Scavenger: A Cloud Service For Optimizing Cost and Performance of ML Training

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Abstract—Cloud computing platforms can provide the computational resources required for training large machine learning models such as deep neural networks. While the pay-as-you-go nature of cloud virtual machines (VMs) makes it easy to spin-up large clusters for training models, it can also lead to ballooning costs. The 100s of virtual machine sizes provided by cloud platforms also makes it extremely challenging to select the “right” cloud cluster configuration for training. Furthermore, the training time and cost of distributed model training is highly sensitive to the cluster configurations, and presents a large and complex tradeoff-space.

In this paper, we develop principled and practical techniques for optimizing the training time and cost of distributed ML model training on the cloud. Our key insight is that both the parallel and statistical efficiency must be considered when selecting the optimum job configuration parameters such as the number of workers and the batch size. By combining conventional parallel scaling concepts and new insights into SGD noise, we develop models for estimating the time and cost on different cluster configurations. Using the repetitive nature of training and our performance models, our Scavenger cloud service can search for optimum cloud configurations in a black-box, online manner. Our approach reduces training times by $2\times$ and costs by more than 50%. Our performance models are accurate to within 2%, and our search imposes only a 10% overhead compared to an ideal oracle-based approach.

I. INTRODUCTION

The discovery of improved machine learning (ML) models has resulted in great advances in computer vision, language and speech processing, scientific computing, and many other areas. These advances are primarily driven by increasingly computationally intensive models, such as deep neural networks (DNNs), being “trained” on large data sets. The ready availability of computing resources is a key enabler of machine learning, and cloud platforms can easily provide these resources. However, current ML techniques and systems are ill-suited for making effective and efficient use of cloud resources, i.e., are not cloud-native.

ML models are trained on large clusters of cloud virtual machines, leading to prohibitive costs, because ML training techniques and frameworks like TensorFlow and PyTorch are oblivious to cost. Moreover, cloud platforms offer 100s of different virtual machine sizes and configurations with different cost/performance tradeoffs, making it extremely challenging to select the “right” type and quantity of cloud resources. Training large ML models on the cloud is thus often performed on sub-optimally configured cloud resources, leading to cost overruns, slow performance, and underutilized resources.

These challenges also exist when optimizing the resource allocation for conventional distributed applications (such as map-reduce data processing) on the cloud [1]. However, model training also has other unique execution and synchronization characteristics and a large array of configuration knobs (such as number of workers and the batch size) which have significant impact on performance and resource efficiency.

In this paper, we present Scavenger, a service for optimizing the cloud training cost and time for ML models. Scavenger is a model-agnostic, black-box, fully online service built using TensorFlow, and searches for good configurations for distributed model training jobs. We use a performance-model guided search across a multi-dimensional configuration space to find the pareto-optimal configurations based on user preferences and constraints. In its search for the best configuration, Scavenger horizontally scales a training job by adding/removing workers, and vertically scales it by changing the batch size.

As a key first step towards understanding and optimizing training time and costs, we develop a new phenomenological performance model for data-parallel distributed model training. Our model uses both conventional parallel scaling concepts such as synchronization overheads, as well as fundamental performance artifacts of Stochastic Gradient Descent (SGD) based optimization. Unlike in classical parallel applications, we find that computation performed by parallel workers doesn’t always compose because of the stochastic nature of the gradient computation. This statistical inefficiency reduces the rate of the job’s forward-progress, and imposes its own tradeoff on time and cost which also depends on the cluster configuration. We measure and consider this statistical inefficiency by using SGD noise (variance of gradients), and show how it can be used as a general scaling indicator.

Scavenger is a fully managed model training service requiring minimal user intervention, prior knowledge, or offline profiling. We use the repetitive and iterative nature of model training to briefly profile the job on different configurations and learn its performance profile by using the scaling indicators. We minimize the overhead of this exploration and search phase by using lightweight model checkpointing, and obtain the cost and time tradeoff curves for different combinations of workers and batch sizes. The performance model is then used to run the reminder of the job on the “best” configuration based on user preferences and constraints.

Our profiling-based strategy of building the performance
model is optimized to reduce the search cost. We obtain a performance profile of an ML model by profiling on only a small subset of configurations. We develop phenomenological first-principles performance models that can be interpolated using linear regression—thus requiring only a partial search. Since Scavenger is a cloud service, it also leverages repeated training of similar models (e.g., part of hyperparameter or neural architecture search), and reuses its learned performance model, to completely eliminate the exploration phase and search costs. Surprisingly, we find that the SGD noise can serve as model-agnostic scaling indicator, and even a “universal” average model can estimate performance with reasonable accuracy without any exploration or pilot jobs.

To the best of our knowledge, Scavenger is the first work which can optimize both cost and time in a fully online manner. We build on recent work for SGD noise based scaling such as [2]–[5], and use it for simple intuitive phenomenological models. By considering both the parallel and statistical efficiency, we are able to accurately predict the training time of a wide range of DNN models with minimal search overhead.

Scavenger is an open-source library built on top of TensorFlow, and provides a practical, online, black-box, model-agnostic service for addressing the crucial problem of cost and performance optimization of distributed machine learning in the cloud. In addition to the practical significance, we make the following research contributions:

1) Our empirical investigation of the cost and time tradeoffs in distributed ML model training reveals the impact of parallel and statistical efficiency.

2) We show how the variance in gradients results in SGD noise, and how it can serve as a reliable scaling indicator for elastic horizontal and vertical scaling.

3) We develop new models for predicting the performance for deep neural networks, which consider both parallel and statistical efficiency, and the aforementioned SGD noise. Our models predict training time and cost for different job configurations (number of workers and batch size), and construct full tradeoff curves and pareto frontiers, with very high accuracy of more than 98%.

4) Our models enable us to search for the optimum job and cluster configuration in a model-agnostic and online manner, and minimize various combinations of cost and time. Our techniques can find the “right” cloud configuration and reduce training time by more than 2× compared to naive configurations.

II. BACKGROUND AND CHALLENGES

In this section, we describe the performance tradeoffs faced by distributed ML training. These observations and insights guide our performance model presented in the next section.

A. Distributed ML Training

Distributed training entails learning the model parameters (or weights) of a model over an input training dataset. A model trains in an iterative-convergent process to minimize a loss function over the dataset by using optimization techniques such as Stochastic Gradient Descent (SGD) [6] and Mini-Batch Gradient Descent [7] or Full Gradient Descent. Since ML training is highly compute intensive, parallelizing it using computational accelerators such as GPUs and TPUs, and through distributed training, is vital [8], [9]. A common parallelization approach is data-parallelism, where training is launched on multiple workers, and each worker learns and updates the model parameters by processing a small batch of the training data [10] at each iteration.

After each iteration, the gradient updates from all workers are aggregated via all-reduce operations to compute the averaged gradients, update model parameters and synchronize the new parameters among the workers [11]. A popular and widely successful data-parallel training approach is the parameter server strategy, where the workers compute the gradients, and parameter servers aggregate and average the gradients from all workers after every iteration and update the model weights. Training a popular image recognition model like ResNet [12], [13] or an attention-based language model like Transformer [14] typically require thousands of iterations until the model’s error converges to the desired low training loss.

Concretely, the training process iteratively computes the model parameters over $K$ workers, each processing a mini-batch of $b$ at iteration $t$ and computing the gradient $\nabla f(x_{k,t})$. The update rule for the model parameters $x$ is given by:

$$x_{t+1} = x_t - \eta \frac{1}{K} \frac{1}{b} \sum_{k=1}^{K} \nabla f(x_{k,t}),$$

where $\eta$ is the learning rate, one of the hyperparameters of the model that is found through empirical search techniques. The global batch size is $B = bK$, and is a crucial job parameter.

Elasticity. Distributed training is resource elastic, which means that the models can be trained on different cluster sizes and configurations, which can also be changed during runtime (i.e., during model training). Training can be horizontally scaled by adjusting the number of workers (i.e., changing $K$ in Equation 1), and vertically scaled by increasing the mini-batch size $b$ on each worker. ML frameworks such as TensorFlow and PyTorch also support model checkpointing, and thus we can adjust the horizontal and vertical scaling dynamically by checkpointing the model state and resuming the training on a different cluster configuration.

![Fig. 1: Running cost and time for different batch sizes and workers for ResNet18 training. Each point along a tradeoff-curve represents 20, 16, 12, 8 workers respectively. Dashed line shows our model prediction.](image_url)
This elasticity makes distributed training a good fit for clouds, since we can easily scale the cluster by adding/removing VMs, and changing the underlying VM size to increase the batch size and intra-worker parallelism. Scavenger makes use of this elasticity in its search for the ideal cloud configuration. However, distributed training has complex and incompletely-understood performance tradeoffs [15] that are affected by the various SGD parameters (such as $K, b$). Simply running more workers and increasing batch size has diminishing returns, as we can see from Figure 1, which shows the running time and cost for training the ResNet18 model. Each point corresponds to a different number of workers for each batch size. We can see that there are diminishing returns, and thus it is not obvious which cluster configuration is the “best”.

B. Horizontal Scaling: Adding Workers

The simplest way of scaling a parallel training job is to add more workers ($K$). Figure 2a shows the decrease in the total training time to reach a fixed accuracy level for three ML models. As the number of workers increases, the training time reduces, but there are diminishing returns. Increasing the workers from 4 to 16 (4×) only reduces the training time from 15 to 5 hours (3×).

Thus, ML training shows parallel inefficiency due to the communication and synchronization overheads. A single model-training iteration consists of a local gradient computation step, and a synchronization step where the gradients are aggregated/averaged. Figure 3 shows the breakdown of this computation and synchronization. Here, the cumulative cluster capacity is same across various $K$, i.e., total CPU cores and memory allocated over all the workers in a cluster is held constant. We can see that increasing the number of workers increases the synchronization time. With parameter servers, more workers means more stragglers, and because bulk synchronous processing is used, this increases the communication costs for everyone. The larger number of workers also increases the work for the parameter servers, which increases the synchronization time further. Figure 3 also shows the breakdown for different batch sizes. We can see that the gradient computation time also increases with increasing batch sizes.

Memory. The VM’s memory size is also an important resource for model training. Insufficient memory forces smaller batch sizes, which reduce the training accuracy and require more iterations and synchronization during model training. Figure 2b shows the final model accuracy reached when training the Transformer model under a strict time deadline. The smaller VMs provide insufficient accuracy, and below 4GB memory, the system (TensorFlow) crashes and makes no forward progress.

C. Vertical Scaling: Increasing Batch Size

One way to reduce the synchronization overheads is to increase the batch size, which reduces the total number of iterations required and increases the parallel efficiency. This is illustrated in Figure 2c, which shows the training-loss for different batch sizes for the ResNet18 model. Larger batch sizes (1024) achieve lower (i.e., better) loss compared to smaller ones. This is also seen for other models in Figure 4, which shows the training time to desired accuracy for $K = 16$.

The gains of compute efficiency with larger $B$ is evident from Fig. 5 where the throughput (i.e., the number of samples processed per second in the training phase) increases as the global batch size increases, then saturates after a knee-point/inflection point. The throughput plateaus after a certain batch size since the CPU utilization of the workers maxes out after the inflection point. In the results shown, the workers used in the cluster are GCP E2-standard VMs with 4 vCPUs and 16 GB memory.

D. Statistical inefficiency

In both horizontal and vertical scaling, parallel training does not scale linearly. The fundamental reason for this non-linear scaling is that not all computing work is effective because of stochastic gradient descent. In conventional parallel applications, all work performed by all workers is equally useful. However, with stochastic gradient descent, the work done by the workers (i.e., the gradients computed) does not fully compose. That is, the total forward progress made (i.e., the decrease in training loss) is not equal to the sum of progress made by the individual workers. We call this the statistical inefficiency of parallel model training, and it reflects how “aligned” the computed gradients of the different workers are. This statistical inefficiency is a fundamental attribute of
SGD (and all optimization algorithms in the SGD family like Adam [16] etc). The statistical inefficiency can be captured by computing the SGD noise, which is the variance in the gradients among the workers [2], [3]. We use a similar variance formulation in our model described in the next section.

Thus adding more workers (increasing K) can increase the divergence between gradients and require more training iterations and increase the overall training time. Similarly, a small batch size means that the gradients are computed on a small subset of data, and are more likely to differ from each other. Thus, larger batch sizes are preferable from a statistical efficiency perspective, but have other tradeoffs: they impose additional memory requirements and communication overheads. Furthermore, increasing batch sizes may have diminishing returns (Figure 5). *Because of gradient noise and statistical inefficiency, throughput (number of training data samples processed per second) is not sufficient to capture the performance, and we need to consider the wall-clock time to reach the desired accuracy level.*

Our performance model is able to capture both the statistical and parallel efficiency associated with different horizontal and vertical scaling configurations, and provide accurate estimates of training time for different configurations which can be used to select the “best” configuration.

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**III. DESIGN**

At a high level, our goal is to find the “best” cloud cluster configuration for a model training job, with minimal information about the ML model, and in an online manner with minimal a priori profiling. We want to minimize the time and cost of training a model to a specified accuracy level.

For optimizing the job configuration for a given ML training job, we first develop an analytical performance model for estimating the total training time (and cost). This performance model is used to compute the time vs. cost tradeoff curves for a job, which can be used to select the “right” cloud cluster based on user preferences and constraints. Our focus is on building simple, practical, and generalizable performance models that do not require offline training, and which can be refined and used with online profiling. Predicting the total training time of ML training is especially challenging due to the statistical inefficiency of distributed SGD. To address this challenge, we use general *SGD noise* indicators, that serve as a proxy for statistical inefficiency (Section III-A). Using these scaling indicators, we develop an analytical *statistical* performance model, which we combine with a more conventional parallel performance model. Finally in Section III-E, we describe how the combined parallel and statistical performance model can be obtained and used in practice.

**A. SGD Noise as a Scaling Indicator**

We have seen that simply adding more resources to a distributed training job doesn’t decrease the training time uniformly. This inefficiency is crucial in cloud environments, since it increases costs without proportional decrease in training time. We seek a general “scaling indicator” which serves as a proxy for the overall parallel efficiency. For example, such a scaling indicator should indicate the scenarios in which adding more resources would not decrease training time, and we should stop scaling. Because we want online cluster optimization, this scaling indicator should also be easily computable at run-time, and be independent of the ML model and cluster size.

For classic parallel applications, the communication and synchronization overheads typically serve as scaling indicators. For example, we can compute the scaling efficiency as the fraction of time spent in communication, and stop scaling if this fraction crosses a threshold. Amdahl’s law and other parallel scaling laws can then use these communication overheads and
Fig. 6: The SGD noise requires some iterations to stabilize, after which it is dominated by the number of workers. Inform us about the performance and scaling properties of the application. Communication and synchronization overheads are also applicable for ML training and can be used to model their parallel efficiency. However, they are not sufficient, because of the statistical inefficiency of parallel ML training.

Just as communication overheads can indicate parallel scaling in conventional parallel applications, are there similar scaling indicators for statistical inefficiency? We seek a general indicator for statistical efficiency that is independent of the model and the execution environment (number and type of workers, etc.). For example, such a scaling signal could indicate the batch size threshold for a given cluster size, beyond which scaling does not significantly reduce the training time.

Fundamentally, the statistical inefficiency arises because of the noise in the gradients computed by the workers. Our main observation is that the SGD noise can be captured by the variance in the gradients computed by the workers, and this serves as a useful general statistical inefficiency indicator. This variance/noise can be computed by:

$$\gamma(t) = \frac{1}{K} \sum_{k=1}^{K} \frac{|g_{t}^{(k)}|^2}{E[|\hat{g}_t|^2]}, \quad \text{(2)}$$

where $g_t^{(k)}$ is gradient computed on worker $k$ at iteration $t$ and $\hat{g}_t$ is the aggregated gradient norm obtained by reducing gradients on the parameter servers in a cluster with $K$ workers. This SGD gradient variance has been investigated previously [2], [3] to understand either batch size scaling or worker scaling, and we generalize it to both types of scaling.

The noise, which is essentially the deviation in the calculated gradients from the “true” gradient, is also a practical scaling indicator. It can be easily computed in the data-parallel parameter server strategy during the model training, i.e., in an online manner. The per-worker and aggregated gradients are collected from all workers and parameter servers respectively. Thus, from equation 2, we can compute the gradient noise by computing the ratio of the mean of the workers’ local gradient norms and the aggregated gradient norm.

In the early training stages, the variance in the gradients is on the same scale as the gradients itself and thus the initial noise is low (Figure 6). As the ML model converges, the gradients approach towards the true gradient, increasing the noise before finally saturating to the number of workers $K$. Since we want to compare the noise for different $K$, we normalize it by $K$, so that it is a true statistical efficiency indicator.

We have observed that the noise is not constant over the course of training, even with a static job configuration. Instead, the noise increases and then stabilizes, as we can see from Figure 6. This is a fundamental artifact of SGD-based optimization, and applicable for all models and configurations. The noise is also affected by the SGD learning rate, and we need to account for the learning-rate schedule. For our cluster optimization, we want to search and select for the right cluster configuration as quickly as possible after the training commences. However since the noise from early training epochs is unreliable, we let the noise stabilize before using it as a scaling indicator. When a job starts, we run it on the starting configuration until the noise stabilizes, and then begin the exploration/search process. This increases the overall profiling and search time, since the early iterations are the “cold start”, but provides reliable noise estimates.

How effective is SGD noise in predicting performance? Figure 7 shows the total training time to the desired accuracy vs. the SGD noise for different global batch sizes $B$. For all the three ML models, the increase in noise leads to an increase in training time. We also observe that smaller batches have higher noise. Thus, the SGD noise can serve as a good indicator of the training time and efficiency. We investigate a deeper relation of noise with statistical efficiency in the our performance model developed in the rest of this section.

**B. Performance and Cost Model**

Our analytical model for the total training time and cost of distributed ML training creates the tradeoff curves (like in Figure 1), and guides the cloud resource allocation policies. We use statistical and parallel scaling indicators which can be obtained by profiling in an online manner during job execution, and do not need a-priori offline profiling. The job’s performance depends on its configuration, which consists of the number of parallel workers, $K$, and the total batch size $B$, and our model predicts the performance for each combination of these configuration parameters. ML training is an iterative process, and the total training time, $T$:

$$T = n_i \tau, \quad \text{(3)}$$

where $n_i$ is the number of iterations to reach the specified model accuracy, and $\tau$ is the per-iteration time. The number of iterations depends on the total number of training epochs $e$:

$$n_i = \frac{eD}{B}, \quad \text{(4)}$$

where $D$ is the fixed dataset size, and $B$ is the global batch size, an important job configuration parameter. The number of epochs to reach the desired model accuracy $e$, is the key
unknown, and depends on many factors such as the model size, 
complexity, and desired accuracy, and the statistical inefficiency.

The other key parameter in Equation 3 is \( \tau \), which is the 
per-iteration time. For a given job configuration, i.e., fixed \((K, B)\), 
the time to process a mini-batch is roughly constant over the 
course of training, because the same gradient computation 
and communication steps are being performed on the same 
mini-batch of identically distributed data.

Finally, the total cost is simply the product of training time, 
the number of workers \( K \), and the per-VM price \( p \):

\[
C = TKp
\]  

We estimate the number of epochs, \( e \) using our statistical 
performance model described in the next subsection. The time 
per iteration \( \tau \), will be estimated using our parallel performance 
model in Section III-D.

**Online Profiling and Searching.** Using the model, we first 
obtain the tradeoff curves in our search or exploration phase. In 
the search phase, we briefly run the job on some configuration, 
observe its parallel and statistical scaling indicators, and 
estimate the time (and thus cost) on that configuration. Only 
a small number of iterations (around 20) are usually required 
for estimating the performance of a given configuration, after 
which we checkpoint the model, and run the job on a different 
configuration. This exploration of the various configurations 
allows us to obtain the full time and cost curves.

With checkpointing, there is no lost work. The search cost 
is running the job briefly on suboptimal configurations, and 
the small overhead of restoring the model from checkpointed 
weights. Selecting the next configuration in the exploration 
phase is done using grid search guided by the optimization 
criteria and constraints on K and B. We refer to this as a full or 
offline search, since we first explore the configuration space, 
and then run the remainder of the job on the best configuration.

To reduce the search cost, our phenomenological statistical 
and parallel performance model also allows us to estimate 
the running time on configurations without even profiling 
on them. That is, we can obtain estimates of \( T \) by profiling 
on only a small sample of K, B configurations, and use our 
phenomenological models to build the rest of the tradeoff curve 
by fitting the learned performance models. This partial or 
online search reduces the search cost significantly. However, 
the drawback is that the estimates of running time due to 
the interpolation/regression can be error-prone, and thus the 
tradeoff curves obtained using the online search can differ 
slightly from the offline search.

Finally, we observe that many jobs train nearly identical 
models as part of hyperparameter tuning, neural architecture 
search, etc. For example, the hyperparameter tuning may involve 
dozens of jobs that train the same model, but with different 
activation functions, weight decay, regularization, etc. In such 
cases, the parallel and statistical efficiency of the job doesn’t 
significantly change. Thus, once a job’s performance model is 
learnt, it can be stored and reused when the same or similar 
model is trained in the future. We can thus avoid the exploratory 
search phase entirely, and this is the no-search scenario. We 
develop full, partial, and no-search techniques for both the 
statistical and parallel performance models.

**C. Statistical Performance Model**

The SGD noise scaling indicator allows us to model the 
statistical performance and the number of epochs required for 
achieving the desired accuracy level. The SGD noise increases 
the total amount of work required, and hence the number of 
iterations and epochs. For a given job configuration \((K, B)\), 
the number of epochs \( e \) is proportional to the SGD noise \( \gamma : 

\[
e \propto \gamma,
\]  

Empirical support for this can be seen in Figure 8, which shows 
the normalized noise plotted against epochs taken to reach a 
specific performance target across various \( K \) and for different 
target accuracy levels.

This linear model can be understood in relation to full 
gradient descent, which has no noise, and the minimal number 
of epochs \( e^* \). Thus for SGD, for a given \( B \), we have \( e = e^* + \theta \gamma \), 
where \( \theta \) is the unknown linear-model parameter which relates 
the noise to the statistical efficiency.

**Full offline search.** We profile the model on each \((K, B)\) 
configuration, and measure the noise \( \gamma_{K, B} \). Note that we are 
primarily interested in the relative performance of various 
configurations. Both \( e^*, \theta \) are properties of the model and not 
fected by \( K, B \). We can obtain them from prior profiling 
rules like those shown in Figure 8. From the figure, we can see 
that the epochs required to reach different accuracy levels is 
not sensitive to the number of workers. In fact, this is a static 
property of the ML model itself, and not influenced by any job 
configuration parameter. Thus, if we have access to any single 
prior execution of the model (under any configuration, and even 
without profiling), then we can estimate the epochs required 
to reach any desired accuracy level. In many cases however, 
we only need to compare the relative performance between 
different configurations, for which we do not need any prior 
execution log, and can compare the epochs of configurations 
based on their observed noise.

The above full search technique already provides significant 
new capabilities for statistical efficiency modeling. We refine 
it with two more powerful insights that reduce the search cost 
associated with the statistical efficiency model further.

**Partial Search.** The linear relation between noise and epochs 
is extremely powerful, and we can enhance it even further 
to model the statistical efficiency using partial search without 
exploring the entire configuration space. We use a finer-grained 
noise model, which allows us to relate it to the batch size.

\[
\gamma_{K, B} \propto \frac{1}{\sqrt{B}}
\]  

This allows us to estimate the noise on different batch sizes 
without even requiring profiling, we can profile and find the 
noise on a small number of different \( B \) values in the search 
phase, and build a linear model of \( \gamma, B \), and use it to interpolate 
the noise for the rest of the unseen values of \( B \). This relation 
between \( \gamma, \sqrt{B} \) is derived from the theoretical properties of 
SGD described in [3]. We empirically show it in Figure 9, which
shows the linear relation between noise and $\frac{1}{\sqrt{B}}$ for different ML models. Our partial search is performed by running the job briefly on the extreme points of $B$, i.e., on the smallest and largest batch size provided by the user, and then fitting a model to Equation 7.

For enhanced accuracy, we can repeat the process for different values of $K$. We have found that it is possible to avoid this per-$K$ profiling and instead use a general/average model for noise and $B$. Surprisingly, the relation between noise and batch size is not very sensitive to the number of workers $K$. Thus, we can simplify statistical efficiency model even further, by using an average model for noise vs. $B$. This average model is also shown in Figure 9 (by the solid line). For a given $B$, we average the estimated noise for various $K$ values. With this averaging, using only the batch size, we can predict the noise, and thus the number of epochs.

**No-search.** If the same or similar ML model is being trained repeatedly, then we can use its noise vs. $B$ relation, and do not additional profiling for modeling its statistical performance.

**D. Parallel Performance Model**

The above statistical performance model provides us the estimate of the number of epochs/iterations required. We now tackle the relatively simpler task of modeling the per-iteration time, using more conventional parallel performance techniques. Our key insight is that ML training is highly repetitive and the performance characteristics of each iteration within a job are nearly identical. This allows us to continue using the profiling based search strategy. Thus the full-search for the parallel performance model simply runs the job for a small number of iterations on all the job configurations of interest.

**Partial-search.** For the partial search, we again use a phenomenological model for iteration time and use an interpolation approach. Each iteration entails computing the gradients, collecting and averaging them, and then synchronizing them between workers via the parameter server. Both these major components can be modeled as follows:

$$\tau = compute\_time + sync\_time.$$ (8)

The gradient computation time on a worker depends on the mini-batch size, $b$:

$$compute\_time \propto b.$$ (9)

The synchronization time is influenced by number of workers:

$$sync\_time \propto K.$$ (10)

Using these relations, we can build a model for the per-iteration time $\tau$ by profiling on the extreme points in the $(K, B)$ configuration space, and then fitting linear models for the computation and synchronization.

**E. Resource allocation policies**

We combine the statistical and parallel performance model for our job configuration and cloud resource allocation policies. We first build the time/cost tradeoff curves using the profiling and modeling. Depending on the prior information available, the search strategy and costs may differ. We have built our system as a service, so future jobs training similar models can be significantly sped-up using their stored performance models and using the partial or no-search policies.

The job configuration search is ultimately determined by the user’s objective and constraints. We support optimizing for time, cost, and also a knee-point based optimization that selects the knee-point of the cost/time curves. We determine the knee of the curve using the kneedle [17] algorithm.

Constraints on the maximum cost and time are provided by the user. This bounds the search space and is also practical. These constraints thus also impose a constraint on the number of worker VMs ($K$), and yield $K_{min}$ and $K_{max}$. The bounds on the batch size are determined by the memory-size of the VM, yielding $B_{max}$. Small batch sizes result in extremely high noise, and thus realistic lower-bounds on $B$ are necessary.

**IV. IMPLEMENTATION**

Scavenger is implemented as a modular extension to TensorFlow, and written in Python in about 2000 lines of code. The training scalability indicators are implemented by extending TensorFlow’s estimator framework [18]. Users simply need to download our TensorFlow distribution (or apply a patch), and no modifications are required to the models or any workflow component. The parameter server computes the SGD noise by computing the gradient norms for all the workers’ updates, and the final norm for the averaged gradient. This approximates the gradient variance, as shown in [2]. The gradient variance can be noisy, and we use exponentially weighted moving average to smoothen the output.

All the scaling indicators: the gradient noise, gradient computation time, and synchronization time, are sent to an external model service on every iteration. The model service uses these scaling indicators to update the performance model if operating in the initial exploratory search mode. The user
can select the full or partial search mode based on the search-cost and performance-model prediction accuracy requirements. By default, we use the partial-search, since its results are comparable to full-search with lower search costs. Scavenger saves all performance models on persistent storage, and the no-search strategy is used if a model has been trained before. Once the tradeoff curves are constructed, we select the best configuration and stop all profiling.

We interface with standard cloud APIs for managing VMs. Our partial-search process starts with the smallest $K, B$ configuration, and then adds more VMs to the cluster to reach the largest configuration. We use lightweight checkpointing: since the parameter server stores the latest model weights, the new workers in a new configuration pull the latest weights from the parameter server and resume training. We switch to different configurations only on iteration boundaries, and thus no work is lost. The existing VMs are always reused, to avoid excessive VM churn and startup/shutdown overheads. Although Scavenger is currently implemented in TensorFlow v1.5, its main components are modular, and need only minimal profiling information from the ML framework.

V. EXPERIMENTAL EVALUATION

Our experimental evaluation is focused on answering the following questions: 1. How effective is gradient noise as an indicator of statistical efficiency? 2. How accurate is our performance and cost model across different job configurations? 3. What are the performance and cost tradeoffs for different cloud computing cost models? 4. What are the time and cost savings achievable with our job configuration and resource allocation policies?

While most work on model training uses GPUs, we perform all evaluation on CPU VMs. GPUs simply reduce the per-iteration time, and all aspects of Scavenger such as the model and service are unaffected by the underlying hardware parallelism. Standard CPU VMs can also be sized in a fine-grained manner and we can configure the VM with arbitrary amounts of CPUs and memory. This allows us to also evaluate weak scaling: the total computing resources across all our cluster configurations are the same, but they are distributed among VMs differently. In contrast, GPUs have fixed and limited memory, and severely limit weak-scaling and batch-size scaling. Furthermore, we only consider the worker cost, and assume that sufficient parameter servers are launched and available. Parameter server allocation is tackled by other systems such as Optimus [19], and is orthogonal and complementary to our work.

A. Cost and time tradeoffs

With the performance model described in Section III, we can predict running cost and time for distributed training for various cluster configurations. Figure 10 shows the cost vs. time trade-offs for ResNet18, ResNet50 and Transformer Base to reach 80%-90% train accuracy and 18.0 BLEU score for various $B$ on Google E2-standard-4 VMs. Each scatter point show results from full runs for each $(K, B)$ configuration and dashed line shows the predicted cost and time with the offline performance model. The rental cost of each worker is $0.13402/hr. Each point on the curve represents a decreasing cluster size, with [20, 16, 12, 8] workers.

We can see that there are clear cost vs. time tradeoffs for each batch size. Here, the per-worker compute hardware is the same, and the per-hour total cluster-price is also proportional to $K$. The largest clusters have highest cost but also lowest running time. Decreasing workers reduces cost slightly but significantly increases running time.

Both the ResNet models (Figures 10a, 10b) have a single inflection/knee-point for all batch sizes, after which we see diminishing returns on cost. With Transformer model in Figure 10c, we observe two inflection points corresponding to clusters 12 and 16 for any $B$. We observed a notable decrease in iteration time from $K=12$ to 16 since for the same per-worker compute hardware and $B$, since larger $K$ implies smaller worker mini-batch size. For example, $B=768$ changes mini-batch size from 64 to 48 when $K$ goes from 12 to 16. Thus, we see a significant training time difference between $K=12$ vs. 16, resulting in two distinct inflection points.

Result: The tradeoff curves can be a crucial tool for judicious resource allocation on the cloud for distributed training.

The dashed lines in Figure 10 shows the cost predicted by our performance model using the full-search strategy, which relies on profiling of the gradient noise and iteration-time performance models by running the model on different configurations for a small number of iterations. Compared to the actual job running time, our offline performance model has an error of only 1-5%, across the entire range of models, workers, and batch sizes.
The starting configurations are \((K_{min}, B_{min})\) and \((K_{min}, B_{max})\), until the gradient noise has stabilized. With exponential moving average smoothing, noise for ResNet18, ResNet50 and Transformer Base stabilized at \(2K, 3K\) and \(10K\) iterations respectively. The total search cost for ResNet18, arising from doing this profiling on extreme configurations was minimal. The overhead of exploring a new configuration (due to checkpoint-restore) is minimal, on average 37 seconds for ResNet18, 40 for ResNet50, and 127 seconds for Transformers. Each configuration is run for around 20 iterations, which takes around 17–35 seconds for our three models. Compared to an “oracle” scenario of running on the optimal configuration all along (bypassing the search phase), our approach increases running time by 0.83 hours, and $0.89 to the final cost. This represents a 13% increase in running time and 9% increase in cost, compared to an oracle approach which runs the job on the optimal configuration from the start. Compared to arbitrary job configuration without our techniques, our running times can be more than 2× lower and costs can be more than 40% lower.

**Result:** The partial search increases job running time by 13% and cost by 9%, even compared to an oracle approach, and is a low-overhead strategy for discovering optimal configurations.

**C. Model Accuracy**

Both the full and partial search are able to accurately predict the total training time, as seen from Figure 11. We evaluate three configurations: partial search (red), full search (green), and a worst-case no-search strategy. The figure shows the distribution of the error of running time prediction vs. the empirical job running time, across different K and B. We see that the average error for partial search is 4% for ResNet and less than 2% for Transformer. The full-search is even better: with an average error of 0.5–3.5% across all models and configurations.

In Figure 11, we also evaluate our no-search strategy in a worst-case scenario. The no-search strategy performance is exactly the same as the full-search scenario, if a near-identical ML model has been trained before. However we also evaluate a “universal” performance model which averages the statistical and performance models over all the three ML models. Even this universal global-average model shows acceptable training-time prediction: the error is in the range of 4–20%. Note that this global-average model does not require any search, has no search costs, nor does it require any prior profiling or pilot runs. It is thus fully online and zero overhead.

The running-time prediction error is not highly significant to our overall objective of discovering optimal configurations. We primarily care about the relative running times, because we only compare configurations and run the job on the best-predicted configuration. It is likely that the best-predicted configuration remains the same even with the higher error, or the sub-optimal configuration chosen due to the errors is very close to the optimal configuration in the trade-off curve.

**Result:** Our performance model can predict training times with a low error of 0.5–3.5%, and only 4% even with partial searching. In the fully-online setting, the error range is 4–20%.

**D. Memory-based pricing**

The cost of training is ultimately determined by the VM cost model. So far, we have looked at conventional on-demand VM pricing, where the VM cost scales linearly according to the number of vCPUs. Scavenger can work with different cost models. We consider VMs that are priced both per CPU and also per GB of memory. Google cloud’s custom-sized VMs approximate this model.

With such a finer-grained cost model, the cost-time tradeoff curves are shown in Figure 12. In this case, the VM memory is allocated according to the batch size such that there is negligible free memory. The cost is proportional to the total memory required (the global batch size \(B\)) and running time.
Fig. 11: Error in predicted training time from actual training time across all job configurations. The error for partial and full search is low. Even the universal model, which doesn’t consider any model-specific details, provides acceptable results.

Fig. 12: Scavenger can work with different VM pricing models. In this figure, VMs are priced based on their memory size, resulting in different cost/time tradeoffs.

Comparing the results in Fig. 10 and 12, we see a shift in the inflection points for all ML workloads. This is expected since the running costs changed as a cluster $K$ training on $B = 1024$ will be pricier than that running $B = 384$, as more memory would be allocated to the former.

VI. RELATED WORK

Our work falls in the category of adapting model training on distributed infrastructure such as shared clusters and cloud platforms [20]. Scavenger uses the noise scale proposed in AdaScale SGD [2], which is similar to the gradient noise model of McCandlish et al. [3]. KungFu [4] and Pollux [5] also use this gradient noise metric for monitoring training performance and dynamically adjusting the resource allocation to minimize it. In addition to elasticity and adaptation mechanisms proposed in these papers, we use a performance profiling based approach that also takes cost into account. KungFu is complementary to our work: we can implement Scavenger’s policies as part of their adaptation-policy framework and mechanisms. Pollux also considers statistical efficiency and similar worker and batch-size tradeoffs, but is not cloud cost aware, and instead provides scheduling policies for shared clusters.

Commercial offerings of “model training as a service”, such as Amazon AWS SageMaker [21], use only rudimentary performance models, and do not use statistical efficiency or pareto-optimal allocation. Searching for hyperparameters is an important cloud workload, and reducing this search cost using parallel search techniques and early stopping provide significant cost and time savings [22]–[24]. Unlike hyperparameter optimization which focuses on reducing the cost of a “bag” of jobs, Scavenger focuses on optimizing the cost and time of a single job. Efficient elasticity mechanisms and policies for ML training [25]–[27] can also be incorporated.

Scheduling and resource allocation in shared clusters is challenging for distributed training because of the complex performance tradeoffs we have identified, and the large computing requirements. In shared private clusters, optimizing the use of limited GPU resources is a key challenge [28]–[31]. In clouds, cost is more important than resource contention.

Modeling distributed ML training poses many challenges because of the heterogeneity of ML models and their performance tradeoffs [32]. Optimus [19] models the throughput and communication costs to allocate workers and parameter servers to jobs on a shared Kubernetes cluster. Cynthia [33] minimizes cloud cost and time by scaling workers and parameter servers using a finer-grained analytical model, but does not consider batch sizes and statistical efficiency. We do not adjust the number of parameter servers and assume that they are suitably provisioned. Optimizing parameter server allocation is part of our future work. Batch-size adaptation can be important for model generalizability and performance, and can benefit from second-order gradient information [34].

VII. CONCLUSION

The training time and cost for large machine learning models is significant, and sensitive to many job and cloud configuration parameters. Scavenger is a cloud service which uses online profiling and new performance models for estimating the training performance on different cloud configurations, with high accuracy of over 95%, and reduces training time by $2\times$. Acknowledgements. This research was supported by the NSF grant 2112606.
REFERENCES


