Introduction to Parallel Programming with MPI

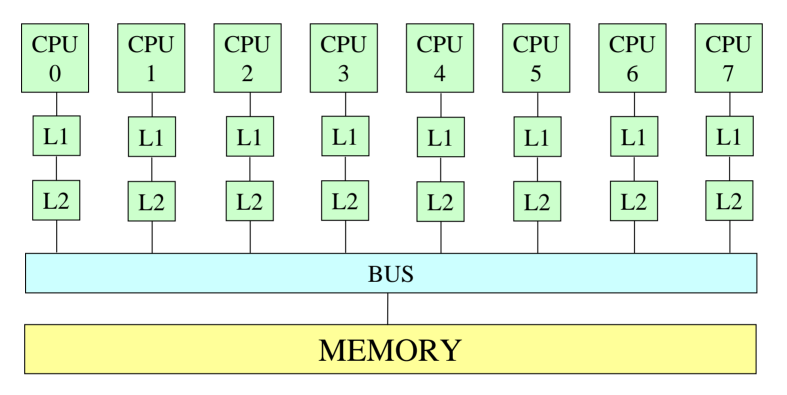
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Why Parallel Computing?

- Want to speed up a calculation.
- Solution:
 - Split the work between several processors.
- How?
 - It depends on the type of parallel computer
 - Shared memory (usually thread-based)
 - Distributed memory (process-based)
 - MPI works on all of them!

Shared memory parallelism



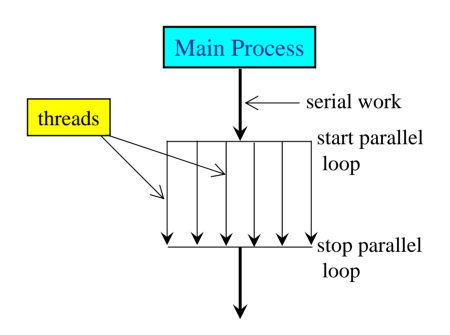
- "Classic" shared memory node
- Renewed interest due to multi-core chips
- Processors communicate through memory

Shared memory parallelism

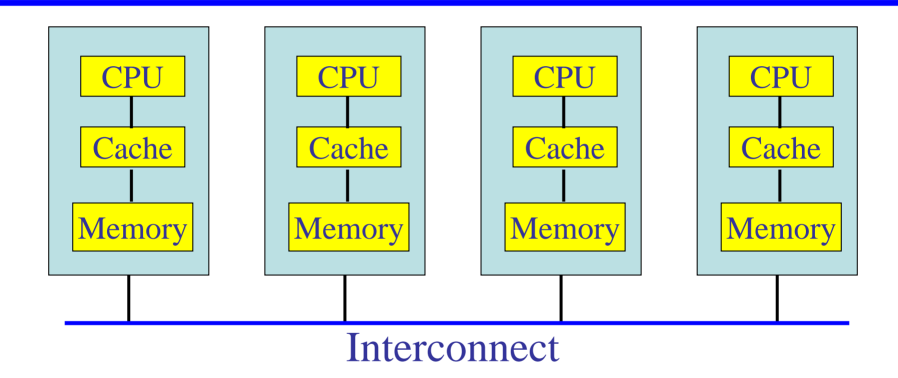
- Program runs inside a single process
- Several "execution threads" are created within that process and work is split between them.
- The threads run on different processors.
- All threads have access to the shared data through shared memory access.
- Must be careful not the have threads overwrite each other's data.

Shared memory programming

- Easy to do loop-level parallelism.
- Compiler-based automatic parallelization
 - Easy but not effective
 - Better to do it yourself with OpenMP
- Coarse-grain parallelism can be difficult
- Becoming important for multicore processors due to very low latency compared to conventional SMP nodes



Distributed memory parallelism



- The processor can only access its local memory
- Data exchange between processors must be explicit

Distributed memory parallelism

- Process-based programming.
- Each process has its own memory space that cannot be accessed by the other processes.
- The work is split between several processes.
- For efficiency, each <u>processor</u> runs a single <u>process</u>.
- Communication between the processes must be explicit, e.g. Message Passing

How to split the work between processors?

 Most widely used method for grid-based calculations:

- DOMAIN DECOMPOSITION

- Split particles in particle-in-cell (PIC) or molecular dynamics codes.
- Split arrays in PDE solvers
- etc...
- Keep it **LOCAL**

What is MPI?

- MPI stands for Message Passing Interface.
- It is a message-passing specification, a standard, for the vendors to implement.
- In practice, MPI is a set of functions (C) and subroutines (Fortran) used for exchanging data between processes.
- An MPI library exists on most, if not all, parallel computing platforms so it is highly portable.

How much do I need to know?

- MPI is small (6 functions)
 - Many parallel programs can be written with just 6 basic functions.
- MPI is large (125 functions)
 - MPI's extensive functionality requires many functions
 - Number of functions not necessarily a measure of complexity
- MPI is just right
 - One can access flexibility when it is required.
 - One need not master all parts of MPI to use it.

How MPI works

• Launch the parallel calculation with:

```
mpirun -np #proc a.out
mpiexec -n #proc a.out
```

- Copies of the same program run on each processor within its own process (private address space).
- Each processor works on a subset of the problem.
- Exchange data when needed
 - Can be exchanged through the network interconnect
 - Or through the shared memory on SMP machines (Bus?)
- Easy to do coarse grain parallelism = <u>scalable</u>

Good MPI web sites

- http://www.llnl.gov/computing/tutorials/mpi/
- http://www.nersc.gov/nusers/help/tutorials/mpi/intro/
- http://www-unix.mcs.anl.gov/mpi/tutorial/gropp/talk.html
- http://www-unix.mcs.anl.gov/mpi/tutorial/
- MPI on Linux clusters:
 - MPICH (http://www-unix.mcs.anl.gov/mpi/mpich/)
 - Open MPI (http://www.open-mpi.org/)

Structure of an MPI program

```
Program mpi code
                                          Fortran 90
  ! Load MPI definitions
    use mpi (or include mpif.h)
                                          Style!...
  ! Initialize MPI
    call MPI Init(ierr)
  ! Get the number of processes
    call MPI Comm size(MPI COMM WORLD, nproc, ierr)
  ! Get my process number (rank)
    call MPI Comm rank(MPI COMM WORLD, myrank, ierr)
    Do work and make message passing calls...
  ! Finalize
    call MPI Finalize(ierr)
end program mpi code
```

Structure of an MPI program

```
#include "mpi.h"
int main( int argc, char *argv[] )
                                       Style!...
  int nproc, myrank;
  /* Initialize MPI */
    MPI Init(&argc,&argv);
  /* Get the number of processes */
   MPI Comm size(MPI COMM WORLD,&nproc);
  /* Get my process number (rank) */
    MPI Comm rank(MPI COMM WORLD,&myrank);
    Do work and make message passing calls...
  /* Finalize */
    call MPI Finalize();
return 0;
```

Compilation

- mpich provides scripts that take care of the include directories and linking libraries
 - mpicc
 - mpiCC
 - mpif77
 - mpif90
- Otherwise, must link with the right MPI library

Makefile

Always a good idea to have a Makefile

```
%cat Makefile
CC=mpicc
CFLAGS=-0
% : %.c
$(CC) $(CFLAGS) $< -0 $@</pre>
```

mpirun and mpiexec

- Both are used for starting an MPI job
- If you don't have a batch system, use mpirun

```
__ mpirun -np #proc -machinefile mfile a.out >& out < in &
```

```
%cat mfile
machine1.princeton.edu
machine2.princeton.edu
machine3.princeton.edu
machine4.princeton.edu
```

• PBS usually takes care of arguments to mpiexec

Batch System: PBS primer

- Submit a job script: qsub script
- Check status of jobs: qstat –a (for all jobs)
- Stop a job: qdel job_id

```
### --- PBS SCRIPT ---
#PBS -l nodes=4:ppn=2,walltime=02:00:00
#PBS -q dque
#PBS -V
#PBS -N job_name
#PBS -m abe
cd $PBS_O_WORKDIR
mpiexec -np 8 a.out
```

Basic MPI calls to exchange data

Point to point: 2 processes at a time

Predefined Communicator: MPI_COMM_WORLD

Collective MPI calls

Collective calls: All processes participate

One process sends to everybody:

```
MPI_Bcast(buffer,count,datatype,root,comm,ierr)
```

All processes send to "root" process and the operation "op" is applied

MPI_Reduce(sendbuf,recvbuf,count,datatype,op,root,comm,ierr)

where **op** = **MPI_SUM**, **MPI_MAX**, **MPI_MIN**, **MPI_PROD**, **etc...**

You can create your own reduction operation with **MPI_Op_create()**

All processes send to everybody and apply the operation "op" (equivalent to an MPI_Reduce followed by an MPI_Bcast)

MPI_Allreduce(sendbuf,recvbuf,count,datatype,op,comm,ierr)

Synchronize all processes

MPI_Barrier(comm,ierr)

More MPI collective calls

All processes send a different piece of data to one single "root" process which gathers everything (messages ordered by index)

All processes gather everybody else's pieces of data

One "root" process send a different piece of the data to each one of the other processes

Each process performs a scatter operation, sending a distinct message to all the processes in the group in order by index.