Scientific Computing RTP

Part 2: Batch processing and HPC

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Shared hardware

- Funded centrally by the university, not individual groups/departments

- Subject to resource restrictions
  - Maximum runtime (currently 48 hours)
  - Maximum number of CPUs, RAM etc per user

- Restrictions are in place to prevent monopolization by single user

- Good codes support checkpoint & restart
  - e.g. dump parameters at final iteration and restart from these
  - e.g. store state of simulation and read on restart
Cluster of Workstations (CoW)

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The Cluster of Workstations (CoW)

- System for batch processing of calculations
- Mostly suitable for serial, low I/O computations
  - Generally available taskfarm queue (shared)
  - Some research groups have their own servers/nodes (not shared)

```bash
#!/bin/bash
#SBATCH --ntasks=1
#SBATCH --mem=512
#SBATCH --time=48:00:00
module load GCC/7.3.0-2.30 OpenMPI/3.1.1 Python/3.6.6
python my_cool_thing.py
```

Submit job script to the CoW, it will run on the next available node in the queue
High Performance Computing

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Orac (2017)

- **Compute nodes**: Lenovo NeXtScale nx360 M5 servers with 2 x Intel Xeon E5-2680 v4 (Broadwell) 2.4 GHz 14-core processors; 28 cores per node; 84 nodes; 2352 cores; 128 GB 2400 MHz DDR4 memory per node

- **Interconnect**: Intel Omni-Path X16, 100 Gbit/s with 1 μs latency

- **Phi node testbed**: Intel Xeon Phi 7250F (Knights Landing) 1.4 GHz 68-core host processors; 4 nodes; 272 cores; 96 GB 2400 MHz DDR4 memory per node

- **OpenPOWER node testbed**: 2 x IBM POWER8 3.259 GHz 8-core processors; 16 cores per node; 1 node; 256 GB DDR4 memory; 4 x NVIDIA P100 GPGPUs (SXM2 NVLink-enabled)
Tinis (2015)

- **Compute nodes**: Lenovo NeXtScale nx360 M5 servers with 2 x Intel Xeon E5-2630 v3 2.4 GHz (Haswell) 8-core processors; 16 cores per node; 203 nodes; 3488 cores; 64 GB DDR4 memory per node / 4 GB per core

- **Interconnect**: QLogic TrueScale InfiniBand, 40GB/s with 1 μs latency

- **GPU nodes**: 8 x NVIDIA Tesla K80 GPU cards; 2 GPU cards per node; 4 GPUs per node; 4 nodes; 64 GB DDR4 memory per node

- **Fat nodes**: 128 x Intel Xeon E7-4809 v3 2.0 GHz Haswell cores; 32 cores per node; 4 nodes; 1 TB DDR3 memory per node; 1 x NVIDIA GRID K2 GPU per node
Parallel code (Python)

- Tinis and Orac are primary for parallel computing

```python
from mpi4py import MPI
import numpy as np

# Number of points to integrate over
n = 1400

comm = MPI.COMM_WORLD
my_rank = comm.Get_rank()
p = comm.Get_size()

i1 = int(n/p) * my_rank # Current rank starts at point number i1
i2 = int(n/p) * (my_rank+1).

# Initialise an MPI communicator
# Get the rank of the current process
# Get the size of the current communicator

# Every rank computes 4/(1+x^2) at points in the segment of [0,1] it has been assigned
i = int(n/p) * (my_rank+1).
tot = 0.0
for i in range(i1,i2):
    x = (i+0.5)/n
    tot += 4.0/(1.0+x**2)
```
Parallel code (Python)

- Parallel code requires *communications*

- Part of the calculation cannot be done in parallel

```python
if (my_rank != 0):
    # Everyone but rank 0 sends to rank 0
    comm.send(tot, dest=0, tag=999)

else:
    # Rank 0 has to receive p-1 messages
    for partner in range(1, p):
        # Add what rank 0 received into the total
        tot += comm.recv(source=partner, tag=999)

    tot = tot/n
    # This is only done by rank 0 - serial
    print("Final result = %s ", tot)  # and prints result
```
Amdahl’s Law

- Consider a code with a *serial fraction* of $F$
- Limits parallel speedup
PX425 – High Performance Computing

- Covers code optimisation and parallel programming in C
- 15 lectures + 5 two-hour workshops starting this week
- OpenMP – multithreaded parallelism for multicore machines
- MPI – message passing parallelism for clusters
- Contact Dr Nick Hine (N.D.M.Hine@Warwick.ac.uk) if wanting to audit or take for credit (7.5 CATS)
Serial jobs on the HPC clusters

- Individual serial runs should be run on a dedicated machine or submitted to the CoW
- Workflows involving hundreds or even thousands of serial runs can and should be submitted to the HPC clusters as a single job

```bash
#!/bin/bash
#SBATCH --nodes=2
#SBATCH --ntasks-per-node=28
#SBATCH --time=08:00:00
#SBATCH --mem-per-cpu=4571

module load parallel intel/2017.4.196-GCC-6.4.0-2.28 impi/2017.3.196 Python/3.6.6

MY_PARALLEL_OPTS="-N 1 --delay .2 -j $SLURM_NTASKS --joblog parallel-$SLURM_JOBID.log"
MY_SRUN_OPTS="-N 1 -n 1 --exclusive"
MY_EXEC="python my_code.py {1}"

parallel $MY_PARALLEL_OPTS srun $MY_SRUN_OPTS $MY_EXEC ::: {0..99}
```
Research Software Engineering

- warwick.ac.uk/rse/
- More detailed talk from Chris Brady/Heather Ratcliffe
Take home message

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