A Global DAG Task Scheduler Using Deep Reinforcement Learning and Graph Convolution Network

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\section*{ABSTRACT}
Parallelization of tasks and efficient utilization of processors are considered important and challenging in operating large-scale real-time systems. Recently, deep reinforcement learning (DRL) was found to provide effective solutions to various combinatorial optimization problems. In this paper, inspired by recent achievements in DRL, we employ DRL techniques for scheduling a directed acyclic graph (DAG) task in which a set of non-preemptive subtasks are specified by precedence conditions among them. We propose a DRL-based priority assignment model for scheduling a DAG task on a multiprocessor system, named GoSu, which adapts a graph convolution network (GCN) to process a complex interdependent task structure and minimize the makespan of a DAG task. Our proposed model makes use of both temporal and structural features in a DAG to effectively learn a priority-based scheduling policy via GCN and policy gradient methods. With comprehensive evaluations, we verify that our model shows comparable performance to several state-of-the-art DAG task scheduling algorithms, and outperforms them by 2\textendash{}3\% in the slowdown of achieved makespans particularly in nontrivial system configurations where workloads are neither too small nor heavy compared to the given number of processors. We also analyze the priority assignment behaviors of our model by leveraging a regression method that imitates the learned policy of the model.

\section*{INDEX TERMS}
Deep reinforcement learning, graph convolution network, policy gradient learning, DAG task.

\section*{I. INTRODUCTION}
In cyber-physical real-time systems, there has been an increasing demand for both high performance and strong timeliness to execute a pipeline of complex functions, e.g., autonomous driving with perception, planning and control. One of the key techniques to meet the demand is to exploit parallelism on a multiprocessor system. A directed acyclic graph (DAG) task has been used to represent the dependencies among a number of task components (subtasks) and to formulate a fine-grained parallel scheduling problem with the interdependent subtasks. Furthermore, as non-preemptive task models can avoid the overhead issue of migration and switching tasks, recently, priority-based non-preemptive scheduling for a DAG task has gained much attention [1]. The problem has been tackled by investigating scheduler techniques with priority assignments, which take a set of subtasks in a single DAG as input and produce a priority order for the subtasks and their non-deterministic execution order [1], [2].

Given a priority order, subtasks can be scheduled to run in parallel on multiple processors subject to the precedence conditions. Existing studies have developed heuristic priority assignment algorithms and analyzed their impact on the goal of minimizing the elapsed time (or makespan) required to complete all the subtasks [1]–[3], as there might be a number...
of different priority orders, e.g., up to \( n! \) where \( n \) is the number of subtasks. Due to the nature of heuristic strategies, they have not been able to establish fundamental design principles for DAG schedulers such as exploiting temporal and graph-structural spatial features under a variety of task configurations and scales, which highly affect prioritization.

In this paper, we present a learning-based priority assignment model for scheduling a single DAG task of multiple interdependent subtasks on a multiprocessor system with a non-preemptive mechanism. The model not only generates a priority order more favorable than existing approaches to achieving the goal but also automatically identifies critical temporal and graph-structural features of a DAG and systematically utilizes the features for priority assignments under various task configurations.

To do so, we use deep reinforcement learning (DRL) techniques, considering the difficulty to collect sufficient supervised labels for optimal priority assignments for DAG tasks. We also consider a large combinatorial problem space in priority assignments. Furthermore, we adapt a graph convolution network (GCN) with both forward-path and inverse-path aggregation to effectively extract temporal (e.g., execution times) and structural (e.g., precedence conditions) features from graph representations. We then present the GoSu (graph convolutional task scheduler) model for which the priority assignment policy using GCN-based embeddings is established through the policy gradient learning with slowdown-based rewards. In GoSu, each subtask is assigned a priority offline, and then during runtime, subtasks are scheduled on the basis of their priority order. This is the same as the fixed-priority (static-priority) scheduling in the field of real-time systems [4] where the highest-priority subtasks that are not yet scheduled are executed first.

The GoSu model consistently shows competitive performance compared to several state-of-the-art DAG scheduling heuristic methods based on fixed-priority assignments (i.e., [1], [2]) under various experimental settings. Furthermore, the model often achieves better performance than those heuristic methods particularly in nontrivial task configurations, e.g., showing a 2~3% gain in the slowdown of achieved makespans for the cases when the number of processors is 3 or 4 with moderate parallelism and when the number of processors is between 3 and 8 with high parallelism (Figure 4(b) and (c)). We also demonstrate that GoSu outperforms the others with a probability of up to 73% for each testing DAG task sample in several nontrivial cases (Figure 5(b) and (c)).

The main contributions of this paper are as follows.

- We present a DRL-based priority assignment model GoSu for scheduling a single DAG task on a multiprocessor system.
- We devise a GCN path aggregation scheme and a slowdown-based reward function specific to the objective of makespan minimization for a DAG task.
- We show performance gains by GoSu through comprehensive evaluations, and provide an analysis on the learned policy by projecting the GoSu model to a linear model via differentiable programming.

The rest of the paper is organized as follows. Section III briefly describes the DAG task scheduling problem and its objective. Section IV presents our proposed encoder-decoder model structure with GCN-based DAG embeddings and Attention-based priority assignments for subtasks in a DAG. Sections V and VI provide the experiment results and analysis, and the review on related work, respectively. Finally, Section VII concludes the study. In addition, Table 1 provides a list of symbols used throughout this paper with three aspects such as DAG tasks, model training, and datasets.
TABLE 2. Summary of related studies: At the top for DAG task scheduling, the "Properties" column describes which properties each method makes explicit use of, i.e., temporal (execution time, deadline) in "Temp." and spatial (precedence conditions) in "Spat." At the middle for DRL-based scheduling, the "Target" column describes the target application domain that each method focuses on. Our work addresses the issue of DAG task scheduling by using both temporal (O in "Temp.") and spatial (O in "Spat.") properties, and shares the similar problem specification with [1], [2] where node-level fixed-priority scheduling is explored; however, unlike those existing studies, GoSu leverages learning-based approaches through DRL and adapts GCN techniques subject to DAG representations. In addition, regarding the "Target" column at the middle, GoSu is a novel DRL-based DAG task scheduling model.

<table>
<thead>
<tr>
<th>DAG task scheduling</th>
<th>Properties</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>Temp.</td>
<td>Spat.</td>
</tr>
<tr>
<td>GEDF [5]</td>
<td>O</td>
<td>X</td>
</tr>
<tr>
<td>DAG-Fluid [3]</td>
<td>O</td>
<td>X</td>
</tr>
<tr>
<td>He19 [2]</td>
<td>X</td>
<td>O</td>
</tr>
<tr>
<td>Zhao20 [1]</td>
<td>O</td>
<td>O</td>
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<table>
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<tr>
<th>DRL-based scheduling</th>
<th>Target</th>
<th>Description</th>
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<tbody>
<tr>
<td>Name</td>
<td></td>
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<tr>
<td>DeepRM [8]</td>
<td>Cluster mgmt.</td>
<td>This model supports a scheduler for long running apps in a cloud environment, enhancing scales (e.g., up to 3K tasks) with recursive MLPs.</td>
</tr>
<tr>
<td>Metis [9]</td>
<td>Cloud computing</td>
<td>This model employs structural embeddings for data parallel tasks (e.g., spark), enabling scalable, online task scheduling in a cluster environment.</td>
</tr>
<tr>
<td>Decima [10]</td>
<td>Data cluster</td>
<td>This model casts the GIPPS problem into combinatorial optimization, employing the sequential decoder structure for priority assignments on a set of periodic tasks.</td>
</tr>
<tr>
<td>Panda [11]</td>
<td>Real-time system</td>
<td>This work verifies the efficiencies and theoretical bounds of global earliest deadline first (EDF) algorithms in DAG scheduling.</td>
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</table>

path are first scheduled and then other nodes that might immediately interfere with the critical path execution are scheduled. Zhao et al. [1] presented a single non-preemptive DAG scheduling framework that partitions nodes in a DAG into providers (i.e., nodes in the critical path) and consumers (nodes not in the critical path), intending to exploit both parallelism and dependency conditions. Given the partitioned nodes, the highest priorities are assigned to the providers, the second-highest priorities to nodes that might block the providers, and the lowest priorities to the other nodes. These prior works concentrate on structural features of DAGs of which the representation needs to be designed according to problem specifications and task configurations. Our work shares the same problem structure with these, focusing on fixed-priority scheduling and producing a priority order to minimize the makespan of a task running on a multiprocessor system. However, unlike these works based on sophisticated engineering procedures, our DRL-based model employs GCNs to extract important features from both individual node and graph-structural data, hence automating the learning of relevant enriched features.

There exist several works on applying DRL for scheduling tasks in the various domains such as cloud computing, networks, and manufacturing systems [8]–[11], [15]–[18]. DeepRM [8] was the first attempt to learn a scheduling policy systematically through DRL. This work introduces a time-slice based framework to solve scheduling problems in cluster management. However, this is limited in scales due to no consideration on permutation-invariant properties, i.e., task sets $\{r_1, r_2, r_3\}$ and $\{r_2, r_1, r_3\}$ are treated differently. Wang et al. [9] presented Metis, a DRL-based scheduler for managing online long-running applications in a cloud computing platform. They tackled the device placement issue in application containers across a cluster of server machines by exploring hierarchical DRL. Mao et al. [10] introduced a DRL-based Scheduling for large-scale cluster task scheduling. They developed sophisticated task and subtask embedding techniques and used REINFORCE algorithm for model training, maximizing service level objectives such as cluster utilization and job completion time. Recently, Lee et al. [11] adopted DRL for the global fixed-priority task scheduling on a multiprocessor system using the Transformer architecture.

While DRL techniques were employed for task scheduling in these studies, as summarized at the bottom of Table 2, the applicability of DRL and GCNs for DAG task scheduling has not been fully investigated. Our work focuses on the learning-based DAG task scheduling model by not only leveraging DRL and adapting GCN techniques but also analyzing the learned priority assignment policy of the model.

With the advancement of deep learning, numerous research works based on DRL were proposed to solve combinatorial optimization problems. The pointer network [19] provided a well-structured mechanism to adopt neural networks for combinatorial problems, e.g., traveling salesman problem (TSP). It establishes the probabilistic representation of a permutation by continuously estimating the probability of selecting a next item, e.g., the next city to visit in TSP. The pointer network was extended in a DRL framework [20]. This extension has been considered effective in that it is often impossible to establish labeled datasets of reliable quality (e.g., those annotated by optimal solutions) for large-scale combinatorial problems. In [21], a learning model based on
Transformer [22] was proposed to solve TSP and several other representative combinatorial problems. In particular, a simple rollout baseline for policy gradient algorithms was introduced to facilitate model training and fast convergence. In the same vein, our work leverages DRL in discrete optimization problem spaces but concentrates on DAG task scheduling with GCNs.

III. DAG TASK SCHEDULING

In this section, we describe the DAG task scheduling problem for which global fixed-priority scheduling (GFPS) [23] schemes can be applied.

In the DAG task scheduling problem with GFPS, we consider a single DAG task that consists of $n$ subtasks and precedence conditions among the subtasks. Considering a system model in which a DAG task runs, we consider a multi-processor platform of $m$ homogeneous processors with non-preemptive scheduling. We also presume that $n$ is much larger than $m$, considering resource limitations. Then, GFPS with a priority order for the $n$ subtasks is able to schedule the $m$ highest-priority subtasks among the subtasks that have not been scheduled but satisfy the precedence conditions in each time slot upon the system of $m$ processors. Since we focus on non-preemptive scheduling, each subtask cannot be preempted by any other subtasks once it starts its execution. The goal of this scheduling is to minimize the makespan of a DAG task. A priority order for the $n$ subtasks corresponds to a mapping of the subtasks to distinct integers $i \in \{1, \ldots, n\}$, and it is fixed. That is, each subtask in a DAG task is assigned such $i$ as its priority so that the precedence conditions in the DAG are satisfied and the makespan can be minimized, when the subtasks are scheduled according to the priority order on $m$ processors in GFPS. We consider a work-conserving scheduler that intends to always utilize all available processors.

Specifically, a DAG task $\tau$ is represented in a task dependency graph $G = (V, E)$ that specifies the precedence conditions among subtasks within $\tau$, where $V$ denotes a set of nodes corresponding to the subtasks (as many as $n$) and $E$ denotes a set of edges corresponding to the precedence conditions. Each node $v_i \in V$ represents a subtask of $\tau$ with worst-case execution time (WCET) $C_i$. Accordingly, we use the terms node and subtask interchangeably. Same as the DAG task scheduling formulation in the real-time system literature [1]–[3], we assume that the DAG specification including WCET $C_i$ is known in advance. Each edge $e_{ij} \in E \subset V \times V$ corresponds to a precedence condition between two subtasks, specifying that $v_j$ cannot run before the completion of $v_i$. That is, $v_i$ is a predecessor of $v_j$, and $v_j$ is a successor of $v_i$.

Furthermore, we use $\text{deg}_{\text{in}}(v_i) = |\{e_{ij} | e_{ij} \in E\}|$ and $\text{deg}_{\text{out}}(v_i) = |\{e_{ij} | e_{ij} \in E\}|$ to denote in-degree and out-degree of node $v_i$, respectively. In a task dependency graph, there might be more than a single source node such that $\text{deg}_{\text{in}}(v_i) = 0$ or a single sink node such that $\text{deg}_{\text{out}}(v_i) = 0$. In the case of multiple source nodes, we add a dummy source of WCET $= 0$ connecting to each of the multiple source nodes. Similarly, in the case of multiple sink nodes, we add a dummy sink of WCET $= 0$ connected to each of the multiple sink nodes. This allows us to consider only the DAG formation with one single source and sink. For example, Figure 1 illustrates a DAG task dependency graph with the precedence conditions among 8 nodes where $v_1$ and $v_8$ are source and sink nodes, respectively. In this example, the edge from $v_1$ to $v_2$ specifies that $v_2$ cannot be executed before the completion of $v_1$.

A path $\lambda = [v_{k_1}, v_{k_2}, \ldots, v_{k_l}]$ is a finite sequence of nodes such that $(v_{k_i}, v_{k_{i+1}}) \in E$ for $i \in \{1, \ldots, k - 1\}$. A path is called complete path if it contains both a source and a sink. A path length is the sum of the execution times of nodes in a path, i.e., $\sum_{v_i \in \lambda} C_i$. A path with the longest path length among complete paths is called critical path. We use $L$ to denote the critical workload, the sum of WCETs of only the nodes in a critical path, while we use $W$ to denote the total workload, the sum of WCETs of all nodes such that $W = \sum_{v_i \in V} C_i$.

For a DAG task $\tau$ of $n$ nodes, our model is learned to generate a priority order for a system of $m$ processors, which maps the nodes in $\tau$,

$$ [v_1, v_2, \ldots, v_n] $$

(1)

to a permutation of distinct integers from 1 to $n$ such as

$$ \pi = [\pi_1, \pi_2, \ldots, \pi_n]. $$

(2)

That is, given such $\pi$, we establish a priority-ordered node list,

$$ [v_{\pi_1}, v_{\pi_2}, \ldots, v_{\pi_n}] $$

(3)

which specifies that node $v_{\pi_1}$ is assigned the highest priority, $v_{\pi_2}$ is assigned the next highest priority, and so on for GFPS upon a system of $m$ processors.

We use makespan $M(\pi, \tau)$ to denote the elapsed time for task $\tau$ by $\pi$-scheduling, which is required to complete $\tau$ when GFPS with priority order $\pi$ is considered. Our model aims to find a priority order $\pi^*$ that minimizes the makespan of $\tau$, allowing efficient use of the computing resources of a system running $\tau$.

$$ \pi^* = \arg\min_{\pi} M(\pi, \tau) $$

(4)

Notice that our model can be used scheduling a periodic DAG task $\tau$ with its deadline $D$ and period $T$, where jobs
Note that throughout this paper, a subscript $\theta$ in an end-to-end fashion by DRL with calculated makespans.

makespan, the model with the encoder and decoder is trained to establish a probability distribution, i.e., $\pi$.

Given the learning objective to minimize the makespan of running each DAG task, so calculated makespans are used as reward signals for updating the model through REINFORCE algorithm [24].

for $\tau$ repeat regularly at the inter-release time of $T$. It is determined that the periodic task $\tau$ is schedulable if the model yields such $\pi^*$ that the time constraint of $\tau$, $M(\pi^*, \tau) \leq D$ holds. In this regard, it is also feasible to use the model for scheduling multiple DAG tasks, because for each DAG task, its WCET can be individually induced by its respective tight makespan bound.

IV. PROPOSED APPROACH

In this section, we describe our scheduling model that takes a DAG task as input and produces a priority order in GFPS for the task. Figure 2 illustrates the overall model structure with encoder and decoder modules. The modules are end-to-end trained through DRL to generate a priority order of a DAG task input, where the learning objective is to minimize the makespan of the task scheduled by the priority order. Specifically, we adapt graph learning with two-way path aggregation in the encoder to effectively extract the relational information in a DAG. We also employ a sequential selection procedure in the decoder to robustly update the ordering probability over timesteps of variable input task sizes.

In the GCN-based encoder, the raw features of individual $n$ nodes (or $n$ subtasks) in a DAG task $\tau$ are first processed via a feed-forward network (FFN). The structural information of $\tau$’s task dependency graph is encoded via the message passing mechanism of a GCN, and then latent vectors of the nodes are generated. With the latent vectors, the Attention-based decoder generates a priority order for the nodes of $\tau$, establishing a probability distribution, i.e., $p_\theta(\pi | \tau)$ over priority orders $\pi$. Given the learning objective to minimize the makespan, the model with the encoder and decoder is trained in an end-to-end fashion by DRL with calculated makespans. Note that throughout this paper, a subscript $\theta$ is used to represent trainable model parameters.

The decoding for a priority order is conducted in a sequential selection procedure of $n$ (time)-steps,

\[
p_\theta(\pi | \tau) = \prod_{t=1}^{n} p_\theta(\pi_t | \pi_1, \ldots, \pi_{t-1}, \tau) \tag{5}
\]

where Attention mechanism [19] is used to calculate the probability of each selection $p_\theta(\pi_t | \pi_1, \ldots, \pi_{t-1}, \tau)$. Each selection at step $t$ assigns the priority $(n-t)$ to a selected node. In general, a problem space for DAG task scheduling can be intractably large, e.g., there are $100!$ different combinations in priority orders for a DAG task with 100 subtasks, so it is not effective to use a single probability distribution on whole possible orders of $n$ subtasks. Rather, it is more feasible to use a sequential procedure to select a subtask with the next-highest priority iteratively, as have been studied in combinatorial optimization problems [19], [20].

In the following, we describe these encoding and decoding procedures based on DRL. We note functions or modules parameterized by $\theta$ using a subscript $\theta$. For example, an affine transform function parameterized by $\theta$ is represented in

\[
\text{Aff}_\theta(x) = W_\theta x + b_\theta. \tag{6}
\]

Note that parameters in different modules denote different parameter sets.

It is noteworthy that while our model is intended for scheduling a single DAG task with many interdependent subtasks, it can be used for scheduling multiple parallel DAG tasks. For a DAG task, the model generates a priority order for its subtasks and yields the makespan bound by the priority order. Then, such makespan bound can be used to estimate the WCET of a DAG task. Given a set of DAG tasks where each task is associated with its respective WCET estimated by the model, it is feasible to adopt multiprocessor real-time task scheduling methods (e.g., [11], [25]).

A. GCN-BASED ENCODER

The encoder learns to represent each node in a DAG task into a fixed-sized vector. Each node has its own execution time, and a set of nodes forms a graph structure with precedence dependency. To reflect both individual node information and graph-structured information, the encoder takes a two-step procedure in that it first transforms individual raw features of each node into vectors through an FFN and then incorporates graph structure into the vectors through a GCN.

1) NODE EMBEDDING

Using an affine transform with the tanh(·) activation, the encoder transforms raw features $x_i$ of node $v_i$ into a latent vector representation via an FFN.

\[
h_i^{(0)} = \tanh (W_\theta^{(0)} x_i + b_\theta^{(0)}) \tag{7}
\]

The raw features used in our implementation are all listed in Table 3. In addition to node-level features such as execution time, the number of incoming edges, and the number of outgoing edges, we also include several graph-level features such as critical and non-critical workloads.

2) GRAPH-LEVEL EMBEDDING

The encoder iteratively updates the generated vector representations with the affine projection above through a
GCN to further incorporate graph-structural features into the vector representations. In principle, the encoding process through a GCN can be seen as iterative message passings along the paths in a DAG. At step $k$, a latent representation $h_i^{(k)}$ of node $v_i$ is updated by aggregating latent representations of $v_i$’s neighbor nodes as shown in Figure 3. Note that in a GCN, $v_i$’s neighbors include directly connected nodes as well as $v_i$ itself. Accordingly, we define

$$N(v_i) = \{v_k|e_{ki} \in E\} \cup \{v_i\}$$ (8)

hereafter.

Then, the GCN message passing can be formulated as

$$h_i^{(k+1)} = \text{Update}(\text{Aggregate}_\theta(\{h_j^{(k)}|v_j \in N(v_i)\}))$$ (9)

where the $\text{Aggregate}$ function accumulates messages from the neighbors of $v_i$, and the $\text{Update}$ function takes the accumulated embedding and performs a nonlinear transformation on the embedding. The representations generated by iterating the above message passing operation $K$ times can contain structural features in that each node is differently aggregated according to graph topology. The representations also preserve node features; at $k = 0$, $h_i^{(0)}$ in Eq. (7) is an initial node embedding from node features that are preserved during GCNs along with structural varieties embedded [26].

For aggregation, we adopt Attention [27] similar to [28]. Specifically, the encoder performs the Attention operations such as

$$\text{Att}_\theta(\{h_j^{(k)}|v_j \in N(v_i)\}) = W_\theta \sum_{v_j \in N(v_i)} \alpha_{ij}h_j^{(k)}$$ (10)

where $\alpha_{ij}$ denotes the Attention coefficient that estimates the importance of node $v_j$ to node $v_i$’s representation. We restrict $\alpha_{ij}$ to satisfy $\sum_j \alpha_{ij} = 1$ and formulate it as

$$\alpha_{ij} = \frac{\exp(a^T_\theta W_\theta h_i + b^T_\theta W_\theta h_j)}{\sum_{v_j \in N(v_i)} \exp(a^T_\theta W_\theta h_i + b^T_\theta W_\theta h_j)}$$ (11)

where $a$ and $b$ are trainable vectors, and $W$ is a trainable matrix.

We exploit several Attention modules (heads) simultaneously and aggregate them to get the final latent representation.

This is useful since different Attention heads can give weights more relevantly to different latent representations ($h$). The $\text{Aggregate}$ function is implemented using multi-head Attention, and is defined as

$$\text{Aggregate}_\theta(\{h_j^{(k)}|v_j \in N(v_i)\}) = (\text{Att}_{\theta,1} \oplus \text{Att}_{\theta,2} \oplus \cdots \oplus \text{Att}_{\theta,H})$$ (12)

where $\oplus$ concatenates vectors, and $\text{Att}_{\theta,1}, \ldots, \text{Att}_{\theta,H}$ are $H$-individual Attention modules. $H$ denotes the number of distinct Attention heads per layer.

We then apply an affine transform followed by the exponential linear unit activation (ELU) [29] for the $\text{Update}$ function. This constructs the Attention network layer for the GCN message passing in Eq. (9).

$$h_i^{(k+1)} = \text{Update}(\text{Aggregate}(h_j^{(k)}|v_j \in N(v_i))) = \text{ELU}(W_\theta (\text{Att}_{\theta,1} \oplus \cdots \oplus \text{Att}_{\theta,H}) + b_\theta)$$ (13)

The result latent vectors $h_i^{(k)}$ are produced through consecutive GCN layers. We refer to those as node embeddings for $k = 3$ in our implementation.

3) TWO-WAY PATH AGGREGATION

In a DAG, an edge $e_{ij}$ normally specifies a predecessor condition such that a node $v_j$ cannot be executed without the termination of a node $v_i$. However, in the message passing, we observe that successor conditions are equally important as predecessor conditions. Thus, we adopt two types of Attention heads, incorporating them in the message-passing loop. This is similar with [22], [28] except we exploit different masks for each Attention head. Specifically, we define inverse neighbors as

$$N^{-1}(v_i) = \{v_k|e_{ik} \in E\} \cup \{v_i\}.$$ (14)

Then, among $H$ Attention heads, we set half of them to be forward-path graph Attention in Eq. (10) and the other half to be inverse-path graph Attention in Eq. (15).

$$\text{Att}_\theta(\{h_j^{(k)}|v_j \in N^{-1}(v_i)\})$$ (15)

This graph path aggregation in a GCN with both forward-path and inverse-path is illustrated in Figure 3. Employing Attention from the two path types enables the model to learn features on predecessor conditions as well.

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**TABLE 3.** Raw node-level features $x_i$ of node $v_i$ and additional graph-level features in DAG task $\tau$: $C_i$, $D_i$ denotes WCET of $v_i$, $W$ denotes total workload of $\tau$, and $L$ denotes critical workload of $\tau$, as described in Section III.

<table>
<thead>
<tr>
<th>Node features</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>normalized execution time in-degree</td>
<td>$C_i/W$, $\log(1+C_i)$</td>
</tr>
<tr>
<td>out-degree</td>
<td>$\deg_{out}(v_i)$</td>
</tr>
<tr>
<td>$v_i$ is source or sink</td>
<td>$0$, $1$</td>
</tr>
<tr>
<td>$v_i$ is in critical path</td>
<td>$0$, $1$</td>
</tr>
<tr>
<td>Graph features</td>
<td>Description</td>
</tr>
<tr>
<td>normalized critical workload</td>
<td>$L/W$</td>
</tr>
<tr>
<td>normalized non-critical workload</td>
<td>$(W-L)/W$</td>
</tr>
</tbody>
</table>
as successor conditions in a DAG while preserving their different relations.

B. SEQUENTIAL DECODER
With the node embeddings (the final latent vectors $h_i^{(k)}$ in Eq. (9)) from the encoder, for a DAG task $\tau$ of $n$ nodes, the decoder sequentially selects nodes to generate an $n$-sized priority order $\pi = [\pi_1, \pi_2, \ldots, \pi_n]$. If node $v_j$ is selected earlier than another $v_j$, then $v_j$ has a higher priority than $v_j$ for $i, j \in \{1, 2, \ldots, n\}$. Thus, $\pi$ corresponds to a priority-ordered node list $[v_{\pi_1}, v_{\pi_2}, \ldots, v_{\pi_n}]$. In the following, we omit superscript $k$ for $h_i$, as it is fixed in the decoder.

1) SEQUENTIAL NODE SELECTION
As formulated in Eq. (5), we decompose the probability function of a priority order into a product of probabilities of node selection in sequence. The decoder chooses a node to have the highest priority by sampling from probability distribution $p_{\theta}(\pi_t | \tau)$. It then chooses another node with the next highest priority (i.e., the highest priority among nodes not chosen yet) by sampling from $p_{\theta}(\pi_2 | \tau, \pi_1)$. For each selection at step $t$, the embedding of a partial priority order ($\pi_1, \ldots, \pi_{t-1}, \tau$) is used to generate the respective conditioned probability. This procedure continues until no node is left, so it comprises $n$-iterative decoding.

It is observed that the order of selected nodes is not important when choosing a node $v_j$ since those are indifferent in that they do not compete with $v_j$ [11]. Thus, we maintain two partitions of nodes, a set of nodes already chosen $O_t$ and a set of nodes that are not chosen yet $L_t$, and exploit the two sets $(O_t, L_t)$ as context information for $t = 1, \ldots, n$. The context information is used to calculate a probability distribution over nodes in $L_t$. In doing so, we aggregate $O_t$ and $L_t$ via nonlinear transformation and obtain their respective vectors $g_t^{(O)}$ and $g_t^{(L)}$ for steps $t = 1, \ldots, n$,

$$ g_t^{(O)} = \sigma_\theta \left( \sum_{j \in O_t} \tanh \left( \text{Att}_\theta(h_j) \right) \right), $$

$$ g_t^{(L)} = \sigma_\theta \left( \sum_{j \in L_t} \tanh \left( \text{Att}_\theta(h_j) \right) \right) $$

(16)

where $\sigma(\cdot)$ is an arbitrary nonlinear activation function.

We also consider the previously selected node $v_{\pi_{t-1}}$ as an important factor to determine its next node and thus make use of its embedding $h_{\pi_{t-1}}$ when making the successive selections. This is consistent with common observation such that fixed-priority task scheduling heuristics share the same pattern that similar priorities are assigned for tasks of similar properties.

$$ g_t^{(\text{last})} = \sigma_\theta(h_{\pi_{t-1}}) $$

(17)

Given the three vectors for the partitioned set and the previously selected node in Eq. (16) and Eq. (17), we combine them into a context vector by

$$ c_t = \text{Att}_\theta(g_t^{(O)} + g_t^{(L)} + g_t^{(\text{last})}). $$

(18)

Then, we exploit the context vector $c_t$ to derive a probabilistic inference for the $r$th subsequent node selection.

$$ p_{\theta}(\pi_t | \pi_1, \ldots, \pi_{t-1}, \tau) = \text{Softmax} \left( C \cdot \tanh(h_i \ W_\theta \ \text{Att}_\theta(c_t, \{h_j | v_j \in L_t\})) \right) $$

for all $v_j \in L_t$.

(19)

We multiply the output of $\tanh(\cdot)$ by an inverse temperature $C$ to confine logits within the range $[-C, C]$, where $C$ is empirically chosen. This derives the probability of priority orders.

$$ p_{\theta}(\pi | \tau) = \prod_{t=1}^n p_{\theta}(\pi_t | \pi_1, \ldots, \pi_{t-1}, \tau) $$

(20)

2) SAMPLING PRIORITY FROM THE DISTRIBUTION
Here we describe our strategy for sampling $\pi_t$ from the distribution at step $t$ in Eq. (19). We can conduct successive selection in a greedy way using

$$ \pi_t = \text{argmax} \left( p_{\theta}(\pi_t | \pi_1, \ldots, \pi_{t-1}, \tau) \right). $$

(21)

It is also possible to use stochastic sampling that randomly draws $\pi_t$ according to the distribution. Note that there are other sampling strategies than those simple approaches, e.g., $A^*$ [30], Beam Search [31], which are computationally expensive and thus are not appropriate for large-scale problem settings.

C. MDP FORMULATION
Here, we discuss the formulation of DAG task scheduling problems from the perspective of a Markov decision process (MDP). In general, an MDP is specified by a tuple with a set of states, a set of actions, a reward function, a state transition probability, and a discount factor.

- **State.** For scheduling a DAG task through sequential selection, a state needs to include information about the subtask partitions that evolve over time (i.e., a set of priority-assigned subtasks $O_t$ and a set of the others or priority-unassigned subtasks $L_t$ as in Eq. (16)). As step $t$ goes on, $O_t$ starts with an empty set and expands to a set containing more subtasks. More formally, according to the probability of priority orders in Eq. (20), a state consists of the embeddings about a DAG task $\tau$ with $n$ subtasks (i.e., $h_1, h_2, \ldots, h_n$) and priority-assigned subtasks; a state at step $t$ is given as

$$ ((h_1, \ldots, h_n), [\pi_1, \pi_2, \ldots, \pi_{t-1}]). $$

(22)

Recall that the embeddings for $\tau$ are calculated by the encoder in Section IV-A, and our model yields an $n$-sized priority order $\pi = [\pi_1, \pi_2, \ldots, \pi_n]$. Through sequential selection on the subtasks by the decoder, at each step $t \leq n$, a partial permutation $[\pi_1, \pi_2, \ldots, \pi_{t-1}]$ is generated. This corresponds to the indices of priority-assigned subtasks and is used as part of the state.
1) LEARNING OBJECTIVE

Action. Upon a state in Eq. (22), the decoder calculates an action to select \( \pi_t \) (i.e., assigning the \( t \)th highest priority to the \( \pi_t \)th subtask in \( \pi \) at step \( t \)). Accordingly, a set of actions corresponds to the distinct integers from 1 to \( n \), and a set of states corresponds to a partial permutation of the subtask embeddings.

Reward. A reward is yielded according to a given objective of scheduling, i.e., minimizing the makespan for a DAG task in Eq. (4). We specifically use the slowdown metric (in Eq. (26)) to evaluate the advantage of specific priority orders over others. The details of our reward design subject to DRL training are described in Section IV-E2.

Transition probability. In this MDP formulation, a transition is assumed to be deterministic in that for a state and action, the next state is determined without randomness. It is because an action for scheduling does not execute a task and it only affects the scheduling strategy and changes a permutation (a priority order) for a task. This setting is common when adopting DRL for large scale combinatorial optimization problems [20].

Discount factor. Given the DAG task specification of finite \( n \) subtasks, we have episodic DRL with terminal states where a full permutation is obtained, and accordingly, we set the discount factor to be 1.

D. COMPLEXITY ANALYSIS

In the encoder, \( K \) iterations of encoding via self-attention with masking for DAG precedence conditions are performed, where \( K \) denotes the number of GCN layers. The self-attention requires \( O(n^2d + nd^2) \) scalar multiplications for a single input of \( n \) subtasks and embedding dimension \( d \). Then, the encoding complexity is \( O(K \times (n^2d + nd^2)) \). In the decoder, for each selection (each time-step), context information for a partitioned set \( \{O_t, L_t\} \) in Eq. (16) and a subtask previously chosen is calculated in \( O(nd^2) \). This is iteratively performed \( n \) times for \( n \) subtasks, requiring \( O(n^2d^2) \).

Therefore, the complexity to infer a priority order for a DAG task input is \( O(n^2d^2) \). Note that the number of GCN layers \( K \) is much smaller than the number of subtasks \( n \) (i.e., as in Table 5) and the embedding dimension \( d \) (\( d = 64 \) in our implementation).

E. DRL TRAINING

As previously explained, the priority assignment for DAG task scheduling is formulated in a probability distribution in Eq. (20), which represents a policy in DRL. In the following, we describe how to establish such a policy through a policy gradient method.

1) LEARNING OBJECTIVE

The objective function \( J \) to learn Eq. (20) is defined as

\[
J = \mathbb{E}_{\pi \sim p_\theta(\pi)} [S(\pi, \tau)] = \sum_{\pi} p_\theta(\pi | \tau) S(\pi, \tau) \tag{23}
\]

where \( \pi \sim p_\theta(\tau) \) specifies that the priority order is sampled from a learned policy, and \( S(\pi, \tau) \) denotes score values which will be explained in Section IV-E2. We update model parameters \( \theta \) by the policy gradient in that differentiating the objective in Eq. (23) derives a gradient update rule as

\[
\nabla_\theta J = \sum_{\pi} \nabla_\theta p_\theta(\pi | \tau) S(\pi, \tau) = \sum_{\pi} p_\theta(\pi | \tau) \nabla_\theta \log(p_\theta(\pi | \tau)) S(\pi, \tau) = \mathbb{E}_{\pi \sim p_\theta} \left[ \nabla_\theta \log(p_\theta(\pi | \tau)) S(\pi, \tau) \right] \tag{24}
\]

using \( \nabla f(x) = f(x) \nabla f(x) \). Specifically, we use the Monte-Carlo stochastic gradient descent method or REINFORCE algorithm [24] to estimate the gradient in Eq. (24) and its average value over a batch of tasks,

\[
\nabla_\theta J = \frac{1}{|B|} \sum_{\tau \in B} p_\theta(\pi | \tau) \nabla_\theta \log(p_\theta(\pi | \tau)) S(\pi, \tau) \tag{25}
\]

where \( B \) is a batch list of tasks randomly sampled from a training dataset.

2) SLOWDOWN-BASED REWARD

As part of the objective function in Eq. (23), the score \( S(\pi, \tau) \) is used to formulate the relevance of priority orders \( \pi \) for task \( \tau \). This score is calculated based on reward values [32] in DRL. Given a DAG representation for parallel task scheduling on a multiprocessor system, our model is intended to learn an optimal priority order for \( \tau \) in terms of the makespan of \( \tau \). The makespan specifies the required time that elapses from the execution of \( \tau \)’s source node to its sink node. Specifically, we implement the slowdown-based reward (score) function \( S(\pi, \tau) \) using normalized makespans for scheduling a task \( \tau \) with a priority order \( \pi \), i.e.,

\[
S(\pi, \tau) = \frac{-M(\pi, \tau)}{M_{LB}(\tau)} \tag{26}
\]

where \( M(\pi, \tau) \) denotes the makespan of \( \tau \) by \( \pi \) and \( M_{LB}(\tau) \) denotes the lower bound of the makespan of \( \tau \) by any priority order. Note that \( \frac{M(\pi, \tau)}{M_{LB}(\tau)} \) represents a slowdown of \( \tau \)’s makespan by \( \pi \) compared to the ideal case for \( \tau \). Since the smaller the slowdown or the makespan, the better the priority order, we define negative rewards based on slowdowns.

The lower bound of the makespan of \( \tau \) upon an \( m \)-processor system is calculated by

\[
M_{LB}(\tau) = L + \max \left( 0, \left[ \frac{1}{m} (W - L \cdot m) \right] \right) \tag{27}
\]

where \( L \) and \( W \) are the critical workload and the total workload of \( \tau \).

Claim: The makespan of a task \( \tau \) is lower bounded by Eq. (27).

Proof: According to the definition of the critical path of a DAG task \( \tau \), the task requires at least the critical workload \( L \) in time to execute. During \( L \), the \( m \)-processor system can process at most \( L \cdot m \) for \( \tau \)’s nodes (in the case when no dependency blocks processing). In that ideal case, if \( \tau \)’s total
workload $W$ is no larger than $L \cdot m$, then $\tau$ can be completed within $L$. Otherwise, there is at least $(W-L \cdot m)$ workload not yet processed until $L$. It requires at least $\frac{1}{m}(W-L \cdot m)$ in time. Summing these two components establishes Eq. (27).

Reward normalization is intended to prevent overweighing of rewards of tasks with high parallelism. Suppose that we have an optimal order $\pi^*$ and another order $\pi$ of lower quality for a task $\tau$. It is observed that the less the parallelism degree of $\tau$, the smaller the makespan difference $M(\pi^*, \tau) - M(\pi, \tau)$. At one extreme, in the case where $\tau$ has no parallelism (e.g., there is only a single path from source to sink), every priority order yields the same makespan (reward). This case is not very meaningful for our model training. For establishing the stability of model training, it is critical to define rewards to be fully dependent on the quality of priority orders, especially when the structure of DAG tasks is complex.

3) BASELINE REDUCTION

In model training, we employ an additional variance reduction technique with baseline [32]. Specifically, we exploit a greedy baseline method, similar to [21], in which a target model $p_\theta$ and another base model $p_\beta$ are used. The two models share the same neural network structure but have distinct $\theta$ and $\beta$ parameters. For a task $\tau$, suppose that we obtain priority orders $\pi$ and $b$ where the former is sampled from $p_\theta$ with stochastic decoding and the latter is sampled from $p_\beta$. We then obtain a reward by $b$, i.e., $B(b, \tau) = -\frac{M(b, \tau)}{M_{LB}(\tau)}$ in Eq. (26). By replacing $S(\pi, \tau)$ in Eq. (24) with $S(\pi, \tau) - B(b, \tau)$, we adopt baseline reduction and hence establish the below.

$$
\nabla_\theta J \equiv \mathbb{E}_{\pi \sim p_\theta} [\nabla_\theta p_\theta(\pi | \tau) [S(\pi, \tau) - B(b, \tau)]]
$$

$$
= \frac{1}{|B|} \sum_{b \in B} \left[ \nabla_\theta \log(p_\theta(\pi | \tau)) \frac{[M(b, \tau) - M(\pi, \tau)]}{M_{LB}(\tau)} \right]
$$

$$
\mathbb{E}_{\pi \sim p_\theta} [\nabla_\theta p_\theta(\pi | \tau)B] = \nabla_\theta \mathbb{E}_{\pi \sim p_\theta}[B] = \nabla_\theta [B] = 0
$$

(28)

We use the base model $p_\beta$, if a makespan calculated by $p_\beta$’s priority order is statistically different from that calculated by $p_\theta$’s priority order. We perform the paired $t$-test on the makespans from the two models to check their difference, e.g., they are different if $p$-value is smaller than 0.01.

Our model training scheme is summarized in Algorithm 1.

V. EVALUATIONS

In this section, we evaluate the performance of our model.

A. EXPERIMENTAL SETTINGS

We describe the experimental settings including data generation, evaluation metrics, and the models in comparison.

Algorithm 1 Model Training of GoSu

// Parameter initialization
Initialze the target model $p_\theta$ with parameters $\theta$
Initialize the baseline model $p_\beta$ with parameters $\beta$
Initialize $\beta \leftarrow \theta$

// Model learning procedure
for $i \leftarrow 1, \ldots, N_{train}$ do

    $\Delta \theta \leftarrow 0$, Sample batch $B$ from training dataset $D$
    for task $\tau$ in $B$ do

        Sample priority order $\pi$ from $p_\theta(\cdot | \tau)$ stochastically
        Sample priority order $b$ from $p_\beta(\cdot | \tau)$ greedily
        $\Delta \theta \leftarrow \Delta \theta + \frac{1}{|B|} \left[ \frac{M(b, \tau) - M(\pi, \tau)}{M_{LB}(\tau)} \right] \nabla_\theta \log(p_\theta(\pi | \tau))$

    end for

// If target is better than baseline
if paired-T Test on $M(\pi, \tau) \neq M(b, \tau)$ then

    // Update baseline
    Update $\beta \leftarrow \theta$

end if

Update $\theta$ with $\Delta \theta$ using Adam Optimizer

end for

TABLE 4. Parameters for task generation with different degrees of parallelism: Unif() denotes uniform distribution.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>$n_{\text{depth}}$</th>
<th>$n_{\text{child}}$</th>
<th>$p_{\text{fork}}$</th>
<th>$p_{\text{pert}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>Unif(2, 5)</td>
<td>Unif(2, 5)</td>
<td>0.5</td>
<td>0.05</td>
</tr>
<tr>
<td>Moderate</td>
<td>Unif(3, 5)</td>
<td>Unif(2, 5)</td>
<td>0.5</td>
<td>0.1</td>
</tr>
<tr>
<td>High</td>
<td>Unif(3, 6)</td>
<td>Unif(2, 6)</td>
<td>0.5</td>
<td>0.15</td>
</tr>
</tbody>
</table>

1) DATASET GENERATION

As there is no publicly available large-scale DAG task datasets with a variety of configurations in real-time task specifications, we use synthetic datasets in which each DAG task is generated according to a nested fork-join task model [33]. This is a widely adopted scheme for analyzing and generating DAG tasks [34], [35]. The task generation algorithm works as follows, similar to [35]. For each node $v_i$ in the layer $k$, its child nodes $v_j$ and edges $e_{ij}$ are generated based on fork probability $p_{\text{fork}}$ where the number of child nodes in layer $k + 1$ are determined by uniform distribution $n_{\text{child}}$. This procedure starts from a source node and repeats for $n_{\text{depth}}$ times, thus creating a DAG of $n_{\text{depth}}$ layers. In addition, the edges of a node pair between the layer $k$ and the layer $k + 1$ are randomly added in the DAG based on perturbation probability $p_{\text{pert}}$. A large perturbation probability leads to a high degree of parallelism. Finally, the edges from nodes in the last layer to the sink node are added.

To perform experiments in various task configurations, we create three datasets with varying degrees of parallelism: Low, Moderate and High. The degree is configured by the aforementioned parameters such as fork probability $p_{\text{fork}}$, perturbation probability $p_{\text{pert}}$, the number of children $n_{\text{child}}$, and depth limit $n_{\text{depth}}$, as shown in Table 4. The characteristics of the datasets are summarized in Table 5.
TABLE 5. Characteristics of task datasets: # nodes and # edges denote the average number of nodes and edges. L and W denote the average workload of critical paths and the average total workload of DAG tasks.

<table>
<thead>
<tr>
<th>Dataset</th>
<th># nodes</th>
<th># edges</th>
<th>L</th>
<th>W</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>26.65</td>
<td>73.70</td>
<td>3284.23</td>
<td>10000</td>
</tr>
<tr>
<td>Moderate</td>
<td>44.80</td>
<td>129.47</td>
<td>1798.21</td>
<td>10000</td>
</tr>
<tr>
<td>High</td>
<td>83.70</td>
<td>339.75</td>
<td>1610.86</td>
<td>10000</td>
</tr>
</tbody>
</table>

2) IMPLEMENTATION

For evaluation, we implement a task scheduling simulator by which the makespan of each DAG task by a given priority order is exactly calculated upon a system of m processors. For implementing the dataset generation module and the models in comparison, we also exploit the open-source implementation provided in [1]. Our implementation is based on Python 3.7.9, PyTorch 1.6.0 [36], and PyTorch-geometric [37]. We train and test the models on a system of an Intel(R) Core(TM) i9-9940X processor with 160G memory, and an NVIDIA Tesla V100 GPU with CUDA 10.1 and cuDNN 7.6.0. In addition, we implement the heuristic algorithms and schedulability tests using Cython [38].

As for a multiprocessor system where DAG tasks are scheduled to run, we set its configuration, such as the number of homogeneous processors m, to be data specific and subsumed in training datasets. That is, for a model learned on specific datasets, each DAG task sampled from the datasets is configured to run upon a system of m processors during model training, and its evaluation system environment is set to have m processors. Thus, each model is evaluated in the same system environment with m processors (e.g., m ∈ {2, 3, 4, 6, 8}) in Figure 4) which it has been trained on.

The models in comparison are two state-of-the-art DAG task scheduling algorithms: He19 [2] and Zhao20 [1].

As described previously, our model aims at minimizing the makespan of individual DAG tasks. Accordingly, we measure the model performance in the slowdown ratio of an achieved makespan to its respective ideal makespan in Eq. (27) and use it as the evaluation metric.

3) MODEL HYPERPARAMETERS

The hyperparameter settings for our model are shown in Table 6. Unless otherwise mentioned, all the hyperparameters are set the same for all experiments.

We generate 10K datasets for each configuration. We use 8K samples for model training, 1K samples for model validation, and 1K samples for evaluation. In the encoder, we set the number of graph convolution layers to \( K = 2 \). For each graph convolution layer, we set the number of heads to 4 where two of them are forward-path Attention modules and the others are inverse-path Attention modules. We set the hidden representation layer to 64. We exploit dropout [39] with probability \( p = 0.1 \). We set the inverse temperature \( C \) to 5. A larger value of \( C \) makes models less exploitative. We set the batch size to 128 and use Adam Optimizer [40], where the learning rate sets to 0.0001. We clip gradients before model update by \((-1, 1)\).

B. PERFORMANCE COMPARISON

For each experiment condition where a dataset is defined by a specific DAG task configuration and the number of processors, we compare the performance of our model with that of other models.

Figures 4(a), 4(b), and 4(c) show the average relative slowdown (i.e., \( \frac{M_\pi(\tau)}{M_{\text{ideal}}(\tau)} \) in Eq. (26)) of achieved makespans with respect to various \( m \) processors \( (m = \{2, 3, 4, 6, 8\}) \) for low, moderate, and high parallelism datasets. As shown, our model achieves comparable performance for all the cases and outperforms He19 [2] and Zhao20 [1], with a relatively large margin of 2~3\\% for the cases when \( m = 3 \) or \( m = 4 \) with the moderate parallelism datasets and when \( m = 3 \sim 8 \) with the high parallelism datasets. This result is consistent with our expectation on DRL-based scheduling approaches such that they learn priority assignment rules tailored for specific environment settings, achieving performance improvement compared to other heuristics upon complex conditions. Notice that the more capable a system with more processors is (e.g., larger \( m \) settings), the less performance impact a specific priority order is likely to have on less or moderate parallelism datasets. This is because most executable subtasks can run immediately regardless of their priority assigned on many available processors. Another extreme case is a single processor system where it necessarily takes the total workload \( W \) in time to complete a DAG task regardless of priority orders.

The cases where the number of processors is 3 or 4 normally correspond to nontrivial configurations in between the cases of highly capable multiprocessor systems and constrained single processor systems in our experimental settings and datasets, rendering our DRL-based model more effective and achieving better makespans and slowdowns.

TABLE 6. Hyperparameter settings in datasets, neural network implementation, and training for our GoSu model.

<table>
<thead>
<tr>
<th>Description</th>
<th>Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dataset</td>
<td></td>
</tr>
<tr>
<td>Num. of train samples</td>
<td>8000</td>
</tr>
<tr>
<td>Num. of validation samples</td>
<td>1000</td>
</tr>
<tr>
<td>Num. of test samples</td>
<td>1000</td>
</tr>
<tr>
<td>Network</td>
<td></td>
</tr>
<tr>
<td>Num. of GCN layers K</td>
<td>2</td>
</tr>
<tr>
<td>Num. of Attention heads H</td>
<td>4</td>
</tr>
<tr>
<td>Embedding dimension d</td>
<td>64</td>
</tr>
<tr>
<td>Parameter initialization</td>
<td>Unif (−( \frac{1}{\sqrt{d}} ), ( \frac{1}{\sqrt{d}} ))</td>
</tr>
<tr>
<td>Dropout probability p</td>
<td>0.1</td>
</tr>
<tr>
<td>Inverse temperature C</td>
<td>5</td>
</tr>
<tr>
<td>Training</td>
<td></td>
</tr>
<tr>
<td>Batch size [B]</td>
<td>128</td>
</tr>
<tr>
<td>Learning rate</td>
<td>0.0001</td>
</tr>
<tr>
<td>Epochs</td>
<td>100</td>
</tr>
<tr>
<td>Optimizer</td>
<td>Adam</td>
</tr>
<tr>
<td>Gradient Clipping</td>
<td>(−1, 1)</td>
</tr>
</tbody>
</table>

1https://github.com/automaticdag/research-dag-scheduling-analysis
in our experiments, we include only the comparison with Zhao20. The data samples for each configuration are divided into three portions. The pink-colored bar indicates the portion of samples for which GoSu’s priority order outperforms (i.e., yielding a tighter makespan than) that of Zhao20 by at least 1% margin in terms of the slowdown in achieved makespans. The gray-colored bar indicates the portion of the opposite case samples. The green-colored bar indicates the portion of the other samples that tie.

It is consistently observed that GoSu performs more competitively in nontrivial configurations; the pink-colored bar increases when the number of processors is either 3 or 4, and its portion is up to 73% in the case when \( m = 3 \) with the moderate and high parallelism datasets. It is interesting that significant performance improvement is made with the systems of 2 and 3 processors for the low parallelism datasets and with the systems of a wider range of processor numbers, from 3 to 8, for the high parallelism datasets. This is because, for the low parallelism datasets, a system of more than 3 processors is likely to enable most executable tasks to run immediately and concurrently, thereby rendering specific priority orders less influential in terms of reducing makespans. For the high parallelism datasets, on the other hand, the same system has much opportunity to minimize makespans of individual tasks by priority orders that are appropriately chosen. This result demonstrates the benefit of GoSu performing steadily in various configurations including nontrivial cases. Through the paired-T tests, we also confirmed that except for the cases of \( m = 8 \) with the low and moderate parallelism datasets and \( m = 2 \) with the high parallelism datasets, our model shows statistically much better performance than the others.

C. COMPARISON OF GCN SCHEMES
In Table 7, we compare the models with different GCN schemes in terms of achieved slowdowns to verify the effect of our GCN processing with both forward-path and inverse-path aggregation. For comparison, GCNs are differently set to have only forward-path aggregation, only inverse-path aggregation, or both, while all the other model hyperparameters are set to the same.

For the low parallelism dataset, all the models show similar performance. However, as the complexity of parallel tasks grows, it is observed that the inverse-path aggregation becomes beneficial. For the moderate parallelism dataset, the model with only forward-path aggregation performs worse than the models with only inverse-path aggregation or both.
TABLE 7. The performance in the slowdown by the models with different GCN schemes: The number of processors is 4, and Both (GoSu) corresponds to the proposed GoSu model that has forward-path and inverse-path aggregation in GCN processing. The other models are set to consider either of two path types only.

<table>
<thead>
<tr>
<th></th>
<th>Low</th>
<th>Moderate</th>
<th>High</th>
</tr>
</thead>
<tbody>
<tr>
<td>Forward-Path Only</td>
<td>1.016</td>
<td>1.033</td>
<td>1.043</td>
</tr>
<tr>
<td>Inverse-Path Only</td>
<td>1.017</td>
<td>1.025</td>
<td>1.034</td>
</tr>
<tr>
<td>Both (GoSu)</td>
<td>1.017</td>
<td>1.024</td>
<td>1.029</td>
</tr>
</tbody>
</table>

For the high parallelism dataset, the model with both performs better than the others. The inverse-path aggregation for node v allows the model to embed the information of nodes blocked by v to the representation of v via message passing. This enables a better scheduling policy for large-scale and highly parallel DAGs.

D. CHARACTERISTICS OF LEARNED POLICY

To inspect what kind of policy the GoSu model learns through DRL, we conduct a regression-based experiment by employing differential programming techniques [41], [42]. It turns out that the experiment results allow us to identify which properties the model learns to value more. Specifically, we construct a linear regression model to produce a pointwise score $s_i$ for a node (subtask) $v_i$ and then fit the regression model to render a score list $[s_1, s_2, \ldots, s_n]$ consistent with the ranking of a priority order $\pi$ generated by GoSu. That is, the linear model is fitted to $[s_\pi(1), s_\pi(2), \ldots, s_\pi(n)]$ which is correctly sorted in the descending order with $\pi$ using Fast-Soft-Sort [41].

For simple analysis, we selectively use the following raw features of individual nodes as input: execution time, out-degree, in-degree, and whether a node is in the critical path (is-critical). A linear model $s_i = w_i x_i + b$ is learned where each element of $w$ is set to be in [0, 1] and $b$ in [0, 10] using clipping. We pose strong L1 regularization 3 on $w$ to make the weights of irrelevant properties be zeros, thus having only weights of significance for scheduling decisions [43], [44].

Figure 6 shows the weight values $w$ of the linear model with respect to different parallelism datasets. If an input feature has a large weight, it can be interpreted as an important factor for the priority assignment policy of GoSu. We observe that the out-degree is most critical for all task configurations. This is consistent with the effect of the inverse-path aggregation in Section V-C, showing that a larger out-degree means the node is more likely to be blocking other nodes and considered important. However, the in-degree does not play an important role, and its weight becomes zero in all task configurations. Another important observation is that the GoSu’s policy varies depending on task configurations. As the parallelism increases, the weight of the is-critical feature increases while those of the out-degree and the execution time decrease. This indicates the adaptability of the DRL-based model for a variety of task configurations. Note that we view that the linear model correctly imitates GoSu since the linear model yields similar but slightly worse (about 6%) performance than GoSu due to its structural limitations and lack of features.

FIGURE 6. Characteristics of learned policy in GoSu: The Y-axis denotes the normalized weight of features in the linear model with respect to different task configurations on the X-axis. The higher property weights (e.g., $w \in [0, 1]$ of a linear model $w_i x_i + b$) of features can be interpreted as having a greater impact on scheduling. This significance of features varies across different task configurations. GoSu can learn an appropriate scheduling policy without exhaustive feature engineering.

E. SCHEDULING MULTI-DAG TASKS

Here, we discuss how to extend our approach for scheduling multi-DAG tasks [35]. Given a periodic DAG task $\tau$ with deadline $D$ and period $T$, we calculate the makespan of $\tau$ using the priority order (i.e., $\pi^*$ in Eq. (4)) produced by our model and set it as $\tau$’s WCET. In this way, we induce WCET estimates of multi-DAG tasks. Then, it is possible to adopt priority assignment methods in real-time task scheduling (e.g., deadline monotonic (DM)) to schedule multi-DAG tasks.

In Figure 7, we shows the performance in schedulability ratio of multi-DAG tasks by three methods calculating WCETs using different single DAG scheduling approaches. We adopt the same DM algorithm and the same schedulability test in [1] for the three methods. The schedulability
ratio represents ratio between the number of schedulable task sets and the number of tested task sets, where each task set consists of 12 individual DAG tasks in this experiment. We set the number of processors to 4 and test 5000 task ratio. As shown, GoSu achieves better performance in schedulability ratio than the other methods. This result implies that tighter makespan bounds induced by GoSu can lead to better performance in scheduling multi-tasks. The performance gain is relatively larger when the utilization is high. This is consistent with the benefits of GoSu particularly for nontrivial cases, as the same scheduling and schedulability test methods are used.

VI. CONCLUSION

In this work, we presented GoSu, a DRL-based priority assignment model for DAG task scheduling, which adapts graph learning to utilize the graph structure of a DAG task. On graph embedding results, our model performs a sequential decoding procedure to obtain a permutation for subtasks in a DAG task. That permutation represents a priority order for task scheduling to minimize the makespan of the DAG task. Through extensive experiments, we demonstrated that GoSu achieves robust performance in the slowdown of DAG tasks, compared to other state-of-the-art priority assignment heuristics. We also showed the adaptability of our DRL-based approach for various configurations by leveraging the rank regression mimicking the policy of GoSu.

The direction of our future work is to develop an integrated learning approach of hierarchical DRL and GCN techniques for large-scale virtual application management problems. As network service chains consist of many virtual network functions, they can be analyzed through GCNs to be mapped on underlying network infrastructures with heterogeneous resources in data centers. While exploiting the structural similarity in the use of graph representation learning between resources in data centers. While exploiting the structural similarity in the use of graph representation learning between

REFERENCES


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