Distributed Machine Learning and the Parameter Server

CS4787 Lecture 23 — Spring 2021

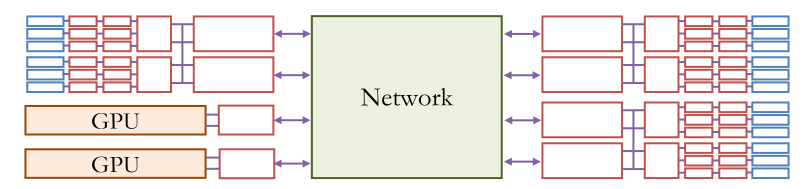
So far, we've been talking about ways to scale our machine learning pipeline that focus on a single machine. But if we *really* want to scale up to *huge* datasets and models, eventually one machine won't be enough.

This lecture will cover methods for using multiple machines to do learning.



Distributed computing basics

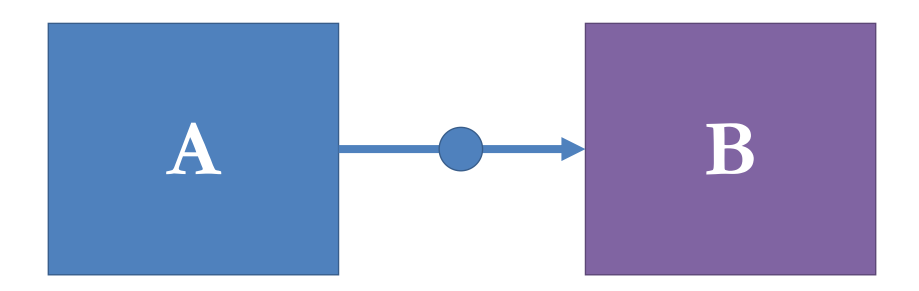
- Distributed parallel computing involves two or more machines collaborating on a single task by communicating over a network.
 - Unlike parallel programming on a single machine, distributed computing requires explicit (i.e. written in software) communication among the workers.



• There are a **few basic patterns of communication** that are used by distributed programs.

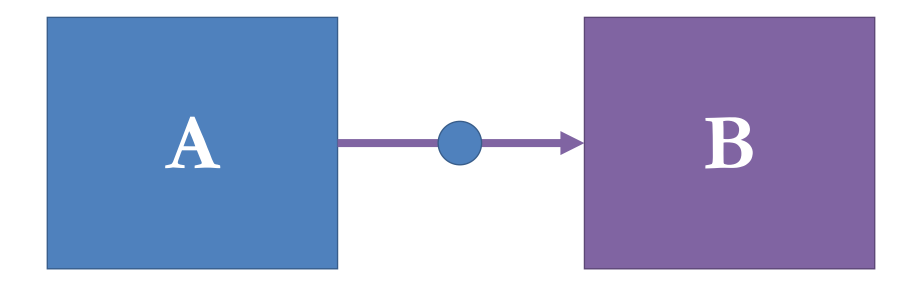
Basic patterns of communication Push

• Machine A sends some data to machine B.



Basic patterns of communication Pull

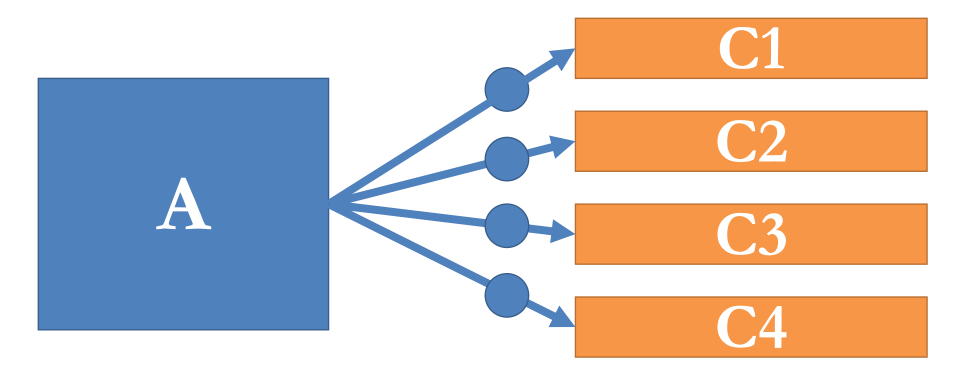
• Machine B requests some data from machine B.



• This differs from push only in terms of who initiates the communication

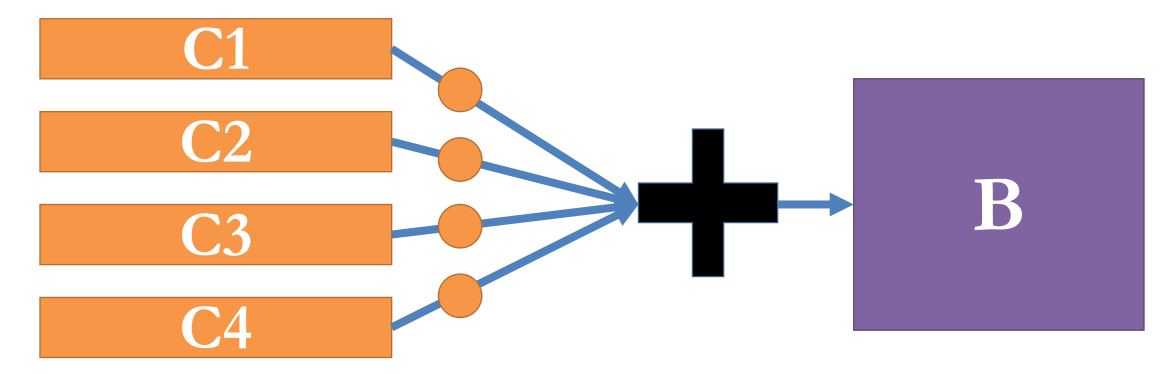
Basic patterns of communication Broadcast

• Machine A sends data to many machines.



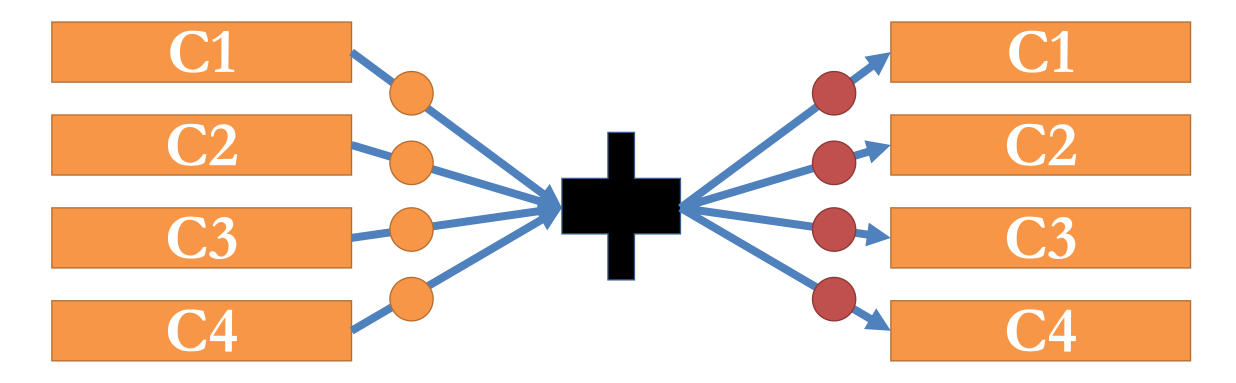
Basic patterns of communication Reduce

• Compute some reduction (usually a sum) of data on multiple machines C1, C2, ..., Cn and materialize the result on one machine B.



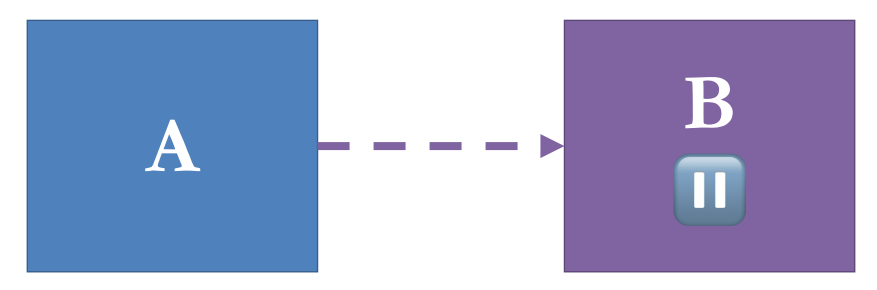
Basic patterns of communication All-Reduce

• Compute some reduction (usually a sum) of data on multiple machines and materialize the result on all those machines.



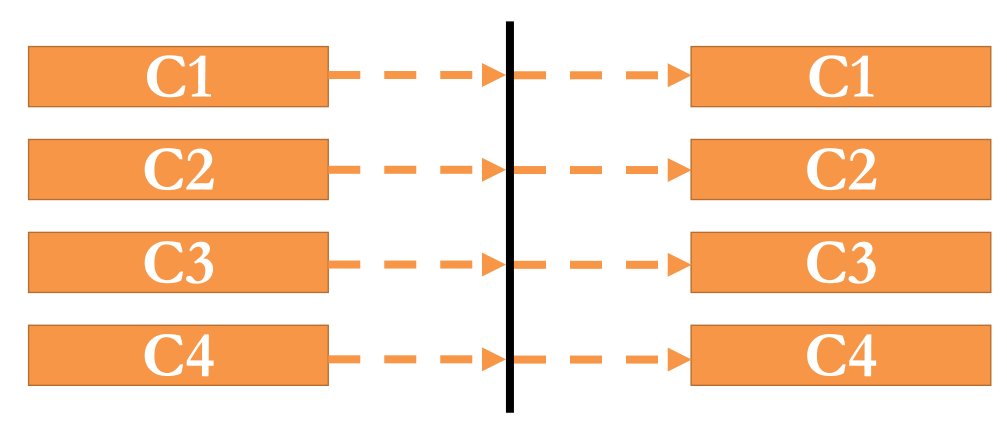
Basic patterns of communication Wait

• One machine pauses its computation and waits on a signal from another machine



Basic patterns of communication Barrier

• Many machines wait until all those machines reach a point in their execution, then continue from there



Patterns of Communication Summary

- **Push.** Machine A sends some data to machine B.
- Pull. Machine B requests some data from machine A.
 - This differs from push only in terms of who initiates the communication.
- Broadcast. Machine A sends some data to many machines C1, C2, ..., Cn.
- Reduce. Compute some reduction (usually a sum) of data on multiple machines C1, C2, ..., Cn and materialize the result on one machine B.
- All-reduce. Compute some reduction (usually a sum) of data on multiple machines C1, C2, ..., Cn and materialize the result on all those machines.
- Wait. One machine pauses its computation and waits for data to be received from another machine.
- Barrier. Many machines wait until all other machines reach a point in their code before proceeding.

Overlapping computation and communication

- Communicating over the network can have high latency
 - we want to hide this latency
- An important principle of distributed computing is overlapping computation and communication
- For the best performance, we want our workers to still be doing useful work while communication is going on
 - rather than having to stop and wait for the communication to finish
 - sometimes called a stall

Running SGD with All-reduce

• All-reduce gives us a simple way of running learning algorithms such as SGD in a distributed fashion.

• Simply put, the idea is to just **parallelize the minibatch**. We start with an identical copy of the parameter on each worker.

• Recall that SGD update step looks like:

$$w_{t+1} = w_t - \alpha_t \cdot \frac{1}{B} \sum_{b=1}^{B} \nabla f_{i_{b,t}}(w_t),$$

Running SGD with All-reduce (continued)

• If there are M worker machines such that $B = M \cdot B'$, then

$$w_{t+1} = w_t - \alpha_t \cdot \frac{1}{M} \sum_{m=1}^{M} \frac{1}{B'} \sum_{b=1}^{B'} \nabla f_{i_{m,b,t}}(w_t).$$

• Now, we assign the computation of the sum when m = 1 to worker 1, the computation of the sum when m = 2 to worker 2, et cetera.

• After all the gradients are computed, we can perform the outer sum with an all-reduce operation.

Running SGD with All-reduce (continued)

- After this all-reduce, the whole sum (which is essentially the minibatch gradient) will be present on all the machines
 - so each machine can now update its copy of the parameters
- Since sum is same on all machines, the parameters will update in lockstep
- Statistically equivalent to sequential SGD!

```
Algorithm 1 Distributed SGD with All-Reduce
  input: loss function examples f_1, f_2, \ldots, number of machines M, per-machine minibatch size B'
  input: learning rate schedule \alpha_t, initial parameters w_0, number of iterations T
  for m=1 to M run in parallel on machine m
      load w_0 from algorithm inputs
      for t = 1 to T do
          select a minibatch i_{m,1,t}, i_{m,2,t}, \ldots, i_{m,B',t} of size B'
          compute g_{m,t} \leftarrow \frac{1}{B'} \sum_{i=1}^{B'} \nabla f_{i_{m,b,t}}(w_{t-1})
          all-reduce across all workers to compute G_t = \sum g_{m,t}
                                                                  m=1
          update model w_t \leftarrow w_{t-1} - \frac{\alpha_t}{M} \cdot G_t
      end for
  end parallel for
  return w_T (from any machine)
```

Same approach can be used for momentum, Adam, etc.

What are the benefits of distributing SGD with all-reduce? What are the drawbacks?

Benefits of distributed SGD with All-reduce

- The algorithm is easy to reason about, since it's statistically equivalent to minibatch SGD.
 - And we can use the same hyperparameters for the most part.
- The algorithm is easy to implement
 - since all the worker machines have the same role and it runs on top of standard distributed computing primitives.

Drawbacks of distributed SGD with all-reduce

- While the communication for the all-reduce is happening, the workers are (for the most part) idle.
- We're not overlapping computation and communication.
 - At least by default
 - We can overlap communication with preprocessing/data augmentation
- The effective minibatch size is growing with the number of machines, and for cases where we *don't* want to run with a large minibatch size for statistical reasons, this can prevent us from scaling to large numbers of machines using this method.

Where do we get the training examples from?

• There are two general options for distributed learning.

Training data servers

- Have one or more non-worker servers dedicated to storing the training examples (e.g. a distributed in-memory filesystem)
- The worker machines load training examples from those servers.
- These servers can handle preprocessing and data augmentation (but usually don't)

Partitioned dataset

• Partition the training examples among the workers themselves and store them locally in memory on the workers.

DEMO

The Parameter Server Model

The Basic Idea

- Recall from the early lectures in this course that a lot of our theory talked about the convergence of optimization algorithms.
 - This convergence was measured by some function over the parameters at time t (e.g. the objective function or the norm of its gradient) that is decreasing with t, which shows that the algorithm is making progress.
- For this to even make sense, though, we need to be able to talk about the value of the parameters at time t as the algorithm runs.
 - E.g. in SGD, we had

$$w_{t+1} = w_t - \alpha_t \nabla f_{i_t}(w_t)$$

Parameter Server Basics Continued

 For a program running on time t is just the value of s DRAM) at that time.

 But in a distributed setting must be done explicitly.

• Each machine will usually time, some of which may have been upwant to do something more complicated

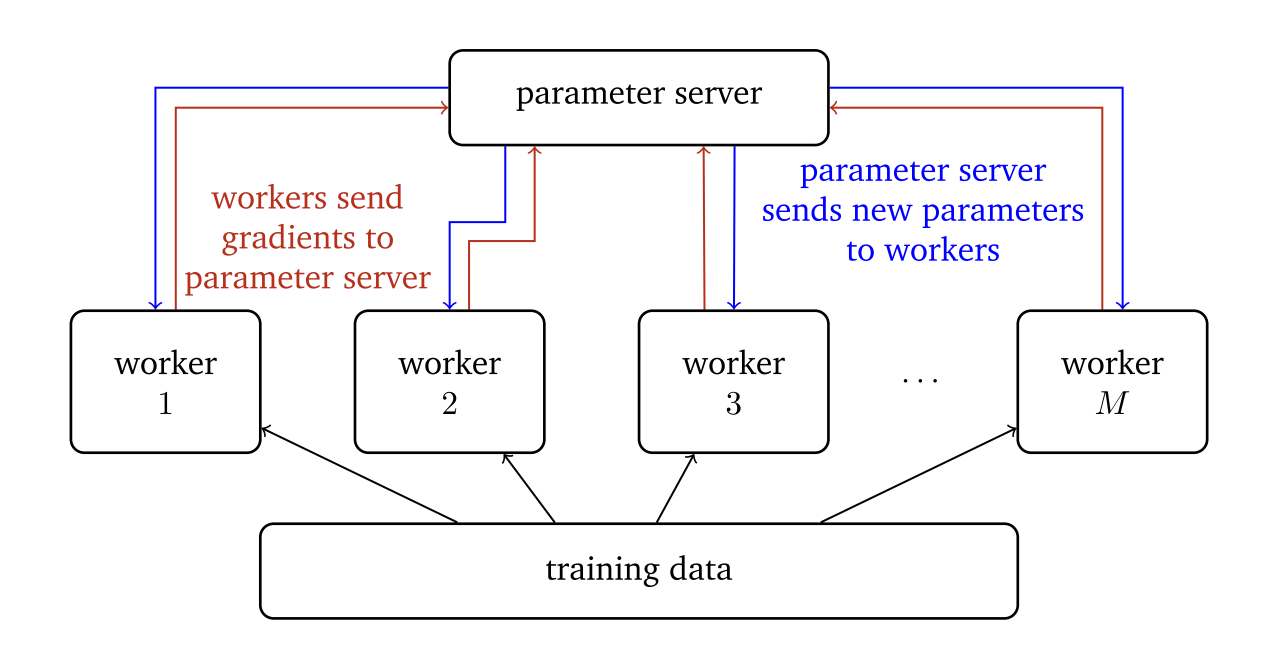
For SGD with all-reduce, we can answer this question easily, since the value of the parameters is the same on all workers (it's guaranteed to be the same by the all-reduce operation). We just appoint this identical shared value to be the value of the parameters at any given time.

reduce.

• This raises the question: when reasoning about a distributed algorithm, what we should consider to be the value of the parameters a given time?

The Parameter Server Model

- The parameter server model answers this question differently by appointing a single machine, the **parameter server**, the explicit responsibility of maintaining the current value of the parameters.
 - The most up-to-date gold-standard parameters are the ones stored in memory on the parameter server.
- The parameter server updates its parameters by using gradients that are computed by the other machines, known as **workers**, and pushed to the parameter server.
- Periodically, the parameter server **broadcasts its updated parameters** to all the other worker machines, so that they can use the updated parameters to compute gradients.



Learning with the parameter server

• Many ways to learn with a parameter server

Synchronous distributed training

• Similar to all-reduce, but with gradients summed on a central parameter server

• Asynchronous distributed training

- Compute and send gradients and add them to the model as soon as possible
- Broadcast updates whenever they are available

Algorithm 2 Asynchronous Distributed SGD with the Parameter Server Model

```
input: loss function examples f_1, f_2, \ldots, number of worker machines M, per-machine minibatch size B'
input: learning rate \alpha, initial parameters w_0, number of iterations per worker T
for m = 1 to M run in parallel on machine m
    load w_{m,0} from the parameter server
    for t = 1 to T do
       select a minibatch i_{m,1,t}, i_{m,2,t}, \dots, i_{m,B',t} of size B' compute g_{m,t} \leftarrow \frac{1}{B'} \sum_{b=1}^{B} \nabla f_{i_{m,b,t}}(w_{m,t-1})
        push gradient g_{m,t} to the parameter server
        receive new model w_{m,t} from the parameter server
    end for
end parallel for
run in parallel on param server
    initialize model w \leftarrow w_0
    loop
        receive a gradient g from a worker
        update model w \leftarrow w - \alpha g
        send w back to the worker
    end loop
end run on param server
return w_T (from any machine)
```

Is this still equivalent to sequential minibatch SGD running on a single machine?

Multiple parameter servers

• If the parameters are too numerous for a single parameter server to handle, we can use multiple parameter server machines.

- We partition the parameters among the multiple parameter servers
 - Each server is only responsible for maintaining the parameters in its partition.
 - When a worker wants to send a gradient, it will partition that gradient vector and send each chunk to the corresponding parameter server; later, it will receive the corresponding chunk of the updated model from that parameter server machine.

• This lets us scale up to very large models!

Other Ways To Distribute

The methods we discussed so far distributed across the minibatch (for all-reduce SGD) and across iterations of SGD (for asynchronous parameter-server SGD).

But there are other ways to distribute that are used in practice too.

Distribution for hyperparameter optimization

- This is something we've already talked about.
- Many commonly used hyperparameter optimization algorithms, such as **grid search and random search**, are very simple to distribute.
 - They can easily be run on many parallel workers to get results faster.

Model Parallelism

- Main idea: **partition the layers** of a neural network among different worker machines.
- This makes each worker responsible for a subset of the parameters.
- Forward and backward signals running through the neural network during backpropagation now also run across the computer network between the different parallel machines.
 - Particularly useful if the parameters won't fit in memory on a single machine.
 - This is very important when we move to specialized machine learning accelerator hardware, where we're running on chips that typically have limited memory and communication bandwidth.

Conclusion and Summary

• Distributed computing is a powerful tool for scaling machine learning

- We talked about two methods for distributed training:
 - Minibatch SGD with All-reduce
 - The parameter server approach
- And distribution can be beneficial for many other tasks too!