XGBoost: Optimization Technique and a Scalable Tree Boosting System for Machine Learning Algorithms

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ABSTRACT

Tree boosting is a highly e ective and widely used machine learning method. In this paper, we describe a scalable end-to-end tree boosting system called XGBoost, which is used widely by data scientists to achieve state-of-the-art results on many machine learning challenges. We propose a novel sparsityaware algorithm for sparse data and weighted quan-tile sketch for approximate tree learning. More importantly, we provide insights on cache access patterns, data compres-sion and sharding to build a scalable tree boosting system. By combining these insights, XGBoost scales beyond billions of examples using far fewer resources than existing systems.

Keywords

Large-scale Machine Learning

INTRODUCTION

Machine learning and data-driven approaches are becom-ing very important in many areas. Smart spam classi ers protect our email by learning from massive amounts of s-pam data and user feedback; advertising systems learn to match the right ads with the right context; fraud detection systems protect banks from malicious attackers; anomaly event detection systems help experimental physicists to nd events that lead to new physics. There are two importan-t factors that drive these successful applications: usage of e ective (statistical) models that capture the complex data dependencies and scalable learning systems that learn the model of interest from large datasets.

Among the machine learning methods used in practice, gradient tree boosting [10]¹ is one technique that shines in many applications. Tree boosting has been shown to give state-of-theart results on many standard classi cation benchmarks [16]. LambdaMART [5], a variant of tree boost-ing for ranking, achieves state-of-the-art result for ranking problems.

¹ Gradient tree boosting is also known as gradient boosting machine (GBM) or gradient boosted regression tree (GBRT)

Besides being used as a stand-alone predictor, it is also incorporated into real-world production pipelines for ad click through rate prediction [15]. Finally, it is the de-facto choice of ensemble method and is used in challenges such as the Netix prize [3].

In this paper, we describe XGBoost, a scalable machine learning system for tree boosting. The system is available as an open source package². The impact of the system has been widely recognized in a number of machine learning and data mining challenges. Take the challenges hosted by the machine learning competition site Kaggle for example. A-mong the 29 challenge winning solutions ³ published at Kag-gle's blog during 2015, 17 solutions used XGBoost. Among these solutions, eight solely used XGBoost to train the mod-el, while most others combined XGBoost with neural net-s in ensembles. For comparison, the second most popular method, deep neural nets, was used in 11 solutions. The success of the system was also witnessed in KDDCup 2015, where XGBoost was used by every winning team

10. Moreover, the winning teams reported that ensemble methods outperform a well-con gured XGBoost by only a small

These results demonstrate that our system gives state-of-the-art results on a wide range of problems. Examples of the problems in these winning solutions include: store sales prediction; high energy physics event classi cation; web text classi cation; customer behavior prediction; motion detec-tion; ad click through rate prediction; malware classi cation; product categorization; hazard risk prediction; massive on-line course dropout rate prediction. While domain depen-dent data analysis and feature engineering play an important role in these solutions, the fact that XGBoost is the consen-sus choice of learner shows the impact and importance of our system and tree boosting.

The most important factor behind the success of XGBoost is its scalability in all scenarios. The system runs more than ten times faster than existing popular solutions on a single machine and scales to billions of examples in distributed or memory-limited settings. The scalability of XGBoost is due to several important systems and algorithmic optimizations. These innovations include: a novel tree learning algorithm is for handling sparse data; a theoretically justi ed weighted quantile sketch procedure enables handling instance weights in approximate tree learning. Parallel and distributed com-puting makes learning faster which enables quicker model ex-ploration. More importantly, XGBoost exploits out-of-core

2https://github.com/dmlc/xgboost

³ Solutions come from of top-3 teams of each competitions.

computation and enables data scientists to process hundred millions of examples on a desktop. Finally, it is even more exciting to combine these techniques to make an end-to-end system that scales to even larger data with the least amount of cluster resources. The major contributions of this paper is listed as follows:

We design and build a highly scalable end-to-end tree boosting system.

We propose a theoretically justi ed weighted quantile sketch for e cient proposal calculation.

We introduce a novel sparsity-aware algorithm for par-allel tree learning.

We propose an e ective cache-aware block structure for outof-core tree learning.

While there are some existing works on parallel tree boost-ing [22, 23, 19], the directions such as out-of-core compu-tation, cache-aware and sparsity-aware learning have not been explored. More importantly, an end-to-end system that combines all of these aspects gives a novel solution for real-world use-cases. This enables data scientists as well as researchers to build powerful variants of tree boosting al-gorithms [7, 8]. Besides these major contributions, we also make additional improvements in proposing a regularized learning objective, which we will include for completeness.

The remainder of the paper is organized as follows. We will rst review tree boosting and introduce a regularized objective in Sec. 2. We then describe the split nding meth-ods in Sec. 3 as well as the system design in Sec. 4, including experimental results when relevant to provide quantitative support for each optimization we describe. Related work is discussed in Sec. 5. Detailed end-to-end evaluations are included in Sec. 6. Finally we conclude the paper in Sec. 7.

TREE BOOSTING IN A NUTSHELL 2.

We review gradient tree boosting algorithms in this sec-tion. The derivation follows from the same idea in existing literatures in gradient boosting. Specicially the second order method is originated from Friedman et al. [12]. We make mi-nor improvements in the reguralized objective, which were found helpful in practice.

2.1 Regularized Learning Objective

For a given data set with n examples and m features

 $D = f(x_i; y_i)g$ $(jDj = n; x_i 2 R^m; y_i 2 R)$, a tree ensem-ble model (shown in Fig. 1) uses K additive functions to predict the output.

$$\begin{array}{ll}
K \\
x_k \\
y^*_i = (x_i) = f_k(x_i); f_k 2 F; \\
= 1 \\
= 1
\end{array}$$
(1)

where $F = ff(x) = w_{q(x)}g(q:R^m ! T; w 2 R^T)$ is the space of regression trees (also known as CART). Here q rep-resents the structure of each tree that maps an example to the corresponding leaf index. T is the number of leaves in the tree. Each fk corresponds to an independent tree structure q and leaf weights w. Unlike decision trees, each regression tree contains a continuous score on each of the leaf, we use wi to represent score on i-th leaf. For a given example, we will use the decision rules in the trees (given by q) to classify

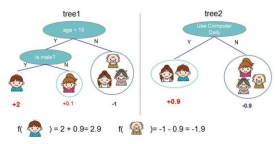


Figure 1: Tree Ensemble Model. The nal predic-tion for a given example is the sum of predictions from each tree.

it into the leaves and calculate the nal prediction by sum-ming up the score in the corresponding leaves (given by w). To learn the set of functions used in the model, we minimize the following regularized objective.

$$L(\) = \begin{array}{c} X & X \\ l(\)_{i} & X \\ i & k \end{array} \tag{2}$$
 where
$$(f) = T + \frac{1}{2} \ kwk^{2}$$

Here l is a di erentiable convex loss function that measures the di erence between the prediction y¹ and the target y_i. The second term penalizes the complexity of the model (i.e., the regression tree functions). The additional regular-ization term helps to smooth the nal learnt weights to avoid over- tting. Intuitively, the regularized objective will tend to select a model employing simple and predictive functions. A similar regularization technique has been used in Regu-larized greedy forest (RGF) [25] model. Our objective and the corresponding learning algorithm is simpler than RGF and easier to parallelize. When the regularization parameter is set to zero, the objective falls back to the traditional gradient tree boosting.

2.2 Gradient Tree Boosting

The tree ensemble model in Eq. (2) includes functions as parameters and cannot be optimized using traditional optimization methods in Euclidean space. Instead, the model is trained in an additive manner. Formally, let $y^{\land}_{i}^{(t)}$ be the prediction of the i-th instance at the t-th iteration, we will need to add ft to minimize the following objective.

$$L^{(t)} = \int_{-1}^{n} (y_i; y^{n})^{(t1)} + f_t(x_i) + (f_t)$$

This means we greedily add the ft that most improves our model according to Eq. (2). Second-order approximation can be used to quickly optimize the objective in the general setting [12].

where $g_i = @_{V^{\Lambda(t)}} l(y_i; y^{\Lambda(t)})$ and $h_i = @_{V^{\Lambda(t)}} l(y_i; y^{\Lambda(t)})$ are rst and second order gradient statistics on the loss function. We can remove the constant terms to obtain the fol-lowing simpli ed objective at step t.

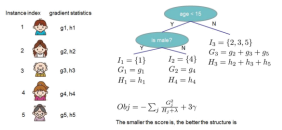


Figure 2: Structure Score Calculation. We only need to sum up the gradient and second order gra-dient statistics on each leaf, then apply the scoring formula to get the quality score.

De ne $I_j = fijq(x_i) = jg$ as the instance set of leaf j. We can rewrite Eq (3) by expanding as follows

For a xed structure q(x), we can compute the optimal weight w_i of leaf j by

$$P_{i2I_{j}\,h_{i}\,+}; \frac{P}{i^{2}I_{j}\,g_{i}\,w_{j}} = 0 \tag{5}$$

and calculate the corresponding optimal value b

Eq (6) can be used as a scoring function to measure the quality of a tree structure q. This score is like the impurity score for evaluating decision trees, except that it is derived for a wider range of objective functions. Fig. 2 illustrates how this score can be calculated.

Normally it is impossible to enumerate all the possible tree structures q. A greedy algorithm that starts from a single leaf and iteratively adds branches to the tree is used instead. Assume that II and IR are the instance sets of left and right nodes after the split. Lettting I = I_L [I_R, then the loss reduction after the split is given

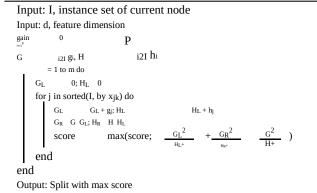
This formula is usually used in practice for evaluating the

2.3 Shrinkage and Column Subsampling

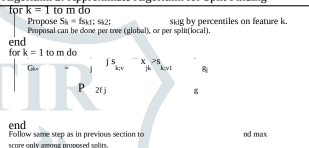
Besides the regularized objective mentioned in Sec. 2.1, two additional techniques are used to further prevent over-tting. The rst technique is shrinkage introduced by Fried-man [11]. Shrinkage scales newly added weights by a factor

after each step of tree boosting. Similar to a learning rate in tochastic optimization, shrinkage reduces the inuence of each individual free and leaves space for future trees to im-prove the model. The second technique is column (feature) subsampling. This technique is used in RandomForest [4,

Algorithm 1: Exact Greedy Algorithm for Split Finding



Algorithm 2: Approximate Algorithm for Split Finding



13], It is implemented in a commercial software TreeNet 4

for gradient boosting, but is not implemented in existing opensource packages. According to user feedback, using col-umn sub-sampling prevents over-tting even more so than the traditional row sub-sampling (which is also supported). The usage of column sub-samples also speeds up computations of the parallel algorithm described later.

SPLIT FINDING ALGORITHMS

Basic Exact Greedy Algorithm

One of the key problems in tree learning is to nd the best split as indicated by Eq (7). In order to do so, a s-plit nding algorithm enumerates over all the possible splits on all the features. We call this the exact greedy algorithm. Most existing single machine tree boosting implementation-s, such as scikit-learn [20], R's gbm [21] as well as the single machine version of XGBoost support the exact greedy algo-rithm. The exact greedy algorithm is shown in Alg. 1. It is computationally demanding to enumerate all the possible splits for continuous features. In order to do so e ciently, the algorithm must rst sort the data according to feature values and visit the data in sorted order to accumulate the gradient statistics for the structure score in Eq (7).

3.2 Approximate Algorithm

The exact greedy algorithm is very powerful since it enumerates over all possible splitting points greedily. However, it is impossible to e ciently do so when the data does not t entirely into memory. Same problem also arises in the dis-

⁴https://www.salford-systems.com/products/treenet

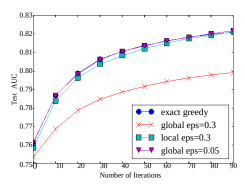


Figure 3: Comparison of test AUC convergence on Higgs 10M dataset. The eps parameter corresponds to the accuracy of the approximate sketch. This roughly translates to 1 / eps buckets in the proposal. We nd that local proposals require fewer buckets, because it re ne split candidates.

tributed setting. To support e ective gradient tree boosting in these two settings, an approximate algorithm is needed.

We summarize an approximate framework, which resem-bles the ideas proposed in past literatures [17, 2, 22], in Alg. 2. To summarize, the algorithm rst proposes candi-date splitting points according to percentiles of feature dis-tribution (a speci c criteria will be given in Sec. 3.3). The algorithm then maps the continuous features into bucket-s split by these candidate points, aggregates the statistics and nds the best solution among proposals based on the aggregated statistics.

There are two variants of the algorithm, depending on when the proposal is given. The global variant proposes all the candidate splits during the initial phase of tree construction, and uses the same proposals for split ending at all levels.

The local variant re-proposes after each split. The global method requires less proposal steps than the local method. However, usually more candidate points are needed for the global proposal because candidates are not re ned after each split. The local proposal re nes the candidates after splits, and can potentially be more appropriate for deeper trees. A comparison of di erent algorithms on a Higgs boson dataset is given by Fig. 3. We nd that the local proposal indeed requires fewer candidates. The global proposal can be as accurate as the local one given enough candidates.

Most existing approximate algorithms for distributed tree learning also follow this framework. Notably, it is also possi-ble to directly construct approximate histograms of gradient statistics [22]. It is also possible to use other variants of bin-ning strategies instead of quantile [17]. Quantile strategy bene t from being distributable and recomputable, which we will detail in next subsection. From Fig. 3, we also nd that the quantile strategy can get the same accuracy as exact greedy given reasonable approximation level.

Our system e ciently supports exact greedy for the single machine setting, as well as approximate algorithm with both local and global proposal methods for all settings. Users can freely choose between the methods according to their needs.

3.3 Weighted Quantile Sketch

One important step in the approximate algorithm is to propose candidate split points. Usually percentiles of a fea-ture are used to make candidates distribute evenly on the da-

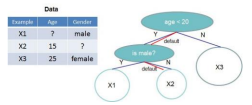


Figure 4: Tree structure with default directions. An example will be classi ed into the default direction when the feature needed for the split is missing.

ta. Formally, let multi-set $D_k = f(x_{1k}; h_1); (x_{2k}; h_2) (x_{nk}; h_n)g$ represent the k-th feature values and second order gradient statistics of each training instances. We can de ne a rank functions $r_k : R ! [0; +1)$ as

$$r_k(z) = \frac{1}{(x;h)2D_k} X h;$$
 (8)

which represents the proportion of instances whose feature value k is smaller than z. The goal is to nd candidate split points fsk1; sk2; sklg, such that

Here is an approximation factor. Intuitively, this means that there is roughly 1= candidate points. Here each data point is weighted by hi. To see why hi represents the weight, we can rewrite Eq (3) as

$$\begin{array}{ccc} n & \\ x & \top_2 \; h_i (f_t(x_i) \; \; g_i \!\!=\!\! h_i)^2 + \; (f_t) + constant; \\ i \!\!=\!\! 1 & \end{array}$$

which is exactly weighted squared loss with labels gi=hi and weights h_i. For large datasets, it is non-trivial to nd can-didate splits that satisfy the criteria. When every instance has equal weights, an existing algorithm called quantile s-ketch [14, 24] solves the problem. However, there is no existing quantile sketch for the weighted datasets. There-fore, most existing approximate algorithms either resorted to sorting on a random subset of data which have a chance of failure or heuristics that do not have theoretical guarantee.

To solve this problem, we introduced a novel distributed weighted quantile sketch algorithm that can handle weighted data with a provable theoretical guarantee. The general idea is to propose a data structure that supports merge and prune operations, with each operation proven to maintain a certain accuracy level. A detailed description of the algorithm as well as proofs are given in the supplementary material⁵ (link in the footnote).

Sparsity-aware Split Finding

In many real-world problems, it is quite common for the input x to be sparse. There are multiple possible causes for sparsity: 1) presence of missing values in the data; 2) frequent zero entries in the statistics; and, 3) artifacts of feature engineering such as onehot encoding. It is impor-tant to make the algorithm aware of the sparsity pattern in the data. In order to do so, we propose to add a default direction in each tree node, which is shown in Fig. 4. When a value is missing in the sparse matrix x, the instance is

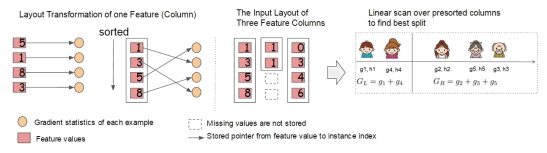


Figure 6: Block structure for parallel learning. Each column in a block is sorted by the corresponding feature value. A linear scan over one column in the block is su cient to enumerate all the split points.

Algorithm 3: Sparsity-aware Split Finding Input: I, instance set of current node Input: $I_k = fi \ 2 \ Ijx_{ik} \ 6 = missingg$ Input: d, feature dimension Also applies to the approximate setting, only collect statistics of non-missing entries into buckets gain 0 for k P G i2I ; g_i,H i2I hi = 1 to m do // enumerate missing value goto right GL 0; HL 0 for j in sorted(I_k , ascent order by x_{jk}) do $H_{L} + h_{i}$ G_L $G_L + g_i$; H_L G_{R} G GL; HR H HL score max(score: end // enumerate missing value goto left GR 0; HR 0 for j in sorted(I_k , descent order by x_{ik}) do G_R $G_R + g_j$; H_R $H_R + h_i$ G_L G GR; HL H HR end end Output: Split and default directions with max gain

classi ed into the default direction. There are two choices of default direction in each branch. The optimal default directions are learnt from the data. The algorithm is shown in Alg. 3. The key improvement is to only visit the non-missing entries I_k . The presented algorithm treats the non-presence as a missing value and learns the best direction to handle missing values. The same algorithm can also be applied when the non-presence corresponds to a user speci ed value by limiting the enumeration only to consistent solutions.

To the best of our knowledge, most existing tree learning algorithms are either only optimized for dense data, or need speci c procedures to handle limited cases such as categor-ical encoding. XGBoost handles all sparsity patterns in a uni ed way. More importantly, our method exploits the s-parsity to make computation complexity linear to number of non-missing entries in the input. Fig. 5 shows the com-parison of sparsity aware and a naive implementation on an Allstate-10K dataset (description of dataset given in Sec. 6). We nd that the sparsity aware algorithm runs 50 times faster than the naive version. This con rms the importance of the sparsity aware algorithm.

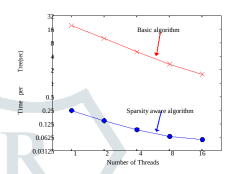


Figure 5: Impact of the sparsity aware algorithm on Allstate-10K. The dataset is sparse mainly due to one-hot encoding. The sparsity aware algorithm is more than 50 times faster than the naive version that does not take sparsity into consideration.

4. SYSTEM DESIGN

4.1 Column Block for Parallel Learning

The most time consuming part of tree learning is to get the data into sorted order. In order to reduce the cost of sorting, we propose to store the data in in-memory units, which we called block. Data in each block is stored in the compressed column (CSC) format, with each column sorted by the corresponding feature value. This input data layout only needs to be computed once before training, and can be reused in later iterations.

In the exact greedy algorithm, we store the entire dataset in a single block and run the split search algorithm by lin-early scanning over the pre-sorted entries. We do the split nding of all leaves collectively, so one scan over the block will collect the statistics of the split candidates in all leaf branches. Fig. 6 shows how we transform a dataset into the format and nd the optimal split using the block structure.

The block structure also helps when using the approximate algorithms. Multiple blocks can be used in this case, with each block corresponding to subset of rows in the dataset. Di erent blocks can be distributed across machines, or s-tored on disk in the out-of-core setting. Using the sorted structure, the quantile nding step becomes a linear scan over the sorted columns. This is especially valuable for lo-cal proposal algorithms, where candidates are generated fre-quently at each branch. The binary search in histogram ag-gregation also becomes a linear time merge style algorithm. Collecting statistics for each column can be parallelized, giving us a parallel algorithm for split nding. Importantly, the column block structure also supports column subsam-pling, as it is easy to select a subset of columns in a block.

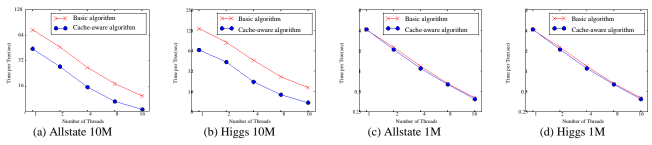


Figure 7: Impact of cache-aware prefetching in exact greedy algorithm. We nd that the cache-miss e ect impacts the performance on the large datasets (10 million instances). Using cache aware prefetching improves the performance by factor of two when the dataset is large.

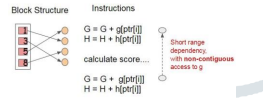


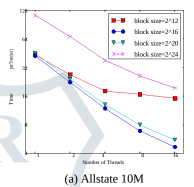
Figure 8: Short range data dependency pattern that can cause stall due to cache miss.

Time Complexity Analysis Let d be the maximum depth of the tree and K be total number of trees. For the exac-t greedy algorithm, the time complexity of original spase aware algorithm is O(Kdkxk₀ log n). Here we use kxk₀ to denote number of nonmissing entries in the training data. On the other hand, tree boosting on the block structure on-ly cost O(Kdkxk₀ + kxk₀ log n). Here O(kxk₀ log n) is the one time preprocessing cost that can be amortized. This analysis shows that the block structure helps to save an ad-ditional log n factor, which is signi cant when n is large. For the approximate algorithm, the time complexity of original algorithm with binary search is O(Kdkxk₀ log q). Here q is the number of proposal candidates in the dataset. While q is usually between 32 and 100, the log factor still introduces overhead. Using the block structure, we can reduce the time to $O(Kdkxk_0 + kxk_0 \log B)$, where B is the maximum num-ber of rows in each block. Again we can save the additional log q factor in computation.

4.2 Cache-aware Access

While the proposed block structure helps optimize the computation complexity of split nding, the new algorithm requires indirect fetches of gradient statistics by row index, since these values are accessed in order of feature. This is a noncontinuous memory access. A naive implementation of split enumeration introduces immediate read/write de-pendency between the accumulation and the non-continuous memory fetch operation (see Fig. 8). This slows down split nding when the gradient statistics do not t into CPU cache and cache miss occur.

For the exact greedy algorithm, we can alleviate the prob-lem by a cache-aware prefetching algorithm. Speci cally, we allocate an internal bu er in each thread, fetch the gra-dient statistics into it, and then perform accumulation in a mini-batch manner. This prefetching changes the direct read/write dependency to a longer dependency and helps to reduce the runtime overhead when number of rows in the is large. Figure 7 gives the comparison of cache-aware vs.



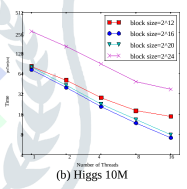


Figure 9: The impact of block size in the approxi-mate algorithm. We nd that overly small blocks re-sults in ine cient parallelization, while overly large blocks also slows down training due to cache misses.

non cache-aware algorithm on the the Higgs and the All-state dataset. We nd that cache-aware implementation of the exact greedy algorithm runs twice as fast as the naive version when the dataset is large.

For approximate algorithms, we solve the problem by choos-ing a correct block size. We de ne the block size to be max-imum number of examples in contained in a block, as this reects the cache storage cost of gradient statistics. Choos-ing an overly small block size results in small workload for each thread and leads to ine cient parallelization. On the other hand, overly large blocks result in cache misses, as the gradient statistics do not t into the CPU cache. A good choice of block size balances these two factors. We compared various choices of block size on two data sets. The results are given in Fig. 9. This result validates our discussion and

System parallel approximate out-of-core global greedy local aware **XGBoost** yes yes yes yes pGBRT no no yes no no yes Spark MLLib no yes no no partially yes H2O no no no partially yes yes scikit-learn yes no no no no no R GBM ves no no no partially no

Table 1: Comparison of major tree boosting systems

shows that choosing 2¹⁶ examples per block balances the cache property and parallelization.

4.3 Blocks for Out-of-core Computation

One goal of our system is to fully utilize a machine's resources to achieve scalable learning. Besides processors and memory, it is important to utilize disk space to handle data that does not t into main memory. To enable out-of-core computation, we divide the data into multiple blocks and store each block on disk. During computation, it is important to use an independent thread to pre-fetch the block into a main memory bu er, so computation can happen in con-currence with disk reading. However, this does not entirely solve the problem since the disk reading takes most of the computation time. It is important to reduce the overhead and increase the throughput of disk IO. We mainly use two techniques to improve the out-of-core computation.

Block Compression The rst technique we use is block compression. The block is compressed by columns, and decompressed on the y by an independent thread when load-ing into main memory. This helps to trade some of the computation in decompression with the disk reading cost. We use a general purpose compression algorithm for com-pressing the features values. For the row index, we substract the row index by the begining index of the block and use a 16bit integer to store each o set. This requires 2¹⁶ examples per block, which is con rmed to be a good setting. In most of the dataset we tested, we achieve roughly a 26% to 29% compression ratio.

Block Sharding The second technique is to shard the data onto multiple disks in an alternative manner. A pre-fetcher thread is assigned to each disk and fetches the data into an in-memory bu er. The training thread then alternatively reads the data from each bu er. This helps to increase the throughput of disk reading when multiple disks are available.

RELATED WORKS

Our system implements gradient boosting [10], which performs additive optimization in functional space. Gradient tree boosting has been successfully used in classi cation [12], learning to rank [5], structured prediction [8] as well as other elds. XGBoost incorporates a regularized model to prevent over tting. This this resembles previous work on regularized greedy forest [25], but simpli es the objective and algorithm for parallelization. Column sampling is a simple but e ective technique borrowed from RandomForest [4]. While sparsity-aware learning is essential in other types of models such as linear models [9], few works on tree learning have considered this topic in a principled way. The algorithm proposed in this paper is the rst uni ed approach to handle all kinds of sparsity patterns.

There are several existing works on parallelizing tree learn-ing [22, 19]. Most of these algorithms fall into the approxi-mate framework described in this paper. Notably, it is also possible to partition data by columns [23] and apply the ex-act greedy algorithm. This is also supported in our frame-work, and the techniques such as cache-aware pre-fecthing can be used to bene t this type of algorithm. While most existing works focus on the algorithmic aspect of paralleliza-tion, our work improves in two unexplored system direction-

s: out-of-core computation and cache-aware learning. This gives us insights on how the system and the algorithm can be jointly optimized and provides an end-to-end system that can handle large scale problems with very limited computing resources. We also summarize the comparison between our system and existing opensource implementations in Table 1.

Quantile summary (without weights) is a classical prob-lem in the database community [14, 24]. However, the ap-proximate tree boosting algorithm reveals a more general problem { nding quantiles on weighted data. To the best of our knowledge, the weighted quantile sketch proposed in this paper is the rst method to solve this problem. The weighted quantile summary is also not speci c to the tree learning and can bene t other applications in data science and machine learning in the future.

END TO END EVALUATIONS

System Implementation 6.1

We implemented XGBoost as an open source package⁶. The package is portable and reusable. It supports various weighted classi cation and rank objective functions, as well as user de ned objective function. It is available in popular languages such as python, R, Julia and integrates naturally with language native data science pipelines such as scikit-learn. The distributed version is built on top of the rabit library for allreduce. The portability of XGBoost makes it available in many ecosystems, instead of only being tied to a speci c platform. The distributed XGBoost runs natively on Hadoop, MPI Sun Grid engine. Recently, we also enable distributed XGBoost on jvm bigdata stacks such as Flink and Spark. The distributed version has also been integrated into cloud platform Tianchi⁸ of Alibaba. We believe that there will be more integrations in the future.

6.2 Dataset and Setup

We used four datasets in our experiments. A summary of these datasets is given in Table 2.

Table 2: Dataset used in the Experiments.

Dataset	n	m	Task
Allstate	10 M	4227	Insurance claim classi cation
Higgs Boson	10 M	28	Event classi cation
Yahoo LTRC	473K	700	Learning to Rank
Criteo	1.7 B	67	Click through rate prediction

ments, we use a randomly selected subset of the data either due to slow baselines or to demonstrate the performance of the algorithm with varying dataset size. We use a su x to denote the size in these cases. For example Allstate-10K means a subset of the Allstate dataset with 10K instances.

The rst dataset we use is the Allstate insurance claim dataset⁹. The task is to predict the likelihood and cost of an insurance claim given di erent risk factors. In the exper-iment, we simpli ed the task to only predict the likelihood of an insurance claim. This dataset is used to evaluate the impact of sparsity-aware algorithm in Sec. 3.4. Most of the sparse features in this data come from one-hot encoding. We randomly select 10M instances as training set and use the rest as evaluation set.

The second dataset is the Higgs boson dataset10 from high energy physics. The data was produced using Monte Carlo simulations of physics events. It contains 21 kinematic prop-erties measured by the particle detectors in the accelerator. It also contains seven additional derived physics quantities of the particles. The task is to classify whether an event corresponds to the Higgs boson. We randomly select 10M instances as training set and use the rest as evaluation set.

The third dataset is the Yahoo! learning to rank challenge dataset [6], which is one of the most commonly used bench-marks in learning to rank algorithms. The dataset contains 20K web search queries, with each query corresponding to a list of around 22 documents. The task is to rank the documents according to relevance of the query. We use the o cial train test split in our experiment.

The last dataset is the criteo terabyte click log dataset¹¹. We use this dataset to evaluate the scaling property of the system in the out-of-core and the distributed settings. The data contains 13 integer features and 26 ID features of user, item and advertiser information. Since a tree based model is better at handling continuous features, we preprocess the data by calculating the statistics of average CTR and count of ID features on the rst ten days, replacing the ID fea-tures by the corresponding count statistics during the next ten days for training. The training set after preprocessing contains 1.7 billion instances with 67 features (13 integer, 26 average CTR statistics and 26 counts). The entire dataset is more than one terabyte in LibSVM format.

We use the rst three datasets for the single machine par-allel setting, and the last dataset for the distributed and out-of-core settings. All the single machine experiments are conducted on a Dell PowerEdge R420 with two eight-core Intel Xeon (E5-2470) (2.3GHz) and 64GB of memory. If not speci ed, all the experiments are run using all the available cores in the machine. The machine settings of the distribut-ed and the out-of-core experiments will be described in the

Table 3: Comparison of Exact Greedy Methods with 500 trees on Higgs-1M data.

Method	Time per Tree (sec)	Test AUC
XGBoost	0.6841	0.8304
XGBoost (colsample=0.5)	0.6401	0.8245
scikit-learn	28.51	0.8302
R.gbm	1.032	0.6224

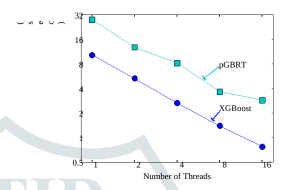


Figure 10: Comparison between XGBoost and pG-BRT on Yahoo LTRC dataset.

Table 4: Comparison of Learning to Rank with 500 trees on Yahoo! LTRC Dataset

Method	Time per Tree (sec)	NDCG@10
XGBoost	0.826	0.7892
XGBoost (colsample=0.5)	0.506	0.7913
pGBRT [22]	2.576	0.7915

corresponding section. In all the experiments, we boost trees with a common setting of maximum depth equals 8, shrink-age equals 0.1 and no column subsampling unless explicitly speci ed. We can nd similar results when we use other settings of maximum depth.

Classification

In this section, we evaluate the performance of XGBoost on a single machine using the exact greedy algorithm on Higgs-1M data, by comparing it against two other common-ly used exact greedy tree boosting implementations. Since scikit-learn only handles non-sparse input, we choose the dense Higgs dataset for a fair comparison. We use the 1M subset to make scikit-learn nish running in reasonable time. Among the methods in comparison, R's GBM uses a greedy approach that only expands one branch of a tree, which makes it faster but can result in lower accuracy, while both scikit-learn and XGBoost learn a full tree. The results are shown in Table 3. Both XGBoost and scikit-learn give bet-ter performance than R's GBM, while XGBoost runs more than 10x faster than scikit-learn. In this experiment, we al-so nd column subsamples gives slightly worse performance than using all the features. This could due to the fact that there are few important features in this dataset and we can bene t from greedily select from all the features.

6.4 Learning to Rank

We next evaluate the performance of XGBoost on the learning to rank problem. We compare against pGBRT [22], the best previously pubished system on this task. XGBoost

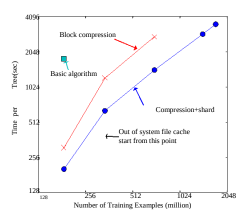


Figure 11: Comparison of out-of-core methods on di erent subsets of criteo data. The missing data points are due to out of disk space. We can nd that basic algorithm can only handle 200M exam-ples. Adding compression gives 3x speedup, and sharding into two disks gives another 2x speedup. The system runs out of le cache start from 400M examples. The algorithm really has to rely on disk after this point. The compression+shard method has a less dramatic slowdown when running out of le cache, and exhibits a linear trend afterwards.

runs exact greedy algorithm, while pGBRT only support an approximate algorithm. The results are shown in Table 4 and Fig. 10. We nd that XGBoost runs faster. Interest-ingly, subsampling columns not only reduces running time, and but also gives a bit higher performance for this prob-lem. This could due to the fact that the subsampling helps prevent over tting, which is observed by many of the users.

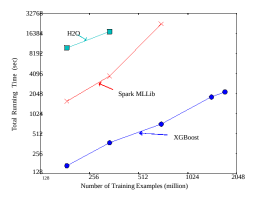
6.5 Out-of-core Experiment

We also evaluate our system in the out-of-core setting on the criteo data. We conducted the experiment on one AWS c3.8xlarge machine (32 vcores, two 320 GB SSD, 60 GB RAM). The results are shown in Figure 11. We can nd that compression helps to speed up computation by factor of three, and sharding into two disks further gives 2x speedup. For this type of experiment, it is important to use a very large dataset to drain the system le cache for a real out-of-core setting. This is indeed our setup. We can observe a transition point when the system runs out of le cache. Note that the transition in the nal method is less dramatic. This is due to larger disk throughput and better utilization of computation resources. Our nal method is able to process 1.7 billion examples on a single machine.

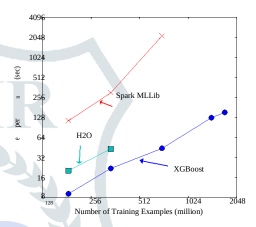
6.6 Distributed Experiment

Finally, we evaluate the system in the distributed setting. We set up a YARN cluster on EC2 with m3.2xlarge ma-chines, which is a very common choice for clusters. Each machine contains 8 virtual cores, 30GB of RAM and two 80GB SSD local disks. The dataset is stored on AWS S3 instead of HDFS to avoid purchasing persistent storage.

We rst compare our system against two production-level distributed systems: Spark MLLib [18] and H2O ¹². We use 32 m3.2xlarge machines and test the performance of the sys-



(a) End-to-end time cost include data loading



(b) Per iteration cost exclude data loading

Figure 12: Comparison of di erent distributed sys-tems on 32 EC2 nodes for 10 iterations on di erent subset of criteo data. XGBoost runs more 10x than spark per iteration and 2.2x as H2O's optimized ver-sion (However, H2O is slow in loading the data, get-ting worse end-to-end time). Note that spark su ers from drastic slow down when running out of mem-ory. XGBoost runs faster and scales smoothly to the full 1.7 billion examples with given resources by utilizing out-of-core computation items with various input size.

Both of the baseline systems are in-memory analytics frameworks that need to store the data in RAM, while XGBoost can switch to out-of-core set-ting when it runs out of memory. The results are shown in Fig. 12. We can nd that XGBoost runs faster than the baseline systems. More importantly, it is able to take advantage of out-of-core computing and smoothly scale to all 1.7 billion examples with the given limited computing re-sources. The baseline systems are only able to handle subset of the data with the given resources. This experiment shows the advantage to bring all the system improvement togeth-er and solve a real-world scale problem. We also evaluate the scaling property of XGBoost by varying the number of machines.

The results are shown in Fig. 13. We can nd XGBoost's performance scales linearly as we add more machines. Importantly, XGBoost is able to handle the entire 1.7 billion data with only four machines. This shows the system's potential to handle even larger data.

¹² www.h2o.ai

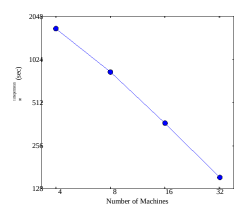


Figure 13: Scaling of XGBoost with di erent num-ber of machines on criteo full 1.7 billion dataset. Using more machines results in more le cache and makes the system run faster, causing the trend to be slightly super linear. XGBoost can process the entire dataset using as little as four machines, and s-cales smoothly by utilizing more available resources.

CONCLUSION

In this paper, we described the lessons we learnt when building XGBoost, a scalable tree boosting system that is widely used by data scientists and provides state-of-the-art results on many problems. We proposed a novel sparsity aware algorithm for handling sparse data and a theoretically justi ed weighted quantile sketch for approximate learning. Our experience shows that cache access patterns, data com-pression and sharding are essential elements for building a scalable end-to-end system for tree boosting. These lessons can be applied to other machine learning systems as well. By combining these insights, XGBoost is able to solve real-world scale problems using a minimal amount of resources.

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