

# EFFICIENT DISTRIBUTION FOR DEEP LEARNING ON LARGE GRAPHS

## ABSTRACT

Graph neural networks (GNN) are compute intensive; thus, they are attractive for acceleration on distributed platforms. We present DeepGalois, an efficient GNN framework targeting distributed CPUs. DeepGalois is designed for efficient communication of high-dimensional feature vectors used in GNN. The graph partitioning engine flexibly supports different partitioning policies and helps the user make tradeoffs among task division, memory usage, and communication overhead, leading to fast feature learning without compromising the accuracy. The communication engine minimizes communication overhead by exploiting partitioning invariants and communication bandwidth in modern clusters. Evaluation on a production cluster for the representative *reddit* and *ogbn* datasets demonstrates that DeepGalois on 32 machines is  $2.5\times$  and  $2.5\times$  faster than that on 1 machine in average epoch time and time to accuracy, respectively. On 32 machines, DeepGalois outperforms DistDGL by  $3.8\times$  and  $10.1\times$  in average epoch time and time to accuracy, respectively.

## 1 INTRODUCTION

Graph neural networks (GNN) (Wang et al., 2019; Huang et al., 2021; Hu et al., 2020b) have been proposed as a powerful approach for feature learning on graphs. GNNs involve intensive computation on high-dimensional feature vectors, which makes it more expensive than conventional graph analytics algorithms. A promising way to accelerate GNN computation is to distribute it on a cluster of CPUs or GPUs. Many distributed GNN systems exist (Zheng et al., 2020; Tripathy et al., 2020; Ma et al., 2019) that tackle the challenging problem of scaling out this computation.

In this paper, we present DeepGalois, an efficient GNN framework targeting distributed CPUs. DeepGalois is designed on top of the parallel graph processing system *Galois* (Nguyen et al., 2013), the customizable graph partitioning framework *CuSP* (Hoang et al., 2019), and the distributed communication substrate *Gluon* (Dathathri et al., 2018). By leveraging the flexibility of the graph partitioning engine in CuSP, DeepGalois users can orchestrate task division, memory usage, and communication overhead to make

the best tradeoff for fast feature learning without compromising accuracy. DeepGalois is the first distributed GNN framework that supports arbitrary vertex-cut partitioning policies; vertex-cuts are essential in scaling out to large clusters (Gill et al., 2018). Gluon is optimized for utilizing the communication bandwidth in modern clusters, and by leveraging Gluon, DeepGalois exploits partitioning invariants, performs local aggregation of messages, and combines messages to minimize communication overhead. DeepGalois adds additional support for vector datatypes in Gluon to support efficient communication for high-dimensional feature vectors.

We evaluate DeepGalois using a wide range of representative graphs on distributed CPUs. Experimental results show that DeepGalois on 32 machines is  $2.5\times$  and  $2.5\times$  faster than that on 1 machine in average epoch time and time to accuracy, respectively. On 32 machines, DeepGalois outperforms DistDGL by  $3.8\times$  and  $10.1\times$  in average epoch time and time to accuracy, respectively.

This work makes the following contributions:

- We present DeepGalois, a scalable GNN framework targeting distributed CPU clusters.
- We provide system support for efficient feature vector communication and flexible graph partitioning to improve scalability.
- Experimental results on up to 32 CPU machines demonstrate that DeepGalois scales well and outperforms the state-of-the-art.

## 2 BACKGROUND

### 2.1 Computation Patterns in Graph Neural Networks

Given an input graph  $G(V, E)$  with vertices  $V$  and edges  $E$  and input features  $h_v^0$  for each  $v \in V$ , a  $k$ -layer GNN learns an output feature vector  $h_v^k$  for each vertex  $v$  that can be used for downstream tasks. In the  $i$ -th layer ( $0 \leq i < k$ ), each vertex *aggregates* feature vectors from its neighbors and *updates* itself with the aggregated neighbor information and its own feature vector  $h_v^i$  to get its feature for layer  $i+1$ ,  $h_v^{i+1}$ . The computation for vertex  $v$  in the  $l$ -th layer follows:

$$m_v^{l+1} = \text{aggregate}(\{h_u^l | u \in \mathcal{N}(v)\}) \quad (1)$$

$$h_v^{l+1} = \text{update}(m_v^{l+1}, h_v^l) \quad (2)$$

$\mathcal{N}(v)$  is the set of  $v$ 's one-hop neighbors. Both functions can use trainable parameters that are learned during GNN training: for example, the update function typically has an associated set of weights that are used to transform the aggregated feature vector using a matrix multiply operation.

Many GNN models have been proposed, including graph convolutional networks (GCN) (Kipf & Welling, 2016), Graph Isomorphism Networks (GIN) (Xu et al., 2019), GraphSAGE (Hamilton et al., 2017), and graph attention networks (GAT) (Veličković et al., 2018). They differ in how they define the GNN's aggregate and update functions.

## 2.2 Existing Graph Neural Network Work

Many techniques exist to support scalable GNN computing. FastGCN (Chen et al., 2018) and GraphSAGE (Hamilton et al., 2017) proposed *mini-batching* and *neighbor sampling* to reduce computation and memory footprints. LGCL (Gao et al., 2018) and GraphSAINT (Zeng et al., 2019; Zeng et al., 2020) proposed *subgraph sampling* which trains GNNs via random-walked sampled subgraphs. GPNN (Liao et al., 2018) adopts two-level (local-global) propagation approach to handle extremely large graphs. Other techniques remove redundant aggregation (Zeng & Prasanna, 2020; Jia et al., 2020b) from common neighbors.

These techniques can be applied on top of distributed CPU/GPU GNN implementations to further accelerate training. Distributed CPU systems include AliGraph (Yang, 2019), Euler (Eul, 2020), DistDGL (Zheng et al., 2020), and AGL (Zhang et al., 2020), and multi-GPU systems include Roc (Jia et al., 2020a), NeuGraph (Ma et al., 2019), and CAGNET (Tripathy et al., 2020). Each system has varying methods of distribution of the GNN computation.

## 2.3 Distributed Vertex Programs

In graph analytics applications, a computation rule called an *operator* (Pingali et al., 2011) is applied repeatedly to a vertex  $v$  and its label until some termination condition is reached. The operator uses  $v$ 's label as well as the labels of  $v$ 's *neighborhood*, which the operator can define arbitrarily (e.g., beyond single-hop neighbors). A *vertex program* is an application where the neighborhood of an operator applied to vertex  $v$  consists only of  $v$ 's immediate neighbors. Many distributed graph analytics systems have been proposed to efficiently distribute the computation of vertex programs across machines such as D-Galois (Dathathri et al., 2018), Lux (Jia et al., 2017), and Gemini (Zhu et al., 2016). In particular, D-Galois is composed of two components that can be reused in any system to distribute vertex programs: the Gluon communication substrate which optimizes communication across machines using structural and temporal invariants of partitioning and the Customizable Streaming Partitioner (CuSP) (Hoang et al., 2019), a library which

partitions the graph.

We observe that *GNN computation is a vertex program*: aggregation followed by update is an operator that reads features (i.e., labels) from immediate neighbors and applies the update function update its own label. The termination condition is the number of layers in the network: this operator will be applied  $k$  times for a  $k$ -layer network.

## 3 DEEPGALOIS

### 3.1 Components of DeepGalois

*Graph Computation*: For computation on each machine, DeepGalois uses *Galois* (Nguyen et al., 2013) which is the state-of-the-art shared memory framework that provides parallel constructs and efficient abstract data structures for graph computations. For matrix multiplications required in GNN computations, DeepGalois uses the *BLAS* library. DeepGalois's modular design allows users to use any such library of their choice.

*Graph Partitioning*: DeepGalois uses a customizable streaming graph partitioner called *CuSP* (Hoang et al., 2019) that provides a simple API to design custom partitioning policies for graphs. CuSP allows the flexibility to customize domain specific partitioning policies for GNN applications; this is especially important for efficient communication (Section 3.3). CuSP provides many graph partitioning policies including edge-cuts, vertex-cuts, and hybrid-cuts, so users of DeepGalois can pick any of these policies for their specific GNN, dataset, and cluster (Gill et al., 2018).

*Synchronization*: Finally, for synchronization of vertex labels (aggregated features in the case of GNNs) among machines in the distributed setting, DeepGalois leverages *Gluon* (Dathathri et al., 2018) which is a partition-aware communication substrate for vertex programs.

The combination of these three components has been shown useful in the past in D-Galois (Dathathri et al., 2018), a state-of-the-art distributed graph *analytics* framework; DeepGalois applies it to the machine learning domain in this work which brings its own set of design challenges.

### 3.2 Partitioning and Data Placement

**Graph Partitioning** In DeepGalois, the *edges* of the graph are uniquely partitioned among all machines, and *proxies* for the endpoints of the edges are created on each machine. A vertex  $v$ 's proxy on a machine has a *cached copy of  $v$ 's feature vector* during GNN computation, and periodic synchronization occurs as necessary to synchronize the proxies of the same vertex on different machines (discussed further in Section 3.3). One of the proxies for a given  $v$  is designated the *master proxy*: it is responsible for determining the canonical feature vector with contribu-

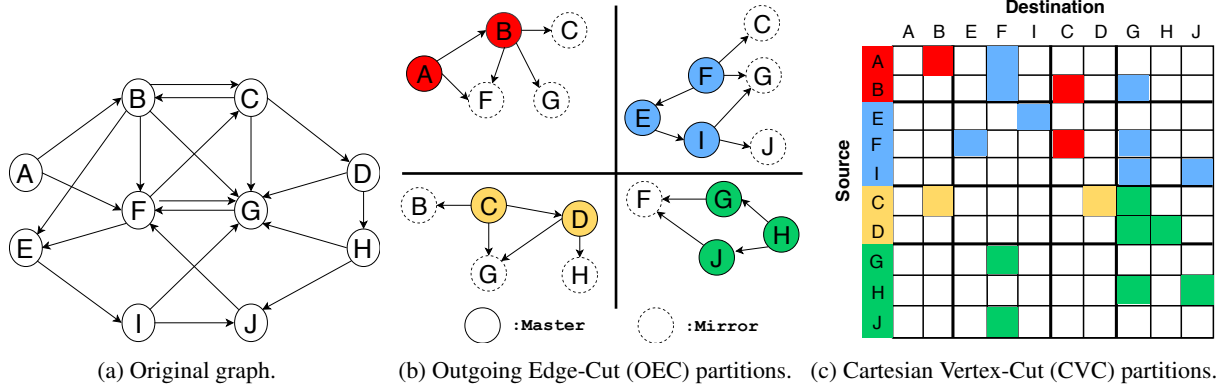


Figure 1. An example of partitioning a graph for four hosts using two different policies. (Hoang et al., 2019)

tions from  $v$ 's *mirror proxies* (i.e., all other proxies) and synchronizing this canonical value among all proxies.

Abstractly, all partitioning policies used by DeepGalois can be expressed with the definition of two heuristics (Hoang et al., 2019): (1) assignment of master proxies and (2) assignment of edges to machines. For example, a simple way to assign master proxies is to split them evenly among all machines in a blocked fashion. An outgoing edge-cut (OEC) would assign edge  $(u, v)$  to the machine that has  $u$ 's master proxy: this results in a partition where the machine with the master proxy ends up with all its outgoing edges. Another example of a more complex policy is a 2D Cartesian vertex cut (CVC) (Boman et al., 2013) which distributes edges according to a 2D block-cyclic distribution of the blocks of the adjacency matrix representing the edges of the graph. Fig. 1 illustrates OEC and CVC. The way the graph is partitioned affects computational load balance as well as the communication patterns during synchronization.

DeepGalois is the first distributed GNN implementation to allow for *arbitrary partitioning* of the graph via CuSP: this allows us to use partitions that scale better than the edge-cuts used in most distributed GNN systems (e.g., DistDGL) as we illustrate in our experimental results.

**Model Placement** As mentioned in Section 2.1, the aggregation and update functions for a GNN may have trainable parameters. We refer to the set of these parameters as the *model* that is trained during GNN execution. DeepGalois, *replicates* parts of the model that are required for carrying out the computation on each machine. For example, in GCN's update function, a vertex's aggregated feature vector is multiplied by a weight matrix to transform it. Since a vector-matrix multiply requires the entire matrix, the entire weight matrix will be replicated on all machines for each layer. The size of these parameters is small compared to the graph size and feature vectors, so full replication does not add much overhead. Keeping the model consistent across

all machines is done by synchronizing the model gradients across the machines during the backward pass of training.

### 3.3 Synchronization of Data

**Aggregation** After the graph is partitioned among machines, each partition is a self-contained subgraph with local proxies. The local application of the aggregation function on a given machine performs *local aggregation* for vertices in the partition on that machine. This does not involve any communication due to the cached copies in the local proxies. In other distributed GNN frameworks like DistDGL (Zheng et al., 2020), vertex features are queried from a server if they are not present locally, so server communication may be required to access vertex features. DeepGalois reduces the communication overhead by performing local aggregation without communication.

To get the full aggregated result after local aggregation, a logical *all-reduce* operation is required in which all partial values on proxies of a given vertex  $v$  are synchronized. This requires that the aggregation operation must be commutative and associative; most GNN aggregation functions satisfy this property. Synchronization using Gluon is a logical *reduction* on the master proxy to produce the final canonical value by reducing the contributions from mirror proxies followed by a logical *broadcast* of the canonical value from the master to mirror proxies. As they may be millions of vertices whose partial values may need to be synchronized, DeepGalois uses Gluon to combine partial values (or messages) for different vertices into a single message per host in each logical reduction or broadcast phases. Such *message aggregation* reduces the communication overhead by better utilizing the communication bandwidth (Dang et al., 2018).

It is important to note that the partitioning policy plays a key role in determining the communication volume during synchronization. Reduction will occur from machine  $a$  to machine  $b$  if  $a$  has a proxy for a vertex  $v$  that host  $b$  owns the master proxy of, and a broadcast occurs from  $b$  to all

machines that have a proxy for  $v$ . It is possible for  $a$  to need to broadcast to all machines in the system if a proxy exists for a vertex it owns on every machine: this does not scale well as the number of machines increases. Therefore, to avoid this problem, it is useful to use a 2D partitioning policy such as CVC which is more *structured*: due to the way edges are placed in CVC, a machine’s communication partners are strictly limited to only a subset of hosts, which allows communication to scale for a higher number of hosts. These communication optimizations based on partitioning policies have been detailed and studied extensively (Dathathri et al., 2018; Gill et al., 2018), and we leverage these findings here for efficient GNN synchronization.

**Model Gradients** During the backward pass of GNN training, each machine calculates only the gradient contributions of its master proxies. A simple sum all-reduce of all gradients among all machines (as all gradients are replicated on all machines) reconstructs the same gradients that would have been derived on a single machine. This ensures that distributed execution does not degrade accuracy.

## 4 EVALUATION

### 4.1 Experimental Setup

We use two platforms for evaluation. The first is a machine with Intel Xeon Gold 5120 2.2 GHz CPUs with 56 cores on 4 sockets and 187GB of DRAM. We refer to this machine as *Ghostwheel* (GW). The second is the Stampede2 supercomputer (Stanzione et al., 2017) at the Texas Advanced Computing Center (TAC); each machine has 48 Intel Skylake cores on 2 sockets and 192GB of DRAM. The machines are connected via a 100 Gb/s Intel Omni-Path network. We run on up to 32 of these machines.

We evaluate three GNN systems. Deep Graph Library (DGL) (Wang et al., 2019) is a library that provides many GNN models as well as data structures to easily build new models. DistDGL (Zheng et al., 2020) is the distributed variant of DGL: it has a subset of the models in DGL. DeepGalois is the system presented in this paper. For our experiments, we evaluate the Graph Convolutional Network model (Kipf & Welling, 2016) and GraphSAGE model (Hamilton et al., 2017) of GNNs on these systems (DistDGL does not have GCN). We evaluate DGL only on Ghostwheel, and we evaluate DistDGL only on Stampede2. We perform vertex classification for these datasets, and accuracy is for single-label vertex classification.

Table 1 shows the input graphs we use for evaluation. *reddit* is a representative graph used in previous works (Jia et al., 2020a; Wang et al., 2019), and the other two datasets are from Open Graph Benchmark (OGB) (Hu et al., 2020a). We were unable to run DistDGL with

	Dataset		
	reddit	ogbn-products	ogbn-papers100M
<b>Vertices</b>	232K	2.4M	111.1M
<b>Edges</b>	11.6M	123.7M	3.2B
$\bar{d}$	50	51	29
<b>Feature</b>	602	100	128
<b>Classes</b>	41	47	172
<b>Train / Val / Test</b>	0.66 / 0.10 / 0.24	0.08 / 0.02 / 0.90	0.78 / 0.08 / 0.14

Table 1. Input graphs.  $\bar{d}$  is the average degree.

		DGL	DeepGalois
<b>GCN</b>	reddit	0.623	0.885
	ogbn-products	3.525	1.712
<b>SAGE</b>	reddit	5.200	1.117
	ogbn-products	6.290	2.763

Table 2. Average epoch time (sec) on ghostwheel based on 200 epoch run (H=16, L=2).

ogbn-papers100M; we use this graph only to show scaling for DeepGalois. For simplicity and consistency, we use a 2-layer GNN with a hidden feature dimension size of 16. Feature dropout and mini-batching are disabled. We average runtimes over 200 epochs except for ogbn-papers100M where we only run for 5 epochs. More experimental setup details are in the appendix.

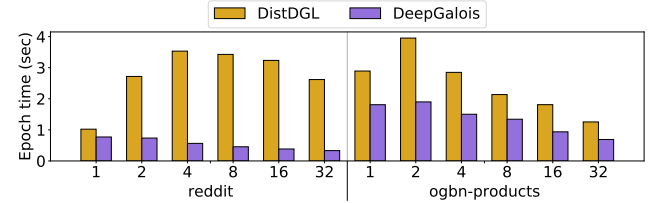


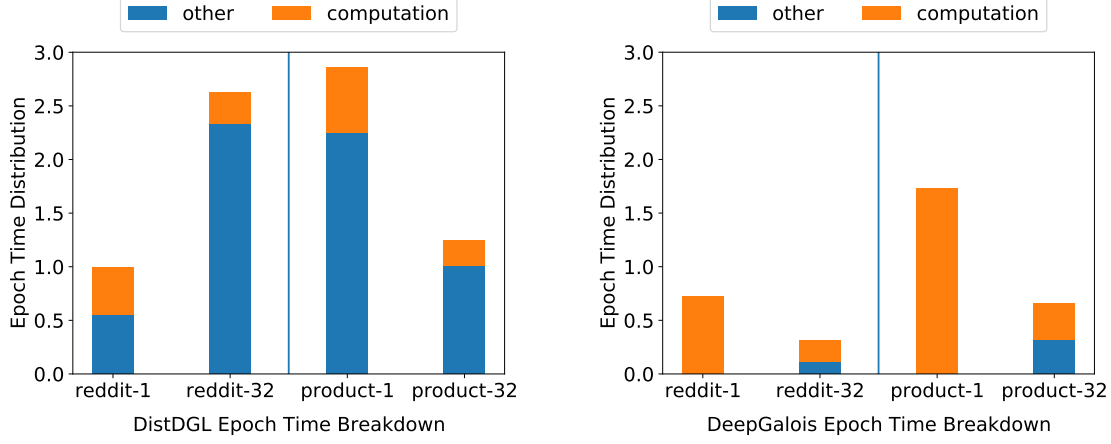
Figure 2. GraphSAGE epoch time in DistDGL and DeepGalois (mini-batching disabled in DistDGL).

### 4.2 Single Machine: DGL and DeepGalois

To show that DeepGalois matches a baseline for performance, we compare runtime with DGL on single-machine. Table 2 shows average epoch time of running 200 epochs. DeepGalois is competitive with DGL, showing that it can match the performance of existing GNN implementations.

Note that DeepGalois is much faster than DGL for GraphSAGE. This is due to an important optimization which orders aggregation and update depending on the dimensions of input embedding  $D_i$  and output embedding  $D_o$ . When  $D_i > D_o$ , update is performed before aggregation. Otherwise, aggregation is performed before update. In this way, aggregation (which is more expensive than update) is always performed on the shorter embeddings. In DGL, this optimization is applied to GCN but not to GraphSAGE because GraphSAGE must support multiple different aggregators, some of which (e.g., LSTM) prevent this optimization from being applied. This can be fixed in DGL by checking the





(a) DistDGL: “other” includes sampling and data copy time. (b) DeepGalois: “other” includes the communication time.

Figure 3. Epoch time breakdown of GraphSAGE on 1- and 32-machines. Note that “computation” involves full graph in DeepGalois since DeepGalois does not do neighbor sampling.

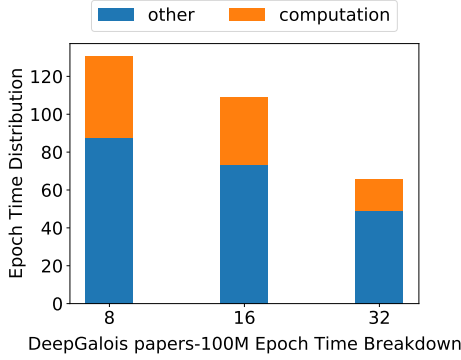


Figure 4. DeepGalois epoch time breakdown of GraphSAGE on papers-100M for 8-, 16- and 32-machines.

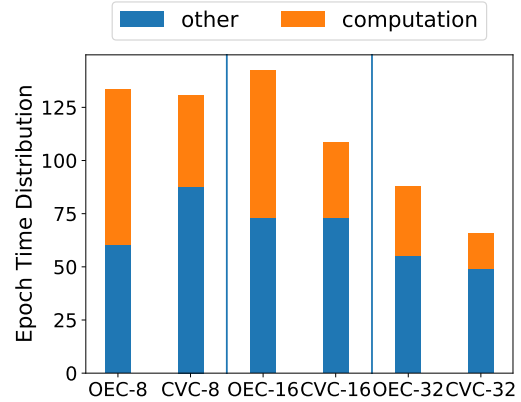


Figure 5. OEC vs. CVC partitioning policy in DeepGalois using ogbn-papers100M on 8-, 16-, 32-machines.

constraint compile time to see if this optimization is applicable. For example, with a mean aggregator, it can be applied.

### 4.3 Distributed Execution: DistDGL vs. DeepGalois

We compare DeepGalois with the state-of-the-art distributed GNN system, DistDGL. Unless specified otherwise, we report results of DeepGalois using CVC partitioning policy, which performs better than OEC. We will compare the two partitioning policies in Section 4.4.

#### 4.3.1 Performance Overview

Fig. 2 compares average training epoch time of DistDGL and DeepGalois on 1-machine to 32-machine for GraphSAGE. We observe that for reddit and products, DeepGalois is constantly faster than DistDGL even though DistDGL does neighbor sampling to reduce computation,

likely due to DeepGalois avoiding the data copying over the network done by DistDGL. Another observation is that DistDGL does not scale well, particularly for reddit. With 32-machine, DistDGL is  $2.6\times$  slower than it is on single-machine, which defeats the purpose of using a distributed cluster. This is also likely due to the significant data copying overhead, which would be non-trivial especially medium sized graphs like reddit where the reduction of computation time can not amortize the communication overhead. In comparison, DeepGalois achieves  $2.3\times$  speedup with 32-machine over 1-machine for reddit.

#### 4.3.2 Performance Breakdown

Fig. 3a and Fig. 3b compares average epoch time distribution of DistDGL and DeepGalois on different number of machines. We observe that there is a non-trivial overhead

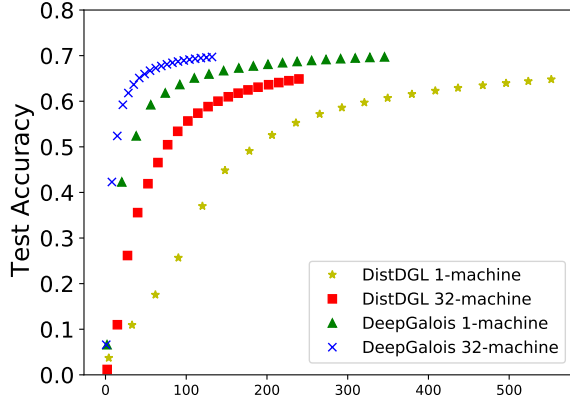


Figure 6. Speed of convergence for DeepGalois and DistDGL on 1-machine and 32-machine using `products`. Each two consecutive points in the figure cover 10 epochs.

in DistDGL to sample and copy the data. For DeepGalois, although it communicates data during each epoch, the communication time in DeepGalois is much less than the data copy time in DistDGL. Although there is some communication overhead in DeepGalois on 32 machines, since the computation time is significantly reduced, we still see substantial speedup over single-machine.

Fig. 4 illustrates the time distribution of DeepGalois on the large graph `papers-100M`. Due to the limited memory capacity on each machine, we can only run it with at least 8 machines. We observe that DeepGalois scales well to 32 machines. Computation time is reduced due to additional compute power. Notice that the communication time also decreases as number of machines increases: even though the total amount of communication increases, communication is overlapped with more machines.

#### 4.4 Scalability Impact of Partitioning Policies

Fig. 5 show the breakdown of DeepGalois average epoch time over `ogbn-papers100M` from 8 machines to 32 machines using OEC and CVC, respectively. We notice that CVC is slower than OEC with a small number of (< 32) machines, which matches observations in a previous study (Gill et al., 2018). However, OEC does have higher computation time than CVC due to load imbalance since OEC assigns all edges of a vertex to the same machine: this may not balance the workload well due to the high-degree vertices. Overall, CVC executes for less time, which is why we choose to use it in practice.

In this experiment, we show that the graph partitioning policy matters: the flexibility provided by CuSP is beneficial for the training speed. It is more important for GNN applications than it is for graph analytics applications to explore the

best graph partitioning policy, as there is much more computation and communication involved in GNN due to vector in GNN instead of scalar in graph analytics. Therefore, carefully balancing workload and reducing communication overhead is critical to the training speed.

#### 4.5 Accuracy Convergence Speed

Fig. 6 shows how the test accuracy changes over time for DistDGL and DeepGalois. We notice that with more machines, both DistDGL and DeepGalois converge faster than single-machine due to reduced epoch time. We observe that DeepGalois converges faster than DistDGL. There are two reasons. First, with the same number of epochs, DeepGalois’s accuracy increases faster, likely because DistDGL uses neighbor sampling but DeepGalois does not. Second, since DeepGalois runs faster than DistDGL, the convergence time is reduced.

## 5 CONCLUSION

This paper presents DeepGalois, a distributed GNN system built using proven distributed graph analytics systems and techniques applied to GNNs. DeepGalois scales as the number of machines grows due to efficient communication that leverages invariants of partitioning policies, and it outperforms DistDGL, a state-of-the-art distributed GNN system. We are in the process of adding GPU support to DeepGalois, and we also plan to add additional layer types such as GAT and techniques such as graph sampling.

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## A MORE EXPERIMENTAL SETUP DETAILS

The test accuracy used to determine the time-to-accuracy speedups in the abstract and intro was 93% for reddit and 60% for ogbn-products.

During partitioning for DeepGalois, we modified CuSP to balance training nodes among machines for reddit and ogbn-products, which had well-defined ranges for training nodes.

We disabled all minibatching in DistDGL in order to do a fair comparison with DeepGalois which does not have minibatching. DistDGL has neighborhood sampling (we could not disable this easily) and inductive training. DeepGalois does not do sampling and does transductive training. The Adam optimizer is used in all three systems with their default parameters in (Dist)DGL and learning rate 0.01, beta1 0.9, beta2 0.999, and epsilon  $1e - 8$  for DeepGalois.

The GitHub commit used for DistDGL runs is [fa5f4d6aa715f25303e497d070c4d7bb75fc47b3](https://github.com/DeepGalois/DeepGalois/commit/fa5f4d6aa715f25303e497d070c4d7bb75fc47b3) (Jan. 6, 2021). For DistDGL, we used the following hyperparameters. Number of trainers was set to 1 (setting it higher seemed to slow down execution). Number of servers was set to 2 (we did not notice much difference varying this number). Minibatching was disabled as mentioned above by setting the batch size to a very high number. The profiler was disabled. The number of workers (equivalent to number of samplers) parameter was set to 0: this was to avoid a bug in which evaluation of the validation and testing set would not work. OMP\_NUM\_THREADS was set to 48 (the number of threads on the cluster it was run on). We used the balance\_train and balance\_edges arguments to the graph partitioner, and the number of partitions created was equal to the number of machines used.