

Advanced MPI

Introduction to Parallel Computing with MPI and OpenMP

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Pack/Unpack



Example: 0501MPlexample-pack

Consider the problem of sending data of different kinds. For example, root should broadcast the following values:

MPI_DOUBLE_PRECISION: SWV(2), Range

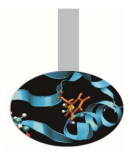
MPI_INTEGER: XYdots, Niter

the broadcasting function should be called 4 times:

```
CALL MPI_BCAST (SWV, 2, MPI_DOUBLE_PRECISION , 0,
    MPI_COMM_WORLD, ierr)
CALL MPI_BCAST (XYdots, 1, MPI_INTEGER, 0, MPI_COMM_WORLD, ierr)
CALL MPI_BCAST (Range, 1, MPI_DOUBLE_PRECISION, 0,
    MPI_COMM_WORLD, ierr)
CALL MPI BCAST (NITER, 1, MPI INTEGER, 0, MPI COMM WORLD, ierr)
```



Pack/Unpack



A better solution is possible, minimizing the use of sending/receiving functions. The four objects:

MPI_DOUBLE_PRECISION: SWV(2), Range ! 8*3 = 24 bytes

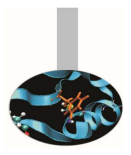
MPI_INTEGER: XYdots, Niter ! 2*4 = 8 bytes

can be packed into a buffer for delivery and unpacked on receiving:

1. root process packs data into the buffer pckd_data, 32 bytes long:



Pack/Unpack



2. Buffered data are distributed: CALL MPI_BCAST (pckd_data, 32, MPI_PACKED, 0, & & MPI_COMM_WORLD, ierr)

3. receiving processes unpack data:

Pos = 0

• •

CALL MPI_UNPACK (pckd_data, 32, pos, SWV, 2, &

& MPI_DOUBLE_PRECISION, MPI_COMM_WORLD, ierr)

CALL MPI_UNPACK (pckd_data, 32, pos, Niter, 1, &

& MPI_INTEGER, MPI_COMM_WORLD, ierr)

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The MPI library enables packing different data in one buffer, that can be sent as a whole. Communication times can thus be reduced. To gather several data in a single buffer the function MPI PACK may be used.

INTERFACE
SUBROUTINE MPI_PACK(inbuf,incount,datatype,outbuf,outsize,position,comm,ierr)
INTEGER, INTENT(IN) :: INCOUNT, DATATYPE, OUTSIZE, COMM
<type>, INTENT(IN) :: INBUF(:)</type>
<type>, INTENT(OUT) :: OUTBUF(:)</type>
INTEGER, INTENT(INOUT) :: POSITION
INTEGER, INTENT(OUT) :: IERR
END SUBROUTINE MPI_PACK
END INTERFACE

INCOUNT elements of type DATATYPE of the buffer INBUF are copied in the buffer OUTBUF from position POSITION (in byte). On exit POSITION has the value of the next free address.



Unpack



The function MPI_UNPACK is used by the receiving processes to extract data from the buffer INBUF.

INTERFACE	
SUBROUTINE MPI_UNPACK (inbuf, insize, position, outbuf, outcount, datat	:ype,
comm, ierr)	
INTEGER, INTENT(IN) :: INSIZE, DATATYPE, OUTCOUNT, COMM	
<type>, INTENT(IN) :: INBUF(:)</type>	
<type>, INTENT(OUT) :: OUTBUF(:)</type>	
INTEGER, INTENT(INOUT) :: POSITION	
INTEGER, INTENT(OUT) :: IERR	
END SUBROUTINE MPI_UNPACK	
END INTERFACE	fortran

int MPI_Unpack(void *inbuf, int insize, int *position, void *outbuf,	
int outcount, MPI_Datatype datatype, MPI_Comm comm)	с/с++







Basic data types

The basic MPI data types are:

If the data to be communicated are structured with etherogeneous members, it may be convenient to define a MPI derived data type. Derived data types are defined using basic data types (and formerly defined derived data types).

 $Dtype = [(typ_0, pos_0), (typ_1, pos_1), ..., (typ_n-1, pos_n-1)]$





Derived data types

To define a derived data type it is required:

- To specify the structure of the new data type, on the basis of previously defined or basic data types.
- To register the new data type

The newly defined data type must be registered to MPI with the following function:

interface	fortran
subroutine mpi_type_commit (mpi_mytype, cod_err)	_
integer, intent (in) :: mpi_mytype ! Il nome del nuovo tipo di dat	ti
integer, intent (out):: cod_err ! codice di errore.	
end subroutine mpi_type_commit	
end interface	

int MPI_Type_commit (MPI_Datatype *mpi_mytype)

с/с++

Once committed the new data type becomes a recognized MPI data type.







Derived data types

Whenever a defined data type is of no use any more, the following function should be called:

interface	fortran
<pre>subroutine mpi_type_free (mpi_mytype, cod_err) integer, intent (in) :: mpi_mytype ! Data type handler integer, intent (out):: cod err ! Error code</pre>	jortran
end subroutine mpi_type_commit	
end interface	

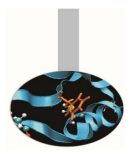
int MPI_Type_free (MPI_Datatype *mpi_mytype)

с/с++

Pending operations will complete normally.





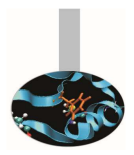


Example: 0502MPlexample-struct

Consider the problem of sending heterogeneous data contained in a structure:

TYPE input_data	struct {
REAL(8) :: SWV(2), Range	<pre>double SWV[2], Range;</pre>
INTEGER :: XYdots, Niter	<pre>int Xydots, Niter;</pre>
END TYPE	<pre>} input data;</pre>





A better solution is possible, minimising the communication calls.

A MPI derived data type is built from blocks of homogeneous elements. For example, if we are interested in communicating the following data structure:

TYPE input_data	struct {
REAL(8) :: SWV(2), Range	<pre>double SWV[2], Range;</pre>
INTEGER :: XYdots, Niter	<pre>int Xydots, Niter;</pre>
END TYPE	<pre>} input data;</pre>

four blocks may be defined, one for each component of the structure:

Block #1: 2 doublesBlock #2: 1 doubleBlock #3: 1 integerBlock #4: 1 integer





The MPI representation of a generic data structure requires three arrays to be defined:

- v_len_blk actual length of each block (in elements)
- v_head starting postion of each block (in bytes)
- v_el_typ data type of the elements in each block

It should be noted that the values in v_head must be given in bytes because the elements of each block may be of different type with different byte extensions.





The data structure can thus be described using the MPI_Type_struct function:

Generic structures

```
num_blk = 4
v_len_blk = [2,1,1,1]
v_head = [0,16,24,28]
v_el_typ = [MPI_DOUBLE_PRECISION, &
    & MPI_DOUBLE_PRECISION, &
    & MPI_INTEGER, MPI_INTEGER]
CALL MPI_Type_struct(num_blk, v_len_blk, v_head, &
        & v_el_typ, new_type, ierr)
CALL MPI_Type_commit(new_type,ierr)
```







The function mpi type struct has the following interface:

```
interface
subroutine mpi_type_struct(num_blk,v_len_blk,v_head,v_el_typ,new_typ,ierr)
integer, intent(in) :: num_blk ! How many blocks
integer, intent(in), dimension(:) :: v_len_blk ! How many elements per block
integer, intent(in), dimension(:) :: v_head ! How many bytes before
! each block
integer, intent(in), dimension(:) :: v_el_typ ! Element type per block
integer, intent(out) :: new_typ ! Data type handler
integer, intent(out) :: ierr ! Error code
end subroutine mpi_type_struct
end interface
```





C language *structures* and Fortran *derived data types* may be easily mapped into *MPI derived data types*.

The programmer must anyway be sure that the relative positions of the structure components are not modified by compiler optimizations: Fortran derived types should contain the instruction SEQUENCE.





Contiguous elements

Example: 0503MPlexample-struct_gather

Consider the problem of distributing a vector of structured data:

TYPE person

SEQUENCE

CHARACTER(80) :: Name, Surname

INTEGER, DIMENSION(3) :: Birth_date

INTEGER :: Position, Id

END TYPE person

```
TYPE(person), dimension(8) :: lteam, team
```

If we would like to distribute the *team* array, we could send it as a numer of elements of type person (or whatever is called the object MPI_Datatype) or as a global object composed of elements of type person.







Contiguous elements

An array of contiguous and homogeneous elements is the simplest derived type to be defined. From element to element there must be no spaces.

```
interface
   subroutine mpi_type_contiguous (num_el, el_type, new_type, ierr)
        integer, intent(in) :: num_el ! How many elements in the array
        integer, intent(in) :: el_type ! Element type
        integer, intent(out) :: new_type ! New data type handler
        integer, intent(out) :: ierr ! Error code
   end subroutine mpi_type_contiguous
   fortran
end interface
```

с/с++

This function defines the new data type starting from an array of num_el elements. All the elements must be of the same (derived) data type.







Contiguous elements

As an other simple example, if

 $El_type = \{ (double, 0), (char, 8) \}$

is a 16 bytes data type and 3 elements of that type are filed in an array, then

New_type =	{(double,	0),	(char,	8)
_	(double,	16),	(char,	24)
	(double,	32),	(char,	40)}

Of course there is no point in *El_type* being a basic MPI type even if it may be as well.







Not contiguous elements

Making things a bit more complicated, the following function is used to define arrays with useful data separated by fixed strides. i.e. arrays may be seen as sequences of identical blocks containing elements to be communicated and elements to be discarded

interface

subroutine mpi_type_vector(num_blk,1	<pre>en_blk,blk_siz,el_typ,new_typ,errcode)</pre>
<pre>integer, intent(in) :: num_blk</pre>	! How many blocks
<pre>integer, intent(in) :: len_blk</pre>	! How many useful elements per block
<pre>integer, intent(in) :: blk_siz</pre>	! Total number of elements per block
<pre>integer, intent(in) :: el_typ</pre>	! Data type of the block elements
<pre>integer, intent(out) :: new_typ</pre>	! New data type handler
integer, intent(out) :: ierr	! Error code
end subroutine mpi_type_vector	
end interface	fortran









Not contiguous elements

It can be noted that the size and the useful length of the blocks is given in number of elements.

As an example if blk_siz=10 and len_blk=7 and the elements are of type MPI_INTEGER, the actual size of each block is 4 x 10 = 40 bytes. But only 4 x 7 = 28 elements are communicated and 4 x (10-7) = 12 bytes are never sent.





More on not contiguous elements

The following function must be used to define arrays with blocks of different dimensions. Two vectors are needed to define the lengths because each block has its own number of useful and discarded elements.

int MPI_Type_indexed(<pre>int num_blk, int v_len_blk[], int v_head[],</pre>	
	MPI_Datatype el_typ, MPI_Datatype *new_typ)	c/c++

Please note that instead of specifying the total length of each block, the starting position of the blocks have to be passed to the function.





More on not contiguous elements

As an example, if we have to describe data structured in three blocks, 3 elements parted each other and containing 5, 13 and 7 elements, the arrays v_len_blk and v_head must be defined as follow:

v_len_blk = (/ 5, 13, 7 /)
v_head = (/ 0, 8, 24 /)

The following function may be used to know the extension of a MPI (either basic or derived) data type:

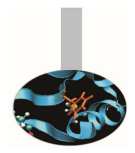
interface	fortran
<pre>subroutine mpi_type_extent (datatype, ext, cod_err)</pre>	
integer, intent(in) :: datatype ! MPI data typ	e
integer, intent(out) :: ext ! Extension (i	n bytes)
integer, intent(out) :: ierr ! Error code	
end subroutine mpi_type_extent	
end interface	

int MPI_Type_extent(MPI_Datatype datatype, MPI_Aint *ext)





Useful functions



The function mpi_type_hvector is similar to mpi_type_vector, but
blk_siz is given in bytes. The function mpi_type_hindexed is alike
mpi_type_indexed, but v_head is measured in bytes.

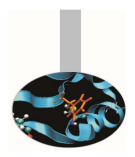
The function mpi_address returns the starting address of an object. It is important for portability issues.

interface	fortran
subroutine mpi_address (obj, address, ierr)	joreren
<pre>integer, intent(in) :: obj</pre>	
integer, intent(out) :: address ! Starting address	
integer, intent(out) :: ierr ! Error code	
end subroutine mpi_address	
end interface	

int MPI Address(void *obj, MPI Aint *address)

с/с++





Example: 0504MPlexample-struct_file

The MPI derived data types may be used not only in communications but in I/O operations also.

In the former example, imagine to substitute data distribution function with collective file writing:

CALL MPI_File_open(MPI_COMM_WORLD, 'team.dat', &

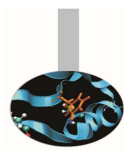
- & MPI_MODE_WRONLY+MPI_MODE_CREATE, &
- & MPI_INFO_NULL, fh, ierr)

CALL MPI_File_write_ordered(fh, lteam, 1, &

& pair_type, status, ierr)

CALL MPI_File_close(fh,ierr)

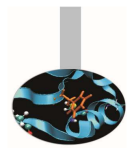




After writing, data is kept on disk and can be recovered when needed:







Basic MPI-IO operations are: open, seek, read, write, close

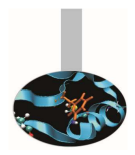
- open/close operations must be issued by all processes on the same file (collective operations)
- MPI read/write functions are similar to send/recv
- a local pointer to the file (individual file pointer) is kept per each process for seek, read, write operations

с/с++

fortran







Before beginning I/O operations the destination file must be connected to the MPI system. This is afforded by the MPI_File_open function. Remember that this function is collective: it must be called by all the processes in a communicator.

interface subroutine MPI_FILE_OPEN(comm, filename, amode, info, fh, ierr) integer, intent(in) :: comm ! Communicator character(*), intent(in) :: filename fortran integer, intent(in) :: amode ! Access mode integer, intent(in) :: info ! Access details integer, intent(out) :: fh ! File handle intent(out) :: ierr ! Error code end subroutine MPI_FILE_OPEN end interface







Few notes about the open function:

- the function is collective; would it be the case just one process has to call it, MPI_COMM_SELF should be used
- the filename must be the same for all involved processes
- if the MPI_Info handler is not used, MPI_INFO_NULL value can be passed







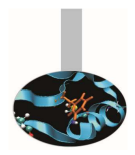
• the access mode value must be the same for all involved processes; some of the most common mode values are:

MPI_MODE_RDONLY	read only
MPI_MODE_RDWR	read/write
MPI_MODE_WRONLY	write only
MPI_MODE_CREATE	create if not existing
MPI_MODE_DELETE_ON_CLOSE	delete on closing

Mode values may be chained with + (plus, Fortran) or | (pipe, C, C++) characters







After the destination file have been used, the MPI_File_close function should be called. Remember that this function also is collective: it must be called by all the processes in the communicator.

interface			
subroutine MPI_FILE_CLOSE(fh, ierr	()		
integer, intent(in) :: fh ! File handle			
intent(out) :: ierr	! Error code	fortran	
end subroutine MPI_FILE_CLOSE		jereren	
end interface			

int MPI_File_close(MPI_File* fh)

с/с++







There are several functions for storing and recovering data to/from disk.

- Some functions are blocking, while others can overlap I/O with computation.
- Besides operations can be collective or individually managed.
- Positioning can be collectively or individually or explicitly managed.
- In the previous example a writing collective function has been used to store data. In this case the operations is automatically managed by MPI and processes do not need to take care about data positioning in the file.
- Instead data retrieving has been accomplished by calling a non collective function with individual pointer.



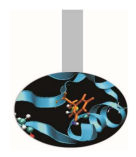




The MPI_File_write_ordered function can be used to store distributed data in the process order.

interface	
subroutine MPI_File_write_ordered(fh, buf, count, &	fortran
& datatype, status, ierr)	
<pre>integer, intent(in) :: fh</pre>	
<pre>integer, intent(in) :: buf ! Data (actual type may vary</pre>	
<pre>integer, intent(in) :: count</pre>	
integer, intent(in) :: datatype ! MPI datatype of data	
integer, intent(out) :: status ! Infos	
intent(out) :: ierr ! Error code	
end subroutine MPI_File_write_ordered	
end interface	Í



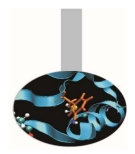


The MPI_File_read function can be used to retrieve data individually, i.e. each process read data indipendently by all others.

interface		
<pre>subroutine MPI_File_read(fh, buf, count,</pre>	datatype, &	fortran
& status, ierr)		
<pre>integer, intent(in) :: fh !</pre>	File handle	
<pre>integer, intent(out) :: buf !</pre>	Data (actual type may vary)	
<pre>integer, intent(in) :: count</pre>	Elements in buffer	
<pre>integer, intent(in) :: datatype !</pre>	MPI datatype of data	
<pre>integer, intent(out) :: status !</pre>	Infos	
intent(out) :: ierr !	Error code	
end subroutine MPI_File_read		
end interface		

int MPI_File_read(MPI_File fh, void *buf, int count, C/C++ MPI_Datatype datatype, MPI_Status *status)





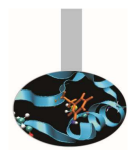
The status object returned by I/O functions may be used to control data movements. In the example the MPI_Get_count has been used to show how many elements have been written in each writing operation.

interface			
<pre>subroutine MPI_Get_count(status, data</pre>	type, count, ierr)	fortran	
integer, intent(in) :: status	! I/O infos		
integer, intent(in) :: datatype	! MPI datatype of data		
intent(out) :: count	! Elements moved		
intent(out) :: ierr	! Error code		
end subroutine MPI_Get_count			
end interface			

int MPI_Get_count(MPI_Status *status, MPI_Datatype datatype, int *count) c/c++







There are many other MPI functions for saving and retrieving data on disk; here is just a list of some.

Blocking, individual file pointer:

- int MPI_File_write (MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)
- int MPI_File_read(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)

and the collective versions:

- int MPI_File_write_all(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)
- int MPI_File_read_all(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)





MPI-IO basics



Blocking, shared file pointer:

- int MPI_File_write_shared(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)
- int MPI_File_read_shared(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)

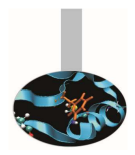
and the collective versions:

- int MPI_File_write_ordered(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)
- int MPI_File_read_ordered(MPI_File fh, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)





MPI-IO basics



Blocking, explicit file offset:

- int MPI_File_write_at(MPI_File fh, MPI_Offset offset, ROMIO_CONST void *buf, int count, MPI_Datatype datatype, MPI_Status *status)
- int MPI_File_read_at(MPI_File fh, MPI_Offset offset, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)

and the collective versions:

- int MPI_File_write_at_all(MPI_File fh, MPI_Offset offset, ROMIO_CONST void *buf, int count, MPI_Datatype datatype, MPI_Status *status)
- int MPI_File_read_at_all(MPI_File fh, MPI_Offset offset, void *buf, int count, MPI_Datatype datatype, MPI_Status *status)

Besides the cited functions there are the non-blocking versions.

Furthermore please note that MPI I/O is unformatted: data are saved on disk as they are stored in memory.





In MPI terminology the process is the computing unit. MPI processes behave following the MIMD model. Each process is an indipendent unit and has its own memory space; it should be thought of as running on its own computing machine.





Every MPI process belongs to one or more MPI group and has its own identification number or rank. MPI ranks are always numbered starting from 0. The 0 process is often called the *master* and usually acts as the boss in master-slave programming model, but it is not mandatory: any other process can act the same.

MPI groups may be generated and destroyed but they are otherwise static.





Example: 0505MPIExample_two_groups

In this example the group related to the default communicator is splitted in two groups using an array of indices:

```
numproc0 = numproc/2;
```

for (i = 0; i < numproc0; i++) ranks0[i] = i;</pre>

MPI_Group_incl(GlobalGroup, numproc0, ranks0, &Group0); MPI_Group_excl(GlobalGroup, numproc0, ranks0, &Group1); MPI Group size(Group1, &numproc1);







Each group has its own handle but it is an opaque object: the programmer can not access its details.

Proper functions must be used to manage group properties:

call mpi_group_size(group, size, ierr)
call mpi_group_rank(group, rank, ierr)

At the beginning all processes belong to the default group, the one associated to the default communicator MPI_COMM_WORLD . All other groups must be explicitly generated.

MPI processes may belong to different groups.







Given a communicator the following function returns the handle of the associated group:

interface
 subroutine mpi_comm_group(comm,group, ierr)
 integer, intent(in) :: comm
 integer, intent(out) :: group, ierr
 end subroutine mpi_comm_group
end interface

fortran

int MPI_Comm_group (MPI_Comm comm, MPI_Group *group)

c/c++





The following function enables generating a new group on the basis of an existing group. The process with rank RANKS (I) in the old group is given rank I in the new group:

interface subroutine mpi_group_incl(group, n, ranks, newgroup, ierr) integer, intent(in) :: group, n, ranks integer, intent(out) :: newgroup, ierr end subroutine mpi_group_incl end interface

fortran

int MPI_Group_incl(MPI_Group group, int n, int *ranks, MPI_Group *newgroup) c/c++





Example:

if GROUP contains 8 processes (numbered from 0 to 7) and the array has values RANKS (1:3) = (1, 5, 2), the instruction

call mpi_group_incl (group, 3, ranks, newgroup, ierr)

fortran

generates the new NEWGROUP with the three processes above numbered.

The following table shows correspondence between the two groups:

Group	Newgroup
1	0
5	1
2	2





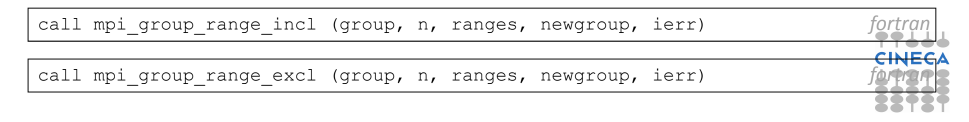


On the contrary in the following function the array RANKS (I) specify the processes of GROUP to be eliminated for building NEWGROUP:

```
interface fortran
subroutine mpi_group_excl(group, n, ranks, newgroup, ierr)
integer, intent(in) :: group, n, ranks
integer, intent(out) :: newgroup, ierr
end subroutine mpi_group_excl
end interface
```

int MPI_Group_excl(MPI_Group group, int n, int *ranks, MPI_Group *newgroup) c/c++

It is also possible to specify a range of indexes, like RANGES (1:N, 1:3). In the following functions the second dimension of the array specifies the first and the last index to be included and the stride.







Example:

If group contains 1000 processes and a new group is to be generated with half the number of the processes, taken from the odd positions, the array may be defined as RANGES (1, 1) = 2, RANGES (1, 2) = 1000, RANGES (1, 3) = 2and the program should issue the following instruction:

call mpi_group_range_excl (group, n, ranges, newgroup, ierr) fortran

The correspondence between the two groups would be:

Group	Newgroup
1	0
3	1
5	2







The operations to manage groups are local and do not involve communications.

The following function may be used to know the rank of each process in two different groups.

RANKS1 (:) are the known ranks of the processes in GROUP1

RANKS2(:) are the related ranks in GROUP2

```
interface
   subroutine mpi_group_translate_ranks(group1, n, ranks1, group2, &
        ranks2, ierr)
        integer, intent(in) :: group1, n, ranks1(:), group2
        integer, intent(out) :: ranks2(:), ierr
        end subroutine mpi_group_translate
        fortran
end interface
```

int MPI_Group_translate (group1, n, ranks1, group2, ranks2, ierr)

c/c++







It is possible to check similarity of two groups:

interface	fortran
<pre>subroutine mpi_group_compare(group1, group2, result, ierr)</pre>	
integer, intent(in) :: group1, group2	
integer, intent(out) :: result, ierr	
end subroutine mpi_group_compare	
end interface	

int MPI_Group_compare (group1, group2, result, ierr)

с/с++

The returned values may be one out of the following:

- MPI_IDENT if the groups have the same processes with identical ranks
- MPI_SIMILAR if the groups have the same processes but unequal ranks
- MPI_UNEQUAL if the groups are different





Communicators



A communicator defines the processes that can communicate each other. Each communicator has its own handle, is an opaque object and can be managed by proper functions only.

The default communicator is named MPI_COMM_WORLD, but in a real program it is often useful to generate additional communicators, to be able to directly manage communications among process subsets.

Communicator handles must always be specified in sending and receiving functions.





Communicators



Example: 0506MPIExample_two_comms

The default communicator is splitted in two sets of processes and an intercommunicator is created:

```
colour = id % 2;
MPI_Comm_split(GlobalComm, colour, id, &LocComm);
if ( colour == 0 ) {
    MPI_Intercomm_create(LocComm, 0, GlobalComm, 1,
        01, &InterComm);
} else {
    MPI_Intercomm_create(LocComm, 0, GlobalComm, 0,
        01, &InterComm);
}
```





fortran

Managing communicators

The following function may be used to generate a new communicator connected to an existing group:

inte	erface
	<pre>subroutine mpi_comm_create(comm, group, newcomm, ierr)</pre>
	integer, intent(in) :: comm, group
	integer, intent(out) :: newcomm, ierr
	end subroutine mpi_comm_create
end	interface

int MPI_Comm_create (MPI_Comm comm, MPI_Group group, MPI_Comm *newcomm) c/c++

- COMM is an existing communicator related to a wider process group; the function must be called by all processes in the COMM communicator
- GROUP is a sub-group of the process group related to COMM
- NEWCOMM is the handle of the newly generated communicator

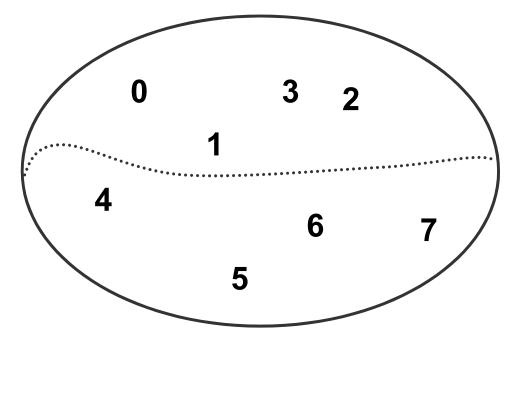






Managing communicators

Suppose there is a communicator connected to a group of 8 processes and 2 new communicators are required by dividing the communicator in two parts as follow:







To accomplish this task all the processes of the existing communicator shal issue the following instructions:

call mpi_comm_rank (comm, rank, ierr)	
call mpi_comm_size (comm, size, ierr)	
color = 2*rank/size	
key = size - rank - 1	
call mpi_comm_split (comm, color, key, newcomm, ierr)	fortran

Each process receives a new communicator handle and will have the rank:

Communicator 1		Communicator 2	
Rank in new group	Rank in old group	Rank in new group	Rank in old group
0	3	0	7
1	2	1	6
2	1	2	5
3	0	3	4

If for some process COLOR=MPI_UNDEFINED, the function MPI_COMM_SPLIT returns NEWCOMM=MPI_COMM_NULL





Once the processes have been separated in several groups it is possible to realize client-server connections by connecting disjoined groups.

Communications between separated groups can only be of point-to-point type: no collective communications are available.





Whenever a new inter-communicator has been created, the sending process must specify the rank of the receiving process (relevant to the other group); the receiving process must specify the rank of the sender (relevant to the other group).

To enable this, while the functions mpi_comm_size, mpi_comm_rank, mpi_comm_group return informations relevant to the local communicator, the functions mpi_comm_remote_size, mpi_comm_remote_group instead return informations on the disjoined intercommunicator group.

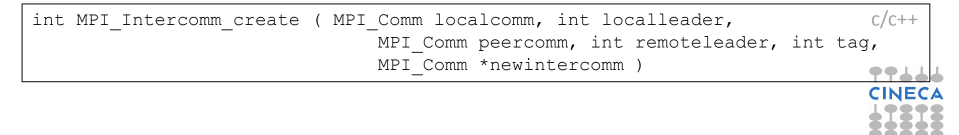


- A communicator connecting disjoined groups is called an intercommunicator and can be generated by calling the function mpi_intercomm_create. This function requires:
- A leading process for each one of two disjoined groups
- An intra-communicator between the two leading processes
- A tag for safe communications between the two leading processes





The following function generates an inter-communicator NEWINTERCOMM between the processes LOCALLEADER and REMOTELEADER of the intracommunicator LOCALCOMM, using TAG and the point-to-point communicator PEERCOMM. It should be noted that REMOTELEADER and PEERCOMM are referred to the local process, while TAG must have the same value for both the processes:





The intra-communicator NEWINTRACOMM may be generated from an intercommunicator INTERCOMM calling the function:

```
interface
    subroutine mpi_intercomm_merge(intercomm, high, newintracomm, ierr)
        integer, intent(in) :: intercomm, high
        integer, intent(out) :: newintracomm, ierr
        end subroutine mpi_intercomm_merge
end interface
```

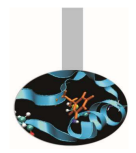
int MPI_Intercomm_merge(MPI_Comm intercomm,int high,MPI_Comm *newintracomm) c/c++

This way two separated groups may be joined. The value of HIGH must be the same for all the processes belonging to the same group. If HIGH = .FALSE. for group 1 and HIGH = .TRUE. for group 2, in the new communicating group the processes are ordered starting from group 1; i.e. the processes in group 2 have a higher rank.

Example: 0507MPIExample_comms_merge (Fortran and C)







In many programs it may be important to arrange the processes in a given topology. MPI enables the definition of topologies, with an explicit support for cartesian topology. This topology may be defined by calling the function:





Example: 0508MPIExample_cart_create

In this example a 2D cartesian topology is created to send a message along horizontal and vertical «bands».

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The MPI_CART_CREATE function returns the new communicator COMM_CART, connected to a grid with NDIMS dimensions. The extent of each dimension must be defined in LDIMS (1:NDIMS) and it is possible to specify periodicity for each dimension. The REORDER variable is used to allow reordering of the processes.

In cartesian topologies the processes are ordered by rows.

Functions dealing with informations and details about the topology associated to a communicator are available.







Given a communicator COMM, the function MPI_TOPO_TEST returns the associated topology:

MPI_GRAPH: graph topology
MPI_CART: cartesian topology
MPI_UNDEFINED: no topology

interface
 subroutine mpi_topo_test(comm, topol, ierr)
 integer, intent(in) :: comm
 integer, intent(out) :: topol, ierr
 end subroutine mpi_topo_test
end interface

int MPI Topo test (MPI Comm comm, int *topol)

с/с++







Given a communicator COMM, with cartesian topology, the function MPI CARTDIM GET returns the number of dimensions

interface	fortran
<pre>subroutine mpi_cartdim_get(comm, ndims, ierr)</pre>	
integer, intent(in) :: comm	
integer, intent(out) :: ndims, ierr	
end subroutine mpi_cartdim_get	
end interface	

int MPI_Cartdim_get (MPI_Comm comm, int *ndims)

С/С++





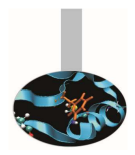


The function MPI_CART_GET, returns the number DIMS (:) of processes in each dimension, the periodicity for each dimension, the process coordinates.

interface	fortran
subroutine mpi_cart_get(comm, maxdims, dims, periods, coords, ierr)	
integer, intent(in) :: comm, maxdims	
integer, intent(out) :: ierr	
<pre>integer, dimension(:), intent(out) :: dims, coords</pre>	
logical, dimension(:), intent(out) :: periods	
end subroutine mpi_cart_get	
end interface	







Given a communicator associated to a cartesian topology and the process coordinates, the following function returns the process rank:

interface	
<pre>subroutine mpi_cart_rank(comm, coords, rank, ierr)</pre>	
integer, intent(in) :: comm	
<pre>integer, dimension(:), intent(in) :: coords</pre>	
integer, intent(out) :: rank, ierr	
end subroutine mpi_cart_rank	
end interface	fortran

int MPI_Cart_rank(MPI_Comm comm, int *coords, int *rank)

C/C++





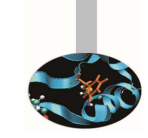
The following function returns the coordinates of a process in a cartesian topology:

interface	
subroutine mpi_cart_coords(comm, rank, maxdims, coords, ierr)	
integer, intent(in) :: comm, rank, maxdims	
<pre>integer, dimension(:), intent(out) :: coords</pre>	
integer, intent(out) :: ierr	
end subroutine mpi_cart_coords	
end interface	fortran

int MPI_Cart_coords(MPI_Comm comm, int rank, int maxdims, int *coords) c/c++







The following function helps defining the length in each dimension for a cartesian topology:

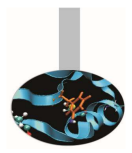
interface	
subroutine MPI_Dims_create(nnodes, ndims, dims)	
integer, intent(in) :: nodes, dims	
<pre>integer, dimension(:), intent(inout) :: dims</pre>	
integer, intent(out) :: ierr	
end subroutine MPI_Dims_create	
end interface	fortran

int MPI_Dims_create(int nnodes, int ndims, int *dims)

c/c++

The dims(:) array is filled only where value is 0. As an example if nnodes=6, ndims=3, dims=[0,3,0], the output is dims=[2,3,1]. Otherwise would it be dims=[0,0,0], the output would be dims=[3,2,1]





Topologies may be useful to send messages along specific directions.

As an example, suppose that every process in a cartesian topology has to send data toward the DIM dimension to a DELTA distance. The following function returns the ranks of the processes SOURCE and DEST

```
interface
    subroutine mpi_cart_shift(comm, dim, delta, source, dest, ierr)
        integer, intent(in) :: comm, dim, delta
        integer, intent(out) :: source, dest, ierr
        end subroutine mpi_cart_shift
end interface
```

int MPI_Cart_shift(MPI_Comm comm, int dim, int delta, int *source, int *dest) c/c++

to be passed to the function

CALL MPI_SENDRECV(SENDBUF,	SENDCOUNT, SENDTYPE, I	DEST, &	
SENDTAG,	RECVBUF, RECVCOUNT, REC	CVTYPE, &	,
SOURCE, R	ECVTAG, COMM, STATUS,	IERROR))



fortran





The following function generates new cartesian topologies by cutting a wider cartesian space along the given dimensions:

Example: if COMM is associated to a cartesian topology with extensions 2x3x4 and REMAIN_DIMS=(.T.,.T.,.F.), four new topologies are generated with extension 2x3.

Each process is returned one communicator handle; the former group is divided into 4 new groups with 6 processes each.



MPI+OpenMP



It is possible to develop parallel programs mixing MPI calls and OpenMP directives.

Intel compilers: mpixxx -openmp -O3 -o nomefile.exe nomefile.xxx

PGI compilers: mpi*XXX* -mp -O3 -o nomefile.exe nomefile.*XXX*

GNU compilers: mpi*xxx*-fopenmp -O3 -o nomefile.exe nomefile.*xxx*

Execution:

export OMP_NUM_THREADS=*threads* mpirun -np 2 -machinefile *mc* -x OMP_NUM_THREADS nomefile.exe

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