# Time-aware Random Walk Diffusion to Improve Dynamic Graph Learning

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#### Abstract

How can we augment a dynamic graph for improving the performance of dynamic graph neural networks? Graph augmentation has been widely utilized to boost the learning performance of GNN-based models. However, most existing approaches only enhance spatial structure within an input static graph by transforming the graph, and do not consider dynamics caused by time such as temporal locality, i.e., recent edges are more influential than earlier ones, which remains challenging for dynamic graph augmentation. In this work, we propose TIARA (Time-aware Random Walk Diffusion), a novel diffusion-based method for augmenting a dynamic graph represented as a discrete-time sequence of graph snapshots. For this purpose, we first design a time-aware random walk proximity so that a surfer can walk along the time dimension as well as edges, resulting in spatially and temporally localized scores. We then derive our diffusion matrices based on the time-aware random walk, and show they become enhanced adjacency matrices that both spatial and temporal localities are augmented. Throughout extensive experiments, we demonstrate that TIARA effectively augments a given dynamic graph, and leads to significant improvements in dynamic GNN models for various graph datasets and tasks.

#### Introduction

Dynamic graphs represent various real-world relationships that dynamically occur over time. Learning such dynamic graphs has recently attracted considerable attention from machine learning communities (Skarding, Gabrys, and Musial 2021; Han et al. 2021), and plays a crucial role in diverse applications such as link prediction (Yang et al. 2021; Pareja et al. 2020), node or edge classification (Xu et al. 2019; Pareja et al. 2020), time-series traffic forecasting (Wu et al. 2020; Guo et al. 2019), knowledge completion (Jung, Jung, and Kang 2021), and pandemic forecasting (Panagopoulos, Nikolentzos, and Vazirgiannis 2021). Over the last years, many researchers have put tremendous effort into developing interesting methods by sophisticatedly fusing GNNs and recurrent neural networks (RNN) or attention mechanisms for continuous-time (Xu et al. 2020; Rossi et al. 2020) and discrete-time (Seo et al. 2018; Pareja et al. 2020; Yang et al. 2021) dynamic graphs.

\*Jinhong Jung is the corresponding author. Copyright © 2023, Association for the Advancement of Artificial Intelligence (www.aaai.org). All rights reserved. With the astonishing progress of GNNs, diverse augmentation techniques (Zhao et al. 2022; Yoo, Shim, and Kang 2022) have been proposed to increase the generalization power of GNN models, especially on a static graph. Previous approaches mainly transform the topological structure of the input graph. For example, drop-based methods stochastically remove a certain number of edges (Rong et al. 2020) or nodes (Feng et al. 2020) at each training epoch in a similar manner to dropout regularization. On the contrary, diffusion methods (Klicpera, Weißenberger, and Günnemann 2019) insert additional edges having weights scored by graph diffusions such as Personalized PageRank (Tong, Faloutsos, and Pan 2006), thereby augmenting a spatial locality around each node and improving graph convolution.

However, the aforementioned techniques assume to augment data within a static graph, and dynamic graph augmentation problem has not yet been comprehensively studied. Unlikely static graphs, dynamic graphs change or evolve over time by their nature; thus, dynamic graph augmentation needs to simultaneously consider temporal dynamics as well as spatial structure. More specifically, as verified in previous works (Rossi et al. 2020; Shin 2017; Lee, Shin, and Faloutsos 2020), real-world dynamic graphs exhibit *temporal locality* indicating that graph objects such as nodes and triangles tend to be more affected by more recent edges than older ones, i.e., edges closer to a specific object in time are more likely to provide important information. Naively applying a static augmentation method to each time step cannot consider such a temporal locality.

In this work, we propose TIARA (<u>Time-a</u>ware <u>Random</u> Walk Diffusion), a novel diffusion-based augmentation method for a discrete-time dynamic graph which is represented by a temporal sequence of graph snapshots. TIARA aims to augment both spatial and temporal localities of each graph snapshot. For this purpose, we design a time-aware random walk that a surfer randomly moves around nodes or a time-axis to measure spatially and temporally localized scores. We then derive time-aware random walk diffusion from the scores, and interpret it as the combination of spatial and temporal augmenters. Our diffusion matrices are used as augmented adjacency matrices for any dynamic GNN models in discrete-time domain. We further adopt approximate techniques such as power iteration and sparsification to reduce a heavy cost for computing the diffusion matrices.

Our contributions are summarized as follows:

- Method. We propose TIARA, a novel and modelagnostic method for dynamic graph augmentation using time-aware random walks. TIARA strengthens not only a spatial locality but also a temporal locality of a dynamic graph so that dynamic GNNs perform better.
- Analysis. We analyze how TIARA augments both spatial and temporal localities (Theorem 1) and complexities of TIARA (Theorem 2) in real dynamic graphs.
- Experiments. We demonstrate that TIARA effectively augments a given dynamic graph, and leads to consistent improvements in GNNs for temporal link prediction and node classification tasks.

The code of TIARA and the datasets are publicly available at https://github.com/dev-jwel/TiaRa.

#### **Related Work**

Augmentation for Static GNNs. Graph augmentation (Zhao et al. 2022) aims to reduce over-fitting for training GNN models by modifying an input graph. DropEdge (Rong et al. 2020) or DropNode (Feng et al. 2020) randomly drop edges or nodes at each epoch. These augment the diversity of the input graph by creating different copies sampled from the graph. GDC (Klicpera, Weißenberger, and Günnemann 2019) adds new edges weighted by a graph diffusion derived from node proximities. GDC boosts a spatial locality of the graph so that a GNN can consider adjacent nodes as well as distant ones during their convolutions, enhancing its representation power. However, most of existing methods are limited to augment dynamic graphs because they do not consider temporal properties.

GNNs and Augmentation for Dynamic Graphs. Dynamic graphs (Kazemi et al. 2020) are categorized as: discrete-time dynamic graphs (DTDG) and continuous-time dynamic graphs (CTDG) where a DTDG is represented as a sequence of graph snapshots with multiple discrete time steps while a CTDG is represented as a set of temporal edges whose time-stamps have continuous values. It is straightforward to convert a CTDG to a DTDG by distributing the continuous-time edges into multiple bins in chronological order, but the reverse is not possible because continuous-time values are generally lacked in most DTDGs (Yang et al. 2021), i.e., models for DTDGs can be applied to CTDGs, but the reverse is rather limited. Hence, we narrow our focus to representation learning on DTDGs.

Dynamic GNNs have rapidly advanced under the framework that closely integrates GNNs and temporal sequence models such as RNNs to capture spatial and temporal relations on dynamic graphs (Skarding, Gabrys, and Musial 2021). GCRN (Seo et al. 2018) uses a GCN to produce node embeddings on each graph snapshot, and then forwards them to an LSTM for modeling temporal dynamics. STAR (Xu et al. 2019) utilizes a GRU combined with spatial and temporal attentions. DySat (Sankar et al. 2020) employs a self-attention strategy to aggregate spatial neighborhood and temporal dynamics. EvolveGCN (Pareja et al. 2020) evolves the parameters of GCNs using RNNs. To consider

hierarchical properties in real graphs, HTGN (Yang et al. 2021) extends the framework to hyperbolic space.

As a related method, MeTA (Wang et al. 2021) adaptively augments a temporal graph based on predictions of a temporal graph network, which perturbs time and removes or adds edges. However, it is difficult to employ MeTA for the aforementioned DTDG models because MeTA is designed for CTDGs requiring continuous-time values.

#### **Preliminaries**

Random Walk with Restart (RWR). Our work is related to RWR which measures node similarity scores which are spatially localized to seed node s (Nassar, Kloster, and Gleich 2015), i.e., scores of nearby nodes highly associated to s are high while those of distant nodes are low. Diffusion methods such as GDC exploit RWR to augment a spatial locality.

Let  $\mathbf{x}_s$  be a vector of RWR scores w.r.t. the seed node s. Given a row-normalized adjacency matrix  $\tilde{\mathcal{A}}$  and a restart probability  $\alpha$ , the vector  $\mathbf{x}_s$  is represented as follows:

$$\mathbf{x}_s = (1 - \alpha) \tilde{\mathbf{A}}^{\top} \mathbf{x}_s + \alpha \mathbf{i}_s \Leftrightarrow \mathbf{x}_s = \alpha \mathbf{L}^{-1} \mathbf{i}_s \Leftrightarrow \mathbf{x}_s = \mathbf{\mathcal{L}}^{\text{rwr}} \mathbf{i}_s$$

where  $\mathbf{L} = \mathcal{I}_n - (1-\alpha)\tilde{\mathcal{A}}^{\top}$  is the random-walk normalized Laplacian matrix, and  $\mathbf{i}_s$  is the *s*-th unit vector. Notice that  $\mathcal{L}^{\text{rwr}} = \alpha \mathbf{L}^{-1}$  is a column-stochastic transition matrix interpreted as a diffusion kernel that diffuses a given distribution such as  $\mathbf{i}_s$  on the graph through RWR.

**Problem Formulation.** A discrete-time dynamic graph (DTDG)  $\mathcal{G}$  is represented as a sequence  $\{\mathcal{G}_1,\cdots,\mathcal{G}_T\}$  of snapshots in a chronological order where T is the number of time steps (Skarding, Gabrys, and Musial 2021). Each snapshot  $\mathcal{G}_t = (\mathcal{V}, \mathcal{E}_t, \mathbf{F}_t)$  is a graph with a shared set  $\mathcal{V}$  of nodes and a set  $\mathcal{E}_t$  of edges at time t where  $n = |\mathcal{V}|$  is the number of nodes.  $\mathbf{F}_t \in \mathbb{R}^{n \times d}$  is an initial node feature matrix where d is a feature dimension, and  $\mathbf{A}_t \in \mathbb{R}^{n \times n}$  denotes the sparse and self-looped adjacency matrix of  $\mathcal{G}_t$ . The node representation learning on the dynamic graph  $\mathcal{G}$  aims to learn a function  $\mathcal{F}_{\Theta}(\cdot)$  parameterized by  $\Theta$  and produce hidden node embeddings  $\mathbf{H}_t \in \mathbb{R}^{n \times d}$  for each time t, represented as:

$$\mathbf{H}_t = \mathcal{F}_{\Theta}(\tilde{\mathbf{A}}_t, \mathbf{F}_t, \mathbf{H}_{t-1}) \tag{1}$$

where  $\bar{\mathcal{A}}_t$  is a normalized adjacency matrix of  $\mathbf{A}_t$ , and  $\mathbf{H}_{t-1}$  contains the latest hidden embeddings before time t. The above framework of Equation (1) is generally adopted in existing methods (Seo et al. 2018; Pareja et al. 2020; Yang et al. 2021) for learning DTDGs where  $\mathcal{F}_{\Theta}(\cdot)$  is usually designed by the combination of GNNs and RNNs.

**Problem 1** (Dynamic Graph Augmentation). Given a temporal sequence  $\{\mathbf{A}_1, \dots, \mathbf{A}_T\}$  of  $\mathcal{G}$ , the problem is to generate a sequence of new adjacency matrices improving the performance of a model  $\mathcal{F}_{\Theta}(\cdot)$ .

## **Proposed Method**

We depict the overall framework of TIARA in Figure 1. Given  $\{\mathbf{A}_1, \cdots, \mathbf{A}_T\}$  of a dynamic graph  $\mathcal{G}$ , our TIARA aims to produce a time-aware random walk diffusion matrix  $\tilde{\mathcal{X}}_t \in \mathbb{R}^{n \times n}$  for each time step t using two diffusion based modules, called *spatial* and *temporal* augmenters.

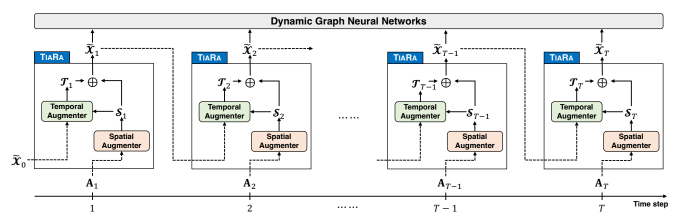


Figure 1: Overall architecture of TIARA. Given the adjacency matrix  $\mathbf{A}_t$  at time t, TIARA outputs a time-aware random walk diffusion matrix  $\tilde{\mathcal{X}}_t$  combined with spatial augmenter  $\mathcal{S}_t$  and temporal augmenter  $\mathcal{T}_t$  after sparsification.

The spatial augmenter enhances a spatial locality of  $\mathbf{A}_t$  using random walks, resulting in a spatial diffusion matrix  $\mathcal{S}_t$ . The temporal augmenter receives the previous  $\tilde{\mathcal{X}}_{t-1}$  that contains information squashed from the initial time to t-1, and then disseminates it through  $\mathcal{S}_t$  at the current t. This leads to a temporal diffusion matrix  $\mathcal{T}_t$  in which a temporal locality is magnified. Finally, TIARA linearly combines  $\mathcal{S}_t$  and  $\mathcal{T}_t$ , and sparsifies to form  $\tilde{\mathcal{X}}_t$ . We replace each adjacency matrix  $\mathbf{A}_t$  with  $\mathcal{X}_t$  for the inputs of dynamic GNN models. If necessary, we simply use edges of the graph in represented by  $\tilde{\mathcal{X}}_t$  without weights, or make the graph undirected by using  $(\tilde{\mathcal{X}}_t + \tilde{\mathcal{X}}_t^\top)/2$  after the sparsification.

#### **Time-aware Random Walk with Restart**

It is limited to directly employ RWR in a dynamic graph because RWR measures only spatially localized scores in a single static graph. In this section, we extend RWR to Timeaware RWR (TRWR) so that TRWR produces node-to-node scores which are spatially and temporally localized.

One idea for TRWR is to virtually connect identical nodes from  $\mathcal{G}_t$  to  $\mathcal{G}_{t+1}$  for each time step t (Huang, Sun, and Wang 2021) as shown in Figure 2. Then, a random surfer not only moves around the current  $\mathcal{G}_t$  but also jumps to the next  $\mathcal{G}_{t+1}$ ; thus, the surfer becomes time-aware. In the beginning, the surfer starts from a seed node s at the initial time step (e.g., t=1). After a few movements, suppose the surfer is at node s in s, then, it takes one of the following actions:

- Action 1) Random walk. The surfer randomly moves to one of the neighbors from node u in the current graph  $\mathcal{G}_t$  with probability  $1 \alpha \beta$ .
- Action 2) Restart. The surfer goes back to the seed node
   s in G<sub>t</sub> with probability α.
- Action 3) Time travel. The surfer does time travel from node u in  $\mathcal{G}_t$  to that node in  $\mathcal{G}_{t+1}$  with probability  $\beta$ .

where  $\alpha$  and  $\beta$  are called restart and time travel probabilities, respectively, and  $0 < \alpha + \beta < 1$ . Note that we do not allow the surfer to move backward from  $\mathcal{G}_{t+1}$  to  $\mathcal{G}_t$  because the future information at time t+1 should be prevented when we make a prediction at time t.

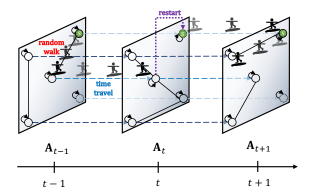


Figure 2: Illustration of how a time-aware random surfer moves around a dynamic graph where dashed arrows indicate virtually connected edges along the time-axis.

Through TRWR, the vector  $\mathbf{x}_t \in \mathbb{R}^n$  of stationary probabilities that the surfer visits each node from the seed node s in  $\mathcal{G}_t$  is recursively represented as follows:

$$\mathbf{x}_{t,s} = \underbrace{(1 - \alpha - \beta)\tilde{\mathbf{A}}_{t}^{\top}\mathbf{x}_{t,s}}_{\text{Random walk}} + \underbrace{\alpha \mathbf{i}_{s}}_{\text{Restart}} + \underbrace{\beta \mathbf{x}_{t-1,s}}_{\text{Time travel}}$$
(2)

where  $\mathbf{i}_s$  is the s-th unit vector of size n.  $\tilde{\mathcal{A}}_t$  is a row-normalized matrix of  $\mathbf{A}_t$  (i.e.,  $\tilde{\mathcal{A}}_t = \mathbf{D}_t^{-1} \mathbf{A}_t$  where  $\mathbf{A}_t$  is a self-looped adjacency matrix and  $\mathbf{D}_t$  is a diagonal out-degree matrix of  $\mathbf{A}_t$ ). If t = 0, we define  $\mathbf{x}_{0,s}$  as  $\mathbf{i}_s$ .

In the above equation, the random walk part propagates scores of  $\mathbf{x}_{t,s}$  over  $\tilde{\mathcal{A}}_t$ . The restart part makes the scores spatially localized around the seed node s, which is controlled by  $\alpha$ . The time travel part injects scores of the previous  $\mathbf{x}_{t-1,s}$  to make  $\mathbf{x}_{t,s}$  temporally localized, which is controlled by  $\beta$ . Notice TRWR extends RWR to a discrete-time dynamic graph, i.e.,  $\beta=0$  leads to RWR scores on each graph snapshot without considering temporal information.

## **Time-aware Random Walk Diffusion Matrices**

In Equation (2),  $\mathbf{x}_{t,s} \in \mathbb{R}^{n \times 1}$  is a column vector of a probability distribution w.r.t. a seed node s. For all seeds  $s \in \mathcal{V}$ ,

we horizontally stack  $\{\mathbf{x}_{t,s}\}$  to form  $\mathcal{X}_t \in \mathbb{R}^{n \times n}$  such that  $\mathbf{x}_{t,s}$  is the s-th column of  $\mathcal{X}_t$ , i.e.,  $\mathbf{x}_{t,s} = \mathcal{X}_t \mathbf{i}_s$ . We call  $\mathcal{X}_t$  a time-aware random walk diffusion matrix at time t. The derivation of  $\mathcal{X}_t$  starts by moving the term of the random walk to the left side in Equation (2) as follows:

$$\left(\mathcal{I}_n - (1 - \alpha - \beta)\tilde{\mathcal{A}}_t^{\top}\right)\mathbf{x}_{t,s} = \alpha \mathbf{i}_s + \beta \mathbf{x}_{t-1,s}$$

Let  $\mathbf{L}_t \coloneqq \mathcal{I}_n - (1 - \alpha - \beta)\tilde{\mathcal{A}}_t^{\top}$  where  $\mathcal{I}_n$  is an  $n \times n$  identity matrix, and  $\mathbf{x}_{t-1,s} = \mathcal{X}_{t-1}\mathbf{i}_s$  as described above. Thus,  $\mathbf{x}_{t,s}$  is written as the following:

$$\mathbf{x}_{t,s} = \mathbf{L}_t^{-1} \left( \alpha \mathbf{i}_s + \beta \mathbf{\mathcal{X}}_{t-1} \mathbf{i}_s \right)$$
$$= \left( \alpha \mathbf{L}_t^{-1} \mathbf{\mathcal{I}}_n + \beta \mathbf{L}_t^{-1} \mathbf{\mathcal{X}}_{t-1} \right) \mathbf{i}_s = \mathbf{\mathcal{X}}_t \mathbf{i}_s$$
(3)

where  $\mathcal{X}_t = \alpha \mathbf{L}_t^{-1} \mathcal{I}_n + \beta \mathbf{L}_t^{-1} \mathcal{X}_{t-1}$  for t > 0, and  $\mathcal{X}_0 = \mathcal{I}_n$  because  $\mathbf{x}_{0,s}$  is defined as  $\mathbf{i}_s$ .

**Spatial and Temporal Augmenters.** We obtain the recurrence relation of  $\mathcal{X}_t$  from Equation (3), and further rearrange it to interpret the process as follows:

$$\mathcal{X}_{t} = \alpha \mathbf{L}_{t}^{-1} \mathcal{I}_{n} + \beta \mathbf{L}_{t}^{-1} \mathcal{X}_{t-1} 
= \frac{\alpha}{\alpha + \beta} \left[ (\alpha + \beta) \mathbf{L}_{t}^{-1} \mathcal{I}_{n} \right] + \frac{\beta}{\alpha + \beta} \left[ (\alpha + \beta) \mathbf{L}_{t}^{-1} \mathcal{X}_{t-1} \right]$$

In the above, we set  $\mathcal{L}_t^{\text{rwr}} = (\alpha + \beta) \mathbf{L}_t^{-1}$  which is the diffusion kernel by RWR on the graph  $\mathcal{G}_t$  where its restart probability is  $\alpha + \beta$ . Let  $\gamma = \beta/(\alpha + \beta)$  where  $0 < \gamma < 1$ ; then,  $\mathcal{X}_t$  is represented as follows:

$$\mathcal{X}_{t} = (1 - \gamma) \underbrace{(\mathcal{L}_{t}^{\text{rwr}} \mathcal{I}_{n})}_{\mathcal{S}_{t}} + \gamma \underbrace{(\mathcal{L}_{t}^{\text{rwr}} \mathcal{X}_{t-1})}_{\mathcal{T}_{t} = \mathcal{S}_{t} \mathcal{X}_{t-1}}$$
(4)

where  $S_t$  is a spatial diffusion matrix, and  $T_t$  is a temporal diffusion matrix.

The meaning of  $\mathcal{S}_t$  is the result of diffusing the s-th column  $\mathbf{i}_s$  of  $\mathcal{I}_n$  through  $\mathcal{L}_t^{\text{twr}}$  for each node s. This is interpreted as the augmentation of a spatial locality of each node through RWR within  $\mathcal{G}_t$ . On the other hand,  $\mathcal{T}_t$  is the result of diffusing  $\mathbf{x}_{s,t-1}$  of  $\mathcal{X}_{t-1}$  through  $\mathcal{L}_t^{\text{twr}}$  for each node s. Note that  $\mathbf{x}_{s,t-1}$  contains the probabilities that the surfer visits each node starting from node s during the travel from the initial time to t-1. Thus, it spreads the past proximities of  $\mathbf{x}_{s,t-1}$  in the current  $\mathcal{G}_t$  through  $\mathcal{L}_t^{\text{twr}}$ , which consequently reflects the temporal information to  $\mathcal{G}_t$ .

The final diffusion matrix  $\mathcal{X}_t$  is a convex combination between  $\mathcal{S}_t$  and  $\mathcal{T}_t$  w.r.t.  $\gamma$ , which is denoted by  $\oplus$  in Figure 1. Notice that  $\mathcal{X}_t$  is a column stochastic transition matrix for every time step t, which is proved in (Lee and Jung 2022), implying that as an augmented adjacency matrix,  $\tilde{\mathcal{A}}_t$  can be replaced with  $\mathcal{X}_t^{\top}$  for the input of GNNs in Equation (1).

**Interpretation.** We further analyze how  $\mathcal{X}_t$  reflects the spatial and temporal information of the input dynamic graph. For this purpose, we first obtain the closed-form expression of  $\mathcal{X}_t$  which is described in Theorem 1.

**Theorem 1.** The closed-form expression of  $\mathcal{X}_t$  is:

$$\mathcal{X}_{t} = (1 - \gamma) \left( \sum_{i=0}^{t-2} \gamma^{i} \mathcal{L}_{t \leftrightarrow t-i}^{\text{rwr}} \right) + \gamma^{t-1} \mathcal{L}_{t \leftrightarrow 1}^{\text{rwr}}$$
 (5)

where  $\mathcal{L}_{j \leftarrow i}^{\text{rwr}} = \mathcal{L}_{j}^{\text{rwr}} \mathcal{L}_{j-1}^{\text{rwr}} \cdots \mathcal{L}_{i}^{\text{rwr}}$  for j > i, and  $\mathcal{L}_{i \leftarrow i}^{\text{rwr}} = \mathcal{L}_{i}^{\text{rwr}}$ .

*Proof.* It is proved by mathematical induction, and the detailed proof is described in (Lee and Jung 2022). □

In the theorem,  $\mathcal{L}_{j \leftarrow i}^{\text{rwr}}$  indicates a random walk diffusion traveling from former time i toward latter time j. According to Equation (5),  $\mathcal{X}_t$  is concisely represented as:

Augmentation of spatial and temporal localities

$$\mathcal{X}_{t} \propto \overbrace{\gamma^{0} \mathcal{L}_{t}^{\text{rwr}} + \gamma^{1} \mathcal{L}_{t \leftarrow t-1}^{\text{rwr}} + \cdots + \gamma^{t-2} \mathcal{L}_{t \leftarrow 2}^{\text{rwr}} + \frac{\gamma^{t-1}}{1 - \gamma} \mathcal{L}_{t \leftarrow 1}^{\text{rwr}}}^{\text{rwr}}$$

$$\leftarrow \text{Emphasized} \qquad \qquad \text{Decayed} \Longrightarrow$$

Note that  $\mathcal{X}_t$  is more affected by the information close to time t than that passed from the distant past. The influence of  $\mathcal{L}_{t\leadsto k}^{\text{rwr}}$  is decayed by  $\gamma$  as time k is further away from time t, while it is emphasized as time k is near to time t where the ratio  $\gamma$  is interpreted as a *temporal decay ratio*. This explanation is consistent with the temporal locality, i.e., the tendency that recent edges are more influential than older ones. Combined with the spatial diffusion  $\mathcal{L}_t^{\text{rwr}}$ , the result of  $\mathcal{X}_t$  augments both spatial and temporal localities in  $\mathcal{G}_t$ .

**Discussion.** TIARA is a generalized version of GDC with PPR kernel to dynamic graphs since TIARA with  $\beta=0$  spatially augments data within a single  $\mathcal{G}_t$  at each time, which is exactly what GDC does. However, GDC does not consider temporal information for its augmentation, and it performs worse than TIARA as shown in Tables 2 and 3.

#### Algorithm for TIARA

Most graph diffusions involve heavy computational cost, especially for a large graph, and result in a dense matrix. The computation of  $\mathcal{X}_t$  also exhibits the same issue, and thus we adopt approximate techniques to alleviate the problem. Including the approximate strategies, the procedure of TIARA is summarized in Algorithm 1 where  $\tilde{\mathcal{X}}_0$  is set to  $\mathcal{I}_n$ .

**Power iteration.** The main bottleneck for obtaining  $\mathcal{X}_t$  is to compute the matrix inversion  $\mathbf{L}_t^{-1}$  of  $\mathcal{L}_t^{\mathrm{rwr}}$  in Equation (4), which requires  $O(n^3)$  time. Instead of directly calculating the inversion, we use power iteration (lines  $9 \sim 15$ ) based on the following (Yoon, Jung, and Kang 2018):

$$\mathbf{L}_{t}^{-1} = \sum_{k=0}^{\infty} c^{k} \left( \tilde{\boldsymbol{\mathcal{A}}}_{t}^{\top} \right)^{k} \approxeq \sum_{k=0}^{K} c^{k} \left( \tilde{\boldsymbol{\mathcal{A}}}_{t}^{\top} \right)^{k}$$

where  $c=1-\alpha-\beta$ , and K is the number of iterations. Let  $\mathbf{M}_t^{(k)}$  be the result after k iterations; then, it is recursively represented as follows:

$$\mathbf{M}_t^{(k)} = \mathbf{\mathcal{I}}_n + c\tilde{\mathbf{\mathcal{A}}}_t^{\top} \mathbf{M}_t^{(k-1)}$$

where  $\mathbf{M}_t^{(0)} = \mathcal{I}_n$  and  $\mathbf{M}_t^{(K)} \cong \mathbf{L}_t^{-1}$ . Note  $\tilde{\mathcal{A}}_t$  is a normalized adjacency matrix (line 1) in which self-loops are added, as traditional GNNs usually do. The approximate error is bounded by  $c^k$ , and converges to 0 as  $k \to \infty$  (Lee and Jung 2022). After that, we set  $\mathcal{L}_t^{\mathrm{rwr}} \leftarrow (1-c)\mathbf{M}_t^{(K)}$  (line 13).

At a glance, each iteration seems to take  $O(n^3)$  time for the matrix multiplication, but it is much faster than that since each snapshot  $\mathcal{G}_t$  is sparse in most cases. More specifically, only a few nodes form edges at each time step in real graphs. We call such nodes *activated* where  $\mathcal{V}_t$  is the set of activated nodes at time t, and  $n_t = |\mathcal{V}_t|$ . In each  $\mathcal{G}_t$ , a surfer can move only between activated nodes, i.e., only pairs of nodes in  $\mathcal{V}_t$  are diffused. As seen in Table 1, the average  $\bar{n}_t$  of  $n_t$  over time is smaller than n except the Brain dataset.

This allows us to do the power iteration on the sub-matrix  $\tilde{\mathcal{A}}_{\mathcal{V}_t} \in \mathbb{R}^{n_t \times n_t}$  of  $\tilde{\mathcal{A}}_t$  for nodes in  $\mathcal{V}_t$  where  $m_t$  is the number of non-zeros of  $\tilde{\mathcal{A}}_{\mathcal{V}_t}$ . Then, an iteration takes  $O(m_t n_t)$  time for a sparse matrix multiplication. Note  $m_t$  is linearly proportional to  $n_t$  in real graphs, i.e.,  $m_t = C_t n_t$  where  $C_t$  is a constant. Let  $\bar{m}_t$  be the average number of edges over time. As seen in Table 1,  $\bar{C}_t = \bar{m}_t/\bar{n}_t$  is smaller than  $\bar{n}_t$ . Thus, each iteration takes  $O(n_t^2)$  time in average; overall, it takes  $O(n_t^2K + nK)$  time and  $O(n_t^2 + n)$  space for  $\mathcal{L}_t^{\text{rwr}}$  (as only  $n_t$  nodes are diffused). More details are provided in (Lee and Jung 2022).

**Sparsification.** Another bottleneck is that  $\mathcal{X}_t$  is likely to be dense by repeatedly multiplying  $S_t X_{t-1}$  (line 4) as time t increases where  $\mathcal{S}_t = \mathcal{L}_t^{\text{rwr}}$ . This could be problematic in terms of space as well as running time, especially for graph convolutions since  $\mathcal{X}_t$  is used as an adjacency matrix. To alleviate this issue, we adopt a sparsification technique suggested in (Klicpera, Weißenberger, and Günnemann 2019). As established in Theorem 1, the graph structure of  $\mathcal{X}_t$  is spatially and temporally localized, which allows us to drop small entries of  $\mathcal{X}_t$ , resulting in the sparse  $\tilde{\mathcal{X}}_t$ . For this, we use a filtering threshold  $\epsilon$  to set values of  $\mathcal{X}_t$  below  $\epsilon$  to zero (line 6). This strategy has two advantages. First, it keeps  $\mathcal{X}_t$ sparse at each time. Second, it reduces the cost for processing  $\mathcal{S}_t \hat{\mathcal{X}}_{t-1}$  as  $\mathcal{S}_t$  and  $\mathcal{X}_{t-1}$  are sparse. After the sparsification, we normalize  $\tilde{\mathcal{X}}_{t-1}$  (line 7) column-wise. As shown in Figure 4, this sparsification makes the augmentation process fast and lightweight with tiny errors while it does not harm predictive accuracy too much, or can even improve.

**Theorem 2** (Complexity Analysis). For each time step t, TIARA takes  $O(n_t n/\epsilon + n_t^2 K)$  time on average, and produces  $\tilde{X}_t$  consuming  $O(n/\epsilon)$  space where n is the number of total nodes,  $n_t$  is the number of activated nodes at time t, K is the number of iterations, and  $\epsilon$  is a filtering threshold.

*Proof.* The proof is provided in (Lee and Jung 2022).  $\Box$ 

**Discussion.** Theorem 2 implies TIARA shows faster than  $O(n^3)$ , and uses space less than  $O(n^2)$  for storing  $\tilde{\mathcal{X}}_t$  in most real dynamic graphs. Nevertheless, its time complexity can reach  $O(n^2)$  for a graph such as the Brain dataset; thus, for larger graphs, its scalability can be limited. However, TIARA is based on matrix operations which are easy-to-accelerate using GPUs, and other diffusion methods such as GDC lie at the same complexity. Furthermore, there are extensive works of efficient RWR computations (Andersen, Chung, and Lang 2006; Jung et al. 2017; Shin et al. 2015; Wang et al. 2017; Hou et al. 2021) and accelerated multiplications of sparse matrices (Srivastava et al. 2020), which can make TIARA scalable. In this work, we focus on effectively augmenting a dynamic graph, and leave further computational optimization on the augmentation as future work.

Algorithm 1: TIARA at time t

**Require:** adjacency matrix  $\mathbf{A}_t$ , previous time-aware diffusion matrix  $\tilde{\mathbf{X}}_{t-1}$ , restart probability  $\alpha$ , time travel probability  $\beta$ , number K of iterations, filtering threshold  $\epsilon$ 

```
Ensure: time-aware diffusion matrix \tilde{\mathcal{X}}_{t}
   1: \tilde{\mathcal{A}}_t \leftarrow \mathbf{D}_t^{-1} \mathbf{A}_t where \mathbf{D}_t = \operatorname{diag}(\mathbf{A}_t \mathbf{1})
  2: \mathcal{L}_t^{\text{rwr}} \leftarrow \text{Power-Iteration}(\tilde{\mathcal{A}}_t, \alpha, \beta, K)
  3: \boldsymbol{\mathcal{S}}_t \leftarrow \boldsymbol{\mathcal{L}}_t^{\text{rwr}}

    ▷ Spatial augmenter

  4: \mathcal{T}_t \leftarrow \mathcal{S}_t \tilde{\mathcal{X}}_{t-1}

    ▷ Temporal augmenter

  5: \mathcal{X}_t \leftarrow (1 - \gamma)\mathcal{S}_t + \gamma \mathcal{T}_t where \gamma = \beta/(\alpha + \beta)
  6: \tilde{\mathcal{X}}_t \leftarrow filter entries of \mathcal{X}_t if their weights are < \epsilon
  7: normalize \mathcal{X}_t column-wise
  8: return \tilde{\mathcal{X}}_t
  9: function POWER-ITERATION(\tilde{\mathcal{A}}_t, \alpha, \beta, K)
                    set c \leftarrow 1 - \alpha - \beta and \mathbf{M}_{t}^{(0)} \leftarrow \mathcal{I}_{n}
10:
                   \begin{aligned} & \text{for } k \leftarrow 1 \text{ to } K \text{ do} \\ & \mathbf{M}_t^{(k)} \leftarrow \mathcal{I}_n + c \tilde{\mathcal{A}}_t^\top \mathbf{M}_t^{(k-1)} \\ & \mathcal{L}_t^{\text{rwt}} \leftarrow (1-c) \mathbf{M}_t^{(K)} \text{ where } \mathbf{M}_t^{(K)} \cong \mathbf{L}_t^{-1} \end{aligned}
12:
13:
                    normalize \mathcal{L}_t^{\text{rwr}} column-wise and return \mathcal{L}_t^{\text{rwr}}
15: end function
```

Table 1: Summary of datasets. n and m are the total numbers of nodes and edges, resp. T and L are the numbers of time steps and labels, resp.  $\bar{n}_t$  and  $\bar{m}_t$  are the average numbers of activate nodes and edges over time, resp.  $\bar{C}_t = \bar{m}_t/\bar{n}_t$ . The first 3 data are used for link prediction, and the others are for node classification.

Datasets	n	m	T	L	$\lfloor \bar{n}_t \rfloor$	$\bar{C}_t$
BitcoinAlpha	3,783	31,748	138	2	105	2.2
WikiElec	7,125	212,854	100	2	354	6.0
RedditBody	35,776	484,460	88	2	2,465	2.2
Brain	5,000	1,955,488	12	10	5,000	32.6
DBLP-3	4,257	23,540	10	3	782	3.0
DBLP-5	6,606	42,815	10	5	1,212	3.5
Reddit	8,291	264,050	10	4	2,071	12.8

#### **Experiment**

In this section, we evaluate TIARA to show its effectiveness for the augmentation problem for dynamic graphs.

### **Experimental Setting**

**Datasets.** Table 1 summarizes 7 public datasets used in this work. BitcoinAlpha is a social network between bitcoin users (Kumar et al. 2016, 2018b). WikiElec is a voting network for Wikipedia adminship elections (Leskovec, Huttenlocher, and Kleinberg 2010). RedditBody is a hyperlink network of connections between two subreddits (Kumar et al. 2018a). For node classification, we use the following datasets evaluated in (Xu et al. 2019). Brain is a network of brain tissues where edges indicate their connectivities. DBLP-3 and DBLP-5 are co-authorship networks extraced from DBLP. Reddit is a post network where two posts were connected if they contain similar keywords.

**Baseline augmentation methods.** We compare TIARA to the following baselines. NONE indicates the result of a model without any augmentation. DROPEDGE is a drop-based method randomly removing edges at each epoch. GDC is a graph diffusion-based method where we use PPR

Table 2: Temporal link prediction accuracy (AUC) where NONE is a result without augmentation, and ▲ (or ▼) indicates improvement (or degradation) compared to None. TIARA shows consistent improvement across most models and datasets.

AUC		BitcoinAlph	a		WikiElec		RedditBody			
	GCN	GCRN	EGCN	GCN	GCRN	EGCN	GCN	GCRN	EGCN	
None	57.3±1.6	80.3±6.0	58.8±1.1	59.9±0.9	72.1±2.4	66.9±3.7	77.6±0.4	88.9±0.3	77.6±0.2	
DROPEDGE GDC MERGE	<b>&gt;</b> 56.3±1.0 <b>&gt;</b> 57.5±1.6 <b>&gt;</b> 66.8±2.6	73.9±2.2 77.3±6.5 93.1±0.4	▼57.4±0.9 ▼57.4±1.2 ▲61.0±9.2	▼50.1±1.0 ▲62.8±0.8 ▲60.6±1.7	56.0±9.3 67.9±1.0 68.4±3.2	V47.9±6.4 V63.1±0.7 V60.7±1.3	73.0±0.4 74.6±0.0 69.7±0.7	777.0±1.7 86.4±0.3 89.8±0.5	71.9±0.7 73.8±0.3 80.3±0.5	
TIARA	▲76.0±1.3	▲94.6±0.8	^77.2±1.4	▲69.0±1.2	^73.4±2.2	▲69.1±0.3	▲80.8±0.6	▲90.2±0.4	▲82.0±0.1	

for this as our approach is based on random walks. MERGE is a simple baseline merging adjacency matrices from time 1 to t when training a model at time t. We apply DROPEDGE and GDC to each snapshot since they are designed for a static graph.

Baseline GNNs. We use GCN (Kipf and Welling 2017), GCRN (Seo et al. 2018) and EvolveGCN (Pareja et al. 2020), abbreviated to EGCN, for performing dynamic graph tasks. We naively apply a static GCN to each graph snapshot for verifying how temporal information is informative. We choose GCRN and EvolveGCN, lightweight and popular dynamic GNN models showing decent performance, to observe practical gains from augmentation. We adopt GCN layers for GCRN's graph convolution. We use the implementation of (Rozemberczki et al. 2021) for GCRN and EGCN. Note that any GNN models following Problem 1 can utilize TIARA because our approach is model-agnostic.

**Training details.** For each dataset, we tune the hyperparameters of all models on the original graph (marked as None) and augmented graphs separately through a combination of grid and random search on a validation set, and report test accuracy at the best validation epoch. For TIARA, we fix K to 100, search for  $\epsilon$  in [0.0001, 0.01], and tune  $\alpha$  and  $\beta$  in (0,1) s.t.  $0<\alpha+\beta<1$ . We use the Adam optimizer with weight decay  $10^{-4}$ , and the learning rate is tuned in [0.01, 0.05] with decay factor 0.999. The dropout ratio is searched in [0,0.5]. We repeat each experiment 5 times with different random seeds, and report the average and standard deviation of test values. We use PyTorch and DGL (Wang et al. 2019) to implement all methods. All experiments were done at workstations with Intel Xeon 4215R and RTX 3090.

Details about the experimental setting are provided in (Lee and Jung 2022).

#### **Temporal Link Prediction Task**

This aims to predict whether an edge exists or not at time t+1 using the information up to time t. As a standard setting (Pareja et al. 2020), we follow a chronological split with ratios of training (70%), validation (10%), and test (20%) sets. We sample the same amount of negative samples (edges) to positive samples (edges) for each time, and use AUC as a representative measure. We set the number of epochs to 200 with early stopping of patience 50.

As shown in Table 2, TIARA consistently improves the performance of dynamic GNN models such as GCRN and

EGCN compared to None (i.e., without augmentation) while static augmentations of Dropede and GDC do not. Tiara also outperforms the static methods on all models and datasets. This indicates it is not beneficial to only spatially augment the graphs for this task. Tiara even improves static GCN, which is competitive with EGCN, implying that effectively and temporally augmented data can even make static GNNs learn dynamic graphs well. In addition, Merge also improves the accuracy of the tested models on many datasets. This confirms the need to utilize temporal information when it comes to dynamic graph augmentation in this task. However, Merge performs worse than Tiara in most cases because Tiara can effectively augment both spatial and temporal localities at once while Merge does not have a mechanism to enhance such localities.

#### **Node Classification Task**

This is to classify a label of each node where a graph and features change over time. Following (Xu et al. 2019), we split all nodes into training, validation, and test sets by the 7:1:2 ratio. We feed node embeddings  $\mathbf{H}_T$  of each model forward to a softmax classifier, and use Macro F1-score because labels are imbalanced in each dataset. We set the number of epochs to 1,000 with early stopping of patience 100.

Table 3 shows TIARA consistently improves the accuracies of GNNs on most datasets. Especially, TIARA significantly enhances the accuracies on the Brain dataset as another diffusion method GDC does, but TIARA shows better accuracy than GDC, implying it is effective to augment a temporal locality for the performance. For the other datasets, TIARA slightly improves each model, but it overall performs better than other augmentations. Note GCN and EGCN are worse than a random classifier of 1/L score (0.25 for  $L\!=\!4$ ) in the Reddit where L is the number of labels, and all tested augmentations fail to beat the score, implying even these augmentations could not boost a poor model in this task.

#### **Effect of Hyperparameters**

We analyze the effects of temporal decay ratio  $\gamma$  and filtering threshold  $\epsilon$  that mainly affect TIARA's results. We fix the number K of iterations to 100 for the power iteration, which leads to sufficiently accurate results for  $\mathcal{L}_t^{\text{rwr}}$ .

Effect of the temporal decay ratio  $\gamma$ . As TIARA's hyperparameters,  $\alpha$  and  $\beta$  should be analyzed, but our preliminary experiments showed that patterns vary by models and

Table 3: Node classification accuracy (Macro F1-score) where NONE is a result without augmentation, and ▲ (or ▼) indicates improvement (or degradation) compared to None. TIARA shows consistent improvement across most models and datasets.

Macro F1	Brain		Reddit			DBLP-3			DBLP-5			
	GCN	GCRN	EGCN	GCN	GCRN	EGCN	GCN	GCRN	EGCN	GCN	GCRN	EGCN
None	44.7±0.8	66.8±1.0	43.4±0.7	18.2±2.9	40.4±1.6	18.6±2.3	53.4±2.6	83.1±0.6	51.3±2.7	69.6±0.9	75.4±0.7	68.5±0.6
DROPEDGE	35.2±1.7	▲67.8±0.6	39.7±1.8	19.4±0.8	▼40.3±1.4	18.0±2.7	▲55.8±1.9	▲84.3±0.6	▲52.4±1.7	▲70.5±0.5	▲75.6±0.7	68.0±0.7
GDC	▲63.2±1.2	▲88.0±1.5	▲67.3±1.3	₹17.5±2.3	▲41.0±1.6	18.5±2.8	▲53.4±2.1	▲84.7±0.5	▲52.8±2.2	▲70.0±0.7	▲75.5±1.2	▲69.1±1.0
MERGE	₹34.4±3.4	63.2±1.6	▲53.0±0.9	▲19.3±3.0	▼39.6±0.8	20.4±3.0	▲54.9±3.1	*83.0±1.4	▲53.3±1.2	▲70.8±0.4	74.5±0.8	▲69.7±1.6
TIARA	▲68.7±1.2	▲91.3±1.0	^72.0±0.6	▲18.4±3.0	41.5±1.5	21.9±1.6	^57.5±2.2	▲84.9±1.6	▲56.4±1.8	^71.1±0.6	^77.9±0.4	^70.1±1.0

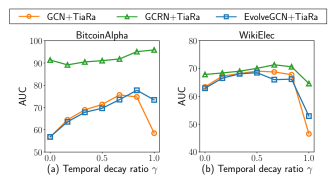


Figure 3: Effect of the temporal decay ratio  $\gamma$ .

datasets in the changes of  $\alpha$  and  $\beta$ . Instead, we narrow our focus to  $\gamma$  in Equation (4) where  $\gamma=\beta/(\alpha+\beta)$ . For this experiment, we vary  $\gamma$  from  $10^{-5}$  to  $1-10^{-5}$  by tweaking  $\alpha$  and  $\beta$  s.t.  $\alpha+\beta$  is fixed to 0.6. Figure 3 shows that too small or large values of  $\gamma$  can degrade link prediction accuracy except GCRN with TIARA in BitcoinAlpha. This implies that it is important to properly mix spatial and temporal information about the performance, which is controlled by  $\gamma$ .

Effect of the filtering threshold  $\epsilon$ . Figure 4 shows the effects of  $\epsilon$  in terms of approximate error, time, space, and accuracy of link prediction in BitcoinAlpha and WikiElec. We fix  $\alpha$  and  $\beta$  to 0.25, and vary  $\epsilon$  from  $10^{-7}$  to  $10^{-2}$  for this experiment.

We measure the approximate error  $\|\Delta \lambda_t\| = \|\lambda_t - \tilde{\lambda}_t\|_2$  of eigenvalues where  $\lambda_t$  and  $\tilde{\lambda}_t$  are vectors of eigenvalues of  $\mathcal{X}_t$  (i.e.,  $\epsilon = 0$ ) and  $\tilde{\mathcal{X}}_t$ , respectively, as similarly analyzed in (Klicpera, Weißenberger, and Günnemann 2019). The right y-axis of Figures 4(a) and (b) is the error, and the left y-axis is the number  $|\mathbf{A}_t|$  of edges in  $\mathbf{A}_t$ . As time t increases, the errors (red and blue lines) remain small, and do not explode, implying errors incurred by repeated sparsifications are not excessively accumulated over time. Rather, the errors tend to be proportional to  $|\mathbf{A}_t|$  at each time.

Figures 4 (c) and (d) show the space measured by  $\sum_t |\mathcal{X}_t|$  (left y-axis) and the augmentation time (right y-axis) of TIARA by  $\epsilon$  between  $10^{-7}$  and  $10^{-2}$ . As the strength of sparsification increase (i.e.,  $\epsilon$  becomes larger), the produced non-zeros and the augmentation time decrease. On the other hand, most of the accuracies remain similar except  $\epsilon = 10^{-2}$ 

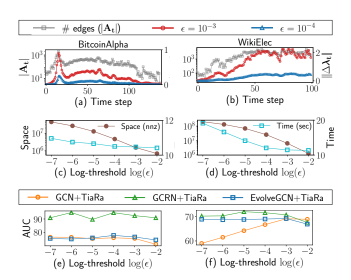


Figure 4: Effect of the filtering threshold  $\epsilon$ .

as shown in Figures 4(e) and (f). Note it is not effective to truncate too many entries (e.g.,  $\epsilon=10^{-2}$ ), or too dense  $\tilde{\mathcal{X}}_t$  can worse the performance as GCN in WikiElec. Thus, the sparsification with proper  $\epsilon$  such as  $10^{-3}$  or  $10^{-4}$  provides a good trade-off between error, time, space, and accuracy.

### **Conclusion**

In this work, we propose TIARA, a novel and modelagnostic diffusion method for augmenting a dynamic graph with the purpose of improvements in dynamic GNN models. We first extend Random Walk with Restart (RWR) to Timeaware RWR so that it produces spatially and temporally localized scores. We then formulate time-aware random walk diffusion matrices, and analyze how our diffusion approach augments both spatial and temporal localities in the dynamic graph. As graph diffusions lead to dense matrices, we further employ approximate techniques such as power iteration and sparsification, and analyze how they are effective for achieving a good trade-off between error, time, space, and predictive accuracy. Our experiments on various real-world dynamic graphs show that TIARA aids GNN models in providing better performance of temporal link prediction and node classification tasks.

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