Out-of-Core GPU Gradient Boosting

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ABSTRACT
GPU-based algorithms have greatly accelerated many machine learning methods; however, GPU memory is typically smaller than main memory, limiting the size of training data. In this paper, we describe an out-of-core GPU gradient boosting algorithm implemented in the XGBoost library. We show that much larger datasets can fit on a given GPU, without degrading model accuracy or training time. To the best of our knowledge, this is the first out-of-core GPU implementation of gradient boosting. Similar approaches can be applied to other machine learning algorithms.

CCS CONCEPTS
• Computing methodologies → Machine learning; Graphics processors; • Information systems → Hierarchical storage management.

KEYWORDS
GPU, out-of-core algorithms, gradient boosting, machine learning

1 INTRODUCTION
Gradient boosting [7] is a popular machine learning method for supervised learning tasks, such as classification, regression, and ranking. A prediction model is built sequentially out of a ensemble of weak prediction models, typically decision trees. With bigger datasets and deeper trees, training time can become substantial.

Graphics Processing Units (GPUs), originally designed to speed up the rendering of display images, have proven to be powerful accelerators for many parallel computing tasks, including machine learning. GPU-based implementations [4, 6, 15] exist for several open-source gradient boosting libraries [3, 10, 14] that significantly lower the training time.

Because GPU memory has higher bandwidth and lower latency, it tends to cost more and thus is typically of smaller size than main memory. For example, on Amazon Web Services (AWS), a p3.2xlarge instance has 1 NVIDIA Tesla V100 GPU with 16 GiB memory, and 61 GiB main memory. On Google Cloud Platform (GCP), a similar instance can have as much as 78 GiB main memory. Training with large datasets can cause GPU out-of-memory errors when there is plenty of main memory available.

XGBoost, a widely-used gradient boosting library, has experimental support for external memory [5], which allows training on datasets that do not fit in main memory 1. Building on top of this feature, we designed and implemented out-of-core GPU algorithms that extend XGBoost external memory support to GPUs. This is challenging since GPUs are typically connected to the rest of the computer system through the PCI Express (PCIe) bus, which has lower bandwidth and higher latency than the main memory bus. A naive approach that constantly swaps data in and out of GPU memory would cause too much slowdown, negating the performance gain from GPUs.

By carefully structuring the data access patterns, and leveraging gradient-based sampling to reduce working memory size, we were able to significantly increase the size of training data accommodated by a given GPU, with minimal impact to model accuracy and training time.

2 BACKGROUND
In this section we review the gradient boosting algorithm as implemented by XGBoost, its GPU variant, and the previous CPU-only external memory support. We also describe the sampling approaches used to reduce memory footprint.

2.1 Gradient Boosting
Given a dataset with $n$ samples $\{(x_i, y_i)\}_{i=1}^n$, where $x_i \in \mathbb{R}^m$ is a vector of $m$-dimensional input features, and $y_i \in \mathbb{R}$ is the label, a decision tree model predicts the label:

$$\hat{y}_i = F(x_i) = \sum_{k=1}^{K} f_k(x_i),$$

where $f_k \in F$, the space of regression trees, and $K$ is the number of trees. To learn a model, we minimize the following regularized objective:

$$\mathcal{L}(F) = \sum_{i} l(\hat{y}_i, y_i) + \sum_{k} \Omega(f_k)$$

where $\Omega(f) = yT + \frac{1}{2}||w||^2$.

Here $I$ is a differentiable loss function, $\Omega$ is the regularization term that penalizes the number of leaves in the tree $T$ and leaf weights $w$, controlled by two hyperparameters $\gamma$ and $\lambda$.

The model is trained sequentially. Let $\hat{y}_i^{(t)}$ be the prediction at the $t$-th iteration, we need to find tree $f_t$ that minimizes:

$$\mathcal{L}^{(t)} = \sum_{i} l(y_i, \hat{y}_i^{(t-1)} + f_t(x_i)) + \Omega(f_t)$$

The quadratic Taylor expansion is:

$$\mathcal{L}^{(t)} = \sum_{i} [l(y_i, \hat{y}_i^{(t-1)}) + g_i f_t(x_i) + \frac{1}{2} h_i f_t^2(x_i)] + \Omega(f_t),$$

where $g_i$ and $h_i$ are first and second order gradients on the loss function with respect to $\hat{y}_i^{(t-1)}$. For a given tree structure $q(x)$, let $I_j = \{i | q(x_i) = j\}$ be the set of samples that fall into leaf $j$. The optimal weight $w_j^*$ of leaf $j$ can be computed as:

$$w_j^* = \frac{\sum_{i \in I_j} g_i}{\sum_{i \in I_j} h_i + \lambda}.$$
and the corresponding optimal loss value is:

$$\hat{L}^{(t)}(q) = \frac{1}{2} \sum_{j=2}^{T} \left( \frac{\sum_{i \in I_j} g_i^2}{\sum_{i \in I_j} h_i + \lambda} + y^T \right).$$  

(7)

When constructing an individual tree, we start from a single leaf and greedily add branches to the tree. Let $I_L$ and $I_R$ be the sets of samples that fall into the left and right nodes after a split, then the loss reduction for a split is:

$$L_{\text{split}} = \frac{1}{2} \left( \frac{\sum_{i \in I_L} g_i^2}{\sum_{i \in I_L} h_i + \lambda} + \frac{\sum_{i \in I_R} g_i^2}{\sum_{i \in I_R} h_i + \lambda} - \frac{\sum_{i \in I} g_i^2}{\sum_{i \in I} h_i + \lambda} \right) - \gamma$$  

(8)

where $I = I_L \cup I_R$.

### 2.2 GPU Tree Construction

The GPU tree construction algorithm in XGBoost [11, 12] relies on a two-step process. First, in a preprocessing step, each input feature is divided into quantiles and put into bins (max_bin defaults to 256). The bin numbers are then compressed into ELLPACK format, greatly reducing the size of the training data. This step is time consuming, but it should only be done once at the beginning of training.

#### Algorithm 1: GPU Tree Construction

Input: $X$: training examples
Output: $\text{tree}$: set of output nodes

```
Input: $g$: gradient pairs for training examples
Output: $\text{tree}$: set of output nodes
```

```
tree $\leftarrow \{\}$
queue $\leftarrow \text{InitRoot()}$

while queue is not empty do

    entry $\leftarrow$ queue.pop()
    tree.insert(entry)

    // Sort samples into leaf nodes
    RepartitionInstances(entry, $X$)

    // Build gradient histograms
    BuildHistograms(entry, $X$, $g$)

    // Find the optimal split for children
    left_entry $\leftarrow \text{EvaluateSplit(entry.left.histogram)}$
    right_entry $\leftarrow \text{EvaluateSplit(entry.right.histogram)}$
    queue.push(left_entry)
    queue.push(right_entry)
```

In the second step, the tree construction algorithm is shown in Algorithm 1. Note that this is a simplified version for single GPU only. In a distributed environment with multiple GPUs, the gradient histograms need to be summed across all GPUs using AllReduce.

### 2.3 XGBoost Out-of-Core Computation

XGBoost has experimental support for out-of-core computation [3, 5]. When enabled, training is also done in a two-step process. First, in the preprocessing step, input data is read and parsed into an internal format, which can be Compressed Sparse Row (CSR), Compressed Sparse Column (CSC), or sorted CSC. Each sample is appended to an in-memory buffer. When the buffer reaches a predefined size (32 MiB), it is written out to disk as a page. Second, during tree construction, the data pages are streamed from disk via a multi-threaded pre-fetcher.

### 2.4 Sampling

In its default setting, gradient boosting is a batch algorithm: the whole dataset needs to be read and processed to construct each tree. Different sampling approaches have been proposed, mainly as an additional regularization factor to get better generalization performance, but they can also reduce the computation needed, leading to faster training time.

#### 2.4.1 Stochastic Gradient Boosting (SGB)

Shortly after introducing gradient boosting, Friedman [8] proposed an improvement: at each iteration a subsample of the training data is drawn at random without replacement from the full training dataset. This randomly selected subsample is then used in place of the full sample to construct the decision tree and compute the model update for the current iteration. It was shown that this sampling approach improves model accuracy. However, the sampling ratio, $f$, needs to stay relatively high, $0.5 \leq f \leq 0.8$, for this improvement to occur.

#### 2.4.2 Gradient-based One-Side Sampling (GOSS)

Ke et al. proposed a sampling strategy weighted by the absolute value of the gradients [10]. At the beginning of each iteration, the top $a \times 100\%$ of training instances with the largest gradients are selected, then from the rest of the data a random sample of $b \times 100\%$ instances is drawn. The samples are scaled by $\frac{1}{\sqrt{\text{dataset-size}}}$ to make the gradient statistics unbiased. Compared to SGB, GOSS can sample more aggressively, only using 10% - 20% of the data to achieve similar model accuracy.

#### 2.4.3 Minimal Variance Sampling (MVS)

Ibragimov et al. proposed another gradient-based sampling approach that aims to minimize the variance of the model. At each iteration the whole dataset is sampled with probability proportional to regularized absolute value of gradients:

$$\hat{g}_i = \sqrt{\gamma_i^2 + \lambda h_i^2},$$

(9)

where $g_i$ and $h_i$ are the first and second order gradients, $\lambda$ can be either a hyperparameter, or estimated from the squared mean of the initial leaf value.

MVS was shown to perform better than both SGB and GOSS, with sampling rate as low as 10%.

### 3 Method

In this section we describe the design of out-of-core GPU-based gradient boosting. Since XGBoost is widely used in production, as much as possible, we try to preserve the existing behavior when adding new features. In external memory mode, we assume the training data is already parsed and written to disk in CSR pages.

#### 3.1 Incremental Quantile Generation

As stated above, GPU tree construction in XGBoost is a two-step process. In the preprocessing step, input features are converted into a quantile representation. Quantiles are cut points dividing the range of each feature into continuous intervals (i.e. bins) with equal probabilities. Algorithm 2 shows the in-core version of quantile sketch.
Algorithm 2: In-Core Quantile Sketch
Input: \( X \): training examples
Output: \( \text{histogram\_cuts} \): cut points for all features
\[
\text{foreach batch in } X \text{ (a single CSR page) do}
\quad \text{CopyToGPU(batch)}
\]
\[
\text{foreach column in batch do}
\quad \text{cuts} \leftarrow \text{FindColumnCuts(batch, column)}
\quad \text{CopyColumnCuts(histogram\_cuts, cuts)}
\]

Algorithm 3: Out-of-Core Quantile Sketch
Input: \( X \): training examples
Output: \( \text{histogram\_cuts} \): cut points for all features
\[
\text{foreach page in } X \text{ do}
\quad \text{foreach batch in page do}
\quad \quad \text{CopyToGPU(batch)}
\quad \quad \text{foreach column in batch do}
\quad \quad \quad \text{cuts} \leftarrow \text{FindColumnCuts(batch, column)}
\quad \quad \quad \text{CopyColumnCuts(histogram\_cuts, cuts)}
\]

Since the existing code already operates in batches and handles the necessary bookkeeping, it is straightforward to extend it to external memory mode with multiple CSR pages, as shown in Algorithm 3.

3.2 External ELLPACK Matrix

Algorithm 4: In-Core ELLPACK Page
Input: \( X \): training examples
Input: \( \text{histogram\_cuts} \): cut points for all features
Output: \( \text{ellpack\_page} \): compressed ELLPACK matrix
\[
\text{AllocateOnGPU(ellpack\_page)}
\quad \text{foreach batch in } X \text{ (a single CSR page) do}
\quad \quad \text{CopyToGPU(batch)}
\quad \quad \text{foreach row in batch do}
\quad \quad \quad \text{foreach column in row do}
\quad \quad \quad \quad \text{bin} \leftarrow \text{LookupBin(histogram\_cuts, column)}
\quad \quad \quad \text{Write(ellpack\_page, bin)}
\]

Once the quantile cut points are found, input features can be converted to bin numbers and compressed into ELLPACK format, as shown in Algorithm 4.

In external memory mode, we assume the single ELLPACK matrix may not fit in GPU memory, thus is broken up into multiple ELLPACK pages and written to disk. Since CSR pages contain a variable number of rows, we cannot pre-allocate these ELLPACK pages. Instead, the CSR pages are accumulated in memory first. When the expected ELLPACK page reaches the size limit, the CSR pages are converted and written to disk, as shown in Algorithm 5.

Algorithm 5: Out-of-Core ELLPACK Pages
Input: \( X \): training examples
Output: \( \text{ellpack\_pages} \): compressed ELLPACK matrix pages
\[
\text{list} \leftarrow \{\}
\quad \text{foreach page in } X \text{ do}
\quad \quad \text{list.append(page)}
\quad \text{if CalculateEllpackPageSize(list) } \geq 32 \text{ MiB then}
\quad \quad \text{AllocateOnGPU(ellpack\_page)}
\quad \quad \text{foreach page in list do}
\quad \quad \quad \text{Write(ellpack\_page, page)}
\quad \quad \text{WriteToDisk(ellpack\_page)}
\quad \text{list} \leftarrow \{\}
\]

// Convert list to ELLPACK and write to disk...

Algorithm 6: Naive Out-of-Core GPU Tree Construction
Input: \( X \): training examples
Input: \( g \): gradient pairs for training examples
Output: \( \text{tree} \): set of output nodes
\[
\text{tree} \leftarrow \{\}
\quad \text{// Loop through all the pages}
\quad \text{queue} \leftarrow \text{InitRoot()}
\quad \text{while queue is not empty do}
\quad \quad \text{entry} \leftarrow \text{queue.pop()}
\quad \quad \text{tree.insert(entry)}
\quad \quad \text{foreach page in } X \text{ do}
\quad \quad \quad \text{// Sort samples into leaf nodes}
\quad \quad \quad \text{RepartitionInstances(entry, page)}
\quad \quad \quad \text{// Build gradient histograms}
\quad \quad \quad \text{BuildHistograms(entry, page, g)}
\quad \quad \text{// Find the optimal split for children}
\quad \quad \quad \text{left\_entry} \leftarrow \text{EvaluateSplit(entry.left\_histogram)}
\quad \quad \quad \text{right\_entry} \leftarrow \text{EvaluateSplit(entry.right\_histogram)}
\quad \quad \quad \text{queue.push(left\_entry)}
\quad \quad \quad \text{queue.push(right\_entry)}
\]

shown in Algorithm 6. However, because of the PCIe bottleneck, this approach performed badly, even slower than the CPU tree construction algorithm.

3.4 Use Sampled Data
To improve the training performance, we implemented gradient-based sampling using MVS. For each iteration, we first sample the gradient pairs. Then the multiple ELLPACK pages are compacted together into a single page, only keeping the rows with non-zero gradients. Algorithm 7 shows this approach.

4 RESULTS
We measured the effectiveness of out-of-core GPU gradient boosting from several dimensions: data size, model accuracy, and training time.
When not sampling the data, the out-of-core GPU algorithm is equivalent to the in-core version. With sampling, the size of the data that can fit on a given GPU is increased. Ideally, this should not change the generalization performance of the trained model. Figure 1 shows the training curves on the Higgs dataset [1]. Models with different sampling rates performed similarly, only dropped slightly when \( f = 0.1 \).

For a more detailed evaluation of MVS, see [9].

### 4.2 Model Accuracy

When not sampling the data, the out-of-core GPU algorithm is equivalent to the in-core version. With sampling, the size of the data that can fit on a given GPU is increased. Ideally, this should not change the generalization performance of the trained model. Figure 1 shows the training curves on the Higgs dataset [1]. Models with different sampling rates performed similarly, only dropped slightly when \( f = 0.1 \).

For a more detailed evaluation of MVS, see [9].

### 4.3 Training Time

For end-to-end training time, the Higgs dataset is used, split randomly 0.95/0.05 for training and evaluation. All the XGBoost parameters use their default value, except that max_depth is increased to 8, and learning_rate is lowered to 0.1. Training is done for 500 iterations. The hardware used is a desktop computer with an Intel Core i7-5820K processor, 32 GB main memory, and an NVIDIA Titan V with 12 GB memory. Table 2 shows the training time and evaluation AUC for the different modes.

Although out-of-core GPU training is slower than the in-core version when sampling is enabled, it is still significantly faster than the CPU-based algorithm.

## 5 DISCUSSION

Faced with the explosive growth of data, GPU proved to be an excellent choice to speed up machine learning tasks. However, the relative small size of GPU memory puts a constraint on how much data can be handled on a single GPU. To train on larger datasets, distributed algorithms can be used to share the workload on multiple machines with multiple GPUs. Setting up and managing a distributed GPU cluster is expensive, both in terms of hardware and networking cost and system administration overhead. It is therefore desirable to relax the GPU memory constraint on a single machine, to allow for easier experimentation with larger datasets.

Because of the PCIe bottleneck, GPU out-of-core computation remains a challenge. A naive implementation that simply spills data over to main memory or disk would likely to be too slow to be useful. If the out-of-core GPU algorithm is slower than the CPU version, then what is the point? Only by pursuing algorithmic changes, as we have done with gradient-based sampling here, can out-of-core GPU computation become competitive. The sampling approach may be applicable to other machine learning algorithms. This is left as possible future work.

Working with XGBoost also presented unique software engineering challenges. It is a popular open source project with many contributors, ranging from students, data scientists, to machine learning software engineers. Code quality varies between different parts of the code base. In order to support the existing users, many of which run XGBoost in production, care must be taken to preserve the current behavior, and plan for breaking changes carefully. Much of the effort during this project was spent on refactoring the code to make it easier to add new behaviors.

## 6 CONCLUSION

In this paper we presented the first ever out-of-core GPU gradient boosting implementation. This approach greatly expands the size of training data that can fit on a given GPU, without sacrificing model accuracy or training time. The source code changes are merged into the open-source XGBoost library. It is available for production use and further research.

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![Figure 1: Training curves on Higgs dataset](image-url)

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