MPI on Cirrus and ARCHER2











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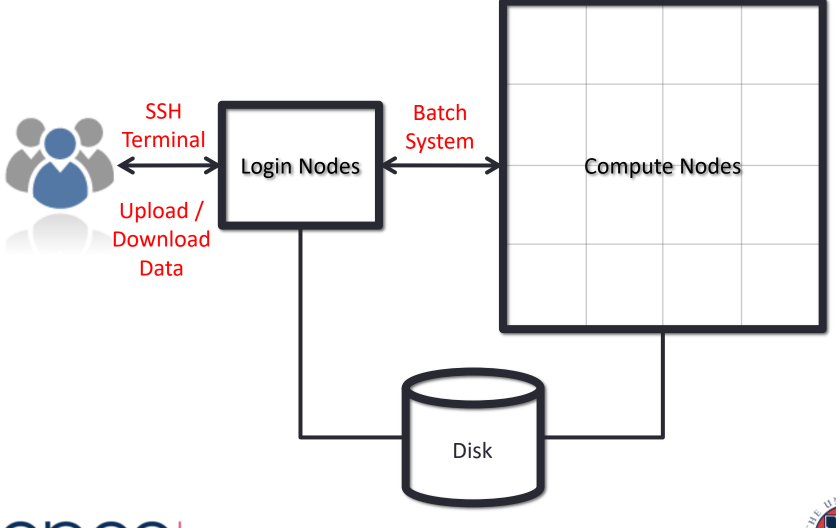


Access

- ARCHER2: ssh -XY user@login.archer2.ac.uk
- Cirrus: ssh -XY user@login.cirrus.ac.uk
- You can access EPCC systems using ssh from anywhere
 - Linux: terminal + command line
 - Mac: Mac terminal + command line + enable an X server (e.g. xquartz) to display any graphics
 - Windows: need to install an ssh program such as MobaXterm
- This gives you access to a login node
 - a few login nodes, users assigned to different nodes depending on system load
 - login nodes shared by all users
 - the many hundreds of compute nodes are accessed exclusively via the SLURM batch sys
 - on ARCHER2, MPI programs can only be run on the compute nodes
- Full instructions on access & login in separate document on course web pages



Typical HPC system layout





Useful files and templates

- Take a copy of MPP-templates.tar
 - see the course web pages
- unpack: tar -xvf MPP-templates.tar
- Crib sheets for MPI programs available on course web pages





Setting up Cirrus environment

- Load the Intel Compilers
 - module load intel-compilers-19
- Load the Message-Passing Toolkit
 - module load mpt
- To automate, add these lines to your ".bash_profile" file

```
[user@cirrus] emacs -nw ~/.bash profile
```





Compiling MPI Programs on Cirrus

• C programmers use: mpicc -cc=icc

• C++ programmers use: mpicxx -cxx=icpc

Fortran programmers use: mpif90

- There is nothing magic about these MPI compilers!
 - simply wrappers which automatically include various libraries etc
 - compilation done by standard (e.g. Intel) compilers
 - · icc, icpc and ifort
- You can use the supplied Makefiles for convenience

```
- make -f Makefile_c
```

- make -f Makefile cc
- make -f Makefile_f90
- Easiest to make a copy of one of these called "Makefile"
 - also need to change the line "MF=" in the Makefile itself





Running interactively on Cirrus

- Timings will not be reliable
 - shared with other users, many more processes than processors
 - but **very useful** during development and for debugging
- mpirun -n 4 ./mpiprog.exe
 - runs your code on 4 processes
- NOTE
 - output might be buffered
 - if your program crashes, you may see no output at all
- It may help to explicitly flush prints to screen
 - fflush(stdout);
 - FLUSH(6)





Running batch jobs on Cirrus

- Run via a batch system
 - Cirrus uses SLURM: you submit a batch script that launches your program
- In MPP-templates/ is a standard batch script: cirrusmpi.job
 - set up to run a program called "hello" on 4 processors
- To run on 4 processors: sbatch cirrusmpi.job
 - runs executable called "hello"
 - output will appear in a file called hello-xxxx.out
 - errors will appear in a file called hello-xxxx.err
 - can follow job progress using squeue or squeue -u <user>
 - e.g. if your username is s1234567 use: squeue -u s1234567
 - full instructions included as comments in the template
- MSc students should alter charging: dc032 -> dc032-user
 - e.g. #SBATCH --account=dc032-s1234567





Cirrus idiosyncrasies

 By default, MPI wrappers are not in your path user@cirrus\$ mpicc
 -bash: mpicc: command not found

- To access correct version: module load mpt
 -defaults to GNU compilers: gcc, g++ and gfortran
 -in SLURM batch system, job launcher is called srun
- Intel compilers: module load intel-compilers-19

 can add these to end of your .bash_profile file in home directory
 to check you have the right version (similarly for mpif90)
 user@cirrus\$ which mpicc
 /opt/hpe/hpc/mpt/mpt-2.22/bin/mpicc

```
-to use Intel C compiler: mpicc -cc=icc
-to use Intel C++ compiler: mpicxx -cxx=icpc
-to use Intel Fortran compiler: mpif90 -fc=ifort
```





Compiling MPI Programs on ARCHER2

- C programmers use cc
- C++ programmers use CC
- Fortran programmers use ftn
- There is nothing magic about these MPI "compilers"!
 - simply wrappers which automatically include various libraries etc
 - compilation done by standard (Cray) compilers
 - clang, clang++ and crayftn
- You can use the supplied Makefiles (C, C++, Fortran) for convenience

```
make -f Makefile_c
make -f Makefile_cc
make -f Makefile f90
```

- Easiest to make a copy of your choice called "Makefile"
 - e.g. cp Makefile_c Makefile
 - also need to change the first line "MF=" in the Makefile itself
 - then you can just type "make"





ARCHER2 idiosyncrasies

- Not possible to run MPI programs directly on login nodes
- Can be a substantial delay in batch queues
 - have dedicated queues for the course for more rapid turnaround
 - or use the short queue for development
- Cannot run from the home file system
 - on logging in, your working directory is /home/ta020/ta020/username/
 - compute nodes can only see the /work/ file system, not /home/
- Recommendation
 - do everything in /work/
 - i.e. change directory to /work/ta020/ta020/username/





Running on ARCHER2 back-end

- Run via a batch system
 - on ARCHER2 we use SLURM
 - submit a script that then launches your program
- In MPP-templates/ is a standard batch script: archer2mpi.job
 - set up to run a program called **hello** in the reserved queue for Wednesday
 - requires editing for different programs or queues on other days
- Submit: sbatch archer2.job
 - runs on 4 processes of a single ARCHER2 node (each node has 128 CPU-cores)
 - output will appear in a file called hello-xxxxxx.out
 - can follow job progress using command: squeue -u \$USER
 - full instructions included as comments in the template
- If there is no reserved queue
 - batch script has instructions on how to use short and standard queues





Compiling MPI Programs on NEXTGenIO

C programmers use: mpicc

• C++ programmers use: mpicxx

Fortran programmers use: mpif90

- There is nothing magic about these MPI compilers!
 - simply wrappers which automatically include various libraries etc
 - compilation done by standard (here, GNU) compilers
 - gcc, g++, gfortran
- You can use the supplied Makefiles for convenience

```
- make -f Makefile c
```

- make -f Makefile cc
- make -f Makefile_f90
- Easiest to make a copy of one of these called "Makefile"
 - also need to change the line "MF=" in the Makefile itself





Running interactively on NEXTGenIO

- Timings will not be reliable
 - shared with other users, many more processes than processors
 - but very useful during development and for debugging
- mpirun -n 4 ./mpiprog.exe
 - runs your code on 4 processes
- NOTE
 - output might be buffered
 - if your program crashes, you may see no output at all
- It may help to explicitly flush prints to screen
 - fflush(stdout);
 - FLUSH(6)





C++ Interface

- MPI is not an OO interface
 - however, can be called from C++
- Originally had different function calls, e.g.

```
- MPI::Intracomm comm;
- ...
- MPI::Init();
- comm = MPI::COMM_WORLD;
- rank = comm.Get_rank();
- size = comm.Get_size();
```

- Compiler is called CC
 - See hello.cc and Makefile_cc

C++ interface is now removed

Must therefore cross-call to C





Documentation

- MPI Standard available online
 - See: http://www.mpi-forum.org/docs/
 - currently version 3.1

- Available in printed form
 - http://www.hlrs.de/mpi/mpi31/

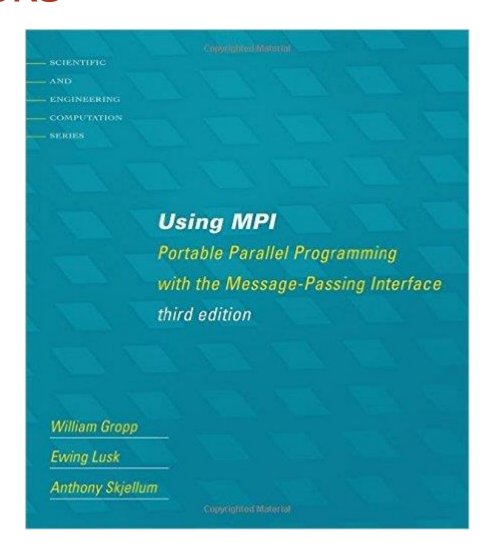


- Man pages available on Cirrus and ARCHER
 - must use the C style of naming: man MPI_Routine_name, e.g.:
 - user@computer\$ man MPI_Init





MPI Books







Exercise: Hello World

The minimal MPI program

- See Exercise 1 on the exercise sheet
- Write an MPI program that prints a message to the screen
- Main purpose is to get you compiling and running parallel programs on cirrus
 - both on login node and on compute nodes via SLURM and sbatch
 - also illustrates the SPMD model and use of basic MPI calls
- We supply some very basic template code
 - you need to add appropriate calls to compute rank and size



