Parallel Programming

Libraries and implementations
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Outline

• How we manage software packages & libraries on ARCHER

• MPI – distributed memory de-facto standard
  - Using MPI

• OpenMP – shared memory de-facto standard
  - Using OpenMP

• Other parallel programming technologies
  - CUDA, OpenCL, OpenACC

• Examples of common scientific libraries
The module environment

Managing software packages and libraries
Module environment

user@eslogin001:~> module list
Currently Loaded Modulefiles:
1) modules/3.2.10.2
2) eswrap/1.3.3-1.020200.1278.0
3) switch/1.0-1.0502.57058.1.58.ari
4) craype-network-aries
5) craype/2.4.2
6) cce/8.4.1
7) cray-libsci/13.2.0
8) udreg/2.3.2-1.0502.9889.2.20.ari
9) rca/1.0.0-2.0502.57212.ari
10) atp/1.8.3
11) PrgE56
12) pbs/12.2.401.141761
13) craype-ivybridge
14) cray-mpich/7.2.6
15) packages-archer
16) bolt/0.6

- The module environment allows you to easily load different packages and manage different versions of packages.

- Via the *module* command
  - *List loaded modules, view available modules, load and unload modules*
Using the module environment

<table>
<thead>
<tr>
<th>Command</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>module avail</code></td>
<td>List available modules</td>
</tr>
<tr>
<td><code>module load fftw</code></td>
<td>Load the fftw module</td>
</tr>
<tr>
<td><code>module unload fftw</code></td>
<td>Unload the fftw module</td>
</tr>
<tr>
<td><code>module load fftw/2.1.5.7</code></td>
<td>Load a specific version of the fftw module</td>
</tr>
<tr>
<td><code>module switch fftw/2.1.5.7 fftw/3.3.4.9</code></td>
<td>Switch to a different version of the fftw module</td>
</tr>
<tr>
<td><code>module swap PrgEnv-cray PrgEnv-gnu</code></td>
<td>Swap the current module environment</td>
</tr>
</tbody>
</table>
MPI Library

Distributed, message-passing programming
Message-passing concepts
What is MPI?

- Message Passing Interface
- MPI is not a programming language
  - There is no such thing as an *MPI compiler*
- MPI is available as a *library* of function/subroutine calls
  - The library implements the MPI standard

- The C or Fortran compiler knows nothing about what MPI actually does
  - Just the prototype/interfaces of the functions/subroutine
  - It is just another library
The MPI standard

• MPI itself is a standard
• Agreed upon by approx 100 representatives from about 40 organisations (the MPI forum)
  - Academics
  - Industry
  - Vendors
  - Application developers

• First standard (MPI version 1.0) drafted in 1993
  - We are currently on version 3
  - Version 4 is being drafted
MPI Libraries

- The MPI forum defines the standard and vendors/open source developers then actually implement this.

- There are a number of different implementations but all should support version 2.0 or 3.0.
  - As with compilers there are variations in implementation details but all features in the standards should work.
  - Examples: MPICH and OpenMPI.
  - Cray-MPICH on ARCHER which implements version 3.1 of the standard (optimised for Cray machines, specifically the interconnect).
Features of MPI

• MPI is a portable library used for writing parallel programs using the message passing model
  - You can expect MPI to be available on any HPC platform you use
  - Aids portability between HPC machines and is trivial to install on local clusters

• Based on a number of processes running independently in parallel
  - The HPC resource provides the command to launch the processes in parallel (i.e. `aprun` or `mpiexec`)
  - Can think of each process as an instance of your executable communicating with other instances
Explicit Parallelism

• In message-passing all the parallelism is explicit
  - The program includes specific instructions for each communication
  - What to send or receive
  - When to send or receive
  - Synchronisation

• It is up to the developer to design the parallel decomposition and implement it
  - How will you divide up the problem?
  - When will you need to communicate between processes?
Supported features

• Point to point communications
  - Communications involving two processes; a sender and receiver
  - Wide variety of semantics involving non-blocking communications
  - Other aspects such as wildcards & custom data types

• Collective communications
  - Communication that involves many processes
  - Implements all the collective communications we saw in the programming models lecture and many more
  - Also supports non-blocking communications and custom data types
Example: MPI Hello World

#include "mpi.h"

int main(int argc, char* argv[]) {
    int size, rank;

    MPI_Init(&argc, &argv);
    MPI_Comm_size(MPI_COMM_WORLD, &size);
    MPI_Comm_rank(MPI_COMM_WORLD, &rank);

    printf("Hello world - I'm rank %d of %d\n", rank, size);

    MPI_Finalize();
    return 0;
}
OpenMP

Shared-memory parallelism using directives
Shared-memory concepts

- Threads “communicate” by having access to the same memory space
  - Any thread can alter any bit of data
  - No explicit communications between the parallel tasks
OpenMP

- Open Multi Processing
  - Application programming interface (API) for shared variable programming
- Set of extensions to C, C++ and Fortran
  - Compiler directives
  - Runtime library functions
  - Environment variables
- Not a library interface like MPI
- Uses directives, which are a special line in the source code with a meaning understood by the compilers
  - Ignored if OpenMP is disabled and it becomes regular sequential code
- This is also a standard (http://openmp.org)
Features of OpenMP

• Directives define parallel regions in the code
  - OpenMP threads are active in these regions and divide the workload amongst themselves

• The compiler needs to understand what OpenMP does
  - It is responsible for producing the parallel code
  - OpenMP supported by all common compilers used in HPC

• Parallelism less explicit than MPI
  - You just specify what parts of the program you want to run in parallel

• OpenMP version 4.5 is the latest version

• Can be used to program the Xeon Phi
Loop-based parallelism

- The most common form of OpenMP parallelism is to parallelise the work in a loop
  - The OpenMP directives tell the compiler to divide the iterations of the loop between the threads

```c
#pragma omp parallel shared(a,b,c) private(i)
{
    #pragma omp for schedule(dynamic) nowait
    for (i=0; i < N; i++) {
        c[i] = a[i] + b[i];
    }
}
```
Addition example

\[ \text{asum} = 0.0 \]

```c
#pragma omp parallel \
shared(a,N) private(i) \
reduction(+:asum)
{
    #pragma omp for
    for (i=0; i < N; i++)
    {
        asum += a[i];
    }
}
printf("asum = \%f\n", asum);
```

Diagram:

- `P0(T0)` to `P0(T1)`
- Loop: `i = istart,istop`
- `myasum += a[i]` end loop

- `asum`
Other parallel programming technologies

Programming accelerators and less common technologies
CUDA

- CUDA is an Application Program Interface (API) for programming NVIDIA GPU accelerators
  - Proprietary software provided by NVIDIA. Should be available on all systems with NVIDIA GPU accelerators
  - Write GPU specific functions called *kernels*
- Launch kernels using syntax within standard C programs
- Includes functions to shift data between CPU and GPU memory
- Similar to OpenMP programming in many ways in that the parallelism is implicit in the kernel design and launch
OpenCL

• An open, cross-platform standard for programming accelerators
  - includes GPUs, e.g. from both NVIDIA and AMD
  - also Xeon Phi, Digital Signal Processors, ...

• Comprises a language + library

• Harder to write than CUDA if you have NVIDIA GPUs
  - but portable across multiple platforms
  - although maintaining performance is difficult
Other parallel implementations

- Partitioned Global Address Space (PGAS)
  - Coarray Fortran, Unified Parallel C, Chapel

- Cray SHMEM, OpenSHMEM
  - Single-sided communication library

- OpenACC
  - Directive-based approach for programming accelerators
Common scientific parallel libraries

Two examples commonly used on HPC machines
PETSc

• Portable Extensible Toolkit for Scientific Computation
  • Suite of data structures & routines for the parallel and scalable solution of PDEs
  • The programmer uses the library framework itself which under the hood will use parallel technologies MPI, OpenMP and/or CUDA.

• Unlike many serial libraries, you the programmer are responsible for performance & scalability.
NetCDF

- Network Common Data Form
  - Self describing, machine independent file data format and implementation that is very common for writing and reading scientific data
- Parallel version supporting parallel IO
  - Multiple processes/threads can read and write to a file concurrently
  - Built on top of MPI
- Many third party tools such as visualisation suites

- Again requires user understanding, both from the programmer and also the user (file configuration options)
Summary
Parallel and scientific libraries

• The module environment is an easy way of managing many different software packages, their dependencies and different versions.

• Distributed memory programmed using MPI
• Shared memory programmed using OpenMP
• GPU accelerators most often programmed using CUDA

• There are very many software packages installed on ARCHER, but scientific libraries often require in-depth knowledge and understanding to get good performance.