What is MPI?

- A message-passing library specification
 - extended message-passing model
 - not a language or compiler specification
 - not a specific implementation or product
- For parallel computers, clusters, and heterogeneous networks
- Full-featured
- Designed to provide access to advanced parallel hardware for end users, library writers, and tool developers
- Credits for Slides: Rusty Lusk, Mathematics and Computer Science Division, Argonne National Laboratory

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Where Did MPI Come From?

- Early vendor systems (Intel's NX, IBM's EUI, TMC's CMMD) were not portable (or very capable)
- Early portable systems (PVM, p4, TCGMSG, Chameleon) were mainly research efforts
 - Did not address the full spectrum of issues
 - Lacked vendor support
 - Were not implemented at the most efficient level
- The MPI Forum organized in 1992 with broad participation by:
 - vendors: IBM, Intel, TMC, SGI, Convex, Meiko
 - portability library writers: PVM, p4
 - users: application scientists and library writers
 - finished in 18 months

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Novel Features of MPI

- <u>Communicators</u> encapsulate communication spaces for library safety
- <u>Datatypes</u> reduce copying costs and permit heterogeneity
- Multiple communication <u>modes</u> allow precise buffer management
- Extensive <u>collective operations</u> for scalable global communication
- <u>Process topologies</u> permit efficient process placement, user views of process layout
- Profiling interface encourages portable tools

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MPI References

- · The Standard itself:
 - at http://www.mpi-forum.org
 - All MPI official releases, in both postscript and HTML
- · Books:
 - Using MPI: Portable Parallel Programming with the Message-Passing Interface, 2nd Edition, by Gropp, Lusk, and Skjellum, MIT Press, 1999.
 Also Using MPI-2, w. R. Thakur
 - MPI: The Complete Reference, 2 vols, MIT Press, 1999.
 - Designing and Building Parallel Programs, by Ian Foster, Addison-Wesley, 1995.
 - Parallel Programming with MPI, by Peter Pacheco, Morgan-Kaufmann, 1997.
- Other information on Web:
 - at http://www.mcs.anl.gov/mpi
 - pointers to lots of stuff, including other talks and tutorials, a FAQ, other MPI pages

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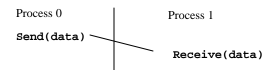
Hello (C)

```
#include "mpi.h"
#include <stdio.h>
int main( argc, argv )
int argc;
char *argv[];
   int rank, size;
   MPI_Init( &argc, &argv );
   MPI_Comm_rank( MPI_COMM_WORLD, &rank );
   MPI_Comm_size( MPI_COMM_WORLD, &size );
   printf( "I am %d of %d\n", rank, size );
   MPI_Finalize();
   return 0;
}
```

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MPI Basic Send/Receive

· We need to fill in the details in



- · Things that need specifying:
 - How will "data" be described?
 - How will processes be identified?
 - How will the receiver recognize/screen messages?
 - What will it mean for these operations to complete?

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Some Basic Concepts

- Processes can be collected into groups
- Each message is sent in a <u>context</u>, and must be received in the same context
 - Provides necessary support for libraries
- A group and context together form a <u>communicator</u>
- A process is identified by its <u>rank</u> in the group associated with a communicator
- There is a default communicator whose group contains all initial processes, called MPI_COMM_WORLD

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MPI Datatypes

- The data in a message to send or receive is described by a triple (address, count, datatype), where
- An MPI datatype is recursively defined as:
 - predefined, corresponding to a data type from the language (e.g., MPI_INT, MPI_DOUBLE)
 - a contiguous array of MPI datatypes
 - a strided block of datatypes
 - an indexed array of blocks of datatypes
 - an arbitrary structure of datatypes
- There are MPI functions to construct custom datatypes, in particular ones for subarrays

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MPI Tags

- Messages are sent with an accompanying user-defined integer tag, to assist the receiving process in identifying the message
- Messages can be screened at the receiving end by specifying a specific tag, or not screened by specifying MPI_ANY_TAG as the tag in a receive
- Some non-MPI message-passing systems have called tags "message types". MPI calls them tags to avoid confusion with datatypes

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MPI Basic (Blocking) Send

MPI_SEND(start, count, datatype, dest, tag, comm)

- The message buffer is described by (start, count, datatype).
- The target process is specified by dest, which is the rank of the target process in the communicator specified by comm.
- When this function returns, the data has been delivered to the system and the buffer can be reused. The message may not have been received by the target process.

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MPI Basic (Blocking) Receive

MPI_RECV(start, count, datatype, source, tag, comm, status)

- Waits until a matching (both source and tag) message is received from the system, and the buffer can be used
- source is rank in communicator specified by comm, or MPI_ANY_SOURCE
- tag is a tag to be matched on or MPI_ANY_TAG
- receiving fewer than count occurrences of datatype is OK, but receiving more is an error
- status contains further information (e.g. size of message)

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MPI is Simple

- Many parallel programs can be written using just these six functions, only two of which are non-trivial:
 - MPI_INIT
 - MPI_FINALIZE
 - MPI_COMM_SIZE
 - MPI_COMM_RANK
 - MPI_SEND
 - MPI_RECV

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Collective Operations in MPI

- Collective operations are called by all processes in a communicator
- MPI_BCAST distributes data from one process (the root) to all others in a communicator
- MPI_REDUCE combines data from all processes in communicator and returns it to one process
- In many numerical algorithms, SEND/RECEIVE can be replaced by BCAST/REDUCE, improving both simplicity and efficiency

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Example: PI in C - 1

```
#include "mpi.h"
#include <math.h>
int main(int argc, char *argv[])
{
   int done = 0, n, myid, numprocs, i, rc;
   double PI25DT = 3.141592653589793238462643;
   double mypi, pi, h, sum, x, a;
   MPI_Init(&argc,&argv);
   MPI_Comm_size(MPI_COMM_WORLD,&numprocs);
   MPI_Comm_rank(MPI_COMM_WORLD,&myid);
   while (!done) {
      if (myid == 0) {
          printf("Enter the number of intervals: (0 quits) ");
          scanf("%d",&n);
      }
      MPI_Bcast(&n, 1, MPI_INT, 0, MPI_COMM_WORLD);
      if (n == 0) break;
```

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Example: PI in C - 2

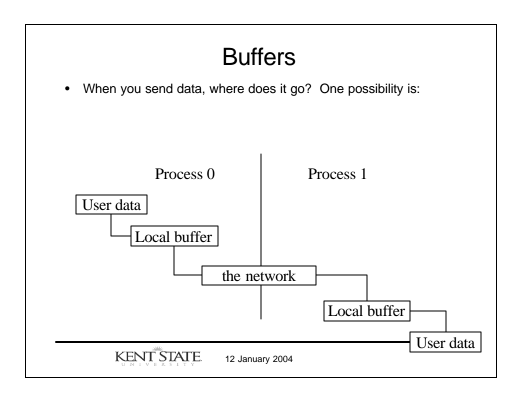
```
h = 1.0 / (double) n;
  sum = 0.0;
  for (i = myid + 1; i <= n; i += numprocs) {</pre>
    x = h * ((double)i - 0.5);
    sum += 4.0 / (1.0 + x*x);
  mypi = h * sum;
  MPI_Reduce(&mypi, &pi, 1, MPI_DOUBLE, MPI_SUM, 0,
             MPI_COMM_WORLD);
  if (myid == 0)
    printf("pi is approximately %.16f, Error is .16f\n",
            pi, fabs(pi - PI25DT));
MPI_Finalize();
return 0;
```

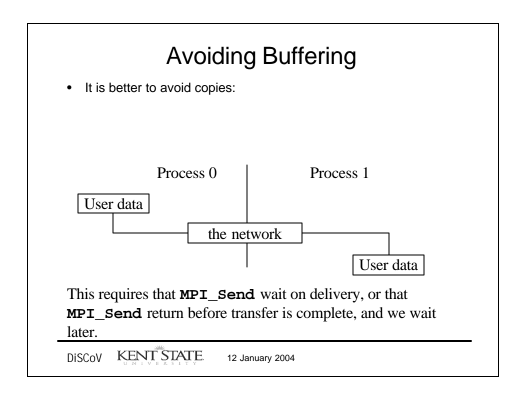
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Alternative Set of 6 Functions

- · Using collectives:
 - MPI_INIT
 - MPI_FINALIZE
 - MPI_COMM_SIZE
 - MPI_COMM_RANK
 - MPI_BCAST
 - MPI_REDUCE

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Blocking and Non-blocking Communication

- So far we have been using *blocking* communication:
 - MPI_Recv does not complete until the buffer is full (available for use).
 - MPI_Send does not complete until the buffer is empty (available for use).
- Completion depends on size of message and amount of system buffering.

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Sources of Deadlocks

- Send a large message from process 0 to process 1
 - If there is insufficient storage at the destination, the send must wait for the user to provide the memory space (through a receive)
- What happens with this code?

Process 0	Process 1	
Send(1)	Send(0)	
Recv(1)	Recv(0)	

• This is called "unsafe" because it depends on the availability of system buffers

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Some Solutions to the "unsafe" Problem

· Order the operations more carefully:

Process 0	Process 1
Send(1)	Recv(0)
Recv(1)	Send(0)

Supply receive buffer at same time as send:

 Process 0	Process 1
Sendrecv(1)	Sendrecv(0)

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More Solutions to the "unsafe" Problem

• Supply own space as buffer for send

	Process 0	Process 1	
	Bsend(1) Recv(1)	Bsend(0) Recv(0)	
Use	non-blocking operation	s:	
_	Process 0	Process 1	
	Isend(1) Irecv(1) Waitall	Isend(0) Irecv(0) Waitall	

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MPI's Non-blocking Operations

 Non-blocking operations return (immediately) "request handles" that can be tested and waited on.

```
MPI_Isend(start, count, datatype,
    dest, tag, comm, request)
MPI_Irecv(start, count, datatype,
    dest, tag, comm, request)
MPI Wait(&request, &status)
```

One can also test without waiting:
 MPI_Test(&request, &flag, status)

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Multiple Completions

It is sometimes desirable to wait on multiple requests:

```
MPI_Waitall(count, array_of_requests,
    array_of_statuses)
MPI_Waitany(count, array_of_requests,
    &index, &status)
MPI_Waitsome(count, array_of_requests,
    array_of_indices, array_of_statuses)
```

There are corresponding versions of test for each of these.

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Communication Modes

- MPI provides multiple modes for sending messages:
 - Synchronous mode (MPI_ssend): the send does not complete until a matching receive has begun. (Unsafe programs deadlock.)
 - Buffered mode (MPI_Bsend): the user supplies a buffer to the system for its use. (User allocates enough memory to make an unsafe program safe.
 - Ready mode (MPI_Rsend): user guarantees that a matching receive has been posted.
 - Allows access to fast protocols
 - undefined behavior if matching receive not posted
- Non-blocking versions (MPI_Issend, etc.)
- MPI_Recv receives messages sent in any mode.

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Other Point-to Point Features

- MPI Sendrecv
- MPI_Sendrecv_replace
- MPI_Cancel
 - Useful for multibuffering
- Persistent requests
 - Useful for repeated communication patterns
 - Some systems can exploit to reduce latency and increase performance

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MPI_Sendrecv

- Allows simultaneous send and receive
- · Everything else is general.
 - Send and receive datatypes (even type signatures) may be different
 - Can use Sendrecv with plain Send or Recv (or Irecvor Ssend_init, ...)
 - More general than "send left"

Process 0 Process 1

SendRecv(1) SendRecv(0)

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Collective Operations in MPI

- Collective operations must be called by all processes in a communicator.
- MPI_BCAST distributes data from one process (the root) to all others in a communicator.
- MPI_REDUCE combines data from all processes in communicator and returns it to one process.
- In many numerical algorithms, SEND/RECEIVE can be replaced by BCAST/REDUCE, improving both simplicity and efficiency.

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MPI Collective Communication

- Communication and computation is coordinated among a group of processes in a communicator.
- Groups and communicators can be constructed "by hand" or using topology routines.
- Tags are not used; different communicators deliver similar functionality.
- · No non-blocking collective operations.
- Three classes of operations: synchronization, data movement, collective computation.

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Synchronization

- MPI_Barrier(comm)
- Blocks until all processes in the group of the communicator comm call it.

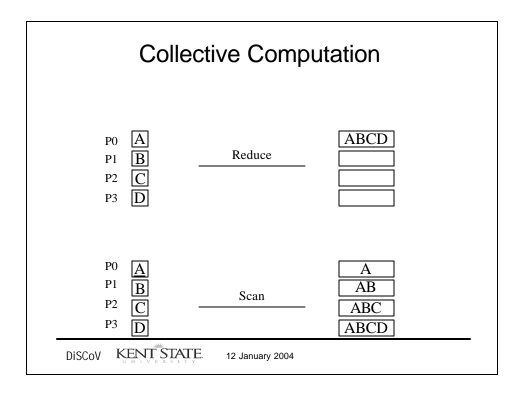
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Synchronization

- MPI_Barrier(comm, ierr)
- Blocks until all processes in the group of the communicator comm call it.

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More Collective Data Movement P0 Allgather В P1 A B C D P2 $A \mid B \mid C \mid D$ P3 P0A0B0C0D0 P1 A1B1C1D1 B0 B1 B2 B3 Alltoall P2 A2B2C2D2 C0 C1 C2 C3 P3 D0D1D2D3 A3 B3 C3 D3 KENT STATE DiSCoV 12 January 2004



MPI Collective Routines

- Many Routines: Allgather, Allgatherv, Allreduce,
 Alltoall, Alltoallv, Bcast, Gather, Gatherv, Reduce,
 Reduce_scatter, Scan, Scatter, Scatterv
- All versions deliver results to all participating processes.
- · V versions allow the hunks to have different sizes.
- Allreduce, Reduce, Reduce_scatter, and Scan take both built-in and user-defined combiner functions.

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MPI Built-in Collective Computation Operations MAXIMUM

MPI_Max
MPI_Min
MPI_Prod
MPI_Sum
MPI_Land
MPI_Lor
MPI_Lor
MPI_Lor

MPI_Lxor Logical exclusive or

MPI_Band Binary andMPI_Bor Binary or

MPI_Bxor Binary exclusive or
 MPI_Maxloc Maximum and location
 MPI_Minloc Minimum and location

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How Deterministic are Collective Computations? In exact arithmetic, you always get the same results

- - but roundoff error, truncation can happen
- MPI does *not* require that the same input give the same output
 - Implementations are encouraged but not required to provide exactly the same output given the same input
 - Round-off error may cause slight differences
- Allreduce does guarantee that the same value is received by all processes for each call
- Why didn't MPI mandate determinism?
 - Not all applications need it
 - Implementations can use "deferred synchronization" ideas to provide better performance

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Defining your own Collective **Operations**

```
Create your own collective computations with:
MPI_Op_create( user_fcn, commutes, &op );
MPI_Op_free( &op );
```

```
user_fcn( invec, inoutvec, len, datatype );
```

The user function should perform:

```
inoutvec[i] = invec[i] op
                           inoutvec[i];
```

for i from 0 to len-1.

The user function can be non-commutative.

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MPICH Goals

- Complete MPI implementation
- Portable to all platforms supporting the message-passing model
- High performance on high-performance hardware
- As a research project:
 - exploring tradeoff between portability and performance
 - removal of performance gap between user level (MPI) and hardware capabilities
- As a software project:
 - a useful free implementation for most machines
 - a starting point for vendor proprietary implementations

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MPICH Architecture

- · Most code is completely portable
- An "Abstract Device" defines the communication layer
- The abstract device can have widely varying instantiations, using:
 - sockets
 - shared memory
 - other special interfaces
 - e.g. Myrinet, Quadrics, InfiniBand, Grid protocols

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Getting MPICH for your cluster

- http://www.mcs.anl.gov/mpi/mpich
- Either MPICH-1 or
- MPICH-2

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What's in MPI-2

- Extensions to the message-passing model
 - Dynamic process management
 - One-sided operations (remote memory access)
 - Parallel I/O
 - Thread support
- · Making MPI more robust and convenient
 - C++ and Fortran 90 bindings
 - External interfaces, handlers
 - Extended collective operations
 - Language interoperability

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MPI as a Setting for Parallel I/O

- Writing is like sending and reading is like receiving
- Any parallel I/O system will need:
 - collective operations
 - user-defined datatypes to describe both memory and file layout
 - communicators to separate application -level message passing from I/O-related message passing
 - non-blocking operations
- I.e., lots of MPI-like machinery

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MPI-2 Status

- Many vendors have partial implementations, especially I/O
- MPICH2 is nearly complete, not completely tested
- Expect completion by Thanksgiving

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Some Research Areas

- MPI-2 RMA interface
 - Can we get high performance?
- Fault Tolerance and MPI
 - Are intercommunicators enough?
- MPI on 64K processors
 - Umm...how do we make this work :)?
 - Reinterpreting the MPI "process"
- MPI as system software infrastructure
 - With dynamic processes and fault tolerance, can we build services on MPI?

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High-Level Programming With MPI

- MPI was designed from the beginning to support libraries
- · Many libraries exist, both open source and commercial
- Sophisticated numerical programs can be built using libraries
 - Solve a PDE (e.g., PETSc)
 - Scalable I/O of data to a community standard file format

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Higher Level I/O Libraries

- Scientific applications work with structured data and desire more self-describing file formats
- netCDF and HDF5 are two popular "higher level" I/O libraries
 - Abstract away details of file layout
 - Provide standard, portable file formats
 - Include metadata describing contents
- For parallel machines, these should be built on top of MPI-IO

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Exercise

- · Jacobi problem in 2 dimensions with 1-D decomposition
 - Explained in class
 - Simple version fixed number of iterations
 - Fancy version test for convergence

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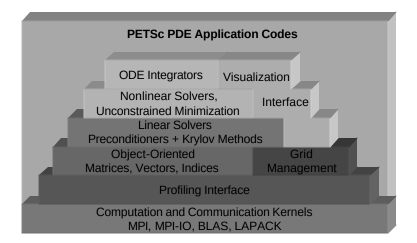
The PETSc Library

- PETSc provides routines for the parallel solution of systems of equations that arise from the discretization of PDEs
 - Linear systems
 - Nonlinear systems
 - Time evolution
- · PETSc also provides routines for
 - Sparse matrix assembly
 - Distributed arrays
 - General scatter/gather (e.g., for unstructured grids)

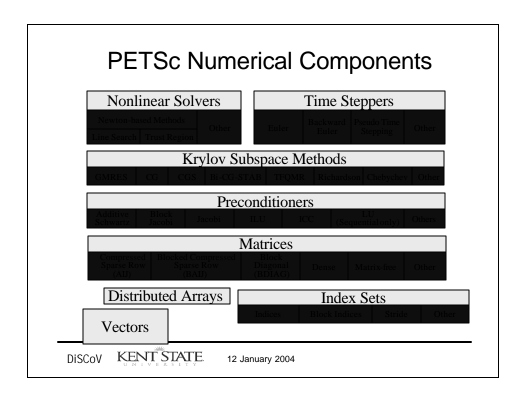
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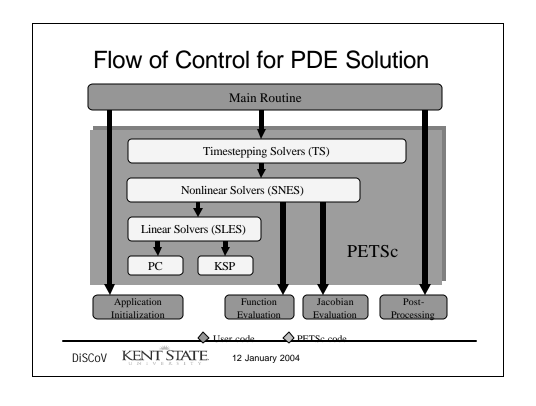
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Structure of PETSc



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Poisson Solver in PETSc

- The following 7 slides show a complete 2-d Poisson solver in PETSc. Features of this solver:
 - Fully parallel
 - 2-d decomposition of the 2-d mesh
 - Linear system described as a sparse matrix; user can select many different sparse data structures
 - Linear system solved with any user-selected Krylov iterative method and preconditioner provided by PETSc, including GMRES with ILU, BiCGstab with Additive Schwarz, etc.
 - Complete performance analysis built-in
- Only 7 slides of code!

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