STRONGHOLD: Fast and Affordable Billion-Scale Deep Learning Model Training

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Abstract—Deep neural networks (DNNs) with billion-scale parameters have demonstrated impressive performance in solving many tasks. Unfortunately, training a billion-scale DNN is out of the reach of many data scientists because it requires high-performance GPU servers that are too expensive to purchase and maintain. We present STRONGHOLD, a novel approach for enabling large DNN model training with no change to the user code. STRONGHOLD scales up the largest trainable model size by dynamically offloading data to the CPU RAM and enabling the use of secondary storage. It automatically determines the minimum amount of data to be kept in the GPU memory to minimize GPU memory usage. Compared to state-of-the-art offloading-based solutions, STRONGHOLD improves the trainable model size by 1.9x~6.5x on a 32GB V100 GPU, with 1.2x~3.7x improvement on the training throughput. It has been deployed into production to successfully support large-scale DNN training.

Index Terms—Deep learning, Distributed training, DNNs training acceleration

I. INTRODUCTION

We are seeing exponential growth of deep neural network (DNN) sizes. For example, in a short span of 3 years, model size has grown by over 1000 folds from around 100 million model weights in 2018 for ELMo [1] or BERT [2] to 175 billion for GPT-3 [3] in 2020 and 530B for MT-NLG [4] in 2021. At the same time, the GPU memory capacity is increased by less than 3 folds per GPU generation. For instance, the latest high-end NVIDIA A100 GPU has 80 GB of memory, which is a 2.5x improvement on the GPU memory capacity over the 32 GB V100 predecessor. As a larger model tends to provide a better learning ability over the smaller counterparts, many future DNNs will continue to integrate more parameters and consume more GPU memory. This will further increase the gap between the memory demand of DNN training and the available GPU memory, making entry into large model training out of reach for many data scientists and academics.

Large DNN training can be achieved by utilizing aggregated computing resources of multiple GPUs through parallel and distributed computation. Examples of such strategies include data [5], model [6] and pipeline [7], [8] parallelisms and by placing the model parameters, gradients and optimizer states across computing devices [9]. However, all these distributed training schemes require sufficient computing resources, which can incur significant infrastructure and operational costs [10]. For example, to efficiently train a 10B parameter model currently requires a computing system with 16 NVIDIA V100 GPUs, costing over $100K to purchase [11]. While such cost may not concern big tech firms, it can place a big financial burden on small businesses and academic organizations.

Efforts have been made to reduce the GPU memory pressure and computing resource requirement for large DNN training. This is done by either trading precision for lower storage space [12] or leveraging CPU memory [13], [14], [15], [16]. The former uses low or mixed precision representations for model states (e.g., model parameters, gradients and optimizer states) to reduce GPU memory consumption, but it can slow down the model convergence speed. The latter approaches reduce the GPU memory requirement by either implementing a software-managed cache or using a dedicated memory allocation scheme. However, most of the techniques for using CPU memory are designed for convolutional neural networks (CNNs), where the memory consumption during training is dominated by the dynamically generated activations rather than the optimizer (e.g., SGD) states. Unfortunately, these techniques are ill-suited for the latest attention-based language models (e.g., Transformer-based DNNs) that have become the de facto approach for building state-of-the-art DNNs [17], [2], [3], where the model and optimizer (e.g., Adam) states rather than activation memory is the memory bottleneck.

ZeRO-Offload [11] and L2L [18] were among the first attempts to leverage CPU memory to train large Transformer-based DNNs. ZeRO-Offload moves optimizer states from the GPU to the CPU memory while keeping the entire model parameters in the GPU memory. It is able to train a model with 13B parameters on a 32GB V100 GPU. As ZeRO-Offload requires storing the entire model parameters in the GPU, the trainable model size is limited by the smallest part of the available GPU and CPU memory capacity. L2L is specifically designed for Transformer-based models, only keeping one Transformer block in the GPU memory by dynamically offloading the model parameters. ZeRO-Infinity [19] adopts a dynamic strategy to leverage the secondary storage (e.g., NVMe) and partitions the model parameters and optimizer states across heterogeneous memory hierarchy. While these approaches can increase the trainable model size, as we will show later in the paper, they come with significant overhead, leading up to 1.76x slowdown (up to 29.2x when NVMe is used). Such magnitude of overhead greatly reduces the practicability of these techniques in training and fine-tuning large DNNs where the training time is a major concern [9].

We present STRONGHOLD, a new approach for leveraging...
heterogeneous memory resources to scale up the trainable model size on GPUs without significantly compromising the training efficiency. The key insight of STRONGHOLD is to keep just a sufficient number of layers and their model parameters in the GPU to avoid GPU stalls during data offloading. Doing so can increase the trainable model size because only a small part of the model states are presented at the GPU memory at any time. By keeping the GPU computation going with overlapped data transfer, this strategy can also minimize and even hide the CPU-GPU communication overhead. To this end, STRONGHOLD implements a software pipeline between the GPU and CPU to dynamically offload the required model states between two memory spaces, without user code refactoring. Unlike ZeRO-Offload, STRONGHOLD does not store all model parameters in the GPU. Instead, it implements a dynamic working window to only store a few model layers in the GPU. It then dynamically moves the required layers and their parameters and the generated gradients between the CPU and the GPU. The STRONGHOLD runtime automatically determines a suitable working window size to ensure that asynchronous CPU-GPU data transfer can be overlapped with GPU computation to hide the data transfer latency.

A key challenge of STRONGHOLD is to decide how many layers are kept in the GPU. Having an unnecessarily large working window would waste the precious GPU memory with little performance gain while using an insufficient working window makes it difficult to overlap the CPU-GPU data transfer with GPU computation to minimize the overhead. We address this challenge by modeling the GPU computation with data transfer and using an analytical model to derive the right working window size based on profiling information collected from a few initial iterations during the warm-up phase.

STRONGHOLD runs multiple concurrent optimizers on the multi-core CPU to parallelize the model parameter update process. Since parameter update, data transfer, and GPU computation are asynchronous processes, they run in parallel to utilize the hardware parallelism. To reduce the GPU memory management overhead, STRONGHOLD employs a user-level GPU memory management scheme for the working window to avoid frequent invocations of the expensive GPU memory operations. We show that by reducing the GPU memory footprint, STRONGHOLD also opens up new opportunities to implement data parallelism within a single GPU. This is achieved by running multiple training workers across concurrently running streaming multiprocessors (SM) of a GPU while keeping one copy of model parallelism. This optimization leads to comparable or even better training performance over expert-tuned implementation when training a billion-scale model.

We implemented STRONGHOLD in PyTorch [20] and evaluated it in single and distributed GPU environments. We compare STRONGHOLD to three state-of-the-art offloading solutions [18], [11], [19]. Experimental results show that STRONGHOLD can support larger DNNs with a higher training throughput than competing schemes. Specifically, STRONGHOLD supports the training of a model with 39.5B and 82.1B parameters on a single 32GB V100 GPU and eight distributed 24GB A10 GPUs respectively without significantly compromising the training efficiency.

We envision STRONGHOLD to be attractive in two scenarios. It is useful in fine-tuning a large pre-trained DNN or using a pre-trained DNN to guide the training of a small model (a.k.a. knowledge distillation [21]) using limited GPU resources. This feature makes large DNN fine-tuning more accessible and affordable to small organizations and data scientists. It is also useful for accelerating model training by reducing cross-node communications or utilizing fine-grained GPU parallelization. For this use case, STRONGHOLD has been deployed to the production environment of Alibaba to support the training of DNNs with hundreds of billions and trillions of parameters.

This paper makes the following contributions:

- A new CPU-GPU offloading framework to scale up the trainable model size (Section III);
- An analytic model to determine the right working window for dynamic DNN training offloading (Section III-C);
- A new fine-grained GPU parallelism to speed up DNN training (Section IV-A).

II. BACKGROUND AND MOTIVATION

A. Deep Learning Model Training

DNN training typically consists of millions of iterations performed across multiple training epochs. Each iteration mainly involves three stages: forward propagation (FP), backward propagation (BP) and parameter update. In the FP stage, a batch of the training samples are passed through the DNN model to compute a loss based on an objective function. In the BP stage, the loss value is propagated reversely through model layers to compute the gradients. In the last stage, an optimizer uses the aggregated gradients to update parameter weights of individual model layers.

The memory consumption during DNN training largely stems from model states and residual states. Model states include model parameters, gradients and optimizer states (i.e., Adam optimizer [22] stores momentums and variances for parameter updating). Residual states include activations (i.e., the intermediate tensors saved for BP stage to produce gradients) and other temporary buffers. When training large DNNs, model states dominate the memory consumption, which can account for 87.5% of the GPU memory footprint when low-precision (i.e., 16-bit precision) is used [9].

Large DNN models are trained with parallelization techniques. For a model that can fit into the device memory for training, data parallelism is commonly used to distribute the training samples across multiple devices to improve the training throughput. When the model cannot fit into the device memory, model parallelism [23], [24] and pipeline parallelism [25], [7] can be leveraged to split the model layers or parameters to make the best use of the memory across multiple devices. While these three parallelism strategies can produce a synergy, model and pipeline parallelism often require additional code refactoring to split a DNN into model and pipeline components.
B. CUDA Streams

Modern GPUs consist of a large number of processing units, which are organized as streaming multiprocessors (SMs). For example, the NVIDIA V100 GPU supports 80 SMs, where each SM has a fixed number of cores. In the CUDA programming model, instructions placed within a single CUDA stream are executed sequentially. However, operations placed in different CUDA streams can be executed concurrently in different hardware SMs. STRONGHOLD uses multiple CUDA streams to accelerate DNN training when possible.

C. Motivation

As a motivation example, consider Figure 1 that shows the performance given by ZeRO-Offload [11] and ZeRO-Infinity [19] - the state-of-the-art offloading solutions for large DNNs. This experiment was conducted on a GPU server with a 32GB NVIDIA V100 GPU and 755GB of DDR4 RAM (see Section V-A). As a reference, we use Megatron-LM, NVIDIA’s heavily-optimized library for Transformer-based model training.

Figure 1a shows the largest trainable model size for each approach, while Figure 1b shows the throughput (i.e., the number of samples processed per second) on a common 1.7B Transformer-based model (the largest model supported by Megatron-LM on our platform). Although techniques like ZeRO-Offload and ZeRO-Infinity can scale up the trainable model size, this comes at the cost of significantly lower throughput and poor efficiency. For example, ZeRO-Offload enables training of a model that is 3x larger than Megatron-LM, but the training throughput on the 1.7B model is 6.7x less than Megatron-LM. By offloading some of the model parameters and states to the secondary NVMe SSD, ZeRO-Infinity (w/ NVMe) scales up the trainable model size by 29x over Megatron-LM, but its throughput drops by over 800x compared to Megatron-LM on a 1.7B model. This poor training efficiency of existing offloading solutions makes them impractical to train large models due to the long training time. STRONGHOLD aims to avoid this pitfall.

III. OUR APPROACH

STRONGHOLD is our open-source framework designed to enable efficient training of large DNNs on single or distributed GPUs. This is achieved by implementing dynamic software prefetching and offloading techniques to only store part of the model states - the main GPU memory consumer for large DNN training - in the GPU memory. STRONGHOLD utilizes the CPU memory and secondary storage to reduce the GPU memory pressure for training large DNNs. By doing so, the trainable model size is no longer bounded by the GPU device memory but the system’s storage capacity. STRONGHOLD is designed to be used without any user code refactoring similar to standard data-parallel training in PyTorch.

STRONGHOLD advances ZeRO-Offload [11], the state-of-the-art static CPU-GPU DNN training offloading framework, by offloading optimizer states onto the CPU side. Like ZeRO-Infinity [19], STRONGHOLD can also leverage secondary storage, but it delivers a higher throughput than ZeRO-Infinity.

A. Overview of STRONGHOLD

Figure 2 gives a high-level overview of the STRONGHOLD dynamic offloading scheme. The idea is to store model states (parameters, gradients and optimizer states) for selected DNN layers in the GPU memory. This is achieved by managing a working window in the GPU, where layer states are dynamically moved between the GPU and CPU memory. Precisely, the STRONGHOLD runtime swaps the already computed layer states to the CPU RAM. Actions ➊ and ➋ indicate offloading layers that have been used during FP and BP from GPU memory to CPU RAM. These actions also trigger ➌ and ➍ to prefetch the future-used layers from CPU RAM to GPU memory.

Figure 1: Performance of Megatron-LM and ZeRO-based solutions measured on a 32GB V100 GPU by the trainable model size (a) and throughput on a 1.7B model (b).

Figure 2: Dynamic model state offloading of STRONGHOLD. STRONGHOLD stores some DNN layers in the GPU memory and swapping out the finished layer states to the CPU RAM. Actions ➊ and ➋ indicate offloading layers that have been used during FP and BP from GPU memory to CPU RAM. These actions also trigger ➌ and ➍ to prefetch the future-used layers from CPU RAM to GPU memory.

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B. Preprocessing

During the model loading stage, STRONGHOLD extracts DNN layers and their execution order from the tensor graph. Most Transformer-based models follow a sequential layer execution order by stacking multiple Transformer blocks (Figure 3a), resulting in a static relationship. Extracting the layer execution order of such model architectures is straightforward. However, there are also other model structures with residual components [26] or gating mechanisms [27], [28] like mixture of experts (MoE) models [29], where the execution path can change dynamically at inference time. For these non-linear structures, STRONGHOLD either offloads all units/layers directly connected to a branch to the GPU working window (if possible), or delays the layer movement until it knows which layer will be computed to avoid GPU out-of-memory (OOM) errors. This can be further improved by leveraging techniques that pre-compute the activated layers [15] to proactively determine which layers to be moved to the GPU working window ahead of time. When loading a model, for each DNN layer, STRONGHOLD also computes the required storage size for the parameter tensors and the associated gradients and optimizer states. This storage size is then used to determine the GPU working window size during FP and BP.

During the first few iterations (5 by default) of model training (i.e., the warm-up phase), the STRONGHOLD runtime profiles the GPU computation time and the data transfer time of model states of each layer. It then uses this information to derive the working window size for later training iterations. At the warm-up phase, STRONGHOLD ensures that the chosen GPU working window size does not cause out-of-memory (OOM) errors by using the storage size information of the DNN layers to compute the right working window size. Despite the initial working window size may not lead to the optimal GPU memory use, the overhead is negligible since the profiling is only performed on the first few iterations. The dynamic working window size derived for later training iterations (see Section III-C) is designed to overlap the CPU-GPU data transfer with the GPU computation while minimizing the GPU memory consumption. We note that the computation performed in the warm-up phase also contribute to the final training outcome, so no computation cycle is wasted.

C. Dynamic GPU Offloading

As a working example, we use a simplified Transformer-based model shown in Figure 3a to illustrate the dynamic GPU working window mechanism of STRONGHOLD. In this subsection, we assume STRONGHOLD does not use secondary storage (which will be discussed later in Section III-G).

The STRONGHOLD runtime maintains a GPU working window with the layer-specific inputs, model parameters, and gradients (for BP). Within STRONGHOLD, the basic offloading unit under data and pipeline parallelism is an entire DNN layer. However, under tensor parallelism, this can be a sliced layer on the GPU defined by the user code. The working window essentially contains GPU buffers for the tensor operator implementations (kernels) and the data that the kernel operates on. CPU-GPU data movement is automatically handled by the STRONGHOLD runtime, which registers callback functions for each layer through the hooking mechanism provided by mainstream deep learning frameworks. STRONGHOLD supports activation checkpointing as long as the working window size is larger than the number of layers between two consecutive checkpoints. Conceptually, this mechanism resembles applying a sliding window to the DNN model along the FP or BP direction, described as follows.

FP stage. As shown in Figure 3b, before executing each layer in the working window, the pre_forward hook function is called to issue an asynchronous load operation to fetch the next layer right outside the current working window from the CPU RAM to the GPU memory (step 1). Next, the GPU performs FP computation as normal on the first layer in the current working window (step 2). At the end of the layer computation, the post_forward hook function invokes another asynchronous operation to move model parameters of the already computed layer back to the CPU RAM (step 3). The computation result will then be passed to the next layer in the working window. At the end of the GPU-CPU transfer, the GPU buffer used by a computed layer will be recycled by a newly fetched layer. After these steps, the working window moves toward the successive layer of the DNN, following the direction of FP. The asynchronous CPU-GPU data transfer takes place concurrently with the GPU computation and will not block the STRONGHOLD runtime.

BP stage. As depicted in Figure 3c, this step moves the working window along the BP direction. Before computing a layer in the working window, the pre_backward hook function invokes an asynchronous operation to fetch the parameters of the layer that is just outside the current working window in the BP direction (step 1). The pre_backward function also issues an asynchronous operation to move the model parameters (and gradients) of the last computed layer in the working window to the CPU (step 2), followed by a call to the optimizer to update the majority of the model parameters on the CPU (step 3). Finally, the GPU computes gradients for the second-last layer of the working window (step 1). Once again, the CPU-GPU data communications run concurrently with the GPU computation and the windows moves towards the BP direction.

D. Modeling Offloading Parameters

STRONGHOLD uses an analytical model to automatically find a suitable GPU working window size during FP and BP. The key here is to find the right window size where the asynchronous CPU-GPU data transfer can overlap with GPU computation to hide the data transfer latency, without oversubscribing the GPU memory.

Notations. We use the following notations to model CPU-GPU offloading. We use $s_{i_f}$ and $s_{i_b}$ to denote the memory consumption of layer $i$ during FP and BP respectively, and
\( t^i_{fp} \) and \( t^i_{bp} \) to denote the GPU computation time on layer \( i \) during FP and BP respectively, \( t^i_{2g} \) and \( t^i_{2c} \) to denote the CPU to GPU and GPU to CPU data transfer respectively for layer \( i \), and \( t_{async} \) and \( t_{opt_{gpu}} \) to denote the overhead on an asynchronous function call and one layer’s parameter update respectively. We let \( N = \sum_{k=0}^{m} S_k \) be the GPU working window size, where \( S_k \) is the model state (parameters and gradients) of layer \( k \) within the \( m \)-layer window.

**FP offloading.** We use the following formulation, \( P_1 \), to ensure layer fetching does not become a bottleneck of FP.

\[
P_1 : \min_{m} m \quad s.t. \quad \sum_{i=0}^{m} t^i_{fp} \geq t^i_{2g}, \tag{1a}
\]

\[
\sum_{i=0}^{m} s^i_{fp} + s^i_{fp} \leq S_{avail}, \tag{1b}
\]

\[
\sum_{i=0}^{m} t^i_{fp} \geq \sum_{i=0}^{m} t^i_{2g} + \sum_{i=0}^{m} t^i_{2c}, \tag{1c}
\]

where \( m \) is the number of layers in the GPU working window, layer \( j \) is the layer outside the current working window along the FP direction, \( s^m_{fp} \) is the buffer size required for fetching the layer outside the current working window (step 1 in Figure 3b), and \( S_{avail} \) is the available GPU memory (by excluding the runtime memory consumption). Here, terms (1b) and (1c) are hard constraints to ensure that the data transfer time is less than the layer computation time, and no GPU OOM will happen. The FP computation time for one layer is \( t^i_{fp} + t^i_{async} \). Term (1d) is a soft constraint to ensure that we can recycle the buffer of the computed layer (steps 2 and 3 in Figure 3b) to further reduce the GPU memory consumption.

**BP offloading.** Like FP, we use the following formula, \( P_2 \), to ensure gradient and parameter offloading does not become the bottleneck during BP:

\[
P_2 : \min_{m} m \quad s.t. \quad \sum_{i=0}^{m} t^i_{bp} \geq t^i_{2c}, \tag{2a}
\]

\[
\sum_{i=0}^{m} s^i_{bp} \leq S_{avail}, \tag{2b}
\]

\[
\sum_{i=0}^{m} t^i_{bp} \geq \sum_{i=0}^{m} t^i_{2g} + \sum_{i=0}^{m} t^i_{2c}, \tag{2c}
\]

\[
\sum_{i=0}^{m} t^i_{bp} + n * 2 t_{async} + \sum_{i=0}^{m} t^i_{opt_{gpu}} \leq \sum_{i=0}^{m} t^i_{fp} + \sum_{i=0}^{m} t^i_{bp} + \sum_{i=0}^{m} t^i_{opt_{gpu}}, \quad k \in [m, n]. \tag{3}
\]

To avoid introducing additional overhead during a training iteration, we need to ensure the computation time incurring by \( P_{1,2} \) during BP is not greater than the time for conventional training, i.e.,

\[
\sum_{i=0}^{m} t^i_{bp} + n * 2 t_{async} + \sum_{i=0}^{m} t^i_{opt_{gpu}} \leq \sum_{i=0}^{m} t^i_{fp} + \sum_{i=0}^{m} t^i_{bp} + \sum_{i=0}^{m} t^i_{opt_{gpu}}, \quad k \in [m, n]
\]

which gives us:

\[
5 n t_{async} \leq \sum_{i=m}^{n} t^i_{opt_{gpu}} \tag{4}
\]

When applying into a DNN where most of the layers are homogeneous with the same number of parameters, e.g., Transformer-based models, GPU computation time can be approximated as \( \sum_{i=m}^{n} t^i_{opt_{gpu}} \approx (n - m) t_{opt_{gpu}} \) and hence:

\[
5 n t_{async} \leq (n - m) t_{opt_{gpu}} \tag{5}
\]

Since the overhead of asynchronous function calls is largely constant regardless of the DNN model size, we can easily satisfy (5) with a deep (i.e., a large \( n \)) or a wide network (i.e., a large \( t_{opt_{gpu}} \) due to more parameters that a layer has).

**Determining the working window size.** For an \( n \)-layer DNN, \( P_{1,2} \) automatically finds a suitable working window size \( m \) that meets all \( P_1 \) and \( P_2 \) across all layers during a
training iteration. It is possible that there is not enough GPU memory to find an optimal $m$ to meet all the constraints. In this scenario, STRONG still uses the largest possible $m$ layers permitted by the available GPU memory to train a large DNN (that would not be possible using conventional training methods) but the training efficiency may be sub-optimal. By default, STRONG finds an available GPU buffer for $m$ layers. This strategy improves the GPU cache locality for Transformer-based models that have a large number of identical layer structures (and computation kernels). However, STRONG also supports having a fixed-size GPU buffer where the number of DNN layers stored can dynamically change, which can be turned on by users to improve GPU memory utilization for DNN models with a heterogeneous layer structure.

As a working example, Figure 4 shows the profiling data of one training iteration when applying STRONG to train a 4B model on a 32GB V100 GPU. The profiling results show that the CPU-directed offloading is largely overlapped by the GPU computation when criteria $P_1$ and $P_2$ set in our analytical models are met. In this case, the communication overhead can be hidden by the GPU computation, suggesting the effectiveness of our analytical model.

E. Offloading Optimization

STRONG utilizes the gRPC module of Ray [30] and concurrent library for communications among parallel CPU workers. As a result, STRONG can support concurrent and asynchronous parameter updates and data transfer. To this end, STRONG maintains a thread pool. All workers are initialized with the model code, and a worker remains idle until a task has been assigned to it through a callback function. By default, STRONG uses all available CPU cores, but the user can change this.

1) Concurrent parameter update: Unlike conventional training schemes (including ZeRO-Offload) that employ a single optimizer for parameter update, STRONG creates multiple optimizers during the model initialization stage. It then dispatches several optimizers to run as asynchronous actors to perform parameter updates on multiple layers simultaneously (step 3 in Figure 3c). This optimization leverages multiple CPU cores to process the parameter updates of multiple layers simultaneously, reducing the chance for the CPU becoming a bottleneck. As parameter updates are performed by the CPU, this process runs concurrently with the GPU computation during BP. By default, STRONG keeps the first few layers of the model (i.e., layers of the first working window) in the GPU memory. Since the last $m$ layers in BP (i.e., the first $m$ layers of the model) remain in the GPU working window before the start of FP, there is no GPU stall when computing BP of the last layers in Figure 3c.

2) Heterogeneous collective communications: For distributed training involved multiple computing devices, gradients communications are realized through collective communications operations like all-scatter and all-gather. With a native deep learning framework (e.g., PyTorch and Tensorflow), only one type of tensors (CPU or CUDA) can participate in collective communications at a time. STRONG lifts this restriction to support concurrent heterogeneous collective communications on CPU and CUDA tensors. This feature is essential for STRONG to support concurrent CPU and GPU processing. This is achieved by extending the low-level collective communication libraries, NVIDIA NCCL [31] and Gloo [32] for GPU and CPU communications, respectively.

3) Runtime memory management: During DNN training, many temporary tensors will be allocated and deallocated. Frequent device memory operations using the native CUDA memory (de)allocation API can result in expensive runtime due to explicit and implicit synchronizations. Frameworks like PyTorch and Tensorflow avoid this issue by reusing the previously allocated buffers through a software caching mechanism. For an $n$-layer DNN, where each layer has $k$ tensors, such a caching mechanism incurs up to $n \times k$ GPU memory allocation operations. After the first training iteration, these $n \times k$ GPU buffers are then reserved by the runtime, which can then be reused for future training iterations. This strategy is ill-suited for our scenarios when the model is too large to be fit into the GPU memory (i.e., the $n \times k$ buffers are beyond the GPU memory capacity). STRONG addresses this issue by employing a user-level software memory management scheme on the CPU and GPU. For an $m$-layer GPU working window, STRONG only needs to incur a one-off $m \times k$ CUDA memory operation at the warm-up stage. Since the working window size, $m$, is smaller than the number of layers (i.e., $m < n$) of the DNN, STRONG reduces the GPU memory footprint while incurring fewer memory allocation operations than existing caching mechanisms.

Specifically, when loading the DNN, STRONG allocates pinned memory on the CPU for each DNN layer. The pinned (or page-locked) memory permits STRONG to asynchronously transfer the CPU data to the GPU using an idle CUDA stream so that the GPU will not be blocked during data transfer. At the same time, STRONG also reserves GPU buffers for layers of the first working window. The reserved buffers will be managed by STRONG in future training iterations. The reserved GPU buffer may grow (but not shrink) if larger buffers are needed once STRONG has determined the working window size after the warm-up stage. When prefetching a layer from the CPU memory to the
GPU (e.g., step 1 in Figure 3b), StrongHold first allocates a free GPU buffer from the reserved GPU memory in a round-robin manner. It then copies the corresponding data content to the corresponding GPU tensor (e.g., through the PyTorch tensor.copy_( ) API). Similarly, when offloading a layer from the GPU to the CPU memory, StrongHold copies the data property back to the corresponding CPU buffer. It then returns the GPU buffer to the StrongHold managed GPU buffer queue. Whenever StrongHold requests or releases device memory, the StrongHold runtime always reuses the reserved GPU memory by overwriting the in-place memory management methods of the layer implementation.

F. Cross-server Communication Optimization

Another benefit of StrongHold is that it can eliminate the cross-server communications introduced by traditional model parallelism in certain cases. For example, if a model cannot fit into the GPU memory under a traditional training method, model parallelism is typically adopted to break the model layers (and their parameters) across multiple GPUs. In contrast, if the same model can fit into the same GPU under StrongHold, we can then use the additional GPUs to run data parallelism training without incurring the synchronization and communication overhead of model parallelism. The reduction of cross-server communications when converting model parallelism to data parallelism for Transformer-based models can be estimated as follows. The communication volume for an n-layer Transformer model is $V_{dp} = (w - 1)w \times (12 \times n \times hd^2 + hd \times vs)$ for w-way data parallelism, and $V_{mp} = (w - 1)w \times n \times bs \times seq \times hd$ for w-way model parallelism. Here, hd, bs, seq, and vs are the the hidden size, batch size, sequence length and vocabulary size, respectively. Furthermore, we obtain the constant number 12 summing $4 \times hd^2$ for attention and $2 \times 4 \times hd^2$ in the feed-forward network in one Transformer block. By converting w-way model parallelism to w-way data parallelism, StrongHold reduces the communication volume by $\frac{V_{mp}}{V_{dp}}$.

Using a typical training setup where the training sentence sequence length is set to 1024 ($seq = 1024$) and vocabulary size is set to $30k$ ($vs = 30K$), we can simplify $\frac{V_{mp}}{V_{dp}}$ as $\frac{bs}{3 \times hd/256 + 30/\pi}$. Let $k = \frac{1}{3 \times hd/256 + 30/\pi}$, we now have $V_{mp}/V_{dp} = k \times bs$. Here, the saving in cross-node communication depends on $n, hd, w, and bs$. By increasing the trainable model size, StrongHold allows one to use a w-way data parallelism to replace the traditional w-way model parallelism. For a 20B model with a typical $bs = 16, n = 50, hd = 4K$, StrongHold halves the communication traffics by comparing to model parallelism (see also Section VI-D2).

G. Utilizing Secondary Storage

Like ZeRO-Infinity, StrongHold provides an option to use NVMe SSDs to further increase the trainable model size. This is achieved by memory-mapping a swap file on the secondary storage to the CPU memory space and using the read/write library to optimize asynchronous bulk read/write requests between the CPU and the device. The support for asynchrony allows StrongHold to overlap the I/O requests with CPU-GPU communication or computation. Since the I/O bandwidth between the CPU and an NVMe SSD (up to 7GB/s for PCIe 4.0) is an order of magnitude slower than the CPU-GPU bandwidth, we do not expect the user to train a large DNN from scratch with this option. It is also not advised to train a large model with this strategy because frequent random reads and writes can increase the chance of NVMe disk failure [33]. However, this option can be useful in fine-tuning a pre-trained model with fewer training iterations.

IV. StrongHold-enabled Optimizations

By reducing the GPU memory footprint, StrongHold not only permits training larger DNNs but also opens up new optimization opportunities. For example, StrongHold allows training a large DNN that previously would otherwise only possible using model parallelism to split the DNN layers across GPU servers. With StrongHold, the distributed GPU servers can be used to run data-parallel training workers, where each computing node holds the entire model parameters. This can massively reduce the communication and synchronization overhead imposed by model parallelism (see also Section VI-D2). Another use case is to support teacher-student based knowledge distillation [21] by supporting large DNN inference on a single GPU (Section VI-D3). The third interesting optimization is to support data parallel training within a single GPU by utilizing multiple CUDA streams (see also Section II-B) to use the GPU parallelism to improve the training throughput, which was not attempted in accelerating DNN training before. This is described in the next subsection.

A. Multi-streamed GPU Execution

As depicted in Figure 5, StrongHold enables the use of multiple CUDA streams to accelerate training if there is enough GPU memory to store the gradients and model inputs for at least two training workers.

Conceptually, this is achieved through applying data parallelism within a single GPU by partitioning the training batch into mini-batches. To this end, StrongHold introduces `executors’ within its runtime to manage the GPU kernel execution context across training workers. An executor is a process of the StrongHold runtime that manages the working window for a training worker. The executor dispatches kernels to run on a CUDA stream. Note that only one copy of the model...
parameters and kernel code is stored in the GPU memory, despite that there may be more than one training worker and working window on the GPU. Furthermore, STRONGHOLD’s strategy of dividing a larger batch into micro-batches to be processed by concurrently running does not affect the model consistency since parameter update takes after the entire batch has been processed, i.e., data-parallel training.

During runtime, we bind each executor to an available CUDA stream and allocate dedicated buffers on the CPU to allow the executor to manage the host-device communications for the corresponding GPU working window. Each executor runs as a multi-staged pipeline to process a mini-batch, and each pipeline stage of an executor can execute a kernel for FP, BP, computation, communication or another user-defined kernel. As depicted in Figure 5, by mapping different concurrent executors onto multiple CUDA streams, we can allow different processing pipelines to run in parallel on a single GPU to improve the throughput. STRONGHOLD uses an all-reduce operation to synchronize the gradients among parallel training workers before performing parameter updates, similar to data parallel training across multiple GPUs (but STRONGHOLD has the advantage of only keeping one copy of model parameters). The number of concurrent streams used is determined during the warm-up phase, where STRONGHOLD computes the GPU memory consumption of the GPU working window size to determine how many CUDA streams to use so that multi-streamed execution does not cause GPU OOM.

STRONGHOLD essentially creates a lightweight parallel execution environment for individual executors within a single user program. This is different from existing GPU virtualization schemes such as NVIDIA MIG (Multi-Instance GPU) [34], which aim to provide an isolated execution environment for different user programs running on the same GPU. MIG is ill-suited for our purpose because CUDA streams or processes in different virtualized environments cannot directly communicate with each other. To implement our idea within MIG would also require each GPU process holds a copy of model parameters, resulting in multiple model parameters being stored on a single GPU, increasing the GPU memory pressure. Our lightweight approach can avoid such pitfalls by only storing one copy of the model parameters on the GPU and enabling CUDA streams to directly communicate with each other through the GPU hardware communication scheme. Since our goal is to accelerate a single user program, we do not need a heavy and strongly isolated execution environment.

V. Evaluation Setup

A. Evaluation Platforms

We evaluate STRONGHOLD on two hardware platforms. Our main evaluation platform is a V100 GPU server with one 32GB NVIDIA Tesla V100 GPU, 2x 24-core Intel Xeon Platinum 8163 CPU at 2.50GHz and 755GB of DDR4 RAM. We also evaluate STRONGHOLD on an 8-node A10 GPU cluster where servers are connected through a 800 Gbps network. Each computing node of the cluster has 2x 64-core Intel Xeon Platinum 8369B CPUs at 2.90 GHz with 1TB of DDR4 RAM, and a 24GB GPUDirect-RDMA-enabled A10 GPU based on the latest Ampere architecture. The platforms run Ubuntu 20.04 operating system with Linux kernel 4.19.91. We use CUDA 11.2 and PyTorch 1.10.2.

B. Workloads

We evaluate STRONGHOLD on GPT-like Transformer-based models [35], [3] – the current de facto approach to building large-scale models [2], [3]. Following the evaluation setup of ZeRO-Offload [11], we vary the hidden dimension of a layer to increase the model width and the number of layers to scale the model depth. Table I lists the model parameters. For each model, we consider a batch size of 2, 4, 8, and 16 per GPU. Note that scaling the depth alone is often not sufficient because it would make training harder to converge [36]. Unless stated, we use training hyperparameters like weight decay and learning rate from [23], [11].

C. Competing Baselines

We compare STRONGHOLD against the following billion-scale model training solutions:

Megatron-LM [37] is NVIDIA’s optimizing library for Transformer-based models. We use Megatron-LM v2.6 as a reference model for the training throughput and trainable model size.

L2L [18] keeps one Transformer layer in the GPU at a time, by sequentially offloading parameters between the GPU and CPU memory. Since L2L still stores the optimizer states on the GPU, it is largely limited by the GPU memory.

ZeRO-Offload [11] statically stores the model states in GPU memory and optimizer states in the CPU RAM. It also utilizes the CPU computation cycle to update the model parameters through a CPU-tuned optimizer.

ZeRO-Infinity [19] utilizes GPU, CPU and NVMe memory. By default, we compare STRONGHOLD against ZeRO-Infinity with CPU RAM instead of using NVMe due to the expensive I/O overhead. In Section VI-C, we compare STRONGHOLD with ZeRO-Infinity when using a 2TB PCIe4.0 NVMe SSD.

ZeRO-2 and ZeRO-3 [9] partition the model states across distributed machines but with a data parallel strategy. We compare STRONGHOLD to the ZeRO-2 and -3 solutions (which are the
core of the DeepSpeed [38] framework) for distributed training in Section VI-D2.

D. Performance Report

We consider trainable model size and throughput in the evaluation. We report the trainable model size by counting the number of model parameters (using FP32 representation) that can be trained without incurring GPU OOM, and report throughput by measuring the number of training samples processed per second. We run each test case 10 times on unloaded servers to measure the metrics, and then report the geometric mean across different runs. In our evaluation, we use layer-wised activation checkpointing [39] that is widely used in large-scale training. We note that the profiling overhead at the warm-up phase accounts for less than 0.5% of the total training time on our evaluation setup. This overhead will be much smaller (and negligible) in a real-life training scenario with a higher number of training iterations. We have included this overhead when computing the throughput of StrongHold. Finally, the throughput and efficiency variance across runs is small, less than 3%.

VI. EXPERIMENTAL RESULTS

In this section, we first show that StrongHold outperforms all the offloading baselines by enabling the training of a 39.5B model on a V100 GPU and an 82.1B model across 8 distributed A10 GPUs using model parallelism (Section VI-A) and giving a higher training throughput (Section VI-B). We then perform further analysis on StrongHold (Section VI-C) - including the use of NVMe - before showcasing the new optimizations enabled by StrongHold (Section VI-D).

A. Trainable Model Size

Figure 6 compares the trainable model size when using CPU RAM only. The min-max bar on the diagram gives the range of measurements for different DNN configurations when varying the hidden dimension and model depth.

1) Single V100 GPU: Figure 6a gives the largest trainable model on a single 32GB V100 GPU. Megatron-LM allows one to train a model with up to 1.7B parameters on a V100 GPU before incurring GPU OOM. L2L and Zero-Offload can expand the trainable size by 3.5x over Megatron-LM, to support a model with around 6B of parameters through CPU-GPU offloading. Moreover, the fine-grained parameter partitions in ZeRO-Infinity (with CPU RAM only) can support a 20.6B model. StrongHold outperforms all baselines, supporting model training with 39.5B parameters and giving comparable training efficiency compared to other offloading scheme. StrongHold is limited by the GPU and CPU memory and the offloading unit size. Therefore, it can support a larger model with larger CPU RAM and GPU memory (or reducing the size of the offloading unit through tensor parallelism). The trainable model size of StrongHold translates to a 6.5x improvement over L2L and Zero-Offload, and an 1.9x improvement over Zero-Infinity. ZeRO-Infinity requires moving the parameters, gradients, and optimizer states to the GPU for runtime model refactoring. This operation requires making a copy of the refactored model parameters, incurring extra GPU memory overhead. StrongHold does not have this overhead, leading to a larger trainable model size. Furthermore, while Zero-Infinity could assure large DNN training, as we will show in Section VI-B, it comes at the cost of poor training efficiency.

2) Distributed GPUs: Figure 6b shows the largest trainable model size across 8 distributed A10 GPU servers using model parallelism. All offloading approaches benefit from additional GPU resources. However, L2L and Zero-Offload give limited improvement on the trainable model size as they are largely constrained by a single GPU memory. By partitioning the model states across heterogeneous devices, ZeRO-Infinity and StrongHold demonstrate stronger scalability by scaling the trainable model size to 56.9B and 82.1B parameters respectively, with StrongHold supports the largest model. In all test cases, StrongHold gives a nearly 100% GPU utilization, with 80% utilization of the theoretical peak bandwidth of the CPU-GPU PCIe or communication network.

B. Training Throughput

Figure 7 compares the training throughput on the largest trainable model size supported by each baseline. StrongHold runs the same model of its counterpart. StrongHold outperforms all baselines (including Megatron-LM thanks to StrongHold’s multi-streamed optimization), achieving 42 ~ 57% hardware performance by delivering 6~9 TFlops on a V100 GPU. The TFLOPS given by StrongHold far exceeds the one delivered by L2L (1.88).
ZeRO-Offload (0.59) and ZeRO-Infinity (0.53). It improves the training throughput by at least 1.1x (up to 3.7x) by better overlapping the CPU-GPU communication. Figure 8a shows the throughput obtained when a common 1.7B model (the largest trainable model supported by Megatron-LM) on a V100 GPU. For this case, L2L delivers only 22.2% of the Megatron-LM throughput because it simply serializes computation with data transfer for each DNN layer. ZeRO-Offload and ZeRO-Infinity achieve less than 57% of the Megatron-LM training efficiency because a large portion of the CPU-GPU data transfer and computation cannot overlap due to their CPU optimizer implementation. **STRONGHOLD** is the only offloading solution that gives an improvement over Megatron-LM. The results show that **STRONGHOLD** can scale up the trainable model size and accelerate DNN training.

### C. Further Analysis

1) **Training efficiency:** Figure 8b shows that **STRONGHOLD** delivers nearly linear training efficiency on a single V100 GPU, using a 1.7B model as the starting point. The training efficiency is a *lower-is-better* metric, measured by the averaged time in performing one training iteration. **STRONGHOLD**’s performance is on par with a perfect linear scaling projection, albeit there are some fluctuations in the scaling trend due to the impact of the GPU working window size on the GPU cache performance. Using the same resources, **STRONGHOLD** can train 25x, 4x, and 20x bigger models in a single-GPU-single-node, multiple-GPU-single-node and multiple-GPU-multiple-node environment, respectively; and it achieves these without significantly compromising the training efficiency. Therefore, **STRONGHOLD** can reduce the number of GPUs required by at least 4x compared to the traditional training method. The resource-saving can also be used to increase data parallelism by 4x to speed up the training time. Based on this scalability projection, we estimate that **STRONGHOLD** can complete the training of 175B GPT-3 using a quarter of the GPUs with a similar training time, or it can speed up the training process by 4x when using the same number of GPUs compared to the conventional distributed training method. This capability can reduce the cost by using either smaller-scale GPU resources or fewer GPU hours, making the training of large DNN more accessible and affordable.

2) **Impact of working window size:** Figure 9 shows how the GPU working window size affects the throughput by running a 1.7B and a 39.5B model on a V100 GPU. In our cases, initially, a larger window can better overlap GPU computation with data transfer, which leads to a higher training throughput. However, the improvement reaches a plateau with a window size of 8. Using a window size greater than 8 does not justify the gain in throughput but will increase the GPU memory pressure. Using the analytical method described in Section III-D, **STRONGHOLD** automatically determines to use a window size of 8 for this model.

3) **Using NVMe:** Figure 10 shows the throughput improvement over ZeRO-Infinity when NVMe is used to scale up the model size on the V100 GPU server. When using NVMe, **STRONGHOLD** and Zero-Infinity can support the training of a model with half trillions of parameters on our V100 GPU server. Compared to Zero-Infinity, **STRONGHOLD** can also better overlap the disk I/O requests with GPU computation, improving the throughput by over 8x.

### D. **STRONGHOLD**-enabled Optimizations

1) **Multi-streamed optimization:** Figure 11 shows performance improvement given by **STRONGHOLD** over Megatron-LM when varying the batch size. By only keeping part of the DNN layers and gradients in the GPU, **STRONGHOLD** reduces the GPU memory footprint by 60%. The reduced memory footprint permits **STRONGHOLD** to use multiple CUDA streams speedup the training process, leading to at least 1.7x (up to 2.1x) speedup over Megatron-LM.

2) **Accelerating distributed training:** Figure 12 compares **STRONGHOLD** against ZeRO-2 and ZeRO-3 in a distributed training setup on the A10 GPU cluster. ZeRO-2 partitions the optimizer states and gradients across parallel processes running on multiple servers while ZeRO-3 partitions the model parameters on top of ZeRO-2 across GPU servers [9]. We apply all approaches to the largest model (3B parameters) that can be supported by ZeRO-2 with a batch size of 1. ZeRO-2
and ZeRO-3 have to partition the model/optimizer states across GPUs due to the GPU memory restrictions, but they introduce extra communication overhead across GPUs and serve nodes. By reducing the GPU consumption, StrongHold does not need to partition the model across GPU servers. Instead, it can run the entire model on a single server to exploit data parallelism across GPU servers. As a result, StrongHold reduces the cross-server communications, leading to over 2.6x throughput improvement over ZeRO. For example, StrongHold reduces the cross-server communication by around 50% on a 20B model (see also Section III-F). This experiment demonstrates another advantage of StrongHold in speeding up distributed deep learning training.

3) Knowledge distillation: StrongHold can also support knowledge distillation [21]. This strategy is widely used to accelerate DNN inference by using a trained large model to guide the training of a smaller but faster DNN. Under this setting, the large DNN only needs to perform FP on training samples to provide layer-wise activations to guide the student model training. Inferencing frameworks like TensorRT [40] are not suitable for this scenario because they do not produce activations of intermediate layers. As can be seen from Figure 13, StrongHold can effectively support large DNN inference for knowledge distillation. Note that as StrongHold only needs to support FP in this scenario, it can support a larger model than when it is used for training that is involved with both FP and BP. It gives similar performance for small DNN inference compared to PyTorch but delivers linear scalability for large DNNs where PyTorch give an OOM error.

4) Optimization breakdown: Figure 14 reports the performance improvement on the V100 GPU platform when turning on a single optimization to a baseline offloading scheme without optimization. By utilizing the multi-core CPU, concurrent parameter update with heterogeneous collective communication techniques (Sections III-E1 and III-E2) give 1.5x throughput improvement. By minimizing the GPU tensor allocation overhead, our memory management optimization (Section III-E3) alone gives a 2.2x throughput improvement. Similarly, by launching multiple kernels concurrently, our multi-streamed optimization (see Section IV-A) offers up to 2x improvement.

VII. RELATED WORK

Data and model parallelisms are two dominant parallelization techniques for large DNN training [41], [38], [42]. The former partitions the training data across multiple GPUs [5], [43], and the latter splits the model layers vertically and then distributes different layers onto GPUs to reduce the memory pressure of the model states on a single GPU [6]. ZeRO [9] partitions the training batch across multiple GPUs, similar to data parallelism, but it further splits the model states across GPUs and uses collective operations to gather the required model parameters. Parallelization can also be achieved by horizontally partitioning a tensor operator across multiple GPUs [23], [44]. StrongHold complements the existing data-parallel approaches by using multiple GPU SMs to achieve fine-grained data parallelism while just keeping one copy of the model parameters across parallel training workers.

A recent line of work adds pipelines into model parallelism by partitioning model layers into parallel stages [7], [8], [45], [46], [47], [48], [49]. In this way, each training batch is divided into micro-batches to be processed by pipeline stages across computing devices. Although StrongHold also follows pipelining, tasks on a StrongHold pipeline are finer-grained, where the model layers stored in a GPU memory can change dynamically during execution.

All the aforementioned strategies utilize the aggregated GPU memory of multiple GPUs to meet the memory requirement of large DNNs. StrongHold is among the recent attempts in scaling up the trainable model size on a single
GPU. This line of research includes activation check-pointing methods, which trade computation for GPU memory saving [39]. This strategy drops the activations after FP and recomputing them from checkpoints during BP. Unfortunately, this comes as the cost of huge overhead for large DNNs because a large number of activations to be recomputed. Compression techniques, such as using low or mixed precision representations for model states, can reduce the memory footprint [12]. However, compression techniques can reduce the training accuracy and slowdown model convergence.

Our work falls under a third approach to utilize external memory like the CPU RAM and NVMe to expand the memory capacity during training [50], [13], [51], [15], [52], [16]. ZeRO-Offload [11] and ZeRO-Infinity [19] are the most closely related work. ZeRO-Offload uses the CPU memory to store gradients and optimizer states. Unlike STRONGOLD, ZeRO-Offload stores the entire model parameters in the GPU memory. As such, ZeRO-Offload is limited by the GPU memory, rather than the external memory like STRONGOLD. ZeRO-Infinity extends ZeRO-Offload through fine-grained model state partitioning and utilizing the secondary storage. When using NVMe, ZeRO-Infinity can greatly increase the trainable model size, but this comes at the cost of prohibitively long training time. By carefully overlapping computation and communication, STRONGOLD significantly improves the training efficiency over ZeRO-Infinity. L2L [18] is a Transformer-specific offloading scheme. It keeps an encoder layer in the GPU memory, by synchronously moving the model parameters the CPU memory to mimic layer-by-layer computation. Unlike STRONGOLD, L2L requires model refactoring and offers poor efficiency due to extensive communication overhead and frequent GPU stalls. The M6 model [53] keeps a fixed number of layers on GPUs, which is specific to the current model, and requires code refactoring.

The work presented in [54] combines rematerialization to trade memory for computation time and offloading to trade memory for data movement. It employs a dynamic programming heuristic to determine the optimal offloading sequence. AxoNN [55] exploits asynchronous and message-driven execution for scheduling parallel training workers to improve GPU utilization and system throughput. Varuna [56] leverages low-priority virtual machines and pipelining to enable low-cost model training over commodity networking. These techniques are complementary to STRONGOLD.

Dorylus exploits the workload characteristics of graph neural networks (GNNs) to use parallel serverless CPU threads for model training [57]. Dorylus and other memory [58] or computation [59] optimization techniques can be used in combination with STRONGOLD to utilize low-cost CPU threads to train GNNs. Furthermore, STRONGOLD can also be used together with asynchronous training [60] to further reduce the waiting time across training epochs, but care must be taken to avoid slowing down model convergence [61].

VIII. CONCLUSION

We have presented STRONGOLD, a new offloading framework to lower the GPU memory consumption for training billion-scale DNNs. STRONGOLD utilizes heterogeneous resources to scale the trainable model size. Compared to existing offloading solutions, STRONGOLD reduces the GPU memory footprint with lower computation overhead. It achieves this by maintaining a compact GPU working window and using data prefetching techniques to overlap data transfer and GPU computation. By reducing the GPU memory consumption, we demonstrate that STRONGOLD enables new optimization to utilize the GPU hardware parallelism to improve performance. We show that STRONGOLD allows the training of a larger DNN model with better training efficiency than the state-of-the-art offloading techniques. Specifically, it enables the training of a DNN with 39.5 billion parameters on a single V100 GPU without changing user code and supports faster model training in a distributed GPU environment.

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