NEURAL APPROXIMATION OF EXTENDED PERSISTENT HOMOLOGY ON GRAPHS

Zuoyu Yan  
Wangxuan Institute of Computer Technology  
Peking University  
yanzuoyu3@pku.edu.cn

Tengfei Ma  
T. J. Watson Research Center  
IBM  
Tengfei.Ma1@ibm.com

Liangcai Gao  
Wangxuan Institute of Computer Technology  
Peking University  
glc@pku.edu.cn

Zhi Tang  
Wangxuan Institute of Computer Technology  
Peking University  
tangzhi@pku.edu.cn

Yusu Wang  
Halıcıoğlu Data Science Institute  
University of California, San Diego  
yusuwang@ucsd.edu

Chao Chen  
Department of Biomedical Informatics  
Stony Brook University  
chao.chen.1@stonybrook.edu

ABSTRACT

Persistent homology is a widely used theory in topological data analysis. In the context of graph learning, topological features based on persistent homology have been used to capture potentially high-order structural information so as to augment existing graph neural network methods. However, computing extended persistent homology summaries remains slow for large and dense graphs. Inspired by recent success in neural algorithmic reasoning, we propose a novel learning method to compute extended persistence diagrams on graphs. The proposed neural network aims to simulate a specific algorithm and learns to compute extended persistence diagrams for new graphs efficiently. Experiments on approximating extended persistence diagrams and several downstream graph representation learning tasks demonstrate the effectiveness of our method. Our method is also efficient; on large and dense graphs, we accelerate the computation by nearly 100 times.

1 INTRODUCTION

In recent years, much effort has been made to improve the expressiveness of graph neural networks (GNNs). Among these methods, a widely used approach is to explicitly inject other high order information, such as graph topological/structural information, into the GNN models (You et al., 2019; Li et al., 2020). To this end, persistent homology (Edelsbrunner et al., 2000; Edelsbrunner & Harer, 2010), which captures topological structures (e.g., connected components and loops) and encodes them in a summary called persistence diagram (PD), has been injected to machine learning pipelines for various graph learning tasks (Zhao & Wang, 2019; Zhao et al., 2020; Hofer et al., 2020; Carrière et al., 2020; Chen et al., 2021; Yan et al., 2021). In particular, it has been found helpful to use the so-called extended persistence diagrams (EPDs) (Cohen-Steiner et al., 2009), which contain richer information than the standard PDs.

Despite the strong learning power of PDs and EPDs, their computation remains a bottleneck in graph learning. In situations such as node classification (Zhao et al., 2020) or link prediction (Yan et al., 2021), one has to compute EPDs on vicinity graphs generated around all the nodes or all possible edges in the input graph. This can be prohibitive especially for large and dense graphs.

Our goal is to develop a learning-based framework to estimate the EPDs for graphs efficiently. Recent works on algorithmic learning on graphs (Veličković et al., 2019; Xhonneux et al., 2021) showed

* Correspondence to Chao Chen, Yusu Wang, and Liangcai Gao
that sequential algorithms such as Dijkstra and Depth-First Search (DFS) can be approximated by GNNs. Inspired by these works, we rewrite the algorithm to compute EPDs into a sequential-like form and propose a novel neural framework to estimate the EPDs whose architecture is aligned with this algorithm. Experiments show that on large and dense graphs, the proposed framework is much faster than the direct computation of EPDs.

Compared with the sequential algorithms proposed in [Xhonneux et al., 2021], the algorithm to compute EPDs requires extra steps. To address these challenges, we propose several modules to approximate these steps. Using these modules, we empirically show that our method achieves a satisfying approximation quality of EPDs on different graphs. To further evaluate the effectiveness of our framework, we perform experiments on two downstream graph representation learning tasks: node classification and link prediction. We show that the deep learning models using the predicted EPDs perform comparably with the architectures using the ground-truth EPDs. In other words, the approximated EPDs do not lose learning power.

To the best of our knowledge, we are the first to directly estimate EPDs on graphs with deep learning models. We note that our method is fundamentally different from previous works learning to directly approximate persistence images, a vectorization of persistence diagrams (Som et al., 2020; Montufar et al., 2020). These methods fail to simulate the computation algorithm closely. Furthermore, their training loss does not respect the special metric of EPDs. Therefore, their output is not satisfying in both approximation quality and learning power. Finally, we observe that our model can be easily transferred to unseen graphs. This is encouraging as we may now generalize topological computation to various challenging real-world graphs without extra effort.

For detailed reading, we refer the readers to a more complete version of the paper: [https://arxiv.org/pdf/2201.12032.pdf](https://arxiv.org/pdf/2201.12032.pdf)

2 Related Works

Learning with Persistent Homology. Based on the theory of algebraic topology [Munkres, 2018], persistent homology [Edelsbrunner et al., 2000; Edelsbrunner & Harer, 2010] extends the classical notion of homology and can capture the topological structures (e.g., loops, connected components) of the input data in a robust (Cohen-Steiner et al., 2007) manner. It has already been combined with various deep learning methods including kernel machines [Reininghaus et al., 2015; Kusano et al., 2016; Carriere et al., 2017], convolutional neural networks [Hofer et al., 2017; Hu et al., 2019; Wang et al., 2020; Zheng et al., 2021], transformers [Zeng et al., 2021], connectivity loss [Chen et al., 2019; Hofer et al., 2019], and graph neural networks [Zhao et al., 2020; Chen et al., 2021; Yan et al., 2021; Zhao & Wang, 2019; Hofer et al., 2020; Carrière et al., 2020].

Neural Algorithm Execution. Many works have studied neural execution in different domains before [Zaremba & Sutskever, 2014; Kaiser & Sutskever, 2015; Kurach et al., 2015; Reed & De Freitas, 2015; Santoro et al., 2018; Yan et al., 2020]. With the rapid development of GNNs in graph represen-
tation learning, learning graph algorithms with GNNs has attracted researchers’ attention \cite{velickovic2019graph,velickovic2020learn,xhonneux2021learning}. These works exploit GNNs to approximate certain classes of graph algorithms, such as parallel algorithms and sequential algorithms.

3 Extended Persistent Homology and Learning Extended Persistence Diagrams

Extended Persistent Homology. Extended persistent homology captures 0-dimensional (connected components) and 1-dimensional (loops) topological structures and summarizes their topological information into the so-called extended persistence diagram (EPD), which is a planar multiset of points, each of which \((b, d)\) corresponds to the information of some homological feature (i.e., components, loops, and their higher dimensional analogs). For example, for the input graph in Figure 1 (a), its output EPD is shown in Figure 1 (b). Further details around extended persistent homology are available in Section A.1 in the appendix.

Computing Extended Persistence Diagrams. \cite{xhonneux2021learning} point out that sequential algorithms (e.g., Dijkstra) can be approximated by GNNs. Inspired by their idea, we rewrite the algorithms to compute EPDs in a sequential-like form that is easier for a GNN-like architecture to “simulate”. The algorithms are listed in the Appendix.

Note that when we perform the standard persistence algorithm via matrix reduction, every simplex in the input simplicial complex will either be a creator (indicating that adding this simplex will create a new family of homology classes/topological features) or a destroyer (indicating that adding this simplex will destroy some existing homology classes/topological features). In the end, the persistence algorithm will pair up these creators and destroyers, and their function values give rise to the persistence points in the resulting persistence diagrams. In our context where the input is a graph which can be viewed as a 1-dimensional simplicial complex (consisting of 0-simplices/vertices \(V\) and 1-simplices/edges \(E\)), the persistence algorithms will pair up these simplices. In other words, to compute EPDs, we will simply find the pairing partners for all edges.

Description of the Sequential algorithm. Here we focus on describing the algorithm to compute 1D EPD. Further details are available in the Appendix. Our sequential algorithm, as shown in the Appendix, exploits the observations from \cite{agarwal2006computational}. Specifically, consider a vertex \(v_i \in V\), and suppose there are \(k\) edges \(e_{i1}, \ldots, e_{ik}\) incident to \(v_i\) with function values higher than \(v_i\) (i.e., the function value of the other endpoint of these edges is higher than \(f(v_i)\)). See Figure 1 (c) for an illustration. For node \(u_1, k = 3\), and the three edges are \(u_1u_3, u_1u_4,\) and \(u_1u_6\). Now imagine we put each such edge in a different component \(C_j, j \in [1, k]\) – we call this upper-edges splitting operation – and start to sweep the graph \(G\) in increasing values of \(a\) but starting at \(f(u_1)\). Then, the first time any of two such components merge will give rise to a new persistence point in the 1D EPD. For instance, in Figure 1 (c), \(C_{14}\) and \(C_{13}\) first merge at \(u_4\), and this will give rise to the brown loop in Figure 1 (a) with \((t_4, t_2)\) as its persistence point. Intuitively, this pairing captures the so-called thinnest cycle basis \cite{agarwal2006computational,cohen2009extending}.

Hence to compute the extended persistence pairing induced by \(v_i\), we can call Algorithm 3 to identify the first time when components containing \(C_i\) s are merged. This can be achieved using the union-find-like data structure to track the components. The main difference from the standard union-find data structure is that as we merge components, each component needs to be represented by the minimum (the vertex in this component with the smallest function value) which is not required in a standard union-find data structure. (The same holds for the algorithm to compute the 0D PD.)

Learning Extended Persistence Diagrams. Considering that every edge in the input graph will give rise to either a 0D ordinary persistence point or a 1D extended persistence point, we transfer the learning of EPDs into a link prediction problem. Specifically, our base architecture follows standard link-prediction architectures \cite{chami2019link,yan2021deeper}. (1) For a input graph, we first use a specially designed GNN model which later we call PDGNN to obtain the node embedding for all these vertices. (2) Subsequently, a MLP (Multi-layer perceptron) is applied to the node embeddings to obtain the persistent pairing information for each edge.

Our specially designed GNN for estimating persistence diagrams, is called PDGNN (Persistence Diagram Graph Neural Network). Compared with the Sequential algorithms proposed in \cite{xhonneux2021learning}, extended persistence diagrams need extra care: (1) the Find-Root algorithm needs
Table 1: Time evaluation on different datasets (seconds)

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Cora</th>
<th>Citeseer</th>
<th>PubMed</th>
<th>Photo</th>
<th>Computers</th>
<th>CS</th>
<th>Physics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Avg. N/E</td>
<td>38/103</td>
<td>16/43</td>
<td>61/190</td>
<td>797/1602</td>
<td>1879/47477</td>
<td>97/431</td>
<td>193/1315</td>
</tr>
<tr>
<td>Fast (Yan et al., 2021)</td>
<td>0.95</td>
<td>0.39</td>
<td>2.15</td>
<td>362.60</td>
<td>1195.66</td>
<td>5.72</td>
<td>24.14</td>
</tr>
<tr>
<td>Gudhi (The GUDHI Project, 2015)</td>
<td>0.44</td>
<td>0.21</td>
<td>1.00</td>
<td>583.55</td>
<td>8585.50</td>
<td>3.00</td>
<td>26.58</td>
</tr>
<tr>
<td>Ours</td>
<td>5.21</td>
<td>4.72</td>
<td>4.78</td>
<td>6.67</td>
<td>7.32</td>
<td>5.18</td>
<td>5.42</td>
</tr>
</tbody>
</table>

to return the minimum of the component, (2) edge operations such as upper-edge splitting. Our PDGNN modifies standard GNNs by (1) implementing the root-finding process by a concatenation of sum aggregation and min aggregation as our message aggregation function; (2) incorporating edge operations such as the upper-edge splitting operation with edge features and edge attention. Further details are available in the Appendix.

4 EXPERIMENTS

In this paper, we thoroughly evaluate the proposed model from 3 different perspectives: approximation quality, transferability, and algorithm efficiency. Due to the page limit, we put the evaluation on approximation quality and transferability to the Appendix, and briefly introduce the results in the following paragraph. As for the evaluation on algorithm efficiency, experiments demonstrate that the proposed method is much faster than the original algorithm, especially on large and dense graphs (shown in Table 1).

**Datasets.** To compute EPDs, we need to set the input graphs and the filter functions. Following the settings of existing state-of-the-art models (Zhao et al., 2020; Yan et al., 2021), for a given graph \( G = (V, E) \), we extract the \( k \)-hop neighborhoods of all the vertices, and extract \(|V|\) vicinity graphs as input graphs. In terms of filter functions, we introduce Ollivier-Ricci curvature (Ni et al., 2018), heat kernel signature, and the node degree as the filter function. The datasets include (1) citation networks including Cora, Citeseer, and PubMed (Sen et al., 2008); (2) Amazon shopping datasets including Photo and Computers (Shchur et al., 2018); (3) coauthor datasets including CS and Physics (Shchur et al., 2018). Details are available in the Appendix.

**Approximation Quality.** In Section A.4.1, we evaluate the approximation error between the predicted diagram and the ground truth diagram and show that the prediction is very close to the original diagram. We also add ablation study to show the effectiveness of all the proposed modules. To understand how much does the approximation error influence downstream tasks, in Section A.4.2, we evaluate the learning power of the predicted diagrams through 2 downstream graph representation learning tasks: node classification and link prediction. We observe that the model using the predicted diagrams performs comparably with the model using the ground truth diagrams. From the two evaluation, we can safely conclude that the predicted diagram is a wonderful substitution of the original EPD in terms of both approximation error and learning power.

**Transferability.** One appealing feature of our method is its transferability. Training on one graph, our algorithm can still approximate EPDs well on another graph. This makes it possible to apply the computationally expensive topological features to a wide spectrum of real-world graphs; we can apply a pre-trained model to large and dense graphs, on which direct EPD computation is infeasible.

We prove the transferability empirically. In Table 4 we adopt the model pre-trained on Photo to predict the EPDs of CS and Physics, and achieve good approximation performance. We provide comprehensive experiments to evaluate the transferability of our methods. See Table 6 and Table 7 in the Appendix for the approximation quality and learning power evaluation of transferred models.

**Algorithm Efficiency.** For a fair and complete comparison, we compare with algorithms from Gudhi (The GUDHI Project, 2015) and from (Yan et al., 2021). We select the first 1000 nodes from Cora, Citeseer, PubMed, Photo, Computers, CS, Physics, and then extract their 2-hop neighborhoods as the input vicinity graphs. We then compute the EPDs and report the time (seconds) used to infer these diagrams.

We list the average nodes and edges of these vicinity graphs in the first line of Table 1. As shown in Table 1, although our model is slower on small datasets like Cora or Citeseer, it is much faster on large and dense datasets. Therefore we can simply use the original algorithm to compute the extended persistence diagrams on small graphs, and use our model to execute extended persistence diagrams
on large graphs. The model can be applied to various graph representation learning works based on persistent homology.

5 CONCLUSION

We propose a learning algorithm to approximate EPDs. Inspired by recent success on neural algorithm execution, we propose a novel GNN with different technical contributions to simulate the computation of EPDs. Experiments show that our method achieves satisfying approximation quality and learning power while being significantly faster than the original algorithm, especially on large and dense graphs. Another strength of our method is the transferability: training on one graph, our algorithm can still approximate EPDs well on another graph. This makes it possible to apply the computationally expensive topological features to a wide spectrum of real-world graphs.

REFERENCES


A APPENDIX

In the Appendix, we provide (1) Introduction of extended persistent homology and its computation; (2) additional experimental details, including introduction on the datasets, and the experimental settings; (3) further experiments, including the evaluation on approximation quality, the evaluation on transferability, and the influence of training samples; (4) experiments on graph classification datasets.

A.1 EXTENDED PERSISTENT HOMOLOGY

We briefly introduce extended persistent homology and refer the readers to (Cohen-Steiner et al., 2009) and (Edelsbrunner & Harer, 2010) for more details.

Ordinary Persistent Homology. Persistent homology captures 0-dimensional (connected components) and 1-dimensional (loops) topological structures and measures their saliency via a scalar function called filter function. Given a input graph \( G = (V, E) \), with node set \( V \) and edge set \( E \), we call all the nodes and edges simplices. Denote by \( X = V \cup E \) the set of all simplices. We define a filter function on all simplices, \( f : X \to \mathbb{R} \). Often, \( f \) is induced by a node-valued function (e.g., node degrees), and further defined on edges as \( f(uv) = \max(f(u), f(v)) \).

Denote by \( X_a \) the sublevel set of \( X \), consisting of simplices whose filter function values \( \leq a \), \( X_a = \{ x \in X | f(x) \leq a \} \). As the threshold value \( a \) increases from \(-\infty \) to \( \infty \), we obtain a sequence of growing spaces, called an ascending filtration of \( X \): \( \emptyset = X_{-\infty} \subseteq \ldots \subseteq X_{\infty} = X \). As \( X_a \) increases from \( \emptyset \) to \( X \), new topological structures gradually appear (born) and disappear (die). For instance, the blue square persistence point at \((t_2, t_3)\) in Figure 1(b) indicates that the connected component \( u_2 \) appears at \( X_{t_2} \) and is merged with the whole connected component at \( X_{t_3} \).

Applying the homology functor to the filtration, we can more precisely quantify the birth and death of topological features (as captured by homology groups) throughout the filtration, and the output is the so-called persistence diagram (PD), which is a planar multiset of points, each of which \((b, d)\) corresponds to the birth and death time of some homological feature (i.e., components, loops, and their higher dimensional analogs). The lifetime \(|d - b|\) is called the persistence of this feature and intuitively measures its importance w.r.t. the input filtration.

Extended Persistent Homology. In the ordinary persistent homology, topology of the domain (e.g., the graph) will be created at some time (has a birth time), but never dies (i.e., with death time being equal to \( +\infty \)). Hence we cannot capture their importance. In the context of graphs, the importance of loops are not captured via the ordinary persistence. To this end, extended persistence (Cohen-Steiner et al., 2009) introduces a descending filtration: \( \emptyset = X_{\infty} \subseteq \ldots \subseteq X_{-\infty} = X \), where \( X^a = \{ x \in X | f(x) \geq a \} \) is a superlevel set. This induces a sequence of homology groups of the form \( H(X) = H(X, X_{\infty}) \to \cdots \to H(X, X_a) \to \cdots \to H(X, X_{-\infty}) = \emptyset \).

When the input domain is a graph, the 1-dimensional EPD consists of \( \beta_1 \) number of persistent points, capturing the birth and death of independent loop features. Here \( \beta_1 \) is the rank of the first homology group; for a connected graph, it is simply the number of independent loops, \( \beta_1 = |E| - |V| + 1 \). A loop will be created during the ascending filtration, but killed during the descending filtration. The birth and death times of the feature correspond to the threshold value \( a \)'s when these events happen. In general, the death time for such loop feature is smaller than the birth time. For example, the red triangle persistence point in Figure 1(b) denotes that the red cycle in Figure 1(a) appears at \( X_{t_2} \) in the ascending filtration and appears again at \( X_{t_1} \) in the descending filtration.

Finally, persistence diagrams live in an infinite-dimensional space equipped with an appropriate metric structure, such as the so-called \( p \)-th Wasserstein distance, or the bottleneck distance (Cohen-Steiner et al., 2007). There have been many works in the literature to vectorize persistence diagrams for downstream analysis. A popular choice is the persistence image (Adams et al., 2017).

A.1.1 COMPUTING EXTENDED PERSISTENCE DIAGRAMS (EPDs)

In this section, we rewrite the algorithms to compute the EPDs for graphs into a form that is easier for GNNs to simulate as inspired by the neural execution work of (Veličković et al., 2019; Xhonneux 2010).
The algorithm for computing the 0D EPDs is well-known, using the so-called union-find data structure (Edelsbrunner & Harer, 2010; Dey & Wang, 2022). The algorithm for computing the 1D EPDs has not been reported before, however, the idea follows from existing work (Agarwal et al., 2006). For simplicity of presentation, we assume that all vertices have distinct function values $f : V \rightarrow \mathbb{R}$, and our goal is to compute the 0D and 1D EPDs $PD_0$ and $PD_1$.

The neural execution of “sequential” algorithms. Xhonneux et al. (2021) point out that sequential algorithms as shown in Algorithm 1 (e.g., Dijkstra) can be approximated by GNNs.

Inspired by their idea, we rewrite the algorithms to compute EPDs in a sequential-like form that is easier for a GNN-like architecture later to “simulate”. In particular, the algorithm for computing 1D extended persistence is shown in Algorithm 2; the algorithm for computing 0D persistence, which is shown in Algorithm 3, is the standard union-find strategy.

High-level description of sequential algorithms. Note that when we perform the standard persistence algorithm via matrix reduction, every simplex in the input simplicial complex will either be a creator (indicating that adding this simplex will create a new family of homology classes/topological features) or a destroyer (indicating that adding this simplex will destroy some existing homology classes/topological features). In the end, the persistence algorithm will pair up these creators and destroyers, and their function values give rise to the birth and death times in the resulting persistence diagrams. In our context where the input is a graph which can be viewed as a 1-dimensional simplicial complex (consisting of 0-simplices/vertices $V$ and 1-simplices/edges $E$), the persistence algorithms will pair up these simplices.

In particular, for an edge (1-simplex) $e \in E$: (1) it can either serve as a destroyer (killing a connected component), in which case it is paired up with a vertex $v$, giving a persistence pairing $(v, e)$ and a corresponding persistence point $(f(v), f(e))$ in the 0-D PD $PD_0$. For example, the persistence point located at $(t_2, t_3)$ in Figure 1(b) denotes that the edge $u_2u_3$ is paired with $u_2$. Alternatively, the edge could be creating a 1-cycle (intuitively a loop in the graph) and is a creator during the ascending filtration. This loop will ultimately be killed during the descending filtration when we take relative homology $H_1(X, X^e)$, when we sweep past some vertex $w$ (and $a = f(w)$). Hence the algorithm will pair up $(e, w)$ which gives rise to a persistence point $(f(e), f(w))$ in the 1-D EPD $PD_1$.

In other words, to compute $PD_0$ and $PD_1$, we will simply find the pairing partners for all edges. We note that in the literature, when the filtration is induced by a filter function (e.g., Dijkstra) can be approximated by GNNs.

Sequencial algorithm to compute $PD_1$. The algorithm to compute $PD_0$ is similar to Algorithm 3. Therefore, here we just describe the algorithm to compute $PD_1$. It turns out that using the observations from (Agarwal et al., 2006), the following procedure will produce the 1D extended persistence capturing loops. Specifically, consider a vertex $v_i \in V$, and suppose there are $k$ edges $e_{i_1}, \ldots, e_{i_k}$ incident to $v_i$ with function values higher than $v_i$ (i.e., the function value of the other endpoint of these edges is higher than $f(v_i)$). See Figure 1(c) for a simple illustration. For node $u_1$, $k = 3$, and the three edges are $u_1u_3$, $u_1u_4$, and $u_1u_6$. Now imagine we put each such edge in a different component $C_i$, $j \in [1, k]$ – we call this upper-edges splitting operation – and start to sweep the graph $G$ in increasing values of $a$ but starting at $f(v_i)$. Then, the first time any two such components merge will give rise to a new persistence point in the 1D extended persistence diagram. For instance, in Figure 1(c), $C_{i_1}$ and $C_{i_1}$ first merge at $u_4$, and this will give rise to the brown loop in Figure 1(a) with $(t_4, t_5)$ as its persistence point. Intuitively, this pairing captures the so-called thinnest cycle basis (Agarwal et al., 2006; Cohen-Steiner et al., 2009).

Hence to compute the extended persistence pairing induced by $v_i$, we can call Algorithm 3 to identify the first time when components containing $C_i$ s are merged. This can be achieved using the union-find-like data structure to track the components. The main difference from the standard union-find data structure is that as we merge components, each component needs to be represented by the minimum (the vertex in this component with the smallest function value) which is not required in a standard
Algorithm 1 Sequential algorithm

1: **Input**: graph $G = (V, E)$, filter function $f$.
2: Initialise-Nodes($V, f$)
3: $Q = \text{Sort-Queue}(V)$
4: while $Q$ is not empty do
5:    $u = Q.\text{pop-min}()$
6:    for $v \in G.\text{neighbors}(u)$ do
7:      $\text{Relax-Edge}(u, v, f)$
8:    end for
9: end while

Algorithm 2 Computation of 1D EPD

1: **Input**: filter function $f$, input graph $G = (V, E)$
2: $V, E = \text{sorted}(V, E, f)$
3: $PD_0 = \text{Union-Find}(V, E, f), PD_1 = \emptyset$
4: for $i \in V$ do
5:    $C_i = \{C_{ij} | (i, j) \in E, f(j) > f(i)\}, E_i = E$
6:    for $C_{ij} \in C_i$ do
7:      $f(C_{ij}) = f(i), E_i = E_i - \{(i, j)\} + \{(C_{ij}, j)\}$
8: end for
9: $PD_i = \text{Union-Find-step}(V + C_i - \{i\}, E_i, f, C_i)$
10: $PD_1 += PD_i$
11: end for
12: **Output**: $PD_0, PD_1$

Algorithm 3 Union-Find-step (Sequential)

1: **Input**: $V, E, f, C_i$
2: $PD_i = \emptyset$
3: for $v \in V$ do
4:    $v.\text{value} = f(v), v.\text{root} = v$
5: end for
6: $Q = \text{Sort}(V), Q = Q - \{v | f(v) < f(i)\}$
7: while $Q$ is not empty do
8:    $u = Q.\text{pop-min}()$
9:    for $v \in G.\text{neighbors}(u)$ do
10:      $pu, pv = \text{Find-Root}(u), \text{Find-Root}(v)$
11:      if $pu \neq pv$ then
12:        $s/l = \text{argmin/argmax}(pu.\text{value}, pv.\text{value})$
13:        $l.\text{root} = s$
14:      if $pu \in V_i$ and $pv \in V_i$ then
15:        $PD_i += \{(u.\text{value}, l.\text{value})\}$
16:      end if
17:    end if
18: end for
19: end while
20: **Function**: Find-Root($u$)
21: $pu = u$
22: while $pu \neq pu.\text{root}$ do
23:    $pu.\text{root} = (pu.\text{root}).\text{root}, pu = pu.\text{root}$
24: end while
25: Return: $pu$
As explained in the previous section, we will pair up all simplices (edges/vertices) in the input graph, \( PD_0 \) in the Appendix. The detailed algorithm/procedures involved can be found in Algorithm 3.

**Complexity.** The computational complexity of the Union-Find algorithm is \( O(|E|\alpha(|E|)) \), where \( \alpha(\cdot) \) is the inverse Ackermann function. Therefore, we need \( O(|V||E|\alpha(|E|)) \) time to compute an EPD using the sequential algorithm described in this section. Note this sequential algorithm is not necessarily the most efficient one. In practice, one may use the quadratic algorithm \( O(|V||E|) \) as in (Yan et al. 2021). We also note that although not formally published, the best known algorithm for EPD computation is quasilinear, \( O(|E|\log |V|) \), using the data structure of mergeable trees (Agarwal et al. 2006; Georgiadis et al. 2011). But this algorithm remains theoretical so far.

**Algorithm 4 Union-Find (Sequential)**

1: **Input:** \( V, E, f \)
2: \( PD_0 = \{\} \)
3: **for** \( v \in V \) **do**
4: \( v.value = f(v), v.root = v \)
5: **end for**
6: \( Q = \text{Sort}(V) \)
7: **while** \( Q \) is not empty **do**
8: \( u = Q.pop-min() \)
9: **for** \( v \in G.neighbors(u) \) **do**
10: \( pu, pv = \text{Find-Root}(u), \text{Find-Root}(v) \)
11: **if** \( pu \neq pv \) **then**
12: \( s/l = \text{argmin/argmax}(pu.value, pv.value) \)
13: \( l.root = s \)
14: \( PD_0 + \{(l.value, u.value)\} \)
15: **end if**
16: **end for**
17: **end while**
18: **Function:** \( \text{Find - Root}(u) \)
19: \( pu = u \)
20: **while** \( pu \neq pu.root \) **do**
21: \( pu.root = (pu.root).root, pu = pu.root \)
22: **end while**
23: **Return:** \( pu \)

**A.2 Learning Extended Persistence Diagrams**

In this section, we will introduce our neural network architecture to approximate the EPDs on graphs. As explained in the previous section, we will pair up all simplices (edges/vertices) in the input graph, instead of outputting only those non-local pairing corresponding to the pairing between critical points in the smooth case. The reason is that in this way, every edge is paired and corresponds to a persistence point, therefore the computation of EPD can be transformed into a link prediction problem on graphs. Furthermore, once such pairings are computed, we can easily remove all local pairings.

**Basic Framework.** As mentioned in Section A.1.1, every edge in the input graph will give rise to either a 0-dim ordinary persistence point or a 1D extended persistence point. Therefore, we transfer the learning of extended persistence diagram into a link prediction problem. Specifically, our base architecture, which is shown in Figure 2, follows standard link-prediction architectures (Chami et al. 2019; Yan et al. 2021): (1) For a input graph \( G = (V, E) \) and a filter function \( f \), we first obtain the initial filter value for all the nodes: \( X = f(V) \in R^{V \times 1} \), and then use a specially designed GNN model which later we call PDGNN \( G \) to obtain the node embedding for all these vertices: \( H = G(X) \in R^{V \times d_H} \). (2) Subsequently, a MLP (Multi-layer perceptron) \( W \) is applied to the node embeddings to obtain the persistent pairing information for each edge \( (u, v) \in E \) via \( PP_{uv} = W([h_u \oplus h_v]) \in \mathbb{R}^2 \) which is the persistence point associated to this edge \( (u, v) \). Here, \( h_u \) and \( h_v \) denote the node embedding for node \( u \) and \( v \), and \( \oplus \) represents the concatenation of vectors.
Figure 2: The basic framework.

**Standard GNN.** A GNN typically learns the node embedding via an iterative aggregation of local graph neighbors. The $k$-th iteration (the $k$-th GNN layer) can be written as:

$$h^k_u = AGG^k(\{MSG^k(h^{k-1}_v), v \in N(u)\}, h^{k-1}_u)$$  \quad (1)

where $h^k_u$ is the node features for node $u$ after $k$-th iterations, and $N(u)$ is the neighborhood of node $u$. In our setting, $h^0_u = x_u$ is initialized to be the filter value of node $u$. Different GNNs have different $MSG$ and $AGG$ functions, e.g., in GIN (Xu et al., 2018), $MSG$ is a MLP followed by an activation function, and $AGG$ is a sum aggregation.

**PDGNN.** We now describe our specially designed GNN for estimating persistence diagrams, called PDGNN (Persistence Diagram Graph Neural Network). Compared with the Sequential algorithms proposed in (Xhonneux et al., 2021), extended persistence diagrams need extra care: (1) the Find-Root algorithm needs to return the minimum of the component, (2) edge operations such as upper-edge splitting. Our PDGNN modifies standard GNNs with the following modules to handle these steps.

**Find-Root Algorithm.** Finding the minimum intuitively suggests using a combination of several local min-aggregations. Considering that the sum aggregation can bring the best expressiveness to GNNs (Xu et al., 2018), we implement the root-finding process by a combination of sum aggregation and min aggregation as our $AGG$ function. To be specific:

$$AGG^k(.) = SUM(.) \bigoplus MIN(.)$$ \quad (2)

**Incorporating edge operations.** As shown in (Veličković et al., 2019, Xhonneux et al., 2021), classic GNNs are not effective in “executing” algorithms operating on edges, such as Relax-Edge subroutine. Furthermore, in Algorithm 2, we also need the upper-edge splitting operation for each vertex. To this end, similar to (Xhonneux et al., 2021), we also use edge features and attention to provide bias using edges. Specifically, we propose the following $MSG$:

$$MSG^k(h^{k-1}_v) = \sigma^k[\alpha^k_{uv}(h^{k-1}_u \bigoplus h^{k-1}_v)W^k]$$ \quad (3)

where $\sigma^k$ is an activation function, $W^k$ is a MLP module, and $\alpha^k_{uv}$ is the edge weight for $uv$. We adopt $PRELU$ as our activation function, and the edge weight proposed in GAT (Veličković et al., 2018) as our edge weight.

Given that the Union-Find-step algorithm should be implemented on all nodes to obtain the EPDs, ideally we need a large GNN model to simulate all the computations. However, these Union-Find-step algorithms are running in parallel on the same graph. There are many overlapping or similar computational steps between computations on different nodes. Therefore we expect a limited-size single GNN model can simultaneously approximate all EPD computations. This is confirmed by empirical evidence in Section 4.
Table 2: Statistics of the node classification datasets

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Classes</th>
<th>Nodes</th>
<th>Edges</th>
<th>Features</th>
<th>Avg degree</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cora</td>
<td>7</td>
<td>2708</td>
<td>5429</td>
<td>1433</td>
<td>2.00</td>
</tr>
<tr>
<td>Citeseer</td>
<td>6</td>
<td>3327</td>
<td>4732</td>
<td>3703</td>
<td>1.42</td>
</tr>
<tr>
<td>PubMed</td>
<td>3</td>
<td>19717</td>
<td>44338</td>
<td>500</td>
<td>2.25</td>
</tr>
<tr>
<td>CS</td>
<td>15</td>
<td>18333</td>
<td>100227</td>
<td>6805</td>
<td>5.47</td>
</tr>
<tr>
<td>Physics</td>
<td>5</td>
<td>34493</td>
<td>282455</td>
<td>8415</td>
<td>8.19</td>
</tr>
<tr>
<td>Computers</td>
<td>10</td>
<td>13381</td>
<td>259159</td>
<td>767</td>
<td>19.37</td>
</tr>
<tr>
<td>Photo</td>
<td>8</td>
<td>7487</td>
<td>126530</td>
<td>745</td>
<td>16.90</td>
</tr>
</tbody>
</table>

A.3 Experimental details

A.3.1 Datasets

In this paper, we exploit real-world datasets including:

1. Citation networks: Cora, Citeseer, and PubMed (Sen et al., 2008) are standard citation networks where nodes denote scientific documents and edges denote citation links.

2. Amazon shopping records: In Photo and Computers (Shchur et al., 2018), nodes represent goods, edges represent that two goods are frequently brought together, and the node features are bag-of-words vectors.

3. Coauthor datasets: In CS and Physics (Shchur et al., 2018), nodes denote authors and edges denote that the two authors co-author a paper.

The detailed statistics are available in Table 2.

To compute EPDs, we need to set the input graphs and the filter functions. Existing state-of-the-art models on node classification (Zhao et al., 2020) and link prediction (Yan et al., 2021) mainly focus on the local topological information of the target node(s). Following their settings, for a given graph $G = (V, E)$ (e.g., Cora), we extract the $k$-hop neighborhoods of all the vertices, and extract $|V|$ vicinity graphs.

In terms of filter functions, we use Ollivier-Ricci curvature (Ni et al., 2018), heat kernel signature with two temperature values (Sun et al., 2009; Hu et al., 2014), and the node degree (Hofer et al., 2020). For an input vicinity graph, we compute 4 EPDs based on the 4 filter functions, and then vectorize them to get 4 persistence images (Adams et al., 2017). Therefore, we can get $4|V|$ EPDs in total.

We note that in the literature, EPDs often contain the extended persistence point of the whole connected component. In our setting, we remove the point because that (1) no edge is paired with the whole connected component; (2) the value is easy to obtain, and does not need an extra prediction.

A.3.2 Experimental Details

In this section, we mainly present the experimental settings on neural execution, as for the setting in downstream graph representation learning tasks, we are consistent with (Zhao et al., 2020; Yan et al., 2021).

Evaluation metrics. Recall that the input of our model is a graph and a filter function, and the output is the predicted diagram. After obtaining the predicted diagram, we vectorize it with persistence image (Adams et al., 2017) and evaluate (1) the 2-Wasserstein ($W_2$) distance between the predicted diagram and the ground truth EPD, (2) the total square error between the predicted persistence image and the ground truth image (persistence image error, denoted as PIE). Considering that our aim is to compute extended persistence diagrams on graphs rather than roughly approximating persistence.

2 Following the settings in (Zhao et al., 2020; Yan et al., 2021), we adopt the Ollivier-Ricci curvature as the graph metric, and the distance to target node(s) as the filter function.

3 Following the settings in (Carrière et al., 2020), we set the temperature $t = 10$ and 0.1 and adopt these two kernel functions as the filter functions.

4 Node degree is used as the initial filter function in (Hofer et al., 2020).
images, we use the $W_2$ distance as the training loss, while the PIE is only used as an evaluation metric. Given an input graph (e.g., Cora, Citeseer, etc.) and a filter function, we extract the $k$-hop neighborhoods of all the vertices and separate these vicinity graphs randomly into 80%/20% as training/test sets. We report the mean $W_2$ distance between diagrams and PIE on different vicinity graphs and all different filter functions.

### Other settings.
Following the settings in (Zhao et al., 2020; Yan et al., 2021), we extract 2-hop neighborhoods of all the nodes in Cora, Citeseer, PubMed, and 1-hop neighborhoods of all the nodes in Photo, Computers, Physics, and CS. In the training process, we only adopt the $W_2$ distance between the predicted diagram and the ground truth diagram as the loss function, while the PIE between the predicted persistence image and the ground truth persistence image only serves as an evaluation metric.

We adopt Adam as the optimizer with the learning rate set to 0.002 and weight decay set to 0.01. We build a 4-layer GNN framework with dropout set to 0.1. In the training process, we set the batch size to 10, and the training epoch to 20. In this paper, we also exploit a 2-layer MLP to transform the node embedding obtained by the GNN to the persistence points on edges. In the whole model, PRELU is adopted as the activation function, the dimension of hidden layers is set to 32, and the dimension of the output persistence image is 25. All the experiments are implemented with two Intel Xeon Gold 5128 processors, 192GB RAM, and 10 NVIDIA 2080TI graphics cards.

Notice that in normal computation of Wasserstein distance between PDs, the persistence points can be paired to the diagonal or the persistence points in the other diagram. However, in the experiments, we observe that with this loss the model may converge to local minima, e.g., all the predicted persistence points are paired to diagonal. Therefore, the predicted points all converge to the diagonal and contain no topological information. In case of these situations, we force the predicted points to pair with the persistence points in the ground truth diagram rather than the diagonal in the training stage, and report the normal $W_2$ distance, that is, to let the predicted points pair with the diagonal in the inference stage.

### A.4 Additional Experiments

#### A.4.1 Approximation Quality

In this section, we evaluate the approximation error between the predicted EPDs and the ground truth EPDs.

**Evaluation metrics.** Recall that the input of our model is a graph and a filter function, and the output is the predicted diagram. After obtaining the predicted diagram, we vectorize it with persistence image (Adams et al., 2017) and evaluate (1) the 2-Wasserstein ($W_2$) distance between the predicted
PI shows the benefit of (Yan et al., 2021), that is, to get the node embedding through these models, and use the Fermi-Dirac weight being helpful. Edge weights help the model focus on the individual Relax-Edge sub-algorithm with GIN (Xu et al. (2018) and GAT (Veličković et al. (2018) as the backbone GNNs.

To show the effectiveness of the modules proposed in Section A.2, we add other baselines with our PEGN (GIN showing the benefit of directly predicting EPDs, we also add the baseline methods (Zhao et al., 2020) and the two target nodes. In comparison with the original extended persistence diagram, we also add decoder (Krioukov et al., 2010; Nickel & Kiela, 2017) to predict whether there is a link between not originally designed for link prediction, therefore we follow the settings in (Chami et al., 2019; SEAL such as (Chami et al., 2019). For link prediction, we compare with several state-of-the-art methods (Kipf & Welling, 2016), popular GNN models including GCN and GAT (+MIN) with SEAL denoting our proposed method, that is, the GNN framework with the AGG function and MSG function proposed in Section A.2. Its strategy is to first predict the EPD, and then convert it into the persistence image. To show the superiority of the strategy, we compare with the strategy from Som et al. (2020); Montufar et al. (2020), i.e., directly approximate the persistence image of the input graph, as a baseline strategy. GIN.PI and GAT.PI denote the baseline strategy with GIN (Xu et al. (2018) and GAT (Veličković et al. (2018) as the backbone GNNs.

To show the effectiveness of the modules proposed in Section A.2, we add other baselines with our proposed strategy. GAT denotes GAT as the backbone GNN. GAT (+MIN) denotes GAT with the new AGG function. Compared with PDGNN, it exploits the original node feature rather than the new edge feature in the MSG function. PDGNN (w/o ew) denotes PDGNN without edge weight. Further experimental settings can be found in the Appendix.

Table 5: AUC-ROC score on various link prediction benchmarks

<table>
<thead>
<tr>
<th>Method</th>
<th>Cora</th>
<th>Citeseer</th>
<th>PubMed</th>
<th>Photo</th>
<th>Computers</th>
</tr>
</thead>
<tbody>
<tr>
<td>GCN</td>
<td>90.5±0.2</td>
<td>82.6±1.9</td>
<td>89.6±3.7</td>
<td>91.8±0.0</td>
<td>87.8±0.0</td>
</tr>
<tr>
<td>GAT</td>
<td>72.8±0.2</td>
<td>74.8±1.5</td>
<td>80.3±0.0</td>
<td>92.9±0.3</td>
<td>86.4±0.0</td>
</tr>
<tr>
<td>HGCN</td>
<td>93.8±0.1</td>
<td>96.6±0.1</td>
<td>96.3±0.0</td>
<td>95.4±0.0</td>
<td>93.6±0.0</td>
</tr>
<tr>
<td>P-GNN</td>
<td>74.1±2.4</td>
<td>73.9±2.6</td>
<td>79.6±0.5</td>
<td>90.9±0.7</td>
<td>88.3±1.0</td>
</tr>
<tr>
<td>SEAL</td>
<td>91.3±5.7</td>
<td>89.8±2.3</td>
<td>92.4±1.2</td>
<td>97.8±1.3</td>
<td>96.8±1.5</td>
</tr>
<tr>
<td>TLC-GNN (True Diagram)</td>
<td>94.9±0.4</td>
<td>95.1±0.7</td>
<td>97.0±0.1</td>
<td>98.2±0.1</td>
<td>97.9±0.1</td>
</tr>
<tr>
<td>TLC-GNN (GIN.PI)</td>
<td>93.0±0.2</td>
<td>92.8±0.6</td>
<td>96.3±0.2</td>
<td>95.8±1.0</td>
<td>96.2±0.3</td>
</tr>
<tr>
<td>TLC-GNN (PDGNN)</td>
<td>95.0±0.3</td>
<td>95.6±0.4</td>
<td>97.0±0.1</td>
<td>98.4±0.6</td>
<td>98.2±0.3</td>
</tr>
</tbody>
</table>

Table 3 reports the approximation error, we observe that PDGNN outperforms all the baseline methods among all the datasets. The comparison between GAT and GAT.PI shows the benefit of predicting EPDs instead of predicting the persistence image. Comparing GAT and GAT (+MIN), we observe the advantage of the new AGG function, which shows the necessity of using min aggregation to approximate the Find-Root algorithm; Comparing GAT (+MIN) and PDGNN, we observe the effectiveness of using the new MSG function to help the model capture information of the separated connected components. Comparing PDGNN (w/o ew) and PDGNN, we find the use of the edge weight being helpful. Edge weights help the model focus on the individual Relax-Edge sub-algorithm operated on every edge.

A.4.2 Downstream Tasks

In this section, we evaluate the performance of the predicted diagrams on 2 graph representation learning tasks: node classification and link prediction. To be specific, we replace the ground truth EPDs in state-of-the-art models based on extended persistent homology (Zhao et al., 2020; Yan et al., 2021) with our predicted diagrams and report the results.

Baselines. We compare our method with various state-of-the-art methods. We compare with popular GNN models including GCN (Kipf & Welling, 2016), GAT (Veličković et al. 2018) and HGCN (Chami et al. (2019). For link prediction, we compare with several state-of-the-art methods such as SEAL (Zhang & Chen (2018) and P-GNN (You et al. (2019). Notice that GCN and GAT are not originally designed for link prediction, therefore we follow the settings in (Chami et al. (2019); Yan et al. (2021); that is, to get the node embedding through these models, and use the Fermi-Dirac decoder (Krioukov et al. (2010) [Nickel & Kiela (2017)] to predict whether there is a link between the two target nodes. In comparison with the original extended persistence diagram, we also add PEGN (Zhao et al. (2020) and TLC-GNN (Yan et al. (2021) as baseline methods. Furthermore, to show the benefit of directly predicting EPDs, we also add the baseline methods PEGN (GIN.PI) and
Table 6: Transferability in terms of different graph structures ($W_2$ distance.)

<table>
<thead>
<tr>
<th></th>
<th>Pre-train</th>
<th>Cora</th>
<th>Citeseer</th>
<th>PubMed</th>
<th>Photo</th>
<th>Computers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-train</td>
<td>0.392</td>
<td>0.279</td>
<td>0.444</td>
<td>0.379</td>
<td>0.404</td>
<td></td>
</tr>
<tr>
<td>Fine-tune</td>
<td>0.348</td>
<td>0.259</td>
<td>0.360</td>
<td>0.380</td>
<td>0.381</td>
<td></td>
</tr>
<tr>
<td>Standard</td>
<td>0.354</td>
<td>0.267</td>
<td>0.344</td>
<td>0.379</td>
<td>0.377</td>
<td></td>
</tr>
</tbody>
</table>

**TLC-GNN (GIN.PI)**, which replace the original persistent homology feature with the output from GIN.PI.

**Evaluation metrics.** For node classification, our setting is the same as [Kipf & Welling, 2016; Veličković et al., 2018; Zhao et al., 2020]. To be specific, we train the GNNs with 20 nodes from each class and validate (resp. test) the GNN on 500 (resp. 1000) nodes. We run the GNNs on these datasets 10 times and report the average classification accuracy and standard deviation. For link prediction, our setting is the same as [Chami et al., 2019; Yan et al., 2021]. To be precise, we randomly split existing edges into 85/5/10% for training, validation, and test sets. An equal number of non-existent edges are sampled as negative samples in the training process. We fix the negative validation and test sets, and randomly select the negative training sets in every epoch. We run the GNNs on these datasets 10 times and report the mean average area under the ROC curve (ROCAUC) scores and the standard deviation.

**Results.** Table 4 and Table 5 summarize the performance of all methods on node classification and link prediction. We observe that PEGN (PDGNN) and TLC-GNN (PDGNN) consistently perform comparably with PEGN and TLC-GNN, showing that the EPDs approximated by PDGNN have the same learning power as the true EPDs. Furthermore, PEGN using the approximated EPDs achieve better or comparable performance with different SOTA methods.

We also discover that PEGN (GIN.PI) and TLC-GNN (GIN.PI) perform much inferior to the original models using the true EPDs. It demonstrates that the large approximation error from GIN.PI lose much of the crucial information which is preserved in PDGNN.

### A.4.3 Experiments on Transferability

In this section, we design experiments to evaluate the transferability of PDGNN in terms of different graph structures. Our aim is that with a pre-trained model, or simply fine-tuning the pre-trained model can let the model predict EPDs on totally unseen graphs. Therefore, we use the models pre-trained on Photo, and report the $W_2$ distance between the predicted diagrams and ground truth EPDs. Notice that we only use Ollivier-Ricci curvature [Ni et al., 2018] as the filter function. The results are shown in Table 6.

In Table 6, “Pre-train” is to directly predict the EPDs with the pre-trained model, and “Fine-tune” is to fine-tune an epoch on the new datasets, and then predict the EPDs. As shown in Table 6, directly predicting the EPDs with the pre-trained model perform comparably with the standard settings among datasets. We also observe that with only a one-epoch fine-tune, the pre-trained model can achieve almost an equal performance compared with the standard setting. It justifies the fine transferability of PDGNN.

The fine transferability makes it possible to apply the computationally expensive topological features to a wide spectrum of real-world graphs; we can potentially apply a pre-trained model to large and dense graphs, on which direct EPD computation is infeasible. In a totally new environment, instead of training the models without pre-training for many epochs, we can simply fine-tune or even directly use the pre-trained model to predict extended persistence diagrams on new graph structures.

### A.4.4 Evaluation on the Influence of Training Samples

In this section, we evaluate the influence of training samples on PDGNN. Our aim is to show that the model can reach an acceptable performance with only a small number of training samples.
Recall that for a given graph, we extract the $k$-hop neighborhoods of all the nodes and randomly select 80% of these vicinity graphs to train PDGNN. For a thorough evaluation, we train PDGNN with 5/10/20/40% vicinity graphs in this experiment and report the $W_2$ distance of persistence diagrams, the PIE of persistence images, and the node classification accuracy (NCA) in Table 7. We also visualize the influence in Figure 3 and Figure 4.

As shown in Figure 3, the training error tends to converge as the training samples gradually increase. Considering that the $W_2$ distance and PIE cannot directly reflect the influence as NCA does, we select a vicinity graph in Cora which is hard for PDGNN to learn and visualize in Figure 4. As shown in the figure, as the number of training samples increases, we find that PDGNN can gradually capture the ground truth persistent points in the up y-axis and the up-right diagonal with much less noise. The number of training samples may help the model learn the hard samples better.

We can also observe that in Table 7, PDGNN can reach a comparable performance on NCA with much fewer training samples. The observation shows that a little perturbation on the persistence image will not influence its structural information very much.

Combining the observation in Section A.4.3 and Section A.4.4, we can draw a conclusion that our model can be easily generalized to other frameworks. PDGNN does not need many training samples to reach an acceptable performance, while it can be easily transferred to totally unseen graphs.

### A.4.5 Experiments on graph classification datasets.

In the experiment part (Section 4), we only consider predicting EPDs in the $k$-hop neighborhoods of the original graphs. Even if these vicinity graphs can be large and dense, there are fundamental structural differences between these vicinity graphs and other real-world graphs. For example, in a $k$-hop neighborhood of node $u$, the distance between a node $u_1$ to the other node $u_2$ is at most $2k$. 

Recall that the $W_2$ distance and PIE cannot directly reflect the influence as NCA does. We select a vicinity graph in Cora which is hard for PDGNN to learn and visualize in Figure 4. As shown in the figure, as the number of training samples increases, we find that PDGNN can gradually capture the ground truth persistent points in the up y-axis and the up-right diagonal with much less noise. The number of training samples may help the model learn the hard samples better.

We can also observe that in Table 7, PDGNN can reach a comparable performance on NCA with much fewer training samples. The observation shows that a little perturbation on the persistence image will not influence its structural information very much.

Combining the observation in Section A.4.3 and Section A.4.4, we can draw a conclusion that our model can be easily generalized to other frameworks. PDGNN does not need many training samples to reach an acceptable performance, while it can be easily transferred to totally unseen graphs.
Figure 4: Visualization on the influence of training samples. We select a vicinity graph in Cora with Ollivier-Ricci curvature as the filter function, and plot the influence of training samples the $W_2$ distance (loss) of EPDs. (a), (b), and (c) denote the prediction of PDGNN with 5/10/20% training samples, (d) denotes the prediction of PDGNN with the standard setting.
Table 8: Statistics and approximation error on the graph classification datasets

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Graphs</th>
<th>Avg Nodes</th>
<th>Avg Edges</th>
<th>$W_2$</th>
<th>PIE</th>
</tr>
</thead>
<tbody>
<tr>
<td>MUTAG</td>
<td>188</td>
<td>17.9</td>
<td>39.6</td>
<td>0.300</td>
<td>3.06e-4</td>
</tr>
<tr>
<td>ENZYMES</td>
<td>600</td>
<td>32.6</td>
<td>124.3</td>
<td>0.299</td>
<td>3.72e-3</td>
</tr>
<tr>
<td>PROTEINS</td>
<td>1113</td>
<td>39.1</td>
<td>145.6</td>
<td>0.194</td>
<td>8.30e-4</td>
</tr>
<tr>
<td>IMDB-BINARY</td>
<td>1000</td>
<td>19.8</td>
<td>193.1</td>
<td>0.176</td>
<td>4.13e-4</td>
</tr>
<tr>
<td>REDDIT-BINARY</td>
<td>2000</td>
<td>429.6</td>
<td>995.5</td>
<td>0.383</td>
<td>1.92e-4</td>
</tr>
</tbody>
</table>

$d(u_1, u_2) \leq d(u_1, u) + d(u_2, u) \leq k + k = 2k$. However, in a real-world graph, the distance can be very long.

**Observation.** Assume that our aim is to find a loop generated by two connected components that originate from the same node. An example is shown in Figure 1(c), in which the two connected components $C_{14}$ and $C_{13}$ are combined when node $u_4$ arises. In every message passing iteration, the Union-Find-step algorithm will only run the Relax-Edge function once. If starting from the connected component $C_{14}$, the algorithm needs 1 GNN layer to find node $u_4$. Similarly, if starting from the connected component $u_{13}$, the algorithm needs 2 GNN layers to find node $u_4$. Therefore, the algorithm needs $\max(1, 2) = 2$ GNN layers to find node $u_4$. In conclusion, to find a loop generated by two connected components $c_1$ and $c_2$, the algorithm needs $\max(d_1, d_2)$ layers to obtain the persistence point, where $d_1$ (resp. $d_2$) denotes the distance from $c_1$ (resp. $c_2$) to the node that the two connected components are combined. Therefore, we can use a $2k$-layer PDGNN to obtain the persistence points of all the potential loops in a $k$-hop neighborhood. In the settings of [Zhao et al., 2020; Yan et al., 2021], $k$ is at most 2. Consequently in Section 4, we set the layer of PDGNN to 4 and achieve a promising approximation result.

In this section, we do further experiments on graph classification datasets, in which we approximate the EPDs on the original graph rather than the vicinity graphs. We exploit the datasets from the TU Dortmund University [Morris et al., 2020], the detailed information of these datasets and the approximation error are all available in Table 8.

Notice that we do not add Ollivier-Ricci curvature as the filter function here, because computing the filter function on all the graphs will bring too much computational cost. Comparing the results from Table 8 and Table 4, we observe that the performance on graph classification datasets is generally worse. There are 2 possible explanations: (1) In the graph classification datasets, the training samples can be very small, e.g., there are only 188 graphs in MUTAG, therefore PDGNN has not well approximated the algorithm. (2) As discussed above, the diameter of these graphs can be large, therefore extra GNN layers are needed to execute the algorithm. However, we discover that the increasing of GNN layers only brings little improvement to the approximation error. This can be due to the well-known vanishing gradient problem in deep learning models.

To evaluate the results more clearly, we also visualize some selected examples in Figure 5. As shown in the figure, in most situations, PDGNN can well execute the EPDs on these graphs, and the $W_2$ distance around 0.3 is generally an acceptable result. However, in certain cases like Figure 5(d), the model only captures a tendency of the EPD. This can be due to the long diameter of the graph.
Figure 5: Visualization of graph classification samples. We select samples from IMDB-BINARY, PROTEINS, ENZYMES, and REDDIT-BINARY, respectively, and report the $W_2$ distance (loss).