Scaling to Multiple Graphics Processing Units (GPUs) in TensorFlow

by Song J Park

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Although accuracies of neural networks are surpassing human performance, training a deep neural network is a time-consuming task due to its increasing high-dimensional parameters. It is not uncommon for the training of deep neural networks to run for a week. Accordingly, the size of neural networks has doubled every 2.4 years, exhibiting an exponential growth from 1958 to 2014. The increasing size of neural network architectures will likely lead to higher computational complexity that will need scalable solutions. To mitigate the computational requirement and maximize throughput, this work focuses on multi-graphics-processing-unit scalability.

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1. Introduction

The interest in artificial neural networks has peaked again in recent years. The popularity of neural networks has fluctuated since the introduction of perceptron by Rosenblatt. Neural networks have undergone a transformation from the early days of perceptron to modern deep neural networks (DNNs). Success, however, was not achievable until the rise of big data and access to highly parallel computing power. In terms of input data, it is recommended that the size of training examples be at least 10 million for DNNs. Hence, availability of labeled data can be a challenge. As for the neural network model complexity, the trend shows that starting from perceptron to GoogLeNet, the size of neural networks has doubled every 2.4 years, exhibiting an exponential growth from 1958 to 2014. The increasing size of neural network architectures will likely lead to higher computational complexity that will need scalable solutions.

DNNs have demonstrated remarkable advances in the following fields: image recognition, natural language processing, object detection, and reinforcement learning. For instance, DNNs now outperform humans in the object recognition ImageNet competition, as human experts mislabel objects due to multiple objects in an image or difficulty in fine-grained species recognition. Although accuracies of neural networks are surpassing human performance, training a DNN is a time-consuming task due to its increasing high-dimensional parameters. It is not uncommon for the training of DNNs to run for a week. Exploring deeper neural networks with higher complexity will require parallel execution and distributed scaling of DNNs’ training algorithms.

Given the compute-intensive nature of DNNs, a powerful computing platform is ideal to mitigate the training time. The weighted sum of a neural network can be concisely represented as a matrix multiplication operation, which means that DNNs’ core computations comprise multiply and add operations. Due to design choices behind graphics processing unit (GPU) architectures, a GPU processor will have a high number of raw processing units for executing multiply and add operations. Therefore, to maximize throughput, this work focuses on multi-GPU scalability.

TensorFlow provided the software framework to perform a scalability analysis of neural networks that can target a multi-GPU system. The objective of this study was to identify the tradeoffs between portability and programmability in TensorFlow. The aim was to study the impact of the flexibility and portability of TensorFlow on programming complexity for multi-GPU implementations.
2. Software Programming Framework

TensorFlow is an open-source library for numerical computations using dataflow graphs originally developed by Google Brain. The TensorFlow tool is widely used for machine learning and deep learning research. TensorFlow features include front-end Python application programming interface (API), auto-differentiation, and visualization. It is highly portable, supporting central processing units (CPUs), GPUs, tensor processing units (TPUs), and ARM architectures running Linux, Windows, iOS, and Android. A notable difference with TensorFlow is its declarative style of programming via dataflow graphs. In essence, TensorFlow separates the definition of computations from actual execution. As such, a source code will be composed of a description portion for a computational graph and a session portion for the execution of operations.

Parallel strategies for training a DNN include model parallelism and data parallelism. In model parallelism, a DNN model is divided and assigned to computational resources. This can free up memory requirements on each computing resource since only a fraction of a DNN model is assigned for each computing device. On the other hand, data parallelism replicates the whole DNN model on all available computing resources, and different batches of training data are assigned to each computing device, referred to as a “tower”. In data parallelism, DNN weights need to be shared across devices, requiring a synchronization protocol. This report focuses on the data parallelism approach on multiple GPUs.

3. Hardware Computing Platforms

DNNs can be targeted for CPUs, GPUs, field-programmable gate arrays (FPGAs), and application-specific integrated circuits (ASICs). Commercial examples of the use cases of GPUs, FPGAs, and ASICs can be found at Amazon, Microsoft, and Google. Multiple tradeoffs exist in this spectrum of silicon hardware alternatives. For a comparison and tradeoff, attributes of interest are limited to throughput, power, flexibility, and development time. Table 1 illustrates the attribute values for a TPU, FPGA, and GPU.

<table>
<thead>
<tr>
<th>Processor</th>
<th>Throughput</th>
<th>Power (W)</th>
<th>Flexibility</th>
<th>Development time</th>
</tr>
</thead>
<tbody>
<tr>
<td>TPU</td>
<td>92 TOPs (INT8) [19]</td>
<td>75</td>
<td>Low</td>
<td>15 months</td>
</tr>
<tr>
<td>FPGA (VU9P)</td>
<td>21 TOPs (INT8) [20]</td>
<td>225</td>
<td>Medium</td>
<td>Several months</td>
</tr>
<tr>
<td>GPU (P100)</td>
<td>18.7 TFLOPs (FP16) [21]</td>
<td>250</td>
<td>High</td>
<td>&lt;1 month</td>
</tr>
</tbody>
</table>

Note: TOP = tera operations per second; TFLOP = tera floating-point operations per second.
For throughput, TPU and FPGA values reflect the precision of 8-bit integer operations, whereas the GPU throughput conveys 16-bit floating-point operations. As expected, the ASIC solution is the winner for throughput performance and low power. However, AS ICs require longer development time and, once designed, the circuitry is fixed for a specific purpose. FPGAs and GPUs have similar characteristics for throughput and power, but GPUs are simpler and faster to program, develop, and modify. To be adaptable as the landscape of deep learning evolves, the flexible option of GPUs was selected for a scalability study.

The latest development in the shift of Nvidia’s compute element architecture is described here for background. Architectural changes to Nvidia’s GPUs in the current product lineup are specifically designed to target mixed precision matrix multiplication, which is at the core of neural networks. Introduced in Volta architecture, Tensor Cores are added to Nvidia’s multiprocessor to enhance artificial intelligence performance. With Tensor Cores, the theoretical throughput of mixed precision (FP16 and FP32) operations are boosted to 125 TFLOPs. In terms of performance, programmability, and flexibility, GPU’s position on this tradeoff space seems balanced and competitive.

4. Experimental Evaluation

Matrix multiplication is at the core of DNN’s training computations. Therefore, matrix multiplications can be a simplified representation of a DNN’s training operations to test multi-GPU scalability. TensorFlow’s multi-GPU scalability was tested on the DGX-1 system consisting of eight Tesla P100 GPUs. TensorFlow will automatically attempt to execute on a GPU if one is available. Moreover, TensorFlow reserves all visible GPU memories at start even though only one GPU is used by default. This behavior was observed for a simple single GPU matrix multiply code. To be explicit in device allocation, TensorFlow allows specification of device with the “tf.device” function. This function illustrates the flexibility of targeting different processing architectures as well as low-level programming in TensorFlow as evidenced by verbose source code for the multi-GPU execution. In the multi-GPU implementation, matrix multiplication was assigned to each GPU then summed using a CPU. Matrix dimensions were on the order of 10,000 by 10,000. Execution times for the various number of GPUs are provided in Table 2, where batch sizes represent the workload assigned to each GPU.
Table 2  Execution time for matrix multiplication on multiple GPUs

<table>
<thead>
<tr>
<th>Batch size</th>
<th>2</th>
<th>8</th>
<th>32</th>
<th>128</th>
<th>512</th>
</tr>
</thead>
<tbody>
<tr>
<td>GPUs</td>
<td>(s)</td>
<td>(s)</td>
<td>(s)</td>
<td>(s)</td>
<td>(s)</td>
</tr>
<tr>
<td>1</td>
<td>4.2</td>
<td>14.1</td>
<td>53.9</td>
<td>213.7</td>
<td>851.8</td>
</tr>
<tr>
<td>2</td>
<td>2.8</td>
<td>7.9</td>
<td>28.0</td>
<td>107.9</td>
<td>427.4</td>
</tr>
<tr>
<td>4</td>
<td>2.6</td>
<td>5.2</td>
<td>15.1</td>
<td>55.2</td>
<td>215.8</td>
</tr>
<tr>
<td>8</td>
<td>2.5</td>
<td>3.0</td>
<td>8.8</td>
<td>30.3</td>
<td>112.1</td>
</tr>
</tbody>
</table>

As depicted in Fig. 1, larger batch sizes led to improved performance toward the linear speed-up. In reference to the factors against parallelism, the major contributing factor seemed to be the startup costs of transferring data to a GPU device, whereas interference and skew were minimal. At the batch size of 2, the low-compute intensity does not warrant the data transfer communication costs to the GPUs.

Fig. 1  Speed-up plot for matrix multiplication targeting multiple GPUs at different batch sizes

CIFAR10 source code was provided by the TensorFlow tutorial package. The number of GPUs can be provided as an argument for running the CIFAR10 code. Execution time was measured to be 888, 561, 533, and 523 s for one, two, four, and eight GPUs, respectively. As indicated in Fig. 2, the speed-up curve flattens after two GPUs. Monitoring the Tesla GPUs via Nvidia tool revealed that the use of the GPU processors was approximately 80%, 65%, 33%, and 17% for one, two, four, and eight GPUs, respectively. These observations indicate that as the number of
GPUs increased, each GPU’s utilization decreased, which resulted in a suboptimal speed-up. Similar suboptimal multi-GPU results for DNNs are reported in Gawande et al.\textsuperscript{24} and Shi et al.\textsuperscript{25} Underutilization of GPU cores can be the result of an insufficient input data pipeline. Future work is to explore efficient input pipeline support within TensorFlow.

![CIFAR10 speed-up plot](image)

**Fig. 2** Speed-up plot for CIFAR10 using multiple GPUs

## 5. Conclusion

The multi-GPU matrix multiplication exhibited almost linear speed-up, whereas the multi-GPU CIFAR10 showed barely any performance improvement after two GPUs. TensorFlow provides flexible, verbose, and low-level programming interface for scaling to multiple GPUs. However, multi-GPU programming in TensorFlow is an involved process, which can be abstracted by a high-level programming language. For instance, Keras is a high-level API that runs on top of TensorFlow and can implement a multi-GPU version with minimal code modification. The tradeoff between hardware portability and verbose programming model was experienced in TensorFlow. To target the broadest audience and applications, TensorFlow seems to sacrifice high-level programming approach for portability.
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<thead>
<tr>
<th>Symbol</th>
<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>API</td>
<td>application programming interface</td>
<td></td>
</tr>
<tr>
<td>ASIC</td>
<td>application-specific integrated circuit</td>
<td></td>
</tr>
<tr>
<td>CPU</td>
<td>central processing unit</td>
<td></td>
</tr>
<tr>
<td>DNN</td>
<td>deep neural network</td>
<td></td>
</tr>
<tr>
<td>FLOP</td>
<td>floating-point operations per second</td>
<td></td>
</tr>
<tr>
<td>FPGA</td>
<td>field-programmable gate array</td>
<td></td>
</tr>
<tr>
<td>GPU</td>
<td>graphics processing unit</td>
<td></td>
</tr>
<tr>
<td>iOS</td>
<td>Apple mobile operating system</td>
<td></td>
</tr>
<tr>
<td>OPS</td>
<td>operations per second</td>
<td></td>
</tr>
<tr>
<td>TFLOP</td>
<td>teraflop</td>
<td></td>
</tr>
<tr>
<td>TOP</td>
<td>tera operations per second</td>
<td></td>
</tr>
<tr>
<td>TPU</td>
<td>tensor processing unit</td>
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