



MPI in Small Bites PPCES 2024

HPC.NRW Competence Network







THE COMPETENCE NETWORK FOR HIGH PERFORMANCE COMPUTING IN NRW.

MPI Concepts

HPC.NRW Competence Network

MPI in Small Bites



MPI in Small Bites – PPCES 2024

INNOVATION THROUGH COOPERATION.

Library Initialization



- MPI is implemented as a library, not a compiler extension
 - Common (non-local) objects need coordinated construction
 - Library needs to be initialized explicitly

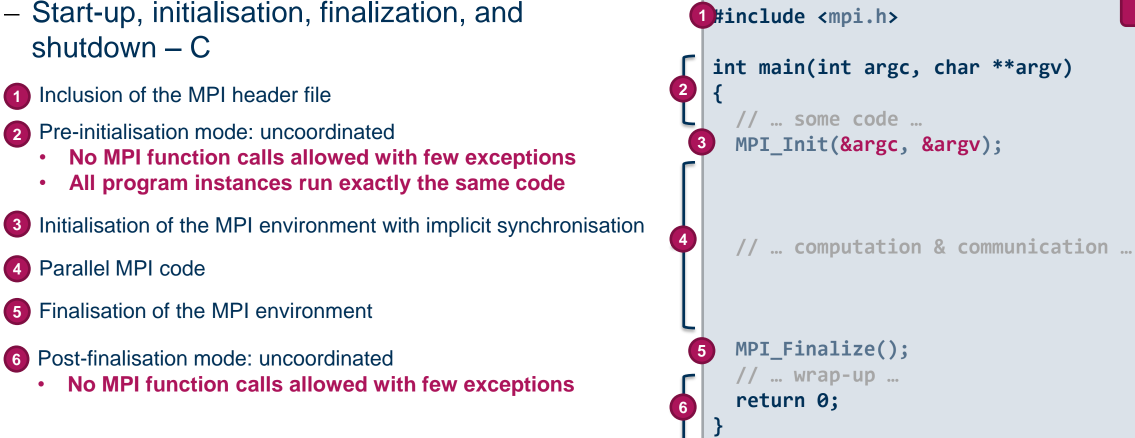
- Multiple methods exist to initialize MPI
 - Classic MPI (pre-MPI 4.0) without threads \rightarrow MPI_Init
 - Classic MPI (pre-MPI 4.0) with threads \rightarrow MPI_Init_thread
 - Covered in another part on hybrid programming
 - New MPI (MPI 4.0) with threads \rightarrow MPI_Session_init
 - Covered in another part on the session model



INNOVATION THROUGH COOPERATION.

MPI in Small Bites – PPCES 2024

Library Initialization (classic MPI – no threads)



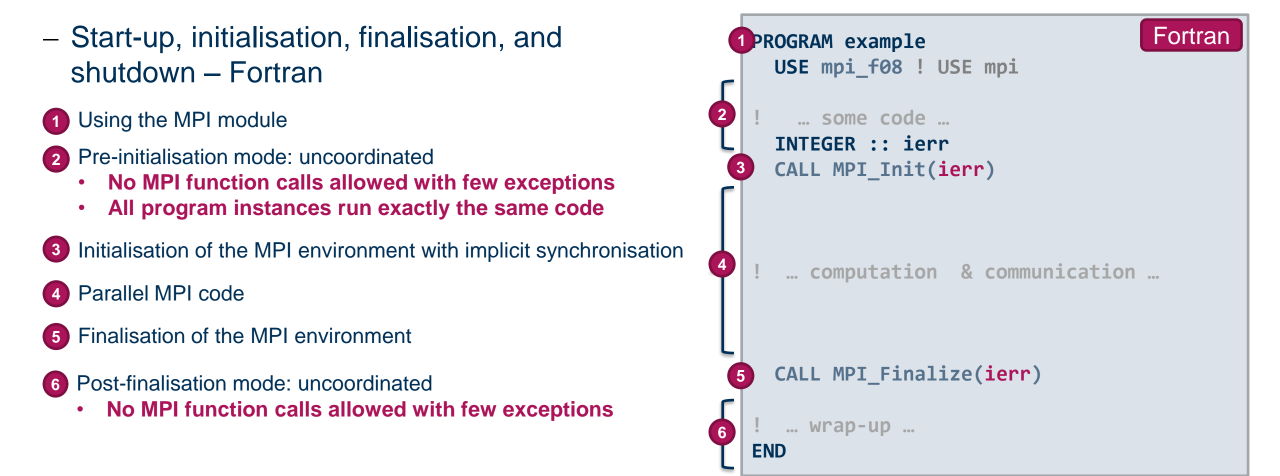




С

Library Initialization (classic MPI – no threads)







INNOVATION THROUGH COOPERATION.

Library Initialization (classic MPI – no threads)



– Initialization:

C: ierr = MPI_Init(&argc, &argv);
Fortran: CALL MPI_Init(ierr)

- Initializes the MPI library and makes the process member of MPI_COMM_WORLD
- [C] Both arguments must be either NULL or they *must* point to the arguments of main()
- May not be called more than once for the duration of the program execution
- Error code as return value in [C] and additional parameter in [F]

- Finalization:

C: ierr = MPI_Finalize(); Fortran: CALL MPI_Finalize(ierr)

- Cleans up the MPI library and prepares the process for termination
- Must be called once before the process terminates
- Having other code after the finalisation call is not recommended

General Structure of an MPI Program



- How many processes are there in total?

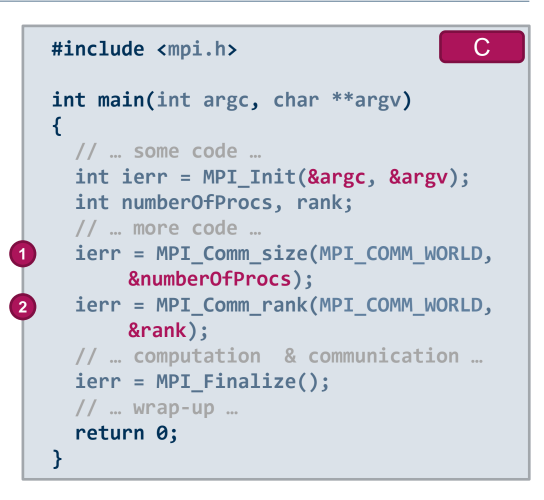
- Who am I?

1 Obtains the number of processes (ranks) in the MPI program

Example: if the job was started with 4 processes, then **numberOfProcs** will be set to 4 by the call

Obtains the identity of the calling process within the MPI program NB: MPI processes are numbered starting from 0

Example: if there are 4 processes in the job, then **rank** receives the value of 0 in the first process, 1 in the second process, etc.





General Structure of an MPI Program



- How many processes are there in total?
- Who am I?
- Obtains the number of processes (ranks) in the MPI program

Example: if the job was started with 4 processes, then **numberOfProcs** will be set to 4 by the call

Obtains the identity of the calling process within the MPI program NB: MPI processes are numbered starting from 0

Example: if there are 4 processes in the job, then **rank** receives the value of 0 in the first process, 1 in the second process, etc.

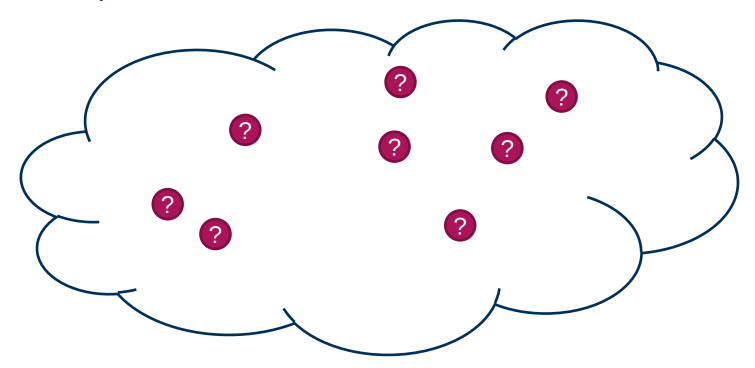








- The processes in any MPI program are initially indistinguishable
- MPI assigns each process a unique identity (rank) in a communication context (communicator)

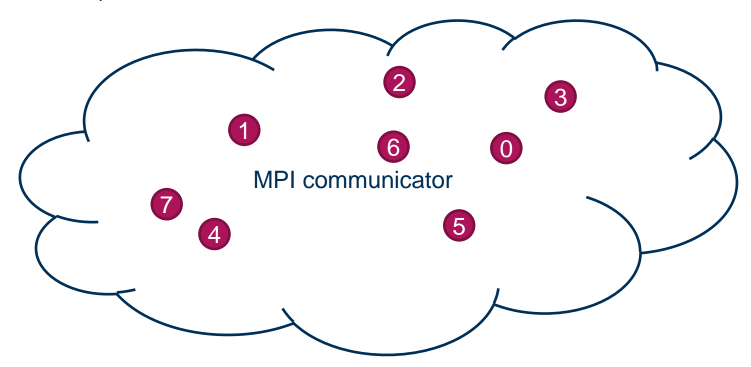








- The processes in any MPI program are initially indistinguishable
- MPI assigns each process a unique identity (rank) in a communication context (communicator)









- The processes in any MPI program are initially indistinguishable (for the user)
- MPI assigns each process a unique identity (rank) in a communication context (communicator)
- Ranks
 - Range from 0 to n-1 (with n processes in the communicator)
 - An MPI process can have different ranks in different communicators
- Communicators
 - Logical contexts where communication takes place
 - Comprises a group of MPI processes with some additional information
 - MPI_COMM_WORLD is implicitly available
 - Comprises all processes initially started with the MPI program



MPI as an SPMD Environment

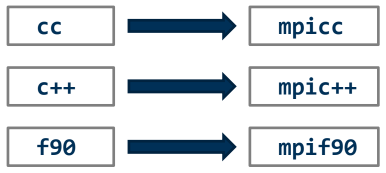


- Provide dynamic identification of all peers
- Who am I and who else is also working on this problem?
- 2. Provide robust mechanisms to exchange data
 - Whom to send data to / From whom to receive the data?
 - How much data?
 - What kind of data?
 - Has the data arrived?
- 3. Provide synchronisation mechanisms
 - Have all processes reached same point in the program execution flow?
- 4. Provide methods to launch and control a set of processes
 - How do we start multiple processes and get them to work together?
- 5. Portability

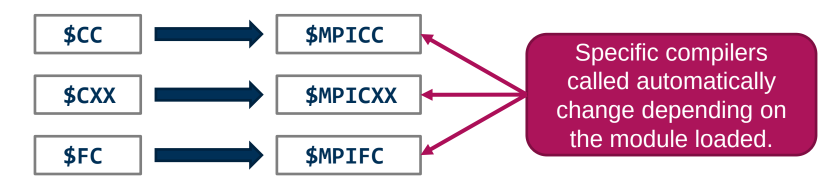
Compiling MPI Programs



- MPI is a typical library with C header files, Fortran modules, etc.
- Most MPI vendors provide convenience compiler wrappers (names are not standardized!)



- On the RWTH Aachen Compute Cluster:





INNOVATION THROUGH COOPERATION.



– RWTH Aachen Cluster defines additional environment variables to minimize confusion

```
cluster:~> $MPICC -show
                                                    # instruct wrapper to show compile line
icc \
-I"/cvmfs/[...]/impi/2021.6.0-intel-compilers-2022.1.0/mpi/2021.6.0/include" \
-L"/cvmfs/[...]/impi/2021.6.0-intel-compilers-2022.1.0/mpi/2021.6.0/lib/release" \
-L"/cvmfs/[...]/impi/2021.6.0-intel-compilers-2022.1.0/mpi/2021.6.0/lib" \
-Xlinker --enable-new-dtags -Xlinker -rpath \
-Xlinker "/cvmfs/[...]/impi/2021.6.0-intel-compilers-2022.1.0/mpi/2021.6.0/lib/release" \
-Xlinker -rpath \
-Xlinker "/cvmfs/[...]/impi/2021.6.0-intel-compilers-2022.1.0/mpi/2021.6.0/lib" \
-lmpifort -lmpi -ldl -lrt -lpthread
cluster:~> echo $MPICC
                                                    # check compiler wrapper name
mpiicc
cluster:~> module purge; module load gompi # switch MPI implementation
cluster:~> echo $MPICC
                                                    # check compiler wrapper name again
mpicc
```





– RWTH Aachen Cluster defines additional environment variables to minimize confusion

```
cluster:~> module purge && module load gompi
cluster:~> echo $MPICC
mpicc
cluster:~> $MPICC -show
gcc \
-I/cvmfs/[...]/OpenMPI/4.1.4-GCC-11.3.0/include \
-L/cvmfs/[...]/OpenMPI/4.1.4-GCC-11.3.0/lib \
-L/cvmfs/[...]/hwloc/2.7.1-GCCcore-11.3.0/lib \
[...]
-Wl,/cvmfs/[...]/libevent/2.1.12-GCCcore-11.3.0/lib -Wl,--enable-new-dtags -lmpi
```





- Most MPI implementations provide a special launcher program:

```
mpiexec -n nprocs ... program <arg1> <arg2> <arg3> ...
```

- Launches nprocs instances of program with command-line arguments arg1, arg2, ...
- The standard specifies the **mpiexec** program, but does not require it:
 - IBM BG/Q: runjob --np 1024 ...
 - SLURM resource manager: srun -n 96 -N 1 ...



Executing MPI Programs



- The launcher often performs more than simply launching processes:
 - Helps MPI processes find each other and establish the world communicator
 - Redirects the standard output of all ranks to the terminal
 - Redirects the terminal input to the standard input of rank 0
 - Forwards received signals (Unix-specific)



MPI as an SPMD Environment



- Provide dynamic identification of all peers
- Who am I and who else is also working on this problem?
- 2. Provide robust mechanisms to exchange data
 - Whom to send data to / From whom to receive the data?
 - How much data?
 - What kind of data?
 - Has the data arrived?
- 3. Provide synchronisation mechanisms
 - Have all processes reached same point in the program execution flow?
 - Provide methods to launch and control a set of processes
 - How do we start multiple processes and get them to work together?

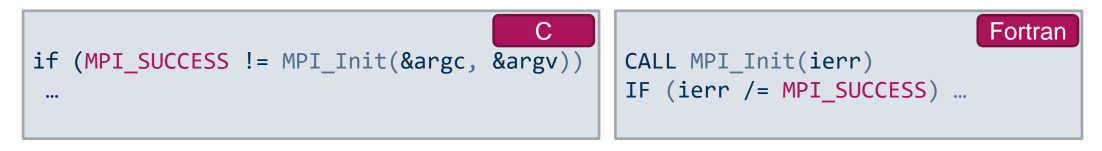


Portability

Error handling



- Error codes indicate the success of the operation:
 - Failure is indicated by error codes other than MPI_SUCCESS



- An MPI error handler is called first before the call returns
 - The default error handler for non-I/O calls aborts the entire MPI program!
 - Error checking in simple programs is redundant
- Actual MPI error code values are implementation specific
 - Use MPI_Error_string to derive human readable information



Handles to Opaque Objects



- MPI objects (e.g., communicators) are referenced via handles
 - Process-local values
 - Cannot be passed from one process to another
 - Objects referenced by handles are opaque
 - Structure is implementation dependent
 - Blackbox for the user
- C (mpi.h)
 - typedef'd handle types: MPI_Comm, MPI_Datatype, MPI_File, etc.



Handles to Opaque Objects II



- Fortran (USE mpi)
 - All handles are INTEGER values
 - Easy to pass the wrong handle type

- Fortran 2008 (USE mpi_f08)
 - Wrapped INTEGER values: TYPE(MPI_Comm), TYPE(MPI_File), etc.
 - The INTEGER handle is still available: comm%MPI_VAL



Datatype Handles



– MPI is a library

- Cannot infer datatypes of supplied buffers at runtime
- User needs to provide additional information on buffer type

- MPI datatype handles tell the MPI library how to:
 - read binary values from the send buffer
 - write binary values into the receive buffer
 - correctly apply value alignments
 - convert between machine representations in heterogeneous environments



Datatype Handles II



- MPI datatypes are handles

- Cannot be used to declare variables of a specific language type
- sizeof(MPI_INT) provides the size of a datatype handle NOT the size of an int in C

– Type Signatures

- Sequence of basic datatypes in a buffer
- Basic datatypes correspond to native language datatypes
- Type Maps
 - Sequence of basic datatypes **AND** their location in a buffer



Basic MPI Datatypes



– MPI provides predefined datatypes for each language binding:

MPI data type	C data type	MPI data type	Fortran data type
MPI_CHAR	char	MPI_INTEGER	INTEGER
MPI_SHORT	short	MPI_REAL	REAL
MPI_INT	int	MPI_REAL8	REAL(KIND=8)
MPI FLOAT	float	MPI_DOUBLE_PRECISION	DOUBLE PRECISION
MPI DOUBLE	double	MPI_COMPLEX	COMPLEX
MPI UNSIGNED INT	unsigned int	MPI_LOGICAL	LOGICAL
		MPI_CHARACTER	CHARACTER(1)
MPI_BYTE	-		
		MPI_BYTE	-
	8 binary digits		
	no conversion		
u	sed for untyped data		



INNOVATION THROUGH COOPERATION.

Local vs. Non-local Procedures



- Non-local procedures may require,
 - during its execution,
 - some specific, semantically-related MPI procedure
 - to be called on another MPI process"

Local procedure are not non-local



MPI Operations



- MPI defines several operations, which are
 - a sequence of steps
 - performed by the MPI library
 - to stablish and enable
 - data transfer
 - and/or synchronization
- Four stages
 - 1. Initialization Resources (argument lists, buffer address, etc.) are handed to the MPI library
 - 2. Starting The operation takes over control of the resources (buffer contents)
 - 3. Completion Return control of the resources (buffer contents)
 - 4. Freeing Return control of the remaining resources



Blocking vs. Non-blocking vs. Asynchronous



- Blocking procedures return when the associated operation is complete locally
 - Any input argument can be safely reused or deallocated
 - Operation may not be completed remotely
- Non-blocking procedures return before associated operation is complete locally
 - One or more additional calls are needed to complete operation
 - Input arguments may not be written or deallocated until operation is complete
- Synchronous operations complete locally only with specific remote intervention
 - Asynchronous operations may complete locally without remote intervention



MPI Communication Paradigms







Point-to-Point Communication

Collective Communication



One-sided Communication

