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# An Integer Linear Programming Approach for Pipelined Model Parallelism

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# An Integer Linear Programming Approach for Pipelined Model Parallelism

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**Abstract:** The training phase in Deep Neural Networks has become an important source of computing resource usage and because of the resulting volume of computation, it is crucial to perform it efficiently on parallel architectures. Even today, data parallelism is the most widely used method, but the associated requirement to replicate all the weights on the totality of computation resources poses problems of memory at the level of each node and of collective communications at the level of the platform. In this context, the model parallelism, which consists in distributing the different layers of the network over the computing nodes, is an attractive alternative. Indeed, it is expected to better distribute weights (to cope with memory problems) and it does not imply large collective communications since only forward activations are communicated. However, to be efficient, it must be combined with a pipelined/streaming approach, which leads in turn to new memory costs. The goal of this paper is to model these memory costs in detail and to show that it is possible to formalize this optimization problem as an Integer Linear Program (ILP).

Key-words: Training, Memory, Model Parallelism, Integer Linear Programming

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# Une approche fondée sur la programmation linéaire pour le parallélisme de modèle

**Résumé :** La phase d'apprentissage dans les réseaux neuronaux profonds est devenue une source importante d'utilisation des ressources de calcul et, en raison du volume de calcul qui en résulte, il est crucial de l'exécuter efficacement sur des architectures parallèles. Aujourd'hui encore, le parallélisme de données est la méthode la plus utilisée, mais l'exigence associée de répliquer tous les poids sur la totalité des ressources de calcul pose des problèmes de mémoire au niveau de chaque nœud et de communications collectives au niveau de la plateforme. Dans ce contexte, le parallélisme de modèle, qui consiste à répartir les différentes couches du réseau sur les nœuds de calcul, est une alternative intéressante. En effet, il est censé mieux répartir les poids (pour faire face aux problèmes de mémoire) et il n'implique pas de grosses communications collectives puisque seules les activations "forward" sont communiquées. Cependant, pour être efficace, elle doit être combinée avec une approche pipelinée/streaming, ce qui entraîne à son tour de nouveaux coûts mémoire. L'objectif de cet article est de modéliser ces coûts de mémoire en détail et de montrer qu'il est possible de formaliser ce problème d'optimisation comme un programme linéaire en nombre entier (ILP).

**Mots-clés :** Apprentissage, Mémoire, Parallélisme de modèle, Programmation linéaire en nombres entiers

# An Integer Linear Programming Approach for Pipelined Model Parallelism

#### February 1, 2022

## 1 Introduction

Deep Neural Network (DNN) training is a long and memory-intensive operation. Indeed, supervised DNN training requires performing numerous forward and backward computations, each on a subset of input data called a mini-batch. In turn, each forward and backward phase involves complex data dependences and induces memory issues. In practice, parallel training is performed both on small groups of GPU machines and on large HPC infrastructures [19, 40], especially because HPC machines offer high-bandwidth and low-latency networks [22, 30, 12].

The first approach to use parallelism at the level of the node is to make the best use of the available multi-core by optimizing the individual compute kernels, which usually consist of tensor computations. This approach has been widely used in the context of GPUs and TPUs and has made the success of frameworks such as TensorFlow [4], PyTorch [35] or NGraph [1].

At a larger scale, the best known approach to parallel DNN training is the so-called data parallel approach. Using data parallelism [43, 34], the model and all associated weights are replicated on all participating nodes. Then, different mini batches are trained in parallel on different nodes. In this framework, all participating nodes execute forward and backward phases in parallel, and thus all compute a gradient for all weights in the network. Synchronization between the nodes takes place at the end of the backward step, and all partial gradients are collected and aggregated through a collective communication such as AllReduce, and then broadcast to all participating nodes by a Broadcast type operation.

The above approach is feasible and scalable as long as two conditions are fulfilled. The first condition is related to communication resources and states that the network should support the collective communications of the weights, without inducing too much idle computing time. The second condition is related to memory and states that each participating node must be able to store all network weights and activations corresponding to the processing of a mini-batch.

In order to limit network issues and lower the size of the collective communication, data parallelism can be combined with compression to limit the size of messages. When used at a very large scale, this approach nevertheless leads to poor performance due to synchronization and communication costs and requires the use of huge mini-batches, which could also have a negative impact on the performance of the training phase [31].

To deal with limitations induced by memory issues, several approaches have been proposed. Indeed, it has been observed that in many cases, large and heavy models are required to reach good classification accuracy.

In general, the memory consumption during the training phase is composed of two main parts [17]. The first source of memory consumption is due to the storage of all the forward activations on each node (i.e. all the outputs of the different stages of the network) until the associated backward operation. This part is directly proportional to the size of the mini-batch itself. The second part is related to the storage of the network parameters (weights) and is directly connected to the size of the model [27]. Using the data parallel approach, these weights must be replicated on each node, and must even be aggregated and disseminated on the network after each parallel training phase in mini-batches.

To reduce the memory requirements related to the storage of forward activations, several approaches have been proposed. Rematerialization [20, 8, 16, 28, 26, 29] consists in deleting from the memory certain activations computed during the forward phase, which then must be recomputed during the backward phase. Another approach [38, 37, 6] is to offload some of the activations from the GPU memory to the CPU memory during the forward phase and then to bring them back in GPU memory during the backward phase when they are actually required.

In order to limit the memory requirements resulting from the storage of network weights, a natural approach is to distribute the different layers of the network over several computation resources. This approach, denoted as model parallelism, has also been advocated in many papers [13, 24, 31, 41]. In this context, each mini-batch is processed by a sequence of processors, and only activations are communicated between processors. This approach is orthogonal to data parallelism and can naturally be combined with it, using several processors to manage the same sub-part of the DNN.

Unfortunately, if images are processed entirely and in sequence, model

parallelism can reduce memory requirements, but it does not accelerate computations because of the shape of the dependencies introduced by backpropagation, as shown in [24, 31]. To obtain some speedup using this approach, it is necessary to process several images in parallel, using a pipelined (or streaming) approach. On the other hand, such approach requires the storage of several models (to ensure that forward and backward computations for the same image use the same weights) and several activations of each type, corresponding to different images being processed in the pipeline.

Most of the literature on this approach combining pipelining and model parallelism [24, 31, 32, 41] focuses on the problem of finding an efficient partitioning of the network, that balances the load between processors and minimizes data exchanges between resources. However, in these approaches, memory issues are considered a posteriori and typically, the solution found without memory constraints is then degraded, in terms of throughput, to fulfill these constraints.

To the best of our knowledge, the analysis of the induced memory requirements proposed in the present paper is much tighter than what has been proposed in the literature. In [39], the authors share several characteristics with the present paper. As in the present paper, the authors propose an ILP based solution for computing non-contiguous allocations of layers. The main advantage of [39] over our approach is that it does not require graph linearization and can be directly applied to general DAGs. The paper addresses both the inference and training phase, but the specificities of the memory issues in the case of training are not addressed. These specificities are in practice crucial in this context, and the careful modeling of memory requirements is the main contribution of the present paper. Finally, since both [39] and our approach are based on integer linear programming, their use is in practice limited to medium size networks (Resnet-50 is the deepest network considered in both papers).

Overall, in this paper, we perform the theoretical analysis of the pipelined model parallelism under memory constraint. Our theoretical contributions are a very careful modeling of induced memory costs, an analysis of the potential benefits of non-contiguous allocations and an Integer Linear Program to compute a non-contiguous solution, that provides significant improvement over the literature for reasonable size networks.

The rest of the document is organized as follows. The related work is presented in Section 2. We introduce the notations and the computational model we use in Section 3. We then propose an Integer Linear Programming (ILP) formulation of the optimization problem, which takes into account all sources of memory consumption in Section 4. The performance of the ILP-based solution, both in terms of solution quality and running time are analyzed in Section 5. Finally, conclusions and perspectives of this work are proposed in Section 6.

## 2 Related Works

Memory consumption is now becoming an important issue in deep learning and encompasses many different aspects. Unfortunately, the memory limitations of current hardware often prevent data scientists from considering larger models, larger images, or larger batches of images. In practice, training is performed automatically and transparently for the user thanks to autograd tools for backpropagation, such as tf.GradientTape in TensorFlow or torch.autograd.backward in PyTorch.

In order to reduce memory needs, one line of research is to design and train memory efficient architectures while aiming at the same performance as state-of-the-art networks. Reversible Neural Networks [18, 9] (RevNet), for example, are designed to allow the backpropagation algorithm to be performed without storing forward activations until the computation of the associated backward operation. Quantization [36, 25] and pruning [21] rather reduce memory consumption at the time of inference by changing weights and/or network activations into binary or quantized variables. Other *ad-hoc* architectures such as MobileNets [23] or ShuffleNet [42] finally try to sparsify the network architecture in order to reduce the size of the model.

Rematerialization is being increasingly employed to reduce memory usage. The use of rematerialization strategies has recently been advocated for DNN in several papers. A direct adaptation of the results on homogeneous chains has been proposed for the case of Recurrent Neural Networks (RNN) in [20]. A further generalization of the results on homogeneous chains enabled the derivation of optimal rematerialization strategies for joint-networks [8], which consist of several homogeneous chains joined together at the end. Some research has attempted to adapt rematerialization strategies to Arbitrary Computation Graphs (ACG) in [16, 28, 26, 29]. For practical use, an implementation of rematerialization exists in PyTorch [2], based on a simple strategy of periodic and one-pass rematerialization that exploits the ideas presented in [11]. Another implementation [3] based on [5] has also been proposed.

Another alternative [38, 37] is to offload some of the activations from the GPU memory to the CPU memory and to bring them back when required during the backward phase. Finally, domain decomposition or spatial paral-

lelism techniques can be used to limit the memory needed to store the forward activations. In [14], dividing large images into smaller ones makes it possible to train the network in parallel on the smaller images (enhanced by a halo), at the cost of additional communications to synchronize parameter updates. A similar strategy has been used in channel and filter parallelism in [15]. Approaches like rematerialization, activation offloading, spatial parallelism and channel parallelism are orthogonal to our approach and could be used in combination to achieve even greater memory savings, although it is out of the scope of the present paper.

The use of model parallelism has been proposed in [24], where the training batch is split into several mini-batches, which are then pipelined through the layers of the network. Once the forward and backward phases have been computed on all these mini-batches, the weights are then updated. This approach is fairly simple to implement but has the disadvantage of leading to a very limited speedup. It has been proposed in [31] to change a bit this training process, by only forcing that for a given image the forward and backward tasks use the same network weights. By weakening the constraint on the training process, PipeDream is able to achieve a much better utilization of processing resources.

An important issue related to memory and the pipelined approach proposed in [31] is the need to keep many copies of the weights. To solve this issue, a strategy has been proposed in [32] to keep only two models in memory. In the present paper, we use the strategy of [32] in a more general framework. Another strategy to keep fewer models in memory is to allow more asynchronous updates. This strategy has been explored in particular in [10], which proposes an intermediate approach to avoid the phenomenon of gradient staleness, though it requires 2 versions of the model weights and 2 version of the gradients, which does not save memory more.

# 3 Model and Notations

#### 3.1 Notations

In this section, we rely on the model presented in [7]. For completeness, we provide it in this report as well.

In what follows we represent each DNNs as a chain so that the task graph corresponding to forward and backward propagation is depicted in Figure 1. This assumption on the shape of the network might require linearization and is used in most of the related literature [11, 31, 32, 10, 41].

Let us model the network as a chain of L layers (each layer could consist

in a single layer or a block of layers in case of prior linearization), numbered from 1 to L. Each layer l is associated both to a forward operation  $F_l$  and a backward operation  $B_l$ . During training, each mini-batch undergoes a forward pass followed by a backward pass through the whole network and the dependences between tasks are depicted in Figure 1. Further, we consider  $F_l$ and  $B_l$  as compute tasks  $T_{l'}$ , where if  $l' \leq L$  then  $T_{l'} = F_{l'}$  and if l' > L then  $T_{l'} = B_{2L-l'+1}$ .

As we are interested in solutions in which the different layers of the network are allocated to potentially different computation resources, we also introduce communication tasks  $T_{l,l+1}^c$ . These communication tasks correspond to sending forward activations or gradients from one computation resource to another, and their cost is 0 between two successive layers allocated to the same resource. We denote by  $a^{(l)}$  the activation tensor produced by  $T_l, l \leq L$  and by  $a^{(2L-l)} = b^{(l)} = \frac{\partial \mathcal{L}}{\partial a^{(l)}}$  the back-propagated intermediate gradient value provided as input of the backward operation  $T_{2L-l}$ .

The sequence of tasks associated with the processing of a minibatch is therefore given by  $T_1, T_{1,2}^c, T_2, \ldots, T_{L-1,L}^c, T_L, T_{L+1}, T_{L+1,L+2}^c, T_{L+2}, \ldots, T_{2L-1,2L}^c, T_{2L}$ .

During the training operation, we assume that the set of input data (typically images) is split into mini-batches of size  $\mathcal{B}$ . In this context, we denote by

- $d_l = u_{F_l}$  the duration of the forward task on the layer l with a mini-batch of size  $\mathcal{B}$
- $d_{2L-l+1} = u_{B_l}$  the duration of the backward task on the layer l with a mini-batch of size  $\mathcal{B}$
- $a_l$  the size (in bytes) of the tensor  $a^{(l)}$  produced by  $T_l, l \leq L$  when applied to a mini-batch of size  $\mathcal{B}$  and
- $a_{l'}$  the size (in bytes) of the tensor  $a^{(l')} = b^{(2L-l')}$  produced by  $T_{l'}, l' \ge L+1$  when applied to a mini-batch of size  $\mathcal{B}$ . In general, we assume



Figure 1: Graph for the Backward Propagation

that tensors  $a^{(l)}$  and  $b^{(l)}$  have the same size, *i.e.*  $\forall l \leq L, a_{2L-l} = a_l$ .

Let us denote by P the total number of available GPUs and let us assume (as in PipeDream) that all pairs of GPUs are connected through a direct link of capacity  $\beta$ . Moreover, we assume that each GPU is equipped with an available memory of size M. This memory is used to store all data required to perform the training operation. More precisely, this memory requirements can have different origins:

- Model weights. Since we are considering model parallelism, we assume that the L layers of the network are split across the P GPUs. If GPU  $P_k$  is in charge of layer l, then it has to store the corresponding weight denoted as  $W_l$ . As we will see, in order to update the weights and to use consistent weights during both the forward and the backward phases,  $P_k$  in practice stores several copies of the weights. In what follows, in order not to add more weight and activation copies, we assume that the processor in charge of a layer is in charge of processing both the forward and backward tasks associated to this layer.
- Activations. Let us now concentrate on activations  $a^{(l)}$ . As depicted in Figure 1, activation  $a^{(l)}$  is produced by  $T_l$  and it must be kept in memory until task  $T_{2L-l}$  consumes it to produce  $b^{(l)}$ . Therefore, a memory of size  $a_l$  must be reserved to store an activation between tasks  $T_l$  and task  $T_{2L-l+1}$ . As we will see, in general, to keep processors busy, we process several batches in parallel. Therefore, several activations corresponding to layer l are kept in memory simultaneously.
- Gradients. Let us now concentrate on gradients  $b^{(l)}$ . As depicted in Figure 1, the gradient  $b^{(l)}$  produced by task  $T_{2L-l}$  is consumed by  $T_{2L-l+1}$  to produce  $b^{(l-1)}$ . Therefore, a memory of size  $a_{2L-l}(=a_l)$  must be reserved to store an activation between tasks  $T_{2L-l}$  and  $T_{2L-l+1}$  to produce  $b^{(l-1)}$ . Thus, gradients are kept in memory for a much shorter time than activations.
- Communication buffers. When tasks  $T_l$  and  $T_{l+1}$  are assigned to different GPUs, a communication takes place, and we assume that for convenience some memory is reserved as a buffer to store  $a^{(l)}$  while it is sent or received. This requires a storage of size  $a_l$  on both GPUs.

#### 3.2 Periodic Schedules and Valid Patterns

In general, throughout this paper, we are interested in finding optimal task allocation (load balancing) and optimal periodic schedules. Both problems could be solved either simultaneously or separately *i.e.* solving the load balancing problem first and then finding an optimal schedule for it. Considering some fixed allocation, it is possible to build various schedules, but we restrict the search to periodic schedules that follow the Pattern conditions defined in Definition 1 (see Figure 2 for an example). In Definition 1, a valid pattern contains exactly one task  $T_l$  of each type l, for  $l \leq 2L$ . The pattern defines both the sequence of tasks that is performed on each computing resource, and the index shifts between the tasks that are processed in the pattern. For example, if  $T_2$  takes place before  $T_1$  in the pattern, then  $T_2$  must necessarily apply to a batch for which  $T_1$  has been processed in the previous copy of the pattern. In practice, we denote by s(l) the shift associated to  $T_l$ , so that previous condition becomes  $s(1) \geq s(2) + 1$ . Hence, during the *i*-th copy of the pattern in the periodic schedule,  $T_l$  operates on mini-batch i + s(l).

**Definition 1** (Valid Pattern). A valid pattern is defined by the following constraints:

- $\forall l \leq 2L, T_l$  is present exactly once in the pattern, on one of the GPUs and it has an exclusive access to the GPU during  $d_l$  time units.
- $\forall l \leq 2L 1, T_{l,l+1}^c$  is present exactly once in the pattern, on one of the communication links. Its duration is 0 if  $T_l$  and  $T_{l+1}$  are located on the same resource. Otherwise, it gets an exclusive access to the link between the GPUs that process  $T_l$  and  $T_{l+1}$  during  $a_l/\beta$  time units.
- task shifts are valid, *i.e.* if the shift for  $T_l$  is denoted by s(l), then  $\forall l' > l$ , if task  $T_{l'}$  starts before the end of  $T_l$ , then the shift for  $T_{l'}$  must satisfy s(l') < s(l).
- In the pattern, on each GPU and on each communication resource, the length of the period is smaller than T
- the starting dates of all tasks and communications should be within the time range of size  ${\cal T}$

**Definition 2** (1F1B). We call all periodical schedules satisfying the pattern described in Definition 1 as 1F1B schedules (1-Forward-1-Backward).

**Lemma 1.** We can restrict the search of optimal periodic schedules to the search of optimal patterns.



Figure 2: Example of valid pattern. The batch indices are in the superscipts near task names

*Proof.* Clearly, it is easy to build the periodic pattern from a valid periodic schedule.

Let us now suppose that we have a valid periodic pattern and let us prove that it can be turned into a valid periodic schedule. To do so, we need to specify, for each task, the index of the mini-batch on which this task operates. In order to determine these indexes, we can use a basic greedy algorithm. Let us suppose that  $T_{2L}$  processes the *i*-th mini-batch during a given period. Then, we can always assume that during this period,  $T_{2L-1,2L}^c$  is performed on mini-batch i+2. Indeed let us assume that  $T_{2L}$  is performed on  $P_k$  and  $T_{2L-1}$ on  $P_{k'}$ ,  $k' \neq k$  (the case where both tasks are scheduled on the same resource is less interesting since the duration of  $T_{2L-1,2L}^c$  is 0 in this case). Since there could a small shift in the periodic pattern between  $P_k$  and  $P_{k'}$ , it might happen that  $T_{2L}$  has already started on  $P_k$  at the time when  $T_{2L-1,2L}^c$  ends. On the other hand, since this shift is always smaller than T by construction, the communicated activation during the current period is available on  $P_k$  2 periods after, so that  $P_k$  is indeed able to process mini-batch i + 2. Using the same reasoning and the same shift of two mini-batches between any two consecutive processing and communication tasks, it is possible to transform any valid periodic pattern into a valid schedule.

Note that our greedy algorithm induces a very large span for the indexes of the mini-batches performed in the same period. We will see later that this large span induces a large memory costs, but that given a periodic pattern, it is possible to build a schedule with minimal memory need (this will be done in Lemma 8).

Similarly to PipeDream, we use the notion of 1F1B schedules from Def-

inition 2. As mentioned earlier, we are interested only in 1F1B schedules, and therefore we look for 1F1B solutions that minimize the period T given a batch-size  $\mathcal{B}$ , or equivalently that maximize the number of trained images per time unit in steady-state, given by  $\frac{\mathcal{B}}{T}$ .

# 4 Integer Linear Program

We present in this section an Integer Linear Program to find a valid pattern with minimum period length. We concentrate on scheduling issues on both computational and communication resources in Section 4.1, then we consider memory related issues in Section 4.2.

We first present the main variables used in this ILP (other variables are introduced later):

- T denotes the period considered in the 1F1B schedule;
- $z_{l,l'}$  is equal to 1 if task  $T_l$  and task  $T_{l'}$  are processed on the same resource, and 0 otherwise (it is implied that  $z_{l,l} = 1$  and  $z_{l,l'} = z_{l',l}$ );
- $\tau_l$  is the starting time of task  $T_l$  in the period ;
- $\tilde{\tau}_l$  is the starting time in the considered period of the communication of the output of  $T_l$ .

We also use  $d_l$  to denote the duration of the task  $T_l$ , *i.e.*  $u_{F_l}$  if  $l \leq L$ and  $u_{B_l}$  if  $l \geq L + 1$ , and  $\tilde{d}_l$  to denote the time needed to communicate the activation produced by  $T_l$ , equal to  $\frac{a_l}{B}$ . In several places, we use a large constant K which needs to be larger than the period, for example we can use  $K = \sum_l d_l + \frac{a_l}{B}$ .

#### 4.1 Communication and Computation Constraints

#### 4.1.1 Limit on the number of resources

In order to provide a limit on the number of resources used, we introduce variable  $f_l$  which is equal to 1 if and only if task  $T_l$  is the lowest-index task processed on its resource. To this end, we consider the following set of constraints:

$$\forall l < l' < l'', \quad z_{l',l''} \ge z_{l,l'} + z_{l,l''} - 1 \tag{1}$$

$$\forall l, \quad f_l \ge 1 - \sum_{l' < l} z_{l,l'} \tag{2}$$

$$\sum_{l} f_l \le P,\tag{3}$$

and we show that they are enough to obtain the following:

**Lemma 2.** Constraints (1)-(3) ensure that at most P resources are used in the schedule.

Proof. Constraint (1) ensures the consistency of the  $z_{l,l'}$  variables: for any l, l', l'', if  $z_{l,l} = 1$  and  $z_{l,l''} = 1$ , then  $z_{l',l''} = 1$ . They can thus be used to define an equivalence relation between tasks, where each class contains tasks which are processed on the same resource. Then, Constraint (2) ensures that  $f_l$  is exactly 1 for the task with the smallest index among all the tasks processed on a given resource, and is trivially satisfied for all other tasks. Therefore, the sum of  $f_l$  provides the total number of allocated resources, and constraint (3) enforces that no more than P resources are used in the schedule. Reciprocally, in any valid solution that uses no more than P resources, there exists an assignment of  $f_l$  variables such that  $\sum_l f_l \leq P$ , *i.e.* the assignment where  $f_l = 1$  for the task with the smallest index processed on the resource and 0 for all other tasks.

In addition, we consider only schedules where forward tasks  $T_{L-l}$  for  $l \leq L$  are placed on the same resource as their respective backward tasks  $T_{L+l+1}$ , so we add the following equation to the Linear Program

$$\forall l < L, \quad z_{L-l,L+l+1} = 1.$$
 (4)

#### 4.1.2 Ordering of Computational Tasks

Let us now consider tasks that are processed on the same resource. In order to enforce that two tasks processed on the same resource cannot overlap, we introduce a set of variables  $w_{l,l'}$  and the following equations, valid for all  $l \neq l'$ :

$$\tau_l - \tau_{l'} + K(1 - z_{l,l'} + w_{l,l'}) \ge d_{l'},\tag{5}$$

$$\tau_{l'} - \tau_l + K(2 - z_{l,l'} - w_{l,l'}) \ge d_l, \tag{6}$$

$$w_{l,l'} \le z_{l,l'}.\tag{7}$$

Throughout the proofs, K is used to define a condition that must be valid as soon as a boolean variable x is equal to 1. The general idea is to use (1-x) \* K in the equation as follows: if x = 1, then (1-x)K = 0 and the rest of the condition must be satisfied. Conversely, if x = 0 then (1-x)K is significantly larger than the other terms and the condition is automatically satisfied, regardless of the value of the other variables. An example of the use of this technique can be found below in Lemma 3.

**Lemma 3.** Constraints (5)-(7) ensure the following:

- If  $T_l$  and  $T_{l'}$  are assigned to the same resource, then either  $T_{l'}$  starts after the end of  $T_l$  (and  $w_{l,l'} = 1$ ), or  $T_l$  starts after the end of  $T_{l'}$  (and  $w_{l,l'} = 0$ ).
- If  $T_l$  and  $T_{l'}$  are not assigned to the same resource, then  $w_{l,l'} = 0$ .

*Proof.* Let us assume that  $T_l$  and  $T_{l'}$  are assigned to the same resource. Then, by definition,  $z_{l,l'} = 1$  and we obtain

$$\tau_{l} - \tau_{l'} + K w_{l,l'} \ge d_{l'},$$
  
 $\tau_{l'} - \tau_{l} + K(1 - w_{l,l'}) \ge d_{l}$   
and  $w_{l,l'} \le 1.$ 

In turn,  $w_{l,l'} = 1$  implies  $\tau_l - \tau_{l'} + K \ge d_{l'}$  and  $\tau_{l'} \ge \tau_l + d_l$ . The first constraint is always true since K is large and the second constraint implies that  $T_{l'}$  starts after the end of  $T_l$ . The proof for  $w_{l,l'} = 0$  is symmetric and is omitted here.

Then, let us assume that  $T_l$  and  $T_{l'}$  are not assigned to the same resource. Then, by definition,  $z_{l,l'} = 0$  and we obtain

$$\begin{aligned} \tau_l - \tau_{l'} + K(1 + w_{l,l'}) &\geq d_{l'} \\ \tau_{l'} - \tau_l + K(2 - w_{l,l'}) &\geq d_l \\ \text{and } w_{l,l'} &\leq 0. \end{aligned}$$

These three constraints are compatible since the first two are always true independently of the value of  $w_{l,l'}$ , by definition of K, and the last one enforces  $w_{l,l'} = 0$ .

#### 4.1.3 Ordering of Communication Tasks

If tasks  $T_l$  and  $T_{l+1}$  are not processed on the same resource, a communication needs to take place for the output of  $T_l$ . We recall that  $a^{(l)}$  denotes the

data (either an activation or a gradient) computed by  $T_l$  and needed by  $T_{l+1}$ . In order to schedule these communications, we define a new set of variables  $\tilde{z}_{l,l'}$ , which play an analogous role to  $z_{l,l'}$  for communication tasks. More precisely, we want  $\tilde{z}_{l,l'} = 1$  if the communications of  $a^{(l)}$  and  $a^{(l')}$  share the same communication link, and  $\tilde{z}_{l,l'} = 0$  otherwise. We prove that this property is enforced by the following equations, for all  $l \neq l'$ :

$$\widetilde{z}_{l,l'} \ge z_{l,l'} + z_{l+1,l'+1} - z_{l,l+1} - 1 \tag{8}$$

$$\widetilde{z}_{l,l'} \ge z_{l,l'+1} + z_{l+1,l'} - z_{l,l+1} - 1,$$
(9)

$$\begin{aligned} &\tilde{z}_{l,l'} \geq z_{l,l'+1} + z_{l+1,l'+1} - z_{l,l+1} - 1, \\ &\tilde{z}_{l,l'} \geq z_{l,l'+1} + z_{l+1,l'} - z_{l,l+1} - 1, \\ &\tilde{z}_{l,l'} \leq 1 - z_{l,l+1} \end{aligned} \tag{9}$$

$$\widetilde{z}_{l,l'} \le 1 - z_{l',l'+1}$$
 (11)

$$\widetilde{z}_{l,l'} \le z_{l,l'} + z_{l+1,l'}$$
 (12)

$$\widetilde{z}_{l,l'} \le z_{l,l'+1} + z_{l+1,l'+1} \tag{13}$$

*Proof.* Constraints (10) and (11) ensure that  $\tilde{z}_{l,l'} = 0$  if any of  $a^{(l)}$  or  $a^{(l')}$  does not require a communication (because the corresponding tasks are processed on the same resource). In the following, we assume that both  $z_{l,l+1}$  and  $z_{l',l'+1}$  are 0, and we denote by  $P_i$  the processor that runs  $T_l$  and by  $P_j$  the processor which runs  $T_{l+1}$ . We consider several cases:

**First case:**  $a^{(l)}$  and  $a^{(l')}$  share the same communication link.

In that case,  $T_{l'}$  must be processed either on  $P_i$  or  $P_i$ , otherwise the communication of  $a^{(l')}$  occupies another link than  $(P_i, P_j)$ . Therefore, constraint (12) simply becomes  $\tilde{z}_{l,l'} \leq 1$ . Similarly,  $T_{l'+1}$  must be processed either on  $P_i$  or on  $P_j$ , so that constraint (13) simply becomes  $\tilde{z}_{l,l'} \leq 1$ .

Since  $z_{l,l+1} = 0$ , constraints (8) and (9) become

$$\widetilde{z}_{l,l'} \ge z_{l,l'} + z_{l+1,l'+1} - 1$$
  
and  $\widetilde{z}_{l,l'} \ge z_{l,l'+1} + z_{l+1,l'} - 1.$ 

Since the communication of  $a^{(l')}$  use the link between  $P_i$  and  $P_j$ ,  $T_{l'}$  and  $T_{l'+1}$  must be processed on these processors. Hence, either  $T_{l'}$  is on  $P_i$  (and  $T_{l'+1}$  is on  $P_j$ ) or  $T_{l'}$  is on  $P_j$  (and  $T_{l'+1}$  is on  $P_i$ ) and therefore, one of the conditions above enforces  $\tilde{z}_{l,l'} = 1$ .

Second case:  $a^{(l)}$  and  $a^{(l')}$  do not share the same communication link.

Let us first focus on constraint (8). We claim that both  $z_{l,l'}$  and  $z_{l+1,l'+1}$ can not be 1 at the same time. Indeed, this would imply that  $T_l$  and  $T_{l'}$  are processed on  $P_i$ , and also that  $T_{l+1}$  and  $T_{l'+1}$  are processed on  $P_j$ , and thus that  $a^{(l)}$  and  $a^{(l')}$  share the same communication link. Since  $z_{l,l+1} = 0$ , the right hand side of constraint (8) is at most 0. Using the same analysis for constraint (9), we prove that the first two constraints simply become  $\tilde{z}_{l,l'} \geq 0$ .

We now prove by contradiction that at least one among constraints (12) and (13) enforces  $\tilde{z}_{l,l'} = 0$ . Indeed,  $z_{l,l'} + z_{l+1,l'} \ge 1$  implies that  $T_{l'}$  is processed either on  $P_i$  or  $P_j$ , and similarly  $z_{l,l'+1} + z_{l+1,l'+1} \ge 1$  implies that  $T_{l'+1}$  is processed either on  $P_i$  or  $P_j$ . Since we assume that  $z_{l',l'+1} = 0$ , having both  $z_{l,l'} + z_{l+1,l'} \ge 1$  and  $z_{l,l'+1} + z_{l+1,l'+1} \ge 1$  is in contradiction with the fact that  $a^{(l)}$  and  $a^{(l')}$  do not use the same link.

Therefore, in this second case, the system of constraints enforces  $\tilde{z}_{l,l'} = 0$ .

Finally, we can ensure a correct ordering of the communications without overlap in a way similar to Lemma 3. We introduce binary variables  $\widetilde{w}_{l,l'}$  together with the following equations, for all  $l \neq l'$ :

$$\widetilde{\tau}_l - \widetilde{\tau}_{l'} + K(1 - \widetilde{z}_{l,l'} + \widetilde{w}_{l,l'}) \ge \widetilde{d}_{l'}$$
(14)

$$\widetilde{\tau}_{l'} - \widetilde{\tau}_l + K(2 - \widetilde{z}_{l,l'} - \widetilde{w}_{l,l'}) \ge d_l \tag{15}$$

$$\widetilde{w}_{l,l'} \le \widetilde{z}_{l,l'} \tag{16}$$

Lemma 4. Constraints (8)-(16) ensure that:

- $\widetilde{z}_{l,l'} = 1$  if and only if both  $a^{(l)}$  and  $a^{(l')}$  need to be communicated and their communications are assigned to the same link.
- In that case, either the communication of  $a^{(l')}$  starts after the end of the communication of  $a^{(l)}$  (and  $\tilde{w}_{l,l'} = 1$ ), or the communication of  $a^{(l)}$  starts after the end of the communication of  $a^{(l')}$  (and  $\tilde{w}_{l,l'} = 0$ ).
- In the opposite case,  $\widetilde{w}_{l,l'} = 0$ .

*Proof.* The proof is similar to the proof of Lemma 3, replacing  $w_{l,l'}$  by  $\tilde{w}_{l,l'}$  and  $z_{l,l'}$  by  $\tilde{z}_{l,l'}$ , and is therefore omitted here.

#### 4.1.4 Period Length

In order to obtain a valid pattern from the variables defined so far, we use without loss of generality the following conventions: the ending times of all tasks and communications are between 0 and T, and task  $T_1$  starts at time 0:

$$\forall l, \quad 0 \le \tau_l + d_l \le T \tag{17}$$

$$\forall l, \quad 0 \le \tilde{\tau}_l + d_l \le T \tag{18}$$

$$\tau_1 = 0 \tag{19}$$

We cannot specify that all *starting* times should be non-negative: as can be seen on Figure 2, in general the patterns on different processors are not aligned to start at the same time. So in order to ensure that each resource is occupied for a duration at most T, we include the following constraints which state that the distance between the ending time and starting time of two tasks assigned to the same resource is at most T:

$$\forall l \neq l', \quad T \ge \tau_l + d_l - \tau_{l'} - K(1 - z_{l,l'})$$
 (20)

$$\forall l \neq l', \quad T \ge \tilde{\tau}_l + \tilde{d}_l - \tilde{\tau}_{l'} - K(1 - \tilde{z}_{l,l'}) \tag{21}$$

**Lemma 5.** Without considering memory constraints, from any valid pattern according to Definition 1, we can obtain values for all variables T,  $\tau_l$ ,  $z_{l,l'}$ ,  $f_l$ ,  $\tilde{\tau}_l$ ,  $\tilde{z}_{l,l'}$ ,  $w_{l,'}$  and  $\tilde{w}_{l,l'}$  which respect equations (1)-(21), and vice-versa.

*Proof.* If all variables respect the constraints (1)-(21), then Lemmas 2, 3 and 4 ensure that the pattern built from the values of  $\tau_l$  and  $z_{l,l'}$  is a valid pattern. Furthermore, constraint (20) ensures that for any l and l' such that  $z_{l,l'} = 1$ , T is no smaller than  $\tau_l + d_l - \tau_{l'}$  and  $\tau_{l'} + d_{l'} - \tau_l$ , depending on which task starts first. Since a forward task is always allocated to the same resource as the respective backward task (Constraint (4)), all used resources process at least two tasks. The same can be said for communication tasks, which ensures that T is a valid period for the constructed pattern.

Reciprocally, let us consider any valid pattern, and assign values to all the variables according to this pattern. As discussed above, this can be done in a way that respects constraints (17)-(19) without loss of generality. The above lemmas ensure that constraints (1)-(16) are satisfied. Since T is a valid period, constraints (20), (21) and (18) are satisfied for any l and l' such that  $z_{l,l'} = 1$ . Finally, if  $z_{l,l'} = 0$ , these constraints are automatically satisfied since K is large.

#### 4.2 Memory Constraints

In this section, we focus on the memory usage induced by the pattern described in previous section. The memory needs have different origins:

- If two successive tasks  $T_l$  and  $T_{l+1}$  are processed on the same resource, the output of  $T_l$  needs to be stored in memory until it is processed by  $T_{l+1}$ . This is addressed in Section 4.2.1.
- The main point of pipelining is that during one period, the forward task  $T_{L-l}$  and its associated backward task  $T_{L+l+1}$  do not operate on

the same mini-batch. This implies that the processor in charge of these operations must store several activations produced by the forward task and not yet consumed by the corresponding backward task. This will be addressed in Section 4.2.2.

- Processors need to store the weights of the layers that they process. This will be addressed in Section 4.2.3.
- When  $T_{l'-1}$  and  $T_{l'}$  are not processed on the same resource, either a forward activation (if l' < L) or a gradient (if l' > L) are received by the resource in charge of  $T_{l'}$  and are kept in memory until the next  $T_{l'}$  is performed. Similarly, when  $T_{l'}$  and  $T_{l'+1}$  are not processed on the same resource, either a forward activation (if l' < L) or a gradient (if l' > L) must be sent by the resource in charge of  $T_{l'}$  and is kept in memory until the associated communication. This will be addressed in Section 4.2.4.

In order to avoid symmetries in the formulation of the Integer Linear Program, we provide a formulation based on tasks and task collocations rather than on processing resources. We therefore compute, for each task  $T_l$ , the amount of memory required at the instant when  $T_l$  is performed respectively by the storage of models  $M_l^{(MOD)}$ , by direct dependencies  $M_l^{(DIR)}$ , by local activations  $M_l^{(ACT)}$  and by external activations and gradients  $M_l^{(EXT)}$ .

#### 4.2.1 Memory for Direct Dependencies

As depicted on Figure 2, the output  $a^{(l')}$  of a forward task  $T_{l'}$  is used twice: first by the next forward task  $T_{l'+1}$ , then by the corresponding backward task  $T_{2L-l'-1}$ . In this section, we account for the memory consumption of  $a^{(l')}$ from  $T_{l'}$  until  $T_{l'+1}$ ; the memory consumption until the backward task will be accounted for in Section 4.2.2.

To evaluate  $M_l^{(DIR)}$ , let us assume that tasks l, l' and l' + 1 are processed on the same resource. We are interested in the following event: the output produced by  $T_{l'}$  occupies the memory of the resource when task  $T_l$  is performed. This event occurs in three possible situations (see Figure 3):

- $T_{l'}$  is processed before  $T_l$ , and  $T_l$  before  $T_{l'+1}$ ;
- $T_l$  is processed before  $T_{l'+1}$ , and  $T_{l'+1}$  before  $T_{l'}$ ;
- $T_{l'+1}$  is processed before  $T_{l'}$ , and  $T_{l'}$  before  $T_l$ .



Figure 3: Different Cases for direct dependencies, where  $T_l$ ,  $T_{l'}$  and  $T_{l'+1}$  are on the same processor.

We therefore need to consider three variables:  $w_{l',l}$ ,  $w_{l,l'+1}$  and  $w_{l'+1,l'}$ . Since the ordering of execution on the resource is a total order, it is clear that all three variables can not be equal to one, and the list above shows that the event occurs if and only if exactly two of these variables are equal to one. We thus introduce a binary variable  $o_{l,l'}$  for all l and l' with  $l \neq l'$ and  $l \neq l' + 1$ , with the following constraint:

$$o_{l,l'} \ge w_{l',l} + w_{l,l'+1} + w_{l'+1,l'} - 1 \tag{22}$$

We obtain the following lemma:

**Lemma 6.** Consider any valid pattern according to Lemma 5, and assume that variables  $o_{l,l'}$  satisfy Constraint (22).

If the output produced by  $T_{l'}$  is present in memory as a direct dependency when task  $T_l$  is performed, then  $o_{l,l'} \ge 1$ . The total amount of memory that is occupied by direct dependencies is at most  $M_l^{(DIR)} = \sum_{l'=1}^{2L} o_{l,l'} a_{l'}$ .

#### 4.2.2 Memory Required for Local Activations

Let us now consider the memory required by the storage of local activations, between the instant when they are used by a forward task and the instant when they are consumed by the associated backward task. To achieve this, we need to analyze precisely the shifts in indices of the mini-batches that are processed during the same period.

As mentioned in the discussion of Definition 1, we observe that two consecutive tasks in the task graph, either  $(T_l, T_{l,l+1}^c)$  or  $(T_{l,l+1}^c, T_{l+1})$ , can operate on the same mini-batch during a given period if they appear in the proper order, *i.e.*  $T_l$  before  $T_{l,l+1}^c$  or  $T_{l,l+1}^c$  before  $T_{l+1}$ . Otherwise, they must



Figure 4: Example schedule (without communications) with paths of different mini-batches. Black arrows point from the end of a task  $T_l$  to the start of  $T_{l+1}$ , and show the value of the associated  $v_l$  variable.



Figure 5: Example schedule where if task  $T_l$  processes mini-batch *i* during the period,  $T_{l+1}$  needs to process mini-batch i - 2. Tasks indicated with dashed lines belong to different periods (the previous one for  $T_l$ , the next one for  $T_{l+1}$ ).

operate on different mini-batches. An example showing the path of different mini-batches for a simple case of two processors and no communication is shown on Figure 4.

When two successive tasks are too far apart in the pattern, it can even happen (in rare cases) that they have to process mini-batches with an index shift of two. Figure 5 shows such a case, which happens if and only if the difference between the end time of  $T_l$  and the start time of  $T_{l+1}$  is more than T.

To evaluate the shifts of indices in the pattern, we introduce new boolean variables associated to task l:  $v_l$  and  $v'_l$  are used to determine the shift between  $T_l$  and  $T^c_{l,l+1}$ , where  $v_l$  is 1 if the shift is at least 1, and  $v'_l$  is 1 if the shift is 2. Variables  $\tilde{v}_l$  and  $\tilde{v}'_l$  have the same meaning for the shift between

 $T_{l,l+1}^c$  and  $T_{l+1}$ . For all l, we include the following constraints:

$$\tau_l + d_l - \widetilde{\tau}_l + K(1 - v_l) \ge 0 \tag{23}$$

$$\widetilde{\tau}_l - (\tau_l + d_l) + K v_l \ge 0 \tag{24}$$

$$\widetilde{\tau}_l + d_l - \tau_{l+1} + K(1 - \widetilde{v}_l) \ge 0 \tag{25}$$

$$\tau_{l+1} - (\widetilde{\tau}_l + \widetilde{d}_l) + K \widetilde{v}_l \ge 0 \tag{26}$$

$$\widetilde{\tau}_l + T - (\tau_l + d_l) + 2Kv_l' \ge 0 \tag{27}$$

$$\tau_l + d_l - (\tilde{\tau}_l + T) + 2K(1 - v'_l) \ge 0$$
 (28)

$$\tau_{l+1} + T - (\widetilde{\tau}_l + d_l) + 2K\widetilde{v}_l' \ge 0 \tag{29}$$

$$\widetilde{\tau}_l + d_l - (\tau_{l+1} + T) + 2K(1 - \widetilde{\upsilon}'_l) \ge 0$$
(30)

**Lemma 7.** Consider any valid pattern according to Lemma 5, and assume that variables  $v_l$ ,  $v'_l$ ,  $\tilde{v}_l$  and  $\tilde{v}'_l$  satisfy Constraints (23)-(30).

If i denotes the mini-batch performed by  $T_L$ , then for any  $0 \le j \le L-1$ , the index of the mini-batch performed by  $T_{L-j}$  is at least  $i + \sum_{l=L-j}^{L-1} (v_l + \widetilde{v}_l + v'_l + \widetilde{v}'_l)$ , and the index of the mini-batch performed by  $T_{L+j+1}$  is at most  $i - \sum_{l=L}^{L+j} (v_l + \widetilde{v}_l + v'_l + \widetilde{v}'_l)$ .

Hence, the number of activations of type  $a^{(L-j-1)}$  that needs to stored at the beginning of the period of this processor is  $\sum_{l=L-j}^{L+j} v_l + \widetilde{v}_l + v'_l + \widetilde{v}'_l$ .

*Proof.* Similarly to Lemma 3, we can show using the definition of K proposed above that constraints (23)-(26) ensure that for any l,  $v_l = 0$  if  $\tau_l + d_l \leq \tilde{\tau}_l$ , and  $v_l = 1$  otherwise; likewise,  $\tilde{v}_l = 0$  if  $\tilde{\tau}_l + \tilde{d}_l \leq \tau_{l+1}$ , and  $\tilde{v}_l = 1$  otherwise. Additionally, constraints (27)-(30) ensure that if  $\tilde{\tau}_l + T < \tau_l + d_l$ , then  $\tilde{v}_l = 1$ , and  $\tilde{v}_l = 0$  otherwise; likewise if  $\tau_{l+1} + T < \tilde{\tau}_l + \tilde{d}_l$ , then  $\tilde{v}'_l = 1$ , and  $\tilde{v}'_l = 0$ otherwise.

The claimed result can be proved by induction. For any  $0 \le j \le L-1$ , let us set k = L - j - 1 and let us assume that the index of the mini-batch performed by  $T_{k+1=L-j}$  is at least  $I = i + \sum_{l=L-j}^{L-1} v_l + \tilde{v}_l + v'_l + \tilde{v}'_l$ . Let us now consider  $T_{k=L-j-1}$ .

The ordering of  $\tilde{\tau}_k$  and  $\tau_{k+1}$  can yield three possible cases:

- If  $\tilde{\tau}_k + \tilde{d}_k \leq \tau_{k+1}$ , then  $\tilde{v}_k = \tilde{v}'_k = 0$  and both computation and communication tasks can process the same mini-batch in the same period: the communication can process mini-batch I.
- If  $\tilde{\tau}_k + \tilde{d}_k > \tau_{k+1}$  and  $\tilde{\tau}_k + \tilde{d}_k \leq \tau_{k+1} + T$ , then  $\tilde{v}_k = 1$  and  $\tilde{v}'_k = 0$ . In this case, similar to the one shown on Figure 4 with tasks  $T_1$  and  $T_2$ , the communication cannot process mini-batch *I*: if it does, the

result arrives too late and task  $T_{k+1}$  is not able to process mini-batch I. However the communication can process mini-batch I + 1.

• If  $\tilde{\tau}_k + \tilde{d}_k > \tau_{k+1} + T$ , then  $\tilde{v}_k = 1$  and  $\tilde{v}'_k = 1$ . In that case, similar to the one shown on Figure 5, the communication can only process mini-batch I + 2.

Therefore, in all cases the index of the mini-batch processed by the communication is  $I + \tilde{v}_k + \tilde{v}'_k$ .

The same reasoning can be applied to the possible shift between the index of the mini-batch corresponding to  $T_k$  and the communication of this activation, this time involving  $v_k$  and  $v'_k$ . We thus prove that the index of the mini-batch performed by  $T_k$  during the current period is at least  $I + \tilde{v}_k + \tilde{v}'_k + v_k + v'_k = i + \sum_{l=L-j-1}^{L-1} (v_l + \tilde{v}_l + v'_l + \tilde{v}'_l)$ , which achieves the proof for forward tasks.

The proof for backward tasks is very similar and is omitted here.  $\Box$ 

As mentioned above, we are interested in  $M_l^{(ACT)}$ , which is the memory consumed by the set of activations at the time when task  $T_l$  is performed, on the processor which computes  $T_l$ . We thus introduce integer variables  $\sigma_{l,'l}$ , equal to the number of activations of type  $a^{(l'-1)}$  stored on the processor that computes  $T_l$ , which satisfy the following constraints:

$$\forall l', l \ \sigma_{l,l'} \le 8Lz_{l,l'} \tag{31}$$

$$\forall l', l \ \sigma_{l,l'} \ge \sum_{m=l'}^{2L-l'} (v_m + v'_m + \widetilde{v}_m + \widetilde{v}'_m) - 8L(1 - z_{l,l'})$$
(32)

Since Lemma 7 only provides the number of activations at the beginning of the period, we also need to account for the following events which may take place between the beginning of the period and instant  $\tau_l$  (i) a forward task  $T_{l'}$ ,  $l' \leq L$  is computed, inducing an extra activation  $a^{(l'-1)}$  in memory, and (ii) a backward task  $T_{2L-l'+1}$ ,  $l' \leq L$  is computed, removing an activation  $a^{(l'-1)}$  from memory.

**Lemma 8.** Consider any valid pattern according to Lemma 5, satisfying Constraints (23)-(32).

The amount of memory occupied when task  $T_l$  is performed by activations required by future backward tasks is  $M_l^{(ACT)}$ :

$$\mathbf{M}_{l}^{(\text{ACT})} = a_{l-1} + a_{l} + \sum_{l'=1}^{L} (\sigma_{l,l'} + w_{l',l} - w_{2L-l'+1,l}) a_{l'-1}.$$

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*Proof.* We first show that  $\sigma_{l,l'}$  is at least the number of replicas for a layer l' derived in Lemma 7 if l' is on the same processor as l, otherwise  $\sigma_{l,l'} = 0$ . Indeed, if  $T_l$  and  $T_{l'}$  share the same resource then  $z_{l',l} = 1$  and since the number of replicas is less than 8L for any layer,  $\sigma_{l,l'} \geq \sum_{m=l'}^{2L-l'} (v_m + v'_m + \tilde{v}_m + \tilde{v}'_m)$ . On the other hand, when  $T_l$  and  $T_{l'}$  are on different resources  $z_{l',l} = 0$  and  $\sigma_{l',l} \leq 0$ .

According to Lemma 7 the value  $\sum_{m=l'}^{2L-l'} (v_m + v'_m + \tilde{v}_m + \tilde{v}'_m)$  represents the number of activations of layer  $a^{(l'-1)}$  stored in the beginning of the period, but this number may vary within the period: it increases by one after each task  $T_{l'}$  ( $F_{l'}$ ) and it decreases by one after task  $T_{2L-l'+1}$  ( $B_{l'}$ ). Thus, the last term in the equation for  $M_l^{(ACT)}$  corresponds to all activations that have been stored before task  $T_l$  and the rest represents the memory needed to perform task  $T_l$ .

#### 4.2.3 Memory Required for the Models

As it was shown in [32], having just two models stored is enough to perform training. Besides, since the computed gradients have the same shape as model weights, they require the same amount of memory. Thus, the equivalent of three copies of the weights are necessary to perform all forward and backward computations with consistent weights. Thus

$$\mathbf{M}_{l}^{(\text{MOD})} = 3 \sum_{l', l' \le L} z_{l, l'} W_{l'}.$$
(33)

#### 4.2.4 Memory Buffer for Communications

Another type of memory usage on a resource  $P_i$  is the buffer memory to store activations or gradients: *incoming* ones, that were computed by another processor and then sent to  $P_i$ , or *outgoing* ones, that were computed on  $P_i$ and need to be sent. We assume that a buffer is allocated to each of these data for the whole duration of the execution, but they are not shared between different data. We introduce binary variables  $b_{l,l'}$  for all l and l', together with the following constraints:

$$b_{l,l'} \ge z_{l,l'} - z_{l,l'+1} \tag{34}$$

$$b_{l,l'} \ge z_{l,l'+1} - z_{l,l'} \tag{35}$$

**Lemma 9.** Consider any valid pattern according to Lemma 5, satisfying Constraints (34) and (35). If a buffer is required for  $a^{(l')}$  on the processor

which computes  $T_l$ , then  $b_{l,l'} = 1$ . Hence, the memory reserved for buffers at the start of task  $T_l$  is at most  $M_l^{(BUF)} = \sum_{l'=1}^{2L} b_{l,l'} a_{l'}$ .

#### 4.2.5 Final Linear Program

We can now define the complete Linear Program for our problem: the objective is to minimize T, subject to Contraints (1)-(35), together with

$$\forall l, \quad \mathbf{M}_{l}^{(\text{MOD})} + \mathbf{M}_{l}^{(\text{ACT})} + \mathbf{M}_{l}^{(\text{DIR})} + \mathbf{M}_{l}^{(\text{BUF})} \leq M$$

$$\forall l, \quad T, \tau_{l}, \tilde{\tau}_{l} \in \mathbb{R}$$

$$\forall l, l' \quad \sigma_{l,l'} \in \mathbb{Z}$$

$$\forall l, \quad f_{l}, v_{l}, v_{l}', \tilde{v}_{l}, \tilde{v}_{l}' \in \{0, 1\}$$

$$\forall l, l', \quad z_{l,l'}, w_{l,l'}, \tilde{z}_{l,l'}, w_{l,l'}, o_{l,l'}, b_{l,l'} \in \{0, 1\}$$

$$(36)$$

**Theorem 1.** The optimal solution of the above Integer Linear Program provides a valid pattern with minimum period, among those whose memory usage is at most M.

# 5 Experimental Results

In this section, we present simulation results obtained for different stateof-the-art **ResNet** neural networks of size 18, 34 and 50, which are widely used for a large range of tasks. In order to perform these simulations, we first perform the profiling of the neural networks to measure the durations and memory costs of the different operations involved in the training. As mentioned in Section 3, this work only considers networks in the shape of adjoint chains as depicted in Figure 1. In the case of **ResNet** networks, a simple linearization approach is enough to transform the neural network computational graphs into chains, by applying a greedy procedure to obtain minimal groups of operations. In each group, the set of predecessors outside the group and the set of successors outside the group are disjoint. Overall, such groups form a sequence of operations with a straightforward order of execution. This approach was used as well in [5].

We implemented the ILP from Section 4 using the CPLEX solver [33]. In all our experiments, the execution time was limited to one hour. In case there is still a gap after one hour, we keep the best current solution computed by the solver. For ResNet-18 up to ResNet-50, the solutions produced were of very good quality (see the discussion below), though the solver was unable to prove its optimality. The results obtained are therefore heuristic in nature. This time limit is reasonable, because the computed solution can be used during the entire training phase associated with a given image and mini-batch sizes, a given computing platform and a given network. It is common for the training to last several hours/days on a parallel platform, which makes this approach acceptable.

To evaluate the quality of the solutions produced by the integer linear program, we compare the results obtained with those of PipeDream [31], which is the state of the art solution for pipelined model parallelism. In practice, PipeDream takes as input the memory limit and the characteristics of the platform, computes the number of batches to be inserted in the pipeline, called NOAM, and finds a partitioning of the network that is used for model parallelism. PipeDream then uses a greedy 1F1B strategy to schedule tasks. In practice, as we have pointed out, since the memory model used to compute the partitioning is over simplified, the calculated NOAM value is generally not optimal either.

Nevertheless, despite these limitations, PipeDream can be used to produce a large number of solutions from which one can then build valid solutions that fulfill the memory constraints. This is the approach we use in the following experiments. For a fairly large number of possible memory targets and possible NOAM values, we produce the allocations computed by PipeDream, simulate the execution of the eager scheduling strategy for 500 batches, and we evaluate a posteriori the actual memory consumption and the period (the inverse of the asymptotic throughput) that can be obtained. We thus obtain a set of (memory, period) pairs that correspond to feasible solutions. Our observations indicate that the solution produced by PipeDream generally consumes much more memory than the target value. Nevertheless, since the execution time of PipeDream is small, obtaining a set of good valid solutions through this "exhaustive" approach is still practical. In Figures 6 to 11, blue dots correspond to the actual memory consumption and observed period of solutions computed with this approach.

Figures 6 to 11 correspond respectively to the networks ResNet-18 (depth 18, with a batch size of 8 and image size 1000), ResNet-34 (depth 34, with a batch size of 32 and image size 224) and ResNet-50 (depth 50, with a batch size of 8 and image size 224), for the case of 4 or 8 GPUs. The red dots correspond to the best solutions found by CPLEX for our ILP after one hour, for different values of the memory limit.

First of all, it can be observed that the solutions returned by our ILP are almost always better than the solutions returned by the exhaustive approach based on PipeDream, even if optimality cannot be guaranteed. The only exception can be seen on Figure 8, where for instance in the case of a memory



Figure 6: Results with ResNet-18 and 4 processors



Figure 7: Results with ResNet-18 and 8 processors

size of 1 GB, PipeDream finds solutions that strictly dominate the one returned by the linear program. It can also be observed that the ILP is able to find better solutions both in cases where memory is scarce (Figures 7 and 11) and where memory is abundant (Figures 6 and 9). When memory is abundant, the ability of the ILP to use non-contiguous partitionings of the networks allows to use this abundant memory to achieve better load-balancing. When memory is scarce, the precise scheduling formulation of the ILP allows to obtain better solutions by reducing memory costs. The solutions produced by the ILP are therefore of very high quality when the size of the network



Figure 8: Results with ResNet-34 and 4 processors



Figure 9: Results with ResNet-34 and 8 processors

is not too large. On larger networks such as ResNet-101 or DenseNet-121 (of respective depths 101 and 121), one hour of execution is sometimes not enough for the ILP to find integral solutions of good quality. In this case, it would be necessary to consider other approaches, using fractional relaxations of the linear program or decoupling partition and scheduling phases. We leave these ideas for future works.



Figure 10: Results with ResNet-50 and 4 processors



Figure 11: Results with ResNet-50 and 8 processors

# 6 Conclusion

In this paper, we consider the possibility of applying model parallelism, which is an attractive parallelization strategy that allows in particular not to replicate all the weights of the network on all the computation resources. Following the ideas proposed in PipeDream [31] we propose to combine pipelining and model parallelism, which allows to obtain a better resource utilization. Then, model parallelism can be combined with data parallelism to improve scalability.

Nevertheless, the combination of pipelining and model parallelism requires to store more activations at the nodes, which in turn causes memory consumption problems. We propose a very fine analysis of the memory costs induced by this combination. We use this modeling first to establish the computational complexity of the problems related to memory constrained pipelined model parallelism. This model also allows us to find a partition of the network that explicitly takes memory costs into account, contrary to what is done in PipeDream. We show that it is possible to formalize the problem of memory constrained throughput optimization as an integer linear program.

Through experiments on medium size networks (Resnet-18 to ResNet-50), we prove that the ILP is able to compute in reasonable time solutions that are better than those computed by PipeDream, by both providing good partitions of the networks and good scheduling strategies. Nevertheless, the computing cost induced by the integer programming approach becomes too large for very deep networks, and therefore, new heuristic solutions are required in this case, which opens interesting perspectives to this work.

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