Introduction to GNU Parallel - Parallelizing Massive Individual Tasks

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Outline

➢ Why GNU Parallel?
➢ Basic Usage of GNU Parallel
  – GNU Parallel Syntax and Options
  – Introducing Running 3 Types of jobs:
    o Serial Tasks
      ✓ Run each blast task in serial
    o Multi-Threaded Tasks
      ✓ Run each blast task using 2 threads
    o Small MPI Jobs
      ✓ Run each MPI Laplacian solver using 4 processes
➢ Proper usage of GNU Parallel
  – Memory consideration
  – Task granularity
What do we want to accomplish?

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

Limitation:
2 GHz Consumer
4 GHz Server

Source:
http://www.cs.columbia.edu/~sedwards/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

Central Processing Unit
- Control Unit
- Arithmetic Logic Unit

Cache

Core

Main Memory (code/data)

Input Device

Output Device
**The von Neumann Architecture**

- **CPU**: contains core and cache components.
- **CC-interconnect**: connects the cores and cache.
- **Cache**: central memory storage between CPU and main memory.
- **Main Memory**: contains a large amount of data and code, accessed by the CPU.
- **I/O**: input/output components for data exchange with the external world.

*Around year 2005*
GNU Parallel

- GNU parallel is a shell tool for executing jobs in parallel using one or more computers (compute nodes).
- A job can be a **single command or a small script** that has to 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: [https://www.gnu.org/software/parallel/](https://www.gnu.org/software/parallel/)
Adding GNU Parallel to Environment

➢ On SuperMike2:
[fchen14@mike421 ~]$ module av gnuparallel # check available version
------------ /usr/local/packages/Modules/3.2.10/modulefiles/apps
gnuparallel/20180222/INTEL-18.0.0
[fchen14@mike1 ~]$ module load gnuparallel/20180222/INTEL-18.0.0 # load
 gnuparallel to your environment
[fchen14@mike1 ~]$ which parallel
/usr/local/packages/gnuparallel/20180222/INTEL-18.0.0/bin/parallel
[fchen14@mike421 ~]$ parallel --version
GNU parallel 20180222
...
➢ Or add the below line to ~/.modules:

    module load gnuparallel/20180222/INTEL-18.0.0
Introduction to GNU Parallel

GNU Parallel Syntax
GNU Parallel Syntax

➢ Reading commands to be run in parallel from an input file:
  parallel [OPTIONS] < CMDFILE

➢ 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:**

```bash
[fchen14@mike421 ~]$ parallel echo :::: A B C
A
B
C
```

```bash
[fchen14@mike421 ~]$ parallel echo :::: `seq 1 3`
1
2
3
```

```bash
[fchen14@mike421 ~]$ parallel echo :::: {A..Z}
A
B
...
Z
```

```bash
[fchen14@mike421 test]$ ls -1 | parallel echo
2013-06-18.tgz
backups.sh
bigmem_test.pbs
...
```

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ARGLIST from file

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

[fchen14@mike421 blast]$ cd /project/fchen14/gpar/blast
[fchen14@mike421 blast]$ cat input.lst | head
data/test10.faa
data/test11.faa
...
[fchen14@mike421 blast]$ head input.lst -n 5 | parallel echo
data/test10.faa
data/test11.faa
data/test12.faa
data/test13.faa
data/test14.faa
[fchen14@mike421 blast]$ parallel echo ::::: input.lst
data/test10.faa
data/test11.faa
data/test12.faa
...

Replacement Strings

➢ ‘{}’ returns a full line read from the input source.

[fchen14@mike421 blast]$ parallel echo {} :::: data/test1.faa
data/test1.faa

➢ ‘{/**}’ removes everything up to and including the last forward slash:

[fchen14@mike421 blast]$ parallel echo {/**} :::: data/test1.faa
test1.faa

➢ ‘{///}’ returns the directory name of input line.

[fchen14@mike421 blast]$ parallel echo {///} :::: data/test1.faa
data

➢ ‘{.}’ removes any filename extension:

[fchen14@mike421 blast]$ parallel echo {.} :::: data/test1.faa
data/test1

➢ ‘{./}’ returns the basename of the input line without extension. It is a combination of ‘{/**}’ and ‘{.}’:

[fchen14@mike421 blast]$ parallel echo {./} :::: data/test1.faa
test1

➢ 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/test1.faa
  - Output file name: output/test1.out

# Process data/test1.faa and send result to output/test1.out
$ parallel echo {} output/{/}.out ::: data/test1.faa
data/test1.faa output/test1.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”:

```bash
#!/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@mike421 misc]$ parallel --wd /project/fchen14/gpar/misc ./cmd_ex.sh :::: A B C
This script uses input: A on mike421:/project/fchen14/gpar/misc
This script uses input: B on mike421:/project/fchen14/gpar/misc
This script uses input: C on mike421:/project/fchen14/gpar/misc
```

➢ Parallize the script using **ARGFILE**:

```
fchen14@mike421 misc]$ cat argfile
A
B
C
```

```
fchen14@mike421 misc]$ parallel --wd /project/fchen14/gpar/misc ./cmd_ex.sh :::: argfile
This script uses input: A on mike421:/project/fchen14/gpar/misc
This script uses input: B on mike421:/project/fchen14/gpar/misc
This script uses input: C on mike421:/project/fchen14/gpar/misc
```

➢ 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 \(\geq 20161022\) to avoid a “Max jobs to run” bug

    ```bash
    [fchen14@mike421 test]$ parallel --version
    GNU parallel 20161022
    ...
    ```

- **-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

- **--slf filename (--sshloginfile filename)**
  - File with sshlogins. The file consists of sshlogins on separate lines. Empty lines and lines starting with '#' are ignored.
  - Look at “man parallel” for more detailed explanation.
  - A typical example on HPC/LONI clusters while running batch jobs:
    --slf $PBS_NODEFILE
  - Recall what is inside $PBS_NODEFILE?

```
[fchen14@mike421 laplace]$ cat $PBS_NODEFILE
mike421
mike421
...
mike421
mike429
mike429
...
mike429
```

16 repeats (cores) on mike421
16 repeats (cores) on mike429
Common OPTIONS --wd

- --wd mydir (--workdir mydir)
  - Designate the working directory of your commands.
  - A typical value can be $PBS_O_WORKDIR
Common OPTIONS --progress

- **--progress**
  - Show progress of computations.
  - 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:
    ```bash
    [fchen14@mike421 ~]$ parallel --progress echo ::: A B C
    ```

    Computers / CPU cores / Max jobs to run
    1:local / 16 / 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 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@mike421 misc]$ parallel --joblog logfile exit ::: 1 2 0 0
[fchen14@mike421 misc]$ cat logfile
```

<table>
<thead>
<tr>
<th>Seq</th>
<th>Host</th>
<th>Starttime</th>
<th>JobRuntime</th>
<th>Send</th>
<th>Receive</th>
<th>Exitval</th>
<th>Signal</th>
<th>Command</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>:</td>
<td>1477514132.358</td>
<td>0.019</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>exit 1</td>
</tr>
<tr>
<td>2</td>
<td>:</td>
<td>1477514132.375</td>
<td>0.003</td>
<td>0</td>
<td>0</td>
<td>2</td>
<td>0</td>
<td>exit 2</td>
</tr>
<tr>
<td>3</td>
<td>:</td>
<td>1477514132.376</td>
<td>0.002</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>exit 0</td>
</tr>
<tr>
<td>4</td>
<td>:</td>
<td>1477514132.377</td>
<td>0.003</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>exit 0</td>
</tr>
</tbody>
</table>
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.
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Serial Jobs Example
Distribute Serial Jobs - Run blast

```bash
#!/bin/bash
#PBS -l nodes=2:ppn=16
#PBS -l walltime=1:00:00
#PBS -A hpc_hpcadmin3
#PBS -q workq
#PBS -N gpl_ser_blast
#PBS -o gpl_ser_blast.out
#PBS -e gpl_ser_blast.err
JOBS_PER_NODE=16
export WDIR=/project/$USER/distribution_workload/blast
cd $WDIR
# create a folder output
mkdir -p output
# the parallel command
parallel --progress \ # shows the progress
  --joblog logfile \ # specify a job logfile
  -j $JOBS_PER_NODE \ # specify the jobs per node
  --slf $PBS_NODEFILE \ # distribute workload among allocated nodes
  --workdir $WDIR \ # specify the working directory of the script
  $PBS_O_WORKDIR/cmd_ser_blast.sh {{}} {{./}} :::: input.lst
# script_to_parallize input output :::: ARGFILE
```

[fchen14@mike421 blast]$ head input.lst
```
data/test10.faa
data/test11.faa
data/test12.faa
data/test13.faa
data/test14.faa
data/test15.faa
...```
The script to Run blast

```bash
#!/bin/bash
FILE=${(eval echo $1)} # process the input $1 when there is bash variable
echo "using input: $FILE"
TIC=`date +%s.%N`
blastp -query $FILE -db db/img_v400_PROT.00 -out output/$2.out \
    -outfmt 7 -max_target_seqs 100 -num_threads 1
TOC=`date +%s.%N`
J1_TIME=`echo "$TOC - $TIC" | bc -l`
echo "This srunner took=$J1_TIME sec using $FILE on $HOSTNAME"
```

$1 from {}

$2 from {/.}
Introduction to GNU Parallel

Multi-Threaded Example
Distribute Multi-Threaded Jobs - blast

➢ Distribute Multi-Threaded jobs is very similar to the pure serial job example, the only difference is JOBS_PER_NODE...
   - JOBS_PER_NODE=CPU_CORES_PER_NODE / NUM_THREADS_PER_JOB
➢ If each job uses 2 threads, each node on SuperMike2 has 16 cores, then
   - JOBS_PER_NODE=16/2=8
➢ PBS script (#PBS comments omitted):

```bash
JOBS_PER_NODE=8
export WDIR=/project/$USER/distribution_workload/blast
cd $WDIR
mkdir -p output
NTHREADS=2
parallel --progress \
  -j $JOBS_PER_NODE \
  --slf $PBS_NODEFILE \
  --workdir $WDIR \
  $PBS_O_WORKDIR/cmd_mt_blast.sh {} {/.} $NTHREADS \
::: input.lst
```

changes needed compared to serial version
The script to Run Multi-Threaded blast

#!/bin/bash
# This is the script we want to parallize and distribute
echo "using input: $1, output $2.out"
FILE=$(eval echo $1)
echo "using input: $FILE"
TIC=`date +%s.%N`
blastp -query $FILE \
    -db db/img_v400_PROT.00 \
    -out output/$2.out \
    -outfmt 7 -max_target_seqs 100 \
    -num_threads $3
TOC=`date +%s.%N`
J1_TIME=`echo "$TOC - $TIC" | bc -l`
echo "This serrun took=$J1_TIME sec using $FILE with $3 threads on $HOSTNAME"
Introduction to GNU Parallel

Multi-Process (MPI) Example
This section describes how to distribute small MPI jobs.

Example problem - Laplacian Solver

- Solves a 2D Laplacian equation on a 2D grid
- Use 4096x4096 2D grid, run 2000 iterations

\[ A_{k+1}(i,j) = \frac{A_k(i-1,j) + A_k(i+1,j) + A_k(i,j-1) + A_k(i,j+1)}{4} \]
### Graphical representation for Jacobi iteration

#### Array: `told`

<table>
<thead>
<tr>
<th></th>
<th>1.0</th>
<th>1.0</th>
<th>1.0</th>
<th>1.0</th>
<th>1.0</th>
<th>1.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.0</td>
<td>0.0</td>
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<td>1.0</td>
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<td>1.0</td>
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</tr>
<tr>
<td>1.0</td>
<td>0.0</td>
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<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
</tr>
</tbody>
</table>

#### Array: `t(new)`

<table>
<thead>
<tr>
<th></th>
<th>1.0</th>
<th>1.0</th>
<th>1.0</th>
<th>1.0</th>
<th>1.0</th>
<th>1.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.0</td>
<td>0.5</td>
<td>0.375</td>
<td>0.344</td>
<td>0.336</td>
<td>0.334</td>
<td>0.333</td>
</tr>
<tr>
<td>1.0</td>
<td>0.375</td>
<td>0.188</td>
<td>0.133</td>
<td>0.117</td>
<td>0.113</td>
<td>0.112</td>
</tr>
<tr>
<td>1.0</td>
<td>0.344</td>
<td>0.133</td>
<td>0.066</td>
<td>0.046</td>
<td>0.04</td>
<td>0.038</td>
</tr>
<tr>
<td>1.0</td>
<td>0.336</td>
<td>0.117</td>
<td>0.046</td>
<td>0.023</td>
<td>0.016</td>
<td>0.013</td>
</tr>
<tr>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
</tr>
</tbody>
</table>

\[ t_{i,j} = 0.25 \times (1.0 + 1.0 + 0.0 + 0.0) = 0.5 \]
\[ dt = \max(t_{i,j} - told_{i,j}) = 0.5 - 0 = 0.5 \]
PBS Script for Distributing MPI Jobs

#!/bin/bash
#PBS -l nodes=2:ppn=16
#PBS -l walltime=1:00:00
#PBS -A hpc_hpcadmin3
#PBS -q workq
#PBS -N gpl_mpi

JOBS_PER_NODE=4  # uses 4 jobs per node, on SuperMike2: 16/4=4
echo "JOBS_PER_NODE="$JOBS_PER_NODE
NPROCS=4        # uses 4 mpi processes per MPI job
WDIR=/project/$USER/distribution_workload/\laplace

parallel --progress \
-j $JOBS_PER_NODE \
--slf $PBS_NODEFILE \
--workdir $WDIR \
$PBS_O_WORKDIR/cmd_mpi.sh {} $NPROCS :::: input.lst
# script_name input num_mpi_process

[fchen14@mike421 laplace]$ cat input.lst
/project/$USER/distribution_workload/\laplace/input/input1
/project/$USER/distribution_workload/\laplace/input/input2
/project/$USER/distribution_workload/\laplace/input/input3
...

[fchen14@mike421 laplace]$ cat \
/project/$USER/distribution_workload/\laplace/input/input1
4096 4096 2 2 0.08 20000 0 0
The Script to Run MPI Laplacian Solver

#!/bin/bash

echo "using input: $1"

TIC=`date +%s.N`

## get an input from $1 (input.lst)
FILE=$(eval echo $1)

param=`cat ${FILE}`

echo param=$param

# get number of mpi process from $2 ($NPROCS)
mpirun -np $2 ./lap_mpi $param

TOC=`date +%s.N`

J1_TIME=`echo "$TOC - $TIC" | bc -l`

echo "This mpirun took=$J1_TIME sec on $HOSTNAME"
Proper usage of GNU Parallel

Distributed Workload with GNU Parallel
Memory Consideration

➢ **Relationship between node memory and cores**
  – Rule of Thumb: cannot exceed the available memory on a node

- **Available node memory 32 GB**
  - 2 GB mem per task, How many tasks per node?

- **Available node memory 32 GB**
  - 6 GB mem per task, How many tasks per node?

- **Avoid this situation, hard to calculate/predict memory usage**
GNU Parallel spawns the next job when one finishes - keeping the CPUs active and thus saving time.

**Without Load Balancing**

- Cpu0
  - Job 1
  - Job 2
  - Job 3
  - Job 4
  - Waste

- Cpu1
  - Job 5
  - Job 6
  - Job 7

- Cpu2
  - Job 8

**With Load Balancing**

- Cpu0
  - Job 1
  - Job 2
  - Job 3
  - Job 4
  - Job 5

- Cpu1
  - Job 6
  - Job 7

- Cpu2
  - Job 8
  - Job 9
  - Job 10
  - Waste

**Execution Time**

- 03/11/2020
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.

Ref: https://en.wikipedia.org/wiki/Granularity_(parallel_computing)
### Components of a Task (Process)

<table>
<thead>
<tr>
<th>Component</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Load program instructions</td>
<td></td>
</tr>
<tr>
<td>Allocate memory space</td>
<td></td>
</tr>
<tr>
<td>Load program data</td>
<td></td>
</tr>
<tr>
<td>Perform Computation on CPU cores</td>
<td></td>
</tr>
<tr>
<td>Release memory, data, other resources</td>
<td></td>
</tr>
</tbody>
</table>

#### Process Start Overhead

- **Real computation work**
- **Process end overhead**
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
Summary

➢ Why need GNU Parallel?
➢ Basic syntax of GNU Parallel and examples
➢ Use it wisely