ZERO REDUNDANCY DISTRIBUTED LEARNING WITH DIFFERENTIAL PRIVACY

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ABSTRACT

Deep learning using large models have achieved great success in a wide range of domains. However, training these models on billions of parameters is very challenging in terms of the training speed, memory cost, and communication efficiency, especially under the privacy-preserving regime with differential privacy (DP). On the one hand, DP optimization has comparable efficiency to the standard non-private optimization on a single GPU, but existing DP distributed learning (such as pipeline parallelism) has suffered from significantly worse efficiency on multiple GPUs. On the other hand, the Zero Redundancy Optimizer (ZeRO) is a state-of-the-art solution to the standard distributed learning, exhibiting excellent training efficiency on large models, but to work compatibly with DP is technically complicated. In this work, we develop a new systematic solution, DP-ZeRO, (I) to scale up the trainable DP model size, e.g. to GPT-100B, (II) to obtain the same computation and communication efficiency as the standard ZeRO, and (III) to enable mixed-precision DP training. Our DP-ZeRO, like the standard ZeRO, has the potential to train models with arbitrary size and is evaluated on the world's largest DP models in terms of the number of trainable parameters.¹

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1 INTRODUCTION

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029 Recent advances in differentially private (DP) deep learning have witnessed the power of large pretrained models, achieving comparable accuracy to state-of-the-art (SOTA) non-private models across 031 computer vision De et al. (2022); Bu et al. (2022a); Mehta et al. (2022); Xie et al. (2018), natural language processing Yu et al. (2021a); Li et al. (2021); Bu et al. (2023a), and many other tasks. 032 Similar to their non-DP counter-parts, it has been observed that larger DP models tend to have 033 better performance. For example, the DP accuracy increases from 83% using RoBERTa-base (123M 034 parameters) to 86% using RoBERTa-large (354M parameters) on GLUE datasets Li et al. (2021); 035 Bu et al. (2023a); Yu et al. (2021a); the DP BLEU score increases from 61 using GPT2-small (124M parameters) to 64 using GPT2-large (800M parameters) on E2E dataset Li et al. (2021); Bu et al. 037 (2023a); a similar trend is also observed using ViT (Base/Large/Huge) up to 600M parameters to achieve state-of-the-art DP accuracy on ImageNet, around 81% at $\epsilon = 8$ Mehta et al. (2022).

Driven by this success and the surge of computational power, it is high time to enable DP deep learning at the same scale of the standard non-DP one, e.g., GPT3-175B (Brown et al., 2020) and LLaMA-63B (Touvron et al., 2023a;b). Specifically, such a DP training system must have high speed and memory efficiency, low communication cost, and the compatibility with general model architectures.

For small to moderately large models (e.g. with less than a billion parameters) that fit within the memory of a single GPU, a range of DP algorithms are feasible, producing the same result at different efficiency. Examples include TensorFlow-privacy Subramani et al. (2021), Opacus Yousefpour et al. (2021); Bu et al. (2022b), ghost clipping (GhostClip) Goodfellow (2015); Li et al. (2021); Bu et al. (2022a), and Book-Keeping (BK) Bu et al. (2023b), among which the BK algorithm has allowed DP optimization to be almost as efficient as the standard one. To be specific, the time/space complexity of BK algorithm is $1.08 \times /1.05 \times$ of the standard optimization on ViT-Large (300M parameters, 147 layers) and $1.03 \times /1.01 \times$ on GPT2-large (800M parameters, 220 layers).

¹Code at https://anonymous.4open.science/r/DP-ZERO-2821 (this is built on fastDP and no identifiers are related to our work).

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| | Summary of selected DP distributed | learning |
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| 054 | Table 1: Summary of selected DP distributed learning. | | | | | | |
|-----|---|--------------------------|------------|--|--------------------------|---|--|
| 055 | Distributed solution | Parallelism | Model | Standard | DP | Remark | |
| 056 | DDP | Data | No | Li et al. | Yousefpour et al. (2021) | unable to fit large model and DP is memory costly | |
| 057 | DDP | Data | No | Frostig et al. (2018) | De et al. (2022) | unable to fit large model and DP is slow | |
| 058 | GP1pe ZeRO | Pipeline Data(&Model) | Yes Yes | Huang et al. (2019) Rajbhandari et al. (2020) | He et al. (2022) Ours | speed & memory & communication efficient | |

To enable the DP distributed learning of these not-too-large models, one can directly use DDP 061 (distributed data parallelism) (Li et al.), where each mini-batch of data is partitioned to smaller 062 micro-batches and each GPU computes one micro-batch with a full copy of the DP model. A line 063 of researches (Yousefpour et al., 2021; De et al., 2022; Kurakin et al., 2022) have reported that 064 DDP with DP usually either incurs huge memory cost due to caching the per-sample gradients, or 065 suffers from $2 - 9 \times$ slower training speed than non-DP optimization De et al. (2022); Bu et al. 066 (2021). While the efficiency issues can be addressed through a better DP algorithm, such as BK, the 067 feasibility issue remains insurmountable because DDP cannot train models that exceed the capacity of one GPU. Notably, the efficiency of BK algorithm is enhanced by two key techniques: mixed 068 ghost norm (computing per-sample gradient norms almost for free) and book-keeping trick (only 069 using one round of full back-propagation, not two rounds as in Li et al. (2021); Bu et al. (2022a)), which are detailed in Appendix A and will also be leveraged in our DP-ZeRO solution. 071

072 As the model size further increases beyond a reasonable bound for one GPU (e.g. 32GB memory, 073 which roughly translates to 2B model training with Adam), the model must be partitioned in addition to the data, e.g. using pipeline parallelism and model parallelism, so that each GPU only holds 074 a partial shard of the model (see Figure 2). In He et al. (2022), DP is combined with pipeline 075 parallelism to fine-tune about 0.1% of GPT3-175B. Yet, the pipeline parallelism can be inefficient 076 due to a non-DP-related issue – the pipeline bubble, where GPUs are idle while waiting for data to 077 process.

079 Generally speaking, more advanced distributed methods such as Zero Redundancy Optimizer Rajbhandari et al. (2020) (ZeRO) and mixed-precision training have not be paired with DP due to the lack of algorithmic advances. In this work, we develop DP-ZeRO, equipping state-of-the-art dis-081 tributed learning solution with DP (see comparison in Table 1), without altering the mathematics of DP optimization. We summarize our contributions as follows². 083

- 1. We propose the zero redundancy distributed learning with differential privacy (DP-ZeRO), demonstrating the same level of communication efficiency, computation efficiency (speed and memory), and scalability (to GPT3 level and hundreds of GPUs) as the standard ZeRO.
- 2. We enable the mixed-precision training with DP by addressing the issue of loss scaling. This solution allows us to reduce the memory cost by roughly 50% and allow significantly faster communication that was previously not enjoyed by DP distributed learning.
 - 3. We enable DP deep learning with more than 1B trainable parameters for the first time. E.g. we are the first to train the full GPT2-XL, ViT-Gigantic, ViT-10B and GPT-100B with DP.
- 4. We will open-source our codebase as the first DP distributed learning library, that automatically applies DP-ZeRO for general tasks (e.g. classification and language understanding), general network architectures (e.g. ResNet, ViT, GPT), and general distributed solutions (including DeepSpeed and FSDP).
- PRELIMINARY 2
- 2.1 DIFFERENTIAL PRIVACY

DP provides a formal privacy guarantee, making it difficult to extract any information from training 103 data. The privacy guarantee is characterized by (ϵ, δ) -DP in Definition 2.1, with smaller (ϵ, δ) 104 representing lower privacy risk. 105

¹⁰⁶ ²We do not report any accuracy results since this is a system design paper, following the norm in the original 107 ZeRO Rajbhandari et al. (2020) and FSDP Zhao et al.. We verify that DP-ZeRO is implemented correctly by comparing to single-GPU DP libraries in Appendix E.



Figure 1: Total/trainable parameters of existing DP models (blue) and ours by DP-ZeRO (red).

Definition 2.1 (Dwork et al. (2006)). A randomized algorithm M is (ε, δ) -DP if, for any two neighborst boring datasets S, S' that differ by one sample and for any event E, we have $\mathbb{P}[M(S) \in E] \leq$ $e^{\varepsilon}\mathbb{P}\left[M\left(S'\right)\in E\right]+\delta.$

In DP deep learning, the gradients are made private by post-processing through per-sample gradient clipping and random noising:

private gradient (*m*-th group):
$$\mathbf{G}_{[m]} := \sum_{i} C_i(R_m) \boldsymbol{g}_{[m],i} + \sigma_{\mathrm{DP}} \| [R_1, \cdots, R_M] \| \cdot \mathcal{N}(0, \mathbf{I}).$$
 (1)

133 Here the gradient of all trainable parameters is partitioned into M groups, i.e. $g_{[m],i}$ is the *i*-th 134 per-sample gradient of the *m*-th group's parameters, where $m \in \{1 \cdots M\}$ is the group index. C_i is 135 the per-sample gradient clipping factor so that $||C_i g_{[m],i}|| \leq R_m$ and R_m is the clipping threshold. 136 That is, DP optimization is enabled when the standard optimizers such as stochastic gradient descent 137 (SGD) and Adam (Kingma & Ba, 2014) update the trainable parameters with the private gradient, instead of the standard gradient $\sum_i g_i$. 138

140 **Mathematical gradient partition** In equation 1, the trainable parameters and their gradients are 141 mathematically partitioned into M groups, e.g. in all-layer clipping, all parameters form one group 142 (M = 1) Abadi et al. (2016); in layer-wise clipping McMahan et al. (2018); He et al. (2022), 143 each layer's parameters form a group (M = number of layers). Empirical evidence and theoretical 144 analysis show that different partitions have the same training speed, though a finer partition (e.g. layer-wise) has lighter memory footprint³. 145

147 **Per-sample gradient clipping** In equation 1, a number of clipping functions $C_i = C(||g_i||; R)$ are 148 available. Most works Abadi et al. (2016); Li et al. (2021); Yu et al. (2021a) use the vanilla clipping $C_i = \min(R/||g_i||, 1)$. Recently, Bu et al. (2023a); Yang et al. (2022) advocate the automatic 149 clipping $C_i = 1/(||g_i|| + 0.01)$ which is hyperparameter-free and comparably accurate. Note if 150 $C_i \equiv 1$, then the clipped gradient reduces to the standard gradient. The main overhead of DP 151 optimization is the computation of per-sample gradient norms. On a single GPU, the mixed ghost 152 clipping Bu et al. (2023b) has reduced the time complexity to < 10% on large models like GPT2. 153

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155 **Privacy accounting** In equation 1, adding Gaussian noise to the clipped gradient protects the privacy that is quantifiable by the privacy accounting theory Abadi et al. (2016); Bu et al. (2020); 156 Dong et al. (2019); Zhu et al. (2022); Gopi et al. (2021); Koskela et al. (2020). The privacy guarantee 157 is increasing in the noise level $\sigma_{\rm DP}$, independent of R_m , learning rate, clipping function and model 158 architectures, with $\sigma_{\rm DP} = 0$ leading to $\epsilon = \infty$ (non-private). 159

³We note that DP optimization under different group-wise clippings can have the same computation and communication efficiency (under the BK algorithm), with or without ZeRO.

162 2.2 ZERO REDUNDANCY OPTIMIZER (ZERO)

164 2.2.1 PARALLEL COMPUTING

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Parallel computing is necessary to train large-scale models and is critical to the optimization efficiency. For models that fit in a single GPU, data parallelism (DataP) can be used to speed up the training by partition the mini-batch of samples into multiple micro-batches. Then, each GPU (holding a full copy of parameters) executes the forward and backward propagation of one micro-batch, from which the parameter gradients are generated and averaged across GPUs to update the trainable parameters. However, for models that do not fit in a single GPU, the model parameters need to be sharded by alternative solutions such as ZeRO Rajbhandari et al. (2020), model parallelism (ModelP) and pipeline parallelism (PipeP).

173 ModelP partitions a model vertically, e.g. using 3 GPUs to store the parameters of one layer. As 174 a consequence, ModelP does not scale efficiently beyond a single node due to fine-grained com-175 putation and expensive communication between layers. Implementation-wise, ModelP frameworks 176 usually require heavy code integration that may not be generalizable in model architectures. In con-177 trast, PipeP partitions a model horizontally across layers, e.g. storing 3 layers in each GPU. Each 178 GPU deals with all micro-batches sequentially, though PipeP can be inefficient due to the pipeline 179 bubble, which is overcome by ZeRO Rajbhandari et al. (2020). ZeRO is an advanced data parallel 180 method that eliminates memory redundancies during the training, and improves the training speed 181 and communication volume proportionally to the number of GPUs. Unlike basic DataP, ZeRO partitions a model's states across GPUs and gather/reduce in a just-in-time manner, thus sustaining the 182 high efficiency of very large model training. Notice that ZeRO can work compatibly with ModelP 183 and optionally offload the model states to CPUs Ren et al. (2021); Rajbhandari et al. (2021). 184

185 2.2.2 MODEL STATE PARTITION

A major part of the training memory is consumed by the model states⁴. ZeRO has three stages (ZeRO1/2/3) that partition these model states by different levels, with lower level being faster but more memory costly. For instance, in Table 2 of (Rajbhandari et al., 2020), ZeRO1/2/3 at most train 7.6/14.4/128B models on 64 V100 GPUs.

We take an example of mixed-precision Adam optimizer to train a model with Ψ_{model} parameters, which maintains a master copy (fp32) of optimizer states. and the half-precision parameters and gradients.

Optimizer state partition The optimizer states are the (master) parameters, variance and momentum, each taking $4\Psi_{model}$ memory. ZeRO1 only applies the optimizer state partition and updates the parameters locally, reducing the memory cost of model states from $16\Psi_{model}$ for basic DataP to $(4 + \frac{12}{N_d})\Psi_{model}$ at each of N_d GPUs.

| layer 1 | | layer 1 | | layer 1 | -GPU 1 |
|---------|--------|---------|---------|---------|---------|
| layer 2 | -GPU 1 | layer 2 | -GPU I | layer 2 | |
| layer 3 | -GPU 2 | layer 3 | | layer 3 | -GPU 2 |
| layer 4 | | -GPU Z | layer 4 | | layer 4 |

Figure 2: Mathematical (left two, same GPU allocation, different accuracy) and hardware (right two, different GPU allocation, same accuracy) gradient partition. Orange and blue are gradient groups $g_{[1]}$ and $g_{[2]}$ in DP optimization equation 1.

Hardware gradient partition In addition to ZeRO1, during the back-propagation, ZeRO2 (and ZeRO3) further partitions the $2\Psi_{model}$ gradients into different GPUs, reducing the memory cost to $(2 + \frac{14}{N_d})\Psi_{model}$ at each GPU. Figure 2 illustrates the difference between the hardware partition and the mathematical partition in Section 2.1.

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 ⁴Another important part of memory consumption is the batch-size-related variables such as the activation tensors, which is instantiated during the forward propagation and independent to DP (which modifies the gradient during the back-propagation).

Parameter partition In addition to ZeRO2, ZeRO3 also partitions the $2\Psi_{model}$ fp16 parameters, further reducing the memory cost to $\frac{16}{N_d}\Psi_{model}$ at each GPU.

2.3 MIXED-PRECISION TRAINING

Mixed-precision training (Micikevicius et al., 2018) performs the forward and backward propagation on the half precision (fp16 or bf16) parameters, activations, and gradients, while performing the model update in full precision (fp32). Compared to the full-precision training, it is capable of saving the memory by $\approx 50\%$ and accelerating the computation by $\approx 20\%$. We note that fp16 has better precision but a limited range: its representable numbers are among $10^{-8} \sim 10^5$, and vice versa for bf16. Therefore, loss scaling is necessary when using fp16 to prevent small gradients from being rounded to zero, thereby preserving the model's accuracy (see Appendix C for details). In contrast, bf16 usually does not need loss scaling since it has the same range as fp32. However, bf16 is not as widely supported as fp16, e.g. only available on NVIDIA Ampere GPUs or above (Nvidia).

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3 DIFFERENTIALLY PRIVATE ZERO

3.1 Algorithm

Our DP-ZeRO algorithm introduces the per-sample gradient clipping and noising to the standard ZeRO (Rajbhandari et al., 2020), while maintaining the efficiency. At high level, an iteration of ZeRO consists of the following steps:

$$\left(\text{ all-gather } \rightarrow \text{ forward } \rightarrow \text{ all-gather } \rightarrow \text{ backward } \rightarrow \text{ reduce} \right)^{\wedge L \text{ layers}} \rightarrow \text{ update(SGD/Adam/...)}$$

The operations in purple are global and require communication among GPUs, whereas the operations in green are locally computed within each GPU. In particular, DP optimization is only different from the standard optimization in the back-propagation, which can be decomposed into

backward =(output gradient \rightarrow clipping factor \rightarrow parameter gradient \rightarrow noising)

To give more details, we consider training a neural network of linear layers using N_d GPUs. We emphasize that the following procedure is sufficiently generic to cover other layer types, such as convolution, embedding, normalization, and so on, which are all supported by DP-ZeRO. The full algorithm is depicted in Figure 3, where we denote the *j*-th micro-batched variables like $a_l^{(j)}$, for $1 \le j \le N_d$.

The forward propagation of DP optimization is the same as that of the standard optimization:

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$$\mathbf{s}_l = \mathbf{a}_l \mathbf{W}_l + \mathbf{b}_l, \mathbf{a}_{l+1} = \phi_l(\mathbf{s}_l).$$
⁽²⁾

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At the *l*-th layer, $a_l \in \mathbb{R}^{BT_l d_l}$ is the layer's input, also known as the activation, $s_l \in \mathbb{R}^{BT_l p_l}$ is the layer's output, $\mathbf{W}_l \in \mathbb{R}^{d_l p_l}$ is the weight, $\mathbf{b}_l \in \mathbb{R}^{p_l}$ is the bias, and ϕ_l is any inter-layer operation such as ReLU, tanh or pooling. We denote *B* as the physical micro-batch size⁵ and T_l as the hidden feature dimension (e.g. sentence length or number of pixels). During the forward propagation, the activations $\{a_l\}$ are computed and stored in the computation graph, and the loss $L = \sum_i L_i$ is derived, where L_i is the per-sample losses. During the back-propagation, the output gradient is first computed based on the previous layer.

$$\frac{\partial L}{\partial \boldsymbol{s}_l} = \frac{\partial L}{\partial \boldsymbol{s}_{l+1}} \frac{\partial \boldsymbol{s}_{l+1}}{\partial \boldsymbol{a}_{l+1}} \circ \frac{\partial \boldsymbol{a}_{l+1}}{\partial \boldsymbol{s}_l} = \frac{\partial L}{\partial \boldsymbol{s}_{l+1}} \mathbf{W}_{l+1} \circ \phi_l'(\boldsymbol{s}_l),$$

in which \circ is element-wise multiplication. Specifically, the use of parameter \mathbf{W}_{l+1} necessitates the all-gather operation when the model is partitioned into multiple GPUs, which is not needed in single GPU training. Next, the activation a_l is used together with $\frac{\partial L}{\partial s_l}$ to compute the parameter gradient:

DP gradient:
$$\frac{\partial \sum_{i} C_{i} L_{i}}{\partial \mathbf{W}_{l}} + \sigma \mathcal{N}(0, \mathbf{I}) = \boldsymbol{a}_{l}^{\top} \operatorname{diag}(C_{1}, \cdots, C_{B}) \frac{\partial L}{\partial \boldsymbol{s}_{l}} + \sigma \mathcal{N}(0, \mathbf{I}).$$

⁵The micro-batch size *B* is the number of samples processed by each GPU, which determines the time and memory efficiency, but not the performance. The logical batch size that determines the performance is $B \times N_d \times$ gradient_accumulation_steps.



Figure 3: Algorithm of DP-ZeRO with mixed-precision training.

Note the standard gradient can be viewed as $C_i = 1, \sigma = 0$. Here the per-sample gradient norm (or the clipping factor C_i) can be computed at small cost, as we have discussed in Section 1.

3.2 TIME EFFICIENCY OF DP-ZERO

The time efficiency of DP-ZeRO consists of two parts: the local computation (including forward and backward propagation) and the global communication (including intra-node and inter-node communication). Given that the only difference between DP-ZeRO and ZeRO is the back-propagation, we claim that DP-ZeRO could enjoy high efficiency on-par with the standard ZeRO when (I) DP back-propagation exhibits a time efficiency comparable to the standard, similar to the single GPU training, and/or (II) the time efficiency of the parts other than back-propagation is not insignificant. We give the time of each part of DP-ZeRO in equation 3 to illustrate our claim.

$$\frac{\text{DP-ZeRO speed}}{\text{standard ZeRO speed}} = \frac{\text{standard back-prop} + \text{forward prop} + \text{communication}}{\text{DP back-prop} + \text{forward prop} + \text{communication}}$$
(3)

To be explicit, we summarize the time complexity in Table 2 and refer to Appendix B for details.

Table 2: Time complexity (measured by float-point operations) of one iteration in distributed learning⁷. We denote Ψ_{train} to be the number of trainable parameters ($\Psi_{\text{train}} = \Psi_{\text{model}}$ in full parameter training), and define B, T below equation 2.

| | forward propagation back-propagation | | communication | | | |
|------------|--------------------------------------|-----------|-------------------|-------------------------|------------------------------|-------------------|
| | activation | attention | output grad | param grad | DP clipping& noise | communication |
| complexity | $2BT\Psi_{model}$ | $O(BT^2)$ | $2BT\Psi_{model}$ | $2BT\Psi_{	ext{train}}$ | $0.666BT\Psi_{\text{train}}$ | $O(\Psi_{model})$ |

In what follows, we analyze the absolute and relative speed (to standard ZeRO) of DP-ZeRO under important settings.

³²¹ ⁷Here 0.666 is figurative and dependent on the DP mechanisms. For example, if we denote d, p as a layer's ³²² input/output dimensions, then the complexity of BK algorithm Bu et al. (2023b) is $6BTpd + 2BT^2(p+d)$. ³²³ $\mathbb{I}(2T^2 < pd)$, which is non-linear in T but empirically the overall slowdown could be $0.666/6 \approx 11\%$. Notice that $\Psi_{\text{train}} \ll \Psi_{\text{model}}$ when most parameters are frozen, as is the case in parameter efficient fine-tuning.

324 3.2.1 NUMBER OF COMPUTATION DEVICES

When scaling from one GPU (zero communication) to one node (multiple GPUs) and to multiple nodes, the communication efficiency decreases sub-linearly (\uparrow communication). On a single node, multiple GPUs can communicate using the high-speed intra-node connections such as NVLink/NVSwitch (Foley & Danskin, 2017; Ishii et al., 2018). On multiple nodes, which are necessary for large models, the inter-node connections are $3 \sim 24 \times$ slower than the intra-node connections (Li et al., 2019; Zhang et al., 2022). In short, DP-ZeRO can be as fast as ZeRO by equation 3 when multiple nodes are employed.

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3.2.2 MEMORY-EFFICIENT DISTRIBUTED LEARNING

335 The communication volume is specific to different distributed algorithms, most of which trade the 336 communication or speed for memory, in order to feasibly train very large models. For example, 337 ZeRO3 (but not ZeRO1/2) needs to all-gather the sharded parameters at each iteration, hence suffer-338 ing from 50% extra communication volume (^ccommunication). Another example is the activation 339 check-pointing (also known as gradient check-pointint Chen et al. (2016)), where a second forward propagation re-computes the expensive activations during back-propagation, though at a 33% 340 slower speed (*forward propagation*). These techniques improve the relative speed of DP-ZeRO but 341 worsens the absolute speed. 342

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3.2.3 PARAMETER EFFICIENT FINE-TUNING

345 Parameter efficient fine-tuning (PEFT), such as LoRA (Hu et al., 2021), Adapter (Houlsby et al., 346 2019), and BiTFiT (Zaken et al., 2022), optimizes a small fraction (e.g. $\Psi_{\text{train}} = 0.1\%\Psi_{\text{model}}$) 347 of model parameters and thus boosts the efficiency of back-propagation and communication (UDP) 348 back-propagation *lack-propagation lack-propagation*. Consequently, (I) the communication vol-349 ume of the gradient can be reduced possibly by $1000 \times$; (II) the local computation can accelerate by 350 50% (Hu et al., 2021; Bu et al., 2022b), which can be seen by treating Ψ_{train} in Table 2 as almost 351 zero; (III) the memory cost is saved on the non-trainable layers, which translates to larger batch size 352 and faster computation. Hence, both relative and absolute speed of DP-ZeRO improve using PEFT.

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3.3 MEMORY EFFICIENCY OF DP-ZERO

We claim that DP-ZeRO is as memory efficient as the standard ZeRO, similar to the single GPU training, when we use (I) the mixed ghost norm trick Bu et al. (2022a; 2023b), instead of GhostClip (Goodfellow, 2015; Li et al., 2021) or per-sample gradient instantiation (Yousefpour et al., 2021); (II) the layer-wise clipping style instead of the all-layer clipping, so that the book-keeping (Bu et al., 2023b) does not store all output gradients; or (III) a large number of GPUs so that the micro-batch size *B* (i.e. per-GPU batch size) is small: specially, when B = 1 in the gradient accumulation, the per-sample gradient is free. We empirically verify our claim in Figures 4 and 7.





378 3.4 MIXED-PRECISION TRAINING WITH DP

380 We now analyze the intricacy in mixed-precision training with 381 DP, which is not unique to DP-ZeRO but present in the general DP optimization. We emphasize that the per-sample gra-382 dient clipping already plays the role of scaling, and hence DP 383 mixed-precision training must not use loss scaling, as illus-384 trated in Table 3. Specifically, in standard mixed-precision 385 training, there are two steps of scaling: (I) scaling up the 386 loss L_i by $10^3 \sim 10^9$ (and consequently the output gradient 387 $\frac{\partial L_i}{\partial \mathbf{s}}$ as well as $\frac{\partial L_i}{\partial \mathbf{W}}$) before the back-propagation, to prevent 388 the underflow where fp16 gradient is too small to be distin-389 guished from 0, and (II) scaling down the parameter gradient 390 $\frac{\partial L_i}{\partial \mathbf{W}}$, by the same factor, after the back-propagation to recover 391 the correct magnitude of gradient. However, in DP mixed-392 precision training, scaling up the loss may cause overflow, 393



Figure 5: Accuracy of mixedprecision training with loss scaling. ViT-large on CIFAR100.

while scaling down the gradient incorrectly over-shrinks the gradient and worsens the performance. See Figure 5 for a real example. We explain this intricacy step-by-step in Appendix C.

Table 3: Illustration of overflow and underflow issues during mixed-precision training (ghost norm).

| | | | | 0 | 1 | U | ί, C |
|----------------------------|---------------------|---|--|---|------------------------|-----------------------|-------------------------|
| loss scale=10 ³ | activation a | output grad $\frac{\partial L}{\partial L}$ | per-samp | le grad norm | clipping factor | param grad | param grad |
| 1055 Scale=10 | activation u_l | (scaled) ∂s_l | $\operatorname{vec}(\boldsymbol{a}_l \boldsymbol{a}_l^{\top})$ | $\operatorname{vec}\left(\frac{\partial L}{\partial s_l} \frac{\partial L}{\partial s_l}^{\top}\right)$ | enpping factor | (not scaled down) | (if scaled down) |
| standard w/o scaling | $10^{-3} \sim 10^2$ | $10^{-8} \sim 10^{1}$ | N/A | N/A | 1 | $10^{-7} \sim 10^{1}$ | $10^{-7} \sim 10^{1}$ |
| standard w/ scaling | $10^{-3} \sim 10^2$ | $10^{-5} \sim 10^4$ | N/A | N/A | 1 | $10^{-4} \sim 10^4$ | $10^{-7} \sim 10^{1}$ |
| DP w/o scaling | $10^{-3} \sim 10^2$ | $10^{-8} \sim 10^{1}$ | $10^{2} \sim 10^{3}$ | $10^{-6} \sim 10^{0}$ | $10^{-3} \sim 10^2$ | $10^{-7} \sim 10^{1}$ | $10^{-7} \sim 10^{1}$ |
| DP w/ scaling | $10^{-3} \sim 10^2$ | $10^{-5} \sim 10^4$ | $10^{2} \sim 10^{3}$ | $10^0 \sim 10^6$ | $10^{-6} \sim 10^{-1}$ | $10^{-7} \sim 10^{1}$ | $10^{-10} \sim 10^{-2}$ |
| | | | | | | | |

Remark 3.1. DP may be violated due to the floating-point vulnerability at any precision Mironov (2012), but the issues can be more severe at half-precision. Yet, it remains an open discussion whether precision-based attacks can non-trivially break the protection of DP in deep learning, since the model is updated through thousands of iterations which may cover the identifiability of data information. In fact, Bu et al. (2024); Yu et al. (2021b); Li et al. (2024) have demonstrated that DP mixed-precision training can empirically preserve the protection against various privacy attacks.

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4 EMPIRICAL PERFORMANCE OF DP-ZERO

412 We evaluate DP-ZeRO on five aspects : model architectures, efficiency, scalability, compatibility 413 with various distributed learning and DP techniques. We use DP-ZeRO to train ResNet (He et al., 414 2016), ViT (Dosovitskiy et al., 2020; Zhai et al., 2022) and GPT (Radford et al., 2019; Brown 415 et al., 2020), which are workhorses in computer vision and language tasks. We report two accuracy experiments in Appendix E.4 for ViT and GPT to show that the performance is consistent across 416 systems and to verify our implementation. We do not aim for higher accuracy with larger models, 417 because the accuracy depends on many training hyperparameters like learning rate and number of 418 epochs that are not relevant to the system designs. In fact, this is the norm of system papers including 419 the standard non-DP ZeRO Rajbhandari et al. (2020) and FSDP Zhao et al., where the focus is on 420 the system efficiency. 421

We measure the time and memory efficiency of DP-ZeRO under settings such as PEFT and multiple precision formats (fp32 or fp16/bf16). We evaluate the scalability of DP-ZeRO in terms of number of GPUs and number of model parameters. Our experiments scale from single node (8 GPUs) to multiple nodes, up to 256 GPUs, and train models up to 100B trainable parameters. Moreover, DP-ZeRO is compatible with mainstream implementations of ZeRO⁸ and with different clipping styles, clipping functions, privacy accountants, and so on. We leave the experimental details in Appendix D. By default, we use AdamW, mixed-precision training, layer-wise clipping style, B = 4, and A100 GPU with 40GB memory, unless otherwise stated.

 ⁸DP-ZeRO is implemented on DeepSpeed (supporting ZeRO1/2/3), FSDP (Zhao et al.) (supporting ZeRO3), MiCS (Zhang et al., 2022) (supporting ZeRO2/3), and any distributed optimizers supported on these libraries.

432 4.1 GENERALITY OF DP-ZERO 433

434 DP-ZeRO is generally applicable to different neural network architectures, clipping styles, and precision formats. We test DP-ZeRO1 on single node and observe that different clipping styles are 435 equally fast, but layer-wise clipping is more memory efficient than all-layer clipping. Comparing to 436 the standard ZeRO, our DP-ZeRO enjoys almost the same speed and memory efficiency, while the 437 gap will be further closed as we move to more advanced stages of ZeRO. 438

4.2 LIGHTER TRAINING OF DP-ZERO



449 Figure 6: Efficiency of DP-ZeRO with lighter training. Left&Middle: $> 2 \times$ speedup with lighter 450 training; ViT-5B, B = 1. Right: DP-ZeRO benefits more than ZeRO from PEFT.

451 DP-ZeRO can employ low-memory optimizers and train on fewer parameters, therefore vastly reduc-452 ing the memory and communication cost. On a single node, we demonstrate that DP-ZeRO actually 453 benefits (more than standard ZeRO and single-GPU training) from lighter training, following from 454 our discussion in Section 3.2. 455

456 **Low-memory optimizers** Low-memory optimizers can boost the training efficiency at the cost of 457 accuracy degradation. For example, SGD requires only 1/3 optimizer states of Adam and significantly saves the memory; 1-bit Adam (Tang et al., 2021) and signSGD compress the gradient and 458 reduce the communication volume up to $32 \times$. 459

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Fewer trainable parameters In the fine-tuning phase, as we analyzed in Section 3.2.3, PEFT improves both the local computation and the communication volume. Hence, DP-ZeRO allows 462 PEFT on ViT and GPT to be $\approx 2 \times$ faster than full fine-tuning, whereas the single GPU acceleration 463 is $\leq 1.5 \times$.

Remark 4.1. We leverage DP-ZeRO3 with SGD to train ViT-10B (full parameters) and ViT-22B 465 (PEFT; 1M trainable parameters) on one node. See Appendix D for configurations. 466

467 4.3 THREE STAGES OF DP-ZERO 468

DP-ZeRO supports all stages of ZeRO under different implementations including DeepSpeed (de-469 fault) and FSDP. 470

471 In Figure 7, the efficiency of DP-ZeRO catches up with the standard ZeRO when we move up the 472 stages. For instance of ViT-Gigantic, the throughput increases from 83% by DP-ZeRO1 to 95-97% 473 by DP-ZeRO3. Following equation 3, we can attribute the relatively fast training of DP-ZeRO to 474 the increase cost of communication, especially in DP-ZeRO3. Additionally, we observe that the throughput of DP-ZeRO1/2 improves to over 95% on 4 nodes, as predicted by Section 3.2.1. Notice 475 that we save DP-ZeRO3 of GPT to Section 4.4 on super-large scale. 476

- 477 478
- 4.4 SCALABILITY OF DP-ZERO

479 We evaluate the scalability of DP-ZeRO3 in terms of large sequence length (2048), large model size 480 $(7 \sim 100B)$, and large number of GPUs (up to 256). We use A100 with 80GB memory, as well as 481 the activation check-pointing and ModelP. 482

In Figure 8 (left), we observe that for a fixed model with 26B trainable parameters, DP-ZeRO is 483 super-linearly scalable to the number of GPUs, achieving > 95% speed of the standard ZeRO. Here 484 super-linearity is a property of ZeRO (see Figure 3 in Rajbhandari et al. (2020)) which allows more 485 GPUs to shard the model states (and reduce the per-GPU memory cost) more aggressively, and to



Figure 7: Efficiency of DP-ZeRO on ViT-Gigantic and GPT2-XL under different implementations.

train faster since the micro-batch size is larger. Furthermore, in Figure 8 (right), for a fixed number of GPUs, DP-ZeRO is linearly scalable to the model size, achieving the same speed and scalability as the standard ZeRO. In short, DP-ZeRO is almost equal to ZeRO in terms of training efficiency in super-large scale.



Figure 8: Scalability of DP-ZeRO3 on 26B model (left) and 256 GPUs (right). The colored lines match the left y-axis. The grey bars match the right y-axis. *max B* means we use the maximum micro-batch in each GPU and *fixed B* means B = 2.

Remark 4.2. In comparison to DP-ZeRO, DataP (DP or standard) at most fits 5B models (Rajbhandari et al., 2020) in 80GB memory, regardless of the number of GPUs. We cannot compare to DP-PipeP in He et al. (2022) because the codebase and experiment details (e.g. number of trainable parameters and sequence length) are not publicly available. Nevertheless, since DP-ZeRO resembles the efficiency of standard ZeRO, the usefulness of DP-ZeRO can be demonstrated by comparing ZeRO to PipeP.

5 DISCUSSION

In this work, we develope DP-ZeRO that enables the optimization of models up to 100B trainable parameters, thus allowing DP distributed learning to be as efficient and scalable as the standard one, for as few as 2 and as many as hundreds of GPUs. We believe this is a significant milestone to pave the path towards DP foundation models, especially for its open-source nature (link to be released).

We emphasize that, since DP only modifies the back-propagation, our DP-ZeRO is orthogonal to
any large-scale training techniques that are not tied to back-propagation: for example, activation
check-pointing, CPU offloading, weight/activation quantization (Dettmers et al., 2023; Xiao et al., 2023), and tensor parallelism (Narayanan et al., 2021) in 3D parallelism (e.g. Megatron).

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A THE BOOK-KEEPING (BK) ALGORITHM

A.1 EFFICIENT COMPUTATION OF PER-SAMPLE GRADIENT NORMS

The mixed ghost norm Bu et al. (2022a) is the state-of-the-art technique to compute the per-sample gradient norm of the **weight**, almost for free. It hybridizes two basic techniques – the per-sample gradient instantiation and the ghost norm – to compute the Frobenius norm of weight gradient,

$$\left\|\boldsymbol{a}_{l,i}^{\top} \frac{\partial L}{\partial \boldsymbol{s}_{l,i}}\right\|^{2} \stackrel{\text{per-sample grad}}{=} \left\|\frac{\partial L_{i}}{\partial \mathbf{W}_{l}}\right\|_{\text{Fro}}^{2} \stackrel{\text{ghost norm}}{=} \operatorname{vec}\left(\frac{\partial L}{\partial \boldsymbol{s}_{l,i}} \frac{\partial L}{\partial \boldsymbol{s}_{l,i}}^{\top}\right) \cdot \operatorname{vec}(\boldsymbol{a}_{l,i} \boldsymbol{a}_{l,i}^{\top})$$
(4)

where "vec" flattens the tensor to an one-dimensional vector. In words, Equation (4) gives two equations that are equivalent mathematically, but significantly different in efficiency:

- $||X||_{\text{Fro}}^2 = ||A^{\top}B||^2$ firstly computes $A^{\top}B$ and then its norm.
- $||X||_{\text{Fro}}^2 = \text{vec}(AA^{\top}) \cdot \text{vec}(BB^{\top})$ firstly computes $AA^{\top}, BB^{\top} \in \mathbb{R}^{TT}$ and then their dot product.

In summary, the mixed ghost norm always applies the cheaper of two techniques at each layer of a neural network.

Finally, we note that the per-sample gradient norm of the **bias** is computed differently. This is because $\partial I = \partial I$

$$\frac{\partial L_i}{\partial b_l} = \mathbf{1}^\top \frac{\partial L}{\partial \boldsymbol{s}_{l,i}}$$

is not actually a product of tensors like $X = A^{\top}B$. In fact, the multiplication with 1 turns out to be a summation along the first dimension, and it suffices to use per-sample gradient instantiation for the bias.

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A.2 BOOK-KEEPING THE OUTPUT GRADIENT

The BK algorithm uses two rounds of back-propagation (though each round only takes half the complexity, hence the total complexity of DP back-propagation matches the non-DP back-propagation). Therefore, output gradients $\frac{\partial L}{\partial s_{l,i}}$ are kept to avoid repeated computation. Notice that the output gradient are relatively cheap to book-keep (see Figure 4 and Figure 5 in Bu et al. (2023b)).

B COMPONENT-WISE TIME COMPLEXITY OF DP-ZERO

⁷⁷⁵ In this section, we explain the time complexity of each part in Table 2, and demonstrate how the complexity can be different under different settings.

Forward propagation: The matrix multiplication during forward propagation results in $2BT\Psi_{model}$ complexity (see Bu et al. (2022a)). Notice that, if the activation check-pointing is used, essentially two rounds of forward propagation take place in one iteration. Hence the time complexity doubles and becomes $4BT\Psi_{model}$.

Back-propagation: This contains two sub-processes: the output gradients are computed at all layers, taking $2BT\Psi_{model}$ complexity; the parameter gradients are computed only at trainable layers (a few if doing PEFT), taking $2BT\Psi_{train}$ complexity. Clearly, in full parameter training, the total is $4BT\Psi_{model}$, and in PEFT, about $2BT\Psi_{model}$.

Attention: The time complexity of attention is $O(BT^2)$ in Vaswani et al. (2017), where *T* is the sequence length (a.k.a. token length). When *T* is large, e.g. training with long context like T = 8192, this cost is prohibitively high. In this regard, a line of researches have proposed linear complexity attention, including but not limited to Wang et al. (2020); Katharopoulos et al. (2020); Shen et al. (2021).

Communication: For algorithms that don't shard the model, such as data parallelism and ZeRO1/2, the communication is only used to send gradients and optimizer states. Hence the communication volume is proportional to the number of trainable parameters $O(\Psi_{\text{train}})$. Otherwise, for algorithms such as ZeRO3 and tensor parallelism, the communication volume is proportional to the number of total parameters $O(\Psi_{\text{model}})$, because the forward propagation needs to gather the parameters from many GPUs. This makes a big difference in PEFT when $\Psi_{\text{train}} \ll \Psi_{\text{model}}$.

C LOSS SCALING IN MIXED-PRECISION TRAINING

800 We write the per-sample gradient with loss scaling S as

$$\frac{\partial C_i L_i}{\partial \mathbf{W}_l} = C_i \frac{1}{S} \cdot \left(\boldsymbol{a}_{l,i}^\top \left(S \cdot \frac{\partial L}{\partial \boldsymbol{s}_{l,i}} \right) \right)$$

This covers the standard gradient ($C_i = 1$) and DP gradient (e.g. $C_i = 1/||g_i||$, computed by the mixed ghost norm in Appendix A), in which S enlarges the output gradient to avoid underflow, and $\frac{1}{S}$ shrinks the parameter gradient to the correct magnitude.

Recall that a standard mixed-precision training (with loss scaling) uses steps $1 \rightarrow 2 \rightarrow 3 \rightarrow 5 \rightarrow 6$ ⁹, or $1 \rightarrow 3 \rightarrow 6$ without loss scaling.

[%]See https://docs.nvidia.com/deeplearning/performance/ mixed-precision-training/index.html#lossscaling.

810 1. Forward propagation (fp16 weights and activations) and get the loss. 811 2. Scaling up the loss by a factor S. 812 813 3. Backward propagation on the scaled loss (fp16 parameters and their gradients). 814 4. Per-sample gradient clipping (sensitivity = 1) and noising for DP. 815 816 5. Scaling down the parameter gradient by a factor of 1/S. 817 6. Update the parameters with their gradients. 818 819 If we follow the same procedure under the DP regime, say using a hook function to be called 820 after back-propagation creates the gradients like in Opacus (Yousefpour et al., 2021), Private-821 Transformers (Li et al., 2021), FastDP (Bu et al., 2023b), then the per-sample clipping factor is 822 scaled up S times so as to normalize the gradient. Hence per-sample gradient clipping has already 823 played the role of scaling down. If we scale down the gradient for a second time, the gradient is 824 incorrectly over-shrunk. This is the case in Yu et al. (2021a) and in the alternative implementation 825 of (Li et al., 2021, Appendix T) (see also Figure 5). To be sure, this approach is still DP, but the performance does not match fp32 DP training correctly, and usually degrades too much to be useful. 826 827 One walk-around is to prevent per-sample gradient clipping to scale down the gradients and let step 828 5 do its job, i.e. $1 \rightarrow 2 \rightarrow 3 \rightarrow 4^* \rightarrow 5 \rightarrow 6$. We note that (Li et al., 2021, Appendix T) follows 829 this path (though no experiment results or codes are available at the time of writing) by modifying 830 step 4: clipping threshold (sensitivity) = S instead of 1, so that the clipped gradient is S times larger 831 than the DP f32 training, to be scaled down by step 5. However, this introduces additional design

Another walk-around is to delete step 5 and let per-sample gradient clipping do its job, i.e. $1 \rightarrow 2 \rightarrow 3 \rightarrow 4 \rightarrow 6$. However, this approach is harder to implement because in the standard process step 2 and 5 are simultaneously enabled or disabled. Also we cannot prevent overflow when using fp16 as we still use step 2.

decisions and does not prevent overflow when using fp16 (due to step 2, see Table 3).

Therefore, we propose to not use loss scaling (or equivalently we set S = 1 statically for all steps) during DP mixed-precision training, i.e. $1 \rightarrow 3 \rightarrow 4 \rightarrow 6$. Although, by not using step 2, we cannot prevent underflow when using fp16, this is much less a problem compared to overflow: underflow (treating small values as 0) makes the training less accurate but does not fail the training like overflow (treating large values as NAN). Lastly, the underflow issue is perfectly mitigated by bf16, which we recommend for DP mixed-precision training whenever possible.

| | | steps | fp16 issue | note | reference |
|----|--------|--------------|------------|---------------------------------|----------------------------|
| st | andard | 136 | underflow | | Micikevicius et al. (2018) |
| st | andard | 12356 | none | | Micikevicius et al. (2018) |
| | DP | 123456 | overflow | incorrect due to over-shrinking | Li et al. (2021) |
| | DP | $1234^{*}56$ | overflow | different clipping threshold | Li et al. (2021) |
| | DP | 1346 | underflow | perfect with bf16 | ours |
| | DP | 12346 | overflow | hard to implement | ours |

Table 4: Mixed-precision training with DP or not.

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EXPERIMENT SETTINGS

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Datasets: To evaluate the efficiency, it suffices to declare the data's dimension (e.g. micro-batch size and feature dimension) without specifying the dataset (though sometimes specifying the dataset means declaring the dimension, e.g. MNIST usually means 28*28 pixels). This is the norm in system papers such as Rajbhandari et al. (2020; 2021); Zhao et al.. In this work, vision models are trained with 224*224 pixels at ImageNet scale; GPT models are trained with sequence length 100, except in Figure 8 where sequence length is 2048.

Figure 5 and Table 3: We train ViT-large (300M parameters) and CIFAR100, 5 epochs, learning rate 5e-4, logical batch size 1000.

Figure 6: To fit as large a model as possible, we set B = 1 and use SGD. We set 48 attention heads, 21 layers, MLP=4*width (also known as embedding dimension), and modify width for all models. For instance, ViT-10B uses width=768 * 22, ViT-22B uses width=768 * 34.

Figure 8: We train AdamW with layer-wise clipping. DP distributed learning is based on MiCS (ZeRO3) using bf16 mixed-precision training. Most of GPT configuration is the same as Touvron et al. (2023a) (Table 2) in terms of embedding dimension, attention heads and number of layers. However, GPT-100B uses the configuration from Brown et al. (2020) (Table 2.1) but a smaller width.

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- E CODEBASE DESIGN
- E.1 WITH FORWARD & BACKWARD HOOKS

Hooks¹⁰ are important functions to enrich the deep learning optimization. To be specific, there are

- 1. forward modular hook (nn.register_forward_hook),
- 2. backward modular hook (nn.register_backward_hook),
- 3. backward tensor hook (tensor.register_hook).

884 DP libraries including Opacus Yousefpour et al. (2021), Private-transformers Li et al. (2021), 885 Private-Vision Bu et al. (2022a), FastDP Bu et al. (2023b;a), FastGradClip Lee & Kifer (2021) 886 and so on, use modular hooks to modify the standard optimization. However, ZeRO libraries in-887 cluding DeepSpeed and FSDP use tensor hooks. This difference in the types of hooks and many other differences (e.g. both ZeRO libraries and DP libraries modify the optimizer's step function) 889 cause non-trivial problems when combining DP with ZeRO. For example, to keep DP optimization 890 as efficient as the standard, it is necessary to not waste time on computing the non-private gradient. However, if we skip such computation, then ZeRO's tensor hook will not be triggered and the corre-891 sponding distributed-learning-related operations cannot carry on. For another example, because DP 892 and ZeRO add different types of hooks, the number of hooks is larger than either optimization and 893 they slows down the training: consider an 100-layer network, each layer with weight and bias (2 ten-894 sors), then DP-ZeRO in this subsection needs 100 modular hooks and 200 tensor hooks, adding to a 895 total of 300 hooks. In addition, the Book-Keeping algorithm (in FastDP) in its original form cannot 896 be implemented together with ZeRO3, because all model states are partitioned including the output 897 gradients which are meant to be book-kept. To work around this requires rewriting the distributed so-898 lution's communication mechanism, and if successful, still requiring additional communication cost 899 during the second back-propagation. Similar problems are present for Opacus and FastGradClip, 900 which instantiates per-sample gradients that will be partitioned in ZeRO2/3 and requires additional 901 communication cost when gathered to create the privatized gradient.

As a consequence, the hooks are fully supported on DP-ZeRO1 and partially supported on DP-ZeRO2/3 under the layer-wise clipping.

E.2 WITHOUT HOOKS

Instead of registering hooks on top of the original (non-DP) back-propagation, we can directly modify the back-propagation following Appendix A: e.g., given the activation and output gradient,

$$\frac{\partial C_i L_i}{\partial \mathbf{W}_l} = \mathbf{a}_{(l),i}^{\top} \frac{\partial L}{\partial \mathbf{s}_{(l),i}} \Big/ \sqrt{\operatorname{vec}\left(\frac{\partial L}{\partial \mathbf{s}_{(l),i}} \frac{\partial L}{\partial \mathbf{s}_{(l),i}}^{\top}\right) \cdot \operatorname{vec}(\mathbf{a}_{(l),i} \mathbf{a}_{(l),i}^{\top})}$$

This approach requires rewriting the back-propagation for each layer type (linear, embedding, convolution, normalization, ...) and can be done at different levels (Pytorch, C++, CUDA kernel).

¹⁰See https://pytorch.org/tutorials/beginner/former_torchies/nnft_ tutorial.html#forward-and-backward-function-hooks.

918 E.3 USER INTERFACE

920 DP-ZeRO can be enabled by one piece of code: after the model is instantiated,

```
922 privacy_engine = PrivacyEngine(model,
923 batch_size=256, sample_size=50000,
924 epochs=3, target_epsilon=3)
```

The codebase is designed not to modify the optimizer, hence DP-ZeRO can work with arbitrary op-timizer. Because of this design, our DP-ZeRO will not distinguish micro-batches. This is different from the gradient accumulation in Opacus (version ==0.x) and Private-Vision, where only the last micro-batch is processed by "optimizer.step()" but all other micro-batches are processed by "opti-mizer.virtual_step()". In other words, the noise $\sigma_{DP}N(0, I)$ is added on the last micro-batch, after the micro-batches are accumulated. But DP-ZeRO adds the noise on each micro-batch equally. Note that the noise level per micro-batch is $\sigma_{\rm DP}/\sqrt{N_d}$ if a random seed is set across N_d GPUs, or $\sigma_{\rm DP}/N_d$ otherwise.

934 E.4 VERIFICATION OF OUR IMPLEMENTATION

DP-ZeRO is implementing the same DP-SGD (hence the same utility) as in Abadi et al. (2016), only
 in a much more scalable and distributed manner than libraries like Private Transformers and Opacus.

To be sure, we provide the results on 5 runs of ViT (image classification) and GPT2 (text generation), at $\epsilon = 8$, with standard deviations in the parenthesis. We compare to other libraries (without mixed precision) and our DP-ZeRO uses mixed precision, in order to validate both (I) the correctness of DP mixed precision and (II) the correctness of our implementation of DP-ZeRO.

| | ViT/CIFAR100/accuracy | GPT2/E2E/BLEU |
|-----------|-----------------------|---------------|
| Opacus | 83.45(0.28) | 63.35(0.50) |
| FastDP | 83.46(0.19) | 64.01(0.54) |
| GhostClip | 83.51(0.33) | 63.71(0.31) |
| DP-ZeRO1 | 83.46(0.13) | 64.03(0.41) |
| DP-ZeRO2 | 83.43(0.09) | 63.72(0.47) |
| DP-ZeRO3 | 83.48(0.18) | 63.92(0.39) |