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# Weight Offloading Strategies for Training Large DNN Models

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#### Abstract

The limited memory of GPUs induces serious problems in the training phase of deep neural networks (DNNs). Indeed, with the recent tremendous increase in the size of DNN models, which can now routinely include hundreds of billions or even trillions of parameters, it is impossible to store these models in the memory of a GPU and several strategies have been devised to solve this problem. In this paper, we analyze in detail the strategy that consists in offloading the weights of some model layers from the GPU to the CPU when they are not used. Since the PCI bus bandwidth between the GPU and the CPU is limited, it is crucial to know which layers should be transferred (offloaded and prefetched) and when. We prove that this problem is in general NP-Complete in the strong sense and we propose a lower bound formulation in the form of an Integer Linear Program (ILP). We propose heuristics to select the layers to offload and to build the schedule of data transfers. We show that this approach allows to build near-optimal weight offloading strategies on realistic size DNNs and architectures. Keywords: Training of DNNs, Scheduling, Offloading Strategies

### 1 Introduction

Training of Deep Neural Networks (DNNs) is a computationally and memory intensive operation. From a computational point of view, there are many parallel strategies that allow to distribute the computation over several resources. These

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strategies exploit the possibility to distribute the data and treat it over several resources (data parallelism) or the DNN itself (model parallelism) or to partition the data tensors along other dimensions (tensor parallelism, filter parallelism,...).

However, contrary to the case of typical HPC applications, such as linear algebra kernels, the nature of the dependencies between the tasks prevents in general to distribute among participating resources the memory necessary to store the intermediate produced data (activations), the model parameters and the optimizer states. For example, in the data parallel approach [28, 6], the same model is replicated on all participating nodes even if the activations are actually distributed. In the model parallel approach [26, 16, 17], on the contrary, the weights are well distributed between the different resources, but to be efficient, the model parallel approach must be pipelined like in Pipedream [16, 3]. In turn, pipelined strategies require to store several models: the number of models to store, which could be large in Pipedream [16], has been reduced to two in [17] or to a constant number using a different approach in [14]. Pipedream strategies also require to significantly increase the amount of activations to store, as shown in [3].

In practice, memory needs are a major reason for the renewal of GPUs and being able to decrease memory requirements can therefore help to limit the carbon impact, since in data centers, the carbon impact associated with the production of computational resources is often more significant than the impact associated with energy consumption [9] during the exploitation phase. These memory requirements come from both the data and the models. Indeed, high dimensional data have a direct impact (usually proportional) on the size of the activations. In addition, the use of deeper models with more parameters has in general an impact on both the size of the model and the number of the activations. Obviously, high dimensional data and deeper neural networks have in general a positive impact on the accuracy, so that it is crucial in practice to design strategies limiting the memory size required on a computational node without changing the model or the data and thus the performance of the model.

Historically, minimizing the memory consumption induced by activations was the first active research axis, because in vision networks, the memory consumption related to activations was the most significant. Strategies based on rematerialization [8, 12, 10] and offloading [25, 13, 1] of activations have been proposed, as well as the combination of these two approaches [2]. Re-materialization consists in deleting some activations from memory during the forward phase and recomputing them when needed during the backward phase, which corresponds to trading memory for computation. Offloading consists in offloading activations from the (small) GPU memory to the (large) CPU memory during the forward phase, and to prefetch them back during the backward phase.

Recently, minimizing memory consumption induced by model size and optimizer states has received more attention, especially due to the explosion of model sizes. For example, in Natural Language Processing (NLP), it is not uncommon to see models with billions parameters [7, 20, 5]. A typical example of such large models is GPT-3 [5], which contains 175 billion parameters, which is about 650 GB using 32-bit floating point numbers for the weights, not counting activations and optimizer states. As there is no GPU with such a memory size and as model parallelism increases the cost of storing activations as soon as more than a few tens of nodes are used [3], it is therefore necessary to rely on strategies to save the memory induced by model parameters.

In the literature, this problem has been addressed in several papers [21, 19, 15]. In these works, the objective is to train the largest possible model, thus to keep in the GPU memory, for each computational task related to a given layer (either forward or backward), only the data that is absolutely necessary for this task. These very aggressive optimization approaches typically lead to keeping all the weights (except those currently used) and all the states of the optimizer in the memory of the CPU, and to transfer the weights (when they are needed) from the CPU to the GPU and the gradients (as soon as they are computed) from the GPU to the CPU. This approach naturally leads to a very intensive use of the PCI bus, whose bandwidth can then become the limiting factor [15]. In the present paper, we consider the problem from an optimization point of view, focusing on the case where the training is performed on a single GPU. More precisely, we try to solve the following problem: given a set of layers defined by their sizes and processing times (forward and backward), given a PCI bus bandwidth and given a memory size, what is the strategy that minimizes the training execution time? Determining such a strategy requires solving both a selection problem (which layers should be transferred) and a scheduling problem (at what time should the transfers to and from the GPU be performed).

The paper is organized as follows. After a review of related works limited to weight offloading strategies (a more complete survey on memory saving techniques can be found in Chapter 2 of [23]) in Section 2, we present the model in Section 3 and complexity results in Section 4. Algorithms to solve the selection problem and the scheduling problems are presented in Section 5. To evaluate the quality of the proposed strategies, a lower bound based on Integer Linear Programming (ILP) is established in Section 6 and we rely on a set of simulations for transformer-based models in Section 7. Concluding remarks and perspectives are given in Section 8.

#### 2 Related works

The methods either targeting the offloading of activations [22, 25, 1, 2], or of any tensor [13, 27, 11]. can be adjusted (though heuristically) to deal with weights in the similar way as activations. The latter group can be directly applied to weights. In [13, 11], the tensors that will not be needed for the longest time are preferred candidates for offloading. Similarly to our approach, AutoSwap [27] relies on priority scores for choosing the best candidates for offloading, though it is limited to the case of only one memory peak, which is enough for activation offloading, but not for weigths. The schedule of transfers is performed in a similar way for all above methods: offloading is performed as soon as possible and prefetching as late as possible, trying to avoid the idle time before the operation requiring the prefetched data. Despite their universality,



Figure 1: Data dependencies induced by the training phase of Sequential DNNs

these methods have a few disadvantages: in particular, they do not fully consider benefits of weight offloading during the backward phase contrary to the present paper.

Several recent works focus on offloading strategies of DNN weights during the training phase to limit memory consumption. The first strategy is proposed in [19] which presents the L2L (Layer-to-Layer) algorithm that we use as a basis for comparison in the Section 7. For a graph typically consisting of a sequence of encoders, a synchronous parameter-server, stored on the CPU, allows to keep a copy of the model weights and to keep on the GPU only the encoder that is being used. A strategy allowing to adapt the size of the mini-batch is then proposed, which allows to use the biggest possible mini-batch given this systematic strategy of layer offloading. Closed-form formulas allowing to estimate the training time (for all identical layers) are proposed. This strategy was then refined in ZeROoffload [21].

Recently, in [15], the authors observed that approaches such as L2L [19] or Zero-Offload [21] generate, despite the optimizations proposed in [21], a very important data traffic on the PCI bus between the CPU and GPU. As its bandwidth is limited, it can become the bottleneck and significantly slow down the whole training process. For this purpose, it is proposed to offload only some layers, the layers from de first half being arbitrarily chosen in [15]. Our contribution in the present paper is based on the same idea, with a formalization of the optimization problem which consists in determining which layers to offload and when to do it. For this, we rely on the simplified model of [19] and we leave for future work the consideration of the optimizer states performed in [21].

#### 3 Model

In this paper, we focus on the case of a DNN that can be written as a sequence of layers (in practice, each layer may consist of arbitrarily complex computations). DNNs are trained using forward-backward propagation that is used to compute gradients of the loss (error of the model) with respect to weights, which are later used by optimization algorithms such as Stochastic Gradient Descent. One elementary iteration of this algorithm is described in Figure 1: it consists in a traversal by a mini-batch (a set of training samples, *e.g.* images, text sequences) of the different layers of the DNN, to compute the loss and then to backpropagate

the gradients to update the network weights. We do not go into the details of the computations that can be found in [4], but it is important for us to study the dependencies and memory requirements at each step. In practice, the *loss* related operation takes very little time and does not incur additional memory costs. For simplicity, in the rest of the paper we neglect this operation and consider that  $B_L$  comes right after  $F_L$ .

**Memory usage of operations** The standard practice is to compute all gradients during the forward-backward propagation, and then update all the weights in an *optimization* step. This means that the weight  $w_i$  of a given layer is used in three occasions during a training step: forward of layer i ( $F_i$ ), backward of layer i ( $B_i$ ), and for the update of  $w_i$  in the optimization step. We propose to perform the update of  $w_i$  during backpropagation, as soon as its gradient is computed, which obviously saves memory.

As a result, each operation  $F_i$  takes as input  $a^{i-1}$  and produces  $a^i$ , which will be used by  $F_{i+1}$  and  $B_{i+1}$ . Each operation  $B_i$  takes as input  $a^{i-1}$  and  $\delta^i$  and produces  $\delta^{i-1}$ , while updating weights  $w_i$  of layer *i* from the following sequence of operations that are considered atomic:

- gradient  $\delta w_i$  is computed from  $a^{i-1}$  and  $\delta^i$ ; once this operation is performed  $a^{i-1}$  can be deleted from memory, but  $w_i$  must remain.
- $\delta^{i-1}$  is computed from  $\delta^i$  and  $w_i$ ; once this operation is performed  $\delta^i$  can be removed from the memory, but  $w_i$  and  $\delta w_i$  must remain in memory.
- new  $w_i$  is computed from old  $w_i$  and  $\delta w_i$ ; then  $\delta w_i$  can be removed from memory (and  $w_i$  can be offloaded), but  $\delta^{i-1}$  must remain in memory to be passed to  $B_{i-1}$ .

In this paper, we assume that activations  $a^i$  are kept in memory from  $F_i$ until  $B_{i+1}$ , so that when computing  $B_i$ , all activations  $|a^j|$ ,  $0 \le j \le i-1$  must reside in memory. We further assume, as in the literature [4], that  $|\delta^i| = |a^i|$ and  $|w_i| = |\delta w_i|$ .

During  $B_i$ , the memory peak is reached when  $\delta^{i-1}$  is computed. We denote  $M_{F_i}$  and  $M_{B_i}$  the memory occupation during  $F_i$  and  $B_i$  (without counting the weights  $w_j$  related to other layers  $j \neq i$ , that can potentially be offloaded and no longer reside in memory). These values are given by:

$$M_{F_i} = |w_i| + \sum_{k \le i} |a^k|$$
 and  $M_{B_i} = 2|w_i| + \sum_{k \le i} |a^k|.$  (1)

Weight offloading We focus on the possibility to offload the weights  $(w_i)$  from the GPU memory to the CPU memory. We assume that there is a fullduplex communication link between the GPU and the CPU with bandwidth  $\beta$ , which means that we can at the same time offload a layer from the GPU and prefetch another one from the CPU. In any case, the communication buffers on



Figure 2: Instance used in the NP-completeness proof of Theorem 1. Node size represents operation times:  $\circ$  for  $u_i = 0$ ,  $\bigcirc$  for  $u_i = 1$ ; the size of the weights is represented by node thickness and written above.

the GPU are reserved from the beginning of a prefetch operation and until the end of an offload operation.

Given that  $w_i$  is only needed for  $F_i$  and  $B_i$ , we can save memory by (a) offloading  $w_i$  after  $F_i$  and prefetch it before  $B_i$ ; or (b) offloading  $w_i$  after  $B_i$  and prefetch it before  $F_i$ . In the following, we denote by offloading a weight  $w_i$  the action of sending  $w_i$  to the memory of the CPU, and deleting it right after. By contrast, the *uploading* operation consists in sending  $w_i$  to the CPU without deleting it on the GPU afterwards.

The problem comes however with an additional complication: only  $B_i$  modifies the value of  $w_i$ . This means that it can be beneficial to upload  $w_i$  in advance before the  $F_i$  operation (for example to benefit from an under-utilized communication link), and to delete it only after the  $F_i$  operation. It also means that between two  $B_i$  operations (on two successive batches), if  $w_i$  is offloaded after  $B_i$ , it is useless to offload it again after  $F_i$ , since that would mean rewriting the same value of  $w_i$  in the CPU memory. Therefore, it is possible that one offload operation corresponds to two prefetch operations, leading to what we call discounted communications. To the best of our knowledge, this discount possibility is not considered in other related works, and we show in Section 7 that it actually significantly improves the performance of the training phase.

We are interested in solving the following problem:

**Problem 1** ( $OFF(L, M_{GPU}, \beta)$ ). Consider a training phase with L-layers network. The operation time of forward and backward phase of the *i*-th layer is denoted as  $u_{F_i}$  and  $u_{B_i}$ . Given a GPU with memory  $M_{GPU}$  and bidirectional bandwidth  $\beta$  between the GPU and main memory, is there a schedule of uploading, delete and prefetching operations so that the training phase can be performed with an execution time at most T?

#### 4 Complexity Results

**Theorem 1.**  $OFF(L, M_{GPU}, \beta)$  is NP-complete in the strong sense.

*Proof.*  $OFF(L, M_{GPU}, \beta)$  clearly belongs to NP: given a schedule of all operations and communications, one can check in polynomial time that the execution is valid and fits within the execution time T.



Figure 3: Instance and schedule used in the NP-completeness proof of Theorem 1.

We prove that this problem is strongly NP-hard and therefore NP-complete by a reduction from the 3-partition problem: given a set of integers  $\{x_0, x_2, \ldots, x_{3m-1}\}$ such that  $\sum_i x_i = mV$ , decide whether it is possible to partition it into m parts  $\{S_1, \ldots, S_m\}$  so that for any  $j \leq m, |S_j| = 3$  and  $\sum_{i \in S_i} x_i = V$ . This problem is known to be NP-complete in the strong sense. Given an instance of 3-partition, we build the following instance of Problem 1:

- L = 5m,  $\beta = V$ ,  $M_{\text{GPU}} = mV + V$ , T = 2m
- $u_{F_i} = 0$  and  $|a_i| = 0$  for all i, except  $u_{F_{2m}} = m$
- $u_{B_i} = 0$  and  $|w_i| = V$  for  $i = 2k 1, k \in \{1, \dots, m\}$
- $u_{B_i} = 1$  and  $|w_i| = 0$  for  $i = 2k, k \in \{1, \dots, m\}$
- $u_{B_i} = 0$  and  $|w_i| = x_{i-2m}$  for  $2m + 1 \le i \le 5m$

We claim that this instance can be scheduled in time T = 2m if and only if the 3-partition instance is positive.

Let us first assume that there exists a solution to the 3-partition instance, i.e. sets  $(S_j)_{1 \le j \le m}$  such that  $\sum_{i \in S_j} x_i = V$ , and let us build a feasible schedule, as described on Figure 3.

Before  $F_1$ , the GPU memory contains all  $w_i$  for  $i \leq 2m$ , thus the available memory is  $M_{\text{GPU}} - \sum_{i=1}^{2m} |w_i| = V$ . The forward operations for the first 2m

layers take no time. With V free memory at the beginning of  $F_{2m}$ , prefetching  $w_{i+2m}$  for  $i \in S_1$  and offloading  $w_1$  can start simultaneously. After 1 time unit, both transfers are over since  $|w_1| = \sum_{i \in S_1} |w_{i+2m}| = V$ . Prefetching  $w_{i+2m}$  for  $i \in S_2$  and offloading  $w_3$  can thus start immediately. By the end of  $F_{2m}$ , all  $w_i$  for i > 2m are prefetched into memory and all  $w_{2k-1}$  for  $k \leq m$  are offloaded.

With  $w_i$  for i > 2m in the memory and V free memory for the gradient,  $F_{2m+1}$  to  $F_{5m}$  and  $B_{5m}$  to  $B_{2m+1}$  can also be performed in no time. After the end of  $B_{2m+1}$ , the free memory is given by  $M_{\text{GPU}} - \sum_{i=2m+1}^{5m} w_i = V$ . During  $B_{2m}$ ,  $w_{2m-1}$  is prefetched, and  $w_{i+2m}$  for  $i \in S_1$  is offloaded, which allows  $B_{2m-1}$  to start without delay. This pattern is repeated until  $B_1$ . After  $B_1$ , all  $w_i$  for  $i \leq 2m$  are in the GPU memory, and deleting  $\delta w_1$  provides V free memory. A new cycle can start from there. In this schedule, all the offloading and prefetching are overlapped with the computations, which yields an execution time for one cycle of  $u_{F_{2m}} + \sum_{k=1}^{m} u_{B_{2k}} = 2m$ .

Let us now assume that there exists a valid schedule of duration T = 2m, i.e. without any idle time on the processing device. Since the operation time between  $F_{2m+1}$  and  $B_{2m+1}$  is 0, all  $w_i$  for i > 2m have to be saved in memory before starting  $F_{2m+1}$  to avoid idle time. The memory usage is then  $\sum_{i=2m+1}^{5m} |w_i| = mV$ . If any  $w_{2k-1}$  for  $k \in \{1, \ldots, m\}$  is also saved in memory, the usage would be mV + V and there will be no space for the gradient computed by backwards. Therefore, all  $w_{2k-1}$  for  $k \in \{1, \ldots, m\}$  must be offloaded before  $F_{2m+1}$ .

To start  $B_{2m-1}$  immediately after  $B_{2m}$ ,  $w_{2m-1}$  has to be prefetched, and there should be at least V free memory for the gradient  $\delta w_{2m-1}$ . Given the state at the end of  $B_{2m+1}$  described above, the schedule needs to prefetch  $w_{2m-1}$ within time  $u_{B_{2m}}$ , and free enough memory to make room for  $\delta w_{2m-1}$ . Denote W the amount of memory offloaded during  $B_{2m}$ . The memory constraint to start  $B_{2m-1}$  without delay implies that  $W \ge V$ , and the bandwidth limitation implies that  $W \le V$ . Together, we have W = V, which means that there exists a set  $S_1$  such that  $\sum_{i \in S_1} |w_{i+2m}| = V, i > 2m$ . By repeating the same reasoning, we can find m subsets  $S_j$  such that  $\sum_{i \in S_j} |w_{i+2m}| = V$ , which gives a solution to the 3-partition problem and completes the proof.

This proof actually shows a stronger result: even if the set of weights to offload is given, finding an optimal schedule is still NP-complete. Indeed, in the above instance, the set of weights to offload provides no information about how to solve the 3-partition instance.

### 5 Greedy algorithm

We now propose a greedy algorithm to solve  $OFF(L, M_{\text{GPU}}, \beta)$ . The algorithm works in two phases: a *selection* phase to find which weights will be offloaded, and a *schedule* phase, which computes how to interleave the prefetching and offloading operations with F and B operations during the training cycle of one batch. **Selection phase** As mentioned in section 3, there are two options for offloading the weight of a layer k: (a) between  $F_k$  and  $B_k$ ; or (b) between  $B_k$  and  $F_k$ . We use (k, d) to represent the offloading choices, where d = 1 represents choice (a) and d = 0 represents choice (b). A solution of the selection algorithm consists of the set of offloading choices (k, d) with  $k \in \{1, \ldots, L\}$  and  $d \in \{0, 1\}$ . For an offloading choice (k, d), we say that it *covers* all the operations that are performed while  $w_k$  is no longer in memory, without taking into account the time taken by the communications: the choice (k, 1) covers  $F_{k+1}, \ldots, F_L, B_L, \ldots, B_{k+1}$ , while (k, 0) covers  $B_{k-1}, \ldots B_1, F_1, \ldots F_{k-1}$ ).

Our algorithm for the selection phase is based on a scoring function which estimates the *benefit* associated to an offloading choice. The algorithm maintains a value  $E_o$  for each operation o (where  $o \in \{F_1, \ldots, F_L, B_L, \ldots, B_1\}$ ), equal to the *excess* memory usage during operation o. Initially, the algorithm assumes that no weight is offloaded, *i.e.*  $E_o = M_o + \sum_i |w_i| - M_{\text{GPU}}$ , where  $M_o$  is defined by eq. (1). An offloading choice (k, d) reduces the excess memory usage of operation o by  $\min(E_o, |w_k|)$ , and its benefit is defined as the total memory saved over all operations o covered by this choice. The *cost* of an offloading choice  $w_k$ is simply the associated communication cost, equal to twice the size  $|w_k|$  since it needs to be communicated back and forth. The *profit* of an offloading choice is then defined as the ratio between its benefit and its cost.

There is one special case for the offloading cost: if one choice (k, d) has already been selected for one layer, then using to the *discount* opportunity described in section 3, selecting the other choice (k, 1 - d) only induces one communication. In that case, the cost is halved, and the profit is thus twice as large.

At the beginning, we assume that all the weights are kept in memory for every operation. As long as the maximum memory cost exceeds the available memory of GPU, the most profitable offloading choice is selected. The updated excess memory costs  $E_o$  will change the benefits of each choice for the next run. The resulting algorithm is described in Algorithm 1.

**Scheduling phase** Once the offloading choices have been selected, another greedy algorithm is used to build an actual schedule. Our algorithm relies on simple ordering rules: prefetching operations are performed in the order in which the data is required: if both  $w_i$  and  $w_{i+1}$  are offloaded after the forward phase,  $w_{i+1}$  will be prefetched before  $w_i$  since it will be needed in the backward phase earlier. Offloading operations are performed in First-In-First-Out (FIFO) order, which means that data produced earlier is offloaded first. There is an exception to this:  $F_k$  does not change  $w_k$ , and thus does not produce data to be offloaded. So if (k, 1) is selected in Algorithm 1, it is possible to schedule the corresponding communication *before* the forward operation. However, this is not required and we use this possibility on a best-effort basis.

Given these rules, our scheduling algorithm described in Algorithm 2 performs each operation as soon as possible. In general, an operation needs to wait for three events: (1) its input data is available (2) its resource is available (either

#### Algorithm 1 Greedy Selection Algorithm

Input: Cost of each operation, memory usage of each layer Output: selection of which layer weights to offload for all operations  $o \in \{F_1, \ldots, loss, B_{loss}, \ldots, B_1\}$  do  $E_o = \sum_i |w_i| + M_o - M_{GPU}$ while  $\max_o(E_o) > 0$  do for all choices (k, d) not yet selected do  $S \leftarrow$  set of operations covered by (k, d)if (k, 1 - d) is chosen then  $profit(k, d) = \cdot \sum_{o \in S} \min(E_o, |w_k|) / |w_k|$ else  $profit(k, d) = \frac{1}{2} \sum_{o \in S} \min(E_o, |w_k|) / |w_k|$ Select the choice  $(k, d) = \arg \max_{k,d} profit(k, d)$   $S \leftarrow$  set of operations covered by (k, d)Recompute  $E_o$  for all  $o \in S$ :  $E_o \leftarrow \max\{E_o - |w_k|, 0\}$ 

the GPU for a computing operation, or the corresponding communication link) (3) enough memory is available on the GPU (not applicable for an offloading operation). We say that an operation is *feasible* at time t if constraint (3) is satisfied. Algorithm 1 ensures that with our ordering rules all operations eventually become feasible. In Algorithm 2,  $Pre_k$  and  $Off_k$  denote prefetch and offload operations of  $w_k$ ,  $T_{ava}(\cdot)$  denotes the available time of a communication link and  $T_{end}(\cdot)$  denotes the ending time of an operation in Algorithm 2.

The scheduling process performs the operations in the order  $B_L, \ldots, B_1$ , followed by  $F_1, \ldots, F_L$ . With this order, it is easier to perform the early, best-effort scheduling of the offloading operations associated with a forward operation, as mentioned above: while the communications related to the backward operations are scheduled, we can keep a waiting list WL of best-effort offload opportunities, and schedule them whenever the communication link is available. However, the scheduling algorithm needs to know which data have already been offloaded or prefetched at the start of  $B_l$ . With our ordering rules, we can try all possibilities by selecting a cutoff point in the sequence of communication operations that needs to be performed, assuming that all communications before the cutoff have been performed. Algorithm 2 returns the solution with the smallest execution time among all possibilities.

The time complexity of this greedy algorithm is  $O(L^3)$ . In practice, it takes less than 1 minute to find a solution for a DNN with up to 128 layers.

#### 6 Lower bound

In this section, we present an original Mixed Integer Linear Programming formulation for the  $OFF(L, M_{GPU}, \beta)$  problem. Given the difficulty of the problem, we do not propose an exact formulation, but one that provides a valid lower

Algorithm 2 Greedy Scheduling Algorithm for all cutting points in the sequence of communications do Assume communications before the cutting point are performed before  $B_L$ for k in  $[L, \ldots, 1]$  do if (k, 1) has been selected then  $\triangleright B_k$  needs Prefetch Schedule  $Pre_k$  at time max $(T_{ava}(Pre))$ , earliest time when  $Pre_k$  is feasible) Schedule  $B_k$  at time  $\max(T_{end}(Pre_k), T_{end}(B_{k+1}))$  $\triangleright B_k$  needs Offload if (k, 0) has been selected then while  $T_{end}(B_k) - T_{ava}(Off) > \min_{k' \in WL} |w_{k'}|$  do Offload min  $|w_{k'}|$  from WL, without deleting it from GPU Schedule  $Off_k$  at time max $(T_{end}(B_k), T_{ava}(Off))$ else if (k, 1) has been selected then  $\triangleright F_k$  needs Offload Append k to WL $\triangleright WL$  is initially empty for k in  $[1, \ldots, L]$  do if (k, 0) has been selected then  $\triangleright F_k$  needs Prefetch Schedule  $Pre_k$  at time max $(T_{ava}(Pre))$ , earliest time when  $Pre_k$  is feasible) Schedule  $F_k$  at time  $\max(T_{end}(Pre_k), T_{end}(F_{k-1}))$  $\triangleright F_k$  needs Offload if (k, 1) has been selected then if  $w_k$  is not yet in CPU memory then Schedule  $Off_k$  at time  $\max(T_{end}(F_k), T_{ava}(Off))$ else Delete  $w_k$  from GPU memory at time  $\max(T_{end}(F_k))$ ,  $T_{ava}(Off)$ Finish communications before the cutting point return the solution with lowest execution time

bound on the makespan: the result of this MILP is not a schedule, because it satisfies only a subset of the constraints of  $OFF(L, M_{GPU}, \beta)$ . This lower bound will be used in Section 7 to evaluate the quality of the schedules produced by the algorithms of Section 5 and the algorithms of the literature.

We number the operations from 1 to 2*L*, starting from  $B_L$ , so that the *L*-th operation is  $B_1$ ,  $F_i$  is the (L + i)-th operation, and  $B_i$  is the (L - i + 1)-th operation. In the following, we use  $\overleftrightarrow_{j=a}^{b}$  to denote a sum from operation *a* to operation *b*, potentially cycling back to 1 if b < a: in that case,  $\overleftrightarrow_{j=a}^{b} X_j = \sum_{i=a}^{2L} X_i + \sum_{i=1}^{b} X_i$ .

 $\sum_{j=a}^{2L} X_j + \sum_{j=1}^{b} X_j.$ For each operation j, our formulation considers the interval between the start of operation j and the start of the next operation. Let us define the following continuous variables:

- for all i,  $O_{i,j}$  represents the amount of data from  $w_i$  offloaded from GPU to CPU (note that this transfer does not change the GPU memory used)
- for all  $i, P_{i,j}$  represents the amount of data from  $w_i$  prefetched from CPU
- for all  $i, D_{i,j}$  represents the amount of data from  $w_i$  deleted from the GPU
- $idle_i$  represents the idle time after operation j

In addition, for each layer i we define three binary variables, that encode offloading choices:

- $D_i^{(1)} = 1$  if  $w_i$  is removed from memory between  $F_i$  and  $B_i$  (similar to the choice (i, 1) in Algorithm 1), and 0 otherwise.
- $D_i^{(0)} = 1$  if  $w_i$  is removed from memory between  $B_i$  and  $F_i$  (similar to the choice (i, 0) in Algorithm 1), and 0 otherwise.
- $O_i^* = 1$  if  $w_i$  is offloaded at some point, and 0 otherwise.

The constraints of our formulation can be classified in four groups:

Idle time. We make sure the communications are finished before going to the next operation, with a special case to ensure that the transfer of  $w_i$  does not take place during its backward:

$$\forall j, \quad u_j + idle_j \ge \sum_i O_{i,j}/\beta \qquad \text{and} \quad u_j + idle_j \ge \sum_i P_{i,j}/\beta \qquad (2)$$

$$\forall i, \quad idle_{L-i+1} \ge O_{i,L-i+1}/\beta \tag{3}$$

**Memory constraints.** Between operations j and j+1,  $(P_{i,j}-D_{i,j})$  indicates the change in memory usage associated with  $w_i$ . We define  $m_{i,k}$  as the memory usage of  $w_i$  at the beginning of operation k. Since  $w_i$  is completely in GPU memory before  $B_i$ , we get  $m_{i,(L-i+1)} = |w_i|$ . We can thus deduce  $m_{i,k} =$  $|w_i| + \sum_{j=L-i+1}^{k-1} P_{i,j} - D_{i,j}$ . Recall that the memory peak during operation kis  $M_k$ , as defined by eq. (1). The global memory constraint is given by

$$\forall k, \sum_{i} m_{i,k} \le M_{\rm GPU} - M_k \tag{4}$$

Validity constraints. We ensure that the cycle is stable, and that  $w_i$  is completely in GPU at the start of  $F_i$  (operation L + i):

$$\forall i, \quad \sum_{j=0}^{2L-1} P_{i,j} - D_{i,j} = 0 \qquad \text{and} \qquad \sum_{j=L-i+1}^{L+i-1} P_{ij} - D_{ij} = 0 \qquad (5)$$

Additionally, we write constraints on  $w_i$  at the beginning of every operation k to ensure that the amount of prefetched data is not higher than the deleted data, and that no data is lost when deleting from GPU memory:

$$\forall i, k, \quad \sum_{j=L-i+1}^{k-1} P_{i,j} - D_{i,j} \ge 0 \quad \text{and} \quad \sum_{j=L-i+1}^{k-1} O_{i,j} + P_{i,j} - D_{i,j} \ge 0 \quad (6)$$

**Binary constraints.** Finally, we add constraints to ensure that the weight of a given layer is either deleted/offloaded completely, or not at all.

$$\forall i, \quad \sum_{j=1}^{2L} O_{i,j} = |w_i| O_i^* \tag{7}$$

$$\forall i, \quad \sum_{j=L+i}^{L-i} D_{i,j} = |w_i| D_i^{(1)} \qquad \text{and} \quad \sum_{j=L-i+1}^{L+i-1} D_{i,j} = |w_i| D_i^{(0)} \qquad (8)$$

Obviously, any solution to  $OFF(L, M_{GPU}, \beta)$  can be transformed into a set of variables that fulfill these contraints. We thus obtain the following lower bound:

**Theorem 2.** The optimal execution time  $T^*$  of problem  $OFF(L, M_{GPU}, \beta)$  is not smaller than the optimal value of the MILP defined by:

$$\min \sum_{j} u_{j} + idle_{j}$$
  
s.t. eqs. (2) to (8)

Note that the opposite is not true: it is not clear that any solution to this MILP can be turned into a valid solution of  $OFF(L, M_{GPU}, \beta)$ . Indeed, the memory constraints (4) are only expressed at the beginning of each operation, instead of during the while interval from the beginning of an operation to the next. The solution of the MILP therefore only provides a lower bound, not the optimal value of  $OFF(L, M_{GPU}, \beta)$ .

#### 7 Simulation results

To estimate time and memory costs when training GPT-2 and Bert models, we rely on the hyper-parameters from Megatron-LM [24]. We test both DNNs with a number of layers varying from 36 to 144 and batch sizes of 16, 32 and 64. For both models, we use the hidden size of 3072 and 24 attention heads. The number of parameters ranges from 4.2B to 16.4B. For a batch size of 16, about 20MB of activations are saved in GPU memory per layer, while about 1GB to 2GB intermediate activations will be recomputed for each layer during the backward. The execution times for the simulations are experimentally obtained on a GPU V100 with 16GB memory. Since a large memory is needed to store the intermediate activations generated in each transformer block,  $M_{GPU}$  used for simulation is smaller than 16GB and depends on the batch size. IBM ILOG CPLEX [18] is used to find the solution for the ILP as described in Section 6.

#### 7.1 Results

Table 1 provides the simulation results using the measurements from real machines. The lower bound defined in Section 6 gives no idle time in all settings. Our greedy algorithm defined in Section 5 shows about 1% of overhead with respect to the lower bound. It demonstrates a significant improvement compared to the overhead of L2L [19], which is 10% to 50% larger than the lower bound. The performance of L2L highly depends on the processing times, that are approximately linear with respect to the batch size. We also notice that offloading weights more than once does not bring a significant change to the performance of the greedy algorithm in these simulations.

#### 7.2 Bandwidth

To study the effect of bandwidth, another set of simulations has been performed for different values of  $\beta$ . The results with the ILP show almost no idle time when the bandwidth is higher than 4GB/s. The comparison between blue and green plots shows that our greedy algorithm is very close to the lower bound for a given bandwidth. The L2L plot indicates that L2L always causes a reduction in throughput compared to our greedy algorithm, less significant when both algorithms need long transfer times. Also, we notice that Greedy<sup>\*</sup> is always slower than Greedy, which means that the discount is useful, especially for certain bandwidth ranges. In some cases, not considering the discount opportunity

Model			Duration (ms)				
Network	Depth	$Batch_{size}$	$\sum_{j} u_{j}$	ILP	Greedy	L2L	$Greedy^*$
GPT-2	74	64	47421	47421	47493	52707	47493
GPT-2	56	64	35553	35553	35625	39542	35625
GPT-2	38	64	23694	23694	23838	26387	23838
GPT-2	74	32	23697	23697	23769	28983	23769
GPT-2	56	32	17762	17762	17834	21751	17834
GPT-2	38	32	11840	11840	11948	14533	11948
GPT-2	74	16	11612	11612	11684	16898	11715
GPT-2	56	16	8697	8697	8769	12686	8800
GPT-2	38	16	5794	5794	5902	8487	5934
Bert	144	64	34486	34486	34499	38379	34499
Bert	96	64	22965	22965	22978	25577	22978
Bert	144	32	17443	17443	17483	21336	17483
Bert	96	32	11617	11617	11657	14230	11657
Bert	144	16	9090	9090	9183	12983	9215
Bert	96	16	6058	6058	6085	8670	6085

Table 1: The bidirectional bandwidth is 12GB/s for both prefetch and offload. Greedy\* represents the Greedy algorithm while weights are not only offloaded once.

can induce up to 5 times more idle time.

### 8 Conclusion

In this paper, we consider GPU memory savings during training, by using the possibility to store the network layers in the CPU memory when they are not used. We propose a sophisticated model for the memory associated with the storage of layer weights, which takes into account all opportunities to save memory: by updating weights as soon as possible and by performing only one offload per forward/backward cycle. We have established the NP-completeness of the layer selection and transfer scheduling problem, and we have proposed an efficient heuristic to solve it in practice. The perspectives opened by this work are numerous. The extension to the offloading of optimizer states and activations are natural perspectives, as well as the possibility to perform weight updates on the CPU. Moreover, the proposed strategy is more efficient when the batch size is large, which suggests combining it with re-materialization approaches to increase the batch size.

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Figure 4: To study the effect of bandwidth, we use the same measurement showed in Table1, while the bandwidth varies from 0.5, 1, 2, 4, 8 and 12 GB/s. The batch size is chosen as 32 or 64.