



Introduction to GNU Parallel

Parallelizing Massive Individual Tasks

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http://www.hpc.lsu.edu/training/archive/tutorials.php







Introduction

- What is GNU Parallel?
 - ✤ A command-line tool used for parallelizing massive individual tasks
- When and Why to use GNU Parallel?
 - Examples: 1 million small MD simulations, 10k protein analysis

Basic Usage of GNU Parallel

- GNU Parallel Syntax and Options
- Introducing Running 3 Types of jobs using GNU Parallel:
 - Serial Tasks Run each LAMMPS task in serial
 - Multi-Threaded Tasks Run each LAMMPS task using multiple threads
 - Small MPI Jobs Run each LAMMPS task using multiple MPI processes

Proper usage of GNU Parallel

- Memory consideration
- Task granularity









What is GNU Parallel? Parallelizing Massive Individual Tasks







Parallelizing Massive Individual Tasks

✤ Parallelize

This refers to the process of dividing a workload into multiple smaller tasks and executing them simultaneously. It allows for better utilization of computational resources, such as multi-core CPUs or multiple nodes in a cluster.





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Individual (Independent)

Each task operates independently of the others. This means there's no dependency between tasks, allowing them to be executed in parallel without waiting for one another.





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Tasks

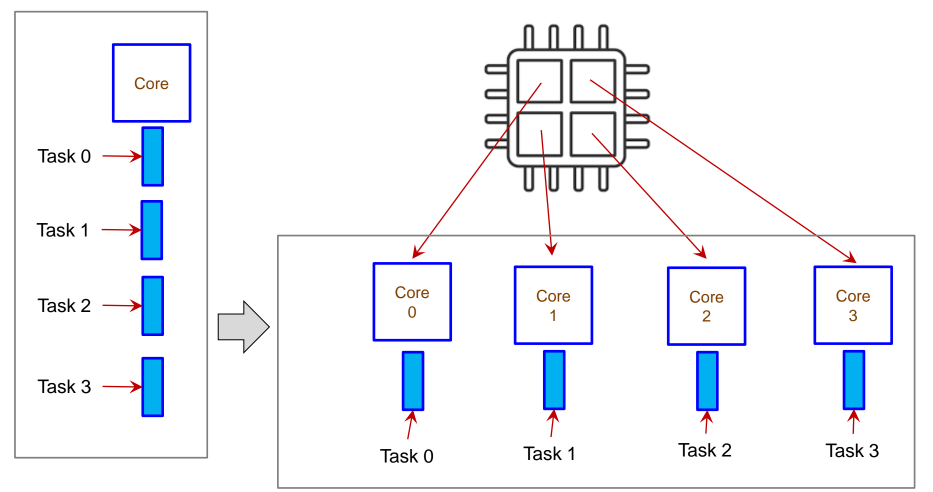
These are the units of work or commands you want to execute. Each task could be a program, script, or function that processes a specific set of inputs.





What do we want to accomplish?

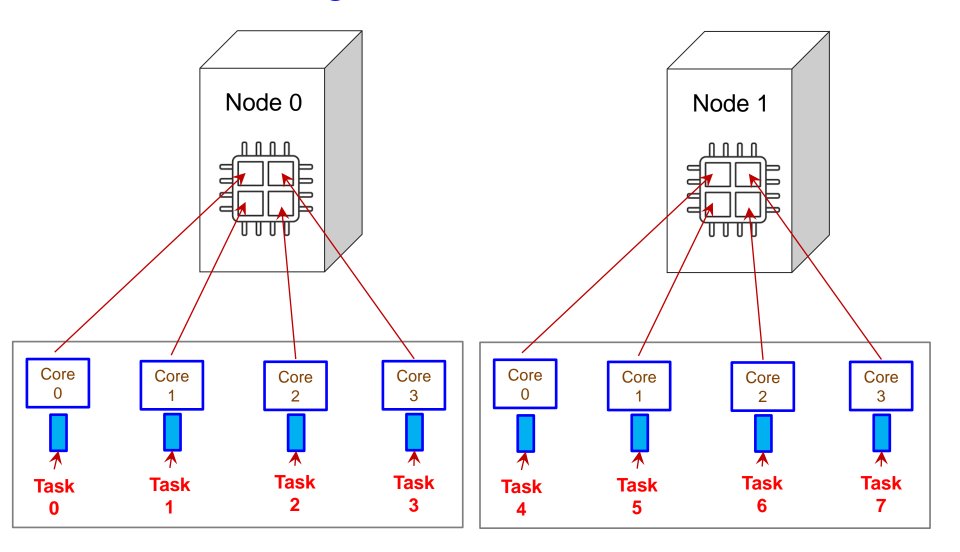
 Parallize lots of small independent tasks on a multi-core platform (compute nodes)







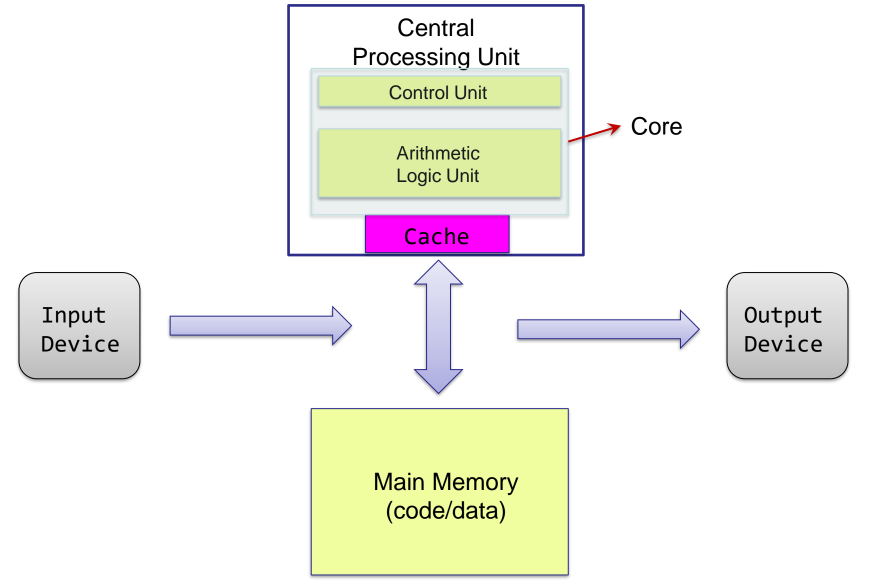
Background and Distribute







The von Neumann Architecture



Introduction to GNU Parallel

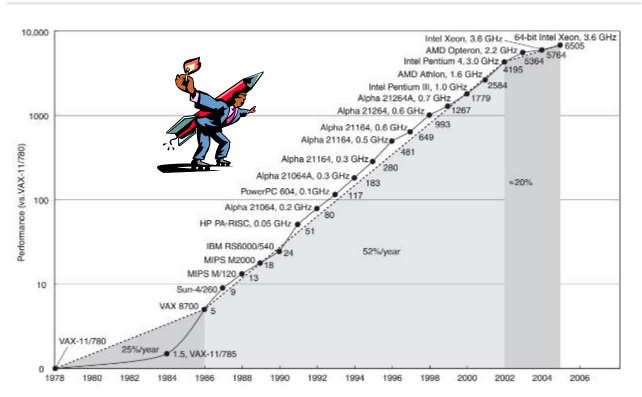




Changing Times

- From 1986 2002, microprocessors were speeding like a rocket, increasing in performance an average of 50% per year.
- Since then, it's dropped to about 20% increase per year.

History of Processor Performance



Moore's Law states that transistor counts on chips double roughly every two years.

Limitation:

2 GHz Consumer 4 GHz Server

Source:

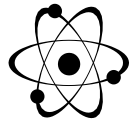
http://www.cs.columbia.edu/~sed wards/classes/2012/3827-spring/





A Little Physics Problem

- Smaller transistors = faster processors.
- Faster processors = increased power consumption.
- Increased power consumption = increased heat.
- Increased heat = unreliable processors.



Solution:

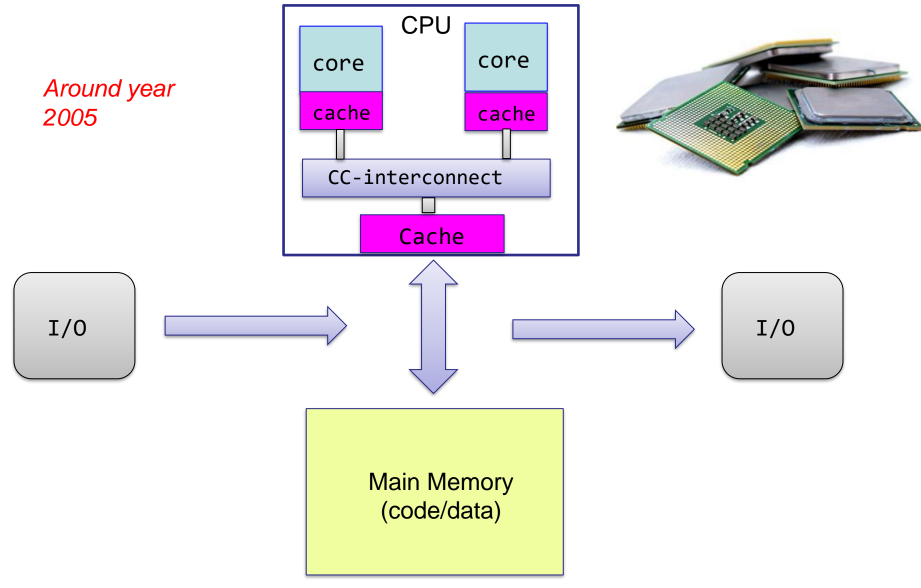
- Move away from single-core systems to multicore processors.
- "core" = central processing unit (CPU)
- Introducing parallelism
 - What if your problem is also not CPU dominant?







The von Neumann Architecture



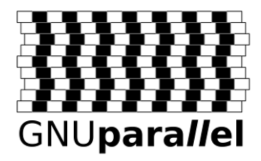
Introduction to GNU Parallel





GNU Parallel

- GNU parallel is a shell tool for executing "embarrassingly parallel" tasks using one or more computers (compute nodes).
- > Terminology:
 - Task: Each small, independent piece of computation work to be finished, e.g., a single genome computation, a single MD simulation
 - Job: The list of tasks to be completed on a set of nodes (cores)
- A task can be a single command or a small script that must be run for each of the lines in the input.
- The typical input is a *list of files*, a list of hosts, a list of users, a list of URLs, or a list of tables.
- See more at: <u>https://www.gnu.org/software/parallel/</u>







Introduction to GNU Parallel

GNU Parallel Syntax



. . .



Adding GNU Parallel to Environment

> On SuperMike3:

[fchen14@mike139 ~]\$ module av parallel ---- /usr/local/packages/Modules/default/modulefiles/linux-rhel8-icelake ---parallel-netcdf/1.12.2/intel-2021.5.0 parallel-netcdf/1.12.2/intel-2021.5.0-intel-mpi-2021.5.1 # load the module to environment [fchen14@mike139 ~]\$ module load parallel/20210922/intel-2021.5.0 [fchen14@mike139 ~]\$ which parallel /usr/local/packages/parallel/20210922/hrsviur/bin/parallel [fchen14@mike139 ~]\$ parallel -version # check GNU parallel version GNU parallel 20210922 Copyright (C) 2007-2021 Ole Tange, http://ole.tange.dk and Free Software Foundation, Inc.

Or add the below line to ~/.modules:

module load parallel/20210922/intel-2021.5.0





GNU Parallel Syntax

- Reading commands to be run in parallel from an input file: parallel [OPTIONS] < CMDFILE</p>
- Reading command arguments on the command line: parallel [OPTIONS] COMMAND [ARGUMENTS] ::: ARGLIST
- Reading command arguments from an input file:

parallel [OPTIONS] COMMAND [ARGUMENTS] :::: ARGFILE



. . .



ARGLIST from command line

> parallel [OPTIONS] COMMAND [ARGUMENTS] ::: ARGLIST

```
> Examples:
    [fchen14@mike139 ~]$ parallel echo ::: A B C
    Α
    В
    С
    [fchen14@mike139 ~]$ parallel echo ::: `seq 1 3`
    1
    2
    3
    [fchen14@mike139 ~]$ parallel echo ::: {A..Z}
    Α
    В
     . . .
    7
    [fchen14@mike139 test]$ ls -1 | parallel echo
    2013-06-18.tgz
    backups.sh
    bigmem test.pbs
```





ARGLIST from file

```
> parallel [OPTIONS] COMMAND [ARGUMENTS] :::: ARGFILE
   [fchen14@mike139 GNU_PARALLEL]$ pwd
   /project/fchen14/GNU PARALLEL
   [fchen14@mike139 GNU_PARALLEL]$ cat input.lst | head
   01.lj
   02.1j
   03.1j
   . . .
   [fchen14@mike139 GNU_PARALLEL]$ head input.lst -n 5 | parallel echo
   01.lj
   02.1j
   03.1j
   04.1j
   05.1j
   [fchen14@mike139 GNU PARALLEL]$ parallel echo :::: input.lst
   01.lj
   02.1j
    . . .
```





Replacement Strings

```
[+chen14@mike139 GNU_PARALLEL]$ parallel echo {.} ::: data/in
data/in
```

`{/.}' returns the basename of the input line without extension. It is a combination of {/} and {.}:

```
[fchen14@mike139 GNU_PARALLEL]$ parallel echo {/.} ::: data/
in
```

See "man parallel" for more detailed explanation.





Replacement String Example

- Print the full path of the input file, and then print the desired output file name, e.g.:
 - Input file: data/lj.in
 - Output file name: output/lj.out

Process data/lj.in and send result to output/lj.out
\$ parallel echo {} output/{/.}.out ::: data/lj.in
data/lj.in output/lj.out





Parallelize Job Script

- GNU parallel is often called as this: cat input_file | parallel command parallel command ::: foo bar
- If command is a script, parallel can be combined into a single file so this will run the script in parallel:

```
parallel [OPTIONS] script [ARGUMENTS] ::: ARGLIST
```

– or

parallel [OPTIONS] script [ARGUMENTS] :::: ARGFILE

See next slide for example...





Parallize Script Example

This is the script we want to parallize "cmd_ex.sh":

```
#!/bin/bash
```

print the input, on which host, which working directory
echo "This script uses input: \$1 on \$HOSTNAME:\$PWD"

Parallize the script using ARGLIST from command line: [fchen14@mike139 GNU_PARALLEL]\$ parallel --wd \$PWD ./cmd_ex.sh ::: A B C This script uses input: A on mike139:/project/fchen14/GNU_PARALLEL This script uses input: B on mike139:/project/fchen14/GNU_PARALLEL This script uses input: C on mike139:/project/fchen14/GNU_PARALLEL

```
Parallize the script using ARGFILE:
```

```
[fchen14@mike139 GNU_PARALLEL]$ cat argfile
A
B
C
[fchen14@mike139 GNU_PARALLEL]$ parallel --wd $PWD ./cmd_ex.sh :::: argfile
This script uses input: A on mike139:/project/fchen14/GNU_PARALLEL
This script uses input: B on mike139:/project/fchen14/GNU_PARALLEL
This script uses input: C on mike139:/project/fchen14/GNU_PARALLEL
```

Can parallize Python/Perl scripts, see "man parallel" for details





Common OPTIONS --jobs (-j)

≻ --jobs N (-j N)

. . .

- Number of jobslots on each machine (node). Run up to N jobs in parallel. 0 means as many as possible. Default is 100% which will run one job per CPU core on each machine.
- On HPC/LONI clusters, N is number of jobslots per node.
- Make sure you use GNU Parallel version >=20161022 to avoid a "Max jobs to run" bug

[fchen14@mike139 test]\$ parallel --version
GNU parallel 20210922

≻ -j +N

- Add N to the number of CPU cores. Run this many jobs in parallel.
- ≻ -j -N
 - Subtract N from the number of CPU cores. Run this many jobs in parallel. If the evaluated number is less than 1 then 1 will be used.





Common OPTIONS --slf (PBS)

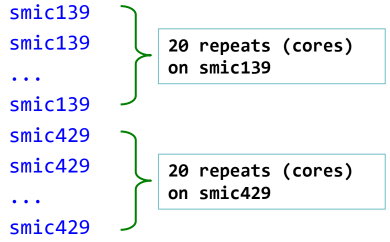
> --slf filename (--sshloginfile filename)

- File with schlogins. The file consists of schlogins on separate lines. Empty lines and lines starting with '#' are ignored.
- Look at "man parallel" for more detailed explanation.
- A typical example on PBS clusters while running batch jobs:

```
--slf $PBS_NODEFILE
```

– Recall what is inside \$PBS_NODEFILE?

```
[fchen14@smic139 laplace]$ cat $PBS_NODEFILE # assume we use SuperMIC
```







Common OPTIONS --slf (Slurm)

- > --slf filename (--sshloginfile filename)
 - To get the sshloginfile on Slurm job session, use the below command: scontrol show hostname \$SLURM_NODELIST > nodefile
 - A typical example on HPC/LONI clusters while running batch jobs:
 -slf nodefile
 - Look at \$SLURM_NODELIST and nodefile

```
# start an interactive job requesting 2 nodes (64x2=128 cores)
[kasetti@mike4 ~]$ srun -N2 -n128 -p workq --cpu-bind none --pty bash
srun: Job is in held state, pending scheduler release
srun: job 39474 queued and waiting for resources
Interactive job 39474 waiting:
srun: job 39474 has been allocated resources # we got mike157 and mike158
[kasetti@mike157 ~]$ echo $SLURM_NODELIST
mike[157-158]
[kasetti@mike157 ~]$ scontrol show hostname $SLURM_NODELIST > nodefile
[kasetti@mike157 ~]$ cat nodefile
```

```
mike157
mike158
```





Common OPTIONS --sshdelay

- If many tasks are started on the same compute node, sshd can be overloaded. On SuperMike3/QB3, some of the tasks might fail to start, e.g., starting all 64/48 tasks at the same time.
- GNU parallel can insert a delay between each task run on the same server:

```
[fchen14@mike139 GNU_PARALLEL]$ parallel --sshdelay 0.1 echo ::: A B C
A
B
C
```





Common OPTIONS --wd

- > --wd mydir (--workdir mydir)
 - Designate the working directory of your commands.
 - A typical value can be:
 - \$PBS_O_WORDIR (PBS)
 - \$SLURM_SUMBIT_DIR (Slurm)





Common OPTIONS -- env

> --env ENV_VAR

- --env to tell GNU parallel to transfer an environment variable to the remote system.
- A typical usage:

```
export OMP_NUM_THREADS=5
```

```
parallel --env OMP_NUM_THREADS cmd ::: ARGLIST
```





Common OPTIONS --progress

--progress

- Show progress of computations. (Not recommended for batch jobs)
- List the computers involved in the task with number of CPU cores detected and the max number of jobs to run.
- After that show progress for each node: number of running jobs, number of completed jobs, and percentage of all jobs done by this computer.
- Example:

```
[fchen14@mike139 ~]$ parallel --progress echo ::: A B C
```

```
Computers / CPU cores / Max jobs to run
1:local / 64 / 3
```

Computer:jobs running/jobs completed/%of started jobs/Average seconds to complete local:3/0/100%/0.0s A local:2/1/100%/1.0s B local:1/2/100%/0.5s C local:0/3/100%/0.3s

See also --bar





Common OPTIONS --joblog

> --joblog logfile

- Creates a record for each completed subjob (task) to be written to LOGFILE, with info on how long they took, their exit status, etc.
- Can be used to identify failed jobs, e.g.:

[fchen14@mike139 misc]\$ parallel --joblog logfile exit ::: 1 2 0 0
[fchen14@mike139 misc]\$ cat logfile

Seq	Host	Starttime	JobRuntime	Send	Receive	Exitval	Signal	Command
1	:	1477514132.358	0.019	0	0	1	0	exit 1
2	:	1477514132.375	0.003	0	0	2	0	exit 2
3	:	1477514132.376	0.002	0	0	0	0	exit 0
4	:	1477514132.377	0.003	0	0	0	0	exit 0





Common OPTIONS --timeout

--timeout secs

- Time out for command. If the command runs for longer than secs (seconds) it will get killed.
- If secs is followed by a % then the timeout will dynamically be computed as a percentage of the median average runtime. Only values > 100% will make sense.
- Useful if you know the command has failed if it runs longer than a threshold.





Introduction to GNU Parallel

Serial Jobs Example





LAMMPS Introduction

- LAMMPS is a classical molecular dynamics code with a focus on materials modeling. <u>https://www.lammps.org/</u>
- You don't need any background in molecular dynamics/LAMMPS to understand today's example.
- > Typical LAMMPS Syntax
 - Serial run

lmp_serial -in in.script

- Multi-Threaded run

env OMP_NUM_THREADS=4 lmp_omp -sf omp -in in.script # use OMP_NUM_THREADS
lmp_omp -sf omp -pk omp 5 -in in.script # use -pk omp to specify threads

- MPI run

srun --overlap -n 4 lmp_mpi -in in.script # Slurm version, --overlap only
needed for interactive job
mpirun -n 4 lmp mpi -in in.script # PBS version

– Custom command:

lmp_serial -var nsteps 200 -in in.script # we defined a custom variable in the input file to let nsteps control total steps to run, The above command will run our input for 200 steps





LAMMPS Input File Used Today Janas malt

# 3d Lennard-Jone	s melt					
variable x index	1					
variable y index	1					
variable z index	1					
variable xx equal	80* <mark>\$x</mark>					
variable yy equal	riable yy equal <mark>80*\$y</mark>					
variable zz equal	80* <mark>\$z</mark>					
units	lj					
atom_style	atomic					
lattice	fcc 0.8442					
region	box block 0 <mark>\${xx}</mark> 0 <mark>\${yy}</mark> 0 <mark>\${zz}</mark>					
create_box	1 box					
create_atoms	1 box					
mass	1 1.0					
velocity all create 1.44 87287 loop geom						
pair_style	lj/cut 2.5					
pair_coeff	1 1 1.0 1.0 2.5					
neighbor 0.3 bin						
<pre>neigh_modify</pre>	delay 0 every 20 check no					
fix	1 all nve					
# \${nsteps} is the parameter passing from LAMMPS command line "-var nste						
run	<pre>\${nsteps}</pre>					

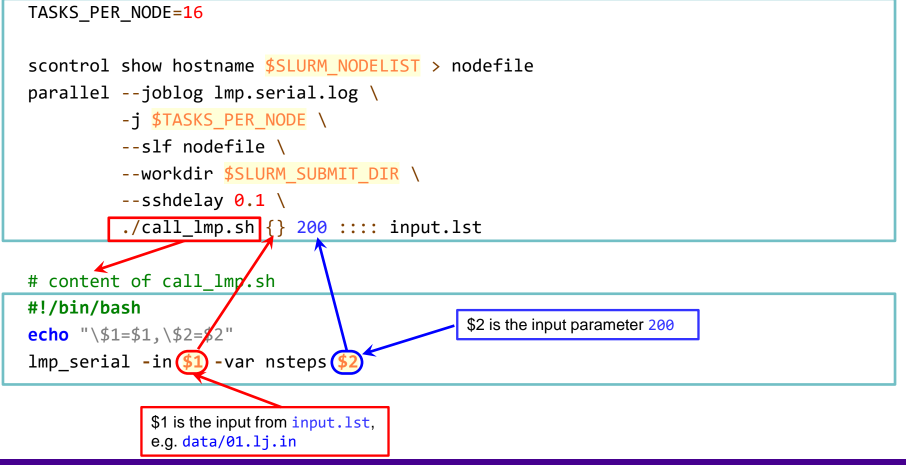


Distribute Serial Jobs LAMMPS (Slurm)

```
#!/bin/bash
#SBATCH -N 2
                                      # request two nodes
#SBATCH -n 128
                                      # specify 128 process
#SBATCH -t 2:00:00
                                       [fchen14@mike2 GNU PARALLEL]$ head input.lst
#SBATCH -p checkpt
                                       data/01.lj.in
                                       data/02.lj.in
#SBATCH - A hpc hpcadmin8
                                       data/03.lj.in
#SBATCH -o gp-serial.out
                                       data/04.lj.in
                                       data/05.lj.in
TASKS PER NODE=16
                                       . . .
SECONDS=0
scontrol show hostname $SLURM NODELIST > nodefile
parallel --joblog lmp.serial.log \
         -j $TASKS PER NODE \
         --slf nodefile \
         --workdir $SLURM SUBMIT DIR \
         --sshdelay 0.1 \
         `which lmp_serial` -in {} -var nsteps 200 :::: input.lst
echo "took $SECONDS sec"
```

In case the task command is too long (complex)

- Use a script for each task to be distributed, example here (call_lmp.sh)
- GNU Parallel will distribute each task script







Distribute Serial Jobs LAMMPS (PBS)

#!/bin/bash

- #PBS -1 nodes=2:ppn=20
- #PBS -1 walltime=1:00:00
- #PBS -q checkpt
- #PBS -A hpc_hpcadmin8
- #PBS -j oe

```
#PBS -o gp-serial-pbs.out
```

```
module load parallel
module load lammps
TASKS_PER_NODE=20
SECONDS=0
```

[fchen14@smic1	GNU_PARALLEL]\$	head	input.lst
data/01.lj.in			
data/02.lj.in			
data/03.lj.in			
data/04.lj.in			
data/05.lj.in			
data/06.lj.in			
•••			

```
cd $PBS_O_WORKDIR
parallel --joblog lmp.serial.log \
    -j $TASKS_PER_NODE \
    --slf $PBS_NODEFILE \
    --workdir $PBS_O_WORKDIR \
    --sshdelay 0.1 \
    `which lmp_mpi` -in {} -var nsteps 200 :::: input.lst
echo "took $SECONDS sec"
```





Introduction to GNU Parallel

Multi-Threaded Example

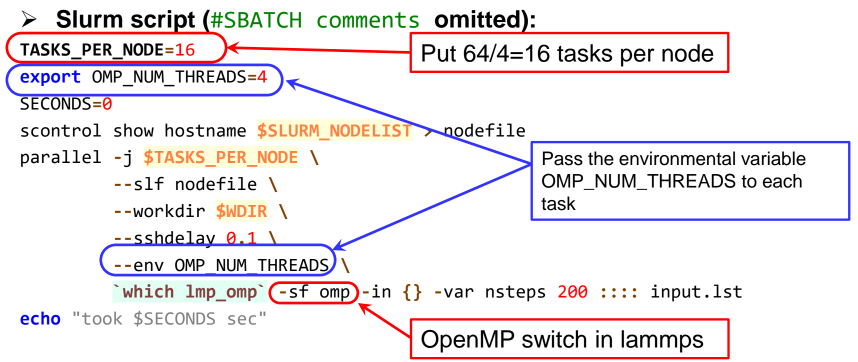
Introduction to GNU Parallel





Distribute Multi-Threaded Jobs

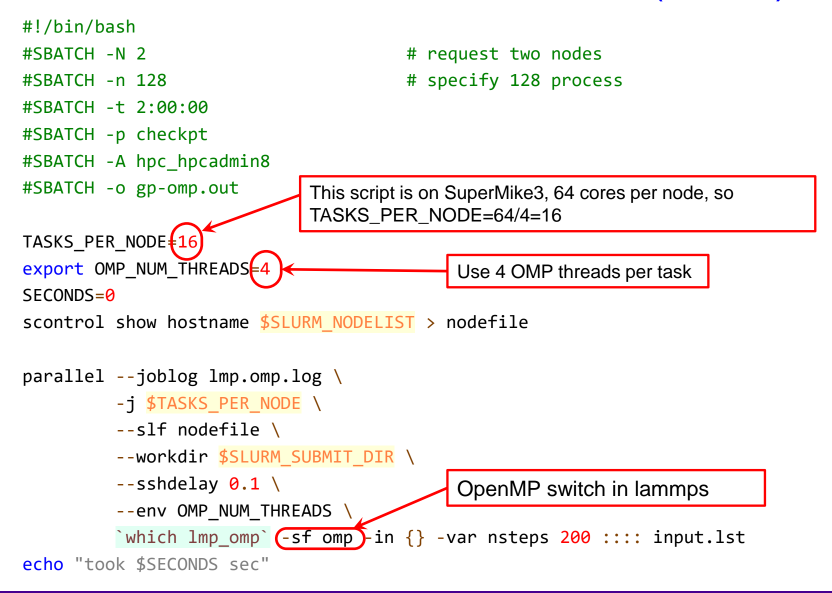
- Distribute Multi-Threaded jobs is very similar to the pure serial job example, the only difference is TASKS_PER_NODE:
 - TASKS_PER_NODE=CPU_CORES_PER_NODE / NUM_THREADS_PER_TASK
- If each job uses 4 threads, each node on SuperMike3 has 64 cores, then
 - TASKS_PER_NODE=64/4=16







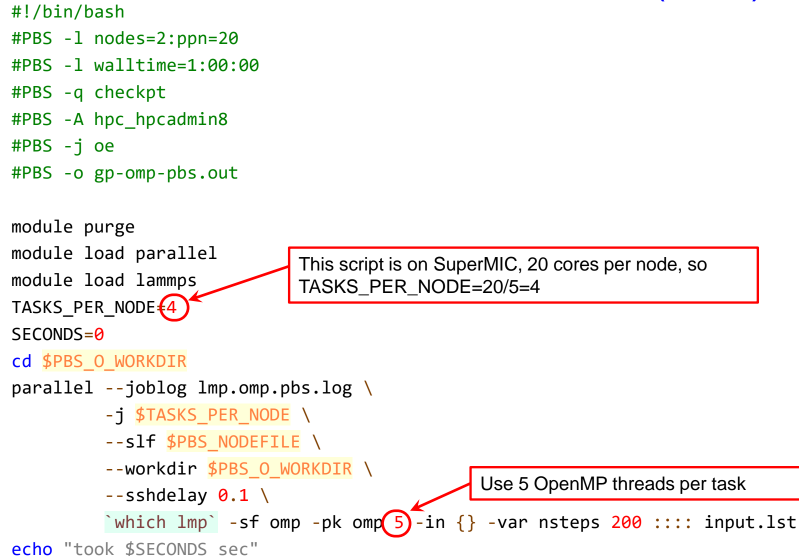
Multi-Threaded LAMMPS (Slurm)







Multi-Threaded LAMMPS (PBS)







Introduction to GNU Parallel

Multi-Process (MPI) Example





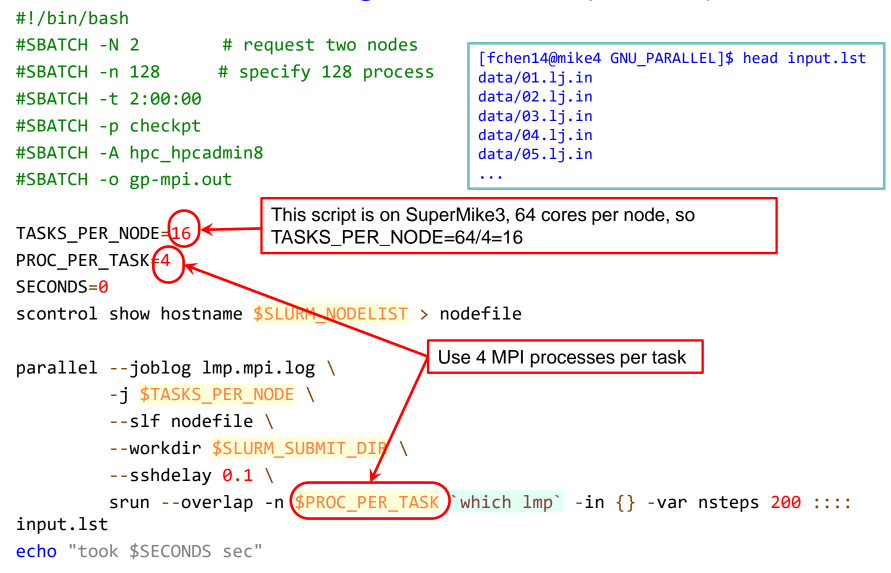
Distribute MPI Jobs - LAMMPS

- > This section describes how to distribute small MPI jobs.
- Example problem LAMMPS MPI
 - Using the same input file, but with multiple MPI process for each task.
 - For simplicity, each MPI process will use only one thread





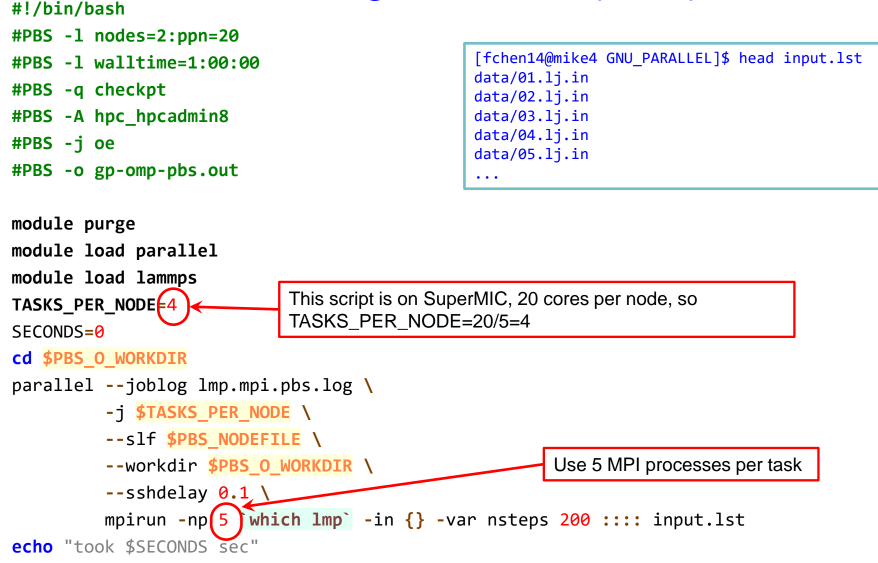
Distributing MPI Jobs (Slurm)







Distributing MPI Jobs (PBS)







Introduction to GNU Parallel

Proper Usage of GNU Parallel





Common Rules

- Don't use more than one node when you are debugging/testing your code
- > Know the performance of single task
- Start with only a few tasks in your input.lst
- After you are comfortable with one node job, start with two node job first before jumping to more than three nodes
- > Typically use no more than 5 nodes.

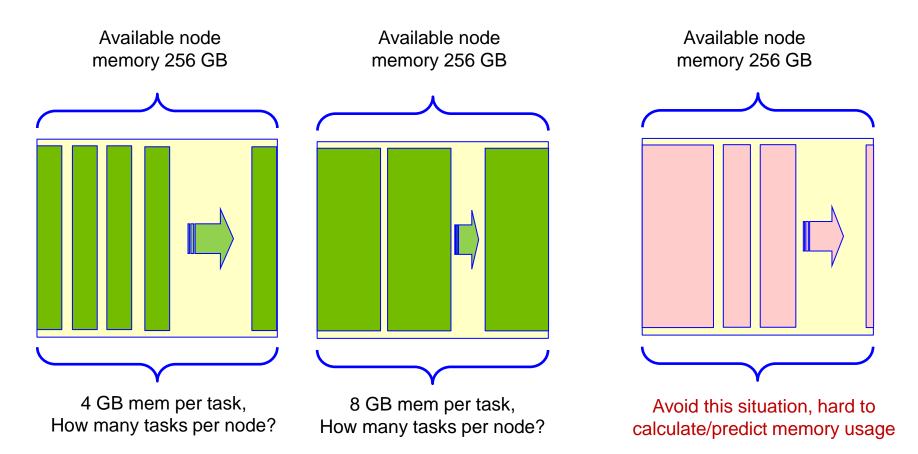




Memory Consideration

Relationship between node memory and cores

- Rule of Thumb: cannot exceed the available memory on a node

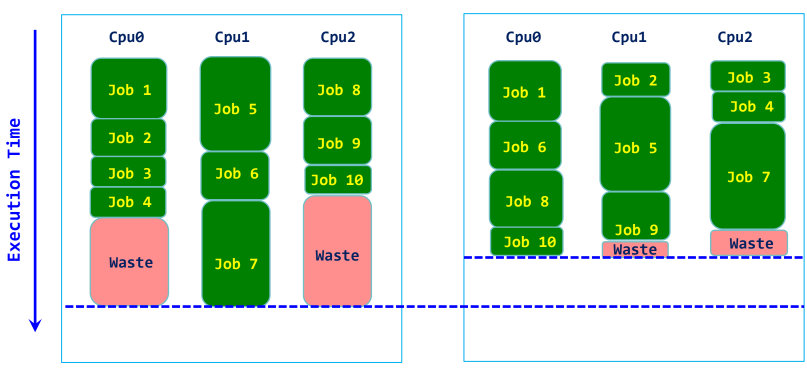






Load Balancing in GNU Parallel

GNU Parallel spawns the next job when one finishes - keeping the CPUs active and thus saving time.



Without Load Balancing

With Load Balancing





Task Granularity

In parallel computing, granularity (or grain size) of a task is a measure of the amount of work (or computation) which is performed by that task.

Impact of granularity on performance

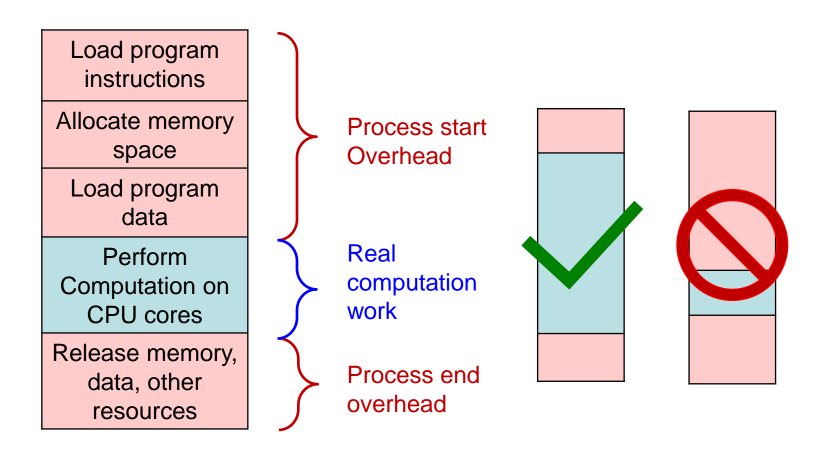
- Using fine grains or small tasks results in more parallelism and hence increases the speedup. However, synchronization overhead, scheduling strategies etc. can negatively impact the performance of fine-grained tasks.
- Simply increasing parallelism alone cannot give the best performance.
- In order to reduce the communication overhead, granularity can be increased. Coarse grained tasks have less communication overhead but they often cause load imbalance. Hence optimal performance is achieved between the two extremes of fine-grained and coarse-grained parallelism.

<u>Ref: https://en.wikipedia.org/wiki/Granularity (parallel computing)</u>





Components of a Task (Process)



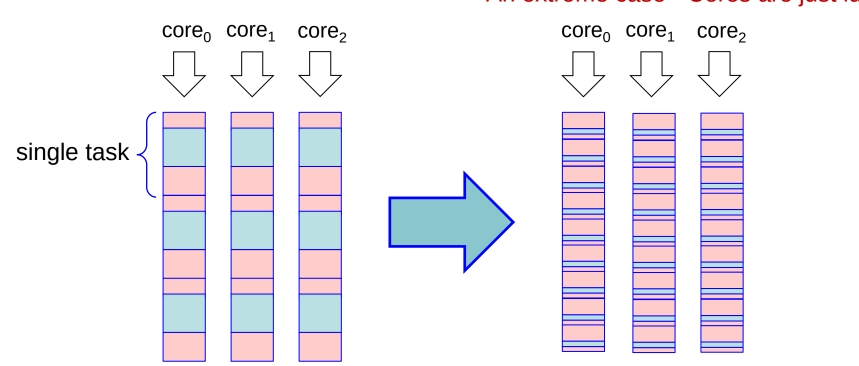


Typical Misuse -Tiny grain size case



Tiny grain size

- E.g., each task takes little time (e.g., less than a second)
- Most time will be spent on overhead



An extreme case - Cores are just idling

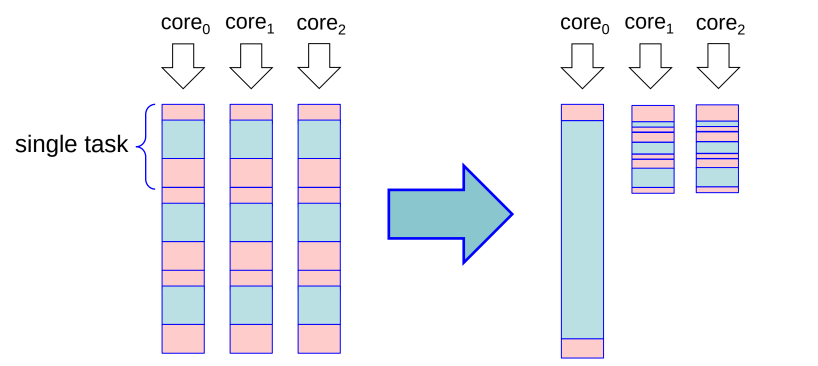


Typical Misuse -Large grain size case



Large grain size

- Some tasks are much longer than the rest
- Load balancing can never be achieved



An extreme case - Load imbalancing







- > Why need GNU Parallel?
- Basic syntax of GNU Parallel and examples
- Ho to use it wisely

- > For more information about GNU Parallel, refer to:
 - <u>https://www.gnu.org/software/parallel/parallel_tutorial.html</u>