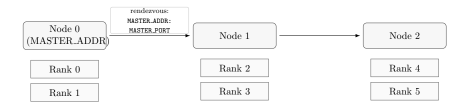
# Multi-Node CPU DDP torchrun on SLURM

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Goal. In this lab, you will run distributed CPU training across multiple SLURM nodes using torch.distributed (Gloo backend), torchrun, and HuggingFace Trainer. You will stage dependencies for offline nodes, verify per-node environments, and interpret structured logs to confirm ranks, world size, and correct rendezvous.

The codes are updated: train\_tiny.py, side\_shell\_pr.sh, and req.sh. This handout explains how they work together, why each design choice was made, and how to validate every step.

# 1 Architecture Overview (What Runs Where)



WORLD\_SIZE = #nodes  $\times$  NPROC\_PER\_NODE. For 3 nodes and 2 procs/node: WORLD\_SIZE = 6. Ranks are global (0..WORLD\_SIZE-1). Each rank runs the same training loop (DDP).

Figure 1: Ranks per node, rendezvous, and world size.

- Hosts. Each "Node" is a SLURM node. The leftmost node is chosen as the rendezvous server: we export MASTER\_ADDR and MASTER\_PORT; PyTorch's c10d store (Gloo on CPU) uses these to let all processes find each other (init\_method="env://").
- Ranks beneath On each node, torchrun spawns NPROC\_PER\_NODE worker processes. Each worker is a rank with a unique global RANK in 0..WORLD\_SIZE 1. In the example: Node 0 has ranks 0-1, Node 1 has 2-3, Node 2 has 4-5.
- World size. WORLD\_SIZE = nnodes  $\times$  nproc\_per\_node. For 3 nodes  $\times$  2 procs/node, WORLD\_SIZE = 6.
- Which IDs exist? Each process gets:
  - RANK (global, unique across all nodes),
  - LOCAL\_RANK (index of the process on its node:  $0..NPROC\_PER\_NODE 1$ ),
  - NODE\_RANK (which node this is: 0..nnodes 1).

These are set by torchrun via environment variables and consumed by init\_process\_group.

• What runs on each rank? train\_tiny.py executes identically on every rank: it loads the tokenizer and data shard, builds the tiny BERT, and performs a forward/backward step. At the end of each step, DDP (via Gloo) averages gradients across all ranks, so the model stays in sync. Only rank 0 saves checkpoints.

• Arrows between nodes. They indicate the DDP control/data connections established after rendezvous at MASTER\_ADDR:MASTER\_PORT. The label sits above the first hop to remind you where rendezvous happens; the actual group spans all nodes.

Node	node₋rank	$local\_rank  o global RANK (this figure)$
Node 0	0	$0 \rightarrow 0, 1 \rightarrow 1$
Node 1	1	$0 \rightarrow 2, 1 \rightarrow 3$
Node 2	2	$0 \to 4, \ 1 \to 5$

Table 1: \*

Mapping of per-node processes to global ranks when nnodes=3, nproc\_per\_node=2.

What is a rank? A rank is a single Python process in the DDP job. DDP assigns:

- Global RANK  $\in \{0, ..., \text{WORLD\_SIZE}-1\}$  unique across the whole job. Rank 0 is the "leader" we use for logging/checkpoints.
- Local LOCAL\_RANK  $\in \{0, ..., NPROC\_PER\_NODE 1\}$  index of the process on its node.
- Node NODE\_RANK  $\in \{0, ..., nnodes 1\}$  which node you are on.

They satisfy

 $\mathtt{WORLD\_SIZE} = \mathtt{nnodes} \times \mathtt{nproc\_per\_node}, \qquad \mathtt{RANK} = \mathtt{NODE\_RANK} \times \mathtt{NPROC\_PER\_NODE} + \mathtt{LOCAL\_RANK}.$ 

In our figure (nnodes=3, nproc\_per\_node=2) the global ranks are: Node  $0 \to RANK\ 0,1$ ; Node  $1 \to RANK\ 2,3$ ; Node  $2 \to RANK\ 4,5$ . All six ranks run identical training loops; after each backward pass, DDP (Gloo backend) averages gradients across all ranks so the models stay in sync. Only RANK 0 saves to ./tiny\_out.

Not to confuse: a rank is not a CPU core or a SLURM task. In this lab we request --ntasks-per-node=1 with SLURM, and then torchrun creates NPROC\_PER\_NODE=2 ranks (processes) per node; each rank may use a few CPU threads (e.g., OMP\_NUM\_THREADS=2).

# 2 Requirements & Offline Staging (req.sh)

#### 2.1 What req.sh sets up and why

- Offline cache root at \$HOME/offline\_repo\_py311: holds wheels or a vendorized pkgs tree so worker nodes do not need internet.
- Pinned versions in requirements\_local.txt:
  - torch==2.3.1+cpu (CPU wheels), transformers==4.41.2, datasets==2.19.0,
  - numpy==1.26.4 (avoids a known formatting issue with datasets+NumPy 2.x),
  - pyarrow==16.1.0, tokenizers==0.19.1, etc.
- Two modes: (a) Online on login node → directly install into pkgs; (b) Offline cluster → pip download wheels to wheels/ then pip install --no-index into pkgs.

#### 2.2 Run it once on the login node

```
bash req.sh
# Verify that you now have:
# $0FFLINE_ROOT/pkgs/ (many site-packages)
# $0FFLINE_ROOT/wheels/ (if you used the offline flow)
```

Listing 1: Prepare local offline/online dependencies

#### 2.3 Why NumPy 1.26.4?

datasets==2.19.0 can trip on NumPy 2.x formatting paths. Pinning to 1.26.4 keeps tokenization/arrow formatting stable. (Our launcher also contains a *defensive* patch that gracefully adapts if a node happens to have NumPy 2.x.)

### 3 The Training Script (train\_tiny.py) — What to Notice

### 3.1 CPU-only safety patch

If CUDA is absent, some utilities (e.g., Accelerate) might try torch.cpu.set\_device(). We monkey-patch it to a no-op on pure CPU nodes to prevent spurious errors:

```
# if not torch.cuda.is_available(): torch.cpu.set_device = lambda *_: None
```

Listing 2: Concept (already in your code, no action needed)

#### 3.2 Distributed init and debugging prints

- Backend: gloo on CPU; nccl on GPU (not used here).
- Rendezvous: env:// uses MASTER\_ADDR, MASTER\_PORT, WORLD\_SIZE, RANK.
- Debug line: every process prints [RANK r] WORLD\_SIZE=w.

#### 3.3 Model & data

- Tokenizer: google/bert\_uncased\_L-2\_H-128\_A-2.
- Tiny BERT config (hidden\_size=64, layers=2, heads=2, intermediate=256, max\_pos=256).
- Dataset: ag\_news (subset to --subset 2000 for fast CPU runs).
- Trainer + DDP: ddp\_find\_unused\_parameters=False, logging\_steps=10.
- Rank 0 saves the model to --out (default ./tiny\_out).

# 4 The Launcher (launch\_tiny.sh) — Phases & Rationale

#### 4.1 Resource allocation (outer salloc)

If not already inside a job, the script re-execs itself under salloc:

- --partition=torch (example partition name),
- --nodes=\$JOB\_NODES (e.g., 10),
- --ntasks-per-node=1 (one launcher task per node),
- --cpus-per-task=4 (room for tokenization/BLAS),
- --time=00:30:00.

#### 4.2 Rendezvous & world size

After allocation:

```
export MASTER_ADDR=$(scontrol show hostnames "$SLURM_NODELIST" | head -n1)
export MASTER_PORT=29500
export NNODES=${SLURM_NNODES:-$(scontrol show hostnames "$SLURM_NODELIST" | wc -1)}
```

**Interpretation:** The first hostname acts as rendezvous server. WORLD\_SIZE = NNODES  $\times$  NPROC\_PER\_NODE.

### 4.3 Preflight visibility check (critical!)

We must ensure each node sees:

- 1. the project directory (\$PROJECT\_DIR),
- 2. the offline packages (\$OFFLINE\_ROOT/pkgs).

The script runs:

```
VIS_REPORT=$(srun --export=ALL -N "$NNODES" -n "$NNODES" bash -lc '
  echo -n "$HOSTNAME "
  [[ -d "'"$PROJECT_DIR"'" ]] && printf "proj=ok " || printf "proj=missing "
  [[ -d "'"$0FFLINE_ROOT"'/pkgs" ]] && printf "pkgs=ok\n" || printf "pkgs=missing\n"
  ')
  echo "$VIS_REPORT" | tee "$HOME/slurm_logs/preflight.$SLURM_JOB_ID.txt"
```

Listing 3: Node visibility report

Read this file first: slurm\_logs/preflight.\$SLURM\_JOB\_ID.txt. You should see lines like hpc07 proj=ok pkgs=ok for every node.

#### 4.4 Conditional staging with sbcast

If any node reports proj=missing or pkgs=missing, the launcher:

- 1. tars \$PROJECT\_DIR and pkgs/,
- 2. broadcasts them to all nodes via sbcast into /tmp,

3. unpacks under a per-job temp root (\$SLURM\_TMPDIR or fallback).

At runtime, each task selects either the shared path or the staged copy:

```
RUN_TMP="${SLURM_TMPDIR:-/tmp/$USER/slurm_$SLURM_JOB_ID}"
RUN_OFFLINE_ROOT="$0FFLINE_ROOT"
RUN_PROJECT_DIR="$PROJECT_DIR"
[[ -d "$RUN_PROJECT_DIR" ]] || RUN_PROJECT_DIR="$RUN_TMP/$(basename "$PROJECT_DIR")"
[[ -d "$RUN_OFFLINE_ROOT/pkgs" ]] || RUN_OFFLINE_ROOT="$RUN_TMP"
```

Listing 4: Per-node runtime roots (inside launcher)

This guarantees all nodes have a working copy without NFS hiccups.

#### 4.5 Environment hygiene (per-node)

Before Python starts, we export environment to make the run self-contained and deterministic:

- PYTHONPATH=RUN\_OFFLINE\_ROOT/pkgs:\$RUN\_PROJECT\_DIR:\$PYTHONPATH
- HF\_HOME=\$HOME/.cache/hf, TRANSFORMERS\_OFFLINE=1, HF\_DATASETS\_OFFLINE=1
- OMP\_NUM\_THREADS=2 (keep BLAS reasonable under DDP)
- TOKENIZERS\_PARALLELISM=false
- PYTORCH\_DIST\_BACKEND=gloo

### 4.6 Sanity probe (per node)

We verify that Python can import torch, transformers, datasets, etc. The launcher prints a one-line summary:

```
srun ... python /tmp/_sanity.py
```

Listing 5: Expect lines like: NODE hpc09 OK -; PY 3.11 torch 2.3.1 tfm 4.41.2 ...

If a node fails here: check that it received staged copies; confirm Python is available; re-run with --exclude=<node> via SALLOC\_OPTS.

#### 4.7 The wrapper and optional NumPy 2.x patch

A tiny wrapper (/tmp/\_run\_wrapper.py) inserts pkgs/ and project path into sys.path, and if NumPy 2.x is detected, applies a safe adapter to the datasets formatter. This keeps older datasets versions working even if a worker node has a newer NumPy by accident.

#### 4.8 The distributed launch

Finally, we call:

```
python -m torch.distributed.run \
    --nnodes="$NNODES" \
    --nproc_per_node="$NPROC_PER_NODE" \
    --rdzv_backend=c10d \
    --rdzv_endpoint="$MASTER_ADDR:$MASTER_PORT" \
    --rdzv_id="$SLURM_JOB_ID" \
```

```
--node_rank="$SLURM_NODEID" \
/tmp/_run_wrapper.py --subset 2000 --epochs 3
```

Listing 6: Key torch.distributed.run arguments

Interpretation: node\_rank is assigned by SLURM, WORLD\_SIZE = nnodes \* nproc\_per\_node, and rendezvous is shared via the master host:port.

# 5 Running the Lab

### 5.1 One-time setup

```
bash req.sh
```

Listing 7: Install deps into offline cache (login node)

### 5.2 Launch (interactive job)

```
bash ~/launch_tiny.sh
# Optional: exclude flaky nodes
# SALLOC_OPTS="--exclude=hpc42" bash ~/launch_tiny.sh
```

Listing 8: Start the distributed job (10 nodes, 2 procs/node)

#### 5.3 What to watch on screen

- 1. Allocation line: [alloc] job=... nodes=...
- 2. Rendezvous line: NNODES=... MASTER\_ADDR=... MASTER\_PORT=...
- 3. Preflight report: each host prints proj=ok pkgs=ok (or triggers staging).
- 4. Sanity probe: each host prints NODE <name> OK -> PY 3.11 torch 2.3.1 ...
- 5. Torchrun banner: nnodes=?, nproc\_per\_node=?, node\_rank=?, rdzv=?
- 6. Training logs: every rank periodically logs via HF Trainer; each process prints [RANK r] WORLD\_SIZE=w.

# 6 Logs and How To Read Them Carefully

#### 6.1 Files generated

All stdout/stderr from srun phases are captured to:

```
$HOME/slurm_logs/<jobname>.<jobid>.<nodename>.<taskid>.out
```

Plus one preflight summary: slurm\_logs/preflight.\$SLURM\_JOB\_ID.txt.

#### 6.2 Minimal checklist

- 1. **Preflight:** Open the preflight file; ensure *every* node is proj=ok pkgs=ok.

  If not: confirm that staging ran (you should see "staging via sbcast" on screen), then check per-node .out to see lines like \$HOSTNAME staged -> /tmp/....
- 2. Sanity probe: In each node's .out, find the NODE <name> OK -> PY ... torch ... tfm ... datasets .... All must say OK.
- 3. World size: In any training .out, grep for [RANK. Confirm that the largest rank equals WORLD\_SIZE-1.

Expected: WORLD\_SIZE = NNODES \* NPROC\_PER\_NODE.

4. Saving: Only rank 0 will save artifacts; look for Saving model to ./tiny\_out.

#### 6.3 Useful greps

```
# Show preflight summary
cat ~/slurm_logs/preflight.$SLURM_JOB_ID.txt

# List all node-task outputs for this job
ls -1 ~/slurm_logs/*.$SLURM_JOB_ID.*.out

# Check ranks and world size
grep -h "\[RANK" ~/slurm_logs/*.$SLURM_JOB_ID.*.out | sort -u

# Find any failure signatures
grep -HiE "Traceback|ERROR|RuntimeError|OSError" ~/slurm_logs/*.$SLURM_JOB_ID.*.out
```

Listing 9: Quick log triage

# 7 Design Choices (Deep Dive)

#### 7.1 Why Gloo + env:// on CPU

Gloo is PyTorch's reliable CPU backend. Using env:// keeps launch simple under SLURM: we export rendezvous variables once and let torchrun propagate them to all ranks.

### 7.2 Why --ntasks-per-node=1 but NPROC\_PER\_NODE=2

We place one SLURM task per node (clean lifecycle and logging), then let torchrun spawn 2 processes per node (total ranks per node = 2). This makes failure handling (--kill-on-bad-exit=1) predictable and reduces SLURM scheduling overhead.

### 7.3 Why OMP\_NUM\_THREADS=2

On CPU, too many BLAS threads per process can degrade performance under DDP. With 2 ranks/node and light models, OMP\_NUM\_THREADS=2 balances tokenization and linear algebra without oversubscription.

### 7.4 Why conditional staging with sbcast

Some nodes may not see shared storage or may mount it inconsistently. Broadcasting a tarball to /tmp ensures a consistent, job-scoped snapshot with low latency.

#### 7.5 Defensive NumPy patch

If a worker accidentally has NumPy 2.x, an adapter in the wrapper rebinds a datasets formatter method to keep np.asarray semantics. This avoids mid-run crashes due to heterogeneous node images.

# 8 Verification, Metrics, and Expected Outputs

# 8.1 Rank banner (from train\_tiny.py)

```
[RANK 0] WORLD_SIZE=20
[RANK 7] WORLD_SIZE=20
...
```

Here, 10 nodes  $\times$  2 procs/node  $\Rightarrow$  WORLD\_SIZE = 20.

#### 8.2 HF Trainer logs

Every logging\_steps steps (e.g., 10), each rank prints a short report. You may see throughput stabilize after the first few steps as tokenizers warm up.

#### 8.3 Saved artifacts (rank 0)

- tiny\_out/config.json, pytorch\_model.bin, tokenizer.json, etc.
- Check modified time to confirm it was produced by the current run.

# 9 Troubleshooting (Most Common Issues)

Symptom	Fix / Explanation
Some nodes show proj=missing or pkgs=missing	This is normal on clusters without shared storage. The launcher will stage via
	sbcast. Confirm per-node staged ->
	/tmp/ messages.
Traceback about datasets and NumPy 2.x	Pin NumPy to 1.26.4 in req.sh. The wrapper
	also applies a safety patch; ensure the wrap-
	per ran (it prints a "patch applied" line).
Address already in use on MASTER_PORT	Change MASTER_PORT in the launcher (e.g.,
	any unreserved port $> 1024$ ).
Ranks hang at init	${\it Check that {\tt MASTER\_ADDR resolves from}}$
	all nodes. Verify firewalls. Ensure
	rdzv_endpoint matches the allocation.
Slow training or CPU spikes	Lower OMP_NUM_THREADS to 1, or reduce
	NPROC_PER_NODE. Confirmcpus-per-task
	leaves headroom.
Model not saved	Only rank 0 saves. Look for [RANK 0] logs.
	Ensure disk write permissions inout.

# 10 Lab Tasks & Checkoff

- 1. Run req.sh. Show your pkgs/ tree exists and includes torch, transformers, datasets, numpy.
- 2. Launch training on 3 nodes with NPROC\_PER\_NODE=2. Capture:
  - Preflight summary (proj=ok pkgs=ok per node).
  - At least one node's sanity probe line.
  - A snippet with [RANK r] WORLD\_SIZE=w proving correct world size.
- 3. Open tiny\_out/ and list saved files (rank 0).
- 4. (Optional) Repeat with 5 nodes and compare time/epoch.

# 11 Reference: Key Variables at a Glance

Name	Set In	Meaning
MASTER_ADDR	launcher	Hostname for rendezvous (first node).
MASTER_PORT	launcher	TCP port for rendezvous (e.g., 29500).
NNODES	launcher	Number of allocated nodes.
NPROC_PER_NODE	launcher	# of ranks per node (torchrun).
WORLD_SIZE	implicit	NNODES * NPROC_PER_NODE.
node_rank	torchrun	Rank of this node among nodes (0-based).
RANK	torchrun	Global rank of a process (0WORLD_SIZE-1).
RUN_PROJECT_DIR	launcher	Per-node project path (shared or staged).
RUN_OFFLINE_ROOT	launcher	Per-node offline root (shared or staged).
OMP_NUM_THREADS	launcher	BLAS threading per process (e.g., 2).
TRANSFORMERS_OFFLINE	launcher	Disables net calls in transformers.
HF_DATASETS_OFFLINE	launcher	Disables net calls in datasets.

# 12 Appendix: Commands You Will Actually Type

#### A. Prepare dependencies

bash req.sh

# B. Start a 10-node, 2-proc/node run

```
bash ~/launch_tiny.sh
# or exclude one bad node:
# SALLOC_OPTS="--exclude=hpc42" bash ~/launch_tiny.sh
```

# C. Inspect logs

```
cat ~/slurm_logs/preflight.$SLURM_JOB_ID.txt
ls ~/slurm_logs/*.$SLURM_JOB_ID.*.out | wc -1
grep -h "\[RANK" ~/slurm_logs/*.$SLURM_JOB_ID.*.out | sort -u | head
```

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