Toward Graph Self-Supervised Learning With Contrastive Adjusted Zooming

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Abstract—Graph representation learning (GRL) is critical for graph-structured data analysis. However, most of the existing graph neural networks (GNNs) heavily rely on labeling information, which is normally expensive to obtain in the real world. Although some existing works aim to effectively learn graph representations in an unsupervised manner, they suffer from certain limitations, such as heavy reliance on monotone contrastiveness and limited scalability. To overcome the aforementioned problems, in light of the recent advancements in graph contrastive learning, we introduce a novel self-supervised GRL algorithm via graph contrastive adjusted zooming, namely, G-Zoom, to learn node representations by leveraging the proposed adjusted zooming scheme. Specifically, this mechanism enables G-Zoom to explore and extract self-supervision signals from a graph from multiple scales: micro (i.e., node level), meso (i.e., neighborhood level), and macro (i.e., subgraph level). First, we generate two augmented views of the input graph via two different graph augmentations. Then, we establish three different contrastiveness on the above three scales progressively, from node, neighboring, to subgraph level, where we maximize the agreement between graph representations across scales. While we can extract valuable clues from a given graph on the micro and macro perspectives, the neighboring-level contrastiveness offers G-Zoom the capability of a customizable option based on our adjusted zooming scheme to manually choose an optimal viewpoint that lies between the micro and macro perspectives to better understand the graph data. In addition, to make our model scalable to large graphs, we use a parallel graph diffusion approach to decouple model training from the graph size. We have conducted extensive experiments on real-world datasets, and the results demonstrate that our proposed model outperforms the state-of-the-art methods consistently.

Index Terms—Contrastive learning, graph neural networks (GNNs), graph representation learning (GRL), self-supervised learning.

I. INTRODUCTION

GRAPH representation learning (GRL) has become a pivotal strategy for analyzing semi-structured graph data in recent years. GRL aims to distill the high-dimensional structural and attributive information from graphs to generate low-dimensional embeddings for nodes or graphs. These learned embeddings can then be used in various downstream tasks such as node classification [1], [2], graph classification [3], and link prediction [4]. It has been applied to many real-world graph datasets from different domains [5] such as social networks [1], geoscience [6], anomaly detection [7], [8] and molecules [9].

Recently, the existing successful GRL approaches are mainly powered by graph neural networks (GNNs) [1], [10], [11], which learn low-dimensional node embeddings via iterative message passing to aggregate topological representations of neighbors. However, most of these methods adopt supervised learning and rely extensively on labeling information, which is both laborious and expensive to collect in the real world. To address this issue, some self-supervised GRL methods based on contrastive learning emerged, e.g., deep graph infomax (DGI) [12], GRACE [13], and graphical mutual information (GMI) [14]. The basic idea of these approaches is to set up pretext tasks without using labeling information to train a GNN encoder for node embeddings’ generation. While DGI [12] focuses on maximizing the mutual information (MI) between node- and graph-level representations, GRACE [13] and GMI [14] extend this MI maximization scheme to contrast node-level representations in two different graph views or enlarge the MI between the hidden representation of nodes and the raw node features of their one-hop neighbors.

Though these methods have achieved promising results, they adopt monotonous contrastive learning on graphs, which only inspects graph-structured data from a single fixed perspective. For example, DGI and MVGRL only conduct contrasting between patch and global representations, i.e., contrasting from a global viewpoint. Without considering the contrastiveness from a macro perspective, other methods such as GRACE, sub-graph contrast (SubG-Con) [15], and GMI only emphasize the local structure of target nodes by maximizing the agreement between patch representations in two augmented views, or between nodes and representations of their neigh-
we can see the object clearly (e.g., dog head). Similarly, in a graph, we can choose a target node and then decide the scope of its contextual information we want to examine.

In general, it is straightforward to find that different perspectives offer different scenes and contextual information. While from the macro perspective, we zoom out observing the entire picture and ignore the detailed information, from the micro perspective, we concentrate on the microscopic element of the data regardless of the broader view. Different from other two perspectives, the meso perspective offers a more flexible viewpoint, enabling us to inspect a specific area with the desired scale. This perspective forces the model to explore the fine-grained information embodied in this scale, which can easily be neglected from the macro-level viewpoint. As such, it complements the macro perspective with overlooked detailed information in the semi-global context. Therefore, it is necessary to include various perspectives on multiple scales when conducting graph contrastive learning.

In G-Zoom, we propose to formulate three different contrastive paths on various scales progressively: node, neighboring, and subgraph levels. Inspired by GRACE and MVGRL, we establish node- and subgraph-level contrastiveness to facilitate discrimination from the micro and macro perspectives. Then, with our proposed adjusted zooming scheme, we design the neighboring-level discrimination paths, which provide a tailored view of contrastiveness. Specifically, two graph views are first sampled with different graph augmentations [e.g., graph sampling (GS) and diffusion]. Then, we input two augmented views into a GNN encoder to get node representations in sampled graphs. At this stage, we can readout subgraph-level representations via graph pooling and obtain neighboring-level representations by aggregating the top-k neighborhood representations of selected central nodes. Finally, we formulate our training loss by summing up the contrastive losses on three different contrasting levels to train the graph encoder for node embedding calculation. Compared with prior arts, G-Zoom not only facilitates multiscale contrasting to provide learnable clues from multiple perspectives for model training but also can be extended to handle large-sized graphs. The experimental results on a variety of datasets have shown the superiority of G-Zoom compared with several state-of-the-art baselines. The main contributions of our work are listed as below.

1) We propose a novel algorithm, namely, G-Zoom, to learn graph representation in an unsupervised manner. To the best of our knowledge, this is the first attempt to conduct graph contrastive learning on multiple scales that crossed three different graph topological levels.

2) We propose the graph adjusted zooming mechanism, which aims to facilitate multiscale graph contrastive learning from various perspectives to enrich the self-supervision signals and overcome the limitations of the existing works.

3) We conduct extensive experiments on real-world datasets to validate the effectiveness of our proposed approach. The results show that our model consistently outperforms the state-of-the-art baselines.
II. RELATED WORK

In this section, we review related works in three areas: GNNs, unsupervised graph representation learning (GRL), and contrastive self-supervised learning.

A. Graph Neural Networks

Under the umbrella of deep neural network [17], GNNs learn node embeddings using both attributive and topological information of the non-Euclidean graph-structured data. GNN is first introduced in [18], which proposes a spectral-based method extending convolution networks to graphs. Then, a series of subsequent spectral-based convolution GNNs have been introduced [19], [20], which adopt filters in light of graph signal processing [21]. Notably, bridging the gap between the spectral-based and spatial-based GNN methods, GCN [1] simplifies graph convolutions by approximating spectral-based graph convolutions with stacked first-order Chebyshev polynomial filters. After that, the spatial-based approaches grow promptly because of their advantages of efficiency and general applicability, e.g., GAT [10] incorporates the attention mechanism [22] to consider the difference of importance among node neighbors rather than simply averaging neighbors’ information. SGC [11] reduces the excess complexity of GCN via removing nonlinearity and collapsing weight matrices between graph convolution layers. Except for SGC, many studies are aiming to improve GNN from different perspectives such as scalability extension [23], [24] and receptive field enlargement [25], [26], [27], [28]. The existing GNN approaches have been successfully applied in various domains, e.g., bankruptcy prediction [8], knowledge graph [29], [30], and traffic prediction [31].

Though there are many off-the-shelf GNNs available, many of them rely extensively on labeling information, which are unable to deal with unlabeled graphs. However, unlabeled graphs are pervasive in the real world. To tackle this problem, we proposed G-Zoom, which uses an adjusted-zooming-based contrastive learning scheme to generate effective node embeddings without the guidance of node labels. In G-Zoom, we select GCN as the kernel of our GNN encoder component. Except for GCN, the encoder component of G-Zoom can be underpinned by any aforementioned GNNs.

B. Unsupervised GRL

Unsupervised GRL aims to learn node embeddings on unlabeled graphs, in which node labels are unavailable. Mainly, there are two traditional ways for unsupervised algorithms to exploit the underlying topological information. One way is adopting an auto-encoder, which consists of an encoder for embedding a graph into latent representation and a decoder for reconstructing the topology of the graph [32], [33]. These autoencoder-based methods heavily rely on the assumption that neighboring nodes have similar representations. The effectiveness of objectives based on this assumption is questionable [15]. Another way conducts model optimization using random-walk (RW)-based objectives [34], [35], [36]. However, these methods are criticized because of having one or multiple issues including overwhelmingly emphasizing structural information embodied in graph proximity, limited scalability, and the inability of learning attributive information [13], [37].

C. Contrastive Self-Supervised Learning

Recently, there are some successful applications of using self-supervised contrastive learning approaches for image processing [38], [39], [40], [41], [42] which have achieved promising results. With the help of defined annotation-free pretext tasks, these methods train models via contrasting positive and negative instance pairs to calculate the contrastiveness loss. Some recent works also attempt to adapt contrastive methods to the graph domain. Inspired by deep infomax [40], DGI [12] pioneers contrastive learning for graphs. Specifically, DGI considers node- and graph-level representations obtained with a readout function as positive pairs and uses graph corruption techniques to conduct negative sampling. On the basis of DGI, multiview GRL (MVGRL) [37] consolidates the patch summary contrastive learning approach of DGI by integrating graph augmentation techniques including GS and graph diffusion to generate multiple augmented graph views. GRACE [13] focuses on node-level contrasting between two augmented views, which are built through attributive and structural modification on the input graph. Graph mutual information (GMI) and SubG-Con aim to encode localized signals by contrasting the central node representation with the representations of their close neighbors.

Although these graph self-supervised approaches fulfill comparable performance to supervised methods, they are still suffering from issues such as monotonous contrastive learning schemes and limited scalability. To be specific, DGI, MVGRL, GRACE, and GMI fail to handle large-scale graphs, and all the aforementioned methods are incapable of encoding information of a graph thoroughly and can only extract information from a single perspective, i.e., micro, macro, or meso. To address these issues, we proposed an adjusted zooming contrastive learning mechanism for comprehensive multiscale contrasting in G-Zoom. Also, G-Zoom can be extended to handle large-scale graphs. The detailed explanation of this extension is shown in Section V-B. Extensive experiments in the following sections have shown the superiority of G-Zoom in GRL.

III. PROBLEM FORMULATION

In this section, we introduce the problem of unsupervised node representation learning. The notations used in this article are summarized in Table I. In this article, we use bold uppercase letters (e.g., $X$) and bold lowercase letters (e.g., $x$) to represent matrices and vectors, respectively.

Let $G = (X, A)$ denote a graph with a node feature matrix $X \in \mathbb{R}^{n \times d}$ and an adjacency matrix $A \in \mathbb{R}^{n \times n}$, in which the value at the $i$th row in the $j$th column of $A$ is 1 if nodes $v_i$ and $v_j$ are connected. Here, $n$ and $d$ denote the number and dimension of nodes in $G$, respectively. In G-Zoom, we leverage a GNN as the backbone encoder to transform the original high-dimensional features into a low-dimensional representation, which has been defined below.

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TABLE I

| Symbols   | Description                                                                 |
|-----------|-----------------------------------------------------------------------------|
| \( \mathcal{G} = (X, A) \)        | A graph with feature matrix                                                  |
| \( X \in \mathbb{R}^{n \times d} \) | The feature matrix of \( \mathcal{G} \)                                  |
| \( A \in \mathbb{R}^{n \times n} \) | The adjacency matrix of \( \mathcal{G} \)                                 |
| \( n \)   | Number of nodes in \( \mathcal{G} \)                                       |
| \( d \)   | Number of dimensions of \( X \)                                             |
| \( d' \)  | Number of dimensions of latent representations                              |
| \( \tilde{G} = (X, \tilde{A}) \) | The diffused version of the input graph \( \mathcal{G} \)                   |
| \( B \)   | Number of nodes in a batch                                                  |
| \( P \)   | Number of nodes in a \( \tilde{G} \)                                       |
| \( v_t \) | The target node                                                             |
| \( k \)   | Number of top-ranked neighbors for \( v_t \)                                |
| \( \mathcal{G}_1, \mathcal{G}_2 \) | The augmented view of \( \mathcal{G} \)                                    |
| \( g_\theta \) | The GNN encoder used in our model                                          |
| \( H_1, H_2 \in \mathbb{R}^{n \times d'} \) | The latent representation of \( \mathcal{G}_1 \) and \( \mathcal{G}_2 \) encoded via \( g_\theta \) |
| \( \tilde{H} \in \mathbb{R}^{n \times d'} \) | Corrupted representation for \( \mathcal{G} \)                             |
| \( \tilde{H} \in \mathbb{R}^{n \times d'} \) | Final embedding matrix for \( \mathcal{G} \)                              |
| \( h_1, h_2 \in \mathbb{R}^{d'} \) | Node-level representation of \( v_t \) in \( H_1, H_2 \)                  |
| \( s_1, s_2 \in \mathbb{R}^{d'} \) | Subgraph-level representation for \( \mathcal{G}_1, \mathcal{G}_2 \)         |
| \( n_1, n_2 \in \mathbb{R}^{d'} \) | Neighboring-level representation of \( v_t \) in \( H_1, H_2 \)             |
| \( \mathcal{D} \) | Discriminator for mutual information evaluation                            |
| \( W^l \) | The \( l \)-th layer trainable weight matrix in \( g_\theta \). If \( l = 1 \), \( W^l \in \mathbb{R}^{d \times d'} \), while \( l > 1 \), \( W^l \in \mathbb{R}^{d' \times d'} \). |
| \( W_D \in \mathbb{R}^{d' \times d'} \) | The trainable weight matrix in \( \mathcal{D} \)                            |

Definition 1 (Graph Neural Networks): Given an attributed graph \( \mathcal{G} = (X, A) \), a typical GNN mainly consists of two components: message aggregation and transformation

\[
\begin{align*}
    m_G^{(l)}[v_t; :) &= \text{aggregate}^{(l)}(H_G^{(l-1)}[v_k; :) : v_k \in \mathcal{N}(v_t)) \\
    H_G^{(l)}[v_t; :) &= \text{transform}^{(l)}(H_G^{(l-1)}[v_t; :) , m_G^{(l)}[v_t; :) )
\end{align*}
\]

(1)

where aggregate and transform are the message aggregation and transformation functions, respectively. \( v_t \) represents the target node whose embedding is to be learned. \( l \) in the above formulas denotes the \( l \)-th layer. \( \mathcal{N}(v_t) \) is the set of neighboring nodes for \( v_t \). \( m_G^{(l)} \) is the aggregated message encapsulating information from the neighboring nodes, whose dimension size is \( n \times d \) when \( l = 1 \) and \( n \times d' \) when \( l > 1 \). In general, a GNN generates the representation of a node by combining representation of its own and its neighbors.

The main focus of this work is to conduct unsupervised node representation learning on graphs, which is defined as follows:

Definition 2 (Unsupervised node representation learning): Provided an attributed graph \( \mathcal{G} = (X \in \mathbb{R}^{n \times d}, A \in \mathbb{R}^{n \times n}) \), where \( n \) is the number of nodes and \( d \) is the feature dimension, our aim is to train an effective GNN \( g_\theta \), which learns node embeddings \( H \in \mathbb{R}^{n \times d'} \) from \( \mathcal{G} = (X, A) \), where \( d' \) is the hidden and the output dimension size, without relying on any labeling information.

Finally, by passing the output embeddings \( H \) to an output classifier, our model is able to handle various downstream tasks, such as node classification.

IV. METHODOLOGY

In this section, we present the proposed G-Zoom algorithm. As shown in Fig. 2, our model consists of two main components, which are graph augmentation and encoding, and contrastive learning with adjusted zooming. First, we apply graph augmentations including graph diffusion and sampling to generate two augmented views based on the input graph. While the first augmented view \( \tilde{G}_1 \) is created by applying GS on the input graph \( \mathcal{G} \) directly, the second congruent view \( \tilde{G}_2 \) is sampled from the diffused graph \( \mathcal{G} \). Then, by feeding these two views to the GNN encoder \( g_\theta \), we map node features to the latent space, denoted as \( H_1 \) and \( H_2 \) for \( \tilde{G}_1 \) and \( \tilde{G}_2 \), respectively. This process is shown in the leftmost part in Fig. 2. After this, we set up different contrastive paths on multiple scales via the proposed adjusted zooming scheme to facilitate the extraction of self-supervision signals for our model training. Given a target node \( v_t \), we can get its top- \( k \)-correlated neighbors by designing a neighborhood register, which is defined in Definition 3 (defined in Section IV-B2).

Then, we aggregate their embeddings through a pooling layer to obtain the neighboring-level representation of \( v_t \), \( n_1^t \) and \( n_2^t \) for the two views. To obtain the subgraph-level representation \( s_1 \) and \( s_2 \) for \( \tilde{G}_1 \) and \( \tilde{G}_2 \), we readout the embeddings of nodes in \( H_1 \) and \( H_2 \) via the same pooling layer. At this stage, we conduct contrastive learning through the adjusted zooming scheme on three perspectives, namely, micro, meso, and macro by contrasting among node-, neighboring-, and subgraph-level representations of the target node \( v_t \) in two augmented views.

The following section will discuss the two main segments of our framework in detail (Sections IV-A to IV-B) and the model training and algorithm of G-Zoom (Section IV-C).

A. Graph Augmentations and Encoding

Recently, there are some successful attempts on self-supervised visual representation learning, which allow encoders to learn effective representations by contrasting different augmented views of images [39]. However, we cannot directly use standard image augmentations such as image cropping, rotation, and distortion, to graphs due to their complex properties, e.g., no spatial locality and fixed node ordering. To address this issue, there are mainly two types of augmentations on graphs: topological and attributive. While the first type of strategy augments the topological space of a graph via GS, edge dropout (ED), or graph diffusion (GD), the latter one conducts the augmentation operation on attributive properties of nodes, such as node feature dropout (NFD), which masks a fraction of node feature dimensions with zeros.

In our model, we mainly apply topological graph augmentations to establish different graph views. The reason is not only...
because there are many unattributed graphs in the real world but also through our experiments (i.e., Section VI-E) we find that attributive augmentations bring limited performance gain to G-Zoom. Specifically, we use GS to generate a congruent view $\tilde{G}_1$ and use the combination of GD and GS to create an incongruent view $\tilde{G}_2$ of the input graph. Note that all the nodes in $\tilde{G}_1$ are the same as $\tilde{G}_2$ since they share the same subsampling scheme. Previous works on graph contrastive learning, such as MERIT [43] and MVGRL [37], suggest that the congruent view with and without GD are expected to provide a global and local viewpoint when using the underlying topological information. Therefore, in our model, we adopt a similar approach to assist G-Zoom in encoding rich local and global information of a graph simultaneously. The detailed explanation of GS, ED, GD, and NFD is shown below.

1) **Graph Sampling:** To ensure an augmented view includes all the neighbors, i.e., all top-k strongly related neighbors, of a batch of target nodes for generating neighboring-level embeddings, we have used a special GS approach. Given a batch with $B$ nodes, $k$ closely related neighbors to be sampled for each target node $v_t$, and the sampled subgraph size $P$, to generate the sampled graph $\tilde{G}$, we first need to include all $B$ target nodes and all their neighbors, whose size is at most $B \times k$ in the sampled subgraph. In a batch, target nodes may share the same neighbors, and thus, the number of $B$ target nodes plus their top-$k$ neighbors is likely to be smaller than $B \times k$. Then, we sample nodes from $\tilde{G}$ excluding $B$ target nodes and their top-$k$ neighbors to enlarge the sampled graph to size $P$. In G-Zoom, to ensure all the target nodes and their neighbors can be included in $\tilde{G}$, the sampled graph size $P$ has to be larger than the number of a batch of target nodes and their top-$k$ neighbors. Also, $P$ has to be smaller than the size of $\tilde{G}$ to ensure that the sampled subgraph is included in the input graph. Then, we randomly sample nodes other than target nodes and their neighbors in the input graph $G$ to complete $\tilde{G}$. An additional advantage of GS is enabling the batch processing to extend the scalability of our model, especially allowing the learning on large-scale graphs.

2) **Edge Dropout:** ED is a type of augmentation placed on the graph topological space. With a predefined probability $p$, ED first randomly removes a $p$ percent of edges in a subgraph $\tilde{G}$, and then adds the same proportion of edges back to it. Note that both adding and dropping edges processes follow i.i.d. uniform distribution.

3) **Graph Diffusion:** We adopt graph diffusion for creating the incongruent view $\tilde{G}_2$ to inject global information to facilitate our different contrastive schemes. GD is proposed to solve two main issues in previous GNNs, which are the limited receptive field for message passing and noisy edges in real-world graphs [25]. Specifically, GD creates a new view $\tilde{G}$ through the spatial message passing, which can distribute node information to a broader neighborhood instead of passing only to the first-hop neighbors. As such, GD enables the model to aggregate information from multihop neighbors and encode richer global information into nodes. This process can be formulated as follows:

$$S = \sum_{a=0}^{\infty} \theta_a T^a$$  \hspace{1cm} (2)

where $\theta_a$ is a weighting term to determine the proportion of local and global information to be encoded. $T \in \mathbb{R}^{n \times n}$ denotes...
the generalized transition matrix. To ensure the convergence, two conditions including \( \theta_a \in [0, 1] \), \( \sum_{a=d}^{+\infty} \theta_a = 1 \) and the eigenvalues of \( T \) are restricted by \( \lambda_i \in [0, 1] \) should be satisfied.

In our study, we adopted the personalized page rank (PPR)-based graph diffusion \( S_{PPR} \). Given an adjacency matrix \( A \in \mathbb{R}^{n \times n} \) and its degree matrix \( D \in \mathbb{R}^{n \times n} \), the transition matrix \( T \) and the weighting coefficient \( \theta_a \) can be formulated as \( T = D^{-1/2} A D^{-1/2} \) and \( \theta_a = \alpha(1 - \alpha)^a \), respectively. Here, \( \alpha \in [0, 1] \) denotes the teleport probability, which determines the tendency of returning to the starting node or teleporting to its neighbors in an RW. \( S_{PPR} \) can be formulated as follows [37]:

\[
S_{PPR} = \alpha(I - (1 - \alpha)D^{-1/2}AD^{-1/2})^{-1}.
\]

Graph diffusion effectively generates an incongruent view with a more informative neighborhood. However, it is problematic when tackling large-scale graphs due to its memory-intensive operation on computing matrix inversion. To overcome this problem and to make G-Zoom scalable to large-scale graphs, we have proposed a rowwise graph diffusion method, which is shown in (12). Specifically, we adopt power iteration (i.e., iteratively doing matrix multiplication) to approximate matrix inversion for diffusion matrix generation.

4) Node Feature Dropout: To augment the attributive information of a graph, NFD randomly masks a fraction of feature dimensions in the feature matrix of \( \tilde{G} \), i.e., \( \tilde{X} \), by defining an augmentation ratio \( p \). In other words, after applying NFD, \( p \) percent of the columns of \( \tilde{X} \) would be assigned zeroes.

After getting \( \tilde{G}_1 \) and \( \tilde{G}_2 \) via different graph augmentations, we feed them into a shared GNN encoder \( g_{θ} \). In our study, for simplicity, we adopt a one-layer GCN [1] as the backbone encoder

\[
\tilde{H}_g^{(l)} = σ \left( \tilde{D}^{-1/2} \tilde{A}^{-1/2} \tilde{H}_g^{(l-1)} \tilde{W}^{(l-1)} \right)
\]

where \( \tilde{A} = A + I \) and \( \tilde{D} \) is the diagonal degree matrix of \( \tilde{A} \). In the above formula, \( l \) denotes the \( l \)th layer, and \( \tilde{W} \) is a learnable weight matrix. We resort to this encoder to map node the underlying topological and attributional information into the latent space, where we have node representations \( \tilde{H}_1 \) and \( \tilde{H}_2 \) for \( \tilde{G}_1 \) and \( \tilde{G}_2 \).

B. Graph Contrastive Learning With Adjusted Zooming

In G-Zoom, we introduce a graph contrastive scheme based on the adjusted zooming, which injects contrastiveness among multiple scales: micro, meso, and macro, each of which represents the node-, neighboring-, and subgraph-level viewpoints.

1) Micro (Node-Level) Contrastiveness: Viewing a graph from the micro perspective helps G-Zoom focus on specific self-information of a node. In this level of contrastiveness, we compare target node embeddings in two graph views, where each view contains the graph topological information on different scales.

After mapping the node features of \( \tilde{G}_1 \) and \( \tilde{G}_2 \) to the latent space via the GNN encoder \( g_{θ} \), we have \( \tilde{H}_1 \) and \( \tilde{H}_2 \) as the node representation matrices of two graph views. This GNN encoder can be replaced with different GNN methods (e.g., GCN [1] and GAT [10]). For simplicity, we only use a one-layer GCN as the backbone graph encoder in our model. As shown in Fig. 3, given a target node \( v_t \), we first maximize the cosine similarity of the positive pair \((\tilde{h}_1^{v_t}, \tilde{h}_2^{v_t})\), i.e., the red solid line, where \( \tilde{h}_1^{v_t} \in \tilde{H}_1 \) and \( \tilde{h}_2^{v_t} \in \tilde{H}_2 \) are the representations of \( v_t \) in two views. With this maximization, we pull the representations of a node closer in two views to learn the invariant patterns. To regularize this discrimination and avoid model collapse, we have defined two negative sampling strategies to consist of our micro contrastiveness, which further exploits the rich contrastive relations within and between \( H_1 \) and \( H_2 \). In Fig. 3, the blue dashed line shows the first type of negative sampling, which considers all the nodes except the target node within a view as negative samples. Similarly, the yellow dashed line indicates the second type of negative sampling, which regards all the nodes excluding the target node in a different view as negative counterparts. The cosine similarity of all these negative pairs needs to be minimized. Thus, the aforementioned process can be formulated as follows:

\[
\mathcal{L}(\tilde{h}_1^{v_t}, \tilde{h}_2^{v_t}) = \log \frac{e^{s(\tilde{h}_1^{v_t}, \tilde{h}_2^{v_t})}}{\sum_{i=1}^{2} \sum_{j=1; j \neq i}^{B} e^{s(\tilde{h}_1^{v_t}, \tilde{h}_j^{v_t})}}
\]

where \( s(\cdot) \) is the cosine similarity, \( B \) denotes the number of target nodes in a batch, and \( v \in [1, 2] \) denotes one of the augmented view. If \( v = 1 \), \( \tilde{h}_1^{v_t} \) represents the representation of node \( i \) in \( \tilde{H}_1 \), whereas when \( v = 2 \), \( \tilde{h}_2^{v_t} \) is the node representation of node \( i \) in \( \tilde{H}_2 \). Taking the argument in an opposite position, i.e., \((\tilde{h}_2^{v_t}, \tilde{h}_1^{v_t})\), the loss \( \mathcal{L}(\tilde{h}_1^{v_t}, \tilde{h}_2^{v_t}) \) is combined with \( \mathcal{L}(\tilde{h}_1^{v_t}, \tilde{h}_2^{v_t}) \) to reinforce the self-supervised signals. The micro-level objective function of our model can be defined as follows:

\[
\mathcal{L}_{\text{micro}} = -\frac{1}{2B} \sum_{V_t=1}^{B} \left( \mathcal{L}(\tilde{h}_1^{v_t}, \tilde{h}_2^{v_t}) + \mathcal{L}(\tilde{h}_1^{v_t}, \tilde{h}_2^{v_t}) \right).
\]
scoring approaches to measure the importance among nodes in a graph: the RW-based and PPR-based approach. The first method determines the importance of a node \(v_t\) to the target node \(v_i\) by creating a randomly generated sequence starting from node \(v_t\) and counting the occurrence frequency of the node \(v_i\) in the sequence. The higher the occurrence frequency of \(v_i\), the more important it is to \(v_t\). On the other hand, the latter approach uses the PPR algorithm, whose formulation is shown in (3).

As shown in Fig. 4, to generate the neighboring-level representations for target nodes in a batch by averagely aggregating the node representations in the register via the pooling layer (i.e., mean pooling).

In the neighborhood register includes the top-5 strongly correlated neighbors for the target node \(v_i\), namely, \(v_a, v_b, v_c, v_d, \) and \(v_e\). With this register, we can calculate the neighboring-level representations for target nodes in a batch by aggregating the node representations in the register via the pooling layer. Specifically, for a target node \(v_i\), we use \(\mathbf{n}_1^t\) and \(\mathbf{n}_2^t\) to denote its neighboring-level representations in \(H_1\) and \(H_2\), respectively. This aggregation process is presented as the yellow and blue boxes in the figure. Then, to establish the neighboring-level contrastiveness, we use the idea of Deep InfoMax [39] to maximize the mutual information between two positive pairs \((\mathbf{h}_i^1, \mathbf{n}_2^1)\) and \((\mathbf{h}_i^2, \mathbf{n}_2^1)\), while pushing away two negative pairs \((\mathbf{h}_i^1, \mathbf{n}_2^0)\) and \((\mathbf{h}_i^1, \mathbf{n}_2^0)\) by minimizing their agreement, where \(\mathbf{h}_i^1\) is the corrupted representation of \(v_t\) extracted from the learned representations \(\tilde{\mathbf{H}}\) of a randomly shuffled feature matrix \(\tilde{\mathbf{X}}\). The contrastive loss for \((\mathbf{h}_i^1, \mathbf{n}_2^1)\) can be formulated as

\[
\mathcal{L}(\mathbf{h}_i^1, \mathbf{n}_2^1) = \sum_{t=1}^{B} \log \mathcal{D}(\mathbf{h}_i^1, \mathbf{n}_2^1) + \log \left(1 - \mathcal{D}(\mathbf{h}_i^1, \mathbf{n}_2^0)\right) \tag{7}
\]

where \(\mathcal{L}(\mathbf{h}_i^1, \mathbf{n}_2^1)\) denotes the neighboring-level contrastiveness loss for \((\mathbf{h}_i^1, \mathbf{n}_2^1)\). Also, \(B\) is the number of nodes in a batch, and \(\mathcal{D}\) is a discriminator, which computes the agreement between the elements in an instance pair to evaluate its mutual information. The formulation of \(\mathcal{D}\) is shown in (13). Similarly, we can obtain the contrastive loss \(\mathcal{L}(\mathbf{h}_i^2, \mathbf{n}_2^1)\).

The overall neighboring-level contrastiveness can be defined by marrying \(\mathcal{L}(\mathbf{h}_i^1, \mathbf{n}_2^1)\) and \(\mathcal{L}(\mathbf{h}_i^2, \mathbf{n}_2^1)\)

\[
\mathcal{L}_{\text{meso}} = -\frac{1}{2B} \left(\mathcal{L}(\mathbf{h}_i^1, \mathbf{n}_2^1) + \mathcal{L}(\mathbf{h}_i^2, \mathbf{n}_2^1)\right). \tag{8}
\]

3) Macro (Subgraph-Level) Contrastiveness: From the macro perspective, our model can have a bird’s-eye view to inspect all the nodes within a sampled graph. At this level, we define a contrastive path by maximizing the agreement between target node embeddings and subgraph-level representations, which can be obtained by aggregating all the node embeddings in the sampled subgraph with size \(P\), which consists of a batch of target nodes, their top-\(k\) neighbors, and a small portion of randomly sampled nodes. This process is expected to extract rich global information from the graph.

As shown in Fig. 5, to obtain the subgraph-level representations (i.e., \(s_1\) for \(H_1\) and \(s_2\) for \(H_2\)), we feed \(H_1\) and \(H_2\) into the pooling layer. Then, we contrast the target node embeddings with subgraph-level representations between different views, i.e., \(\mathbf{h}_1^1\) versus \(s_2\) and \(\mathbf{h}_1^2\) versus \(s_1\). The contrastive loss for the
instance pair \((h_1^t, s_2^t)\) can be formulated as
\[
L(h_1^t, s_2^t) = \sum_{i=1}^{B} \log D(h_1^t, s_2^t) + \log \left(1 - D(h_1^t, s_2^t) \right) \tag{9}
\]
and we can calculate the loss \(L(h_1^t, s_2^t)\) in a similar way. By merging these two losses, we can get the objective function of our subgraph-level contrastiveness
\[
L_{\text{Macro}} = -\frac{1}{2B} \left( L(h_1^t, s_2^t) + L(h_1^t, s_1^t) \right). \tag{10}
\]

C. Model Optimization and Algorithm
\[
L = \alpha L_{\text{micro}} + \beta L_{\text{meso}} + \gamma L_{\text{macro}}. \tag{11}
\]

Algorithm 1 Overall Procedure of G-Zoom

| Input | Output |
|-------|--------|
| Input Graph: \(G\); Diffused Graph: \(\tilde{G}\); Batch Size: \(B\); Number of top neighbors for target node \(v_t\); \(k\); Neighborhood Register: \(R\); Sample Size: \(P\); Maximum training epochs: \(E\); | Final representations \(\tilde{H}\); |

1. //Model Training;
2. for \(e = 1 \text{ to } E\) do
3. //Batch sampling;
4. Sample \(B\) target nodes \(V\) to create a batch;
5. Create \(\tilde{G}_1\) and \(\tilde{G}_2\) from \(G\) and \(\tilde{G}\), respectively, as described in section IV-A1;
6. //Obtain node-, neighboring-, and subgraph-level representations;
7. for \(v_t\) in \(V\) do
8. Obtain \(h_1^t\) and \(h_2^t\) via (4);
9. Obtain \(n_1^t\) and \(n_2^t\) by aggregating neighbors stored in \(R\);
10. Obtain \(s_1\) and \(s_2\) by aggregating all the nodes in \(\tilde{G}_1\) and \(\tilde{G}_2\);
11. //Compute Loss;
12. Calculate \(L_{\text{micro}}\) via (6);
13. Calculate \(L_{\text{meso}}\) via (8);
14. Calculate \(L_{\text{macro}}\) via (10);
15. Calculate \(L\) via (11);
16. //Update parameters;
17. Update trainable matrix \(W\) in encoder \(g_0\) and \(W_D\) in discriminator shown in (13);
18. //Inference;
19. Obtain final embeddings \(\tilde{H} = g_0(G) + g_0(\tilde{G})\).

To train our model and learn effective node embeddings for downstream tasks, we combine the contrastiveness loss from the aforementioned three perspectives to define the overall objective function of G-Zoom, where \(\alpha\), \(\beta\), and \(\gamma\) are three weighting terms for controlling the contribution of each contrastiveness loss in model training. During the training process, we aim to minimize the overall loss \(L\). For model inference, to obtain the final representation of a target node \(v_t\), we combine \(h_1^t\) and \(h_2^t\), i.e., \(h_1^t + h_2^t \in \mathbb{R}^d\), to get the final node representation \(\tilde{h}\).

Algorithm: The overall procedure of G-Zoom is depicted in Algorithm 1. From Algorithm 1, we can see that the overall procedure consists of two stages: model training and inference. During model training, within every epoch, we create two augmented graph views \(\tilde{G}_1\) and \(\tilde{G}_2\) with the sample size \(P\) as the inputs of our GNN encoder \(g_0\) in three steps. First, we sample \(B\) target nodes as the base of graph views. Then, we put all these target node top-\(k\) neighbors in the sampled views. Finally, we enlarge these two views to size \(P\) by adding randomly sampled nodes from the input graph \(G\) excluding the selected \(B\) target nodes and their top-\(k\) neighbors.

After this, we feed \(\tilde{G}_1\) and \(\tilde{G}_2\) to \(g_0\), which outputs \(H_1\) and \(H_2\). Then, by leveraging the neighborhood register and pooling layer, we construct three different contrastive paths in the latent space from various perspectives (e.g., micro, meso, and macro). Finally, we have our overall loss \(L\) by combining three levels of contrastiveness. After this, a backward propagation with the gradient descent algorithm fuels the parameter updating process for \(g_0\).

At the inference stage, summing up the result representations of input graph \(G\) and the diffused graph \(\tilde{G}\) generates the final embedding matrix \(\tilde{H}\).

V. COMPLEXITY ANALYSIS AND FURTHER EXTENSION

We provide an analysis of our algorithm G-Zoom and how to extend G-Zoom to handle large-scale graphs.

A. Time Complexity of G-Zoom

In this section, we analyze the time complexity of G-Zoom by considering two main components: graph augmentation and contrastive learning process. Specifically, we train our model at most \(E\) times, and within every epoch, we conduct subsampling on the input graph once, whose time complexity is \(O(P)\), where \(P\) is the sample size. Therefore, the total complexity for this component is \(O(EP)\).

For the contrastive learning process, its time complexity is mainly contributed by the GNN encoder, the infoNCE loss computation, and the discriminator. The time complexity of the GNN encoder is \(O(\mathcal{E})\) with a sparse aware matrix product, where \(\mathcal{E}\) is the number of edges of a graph. In our method, with a one-layer GCN, the time complexity of the GNN component is \(O(\mathcal{E}_1 + \mathcal{E}_2)\), where \(\mathcal{E}_1\) is the number of edges for the original sampled graph, and \(\mathcal{E}_2\) represents the diffused one. With diffusion, \(\mathcal{E}_2\) is significantly larger than \(\mathcal{E}_1\). Then, we compute the infoNCE loss for micro-level contrastiveness, whose time complexity is \(O(B^2d')\), as shown in (5), where \(B\) is the batch size of target nodes. As shown in (13), we adopt a bilinear discriminator for calculating the other two losses (i.e., meso and macro levels). In our methods, in each training epoch, we need to process the two-level losses using (13). As shown in (7) and (9), each of these two losses requires \(O(2Bd')\) for computation. Therefore, calculating these two-level losses requires \(O(4Bd')\). To sum up, in each training epoch, the time complexity of processing all three contrastiveness is \(O(4Bd' + B^2d')\). Also, as shown
in (6), (8), and (10), we combine two losses to reinforce the self-supervised signals, and the time complexity of the computation should be doubled to $O(8Bd' + 2B^2d')$. By combining the two main components, the overall complexity is $O(E(P + E_1 + E_2 + 8Bd' + 2B^2d'))$. As we adopted Big O notation, we ignore the constant, and the final time complexity becomes $O(E(P + E_1 + E_2 + Bd' + B^2d'))$.

**B. Extending G-Zoom to Large-Scale Graphs**

To fulfill the applications in the real world and increase the scalability of G-Zoom, we extend our proposed algorithm to large-scale graphs. Most self-supervised GNN studies cannot handle large input graphs mainly due to two reasons. First, they feed the entire graph as the input for model training, e.g., DGI [12], GRACE [13], and GMI [14]. In this case, the capacity of a model is constrained by the input graph size. Second, some of them adopt a nonexpansible graph diffusion approach as their structural graph augmentation, e.g., MVGRL [37].

To overcome the aforementioned two issues, we first use a tailored sub-sampling sampler, which has been introduced in Section IV-A1, to create subgraphs. This sampler not only alleviates the restriction of the input graph size in model training but can also sufficiently include the important features of the initial graph. To prove the effectiveness of this sampler, we present the theoretical analysis in Appendix A.

To tackle the second issue, we apply a scalable graph diffusion method to create a diffused graph $\tilde{G}$ and then use the aforementioned sub-sampling strategy to generate the augmented view $\tilde{G}_2$. Without additional graph diffusion operations, $\tilde{G}_1$ is obtained by GS. Specifically, we adopt a rowwise power-iteration-based diffusion approach with the PPR kernel instead of getting the final diffused graph at once by applying (3), we calculate the importance score node by node, and then concatenate all the output importance vectors together to form the final diffused graph. There are two advantages of applying this rowwise diffusion approach.

1) Parallel the computation of diffusion matrix, which significantly speeds up the diffusion process on graphs.
2) Alleviate the resource burdens of matrix inversion when calculating the graph diffusion matrix, which only supports the operations on dense matrices.

With power iteration, we can approximate the final diffusion by iteratively conducting matrix multiplication rather than computing matrix inversion. This process can be formulated as

$$\pi_{ppr}(v_i) = (1 - \alpha)i + \alpha \tilde{A}_i \pi_{ppr}(v_i)$$

where $v_i$ is a node, $i$ is a one-hot identity vector, in which only the $i$th row entry is one, and all the remaining entries are 0. $\pi_{ppr}(x)$ is the importance vector that stores all the nodes’ importance to $v_i$, $\alpha$ is a teleport probability determining the tendency of teleporting back to node $i$, and $\tilde{A}$ is the normalized adjacency matrix $A\textbf{D}^{-1}$, where $D$ is the diagonal matrix. To show the impact of using the subsampling and the proposed scalable graph diffusion method, we compare the performance of G-Zoom with two variants, which remove the subsampling or use the scalable graph diffusion in Section VI-D.

### VI. Experiment

In this section, we have conducted experiments to present the effectiveness of G-Zoom. First, we introduce five benchmark datasets and experimental setup in Sections VI-A and VI-B. After this, the overall comparisons, ablation study, parameter sensitivity analysis, and visualization are illustrated from Sections VI-C to VI-G.

#### A. Datasets

We have selected five benchmark datasets varying with sizes, including Cora, Citeseer, Coauthor CS, Coauthor Physics, and ogbn-arxiv for experiments. Within each dataset, nodes represent scientific publications and are connected via citations. As for attributive information, while the first four datasets carry the bag-of-word representation, the feature vector of ogbn-arxiv is obtained by averaging the word embeddings of paper title and abstract. The statistics of this dataset are summarized in Table II.

As shown in the table above, the first two datasets only have thousands of nodes and edges, whereas Coauthor CS and Coauthor Physics have over 10,000 vertices and edges. Retrieving from open graph benchmark (OGB) [44], ogbn-arxiv is a large-scale graph dataset with over a hundred thousand nodes and a million edges, which is typically used for evaluating the robustness and scalability of the graph models.

#### B. Experiment Setup

In this section, we discuss specific module designs and hyperparameter settings in G-Zoom. In our experiment, for simplicity, we adopt a one-layer GCN as the backbone graph encoder in our model, as shown in (4). It is worth noting that our graph encoder can also be replaced by other GNN-based candidate models, such as GAT and SGC.

For the discriminator used in meso and macro contrastiveness, we consider a simple bilinear scoring function [45] as the discriminator $D$ in our framework, which can be formulated as

$$\mathcal{D}(h_i, x_j) = \sigma(h_i^T W_D x_j)$$

where $(h_i \in \mathbb{R}^{d'}, x_j \in \mathbb{R}^{d'})$ are a pair of representation to be scored, $\sigma$ is a nonlinear activation function and $W_D \in \mathbb{R}^{d' \times d'}$ represents a trainable matrix.

For hyperparameter settings, all the experiments are implemented using PyTorch [46]. In the training phase, we set the batch size $B$ to 200 and 400 for Cora and Citeseer, respectively, while $B$ for the three remaining datasets is set to 500. For the number of strongly related neighbors $k$ of $B$
target nodes to be collected in the sampled graph, we define $k$ as 100 for all the datasets except ogbn-arxiv where $k = 10$. As discussed in Section IV-A1, the sample graph size $P$ has to be larger than the number of $B$ and their top-$k$ neighbors. As such, $P$ is set to be larger than $B \times k$ to ensure $P$ meets this requirement. Specifically, we set 3000 for the first two datasets, while $P$ is 7000 for Coauthor CS, Coauthor Physics, and ogbn-arxiv. To improve the training efficiency, we use an early stopping mechanism to detect the model convergence and halt the training process based on a predefined patience value. The training epochs are fixed to at most 3000 for all the datasets. The patience is defined as 50 for the first two datasets and 200 for Coauthor datasets and ogbn-arxiv. Except for ogbn-arxiv, whose learning rate is 0.0001, all the datasets have the learning rate tuned to 0.001. Also, for the weighting terms $\alpha$, $\beta$, and $\gamma$ used in loss calculation as presented in (11), they are all set to 1 except for Coauthor Physics, where $\alpha$ is set to 0.8. Therefore, in the experiment, the three-level contrastiveness contributes almost equally to our model training.

### C. Performance Comparison of G-Zoom and Baselines

In this section, we compare the performance of G-Zoom in the aforementioned five datasets with nine baselines, including four supervised and five self-supervised baselines. We adopt linear node classification as the evaluation protocol to validate the expressiveness of the learned node representations.

1) **Supervised GRL Baselines:**

1. GCN [1] adapts the traditional convolution network used in image processing to graph-structured data.
2. GAT [10] adopts an attention mechanism to consider weights of neighbors in information aggregation instead of simply averaging their representations.
3. SGC [11] reduces the complexity of GCN by removing its nonlinearities and collapsing weight matrices.
4. CG3 [47] uses both data similarities and graph structure to enrich the supervision signals in graph-based semi-supervised learning.

2) **Self-Supervised GRL Baselines:**

1. DGI [12] adapts the idea of MI maximization from image processing [39] to graphs. Specifically, DGI implements a patch summary maximization strategy (i.e., maximizing MI between node- and graph-level representations).
2. GMI [14] focuses on maximizing MI between input graphs and high-level hidden representations based on node features and topological structure.
3. MVGRL [37] uses graph augmentation techniques to generate multiple views and maximize the mutual information of the output node-level and readout subgraph-level representations between different views.
4. GRACE [13] also applies graph augmentation for creating two augmented views. Unlike the aforementioned self-supervised GRL baselines, it defines an InfoNCE contrastiveness loss, which contrasts embeddings at the node level from two different augmented views.
5. SubG-Con [15] defines a self-supervised graph contrastive learning strategy, which extracts regional structure information from the correlation between central nodes and their sampled subgraphs (i.e., a subgraph built by central nodes and its close neighbors).

The experimental result is shown in Table III. Here, we adopt classification accuracy as the evaluation metrics. Every data entry in the table is the averaged result and associated standard deviation over ten individual runs. According to this table, we have made the following observations.

1) On all the datasets, our proposed approach G-Zoom has surpassed all the baselines and achieved the best performance. Notably, G-Zoom exceeds the best result of its self-supervised GRL counterparts by more than 1% on Cora and around 1% on CiteSeer and Coauthor CS. This is mainly because G-Zoom successfully extracts useful clues from different perspectives through contrastive learning via the proposed adjusted zooming scheme, by which G-Zoom can inspect graphs comprehensively and learn effective embeddings.

2) G-Zoom successfully handles a large-scale dataset (i.e., ogbn-arxiv), and achieves the best performance on the large-scale dataset. At the same time, almost all the self-supervised GRL baselines except for SubG-Con failed to handle it within the limited GPU memory budget. This is because G-Zoom adopts a tailored

### Table III

| Information Used | Method   | Cora  | CiteSeer | Coauthor Physics | Coauthor CS | ogbn-arxiv |
|------------------|----------|-------|----------|------------------|-------------|------------|
| $X, A, Y$        | GCN      | 81.5  | 70.3     | 95.7 ± 0.2       | 93.0 ± 0.3  | 68.15 ± 0.1 |
| $X, A, Y$        | GAT      | 83.0 ± 0.7 | 72.5 ± 0.7 | 95.5 ± 0.2       | 92.3 ± 0.2  | 68.85 ± 0.1 |
| $X, A, Y$        | SGC      | 81.0 ± 0.0 | 71.9 ± 0.1 | 95.8 ± 0.1       | 92.7 ± 0.1  | 68.91 ± 0.1 |
| $X, A, Y$        | CG3      | 83.4 ± 0.7 | 73.6 ± 0.8 | OOM              | 92.3 ± 0.2  | OOM        |
| $X, A$           | DGI      | 81.7 ± 0.6 | 71.5 ± 0.7 | 94.5 ± 0.5       | 92.2 ± 0.6  | OOM        |
| $X, A$           | GMI      | 82.7 ± 0.2 | 73.0 ± 0.3 | OOM              | OOM         | OOM        |
| $X, A$           | MVGRL    | 82.9 ± 0.7 | 72.6 ± 0.7 | 95.3 ± 0.1       | 92.1 ± 0.1  | OOM        |
| $X, A$           | GRACE    | 80.0 ± 0.4 | 71.7 ± 0.6 | 95.2 ± 0.2       | 92.8 ± 0.1  | OOM        |
| $X, A$           | SubG-Con | 83.5 ± 0.5 | 73.2 ± 0.2 | 96.2 ± 0.4       | 94.0 ± 0.6  | 55.7 ± 0.2 |
| $X, A$           | G-Zoom   | 84.7 ± 0.4 | 74.2 ± 0.3 | 96.6 ± 0.3       | 94.9 ± 0.3  | 70.1 ± 0.1 |
TABLE IV
PERFORMANCE COMPARISON OF G-ZOOM AND ITS SIX VARIANTS ON FIVE BENCHMARK DATASETS

| Loss       | Cora      | CiteSeer    | Coauthor CS | Coauthor Physics | ogbn-arxiv |
|------------|-----------|-------------|-------------|------------------|------------|
| G-ZoomMacro | 83.4 ±0.4 | 73.9 ±0.4   | 92.9 ±0.7   | 96.0 ±0.3        | 69.4 ±0.1  |
| G-ZoomMeso | 83.7 ±0.7 | 73.9 ±0.6   | 93.1 ±0.6   | 95.6 ±0.2        | 69.4 ±0.1  |
| G-ZoomMicro | 77.31 ±1.2| 66.7 ±1.1   | 89.2 ±0.9   | 95.6 ±0.3        | 69.9 ±0.1  |
| G-ZoomMicro | 83.7 ±0.5 | 73.8 ±0.4   | 92.8 ±0.4   | 95.4 ±0.3        | 69.5 ±0.1  |
| G-ZoomMeso | 84.3 ±0.7 | 73.3 ±0.4   | 93.9 ±0.6   | 96.3 ±0.3        | 70.1 ±0.1  |
| G-ZoomMacro | 84.4 ±0.5 | 73.2 ±0.4   | 94.0 ±0.5   | 96.2 ±0.2        | 70.2 ±0.1  |
| G-Zoom     | 84.7 ±0.4 | 74.2 ±0.3   | 94.9 ±0.3   | 96.6 ±0.3        | 70.1 ±0.1  |

D. Ablation Study

As mentioned in Section IV-C, G-Zoom establishes the contrastive paths from three perspectives, namely, micro, meso, and macro, to encode both localized and global information comprehensively. To shed light on our proposed adjusted zooming-based contrastiveness, we present the experimental results of G-Zoom variants which only adopt a specific level of loss or use the combination of two contrastive losses on five benchmark datasets, as shown in Table IV. For better illustration, we use G-ZoomMicro, G-ZoomMeso, and G-ZoomMacro to denote G-Zoom variants without considering the micro, meso, and macro perspectives, respectively. Also, G-ZoomMicro, G-ZoomMeso, and G-ZoomMacro represent the reduced models using only one level of contrastiveness. From Table IV, we observe that G-ZoomMeso and G-ZoomMacro generally surpass variants with only one level of contrastiveness. This indicates that adding G-ZoomMeso or G-ZoomMacro improves the model performance, to which the inclusion of two distinct viewpoints contributes. While the meso or macro viewpoint can provide rich global information, the micro viewpoint extracts the most fine-grained localized information from node–node comparison.

Also, we can see that without either the meso or macro contrastiveness, G-Zoom cannot achieve the best performance in four out of five datasets, which indicates that the combination of these two components is effective. We conjecture this is because the meso module can force the model to explore the fine-grained information embodied in a specific scale of a graph, which can be easily neglected from a macro-level viewpoint.

To evaluate the impact of the subsampling and the proposed graph diffusion technique (defined in Section V-B) on G-Zoom, we compare two model variants G-ZoomSample (i.e., G-Zoom without subsampling) and G-ZoomPower (i.e., G-Zoom with power iteration diffusion) with G-Zoom. The experimental result is reported in Table V. From the table, we observe that without subsampling, the performance of G-Zoom degrades. We conjecture this is because the subsampling creates different subgraphs in each iteration, which increases the difficulty of the self-supervised learning tasks with changing topology of the input graph. Thus, subsampling can improve the model performance. In addition, using the power-iteration-based diffusion, the performance of G-ZoomPower is comparatively lower than G-Zoom with the original graph diffusion. This is probably because the power iteration graph diffusion is an approximation of the original diffusion. Thus, it may not be as comprehensive as the original diffusion in global information extraction.

E. Augmentation

To find the most appropriate augmentation scheme for G-Zoom, we have conducted experiments using eight different graph augmentations, which can be categorized into two types: “Structural” and “Structural+Attributive.” Note that there is no attributive-only augmentation scheme in our model because to ensure the scalability of G-Zoom, we have to apply GS as a structural augmentation technique to create subgraphs as the inputs of graph encoder.

The results of the experiment are presented in Table VI. As shown in the table, we can see that the most effective augmentation scheme is GS/GS+GD, which has achieved the best performance on all the benchmark datasets. Except for its effectiveness, GS/GS+GD is also comparatively more efficient in computation than most augmentation schemes. Without

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integrating NFD in an augmentation scheme negatively affects much of the underlying linkages. Also, it is worth noting that and thus applying ED on the diffused graph may corrupt too much the diffused graph has much more edges than the input graph, in messaging passing by densifying a graph. As a result, the graph diffusion operation is extending the receptive field ED on diffused graph corrupts the augmented view too much.

The graph diffusion operation is extending the receptive field and the memory consumption of G-Zoom is mainly controlled by the batch size $B$ and sample size $P$. The uses a small-size subgraph generation approach for GRL. The algorithm, we report the memory consumption of G-Zoom for choosing the number of closest neighbors for each target node on five benchmark datasets. Thus, it can handle large datasets.

### F. Memory Efficiency

To quantitatively compare the memory required by each algorithm, we report the memory consumption of G-Zoom and the baselines on five benchmark datasets. Specifically, we run the baselines based on their default setting, and set the batch size $B$ and the sample size $P$ to 200/2000 for the first two datasets and 500/5000 for the remaining datasets when running G-Zoom. As shown in Table VII, the memory consumption of G-Zoom is the lowest in three out of five datasets of all the methods, which clearly indicates the memory efficiency of G-Zoom. Another algorithm that can successfully complete all the tasks is SubG-Con, as it uses a small-size subgraph generation approach for GRL. The memory consumption of G-Zoom is mainly controlled by the batch size $B$ and the sample size $P$. With lower $B$ and $P$, the memory consumption of G-Zoom can be even lower, and thus it can handle large datasets.

### G. Parameter Study

In this section, we investigate the influences of four key parameters on the performance of our proposed model, which are batch size $B$, sample size $P$, hidden size $d'$, and parameter $k$ for choosing the number of closest neighbors for each target node on five benchmark datasets.

1) **Batch Size $B$ and Sample Size $P$**: In this experiment, we change the value of $B$ and $P$ simultaneously to see how these two parameters affect the performance of G-Zoom, and the experimental results are reported in Fig. 6(a). As illustrated in Section IV-A, increasing $B$ requires larger $P$ since the subgraph with size $P$ needs to contain $B$ nodes and their top-$k$ neighbors. Therefore, we choose five $B$ values from 100 to 500, incremented by 100, and their corresponding $P$ from 1000 to 5000, incremented by 1000. As shown in Fig. 6(a), changing $B$ and $P$ has different impacts on model performance on other datasets. We can observe that compared with small graphs such as Cora and CiteSeer, adopting large $B$ and $P$ on bigger datasets such as ogbn-arxiv, coauthor CS, and Physics in G-Zoom achieve improved results. This is probably because increasing $B$ and $P$ can help better encode global information in large graphs, while a small size $P$ is enough to contain a large proportion of nodes in small datasets. Thus, choosing appropriate $B$ and $P$ for G-Zoom is based on the size of datasets to be trained on.

2) **Hidden Size $d'$**: We investigate the sensitivity of the hidden size $d'$ for our proposed framework G-Zoom. Here, we choose 64, 128, 256, 512, and 1024 as the hidden size of our model to be explored to see how increasing $d'$ impacts the accuracy of G-Zoom on benchmark datasets. From Fig. 6(b), there is an obvious upward trend on all the datasets except for Cora, which indicates $d'$ has a positive relationship with the performance of G-Zoom. The possible reason is that rising $d'$ will increase the number of trainable parameters, which extends the expressive ability of our model. However, without sufficient data, too many parameters may cause the overfitting problem, which degrades the performance of a model [48]. This may explain why with a large $d'$, G-Zoom suffers from
lower performance on Cora since it contains a limited number of nodes and edges.

3) Neighborhood Sampling Size $k$: To explore the significance of $k$ on different datasets, we conducted experiments using four different $k$ values, which are 1, 10, 100, and 1000. Our results are demonstrated in Fig. 6(c), where we have a similar trend over all the datasets: The expressiveness of the model has been boosted by increasing $k$ from 1 to 10 or 100. However, a large number of sampled neighbors (e.g., more than 100) hurts the performance.

H. Visualization

For an intuitive illustration, we visualize the learned embeddings of four baselines and our method (i.e., GCN, DGI, GRACE, SubG-Con, and G-Zoom) on CiteSeer via using the t-SNE method [49]. The visualized representation of these embeddings is given in Fig. 7. As can be observed, the 2-D projection of embeddings generated by G-Zoom exhibits more discernible clusters than embeddings learned by other approaches. Qualitatively, visualization demonstrates the effectiveness of G-Zoom in graph self-supervised learning. To verify this, we further added a new experiment, which applied K-mean clustering to the learned representations of the five GCL methods on CiteSeer and used the normalized mutual information (NMI) as the metric to evaluate the clustering goodness. The experimental results are shown in the caption of each subfigure in Fig. 7. From the results, it can be seen that the learned representations generated by G-Zoom achieved the highest NMI score, indicating better performance.

VII. CONCLUSION

In this article, we have proposed a novel framework G-Zoom, a graph self-supervised learning approach powered by contrastive adjusted zooming. In our framework, G-Zoom consists of two components: augmented graph encoding and adjusted zooming powered contrastive learning. While the former component can generate multiple augmented views, the latter one facilitates the construction of underlying contrastive paths from three viewpoints: micro, meso, and macro. G-Zoom can effectively extract both local and global self-supervision signals of a graph by progressively inspecting it from the finest details to the coarse landscape. Extensive experiments have demonstrated the effectiveness and superiority of our model over existing self-supervised, as well as some supervised GRL baselines by large margins. In the future, we plan to exploit automatic machine learning techniques [50] to search for the best neural architecture for GRL.

APPENDIX A

THEORETICAL ANALYSIS OF THE PROPOSED SAMPLER

To prove the effectiveness of the proposed sampler, we present the theoretical analysis as follows:

Theorem 1: Given a graph $\mathcal{G} = (X \in \mathbb{R}^{n \times d}, A \in \mathbb{R}^{n \times n})$, the number of nodes in a sampled batch $B$, and the number of training epochs $E$, the probability of a node $i$ being sampled from $\mathcal{G}$ is $M$, if $E$ becomes infinite, we have

$$ P(i) = \lim_{E \to \infty} \left(1 - \frac{1}{M}ight)^{BE} = 0 \quad (14) $$

where $P(i)$ is the probability of a node $i$ not being sampled after $E$ epochs.

According to Theorem 1, we can see that when $E$ becomes larger, the probability of a node $i$ not being sampled will gradually tend toward zero. As such, we can derive that with a large $E$, our sampler can guarantee all the nodes can be sampled, which means the important features of the initial graph will not be lost during training. To validate this, we conduct an experiment to show how $P(i)$ changes when training epochs $E$ increase for five benchmark datasets,
as shown in Fig. 8, where we can see the probability of a node not being sampled quickly converged to 0 when $E$ increases. Also, the smaller the size of the dataset, the faster the convergence proceeds.

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