Orca: Scalable Temporal Graph Neural Network Training with Theoretical Guarantees

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Representation learning over dynamic graphs is critical for many real-world applications such as social network services and recommender systems. Temporal graph neural networks (T-GNNs) are powerful representation learning methods and have achieved remarkable effectiveness on continuous-time dynamic graphs. However, T-GNNs still suffer from high time complexity, which increases linearly with the number of timestamps and grows exponentially with the model depth, causing them not scalable to large dynamic graphs. To address the limitations, we propose Orca, a novel framework that accelerates T-GNN training by non-trivially caching and reusing intermediate embeddings. We design a brand new and optimal cache replacement algorithm, called MRD, under a practical cache limit. MRD not only improves the efficiency of training T-GNNs by maximizing the number of cache hits but also reduces the approximation errors by avoiding keeping and reusing extremely stale embeddings. Meanwhile, we develop profound theoretical analyses of the approximation error introduced by our reuse schemes and offer rigorous convergence guarantees. Extensive experiments have validated that Orca can obtain two orders of magnitude speedup over the state-of-the-art baselines while achieving higher precision on large dynamic graphs.

CCS Concepts: • Information systems \rightarrow Data management systems; • Theory of computation \rightarrow Dynamic graph algorithms.

Additional Key Words and Phrases: temporal graph neural networks, cache replacement

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1 INTRODUCTION

Dynamic graphs serve as cornerstones in a broad spectrum of domains, including social network services, citation network analysis, e-commerce, and recommender systems. Real-world dynamic graphs can be generally modeled as temporal interactions between nodes. For instance, in social networks like Reddit, users interact with each other by leaving a comment or becoming a follower;

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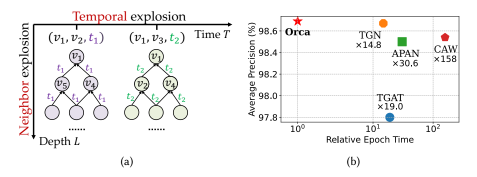


Fig. 1. (a) An illustration of the computational cost of T-GNNs, which increases linearly with the number of timestamps (i.e., temporal explosion) and grows exponentially with the model depth (i.e., neighbor explosion). (b) Comparison of Orca with the state-of-the-art baselines on the Reddit dataset [35].

in an e-commerce platform, customers interact with items by making a purchase. The above applications require learning representation on evolving graphs, where nodes and edges are dynamically updated, to support downstream tasks such as temporal link prediction or recommendation. However, the majority of previous representation learning methods [25, 28, 47, 73] assume that the input graph is *static*. Although it would be possible to apply static graph methods to dynamic graphs by ignoring the evolutionary patterns of dynamic structures, such solutions have been shown to be suboptimal [50, 70].

Temporal graph neural networks (T-GNNs) [35, 50, 65, 66, 70] are state-of-the-art methods for learning representations on dynamic graphs. In general, T-GNNs mimic the message passing of static GNNs [28, 62] by *temporal sampling* and *recursive neighborhood aggregation*. Specifically, TGAT [70] encodes continuous time using random Fourier features and applies attention modules for neighborhood aggregation; TGN [50] is a generic T-GNN framework that unifies several previous works [35, 59, 70] as special cases; CAW [66] further improves the inductive performance through anonymized random walks. It has been shown in [50, 66] that T-GNNs can outperform the static graph methods [25, 28, 34, 47, 62] and snapshot-based models [23, 26, 46] by a large margin in average precision.

Unfortunately, the high time complexity of T-GNNs hinders their applicability to large dynamic graphs. For instance, it takes over 12 hours for CAW [50] to finish one epoch of training on a large dynamic graph (7.8 million interactions) with an NVIDIA RTX 2080 Ti GPU; T-GNNs including JODIE [35] and TGN [50] cannot scale to large graphs due to high memory usage. Consequently, existing methods are typically evaluated on small dynamic graphs.

The computational bottlenecks of T-GNNs lie in $temporal\ explosion$ and $neighbor\ explosion$. As shown in Figure 1a, given a batch of two edges (v_1, v_2, t_1) and (v_1, v_3, t_2) , T-GNNs generally compute temporal embeddings of target nodes at t_1 and t_2 with two time-dependent computation graphs. On the one hand, unlike static GNNs where the representation of a node is computed only once, T-GNNs need to calculate the $intermediate\ embeddings$ of the same node at $all\ the\ involved\ timestamps$ in a data batch. For instance, the embeddings of node v_4 at an intermediate layer are computed twice. We refer to this problem as $temporal\ explosion$. On the other hand, T-GNNs suffer from $neighbor\ explosion$, which means that the computation graph of T-GNNs grows exponentially with the model depth due to recursive neighborhood aggregation [50]. The above two factors make it challenging to scale T-GNNs to large dynamic graphs.

To solve the above challenges, a straightforward approach would be to cache the embeddings at each intermediate layer and accelerate model training by judiciously reusing these cached results. Take Figure 1a as an example. After executing the computation graph at t_1 , we cache the intermediate embedding of node v_4 on GPU. Then at t_2 , we avoid recursive neighborhood aggregation for the node v_4 by reusing its cached embedding. When new data batches come, we can also reuse historical embeddings computed in previous batches. In this way, we effectively reduce the computational cost of T-GNNs and thus accelerate model training. However, such a caching and reuse scheme could have the following shortcomings.

- Limited GPU memory. Ideally, we can maximize the training efficiency by caching and reusing all the intermediates. However, this is impractical for large dynamic graphs due to limited GPU memory. For instance, it takes 32 GB of memory to cache all the intermediates for a 3-layer T-GNN with the embedding size 1024 on a 4M-node graph. Besides, edge features, model parameters, and gradients consume extra space. In contrast, a high-end GeForce RTX 3090 Ti GPU is equipped with only 24 GB of memory.
- **Performance drop.** While prior works on static GNNs [19, 45] simply cache all the intermediates, this practice is unfavorable for large dynamic graphs. Since model parameters and the graph structure evolve over time, cached embeddings would get stale. Reusing an incredibly stale embedding could introduce huge approximation errors and thus hurt model performance badly. The staleness problem can be especially serious on real-world dynamic graphs where the average frequency of updating each node's embedding is quite low.
- **Gradient explosion.** The computation graphs of vanilla T-GNNs at different timestamps are independent. However, sharing intermediate embeddings across multiple timestamps complicates the computation graph. It could lead to exploding gradients when updating model parameters in back-propagation, thus severely hurting model accuracy. According to our experiments, exploding gradients can degrade the accuracy of a 2-layer T-GNN by 4% on the Wikipedia dataset [5].

In this work, we propose Orca, a scalable caching-based framework for temporal graph neural networks. Orca solves the temporal explosion and neighbor explosion problems by sharing newly computed intermediates across multiple timestamps in a data batch and by reusing previously cached historical embeddings, respectively. The technical challenge is how to manage caches such that we can effectively reduce the recomputation cost without compromising model accuracy. We realize that the classical cache replacement policies like LRU [15] tend to cache all the newly computed embeddings even if they may not be reused in the near future. Thus, the caching policies could suffer from low cache hit ratios and stale embeddings that would cause large approximation errors. To address the problems, we have introduced the novel concept reuse distance for intermediate embeddings by analyzing the distribution of subsequent training batches. We then develop a caching policy MRD that can maximize the number of cache hits for offline T-GNN training and implicitly filter out extremely stale embeddings. We further prove that our proposed reuse schemes have a bounded approximation error on the model outputs. Moreover, Orca addresses the exploding gradient issue with an effective gradient blocking strategy. Although our gradient blocking and reuse schemes introduce approximate gradient values, we provide rigorous convergence guarantees for our proposed techniques.

In summary, we have made the following contributions:

- We present Orca, a framework that accelerates T-GNN training by caching and reusing intermediates. Orca reduces the time complexity of T-GNNs from $O(|E| \cdot k^L)$ to $O(|E| \cdot kL)$, where |E| is the batch size, L is the model depth, and k is the neighbor size.
- We formulate the cache replacement problem for training T-GNNs under practical cache limits and propose an optimal caching policy MRD. It not only improves the training efficiency of

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T-GNNs by maximizing the number of cache hits but also boosts model performance by resolving the staleness problem.

- We formulate the cache replacement problem for training T-GNNs under practical cache limits
 and propose an optimal caching policy MRD. It not only improves the training efficiency of
 T-GNNs by maximizing the number of cache hits but also boosts model performance by resolving
 the staleness problem.
- We propose a novel gradient blocking strategy to avoid exploding gradients caused by reusing intermediates, thus stabilizing model updates. Moreover, we theoretically investigate that T-GNNs trained with the approximate gradients can converge to a local optimum as vanilla T-GNNs do.
- We develop theoretical guarantees on the approximation errors introduced by reusing intermediates. Our theory indicates that the errors of approximate temporal embeddings can be bounded as long as the cached embeddings do not get too stale.

The remainder of this paper is organized as follows. We introduce the technical background in Section 2. Then, we present our Orca framework in Section 3 and study Orca under cache limits in Section 4. We develop approximation and convergence guarantees in Section 5. Finally, we conduct experiments in Section 6, review the related work in Section 7, and conclude this paper in Section 8.

2 BACKGROUND

In this section, we introduce the background of representation learning on dynamic graphs and a generic T-GNN architecture. Table 1 summarizes the major notations used throughout this paper.

2.1 Dynamic Graphs

In this work, we focus on *continuous-time* dynamic graphs (CTDGs) due to their generality and flexibility. CTDGs can come with edge features [35] or simply be non-attributed [1, 2]. Without loss of generality, we define the CTDG as follows.

DEFINITION 1 (CONTINUOUS-TIME DYNAMIC GRAPH (CTDG)). A CTDG is represented as a sequence of interaction events $G = \{\alpha(t_1), \alpha(t_2), \dots\}$ arranged in increasing order of time. Each event is a tuple $\alpha(t) = (v_i, v_j, e_{ij}(t), t)$ representing a (directed) temporal edge, where v_i and v_j are nodes, $e_{ij}(t)$ is an edge feature vector, and $t \in \mathbb{N}^+$ is the timestamp at which the interaction happens.

In social networks, for instance, a user v_i can follow, unfollow, or comment on another user v_j at timestamp t. Here, different behaviors are represented by different edge feature vectors $e_{ij}(t)$.

2.2 Embedding Learning on Dynamic Graphs

Embedding learning (also known as representation learning) problems on dynamic graphs typically follow an *encoder-decoder* framework [27]. Given an interaction event $\alpha(t) = (v_i, v_j, e_{ij}(t), t)$, an encoder function $f: \alpha(t) \to h_i(t), h_j(t)$ maps the event features to node embeddings, where $h_i(t) \in \mathbb{R}^d$ and $h_j(t) \in \mathbb{R}^d$ are temporal embeddings of node v_i and v_j at timestamp t, respectively. A decoder function (e.g., MLP [49]) then takes these temporal embeddings as input and makes prediction for downstream tasks such as link prediction [50, 65, 66] or dynamic node classification [35, 65].

2.3 Temporal Graph Neural Networks

Temporal graph neural networks (T-GNNs) are powerful models for learning temporal embeddings on CTDGs. Existing T-GNNs [50, 59, 70] follow a *generic* architecture and usually maintain a state vector $s_i(t)$ for each node v_i in the graph. The state vector $s_i(t)$ evolves with time and summarizes all the interactions that involve node v_i until timestamp t. Given a new interaction $\alpha(t) = (v_i, v_j, e_{ij}(t), t)$, an L-layer T-GNN works in the following two sequential steps, i.e., *temporal message passing* and *state update*.

Notation	Description							
G	a continuous-time dynamic graph							
$\alpha(t)$	an interaction that occurs at timestamp <i>t</i>							
v_i	a node in the dynamic graph							
$e_{ij}(t)$	feature of the edge occurring between v_i and v_j at t							
E	a batch of interactions events							
В	a set of target nodes that appear in <i>E</i>							
L	model depth							
k	limit on the sampled neighbor size							
M^{I}	cache at the <i>l</i> -th layer							
m	cache size limit at each layer							
$N_i^l(t)$	sampled neighbors of node v_i at layer l and time t							
$g_i^l(t)$	aggregated neighborhood of node v_i at layer l and time t							
$h_i^l(t)$	exact embedding of node v_i at layer l and time t							
$\tilde{h}_i^l(t)$	approximate embedding of v_i at layer l and time t							
$ar{h}_i^l$	cached embedding of node v_i at layer l ($\bar{h}_i^l \in M^l$)							
$s_i(t)$	the state vector of node v_i at time t							

Table 1. Summary of major notations.

• **Temporal Message Passing.** In this step, T-GNN computes the temporal embedding $h_i^L(t)$ of node v_i by *recursively aggregating* its *L*-hop neighborhood information as follows:

$$N_i^l(t) = \mathsf{SAMPLE}(G, v_i, t), \tag{1}$$

$$g_i^l(t) = \mathsf{AGGREGATE}(\{(h_j^{l-1}(t), e_{ij}(\tau), \phi(t-\tau))|$$

$$(j,\tau) \in N_i^l(t)\}, \forall l = 1, \dots, L,$$
(2)

$$h_i^l(t) = \text{COMBINE}(h_i^{l-1}(t), g_i^l(t)), \forall l = 1, ..., L,$$
 (3)

$$h_i^0(t) = s_i(t^-).$$
 (4)

At each layer l, T-GNN first samples from graph G a set of temporal neighbors $N_i^l(t)$, where each $(j,\tau) \in N_i^l(t)$ indicates that node v_j interacted with node v_i at timestamp τ ($\tau < t$) (Eq.1). Then, it aggregates the sampled neighborhood information (e.g., edge features and time encoding) of node v_i (Eq.2). Here, the function $\phi(\cdot)$ encodes the delta time between the current interaction $\alpha(t)$ and a sampled interaction $\alpha(\tau)$. Thus, the computation graphs of T-GNNs are time-dependent. A special case is that T-GNN aggregates the latest state vectors before t when computing the first layer embeddings (Eq.4). Finally, T-GNN computes the l-th layer output $h_i^l(t)$ by combing the previous layer output $h_i^{l-1}(t)$ and the aggregated neighborhood information $g_i^l(t)$ (Eq.3).

• **State Update.** After receiving a new event $\alpha(t) = (v_i, v_j, e_{ij}(t), t)$, T-GNN updates the state vectors of nodes v_i and v_i . Specifically, the state vector of node v_i is updated as follows:

$$s_i(t) = \text{UPDATE}(s_i(t^-), s_i(t^-), e_{ij}(t), t),$$
 (5)

where $s_i(t^-)$ and $s_j(t^-)$ are the latest state vectors before time t.

2.4 Training T-GNNs

T-GNNs are trained using batches of interactions. Given a new batch of interactions *E*, we first compute temporal embeddings of nodes appeared in these edges through *forward propagation*.

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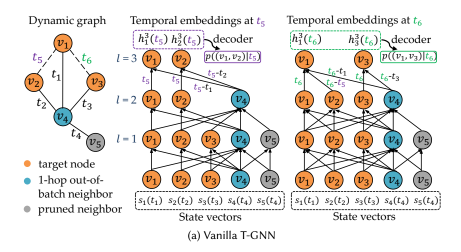


Fig. 2. An example of training a 3-layer T-GNN for a link prediction task. In the dynamic graph, the dotted lines with timestamps t_5 and t_6 ($t_5 < t_6$) represent a new batch of two edge interactions, while the solid lines with timestamps indicate historical interactions. To predict the existence of these two links, a T-GNN computes temporal embeddings of target nodes at t_5 and t_6 with two independent computation graphs, where each edge is associated with a time encoding. The temporal embeddings are then decoded to get link probabilities $p((v_1, v_2)|t_5)$ and $p((v_1, v_3)|t_6)$. Based on the predicted probabilities and the labels of the newly arrived edges, parameters of the T-GNN are updated via stochastic gradient descent (SGD).

Then, we use supervision signals from a downstream task to update model parameters via *backward propagation*. We illustrate the training procedure of T-GNNs as follows.

Example 1. Figure 2 shows an example of training a T-GNN for a link prediction task. There comes a new mini-batch of two interactions $\alpha(t_5)=(v_1,v_2,e_{12}(t_5),t_5)$ and $\alpha(t_6)=(v_1,v_3,e_{13}(t_6),t_6)$. In forward propagation, the T-GNN model computes temporal embeddings of the target nodes $\{v_1,v_2,v_3\}$ at t_5 and t_6 with two time-dependent computation graphs, where each edge is associated with a time encoding. The T-GNN model incorporates the new coming event $\alpha(t_5)$ when calculating the embedding $h_1^3(t_6)$ since $\alpha(t_5)$ happens before time t_6 , while the calculation of $h_1^3(t_5)$ is purely based on historical interactions before time t_5 . The temporal embeddings $h_1^3(t_5)$ and $h_2^3(t_5)$ are then fed into a decoder module to predict the existence of the true links and negatively sampled false links. Finally, the parameters of T-GNN are updated by stochastic gradient descent (SGD) [52] based on a predefined loss function.

Note that in the above example, node v_4 is sampled as a neighbor node at different timestamps. Then, T-GNN need to calculate its intermediate embeddings in two independent computation graphs. Compared to static GNNs, the computational cost of T-GNNs scales not only with the number of layer L but also with the number of timestamps. Hence, T-GNNs are much more computationally expensive, and it is difficult to scale them to large dynamic graphs.

Time Complexity. Let k be the maximum number of neighbors sampled for each target node in Eq.1. Then, given an L-layer T-GNN, there are $O(k^{L-l})$ ($1 \le l \le L$) nodes at the l-th layer in the computation graph. To calculate the representation of each node, a T-GNN needs to aggregate the neighborhood information of at most k neighbors. Therefore, the time complexity of T-GNNs is $O(|E| \cdot \sum_{l=1}^{L} k \cdot k^{L-l}) = O(|E| \cdot k^{L})$, where |E| is the batch size.

Algorithm 1: Orca via push-and-pull

```
input : Batch of interactions E, caches \{M^l\}_{l=1}^{L-1} ouput : Approximate temporal embeddings

1 // first layer

2 for \alpha(t) = (v_i, v_j, e_{ij}(t), t) \in E do

3 h_i^1(t), h_j^1(t) \leftarrow temporal message passing (Eq.1-Eq.4)

4 push h_i^1(t), h_j^1(t) to the cache M^1

5 // subsequent layers

6 for l = 2, \dots, L do

7 for \alpha(t) = (v_i, v_j, e_{ij}(t), t) \in E do

8 pull 1-hop neighbor embeddings of v_i, v_j from M^{l-1}

9 h_i^l(t), h_j^l(t) \leftarrow approximate temporal message passing (Eq.6-Eq.8)

10 if l \neq L then

11 push h_i^l(t), h_j^l(t) to the cache M^l
```

3 THE ORCA FRAMEWORK

As discussed in Section 2.4, the training cost of T-GNNs increases linearly with the number of unique timestamps (i.e., *temporal explosion*) and grows exponentially with the model depth *L* (i.e., *neighbor explosion*). To address the computational bottlenecks, we propose Orca, a scalable caching-based framework for temporal graph neural networks. Orca accelerates T-GNN training by caching and reusing intermediate embeddings in *forward propagation* (Section 3.1). Such an aggressive reuse scheme may cause severe accuracy drop due to the gradient explosion issue in *backward propagation*. Orca resolves this problem through a novel gradient blocking strategy (Section 3.2).

3.1 Acceleration via Reusing Embeddings

The basic idea of Orca is to aggressively prune the computation graph via a *push-and-pull* strategy. Without loss of generality, we consider an L-layer ($L \ge 2$) T-GNN with caches M^1, \ldots, M^{L-1} . Here, M^l dynamically maintains a set of intermediate embeddings that are computed at the l-th layer of the T-GNN. In this section, we assume that the storage space is big enough to accommodate all the intermediate embeddings. We will remove this assumption and study Orca under practical cache size limit in Section 4.

Given a batch of interaction events E, Orca works as described in Algorithm 1. Specifically, the first layer is treated as a special case, where Orca aggregates neighborhood features for target nodes v_i and v_j as vanilla T-GNN (line 3). This is because both Orca and vanilla T-GNN aggregate features based on state vectors without reusing historical embeddings. The difference is that Orca then pushes the first layer embeddings $h_i^1(t), h_j^1(t)$ to the cache M^1 (line 4). For a subsequent layer $l=2,\ldots,L$, Orca generally works in three steps: ① pull cached neighbor embeddings from the cache M^{l-1} (line 8), ② approximate the current layer output $\tilde{h}_i^l(t)$ with pulled embeddings (line 9), and ③ push the newly computed embeddings to the cache M^l ($l\neq L$) (lines 10-11). We elaborate the above three steps as follows.

• **Pull.** Given a target node v_i at timestamp t, Orca samples its 1-hop temporal neighbors $N_i^l(t)$ before feature aggregation (Eq.1). For each sampled neighbor node v_j , Orca pulls the corresponding cached embedding \bar{h}_i^{l-1} if there is one in the cache M^{l-1} . Otherwise, Orca uses zero embeddings

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for *cold-start* [54] nodes whose intermediate representations have not been cached yet. In this way, Orca prunes the computation graphs by *avoiding recursively computing the embedding for each neighbor node*. Note that the pulled embeddings are either computed in prior training iterations or newly computed in the current batch. Therefore, Orca not only reuses *historical* embeddings but also shares the *newly computed* embeddings across *all the possible timestamps* in a training data batch.

Approximate Message Passing. In analogous to the vanilla T-GNN, Orca conducts approximate
message passing based on cached embeddings as follows:

$$\begin{split} \tilde{g}_i^l(t) &= \mathsf{AGGREGATE}(\{(\bar{h}_j^{l-1}, e_{ij}(\tau), \phi(t-\tau)) \\ &| (j,\tau) \in N_i^l(t)\}), \forall l=2,\dots,L, \end{split} \tag{6}$$

$$\tilde{h}_i^l(t) = \text{COMBINE}(\tilde{h}_i^{l-1}(t), \tilde{g}_i^l(t)), \forall l = 2, \dots, L,$$

$$(7)$$

$$\tilde{h}_{i}^{1}(t) = h_{i}^{1}(t). \tag{8}$$

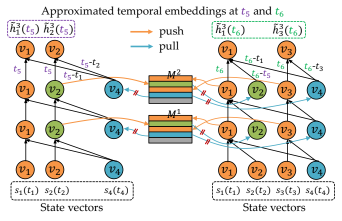
Specifically, Orca approximates the neighborhood information $\tilde{g}_i^l(t)$ by aggregating the pulled embeddings (Eq.6). Then, it combines the previous layer embedding $\tilde{h}_i^{l-1}(t)$ with $\tilde{g}_i^l(t)$ to get the l-th layer output $\tilde{h}_i^l(t)$ (Eq.7). We theoretically study the approximation errors introduced by reusing intermediates in Section 5.

• **Push.** After obtaining the representation $\tilde{h}_i^l(t)$ at the intermediate layer l ($l \neq L$), Orca pushes it to the cache M^l and *replaces* the previously cached intermediate of node v_i (lines 10-11). Note that the node v_i may appear in multiple interactions in the current batch E. For example, in Figure 3, node v_1 appears as a target node at timestamps t_5 and t_6 ($t_5 < t_6$). In such a case, only the intermediate embedding $\tilde{h}_1^l(t_6)$ with the latest timestamp t_6 will be kept in the cache M^l since the cache maintains at most one intermediate embedding for each node.

Example 2. Consider the link prediction task in Figure 2. Figure 3 shows how our push-and-pull strategy accelerates T-GNN training. Take the calculation process of $\tilde{h}_1^3(t_6)$ for example. On the one hand, after obtaining the intermediate representation $\tilde{h}_1^l(t_6)$ (l=1,2), Orca pushes it to the cache M^l so that it can be reused in subsequent training batches. On the other hand, Orca prunes the computation graph by pulling and reusing the cached embeddings of neighbor nodes v_2 and v_4 when calculating the (l+1)-th layer output $\tilde{h}_1^{l+1}(t_6)$. The pulled embedding \tilde{h}_2^l of node v_2 is exactly the newly computed representation $\tilde{h}_2^l(t_5)$, while the pulled embedding \tilde{h}_4^l was calculated in a previous batch.

Complexity. The time complexity of Algorithm 1 is $O(|E| \cdot kL)$, where k is the limit on the size of sampled neighbor set, and |E| is the batch size. In contrast, the vanilla T-GNN has a time complexity $O(|E| \cdot k^L)$. Therefore, our Orca framework is exponentially faster. Additionally, Algorithm 1 caches the embeddings of each node at each intermediate layer. As a result, its storage cost is $O(|V| \cdot (L-1))$, where V is the number of nodes in the dynamic graph.

Reuse Chance Analysis. The reuse chances of Orca come from two sources: out-of-batch neighbor nodes (e.g., node v_4 in Figure 3) and in-batch neighbor nodes (e.g., node v_2 in Figure 3). For out-of-batch neighbor nodes, Orca avoids neighbor explosion by directly reusing the historical embeddings computed in previous iterations. For in-batch neighbor nodes, Orca avoids temporal explosion by sharing the newly computed embeddings across multiple timestamps. According to Figure 8, there can be more in-batch neighbors than out-of-batch neighbors due to the temporal locality of real-world dynamic graphs.



(a) Orca via push-and-pull

Fig. 3. Orca accelerates model training by reusing cached embeddings. After obtaining intermediate results at the l-th layer ($1 \le l < L$), Orca first pushes the embeddings of target nodes $\{v_1, v_2, v_3\}$ to the cache M^l and then pulls embeddings of the neighbor nodes v_2 and v_4 from the cache M^l to calculate the (l+1)-th layer output. The green nodes indicate that the newly computed intermediates of the in-batch neighbor node v_2 are shared across multiple timestamps, while the blue nodes suggest that the previously cached historical embeddings of the in-batch neighbor node in0 are updated via SGD. Particuarly, as shown by the red slash symbols, gradients will not propagate through the reused embeddings in order to avoid gradient explosion.

The Staleness Problem. In contrast to classical caching problems in database systems, reusing previously cached intermediates to accelerate training in T-GNNs inevitably leads to approximation errors. During training, model parameters and graph structure keep changing. As a result, the input features, neighbor set, and model parameters required to calculate intermediate embeddings for a given target node vary between different training batches. Thus, previously computed intermediates may become outdated relative to the current batch, potentially introducing errors. To address the staleness problem, we propose a cache replacement algorithm to filter out those extremely stale embeddings in Section 4. Moreover, we theoretically investigate how the staleness problem affects model outputs and the convergence rate in Section 5.

3.2 Effective Training via Gradient Blocking

Although our push-and-pull scheme (Algorithm 1) reduces the computational cost of T-GNNs in *forward propagation*, it brings new challenges to gradient update in *backward propagation*. To achieve stable and effective model training, the critical problem here is how to manage gradient flows with cached embeddings. We discuss this problem in two cases, i.e., reusing *newly computed* embeddings and reusing *historical* embeddings.

Gradient Flow with Newly Computed Embeddings. Consider a newly computed embedding $\bar{h}_i^l \in M^l$ that is shared across multiple timestamps in forward propagation. By the chain rule of differentiation, gradients from multiple links can propagate through it in back-propagation. However, such a gradient flow scheme may cause *exploding gradients* [29], thus hurting model accuracy. For instance, in Figure 3, the embedding $h_2^1(t_5)$ is used to compute $\tilde{h}_2^2(t_5)$ and $\tilde{h}_1^2(t_6)$. Therefore, in back-propagation, gradients from these two links will accumulate on $h_2^1(t_5)$ and may produce a large gradient value (i.e., exploding gradient). In contrast, for the vanilla T-GNN

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Algorithm 2: Orca via reuse-or-recompute

```
input: Batch of events E, caches \{M_{T-1}^l\}_{l=1}^{L-1}, cache size m ouput: Approximate temporal embeddings

1 B_T = \{(v_i, t), (v_j, t) | (v_i, v_j, e_{ij}(t), t) \in E_T\} // target nodes

2 \tilde{H}_{B_T}^L \leftarrow \text{Recursion}(B_T, L, m) // model output

3 return \tilde{H}_{B_T}
```

Algorithm 3: Recursion

```
input : Target nodes B, depth l, cache size m
ouput: Approximate embeddings \tilde{H}_B^l

1 if l=1 then

2 \lfloor H_B^l \leftarrow temporal message passing (Eq.1-Eq.4)

3 else

4 \vert O \leftarrow sampled out-of-batch neighbor nodes of B

5 \vert U \leftarrow uncached neighbor nodes

6 \vert B' \leftarrow B \cup U

7 \vert \tilde{H}_{B'}^{l-1} \leftarrow Recursion(B', l-1, m)

8 \vert \tilde{H}_{O}^{l-1}, M_{I}^{l-1} \leftarrow push_and_pull (M_{I-1}^{l-1}, \tilde{H}_{B'}^{l-1}, O, m)

9 \vert \tilde{H}_B^l \leftarrow approximate message passing (Eq.6-Eq.8)
```

in Figure 2, the embedding $h_2^1(t_5)$ will not partipate in the computation at other timestamps. Hence, the exploding gradient issue will not occur on vanilla T-GNNs. According to Figure 6, the gradient explosion issue can be especially severe if embeddings are aggressively shared across many timestamps in a data batch.

Gradient Flow with Historical Embeddings. Given a historical embedding $\bar{h}_i^l \in M^l$ that is reused in forward propagation, it is impractical for the gradient reaching \bar{h}_i^l to propagate through it in back-propagation. The reason is that the embedding \bar{h}_i^l was computed in a previous batch, and the computation graph for calculating it was already discarded after finishing model training on that batch. Moreover, recursively propagating gradients across multiple training batches can also cause unstable model updates.

Gradient Blocking. Based on the above discussions, we propose to block all the gradient flows reaching the reused embeddings in back-propagation. Specifically, as shown by the red slash symbols, we do not allow gradients to propagate through the reused embeddings. Hence, gradients that originate from different timestamps will not accumulate anymore. In this way, we can effectively avoid exploding gradients, thus stabilizing model training. Although our blocking approach results in incomplete gradient values, we show later through Theorem 6 that the approximate gradient is asymptotically unbiased. Therefore, T-GNNs trained with such approximate gradients can still converge to a local optimum.

4 ORCA UNDER CACHE SIZE LIMIT

Note that Algorithm 1 assumes that the intermediates of all the nodes in a dynamic graph can be cached. In this section, we scale Orca to large dynamic graphs with limited cache size. The

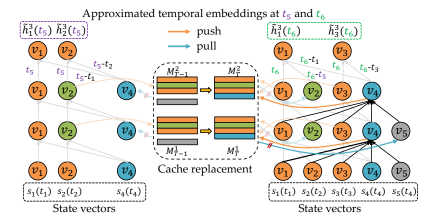


Fig. 4. Orca via reuse-or-recompute under cache size limit. Consider the link prediction task in Figure 2 and our Orca with cache size constraint m = 4. Assuming the historical embeddings of node v_4 are not cached, Orca needs to recompute its intermediate representations. Compared to the case without cache limit (Figure 3), the extra computation and data flow are highlighted.

motivation for restricting the cache size is to fit intermediates in limited GPU memory and mitigate the staleness problem of cached embeddings. We introduce our *reuse-or-recompute* scheme in Section 4.1, formally define the cache replacement problem for T-GNNs in Section 4.2, develop an optimal caching policy called MRD in Section 4.3, and prove the optimality of MRD in Section 4.4.

4.1 Reuse-or-Recompute

To compute the embedding of a target node v, Orca adopts a *reuse-or-recompute* scheme under cache limit. The basic idea is to reuse intermediates of v's neighbor nodes if they are cached; otherwise, Orca recomputes the corresponding representations.

Algorithm 2 depicts our reuse-or-recompute algorithm, which recursively invokes Algorithm 3 to compute the representations at each layer. Specifically, given a set of target nodes B at layer l, we first collect the corresponding 1-hop out-of-batch neighbor nodes O and uncached 1-hop neighbor nodes $U \subseteq O$ (lines 4-5 in Algorithm 3). Then, we calculate the (l-1)-layer representations of $B' = B \cup U$ by recursion (line 7). We next push the newly computed intermediate embeddings $\tilde{H}_{B'}^{l-1}$ into the cache M_T^{l-1} and pull the cached embeddings \tilde{H}_O^{l-1} of out-of-batch neighbor nodes O (line 8). Meanwhile, the cache is updated with $\tilde{H}_{B'}^{l-1}$ following the predefined cache size limit m (see Section 4.3). Finally, we get the l-th layer output by approximate message passing (line 9). Figure 4 gives an example of how our reuse-or-recompute scheme works.

Time Complexity. Compared to Algorithm 1, it takes extra time for Algorithm 2 to compute the intermediates of *out-of-batch neighbor nodes* whose historical embeddings are not cached. Let n be the total number of such uncached nodes when training a T-GNN on the batch E. Then, the time complexity of Algorithm 2 is $O(|E| \cdot kL + nk)$. We will empirically show in Section 6.2 that the recomputation cost of Algorithm 2 can be marginal since the number of our-of-batch neighbors can be smaller than that of in-batch neighbors.

4.2 Cache Replacement Problem

A critical issue in Algorithm 3 is dynamically managing the caches so that we can mitigate the staleness problem of cached intermediates and enhance the efficiency of *offline* model training

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by minimizing the amount of recomputation. In offline T-GNN training, we consider a sequence of batches E_1, E_2, \ldots, E_T , where each batch E_t contains a set of interactions. Let B_t be the set of target nodes that appear in E_t and O_t be the set of 1-hop out-of-batch neighbors sampled for B_t . Orca needs to recompute the representations of uncached nodes U_t ($U_t \subseteq O_t$) and then updates the cache with the newly computed intermediates of $B_t \cup U_t$. Given the above preliminaries, we formally define the cache replacement problem as follows.

DEFINITION 2 (CACHE REPLACEMENT PROBLEM). Consider a sequence of offline training batches E_1, \ldots, E_T . Let O_t be the set of 1-hop out-of-batch neighbors sampled for the target nodes B_t . Denote by M_t the set of cached embeddings after finishing model training on the batch E_t . We aim to find a cache replacement strategy $f(\cdot)$ that maximizes the number of cache hits under the cache size limit m:

$$\max_{f} \sum_{t=1}^{T} \sum_{v_{i} \in O_{t}} 1\{\bar{h}_{i} \in M_{t-1}\} = \min_{f} \sum_{t=1}^{T} |U_{t}|,$$

$$M_{t} = f(M_{t-1}, H_{B_{t}} \cup H_{U_{t}}), \sum_{\bar{h}_{i} \in M_{t}} cost(\bar{h}_{i}) \leq m, \forall t = 1, \dots, T,$$
(9)

where H_{B_t} and H_{U_t} are the newly computed embeddings of the target nodes B_t and uncached 1-hop neighbor nodes $U_t \subseteq O_t$, respectively.

Without loss of generality, we drop the superscript of layer l in the above problem definition since we cache intermediate results of *the same set of nodes* at different layers. In real-world applications, models are often updated *offline* and then deployed to production if they pass the A/B test [6, 12, 38, 67]. Our basic idea is to minimize the amount of recomputation by maximizing the total number of cache hits in all the offline training batches. In practice, we consider m as the maximum number of items allowed and set the cost of each cached embedding \bar{h}_i to be $cost(\bar{h}_i) = 1$. Hardness. The cache replacement problem (CRP) is a variant of the general caching problem [7, 9] and the multidimensional knapsack problem [20]. CRP is NP-hard in the *general* form, where different items can have different storage costs. However, in T-GNNs, all the intermediate representations at a specific layer l have the same dimensionality. Therefore, we consider the *uniform* case and then develop an optimal cache replacement algorithm in Section 4.3.

4.3 Minimum Reuse Distance Algorithm

In this section, we develop an *optimal* algorithm for the cache replacement problem. Given an intermediate embedding h_v , we define its *maximum active time* and *reuse distance* as follows.

DEFINITION 3 (MAXIMUM ACTIVE TIME). Consider an intermediate embedding h_v of node v that is created at the data batch E_i . Let E_j (i < j) be the latest batch where the node v appears as a target node (i.e., $v \in B_i$). Then, the maximum active time of the intermediate h_v is $\gamma(h_v) = j - 1$.

As the node v appears in batch E_j as a target node, our Orca will compute its latest intermediate representations. Hence, the historical embedding h_v computed at the previous batch E_i will never be reused when training a model on the batch E_j and afterward.

DEFINITION 4 (REUSE DISTANCE). Consider an intermediate embedding h_v of node v after updating model on the batch E_i . Let E_j (i < j) be the latest batch where the node v appears as a 1-hop out-of-batch neighbor node (i.e., $v \in O_j$). Then, the reuse distance of h_v right after finishing model training on the batch E_i is

$$\Delta(h_v, i) = \begin{cases} j - i, j \le \gamma(h_v) \\ \infty, j > \gamma(h_v). \end{cases}$$

Algorithm 4: Minimum reuse distance (MRD) algorithm

```
input :Offline training batches \{E_t\}_{t=1}^T, initial cache M_0, cache size m ouput: Cache plans \{M_t\}_{t=1}^T

1 for t = 1, ..., T do //data preparation

2 B_t \leftarrow target nodes in the batch E_t

3 N_t \leftarrow sampled neighbor nodes of B_t (Eq.1)

4 O_t \leftarrow N_t \setminus B_t

5 for t = 1, ..., T do //generate cache plans

6 M' \leftarrow M_{t-1} \cup B_t \cup O_t //candidate set

7 M_t \leftarrow nodes in M' with the top-m minimum reuse distance (ties are broken arbitrarily)

8 return \{M_t\}_{t=1}^T
```

Based on the above definition, we propose the minimum reuse distance (MRD) algorithm that can generate optimal cache plans. As shown in Algorithm 4, the basic idea of MRD is to cache intermediates that are most likely to be reused in the near future. Given a sequence of offline training batches, MRD first extracts the target nodes B_t and out-of-batch neighbor nodes O_t (lines 1-4). The sampling results (line 3) will be reused in offline training. Then, in each iteration t, MRD selects nodes with the top-m minimum reuse distance from the candidate set $M_{t-1} \cup B_t \cup O_t$ with ties broken arbitrarily (lines 5-7). Note that MRD only generates cache plans instead of a set of cached embeddings. Based on these plans, we dynamically update caches at all the intermediate layers in the offline training stage.

Time Complexity. The time complexity of the data preparation step in Algorithm 4 is dominated by the sampling operation (line 3). The sampling cost can be amortized by reusing the sampled results in the offline training stage. For each node in the candidate set, it takes O(1) time to get the reuse distance (line 6). Then, it takes O(|M'|) time to select the top-m nodes (line 7). Therefore, the time complexity of Algorithm 4 is $O(T \cdot (|E| \cdot c + |M|))$, where |E| is the maximum batch size, c is the sampling cost given a target node, and |M| is the maximum candidate size.

Comparison with Classical Caching Policies. Note that the classical cache replacement policies like LRU [15] and 2Q [32] are designed for the single-point query. In contrast, MRD can make a better cache plan by analyzing the dependencies between different items in a data batch and the distribution of subsequent training batches. On the other hand, LRU tends to cache all the newly computed embeddings even if they may not be reused in the near future. Thus, LRU could fill the cache with stale embeddings of some cold nodes. Instead, by its principle, our MRD policy does not cache intermediates with large reuse distances (i.e., embeddings of cold nodes) in the first place. Therefore, MRD can implicitly avoid keeping and reusing extremely stale embeddings that would introduce large approximation errors. We will empirically compare MRD with classical caching policies in Section 6.3.

4.4 Proof of Optimality

We follow the dynamic programming technique [51] to prove that our MRD policy (Algorithm 4) is optimal in maximizing the number of cache hits. Given a cache M_{t-1} , let $J_t(M_{t-1})$ be the maximum possible number of cache hits starting with the batch E_t . $J_t(M_{t-1})$ is recursively defined as

$$J_t(M_{t-1}) = \sum_{v_i \in O_t} \mathbf{1}\{\bar{h}_i \in M_{t-1}\} + \max_{M_t \in C_t(M_{t-1})} J_{t+1}(M_t), \tag{10}$$

$$C_t(M_{t-1}) = \{ M \subseteq M_{t-1} \cup H_{B_t} \cup H_{U_t} : |M| = m \}, \tag{11}$$

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where $C_t(M_{t-1})$ is the set of candidate cache states.

Consider two caches M_{t-1} and M'_{t-1} ($|M_{t-1}| = |M'_{t-1}| = m$), we define the *potential advantage* of M'_{t-1} over M_{t-1} as

$$d_t(M_{t-1}, M'_{t-1}) = \min_{a} \{ \bar{h} \in M_{t-1} | \Delta(\bar{h}, t-1) > \Delta(g(\bar{h}), t-1) \}, \tag{12}$$

where $g: M_{t-1} \to M'_{t-1}$ is a bijection between items in the two cache states M_{t-1} and M'_{t-1} . $d_t(M_{t-1}, M'_{t-1})$ compares the reuse distance of all the items in M_{t-1} and M'_{t-1} . A larger $d_t(M_{t-1}, M'_{t-1})$ value indicates that items in M'_{t-1} generally have smaller reuse distance and thus may lead to more cache hits than M_{t-1} . We next prove the following lemma.

LEMMA 3.
$$J_t(M'_{t-1}) - J_t(M_{t-1}) \le d_t(M_{t-1}, M'_{t-1})$$
 for all $1 \le t \le T+1$ and $|M_{t-1}| = |M'_{t-1}|$.

PROOF. If t=T+1, $J_t(M'_{t-1})=J_t(M_{t-1})=0$ and $d_t(M_{t-1},M'_{t-1})=0$. Therefore, the inequality holds for the special case t=T+1. If $t\leq T$, we proceed by induction. Assume that $J_{t+1}(M'_t)-J_{t+1}(M_t)\leq d_{t+1}(M_t,M'_t)$ for all $|M_t|=|M'_t|$. Let $M'_t\in C_t(M'_{t-1})$ be chosen by an optimal strategy and $M_t\in C_t(M_{t-1})$ be chosen by our MRD algorithm, respectively. We study the relationship between $d_{t+1}(M_t,M'_t)$ and $d_t(M_{t-1},M'_{t-1})$ by analyzing whether the historical embedding \bar{h}_i of each node $v_i\in O_t$ is cached.

- $\bar{h}_i \in M_{t-1}$ and $\bar{h}_i \in M'_{t-1}$. After reusing \bar{h}_i , we can either keep \bar{h}_i or evict it. Since MRD evicts items with the largest reuse distance, then $d_{t+1}(M_t, M'_t) \leq d_t(M_{t-1}, M'_{t-1})$.
- $\bar{h}_i \notin M_{t-1}$ and $\bar{h}_i \notin M'_{t-1}$. After recomputing h_i , we can either cache h_i or discard it. Since MRD evicts items with the largest reuse distance, then $d_{t+1}(M_t, M'_t) \leq d_t(M_{t-1}, M'_{t-1})$.
- $\bar{h}_i \in M_{t-1}$ and $\bar{h}_i \notin M'_{t-1}$. Before updating model with the batch E_t , \bar{h}_i contributes to the advantage of M_{t-1} by 1 in Eq.12. After that, M_{t-1} loses its relative advantage by 1. Then, $d_{t+1}(M_t, M'_t) \le d_t(M_{t-1}, M'_{t-1}) + \sum_{v_i \in O_t} \mathbf{1}\{\bar{h}_i \in M_{t-1} \text{ and } \bar{h}_i \notin M'_{t-1}\}$.
- $\bar{h}_i \notin M_{t-1}$ and $\bar{h}_i \in M'_{t-1}$. Before updating model with the batch E_t , \bar{h}_i contributes to the advantage of M'_{t-1} by 1 in Eq.12. After that, M'_{t-1} loses its relative advantage by 1. Then, $d_{t+1}(M_t, M'_t) \le d_t(M_{t-1}, M'_{t-1}) \sum_{v_i \in O_t} 1\{\bar{h}_i \notin M_{t-1} \text{ and } \bar{h}_i \in M'_{t-1}\}$.

Summarize all the above four cases, we get

$$d_{t+1}(M_t, M'_t) + \sum_{v_i \in O_t} \mathbf{1}\{\bar{h}_i \in M'_{t-1}\}$$

$$\leq d_t(M_{t-1}, M'_{t-1}) + \sum_{v_i \in O_t} \mathbf{1}\{\bar{h}_i \in M_{t-1}\}.$$
(13)

Based on the above discussion, we can obtain

$$J_{t}(M'_{t-1}) = \sum_{v_{i} \in O_{t}} 1\{\bar{h}_{i} \in M'_{t-1}\} + J_{t+1}(M'_{t})$$

$$\leq \sum_{v_{i} \in O_{t}} 1\{\bar{h}_{i} \in M'_{t-1}\} + J_{t+1}(M_{t}) + d_{t+1}(M_{t}, M'_{t})$$

$$\leq \sum_{v_{i} \in O_{t}} 1\{\bar{h}_{i} \in M_{t-1}\} + J_{t+1}(M_{t}) + d_{t}(M_{t-1}, M'_{t-1})$$

$$\leq J_{t}(M_{t-1}) + d_{t}(M_{t-1}, M'_{t-1}), \tag{14}$$

where the first inequality follows the inductive assumption, the second inequality follows Eq.13, and the last inequality follows the definition of $J_t(M_{t-1})$. The proof of Lemma 3 is completed. \Box

Theorem 4. The MRD algorithm is optimal in maximizing the number of cache hits for the cache replacement problem (Definition 2).

PROOF. Consider a cache M_{t-1} . Let M_t and M'_t be the successive cache states selected by our MRD algorithm and the optimal algorithm, respectively. Since M_t minimizes $d_{t+1}(M_t, M'_t)$ over the candidate set $C_t(M_{t-1})$, then $d_{t+1}(M_t, M'_t) = 0$. Finally, we get

$$0 \le J_{t+1}(M_t') - J_{t+1}(M_t) \le d_{t+1}(M_t, M_t') = 0, \tag{15}$$

where the first inequality follows the optimality of M'_t and the second inequality follows Lemma 3. Then, Eq.15 indicates that our MRD algorithm is optimal.

5 THEORETICAL ANALYSIS

In this section, we develop theoretical analysis on the approximation error introduced by our reuse schemes in Section 5.1, and offer convergence guarantees for our Orca framework in Section 5.2.

5.1 Approximation Guarantee

We propose a *unified* theoretical framework for bounding the approximation error introduced by *cache-unlimited* Orca (Algorithm 1) and *cache-limited* Orca (Algorithm 2). Specifically, given an L-layer T-GNN and a target node v_i at timestamp t, we aim to bound the approximation error between the exact embedding $h_i^L(t)$ and the approximate embedding $\tilde{h}_i^L(t)$.

Layer-wise Approximation Error. Let $f_1(\cdot) = \mathsf{COMBINE}(\cdot)$ and $f_2(\cdot) = \mathsf{AGGREGATE}(\cdot)$ be λ_1 and λ_2 Lipschitz continuous, i.e., $\|f_1(x_1) - f_1(x_2)\| \le \lambda_1 \|x_1 - x_2\|$ and $\|f_2(y_1) - f_2(y_2)\| \le \lambda_2 \|y_1 - y_2\|$. The Lipschitz properties are widely used in model analysis. We refer readers to [8, 19, 21, 63] for the Lipschitz constants of various combination and aggregation functions used in GNNs. Based on the Lipschitz property, we have the following upper bound on the l-th layer approximation error:

$$||h_{i}^{l}(t) - \tilde{h}_{i}^{l}(t)|| = ||f_{1}(h_{i}^{l-1}(t), g_{i}^{l}(t)) - f_{1}(\tilde{h}_{i}^{l-1}(t), \tilde{g}_{i}^{l}(t))||$$

$$\leq \lambda_{1}(||h_{i}^{l-1}(t) - \tilde{h}_{i}^{l-1}(t)|| + ||g_{i}^{l}(t) - \tilde{g}_{i}^{l}(t)||).$$

$$(16)$$

Moreover, since the aggregation function $f_2(\cdot)$ is λ_2 -Lipschitz continuous, we can obtain the following upper bound on the approximation error of the aggregated neighborhood feature:

$$||g_{i}^{l}(t) - \tilde{g}_{i}^{l}(t)|| \leq \lambda_{2} \sum_{(j,\tau) \in R_{i}^{l}(t)} ||h_{j}^{l-1}(\tau) - \tilde{h}_{j}^{l-1}|| + \lambda_{2} \sum_{(j,\tau) \in N_{i}^{l}(t) \setminus R_{i}^{l}(t)} ||h_{j}^{l-1}(\tau) - \tilde{h}_{j}^{l-1}(t)||,$$

$$(17)$$

where $N_i^l(t)$ is a set of sampled temporal neighbors, $R_i^l(t) \in N_i^l(t)$ denotes the neighbors whose intermediate embeddings are cached and reused, and $N_i^l(t) \setminus R_i^l(t)$ refers to the nodes whose embeddings are newly computed.

For any node v_i at timestamp t, assume that the error of the approximate embedding $\tilde{h}_i^{l-1}(t)$ and the staleness of the cached embedding \bar{h}_i^{l-1} are bounded, i.e., $\|h_i^{l-1}(t) - \tilde{h}_i^{l-1}(t)\| \leq \delta^{(l-1)}$ and $\|h_i^{l-1}(t) - \tilde{h}_i^{l-1}\| \leq \epsilon^{(l-1)}$. Let k be the maximum number of neighbors sampled in Eq.1. Plug Eq.17 into Eq.16, then

$$||h_{i}^{l}(t) - \tilde{h}_{i}^{l}(t)|| \leq \lambda_{1}(||h_{i}^{l-1}(t) - \tilde{h}_{i}^{l-1}(t)|| + ||g_{i}^{l-1}(t) - \tilde{g}_{i}^{l-1}(t)||)$$

$$\leq \lambda_{1}(\delta^{(l-1)} + k\lambda_{2}(\delta^{(l-1)} + \epsilon^{(l-1)}))$$

$$= \lambda_{1}(1 + k\lambda_{2})\delta^{(l-1)} + k\lambda_{1}\lambda_{2}\epsilon^{(l-1)} = \delta^{(l)}.$$
(18)

The above recursion formula indicates that the l-th layer approximation error $\delta^{(l)}$ is composed of the (l-1)-layer approximation error $\delta^{(l-1)}$ and the staleness $\epsilon^{(l-1)}$ of cached embeddings. There is one special case. The 0-layer embedding $h_i^0(t)$ corresponds to the state vector $z_i(t^-)$ of node v_i , i.e., $h_i^0(t) = z_i(t^-)$. On the other hand, the stale embedding $\bar{h}_i^0(t)$ and the approximate embedding $\tilde{h}_i^0(t)$ correspond to the stale state vector $\bar{z}_i(t^-)$, i.e., $\bar{h}_i^0(t) = \bar{h}_i^0(t) = \bar{z}_i(t^-)$. Hence, the input error can be bounded by

$$||h_i^0(t) - \tilde{h}_i^0(t)|| = ||h_i^0(t) - \bar{h}_i^0(t)|| = ||z_i(t^-) - \bar{z}_i(t^-)|| \le \epsilon^{(0)}.$$
(19)

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Since Orca updates the state vectors of all the target nodes as the vanilla T-GNN, there is no staleness in node states, i.e., $\epsilon^{(0)} = 0$.

The Upper Bound. Based on the recursion formula Eq.18, we stack all the layers together and develop the following theorem, which holds with and without cache limit.

Theorem 5. Consider an L-layer T-GNN with COMBINE(·) and AGGREGATE(·) functions that are λ_1 and λ_2 continuous, respectively. Assume that for any node v_i at timestamp t, its cached embedding \bar{h}_i^{l-1} has bounded staleness, i.e., $\|h_i^{l-1}(t) - \bar{h}_i^{l-1}\| \le \epsilon^{(l-1)}$. Then, the output error of the approximate temporal embedding $\tilde{h}_i^L(t)$ is bounded by

$$||h_i^L(t) - \tilde{h}_i^L(t)|| \le \sum_{l=1}^{L-1} k(1 + k\lambda_2)^{L-l-1} \lambda_1^{L-l} \lambda_2 \epsilon^{(l)}, \tag{20}$$

where k is the limit on the sampled neighbor size in Eq.1.

Proof. Eq.20 can be obtained by unrolling the recursive approximation error according to Eq.18 and Eq.19. Then, the proof of Theorem 5 is completed. \Box

Implication. As have discussed in Section 3.1 and Section 4.3, since model parameters and the graph structure keep changing, intermediates computed in previous training iterations may become out-of-date and introduce huge approximation errors. Theorem 5 suggests that more stale embeddings are more likely to hurt model accuracy. The staleness problem can be severe on large dynamic graphs where the average frequency of updating each node's embedding is quite low. Fortunately, our cache replacement policy MRD (Algorithm 4) can implicitly prevent the cache from keeping extremely stale embeddings that would hurt model accuracy. Therefore, cache-limited Orca, equipped with our MRD algorithm, is supposed to achieve better predictive performance than cache-unlimited Orca on large dynamic graphs.

5.2 Convergence Guarantee

In this section, we analyze how our Orca framework affects the convergence rate of model training. T-GNNs are updated using batches of interactions. Given a new data batch B_t , the parameters of a vanilla T-GNN are updated through $W_t = W_{t-1} - \eta \nabla \mathcal{L}(W_{t-1})$, where $\mathcal{L}(\cdot)$ is a loss function, $\nabla \mathcal{L}(W_{t-1})$ is the gradient obtained on the batch B_t with parameters W_{t-1} , and η is the learning rate. Similarly, Orca updates model parameters through $W_t = W_{t-1} - \eta \nabla \tilde{\mathcal{L}}(W_{t-1})$, where $\nabla \tilde{\mathcal{L}}(W_{t-1})$ denotes the approximate gradient caused by our gradient blocking and reuse schemes. Given the above preliminaries, we work out the convergence guarantee for our Orca framework as follows.

Theorem 6. Consider a T-GNN updated by Orca through gradient descent $W_t = W_{t-1} - \eta \nabla \tilde{\mathcal{L}}(W_{t-1})$. Assume that (1) the loss function $\mathcal{L}(\cdot)$ is ρ -smooth, (2) the activation function in T-GNN is Lipschitz continuous, (3) the gradients of model parameters have bounded Frobenius norm, and (4) the distribution of nodes whose embeddings are newly computed by Orca follows the distribution of nodes whose embeddings are computed by vanilla T-GNN. Then, after training the T-GNN for T iterations, we have

$$\frac{1}{T} \sum_{t=1}^{T} \mathbb{E} \|\nabla \mathcal{L}(W_{t-1})\|_F^2 = O(1/\sqrt{T})$$
(21)

by setting the learning rate $\eta = \min(\frac{1}{\rho}, \frac{1}{\sqrt{T}})$.

The above theorem demonstrates that a T-GNN trained through our Orca framework can converge to a local optimum as the vanilla T-GNN does with convergence rate $O(1/\sqrt{T})$. Specifically, Orca introduces the approximate gradient $\nabla \tilde{\mathcal{L}}$ while vanilla T-GNN is updated via the exact gradient $\nabla \mathcal{L}$. Theorem 6 shows that a T-GNN trained with the approximate gradient $\nabla \tilde{\mathcal{L}}$ can eventually achieve a small exact gradient $\nabla \mathcal{L}$, i.e., the gradient that is obtained without reusing intermediates or gradient blocking. This theoretical finding indicates that the approximate gradient $\nabla \tilde{\mathcal{L}}$ computed

Table 2. Summary of datasets used in experiments. |V| and |E| denote the number of nodes and temporal links, respectively. d_v and d_e denote the dimensionality of node and edge features, respectively. The last column indicates whether the corresponding dataset is a bipartite graph.

Dataset	V	<i>E</i>	d_v	d_e	Bipartite
MOOC [35]	7,144	411,749	172	4	True
Wikipedia [35]	9,227	157,474	172	172	True
Reddit [35]	10,984	672,447	172	172	True
AskUbuntu [1]	159,316	964,437	172	0	False
SuperUser [2]	194,085	1,443,339	172	0	False
Wiki-Talk [4]	1,140,149	7,833,140	172	0	False

by Orca is asymptotically unbiased as the number of training iterations $T \to \infty$. Note that the assumptions in Theorem 6 are common and widely used in the convergence analysis [13, 45, 64]. We defer the proof to our technical report [3] due to the page limit.

6 EXPERIMENTS

We design experiments to answer the following questions. (1) How does our Orca perform in terms of training efficiency and model accuracy when compared to state-of-the-art T-GNNs(Section 6.2)? (2) How does our MRD algorithm perform compared to classical caching policies (Section 6.3)? (3) Can our gradient blocking strategy effectively avoid exploding gradients and thus stabilize model training (Section 6.4)? (4) How does the performance of Orca change with the model depth L and the cache size limit m (Section 6.5)? (5) What is the breakdown of reuse chances on real-world dynamic graphs (Section 6.5)?

6.1 Experimental Setup

Datasets. Table 2 summarizes the datasets used in experiments. The first three datasets are common benchmarks for evaluating T-GNNs [35, 50, 66, 70]. Specifically, the MOOC dataset consists of student-course interactions; the Wikipedia dataset records one month of edits made by users on Wikipedia pages; the Reddit dataset contains one month of posts made by users on subreddits. Besides, AskUbuntu, SuperUser, and Wiki-Talk consist of timestamped user-user interactions collected over six years from the corresponding platforms. Since these datasets mainly contain temporal interactions among nodes, we use link prediction as the downstream task. We chronologically split all the graphs in Table 2 into training (70%), validation (15%), and test (15%) sets. To test model performance on unseen nodes, we randomly sample 10% of nodes from each dataset and mask them during model training as in [70]. Since node features are absent, we generate 172-dimensional zero feature vectors for all the datasets following previous works [35, 70].

Compared Models. We apply our Orca framework to TGN [50] and compare our techniques with the state-of-the-art T-GNNs.

- **Orca-P** caches intermediate embeddings without cache limit and prunes the computation graphs via the **p**ush-and-**p**ull technique(Algorithm 1).
- **Orca-R** handles large dynamic graphs via the <u>reuse-or-recompute</u> approach (Algorithm 2) and employs the MRD policy (Algorithm 4) to generate offline cache plans.
- JODIE [35] employs double RNN models to update the representations of both nodes in each edge interaction.
- TGAT [70] encodes continuous time using random Fourier features and mimics the message passing of static GNNs.

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• TGN [50] provides a generic framework that unifies the prior works [35, 59, 70] as special cases.

- CAW [66] adopts anonymized random walks for temporal sampling and aggregates sampled results using attention modules.
- APAN [65] decouples model inference and graph computation and accelerates inference using asynchronous message propagation.

Evaluation Metrics. As for effectiveness, we compare models' average precision (AP) on the test set. More specifically, we consider two types of tasks: *transductive* learning and *inductive* learning. The transductive task examines model performance on nodes that have been observed in training. In contrast, the inductive task tests the learning ability of models on unseen nodes. As for efficiency, we evaluate the per-epoch training time (i.e., epoch time) and the total training time till convergence (i.e., convergence time) of models. Following [50, 66, 70], we say a model converges if its validation performance does not increase for five epochs.

Training Configurations. Our experimental configurations generally follow the previous works [50, 66, 70]. In experiments, we focus on link prediction tasks that forecast whether an interaction happens between two given nodes at a future timestamp. Since the original datasets only contain true observations, we randomly sample an equal number of false links. For all the datasets, we set the batch size to 200 for model training, validation, and testing. Moreover, we set the maximum number of training epochs to 50 with early stopping. Specifically, we stop model training if the validation performance does not increase for five epochs. As for hyperparameters, we tune the learning rate for each method over the space $\{1, 0.1, 0.01, 0.001\} \times 10^{-4}$. We conducted experiments on a server with 72 Intel(R) Xeon(R) 2.60 GHz CPUs, 8 GeForce RTX 2080 GPUs, and 256 GB of memory. Each experiment was repeated five times, and the average results were reported.

Implementation of Models. Orca-P and Orca-R are built on top of the generic framework [50]. Given a target node at time t, our models sample its top-k recent neighbors (Eq.1), aggregate the neighborhood information with a multi-head attention module (Eq.6), and then combine the intermediate results using a 2-layer MLP model (Eq.7). The critical difference is that Orca uses intermediate embeddings as key values in attention modules, while TGN uses state vectors. For the baselines, we adapt the officially released codes to our evaluation pipeline. To achieve a fair comparison, we set the maximum neighbor size k = 10 for APAN, TGAT, TGN, Orca-P, and Orca-R. For each target node in CAW, we sample 64 length-2 anonymized random walks as it is the default setting in the original implementation.

6.2 Key Results

Table 3. Comparison of T-GNN models on small dynamic graphs. We report the performance in transductive average precision (%), inductive average precision (%), and convergence time (s). Besides, we report the number of epochs required for models to converge in parentheses. The best and second-best results are highlighted in bold and underlined, respectively.

Model	Wikipedia			Reddit			MOOC		
	Trans	Induct	Convergence (#)	Trans	Induct	Convergence (#)	Trans	Induct	Convergence (#)
JODIE	95.16	93.13	2356.4 (18)	95.83	93.20	9243.64 (14)	83.26	81.77	5160.30 (15)
TGAT	94.26	92.88	2881.4 (29)	97.80	96.08	11933.8 (21)	70.22	70.83	7838.5 (25)
TGN	98.58	98.05	2182.7 (26)	98.66	97.55	11528.9 (26)	88.88	88.17	5086.83 (21)
APAN	96.41	96.06	1605.0 (21)	98.50	97.38	16431.8 (18)	87.02	86.74	3374.1 (14)
CAW	98.18	98.24	10175.5 (13)	98.54	97.97	75585.1 (16)	80.60	80.18	34063.9 (14)
Orca-P	98.62	98.09	148.81 (23)	98.68	97.54	597.2 (20)	89.34	89.33	303.1 (18)

Table 4. Comparison of T-GNN models on large dynamic graphs. We report the performance in transductive average precision (%), inductive average precision (%), and convergence time (s). Besides, we report the number of epochs required for models to converge in parentheses. The best and second-best results are highlighted in bold and <u>underlined</u>, respectively. The Orca-R models were implemented with a cache limit m = 1,000. "—" denotes CUDA memory error, and "~" denotes time limit exceed such that we cannot finish one epoch of model training in 12 hours.

Model	AskUbuntu			SuperUser			Wiki-Talk		
	Trans	Induct	Convergence (#)	Trans	Induct	Convergence (#)	Trans	Induct	Convergence (#)
JODIE	_	-	_	_	-	_	_	-	_
TGAT	87.57	84.21	11728.5 (21)	86.40	83.12	17129.3 (19)	91.72	85.38	189420.8 (34)
TGN	94.51	92.73	36202.6 (20)	93.18	91.76	81747.6 (24)	-	-	_
APAN	89.17	88.43	21606.1 (14)	87.07	85.50	48724.2 (19)	~	~	~
CAW	90.87	90.63	61903.7 (14)	88.92	88.32	111744.8 (15)	~	~	~
Orca-P	94.58	93.02	715.7 (18)	93.53	91.93	1309.6 (21)	95.57	90.02	8575.4 (18)
Orca-R	95.57	93.06	972.3 (21)	94.49	92.06	1173.5 (17)	95.79	90.37	9929.0 (19)

Table 3 and Table 4 compare our methods with the state-of-the-art baselines on small and large dynamic graphs, respectively. We omit the results of Orca-R in Table 3 since we can easily cache all the intermediates on small graphs. Moreover, we set the cache limit m=1,000 for Orca-R on large graphs since Orca-R can *already achieve considerable cache hits with such a small cache*. For a fair comparison, all the models except JODIE are implemented in 2 layers. JODIE cannot go deep since it does not involve recursive neighbor aggregation. In the following, we compare the methods in terms of efficiency, effectiveness, and convergence rate.

Time Efficiency of Orca. As shown in Table 3 and Table 4, our Orca-P and Orca-R are *one or two orders of magnitude faster* than the compared baselines. Notably, compared to Orca-P, the recomputation cost of Orca-R is marginal on large dynamic graphs. This observation indicates that our optimal cache replacement algorithm MRD (Algorithm 4) can achieve high cache hit ratios even under a small cache limit m=1,000. In principle, Orca-P and Orca-R are much faster because they avoid *temporal explosion* and *neighbor explosion* by sharing *newly computed* embeddings across multiple timestamps and by reusing *previously cached* historical embeddings, respectively. Although JODIE is computationally simple, it partitions the dynamic graph into very small batches to avoid overlapping between interactions. Small batches make GPU resources underutilization, thus slowing down model training. TGAT and TGN follow the generic framework described in Section 2.3. They cannot resolve the temporal and neighbor explosion problems such that the time complexity still increases linearly with the number of unique timestamps and grows exponentially with the number of layers. Besides, CAW and APAN are sensitive to the amount of available CPU resources and RAM. Table 4 shows that they cannot even finish one epoch of training in 12 hours on the Wiki-Talk dataset.

Memory Efficiency of Orca. Orca-P and Orca-R can also reduce memory consumption during model training. Although these two algorithms cache intermediates in GPU memory, our reuse schemes *save not only the computational cost but also the memory required to finish the corresponding computations.* In contrast, the memory cost of the baselines still scales with the number of nodes in the computation graph. As shown by the "—" marks in Table 4, JODIE and TGN cannot scale to large dynamic graphs due to CUDA memory error. Our Orca-R can scale to large graphs without incurring memory error by setting a practical cache limit.

Convergence Rate of Orca. Table 3 and Table 4 show the number of epochs required for different models to converge. Compared to the baselines that do not reuse intermediates, Orca-P and

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Orca-R demonstrate better generalization performance while having no noticeable impact on the convergence rate. This finding is consistent with our theoretical analysis (Theorem 6) on the convergence guarantee of the Orca framework. Although our gradient blocking and reuse schemes cause approximated gradients, Theorem 6 shows that the gradient is asymptotically unbiased. Therefore, Orca-P and Orca-R can converge to a local optimum as vanilla T-GNNs do.

Effectiveness of Orca. Table 3 compares models' average precision (AP) in both transductive and inductive settings. Orca-P matches or outperforms all the compared baselines in the transductive setting. CAW is specifically designed for *inductive* representation learning on dynamic graphs. Although it achieves better inductive AP than Orca-P on Wikipedia and Reddit datasets, our Orca-P is more efficient with only 0.15%-0.43% accuracy loss. Table 4 shows model performance on large dynamic graphs. Surprisingly, our Orca-P and Orca-R outperform all the baselines, which do not reuse intermediates, in both transductive and inductive settings. This finding suggests that although our reuse schemes introduce approximation errors, slight noises could overcome the overfitting problem and thus improve models' generalization ability [31, 45]. Furthermore, Orca-R achieves higher AP than Orca-P in all the cases. The reason is that Orca-R is equipped with our MRD algorithm. MRD can *avoid keeping and reusing incredibly stale embeddings* that would introduce significant approximation errors, thus improving model precision.

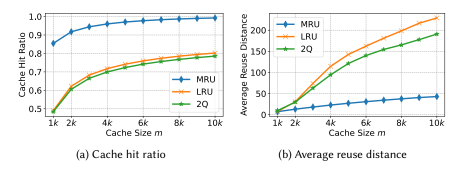


Fig. 5. Comparison of caching policies on AskUbuntu.

6.3 Comparison of Cache Replacement Policies

Cache Hit Ratio and Reuse distance. Figure 5 compares our MRD algorithm with the classical cache replacement policies 2Q [32] and LRU [15]. Since MRD is optimal in maximizing the number of cache hits, it achieves higher cache hit ratios. In contrast, LRU and 2Q tend to cache all the newly computed embeddings even if they may not be reused in the near future. On the other hand, MRD leads to significantly smaller reuse distance with the increase of cache size. Here, the reuse distance of a cached embedding is defined as the number of passed training iterations from its being cached to its being reused. Hence, smaller reuse distance indicates less staleness of the cached embedding. Training Efficiency. Table 5 further demonstrates the overhead of different caching policies and how they affect the model training time till convergence. Since MRD achieves higher cache hit ratios than LRU and 2Q, training Orca-R models with MRD is generally faster. Besides, MRD only takes 4.06 seconds to make cache plans on AskUbuntu with a cache limit of 5,000. The preparation overhead is negligible compared to the total training cost of 823.8 seconds. Therefore, our MRD policy itself is efficient and accelerates Orca-R by reducing the amount of recomputation.

Model Effectiveness. Table 5 also shows that MRD can lead to better model effectiveness than LRU and 2Q on AskUbuntu. The same conclusion can be drawn from other datasets. This is consistent

Table 5. Training Orca-R with various caching policies on AskUbuntu when the cache limit is 5k. We report the preparation overhead (s) of caching policies, convergence time (s) of models, transductive AP (%), and inductive AP (%).

Algorithm	Preparation	Convergence	Trans AP	Induct AP
LRU	1.35	988.8	95.32	92.97 92.94
2Q	1.90	1032.6	95.25	, 2, , ,
MRD	4.06	823.8	95.73	93.06

with our observation in Figure 5b, which demonstrates that MRD results in significantly smaller average reuse distance. Therefore, training Orca-R model with MRD can achieve better predictive performance by filtering out those stale embeddings that could introduce large errors.

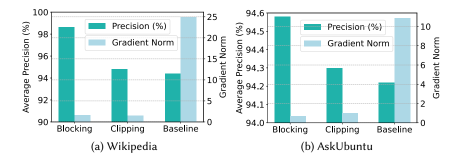


Fig. 6. Training Orca-P with different gradient update strategies on Wikipedia and AskUbuntu.

6.4 Comparison of Gradient Update Strategies

Model Effectiveness. In Section 3.2, we have discussed that simply reusing intermediates could cause the exploding gradient issue. Figure 6 further investigates the impact of various gradient update strategies on the Frobenius norm of gradients and the performance of Orca-P. Note that the simple baseline, which does not attempt to reduce gradient values, indeed suffers from exploding gradients and thus cannot achieve satisfactory model precision. On the other hand, the gradient clipping method [75], a common practice for avoiding gradient explosion, can project gradient values into a normal range. However, our gradient blocking approach is more effective than gradient clipping as the latter only results in a marginal gain in model precision. Simply clipping gradient values cannot prevent gradients from accumulation. Therefore, gradients from multiple timestamps would still aggregate and cause unstable model updates.

Impact of Neighbor Distribution. We observe that the gradient explosion issue on Wikipedia is more severe than that on AskUbuntu. As shown in Figure 6, the gradient norm of the simple baseline on Wikipedia and AskUbuntu is 24.9 and 10.8, respectively. The phenomenon is caused by the difference in these two data distributions. Specifically, in-batch neighbors account for 77.9% of the total 1-hop neighbors on Wikipedia, while this value is 61.2% on AskUbuntu. As a result, the newly computed embeddings are aggressively shared across more timestamps in a data batch when training Orca-P on Wikipedia than those on AskUbuntu. By the chain rule of differentiation, gradients from multiple timestamps would aggregate, resulting in gradient explosion. The above

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observation again validates our analysis in Section 3.2 that simply reusing intermediates can cause exploding gradients.

6.5 Other Ablation Studies

Impact of cache size m. Figure 7a demonstrates the cache hit ratios achieved by MRD on four dynamic graphs. Note that MRD can lead to 100% cache hits on Wikipedia with a tiny cache size m = 1.5k. For the other three large dynamic graphs, MRD achieves a cache hit ratio of about 90% with a cache size m = 2k. Particularly, the training set of Wiki-Talk involves 624,057 different nodes. The cache size m = 2k only accounts for 0.32% of the unique training nodes. This is because dynamic graphs usually follow the *power-law distribution*, i.e., a few nodes appear in most temporal interactions. Therefore, we can achieve a high cache hit ratio by caching intermediates of some frequently occurring nodes.

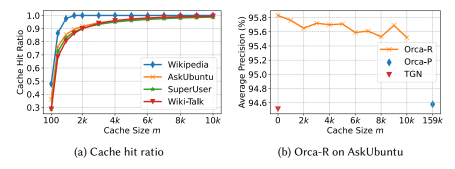


Fig. 7. (a) Cache hit ratios achieved by MRD varying cache size m. (b) Average precision of Orca-R varying cache size m. Since Orca-R can share newly computed embeddings across timestamps without caching, m = 0 indicates that only intermediates of in-batch neighbors are reused.

Figure 7b shows the average precision of Orca-R on AskUbuntu. As the cache size m increases, the performance of Orca-R slightly decreases but is consistently better than Orca-P and TGN. The performance drop is in line with our expectation because a larger cache maintains more stale embeddings and increases the reuse distance, thus introducing larger approximation errors. Note that Orca-R with m=10k achieves almost 100% cache hits but still outperforms Orca-P, which caches all the historical embeddings, by a large margin. This finding suggests that the performance gap between Orca-P and Orca-R is caused by a few incredibly unreliable historical intermediates. Moreover, both Orca-P and Orca-R beat the baseline TGN. As discussed in Section 6.2, this is because our reuse scheme can act as *regularization* techniques [22, 31, 39, 71] and therefore improve models' generalization performance by overcoming the *overfitting* issue [19, 45].

Impact of model depth *L*. Figure 8a compares the epoch time of our Orca-P and Orca-R with baselines on the Wikipedia dataset. Specifically, the epoch time of Orca-P remains stable as we increase the number of layers *L*. In contrast, the cost of Orca-R increases faster as the number of recomputed representations grows exponentially with the number depth *L*. Although Orca-R incurs extra recomputation cost, it is still significantly faster than the compared baselines. Moreover, with the increase of model depth *L*, TGAT, TGN, and CAW fail due to CUDA memory error. In contrast, our Orca-P and Orca-R are memory efficient and scalable. In general, the *efficiency advantage of our methods over baselines increases with the model depth L*.

Reuse chance breakdown. Figure 8b shows the total number of 1-hop in-batch neighbors (e.g., v_2 in Figure 3) and 1-hop out-of-batch neighbors (e.g., v_4 in Figure 3) during model training. All the six

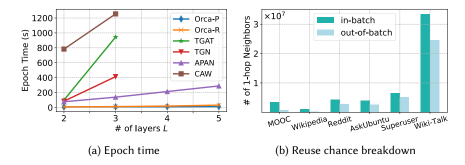


Fig. 8. (a) Epoch time of models varying the number of layers on Wikipedia. Orca-R is implemented with per-layer cache limit m=500. Some results of baselines are absent due to CUDA memory error. (b) The total number of 1-hop in-batch and out-of-batch neighbors during model training.

dynamic graphs exhibit a high degree of *temporal locality*, i.e., the number of in-batch neighbors exceeds that of out-of-batch neighbors. This observation also indicates that the dynamic graphs follow the *power-law distribution*, where some common neighbor nodes frequently appear in different data batches. Orca prunes the computation graph by reusing the newly computed embeddings of in-batch neighbors and the cached intermediates of out-of-batch neighbors. According to the reuse scheme, Figure 8b suggests that *in-batch neighbors contribute more reuse chances* to Orca.

7 RELATED WORK

Representation Learning on Dynamic Graphs. Prior works [23, 24, 40, 46, 53, 55, 72] mainly focus on *discrete-time dynamic graphs* (DTDGs) that are represented by graph snapshots taken at different time intervals. The random walk-based methods [40, 72] incrementally sample time-dependent walks and minimize the pairwise distance of learned representations of adjacent vertices in each walk. Another line of works [46, 53, 55] encodes the structural information of each graph snapshot using a GNN model and then learns the temporal dynamics of consecutive snapshots using a sequence model such as RNN. However, all the DTDG-based solutions need to aggregate graph snapshots with a predefined time granularity. Thus, they are not suitable for many real-world applications where interactions can happen at various time granularities.

T-GNNs [35, 50, 59, 65, 66, 70] have been developed to learn representations on *continuous-time dynamic graphs* (CTDGs) that are represented by sequences of timestamped edges. Given a new edge, the sequence-based approach JODIE [35] updates representations of the two nodes involved in the edge using RNNs. DyRep [59] further considers 2-hop neighborhood information for updating node representations. TGAT [70] encodes continuous timestamps using random Fourier features and mimics the message passing of static GNNs. TGN [50] is a generic T-GNN framework that includes [35, 59, 70] as special cases. To improve the inductive performance on unseen nodes, CAW [66] adopts an anonymization strategy that replace node identities with the number of node occurrences based on a set of sampled walks. In addition, APAN [65] decouples model inference and graph computation and accelerates model inference through asynchronous message propagation. **Reuse in Static GNNs.** A few recent works have considered using historical embeddings to minimize sampling variance [13, 16] or accelerate computation [19, 37, 45, 58] for *static* GNNs. To avoid neighbor explosion, VR-GCN [13] reuses cached intermediate embeddings in forward propagation and develops a control variate based neighbor sampling algorithm. MVS-GNN [16] further improves the sampling efficiency and reduces the variance of mini-batch construction. On

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the other hand, GNNAutoScale [19] prunes the computation graph by reusing historical embeddings of 1-hop out-of-batch neighbors. IGLU [45] adopts a more aggressive caching strategy where the intermediate results are lazily updated at regular intervals. Dorylus [58] reduces the communication cost in distributed GPU servers by caching the embeddings of boundary nodes. CacheGNN [37] accelerates the inference of static GNNs over graph snapshots. In contrast to the above works, we improve the training efficiency of T-GNNs under practical cache limits.

Reuse in ML Systems. Materialization and reuse in ML systems are reminiscent of the classical view materialization problems [14, 30, 41] in database systems. Such techniques have been adopted in a wide range of ML lifecycle management systems such as data versioning [10, 11], model management [42, 61], feature selection [44, 56, 57, 74], model training [43, 64], model diagnosis [60], and model serving [17, 33, 36]. Specifically, HELIX [68, 69] and Collaborative Optimizer [18] optimize the execution cost across ML pipelines by caching and reusing intermediates. LIMA [48] further provides more fine-grained lineage tracing and reuse inside ML systems. Note that the above works cannot be applied to T-GNNs because they do not support approximate reuse. Moreover, HET [43] reduces the communication overhead of training large embedding models by caching hot embeddings on local machines with consistency guarantees. In contrast to prior works, Orca captures the approximate reuse chances for T-GNN training with theoretical guarantees.

8 CONCLUSION

In this paper, we present Orca, a scalable caching-based framework that significantly reduces the computational cost of T-GNNs on dynamic graphs. To scale Orca to large dynamic graphs without compromising model accuracy, we propose an optimal cache replacement algorithm under practical cache limits. Moreover, we develop theoretical analyses of the approximation errors introduced by our reuse mechanisms and offer rigorous convergence guarantees. Extensive experiments have validated that Orca can be two orders of magnitude faster than the state-of-the-art baselines while achieving higher precision on various dynamic graphs.

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A PROOF OF THEOREM 6

We finish the proof of Theorem 6 in four steps. (1) We first bound the change of model parameters during one training epoch. (2) We next show that the staleness of cached embeddings and the approximation error of newly computed embeddings can be bounded if the change of model parameters can be bounded. (3) Then, we relate the bias in approximated gradients to the staleness of reused embeddings. (4) Finally, we establish the convergence guarantee using properties like smoothness and Lipschitz continuous.

A.1 Preliminaries

Generic T-GNN Architecture. The analysis of gradient update and convergence rate is always model-dependent. Without loss of generality, we consider vanilla T-GNNs in the most generic form as follows:

$$Z^{l} = P^{l}H^{l-1}W^{l}, \quad H^{l} = \sigma(Z^{l}), \quad l = 1, \dots, L,$$
 (22)

where H^l is the embeddings at the l-layer, W^l is the learnable weight matrix used to compute the l-th layer embeddings, P^l is a weighted adjacency matrix for feature aggregation, and $\sigma(\cdot)$ is an activation function. For ease of simplicity, we omit edge features and time encodings in Eq.22. However, as we will show later, our theoretical analysis can be easily extended to incorporate edge and time features. Similarly, we can express Orca-P and Orca-R models in the following generic form:

$$\tilde{Z}^{l} = (\tilde{P}^{l}\tilde{H}^{l-1} + \bar{P}^{l}\tilde{H}^{l-1})W^{l}, \quad \tilde{H}^{l} = \sigma(\tilde{Z}^{l}), \quad l = 1, \dots, L,$$
 (23)

where \tilde{H}^l and \bar{H}^l denote the newly computed embeddings and reused embeddings, respectively. Besides, \tilde{P} and \bar{P} denote matrices used to aggregate \tilde{H}^{l-1} and \bar{H}^{l-1} features, respectively.

It is worth noting that the exact embeddings H^l computed by vanilla T-GNN and the approximated embeddings \tilde{H}^l computed by Orca can have different shapes. This is because Orca prunes the computation graph by calculating the embeddings for fewer nodes. Let row(H) denotes the number

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of rows in the matrix H. Then, $\operatorname{row}(H^l) = \operatorname{row}(\tilde{H}^l)$ if l = L. Otherwise, $\operatorname{row}(H^l) \ge \operatorname{row}(\tilde{H}^l)$. It is generally difficult to compare the gradients of embedding matrices in different shapes. For ease of analysis, we rewrite Orca as follows:

$$\tilde{Z}^{l} = (\tilde{P}^{l}(\tilde{H}^{l-1} - \bar{H}^{l-1}) + P\bar{H}^{l-1})W^{l}, \quad \tilde{H}^{l} = \sigma(\tilde{Z}^{l}), \tag{24}$$

where $\operatorname{row}(\tilde{H}^{l-1}) = \operatorname{row}(H^{l-1}) = \operatorname{row}(H^{l-1})$, and \tilde{P}^l is an adjacency matrix used to aggregate the features of newly computed embeddings \tilde{H}^{l-1} . Specifically, in Eq.24, we conceptually augment the computation graph of Orca so that Orca computes embeddings for the same set of nodes as vanilla T-GNNs at each layer l ($l=1,\ldots,L$). More importantly, we can remove the impact of extra conceptually computed embeddings by setting the corresponding terms in the adjacency matrix \tilde{P}^l to zero. As a result, Eq.23 and Eq.24 are mathematically equivalent and produce the same temporal embeddings.

Updating T-GNN via Gradient Descent. T-GNNs are updated using batches of interactions. Consider a batch of target nodes B_{τ} , where $(v_i, t) \in B_{\tau}$ denotes node v_i at timestamp t. For each $(v_i, t) \in B_{\tau}$, an L-layer vanilla T-GNN computes its temporal embedding $h_i^L(t)$ and then gets the loss value $f(h_i^L(t), y_i(t))$, where $y_i(t)$ is the true label of node v_i at t, and $f(\cdot, \cdot)$ is a loss function. Finally, parameters of the vanilla T-GNN are updated through

$$W_{\tau} = W_{\tau-1} - \eta \nabla \mathcal{L}(W_{\tau-1}), \tag{25}$$

where $\nabla \mathcal{L}(W_{\tau-1}) = \frac{1}{|B_{\tau}|} \sum_{(v_i,t) \in B_{\tau}} \nabla f(h_i^L(t), y_i(t))$ is the gradient obtained on the data batch B_{τ} with parameters $W_{\tau-1}, W_{\tau-1} = [W_{\tau-1}^1, \dots, W_{\tau-1}^L]$ is the model parameters obtained after updating T-GNN on the batch $B_{\tau-1}$, and η is the learning rate.

Similarly, for each target node $(v_i,t) \in B_\tau$, an L-layer Orca-P or Orca-R model computes its temporal embedding $\tilde{h}_i^L(t)$ and then gets the loss value $f(\tilde{h}_j^i(t),y_i(t))$. Finally, parameters of our Orca model are updated through

$$W_{\tau} = W_{\tau-1} - \eta \nabla \tilde{\mathcal{L}}(W_{\tau-1}), \tag{26}$$

where $\nabla \tilde{\mathcal{L}}(W_{\tau-1}) = \frac{1}{|B_{\tau}|} \sum_{(v_i,t) \in B_{\tau}} \nabla f(\tilde{h}_i^L(t), y_i(t))$ is the approximated gradient obtained on the batch B_{τ} with parameters $W_{\tau-1}$.

Backpropagation Rules. Let $f_i(t) = f(h_i^L(t), y_i(t))$ and $\tilde{f}_i(t) = f(\tilde{h}_i^L(t), y_i(t))$. Then, for vanilla T-GNNs in the form of Eq.22, we have the following backpropagation rules:

$$\nabla_{H^{l-1}} f_i(t) = (P^l)^T \nabla_{Z^l} f_i(t) (W^l)^T$$

$$\nabla_{W^l} f_i(t) = (P^l H^{l-1})^T \nabla_{Z^l} f_i(t)$$

$$\nabla_{Z^l} f_i(t) = \sigma'(Z^l) \circ \nabla_{H^l} f_i(t).$$
(27)

Moreover, for Orca-P and Orca-R in the form of Eq.24, we have the following backpropagation rules:

$$\nabla_{\tilde{H}^{l-1}}\tilde{f}_{i}(t) = (\tilde{P}^{l})^{T}\nabla_{\tilde{Z}^{l}}\tilde{f}_{i}(t)(W^{l})^{T}$$

$$\nabla_{W^{l}}\tilde{f}_{i}(t) = (\tilde{P}^{l}(\tilde{H}^{l-1} - \tilde{H}^{l-1}) + P^{l}\tilde{H}^{l-1})^{T}\nabla_{\tilde{Z}^{l}}\tilde{f}_{i}(t)$$

$$\nabla_{\tilde{Z}^{l}}\tilde{f}_{i}(t) = \sigma'(\tilde{Z}^{l}) \circ \nabla_{\tilde{H}^{l}}\tilde{f}_{i}(t).$$
(28)

Note that we do not calculate partial dervatives with repect to the cached embeddings \bar{H}^l (l = 1, ..., L-1) in the above rules. This is because Orca adopts a gradient blocking strategy so that gradients will not propagate through the cached and reused embeddings.

A.2 Bounding the Parameter Change

We bound the change of model parameters in one training epoch. Specifically, the parameters of T-GNN are updated in severe training epochs, where each epoch consists of a sequence of training batches. Let the gradient of model parameters be bounded, i.e., $\|\nabla \tilde{\mathcal{L}}(W_{\tau})\|_F \leq G$ and $\|\nabla \mathcal{L}(W_{\tau})\|_F \leq G$. Besides, let n be the number of bathes in an epoch. Then, for any two weight matries W_{τ} and $W_{\tau'}$ ($\tau < \tau'$) in an epoch, we can bound the change of model parameters by

$$||W_{\tau} - W_{\tau'}||_{F} \leq \sum_{z=\tau}^{\tau'-1} ||W_{z} - W_{z+1}||_{F} = \sum_{z=\tau}^{\tau'-1} ||\eta \nabla \tilde{\mathcal{L}}(W_{z})||_{F}$$

$$\leq (\tau' - \tau)\eta G \leq nG\eta,$$
(29)

where η is the learning rate.

A.3 Bounding the Staleness

According to the lemma 1 in [13], the staleness of cached embeddings can be bounded by a constant multiplied by the change of parameters. Moreover, we have already shown in Eq.29 that the parameter change $||W_i - W_j||_F \le nG\eta$. Therefore, for l = 1, ..., L, there exists a constant k_s such that:

$$\begin{aligned} \|h_i^l(t) - \bar{h}_i^l\|_2 &\leq k_s \eta, \quad \|h_i^l(t) - \tilde{h}_i^l(t)\|_2 \leq k_s \eta \\ \|z_i^l(t) - \bar{z}_i^l\|_2 &\leq k_s \eta, \quad \|z_i^l(t) - \tilde{z}_i^l(t)\|_2 \leq k_s \eta, \end{aligned}$$
(30)

where $h_i^l(t)$ is the exact embedding, $\tilde{h}_i^l(t)$ is the approximated embedding computed by Orca, \bar{h}_i^l is the cached embedding, and $z_i^l(t), \bar{z}_i^l(t), \bar{z}_i^l$ denote the corresponding features before activation. In the following analysis, we denote

$$\epsilon = k_{\rm s} \eta \tag{31}$$

as the upper bound on the staleness and approximation error for notation convention.

A.4 Bounding the Gradient Bias

Bounded Frobenius Norm. Assume that the weight matrices, adjacency matrices, embedding matrices, and gradient matrices have bounded Frobenius norm, i.e., there exists a constant *S* so that

$$||P^{l}||_{F} \leq S, \quad ||W^{l}||_{F} \leq S, \quad ||\tilde{H}^{l}||_{F} \leq S, \quad ||\sigma'(\tilde{Z}^{l})||_{F} \leq S,$$

$$||\nabla_{H^{l}} f_{i}(t)||_{F} \leq S, \quad ||\nabla_{Z^{l}} f_{i}(t)||_{F} \leq S, \quad ||\nabla_{\tilde{Z}^{l}} \tilde{f}_{i}(t)||_{F} \leq S.$$
(32)

Note that bounded norm is a common assumption and widely used in the convergence analysis [13, 45, 64].

Continuous and Smoothness Properties. Let the loss function $f(h^L,y)$ be λ_{loss} -Lipschitz continuous and ρ_{loss} -smooth with respect to the temporal embedding h^L , i.e., $|f(h^L,y) - f(\tilde{h}^L,y)| \le \|h^L - \tilde{h}^L\|_2$ and $\|\nabla_{h^L} f(h^L,y) - \nabla_{\tilde{h}^L} f(\tilde{h}^L,y)\|_2 \le \rho_{loss} \|h^L - \tilde{h}^L\|_2$. Moreover, assume that the activation function $\sigma(\cdot)$ is ρ_{act} -smooth, i.e., $\|\sigma'(Z) - \sigma(\tilde{Z})\|_F \le \rho_{act} \|Z - \tilde{Z}\|_F$.

Bounding $\mathbb{E}\|\nabla_{Z^L}f_i(t) - \nabla_{\tilde{Z}^L}\tilde{f}_i(t)\|_F$. We prove by induction that there exists a constant k_Z^l (l = 1, ..., L) such that

$$\mathbb{E}\|\nabla_{Z^l}f_i(t) - \nabla_{\tilde{Z}^l}\tilde{f}_i(t)\|_F \le k_Z^l\epsilon. \tag{33}$$

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We start from the case l = L. Since the loss function $f(h^L, y)$ is ρ_{loss} -smooth, then

$$\|\nabla_{H^{L}}f_{i}(t) - \nabla_{\tilde{H}^{L}}\tilde{f}_{i}(t)\|_{F}$$

$$= \|\nabla_{h_{i}^{L}(t)}f(h_{i}^{L}(t), y_{i}(t)) - \nabla_{\tilde{h}_{i}^{L}(t)}f(\tilde{h}_{i}^{L}(t), y_{i}(t))\|_{2}$$

$$\leq \rho_{loss}\|h_{i}^{L}(t) - \tilde{h}_{i}^{L}(t)\|_{2} \leq \rho_{loss}\epsilon.$$
(34)

Besides, by Eq.30 and Eq.31,

$$||Z^{L} - \tilde{Z}^{L}||_{F} = ||z_{i}^{L}(t) - \tilde{z}_{i}^{L}(t)||_{2} \le \epsilon.$$
(35)

Then, we have

$$\|\nabla_{Z^{L}}f_{i}(t) - \nabla_{\tilde{Z}^{L}}\tilde{f}_{i}(t)\|_{F}$$

$$= \|\sigma'(Z^{L}) \circ \nabla_{H^{L}}f_{i}(t) - \sigma'(\tilde{Z}^{L}) \circ \nabla_{\tilde{H}^{L}}\tilde{f}_{i}(t)\|_{F}$$

$$\leq \|\sigma'(Z^{L}) \circ \nabla_{H^{L}}f_{i}(t) - \sigma'(\tilde{Z}^{L}) \circ \nabla_{H^{L}}f_{i}(t)\|_{F}$$

$$+ \|\sigma'(\tilde{Z}^{L}) \circ \nabla_{H^{L}}f_{i}(t) - \sigma'(\tilde{Z}^{L}) \circ \nabla_{\tilde{H}^{L}}\tilde{f}_{i}(t)\|_{F}$$

$$\leq \|\sigma'(Z^{L}) - \sigma'(\tilde{Z}^{L})\|_{F} \cdot \|\nabla_{H^{L}}f_{i}(t)\|_{F}$$

$$+ \|\sigma'(\tilde{Z}^{L})\|_{F} \cdot \|\nabla_{H^{L}}f_{i}(t) - \nabla_{\tilde{H}^{L}}\tilde{f}_{i}(t)\|_{F}$$

$$\leq S\rho_{act}\epsilon + S\rho_{loss}\epsilon$$

$$\leq S(\rho_{act} + \rho_{loss})\epsilon = k_{Z}^{L}\epsilon.$$
(36)

Hence, Eq.33 holds for l=L, where $k_Z^L=S(\rho_{act}+\rho_{loss})$. If the statement holds for l+1, we get

$$\|\nabla_{Z^{l}}f_{i}(t) - \nabla_{\tilde{Z}^{l}}\tilde{f}_{i}(t)\|_{F}$$

$$\leq \|\sigma'(Z^{l}) \circ (P^{l+1})^{T}\nabla_{Z^{l+1}}f_{i}(t)(W^{l+1})^{T}$$

$$-\sigma'(\tilde{Z}^{l}) \circ (\tilde{P}^{l+1})^{T}\nabla_{\tilde{Z}^{l+1}}\tilde{f}_{i}(t)(W^{l+1})^{T}\|_{F}$$

$$\leq \|\sigma'(Z^{l}) \circ (P^{l+1})^{T}\nabla_{Z^{l+1}}f_{i}(t)$$

$$-\sigma'(\tilde{Z}^{l}) \circ (\tilde{P}^{l+1})^{T}\nabla_{\tilde{Z}^{l+1}}\tilde{f}_{i}(t)\|_{F} \cdot \|(W^{l+1})^{T}\|_{F}$$

$$\leq S(\|(\sigma'(Z^{l}) - \sigma'(\tilde{Z}^{l})) \circ (P^{l+1})^{T}\nabla_{Z^{l+1}}f_{i}(t)\|_{F}$$

$$+ \|\sigma'(\tilde{Z}^{l}) \circ (P^{l+1})^{T}(\nabla_{Z^{l+1}}f_{i}(t) - \nabla_{\tilde{Z}^{l+1}}\tilde{f}_{i}(t))\|_{F}$$

$$+ \|\sigma'(\tilde{Z}^{l}) \circ (P^{l+1} - \tilde{P}^{l+1})^{T}\nabla_{\tilde{Z}^{l+1}}\tilde{f}_{i}(t)\|_{F}).$$
(37)

Note that the randomness in Eq.37 comes from the adjacency matrix \tilde{P}^{l+1} . It captures which embeddings are newly computed. Assume that the distribution of nodes whose embeddings are newly computed by Orca follows the distribution of nodes whose embeddings are computed by the vanilla T-GNN, i.e., $\mathbb{E}[\tilde{P}^{l+1}] = P^{l+1}$. Then,

$$\mathbb{E}\|\nabla_{Z^{l}}f_{i}(t) - \nabla_{\tilde{Z}^{l}}\tilde{f}_{i}(t)\|_{F}$$

$$\leq S \cdot \mathbb{E}\|\sigma'(Z^{l}) - \sigma'(\tilde{Z}^{l})\|_{F} \cdot \|(P^{l+1})^{T}\|_{F} \cdot \|\nabla_{Z^{l+1}}f_{i}(t)\|_{F}$$

$$+ \|\sigma'(\tilde{Z}^{l})\|_{F} \cdot \|(P^{l+1})^{T}\|_{F} \cdot \|\nabla_{Z^{l+1}}f_{i}(t) - \nabla_{\tilde{Z}^{l+1}}\tilde{f}_{i}(t)\|_{F}$$

$$+ \|\sigma'(\tilde{Z}^{l})\|_{F} \cdot \|(P^{l+1} - \tilde{P}^{l+1})^{T}\|_{F} \cdot \|\nabla_{\tilde{Z}^{l+1}}\tilde{f}_{i}(t)\|_{F}]$$

$$\leq S(\rho_{act}\epsilon\sqrt{\text{row}(Z^{l})} \cdot S^{2} + S^{2} \cdot k_{Z}^{l+1}\epsilon + 0)$$

$$= S^{3}(\rho_{act}\sqrt{\text{row}(Z^{l})} + k_{Z}^{l+1})\epsilon = k_{Z}^{l}\epsilon,$$
(38)

where $k_Z^l = S^3(\rho_{act}\sqrt{\text{row}(Z^l)} + k_Z^{l+1})$. Eq.33 holds by induction.

Bounding $\mathbb{E}\|\nabla_{W^l}f_i(t) - \nabla_{W^l}\tilde{f}_i(t)\|_F$. We next bound the bias in partial dervatives with respect to weight matrices W^l :

$$\mathbb{E}\|\nabla_{W^{l}}f_{i}(t) - \nabla_{W^{l}}\tilde{f}_{i}(t)\|_{F}$$

$$=\mathbb{E}\|(P^{l}H^{l-1})^{T}\nabla_{Z^{l}}f_{i}(t) - (\tilde{P}^{l}(\tilde{H}^{l-1} - \tilde{H}^{l-1}) + P^{l}\tilde{H}^{l-1})^{T}\nabla_{\tilde{Z}^{l}}\tilde{f}_{i}(t)\|_{F}$$

$$\leq \mathbb{E}\|((P^{l}H^{l-1})^{T} - (\tilde{P}^{l}(\tilde{H}^{l-1} - \tilde{H}^{l-1}) + P^{l}\tilde{H}^{l-1})^{T}\nabla_{Z^{l}}f_{i}(t)\|_{F}$$

$$+ \mathbb{E}\|(\tilde{P}^{l}(\tilde{H}^{l-1} - \tilde{H}^{l-1}) + P^{l}\tilde{H}^{l-1})^{T}(\nabla_{Z^{l}}f_{i}(t) - \nabla_{\tilde{Z}^{l}}\tilde{f}_{i}(t))\|_{F}$$

$$\leq \mathbb{E}\|(P^{l}H^{l-1})^{T} - (\tilde{P}^{l}(\tilde{H}^{l-1} - \tilde{H}^{l-1}) + P^{l}\tilde{H}^{l-1})^{T}\|_{F} \cdot S$$

$$+ \mathbb{E}\|(\tilde{P}^{l}(\tilde{H}^{l-1} - \tilde{H}^{l-1}) + P^{l}\tilde{H}^{l-1})^{T}\|_{F} \cdot k_{Z}^{l}\epsilon$$

$$\leq S\sqrt{\text{row}(H^{l-1})} \cdot \epsilon \cdot S + S^{2} \cdot k_{Z}^{l}\epsilon$$

$$\leq S^{2}(\sqrt{\text{row}(H^{l-1})} + k_{Z}^{l})\epsilon = k_{W}^{l}\epsilon,$$
(39)

where $k_W^l = S^2(\sqrt{\text{row}(H^{l-1})} + k_Z^l)$.

Bounding $\mathbb{E}\|\nabla \mathcal{L}(W) - \nabla \tilde{\mathcal{L}}(W)\|_F$. Finally, we bound the difference between the approximated gradient obtained by Orca and the exact gradient obtained by the vanilla T-GNN:

$$\mathbb{E}\|\nabla \mathcal{L}(W) - \nabla \tilde{\mathcal{L}}(W)\|_{F}$$

$$\leq \frac{1}{|B|} \sum_{l=1}^{L} \mathbb{E}\|\nabla_{W^{l}} f_{i}(t) - \nabla_{W^{l}} \tilde{f}_{i}(t)\|_{F}$$

$$\leq \frac{1}{|B|} \sum_{l=1}^{L} k_{W}^{l} \epsilon = k_{W} \epsilon,$$

$$(40)$$

where $k_W = \frac{1}{|B|} \sum_{l=1}^{L} k_W^l$, and the second inequality follows Eq.39.

A.5 Convergence Guarantee

We next establish the convergence guarantees for Orca using properties like smoothness and Lipschitz continuous. Assume that the loss function $\mathcal{L}(W)$ is ρ -smooth, i.e.,

$$|\mathcal{L}(W_2) - \mathcal{L}(W_1) - \langle \nabla \mathcal{L}(W_1), W_2 - W_1 \rangle| \le \frac{\rho}{2} ||W_2 - W_1||_F^2.$$
(41)

Let $\delta_i = \nabla \tilde{\mathcal{L}}(W_i) - \nabla \mathcal{L}(W_i)$. By the smoothness property, we have

$$\mathcal{L}(W_{i+1}) \leq \mathcal{L}(W_i) + \langle \nabla \mathcal{L}(W_i), W_{i+1} - W_i \rangle + \frac{\rho}{2} \eta^2 \|W_{i+1} - W_i\|_F^2$$

$$= \mathcal{L}(W_i) - \eta \langle \nabla \mathcal{L}(W_i), \nabla \tilde{\mathcal{L}}(W_i) \rangle + \frac{\rho}{2} \eta^2 \|\nabla \tilde{\mathcal{L}}(W_i)\|_F^2$$

$$= \mathcal{L}(W_i) - \eta \langle \nabla \mathcal{L}(W_i), \delta_i + \nabla \mathcal{L}(W_i) \rangle + \frac{\rho}{2} \eta^2 \|\delta_i + \nabla \mathcal{L}(W_i)\|_F^2$$

$$= \mathcal{L}(W_i) - (\eta - \rho \eta^2) \langle \nabla \mathcal{L}(W_i), \delta_i \rangle$$

$$- (\eta - \frac{\rho \eta^2}{2}) \|\nabla \mathcal{L}(W_i)\|_F^2 + \frac{\rho}{2} \eta^2 \|\delta_i\|_F^2.$$

$$(42)$$

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Let the Frobenius norm of gradients be bounded by a constant, i.e., $\|\nabla \mathcal{L}(W_i)\|_F \leq G$ and $\|\nabla \tilde{\mathcal{L}}(W_i)\|_F \leq G$. By Eq.31 and Eq.40,

$$\mathbb{E}\langle \nabla \mathcal{L}(W_i), \delta_i \rangle \le \|\nabla \mathcal{L}(W_i)\|_F \cdot \mathbb{E}\|\delta_i\|_F \le Gk_W k_s \eta = k_1 \eta, \tag{43}$$

$$\mathbb{E}\|\delta_i\|_F^2 \le \|\nabla \tilde{\mathcal{L}}(W)\|_F^2 + \nabla \mathcal{L}(W)\|_F^2 \le 2G^2 = k_2,\tag{44}$$

where $k_1 = Gk_Wk_s$. Plug Eq.43 and Eq.44 into Eq.42, and sum up the loss values, then

$$(\eta - \frac{\rho \eta^{2}}{2}) \sum_{i=1}^{T} \mathbb{E} \|\nabla \mathcal{L}(W_{i-1})\|_{F}^{2}$$

$$\leq \mathcal{L}(W_{0}) - \mathcal{L}(W^{*}) + k_{1}T(\eta - \rho \eta^{2})\eta + \frac{k_{2}T\rho \eta^{2}}{2},$$
(45)

where W^* denotes a set of local optimal parameters, and T is the number of training iterations. Furthermore, taking the learning rate $\eta = \min(\frac{1}{\rho}, \frac{C}{\sqrt{T}})$, then we get

$$\frac{1}{T} \sum_{i=1}^{T} \mathbb{E} \|\nabla \mathcal{L}(W_{i-1})\|_{F}^{2}$$

$$\leq 2 \frac{\mathcal{L}(W_{0}) - \mathcal{L}(W^{*}) + k_{1}T(\eta - \rho\eta^{2})\eta + \frac{k_{2}T\rho\eta^{2}}{2}}{T\eta(2 - \rho\eta)}$$

$$\leq 2 \frac{\mathcal{L}(W_{0}) - \mathcal{L}(W^{*}) + k_{1}T(\eta - \rho\eta^{2})\eta + \frac{k_{2}T\rho\eta^{2}}{2}}{T\eta}$$

$$\leq \frac{2(\mathcal{L}(W_{0}) - \mathcal{L}(W^{*}))}{T\eta} + 2k_{1}(\eta - \rho\eta^{2}) + k_{2}\rho\eta.$$
(46)

We next discuss the value of C in two cases. If $\sqrt{T} > \rho$, we set C = 1. Hence, $\eta = \frac{1}{\sqrt{T}}$, and

$$\frac{1}{T} \sum_{i=1}^{T} \mathbb{E} \|\nabla \mathcal{L}(W_{i-1})\|_{F}^{2}
\leq \frac{2(\mathcal{L}(W_{0}) - \mathcal{L}(W^{*}))}{\sqrt{T}} + \frac{2k_{1}}{\sqrt{T}} + \frac{k_{2}\rho}{\sqrt{T}}
\leq 2\frac{\mathcal{L}(W_{0}) - \mathcal{L}(W^{*}) + k_{1} + k_{2}\rho}{\sqrt{T}}.$$
(47)

If $\sqrt{T} < \rho$, we set $C = \frac{\sqrt{T}}{\rho}$. It is clear that $\frac{1}{\rho} \le C \le 1$. Hence,

$$\frac{1}{T} \sum_{i=1}^{T} \mathbb{E} \|\nabla \mathcal{L}(W_{i-1})\|_{F}^{2}
\leq \frac{2(\mathcal{L}(W_{0}) - \mathcal{L}(W^{*}))}{C\sqrt{T}} + \frac{2k_{1}C}{\sqrt{T}} + \frac{k_{2}\rho C}{\sqrt{T}}
\leq 2\frac{\rho(\mathcal{L}(W_{0}) - \mathcal{L}(W^{*})) + k_{1} + k_{2}\rho}{\sqrt{T}}.$$
(48)

Based on Eq.47 and Eq.48, we have

$$\frac{1}{T} \sum_{i=1}^{T} \mathbb{E} \|\nabla \mathcal{L}(W_{i-1})\|_F^2 = O(1/\sqrt{T})$$
(49)

if we set the learning rate $\eta=\min(\frac{1}{\rho},\frac{1}{\sqrt{T}}).$ Then, the proof of Theorem 6 is completed.

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