Scaling Neural Machine Translation

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Abstract

Sequence to sequence learning models still require several days to reach state of the art performance on large benchmark datasets using a single machine. This paper shows that reduced precision and large batch training can speedup training by nearly 5x on a single 8-GPU machine with careful tuning and implementation.1 On WMT’14 English-German translation, we match the accuracy of Vaswani et al. (2017) in under 5 hours when training on 8 GPUs and we obtain a new state of the art of 29.3 BLEU after training for 85 minutes on 128 GPUs. We further improve these results to 29.8 BLEU by training on the much larger Paracrawl dataset. On the WMT’14 English-French task, we obtain a state-of-the-art BLEU of 43.2 in 8.5 hours on 128 GPUs.

1 Introduction

Neural Machine Translation (NMT) has seen impressive progress in the recent years with the introduction of ever more efficient architectures (Bahdanau et al., 2015; Gehring et al., 2017; Vaswani et al., 2017). Similar sequence-to-sequence models are also applied to other natural language processing tasks, such as abstractive summarization (See et al., 2017; Paulus et al., 2018) and dialog (Sordoni et al., 2015; Serban et al., 2017; Dusek andJurcícˇek, 2016).

Currently, training state-of-the-art models on large datasets is computationally intensive and can require several days on a machine with 8 high-end graphics processing units (GPUs). Scaling training to multiple machines enables faster experimental turn-around but also introduces new challenges: How do we maintain efficiency in a distributed setup when some batches process faster than others (i.e., in the presence of stragglers)? How do larger batch sizes affect optimization and generalization performance? While stragglers primarily affect multi-machine training, questions about the effectiveness of large batch training are relevant even for users of commodity hardware on a single machine, especially as such hardware continues to improve, enabling bigger models and batch sizes.

In this paper, we first explore approaches to improve training efficiency on a single machine. By training with reduced floating point precision we decrease training time by 65% with no effect on accuracy. Next, we assess the effect of dramatically increasing the batch size from 25k to over 400k tokens, a necessary condition for large scale parallelization with synchronous training. We implement this on a single machine by accumulating gradients from several batches before each update. We find that by training with large batches and by increasing the learning rate we can further reduce training time by 40% on a single machine. Finally, we parallelize training across 16 machines and find that we can reduce training time by an additional 90% compared to a single machine.

Our improvements enable training a Transformer model on the WMT’16 En-De dataset to the same accuracy as Vaswani et al. (2017) in just 32 minutes on 128 GPUs and in under 5 hours on 8 GPUs. This same model trained to full convergence achieves a new state of the art of 29.3 BLEU in 85 minutes. These scalability improvements additionally enable us to train models on much larger datasets. We show that we can reach 29.8 BLEU on the same test set in less than 10 hours when trained on a combined corpus of WMT and Paracrawl data containing ∼150M sentence pairs (i.e., over 30x more training data). Similarly, on the WMT’14 En-Fr task we obtain a state of the art BLEU of 43.2 in 8.5 hours on 128 GPUs.

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1Our implementation is available at: https://www.github.com/pytorch/fairseq
2 Related Work

Previous research considered training and inference with reduced numerical precision for neural networks (Simard and Graf, 1993; Courbariaux et al., 2015; Sa et al., 2018). Our work relies on half-precision floating point computation, following the guidelines of Micikevicius et al. (2018) to adjust the scale of the loss to avoid underflow or overflow errors in gradient computations.

Distributed training of neural networks follows two main strategies: (i) model parallel evaluates different model layers on different workers (Coates et al., 2013) and (ii) data parallel keeps a copy of the model on each worker but distributes different batches to different machines (Dean et al., 2012). We rely on the second scheme and follow synchronous SGD, which has recently been deemed more efficient than asynchronous SGD (Chen et al., 2016). Synchronous SGD distributes the computation of gradients over multiple machines and then performs a synchronized update of the model weights. Large neural machine translation systems have been recently trained with this algorithm with success (Dean, 2017; Chen et al., 2018).

Recent work by Puri et al. (2018) considers large-scale distributed training of language models (LM) achieving 109x scaling with 128 GPUs. Compared to NMT training, however, LM training does not face the same challenges of variable batch sizes. Moreover, we find that large batch training requires warming up the learning rate, whereas their work begins training with a large learning rate. There has also been recent work on using lower precision for inference only (Quinn and Ballesteros, 2018).

Another line of work explores strategies for improving communication efficiency in distributed synchronous training setting by abandoning “stragglers,” in particular by introducing redundancy in how the data is distributed across workers (Tandon et al., 2017; Ye and Abbe, 2018). The idea rests on coding schemes that introduce this redundancy and enable for some workers to simply not return an answer. In contrast, we do not discard any computation done by workers.

3 Experimental Setup

3.1 Datasets and Evaluation

We run experiments on two language pairs, English to German (En–De) and English to French (En–Fr). For En–De we replicate the setup of Vaswani et al. (2017) which relies on the WMT’16 training data with 4.5M sentence pairs; we validate on newstest13 and test on newstest14. We use a vocabulary of 32K symbols based on a joint source and target byte pair encoding (BPE; Sennrich et al. 2016). For En–Fr, we train on WMT’14 and borrow the setup of Gehring et al. (2017) with 36M training sentence pairs. We use newstest12+13 for validation and newstest14 for test. The 40K vocabulary is based on a joint source and target BPE factorization.

We also experiment with scaling training beyond 36M sentence pairs by using data from the Paracrawl corpus (ParaCrawl, 2018). This dataset is extremely large with more than 4.5B pairs for En–De and more than 4.2B pairs for
En–Fr. We rely on the BPE vocabulary built on WMT data for each language pair and explore filtering this noisy dataset in Section 4.5. We measure case-sensitive tokenized BLEU with multi-bleu.pl and de-tokenized BLEU with SacreBLEU (Post, 2018). All results use beam search with a beam width of 4 and length penalty of 0.6, following Vaswani et al. 2017. Checkpoint averaging is not used, except where specified otherwise.

### 3.2 Models and Hyperparameters

We use the Transformer model (Vaswani et al., 2017) implemented in PyTorch in the fairseq-py toolkit (Edunov et al., 2017). All experiments are based on the “big” transformer model with 6 blocks in the encoder and decoder networks. Each encoder block contains a self-attention layer, followed by two fully connected feed-forward layers with a ReLU non-linearity between them. Each decoder block contains self-attention, followed by encoder-decoder attention, followed by two fully connected feed-forward layers with a ReLU between them. We include residual connections (He et al., 2015) after each attention layer and after the combined feed-forward layers, and apply layer normalization (Ba et al., 2016) after each residual connection. We use word representations of size 1024, feed-forward layers with inner dimension 4,096, and multi-headed attention with 16 attention heads. We apply dropout (Srivastava et al., 2014) with probability 0.3 for En-De and 0.1 for En-Fr. In total this model has 210M parameters for the En-De dataset and 222M parameters for the En-Fr dataset.

Models are optimized with Adam (Kingma and Ba, 2015) using $\beta_1 = 0.9$, $\beta_2 = 0.98$, and $\epsilon = 1e-8$. We use the same learning rate schedule as Vaswani et al. (2017), i.e., the learning rate increases linearly for 4,000 steps to $5e-4$ (or $1e-3$ in experiments that specify $2x\ 1r$), after which it is decayed proportionally to the inverse square root of the number of steps. We use label smoothing with 0.1 weight for the uniform prior distribution over the vocabulary (Szegedy et al., 2015; Pereyra et al., 2017).

All experiments are run on DGX-1 nodes with 8 NVIDIA V100 GPUs interconnected by Infiniband. We use the NCCL2 library and torch.distributed for inter-GPU communication.

### 4 Experiments and Results

In this section we present results for improving training efficiency via reduced precision floating point (Section 4.1), training with larger batches (Section 4.2), and training with multiple nodes in a distributed setting (Section 4.3).

#### 4.1 Half-Precision Training

NVIDIA Volta GPUs introduce Tensor Cores that enable efficient half precision floating point (FP) computations that are several times faster than full precision operations. However, half precision drastically reduces the range of floating point values that can be represented which can lead to numerical underflows and overflows (Micikevicius et al., 2018). This can be mitigated by scaling values to fit into the FP16 range.

In particular, we perform all forward-backward computations as well as the all-reduce (gradient synchronization) between workers in FP16. In contrast, the model weights are also available in full precision, and we compute the loss and optimization (e.g., momentum, weight updates) in FP32 as well. We scale the loss right after the forward pass to fit into the FP16 range and perform the backward pass as usual. After the all-reduce of the FP16 version of the gradients with respect to the weights we convert the gradients into FP32 and restore the original scale of the values before updating the weights.

In the beginning stages of training, the loss needs to be scaled down to avoid numerical overflow, while at the end of training, when the loss is small, we need to scale it up in order to avoid numerical underflow. Dynamic loss scaling takes care of both. It automatically scales down the loss when overflow is detected and since it is not possible to detect underflow, it scales the loss up if no overflows have been detected over the past 2,000 updates.

To evaluate training with lower precision, we first compare a baseline transformer model trained on 8 GPUs with 32-bit floating point (Our reimplementation) to the same model trained with 16-
Table 1: Training time (min) for reduced precision (16-bit), cumulating gradients over multiple backwards (cumul), increasing learning rate (2x lr) and computing each forward/backward with more data due to memory savings (5k tkn/gpu). Average time (excl. validation and saving models) over 3 random seeds to reach validation perplexity of 4.32 (2.11 NLL). Cumul=16 means a weight update after accumulating gradients for 16 backward computations, simulating training on 16 nodes. WMT En-De, newstest13.

| model                        | # gpu | bsz  | cumul | BLEU updates | tkn/sec | time   | speedup |
|------------------------------|-------|------|-------|---------------|---------|--------|---------|
| Vaswani et al. (2017)        | 8×P100| 25k  | 1     | 26.4          | 300k    | ∼25k   | ∼5,000  |         |
| Our reimplemention            | 8×V100| 25k  | 1     | 26.4          | 192k    | 54k    | 1,429   | reference |
| +16-bit                      | 8     | 25k  | 1     | 26.7          | 193k    | 143k   | 495     | 2.9x     |
| +cumul                       | 8     | 402k | 16    | 26.7          | 13.7k   | 195k   | 447     | 3.2x     |
| +2x lr                       | 8     | 402k | 16    | 26.5          | 9.6k    | 196k   | 311     | 4.6x     |
| +5k tkn/gpu                  | 8     | 365k | 10    | 26.5          | 10.3k   | 202k   | 294     | 4.9x     |
| 16 nodes (from +2x lr)       | 128   | 402k | 1     | 26.5          | 9.5k    | 1.53M  | 37      | 38.6x    |
| +overlap comm+bwd            | 128   | 402k | 1     | 26.5          | 9.7k    | 1.82M  | 32      | 44.7x    |

Figure 2: Accumulating gradients over multiple forward/backward steps speeds up training by: (i) reducing communication between workers, and (ii) saving idle time by reducing variance in workload between GPUs.

4.2 Training with Larger Batches

Large batches are a prerequisite for distributed synchronous training, since it averages the gradients over all workers and thus the effective batch size is the sum of the sizes of all batches seen by the workers.

Figure 1 shows that bigger batches result in slower initial convergence when measured in terms of epochs (i.e. passes over the training set). However, when looking at the number of weight updates (i.e. optimization steps) large batches converge faster (Hoffer et al., 2017). These results support parallelization since the number of steps define the number of synchronization points for synchronous training.

Training with large batches is also possible on a single machine regardless of the number of GPUs or amount of available memory; one simply iterates over multiple batches and accumulates the resulting gradients before committing a weight update. This has the added benefit of reducing communication and reducing the variance in workload between different workers (see Figure 2), leading to a 36% increase in tokens/sec (Table 1, cumul). We discuss the issue of workload variance in more depth in Section 5.

Increased Learning Rate: Similar to Goyal et al. (2017) and Smith et al. (2018) we find that training with large batches enables us to increase the learning rate, which further shortens training time even on a single node (2x lr).

Memory Efficiency: Reduced precision also decreases memory consumption, allowing for larger sub-batches per GPU. We switch from a maximum of 3.5k tokens per GPU to a maximum of 5k tokens per GPU and obtain an additional 5% speedup (cf. Table 1; 2x lr vs. 5k tkn/gpu).

Table 1 reports our speed improvements due to reduced precision, larger batches, learning rate increase and increased per-worker batch size. Overall, we reduce training time from 1,429 min to 294 min to reach the same perplexity on the same hardware (8x NVIDIA V100), i.e. a 4.9x speedup.
Gradient sync.  
Forward  
Sync after backward  
time  
gpu1  
gpu4  
Overlap sync with backward  
time  
gpu1  
gpu4  
Backward  
Idle  

Figure 3: Illustration of how the backward pass in back-propagation can be overlapped with gradient synchronization to improve training speed.

4.3 Parallel Training

While large batch training improves training time even on a single node, another benefit of training with large batches is that it is easily parallelized across multiple nodes (machines). We run our previous 1-node experiment over 16 nodes of 8 GPUs each (NVIDIA V100), interconnected by Infiniband. Table 1 shows that with a simple, synchronous parallelization strategy over 16 nodes we can further reduce training time from 311 minutes to just 37 minutes (cf. Table 1; 2x lr vs. 16 nodes).

However, the time spent communicating gradients across workers increases dramatically when training with multiple nodes. In particular, our models contain over 200M parameters, therefore multi-node training requires transferring 400MB gradient buffers between machines. Fortunately, the sequential nature of back-propagation allows us to further improve multi-node training performance by beginning this communication in the background, while gradients are still being computed for the mini-batch (see Figure 3). Back-propagation proceeds sequentially from the top of the network down to the inputs. When the gradient computation for a layer finishes, we add the result to a synchronization buffer. As soon as the size of the buffer reaches a predefined threshold we synchronize the buffered gradients in a background thread that runs concurrently with back-propagation down the rest of the network. Table 1 shows that by overlapping gradient communication with computation in the backwards pass, we can further reduce training time by 15%, from 37 minutes to just 32 minutes (cf. Table 1; 16 nodes vs. overlap comm+bwd).

We illustrate the speedup achieved by large batches and parallel training in Figure 4.

4.4 Results with WMT Training Data

We report results on newstest14 for English-to-German (En-De) and English-to-French (En-Fr). For En-De, we train on the filtered version of WMT’16 from Vaswani et al. (2017). For En-Fr, we follow the setup of Gehring et al. (2017). In both cases, we train a “big” transformer on 16 nodes and average model parameters from the last 10 checkpoints (Vaswani et al., 2017). Table 2 reports 29.3 BLEU for En-De in 1h 25min and 43.2 BLEU for En-Fr in 8h 32min. We therefore establish a new state-of-the-art for both datasets, excluding settings with additional training data (Kutylowski, 2018). In contrast to Table 1, Table 2 reports times to convergence, not times to a specific validation likelihood.
4.5 Results with WMT & Paracrawl Training

Fast parallel training lets us additionally explore training over larger datasets. In this section we consider Paracrawl (ParaCrawl, 2018), a recent dataset of more than 4B parallel sentences for each language pair (En-De and En-Fr).

Previous work on Paracrawl considered training only on filtered subsets of less than 30M pairs (Xu and Koehn, 2017). We also filter Paracrawl by removing sentence-pairs with a source/target length ratio exceeding 1.5 and sentences with more than 250 words. We also remove pairs for which the source and target are copies (Ott et al., 2018). On En–De, this brings the set from 4.6B to 700M. We then train a En–De model on a clean dataset (WMT’14 news commentary) to score the remaining 700M sentence pairs, and retain the 140M pairs with best average token log-likelihood. To train an En–Fr model, we filter the data to 129M pairs using the same procedure.

Next, we explored different ways to weight the WMT and Paracrawl data. Figure 5 shows the validation loss for En-De models trained with different sampling ratios of WMT and filtered Paracrawl data during training. The model with 1:1 ratio performs best on the validation set, outperforming the model trained on only WMT data. For En-Fr, we found a sampling ratio of 3:1 (WMT:Paracrawl) performed best.

Test set results are given in Table 3. We find that Paracrawl improves BLEU on En–De to 29.8 but it is not beneficial for En–Fr, achieving just 42.1 vs. 43.2 BLEU for our baseline.

5 Analysis of Stragglers

In a distributed training setup with synchronized SGD, workers may take different amounts of time to compute gradients. Slower workers, or stragglers, cause other workers to wait. There are several reasons for stragglers but here we focus on the different amounts of time it takes to process the data on each GPU.

In particular, each GPU typically processes one sub-batch containing sentences of similar lengths, such that each sub-batch contains at most 3.5k tokens.
speed, Figure 6 shows the amount of time required to process the 44K sub-batches in the En-De training data. There is large variability in the amount of time to process sub-batches with different characteristics: the mean time to process a sub-batch is 0.11 seconds, the slowest sub-batch takes 0.228 seconds and the fastest 0.049 seconds. Notably, there is much less variability if we only consider batches of a similar shape (e.g., batches where $23 \leq \text{src len} \approx \text{tgt len} \leq 27$).

Unsurprisingly, constructing sub-batches based on a maximum token budget as just described exacerbates the impact of stragglers. In Section 4.2 we observed that we could reduce the variance between workers by accumulating the gradients over multiple sub-batches on each worker before updating the weights (see illustration in Figure 2). A more direct, but naïve solution is to assign all workers sub-batches with a similar shape. However, this increases the variance of the gradients across batches and adversely affects the final model. Indeed, when we trained a model in this way, then it failed to converge to the target validation perplexity of 4.32 (cf. Table 1).

As an alternative, we construct sub-batches so that each one takes approximately the same amount of processing time across all workers. We first set a target for the amount of time a sub-batch should take to process (e.g., the 90th percentile in Figure 6) which we keep fixed across training. Next, we build a table to estimate the processing time for a sub-batch based on the number of sentences and maximum source and target sentence lengths. Finally, we construct each worker’s sub-batches by tuning the number of sentences until the estimated processing time reaches our target. This approach improves single-node throughput from 143k tokens-per-second to 150k tokens-per-second, reducing the training time to reach 4.32 perplexity from 495 to 479 minutes (cf. Table 1, 16-bit). Unfortunately, this is less effective than training with large batches, by accumulating gradients from multiple sub-batches on each worker (cf. Table 1, cumul, 447 minutes). Moreover, large batches additionally enable increasing the learning rate, which further improves training time (cf. Table 1, 2x lr, 311 minutes).

6 Conclusions

We explored how to train state-of-the-art NMT models on large scale parallel hardware. We investigated lower precision computation, very large batch sizes (up to 400k tokens), and larger learning rates. Our careful implementation speeds up the training of a big transformer model (Vaswani et al., 2017) by nearly 5x on one machine with 8 GPUs.

We improve the state-of-the-art for WMT’14 En-Fr to 43.2 vs. 41.5 for Shaw et al. (2018), training in less than 9 hours on 128 GPUs. On WMT’14 En-De test set, we report 29.3 BLEU vs. 29.2 for Shaw et al. (2018) on the same setup, training our model in 85 minutes on 128 GPUs. BLEU is further improved to 29.8 by scaling the training set with Paracrawl data.

Overall, our work shows that future hardware will enable training times for large NMT systems that are comparable to phrase-based systems (Koehn et al., 2007). We note that multi-node parallelization still incurs a significant overhead: 16-node training is only ~10x faster than 1-node training. Future work may consider better batching and communication strategies.

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