Taming Unbalanced Training Workloads in Deep Learning with Partial Collective Operations

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Abstract
Load imbalance pervasively exists in distributed deep learning training systems, either caused by the inherent imbalance in learned tasks or by the system itself. Traditional synchronous Stochastic Gradient Descent (SGD) achieves good accuracy for a wide variety of tasks, but relies on global synchronization to accumulate the gradients at every training step. In this paper, we propose eager-SGD, which relaxes the global synchronization for decentralized accumulation. To implement eager-SGD, we propose to use two partial collectives: solo and majority. With solo allreduce, the faster processes contribute their gradients eagerly without waiting for the slower processes, whereas with majority allreduce, at least half of the participants must contribute gradients before continuing, all without using a central parameter server. We theoretically prove the convergence of the algorithms and describe the partial collectives in detail. Experiments are conducted on a variety of neural networks and datasets. The results on load-imbalanced environments show that eager-SGD achieves 2.64× speedup (ResNet-50 on ImageNet) over the asynchronous centralized SGD, and achieves 1.29× speedup (ResNet-50 on ImageNet) and 1.27× speedup (LSTM on UCF101) over the state-of-the-art synchronous decentralized SGDs, without losing accuracy.

CCS Concepts • Theory of computation → Parallel algorithms; • Computing methodologies → Neural networks;

Keywords stochastic gradient descent, distributed deep learning, eager-SGD, workload imbalance, collective operations

1 Motivation
Deep learning models are on a steep trajectory to becoming the most important workload on parallel and distributed computer systems. Early convolutional networks demonstrated groundbreaking successes in computer vision, ranging from image classification to object detection [30, 52]. More recent developments in recurrent and transformer networks enable impressive results in video classification, natural language processing for machine translation, question answering, text comprehension, and synthetic text generation. The latter models contain more than 1.5 billion parameters and take weeks to train [15, 45]. Other demanding neural networks are trained on the largest supercomputers to achieve scientific breakthroughs [35, 41]. Furthermore, the models are growing exponentially in size, OpenAI is predicting a 10x growth each year [3] potentially leading to artificial general intelligence. In order to support this development, optimizing the training procedure is most important.

The training procedure of deep learning is highly parallel but dominated by communication [10]. Most parallel training schemes use data parallelism where full models are trained with parts of the dataset and parameters are synchronized at the end of each iteration. The total size of allreduce grows with the model size, which ranges from a few megabytes [30] to several gigabytes [45] and grows quickly. The allreduce operation is not atomic and it can be split into layer-wise reductions, which can easily be overlapped with the layer computation using non-blocking collectives [5, 25]. Yet, the optimal scaling of an allreduce of size $S$ is at best $O(\log P + S)$ in $P$ processes [26, 43, 47]. Thus, growing process counts will reduce the parallel efficiency and eventually make the reduction a scaling bottleneck.

The communication aspects of deep learning have been investigated in many different contexts [47, 51], see the survey for an overview [10]. In this work, we identify load imbalance as an additional barrier to scalability. When some processes finish the computation later than others, all processes will wait for the last one at the blocking allreduce function. Load imbalance can be caused by the system itself, for example, when training on multi-tenant cloud systems [31, 32, 49] or by system or network noise [27, 28] in high-performance machines. A second, and more prominent cause of imbalance is inherent imbalance in the computation that causes varying load across different processes. While noise from the system is generally low on well-maintained HPC machines [28], the inherent load imbalance of the training workloads cannot
Our main contributions are:

- A detailed analysis of workload imbalance in deep learning training.
- Definition and implementation of partial collectives, specifically majority and solo allreduce.
- Eager-SGD for asynchronous decentralized distributed training of neural networks with proof of convergence.
- An experimental study of convergence and training speed for multiple networks, achieving 1.27× speedup over synchronous SGD on a video classification task without losing accuracy.

2 Load-Imbalance in Deep Learning

Load imbalance widely exists in the training of deep learning models, which can be caused by either the applications or the system itself [27, 28, 31, 32, 49].

2.1 Video Processing

Long short-term memory (LSTM) [23] is a type of unit cell in Recurrent Neural Networks (RNN). In video classification tasks, LSTMs are used [7, 18, 60] to process a sequence of frames for a video as input (optionally following convolutional neural networks that preprocess the images to features), and output a probability distribution over a set of classes. Due to the recurrent structure of the network, the computational overhead is proportional to the number of frames in the input video.

Fig. 2a shows the video length distribution (the number of frames) over all 9,537 videos in the training dataset of UCF101 [53]. The video length is distributed between 29 and 1,776 frames, with a mean frame count of 187 and standard deviation of 97. Fig. 2b shows the runtime distribution over the 1,192 sampled batches in two epochs to train a 2,048-wide single-layer LSTM model on video frame features. As is standard in variable-length training, videos with similar lengths are grouped into buckets for performance. The runtime is distributed from 201 ms to 3,410 ms, with a mean runtime of 1,235 ms and standard deviation of 706 ms. These statistics above show that training an LSTM model for video classification exhibits inherent load imbalance.

2.2 Language Processing

Transformers [57] are sequence-to-sequence models that translate a sequence of words from one language to another. Different from RNN, a Transformer network replaces the recurrent structure with an attention mechanism. To train the Transformer model, the computation overhead increases with the length of the input (and output) sentences. Typically, the sentences in the training dataset for a language model have various lengths, and thus the workload is unbalanced across different batches. Fig. 3 shows the runtime distribution over the 20,653 randomly sampled batches in 1/3 epoch to train a Transformer on the WMT16 dataset. The runtime is
model [21] on ImageNet [14], on a standard Google Cloud instance (n1-standard-16 with 2x Nvidia V100 GPUs). The runtime is distributed from 399 ms to 1,892 ms with a mean of 454 ms and standard deviation of 116 ms. Since ResNet-50 on ImageNet has the same input size for different batches, the load imbalance is caused mainly by the system. Compared with imbalanced applications (e.g., Transformer, LSTM), the load imbalance on cloud servers is relatively light.

3 Distributed Deep Learning

Deep neural networks are continuously differentiable functions that are composed of multiple operators, representable by a directed acyclic graph [36]. The gradient of those functions can be computed using the backpropagation algorithm [37], processing the nodes in the DAG in a reverse topological order. Deep learning frameworks, such as TensorFlow [1], typically execute parallel operations in the DAG in arbitrary order.

Algorithm 1 Distributed Minibatch SGD

```
1: for \( t = 0 \) to \( T \) do
2: \( \bar{x}, \bar{y} \leftarrow \) Sample \( B \) elements from dataset
3: \( w_t \leftarrow \) Obtain parameters from global view
4: \( \bar{z} \leftarrow f(w_t, \bar{x}, \bar{y}) \)
5: \( G_t \leftarrow \frac{1}{i=1} \nabla f(w_t, \bar{z}) \)
6: \( \Delta w \leftarrow U(G_t, w_0, \ldots, w_T, t) \)
7: Update global view of parameters to \( w_t + \Delta w \)
8: end for
```

Supervised deep neural network training typically involves first-order optimization in the form of Stochastic Gradient Descent (SGD) [48]. SGD optimizes the expected loss value over the “true” distribution of input samples by descending in the direction of a random subset of the training samples (minibatch). In a distributed data-parallel setting, the SGD algorithm (Algorithm 1) consists of multiple learner processes, each of which updates a global view of the parameters \( w \) according to a different random minibatch at the same time. Given an update rule \( U \) and local minibatch of size \( B \), the learners modify the global view of the parameters by using an average of the gradients \( G_t \) obtained by the agents.

A straightforward manner to maintain a global view is using a Parameter Server (PS) architecture [13], where one or several nodes assume the role of a PS, broadcasting up-to-date weights (line 3) to learners prior to each step and aggregating gradients from them (line 7). This enables the PS to asynchronously update the global view [46], or require a fraction of learners to send gradients before progressing to the next step [22].

As the PS model is generally not scalable, another mode of operation implements SGD using collective operations. In such implementations, accumulating the gradients (line 7) is done via an allreduce operation, where each learner
contains its own local view of the weights [10]. Horovod [51] is one such implementation over the TensorFlow framework, which also fuses several allreduce operations into one in order to reduce overhead. However, due to the arbitrary order of execution imposed by the frameworks, Horovod uses a master process for negotiation communication (achieving consensus on which parameters are sent).

A more scalable method, used in the Deep500 DSGD optimizer [9], is to ensure an order of communication execution by adding control dependencies into the computation DAG, as shown in Fig. 5. In the backward pass, the allreduce operations are executed in a specific order after finishing the local gradient computation. We use the same method when implementing eager-SGD. Note that synchronizing gradient order can be avoided completely using non-blocking collectives [42]. In this mode, each gradient communication message is assigned to an agreed-upon numeric tag, and multiple allreduce operations may be in-flight concurrently. Prior to updating the local view of the weights, a waitall command must be issued. All in all, these approaches reduce overhead in imbalanced loads by overlapping communication and computation, but do not mitigate it completely.

![Figure 5. Adding control dependency in the computation DAG, using a block of ResNet-50 as an example.](image)

4 Partial Collective Operations

A collective communication involves a set of processes that cooperate to progress their internal state. Some of these operations, e.g., allreduce, implicitly synchronize the participants: the operation cannot terminate before the slowest process joins it. We define these collectives as synchronous and introduce a new class of partial collectives that relax the synchronization. We now discuss two variants of partial collectives: solo and majority.

4.1 Solo Collectives

A solo collective [17] is a wait-free operation, which forces the slow processes to execute the collective as soon as there is one process executing it. This process, called initiator, is in charge of informing the others to join the collective. While solo collectives remove the synchronization delays, they change the semantics of collective operations, which may now be completed by using stale data from the slow processes.

4.1.1 Schedule Activation

We define a schedule as a set of operations that a process executes in order to globally progress the collective operation. In particular, a schedule is a directed acyclic graph (DAG) where the vertices are operations and the edges are happens-before dependencies among them. We define the following operations:

- Point-to-point communications: sends and receives.
- Computations: simple computations defined between two arrays of data items. The type of the data items is defined according to the MPI basic types [42].
- Non-operations (NOP): complete immediately and are only used to build dependencies.

Operations can be dependent on zero, one, or more other operations (with and or logic) of the same schedule.

The main difference between synchronous and solo collectives is the time at which processes activate (i.e., starts executing) their schedule. For synchronous collectives, the schedule is executed only when a process reaches the collective function call (e.g., MPI_Allreduce). We define this activation as internal. For solo collectives, an external activation is also possible: the processes start executing the schedule because of an activation message received from the initiator, which starts broadcasting it immediately after the internal activation of its schedule. In particular, a solo collective is composed of two schedules: one for broadcasting the activation and the other one for executing the collective operation.

In a solo collective, any process can become the initiator, hence any process must be capable of broadcasting the activation message. The activation broadcast is implemented as a modified version of the recursive doubling communication scheme: this is equivalent to the union of \( P \) binomial trees (optimal for small message broadcast, like the activation) rooted at the different nodes.
**Activation example**  Fig. 6 shows a solo allreduce example. On the left, we show the global communications view that is split in two phases: activation and allreduce. The highlighted communication shows the activation path if the initiator is, e.g., process P3. For the allreduce, we use a recursive doubling implementation. Note that any collective implementation that can be expressed as a schedule can be linked to the activation phase. On the right we show the internal schedule of process P3. An internal activation (i.e., P3 making the function call explicitly) translates in the execution of NOP 0 (N0): this leads to the send operations S0 and S1 being fired to start broadcasting the activation message and to the execution of N1, which signals the activation of the allreduce schedule. Alternatively, if P3 is not the initiator, it will receive a message in receive R0 or R1: if the activation is received by R0, then P3 has to forward the activation message to P1 with send S1 (i.e., P3 is an internal node of the activation binomial tree). Also in this case NOP N1 will be executed, leading to the execution of the allreduce schedule.

**Multiple initiators**  Multiple processes may join the collective at the same time: in this case we need to ensure that the collective is executed only once. To address this issue, we set the operations to be consumable, meaning that the same operation cannot be executed twice. For example, let us assume that nodes P2 and P3 reach their internal activation at the same time. When P3 receives the activation message from P2 (i.e., through R0) there are two possible cases: 1) S1 is still not consumed and then it is executed; 2) S1 has been fired due to the internal activation and will not be executed a second time. NOPs are also consumable, hence N1 (i.e., the activation) can be executed only once.

**Persistent schedules**  Processes can be asked to join a solo collective only once before they reach their internal activation: once the schedule is executed, it needs to be re-created by the application in order to be executed again. To enable multiple asynchronous executions of solo collectives, we introduce persistent schedules. Such schedules transparently replicate themselves once executed, able to serve a new solo collective without requiring application intervention. Multiple executions of the same solo collective overwrite the data in the receive buffer, which always contains the value of the latest execution.

4.2 Majority Collectives

An issue of solo collectives is that if one or few processes become the initiator, as in solo collectives. Instead, at each execution of a persistent schedule, the processes designate an initiator by randomly selecting a rank (consensus is achieved by using the same seed for all the processes). When a process joins the collective (i.e., internal activation), it checks whether it is the designated initiator: only in that case it keeps running the internal activation followed by the actual collective schedule.

We now discuss how the above described implementation can provide a statistical guarantee that at least half of the processes on average contribute to the collective. Suppose the same collective operation is called by many iterations, such as in model training. We sort all the P processes by the time they reach a collective operation. Since the probability that any process is specified as the initiator is equal to 1/P, the expectation of the randomly specified initiator is the P/2-th process among the sorted processes, namely on average half of the processes reach the collective operation earlier than the initiator. For a workload distribution with one mode and a tail, such as in Figs. 2, 3, and 4, the probability that part of the processes reach the collective at a similar time to the initiator is high; then, more than half of the processes on average actively participate in the operation.

4.3 Asynchronous Execution by Library Offloading

The schedule of a partial collective can be asynchronously executed with respect to the application. We develop fflib2, a communication library that allows to express communication schedules and offload their execution to the library itself. The schedule execution can take place on the application thread (i.e., when the application enters the library), or on an auxiliary thread. Once the application creates and commits a schedule, the library starts executing all the operations that have no dependencies. The remaining ones are executed as their dependencies are satisfied.

4.4 Discussion

Offloading the schedule execution to the network interface card (NIC) can provide different advantages such as asynchronous execution, lower latency, and streaming processing. Di Girolamo et al. [17] show how solo collectives can be offloaded to Portals 4 [8] NICs by using triggered operations. This approach is limited by the amount of NIC resources that bounds the number of times a persistent schedule can be executed without application intervention. This limit can be removed by implementing the schedule execution with the sPIN programming model [24], which allows to execute user-defined code on the NIC. A sPIN implementation of fflib2 would then be able to replicate the schedule on-the-fly upon completion.
Algorithm 2 Eager-SGD

1: $b$ is local batchsize for $P$ processes
2: for $t = 0$ to $T$ do
3:   $\hat{x}, \hat{y} \leftarrow$ Each process samples $b$ elements from dataset
4:   $\hat{z} \leftarrow \ell (w_t, \hat{x}, \hat{y})$
5:   $G_{t}^{\text{local}} \leftarrow \frac{1}{P} \sum_{p=0}^{P} \nabla \ell (w_t, \hat{z})$
6:   $G_{t}^{\text{global}} \leftarrow \frac{1}{P} \partial$allreduce$\left(G_{t}^{\text{local}}\right)$
7:   $\Delta w \leftarrow U\left(G_{t}^{\text{global}}, w_{t-1}\right)$
8:   $w_{t+1} \leftarrow w_t + \Delta w$
9: end for

5 Eager-SGD algorithm

Algorithm 2 illustrates the main procedure of eager-SGD. Instead of calling a synchronous allreduce in the distributed optimizer (Fig. 5) to accumulate the gradients, eager-SGD uses the partial allreduce operations (Line 7). Either solo or majority allreduce can be used depending on the severity of load imbalance.

Fig. 7 presents an example of how eager-SGD works with partial collectives, in which $w_{t}^{p}$ and $G_{t}^{p}$ represent the weights and the gradients calculated on process $p$ at training step $t$, respectively, and $U \left(G, w\right)$ represents the update rule. In step $t$, suppose process P1 is faster than process P0. P1 finishes the computation of $G_{t}^{1}$ and then triggers the partial allreduce operation. Since P0 does not finish the computation of $G_{t}^{0}$ at this time, it only passively contributes null gradients $G_{null}$ to the partial allreduce at step $t$. After P0 finishes the computation of $G_{t}^{0}$, it finds out that the partial allreduce at step $t$ is already finished by checking the results in the receive buffer. P0 updates the weights of step $t + 1$ using $G_{t}^{1}$ stored in the receive buffer of the partial allreduce and $G_{0}^{0}$ becomes the stale gradients. The stale gradient $G_{t}^{0}$ is then stored in the send buffer. If P0 does not catch up with P1 at step $t + 1$, P0 will passively participate in the partial allreduce again and contribute $G_{t}^{0}$. If P0 catches up with P1 at step $t + 1$ (as in the case shown in Fig. 7), P0 will add $G_{t}^{1}$ and $G_{t+1}^{1}$ (calculated in step $t + 1$ together, and contribute the accumulated gradients $G_{t+1}^{1}$ to the partial allreduce; P0 resets the send buffer to $G_{null}$ after finishing allreduce.

In severe load imbalance situations, some slower processes may lag behind by more than one step. The data in the receive buffer of the partial allreduce will then be overwritten and only the latest data in the receive buffer can be seen, which results in different weights on different processes. This may result in slightly lower accuracy as shown in Section 7.2.2. Thus, we periodically synchronize the models across all processes to eliminate the side effect. Since we only synchronize the models every few epochs, the overhead can be ignored.

6 Correctness and Convergence Guarantees

6.1 System Model

We prove that, under a reasonable set of modeling assumptions, the eager-SGD algorithm will still converge. We assume a system with $P$ asynchronous processors indexed as $i \in \{0, 1, \ldots, P-1\}$, which take steps at different speeds.

For simplicity, we break down the execution at each processor into steps: at step $t$, we assume that each processor $i$ has collected a local view of the parameters, which we denote by $w_{i}^{t}$. We then proceed as follows: the processor computes the gradient $G_{t}^{i}$ on a randomly sampled mini-batch, with respect to the local view $w_{i}^{t}$, and enters an partial-allreduce for the step, whose goal is to attempt to communicate its current parameter updates to other processors. At the end of this, the process obtains its next view of the parameters $w_{t+1}^{i}$, which it will use in the following step $t + 1$.

From a global perspective, we can split the execution in serial fashion into rounds, where each round can be mapped to the partial-allreduce of corresponding index. Without loss of generality, we assume that each processor participates in each round $t$, since it eventually submits an update to the corresponding partial-allreduce, which we denote by ADS($t$), for asynchronous distributed sum. However, its update may or may not be delivered to the other processors. Each partial-allreduce has the following semantics:

- **Invocation** Each process $i$ proposes a $d$-dimensional vector $R_{t}$, corresponding to its current proposed update, to ADS($t$).
- **Response** Each process $i$ receives a tuple $(U_{t}, s_{t}^{i})$, where $U_{t}$ is the $d$-dimensional update to the parameter set corresponding to round $t$, as decided by the shared object ADS($t$), and $s_{t}^{i}$ is a boolean stating whether the update by process $i$ has been included in $U_{t}$.
We can therefore rephrase the algorithm as having each process invoke the ADS\(t\) object in each round, with its current update. If its update is not "accepted" (\(s_i^t = \text{false}\)) then the processor simply adds it to its update in the next iteration. The ADS objects we implement provide the following guarantees.

**Lemma 6.1.** Each ADS object ensures the following:

1. **(Liveness)** The ADS\(t\) object eventually returns an output at every invoking process.
2. **(Safety)** The output is consistent, in the sense that (1) it is a correct average of a subset of the proposed updates in the round; (2) the returned bits reflect its composition; and (3) the output is the same at every invoking process.
3. **(Quorum Size)** The subset of proposed updates included in the output is of size \(Q \geq 1\), where \(Q\) is a lower bound parameter ensured by the algorithm.
4. **(Staleness Bound)** There exists a bounded parameter \(\tau\) such that any update by a process can be rejected by the ADS objects for at most \(\tau\) consecutive rounds from the time it was generated before being accepted.

**Proof.** The proof of the above properties follows directly from the structure of the partial-all-reduce algorithm, and is therefore skipped. \(\square\)

### 6.2 Convergence Proof

We now show that these properties are sufficient for eager-SGD to ensure convergence for a standard class of smooth non-convex objectives. In the following, all norms are \(\ell_2\)-norms, unless otherwise stated.

**Assumption 1 (Loss Function).** We assume that our objective loss function \(f : \mathbb{R}^d \rightarrow \mathbb{R}\) satisfies the following standard properties:

- **(Lower Bound)** The function \(f\) is bounded from below, that is, there exists a finite value \(m\) such that, \(\forall \bar{x} \in \mathbb{R}^d, f(\bar{x}) \geq m\).
- **(Smoothness)** The function \(f\) is \(L\)-smooth, i.e.

\[
\forall \bar{x}, \bar{y} \in \mathbb{R}^d, \|\nabla f (\bar{x}) - \nabla f (\bar{y})\| \leq L\|\bar{x} - \bar{y}\| \text{ for } L > 0.
\]

Further, we make the following standard assumptions about the gradients generated by the nodes:

**Assumption 2 (Gradients).** For any round \(t\) and processor \(i\), the gradients \(G^t_i\) generated by the processes satisfy the following, where expectations are taken with respect to the random data sampling at round \(t\):

- **(Unbiasedness)** The stochastic gradients are unbiased estimators of the true gradients:

\[
\forall \bar{x} \in \mathbb{R}^d, \mathbb{E}\left[G^t_i(\bar{x})\right] = \nabla f(\bar{x}).
\]

- **(Second Moment Bound)** There exists a constant \(M\) such that

\[
\forall \bar{x} \in \mathbb{R}^d, \mathbb{E}\left[\|G^t_i(\bar{x})\|^2\right] \leq M^2.
\]

**Analytic View of the Algorithm.** Let us fix a global round \(t + 1\), and consider the view of an arbitrary process \(i, w^t_i\) at the beginning of this round. Recall that this view consists of the view returned by the object ADS\(t\). Therefore, by Lemma 6.1, this view must include the sum of at least \(Q\) distinct gradients generated in each previous round, possibly together with some additional gradients, some of which are included in their corresponding round, and some of which are delayed. Conversely, if we consider the gradients which have been proposed to ADS objects by all nodes by time \(t\) and are not included in this view, we have that there can be at most \(P - Q\) such gradients for any previous round, up to maximum time \(t\) in the past. We formalize this observation as follows.

Define recursively the auxiliary random variable \(\Lambda_t\) such that for every round \(t \geq 0\),

\[
\Lambda_{t+1} = \Lambda_t - \frac{\alpha}{\bar{p}} \sum_{i=0}^{t-1} G^t_i(w^t_i),
\]

where \(\alpha > 0\) is the learning rate, which we assume to be constant. Without loss of generality, we set \(\Lambda_0 = 0\). Intuitively, \(\Lambda_t\) would like to follow the "clean" SGD iteration, by including all the gradients generated by the end of round \(t\). However, one technical issue is that these gradients are generated not with respect to the model \(\Lambda_{t-1}\) (which would allow us to perform a standard SGD analysis) but with respect to the partial views \(w^t_i\). We will overcome this obstacle by leveraging the fact that the partial view \(w^t_i\) cannot be too far from \(\Lambda_t\). More precisely, the discussion in the previous paragraph implies:

**Lemma 6.2.** For any \(t \geq 0\) and process \(i\), we have:

\[
\mathbb{E}[\|\Lambda_t - w^t_i\|^2] \leq \alpha^2 \tau M^2 (P - Q)/P^2.
\]

**Proof.** Let \(\delta^t_i\) be a binary indicator random variable that is true if the gradient generated by process \(j\) at iteration \(t\) is not delivered by the ADS\(t\) object. Then, we have that:

\[
\|\Lambda_t - w^t_i\|^2 = \|\sum_{t=1}^{\infty} \alpha \sum_{j=1}^{P} \delta^t_j G^t_j/P\|^2 \leq \sum_{t=1}^{\infty} \left(\alpha^2/P\right) \sum_{j=1}^{P} \delta^t_j\|G^t_j\|^2 \leq \sum_{t=1}^{\infty} (\alpha^2/P) \sum_{j=1}^{P} \delta^t_j\|G^t_j\|^2,
\]

where we have used the properties stated in Lemma 6.1 (in particular the Staleness Bound), and the triangle inequality. Next, we notice that (1) the expected squared norm of each of the missing gradients is bounded by \(M^2\) (by the second moment bound), and that (2) there can be at most \(P - Q\) delayed gradients from each round (by the Quorum Size
bound). This finally implies the claimed inequality:
\[
\mathbb{E}[\|\Lambda_t - w_t^i\|^2] \leq (\alpha^2 / P^2) \sum_{i=1}^r \sum_{j=1}^P \delta^i_j \mathbb{E}[\|G_i^j\|^2] \tag{4}
\]
\[
\leq \alpha^2 \tau M^2 (P - Q) / P^2. \tag{5}
\]

**Convergence Bound.** Finally, we put all the machinery together to obtain the following convergence bound:

**Theorem 6.3 (Eager-SGD Convergence).** Consider an arbitrary objective function \( f \) and gradient sampling scheme satisfying Assumptions 1 and 2. Fix the success parameter \( \epsilon > 0 \). Then, if we execute the eager-SGD algorithm for constant learning rate value

\[
\alpha \leq \min \left( \frac{\sqrt{\epsilon} P}{12L^2 r M^2 (P - Q)} , \frac{\sqrt{\epsilon} P}{4L^2 M^2 (P - Q)} , \frac{\epsilon}{12M^2 L} \right)
\]

for \( T = \Theta \left( \frac{(f(w_0) - m)}{\alpha \epsilon} \right) \) iterations, we are guaranteed to reach some iterate \( w_t^* \) with \( 1 \leq t \leq T \) such that

\[
\mathbb{E}[\|\nabla f(w_t^*)\|^2] \leq \epsilon.
\]

**Proof.** We begin from the definition of \( \Lambda_t \):

\[
\Lambda_{t+1} = \Lambda_t - \frac{\alpha}{P} \sum_{i=0}^{P-1} G_i^j (w_t).
\]

We will first prove the above statement for the iterate \( \Lambda_t \), and then will extend the proof for \( w_t \). For simplicity, let us denote \( G_t = \sum_{i=0}^{P-1} G_i^j (w_t) \). We can use the Taylor expansion of \( f(\Lambda_{t+1}) \) around \( \Lambda_t \) and the smoothness condition to obtain the following inequality:

\[
f(\Lambda_{t+1}) \leq f(\Lambda_t) + (\Lambda_{t+1} - \Lambda_t)^T \nabla f(\Lambda_t) + \frac{L}{2} \|\Lambda_{t+1} - \Lambda_t\|^2
\]

\[
= f(\Lambda_t) - \alpha \nabla f(\Lambda_t)^T \nabla f(\Lambda_t) + \frac{\alpha^2 L}{2P^2} ||G_t||^2 + \alpha (\nabla f(\Lambda_t) - G_t / P)^T \nabla f(\Lambda_t).
\]

We can therefore apply the expectation with respect to the random sampling at step \( t \), the second moment bound assumption:

\[
\mathbb{E} [f(\Lambda_{t+1})|\Lambda_t] \leq f(\Lambda_t) - \alpha \mathbb{E}[\nabla f(\Lambda_t)]^2 + \frac{\alpha^2 L}{2} M^2
\]

\[
+ \alpha \mathbb{E}[\nabla f(\Lambda_t) - \nabla f(w_t^*)] \mathbb{E}[\nabla f(\Lambda_t)].
\]

To bound the last term, we can now apply the Cauchy-Schwarz inequality and the fact that the gradients are \( L \)-Lipschitz:

\[
\mathbb{E} [f(\Lambda_{t+1})|\Lambda_t] \leq f(\Lambda_t) - \alpha \|\nabla f(\Lambda_t)\|^2 + \frac{\alpha^2 L}{2} M^2
\]

\[
+ \alpha L \|\Lambda_t - w_t^i\| \|\nabla f(\Lambda_t)\|.
\]

To further bound the last term, we can apply the classic inequality \( a^2 + b^2 \geq 2ab \) together with Lemma 6.2 to obtain:

\[
\mathbb{E} [f(\Lambda_{t+1})|\Lambda_t] \leq f(\Lambda_t) - \alpha \|\nabla f(\Lambda_t)\|^2 + \frac{\alpha^2 L}{2} M^2
\]

\[
+ \alpha \|\nabla f(\Lambda_t)\|^2 / 2 + \frac{\alpha^2 L^2 M^2 (P - Q)}{2P^2}.
\]

Rearranging terms and taking total expectation:

\[
\mathbb{E} [\|\nabla f(\Lambda_t)\|^2] \leq \frac{2\mathbb{E} [f(\Lambda_t) - f(\Lambda_{t+1})]}{\alpha} + \frac{\alpha M^2 L}{2} + \frac{\alpha^2 L^2 M^2 (P - Q)}{2P^2}.
\]

Summing across all \( t \) and dividing by \( T \), we get:

\[
\min_{1 \leq t \leq T} \mathbb{E} [\|\nabla f(\Lambda_t)\|^2] \leq \frac{1}{T} \sum_t \mathbb{E} [\|\nabla f(\Lambda_t)\|^2] \leq \frac{2(f(\Lambda_0) - m)}{\alpha T} + \frac{\alpha M^2 L + \alpha^2 L^2 M^2 (P - Q)}{2P^2}.
\]

We now study the set of conditions for each of the three RHS terms to be less than \( \epsilon / 12 \). We have that it is sufficient for the following three conditions to hold:

1. \( T \geq \frac{24 f(\Lambda_0) - m}{\alpha \epsilon} \);
2. \( \alpha \leq \frac{12M^2 L}{\sqrt{\epsilon} P} \);
3. \( \alpha \leq \frac{\epsilon}{12M^2 L} \).

All these conditions hold by assumption from the theorem statement. We have therefore obtained that there exists \( t^* \) such that \( \|\nabla f(\Lambda_{t^*})\|^2 \leq \epsilon / 4 \). However, by smoothness and Lemma 6.2 we know that

\[
\mathbb{E}[\|\nabla f(\Lambda_{t^*}) - \nabla f(w_t^*)\|^2] \leq \alpha^2 L^2 M^2 (P - Q) / P^2 \leq \epsilon / 4,
\]

where we have used the assumption in the theorem statement on the upper bound on \( \alpha \). Finally, we can apply the classic inequality \( \|a + b\|^2 \leq 2\|a\|^2 + \|b\|^2 \) to obtain that

\[
\mathbb{E}[\|\nabla f(w_t^*)\|^2] \leq \epsilon.
\]

\[ \square \]

**Discussion** We make the following observations regarding the bound. First, we note that, since we analyze non-convex objectives, we must settle for a weaker notion of convergence than in the convex case (where we can prove convergence to a global minimum); specifically, we prove that, for a given sequence of learning rates, the algorithm will converge to a point of negligible gradient. Second, we note the dependence in \( \sqrt{T} \) and \( \sqrt{(P - Q)} \) for the number of iterations to convergence, i.e.:

\[
T \geq \Theta \left( \frac{(f(w_0) - m) \sqrt{T(P - Q)}}{P \epsilon^{3/2}} \right).
\]

Thus, we would like the maximum delay and the number of "missed" gradients per round to be minimized. However, obviously, having no stragglers would imply higher synchronization cost. This suggests that, in practice, the algorithm
should trade off the additional performance cost of synchronization with the slower convergence due to delayed gradient information.

7 Evaluation

Experiments are conducted on the CSCS Piz Daint supercomputer with Cray Aries interconnect. Each XC50 compute node contains a 12-core Intel Xeon E5-2690 CPU with 64 GiB RAM, and one NVIDIA Tesla P100 GPU. The communication library is Cray MPICH 7.7.2. We use one MPI process per node and utilize the GPU for acceleration in all following experiments. First, we evaluate the performance of the partial collective operations using a microbenchmark. Then, we use the different neural networks summarized in Table 1 to compare our eager-SGD with the state-of-the-art synchronous SGD implementations (Horovod [51] and Deep500 [9]), the asynchronous centralized SGD [1], and the gossip-based SGD [4, 39], under simulated and real workload imbalance environments.

![Figure 8. Microbenchmark used to test the latency of the collective operations.](image)

```c
sendbuff={0}; recvbuff={0}; //initialization
pid = process ID; psize = processes number;
for(i = 0; i < psize - 1; i++) {
    sendbuff = {0}; recvbuff = {0}; //reset to dummy values
}
begin = MPI_Wtime();
call MPI/Solo/Majority_Allreduce(sendbuff, recvbuff);
while (sendbuff == -1) {
    latency = MPI_Wtime() - begin;
    average_latency = sum(latency)/psize;
    MPI_Barrier(); //synchronize before the next iteration
    sendbuff = [0]; recvbuff = [0]; //reset to dummy values

Figure 9. Average latency comparison between MPI_Allreduce and partial allreduce running on 32 processes by 64 iterations. Processes are linearly skewed by injecting load imbalance from 1 ms to 32 ms.

For the partial collective operations, we refer to the initiator together with the processes that arrive at the operation before the initiator as the active processes, which contribute the latest data (line 5 in Fig. 8). The other processes contribute null values (line 13 in Fig. 8). For solo allreduce, since the fastest process is the initiator and all the processes are fully skewed, the Number of Active Processes (NAP) is around 1, as shown in Fig. 9. For majority allreduce, since the initiator is randomly specified, the expectation of NAP is half of the total processes. On average 16 out of 32 processes for majority allreduce are active processes, which means half of the processes contribute the latest data when the processes are fully skewed.

7.2 Throughput and Convergence with Simulated Workload Imbalance

We use three networks shown in Table 1, including a multilayer perceptron (MLP), ResNet-32, and ResNet-50, to evaluate the performance of eager-SGD with simulated workload imbalance. From the application perspective, these three networks have balanced workload during the distributed training, since each batch has equivalent workload. We manually inject delays to simulate the load imbalance environment caused by the training system, as discussed in Section 2.3.

7.2.1 Hyperplane Regression, Light Load Imbalance

We generate both training and validation datasets for a 8,192-dimensional hyperplane regression using the equation: $y =$
We use 8 processes with the total batch size of 2,048 to train a SGD, respectively. We observe that the more severe the load imbalance, the worse the performance of synch-SGD because of the synchronization overhead. On the other hand, the performance of eager-SGD is stable. Given that the throughput on a single GPU node with batch size of 2,048 is 0.64 steps/s, eager-SGD with 400 ms load imbalance injection still achieves 3.8x speedup in strong scaling on 8 GPU nodes.

Fig. 10b presents the validation loss (mean squared error) as a function of the training time, which shows that eager-SGD using solo allreduce converges with equivalent loss value (around 4.7) to synch-SGD but significantly reduces the training time. Since the processes are not severely skewed and the stale gradients are added to the next training iteration (as discussed in Section 5), using solo allreduce is enough for convergence. When using majority allreduce, the throughput of eager-SGD is lower than using solo allreduce (1.64 step/s vs 1.37 step/s with 200 ms load imbalance injection).

### Table 1. Neural networks used for evaluation.

| Tasks                  | Models                      | Parameters | Train data size     | Batch size | Epochs | Processes |
|------------------------|-----------------------------|------------|---------------------|------------|--------|-----------|
| Hyperplane regression  | One-layer MLP               | 8,193      | 32,768 points       | 2,048      | 48     | 8         |
| Cifar-10               | ResNet-32 [21]              | 467,194    | 50,000 images       | 512        | 190    | 8         |
| ImageNet [14]          | ResNet-50 [21]              | 25,559,081 | 1,281,167 images    | 8,192      | 90     | 64        |
| UCF101 [53]            | Inception+LSTM [61]         | 34,663,523 | 9,537 videos        | 128        | 50     | 8         |

Fig. 11a presents the throughput comparison between synch-SGD (Horovod and Deep500) and eager-SGD using solo allreduce. With 300 and 460 ms load imbalance injection, eager-SGD achieves 1.25x and 1.29x speedup over Deep500, respectively; 1.14x and 1.27x speedup over Horovod, respectively. Given that the throughput of a single GPU node with batch size of 128 is 1.56 steps/s, eager-SGD running on 64 processes with 460 ms load imbalance injection still achieves 49.8x speedup in weak scaling.

**7.2.2 ResNet-50 on ImageNet, Light Load Imbalance**

Residual Network (ResNet) [21] is widely used in computer vision tasks. To evaluate the performance of eager-SGD, we use 64 processes with a total batch size of 8,192 to train ResNet-50 on ImageNet for 90 epochs. To simulate the load imbalance environment, we randomly select 4 processes out of the 64 processes at every training step to inject a certain amount of delay, according to the performance variability on Cloud machines discussed in Section 2.3.

Fig. 11b and Fig. 11c present the Top-1 train and test accuracy as a function of the training time, respectively. We train the model three times for each SGD, and obtain stable accuracy results. For top-1 accuracy, Deep500 achieves 79.1% train accuracy and 75.7% test accuracy, Horovod achieves 79.0% train accuracy and 75.8% test accuracy, while eager-SGD using solo allreduce achieves 78.4% train accuracy and 75.2% test accuracy on average over different load imbalance injections. Note that without model synchronization at every 10 epochs, the top-1 test accuracy of eager-SGD decreases.
(a) Throughput comparison. Each point is at the boundary of one epoch.

(b) Top-1 training accuracy. Each point is at the boundary of one epoch.

(c) Top-1 test accuracy. Each point is at the boundary of every 10 epochs.

Figure 11. Comparisons between synch-SGD and eager-SGD for ResNet-50 on ImageNet using 64 processes. "synch/eager-SGD-300/460" represent 300/460 ms load imbalance injection, respectively.

Table 2. Throughput comparison with the asynchronous centralized SGD and the gossip-based SGDs for ResNet-50 on ImageNet, using 64 processes (total batch size 8,192) under load imbalance environment.

| SGDs       | Asynch-PS [1] | D-PSGD [39] | SGP [4] | eager-SGD |
|------------|---------------|-------------|---------|------------|
| step/s     | 0.45          | 0.94        | 1.02    | 1.19       |

Table 2 presents the throughput comparison with the asynchronous centralized SGD and the gossip-based SGDs for ResNet-50 on ImageNet. We randomly select 4 processes out of the 64 processes at every training step and inject 460 ms delay for each selected process. Asynch-PS [1] is the asynchronous Parameter-Sever-based (centralized) SGD provided by TensorFlow. The throughput of Asynch-PS is the lowest among all the SGD variants because of the performance bottleneck on the server. Compared with Asynch-PS, eager-SGD achieves 2.64× speedup. D-PSGD [39] and SGP [4] are gossip-based SGDs, which do not use global collective communication primitives, such as Allreduce. Alternatively, each process only communicates with its neighbors (two neighbors for D-PSGD and SGP). However, all the processes need to finish the communications of the current step before going to the next step. The Overlap SGP [4] can mitigate the synchronization effect by overlapping communication and computation. According to the parameter setup in [4], we configure SGP to use one step of gradient computation to overlap the communication, namely the communication synchronization is delayed by one step. Using communication topology optimizations [4, 40], each process can globally propagate its local update using $O(\log P)$ steps. Note that eager-SGD only uses $O(1)$ step to globally propagate the local update. As shown in Table 2, eager-SGD also outperforms the gossip-based SGDs because of the feature of asynchrony.

7.2.3 ResNet-32 on Cifar-10, Severe Load Imbalance
To test the robustness of eager-SGD, we train ResNet-32 on Cifar-10 with 8 processes for 190 epochs in a severe load imbalance environment. All 8 processes are skewed by injecting load imbalance from 50 ms to 400 ms at every training step. The injection amount over the processes is shifted after each step. Fig. 12 presents the test accuracy as a function of the training time. Eager-SGD using solo allreduce has the highest training speed but with lower test accuracy. Solo allreduce only waits for the fastest process to inform the other processes to participate in allreduce, but most of them will contribute stale gradients. Majority allreduce can solve
the lower accuracy problem caused by solo allreduce, which achieves approximately equivalent accuracy to synch-SGD with 1.29x speedup. The results demonstrate that eager-SGD using majority allreduce is tolerant to severe load imbalance.

7.3 Case Study: Video Classification

As discussed in Section 2.1, LSTM on UCF101 for video classification has inherent workload imbalance because of different workload for different batches. We use Inception v3 [55], a CNN model, to extract a 2,048-wide feature from each frame of the videos, and then pass the sequences of features to an LSTM model. The training time reported in the paper is only for the LSTM model, not including the preprocessing time using Inception v3.

To evaluate eager-SGD, we use 8 processes with a total batch size of 128 to train LSTM on UCF101 for 50 epochs (more information is in Table 1). Fig. 13a and Fig. 13b present the train and test accuracy as a function of the training time, respectively. For each SGD, we train the model four times and plot the curves using the average values. Colored areas around the curves are confidence intervals with the boundaries representing the standard deviation. Although eager-SGD using solo allreduce achieves 1.64x speedup over Horovod, it has lower accuracy. Eager-SGD (solo) achieves on average 60.6% (up to 70.4%) top-1 test accuracy while Horovod achieves on average 69.6%. This is because the workload of the video model is severely unbalanced, and solo allreduce introduces too many stale gradients. In contrast, eager-SGD using majority allreduce achieves 1.27x speedup over Horovod with equivalent accuracy. For example, Horovod achieves on average 69.6% top-1 test accuracy (up to 72.2%) and 90.4% top-5 test accuracy (up to 91.9%), while eager-SGD using majority allreduce achieves on average 69.7% top-1 test accuracy (up to 72.8%) and 90.0% top-5 test accuracy (up to 91.7%). Train accuracy results (in Fig. 13a) show a similar trend as the test accuracy. Horovod achieves on average 86.1% top-1 train accuracy and 96.6% top-5 train accuracy, while eager-SGD using majority allreduce achieves on average 86.7% top-1 train accuracy and 96.1% top-5 train accuracy. All the accuracy results are consistent with that claimed in recent work [61]. The training speed and accuracy for Deep500 (not plotted in figures) are similar to Horovod. The results show that majority allreduce, with its statistical guarantee, is sufficient to both speed up training and achieve good accuracy.

The total training time for 50 epochs using a single GPU node with batch size of 16 and 128 is 34,301 seconds and 6,314 seconds, respectively. In weak scaling, Synch-SGD (Horovod) and eager-SGD using majority allreduce achieve 3.72x and 4.71x speedup on 8 GPU nodes, respectively. In strong scaling, synch-SGD and eager-SGD using majority allreduce do not have speedup on 8 GPU nodes; in contrast, eager-SGD using solo allreduce achieves 1.12x speedup on 8 GPU nodes in strong scaling, but with lower accuracy. Note that increasing the batch size can further improve the speedup in strong scaling for eager-SGD. However, large batch sizes commonly
need carefully-tuned learning rate schedules to achieve good accuracy [59], which is out of scope.

8 Related Work

Deep learning Parameter Server SGD implementations use synchronous [20, 38], asynchronous [11, 13], stale- [22, 62], and approximate-synchronous [29] SGD, where the latter two limit the age and insignificance of the gradients, respectively. For synchronous Parameter Server SGD, communication granularity and scheduling optimizations [33] are studied to better overlap communication and computation. In a decentralized setting, asynchrony is achieved by performing communication on an explicit subset of the learners, e.g., forming a ring [40] or a general graph [58]; or on a random subset using Gossip Algorithms [12, 34]. These modes achieve some degree of asynchrony, but take $O(P)$ or $O(\log P)$ (for ring or gossip-based schemes, respectively) update steps to disseminate the information to all $P$ learners.

To the best of our knowledge, this is the first work that implements asynchronous and stale-synchronous decentralized SGD where the messages propagate to all nodes in one step.

Collective communication Several algorithms can be used to implement allreduce operations, and the optimal algorithm depends on network topology, number of processes, and message size [56]. For large message sizes and large number of processes, practical implementations employ the ring-allreduce [19] or the Rabenseifner’s Algorithm [44]. Independently from the specific algorithm, the semantic of the allreduce implies processes synchronization. With eager-SGD we relax this synchronization by using solo and majority allreduce operations.

9 Conclusions

In this work, we show that load imbalance is prevalent in deep learning problems with variable-length inputs, and increasingly becoming an issue in cloud systems. To that end, we propose eager-SGD: an asynchronous decentralized version of SGD where fast processes contribute gradients without waiting for slow processes. Using the resilience of machine learning to bounded error, we implement two variants of partial collectives — solo and majority allreduce — enabling this behavior without a central parameter server.

The experimental results reaffirm our theoretical analysis, showing that eager-SGD using solo allreduce speeds up the training process (1.29× and 2.64× faster than the synchronous decentralized SGD and the asynchronous centralized SGD on ImageNet, respectively) in light load imbalance environments. As the load imbalance increases, the convergence rate of solo allreduce degrades, in which case majority allreduce speeds up the training process (1.27× faster than the synchronous decentralized SGD on UCF101) yet desirable generalization.

The research can extend in different directions. Firstly, the promising results make eager-SGD attractive for other applications as well, such as language models and object detection. Secondly, in order to provide different quorum sizes, it is possible to construct a spectrum between solo, majority, and full collectives. Lastly, partial collectives can be used for other algorithms beyond SGD.

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A Artifact Appendix

A.1 Abstract
We provide source code of eager-SGD and scripts to run experiments from the paper. This artifact is run on the Piz Daint supercomputer. This artifact supports the paper by making it possible to reproduce the figures and numbers in this paper, and it can be validated by comparing the figures and results that this artifact generates with the data from the paper.

A.2 Artifact check-list (meta-information)
- Algorithm: Eager Stochastic Gradient Descent (eager-SGD)
- Compilation: cmake, g++, Python 3.6
- Data set: ImageNet, CIFAR-10, UCF101
- Run-time environment: Cray MPICH 7.7.2, TensorFlow-gpu r1.11, Horovod, mpi4py, Keras
- Hardware: Piz Daint Supercomputer (Intel Xeon E5-2690 CPU, NVIDIA Tesla P100 GPU)
- Execution: SLURM job scheduler on Piz Daint
- Metrics: Execution time, training throughput, loss values, Top1 and Top5 accuracy
- Output: TXT files and Figures
- Experiments: Use the provided scripts in the artifact to build, schedule jobs, and generate figures
- How much disk space required (approximately)?: 400 GB
- How much time is needed to prepare workflow (approximately)?: Assuming access to Piz Daint, 30 minutes
- How much time is needed to complete experiments (approximately)?: About 90 hours if each job can be scheduled to run immediately. Considering the job queuing time, it may take one week.

A.3 Description

A.3.1 How delivered
Via Dropbox: https://www.dropbox.com/s/3k8xgw0rh0s0m7j/eager-SGD-artifact.zip?dl=0

A.3.2 Hardware dependencies
This artifact uses Piz Daint supercomputer.

A.3.3 Software dependencies
To run the experiments, Python 3.6, TensorFlow-GPU, Horovod, mpi4py, MPICH, and Keras are required. To plot out the figures, RStudio and ggplot2 are required.

A.3.4 Data sets
Download the training and validation images of ImageNet from http://www.image-net.org/challenges/LSVRC/2010/downloads. Download the binary version of CIFAR-10 from https://www.cs.toronto.edu/~kriz/cifar-10-binary.tar.gz. Download UCF101 from https://wwwcrcv.ucf.edu/data/UCF101/UCF101.rar

A.4 Installation
1. Download the artifact and move it to your personal $WORK directory. Extract the artifact using unzip.
2. Install the dependent Python modules.

A.5 Experiment workflow
To run experiments, users run the provided scripts that will schedule runs of the executable on Piz Daint via sbatch. To run jobs on Piz Daint, one must put them on a queue and wait until they are scheduled. Once these experiments finish, the results of execution time, loss values, Top1 and Top5 accuracy will be output. We provide scripts that will compile these output results into TXTs using Python, and from these TXTs, we have included scripts that will use R to create the figures that we used in the paper.

A.6 Evaluation and expected result
Users are expected to reproduce the results in this paper, specifically generating the figures in Section 7. Different versions of MPICH, TensorFlow-gpu, Horovod, and mpi4py may lead to slightly variant results compared with the numbers reported in the paper, but this does not affect the general trends claimed in the paper.

1. Evaluate solo and majority allreduce and generate Fig. 9.
   - cd $WORK/eager-SGD-artifact/eager-SGD
   - pip install $requirements.txt
   - Compile ffmpeg2 and set the environment variable.
   - cd $WORK/eager-SGD-artifact/eager-SGD/ffmpeg2/lib
   - cmake .. && make
   - export LD_LIBRARY_PATH=$WORK/eager-SGD-artifact/eager-SGD/ffmpeg2/lib:$LD_LIBRARY_PATH
   - Configure a CMakelist file which will be used for compiling the customized Tensorflow operators.
   - mpm $WORK/eager-SGD-artifact/eager-SGD/ffmpeg2/deep500/deep500/frameworks/tensorflow/custom_operators_CMakelists.txt
   - Update include_directories and link_directories by the path where ffmpeg2 is installed. Set TENSORFLOW_PATH by the path where TensorFlow is installed.
   - export PYTHONPATH=$PYTHONPATH:$WORK/eager-SGD-artifact/eager-SGD-artifact/eager-SGD/ffmpeg2/deep500/

2. Train hyperplane regression and generate Fig. 10.
   - cd $WORK/eager-SGD-artifact/test-scripts/hyperplane-scripts
   - Submit the jobs.
   - mpm ./sbatch_jobs.sh
   - It may take about 10 minutes to finish the jobs, and then outputs majority.txt, solo.txt, and mpi.txt. Next, run the file statistics_summary.py to read the output files and calculate the mean and standard deviation for the latency and results.
   - python statistics_summary.py
   - Now it should generate latency_results_summary.txt. Next, run the file ./plotRstudio/plot-Figure9.R (RStudio and ggplot2 are required) to generate Fig. 9. Detailed steps are stated in ./plotRstudio/ReadMe.

The expected results are as follows:
   a) For the average latency, Solo_Allreduce < Majority_Allreduce < MPI_Allreduce, where "<" means "less than".
   b) For the average results, Solo_Allreduce < Majority_Allreduce < MPI_Allreduce, where "<" means "less than".
A.7 Experiment customization

Users can modify the scripts in the subdirectories of $WORK/eager-SGD-artifact/test-scripts to customize the experiments.

Modify the line of #SBATCH --nodes=<number-of-nodes> to change the number of nodes (processes). Modify the input parameter --bs=<batch-size-per-node> to change the batch size per node. Note that the total-batch-size = number-of-nodes * batch-size-per-node, which means the change of the number of nodes and the batch size per node would change the total batch size. However, different total batch sizes may lead to different results for the train and test accuracy.

Modify the line of #SBATCH --time=<time-quota> to change the time limit to run the job.

Users can also use eager-SGD to train other TensorFlow-based models that are not listed in the paper. To achieve this, simply wrap the TensorFlow optimizer using the eager-SGD optimizer, and then use the eager-SGD optimizer instead to train the model.

A.8 Notes

Some jobs may not be scheduled to run for a surprising long time because of the busy use of the machine. In this case, they may be automatically cancelled by the job scheduler. This is usually resolved by rescheduling the cancelled jobs using sbatch.

A.9 Methodology

Submission, reviewing and badging methodology: http://cTuning.org/ae/submission-20190109.html
http://cTuning.org/ae/reviewing-20190109.html
https://www.acm.org/publications/policies/artifact-review-badging