step through pre-multiplying  $X_l$  by  $\hat{D}^{-\frac{1}{2}} \hat{A} \hat{D}^{-\frac{1}{2}}$ . Consider a node with a feature vector corresponding to the  $i$ -th row in  $X_l$ . The aggregation output, controlled by the  $i$ -th row in  $\hat{D}^{-\frac{1}{2}} \hat{A} \hat{D}^{-\frac{1}{2}}$ , is a weighted sum of the feature vectors of all of its adjacent nodes, including the node itself. We can see that the operation is equivalent to having a local filter for each node, whose receptive field consists of the node itself and all its neighboring nodes. As is common that nodes in a generic graph have different numbers of adjacent nodes, the receptive field size varies, resulting in different local filters. This is a key difference from the regular convolutional operation, where the same local filter is applied to scan each position in grid-like data. Moreover, while using local filters of different sizes for graph data seems reasonable, it is worth noting that there is no trainable parameter in  $\hat{D}^{-\frac{1}{2}} \hat{A} \hat{D}^{-\frac{1}{2}}$ . In addition, each adjacent node receives the same weight in the weighted sum, which makes it a simple average. While CNNs achieve the power of automatic feature extraction by learning the weights in local filters, this non-trainable aggregation operation in GCNs limits the capability of CNNs on generic graph data.

From this perspective, graph attention networks (GATs) [28] tried to enable learnable weights when aggregating neighboring feature vectors by employing the attention mechanism [1, 27]. Like GCNs, each node still has a local filter with a receptive field covering the node itself and all of its adjacent nodes. When performing the

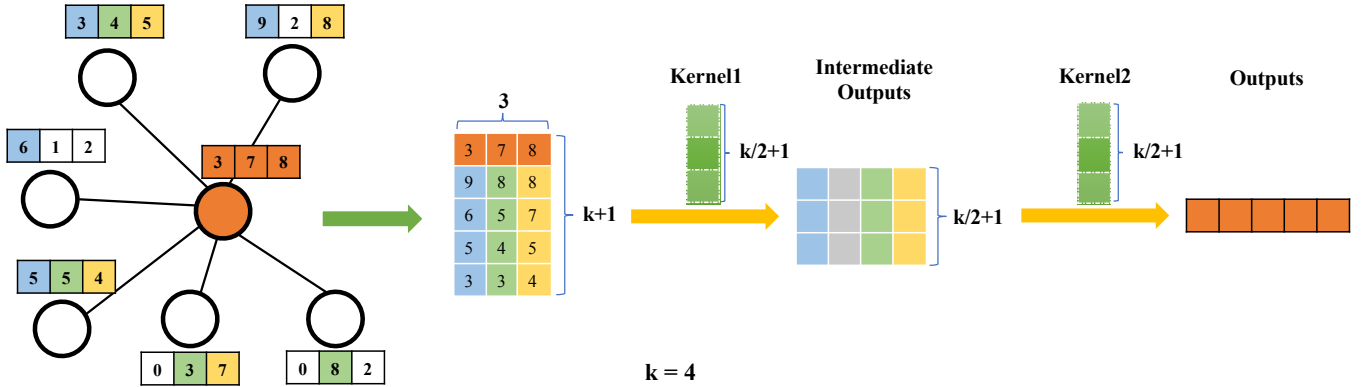


Figure 2: An illustration of a learnable graph convolutional layer (LGCL). We consider a node with 6 adjacent nodes. Each node has three features, represented by a 3-component feature vector. This layer selects  $k = 4$  nodes in the neighborhood and employs a 1-D CNN to produce a new vector representation of five features for the central node, color-coded in orange. The left part describes the process of selecting the  $k$ -largest values for each feature from neighboring nodes. It can be seen from the graph that there are 6 neighbors. Since  $k = 4$ , for each feature, four largest values are selected from the neighborhood based on the ranking. For example, the results of this selection process for the first feature is  $\{9, 6, 5, 3\}$  out of  $\{9, 6, 5, 3, 0, 0\}$ . By repeating the same process for the other two features, we obtain  $(k + 1)$  3-component feature vectors, including that of the orange node itself. Concatenating them gives a 1-D data of grid-like structure, which has  $(k + 1)$  positions and 3 channels. Afterwards, a 1-D CNN is applied to generate the final feature vector. Specifically, we use two convolutional layers with a kernel size of  $(k/2 + 1)$  and without padding. The numbers of output channels are 4 and 5, respectively. In practice, the 1-D CNN can be any CNN model, as long as the final output is a vector, serving as the new feature representation of the central node.

weighted sum of feature vectors, each neighbor receives a different weight by measuring the correlation between its feature vector and that of the central node. Mathematically, for a node  $i$  and one of its adjacent nodes  $j$ , the correlation measurement process between layer  $l$  and  $l + 1$  is given by

$$\begin{aligned} e_l^{i,j} &= a_l(W_l x_l^i, W_l x_l^j) \\ \alpha_l^{i,j} &= \text{softmax}(e_l^{i,j}), \end{aligned} \quad (2)$$

where  $x_l^i$  and  $x_l^j$  represent the corresponding feature vectors, *i.e.*, the  $i$ -th and  $j$ -th row in  $X_l$ , respectively,  $W_l$  is a shared linear transformation and  $a_l$  represents a single-layer feed-forward neural network,  $\alpha_l^{i,j}$  is the weight for node  $j$  in the feature aggregation operation of node  $i$ . Although in this way, GATs provide different and trainable weights to different adjacent nodes, the learning process differs from that of regular CNNs where weights in local filters are learned directly. Also, the attention mechanism requires extra computation between a node and all of its adjacent nodes, which will cause memory and computational resource problems in practice.

Unlike these prior models, which modified the regular convolutional operations to fit them for generic graph data, we instead propose to transform graphs into grid-like data to enable the use of CNNs directly. This idea was previously explored in [20]. However, the transformation in [20] is implemented in the preprocessing process while our method includes the transformation in the networks. Additionally, we introduce a sub-graph training method in this work, which is a simple yet effective approach to allow large-scale training.

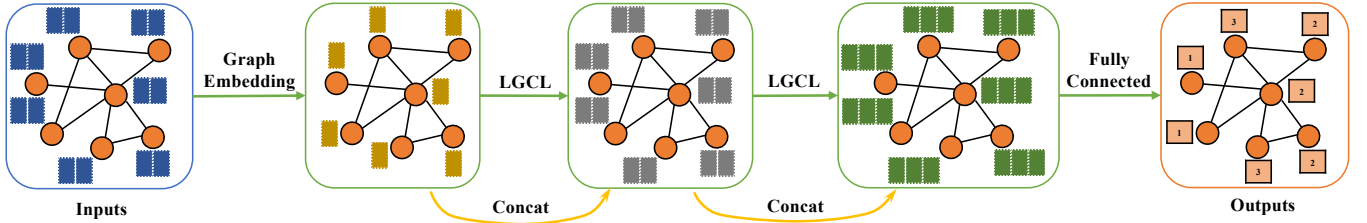
### 3 METHODS

In this section, we introduce the learnable graph convolutional layer (LGCL) and the sub-graph training strategy on generic graph data. Based on these developments, we propose the large-scale learnable graph convolutional networks (LGCNs).

#### 3.1 Challenges of Applying Convolutional Operations on Graph Data

In order to apply regular convolutional operations on graphs, we need to overcome two main challenges that are caused by two major differences between generic graphs and grid-like data. First, the number of adjacent nodes usually varies for different nodes in a generic graph. Second, we cannot order the neighboring nodes in generic graphs, since there is no ranking information among them. For example, in a social network, each person in the network can be seen as a node and the edges represent friendships between people. Obviously, the number of adjacent nodes differs for each node since people can have different numbers of friends. Meanwhile, it is hard to order these friends without additional information for ranking.

Note that grid-like data can be viewed as a special type of graph data, where each node has a fixed number of ordered neighbors. As convolutional operations apply directly on grid-like data such as images, we analyze why the two characteristics mentioned above are necessary to performing regular convolutions. To see the need of having a fixed number of adjacent nodes with ranking information, consider a convolutional filter with a size of  $3 \times 3$  scanning an image. We think of the image as a special graph by thinking of each pixel as a node. During the scan, the computation involves a central node with  $3 \times 3 - 1 = 8$  adjacent nodes each time. These



**Figure 3: An illustration of the proposed learnable graph convolutional network (LGCN). In this example, the nodes in the input have two features. The input feature vectors are transformed into low-dimensional representations using a graph embedding layer. After that, we stack two LGCL layers with skip concatenation connections to refine the feature vectors of each node. Finally, a fully-connected layer is used for node classification. There are three different classes in this example.**

8 nodes become neighbors of the central node by having edges connecting them in the special graph. Meanwhile, we can order these neighboring nodes by their relative positions with respect to the central node. This is crucial to convolutional operations since the correspondence between weights in the filter and nodes in the graph must be maintained during the scan. For instance, in the example above, the upper left weight in the  $3 \times 3$  filter should always be multiplied with the neighboring node at the top left of the central node. Without such ranking information, the outputs of convolution operations are no longer deterministic. We can see from the above discussions that it is challenging to directly apply regular convolutional operations on generic graph data. To address these two challenges, we propose an approach to transform generic graphs into grid-like data.

### 3.2 Learnable Graph Convolutional Layers

To enable the use of regular convolutional operations on generic graphs, we propose the learnable graph convolutional layer (LGCL). Following the notations defined in Section 2, the layer-wise propagation rule of LGCL is formulated as

$$\begin{aligned}\tilde{X}_l &= g(X_l, A, k), \\ X_{l+1} &= c(\tilde{X}_l),\end{aligned}\quad (3)$$

where  $A$  is the adjacency matrix,  $g(\cdot)$  is an operation that performs the  $k$ -largest node selection to transform generic graphs to data of grid-like structures, and  $c(\cdot)$  denotes a regular 1-D CNN that aggregates neighboring information and outputs a new feature vector for each node. We discuss  $g(\cdot)$  and  $c(\cdot)$  separately below.

**$k$ -largest Node Selection.** We propose a novel method known as the  $k$ -largest node selection to achieve the transformation from graphs to grid-like data, where  $k$  is a hyper-parameter of LGCL. After this operation, each node aggregates neighboring information and is represented in a 1-D grid-like format with  $(k+1)$  positions. The transformed data is then fed into a 1-D CNN to generate the updated feature vector.

Suppose  $X_l \in \mathbb{R}^{N \times C}$  with row vectors  $x_l^1, x_l^2, \dots, x_l^N$ , representing a graph of  $N$  nodes where each node has  $C$  features. We are given the adjacency matrix  $A \in \mathbb{N}^{N \times N}$  and a fixed  $k$ . Now consider a specific node  $i$  whose feature vector is  $x_l^i$  and it has  $n$  neighboring nodes. Through a simple look-up operation in  $A$ , we can obtain the indices of these adjacent nodes, say  $i_1, i_2, \dots, i_n$ . Concatenating the corresponding feature vectors  $x_l^{i_1}, x_l^{i_2}, \dots, x_l^{i_n}$  outputs a matrix

$M_l^i \in \mathbb{R}^{n \times C}$ . Without the loss of generalization, assume that  $n \geq k$ . If  $n < k$  in practice, we can pad  $M_l^i$  using columns of zeros. The  $k$ -largest node selection is conducted on  $M_l^i$ ; that is, for each column, we rank the  $n$  values and select  $k$ -largest values. This gives us a  $k \times C$  output matrix. As the columns in  $M_l^i$  represent features, the operation is equivalent to selecting  $k$ -largest values for each feature. By inserting  $x_l^i$  in the first row, the output becomes  $\tilde{M}_l^i \in \mathbb{R}^{(k+1) \times C}$ . This is illustrated in the left part of Figure 2. By repeating this process for each node,  $g(\cdot)$  transforms  $X_l$  to  $\tilde{X}_l \in \mathbb{R}^{N \times (k+1) \times C}$ .

Note that  $\tilde{X}_l$  can be viewed as a 1-D grid-like structure by considering  $N$ ,  $(k+1)$ , and  $C$  as the batch size, the spatial size, and the number of channels, respectively. Therefore, the  $k$ -largest node selection function  $g(\cdot)$  successfully achieves the transformation from generic graphs to grid-like data. The operation makes use of the natural ranking information among real numbers and forces each node to have a fixed number of ordered neighbors.

**1-D Convolutional Neural Networks.** As discussed in Section 3.1, regular convolutional operations can be directly applied on grid-like data. As  $\tilde{X}_l \in \mathbb{R}^{N \times (k+1) \times C}$  is 1-D, we employ a 1-D CNN model  $c(\cdot)$ . The basic functionality of LGCL is to aggregate adjacent information and update the feature vector for each node. Consequently, it requires  $X_{l+1} \in \mathbb{R}^{N \times D}$ , where  $D$  is the dimension of the updated feature space. The 1-D CNN  $c(\cdot)$  should take  $\tilde{X}_l \in \mathbb{R}^{N \times (k+1) \times C}$  as input and output a matrix of dimension  $N \times D$ , or equivalently,  $N \times 1 \times D$ . Basically,  $c(\cdot)$  reduces the spatial size from  $(k+1)$  to 1.

Note that  $N$  is considered as the batch size, which is not related to the design of  $c(\cdot)$ . As a result, we focus on only one data sample, i.e., one node in the graph. Taking the example above, for node  $i$ , the transformed output is  $\tilde{M}_l^i \in \mathbb{R}^{(k+1) \times C}$ , which serves as the input to  $c(\cdot)$ . Due to the fact that any regular convolutional operation with a filter size larger than one and no padding reduces the spatial size, the simplest  $c(\cdot)$  has only one convolutional layer with a filter size of  $(k+1)$  and no padding. The numbers of input and output channels are  $C$  and  $D$ , respectively. Meanwhile, any multi-layer CNN can be employed, provided its final output has the dimension of  $1 \times D$ . The right part of Figure 2 illustrates an example of a two-layer CNN. Again, applying  $c(\cdot)$  for all the  $N$  nodes outputs  $X_{l+1} \in \mathbb{R}^{N \times D}$ . In summary, our LGCL transforms generic graphs to grid-like data using the proposed  $k$ -largest node selection and applies a regular 1-D CNN to perform feature aggregation and refine the feature vector for each node.

**Algorithm 1** Sub-Graph Selection Algorithm

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**Input:** Adjacency matrix  $A$ , Number of nodes  $N$ , Sub-graph size  $N_s$ , Initial number of nodes  $N_{init}$ , Maximum number of nodes expanded per iteration  $N_m$

**Output:** A set of nodes  $S$  as a sub-graph

```

1:  $S = \phi$ 
2:  $initNodes = \text{sample } N_{init} \text{ nodes from } N \text{ nodes.}$ 
3:  $S = S \cup initNodes$ 
4:  $newAddNodes = initNodes$ 
5: while  $\text{size}(S) < N_s$  and  $\text{size}(newAddNodes) \neq 0$  do
6:    $candidateNodes = \text{BFS}(newAddNodes, A)$ 
    $\triangleright$  Obtain first-order neighboring nodes of  $newAddNodes$ 
7:    $newAddNodes = candidateNodes \setminus S$ 
8:   if  $\text{size}(newAddNodes) > N_m$  then
9:      $newAddNodes = \text{sample } N_m \text{ nodes from } newAddNodes$ 
10:  end if
11:  if  $\text{size}(newAddNodes) + \text{size}(S) > N_s$  then
12:     $N_r = N_s - \text{size}(S)$ 
13:     $newAddNodes = \text{sample } N_r \text{ nodes from } newAddNodes$ 
14:  end if
15:   $S = S \cup newAddNodes$ 
16: end while
17: return  $S$ 

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### 3.3 Learnable Graph Convolutional Networks

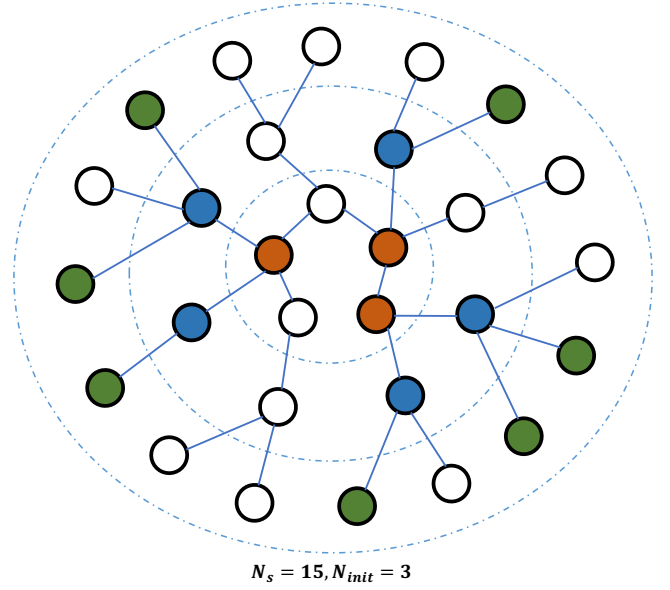
It is known that deeper networks usually yield better performance. However, prior deep models on graphs like GCNs only have two layers. While they suffer from performance loss when going deeper [15], our LGCL enables a deeper design, resulting in the learnable graph convolutional networks (LGCNs) for graph node classification. We build LGCNs based on the architecture of densely connected convolutional networks (DCNNs) [12, 13], which achieved state-of-the-art performance in the ImageNet classification challenge [16].

In LGCNs, we first apply a graph embedding layer to produce low-dimensional representations of nodes, since the original inputs are usually very high-dimensional feature vectors in some graph dataset, such as the Cora [23]. The graph embedding layer is essentially a linear transformation in the first layer expressed as

$$X_1 = X_0 W_0, \quad (4)$$

where  $X_0 \in \mathbb{R}^{N \times C_0}$  represents the high-dimensional input and  $W_0 \in \mathbb{R}^{C_0 \times C_1}$  changes the dimension of feature space from  $C_0$  to  $C_1$ . As a result,  $X_1 \in \mathbb{R}^{N \times C_1}$  and  $C_1 < C_0$ . Alternatively, a GCN layer can be used for graph embedding. As illustrated in Section 2, the number of training parameters in a GCN layer is equal to that of a regular graph embedding layer.

After the graph embedding layer, we stack multiple LGCLs, according to the complexity of the graph data. As each LGCL only aggregates information from first-order neighboring nodes, *i.e.*, direct neighboring nodes, stacked LGCLs can collect information from a larger set of nodes, which is commonly done in regular CNNs. In order to promote the model performance and facilitate the training process, we apply skip connections to concatenate the inputs and outputs of LGCLs. Finally, a fully-connected layer is used before the softmax function for final predictions.



**Figure 4: An example of the sub-graph selection process.** We start with  $N_{init} = 3$  randomly sampled nodes and obtain a sub-graph of  $N_s = 15$  nodes. In the first iteration, we use BFS to find all the first-order neighboring nodes of the 3 initial nodes (orange), excluding themselves. Among these nodes, we randomly select  $N_m = 5$  nodes (blue). In the next iteration, we select  $N_m = 7$  nodes from neighbors of the blue nodes, excluding previously selected nodes. Note that  $N_m$  changes for the two iterations, which is a flexible choice in practice. After two iterations, we have selected  $3 + 5 + 7 = 15$  nodes and obtained a required sub-graph. These nodes, along with the corresponding adjacency matrix, will form the input to the LGCL in a training iteration.

Following the design principle of LGCNs,  $k$  and the number of stacked LGCLs are the most important hyper-parameters. The average degree of nodes in the graph can be a good reference for selecting  $k$ . Meanwhile, the number of LGCLs should depend on the complexity of tasks, such as the number of classes, the number of nodes in a graph, etc. More complicated tasks require deeper models.

### 3.4 Sub-Graph Training on Large-Scale Data

Most prior deep models on graphs suffer from another limitation. In particular, during training the inputs are the feature vectors of all the nodes along with the adjacency matrix of the whole graph, whose sizes become large for large graph data. These prior models work properly on small-scale graphs. However, for large-scale graphs, those methods usually result in excessive memory and computational resource requirements, which limit the practical applications of these models.

Similar problems also happen for deep neural networks on other types of data, such as grid-like data. For example, deep models on image segmentation usually use randomly cropped patches when dealing with large images. Motivated by this strategy, we intend

**Table 1: Summary of datasets used in our experiments [30, 31]. The Cora, Citeseer, and Pubmed datasets are used for transductive learning experiments, while the PPI dataset is for inductive learning experiments. The degree attribute listed is the average node degree of each dataset, which helps the selection of the hyper-parameter  $k$  in LGCLs.**

Dataset	#Nodes	#Features	#Classes	#Training Nodes	#Validation Nodes	#Test Nodes	Degree	Setting
Cora	2708	1433	7	140	500	1000	4	Transductive
Citeseer	3327	3703	6	120	500	1000	5	Transductive
Pubmed	19717	500	3	60	500	1000	6	Transductive
PPI	56944	50	121	44906 (20 graphs)	6514 (2 graphs)	5524 (2 graphs)	31	Inductive

to randomly “crop” a graph to obtain smaller graphs for training. However, while a rectangular patch of an image naturally maintains neighboring information among pixels, how to handle irregular connections between nodes in a graph remains challenging.

In this work, we propose a sub-graph selection algorithm to address the memory and computational resource problems on large-scale graph data, as shown in Algorithm 1. Given a graph, we first sample some initial nodes. Starting from them, we use the Breadth-First-Search (BFS) algorithm to expand adjacent nodes into the sub-graph iteratively. With multiple iterations, high-order neighboring nodes of the initial nodes are included. Note that we use a single parameter  $N_m$  in Algorithm 1 for simplicity. In practice, we can set  $N_m$  to different values for each iteration. Figure 4 provides an example of the sub-graph selection process.

With such randomly “cropped” sub-graphs, we are able to train deep models on large-scale graphs. In addition, we can take advantage of the mini-batch training strategy to accelerate the learning process. In each training iteration, we can use the proposed sub-graph selection algorithm to sample several sub-graphs and put them in a mini-batch. The corresponding feature vectors and adjacency matrices form the inputs to the networks.

## 4 EXPERIMENTAL STUDIES

In this section, we evaluate our proposed large-scale learnable graph convolutional networks (LGCNs) on node classification tasks under both transductive and inductive learning settings. In addition to comparisons with prior state-of-the-art models, some performance studies are performed to investigate how to choose hyper-parameters. Experiments are also conducted to analyze the training strategy based on the proposed sub-graph selection algorithm. Experimental results show that LGCNs yield improved performance, and the sub-graph training is much more efficient than whole-graph training. Our code is publicly available<sup>1</sup>.

### 4.1 Datasets

In our experiments, we focus on node classification tasks under both transductive and inductive learning settings.

**Transduction Learning.** Under the transductive setting, the unlabeled testing data are accessible and available during training. To be specific, for node classification, only a part of nodes in the graph are labeled. The testing nodes, which are also in the same graph, are accessible during training, including their features and connections, except for the labels. This means the training process knows about the graph structure that contains testing nodes. We

use three standard benchmark datasets for transductive learning experiments; those are the Cora, Citeseer, and Pubmed [23], as summarized in Table 1. These three datasets are citation networks with nodes and edges representing documents and citations, respectively. The feature vector of each node corresponds to a bag-of-words representation for a document. For these three datasets, we employ the same experimental settings as those in GCN [15]. For each class, 20 nodes are used for training, 500 nodes are used for validation and 1,000 nodes are used for testing.

**Inductive Learning.** For inductive learning, the testing data are not available during training, which means the training process does not learn about the structure of test graphs. In inductive learning tasks, we usually have different training, validation, and testing graphs. During training, the model only use the training graphs without access to validation and testing graphs. We use the protein-protein interaction (PPI) dataset [31], which contains 20 graphs for training, 2 graphs for validation, and 2 graphs for testing. Since the graphs for validation and testing are separate, the training process does not use them. There are 2,372 nodes in each graph on average. Each node has 50 features including positional, motif genes and signatures. Each node has multiple labels from 121 classes.

### 4.2 Experimental Setup

We describe the experimental setup under both transductive and inductive learning settings.

**Transduction Learning.** In transductive learning tasks, we employ the proposed LGCN models as illustrated in Figure 3. Since transductive learning datasets employ high-dimensional bag-of-words representations as feature vectors of nodes, the inputs go through a graph embedding layer to reduce the dimension. Here, we use a GCN layer as the graph embedding layer. The dimension of the embedding output is 32. Then we apply LGCLs, each of which uses  $k = 8$  and produces 8-component feature vectors. For the Cora, Citeseer, and Pubmed, we stack 2, 1, and 1 LGCLs, respectively. We use concatenation in skip connections. Finally, a fully-connected layer is used as a classifier to make predictions. Before the fully-connected layer, we perform a simple sum to aggregate feature vectors of adjacent nodes. Dropout [25] is applied on both input feature vectors and adjacency matrices in each layer with rates of 0.16 and 0.999, respectively. All LGCN models in transductive learning tasks use the sub-graph training strategy. The sub-graph size is set to 2,000.

**Inductive Learning.** For inductive learning, the same LGCN model as above is used except for some hyper-parameters. For the graph embedding layer, the dimension of output feature vectors

<sup>1</sup><https://github.com/divelab/lgcnn/>



**Table 2: Results of transductive learning experiments in terms of node classification accuracies on the Cora, Citeseer, and Pubmed datasets.  $\text{LGCN}_{sub}$  denotes the LGCN model using the sub-graph training strategy.**

Models	Cora	Citeseer	Pubmed
DeepWalk [21]	67.2%	43.2%	65.3%
Planetoid [30]	75.7%	64.7%	77.2%
Chebyshev [4]	81.2%	69.8%	74.4%
GCN [15]	81.5%	70.3%	79.0%
$\text{LGCN}_{sub}$ (Ours)	<b><math>83.3 \pm 0.5\%</math></b>	<b><math>73.0 \pm 0.6\%</math></b>	<b><math>79.5 \pm 0.2\%</math></b>

**Table 3: Results of inductive learning experiments in terms of micro-averaged F1 scores on the PPI dataset.**

Models	PPI
GraphSAGE-GCN [9]	0.500
GraphSAGE-mean [9]	0.598
GraphSAGE-pool [9]	0.600
GraphSAGE-LSTM [9]	0.612
$\text{LGCN}_{sub}$ (Ours)	<b><math>0.772 \pm 0.002</math></b>

is 128. We stack two LGCLs with  $k = 64$ . We also employ the sub-graph training strategy, with sub-graph initial node size equal to 500 and 200. Dropout with a rate of 0.9 is applied in each layer.

For both transductive and inductive learning LGCN models, the following configurations are shared. For all layers, only the identity activation function is used, which means no nonlinearity is involved in the networks. In order to avoid over-fitting, the  $L_2$  regularization with  $\lambda = 0.0005$  is applied. For training, the Adam optimizer [14] with a learning rate of 0.1 is used. Weights in LGCNs are initialized by the Glorot initialization [7]. We employ the early stopping strategy based on the validation accuracy and train 1,000 epochs at most.

### 4.3 Analysis of Results

The experimental results are summarized in Tables 2 and 3 for transductive and learning settings, respectively.

**Transduction Learning.** For transductive learning experiments, we report node classification accuracies as in [15]. Table 2 provides the comparisons with other graph models. According to the results, our LGCN models achieve better performance over the current state-of-the-art GCNs by a margin of 1.8%, 2.7%, and 0.6% on the Cora, Citeseer, and Pubmed datasets, respectively.

**Inductive Learning.** For inductive learning experiments, we report micro-averaged F1 scores like [9]. From table 3, we can observe that our LGCN model outperforms GraphSAGE-LSTM by a margin of 16%. Without observing the structure of test graphs in training, the LGCN model still achieves good generalization.

The results above show that the proposed LGCN models on generic graphs consistently yield new state-of-the-art performance in node classification tasks on different datasets. These results demonstrate the effectiveness of applying regular convolutional operations on transformed graph data. In addition, the proposed

**Table 4: Results of transductive learning experiments for comparing the  $\text{LGCN}_{sub}$  and GCN layers on the Cora, Citeseer, and Pubmed datasets. Using the network architecture of  $\text{LGCN}_{sub}$ , we replace LGCLs by GCN layers, resulting in the  $\text{LGCN}_{sub}$ -GCN model.**

Models	Cora	Citeseer	Pubmed
$\text{LGCN}_{sub}$ -GCN	$82.2 \pm 0.5\%$	$71.1 \pm 0.5\%$	$79.0 \pm 0.2\%$
$\text{LGCN}_{sub}$ (Ours)	<b><math>83.3 \pm 0.5\%</math></b>	<b><math>73.0 \pm 0.6\%</math></b>	<b><math>79.5 \pm 0.2\%</math></b>

transformation approach through the  $k$ -largest node selection is shown to be effective.

### 4.4 LGCL versus GCN Layers

It may be argued that our LGCN models employ a deeper network architecture than GCNs, which could explain the improved performance. However, the performance of GCNs is reported to decrease when going deeper by stacking more layers. In addition, we conduct another experiment by replacing all LGCLs in LGCN models by GCN layers, denoted as  $\text{LGCN}_{sub}$ -GCN model. All the other settings remain the same in order to ensure the fairness of the comparisons. Table 4 provides the comparison results between  $\text{LGCN}_{sub}$  and  $\text{LGCN}_{sub}$ -GCN. The results show that  $\text{LGCN}_{sub}$  has better performance than  $\text{LGCN}_{sub}$ -GCN, which indicates that the LGCL is more effective than the GCN layer.

### 4.5 Sub-Graph versus Whole-Graph Training

For the experiments above, we use the sub-graph training strategy to learn the LGCN models, which aims at saving memory and training time. However, since the sub-graph selection algorithm samples some nodes as a sub-graph from the whole graph, it means that the models trained in this way do not learn about the structure of whole graph during training. Meanwhile, in transductive learning tasks, the information of testing nodes may be ignored, which raises the risk of performance loss. To address this concern, we perform experiments on transductive learning tasks to compare the sub-graph training strategy with the previous whole-graph training strategy. Through the experiments, we show the advantages of using the sub-graph training strategy, with negligible loss in terms of model performance.

For the sub-graph selection process described in Algorithm 1, the algorithm starts with some initial nodes that are randomly selected. In transductive learning tasks, we sample initial nodes only from the nodes with training labels to make sure that training can be conducted. To be specific, we sample 140, 120, and 60 initial nodes when selecting the sub-graph for the Cora, Citeseer, and Pubmed datasets, respectively. For each iteration in the sub-graph selection algorithm, we do not set  $N_m$  to limit the number of nodes expanded into the sub-graph. The maximum number of nodes in the sub-graph is set to 2,000 for all the three datasets, which is an feasible size for our GPUs in hand.

For comparison, we perform experiments using the same LGCN models, but train them using the same whole-graph training strategy as GCNs, which means the inputs are representations of the entire graph. We denote such models as  $\text{LGCN}_{whole}$ , compared

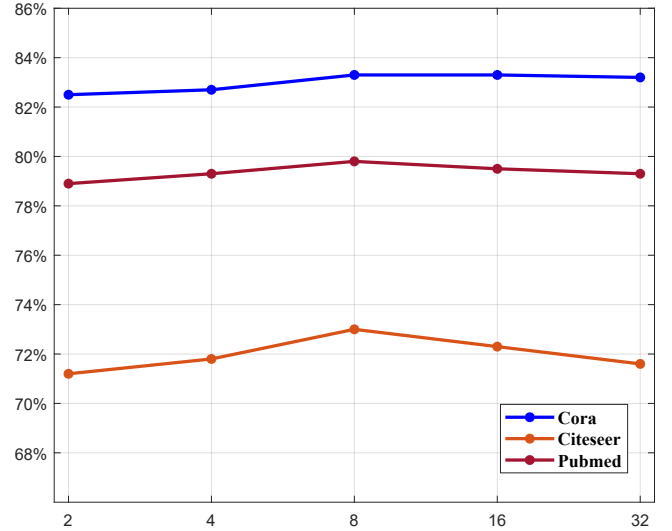
**Table 5: Results of transductive learning experiments for comparing the sub-graph training and whole-graph training strategies on the Cora, Citeseer, and Pubmed datasets. For comparison, we conduct experiments on LGCNs that employ the same whole-graph training strategy as GCNs, denoted as  $\text{LGCN}_{\text{whole}}$ .**

		<b>Cora</b>	<b>Citeseer</b>	<b>Pubmed</b>
<b>GCN</b>	# Nodes	2708	3327	19717
	Accuracy	81.5%	70.3%	79.0%
	Time	<b>7s</b>	4s	38s
<b><math>\text{LGCN}_{\text{whole}}</math></b>	# Nodes	2708	3327	19717
	Accuracy	$83.8 \pm 0.5\%$	$73.0 \pm 0.6\%$	$79.5 \pm 0.2\%$
	Time	58s	30s	1080s
<b><math>\text{LGCN}_{\text{sub}}</math></b>	# Nodes	644	442	354
	Accuracy	$83.3 \pm 0.5\%$	$73.0 \pm 0.6\%$	$79.5 \pm 0.2\%$
	Time	14s	<b>3.6s</b>	<b>2.6s</b>

to  $\text{LGCN}_{\text{sub}}$  with the sub-graph training strategy. The comparing results of these two models with GCNs are provided in Table 5. The number of nodes reported represents how many nodes are used for one iteration of training. The time reported here is the training time for running 100 epochs using a single TITAN Xp GPU.

It can be seen that the actual numbers of nodes in the training sub-graph for the Cora, Citeseer, and Pubmed datasets are 644, 442, and 354, respectively, which are far smaller than the maximum sub-graph size of 2,000. This indicates that the nodes in the Cora, Citeseer, and Pubmed datasets are sparsely connected. Specifically, starting from several initial nodes with training labels, only a small set of nodes will be selected by expanding neighboring nodes to form connected sub-graphs. While these datasets are usually considered as a single large graph, the whole graph is actually composed of several separate sub-graphs that have no connection to each other. The sub-graph training strategy takes advantage of this fact and makes efficient use of the nodes with training labels. Since only the initial nodes have training labels and all their connectivity information is included in the selected sub-graphs, the amount of information loss in the sub-graph training is minimized, resulting in negligible performance loss. This is demonstrated by comparing the node classification accuracies of  $\text{LGCN}_{\text{sub}}$  and  $\text{LGCN}_{\text{whole}}$ . According to the results,  $\text{LGCN}_{\text{sub}}$  models only have a subtle performance loss of 0.5% on the Cora dataset, while yielding the same performance on the Citeseer and Pubmed datasets, as compared to the  $\text{LGCN}_{\text{whole}}$  models.

After investigating the risk of performance loss, we point out the great advantages of the sub-graph training strategy in terms of training speed. By using the sub-graph training,  $\text{LGCN}_{\text{sub}}$  models take a sub-graph of fewer nodes as inputs in contrast to the whole graph, which is expected to greatly promote the training efficiency. It can be seen from the results in Table 5 that the improvement is outstanding. Although GCNs require simpler computation, its running time is much longer than that of LGCN models on large-scale graph datasets like the Pubmed. Powerful deep models are usually used on large-scale data, which makes the sub-graph training strategy useful in practice. The sub-graph training strategy enables



**Figure 5: Results of using different values of hyper-parameter  $k$  in LGCN models. On the Cora, Citeseer, and Pubmed datasets, we employ the same experimental setups described in Section 4.2. We adjust the value of  $k$  in  $\text{LGCN}_{\text{sub}}$  and report node classification accuracies in this figure. It can be seen that  $k = 8$  achieves the best performance for these datasets.**

using more complex layers such as the proposed LGCLs without the concern of long training time. As a result, our large-scale LGCNs with the sub-graph training strategy are not only effective but also very efficient.

#### 4.6 Performance Study of $k$

As described in Section 3.3, the average degree of nodes in graph can be helpful when choosing the hyper-parameter  $k$  in LGCNs. In this part, we conduct experiments to show how different values of  $k$  affect the performance of LGCN models. We vary the value of  $k$  in LGCLs and observe the node classification accuracies on the Cora, Citeseer, and Pubmed datasets. The values of  $k$  are selected from 2, 4, 8, 16, and 32, which cover a reasonable range of integer values.

Figure 5 plots the performance change of LGCN models under different values of  $k$ . As demonstrated in the figure, the LGCN models achieve the best performance on all the three datasets when choosing  $k = 8$ . In the Cora, Citeseer, and Pubmed datasets, the average node degrees are 4, 5, and 6, respectively. This indicates that the best  $k$  is usually a bit larger than the average node degree in the dataset. When  $k$  is too large, the performance of LGCN models decreases. A possible explanation is that if  $k$  is much larger than the average node degree in the graph, too many zero padding is used in the  $k$ -largest node selection process, which compromises the performance of the following 1-D CNN models. For the inductive learning task on the PPI dataset, we also explore different values of  $k$ . The best performance is given by  $k = 64$  while the average node degree is 31. This is consistent with our results above.



## 5 CONCLUSIONS AND FUTURE WORK

In this work, we propose the learnable graph convolutional layer (LGCL), which transforms generic graphs to data of grid-like structures and enables the use of regular convolutional operations. The transformation is conducted through a novel  $k$ -largest node selection process, which uses the ranking between node feature values. Based on our LGCL, we build deeper networks, known as learnable graph convolutional networks (LGCNs), for node classification tasks on graphs. Experimental results show that the proposed LGCN models yield consistently better performance than prior methods under both transductive and inductive learning settings. Our LGCN models achieve new state-of-the-art results on four different datasets, demonstrating the effectiveness of LGCLs.

In addition, we propose a sub-graph selection algorithm, resulting in the sub-graph training strategy, which can solve the problem of excessive requirements for memory and computational resources on large-scale graph data. With the sub-graph training, the proposed LGCN models are both effective and efficient. Our experiments indicate that the sub-graph training strategy brings a significant advantage in terms of training speed, with a negligible amount of performance loss. The new training strategy is very useful as it enables the use of more complex models efficiently.

Based on this work, we discuss several possible directions for future work. First, our methods mainly address the node classification problems. In practice, many other interesting tasks can be formulated as graph classification problems, where each graph has a label. While they are similar to image classification tasks, current graph convolutional methods, including ours, are not able to perform down-sampling on graphs, like the pooling operations on image data. We need a layer to reduce the number of nodes effectively, which is necessary for graph classification. Second, our methods are mainly applied to generic graph data like citation networks. For other data like text, our methods may also be helpful, since we can treat text data as graphs. We will explore these directions in the future.

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