

Introduction to MPI+OpenMP hybrid programming

SuperComputing Applications and Innovation Department





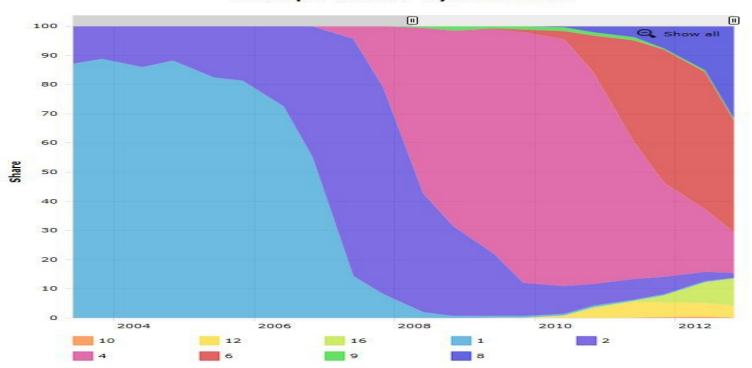
Basic concepts







Cores per Socket - Systems Share









- In a nutshell:
 - memory per core decreases
 - memory bandwidth per core decreases
 - number of cores per socket increases
 - single core clock frequency decreases

 Programming model should follow the new kind of architectures available on the market: what is the most suitable model for this kind of machines?







- Distributed parallel computers rely on MPI
 - strong
 - consolidated
 - standard
 - enforce the scalability (depending on the algorithm) up to a very large number of tasks
- but... is it enough when memory is such small amount on each node?

Example: Bluegene/Q has 16GB per node and 16 cores. Can you imagine to put there more than 16MPI (tasks), i.e. less than 1GB per core?







- On the other side, OpenMP is a standard for shared memory systems
- Pthreads execution models is a lower-level alternative, but OpenMP is often a better choice for HPC programming
- OpenMP is robust, clear and sufficiently easy to implement but
 - depending on the implementation, typically the scaling on the number of threads is much less effective than the scaling on number of MPI tasks
- Putting together MPI with OpenMP could permit to exploit the features of the new architectures, mixing these paradigms



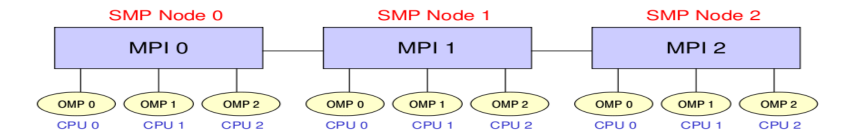




Hybrid model: MPI+OpenMP

- In a single node you can exploit a shared memory parallelism using OpenMP
- Across the nodes you can use MPI to scale up

Example: on a Bluegene/Q machine you can put 1 MPI task on each node and 16 OpenMP threads. If the scalability on threads is good enough, you can use all the node memory.









Pure MPI Pro:

- High scalability
- High portability
- No false sharing
- ❖ Scalability out-of-node

Pure MPI Con:

- * Hard to develop and debug.
- **Explicit communications**
- Coarse granularity
- Hard to ensure load balancing









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Pure OpenMP Pro:

Easy to deploy (often)

Low latency

Implicit communications

Coarse and fine granularity

Dynamic Load balancing

Pure OpenMP Con:

Only on shared memory machines

Intranode scalability

Possible data placement problem

Undefined thread ordering







- Conceptually simple and elegant
- Suitable for multicore/multinodes architectures
- Two-level hierarchical parallelism
- In principle, you can alleviate problems related to the scalability of MPI, reducing the number of tasks and network flooding







- OpenMP introduces fine granularity parallelism
- Loop-based parallelism
- Task construct (OpenMP 3.0): powerful and flexible
- Load balancing can be dynamic or scheduled
- All the work is in charge to the compiler
- No explicit data movement







- Using a hybrid approach means to balance the hierarchy between MPI tasks and thread.
- MPI in most cases (but not always) occupy the upper level respect to OpenMP
 - usually you assign *n* threads per MPI task, not m MPI tasks per thread
- The choice about the number of threads per MPI task strongly depends on the kind of application, algorithm or kernel. (this number can change inside the application)
- There's no golden rule. More often this decision is taken a-posteriori after benchmarks on a given machine/architecture







- Using a hybrid approach MPI+OpenMP can lower the number of MPI tasks used by the application.
- Memory footprint can be alleviated by a reduction of replicated data on MPI level
- Speed-up limited due algorithmic issues can be solved (because you're reducing the amount of communication)





Reality is bitter

- In real scenarios, mixing MPI and OpenMP, sometimes, can make your code slower
 - If you exceed with the number of OpenMP threads you can encounter problems with locking of resources
 - Sometimes threads can stay in a idle state (spin) for a long time
 - Problems with cache coherency and false sharing
 - Difficulties in the management of variables scope





Cache coherency and false sharing

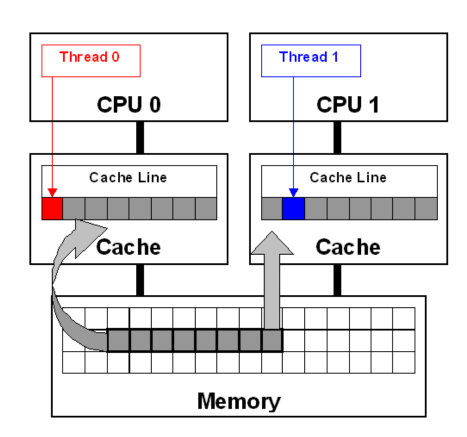
- It is a side effects of the cache-line granularity of cache coherence implemented in shared memory systems.
- The cache coherency implementation keep track of the status of cache lines by appending *state bits to* indicate whether data on cache line is still valid or outdated.
- Once the cache line is modified, cache coherence notifies other caches holding a copy of the same line that its line is invalid.
- If data from that line is needed, a new updated copy must to be fetched.





```
#pragma omp parallel for
shared(a) schedule(static,1)
for (int i=0; i<n; i++)
    a[i] = i;</pre>
```











- The most simple recipe is:
 - start from a serial code and make it a MPI-parallel code
 - implement for each of the MPI task a OpenMP-based parallelization
- Nothing prevents to implement a MPI parallelization inside a OpenMP parallel region
 - in this case, you should take care of the thread-safety
- To start, we will assume that only the master thread is allowed to communicate with others MPI tasks





A simple hybrid code (correct?)

```
call MPI INIT (ierr)
call MPI COMM RANK (...)
call MPI COMM SIZE (...)
... some computation and MPI communication
call OMP SET NUM THREADS(4)
!SOMP PARALLEL
!$OMP DO
 doi=1,n
    ... computation
 enddo
!$OMP END DO
!$OMP END PARALLEL
... some computation and MPI communication
call MPI FINALIZE (ierr)
```







Advantages:

- Simplest hybrid parallelization (easy to understand and to manage)
- No message passing inside a SMP node

Disadvantages:

- All other threads are sleeping during MPI communications
- Thread-safe MPI is required







- MPI_INIT_THREAD (required, provided, ierr)
 - IN: required, desired level of thread support (integer).
 - OUT: provided, provided level (integer).
 provided may be less than required.
- Four levels are supported:
 - MPI_THREAD_SINGLE: Only one thread will run. Equals to MPI_INIT.
 - MPI_THREAD_FUNNELED: processes may be multithreaded, but only the main thread can make MPI calls (MPI calls are delegated to main thread)
 - MPI_THREAD_SERIALIZED: processes could be multi-threaded More than one thread can make MPI calls, but only one at a time.
 - MPI_THREAD_MULTIPLE: multiple threads can make MPI calls, with no restrictions.







- The various implementations differ in levels of thread-safety
- If your application allows multiple threads to make MPI calls simultaneously, without MPI_THREAD_MULTIPLE, is not thread-safe
- Using OpenMPI, you have to use -enable-mpi-threads at configure time to activate all levels
 - see more later
- Higher level corresponds to higher thread-safety. Use the required safety needs.





MPI_THREAD_SINGLE

- There are no additional user thread in the system
 - E.g., there are no OpenMP parallel regions
 - MPI_Init_thread with MPI_THREAD_SINGLE is fully equivalent to MPI_Init

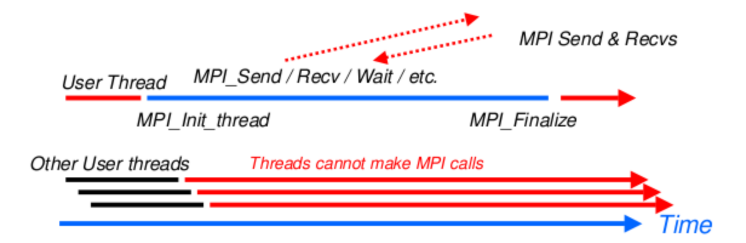
```
int main(int argc, char ** argv)
  int buf[100];
  MPI_Init(&argc, &argv);
  MPI_Comm_rank(MPI_COMM_WORLD, &rank);
  for (i = 0; i < 100; i++)
    compute(buf[i]);
  /* Do MPI stuff */
  MPI Finalize();
  return 0:
```





MPI_THREAD_FUNNELED

- It adds the possibility to make MPI calls inside a parallel region, but only the master thread is allowed to do so
 - All MPI calls are made by the master thread
 - The programmer must guarantee that!







MPI_THREAD_FUNNELED

```
int main(int argc, char ** argv)
int buf[100], provided;
MPI Init thread(&argc, &argv, MPI THREAD FUNNELED, &provided);
if (provided < MPI THREAD FUNNELED) MPI Abort(MPI COMM WORLD,1);
#pragma omp parallel for
for (i = 0; i < 100; i++)
   compute(buf[i]);
/* Do MPI stuff */
MPI_Finalize();
return 0;
```







- MPI function calls can be: outside a parallel region or in a parallel region, enclosed in "omp master" clause
- There is no synchronization at the end of a "omp master" region, so a barrier is needed before and after to ensure that data buffers are available before/after the MPI communication

!\$OMP BARRIER !\$OMP MASTER call MPI_Xxx(...) !\$OMP END MASTER !\$OMP BARRIER

#pragma omp barrier
#pragma omp master
 MPI_Xxx(...);
#pragma omp barrier

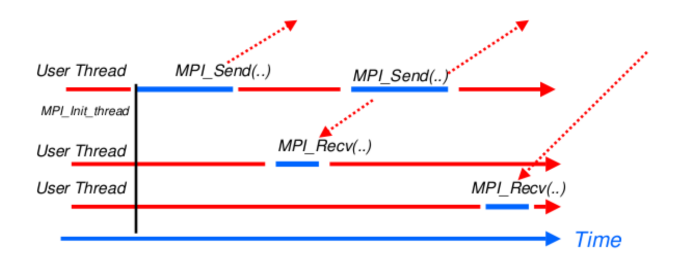






MPI_THREAD_SERIALIZED

- Multiple threads may make MPI calls, but only one at a time:
 - MPI calls are **not** made concurrently from two distinct threads. MPI calls are "serialized"
 - The programmer must guarantee that!







MPI_THREAD_SERIALIZED

```
int main(int argc, char ** argv)
int buf[100], provided;
MPI_Init_thread(&argc, &argv, MPI_THREAD_SERIALIZED, &provided);
if (provided < MPI_THREAD_SERIALIZED) MPI_Abort(MPI_COMM_WORLD,1);
#pragma omp parallel for
for (i = 0; i < 100; i++) {
  compute(buf[i]);
#pragma omp critical
  /* Do MPI stuff */
MPI Finalize(); Return 0;
```







- MPI calls can be outside parallel regions, or inside, but enclosed in a "omp single" region (it enforces the serialization) or "omp critical" or ...
- Again, a starting barrier may be needed to ensure data consistency
 - But at the end of *omp single* there is an automatic barrier
 - Unless nowait is specified

!\$OMP BARRIER !\$OMP SINGLE call MPI_Xxx(...) !\$OMP END SINGLE

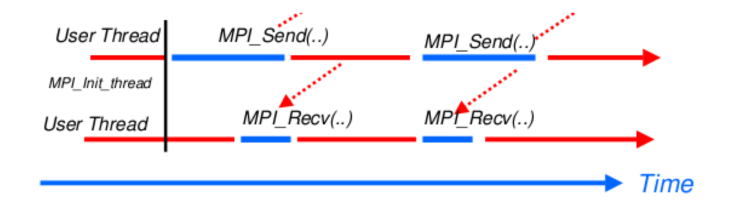
#pragma omp barrier
#pragma omp single
 MPI_Xxx(...);







- It is the most flexible mode, but also the most complicate one
- Any thread is allowed to perform MPI communications, without any restrictions.







MPI_THREAD_MULTIPLE

```
int main(int argc, char ** argv)
int buf[100], provided;
MPI_Init_thread(&argc, &argv, MPI_THREAD_MULTIPLE, &provided);
if (provided < MPI THREAD MULTIPLE) MPI Abort(MPI COMM WORLD,1);
#pragma omp parallel for
for (i = 0; i < 100; i++) {
  compute(buf[i]);
  /* Do MPI stuff */
MPI Finalize();
return 0;
```







- Ordering: when multiple threads make MPI calls concurrently the outcome will be as if the calls executed sequentially in some (any) order
 - Ordering is maintained within each thread
 - User must ensure that collective operations on the same communicator windows or file handle are correctly ordered among threads
 - E.g. cannot call a broadcast on one thread and a reduce on another thread on the same communicator
 - It is the user's responsibility to prevent races when threads in the same application post conflicting MPI calls
 - E.g. accessing an info object from one thread and freeing it from another thread
- Blocking: Blocking MPI calls will block only the calling thread and will not prevent other threads from running or executing MPI functions







- Incorrect example with collectives
 - P0 and P1 can have different ordering of Bcase on Barrier
 - Here the user must use some kind of synchronization to ensure that either thread 1 or thread 2 gets scheduled first on both processes
 - Otherwise a broadcast may get matched with a barrier on the same communicator, which is not allowed in MPI

	Process 0	Process 1
Thread 0	MPI_Bcast(comm)	MPI_Bcast(comm)
Thread 1	MPI_Barrier(comm)	MPI_Barrier(comm)







- Incorrect example with object Management
 - The user has to make sure that one thread is not using an object while another thread is freeing it
 - This is essentially an ordering issue; the object might get freed before it is used

	Process 0	Process 1
Thread 0	MPI_Bcast(comm)	MPI_Bcast(comm)
Thread 1	MPI_Comm_free(comm)	MPI_Comm_free(comm)







- Correct example with point-to-point
 - An implementation must ensure that the example below never deadlocks for any ordering of thread execution
 - That means the implementation cannot simply acquire a thread lock and block within an MPI function. It must release the lock to allow other threads to make progress

	Process 0	Process 1
Thread 0	MPI_Recv(src=1)	MPI_Recv(src=0)
Thread 1	MPI_Send(dst=1)	MPI_Send(dst=0)







Funneled

- All threads but the master are sleeping during MPI communications
- Only one thread may not be able to lead up to max inter-node bandwidth

Pure MPI

Each CPU can lead up max inter-node bandwidth

Hints: overlap as much as possible communications and computations





Overlap communications and computations

- In order to overlap communications with computations, the first step is using MPI_THREAD_FUNNELED mode
- While the master threads (a master thread for each MPI rank) are exchanging data, the other threads performs computation
- The tricky part is separating code that can run before or after the data exchanged are available

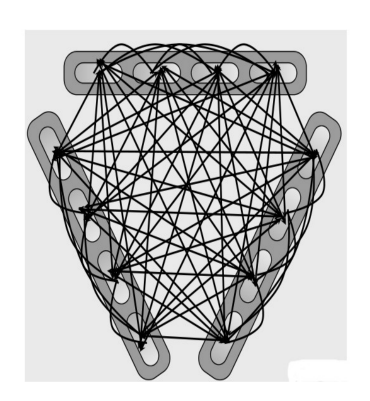
```
!$OMP PARALLEL
if (thread_id==0) then
call MPI_xxx(...)
else
do some computation
endif
!$OMP END PARALLEL
```





- MPI collectives are highly optimized
- Several point-to-point communication in one operations
- They can hide from the programmer a huge volume of transfer (MPI_Alltoall generates almost 1 million point-topoint messages using 1024 cores)
- There is no non-blocking (no longer the case in MPI 3.0)



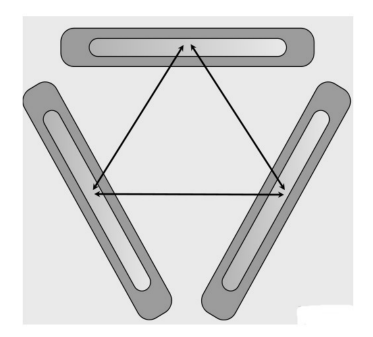






- Better scalability by a reduction of both the number of MPI messages and the number of process. Tipically:
- for all-to-all communications, the number of transfers decrease by a factor #threads^2
- the length of messages increases by a factor #threads
- Allow to overlap communication and computation.





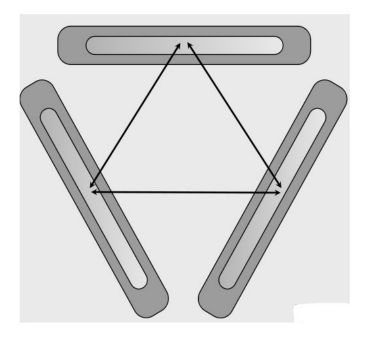




Restrictions:

- In MPI_THREAD_MULTIPLE mode is forbidden at any given time two threads each do a collective call on the same communicator (MPI_COMM_WORLD)
- 2 threads calling each a MPI_Allreduce may produce wrong results
- Use different communicators for each collective call
- Do collective calls only on 1 thread per process (MPI_THREAD_SERIALIZED mode should be fine)











- Introduction of OpenMP into existing MPI codes includes OpenMP drawbacks (synchronization, overhead, quality of compiler and runtime...)
- A good choice (whenever possible) is to include into the MPI code a multithreaded,
 optimized library suitable for the application.
- BLAS, LAPACK, MKL (Intel), FFTW are well known multithreaded libraries available in the HPC ecosystem.
 - Some libraries create their own threads: must be called outside our "omp parallel" regions
 - Otherwise, check "how much" the the library is thread-safe, if at least can be called by the master thread of your omp region (MPI_THREAD_FUNNELED)
 - Look carefully at the doc of your library, e.g.
 http://www.fftw.org/doc/Usage-of-Multi_002dthreaded-FFTW.html







BGQ benchmark example

Up to 64 hardware threads per process are available on bgq (SMT)

Huge simulation, 30000x30000 points. Stopped after 100 iterations only for timing purposes.

Number of threads / processes	MPI+OpenMP (TOT= 64 MPI, 1PPN) MPI_THREAD_MULTIPLE version Elapsed time (sec.)	MPI ONLY (TOT= 1024 MPI, 16,32,64 ppn Elapsed time (sec.)
1	78.84	N.A
4	19.89	N.A
8	10.33	N.A
16	5.65	5.98
32	3.39	7.12
64	2.70	12.07







- Better scalability by a reduction of both the number of MPI messages and the number of processes involved in collective communications and by a better load balancing.
- Better adequacy to the architecture of modern supercomputers while MPI is only a flat approach.
- Optimization of the total memory consumption (through the OpenMP shared-memory approach, savings in replicated data in the MPI processes and in the used memory by the MPI library itself).
- Reduction of the footprint memory when the size of some data structures depends directly on the number of MPI processes.
- It can remove algorithmic limitations (maximum decomposition in one direction for example).







Applications that can benefit from hybrid approach:

- Codes having limited MPI scalability (through the use of MPI_Alltoall for example).
- Codes requiring dynamic load balancing
- Codes limited by memory size and having many replicated data between MPI processes or having data structures that depends on the number of processes.
- Inefficient MPI implementation library for intra-node communication.
- Codes working on problems of fine-grained parallelism or on a mixture of fine and coarse-grain parallelism.
- Codes limited by the scalability of their algorithms.







- Hybrid programming is complex and requires high level of expertise.
- Both MPI and OpenMP performances are needed (Amdhal's law apply separately to the two approaches).
- Savings in performances are not guaranteed (extra additional costs).





PGAS programming model

- PGAS: Partitioned Global Address Space
 - a programming model somehow alternative to MPI and OpenMP models
 - but possibly mixable
 - Tries to combine SPMD (MPI) power with data referencing semantics of shared memory systems
- MPI → Library ; OpenMP → Annotations ; PGAS → Language
- PGAS is the basis of Unified Parallel C and Coarray Fortran
 - UPC: Unified Parallel C http://upc.lbl.gov/
 - CAF: Coarray Fortran, included in Fortran 2008 standard. Implementation available on common compilers (check also https://opencoarrays.org)
- A promising approach...





A touch of CAF

```
program Hello World
 implicit none
 integer :: i ! Local variable
 character(len=20) :: name[*]! scalar coarray, one
"name" for each image.
 ! Note: "name" is the local variable while
"name[<index>]" accesses the
 ! variable in a specific image; "name[this image()]" is
the same as "name".
 ! Interact with the user on Image 1; execution for all
others pass by.
 if (this image() == 1) then
  write(*,'(a)',advance='no') 'Enter your name: '
  read(*,'(a)') name
```

```
! Distribute information to other images
  do i = 2, num images()
   name[i] = name
  end do
 end if
 sync all! Barrier to make sure the data have arrived.
 ! I/O from all images, executing in any order, but
 ! each record written is intact.
 write(*,'(3a,i0)') 'Hello ',trim(name), &
                  'from image', this image()
end program Hello world
```





Implementations and cluster usage notes





MPI implementations and thread support

- An implementation is not required to support levels higher than
 MPI_THREAD_SINGLE, that is, an implementation is not required to be thread safe
- Most MPI implementations support a very low default thread support
 - Usually no support (MPI_THREAD_SINGLE)
 - And probably MPI_THREAD_FUNNELED (even if not explicitly)
 - Which (usually) is ok for cases where MPI communications are called outside of OMP parallel regions (where MPI_THREAD_FUNNELED would be strictly required)
 - Implementations with thread support are more complicated, error-prone and sometimes slower
- Checking the MPI_Init_thread provided support is a good programming practice





MPI implementations: check thread support

```
required = MPI_THREAD_MULTIPLE;
ierr = MPI_Init_thread(&argc,&argv,required,&provided);
if(required != provided) {
  if(rank == 0) { fprintf(stderr,"incompatible MPI thread support\n");
    fprintf(stderr,"required,provided: %d %d\n", required,provided); }
  ierr = MPI_Finalize(); exit(-1);
call MPI_Init_thread(required, provided, ierr)
if(provided /= required) then
  if(rank == 0) then
    print*,'Attention! incompatible MPI thread support'
    print*,'THREAD support required, provided: ',required, provided
  endif
  call MPI Finalize(ierr); STOP
endif
```





MPI implementations and thread support / 2

- Beware: the lack of thread support may result in subtle errors (not always clear)
- OpenMPI (1.8.3): when compiling OpenMPI there is a configure option to specify:
 - --enable-mpi-thread-multiple
 - [Enable MPI_THREAD_MULTIPLE support (default: disabled)]
- IntelMPI (5.0.2): both thread safe/non-thread safe versions are available:
 - Specify the option -mt_mpi when compiling your program to link the thread safe version of the Intel(R) MPI Library
 - E.g. mpif90 -mt_mpi -qopenmp main.f90





MPI and resource allocation

- The problem: how to distribute and control the resource allocation and usage (MPI processes/OMP threads) within a resource manager
 - How to use all the allocated resources: if n cores per node have been allocated how to run my program using all of that cores
 - Not less, maximize performance
 - Not more, do not interact with jobs of other users
 - How to use at its best the resource
 - optimal mapping between MPI processes/OMP threads and physical cores





MPI and resource allocation / 2

- Most MPI implementations allow a tight integration with resource managers in order to ease the usage of the requested resources
- The integration may enforce some constraints or just give hints to the programmer
 - Strict: only the requested physical resources can be used by my job
 - Soft: the programmer may use resources not explicitly allocated but only forcing the default settings





Cineca Marconi: PBSPro and MPI

- Cineca Marconi cluster assign resources using PBSProfessional 13
 - Strict mode is guaranteed through *cgroups* (control groups, a feature of Linux kernel which limits, isolate the usage of resources CPU, memory, I/O, network, etc.. of a process group). The processes can utilize only the resources assigned by the scheduler (minimizing the interaction with other jobs simultaneously running)
- OpenMPI and IntelMPI implementations available, both supporting PBS integration
 - module load autoload openmpi/1.10.3--gnu--6.1.0
 - module load autoload openmpi/1.10.3-threadmultiple--gnu--6.1.0
 - module load autoload intelmpi/2017--binary





Review on PBS select option

- -lselect=<n_chunks>:ncpus:<n_cores>:mpiprocs:<n_mpi>
 - Beware: n_chunks usually means n_nodes but this is not guaranteed provided that two or more chunks of n_cores can be allocated on a single node
 - mpiprocs must be well understood, let us examine the file \$PBS_NODEFILE created by PBS listing the allocated resources for some example cases
 - 6 MPI processes per chunk (node) and 6 cores reserved:useful for pure MPI runs

		node007	node003	node001	node015
		node007	node003	node001	node015
	ncpus=6	node007	node003	node001	node015
		node007	node003	node001	node015
		node007	node003	node001	node015
		node007	node003	node001	node015





Review on PBS select option / 2

• 1 MPI process per chunk (node), but 6 cores reserved, useful for hybrid case (OMP_NUM_THREADS=6)

select=4
ncpus=6
mpiprocs=1

node007	node003	node001	node015
---------	---------	---------	---------

• 2 MPI processes per chunk (node), but 6 cores reserved, again for hybrid cases (OMP_NUM_THREADS=3)

select=4 ncpus=6 mpiprocs=2

node007	node003	node001	node015
node007	node003	node001	node015







OpenMPI and IntelMPI

- Resource allocation is not aware of the MPI implementation which will be used
 - But MPI implementations give a different meaning and enforce different constraints
 - We will discuss OpenMPI and IntelMPI implementations
 - Basically, mpirun <executable> runs MPI processes taken from \$PBS_NODEFILE
 - But...
- IntelMPI allows to run a number of MPI processes even larger than the requested mpiprocs (I.e.the number of lines of the file \$PBS_NODEFILE)
 - But if the PBS strict mode is active, the user will use only the reserved cores
 - If the number of MPI processes is larger than ncpus, we are experimenting the so called oversubscribing
- OpenMPI imposes mpiprocs as the upper limit for the number of MPI processes which can be run
 - Since mpiprocs<ncpus, no oversubscribing is possible





OpenMPI and IntelMPI

- It is possible to modify how the \$PBS_NODEFILE is interpreted by mpirun
- OpenMPI:
 - -npernode <n>
 - With -npernode it is possible to specify how many MPI processes to run on each node
- IntelMPI has the options :
 - -perhost <n> || -ppn <n>
 - but these do not work at least on our installations
 - Since IntelMPI allows to run more processes than mpiprocs declared to PBS, you can specify mpiprocs=1 and then launch more processes which will be allocated roundrobin





OpenMPI and IntelMPI

- It is also possible to completely override the \$PBS_NODEFILE manually imposing a MPI machine file
 - Some limitations still apply
 - IntelMPI:
 - -machine {name} | -machinefile {name}
 - OpenMPI
 - -machinefile {machinefile} || --machinefile {machinefile} || -hostname {machinefile}





Where is my job running?

- It may seem a silly question but it is not
- How to check which resources have been allocated for my job
 - qstat -n1 <job_id> (for completed jobs add the -x flag)
 - qstat -f <job_id> |grep exec_host (for completed jobs add the -x flag)
 - Cat \$PBS_NODEFILE (during the job execution, to be run by the master assigned node, e.g.
 in the batch script)
- How to check where am I actually running? Yes, it may be different from the previous point
 - mpirun <mpirun options> hostname
 - change the source code calling MPI_Get_processor_name
- Am I really using all the requested cores?
 - See more later





NUMA architectures

- Many current architectures used on HPC clusters are NUMA
 - Non-uniform memory access (NUMA) is a computer memory design used in multiprocessing, where the memory access time depends on the memory location relative to the processor. Under NUMA, a processor can access its own local memory faster than non-local memory (memory local to another processor or memory shared between processors)
- How to optimize my code and my run?
 - Numactl Linux command to control NUMA policy for processes or shared memory
 - OpenMPI thread assignment
 - MPI process assignement





OpenMP thread assignment

- Intel compiler: set KMP_AFFINITY to compact/scatter (other options available)
 - Specifying compact assigns the OpenMP* thread <n>+1 to a free thread context as close
 as possible to the thread context where the <n> OpenMP* thread was placed.
 - Specifying scatter distributes the threads as evenly as possible across the entire system.
 scatter is the opposite of compact
 - add the modifier "verbose" to have a list of the mapping (threads vs core). For example: export KMP_AFFINITY=verbose,compact
 - More info at: https://software.intel.com/en-us/node/522691
- GCC OpenMP uses the lower level variable GOMP_CPU_AFFINITY.
 - For example, GOMP_CPU_AFFINITY="0 3 1-2 4-15:2"





OpenMP thread assignment / 2

- OpenMP v4.5 provides OMP_PLACES and OMP_PROC_BIND (it's standard!)
 - OMP_PLACES: Specifies on which CPUs the threads should be placed. The thread
 placement can be either specified using an abstract name or by an explicit list of
 the places. Allowed abstract names: threads, cores and sockets
 - OMP_PROC_BIND: specifies whether theads may be moved between CPUs. If set to TRUE, OpenMP theads should not be moved; if set to FALSE they may be moved. Use a comma separated list with the values MASTER, CLOSE and SPREAD to specify the thread affinity policy for the corresponding nesting level.
 - More from OpenMP recent standards
- Thread assignment can be crucial for manycore architectures (Intel PHI)







OpenMP thread assignment / 4

Example (on a machine with 2 10-core sockets):
 export OMP_PLACES=cores;

export OMP_PROC_BIND= master	export OMP_PROC_BIND= close	export OMP_PROC_BIND= spread
• t0 -> c0	• t0 -> c0	• t0 -> c0
• t1 -> c0	• t1 -> c1	• t1 -> c10
• t2 -> c0	• t2 -> c2	• t2 -> c1
• t3 -> c0	• t3 -> c3	• t3 -> c11







- OpenMPI: tuning possible via options of mpirun
 - main reference: https://www.open-mpi.org/doc/v1.8/man1/mpirun.1.php
 - beware: mpirun automatically binds processes as of the start of the v1.8 series.
 Two binding patterns are used in the absence of any further directives:
 - Bind to core: when the number of processes is <= 2
 - Bind to socket: when the number of processes is > 2
 - more intuitive syntax still works but deprecated-npernode <n> / -npersocket <n>





- But --map-by function is more complete and can reproduce the above cases
 --map-by ppr:<n>:node / --map-by ppr:<n>:socket
- For process binding:
 - --bind-to <node/socket/core/...>
- To order processes (rank in round-robin fashion according to the specified object)
 --rank-by <foo>
- --report-bindings: useful option to see what is happening wrt bindings
 mpirun --map-by ppr:1:socket --rank-by socket --bind-to socket --report-bindings -np
 \$PROCS ./bin/\$PROGRAM





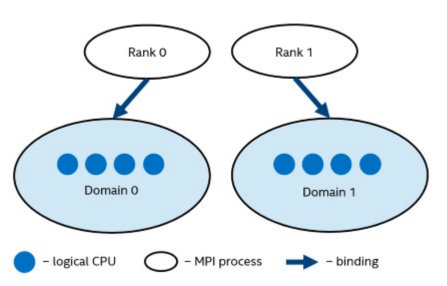
- "If your application uses threads, then you probably want to ensure that you are either not bound at all (by specifying --bind-to none), or bound to multiple cores using an appropriate binding level or specific number of processing elements per application process."
- mpirun --map-by ppr:1:socket:PE=10 --rank-by socket --bind-to socket --report-bindings
 <PROGRAM> (hybrid, 1 process per socket, 10 threads per process)
- mpirun --map-by ppr:2:socket:PE=5 --rank-by socket --bind-to socket --report-bindings
 <PROGRAM>
 - r0 -> c0-4
 - r1 -> c10-14
 - r2 -> c5 9
 - r3 -> c15-19





- IntelMPI: tuning possible via environment variables
 - main reference: https://software.intel.com/en-us/node/528816
- I_MPI_PIN: turn on/off process pinning. Default is on
- I_MPI_PIN_MODE: choose the pinning method
- I_MPI_PIN_PROCESSOR_LIST: one-to-one pinning, define a processor subset and the mapping rules for MPI processes within this subset.
- I_MPI_PIN_DOMAIN: one-to-many pinning, control process pinning for hybrid Intel MPI Library applications
- mpirun -print-rank-map to know the process binding or activate I_MPI_DEBUG=5







- I_MPI_PIN_DOMAIN defines the logical processor in each domain
- If present, it overrides I_MPI_PROCESSOR_LIST
- Different ways to define the domains are allowed
 - Multi-core shape: core, socket, node,...
 - Explicit shape:
 - omp (recommended for hybrid jobs), domain size is equal to OMP_NUM_THREADS
 - auto, domain size is n_procs/n_ranks
 - Explicit domain mask
 - e.g. [0xfc00, 0x3ff] (to understand convert to binary [1111111110000000000, 000000000111111111])
- E.g.
 - mpirun -np 1 -env I_MPI_PIN_DOMAIN omp -env
 I MPI PIN DEBUG 5 ./a.out





References

- MPI documents
 - http://www.mpi-forum.org/docs/
- OpenMP specifications
 - http://openmp.org/wp/openmp-specifications/
- SuperComputing 2015:
 - http://www.mcs.anl.gov/~thakur/sc15-mpi-tutorial/l
- Intel & GNU compilers

