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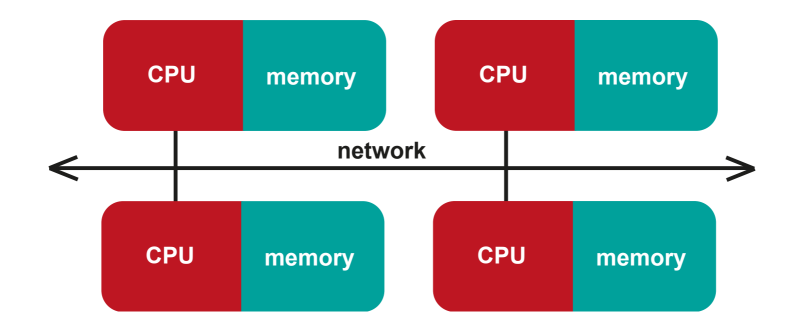
Why go parallel?

- Save time (faster rate of computation).
- Solve larger problems that don't fit on memory of single machine.
- Do different things at the same time.
- Make better use of computing resources (e.g. parallel hardware).

• Increasingly used outside of traditional physics/chemistry simulations (e.g. big data).

What is MPI?

- Message Passing Interface the standard library specification for message passing programs.
- Distributed memory programming model:



- Common implementations: **OpenMPI**, MPICH, Intel MPI, ...
- Usually used with C, C++ or Fortran.
- Download from: http://www.open-mpi.org/

Fundamentals

• All processes execute *exactly the same code*.

• Each process can access only its own memory/variables.

• Code must be specially written with this in mind.

Demo problem: 1D heat equation

Solve

$$u_t = k u_{xx},$$
 $u(x,0) = g(x) = e^{-100x^2}$ on $D = \{ -1 < x < 1, \ t > 0 \}.$

• Discretise *D* with a grid

$$x_j = -1 + j\Delta x,$$
 $t_n = n\Delta t,$ $U^{j,n} = u(x_j, t_n)$

• Use a simple explicit finite-difference scheme

$$\frac{U^{j,n+1} - U^{j,n}}{\Delta t} = k \frac{U^{j-1,n} - 2U^{j,n} + U^{j+1,n}}{(\Delta x)^2}$$

$$\implies U^{j,n+1} = U^{j,n} + \frac{k\Delta t}{(\Delta x)^2} \left(U^{j-1,n} - 2U^{j,n} + U^{j+1,n} \right)$$
Supply conditions.

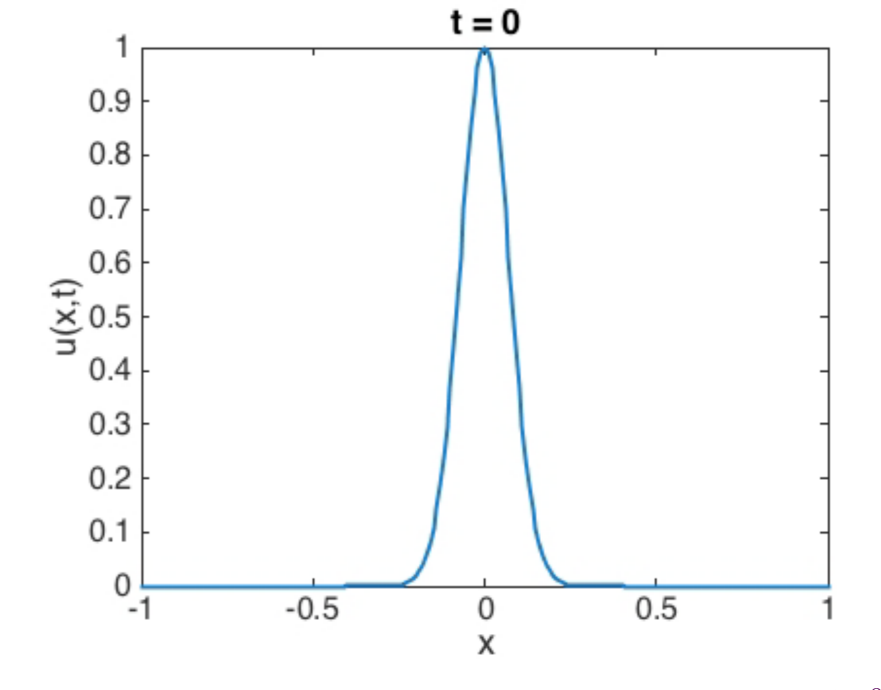
Boundary conditions

$$U^{0,n} = U^{2,n}, \quad U^{nx+1,n} = U^{nx-1,n}$$

Exact solution

$$u(x,t) = \sqrt{\frac{a}{a+4kt}} \exp\left(-\frac{x^2}{a+4kt}\right)$$

$$a = \frac{1}{100}, \ k = 1$$



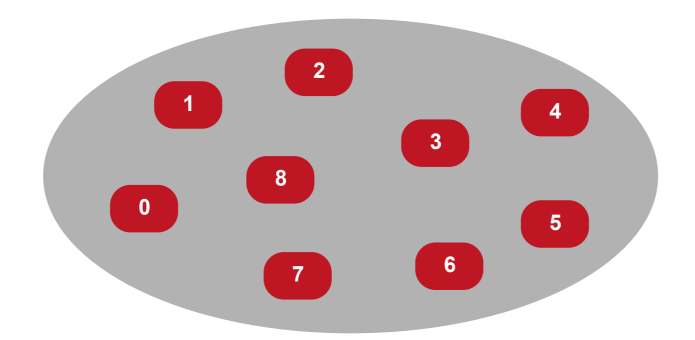
Demo code

- heat.cpp (C++) or heat.f90 (Fortran 90)
- To run from a terminal:

- On the maths network, the path is /usr/lib64/openmpi/bin/
- Main differences between MPI in C++ and Fortran:
 - 1. C++ is case sensitive (e.g. MPI Comm size).
 - 2. Fortran subroutines all have an extra output argument ierr.
 - 3. Special types like MPI_Comm are all just integer in Fortran.
 - 4. Some arguments need to be *references* in C++.
 - 5. Some defined constants like MPI_DOUBLE are different.

Initialising MPI

- **#include <mpi.h>** include file, always required.
- MPI_Init(NULL, NULL) always required, call first and only once.
- MPI_Finalize() must be last MPI call.



- MPI_Comm_size(MPI_COMM_WORLD, &nProcs) returns the number of processes in the global "communicator".
- MPI_Comm_rank(MPI_COMM_WORLD, &myRank) returns the "rank" of the current process. this tells you your identity.

Creating a virtual topology

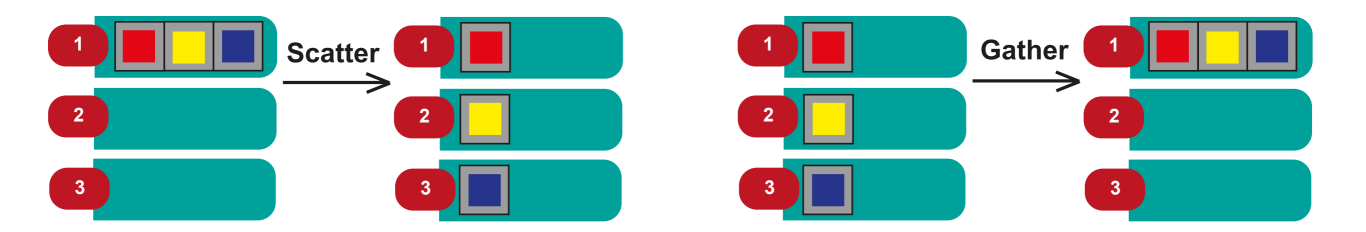
• A "communicator" is a group of processes that can communicate with each other. We use it to define a "virtual topology".



- MPI_Cart_create(MPI_COMM_WORLD,n,dims,periods, reorder,&comm)
 - make a Cartesian communicator comm with n directions, where:
 - dims number of processes in each direction;
 - periods whether periodic boundaries in each direction;
 - reorder whether ranking may be reordered.
- MPI_Comm_rank(comm, &myRank) get rank in this communicator.
- MPI_Cart_shift(comm,dirn,displ,&prvRank,&nxtRank)
 - return the ranks of nearby processes in the coordinate direction dirn, at distance displaway.

Collective communications

- Each process needs different initial conditions: we create a global **xGlob** array in process 0, then distribute chunks to the others.
- MPI_Scatter(&sendbuf, sendcnt, sendtype, &recvbuf, recvcnt, recvtype, root, comm)
 - root is the rank of the sender, sendbuf is the global array, recybuf is the local one.



- MPI_Gather(&sendbuf, sendcnt, sendtype, &recvbuf, recvcnt, recvtype, root, comm)
 - the inverse of MPI_Scatter: we use it at the end to output u.

Point-to-point communications

- MPI_Send(&buf,count,datatype,dest,tag,comm) send from the buffer buf (with count values) to rank dest.
- MPI_Recv(&buf,count,datatype,source,tag,comm,&stat) receive count values in the buffer buf from rank source.
 - These are "blocking" operations:
 - MPI_Send returns when buf is ready for re-use. (This doesn't mean that the message has been received; it may be stored in a "system buffer".)
 - MPI_Recv returns when the requested data has reached the receiving buf and can be used.

Point-to-point communications

- MPI_Send(&buf,count,datatype,dest,tag,comm) send from the buffer buf (with count values) to rank dest.
- MPI_Recv(&buf,count,datatype,source,tag,comm,&stat) receive count values in the buffer buf from rank source.

- a message can only be received if its tag (an integer) matches that in the MPI_Recv statement (or if it is set to MPI_ANY_TAG).
- the structure stat contains the source of the message and the tag: stat.MPI SOURCE and stat.MPI TAG.

Things to be aware of

- There is no requirement that each MPI_Send is matched by an equivalent MPI Recv (or alternative command).
 - There is *no overtaking*: if two messages are send from the same sender to the same destination, and both match an MPI_Recv statement, then the first message to be sent will be received first.
 - If two processes A and B both send messages to process C that match a single receive statement, then *only one* of the sends will complete.
- A process can get *stuck* on an MPI_Recv if no corresponding message is sent.

Deadlock!

- It is possible to reach "deadlock" where two (or more) processes are each waiting for the other to send or receive.
- For example, suppose there is a *circular* topology and we call:

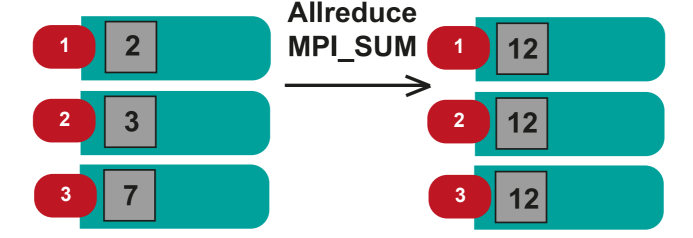
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MPI_Send(&a,1,MPI_INT,nxtRank,tag,comm);
MPI_Recv(&b,1,MPI_INT,prvRank,tag,comm,&stat);
...
```

- This is unsafe; if the system buffer runs out of memory (not under our control), then all of the processes could be stuck waiting to send.
- We can see this if we use the "synchronous send" MPI_Ssend,
 which waits until the destination process has started to receive the
 message before returning.

Collective computations

- MPI has routines for collective computation, or "reduction".
- MPI_Allreduce(&sendbuf,&recvbuf,count,datatype, op,comm)
 - perform the global operation op using sendbuf from each process, and broadcast the overall result to every recybuf.
- Operations include MPI SUM, MPI PROD, MPI MAX, MPI MIN, ...

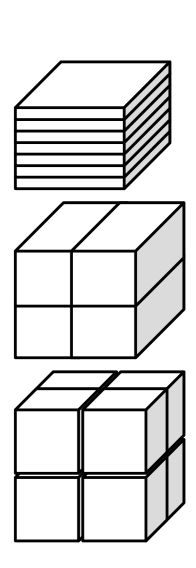
$$\frac{\mathrm{d}}{\mathrm{d}t} \int_{-1}^{1} u \, \mathrm{d}x = \int_{-1}^{1} k u_{xx} \, \mathrm{d}x = k \left[u_{x} \right]_{-1}^{1} = 0$$



Timing

• MPI_Wtime() - returns wall-clock time on the individual processor.

- Improving speed/scaling with number of processors:
 - increase fraction of program that can be parallelized;
 - balance the workload;
 - minimize communication time (decrease both amount of data and number of calls);
 - use collective communications wherever possible.



Further reading

- A useful tutorial and reference for other commands: https://computing.llnl.gov/tutorials/mpi/
- Another tutorial (more advanced topics) with examples:
 http://mpitutorial.com
- The official standard:
 http://mpi-forum.org/docs/docs.html