ZNN – A Fast and Scalable Algorithm for Training 3D Convolutional Networks on Multi-Core and Many-Core Shared Memory Machines

Aleksandar Zlateski∗, Kisuk Lee†

∗Electrical Engineering and Computer Science Dept.
†Brain and Cognitive Sciences Dept.
Massachusetts Institute of Technology
Cambridge, MA 02139 USA
∗zlateski@mit.edu, †kisuklee@mit.edu

H. Sebastian Seung
Princeton Neuroscience Institute
and Computer Science Dept.
Princeton University
Princeton, NJ 08540 USA
sseung@princeton.edu

Abstract—Convolutional networks (ConvNets) have become a popular approach to computer vision. It is important to accelerate ConvNet training, which is computationally costly. We propose a novel parallel algorithm based on decomposition into a set of tasks, most of which are convolutions or FFTs. Applying Brent’s theorem to the task dependency graph implies that linear speedup with the number of processors is attainable within the PRAM model of parallel computation, for wide network architectures. To attain such performance on real shared-memory machines, our algorithm computes convolutions converging on the same node of the network with temporal locality to reduce cache misses, and sums the convergent convolution outputs via an almost wait-free concurrent method to reduce time spent in critical sections. We implement the algorithm with a publicly available software package called ZNN. Benchmarking with multi-core CPUs shows that ZNN can attain speedup roughly equal to the number of physical cores. We also show that ZNN can attain over 90x speedup on a many-core CPU (Xeon Phi TM Knights Corner). These speedups are achieved for network architectures with widths that are in common use. The task parallelism of the ZNN algorithm is suited to CPUs, while the SIMD parallelism of previous algorithms is compatible with GPUs. Through examples, we show that ZNN can be either faster or slower than certain GPU implementations depending on specifics of the network architecture, kernel sizes, and density and size of the output patch. ZNN may be less costly to develop and maintain, due to the relative ease of general-purpose CPU programming.

I. INTRODUCTION

A standard formulation of supervised learning starts with a parametrized class of mappings, a training set of desired input-output pairs, and a loss function measuring deviation of actual output from desired output. The goal of learning is to minimize the average loss over the training set. A popular minimization method is stochastic gradient descent. For each input in sequence, the parameters of the mapping are updated in minus the direction of the gradient of the loss with respect to the parameters. Here we are concerned with a class of mappings known as convolutional networks (ConvNets).

Significant effort has been put into parallelizing ConvNet learning on GPUs, as in the popular software packages Caffe [1], Torch [2] and Theano [3]. ConvNet learning has also been distributed over multiple machines [4]. However, there has been relatively little work on parallelizing ConvNet learning for single shared memory CPU machines.

Here we introduce a software package called ZNN, which implements a novel parallel algorithm for ConvNet learning on multi-core and many-core CPU machines. ZNN implements 3D ConvNets, with 2D as a special case. ZNN can employ either direct or FFT convolution, and chooses between the two methods by autotuning each layer of the network. FFT convolution was previously applied to 2D ConvNets running on GPUs [5], [6], and is even more advantageous for 3D ConvNets on CPUs.

As far as we know, ZNN is the first publicly available software that supports efficient training of sliding window max-pooling ConvNets, which have been studied by [7], [8], [9].

There is related work on using Xeon Phi TM for supervised deep learning [10], and unsupervised deep learning [11].

II. COMPUTATION GRAPH

We define a ConvNet using a directed acyclic graph (DAG), called the computation graph (Fig. 1). Each node represents a 3D image, and each edge some image filtering operation. (2D images are a special case in which one of the dimensions has size one.) If multiple edges converge on a node, the node sums the outputs of the filtering operations represented by the edges.
The set of weights of the linear combination is called the kernel. If the input image has size $n^3$ and the kernel has size $k^3$, then the output image has size $n^3 = (n-k+1)^3$. Image size decreases because an output voxel only exists when the sliding window is fully contained in the input image.\footnote{This is known as a \textit{valid} convolution in MATLAB.} The convolution is allowed to be sparse, meaning that only every $s$th image voxel (in every dimension) within the sliding window enters the linear combination.

\textbf{Max-pooling} divides an image of size $n^3$ into blocks of size $p^3$, where $n$ is divisible by $p$. The maximum value is computed for each block, yielding an image of size $(n/p)^3$.

\textbf{Max-filtering} The maximum within a sliding window is computed for each location of the window in the image. For a window of size $k^3$ and an input image of size $n^3$, the output image has size $(n-k+1)^3$. 3D max-filtering can be performed by sequential 1D max-filtering of $n^2$ arrays in each of the three directions. For each array we keep a heap of size $k$ containing the values inside the 1D sliding window. Each element of the array will be inserted and removed at most once, each operation taking $\log k$. For each position of the sliding window the top of the heap will contain the maximum value.

\textbf{Transfer function} adds a number called the bias to each voxel of the image and then applies a nonlinear function to the result. The nonlinear function is typically nondecreasing. Common choices are the logistic function, the hyperbolic tangent and half-wave rectification.

The computational complexities of max-pooling, max-filtering, and transfer function are shown in Table I.

For ConvNets in common use, the computation graph has the following properties:

- All convergent edges are convolutions; if a node has a sole incoming edge, the edge represents a nonlinear filtering operation.
- Nodes with convergent edges are not adjacent in the graph, but are separated from each other by nonlinear filtering edges. This is a reasonable constraint, because a composition of two convolutions can be collapsed into a single convolution, thereby simplifying the graph.
- The graph has a layered organization in which all edges in a layer represent operations of the same type.

$\begin{array}{|c|c|c|c|}
\hline
\text{Pass} & \text{Pooling} & \text{Filtering} & \text{Transfer function} \\
\hline
\text{Forward} & f \cdot n^3 & f \cdot 6n^3 \log k & f \cdot n^3 \\
\text{Backward} & f \cdot n^3 & f \cdot n^3 & f \cdot n^3 \\
\text{Update} & - & - & f \cdot n^3 \\
\hline
\end{array}$

\textbf{TABLE I:} Number of floating point operations (FLOPs) required by a layer with $f$ nodes that all perform the same nonlinear filtering operation (max-pooling, max-filtering, or transfer function).

For convenience, the discussion below will assume that images and kernels have isotropic dimensions, though this restriction is not necessary for ZNN. The image filtering operations are of the four following types.

\textbf{Convolution} A weighted linear combination of voxels within a sliding window is computed for each location of the window in the image. The convolution is allowed to be sparse, meaning that only every $s$th image voxel (in every dimension) within the sliding window enters the linear combination.

\textbf{Max-pooling} divides an image of size $n^3$ into blocks of size $p^3$, where $n$ is divisible by $p$. The maximum value is computed for each block, yielding an image of size $(n/p)^3$.

\textbf{Max-filtering} The maximum within a sliding window is computed for each location of the window in the image. For a window of size $k^3$ and an input image of size $n^3$, the output image has size $(n-k+1)^3$. 3D max-filtering can be performed by sequential 1D max-filtering of $n^2$ arrays in each of the three directions. For each array we keep a heap of size $k$ containing the values inside the 1D sliding window. Each element of the array will be inserted and removed at most once, each operation taking $\log k$. For each position of the sliding window the top of the heap will contain the maximum value.

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- The graph has a layered organization in which all edges in a layer represent operations of the same type.

ZNN works for general computation graphs, whether or not they possess the above properties.

\section{Backpropagation Learning}

The backpropagation algorithm is a way of calculating the gradient of the loss function with respect to the trainable parameters in a ConvNet, the kernels and biases. For each input, the calculation proceeds in several phases:

1. Obtain an input and desired output from the training set.
2. \textit{Forward pass} - compute the actual output of the ConvNet from the input.
3) Compute the gradient of the loss function with respect to the actual output.
4) **Backward pass** - Compute the gradient of the loss function with respect to the voxels of the output image at each node.
5) **Weight update** - Compute the gradient of the loss function with respect to the kernels and biases, and update these parameters in the direction of minus the gradient.

The forward pass has already been described above. ZNN implements several possibilities for the loss function, such as the Euclidean distance between the actual and desired outputs.

### A. Backward pass

It turns out that the backward pass can be represented by another graph that looks the same as the forward computation graph, except that the direction of every edge is reversed. The output nodes of the forward graph become the input nodes of the backward graph, and are initialized with the gradient of the loss function with respect to the voxels of the output nodes of the forward graph. The nodes of the backward graph are associated with their own images, which are distinct from the ones associated with the nodes of the forward graph.

Every edge in the backward graph is multiplication by the transpose of the Jacobian matrix of the operation represented by the corresponding edge in the forward computation graph. The four edge operations in the forward graph become the following four edge operations in the backward graph.

**Convolution Jacobian** Convolution in the forward pass becomes convolution in the backward pass. The kernel is the same, except that it is reflected along all three dimensions. If the input image has size \( n^3 \) and the kernel has size \( k^3 \), then the output image has size \( n'^3 = (n + k - 1)^3 \). Image size increases because an output voxel exists whenever the sliding window has some overlap with the input image.\(^2\)

**Max-pooling Jacobian** Within each block, all voxels are zeroed out except for the one that was identified as the maximum within that block in the forward pass. An image of size \( n'^3 \) is expanded into an image of size \( n^3 p^3 \).

**Max-filtering Jacobian** Every element of an image of size \( n'^3 = (n + p - 1)^3 \) is initialized to zero. For each position of the sliding window the appropriate value of the input is accumulated to the position from which the maximum element was selected for that window in the forward pass.

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**Transfer function Jacobian** Every voxel of a backward image is multiplied by the derivative of the transfer function for the corresponding voxel in the forward image.

### B. Weight update

After the forward and backward passes are complete, there are “forward images” at the nodes of the forward computation graph, and “backward images” at the nodes of the backward computation graph (except the input nodes). These are used to update the kernels and biases as follows.

**Kernel update** For a convolution going from node \( a \) to node \( b \) in the forward graph, the gradient of the loss with respect to the kernel is computed by convolving the reflected forward image at node \( a \) with the backward image at node \( b \). A valid convolution is performed, yielding an image the same size as the kernel. This is multiplied by a small “learning rate parameter,” and then subtracted from the kernel.

**Bias update** For a bias at node \( a \), the gradient of the loss is calculated as the sum of all voxels in the backward image at node \( a \). The scalar result is multiplied by a small “learning rate parameter,” and then subtracted from the bias.

### IV. Direct vs. FFT Convolution

For a single convolution of an image of size \( n^3 \) with a kernel of size \( k^3 \), it is well-known that the FFT method (complexity \( \mathcal{O}(n^3 \log n) \)), becomes more efficient than the direct method (complexity \( \mathcal{O}(n^3 k^3) \)), for sufficiently large kernel sizes. The crossover point of equal complexity satisfies \( k^3 \sim \log n \). It is less well-known that the FFT-direct crossover occurs at smaller kernel sizes for a ConvNet than for a single convolution. This is because the FFT of an image at a node can be shared by edges at that node (see Table II). ZNN performs layerwise auto-tuning to choose between FFT-based or direct convolution for each layer.

Complexity can be further reduced by memoizing the FFTs of images and kernels obtained during the forward pass for reuse during the backward pass and weight update. This possibility was previously noted in passing but not implemented due to limited onboard GPU memory. The reduction in complexity is approximately a third (Table II).

### V. Task Dependency Graph

The entire gradient learning calculation can be represented by the task dependency graph (Fig. 3). Each node represents one of the four forward operations (convolution, max-pooling/filtering, transfer function), four backward operations (Jacobians), or two update operations (kernel, bias) described above. Two additional tasks interface with the training set. The data provider obtains a training sample used for a single round of training, and the loss gradient calculates the gradient of the loss with respect to the network output.

The edges of the task dependency graph represent dependencies. The forward task of an edge \( e = (u, v) \) in the computation graph depends on forward pass tasks of all edges \( u \).

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\(^2\)This is known as a full convolution in MATLAB.
layered architectures in which every convolutional layer is able speedup,” because it depends on the idealized assumptions. We will refer to the right hand side as the “theoretically achievable speedup.” This amounts to a speedup of at least

We will estimate the theoretically achievable speedup for layered architectures in which every convolutional layer is fully connected. As before, time complexity is measured in number of floating point instructions. We can already estimate $T_1$ by summing the times in Tables I and II for each layer of the network. To estimate $T_\infty$, we analyze the following algorithm employing an infinite number of processors. (1) Move sequentially through the layers, and perform all forward tasks in each layer in parallel. (2) Compute the loss gradient for all output nodes in parallel. (3) Move sequentially backward through the layers, and perform all backward tasks in each layer in parallel. (4) Perform the weight updates for all kernels and biases in parallel.

Since the layers are done sequentially, the total time for the forward pass is the sum of contributions from each layer (convolutional, transfer function, or max pooling/softplus) as specified in Tables III and IV. The time for the backward pass is calculated similarly. Since all kernel and bias updates are done in parallel, the total update time is the maximum of the individual update times, as specified in Tables III and IV. The sum of forward, backward, and update times yields the time complexity of one gradient learning iteration.

Most of the formulas in the tables do not depend on the particular value of $\epsilon$. This is because any error in a layer is also dependent on the error in the layer that feeds it. This behavior is consistent with Eq. (2). We expect $S_\infty$ to approach a fixed fraction of $S_\infty$ when $S_\infty \approx P$. Since $S_\infty$ scales like $f^2$ (neglecting the logarithmic factor due to $T_\infty$), this should happen when $f^2 \approx P$. The power of two means that the theoretically achievable speedup approaches its maximum value even for networks with rather modest widths.

### Table II: Computational complexity of a fully connected convolutional layer, which maps $f$ input images to $f'$ output images using $f$ kernels. FFT complexity for an $n \times n \times n$ image is assumed to be $Cn^3 \log n^3$. Complexity is measured in number of floating point operations.

| Pass | Direct | FFT-based | FFT-based (Memoized) |
|------|--------|-----------|----------------------|
| Forward | $f \cdot f \cdot n^3 \cdot k^3$ | $3Cn^3 \log n [f' + f + f' \cdot f] + 4f' \cdot f \cdot n^3$ | $3Cn^3 \log n [f' + f + f' \cdot f] + 4f' \cdot f \cdot n^3$ |
| Backward | $f \cdot f \cdot n^3 \cdot k^3$ | $3Cn^3 \log n [f' + f + f' \cdot f] + 4f' \cdot f \cdot n^3$ | $3Cn^3 \log n [f' + f + f' \cdot f] + 4f' \cdot f \cdot n^3$ |
| Update | $f \cdot f \cdot n^3 \cdot k^3$ | $3Cn^3 \log n [f' + f + f' \cdot f] + 4f' \cdot f \cdot n^3$ | $3Cn^3 \log n [f' \cdot f] + 4f' \cdot f \cdot n^3$ |
| Total | $3f' \cdot f \cdot n^3 \cdot k^3$ | $9Cn^3 \log n [f' + f + f' \cdot f] + 12f' \cdot f \cdot n^3$ | $6Cn^3 \log n [f' + f + f' \cdot f] + 12f' \cdot f \cdot n^3$ |

### Table IV: Time required to perform pooling, filtering and transfer function on a full layer with infinite number of processors available.

| Pass | Pooling | Filtering | Transfer function |
|------|---------|-----------|-------------------|
| Forward – $T_{\infty}^{\text{pool}}$ | $n^3$ | $6n^3 \log k$ | $n^3$ |
| Backward – $T_{\infty}^{\text{bwd}}$ | $n^3$ | $n^3$ | $n^3$ |
| Update – $T_{\text{update}}^{\text{parallel}}$ | $-\sigma$ | $-\sigma$ | $n^3$ |

The constant $C$ for the FFT operations is assumed to be 5.
Fig. 3: Task dependency graph. Each edge of the computation graph (Fig. 1) generates multiple nodes of the task dependency graph, corresponding to forward (circle), backward (square), and update (diamond) tasks. Node colors indicate transfer function (green), convolution (red), pooling/filtering (blue), input provider (orange), loss gradient (dark red).

TABLE III: Time required to perform operations on fully connected convolutional layers with infinite number of processors available.

| Pass          | Direct                                           | FFT-based                   | FFT-based (Memoized)       |
|---------------|--------------------------------------------------|-----------------------------|-----------------------------|
| Forward – $T_{fwd}^\infty$ | $n^3 \cdot k^3 + n^3 \lceil \log_2 f \rceil$ | $6Cn^3 \log n + 4n^3 \lceil \log_2 f \rceil$ | $6Cn^3 \log n + 4n^3 \lceil \log_2 f \rceil$ |
| Backward – $T_{bwd}^\infty$ | $n^3 \cdot k^3 + n^3 \lceil \log_2 f' \rceil$ | $6Cn^3 \log n + 4n^3 \lceil \log_2 f' \rceil$ | $6Cn^3 \log n + 4n^3 \lceil \log_2 f' \rceil$ |
| Update – $T_{update}^\infty$ | $n^3 \cdot k^3$ | $6Cn^3 \log n + 4n^3$ | $3Cn^3 \log n + 4n^3$ |

Fig. 4: Theoretically achievable speedup (2) using (a) direct convolution (b) FFT-based convolution with memoizing enabled. Multiple lines of the same color represent networks with different depths, ranging from 4 to 40. The kernels in all the networks have size of $5^3$.

VI. TASK SCHEDULING AND EXECUTION

Brent’s theorem guarantees the existence of a parallel algorithm that achieves a large speedup for training wide ConvNets. We now turn to the problem of designing a parallel algorithm that actually achieves large speedup in practice. Since Brent’s theorem assumes no synchronization and communication overhead, we design our algorithm to minimize synchronization overhead and increase temporal locality of computation to reduce cache misses.

The central quantity in our algorithm is a global priority queue that contains tasks that are ready to be executed together with their priority. A predetermined number of workers will then execute the tasks from the global queue.

A. Priority queue

Tasks are placed on a global queue when all non-update dependencies are satisfied (only tasks with update task as requirements are forward tasks). The rationale behind this design choice is that if a forward task is scheduled for execution without the required update task being done, we will force execution of the update task followed by the forward task that requires the result of the update, hence increase the memory locality.

The tasks on the queue are sorted by priority. Priorities are chosen to increase temporal locality of the computation and minimize the latency of the computation. We introduce two unique strict orderings of the nodes in the ConvNet’s computation graph based on the longest distance, in decreasing order, to any output and input node respectively. Nodes with the same distance will be ordered in some unique way. The priority of the forward task of an edge $e = (u, v)$ will be equal to position of the output node $v$ in the ordering based on the distance to the output nodes, and similarly the priority of the backward task will be equal to the ordering of $u$ based on the distance to the input nodes. This ensures that we prioritize tasks with the longest path to a sink node in the task dependency, which should intuitively favor lower latency schedules. The strict ordering of the tasks with the same distance increases temporal locality by assuring that when multiple tasks with the same distance are scheduled we prefer to execute ones computing 3D images that have to be accumulated in the same sum, thus increasing the probability
of the memory accessed being in the cache.

The update tasks will have the lowest priority of all tasks. Their execution will be forced when their result is required for the forward pass, which increases cache locality as the result will be used immediately. The only other time the update tasks will be executed is if there’s no other forward or backward tasks ready to be executed.

B. Task Execution

The tasks are executed by \(N\) workers. Each worker picks up and executes a task with the highest priority on the queue. **Forward task** algorithm is shown in Algorithm 1. The main functionality of the forward task is to apply the appropriate \(\text{FORWARD-TRANSFORM}\) on the given input image \(I\) and accumulate the result to the sum stored in the output node. The task that adds the last image to the sum then queues all dependent forward tasks for execution.

The main functionality of the forward task is shown in the procedure \(\text{DO-FORWARD}\). However such procedure can only be executed when the update task from the previous round has been completed. This is ensured by creating a new subtask containing the main functionality and calling the \(\text{FORCE}\) method.

**Algorithm 1** Executing a forward task

\[
\begin{align*}
\text{FORWARD-TASK}(e, I) \\
1 & \text{t} = \text{CREATE-TASK}(\text{DO-FORWARD}, e, I) \\
2 & \text{FORCE}(e, \text{update_task}, t)
\end{align*}
\]

\[
\begin{align*}
\text{DO-FORWARD}(e = (u, v), I) \\
1 & I^{\text{out}} = e.\text{FORWARD-TRANSFORM}(I) \\
2 & \text{if ADD-TO-SUM}(v, \text{fwd_sum}, I^{\text{out}}) \\
3 & \quad v.I_f = \text{GET-SUM}(v, \text{fwd_sum}) \\
4 & \quad \text{for } e' \in v.\text{out_edges} \\
5 & \quad \quad t = \text{CREATE-TASK}(\text{FORWARD-TASK}, e', v, I_f) \\
6 & \quad \text{ENQUEUE}(e'.\text{fwd_priority}, t)
\end{align*}
\]

The \(\text{FORCE}\) function receives an update task and a forward subtask as parameters. The goal of the function is to execute the forward subtask but also make sure that the update task has been completed. In order to do that the method first examines the state of the update task which can be one of the following (Note that the \(\text{FORCE}\) is called from the thread scheduled to execute the appropriate forward task).

1) **Completed** - the execution of the update task has been completed; in this case the calling thread just executes the forward subtask.

2) **Queued** - the update task is on the queue waiting to be scheduled for execution; in this case the update task is removed from the queue, and the calling thread executes both the update task and the forward subtask.

3) **Executing** - the update task is currently being executed by some other thread; in this case the forward subtask gets attached to the update task. This flags the thread executing the update task to execute the forward subtask as soon as the update task is completed. The calling thread then returns and picks up another task for execution.

Such design ensures that no thread is ever waiting for completion of an update task, but rather executes the required update task itself, or delegates the forward subtask to the thread currently executing the update task.

**Algorithm 2** Executing a backward task

\[
\begin{align*}
\text{BACKWARD-TASK}(e = (u, v), I) \\
1 & I^{\text{out}} = e.\text{BACKWARD-TRANSFORM}(I) \\
2 & \text{if } e.\text{is_trainable} \\
3 & \quad I_f = u.\text{fwd_image} \\
4 & \quad e.\text{update_task} = \text{CREATE-TASK}(\text{UPDATE}, e, I_f, I) \\
5 & \quad \text{ENQUEUE}(e.\text{update_priority}, e.\text{update_task}) \\
6 & \text{if ADD-TO-SUM}(u.\text{bwd_sum}, I^{\text{out}}d) \\
7 & \quad u.I_b = \text{GET-SUM}(u.\text{bwd_sum}) \\
8 & \quad \text{for } e' \in u.\text{in_edges} \\
9 & \quad \quad t = \text{CREATE-TASK}(\text{BACKWARD-TASK}, e', u, I_b) \\
10 & \quad \text{ENQUEUE}(e'.\text{bwd_priority}, t)
\end{align*}
\]

The \(\text{FORCE}\) function receives an update task and a forward subtask as parameters. The goal of the function is to execute the forward subtask but also make sure that the update task has been completed. In order to do that the method first examines the state of the update task which can be one of the following (Note that the \(\text{FORCE}\) is called from the thread scheduled to execute the appropriate forward task).

1) **Completed** - the execution of the update task has been completed; in this case the calling thread just executes the forward subtask.

2) **Queued** - the update task is on the queue waiting to be scheduled for execution; in this case the update task is removed from the queue, and the calling thread executes both the update task and the forward subtask.

3) **Executing** - the update task is currently being executed by some other thread; in this case the forward subtask gets attached to the update task. This flags the thread executing the update task to execute the forward subtask as soon as the update task is completed. The calling thread then returns and picks up another task for execution.

Such design ensures that no thread is ever waiting for completion of an update task, but rather executes the required update task itself, or delegates the forward subtask to the thread currently executing the update task.

**Algorithm 3** Executing an update task

\[
\begin{align*}
\text{UPDATE}(e, I_f, I_b) \\
1 & G = e.\text{COMPUTE-GRADIENT}(I_f, I_b) \\
2 & e.\text{params} = e.\text{params} - e.\eta \cdot G \\
3 & \text{if this.fwd_subtask} \\
4 & \quad t = \text{this.fwd_subtask} \\
5 & \quad this.fwd_subtask = \text{NIL} \\
6 & \text{EXECUTE}(t)
\end{align*}
\]

VII. SYNCHRONIZATION ISSUES

It is important to minimize the amount of time spent in critical sections – parts of the code that can be only executed by a single thread at a time. The main three points in the algorithm that require synchronization are memory management (allocation/deallocation), operations on the global task queue and concurrent summations.
A. Queue operations

The operations on the global task priority queue have to be synchronized. The queue is implemented as a heap of lists lowering the complexity of insertion and deletion from \( \log N \) to \( \log K \), where \( N \) is the total number of tasks in the queue and \( K \) is the number of distinct values for the priority of the tasks inside the queue. Depending on the network structure, this number can be much smaller than the total number of tasks in the queue, which is especially true for wide networks.

B. Wait-free concurrent summation

When multiple edges converge on the same node in the computation graph, it means that multiple convolutions executed in parallel need to add their results to the same accumulated sum. The additions have to be synchronized; only one thread is allowed to change the sum. The naive strategy, waiting until all other threads have finished adding their images to the sum, would lead to critical section time that scales linearly with the image size \( n^3 \). We propose a novel method that eliminates the dependence on image size by performing only pointer operations inside the critical section, which works as follows.

Suppose that multiple threads are executing ADD-TO-SUM in Algorithm 4. For each thread, \( v \) points to a different 3D image. We would like the pointer to the sum of all these images to be stored in the object \( S \) when the computation terminates. This is accomplished by having each thread repeatedly try to reset the pointer to the sum stored in \( S.\text{sum} \) to point to \( v \) instead. If the thread succeeds, it stops working. If the thread fails, it adds the value pointed to by \( S.\text{sum} \) to the location referenced by \( v \), and sets the pointer to NIL. Every thread continues to work until it succeeds. Once the last thread succeeds, \( S \) will contain the correct answer. Note that this algorithm does the time-consuming additions outside the critical section (lines 5-11).

Algorithm 4 Wait-free concurrent summation algorithm

```
ADD-TO-SUM(S, v)
1 v' = NIL
2 last = FALSE
3 while TRUE
4   ACQUIRE(S.lock)
5     if S.sum == NIL
6       S.sum = v
7       v = NIL
8       S.total = S.total + 1
9       last = (S.total == S.required)
10      else v' = S.sum
11     S.sum = NIL
12    RELEASE(S.lock)
13   if v == NIL
14      return last
15   else ADD-TO(v, v') // v = v + v'
```

C. Memory management

ZNN implements two custom memory allocators. These are designed to be faster than standard memory management routines, at the cost of using more memory. One custom allocator is dedicated to 3D images, which are usually large, and the other is dedicated to small objects used in auxiliary data structures. Both allocators maintain 32 distinct pools of memory chunks. Each pool \( i \), \( i \in 0 \ldots 31 \) contains chunks of sizes of \( 2^i \). Lock-free queues, as described in [18] and implemented as a part of the boost [19] library are used to implement the pool operations. The only difference between the allocators is the memory alignment—the 3D image memory allocator ensures proper memory alignment for utilizing SIMD instructions. No memory is shared between the two allocators.

When a chunk of memory of size \( s \) is requested, first \( s \) is rounded up to the nearest power of 2. The appropriate pool is examined for available memory chunks. If there’s an available chunk we return it and remove it from the pool. If no chunks are available we allocate one from the system and return it.

When de-allocating a chunk memory, it is simply added to the appropriate pool, and no memory is ever returned to the system. This means that the memory usage of our program can never decrease. In practice, as the ConvNet training consist of a single loop performing the same work, our memory usage peaks after a few rounds.

In the worst case this strategy can lead to near \( 2\times \) memory usage overhead; however the available memory to the CPU is rarely a limiting factor in training a network. In the future, we might consider implementing more advanced memory allocators, such as ones with thread-local pools in addition to the global pool, or ones with higher granularity of available chunk sizes to reduce the size overhead.

VIII. Scalability

We performed measurements of the speedup achieved by our proposed parallel algorithm relative to the serial algorithm, using the CPU systems listed in Table V. Amazon EC2 instances with 8 and 18 cores (c4.4xlarge and c4.8xlarge) were chosen for benchmarking, because they are readily available to anyone. A 4-way CPU system was included because it has 40 cores, though this is a relatively specialized piece of hardware. For an even larger number of cores, we also benchmarked the Xeon Phi [20] with Intel MKL (version 11.2) libraries for FFTs and direct convolution.

| CPU          | Frequency | Cores/Threads |
|--------------|-----------|---------------|
| Intel® Xeon® E5-2666 v3 | 2.9 GHz   | 8 cores/16 threads |
| Intel® Xeon® E5-2666 v3 | 2.9 GHz   | 18 cores/36 threads |
| Intel® Xeon® E7-4850 | 2.0 GHz   | 40 cores/80 threads |
| Intel® Xeon Phi™ 5110P | 1.053 GHz | 60 cores/240 threads |

TABLE V: Machines used for the experiments
linear transfer function layer (T), and two $2 \times 2$ max-filtering layers (2nd and 4th). The sequence of layer types was CTM-CTMCTCT. The output patch size was $48 \times 48$.

The ZNN measurements were performed by first running the gradient learning algorithm for 5 warm-up rounds and then averaging the time required for the next 50 rounds. The GPU measurements were averaged over 100 rounds.

2D ConvNets were implemented as a special case of 3D ConvNets, by setting one of the dimensions to have size one. The width of the ConvNets was varied as described below. FFT convolution was employed for 2D, and direct convolution for 3D to illustrate the use of both methods; reversing this yields similar results. Other network architectures and kernel sizes also yield similar results.

Fig. 5 shows speedup attained by various CPUs as a function of two parameters, number of worker threads and network width. Each graph shows the result of varying the number of workers while network width is held fixed. To achieve near maximal possible speedup ZNN requires sufficiently wide networks ($\geq 30$ for multicore CPUs and $\geq 80$ for the manycore CPU) and sufficiently many worker threads (number of hyperthreads for multicore and number of hardware threads for manycore). The value of the maximal speedup is equal to the number of cores or a bit larger (maximal height of graphs).

For a wide network on multicore CPUs, speedup increases linearly until the number of worker threads equals the number of cores. After that the increase continues at a slower rate. For wide networks on Xeon Phi™, speedup increases linearly until the number of worker threads equals the number of cores, then more slowly until double that number, and then even slower until the number of hardware threads. The maximal achieved speedups for networks of different widths are shown in Figs. 6 and 7.

IX. CPU vs. GPU

While the preceding results show that ZNN can efficiently utilize CPUs, it is also important to know how the resulting performance compares to GPU implementations of ConvNet learning. Therefore, we benchmarked ZNN against Caffe and Theano, two popular GPU implementations. Comparison can be tricky because CPU and GPU implementations by definition cannot be run on the same hardware.

We chose to run Caffe and Theano on a Titan X GPU, and ZNN on an 18 core Amazon EC2 instance (c4.8xlarge). We chose this particular comparison, because the alternatives seemed unfair. For example, we could have run ZNN on specialized hardware with more CPU cores than the EC2 instance. This comparison seemed unfair because the specialized hardware would have been much more costly than Titan X and less accessible than Amazon EC2. Also, we could have used GPU instances from Amazon EC2, but these are currently much slower than Titan X ($3 \times$ or more on our benchmarks) and have half the onboard RAM.

For Caffe, both default and cuDNN implementations were used. For 3D ConvNets we only used Theano, as the official release of Caffe still does not support 3D ConvNets. Our Caffe and Theano code is publicly available in the ZNN repository.

ZNN used FFT convolution for both 2D and 3D, as this was found to be optimal by the auto-tuning capability of ZNN. Caffe and Theano used direct convolution.

Our ConvNets contained 6 fully-connected convolutional (C) layers, each followed by a rectified linear transfer function
layer (T), and two max-pooling (P) layers, either $2 \times 2$ or $2 \times 2 \times 2$. The sequence of the layer types was CTPT- PCTCTCTCT. All networks had width 40, while the sizes of the kernels and the output patch varied.

All benchmark times were for “sparse training,” meaning that the ConvNet is used to produce predictions for pixels in the output patch that form a lattice with period 4 in every dimension. The loss of predicted output pixels is due to the two layers of max-pooling.

As noted before, ZNN can also perform “dense training,” meaning that the ConvNet is used to produce predictions for every pixel in the output patch by applying the ConvNet to a window that slides across every “valid” location in the input patch. Requiring Caffe or Theano to perform dense training could have been accomplished by computing 16 sparse outputs in 2D and 64 in 3D to assemble a dense output. This method is very inefficient and would have been no contest with ZNN.

A. Speed

The comparison of 2D ConvNets is shown in Fig. 8. ZNN is faster than Caffe and Theano for sufficiently large kernels ($30 \times 30$ or larger). This makes sense because FFT convolution (ZNN) is more efficient than direct convolution (Caffe and Theano) for sufficiently large kernels.

Such large kernels are not generally used in practice, so ZNN may not be competitive with GPU implementations for 2D networks. On the other hand, ZNN opens up the possibility of efficiently training networks with large kernels, and these might find some practical application in the future.

The comparison of 3D ConvNets is shown in Fig. 9. ZNN is comparable to Theano even for modest kernel sizes of $5 \times 5 \times 5$ and outperforms Theano for kernel sizes of $7 \times 7 \times 7$ and greater. Such kernel sizes are currently relevant for practical applications [21]. The comparison makes sense, because we expect the crossover point for complexity of FFT vs. direct convolution to occur for smaller (linear) kernel sizes in 3D.

B. Memory

Working memory is another computational resource that is important for training ConvNets. Given the limited amount of onboard GPU memory, we were unable to use Theano to train 3D networks with kernel sizes larger than $7 \times 7 \times 7$. We were also unable to use Caffe to train many 2D networks (see missing bars in Fig. 8).

ZNN enables training of larger networks mostly because a typical CPU system has much more RAM than even a top GPU. Titan X, for example, has just 12 GB of onboard RAM. Additionally, ZNN can achieve even higher speed by using extra RAM space, as in the case of FFT memoization. When using FFT-based convolutions, with the memoization disabled, ZNN is more efficient in its usage of RAM than the proposed GPU methods. The memory overhead of the methods proposed

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**Fig. 5**: Speedup versus number of threads for 2D (first row) and 3D (second row) achieved on machines in Table V (left-to-right columns). In each graph, the lines are for network widths 5, 10, 15, 20, 25, 30, 40, 50, 60, 80, 100, 120, from bottom to top.

**Fig. 9**: Comparison of ZNN and Theano for 3D ConvNets. The charts from the top down have kernel sizes of $3^3$, $5^3$ and $7^3$. 

- The comparison of 2D ConvNets is shown in Fig. 8. ZNN is faster than Caffe and Theano for sufficiently large kernels ($30 \times 30$ or larger). This makes sense because FFT convolution (ZNN) is more efficient than direct convolution (Caffe and Theano) for sufficiently large kernels.
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in [5, 6] could be very high as it is proportional to the number of kernels in a layer. In contrast ZNN’s memory overhead is proportional to the number of workers.

X. IMPLEMENTATION DETAILS

ZNN is implemented in C++ and is publicly available under the GPL2 license (https://github.com/zlateski/znn-release). It can use either fflw or intel MKL for FFTs and either provided code or intel MKL libraries for direct convolution. Using fflw instead of MKL yields same scalability but lower absolute performances due to the differences in single thread performances of the two libraries. The repository also provides alternative scheduling strategies such as simple FIFO or LIFO as well as some more complex ones based on work stealing [23]. The alternative scheduling strategies achieve noticeably lower scalability than the one proposed in the paper for most networks. However, some very specific networks might benefit from alternative scheduling algorithms. Future work can include automatic detection of the best scheduling strategy.

XI. CONCLUSIONS

ZNN achieves high performances by efficiently utilizing the available CPUs. We expect an increase in the number of cores per chip (or Xeon PhiTM card) in the future, making ZNN even more practical. In fact, we have already used ZNN to achieve state of the art results in boundary detection [23] and computation of dendritic arbor densities [24].

Having a large amount of RAM available to the CPU, ZNN can efficiently train very large ConvNets with large kernels. ZNN allows for easy extensions and can efficiently train a ConvNet with an arbitrary topology, allowing for new research.

Unlike the ZNN’s task parallelization model, the current GPU implementations employ SIMD parallelism to perform computation on one whole layer at a time, thus limiting the network structure. Mainly, the computation is parallelized such that a single thread computes the value of a single voxel of an output image. Libraries like cuDNN provide optimized primitives for fully connected convolutional layers by reducing all the required convolutions in the layer to a matrix multiplication, which is then parallelized on the GPU.

Extending the functionality requires the user to provide a parallelized implementation of the new layer type, which typically requires great knowledge of GPU programming, and might take a long time. Contrary to that, ZNN’s task parallelism allows for easy extensions by simply providing serial functions for the forward and backward pass, as well as the gradient computation, if required. ZNN’s repository contains some sample extensions providing functionality of dropout [25] and multi-scale [14, 16] networks.

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