A Provable Framework of Learning Graph Embeddings via Summarization

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Abstract
Given a large graph, can we learn its node embeddings from a smaller summary graph? What is the relationship between embeddings learned from original graphs and their summary graphs? Graph representation learning plays an important role in many graph mining applications, but learning embeddings of large-scale graphs remains a challenge. Recent works try to alleviate it via graph summarization, which typically includes the three steps: reducing the graph size by combining nodes and edges into supernodes and superedges, learning the supernode embedding on the summary graph and then restoring the embeddings of the original nodes. However, the justification behind those steps is still unknown.

In this work, we propose GELSumm, a well-formulated graph embedding learning framework based on graph summarization, in which we show the theoretical ground of learning from summary graphs and the restoration with the three well-known graph embedding approaches in a closed form. Through extensive experiments on real-world datasets, we demonstrate that our methods can learn graph embeddings with matching or better performance on downstream tasks.

This work provides theoretical analysis for learning node embeddings via summarization and helps explain and understand the mechanism of the existing works.

Introduction
Graph representation learning has attracted much research interest in recent years due to its success in various applications including biology, computer vision, text classification, and more. However, learning node representations of large graphs still remains a problem due to high computational complexity and memory usage. To overcome this problem, some researchers resort to graph summarization methods. They first summarize (some may use the term “coursening”) the input graph into a smaller summary graph by grouping subsets of nodes into supernodes, and then learn supernode embeddings of the summary graph. After that, these embeddings are projected back to the original nodes embeddings with an additional refinement step. Since the size of the graph is decreased, the running time and memory cost are largely reduced. Representative works include HARP (Chen et al. 2018), MILE (Liang, Gurukar, and Parthasarathy 2018), and GraphZoom (Deng et al. 2020).

A key limitation of the existing summarization-based solutions is the lack of theoretical analysis. Existing methods summarize the input graph in a heuristic way and then empirically restore the embeddings of the original nodes from the supernode embeddings, without deeply exploring the connection between them. Thus, there are no theoretical studies of the underlying mechanism. Due to the lack of theoretical study, the following questions arise naturally:

• What is the theoretical mechanism behind learning node embedding via summarization?
• What is the connection between node embeddings of original graphs and summary graphs?
• Which summarization method should we use, and how should we restore embeddings correspondingly with the theoretical ground?

In this work, we aim to answer the above questions and fill the theoretical gap. Specifically, we theoretically show that applying three node embedding methods (DeepWalk, LINE, and GCN) on summary graphs is equivalent to applying them on a reconstructed graph based on the configuration model. As a result, original node embeddings can be ap-
proximated by the embeddings of summary graphs through a simple linear transformation. These theoretical results help explain the empirical success of the existing works, and bring more knowledge and understanding about this problem.

Based on the theory, we propose GELS\textsubscript{UMM}, a novel graph embedding learning framework based on summarization, which aims to learn node embeddings of the input graph from a smaller summary graph, while also providing theoretical justification. We evaluate it on several real-world datasets and show that GELS\textsubscript{UMM} can learn high quality node embeddings with similar or even better performance efficiently.

In summary, our contributions include:

• **Theory.** We analyze three node embedding learning methods, DeepWalk, LINE and GCN, and theoretically show that embeddings learned on original graphs by them can be approximated by that on summary graphs in a closed-form, as summarized in Table 1.

• **Framework.** Based on the derived theory, we propose a novel and well-formulated framework, GELS\textsubscript{UMM}, which learns node embeddings for the original graph efficiently from a smaller summary graph.

• **Extensive Experiments.** We conduct thorough empirical analysis on multiple real-world datasets, and demonstrate that our proposed GELS\textsubscript{UMM} can learn high-quality node embeddings using much less time.

**Related Work**

**Graph representation learning.** Graph representation learning (Cai, Zheng, and Chang 2018; Goyal and Ferrara 2018) aims to map each node in graphs into a low-dimensional vector (called embedding or representation) which captures the structural information. The learned latent embeddings can then be fed into machine learning and data mining algorithms for various downstream tasks, such as node classification and link prediction.

A few representative examples from the rich literature in graph representation learning include: DeepWalk (Perozzi, Al-Rfou, and Skiena 2014), node2vec (Grover and Leskovec 2016), LINE (Tang et al. 2015), and graph neural network (GNN) methods (Zhou et al. 2019; Wu et al. 2021), such as GCN (Kipf and Welling 2017), GraphSAGE (Hamilton, Ying, and Leskovec 2017) and GAT (Veličković et al. 2018), which adopt a message-passing framework and update the node embeddings based on their neighbors’ representations recursively.

Although graph representation learning methods are successful, their lack of scalability and efficiency is an important problem. To tackle this problem, some works employ sampling techniques, including layer sampling (Chen, Zhu, and Song 2018; Chen, Ma, and Xiao 2018; Huang et al. 2018) and subgraph sampling (Ying et al. 2018; Chiang et al. 2019; Zeng et al. 2020).

**Embedding learning by summarization.** Another way to improve the scalability of node embedding learning methods is via graph summarization (Yan et al. 2019; Liu et al. 2018).

The typical approach is to coarsen the original graph into a smaller summary graph, and apply representation learning methods on it to obtain intermediate supernode embeddings. The embeddings of the original nodes are then restored from supernode embeddings with a further refinement step. For example, HARP (Chen et al. 2018) finds a series of smaller graphs which preserve the global structure of the input graph, and learns representations hierarchically. HSRL (Fu, Hou, and Yao 2019) learns embeddings on multi-level summary graphs, and concatenates them to restore original embeddings. MILE (Liang, Gurukar, and Parthasarathy 2018) repeatedly coarsens the input graph into smaller ones using a hybrid matching strategy, and finally refines the embeddings via GCN to obtain the original node embeddings. GPA (Lin et al. 2020) uses METIS (Karypis and Kumar 1998) to partition the graphs, and smooths the restored embeddings via a propagation process. GraphZoom (Deng et al. 2020) employs an extra graph fusion step to combine the structural information and feature information, and then uses a spectral coarsening method to merge nodes based on their spectral similarities. Embeddings are then refined by a graph filter to ensure feature smoothness. (Fahrbach et al. 2020) learns embeddings of the given subset of nodes by coarsening the remaining nodes, which is not capable to learn embeddings of the remaining ones.

These methods, however, are based on heuristic designs and lack theoretical formulation and analysis.

**Preliminary**

In this section, we introduce two topics, graph representation learning and graph summarization. Table 2 gives the most frequently used symbols in the paper.

### Symbol and Definition

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \mathcal{G} = (\mathcal{V}, \mathcal{E}) )</td>
<td>Original graph with nodeset ( \mathcal{V} ) and edgset ( \mathcal{E} )</td>
</tr>
<tr>
<td>( \mathcal{G}_s = (\mathcal{V}_s, \mathcal{E}_s) )</td>
<td>Summary graph with supernodes ( \mathcal{V}_s ) and superedges ( \mathcal{E}_s )</td>
</tr>
<tr>
<td>( \mathcal{G}_r = (\mathcal{V}, \mathcal{E}_r) )</td>
<td>Reconstructed graph with nodeset ( \mathcal{V} ) and edgset ( \mathcal{E}_r )</td>
</tr>
<tr>
<td>( v_i )</td>
<td>Node ( i ) in the original graph ( \mathcal{G} )</td>
</tr>
<tr>
<td>( S_p )</td>
<td>Supernode ( p ) in the summary graph ( \mathcal{G}_s )</td>
</tr>
<tr>
<td>( d_i, d_p^{(s)} )</td>
<td>Degree of node ( i ) and supernode ( p )</td>
</tr>
</tbody>
</table>

#### Table 2: Major Symbols and Definitions.

Within our framework, we theoretically analyze DeepWalk, LINE, and GCN, so we discuss them in more detail here.
**DeepWalk and LINE**  DeepWalk (Perozzi, Al-Rfou, and Skiena 2014) is an unsupervised graph representation learning method inspired by the success of word2vec (Mikolov et al. 2013b,a) in text embedding. It generates random walk sequences and treats them as sentences that are later fed into a skip-gram model with negative sampling to learn latent node representations.

It has been proved in (Qiu et al. 2018) that DeepWalk is implicitly approximating and factorizing the following matrix:

\[
M := \log (\text{vol}(\mathcal{G})) \left( \frac{1}{T} \sum_{\tau=1}^{T} (\mathbf{D}^{-1} \mathbf{A})^\tau \right) \mathbf{D}^{-1} - \log b, \tag{1}
\]

where \(T\) and \(b\) are the context window size and the number of negative samples in DeepWalk, respectively.

LINE (Tang et al. 2015) learns embeddings by optimizing a carefully designed objective function that aims to preserve both the first-order and second-order proximity. Though LINE and DeepWalk appear to be different, it has been shown that LINE is also equivalent to factorizing a similar matrix to Eq. (1) and is a special case of DeepWalk for \(T = 1\) (Qiu et al. 2018):

\[
M := \log (\text{vol}(\mathcal{G})) \mathbf{D}^{-1} \mathbf{A} \mathbf{D}^{-1} - \log b. \tag{2}
\]

**GCN**  GCN (Kipf and Welling 2017) is a graph neural network model transferring traditional convolution neural network to non-Euclidean graph data. In each layer of GCN, node features are propagated based on a first-order approximation of spectral convolutions on graphs:

\[
\mathbf{E}^{(k+1)} = \sigma (\mathbf{D}^{-\frac{1}{2}} \hat{\mathbf{A}} \mathbf{D}^{-\frac{1}{2}} \mathbf{E}^{(k)} \mathbf{W}), \tag{3}
\]

where \(\mathbf{E}^{(k)}\) are the node embeddings at the \(k\)-th layer, \(\mathbf{E}^{(0)} = \mathbf{X}\) is the input node feature matrix, \(\mathbf{W}^{(k)}\) is a learnable weight matrix at the \(k\)-th layer, \(\hat{\mathbf{A}} = \mathbf{A} + \mathbf{I}\) is the augmented adjacency matrix with self-loops (I is the identity matrix), and \(\mathbf{D}\) is the corresponding augmented degree matrix. Finally, \(\sigma(\cdot)\) is the non-linear ReLU operation as an activation function, i.e., \(\sigma(x) = \max(0, x)\).

**Graph Summarization**

Given an input graph \(\mathcal{G} = (\mathcal{V}, \mathcal{E})\) with \(n = |\mathcal{V}|\) nodes, graph summarization aims to find a smaller summary graph \(\mathcal{G}_s = (\mathcal{V}_s, \mathcal{E}_s)\) (with \(n_s = |\mathcal{V}_s|\) nodes) that preserves the structural information of the original graph. The supernode set \(\mathcal{V}_s\) forms a partition of the original node set \(\mathcal{V}\) such that every node \(v \in \mathcal{V}\) belongs to exactly one supernode \(S \in \mathcal{V}_s\). The supernodes are connected via superedges \(\mathcal{E}_s\), which are weighted by the sum of original edges between the constituent nodes. That is, superedge \(\mathbf{A}_s(p, q)\) between supernodes \(S_p, S_q\) is defined as:

\[
\mathbf{A}_s(p, q) := \sum_{v_i \in S_p} \sum_{v_j \in S_q} \mathbf{A}(i, j).
\]

The adjacency matrix of the summary graph can be formulated using a membership matrix \(\mathbf{P} \in \mathbb{R}^{n_s \times n}\) as: \(\mathbf{A}_s = \mathbf{P} \mathbf{A} \mathbf{P}^T\), where

\[
\mathbf{P}(p, i) = \begin{cases} 
1 & \text{if } v_i \in S_p \\
0 & \text{otherwise}.
\end{cases}
\]

Given the summary graph \(\mathcal{G}_s\), the original graph \(\mathcal{G}\) can be approximated with the reconstructed graphs \(\mathcal{G}_r\) with adjacency matrix \(\mathbf{A}_r\) defined as:

\[
\mathbf{A}_r = \mathbf{Q} \mathbf{A} \mathbf{Q}^T, \tag{4}
\]

where \(\mathbf{Q} \in \mathbb{R}^{n \times n_s}\) is the reconstruction matrix. Note that \(\mathbf{A}_r\) can be seen as a low-rank approximation of the original \(\mathbf{A}\).

A natural way to formulate a summarization approach is by minimizing the difference between \(\mathbf{A}\) and \(\mathbf{A}_r\), so summarization and reconstruction are closely related. Specifically, in this work, we consider the configuration-based reconstruction scheme (Zhou et al. 2021), which adopts the configuration-based model as null model. In that case, reconstructed edge weights are proportional to degrees of endpoints. \(\mathbf{Q}\) and \(\mathbf{A}_r\) are defined as:

\[
\mathbf{Q}(i, p) = \begin{cases} 
\frac{d_p}{d_{p}^{(s)}} & \text{if } v_i \in S_p \\
0 & \text{otherwise}
\end{cases}, \tag{5}
\]

\[
\mathbf{A}_r(i, j) = \frac{d_{p}^{(s)}}{d_{p}^{(s)}} \mathbf{A}_s(p, q) \frac{d_q}{d_q^{(s)}} \quad v_i \in S_p, v_j \in S_q. \tag{6}
\]

\(d_p^{(s)}\) is defined as \(d_p^{(s)} = \sum_{v_i \in S_p} d_i\).

We will later show that, this reconstruction scheme plays an important role in our theoretical analysis.

**Proposed methods**

Given the problem definition, we now give our theoretical analysis and propose GELSUMM framework.

**Theory**

In this section, we theoretically reveal the mechanism behind the approach of learning embeddings on summary graphs. In short, we show that running three embedding methods (DeepWalk, LINE and GCN) on a summary graph is equivalent to running them on an approximate configuration-based reconstructed graph.

**Approximating kernel matrices**  We begin our derivation with the following kernel matrix, which appears in three node embedding learning methods DeepWalk, LINE and GCN. By comparing the core function of DeepWalk, LINE and GCN, we observe that a common kernel matrix can be extracted as:

**Definition 1** (Kernel Matrix). DeepWalk, LINE, and GCN are based on the following generalized kernel matrix:

\[
\mathcal{K}(\mathcal{G}) := (\mathbf{D}^{-c} \mathbf{A} \mathbf{D}^{-1+c})^\tau \mathbf{D}^{1-2c}, \tag{7}
\]

where \(0 \leq c \leq 1\) and \(\tau\) is a positive integer. \(\mathbf{A}\) and \(\mathbf{D}\) are adjacency matrix and degree matrix of \(\mathcal{G}\) respectively.

**Remark.** For \(c = 1\), we get \(\mathcal{K}(\mathcal{G}) = (\mathbf{D}^{-1} \mathbf{A})^\tau \mathbf{D}^{-1}\) appears in DeepWalk and LINE (see Eq. (1),(2)). For \(c = \frac{1}{2}\), we get \(\mathcal{K}(\mathcal{G}) = (\mathbf{D}^{-\frac{1}{2}} \mathbf{A} \mathbf{D}^{-\frac{1}{2}})^\tau\) appears in GCN (see Eq. (3)).

As we show next in Lemma 1, under the configuration-based reconstruction scheme (see Eq. (5) and (6)), this kernel matrix on the original graph, \(\mathcal{K}(\mathcal{G})\), can be approximated with the same kernel matrix on the summary graph, \(\mathcal{K}(\mathcal{G}_s)\), in a closed form.
**Lemma 1.** Given $A_r$ (reconstructed by the configuration-based scheme) as a low-rank approximation of the original adjacency matrix $A$, the kernel matrix of $G$ can be approximated by that of $G_s$:

$$K(G) \approx (D^{-c}A,D^{-1+c})^T D^{1-2c}$$

$$= R (D_s^{-c}A_s D_s^{-1+c})^T D_s^{1-2c} R^T$$

(8)

where $R \in \mathbb{R}^{n \times n_s}$ is the restoration matrix:

$$R(i,p) = \begin{cases} \frac{d_i}{d_{S_p}}(1-c) & \text{if } v_i \in S_p \\ 0 & \text{otherwise} \end{cases}$$

which is closely related to the configuration-based reconstruction matrix $Q$ given in (5).

**Proof.** See appendix.

From this general form of the reconstruction matrix, we obtain specific cases for DeepWalk, LINE, and GCN in the next corollaries.

**Corollary 1.** Based on Dfn. 1 of the kernel matrix, $c = 1$ corresponds to DeepWalk and LINE. In this case, (8) becomes:

$$K(G) \approx (D^{-1}A)^T D^{-1} \approx (D^{-1}A_r)^T D^{-1}$$

$$= R (D_s^{-1}A_s)^T D_s^{-1} R^T,$$

where

$$R(i,p) = \begin{cases} 1 & \text{if } v_i \in S_p \\ 0 & \text{otherwise} \end{cases}$$

(10)

**Corollary 2.** Based on Dfn. 1 of the kernel matrix, $c = \frac{1}{2}$ corresponds to GCN. In this case, (8) becomes:

$$K(G) \approx D^{-\frac{1}{2}} A D^{-\frac{1}{2}} \approx D^{-\frac{1}{2}} A_r D^{-\frac{1}{2}}$$

$$= R \left( D_s^{-\frac{1}{2}} A_s D_s^{-\frac{1}{2}} \right) R^T,$$

(11)

where

$$R(i,p) = \begin{cases} \frac{d_i}{d_{S_p}} & \text{if } v_i \in S_p \\ 0 & \text{otherwise} \end{cases}$$

(12)

Note that $R^T R = I$ and $R^\dagger = R^T$ ($R^\dagger$ denotes the pseudo-inverse of $R$) in Corollary 2, which is important in our analysis of GCN.

**Approximating DeepWalk / LINE.** Based on Corollary 1, we now discuss how to approximate the DeepWalk and LINE node embeddings for the original nodes. Since LINE is a special case of DeepWalk, we focus on the former; similar conclusions can be easily drawn for LINE.

**Theorem 1.** Embeddings learned by DeepWalk on the original graph $G$, $E$, can be approximated by embeddings learned by DeepWalk on the summary graph $G_s$, $E_s$, using the restoration matrix $R$ in (10), i.e.,

$$E \approx R E_s$$

(13)

**Proof.** See appendix.

According to Theorem 1 and the definition of $R$ matrix (if $v_i \in S_p$), we can conclude that nodes in the same supernode get the same embeddings after the restoration. This approach, in exactly the way how related works (including HARP, MILE and GraphZoom) restore the embeddings. Thus, Theorem 1 provides a theoretical interpretation for the restoration step of existing methods.

**Approximating GCN.** Given the embeddings $E_s$ learned by GCN on the summary graph which are usually the output of the last convolution layer, i.e., $E_s = E_s^{(K)}$, we can approximate original node embeddings $E$, as stated in the following theorem.

**Theorem 2.** Embeddings learned by GCN on the original graph $G$ can be approximated by embeddings learned by GCN on the summary graph $G_s$ with initial features $X_s := R^T X$, using the restoration matrix $R$ defined in (12), in a least-square approximation perspective:

$$E \approx R E_s$$

(14)

And suppose that the GCN model on the summary graph and the original graph share the same model parameter, then the reconstruction error is bounded by:

$$\|E - R E_s\| \leq C \sqrt{2 KL(A||A_r)} \left( \prod_{i=0}^{K-1} \left\|W(i)\right\| \right) \|E(0)\|$$

(15)

where $C = \sqrt{\sum_i (1 - \lambda_i)^2}$ is a constant only depends on the input graph. $\lambda_i$ is the $i$-th eigenvalue of the normalized Laplacian matrix $L$ (defined as $L = I - D^{-\frac{1}{2}} A D^{-\frac{1}{2}}$). And $KL(A||A_r)$ is the KL-divergence between $A$ and $A_r$ (see Appendix), which is included in the objective function of summarization algorithm we use.

**Proof.** See appendix.

**Algorithms.**

From the theory we develop above, learning embeddings on a summary graph with restoration is equivalent to learning embeddings on a graph reconstructed by the configuration model. Thus, the closer the original graph and the reconstructed one is, the better the embeddings are. This fact motivates us to consider the configuration-based reconstruction scheme in the summarization step.

Specifically, we use a graph summarization method DPGS (Zhou et al. 2021) which utilize the configuration-based reconstruction method. It is based on MDL (Minimum Description Length (Rissanen 1978)) principle and aims to minimize the total description length of both the model cost and configuration-based reconstruction error. More details of the summarization algorithm are provided in appendix.

Subsequently, we propose GELSUMM, which aims to learn embeddings for large graphs by summarization. It consists of four steps:

1. **(S1) Summarization.** First, given the input graph $G$, we use a configuration-based summarization method (Zhou et al. 2021) to obtain a summary graph $G_s$.
Table 3: Node classification results (DeepWalk & LINE). Average running times and accuracy scores over 10 runs are reported. “orig” represents the results on original graphs. r stands for the relative node size. The running time includes the summarization time, the embedding learning time and the refinement time. It can be observed that GELSUMM-DeepWalk and GELSUMM-LINE obtain better results than original DeepWalk and LINE using much less time.

<table>
<thead>
<tr>
<th></th>
<th>DeepWalk</th>
<th>LINE</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>orig</td>
<td>r=0.6</td>
</tr>
<tr>
<td>Cora</td>
<td>acc (%)</td>
<td>72.11</td>
</tr>
<tr>
<td>Citeseer</td>
<td>acc (%)</td>
<td>46.24</td>
</tr>
<tr>
<td></td>
<td>time (secs)</td>
<td>111.022</td>
</tr>
<tr>
<td>Pubmed</td>
<td>acc (%)</td>
<td>72.35</td>
</tr>
<tr>
<td></td>
<td>time (secs)</td>
<td>875.838</td>
</tr>
<tr>
<td>Flickr</td>
<td>acc (%)</td>
<td>53.19</td>
</tr>
<tr>
<td></td>
<td>time (secs)</td>
<td>6142.00</td>
</tr>
</tbody>
</table>

Table 4: Dataset statistics.

| Dataset   | |V| | |E| | Type |
|-----------|---|---|---|---|---|
| Cora      | 2,708 | 5,278 | Citation |
| Citeseer  | 3,327 | 4,552 | Citation |
| Pubmed    | 19,717 | 44,324 | Citation |
| Flickr    | 89,250 | 899,756 | Social |
| Reddit    | 232,965 | 11,606,919 | Social |
| Amazon2M  | 2,449,029 | 61,859,076 | Co-Purchasing |

Algorithm 1: GELSUMM-DeepWalk/LINE/GCN

Input: Input graph $G$, power of filter $k$, embedding method $M$

Output: Original embeddings $E$

1. $G_s ↣$ Run the configuration-based summarization on $G$
2. $E_s ↣ M(G_s)$
3. $E ↣ R · E_s$.
4. $E ↣ (\tilde{D}^{-\frac{1}{2}} A \tilde{D}^{-\frac{1}{2}})^k E$.
5. return $E$

- **(S2) Embedding Learning.** Next, the chosen embedding learning method $M$ (DeepWalk, LINE or GCN), is employed on the summary graph $G_s$ to obtain supernode embeddings $E_s = M(G_s)$.
- **(S3) Restoration.** Then, we restore the embeddings of the original nodes, $E$, from $E_s$ using the restoration matrix $R$, i.e., $E ↣ R · E_s$. The definition of $R$ varies depending on the method $M$ used in step (S2).
- **(S4) Refinement.** Finally, in order to further refine the restored embeddings $E$, we apply $k$ times a low-pass smoothing filter, $\tilde{D}^{-\frac{1}{2}} A \tilde{D}^{-\frac{1}{2}}$ to smooth the restored embedding.

Based on the steps described in previous sections and Theorems 1 and 2, we propose three corresponding algorithms: GELSUMM-DeepWalk, GELSUMM-LINE, and GELSUMM-GCN, described in Algorithm 1.

First, a summary graph $G_s$ is obtained by the configuration-based graph summarization algorithm (S1), and then fed into a base embedding learning algorithm to learn embeddings $E_s$ (S2). Original embeddings are then restored from $E_s$ using corresponding matrix $R$ (S3), and further refined by a smoothing operator (S4). The refinement step is efficient and its time cost is ignorable compared to the embedding learning time. Note that in GELSUMM-GCN, self-loops are added to $G$, since GCN uses augmented adjacency matrix $\tilde{A} = A + I$.

To learn supernode embeddings using GCN, we need to assign labels to supernodes. In this paper, we adopt a majority-vote method that assigns the dominating label with the maximum count, i.e., $\text{label}(S_p) = \arg \max_i |\{v_i | v_i \in S_p, \text{label}(v_i) = l\}|$. That is, the label of a supernode is the dominating label of its constituent nodes.

### Experiments

In this section, we conduct extensive experiments to answer the following questions:

- **Q1 Effectiveness of GELSUMM-DeepWalk/LINE**: Is GELSUMM-DeepWalk and GELSUMM-LINE able to learn high-quality embeddings efficiently from summary graphs compared to baselines?
- **Q2 Effectiveness of GELSUMM-GCN**: Is GELSUMM-GCN able to learn high-quality embeddings for semi-supervised node classification task?
- **Q3 Scalability**: Is GELSUMM scalable and can be applied to large graphs with up to 2 million nodes?

### Experiment Setup

All experiments are conducted on a machine with a 2.40 GHz Intel Xeon E5-2640 CPU, a Tesla K80 GPU, and 128 GB RAM. For the summarization algorithm, we use the source code released by (Zhou et al. 2021). We implement DeepWalk in Python and use LINE’s code released by the authors\(^2\).

\(^1\tilde{A} = A + I\) is the augmented adjacency matrix and $\tilde{D}$ is the corresponding degree matrix.

\(^2\)https://github.com/tangjianpku/LINE
Datasets. Datasets used in our experiments contain three citation networks, two social networks, and one co-purchasing network. The statistics of them are listed in Table 4. In the first three citation networks, nodes represent documents and edges are citation among them. Bag-of-word vectors are used as node features. For Flickr dataset, each node is an image on the Flickr website, and if two images share some properties, there is an link between them. For Reddit dataset, each node is a post on Reddit, and edges represent co-comment by the same user. The largest dataset Amazon2M, a product network linked by co-purchase behaviors, is tested for scalability. We use the widely-used dataset splits as in (Kipf and Welling 2017; Hamilton, Ying, and Leskovec 2017).

Q1 Effectiveness of GELSUMM-DeepWalk/LINE

In this experiment, we evaluate GELSUMM-DeepWalk & GELSUMM-LINE on two downstream tasks: node classification and link prediction (placed in appendix due to space limit). Four datasets Cora, Citeseer, Pubmed and Flickr are used in this experiments. Reddit is not included since running representation learning methods on it raises an OOM(out of memory) error on our machine.

Node Classification In node classification task, we summarize the original graph into several summary graphs with different relative sizes (from 0.6 to 0.2), and run DeepWalk/LINE on summary graphs. The restored embeddings are used to train a Logistic Regression classifier for node classification.

Hyperparameters: Following (Perozzi, Al-Rfou, and Skiena 2014), in DeepWalk, we sample $\cdot |V|$ random walk sequences. The walk length is set to 80 and the window size is set to 10. In LINE, we set the number of negative samples to 5. The embedding dimension is set to 128 for all methods. For GELSUMM-DeepWalk and GELSUMM-LINE, $k$ is set to 2. We give analysis of the effect of paramter $k$ in the appendix.

Results. The experimental results (mean accuracy score and running time of 10 runs) are shown in Table 3. We observe that GELSUMM-DeepWalk and GELSUMM-LINE can match or outperform original DeepWalk and LINE in all datasets. For example, on Citeseer dataset, GELSUMM-DeepWalk get a 2.84% performance increase using 8.78× less time, and GELSUMM-LINE get a 4.77% accuracy increase using 5.03× less time.

Comparison with existing methods. We also compare our GELSUMM framework to the state-of-the-art methods HARP, MILE and GraphZoom. In this experiment, we use node classification task as the base task and choose DeepWalk as the base embedding learning method. For baseline methods, we use the source codes released by the authors off-the-shelf. For MILE and GraphZoom, input networks are summarized at different coarsening levels to produce summary graph with similar sizes. We provide sizes of summary graphs obtained by baseline methods in the appendix. For our GELSUMM, the relative sizes are $[0.6, 0.4, 0.2, 0.1]$ to match the summary sizes.

Running time and classification accuracy (average of 10 runs) are displayed in Figure 1. On all datasets, HARP gets worse performance than that on original graphs. MILE keeps relative good performance when the coarsening level is low, but the performance drops when the coarsening level goes higher. Our GELSUMM can obtain promising results with much less time. Compared to GraphZoom, our GELSUMM gets higher performance on Cora dataset, and gets comparable performance using similar time on Citeseer, Pubmed and Flickr datasets. Furthermore, compared to the baselines, our GELSUMM provides more theoretical analysis and interpretation.

Q2 Effectiveness of GELSUMM-GCN:

In this experiment, we test GELSUMM-GCN on semi-supervised node classification task. We train GCN on summary graphs using majority-label assignment. That is, assigning the supernode label as the maximum-count label within it. The restored node embeddings are used for classification after refinement.

To validate the effect of our theoretically-grounded restoration method, we also report the classification accuracies.
Figure 1: Comparison with the baselines. “orig” represents the results on original graphs. Our algorithm obtains promising performance using much less time.

Figure 2: Embeddings of part of nodes from 4 classes learned via GELSUMM on Amazon2M datasets with summarization ratio 0.2 and 0.1. Vertex colors represent its category. Accuracies are given in the captions. Embeddings are projected into 2-dim space via t-SNE. Representations of different classes are well-separated.

Q3 Scalability

To test the scalability of our GELSUMM framework, we test on a large Amazon2M dataset, which contains over 2.4 million nodes. Running node embeddings methods on the original graph is not applicable due to memory limit. We apply GELSUMM on summary graphs with summarization ratio 0.2 and 0.1 respectively to get the original embeddings. To show the quality of learned embeddings, we randomly choose some nodes from 4 classes and visualize their embeddings via t-SNE, which is shown in Figure 2. It can be seen that nodes from different classes are well-separated, demonstrating that our GELSUMM can learn matchable embeddings of original nodes in large-scale graphs with limited memory and less time.

Conclusion

In this work, we propose a novel and well-formulated framework, GELSUMM, for learning node embeddings using a summarization approach. By theoretically showing that a specific form of kernel matrix—which appears in DeepWalk, LINE and GCN—can be approximated under a configuration-based reconstruction scheme on the summary graph, we propose three algorithms to learn node embeddings on the summary graph with a closed-form solution. Our study helps understand the empirical success of current related methods, and provides theoretical insights for future works on this problem. Future directions include introducing more embedding methods into our framework, and extend to more graph mining tasks.

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References

and applications. IEEE Transactions on Knowledge and Data Engineering, 30(9): 1616–1637.


Deng, C.; Zhao, Z.; Wang, Y.; Zhang, Z.; and Feng, Z. 2020. GraphZoom: A Multi-level Spectral Approach for Accurate and Scalable Graph Embedding. In ICLR.


Fu, G.; Hou, C.; and Yao, X. 2019. Learning topological representation for networks via hierarchical sampling. In 2019 International Joint Conference on Neural Networks (IJCNN), 1–8. IEEE.


Kipf, T. N.; and Welling, M. 2017. Semi-Supervised Classification with Graph Convolutional Networks. In ICLR.


NT, H.; and Maehara, T. 2019. Revisiting Graph Neural Networks: All We Have is Low-Pass Filters. arXiv:1905.0550.


