

# TRAINING DEEP LEARNING MODELS WITH FULLY SHARDED DATA PARALLEL

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# **SUMMARY**

- Parallelism Strategies
- Fully Sharded Data Parallel (FSDP)
- Algorithm Overview
- Communication Optimizations
- Model Initialization
- Sharding Strategies

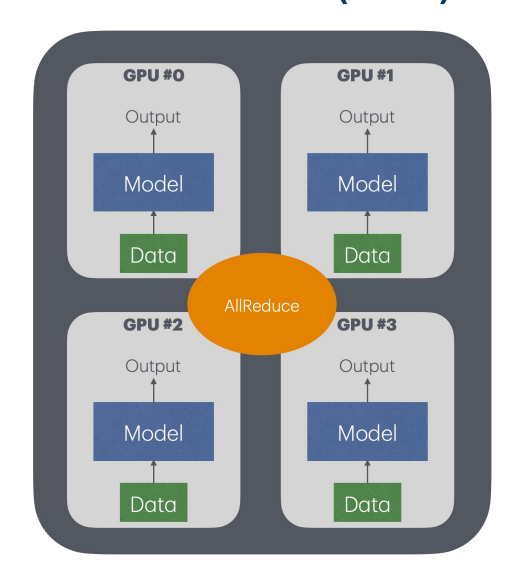


# **SUMMARY**

- FSDP Interoperability
- Limitations
- Conclusions
- Reference

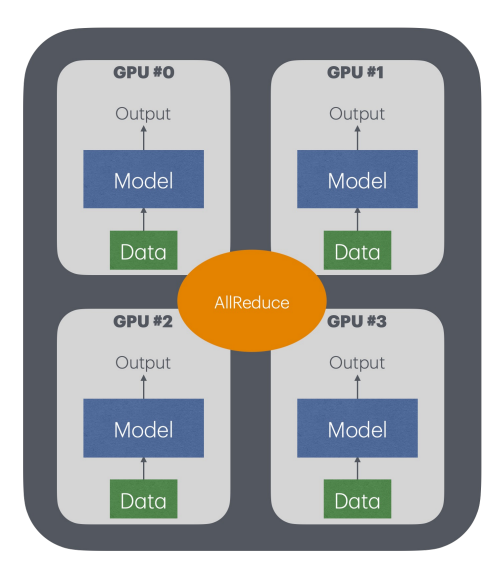


# DISTRIBUTED DATA PARALLEL (DDP)





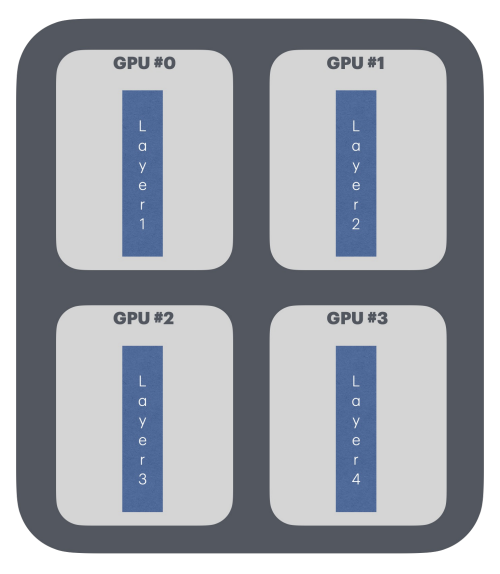
# DISTRIBUTED DATA PARALLEL (DDP)



- Inadequate for supporting large models
- Likely encounter out-of-memory errors on each device

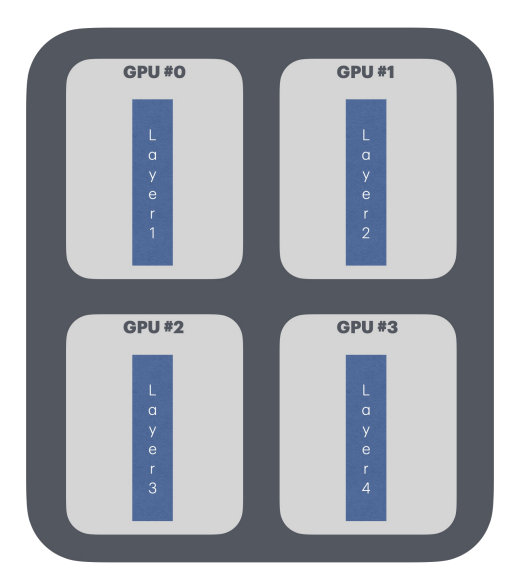


# PIPELINE PARALLELISM





# PIPELINE PARALLELISM



 The entire model must be structured in sequential layers or stages, which might not be feasible for all types of models.



# OTHER PARALLELISM STRATEGIES

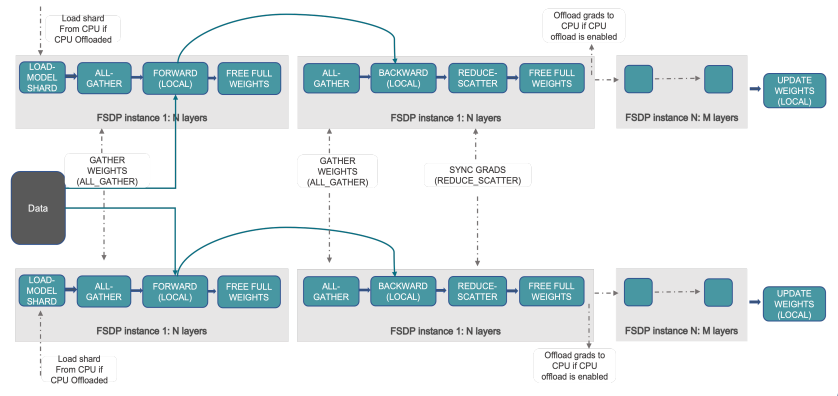
- Tensor Parallelism
- Megatron-LM
- MiCS parallelism

• ..



# **FULLY SHARDED DATA PARALLEL (FSDP)**

- Distributed training technique used to scale deep learning model training across multiple GPUs by sharding the model's parameters.
- Integrates efficiently with PyTorch and optimizes communication and memory usage, making it effective for training extremely large models in a distributed environment.





# **ALGORITHM OVERVIEW**

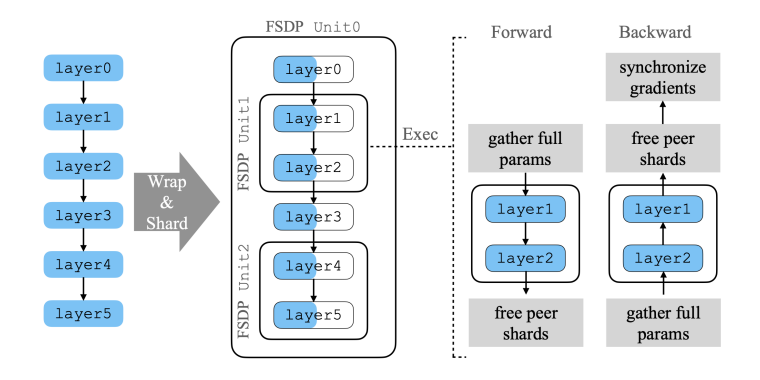
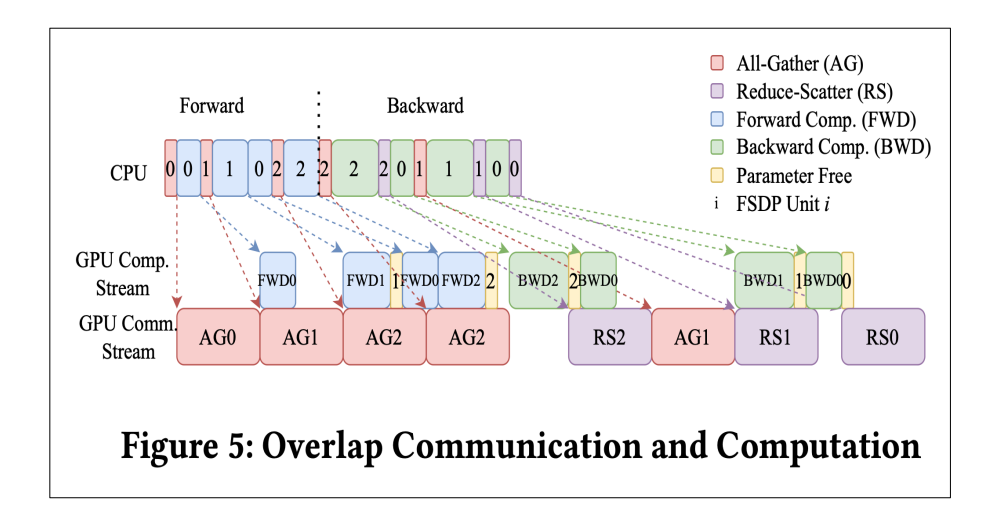
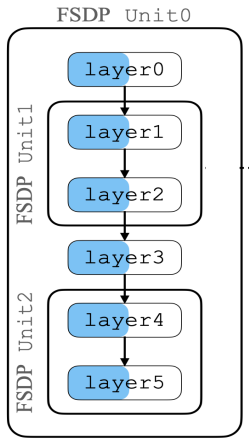


Figure 1: FSDP Algorithm Overview



# **COMMUNICATION OPTIMIZATIONS**







## **DEFERRED INITIALIZATION**

#### **Challenge #1:**

PyTorch required full materialization of the entire model on one device, how to create a model instance
without materializing any tensor storage, postponing initialization until a storage on a concrete device is
attached to the tensor.



#### Solution #1:

- Model parameters are initially allocated on a simulated or "fake" device.
- All parameter initialization operations are recorded during this simulated phase.
- When parameters move to a GPU, recorded initialization operations are replayed.
- This method creates a model instance without consuming GPU memory initially.



# **MODEL INITIALIZATION**

#### **Challenge #2**

 How to ensure accurate initialization of model parameters when the model is too large to fit on a single GPU.



#### Solution #2:

- FSDP initializes and shards one unit at a time.
- With deferred initialization, FSDP decomposes the model into units, moves each unit to a GPU sequentially, and replays tensor initialization operations.



# HANDS-ON CODING



# **GOOD PRACTICE**

Always store your code in the project folder.

```
mkdir /p/project/training2426/$USER
cd /p/project/training2426/$USER
git clone https://github.com/sab148/nxtaim-distributed-models
```

• Store data in the scratch directory for faster I/O access. Files in scratch are deleted after 90 days of inactivity.

ls /p/scratch/training2426/

- Store the data in `\$DATA\_dataset` for a more permanent location.
  - This location is not accessible by compute nodes.
  - You have to Join the project in order to store and access data <a href="https://judoor.fz-juelich.de/projects/datasets/">https://judoor.fz-juelich.de/projects/datasets/</a>



# LET'S PARALLELIZE THE CODE WITH DISTRIBUTED DATA PARALLEL (DDP)



#### Remove line 180 and add this code snippet at line 172

```
torch.distributed.init_process_group(backend='nccl')
local_rank = int(os.getenv('LOCAL_RANK'))
rank = int(os.environ['RANK'])
device = torch.device('cuda', local_rank)
torch.cuda.set_device(device)
torch.random.manual seed(args.seed + torch.distributed.get rank())
```



#### Add this snippet at line 111

```
train_sampler = torch.utils.data.distributed.DistributedSampler(
           train dset,
           shuffle=True,
           seed=args.seed,
       train_dset = torch.utils.data.DataLoader(
           train dset,
           batch_size=args.batch_size,
           sampler=train_sampler,
 Don'
           num_workers=args.train_num_workers,
forge
this lir
           pin_memory=True,
           persistent_workers=args.train_num_workers > 0,
```



```
# Ensures that each process gets a unique subset of the data.
valid_sampler = torch.utils.data.distributed.DistributedSampler(valid_dset)

valid_dset = torch.utils.data.DataLoader(
    valid_dset,
    batch_size=args.batch_size,
    # Uses the distributed sampler to ensure each process gets a different subset of the data.
    sampler=valid_sampler,
    num_workers=args.valid_num_workers,
    # Use pinned memory on GPUs for faster device—copy.
    pin_memory=True,
    persistent_workers=args.valid_num_workers > 0,
)
```

# Here as well

```
# Ensures that each process gets a unique subset of the data.
test_sampler = torch.utils.data.distributed.DistributedSampler(test_dset)

test_dset = torch.utils.data.DataLoader(
    test_dset,
    batch_size=args.batch_size,
    # Uses the distributed sampler to ensure each process gets a different subset of the data.
sampler=test_sampler,
    # Use pinned memory on GPUs for faster device-copy.
    pin_memory=True,
)
```



#### Add this snippet at line 210

```
# Wraps the model in a DistributedDataParallel (DDP) module to parallelize the training
# across multiple GPUs.
model = torch.nn.parallel.DistributedDataParallel(
    model,
    device_ids=[local_rank],
)
```



#### Add at line 227

# sets the current epoch for the dataset sampler to ensure proper data shuffling in each
epoch
train\_dset.sampler.set\_epoch(epoch)



#### Add this line of code at line 242 and line 176

```
# Obtain the global average loss.
torch.distributed.all_reduce(loss, torch.distributed.ReduceOp.AVG)
```



#### Add those utility functions at line 85

```
functools.lru_cache(maxsize=None)
def is_root_process():
    """Return whether this process is the root process."""
    return torch.distributed.get rank() == 0
def print0(*args, **kwargs):
    """Print something only on the root process."""
    if is_root_process():
        print(*args, **kwargs)
def save0(*args, **kwargs):
    """Pass the given arguments to `torch.save`, but only on the root
    process.
    if is root process():
        torch.save(*args, **kwargs)
```



#### Replace print with print0 at line 269

```
print0(f'[{epoch}/{args.epochs}; {i}] loss: {loss:.5f}')
```

#### Replace print with print0 at line 276

```
print0(f'[{epoch}/{args.epochs}; {i}] valid loss: {valid_loss:.5f}')
```

#### Replace print with print0 at line 290

```
print0('Finished training after', end_time - start_time, 'seconds.')
```

#### Replace print with print0 at line 296

```
print0('Final test loss:', test_loss)
```



#### Replace torch.save at line 283

```
# Replace torch.save function by the utility function to save the model.
save0(model, 'model-best.pt')
```

#### Replace torch.save at line 300

```
# Replace torch.save function by the utility function to save the model.
save0(model, 'model-final.pt')
```



#### Change number of node at line 3 in run\_to\_distributed\_training.sh file

```
#SBATCH --nodes=4
```

#### Add in run\_to\_distributed\_training.sh file at line 17



#### Add in run\_to\_distributed\_training.sh file at line 41

```
srun --cpu_bind=none bash -c "torchrun \
--nnodes=$SLURM_NNODES \
--rdzv_backend c10d \
--nproc_per_node=gpu \
--rdzv_id $RANDOM \
--rdzv_endpoint=$MASTER_ADDR:$MASTER_PORT \
--rdzv_conf=is_host=\$(if ((SLURM_NODEID)); then echo 0; else echo 1; fi) \
to_distributed_training.py "
```



# PARALLELIZE THE CODE WITH FULLY SHARDED DATA PARALLEL (FSDP)



#### Replace DistributedDataParallel with FullyShardedDataParallel at line 231

```
my_auto_wrap_policy = functools.partial(
   size_based_auto_wrap_policy, min_num_params=20000
model = fsdp.FullyShardedDataParallel(
   model,
   device id=local rank,
   auto_wrap_policy=my_auto_wrap_policy,
```



#### Add this function at line 85

```
def save_model(
    checkpoint_type,
    model,
    rank,
    save_dir='model-final.pt',
    optim dir="optimizer-final.pt",
    optimizer=None
):
    if checkpoint_type == 'full':
        model_checkpointing.save_model_checkpoint(model, save_dir, rank)
    if optimizer is not None:
        model_checkpointing.save_optimizer_checkpoint(model, optimizer, optim_dir, rank)
    elif checkpoint type == 'sharded':
        if optimizer is not None:
            model_checkpointing.save_model_and_optimizer_sharded(model, save_dir, rank,
    optim=optimizer)
        else:
            model_checkpointing.save_model_and_optimizer_sharded(model, save_dir, rank)
    elif checkpoint_type == 'local':
        model_checkpointing.save_distributed_model_checkpoint(model, save_dir, rank)
        if optimizer is not None:
            model_checkpointing.save_optimizer_checkpoint(model, optimizer, optim_dir, rank)
```



#### Replace save0 at line 312

```
# Replace the following line with the utility function to save the model.
save_model(args.save_model_opt, model, rank, "model-best.pt", "optimizer-best.pt",
optimizer=opt)
```

#### Replace save0 at line 329

```
# Replace the following line with the utility function to save the model. save_model(args.save_model_opt, model, rank, optimizer=opt)
```



#### Export TORCH\_LOGS at line 37 in run\_to\_distributed\_training.sh file

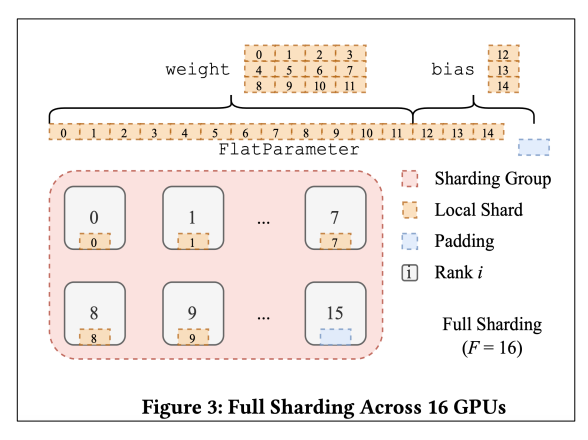
export TORCH\_LOGS='-torch.distributed.checkpoint.\_dedup\_tensors'



# **SHARDING STRATEGIES**



# **SHARDING STRATEGIES**



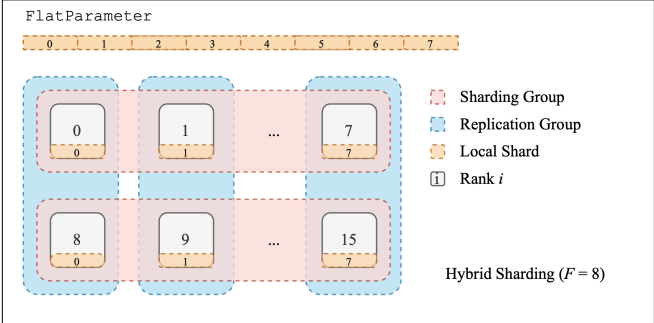


Figure 4: Hybrid Sharding on 16 GPUs: GPUs are configured into 2 sharding groups and 8 replication groups



# SHARDING STRATEGIES

#### Sharding strategies for distributed training by FullyShardedDataParallel

- FULL\_SHARD: Parameters, gradients, and optimizer states are sharded. Default strategy.
- SHARD\_GRAD\_OP: For the parameters, this strategy unshards before the forward, does not reshard them after the forward, and only reshards them after the backward computation
- NO SHARD: Similar to PyTorch's DistributedDataParallel API.
- HYBRID\_SHARD: Apply FULL\_SHARD within a node, and replicate parameters across nodes.
- \_HYBRID\_SHARD\_ZERO2: Apply SHARD\_GRAD\_OP within a node, and replicate parameters across nodes. This is like HYBRID\_SHARD, except this may provide even higher throughput since the unsharded parameters are not freed after the forward pass, saving the all-gathers in the pre-backward.



#### Pass this argument at line 260

```
# Wraps the model in FullyShardedDataParallel (FSDP) module using the specified GPU and
automatic wrapping policy for submodules.
model = fsdp.FullyShardedDataParallel(
    model,
    device_id=local_rank,
    auto_wrap_policy=my_auto_wrap_policy,
    sharding_strategy=fsdp.ShardingStrategy.HYBRID_SHARD
)
```



### **FSDP INTEROPERABILITY**

#### **Combination with Pipeline Parallelism**

- Integration: FSDP can wrap each stage of a pipeline parallel model.
- Challenge with Micro-Batches: Pipeline parallelism involves splitting input batches into micro-batches, requiring frequent unsharding and resharing of model parameters, which can lead to high communication overhead.
- **Optimization**: FSDP offers alternative sharding strategies that keep parameters unsharded post-forward pass, reducing the need for AllGather communications after each micro-batch.
- **Memory Efficiency**: Even though entire pipeline stage parameters are stored on the GPU device, FSDP minimizes memory usage by still sharding gradients and optimizer states.



#### **FSDP INTEROPERABILITY**

#### **Combination with Tensor Parallelism**

- **Distinctive Approach**: Unlike FSDP, tensor parallelism maintains parameters sharded during computations to fit large sub-modules within GPU memory constraints.
- 2D Parallelism in PyTorch: PyTorch's parallelize\_module feature can create a 2D parallel structure by combining tensor parallelism and FSDP.
- **Device Organization**: Devices are organized into a 2D mesh, with DTensor managing tensor parallelism on one dimension (typically intra-node for higher bandwidth) and FSDP handling sharded data parallelism on the other dimension (inter-node).
- Communication Strategy: This setup optimizes communication paths, keeping tensor-parallel operations within the node and FSDP communications between nodes to efficiently manage network bandwidth and computational delays.



# **LIMITATIONS**

#### **Mathematical Equivalence:**

- Challenges with Optimizer Computation: FSDP's sharding mechanism can alter the original data layout of parameters, impacting computations that depend on unsharded values, such as vector norms or complex optimizer functions.
- **Impact on Optimizer Efficiency**: Using padding, uneven sharding, or extra communication to maintain optimizer accuracy can negatively affect performance.
- **Research Area**: Properly integrating sophisticated optimizer computations with parameter sharding remains an open and active area of research.

#### **Shared Parameters:**

- **Management of Shared Parameters**: FSDP must avoid flattening shared parameters into multiple segments to prevent errors related to tensor storage or size mismatches.
- **Strategic Sharding**: It's recommended to structure FSDP units so that shared parameters are always accessible and unsharded across all uses by placing them in the lowest-common-ancestor unit.
- Ongoing Improvements: Methods to handle shared parameters more efficiently are under investigation to optimize their management without extended periods of being unsharded.

# CONCLUSIONS

- FSDP is a distributed training method in PyTorch that shards model parameters and optimizer states across GPUs to maximize memory efficiency and scalability.
- FSDP enables model initialization on a dummy device, which is then replayed on GPUs, supporting large models. This is called deferred initialization.
- FSDP decomposes the model into units and materializes the parameters of one unit at a time.
- FSDP offers different sharding strategies.
- FSDP can be combined with pipeline parallelism or tensor parallelism.



# **REFERENCE**

• Zhao, Yanli, et al. "Pytorch fsdp: experiences on scaling fully sharded data parallel." arXiv preprint arXiv:2304.11277 (2023).



# Thank you!

