HarpLDA+: Optimizing Latent Dirichlet Allocation for Parallel Efficiency

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Abstract—Latent Dirichlet Allocation (LDA) is a widely used machine learning technique in topic modeling and data analysis. Training large LDA models on big datasets involves dynamic and irregular computation patterns and is a major challenge to both algorithm optimization and system design. In this paper, we present a comprehensive benchmarking of our novel synchronized LDA training system HarpLDA+ based on Hadoop and Java. It demonstrates impressive performance when compared to three other MPI/C++ based state-of-the-art systems, which are LightLDA, F+NomadLDA, and WarpLDA. HarpLDA+ uses optimized collective communication with a timer control for load balance, leading to stable scalability in both shared-memory and distributed systems. We demonstrate in the experiments that HarpLDA+ is effective in reducing synchronization and communication overhead and outperforms the other three LDA training systems.

I. INTRODUCTION

Latent Dirichlet Allocation (LDA) [1] is a widely used machine learning technique in topic modeling and data analysis. LDA training is an iterative process, which starts from a randomly initialized model with parameters to learn, iteratively computing and updating the model until it converges. A major challenge of scaling is due to the fact that computation is irregular and the model size can be huge. In the meantime, parallel workers need to synchronize the model continually.

State-of-the-art LDA training systems (trainers) are implemented to handle billions of documents, hundreds of billion tokens, millions of topics and millions of unique tokens. However, the pros and cons of different approaches in the existing tools are often hard to explain because of the many trade-offs between effectiveness (contributions to converge) and efficiency (computational cost) of model updates. Note that model update efficiency should be distinguished from the traditional parallel efficiency of speedup over parallelism.

One of the popular approaches is to decrease the time complexity of the computation by introducing approximations. Another widely used idea is to reduce the synchronization overhead by using an asynchronous parallel system working on a stale model, where the trainer is not using the latest model data. Although these approaches improve model update efficiency, they are done at the cost of the model update effectiveness for convergence. Our approach however, is to use a synchronized system and optimize LDA trainers from a different perspective. The aim is to preserve the effectiveness and at the same time improve parallel efficiency by reducing the synchronization overhead.

Our main contributions can be summarized as follows:
• Review state-of-the-art LDA training systems and summarize their design features.
• Propose new mechanisms to reduce overhead in synchronized systems, dynamic scheduling for shared memory subsystems and Timer Control for distributed systems.
• Implement HarpLDA+ based on Hadoop while demonstrating excellent performance and scalability.
• Summarize our system design approach and its implications for other machine learning algorithms.

In this paper, Section II introduces the background of the LDA algorithm and related work, while Section III analyzes the architecture and parallel efficiency of existing solutions. Section IV describes our system design and implementation details of HarpLDA+ and Section V presents experimental results coupled with a performance analysis. Finally, Section VI draws conclusions and discusses future work.

II. LDA ALGORITHM AND RELATED WORK

A. LDA with Collapsed Gibbs Sampling

Figure 1: Latent Dirichlet Allocation. N and M are sufficient statistics for the probability distribution \( \phi \) and \( \theta \) respectively in the original graphical model.

LDA is a topic modeling technique to discover latent structures inside data. As shown in Fig. 1, data is represented
as a collection of documents, where each document is a bag of words. LDA models each document as a mixture of latent topics and each topic as a multinomial distribution over words.

Many algorithms have been proposed to estimate the parameters for the LDA model. Collapsed Gibbs Sampling (CGS) [2], a Markov chain Monte Carlo (MCMC) algorithm, is commonly used for large scale LDA training. In the MCMC framework, samples can be drawn according to the unknown posterior distribution by a carefully designed transition function that visits the whole parameter space. Gibbs sampling is one such design that visits the parameter space from one dimension to the other. For each iteration, it fixes all the states of other dimensions and only updates the current visiting one.

In CGS, each training data point or token is assigned to a random topic denoted as $z_{ij}$ at initialization. Then it begins to reassign topics to each token at position $i$ in document $j$, $x_{ij} = w$, by sampling from a multinomial distribution of a conditional probability of $z_{ij}$ as shown below:

$$p(z_{ij} = k | z_{-ij}, x, \alpha, \beta) \propto \frac{N_{wk} + \beta}{\sum_w N_{wk} + V\beta} \left( M_{kj}^\alpha + \alpha \right)$$

(1)

Here superscript $-ij$ indicates that the corresponding token is excluded. $V$ is the vocabulary size, $N_{wk}$ is the token count of word $w$ assigned to topic $k$ in $K$ topics, and $M_{kj}$ is the token count of topic $k$ assigned in document $j$. The matrices $Z$, $N$ and $M$, form the model to be learned. Hyper-parameters $\alpha$ and $\beta$ control the topic density in the final model. The model gradually converges during the process of iterative sampling.

Although CGS generally requires a large number of iterations to converge, it is memory efficient and therefore scalable for large models. In this paper, we focus on the LDA trainers under the CGS algorithm family.

### B. Related Work on Parallel LDA-CGS

Gibbs sampling in LDA-CGS is a strictly sequential process. Approximate Distributed LDA (AD-LDA)[3] proposed to relax the requirement of sequential sampling of topic assignments based on the observation that the dependence between the update of one topic assignment $z_{ij}$ and the update of any other topic assignment $z_{ij'}$ is weak. In AD-LDA, the distributed approach is to partition the training data for different workers, run local CGS training and synchronize the model by merging back to a single and consistent set of $N$. PLDA [4] implemented the AD-LDA algorithm in both MPI and MapReduce, where the Allreduce operation was used for synchronization.

A synchronized algorithm that requires global synchronization at each iteration sometimes may not seem feasible or efficient; therefore, an asynchronous solution becomes the alternative choice. Async-LDA [5] extended AD-LDA to an asynchronous solution by a gossip protocol. YahooLDA [6][7] was the first production level LDA trainer. The mechanism is an asynchronous reconciliation of the model, one word at a time for all samplers. Furthermore, Parameter Server [8] was introduced as a general framework that scaled to thousands of nodes. Another advancement was presented by [9]. It proposed a “mixed” approach SSP (Stale Synchronous Parallel), which is a parameter server that can limit the maximum age of the staleness.

Some researchers have investigated synchronized algorithms. For instance, a novel data partitioning scheme [10] was proposed to avoid memory access conflicts on GPUs. The basic idea is to partition the training data into blocks, where all samplers start from the diagonal blocks and then shift to the right neighbor all together. In contrast, a general machine learning framework Petuum Strads [11][12] extended this idea, where parameters of the ML program were partitioned for different workers. As this type of communication pattern observed in the asynchronous trainers is hard to optimize, synchronized designs were proposed with collective communication operators [13] which achieved better performance. Finally, F+NomadLDA [14] was introduced based on the idea of NOMAD [15], in which each variable (one row of $N$) becomes the basic unit to be scheduled, and the ownership of a variable is asynchronously transferred between workers in a decentralized fashion.

Other research involves optimization to the sampling algorithm. According to Equation (1), a naive implementation involves drawing a sample from a discrete distribution which contains two steps: first calculate the probability of each event as $p(z_{ij} = k), k \in K$, secondly generate a random number uniformly from $[0,1)$ and search linearly along the array of the probabilities, stopping when the accumulation of probability mass is greater than or equal to the random number. The time complexity for this is $O(K)$. SparseLDA [16] decomposed the numerator of Equation (1) into three parts: $\alpha \beta, \beta * M_{kj}$ and $N_{wk}(M_{kj} + \alpha)$. The first part is a constant; while the second part is non-zero only when $M_{kj}$ is non-zero, and the third part is non-zero only when $N_{wk}$ is non-zero. Both the probability calculation and search part can benefit from utilizing the characteristics of this sparseness pertaining to the model. When using this feature, the computation time complexity drops to $O(K_d + K_w)$, which is equivalent to the average number of non-zero items in column of $M$ and row of $N$ and is typically much smaller than $K$. F+NomadLDA provides an optimization on the search part by using a $O(log K)$ binary tree search instead of a $O(K)$ linear search by a tree data structure. Based on Alias Table which allows us to draw subsequent samples from the same distribution in $O(1)$ time, Alias-LDA [17] uses the Metropolis Hasting (MH) sampling process to draw each sample correctly from the stale alias table and achieves $O(K_d)$ complexity. LightLDA [18] extends the Alias-LDA idea by decomposing Equation (1) into two
Table I: System Architectures of LDA Trainers. AllReduce, works on a stale model and does synchronization on model replicas. Asynchronous, works on local replicas and synchronizes them in a best effort through a group of parameter servers. Rotation, works on distributed model partitions and the model partitions ‘rotate’ among the workers while at the same time keeping model updates conflict free.

<table>
<thead>
<tr>
<th>Trainer</th>
<th>Sampler</th>
<th>Sampling Time Complexity</th>
<th>Intra-node Design</th>
<th>Inter-node Design</th>
<th>Comm</th>
<th>Model</th>
</tr>
</thead>
<tbody>
<tr>
<td>PLDA</td>
<td>PlainLDA</td>
<td>$O(K)$</td>
<td>Allreduce</td>
<td>Allreduce</td>
<td>stale</td>
<td></td>
</tr>
<tr>
<td>YahooLDA</td>
<td>SparseLDA</td>
<td>$O(K_d + K_w)$</td>
<td>Allreduce</td>
<td>Asynchronous</td>
<td>async</td>
<td>stale</td>
</tr>
<tr>
<td>StratsLDA</td>
<td>SparseLDA</td>
<td>$O(K_d + K_w)$</td>
<td>Allreduce</td>
<td>Rotation</td>
<td>async</td>
<td>stale</td>
</tr>
<tr>
<td>LightLDA</td>
<td>MH</td>
<td>$O(1)$</td>
<td>Asynchronous</td>
<td>Asynchronous</td>
<td>async</td>
<td>stale</td>
</tr>
<tr>
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<td>F+Tree</td>
<td>$O(\log K_d + \log K_w)$</td>
<td>Rotation</td>
<td>Rotation</td>
<td>async</td>
<td>latest</td>
</tr>
<tr>
<td>WarpLDA</td>
<td>MH</td>
<td>$O(1)$</td>
<td>DelayUpdates</td>
<td>Rotation</td>
<td>collective</td>
<td>stale</td>
</tr>
<tr>
<td>HarpLDA+</td>
<td>SparseLDA</td>
<td>$O(K_d + K_w)$</td>
<td>Rotation</td>
<td>Rotation</td>
<td>collective</td>
<td>latest</td>
</tr>
</tbody>
</table>

III. PARALLEL DESIGN PRINCIPLES

A. Parallel Efficiency

Parallelizing a sequential algorithm inevitably introduces overhead. Parallel efficiency can be measured by Speedup, which is defined as parallel performance over original sequential performance in parallel processing, where we have:

$$Speedup = \frac{T_1}{T_P} = \frac{1}{f + s + \frac{f}{P}} \quad (2)$$

With $P$ workers, $f$ is the serial portion that cannot be parallelized and parallelism introduces a time overhead of $s \times T_1$. In parallel CGS, $f$ is small, so the overhead time of $s$ becomes the major issue in order to achieve good parallel efficiency. Communication overhead comes from the additional cost of moving data among the parallel workers. In a shared memory system, this overhead is generally ignored with the assumption of a uniform memory access cost. In a distributed system, asynchronous communication or pipelining can be used to overlap communication with computation and reduce this overhead. Synchronization overhead comes from the additional cost of coordinating parallel workers that reach the same state in order to finish a task together. Asynchronous trainers try to reduce this type of overhead by avoiding a global consensus, relaxing the consistency of the model and working in an independent fashion. Synchronized trainers, however, will face the issue of load imbalance, which is a major source of synchronization overhead.

Some data partitioning algorithms have been proposed that aim to improve load balancing for LDA training. For example, random permutations on the document usually give good results. Some algorithms partition the word-topic model, whereas randomized algorithms do not perform as well as greedy algorithms [13][19] since the word frequency follows the power-law distribution. Unfortunately, even optimal partitioning algorithms cannot completely solve the load imbalance problem. Sampling algorithms may perform differently on the same number of tokens with different distributions. In practice, variations of node performance and stragglers are not uncommon even in homogeneous HPC clusters.

B. System Architectures

Machine learning algorithms can generally tolerate some kind of staleness in the model. Using stale models in computation can degrade the convergence rate but potentially boost the system efficiency because it relaxes the constraints for system design. For example, the sum of the topic count $\sum_w N_{wk}$ in the denominator of Equation (1) is hard to keep in strict consistency while in a parallel situation, because using locks on data being frequently accessed will give poor performance. A typical solution is to remove the locks and keep using a local copy of the model. Furthermore, synchronizing the model at the end of each epoch is sufficient as the deviations are small. However, the decision of whether to use stale values of $N_{wk}$ and $M_{kj}$ in the numerator of Equation (1) is more sensitive to model convergence.

In order to better present HarpLDA+’s design, we summarize the features and parallel architectures of current CGS trainers in a model centric view (see Table I).

IV. HARPLDA+: DESIGN AND IMPLEMENTATION

HarpLDA+ builds upon Harp\(^1\), which includes a Java collective communication library released as a plugin for Hadoop. While our previous work [13] optimizes communications in different architectures, HarpLDA+ focuses on the Rotation architecture and reducing the synchronization overhead.

\(^{1}\)https://dsc-spidal.github.io/harp/
A. Programming Model Based on Collective Communication

Collective communication within a Rotation design is easy to program. For each iteration, all workers concurrently sample on a local training data partition with a local model split without conflicts in model updates. Afterwards, a call to a collective communication operator ‘rotate’ is made, in order to exchange the model partitions globally. (see Algorithm 1)

Algorithm 1: HarpLDA+ Parallel Pseudo Code

```plaintext
input : training data X, P workers, model A^0, number of iterations T
output: A^T
1 parallel for worker p ∈ [1, P] do
  2 for t = 1 to T do
      // initialize: model A^0 is A^{t-1}
      3 for i = 1 to P do
          // update local model split by sampling on local training data
          4 A_{p^i}^t = Sampling(X_p, A_{p^i}^{t-1})
          // synchronization to exchange model splits
          5 rotate(A_{p^i}^t)
```

A concrete scheduling strategy is built into the ‘rotate’ operator. So long as each model split is owned by only one worker, the scheduling strategy guarantees to be conflict free. For instance, when a rotate call returns, all the workers can continue sampling concurrently without causing conflicts when updating the model. A default strategy shifts the model splits to their neighbor nodes (see Fig. 2a). Selecting the neighbor on a random permutation of the node list is also easy to implement. Furthermore, a priority based scheduler and a work load based scheduler can be implemented in this framework without losing the simplicity of the programming model.

Algorithm 1 presents a general framework for scheduling, where multi-threading and distributed parallelism can adopt the same procedure. We can however improve it to reduce synchronization overhead, leveraging the computation characteristics in these two different environments.

B. Dynamic Scheduling in Shared Memory systems

Dynamic scheduling provides a low cost solution to remove synchronization overhead in the shared memory system. It dynamically assigns workload to idle threads in order to increase the throughput of tasks. This is an effective solution, which is also used in parallel matrix factorization for recommender systems [20].

In Fig. 2b, training data is partitioned into blocks, with the row partition using a random permutation of document id and the column partition using a greedy algorithm based on word frequency. Indexes are constructed during the initialization phase in order to build the map from word id to the related documents appearing in each block. In this case, the minimal unit for scheduling is a block. Furthermore, the number of partitions is larger than the number of threads, which means that there are always ‘free’ rows and columns when one thread finishes its current task. In this example, thread 1 finishes first, then the scheduler selects a new block randomly from the ‘free’ blocks, which are the white blocks in the figure. Because thread 2 and thread 3 are still working, the rows and columns are occupied accordingly as denoted by the gray blocks. In a shared memory system, the scheduler does not move data but instead assigns data addresses of selected free blocks to the idle threads. The wait time of the threads are bounded by the overhead of the scheduler. In the case the number of threads is P and the number of splits is L, so a L×L matrix maintains a two level status: free, or finished. The scheduler can randomly select a free block by scanning the matrix with time complexity of no more than O(L^2). The larger the L, the lower the number of conflicts and lower the wait time, but the more overhead introduced by the scheduler itself. Thus, there is a trade-off. By experimentation, we found that L = √2P is a good choice in most cases.

C. Pipelining and Timer Control in Distributed Systems

In distributed systems, the cost of data movement cannot be ignored. As shown in Fig. 2a, each worker holds a static row partition of the training data and corresponding document related model. Only the word-topic model partitions move among the workers. To reduce the synchronization overhead, the first step is to reduce the overhead of the communication inside the rotate operator. Pipelining is a broadly used technique to solve this kind of problem, by overlapping I/O threads with computing threads. First, each block is split further into two slices horizontally, and the
inner loop of algorithm 1 is modified to become a loop over each slice. Consequently, the original rotate call becomes two rotate calls on each slice. As long as the communication time is less than the computing time spent on one slice, the pipeline will be effective in hiding the overhead from communication.

Another overhead of a rotate call is the time taken waiting for all workers to complete their computation. In the presence of load imbalance, all workers wait for the slowest one to finish. To solve this problem, we first discuss the sampling order of the Gibbs Sampling Algorithms. We note that LDA trainers can use two common scan orders: random scan and deterministic scan. For a Gibbs sampler, the usual deterministic-scan order proceeds by updating first \( x_1 \), then \( x_2 \), then \( x_3, \ldots, x_d \) and back to \( x_1 \), visiting the state space \( X \) by a sequential order. Another random scan version usually proceeds at each iteration by choosing \( i \) uniformly from \( 1, 2, \ldots, d \). It has been demonstrated [21] that the sampling order affects the convergence rate of different models but a deterministic scan is commonly used to gain the benefits of hardware locality. This is the situation in current LDA trainers, in which sampling occurs over document or over word on \( Z \) via deterministic scan. Generally, the order with a better memory cache hit rate gives a better performance. For large datasets with \( V \ll D \), word order is better. It is hard to achieve good performance with a pure random scan due to the cache miss issues. However, HarpLDA+ uses a quasi-random order. The dynamic scheduler picks a block uniformly from the free block list. While inside the block, we still keep the word sampling order. Although no significant performance difference is observed for the different sampling orders in LDA-CGS, we found that the random sampling order provides a natural solution for load balancing.

![Diagram of Timer to Control the Synchronization Point](image)

Figure 3: Timer to Control the Synchronization Point

For a given worker in Fig. 3 the white space between two rotate operations denotes the wait time between the end of the computation and the start of synchronization. If we adjust the synchronization point ahead of the computation finishes, the gap of wait time can be closed. Under the deterministic scan order, the adjustment is harder to achieve due to the housekeeping work needed and the original scan order is lost. For a random scan, this adjustment does not change the property of the uniform random selection of blocks. The third rotate call demonstrates this mechanism in Fig. 3.

We further propose a simple solution for the LDA-CGS trainer. Each sampler only works for the same period of time and then the samplers do synchronization all together. They all use a timer to control the synchronization point rather than waiting until all the blocks to finish. Because the model size shrinks and the computation time drops during the process of convergence, we’ve designed an auto-tuning mechanism to set the value of the timer for each iteration in HarpLDA+.

First, the timer works best when the communication can be fully overlapped by computation, where the computation time or the number of training data points being processed should have a lower bound \( L \). Secondly, we make sure that all workers stop at the same time before any of them complete their work. This implies an upper bound \( H \). \( L \) and \( H \) are set as input parameters. In normal cases, \( L = 40\% \), \( H = 80\% \) are good choices.

We set up heuristic rules to automatically determine the values of timer \( t_i \) based on the \( L, H \) settings.

- Rule 1: During the first iteration, we set the timer to a constant \( t_0 \), and obtain the processing ratio \( R_0 \) for each worker at the end of the iteration.
- Rule 2: When \( R_i \) is found to be smaller than \( L \), adjust \( t_{i+1} = t_i \times 2 \) in order to quickly catch up. (In the first iteration, repeat this step until \( R_{i+1} \) is in the range of \( L \) and \( H \)).
- Rule 3: When \( R_i \) is found to be larger than \( H \), \( t_{i+1} \) will be reduced in half.

D. Other Implementation Issues

For a high performance parallel LDA trainer, besides the key factor of the original sampling algorithm and the parallel system design, implementation details may also be important. HarpLDA+ is a Java application, where primitive data types are used in critical data structures. For instance, we found that using primitive arrays with array indexing for the model matrix is significantly faster than using a hashmap in HarpLDA+. Furthermore, minor improvements for SparseLDA are very helpful. Topic counts are sorted periodically to reduce the linear search time of sampling. Caching is also used to avoid repeat calculations. When sampling multiple tokens with the same word and document, the topic probabilities calculated for the first token are reused for the tokens that follow.

V. EXPERIMENTAL EVALUATION

A. Setup of Experiments

Five datasets are used in the experiments (see Table II), which are open datasets in related work. Here, pubmed2m is a subset of Pubmed Dataset\(^2\), cluweb30b is a subset of

\(^2\)https://archive.ics.uci.edu/ml/datasets/bag+of+words
Figure 4: Single Node Performance with 3 rows corresponding to 3 datasets nytimes ($K = 1000$), pubmed2m ($K = 1000$), enwiki ($K = 10000$) respectively. The columns are different observables with columns 1 to 3 reporting sequential single thread measurements and column 4 with 32 threads. Column 5 plots the results against different thread counts. Each column corresponds to an evaluation metric denoted in the title.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Docs</th>
<th>Vocabulary</th>
<th>Tokens</th>
<th>DocLen</th>
</tr>
</thead>
<tbody>
<tr>
<td>nytimes</td>
<td>299K</td>
<td>101K</td>
<td>99M</td>
<td>332/178</td>
</tr>
<tr>
<td>pubmed2m</td>
<td>2M</td>
<td>126K</td>
<td>149M</td>
<td>74/33</td>
</tr>
<tr>
<td>enwiki</td>
<td>3.7M</td>
<td>1M</td>
<td>1B</td>
<td>293/523</td>
</tr>
<tr>
<td>bigram</td>
<td>3.8M</td>
<td>20M</td>
<td>1.6B</td>
<td>437/767</td>
</tr>
<tr>
<td>clueweb30b</td>
<td>76M</td>
<td>1M</td>
<td>29B</td>
<td>392/532</td>
</tr>
</tbody>
</table>

Table II: Datasets for LDA Training, where DocLen represents mean and std. dev. values of document length

ClueWeb09 Dataset\(^3\), enwiki is built from English articles from Wikipedia and bigram is a bigram version of enwiki dataset.

<table>
<thead>
<tr>
<th>trainer</th>
<th>language</th>
<th>threading</th>
<th>communication</th>
</tr>
</thead>
<tbody>
<tr>
<td>LightLDA</td>
<td>C++</td>
<td>Pthread</td>
<td>Zeromq+MPI</td>
</tr>
<tr>
<td>NomadLDA</td>
<td>C++</td>
<td>Intel TBB</td>
<td>MPI</td>
</tr>
<tr>
<td>WarpLDA</td>
<td>C++</td>
<td>OpenMP</td>
<td>MPI</td>
</tr>
<tr>
<td>HarpLDA+</td>
<td>Java</td>
<td>Java Thread</td>
<td>Harp Collective</td>
</tr>
</tbody>
</table>

Table III: Trainers for Experiments

We select four state-of-the-art CGS trainers for comparison in Table III. They represent the different system designs described in Section III-B. Also different versions of HarpLDA+ are included, e.g., Harp-nods without dynamic scheduling, and Harp-notimer without timer control.

To evaluate the performance, we use the following metrics. Firstly, we choose Model log likelihood of the word-topic model to represent the status of convergence. Secondly, we select three main evaluation metrics as follows: 1) Convergence rate evaluates the effectiveness of the algorithm by depicting the relationship between convergence level and model update count. 2) Throughput evaluates the efficiency in a system view by measuring the model update counts per second. 3) Convergence speed is the metric to evaluate a trainer’s overall performance, which depicts the relation between convergence level and training time (initialization time excluded). It represents the overall performance resulting from the combination of efficiency and effectiveness of model updates.

In regards to hardware configuration, all experiments are conducted on a 128-node Intel Haswell cluster at Indiana University. Among them, 32 nodes each have two 18-core Xeon E5-2699 v3 processors (36 cores in total), and 96 nodes each have two 12-core Xeon E5-2670 v3 processors (24 cores in total). All the nodes have 128 GB memory and are connected by QDR InfiniBand. As for the software configuration, all C++ trainers are compiled with gcc 4.9.2 and -O3 compilation optimization. HarpLDA+ compiles with Java 1.8.0 64 bit Server VM and runs on Hadoop 2.6.0. The MPI runtime is mvapich2 2.3a for F+NomadLDA and mpich2 3.0.4 for LightLDA. For MH trainers, we set the MH step parameter to 1 for WarpLDA, select the best one for LightLDA, 16 for clueweb30b and 4 in the other datasets. We set the hyper-parameters $\alpha = 50/K$ and $\beta = 0.01$ in all the experiments. The setting of experiment runs with $n$...
nodes and \( m \) threads on each node and is denoted as \( n \times m \) in the following diagrams. On a similar note, \( K \) signifies the number of topics used in the LDA trainers.

**B. Experimental Results**

1) **Performance of Sequential Algorithm:** We first analyze the performance of the sampling algorithm by evaluating the trainers in a single thread setup.

As shown in Fig. 4, in column 1 of **Convergence Speed**, WarpLDA is the fastest trainer due to its successful trade-off between the efficiency and effectiveness of updates. Furthermore, F+NomadLDA is faster than HarpLDA+ and demonstrates even better performance than WarpLDA in the case of large \( K \). For this assessment, LightLDA is consistently the slowest.

Column 2 of **Convergence Rate** shows that F+NomadLDA, a standard SparseLDA sampler, always runs the fastest, HarpLDA+ is a bit slower due to the caching of the model of identical words. Both MH samplers, LightLDA and WarpLDA are significantly slower since they are an approximation for the original CGS, while WarpLDA is the slowest because of its update delay strategy making each update much less effective. The rank of convergence rate is consistent in parallel versions of these trainers.

In column 3 of **Throughput**, WarpLDA shows much better throughput than the others due to optimization of memory use obtained from removing the random matrix access. Among the others, F+NomadLDA performs slightly better.

2) **Intra-node Parallel Efficiency:** We run a test by increasing number of threads in order to evaluate its impact on performance. As seen in Fig. 4, column 4, the convergence speed at 32 threads shows that the performance rank of F+NomadLDA drops, and HarpLDA+ takes its place and runs as well as WarpLDA at \( K = 1000 \) and exceeds it’s performance for \( K = 10000 \). In Fig. 4 column 5 of **Speedup in Throughput**, HarpLDA+ demonstrates the best parallel efficiency which explains the boost of its performance from a single thread to a large number of threads.

**Concurrence Analysis by VTune Amplifier**

In order to further breakdown the actual working time, we add thread level logs to record the actual sampling time in each iteration. See Fig. 5a, F+NomadLDA has a very large CV level depicting serious problems with load imbalance. Fig. 5b, F+NomadLDA again shows a high overhead ratio. LightLDA is better but still larger than 10%. In contrast, HarpLDA+ presents a relatively large initial overhead, which diminishes over time. This is due to a fixed timer being set to 1 second in the first iteration, while constant overheads...

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<table>
<thead>
<tr>
<th>Trainer</th>
<th>Effective CPU Time</th>
<th>Spin Overhead</th>
<th>Wait Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>WarpLDA</td>
<td>0.91</td>
<td>0</td>
<td>0.09</td>
</tr>
<tr>
<td>NomadLDA</td>
<td>0.75</td>
<td>0.24</td>
<td>0</td>
</tr>
<tr>
<td>LightLDA</td>
<td>0.25</td>
<td>0</td>
<td>0.74</td>
</tr>
<tr>
<td>HarpLDA+</td>
<td>0.98</td>
<td>0</td>
<td>0.02</td>
</tr>
<tr>
<td>Harp-nods</td>
<td>0.75</td>
<td>0</td>
<td>0.24</td>
</tr>
</tbody>
</table>

Table IV: Time Breakdown by VTune Concurrency Analysis. **Effective Time** is CPU time spent in the user code, **Spin time** is wait time during which the CPU is busy, and **Overhead time** is CPU time spent on the overhead of known synchronization and threading libraries. **Wait Time** occurs when software threads are waiting due to APIs that block or cause synchronization. The enwiki dataset is used in experiments (\( K = 1000 \)) and runs on 32 threads of a single node.

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Figure 6: Distributed Performance. Columns 1 to 4, enwiki ($K = 10000$) with 8x16(nodes×threads) in row 1, clueweb30b ($K = 5000$) with 40×16 in row 2. Column 5 shows plots speedup versus node count with 16 threads each. The initial word-topic model size is 2.6GB for enwiki and 12.7GB for clueweb30b.

of hundreds of milliseconds make the overhead ratio appear high. As seen in the charts, HarpLDA+ demonstrates the best load balance and a small overhead. WarpLDA is excluded in this experiment because it is implemented with OpenMP and thread log cannot be added.

3) Distributed Parallel Efficiency: In this section, we test the LDA trainers in distributed mode. WarpLDA is not included because the official source code release does not support distributed mode. Moreover, we expect that the distributed design presented in its paper might not scale well because of the need to exchange a much larger model $Z$ in each iteration among all the workers. F+NomadLDA runs on an InfiniBand network directly supported by mpich2, but lightlda runs on IPoIB (TCP/IP protocol on InfiniBand network) supported by mpich2, and as a Java application, HarpLDA+ runs on IPoIB too. This means F+NomadLDA can potentially utilize a bandwidth which is at least two times larger in these experiments and is easier to scale.

As shown in Fig. 6, column 1 represents convergence speed, where HarpLDA+ has the best overall performance. In column 2 of Convergence Time Ratio, we calculate the speedup in time of other trainers with respect to HarpLDA+, defined as the ratio of the training time to reach a given Model Log-Likelihood which is the abscissa of the graph.

HarpLDA+ is more than 6x faster than LightLDA, 2x faster than NomadLDA and about 50% slower when timer control is not used. In columns 3 and 4, Load Balance of Computation and Overhead, HarpLDA+ demonstrates significant differences from Harp-notimer, which is the factor behind the boost in performance.

LightLDA, as an asynchronous approach, has less problems of load imbalance than the synchronized approaches. The default staleness is set to one which can tolerate any performance undulations only if the lag of the local model replica is less than two iterations. This mechanism is effective in order to provide stability and good scalability for different cluster configurations. This is showcased in column 5. On the other hand, LightLDA has a lower convergence rate stemmed from its asynchronous design and is less efficient during the multi-threading parallel implementation.

F+NomadLDA is observed to have the most load imbalance problems and also exhibits a very high overhead ratio. In the enwiki 10K experiment, the overhead even reaches 90%, i.e., most of the workers are waiting for data. When using a rotation architecture and running directly on the InfiniBand network, this result is not expected. One possible reason for this is the task granularity. It takes very small granularity to schedule on each column of the word-topic model, which seems to have a large overhead, especially when $K$ increases to a large number.

4) Communication Intensive Case Study: The following experiment runs on the bigram dataset, which has a 20 million vocabulary size that is used to test the special communication intensive case in the distributed mode. We set the parameter of the bound in HarpLDA+ to [150%, 350%] to overlap the communication time in this special case, i.e., the dynamic scheduler keeps assigning those sampled blocks to free threads until the timer is timeout. This trainer is named Harp-repeat.

Fig. 7d shows that when the communication time dominates in the training process, all the trainers have a large overhead ratio. In LightLDA, as SSP forces the workers to keep the staleness of the local model within a range, the problem of the wait time comes back. Harp-repeat significantly decreases the overhead and increases the throughput, but at the same time, the effectiveness drops in 7b, to be worse than LightLDA. Hence, harp-repeat does not gain in overall performance. In contrast, Harp-notimer retains the best overall performance. Further optimization should focus on how to decrease the size of the model that needs to be
exchanged.

5) Straggler Case Study: The notion of a straggler is a situation in cloud computations, where some nodes are significantly slower than others in a job for different reasons. In our experiment on the HPC cluster, we also encounter stragglers more often than expected.

For Convergence Speed in Fig. 9b, when $K$ increases to 1 million, LightLDA runs much faster than HarpLDA+ due to time complexity of $O(1)$ in the MH sampling algorithm. However this only happens at the beginning of the training phase. Afterwards it slows down and is surpassed by HarpLDA+ because of its ineffectiveness of computation, despite using a very large MH step parameter such as 128 in Fig. 9. In this big model experiment, HarpLDA+ demonstrates impressive performance, given that the algorithm has a time complexity of $O(K_d + K_w)$ while that of LightLDA is $O(1)$.

VI. CONCLUSIONS AND FUTURE WORK

In this paper, we investigate the system design of large scale LDA trainers with a focus on parallel efficiency. Based on these, we introduce HarpLDA+, selecting the Rotation architecture and proposing a new synchronized LDA training system with reduced overhead. This entails a two level parallelism design, in which a dynamic scheduler is used for multi-threading, while rotation with timer control is used for distributed parallelism. Through extensive experiments, we demonstrate that the HarpLDA+ outperforms the other approaches in scaling and stability. We note that all 4 training systems being evaluated are compared on the identical Haswell cluster with the same configuration.

From HarpLDA+, we’ve gained useful insights in designing a large scale Machine Learning system.

- Optimization of a sequential algorithm does not necessarily lead to high performance parallel systems. Implementation details including programming languages and high performance off-the-shelf communication libraries do not always guarantee good performance as
seen in Tables I and III. The choices of data structures and parallel system design are critical for good performance in our Java HarpLDA+, as shown in Figures 4 and 6.

- Asynchronous parallel designs are favorable for scalability and robustness. However, with increasing parallelism and computation capacity provided by manycore and GPU servers, synchronized parallel designs can achieve better performance on a moderate sized cluster for big data problems in Table II.

Incorporating more parallelism, such as vectorization, into LDA trainers can be further explored in future work. Also, the similarity between LDA-CGS trainer and MF-SGD trainer implies some intrinsic relationship between these two large families of Machine Learning algorithms, which are potential future directions to explore.

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