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Distributed Asynchronous Online Learning for Natural Language Processing

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Abstract

Recent speed-ups for training large-scale models like those found in statistical NLP exploit distributed computing (either on multicore or “cloud” architectures) and rapidly converging online learning algorithms. Here we aim to combine the two. We focus on distributed, “mini-batch” learners that make frequent updates asynchronously (Nedic et al., 2001; Langford et al., 2009). We generalize existing asynchronous algorithms and experiment extensively with structured prediction problems from NLP, including discriminative, unsupervised, and non-convex learning scenarios. Our results show asynchronous learning can provide substantial speed-ups compared to distributed and single-processor mini-batch algorithms with no signs of error arising from the approximate nature of the technique.

1 Introduction

Modern statistical NLP models are notoriously expensive to train, requiring the use of general-purpose or specialized numerical optimization algorithms (e.g., gradient and coordinate ascent algorithms and variations on them like L-BFGS and EM) that iterate over training data many times. Two developments have led to major improvements in training time for NLP models:

- **online** learning algorithms (LeCun et al., 1998; Crammer and Singer, 2003; Liang and Klein, 2009), which update the parameters of a model more frequently, processing only one or a small number of training examples, called a “mini-batch,” between updates; and

- **distributed** computing, which divides training data among multiple CPUs for faster processing between updates (e.g., Clark and Curran, 2004).

Online algorithms offer fast convergence rates and scalability to large datasets, but distributed computing is a more natural fit for algorithms that require a lot of computation—e.g., processing a large batch of training examples—to be done between updates. Typically, distributed online learning has been done in a synchronous setting, meaning that a mini-batch of data is divided among multiple CPUs, and the model is updated when they have all completed processing (Finkel et al., 2008). Each mini-batch is processed only after the previous one has completed.

Synchronous frameworks are appealing in that they simulate the same algorithms that work on a single processor, but they have the drawback that the benefits of parallelism are only obtainable within one mini-batch iteration. Moreover, empirical evaluations suggest that online methods only converge faster than batch algorithms when using very small mini-batches (Liang and Klein, 2009). In this case, synchronous parallelization will not offer much benefit.

In this paper, we focus our attention on asynchronous algorithms that generalize those presented by Nedic et al. (2001) and Langford et al. (2009). In these algorithms, multiple mini-batches are processed simultaneously, each using potentially different and typically stale parameters. The key advantage of an asynchronous framework is that it allows processors to remain in near-constant use, preventing them from wasting cycles waiting for other processors to complete their portion of the current mini-batch. In this way, asynchronous algorithms allow more frequent parameter updates, which speeds convergence.

Our contributions are as follows:

- We describe a framework for distributed asynchronous optimization (§5) similar to those described by Nedic et al. (2001) and Langford et al. (2009), but permitting mini-batch learning. The prior work contains convergence results for asynchronous online stochastic gradient descent
for convex functions (discussed in brief in §5.2).

• We report experiments on three structured NLP tasks, including one problem that matches the conditions for convergence (named entity recognition; NER) and two that depart from theoretical foundations, namely the use of asynchronous stepwise EM (Sato and Ishii, 2000; Cappé and Moulines, 2009; Liang and Klein, 2009) for both convex and non-convex optimization.

• We directly compare asynchronous algorithms with multiprocessor synchronous mini-batch algorithms (e.g., Finkel et al., 2008) and traditional batch algorithms.

• We experiment with adding artificial delays to simulate the effects of network or hardware traffic that could cause updates to be made with extremely stale parameters.

• Our experimental settings include both individual 4-processor machines as well as large clusters of commodity machines implementing the MapReduce programming model (Dean and Ghemawat, 2004). We also explore effects of mini-batch size.

Our main conclusion is that, when small mini-batches work well, asynchronous algorithms offer substantial speed-ups without introducing error. When large mini-batches work best, asynchronous learning does not hurt.

2 Optimization Setting

We consider the problem of optimizing a function $f : \mathbb{R}^d \rightarrow \mathbb{R}$ with respect to its argument, denoted $\theta = (\theta_1, \theta_2, \ldots, \theta_d)$. We assume that $f$ is a sum of $n$ convex functions (hence $f$ is also convex):\footnote{We use “convex” to mean convex-up when minimizing and convex-down, or concave, when maximizing.}

$$f(\theta) = \sum_{i=1}^{n} f_i(\theta)$$

We initially focus our attention on functions that can be optimized using gradient or subgradient methods. Log-likelihood for a probabilistic model with fully observed training data (e.g., conditional random fields; Lafferty et al., 2001) is one example that frequently arises in NLP, where the $f_i(\theta)$ each correspond to an individual training example and the $\theta$ are log-linear feature weights. Another example is large-margin learning for structured prediction (Taskar et al., 2005), which can be solved by subgradient methods (Ratliff et al., 2006).

For concreteness, we discuss the architecture in terms of gradient-based optimization, using the following gradient descent update rule (for minimization problems):\footnote{We use the term “gradient” for simplicity, but subgradients are sufficient throughout.}

$$\theta^{(t+1)} = \theta^{(t)} - \eta^{(t)} g(\theta^{(t)})$$

where $\theta^{(t)}$ is the parameter vector on the $t$th iteration, $\eta^{(t)}$ is the step size on the $t$th iteration, and $g : \mathbb{R}^d \rightarrow \mathbb{R}^d$ is the vector function of first derivatives of $f$ with respect to $\theta$:

$$g(\theta) = \left\{ \frac{\partial f_1}{\partial \theta_1}(\theta), \frac{\partial f_2}{\partial \theta_2}(\theta), \ldots, \frac{\partial f_n}{\partial \theta_n}(\theta) \right\}$$

We are interested in optimizing such functions using distributed computing, by which we mean to include any system containing multiple processors that can communicate in order to perform a single task. The set of processors can range from two cores on a single machine to a MapReduce cluster of thousands of machines.

Note our assumption that the computation required to optimize $f$ with respect to $\theta$ is, essentially, the gradient vector $g(\theta^{(t)})$, which serves as the descent direction. The key to distributing this computation is the fact that $g(\theta^{(t)}) = \sum_{i=1}^{n} g_i(\theta^{(t)})$, where $g_i(\theta)$ denotes the gradient of $f_i(\theta)$ with respect to $\theta$. We now discuss several ways to go about distributing such a problem, culminating in the asynchronous mini-batch setting.

3 Distributed Batch Optimization

Given $p$ processors plus a master processor, the most straightforward way to optimize $f$ is to partition the $f_i$ so that for each $i \in \{1, 2, \ldots, n\}$, $g_i$ is computed on exactly one “slave” processor. Let $I_j$ denote the subset of examples assigned to the $j$th slave processor ($\bigcup_{j=1}^{p} I_j = \{1, \ldots, n\}$ and $j \neq j' \Rightarrow I_j \cap I_{j'} = \emptyset$). Processor $j$ receives the examples in $I_j$ along with the necessary portions of $\theta^{(t)}$ for calculating $g_{i_{j'}}(\theta^{(t)}) = \sum_{i \in I_{j'}} g_i(\theta^{(t)})$. The result of this calculation is returned to the master processor, which calculates $g(\theta^{(t)}) = \sum_{j} g_{i_{j}}(\theta^{(t)})$ and executes Eq. 2 (or something more sophisticated that uses the same information) to obtain a new parameter vector.

It is natural to divide the data so that each processor is assigned approximately $n/p$ of the training examples. Because of variance in the expense
of calculating the different \( g_i \), and because of unpredictable variation among different processors’ speed (e.g., variation among nodes in a cluster, or in demands made by other users), there can be variation in the observed runtime of different processors on their respective subsamples. Each iteration of calculating \( g \) will take as long as the longest-running among the processors, whatever the cause of that processor’s slowness. In computing environments where the load on processors is beyond the control of the NLP researcher, this can be a major bottleneck.

Nonetheless, this simple approach is widely used in practice; approaches in which the gradient computation is distributed via MapReduce have recently been described in machine learning and NLP (Chu et al., 2006; Dyer et al., 2008; Wolfe et al., 2008). Mann et al. (2009) compare this framework to one in which each processor maintains a separate parameter vector which is updated independently of the others. At the end of learning, the parameter vectors are averaged or a vote is taken during prediction. A similar parameter-averaging approach was taken by Chiang et al. (2008) when parallelizing MIRA (Crammer et al., 2006). In this paper, we restrict our attention to distributed frameworks which maintain and update a single copy of the parameters \( \theta \). The use of multiple parameter vectors is essentially orthogonal to the framework we discuss here and we leave the integration of the two ideas for future exploration.

4 Distributed Synchronous Mini-Batch Optimization

Distributed computing can speed up batch algorithms, but we would like to transfer the well-known speed-ups offered by online and mini-batch algorithms to the distributed setting as well. The simplest way to implement mini-batch stochastic gradient descent (SGD) in a distributed computing environment is to divide each mini-batch (rather than the entire batch) among the processors that are available and to update the parameters once the gradient from the mini-batch has been computed. Finkel et al. (2008) used this approach to speed up training of a log-linear model for parsing. The interaction between the master processor and the distributed computing environment is nearly identical to the distributed batch optimization scenario. Where \( M(t) \) is the set of indices in the mini-batch processed on iteration \( t \), the update is:

\[
\theta(t+1) \leftarrow \theta(t) - \eta(t) \sum_{i \in M(t)} g_i(\theta(t))
\]

(4)

The distributed synchronous framework can provide speed-ups over a single-processor implementation of SGD, but inevitably some processors will end up waiting for others to finish processing. This is the same bottleneck faced by the batch version in §3. While the time for each mini-batch is shorter than the time for a full batch, mini-batch algorithms make far more updates and some processor cycles will be wasted in computing each one. Also, more mini-batches imply that more time will be lost due to per-mini-batch overhead (e.g., waiting for synchronization locks in shared-memory systems, or sending data and \( \theta \) to the processors in systems without shared memory).

5 Distributed Asynchronous Mini-Batch Optimization

An asynchronous framework may use multiple processors more efficiently and minimize idle time (Nedic et al., 2001; Langford et al., 2009). In this setting, the master sends \( \theta \) and a mini-batch \( M_k \) to each slave \( k \). Once slave \( k \) finishes processing its mini-batch and returns \( g_{M_k}(\theta) \), the master immediately updates \( \theta \) and sends a new mini-batch and the new \( \theta \) to the now-available slave \( k \). As a result, slaves stay occupied and never need to wait on others to finish. However, nearly all gradient components are computed using slightly stale parameters that do not take into account the most recent updates. Nedic et al. (2001) proved that convergence is still guaranteed under certain conditions, and Langford et al. (2009) obtained convergence rate results. We describe these results in more detail in §5.2.

The update takes the following form:

\[
\theta(t+1) \leftarrow \theta(t) - \eta(t) \sum_{i \in M(\tau(t))} g_i(\theta(\tau(t)))
\]

(5)

where \( \tau(t) \leq t \) is the start time of the mini-batch used for the \( t \)th update. Since we started processing the mini-batch at time \( \tau(t) \) (using parameters \( \theta(\tau(t)) \)), we denote the mini-batch \( M(\tau(t)) \). If \( \tau(t) = t \), then Eq. 5 is identical to Eq. 4. That is, \( t - \tau(t) \) captures the “ staleness” of the parameters used to compute the gradient for the \( t \)th update.

Asynchronous frameworks do introduce error into the training procedure, but it is frequently the case in NLP problems that only a small fraction of parameters is needed for each mini-batch
of training examples. For example, for simple word alignment models like IBM Model 1 (Brown et al., 1993), only parameters corresponding to words appearing in the particular subsample of sentence pairs are needed. The error introduced when making asynchronous updates should intuitively be less severe in these cases, where different mini-batches use small and mostly non-overlapping subsets of θ.

5.1 Implementation

The algorithm sketched above is general enough to be suitable for any distributed system, but when using a system with shared memory (e.g., a single multiprocessor machine) a more efficient implementation is possible. In particular, we can avoid the master/slave architecture and simply start p threads that each compute and execute updates independently, with a synchronization lock on θ. In our single-machine experiments below, we use Algorithm 1 for each thread. A different random seed (r) is passed to each thread so that they do not all process the same sequence of examples. At completion, the result is contained in θ.

5.2 Convergence Results

We now briefly summarize convergence results from Nedic et al. (2001) and Langford et al. (2009), which rely on the following assumptions: (i) The function f is convex. (ii) The gradients g_t are bounded, i.e., there exists $C > 0$ such that $\|g_t(\theta(t))\| \leq C$. (iii) $\exists$ (unknown) $D > 0$ such that $t - \tau(t) < D$. (iv) The stepsizes $\eta(t)$ satisfy certain standard conditions.

In addition, Nedic et al. require that all function components are used with the same asymptotic frequency (as $t \to \infty$). Their results are strongest when choosing function components in each mini-batch using a “cyclic” rule: select function $f_i$ for the $k$th time only after all functions have been selected $k - 1$ times. For a fixed step size $\eta$, the sequence of function values $f(\theta^{(i)})$ converges to a region of the optimum that depends on $\eta$, the maximum norm of any gradient vector, and the maximum delay for any mini-batch. For a decreasing step size, convergence is guaranteed to the optimum. When choosing components uniformly at random, convergence to the optimum is again guaranteed using a decreasing step size formula, but with slightly more stringent conditions on the step size.

Langford et al. (2009) present convergence rates via regret bounds, which are linear in $D$. The convergence rate of asynchronous stochastic gradient descent is $O(\sqrt{T \bar{D}})$, where $T$ is the total number of updates made. In addition to the situation in which function components are chosen uniformly at random, Langford et al. provide results for several other scenarios, including the case in which an adversary supplies the training examples in whatever order he chooses.

Below we experiment with optimization of both convex and non-convex functions, using fixed step sizes and decreasing step size formulas, and consider several values of $D$. Even when exploring regions of the experimental space that are not yet supported by theoretical results, asynchronous algorithms perform well empirically in all settings.

5.3 Gradients and Expectation-Maximization

The theory applies when using first-order methods to optimize convex functions. Though the function it is optimizing is not usually convex, the EM algorithm can be understood as a hillclimber that transforms the gradient to keep $\theta$ feasible; it can also be understood as a coordinate ascent algorithm. Either way, the calculations during the E-step resemble $g(\theta)$. Several online or mini-batch variants of the EM algorithm have been proposed, for example incremental EM (Neal and Hinton, 1998) and online EM (Sato and Ishii, 2000; Cappé and Moulines, 2009), and we follow Liang and Klein (2009) in referring to this latter algorithm as stepwise EM. Our experiments with asynchronous minibatch updates include a case where the log-likelihood $f$ is convex and one where it is not.

\begin{algorithm}
\caption{Procedure followed by each thread for multi-core asynchronous mini-batch optimization. $\theta$ is the single copy of the parameters shared by all threads. The convergence criterion is left unspecified here.}
\begin{algorithmic}
\Procedure{Algorithm 1}{
\State \textbf{Input:} number of examples $n$, mini-batch size $m$, random seed $r$ \nl \State $\theta_0 \leftarrow \theta$ \nl \State $\text{seedRandomNumberGenerator}(r)$ \nl $\text{while}$ \text{converged}(\theta) = \text{false} \nl \State $g \leftarrow 0$ \nl \For {$j \leftarrow 1$ \textbf{to} $m$} \nl \State $k \sim \text{Uniform}\{1, \ldots, n\}$ \nl \State $g \leftarrow g + g_k(\theta_t)$ \nl \EndFor \nl \State $\text{acquireLock}(\theta)$ \nl \State $\theta \leftarrow \text{updateParams}(\theta, g)$ \nl \State $\theta_t \leftarrow \theta$ \nl \State $\text{releaseLock}(\theta)$ \nl $\text{end}$
\end{algorithmic}
\end{algorithm}
### 6 Experiments

We performed experiments to measure speed-ups obtainable through distributed online optimization. Since we will be considering different optimization algorithms and computing environments, we will primarily be interested in the wall-clock time required to obtain particular levels of performance on metrics appropriate to each task. We consider three tasks, detailed in Table 1.

For experiments on a single node, we used a 64-bit machine with two 2.6GHz dual-core CPUs (i.e., 4 processors in all) with a total of 8GB of RAM. This was a dedicated machine that was not available for any other jobs. We also conducted experiments using a cluster architecture running Hadoop 0.20 (an implementation of MapReduce), consisting of 400 machines, each having 2 quad-core 1.86GHz CPUs with a total of 6GB of RAM.

#### 6.1 Named Entity Recognition

Our NER CRF used a standard set of features, following Kazama and Torisawa (2007), along with token shape features like those in Collins (2002) and simple gazetteer features; a feature was included if and only it occurred at least once in training data (total 1.3M). We used a diagonal Gaussian prior with a variance of $1.0$ for each weight.

We compared SGD on a single processor to distributed synchronous SGD and distributed asynchronous SGD. For all experiments, we used a fixed step size of 0.01 and chose each training example for each mini-batch uniformly at random from the full data set. We report performance by plotting test-set accuracy against wall-time over 12 hours.4

#### Comparing Synchronous and Asynchronous Algorithms

Figure 1 shows our primary result for the NER experiments. When using all four available processors, the asynchronous algorithm converges faster than the other two algorithms. Error due to stale parameters during gradient computation does not appear to cause any more various fixed step sizes and decreasing step size schedules, and found a fixed step size to work best for all settings.

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3In preliminary experiments, we experimented with various fixed step sizes and decreasing step size schedules, and found a fixed step size to work best for all settings.

4Decoding was performed offline (so as not to affect measurements) with models sampled every ten minutes.
tion in performance than experienced by the synchronous mini-batch algorithm. Note that the distributed synchronous algorithm and the single-processor algorithm make identical sequences of parameter updates; the only difference is the amount of time between each update. Since we save models every ten minutes and not every \( \mu \)th update, the curves have different shapes. The sequence of updates for the asynchronous algorithm, on the other hand, actually depends on the vagaries of the computational environment. Nonetheless, the asynchronous algorithm using 4 processors has nearly converged after only 2 hours, while the single-processor algorithm requires 10–12 hours to reach the same \( F_1 \).

**Varying the Number of Processors** Figure 2 shows the improvement in convergence time by using 4 vs. 2 processors for the synchronous (top) and asynchronous (bottom) algorithms. The additional two processors help the asynchronous algorithm more than the synchronous one. This highlights the key advantage of asynchronous algorithms: it is easier to keep all processors in constant use. Synchronous algorithms might be improved through load-balancing; in our experiments here, we simply assigned \( m/p \) examples to each processor, where \( m \) is the mini-batch size and \( p \) is the number of processors. When \( m = p \), as in the 4-processor curve in the upper plot of Figure 2, we assign a single example to each processor; this is optimal in the sense that no other scheduling strategy will process the mini-batch faster. Therefore, the fact that the 2-processor and 4-processor curves are so close suggests that the extra two processors are not being fully exploited, indicating that the optimal load balancing strategy for a small mini-batch still leaves processors under-used due to the synchronous nature of the updates.

The only bottleneck in the asynchronous algorithm is the synchronization lock during updating, required since there is only one copy of \( \theta \). For CRFs with a few million weights, the update is typically much faster than processing a mini-batch of examples; furthermore, when using small mini-batches, the update vector is typically sparse.\(^5\) For all experimental results presented thus far, we used a mini-batch size of 4. We experimented with additional mini-batch sizes of 1 and 8, but there was very little difference in the resulting curves.

**Artificial Delays** We experimented with adding artificial delays to the algorithm to explore how much overhead would be tolerable before parallelized computation becomes irrelevant. Figure 3 shows results when each processor sleeps with 0.25 probability for a duration of time between computing the gradient on its mini-batch of data and updating the parameters. The delay length is chosen from a normal distribution with the means (in seconds) shown and \( \sigma = \mu/5 \) (truncated at zero). Since only one quarter of the mini-batches have an artificial delay, increasing \( \mu \) increases the average parameter “staleness”, letting us see how the asynchronous algorithm fares with extremely stale parameters.

The average time required to compute the gradient for a mini-batch of 4 is 0.62 seconds. When the average delay is 1.25 seconds (\( \mu = 5 \)), twice the average time for a mini-batch, the asynchronous algorithm still converges faster than the single-node algorithm. In addition, even with substantial delays of 5–10 times the processing time for a mini-batch, the asynchronous algorithm does not fail but proceeds steadily toward convergence.

The practicality of using the asynchronous algorithm depends on the average duration for a mini-batch and the amount of expected additional overhead. We attempted to run these experiments on

\(^5\)In a standard implementation, the sparsity of the update will be nullified by regularization, but to improve efficiency in practice the regularization penalty can be accumulated and applied less frequently than every update.
Experiments on a Single Machine

We followed Liang and Klein (2009) in using synchronous (mini-batch) stepwise EM on a single processor for this task. We used the same learning rate formula \( \eta(t) = (t + 2)^{-q} \), with \( 0.5 < q \leq 1 \). We also used asynchronous stepwise EM by using the same update rule, but gathered sufficient statistics on 4 processors of a single machine in parallel, analogous to our asynchronous method from §5. Whenever a processor was done gathering the expected counts for its mini-batch, it updated the sufficient statistics vector and began work on the next mini-batch.

We used the sparse update described by Liang and Klein, which allows each thread to make additive updates to the parameter vector and to separately-maintained normalization constants without needing to renormalize after each update. When probabilities are needed during inference, normalizers are divided out on-the-fly as needed.

We made 10 passes of asynchronous stepwise EM to measure its sensitivity to \( q \) and the mini-batch size \( m \), using different values of these hyperparameters \( \{0.5, 0.7, 1.0\}; \ m \in \{5000, 10000, 50000\} \), and selected values that maximized log-likelihood \( q = 0.7, m = 10000 \).

Experiments on MapReduce

We implemented the three techniques in a MapReduce framework. We implemented batch EM on MapReduce by converting each EM iteration into two MapReduce jobs: one for the E-step and one for the M-step.\(^6\) For the E-step, we divided our data into 24 map tasks, and computed expected counts for the source-target parameters at each mapper. Next, we summed up the expected counts in one reduce task. For the M-step, we took the output from the E-step, and in one reduce task, normalized each source-target parameter by the total count for the source word.\(^7\) To gather sufficient statistics for synchronous stepwise EM, we used 6 mappers and one reducer for a mini-batch of size 10000. For the asynchronous version, we ran four parallel asynchronous mini-batches, the sufficient statistics being gathered using MapReduce again for each mini-batch with 6 map tasks and one reducer.

Results

Figure 4 shows log-likelihood for the English→French direction during the first 80 minutes of optimization. Similar trends were observed for the French→English direction as well as for convergence in AER. Table 2 shows the AER at the end of 20 iterations of EM for the same settings.\(^8\) It takes around two hours to finish 20 iterations of batch EM on a single machine, while it takes more than 8 hours to do so on MapReduce. This is because of the extra overhead of transferring \( \theta \) from a master gateway machine to mappers, from mappers to reducers, and from reducers back to the master. Synchronous and asynchronous EM suffer as well.

From Figure 4, we see that synchronous and asynchronous stepwise EM converge at the same rate when each is given 4 processors. The main difference between this task and NER is the size of the mini-batch used, so we experimented with several values for the mini-batch size \( m \). Figure 5 shows the results. As \( m \) decreases, a larger fraction of time is spent updating parameters; this slows observed convergence time even when using the sparse update rule. It can be seen that, though synchronous and asynchronous stepwise EM converge at the same rate with a large mini-batch size \( m = 10000 \), asynchronous stepwise

\(^6\)The M-step could have been performed without MapReduce by storing all the parameters in memory, but memory restrictions on the gateway node of our cluster prevented this.

\(^7\)For the reducer in the M-step, the source served as the key, and the target appended by the parameter’s expected count served as the value.

\(^8\)Note that for wall time comparison, we sample models every five minutes. The time taken to write these models ranges from 30 seconds to a minute, thus artificially elongating the total time for all iterations.
EM converges faster as \( m \) decreases. With large mini-batches, load-balancing becomes less important as there will be less variation in per-mini-batch observed runtime. These results suggest that asynchronous mini-batch algorithms will be most useful for learning problems in which small mini-batches work best. Fortunately, however, we do not see any problems stemming from approximation errors due to the use of asynchronous updates.

### 6.3 Unsupervised POS Tagging

Our unsupervised POS experiments use the same task and approach of Liang and Klein (2009) and so we fix hyperparameters for stepwise EM based on their findings (learning rate \( \eta^{(t)} = (t+2)^{-0.7} \)). The asynchronous algorithm uses the same learning rate formula as the single-processor algorithm. There is only a single \( t \) that is maintained and gets incremented whenever any thread updates the parameters. Liang and Klein used a mini-batch size of 3, but we instead use a mini-batch size of 4 to better suit our 4-processor synchronous and asynchronous architectures.

Like NER, we present results for unsupervised tagging experiments on a single machine only, i.e., not using a MapReduce cluster. For tasks like POS tagging that have been shown to work best with small mini-batches (Liang and Klein, 2009), we did not conduct experiments with MapReduce due to high overhead per mini-batch.

For initialization, we followed Liang and Klein by initializing each parameter as \( \theta_i \sim e^{1+a_i} \), \( a_i \sim \text{Uniform}([0, 1]) \). We generated 5 random models using this procedure and used each to initialize each algorithm. We additionally used 2 random seeds for choosing the ordering of examples,\(^9\) resulting in a total of 10 runs for each algorithm. We ran each for six hours, saving models every five minutes. After training completed, using each model we decoded the entire training data using posterior decoding and computed the log-likelihood. The results for 5 initial models and two example orderings are shown in Figure 6. We evaluated tagging performance using many-to-1 accuracy, which is obtained by mapping the HMM states to gold standard POS tags so as to maximize accuracy, where multiple states can be mapped to the same tag. This is the metric used by Liang and Klein (2009) and Johnson (2007), who report figures comparable to ours. The asynchronous algorithm converges much faster than the single-node algorithm, allowing a tagger to be trained from the Penn Treebank in less than two hours using a single machine. Furthermore, the 4-processor synchronous algorithm improves only marginally.

\(^9\)We ensured that the examples processed in the sequence of mini-batches were identical for the 1-processor and 4-processor versions of synchronous stepwise EM, but the asynchronous algorithm requires a different seed for each processor and, furthermore, the actual order of examples processed depends on wall times and cannot be controlled for. Nonetheless, we paired a distinct set of seeds for the asynchronous algorithm with each of the two seeds used for the synchronous algorithms.
over the 1-processor baseline.

The accuracy of the asynchronous curves often decreases slightly after peaking. We can surmise from the log-likelihood plot that the drop in accuracy is not due to the optimization being led astray, but probably rather due to the complex relationship between likelihood and task-specific evaluation metrics in unsupervised learning (Merialdo, 1994). In fact, when we examined the results of synchronous stepwise EM between 6 and 12 hours of execution, we found similar drops in accuracy as likelihood continued to improve. From Figure 6, we conclude that the asynchronous algorithm has no harmful effect on learned model’s accuracy beyond the choice to optimize log-likelihood.

While there are currently no theoretical convergence results for asynchronous optimization algorithms for non-convex functions, our results are encouraging for the prospects of establishing convergence results for this setting.

7 Discussion

Our best results were obtained by exploiting multiple processors on a single machine, while experiments using a MapReduce cluster were plagued by communication and framework overhead.

Since Moore’s Law predicts a continual increase in the number of cores available on a single machine but not necessarily an increase in the speed of those cores, we believe that algorithms that can effectively exploit multiple processors on a single machine will be increasingly useful. Even today, applications in NLP involving rich-feature structured prediction, such as parsing and translation, typically use a large portion of memory for storing pre-computed data structures, such as lexicons, feature name mappings, and feature caches. Frequently these are large enough to prevent the multiple cores on a single machine from being used for multiple experiments, leaving some processors unused. However, using multiple threads in a single program allows these large data structures to be shared and allows the threads to make use of the additional processors.

We found the overhead incurred by the MapReduce programming model, as implemented in Hadoop 0.20, to be substantial. Nonetheless, we found that asynchronously running multiple MapReduce calls at the same time, rather than pooling all processors into a single MapReduce call, improves observed convergence with negligible effects on performance.

8 Conclusion

We have presented experiments using an asynchronous framework for distributed mini-batch optimization that show comparable performance of trained models in significantly less time than traditional techniques. Such algorithms keep processors in constant use and relieve the programmer from having to implement load-balancing schemes for each new problem encountered. We expect asynchronous learning algorithms to be broadly applicable to training NLP models.

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