

# Lecture 19: Accelerating and scaling DNN training (GPU)

Data C182 (Fall 2024). Week 12. Tuesday Nov 12th, 2024

Speaker: Eric Kim

### Announcements

- HW03 ("Transformers + NLP") out! Due: Fri Nov 22nd 11:59 PM PST
  - Please start early!
- Midterm finalized stats (post regrade requests)

# Today's lecture

- (Part 1) Guest talk by William Chen (TA)!
  - Topic: Robotics + DL
- (Part 2) GPU, multi-GPU, multi-Node training
- GPU tour: CUDA, cuDNN, NCCL (all\_reduce)

### What is a GPU?

- GPU: "Graphics Processing Unit"
- A separate hardware device that connects to your computer. Aka "discrete graphics card"
- Reason to use them: if you can express your computation in a "GPU-friendly" way, then you can have much higher compute **throughput** than CPUs
  - Typical applications: gaming, photo/video editing, video streaming
- CPU benefits: flexible, low-latency, accessible (every computer has a CPU!)
- GPU benefits: (very) high-throughput



Pictured: my GeForce RTX 3080 Ti!

### What is a GPU?

- Two classes of GPUs:
  - Gaming/commodity. Aka what you buy for your gaming PC.
    - Ex: Nvidia GeForce RTX 4080: >\$999
  - Data center. Aka what big companies like Google/Meta/OpenAl train/serve their DNN models on.
    - Ex: Nvidia H100 GPU: ~\$25k per card.
    - Most heavy-duty DNN train machines have 8 GPUS, so ~\$200K per machine.
      - Aka Amazon Cloud + Nvidia is making the big bucks right now!



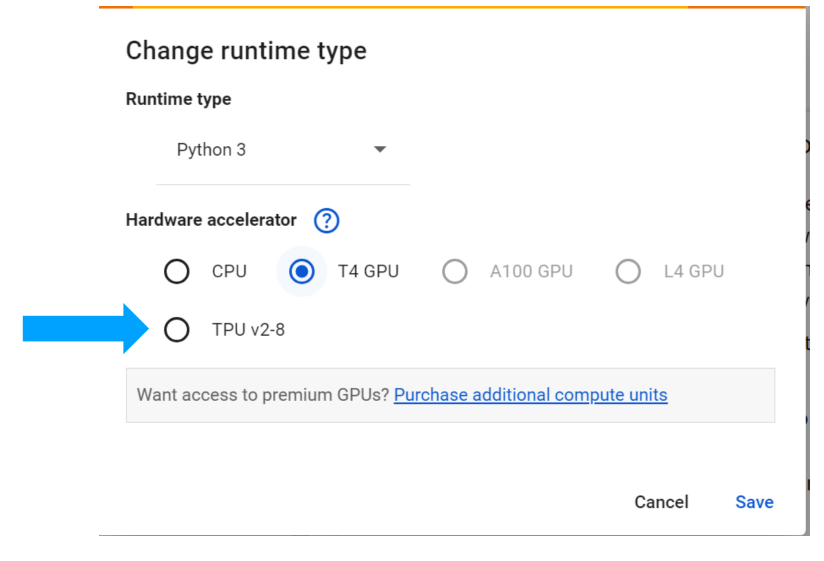
Pictured: Nvidia DGX-2 data center machine [link] at Nvidia's booth for CVPR 2018 (Salt Lake City!). Has 16 Tesla V100 GPUs, likely worth several hundreds of thousands of USD!

# Major GPU types

- As of 2024: in this course (and ~99% of ML/AI): we use Nvidia GPUs (CUDA, cuDNN).
- DNN frameworks like pytorch have excellent support for Nvidia GPUs (CUDA)
- TPU ("Tensor Processing Unit"): A special type of "Al accelerator" hardware built by Google [link]
  - Ex: Tensorflow, Google papers often use TPUs instead of Nvidia GPUs
  - In Colab, you can choose either Nvidia GPU (eg T4) or a TPU.
- AMD is trying to enter the Al/ML market too (IMO, difficult to break in at the moment)



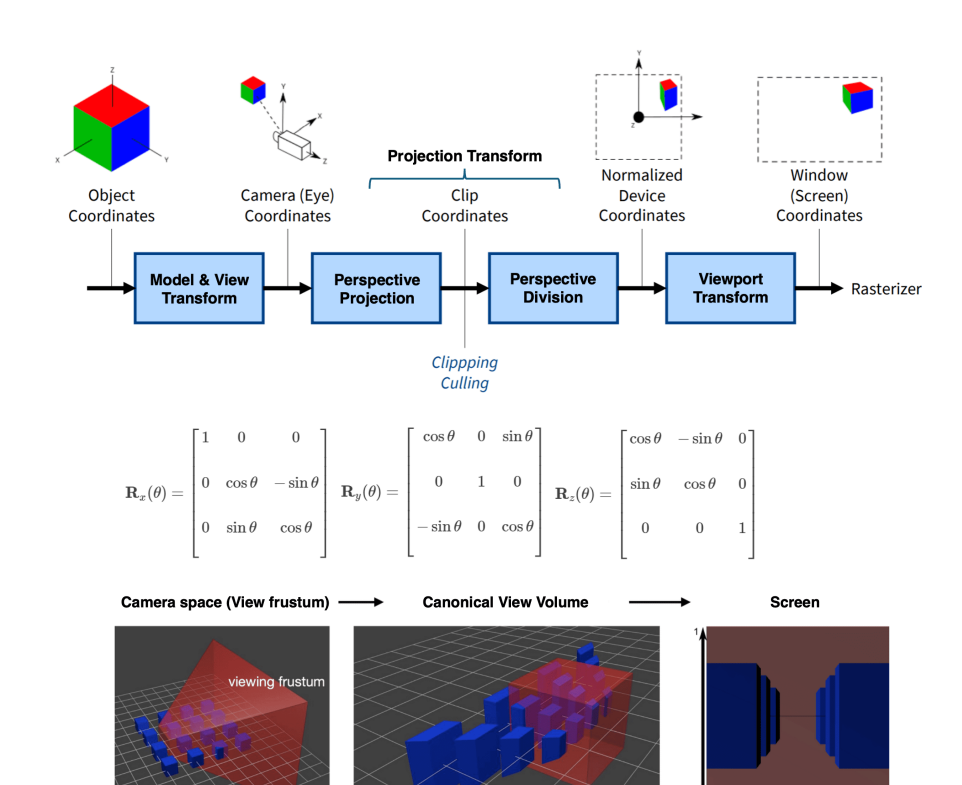


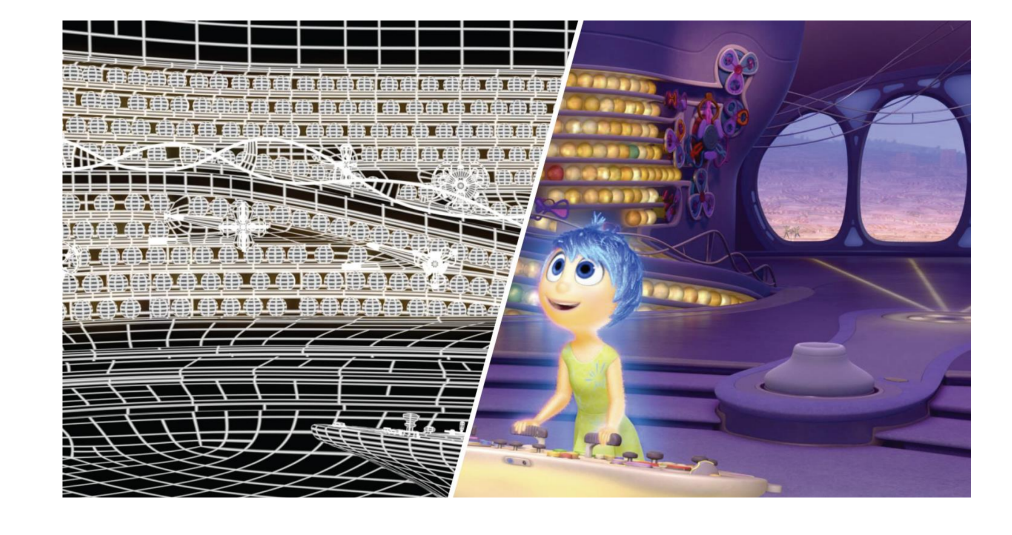


Colab: T4 is Nvidia GPU, "TPU" is Google TPU

### Original GPU motivation

- Original intent: accelerate graphics
   processing, eg for computer graphics (CGI, Pixar) and video games (real-time graphics)
  - Interested? Take CS 184! [link] Computer graphics and computer vision have a healthy relationship, lots of interesting overlap
- Motivation: graphics ultimately boils down to matrices and vectors. Need to accelerate matrix/vector computation, as the CPU wasn't enough back then (and still isn't now)
  - Ex: a 3D point is represented as a 3D\* vector, rotations/translations/scales are represented as 3x3 matrices.





### What do GPUs excel at?

- GPUs can do a LOT of parallel computation, much more so than CPUs!
  - CPU: Typically has 4-16 cores (up to 8-32 active threads with hyperthreading)
  - GPU: A Tesla P100 GPU has 56 "Streaming Multiprocesesors", each with 2048 active threads (up to 114688 active threads!)
- If your computation can be easily parallelized, then GPUs are very good
- Fortunately, nearly all DNN code falls under this category!
  - Matrix/vector calculations (Linear, Conv2d, Relu, Softmax, etc.)
- (2017) "Tensor Cores" [link] are an Nvidia hardware feature that further accelerates certain DNN operations, particularly for lower-precision datatypes like float16 ("mixed precision training")

### GPUs for ML

- AlexNet (2012) was an early (successful!)
   example of training a ConvNet on GPU hardware
- GPU acceleration caught on: DNN libraries began offering "first class" GPU support
- Caffe (2014): Developed at UC Berkeley. [link]
- Caffe2 (2017): Developed at Facebook. [link]
  - Note: basically deprecated in favor of pytorch
- Tensorflow (2015): Developed at Google [link]
- Pytorch (2016+): Developed at Facebook [link]

As of 2024, Pytorch and Tensorflow are the top DNN frameworks in use at both industry and academia. Personally: I prefer pytorch, but each has their strengths and weaknesses.

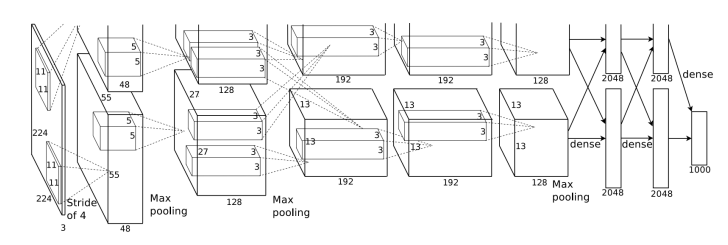


Figure 2: An illustration of the architecture of our CNN, explicitly showing the delineation of responsibilities between the two GPUs. One GPU runs the layer-parts at the top of the figure while the other runs the layer-parts at the bottom. The GPUs communicate only at certain layers. The network's input is 150,528-dimensional, and the number of neurons in the network's remaining layers is given by 253,440–186,624–64,896–64,896–43,264–4096–4096–1000.







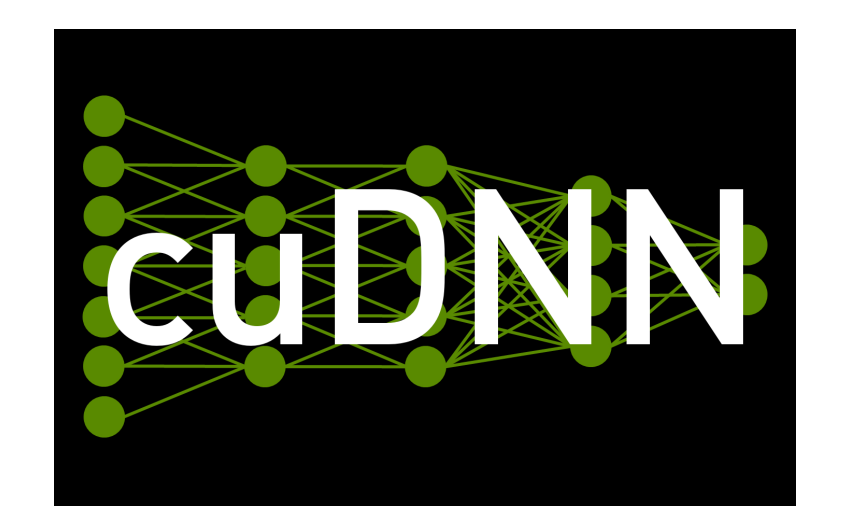
### Nvidia: CUDA

- CUDA: Compute Unified Device Architecture
- Low-level library that tells the GPU how to compute your desired code.
- CUDA code is basically C code
- Main idea: you write CUDA code that expresses your computation in a "parallelizable" way, to effectively utilize the GPU's many execution threads
  - Writing performant parallel code is an art! 99% of the time ML devs don't have to worry about this

```
__global__
void add(int n, float * x, float * y) {
  int index = threadIdx.x;
  int stride = blockDim.x;
  for (int i = index; i < n; i += stride)
    y[i] = x[i] + y[i];
}</pre>
```

### Nvidia cuDNN

- cuDNN: an Nvidia library built on top of CUDA to provide high-performance DNN kernels (like optimized linear forward/backward, conv2d fwd/bkwd, etc)
- DNN frameworks like pytorch, tensorflow ultimately compile down to CUDA code (often via cuDNN calls) which is what actually runs on your GPU!

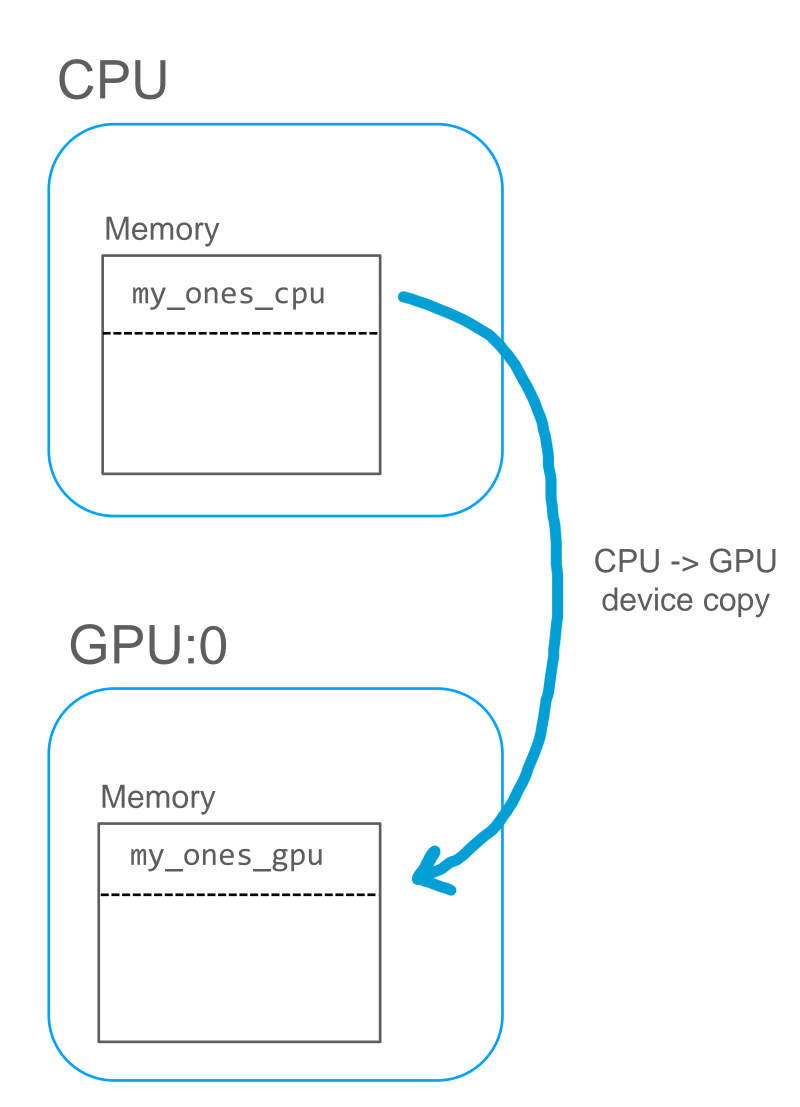


# Pytorch + GPUs

- Pytorch makes it easy to use GPUs in your pytorch code!
- Terminology: all Tensors live on a `torch.device` [link]
  - By default: CPU device: `torch.device("cpu")`
  - GPU (cuda): torch.device("cuda")
    - (If your machine has multiple GPUs) torch.device("cuda:0"), torch.device("cuda:1"), ...
- Main principle: when doing an operation involving two tensors, both tensors must be on the same device!

### Pytorch example code: Tensor devices

```
# create tensor directly on cuda device
my_zeros_cuda = torch.zeros(size=[2, 3], device=torch.device("cuda:0"))
# create tensor on CPU, but move it to GPU (CPU -> GPU copy)
my_ones_cpu = torch.ones(size=[2, 4])
my_ones_cuda = my_ones_cpu.to(device=torch.device("cuda:0"))
# if you have multiple gpus, can send a tensor to a specific one
my_ones_cuda_device0 = my_ones_cpu.to(device=torch.device("cuda:0"))
my ones cuda device1 = my ones cpu.to(device=torch.device("cuda:1"))
# finally, can send a tensor from GPU back to CPU
my_ones_cuda_to_cpu = my_ones_cuda.to(device=torch.device("cpu"))
# Beware: when doing ops between two tensors, they both most be on the
same device!
# RuntimeError: Expected all tensors to be on the same device, but found
at least two devices, cuda:0 and cpu!
my_ones_cpu + my_ones_cuda # ERROR
```



# Pytorch example code: module.to(device)

```
import torch, torch.nn as nn
class Autoencoder(nn.Module):
   def __init__(self, out_channels_first: int = 16):
       super().__init__()
       self.encoder = nn.Sequential( # like the Composition layer you built
           nn.Conv2d(1, out_channels_first, 3, stride=2, padding=1),
           nn.ReLU(),
           nn.Conv2d(out_channels_first, 32, 3, stride=2, padding=1),
           nn.ReLU(),
           nn.Conv2d(32, 64, 7)
       self.decoder = nn.Sequential(
           nn.ConvTranspose2d(64, 32, 7),
           nn.ReLU(),
           nn.ConvTranspose2d(32, out_channels_first, 3, stride=2, padding=1, output_padding=1),
           nn.ReLU(),
           nn.ConvTranspose2d(out_channels_first, 1, 3, stride=2, padding=1, output_padding=1),
           nn.Sigmoid()
   def forward(self, x):
       x = self.encoder(x)
       x = self.decoder(x)
       return x
def print_statedict_info(module: torch.nn.Module):
   for param_key, param_val in module.state_dict().items():
       print(f"param_key={param_key}: {param_val.shape}, device={param_val.device}")
model a = Autoencoder()
print("CPU module")
print_statedict_info(model_a)
model_a_gpu = model_a.to(device=torch.device("cuda:0"))
print("GPU module")
print statedict info(model a gpu)
```

Takeaway: model.to(device=device) copies all of the model parameters to the target device (eg all weight/bias params)

```
(venv) PS
C:\Users\Eric\teaching\data_c182_fall2024\src\lectures\lecture19> python
.\module_gpu_demo.py
CPU module
param_key=encoder.0.weight: torch.Size([16, 1, 3, 3]), device=cpu
param_key=encoder.0.bias: torch.Size([16]), device=cpu
param_key=encoder.2.weight: torch.Size([32, 16, 3, 3]), device=cpu
param_key=encoder.2.bias: torch.Size([32]), device=cpu
param_key=encoder.4.weight: torch.Size([64, 32, 7, 7]), device=cpu
param_key=encoder.4.bias: torch.Size([64]), device=cpu
param_key=decoder.0.weight: torch.Size([64, 32, 7, 7]), device=cpu
param_key=decoder.0.bias: torch.Size([32]), device=cpu
param_key=decoder.2.weight: torch.Size([32, 16, 3, 3]), device=cpu
param_key=decoder.2.bias: torch.Size([16]), device=cpu
param_key=decoder.4.weight: torch.Size([16, 1, 3, 3]), device=cpu
param_key=decoder.4.bias: torch.Size([1]), device=cpu
GPU module
param_key=encoder.0.weight: torch.Size([16, 1, 3, 3]), device=cuda:0
param_key=encoder.0.bias: torch.Size([16]), device=cuda:0
param_key=encoder.2.weight: torch.Size([32, 16, 3, 3]), device=cuda:0
param_key=encoder.2.bias: torch.Size([32]), device=cuda:0
param_key=encoder.4.weight: torch.Size([64, 32, 7, 7]), device=cuda:0
param_key=encoder.4.bias: torch.Size([64]), device=cuda:0
param_key=decoder.0.weight: torch.Size([64, 32, 7, 7]), device=cuda:0
param_key=decoder.0.bias: torch.Size([32]), device=cuda:0
param_key=decoder.2.weight: torch.Size([32, 16, 3, 3]), device=cuda:0
param_key=decoder.2.bias: torch.Size([16]), device=cuda:0
param_key=decoder.4.weight: torch.Size([16, 1, 3, 3]), device=cuda:0
param_key=decoder.4.bias: torch.Size([1]), device=cuda:0
```

### GPU memory: limited resource

- Beware: GPUs have a limited amount of memory. Exceeding GPU memory will lead to your train run being killed with a "GPU out of memory" error!
- Question: when training a DNN model, what are the main uses of GPU memory?
  - Model weights. Ex: Linear's weight/bias parameters.
  - Intermediate activations. If your model has N Conv2d's, then there will be N activation feature maps that pytorch has to keep track of that uses up GPU memory!
    - Scales linearly with your batch\_size!
  - Gradients. Calculated during backwards()
  - Additional optimizer state. Ex: Adam requires `2\*num\_model\_params` additional values (gradient moving avg, squared gradient moving avg)

### Training model on GPU: Change 1/2

(post net.to) conv1.weight.device: cuda:0

- Taking a CPU pytorch training code and migrating it to the GPU is (often) very easy, a one-line(s) change!
- Change 1: Move model (eg its model parameters) to the GPU device

```
# Create model (on CPU first, by default)
net = Net(hidden num chans=model hidden num chans)
# Move model to GPU (if available to pytorch)
device_gpu_maybe = torch.device("cuda:0") if torch.cuda.is_available() else torch.device("cpu")
print(f"(pre net.to) GPU max_memory_allocated: {torch.cuda.max_memory_allocated() / 1e6} MB")
net_gpu_maybe = net.to(device=device_gpu_maybe)
print(f"(post net.to) GPU max_memory_allocated: {torch.cuda.max_memory_allocated() / 1e6} MB")
                                                                                           Note:
(pre net.to) conv1.weight.device: cpu
(pre net.to) GPU max memory allocated: 0.0 MB
(post net.to) GPU max_memory_allocated: 0.481792 MB
```

`torch.cuda.max\_memory\_allocated()` tells us how much GPU memory we've used. This tells us that our model parameters takes up 0.48 MB of GPU memory. Neat!

# Training model on GPU: Change 2/2

 Change 2: move all model inputs (including targets/labels!) to the GPU before calling forward

device"

```
def train_model(model: torch.nn.Module, optimizer, criterion, trainloader, num_epochs: int, device:
torch.device) -> torch.Tensor:
    for epoch in range(num_epochs): # loop over the dataset multiple times
        running loss = 0.0
        for ind_batch, data in enumerate(trainloader, 0):
            # get the inputs; data is a list of [inputs, labels]
            # Note: dataloader outputs inputs, labels as CPU tensors
            inputs, labels = data
            inputs = inputs.to(device=device)
                                                    'device' is our GPU device (or CPU
            labels = labels.to(device=device)
                                                     device if we don't have a GPU!)
            # zero the parameter gradients
            optimizer.zero_grad()
            # forward + backward + optimize
                                                      Since 'model' is on GPU, and
                                                      inputs, labels are on GPU, we
            outputs = model(inputs)
                                                     won't have errors like "mismatch
            loss = criterion(outputs, labels)
```

Implementation tip: this code is "device agnostic", in that it works for both CPU and GPU contexts (just pass in device=torch.device("cpu") or device=torch.device("cuda:0")).

This is the way!

# Demo: pytorch CPU vs GPU (single GPU)

Demo: gpu\_train\_example.py

```
(venv) PS
C:\Users\Eric\teaching\data_c182_fall2024\src\lectures\lec
ture19> python .\gpu_train_example.py
Files already downloaded and verified
Files already downloaded and verified
batchsize=64, model hidden num chans=128,
num_dataloader_workers=2
(CPU) Begin training (120078 model params)
(CPU) Finished Training (33.91305661201477 secs,
1474.3584033733462 imgs/sec)
(pre net.to) GPU max_memory_allocated: 0.0 MB
(post net.to) GPU max_memory_allocated: 0.481792 MB
(GPU) Begin training (120078 model params)
(post train_model) GPU max_memory_allocated: 96.306688 MB
(GPU) Finished Training (9.31104588508606 secs,
5369.966018542273 imgs/sec)
```

Device	Batchsize	Train throughput
CPU	64	1474 imgs/sec
GPU	64	5369 imgs/sec

# Demo: pytorch CPU vs GPU (single GPU)

- Interestingly: the CPU doesn't always outperform the GPU! (here it does, but there are settings where it doesn't)
- A few rules of thumb:
  - Model should be big enough
  - Batchsize needs to be big enough too!
- Reason: too small model/batchsize means you spend most of your time doing CPU<->GPU communication, leading to poor GPU utilization
- GPU's ideal computing mode: operate on large data "all at once" (aka large batches), rather than small data one-at-a-time.

Device	Batchsize	Train throughput
CPU	64	1474 imgs/sec
GPU	64	5369 imgs/sec
CPU	4	1350 imgs/sec
GPU	4	1633 imgs/sec
CPU	2	860 imgs/sec
GPU	2	937 imgs/sec

### Multi-GPU

- When working with large models + large datasets, we want to keep scaling up, ideally by throwing more compute (aka hardware, aka \$) at the problem
  - Horizontal scaling: add more compute (machines, GPUs)
  - Vertical scaling: make each individual GPU faster
- Both are important! But at a given time, horizontal scaling is quicker+easier to do

### Multi-GPU

- Fortunately, we can easily\* attach multiple GPUs to a single machine at a time
- Example: AWS EC2 p4d.24xlarge cloud instance type has 8 NVIDIA A100 Tensor Core GPUs, each with 40 GB GPU memory (quite large as of 2024!)
  - This is what many people in industry use to train large models (including my team). As of 2024-11, costs ~\$290k per year to rent one!
- Engineering Question: how to effectively utilize multiple GPUs on a single machine for training DNN models?
- For simplicity: assume that all GPUs on a single machine are all the same type, eg same exact model (in practice this is true 99% of the time)

### Multi-GPU: Scenario 1

- Suppose our machine has 8 GPUs, and our DNN model (and activations) can fit on a single GPU.
- Question: if we wanted to maximize training throughput, what's one way we can utilize the 8 GPUs?
- Answer: load the model onto all 8
   GPUs separately, and have each
   GPU do their own forward/backward
   passes in parallel!

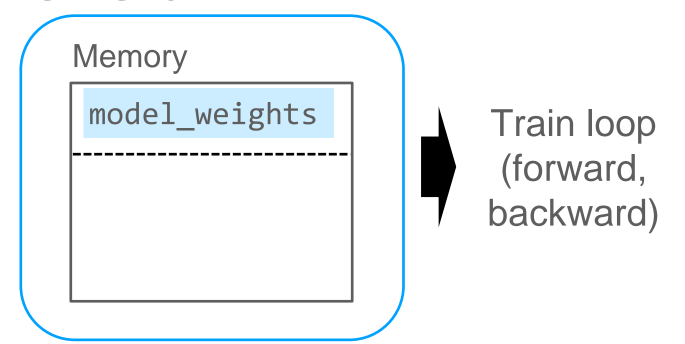
Memory GPU:0 model\_weights Train loop Memory (forward, backward) model\_weights GPU:2 Memory model\_weights Train loop (forward, backward) GPU:3 Memory model\_weights Train loop (forward, backward)

GPU:1

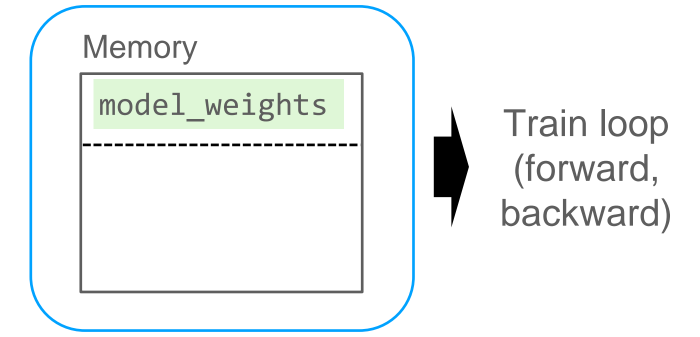
Implementation Note: each GPU trainer gets its own training Dataloader. Take care to ensure that each Dataloader splits up the training dataset appropriately (ex: don't want all N GPU workers to train on the same batches!)

### Multi-GPU: Scenario 1

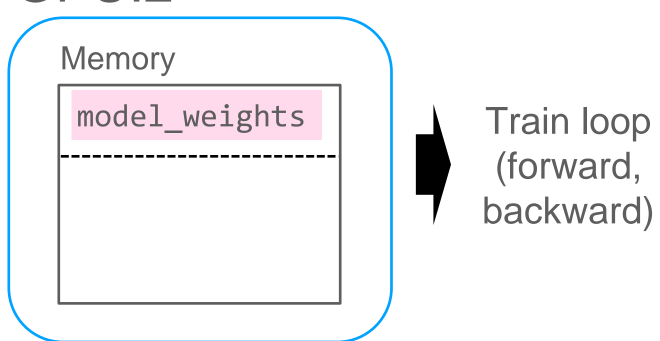
#### GPU:0



#### GPU:1



#### GPU:2



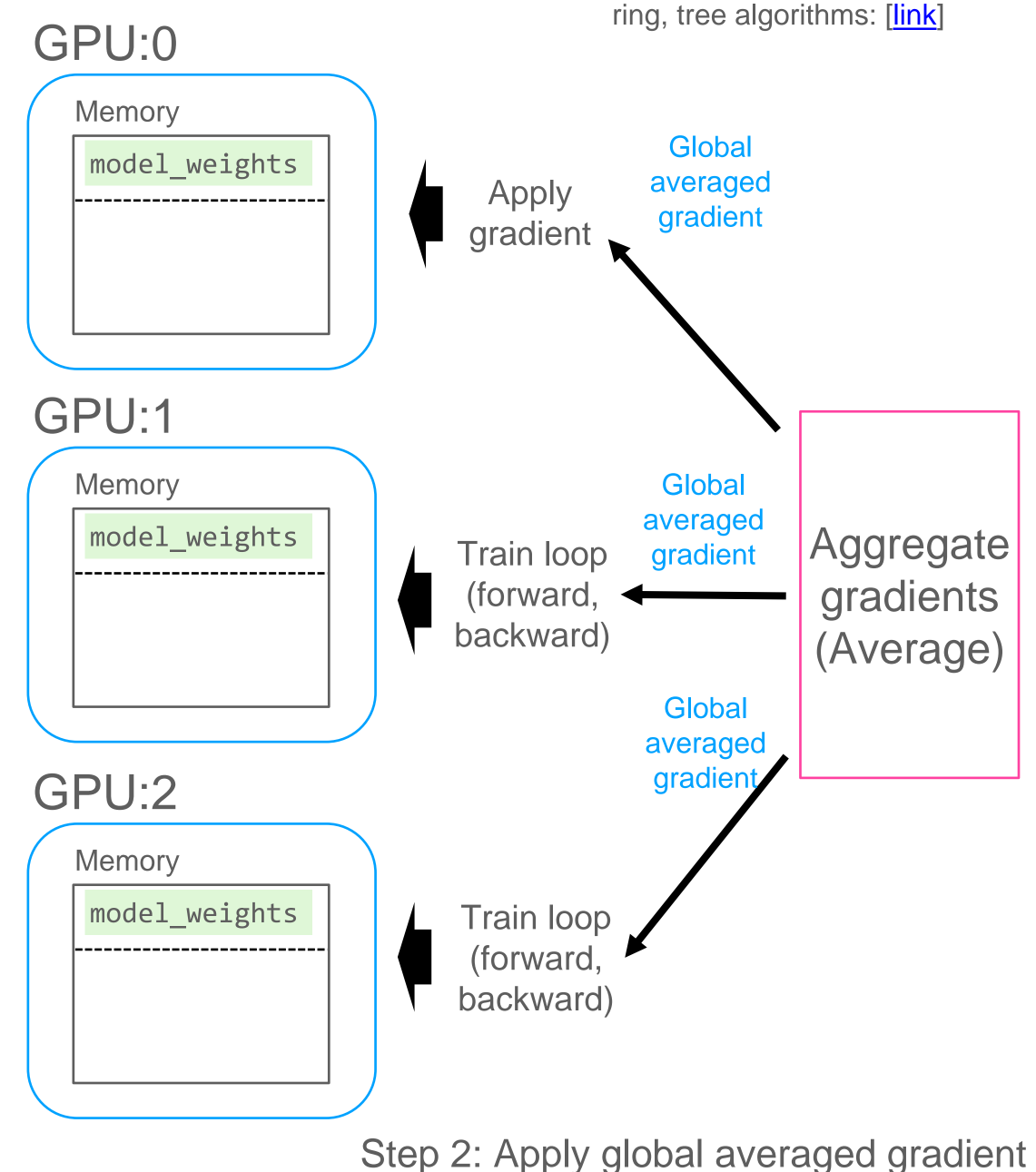
Question: suppose each GPU worker has different training dataset splits (aka each GPU worker operates on different train batches).

After N training batches, will the model parameters be the same across all GPU workers?

Answer: Nope! Each GPU worker's train loop will process different batches, thus the gradients for GPU:0 will be different than GPU:1, resulting in different model weights for each worker. This complicates things!
Question: how can we ensure that, after each training batch (for all N GPUs), the model\_weights are the same ("in sync") for all N GPUs?

Answer: one popular way is to aggregate all of the GPU worker's gradients (average), then have each GPU worker apply the same aggregated ("global average") gradient. Ensures that model\_weights is the same on all GPU workers!

Multi-GPU: Aggregate gradients GPU:0 Memory **Gradient for** model\_weights Train loop GPU:0 (forward, backward) GPU:1 Memory **Gradient for** model\_weights Aggregate GPU:1 Train loop gradients (forward, backward) (Average) **Gradient for** GPU:2 GPU:2 Memory model\_weights Train loop (forward, backward) Step 1: Aggregate incoming gradients



Note: there exist other more efficient,

cleverer ways of optimizing this. Ex: NCCL's

# Pytorch: DistributedDataParallel (DDP)

- Enter: torch.nn.parallel.DistributedDataParallel (or DDP) [link]
- DDP does exactly what we proposed! Consists of two parts:
  - Model copy. copy the initial model weights to each GPU, via `.to(device)`
  - Gradient synchronization. adds a "hook" to the backwards() pass to calculate the aggregated gradients across all GPUs (via averaging)
    - This ensures that all GPUs use the same gradients for each step, which ensures that each GPU always has the same model weights
- Each GPU processes their own batches independently
- For 8 GPUs, you've effectively increased your batchsize by 8x!

### Terminology: world\_size, rank

- world\_size: the total number of workers. Ex: total number of GPUs
- rank: an integer between [0, world\_size-1] (inclusive).
- Setup: each GPU will be assigned a "rank" and the "world\_size", and will use this metadata to perform its job
- Ex: Suppose world\_size is 8.
- A GPU worker assigned rank=0 will use GPU0: `torch.device("cuda:0")`
  - Rank=1 will use GPU1: `torch.device("cuda:1")`
  - •
  - Rank=7 will use GPU7: `torch.device("cuda:7")`

# DDP in pytorch

if \_\_name\_\_ == "\_\_main\_\_":

demo\_basic()

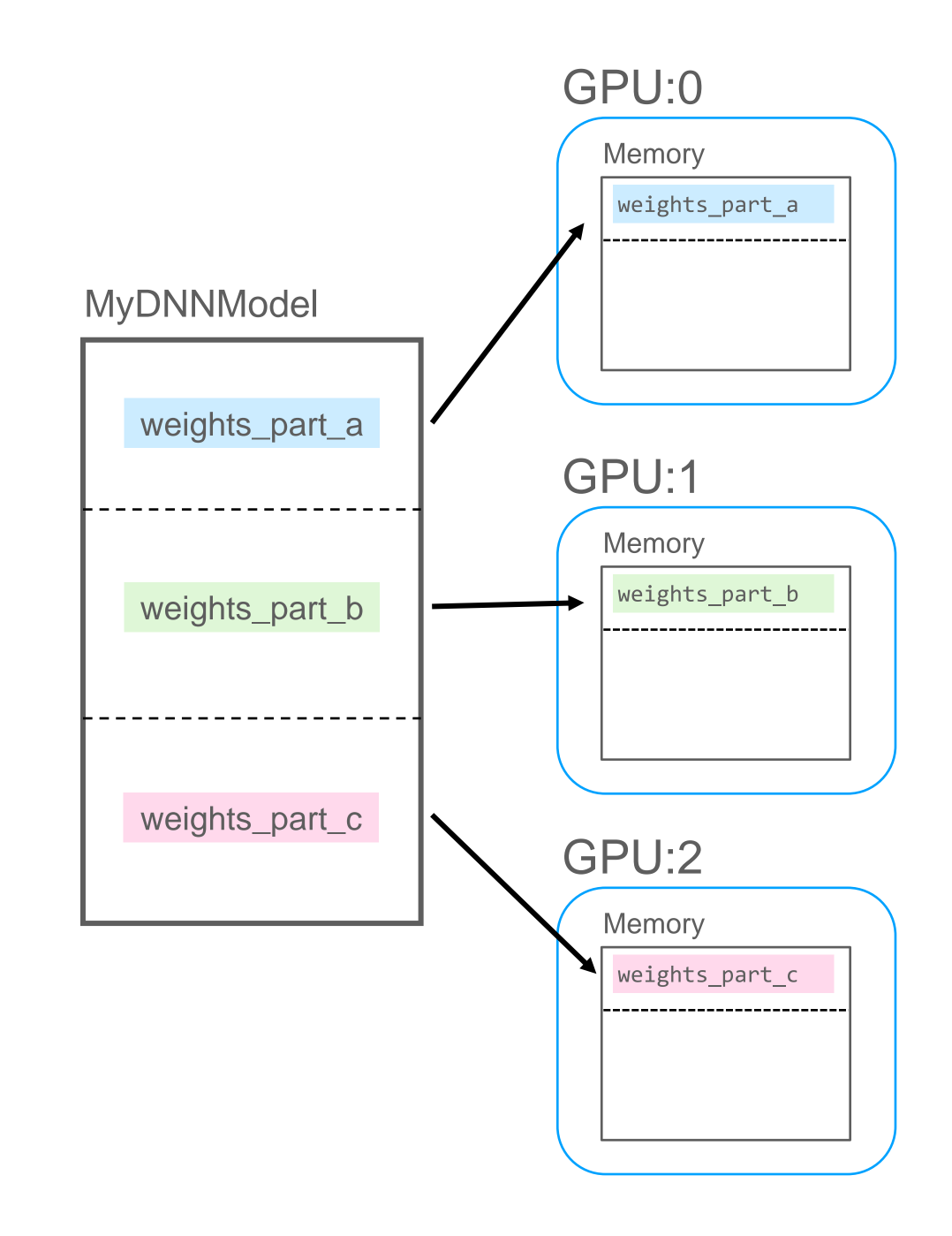
```
import torch.distributed as dist
import torch.nn as nn
                                                                      # use torch_elastic to launch script N times, one
import torch.optim as optim
from torch.nn.parallel import DistributedDataParallel as DDP
                                                                      # for each GPU worker
class ToyModel(nn.Module):
                                                                      # --nnodes: number of nodes. For us, 1 machine
   def __init__(self):
                                                                      # --nproc_per_node: number of GPUs. For us: 8
      super(ToyModel, self).__init__()
      self.net1 = nn.Linear(10, 10)
                                                                      torchrun --nnodes=1 --nproc_per_node=8 --rdzv_id=100 --
      self.relu = nn.ReLU()
                                                                      rdzv_backend=c10d --rdzv_endpoint=$MASTER_ADDR:29400
      self.net2 = nn.Linear(10, 5)
   def forward(self, x):
                                                                      elastic_ddp.py
      return self.net2(self.relu(self.net1(x)))
def demo_basic():
                                                                                Boilerplate code to initialize
    dist.init_process_group("nccl")
                                                                                distributed backend (NCCL)
    rank = dist.get rank()
    print(f"Start running basic DDP example on rank {rank}.")
    # create model and move it to GPU with id rank
                                                                                `device_id` is the assigned GPU
    device_id = rank % torch.cuda.device_count()
                                                                               device to use: f"cuda:{device_id}"
    model = ToyModel().to(device_id)
                                                                               Wrap model on DDP()
    ddp_model = DDP(model, device_ids=[device_id])
    loss fn = nn.MSELoss()
    optimizer = optim.SGD(ddp_model.parameters(), lr=0.001)
                                                                                Can use 'ddp_model' just like
    optimizer.zero_grad()
                                                                               regular 'model'! Very convenient.
    outputs = ddp_model(torch.randn(20, 10))
    labels = torch.randn(20, 5).to(device id)
    loss_fn(outputs, labels).backward()
    optimizer.step()
    dist.destroy_process_group()
    print(f"Finished running basic DDP example on rank {rank}.")
                                                                                             For more info, see: https://pytorch.org/tutorials/intermediate/ddp_tutorial.html
```

### DDP: sync and all\_reduce (NCCL)

- An important part of DDP is the gradient synchronization
- For N GPUs, each GPU will independently perform their own "local" forward/backwards pass to calculate N different gradient updates
- DDP's job is to take all N gradient updates, average them, then transmit this aggregated gradient to all N GPUs so that each GPU can perform their weight updates.
- Common name for this operation: "all\_reduce"
- Pytorch supports dispatching the all\_reduce call to a variety of libraries [link]
  - For Nvidia GPUs: use NCCL [link]

### Multi-GPU: Scenario 2

- Suppose our machine has 8 GPUs, and our DNN model (and activations) can't fit on a single GPU.
- Question: how can we utilize the 8 GPUs to train our DNN model?
- Answer: split the model up across multiple GPUs!
- Question: what downsides can you think of?
- Answer: slower training, due to additional cross-GPU communication



# Pytorch: FullyShardedDataParallelism (FSDP)

- Pytorch's FSDP implements this "model sharding" idea [link]
- Implementation challenge: how to distribute model parameters (eg layers) in a way that minimizes cross-gpu device copies?
- Heuristic: FSDP works well for very large models, eg when model weights are a substantial fraction of available GPU memory
- FSDP is also handy to increase total batchsize at the expense of some efficiency

### DDP vs FSDP

Name	Impact on: GPU memory	Impact on: Train throughput	When to use?
DistributedDataParallel (DDP)	For N GPUs, model weights are duplicated N times (waste). Con!	Each GPU does forward/backward independently: no cross-GPU communication required (except for all_reduce). Pro!	If your model+activations comfortably fits in GPU memory AND you are happy with your current batch_size
FullyShardedDataParallel (FSDP)	Model weights are instantiated only once (split across N GPUs).  Pro!	Training throughput is worse due to cross-GPU communication. Con!	If your model can't fit on a single GPU: MUST use FSDP. Or: if you need a higher batchsize but it won't fit with DDP: try FSDP!

### Multi-Node, Multi-GPU: DDP

- Suppose we have M machines, each with N GPUs. How to effectively utilize this for training DNN models?
  - Fortunately, the earlier principles generalize quite nicely!

#### DDP:

- M=1 machine, N GPUs: all\_reduce across GPUs (within a single machine)
- M>1 machines, N GPUs each: all\_reduce across ALL GPUs
  - Involves cross-machine communication during gradient sync!

#### FSDP

- M=1 machine, N GPUs: split model parameters across N GPUs
- M>1 machines, N GPUs each: split model parameters across M\*N GPUs
  - Involves cross-machine comm during forward/backward

# DDP in pytorch (multi-node)

```
import torch
import torch.distributed as dist
import torch.nn as nn
import torch.optim as optim
from torch.nn.parallel import DistributedDataParallel as DDP
class ToyModel(nn.Module):
   def __init__(self):
       super(ToyModel, self).__init__()
       self.net1 = nn.Linear(10, 10)
       self.relu = nn.ReLU()
       self.net2 = nn.Linear(10, 5)
   def forward(self, x):
       return self.net2(self.relu(self.net1(x)))
def demo_basic():
    dist.init process group("nccl")
    rank = dist.get_rank()
    print(f"Start running basic DDP example on rank {rank}.")
    # create model and move it to GPU with id rank
    device id = rank % torch.cuda.device count()
    model = ToyModel().to(device_id)
    ddp_model = DDP(model, device_ids=[device_id])
    loss fn = nn.MSELoss()
    optimizer = optim.SGD(ddp_model.parameters(), lr=0.001)
    optimizer.zero_grad()
    outputs = ddp_model(torch.randn(20, 10))
    labels = torch.randn(20, 5).to(device_id)
    loss_fn(outputs, labels).backward()
    optimizer.step()
    dist.destroy_process_group()
    print(f"Finished running basic DDP example on rank {rank}.")
if __name__ == "__main__":
    demo basic()
```

```
# use torch_elastic to launch script N times, one
# for each GPU worker
# --nnodes: number of nodes. For us, 4 machines
# --nproc_per_node: number of GPUs. For us: 8
torchrun --nnodes=4 --nproc_per_node=8 --rdzv_id=100 --
rdzv_backend=c10d --rdzv_endpoint=$MASTER_ADDR:29400
elastic_ddp.py
```

Now, `rank` spans multiple machines. Ex: if each machine has 8 GPUs, then: Rank=[0, 1, ..., 7]: Machine 0 Rank=[8, 9, ..., 15]: Machine 1

Other than that, everything is nearly the same! Pytorch set things up nicely so that very little has to change when scaling from M=1 machines to M=4 machines. The world\_size/rank abstraction makes things very convenient:)

### Example: "Train ImageNet in 1 hour"

- In "Accurate, Large Minibatch SGD: Training ImageNet in 1 Hour" (2018) [link] the authors, using a large distributed GPU cluster, trained a ResNet-50 model on ImageNet-1k "from scratch" in one hour. A neat accomplishment for that time!
  - Hardware: 256 GPUs ("Big Basin" [link] GPU cluster internal to Facebook, 16GB GPU mem per card. Nvidia Tesla P100).
- Distributed training setup: basically DDP (they used Caffe2, not pytorch, but same idea)

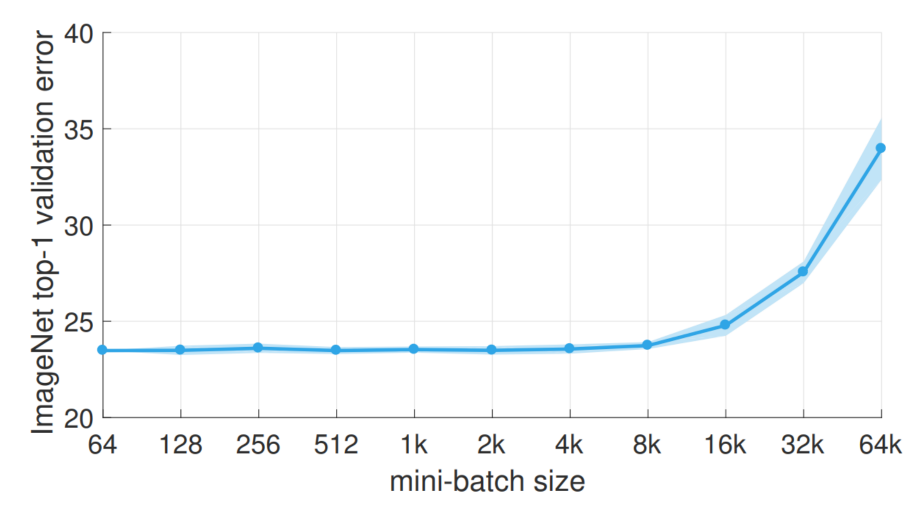


Figure 1. **ImageNet top-1 validation error** *vs.* **minibatch size.** Error range of plus/minus *two* standard deviations is shown. We present a simple and general technique for scaling distributed synchronous SGD to minibatches of up to 8k images *while maintaining the top-1 error of small minibatch training*. For all minibatch

### Learning rate: Linear scaling rule

- Learning: when scaling up the number of GPUs (aka increasing the effective batchsize), one must adjust the learning rate accordingly ("linear scaling rule").
  - Rule: double the batchsize -> double the learning rate.

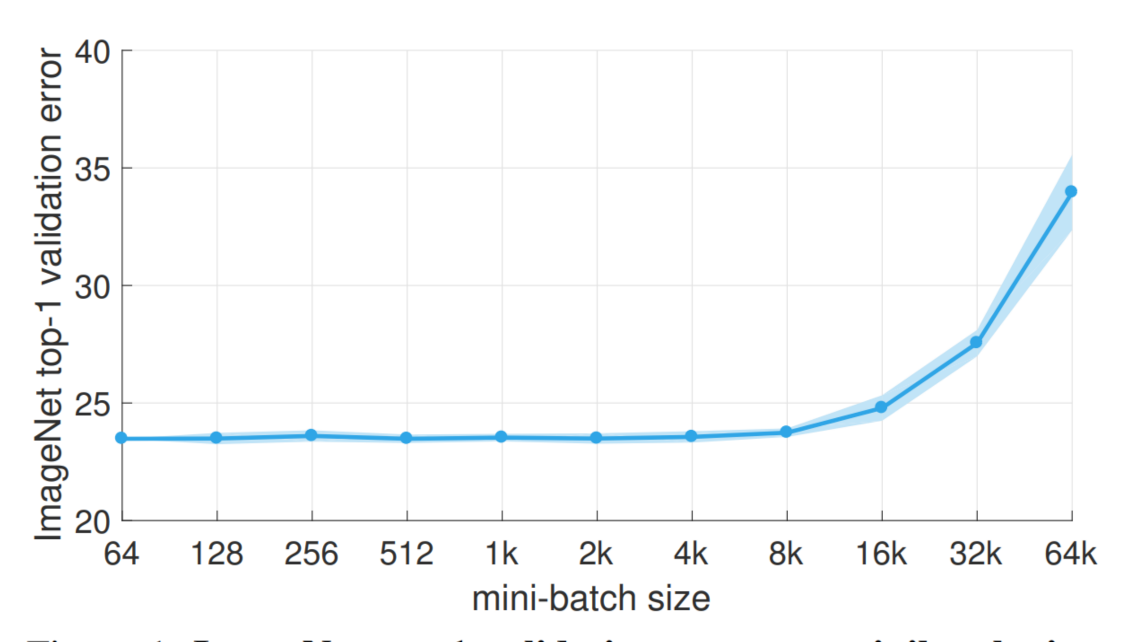


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# Gradient quality vs num steps?

- Observation: if you keep the number training epochs fixed, then increasing the batchsize leads to fewer model updates.
  - Higher batchsize -> higher quality gradient updates, but fewer parameter updates
  - Lower batchsize -> noisier gradient updates, but more parameter updates.
- What is best? Paper's answer: higher batchsize AND higher learning rate.
  - ...to a point. Beyond batchsize=8k, classification error starts increasing.

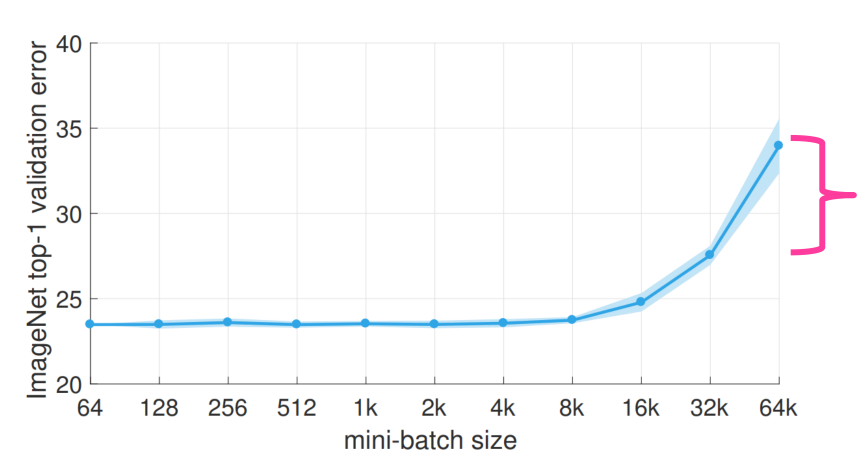


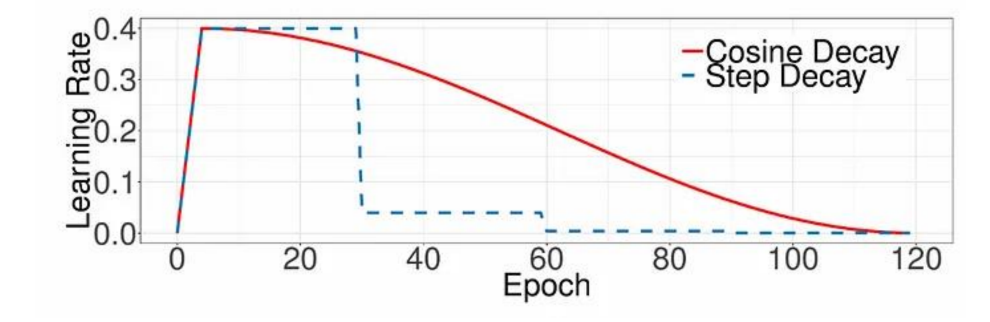
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### "Train ImageNet in 1 hour": Hardware advances

- A sign that GPU hardware (and DNN libraries + distributed training frameworks)
  is advancing quickly
- ...And, a hint that ImageNet-1k is starting to feel small!
- (later in Aug 2018, someone showed we can train ImageNet in 18 mins for \$40 using AWS cloud! [link])
- In 2024, I bet things are even faster + cheaper! Technology marches on...

### Tangent: Learning rate schedules

- So far in this class, we've used a single learning rate. In practice, it's better to use learning rate schedules
- Start learning rate small, then gradually ramp it up to a larger value (eg the first ~100 iterations)
  - Intuition: starting learning rate too high often leads to training divergence (eg NaN losses). Thus, we start it low to get the model weights in a "healthy" region, then slowly increase the learning rate
- Over the course of training, decay the learning rate
  - Intuition: during early parts of training, model needs to make big steps (high LR). But, near the end of training, model is focusing on finer-grained details (small LR).



# (unused) CUDA example: element-wise vector addition

Consider: elementwise-vector addition. Given two vectors x, y (with shape=[N]), output (x+y). This is an "embarrassingly parallel" problem: chunk up the input vectors into K chunks, and process each chunk independently in parallel!

```
__global__
void add(int n, float * x, float * y) {
  int index = threadIdx.x;
  int stride = blockDim.x;
  for (int i = index; i < n; i += stride)
    y[i] = x[i] + y[i];
}</pre>
```

Ex: for vector element-wise addition, one way to do it ("embarassingly parallel"):

- Break the input vector into numBlocks chunks
- within each block, have a separate thread perform a single `y[i] = a[i] + b[i]`

Here, we divide the input array into blocks. Within each block, we assign threads to each block element.

