

MPI

More of the Story

Timothy H. Kaiser, PH.D.

tkaiser@mines.edu



Outline

- Review
- Types
- Wildcards
- Using Status and Probing
- Asynchronous Communication, first cut
- Global communications
- Advanced topics
 - "V" operations
 - Derived types
 - Communicators

<http://geco.mines.edu/workshop>

Six basic MPI calls

MPI_INIT

Initialize MPI

MPI_COMM_RANK

Get the processor rank

MPI_COMM_SIZE

Get the number of processors

MPI_Send

Send data to another processor

MPI_Recv

Get data from another processor

MPI_FINALIZE

Finish MPI

Send and Receive Program Fortran

```
program send_receive
include "mpif.h"
integer myid,ierr,numprocs,tag,source,destination,count
integer buffer
integer status(MPI_STATUS_SIZE)
call MPI_INIT( ierr )
call MPI_COMM_RANK( MPI_COMM_WORLD, myid, ierr )
call MPI_COMM_SIZE( MPI_COMM_WORLD, numprocs, ierr )
tag=1234; source=0; destination=1; count=1
if(myid .eq. source)then
    buffer=5678
    Call MPI_Send(buffer, count, MPI_INTEGER,destination,&
        tag, MPI_COMM_WORLD, ierr)
    write(*,*)"processor ",myid," sent ",buffer
endif
if(myid .eq. destination)then
    Call MPI_Recv(buffer, count, MPI_INTEGER,source,&
        tag, MPI_COMM_WORLD, status,ierr)
    write(*,*)"processor ",myid," got ",buffer
endif
call MPI_FINALIZE(ierr)
stop
end
```

Send and Receive Program C

```
#include <stdio.h>
#include "mpi.h"
int main(int argc,char *argv[ ])
{
    int myid, numprocs, tag,source,destination,count, buffer;
    MPI_Status status;
    MPI_Init(&argc,&argv);
    MPI_Comm_size(MPI_COMM_WORLD,&numprocs);
    MPI_Comm_rank(MPI_COMM_WORLD,&myid);
    tag=1234;   source=0;   destination=1;   count=1;
    if(myid == source){
        buffer=5678;
        MPI_Send(&buffer,count,MPI_INT,destination,tag,MPI_COMM_WORLD);
        printf("processor %d sent %d\n",myid,buffer);
    }
    if(myid == destination){
        MPI_Recv(&buffer,count,MPI_INT,source,tag,MPI_COMM_WORLD,&status);
        printf("processor %d got %d\n",myid,buffer);
    }
    MPI_Finalize();
}
```

MPI Types

- MPI has many different predefined data types
- Can be used in any communication operation

Predefined types in C

C MPI Types	
MPI_CHAR	signed char
MPI_SHORT	signed short int
MPI_INT	signed int
MPI_LONG	signed long int
MPI_UNSIGNED_CHAR	unsigned char
MPI_UNSIGNED_SHORT	unsigned short int
MPI_UNSIGNED	unsigned int
MPI_UNSIGNED_LONG	unsigned long int
MPI_FLOAT	float
MPI_DOUBLE	double
MPI_LONG_DOUBLE	long double
MPI_BYTE	-
MPI_PACKED	-

Predefined types in Fortran

Fortran MPI Types

MPI_INTEGER	INTEGER
MPI_REAL	REAL
MPI_DOUBLE PRECISION	DOUBLE PRECISION
MPI_COMPLEX	COMPLEX
MPI_LOGICAL	LOGICAL
MPI_CHARACTER	CHARACTER(1)
MPI_BYTE	-
MPI_PACKED	-

Wildcards

- Allow you to not necessarily specify a tag or source
- Example

```
MPI_Status status;  
int      buffer[5];  
int      error;  
error = MPI_Recv(&buffer[0], 5, MPI_INT,  
                  MPI_ANY_SOURCE, MPI_ANY_TAG,  
                  MPI_COMM_WORLD, &status);
```

- **MPI_ANY_SOURCE** and **MPI_ANY_TAG** are wild cards
- Status structure is used to get wildcard values

Status

- The status parameter returns additional information for some MPI routines
 - Additional Error status information
 - Additional information with wildcard parameters
- C declaration : a predefined struct
 - **MPI_Status status;**
- Fortran declaration : an array is used instead
 - **INTEGER STATUS(MPI_STATUS_SIZE)**

Accessing status information

- The tag of a received message
 - C : `status.MPI_TAG`
 - Fortran : `STATUS(MPI_TAG)`
- The source of a received message
 - C : `status.MPI_SOURCE`
 - Fortran : `STATUS(MPI_SOURCE)`
- The error code of the MPI call
 - C : `status.MPI_ERROR`
 - Fortran : `STATUS(MPI_ERROR)`
- Other uses...

MPI_Probe

- MPI_Probe allows incoming messages to be checked without actually receiving .
 - The user can then decide how to receive the data.
 - Useful when different action needs to be taken depending on the "who, what, and how much" information of the message.

MPI_Probe

- C
 - **int MPI_Probe(source, tag, comm, &status)**
- Fortran
 - **MPI_PROBE(SOURCE, TAG, COMM, STATUS, IERROR)**
- Parameters
 - Source: source rank, or **MPI_ANY_SOURCE**
 - Tag: tag value, or **MPI_ANY_TAG**
 - Comm: communicator
 - Status: status object

MPI_Probe example (part I) f_ex02.f

```
! How to use probe and get_count
! to find the size of an incoming message
program probe_it
include 'mpif.h'
integer myid,numprocs
integer status(MPI_STATUS_SIZE)
integer mytag,icount,ierr,iray(10)
call MPI_INIT( ierr )
call MPI_COMM_RANK( MPI_COMM_WORLD, myid, ierr )
call MPI_COMM_SIZE( MPI_COMM_WORLD, numprocs, ierr )
mytag=123; iray=0; icount=0
if(myid .eq. 0)then
! Process 0 sends a message of size 5
icount=5
iray(1:icount)=1
call MPI_SEND(iray,icount,MPI_INTEGER,
             1,mytag,MPI_COMM_WORLD,ierr)
&
endif
```

MPI_Probe example (part 2)

```
if(myid .eq. 1)then
! process 1 uses probe and get_count to find the size
call mpi_probe(0,mytag,MPI_COMM_WORLD,status,ierr)
call mpi_get_count(status,MPI_INTEGER,icount,ierr)
write(*,*)"getting ", icount, " values"
call mpi_recv(iray,icount,MPI_INTEGER,0,
             mytag,MPI_COMM_WORLD,status,ierr)
&
endif
write(*,*)iray
call mpi_finalize(ierr)
stop
End
```

MPI_BARRIER

- Blocks the caller until all members in the communicator have called it.
- Used as a synchronization tool.
- C
 - **MPI_Barrier(comm)**
- Fortran
 - **Call MPI_BARRIER(COMM, IERROR)**
- Parameter
 - Comm communicator (**MPI_COMM_WORLD**)

Asynchronous Communication

- Asynchronous send: send call returns immediately, send actually occurs later
- Asynchronous receive: receive call returns immediately. When received data is needed, call a wait subroutine
- Asynchronous communication used in attempt to overlap communication with computation (usually doesn't work)
- Can help prevent deadlock (not advised)

Asynchronous Send with MPI_Isend

- C
 - **MPI_Request request**
 - **int MPI_Isend(&buffer, count, datatype, dest, tag, comm, &request)**
- Fortran
 - **Integer REQUEST**
 - **MPI_ISEND(BUFFER, COUNT, DATATYPE, DEST, TAG, COMM, REQUEST, IERROR)**
- Request is a new output Parameter
- Don't change data until communication is complete

Asynchronous Receive with MPI_Irecv

- C
 - `MPI_Request request;`
 - `int MPI_Irecv(&buf, count, datatype, source, tag, comm, &request)`
- Fortran
 - `Integer request`
 - `MPI_IRECV(BUF, COUNT, DATATYPE, SOURCE, TAG, COMM, REQUEST, IERROR)`
- Parameter Changes
 - Request: communication request
 - Status parameter is missing
 - Don't use data until communication is complete

MPI_Wait used to complete communication

- Request from Isend or Irecv is input
- The completion of a send operation indicates that the sender is now free to update the data in the send buffer
- The completion of a receive operation indicates that the receive buffer contains the received message
- **MPI_Wait** blocks until message specified by "request" completes

MPI_Wait used to complete communication

- C
 - `MPI_Request request;`
 - `MPI_Status status;`
 - `MPI_Wait(&request, &status)`
- Fortran
 - `Integer request`
 - `Integer status(MPI_STATUS_SIZE)`
 - `MPI_WAIT(REQUEST, STATUS, IERROR)`
- `MPI_Wait` blocks until message specified by "request" completes

MPI_Test

- Similar to MPI_Wait, but does not block
- Value of flags signifies whether a message has been delivered
- C
 - **int flag**
 - **int MPI_Test(&request, &flag, &status)**
- Fortran
 - **LOGICAL FLAG**
 - **MPI_TEST(REQUEST, FLAG, STATUS, IER)**

Non blocking send example

```
call MPI_Isend (buffer,count,datatype,dest,  
tag,comm, request, ierr)
```

10 continue

Do other work ...

```
call MPI_Test (request, flag, status, ierr)  
if (.not. flag) goto 10
```

Exercise 3 :Asynchronous Send and Receive

- Write a parallel program to send and receive data using `MPI_Isend` and `MPI_Irecv`
 - Initialize MPI
 - Have processor 0 send an integer to processor 1
 - Have processor 1 receive an integer from processor 0
 - Both processors check on message completion
 - Quit MPI

MPI Broadcast call: MPI_Bcast

- All nodes call MPI_Bcast
- One node (root) sends a message all others receive the message
- C
 - **`MPI_Bcast(&buffer, count, datatype, root, communicator);`**
- Fortran
 - **`call MPI_Bcast(buffer, count, datatype, root, communicator, ierr)`**
- Root is node that sends the message

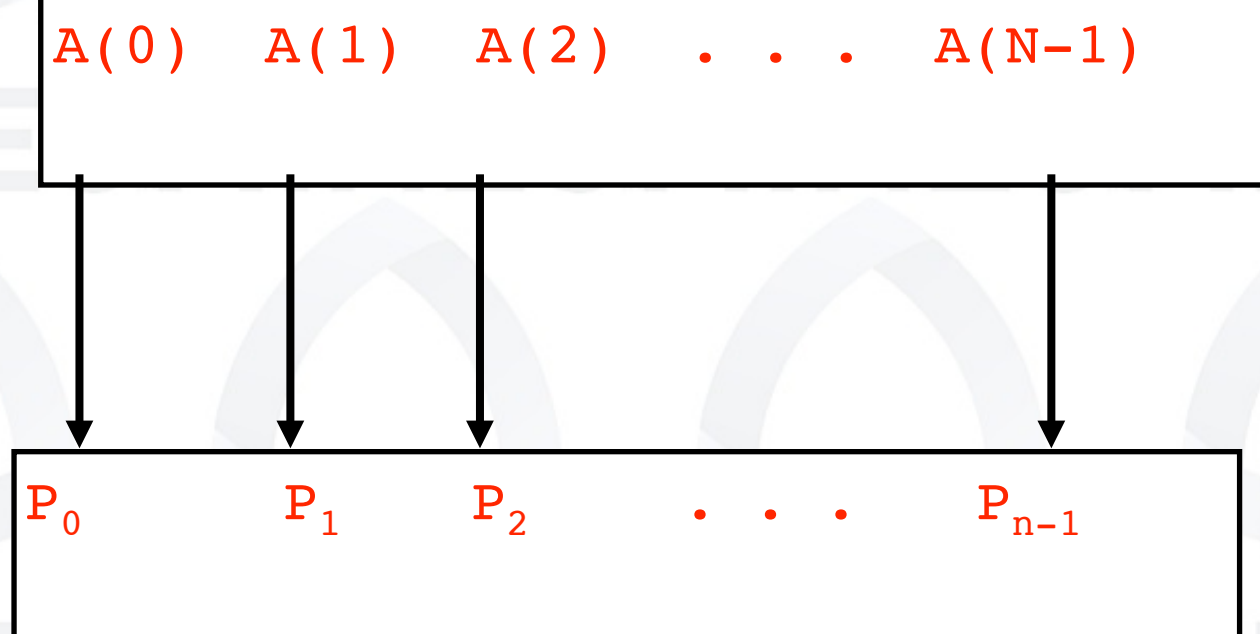
Exercise 4 : Broadcast

- Write a parallel program to broadcast data using MPI_Bcast
 - Initialize MPI
 - Have processor 0 broadcast an integer
 - Have all processors print the data
 - Quit MPI

Scatter Operation using MPI_Scatter

- Similar to Broadcast but sends a section of an array to each processors

Data in an array on root node:



Goes to processors:

MPI_Scatter

- C

- `int MPI_Scatter(&sendbuf, sendcnts, sendtype,
&recvbuf, recvcnts, recvtype, root, comm);`

- Fortran

- `MPI_Scatter(sendbuf,sendcnts,sendtype,
recvbuf,recvcnts,recvtype,root,comm,ierror)`

- Parameters

- Sendbuf is an array of size (number processors*sendcnts)
 - Sendcnts number of elements sent to each processor
 - Recvcnts number of elements obtained from the root processor
 - Recvbuf elements obtained from the root processor, may be an array

Scatter Operation using MPI_Scatter

- Scatter with Sendcnts = 2

Data in an array on root node:

A(0)	A(2)	A(4)	...	A(2N-2)
A(1)	A(3)	A(5)	...	A(2N-1)

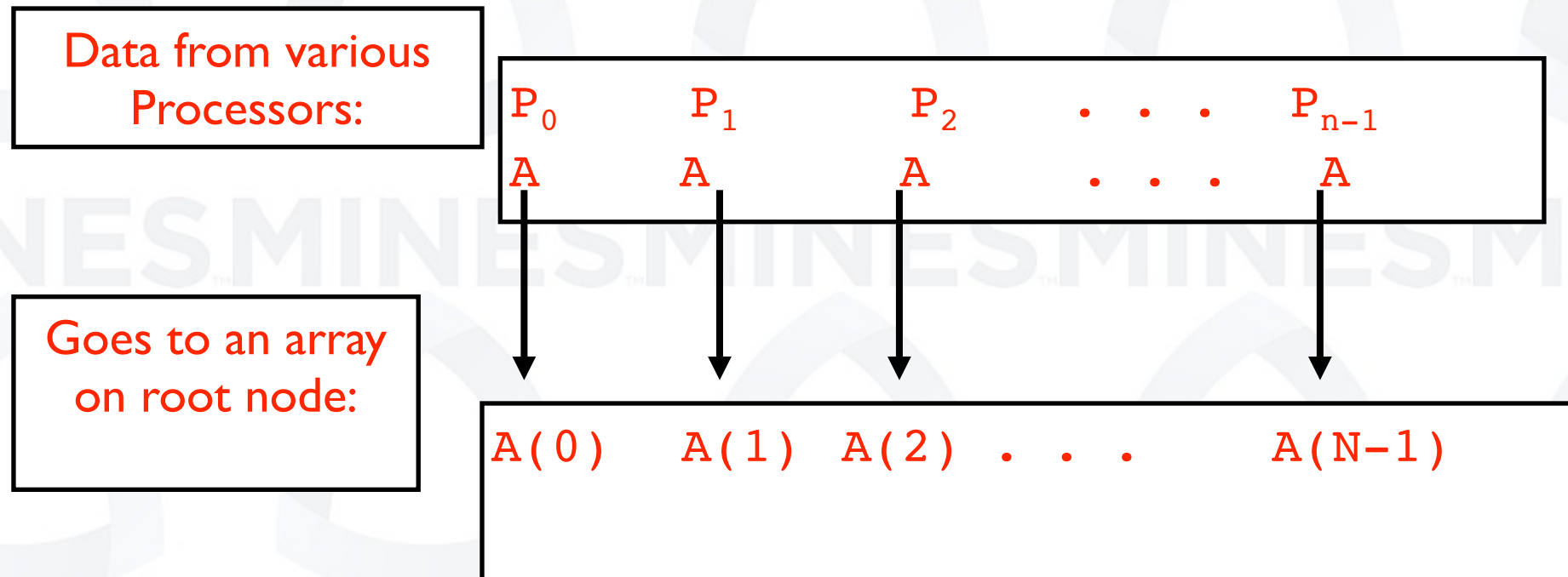


Goes to processors:

P ₀	P ₁	P ₂	...	P _{n-1}
B(0)	B(0)	B(0)		B(0)
B(1)	B(1)	B(1)		B(1)

Gather Operation using MPI_Gather

- Used to collect data from all processors to the root, inverse of scatter
- Data is collected into an array on root processor



MPI_Gather

- C

- **int MPI_Gather(&sendbuf, sendcnts, sendtype, &recvbuf, recvcnts, recvtype, root, comm);**

- Fortran

- **MPI_Gather(sendbuf, sendcnts, sendtype, recvbuf, recvcnts, recvtype, root, comm, ierror)**

- Parameters

- Sendcnts # of elements sent from each processor
- Sendbuf is an array of size sendcnts
- Recvcnts # of elements obtained from each processor
- Recvbuf of size Recvcnts*number of processors

Exercise 5 : Scatter and Gather

- Write a parallel program to scatter real data using `MPI_Scatter`
- Each processor sums the data
- Use `MPI_Gather` to get the data back to the root processor
- Root processor sums and prints the data

Reduction Operations

- Used to combine partial results from all processors
- Result returned to root processor
- Several types of operations available
- Works on single elements and arrays

MPI routine is MPI_Reduce

- C
 - `int MPI_Reduce(&sendbuf, &recvbuf,
count, datatype, operation,root,
communicator)`
- Fortran
 - `call MPI_Reduce(sendbuf, recvbuf,
count, datatype, operation,root,
communicator, ierr)`
- Parameters

Operations for MPI_Reduce

MPI_MAX	Maximum
MPI_MIN	Minimum
MPI_PROD	Product
MPI_SUM	Sum
MPI_LAND	Logical and
MPI_LOR	Logical or
MPI_LXOR	Logical exclusive or
MPI_BAND	Bitwise and
MPI_BOR	Bitwise or
MPI_BXOR	Bitwise exclusive or
MPI_MAXLOC	Maximum value and location
MPI_MINLOC	Minimum value and location

Global Sum with MPI_Reduce

C

```
double sum_partial, sum_global;
sum_partial = ...;
    ierr = MPI_Reduce(&sum_partial, &sum_global,
                      1, MPI_DOUBLE_PRECISION,
                      MPI_SUM, root,
                      MPI_COMM_WORLD);
```

Fortran

```
double precision sum_partial, sum_global
sum_partial = ...
call MPI_Reduce(sum_partial, sum_global,
                1, MPI_DOUBLE_PRECISION,
                MPI_SUM, root,
                MPI_COMM_WORLD, ierr)
```

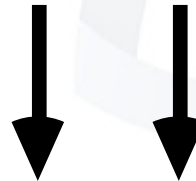
Exercise 6 : Global Sum with MPI_Reduce

- Write a program to sum data from all processors

Global Sum with MPI_Reduce

2d array spread across processors

	X(0)	X(1)	X(2)
NODE 0	A0	B0	C0
NODE 1	A1	B1	C1
NODE 2	A2	B2	C2



	X(0)	X(1)	X(2)
NODE 0	A0+A1+A2	B0+B1+B2	C0+C1+C2
NODE 1			
NODE 2			

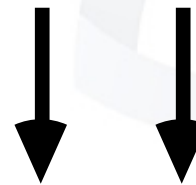
All Gather and All Reduce

- Gather and Reduce come in an "ALL" variation
- Results are returned to all processors
- The root parameter is missing from the call
- Similar to a gather or reduce followed by a broadcast

Global Sum with MPI_AllReduce

2d array spread across processors

	X(0)	X(1)	X(2)
NODE 0	A0	B0	C0
NODE 1	A1	B1	C1
NODE 2	A2	B2	C2



	Y(0)	Y(1)	Y(2)
NODE 0	A0+A1+A2	B0+B1+B2	C0+C1+C2
NODE 1	A0+A1+A2	B0+B1+B2	C0+C1+C2
NODE 2	A0+A1+A2	B0+B1+B2	C0+C1+C2

All to All communication with MPI_Alltoall

- Each processor sends and receives data to/from all others
- C
 - `int MPI_Alltoall(&sendbuf, sendcnts, sendtype,
&recvbuf, recvcnts, recvtype, root, MPI_Comm);`
- Fortran
 - `call MPI_Alltoall(sendbuf, sendcnts, sendtype,
recvbuf, recvcnts, recvtype, root, comm, ierror)`

All to All with MPI_Alltoall

- Parameters
 - Sendcnts # of elements sent to each processor
 - Sendbuf is an array of size sendcnts
 - Recvcnts # of elements obtained from each processor
 - Recvbuf of size Recvcnts*number of processors
- Note that both send buffer and receive buffer must be an array of size of the number of processors

Things Left

- “V” operations
- Communicators
- Derived typed
- Parallel IO

The dreaded “V” or variable or operators

- A collection of very powerful but difficult to setup global communication routines
- `MPI_Gatherv`: Gather different amounts of data from each processor to the root processor
- `MPI_Alltoallv`: Send and receive different amounts of data form all processors
- `MPI_Allgatherv`: Gather different amounts of data from each processor and send all data to each
- `MPI_Scatterv`: Send different amounts of data to each processor from the root processor
- We discuss `MPI_Gatherv` and `MPI_Alltoallv`

MPI_Gatherv

- C
 - `int MPI_Gatherv (&sendbuf, sendcnts, sendtype, &recvbuf, &recvcnts, &rdispls, recvtype, root, comm);`
- Fortran
 - `MPI_Gatherv (sendbuf, sendcnts, sendtype, recvbuf, recvcnts, rdispls, recvtype, root, comm, ierror)`
- Parameters:
 - **Recvnts** is now an array
 - **Rdispls** is a displacement

MPI_Gatherv

- Recvcnts
 - An array of extent Recvcnts(0:N-1) where Recvcnts(N) is the number of elements to be received from processor N
- Rdispls
 - An array of extent Rdispls(0:N-1) where Rdispls(N) is the offset, in elements, from the beginning address of the receive buffer to place the data from processor N
- Typical usage

```
recvcnts=...
rdispls(0)=0
do I=1,n-1
    rdispls(I) = rdispls(I-1) + recvcnts(I-1)
enddo
```

MPI_Gatherv Example

- This program shows how to use **MPI_Gatherv**. Each processor sends a different amount of data to the root processor.
- We use **MPI_Gather** first to tell the root how much data is going to be sent.

MPI_Alltoallv

- Send and receive different amounts of data from all processors
- C
 - `int MPI_Alltoallv (&sendbuf, &sendcnts,
&sdispls, sendtype, &recvbuf, &recvcnts,
&rdispls, recvtype, comm);`
- Fortran
 - `Call MPI_Alltoallv(sendbuf, sendcnts,
sdispls, sendtype, recvbuf, recvcnts,
rdispls, recvtype, comm, ierror);`

MPI_Alltoallv

- We add **sdispls** parameter
 - An array of extent **sdispls**(0:N-1) where **sdispls**(N) is the offset, in elements, from the beginning address of the send buffer to get the data for processor N
- Typical usage

```
recvnts=...
Sendcnts=...
rdispls(0)=0
Sdispls(0)=0
do I=1,n-1
    rdispls(I) = rdispls(I-1) + recvnts(I-1)
    sdispls(I) = sdispls(I-1) + sendcnts(I-1)
Enddo
```

MPI_Alltoallv example

- Each processor send/rec a different and random amount of data to/from otherprocessors.
- We use MPI_Alltoall first to tell how much data is going to be sent.

Derived types

- C and Fortran 90 have the ability to define arbitrary data types that encapsulate reals, integers, and characters.
- MPI allows you to define message data types corresponding to your data types
- Can use these data types just as default types

Derived types, Three main classifications:

- Contiguous Vectors: enable you to send contiguous blocks of the same type of data lumped together
- Noncontiguous Vectors: enable you to send noncontiguous blocks of the same type of data lumped together
- Abstract types: enable you to (carefully) send C or Fortran 90 structures, don't send pointers

Derived types, how to use them

- Three step process
 - Define the type using
 - `MPI_TYPE_CONTIGUOUS` for contiguous vectors
 - `MPI_TYPE_VECTOR` for noncontiguous vectors
 - `MPI_TYPE_STRUCT` for structures
 - Commit the type using
 - `MPI_TYPE_COMMIT`
 - Use in normal communication calls
 - `MPI_Send(buffer, count, MY_TYPE, destination, tag, MPI_COMM_WORLD, ierr)`

MPI_TYPE_CONTIGUOUS

- Defines a new data type of length count elements from your old data type
- C
 - `MPI_TYPE_CONTIGUOUS(int count, old_type,
&new_type)`
- Fortran
 - `Call MPI_TYPE_CONTIGUOUS(count, old_type,
new_type, ierror)`
- Parameters
 - `old_type`: your base type
 - `New_type`: a type count elements of `Old_type`

MPI_TYPE_VECTOR

- Defines a datatype which consists of `count` blocks each of length `blocklength` and `stride` displacement between blocks
- C
 - `MPI_TYPE_VECTOR(count, blocklength, stride, old_type, *new_type)`
- Fortran
 - Call `MPI_TYPE_VECTOR(count, blocklength, stride, old_type, new_type, ierror)`
- We will see examples later

MPI_TYPE_STRUCT

- Defines a MPI datatype which maps to a user defined derived datatype
- C
 - `int MPI_TYPE_STRUCT(count,
&array_of_blocklengths,
&array_of_displacement, &array_of_types,
&newtype);`
- Fortran
 - `Call MPI_TYPE_STRUCT(count,
array_of_blocklengths,
array_of_displacement, array_of_types,
newtype,ierror)`

MPI_TYPE_STRUCT

- Parameters:
 - [IN count] # of old types in the new type (integer)
 - [IN array_of_blocklengths] how many of each type in new structure (integer)
 - [IN array_of_types] types in new structure (integer)
 - [IN array_of_displacement] offset in bytes for the beginning of each group of types (integer)
 - [OUT newtype] new datatype (handle)
- `Call MPI_TYPE_STRUCT(count, array_of_blocklengths, array_of_displacement, array_of_types, newtype, ierror)`

Derived Data type Example

Consider the data type or structure consisting of

- 3 MPI_DOUBLE_PRECISION
- 10 MPI_INTEGER
- 2 MPI_LOGICAL

Creating the MPI data structure matching this C/Fortran structure is a three step process

Fill the descriptor arrays:

- B - blocklengths

- T - types

- D - displacements

Call `MPI_TYPE_STRUCT` to create the MPI data structure

Commit the new data type using `MPI_TYPE_COMMIT`

Derived Data type Example

- Consider the data type or structure consisting of
 - 3 MPI_DOUBLE_PRECISION
 - 10 MPI_INTEGER
 - 2 MPI_LOGICAL
- To create the MPI data structure matching this C/Fortran structure
 - Fill the descriptor arrays:
 - B - blocklengths
 - T - types
 - D - displacements
 - Call MPI_TYPE_STRUCT

Derived Data type Example (continued)

! t contains the types that
! make up the structure

t(1)=MPI_DOUBLE_PRECISION

t(2)=MPI_INTEGER

t(3)=MPI_LOGICAL

! b contains the number of each type

b(1)=3;b(2)=10;b(3)=2

! d contains the byte offset of
! the start of each type

d(1)=0;d(2)=24;d(3)=64

call MPI_TYPE_STRUCT(3,b,d,t,
MPI_CHARLES,mpi_err)

MPI_CHARLES is our new data type

MPI_Type_commit

- Before we use the new data type we call **MPI_Type_commit**
- C
 - **MPI_Type_commit(MPI_CHARLES)**
- Fortran
 - Call **MPI_Type_commit(MPI_CHARLES,ierr)**

Communicators

- A communicator is a parameter in all MPI message passing routines
- A communicator is a collection of processors that can engage in communication
- `MPI_COMM_WORLD` is the default communicator that consists of all processors
- MPI allows you to create subsets of communicators

Why Communicators?

- Isolate communication to a small number of processors
- Useful for creating libraries
- Different processors can work on different parts of the problem
- Useful for communicating with "nearest neighbors"

MPI_Comm_create

- **MPI_Comm_create** creates a new communicator `newcomm` with group members defined by a group data structure.
- C
 - `int MPI_Comm_create(old_comm, group, &newcomm)`
- Fortran
 - `Call MPI_COMM_CREATE(COMM, GROUP, NEWCOMM, IERROR)`
- How do you define a group?

MPI_Comm_group

- Given a communicator, **MPI_Comm_group** returns in group associated with the input communicator
- C
 - **int MPI_Comm_group(comm, &group)**
- Fortran
 - **Call MPI_COMM_GROUP(COMM, GROUP, IERROR)**
- MPI provides several functions to manipulate existing groups.

MPI_Group_incl

- MPI_Group_incl creates a group **new_group** that consists of the n processes in **old_group** with ranks rank[0],..., rank[n-1]
- C
 - `int MPI_Group_incl(group,
n,&ranks,&new_group)`
- Fortran
 - `Call MPI_GROUP_INCL(GROUP, N, RANKS,
NEW_GROUP, IERROR)`

MPI_Group_incl

- Fortran
 - **Call MPI_GROUP_INCL(old_GROUP, N, RANKS, NEW_GROUP, IERROR)**
- Parameters
 - old_group: your old group
 - N: number of elements in array ranks (and size of new_group) (integer)
 - Ranks: ranks of processes in group to appear in new_group (array of integers)
 - New_group:new group derived from above, in the order defined by ranks

MPI_Group_excl

- MPI_Group_excl creates a group of processes **new_group** that is obtained by deleting from **old_group** those processes with ranks `ranks[0], ..., ranks[n-1]`
- C
 - `int MPI_Group_excl(old_group, n, &ranks,
MPI_Group &new_group)`
- Fortran
 - `Call MPI_GROUP_EXCL(OLD_GROUP, N, RANKS,
NEW_GROUP, IERROR)`

MPI_Comm_split

- Provides a short cut method to create a collection of communicators
- All processors with the "same color" will be in the same communicator
- Index gives rank in new communicator
- Fortran
 - `call MPI_COMM_SPLIT(OLD_COMM, color, index,
NEW_COMM, mpi_err)`
- C
 - `MPI_Comm_split(OLD_COMM, color, index, &NEW_COMM)`

MPI_Comm_split

- Split odd and even processors into 2 communicators

```
Program comm_split
include "mpif.h"
Integer color,zero_one
call MPI_INIT( mpi_err )
call MPI_COMM_SIZE( MPI_COMM_WORLD, numnodes, mpi_err )
call MPI_COMM_RANK( MPI_COMM_WORLD, myid, mpi_err )
color=mod(myid,2) !color is either 1 or 0
call MPI_COMM_SPLIT(MPI_COMM_WORLD,color,myid,NEW_COMM,mpi_err)
call MPI_COMM_RANK( NEW_COMM, new_id, mpi_err )
call MPI_COMM_SIZE( NEW_COMM, new_nodes, mpi_err )
Zero_one = -1
If(new_id==0)Zero_one = color
Call MPI_Bcast(Zero_one,1,MPI_INTEGER,0, NEW_COMM,mpi_err)
If(zero_one==0)write(*,*)"part of even processor communicator"
If(zero_one==1)write(*,*)"part of odd processor communicator"
Write(*,*)"old_id=", myid, "new_id=", new_id
Call MPI_FINALIZE(mpi_error)
End program
```

MPI_Comm_split example output

- Note, I have sorted the output

```
[mbpro:~] tkaiser% mpiexec -np 8 split.exe | sort
old_id= 0 new_id= 0
old_id= 1 new_id= 0
old_id= 2 new_id= 1
old_id= 3 new_id= 1
old_id= 4 new_id= 2
old_id= 5 new_id= 2
old_id= 6 new_id= 3
old_id= 7 new_id= 3
part of even processor communicator
part of odd processor communicator
[mbpro:~] tkaiser%
```

MPI_Comm_split output with task labels

- Split odd and even processors into 2 communicators

0: part of even processor communicator

0: old_id= 0 new_id= 0

2: part of even processor communicator

2: old_id= 2 new_id= 1

1: part of odd processor communicator

1: old_id= 1 new_id= 0

3: part of odd processor communicator

3: old_id= 3 new_id= 1

Group and Communicator example

This program is designed to show how to set up a new communicator. We set up a communicator that includes all but one of the processors, The last processor is not part of the new communicator, `TIMS_COMM_WORLD`.

We use the routine `MPI_Group_rank` to find the rank within the new communicator. For the last processor the rank is `MPI_UNDEFINED` because it is not part of the communicator. For this processor we call `get_input`. The processors in `TIMS_COMM_WORLD` pass a token between themselves in the subroutine `pass_token`. The remaining processor gets input, `i`, from the terminal and passes it to processor 1 of `MPI_COMM_WORLD`. If `i > 100` the program stops.