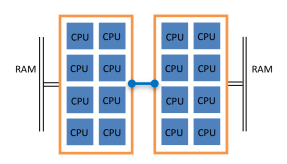
# **Hybrid Parallel Overview**

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Based on materials developed by CAC and TACC

# RAM Arrangement on Stampede

- Many nodes → <u>distributed memory</u>
  - each node has its own local memory
  - not directly addressable from other nodes
- Multiple sockets per node
  - each node has 2 sockets (chips)
- Multiple cores per socket
  - each socket (chip) has 8 cores
- Memory spans all 16 cores → <u>shared memory</u>
  - node's full local memory is addressable from any core in any socket
- Memory is attached to sockets
  - 8 cores sharing the socket have fastest access to attached memory
  - we are ignoring any attached MIC coprocessors for the moment...



# **Dealing with NUMA**

How do we deal with NUMA (Non-Uniform Memory Access)?

Standard models for parallel programs assume a uniform architecture –

- Threads for shared memory
  - parent process uses pthreads or OpenMP to fork multiple threads
  - threads share the same virtual address space
  - also known as SMP = Symmetric MultiProcessing
- Message passing for distributed memory
  - processes use MPI to pass messages (data) between each other
  - each process has its own virtual address space

If we attempt to combine both types of models -

- Hybrid programming
  - try to exploit the whole shared/distributed memory hierarchy

## Why Hybrid? Or Why Not?

### Why hybrid?

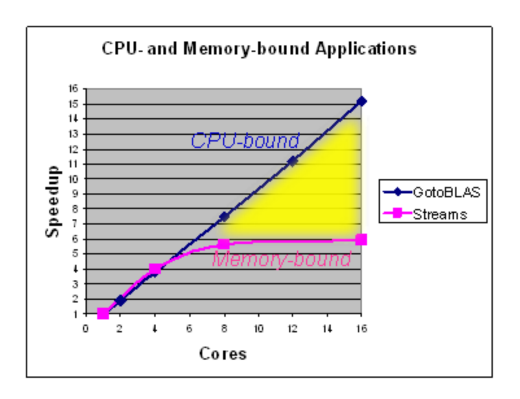
- Eliminates domain decomposition <u>at node level</u>
- Automatic memory coherency <u>at node level</u>
- Lower (memory) latency and data movement within node
- Can synchronize on memory instead of barrier
- Can be easier to incorporate coprocessors

### Why not hybrid?

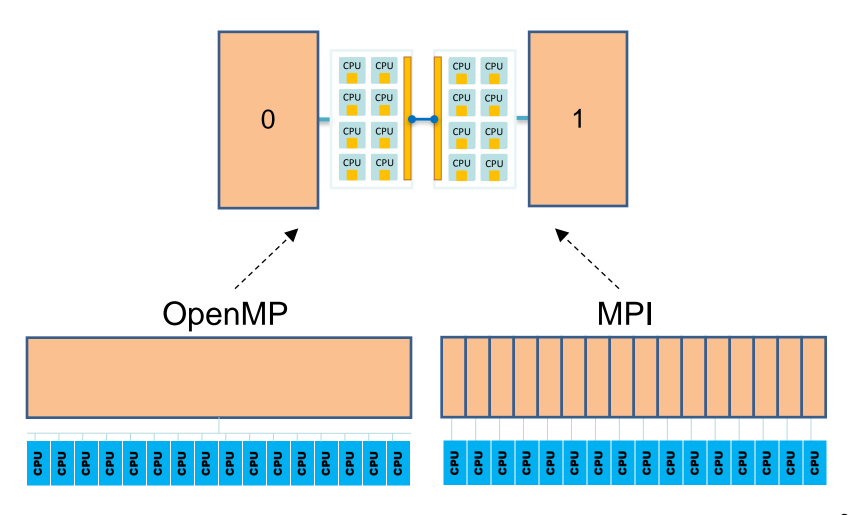
- An SMP algorithm created by aggregating MPI parallel components on a node (or on a socket) may actually run slower
- Possible waste of effort

## **Motivation for Hybrid**

- Balance the computational load
- Reduce memory traffic, especially for memory-bound applications



## Two Views of a Node



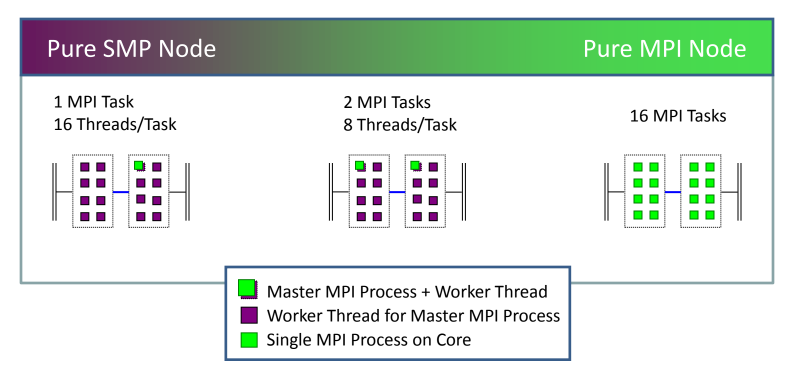
# Two Views = Two Ways to Write Parallel Programs

- OpenMP (or pthreads) only
  - launch one process per node
  - have each process fork one thread (or maybe more) per core
  - share data using shared memory
  - can't share data with a different process (except maybe via file I/O)
- MPI only
  - launch one process per core, on one node or on many
  - pass messages among processes without concern for location
  - (maybe create different communicators intra-node vs. inter-node)
  - ignore the potential for any memory to be shared
- With hybrid OpenMP/MPI programming, we want each MPI process to launch multiple OpenMP threads that can share local memory

# Some Possible MPI + Thread Configurations

- Treat each node as an SMP
  - launch a single MPI process per node
  - create parallel threads sharing full-node memory
  - typically want 16 threads/node on Stampede, e.g.
- Treat each socket as an SMP
  - launch one MPI process on each socket
  - create parallel threads sharing same-socket memory
  - typically want 8 threads/socket on Stampede, e.g.
- No SMP, ignore shared memory (all MPI)
  - assign an MPI process to each core
  - in a master/worker paradigm, one process per node may be master
  - not really hybrid, may at least make a distinction between nodes

## **Creating Hybrid Configurations**



To achieve configurations like these, we must be able to:

- Assign to each process/thread an affinity for some set of cores
- Make sure the allocation of memory is appropriately matched

## **NUMA Operations**

Where do processes, threads, and memory allocations get assigned?

- If memory were completely uniform, there would be no need to worry about questions like, "where do processes go?"
- Only for NUMA is the placement of processes/threads and allocated memory (NUMA control) of any importance

### The default NUMA control is set through policy

- The policy is applied whenever a process is executed, or a thread is forked, or memory is allocated
- These are all events that are directed from within the kernel

NUMA control is managed by the kernel.

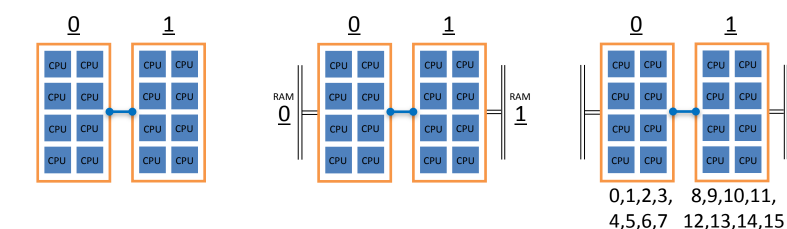
NUMA control can be changed with numactl.

# **Process Affinity and Memory Policy**

- One would like to set the affinity of a process for a certain socket or core, and the allocation of data in memory relative to a socket or core
- Individual users can alter kernel policies
   (setting Process Affinity and Memory Policy == PAMPer)
  - users can PAMPer their own processes
  - root can PAMPer any process
  - careful, libraries may PAMPer, too!
- Means by which Process Affinity and Memory Policy can be changed:
  - 1. dynamically on a running process (knowing process id)
  - 2. at start of process execution (with wrapper command)
  - 3. within program through F90/C API

## Using numactl, at the Process Level

### numactl <option socket(s)/core(s)> ./a.out



For a Process: Socket Control	For a Process's Memory: Socket Control	For a Process: Core Control
socket assignment	memory allocation -l -ipreferred -m (local, interleaved, preferred, mandatory)	core assignment

## **Quick Guide to numactl**

Socket Affinity	-N	{0,1}	Execute process on cores of this (these) socket(s) only.
Memory Policy	-1	no argument	Allocate on current socket; fallback to any other if full.
Memory Policy	-i	{0,1}	Allocate round robin (interleave) on these sockets. No fallback.
Memory Policy	preferred=	{0,1} select one	Allocate on this socket; fallback to any other if full.
Memory Policy	-m	{0,1}	Allocate only on this (these) socket(s). No fallback.
Core Affinity	-C	{0,1,2,3,4,5,6,7,8,9,10,11,12,13,14,15}	Execute process on this (these) core(s) only.

### **SMP Nodes**

#### Hybrid batch script for 16 threads/node

- Specify total MPI tasks to be started by batch
- Specify total nodes equal to tasks
- Set number of threads for each process
- PAMPering at job level
  - controls behavior (e.g., process-core affinity) for ALL processes
  - no simple/standard way to control thread-core affinity with numactl

job script (Bourne shell)	job script (C shell)	
#SBATCH -n 2 -N 2	#SBATCH -n 2 -N 2	
	• • •	
export OMP_NUM_THREADS=16	setenv OMP_NUM_THREADS 16	
•••	• • •	
ibrun numactl -i all ./a.out	ibrun numactl -i all ./a.out	

## **SMP Sockets**

### Hybrid batch script for 2 tasks/node, 8 threads/task

- Specify total MPI tasks to be started by batch
- Specify total nodes equal to tasks/2 (so 2 tasks/node)
- Set number of threads for each process
- PAMPering at process level, must create script to manage affinity
  - tacc\_affinity script pins tasks to sockets, ensures local memory allocation
  - use it as a numactl starting point if it's not quite right for your application

job script (Bourne shell)	job script (C shell)
#SBATCH -n 4 -N 2	#SBATCH -n 4 -N 2
•••	• • •
export OMP_NUM_THREADS=8	setenv OMP_NUM_THREADS 8
•••	• • •
ibrun tacc_affinity ./a.out	ibrun tacc_affinity ./a.out

## **Basic Hybrid Program Template**

Start with MPI initialization

(Serial regions are executed by the master thread of the MPI process)

Create OMP parallel regions within each MPI process

- MPI calls may be allowed here too
- MPI rank is same for all threads

Call MPI in single-threaded regions

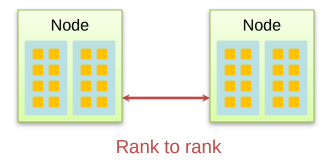
Finalize MPI

```
MPI Init
MPI Call
   OMP parallel
   MPI Call
   end parallel
MPI Call
MPI Finalize
```

## Types of MPI Calls Among Threads

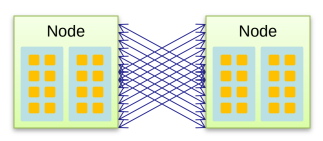
#### Single-threaded messaging

- Call MPI from a serial region
- Call MPI from a single thread within a parallel region



#### Multi-threaded messaging

- Call MPI from multiple threads within a parallel region
- Requires an implementation of MPI that is thread-safe



rank-thread ID to rank-thread ID

# Multiple Threads Calling MPI

- Thread ID as well as rank can be used in communication
- Technique is illustrated in multi-thread "ping" (send/receive) example

# **Example: Multiple Threads Calling MPI**

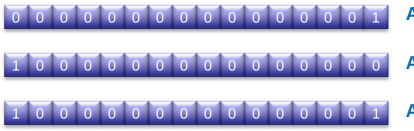
```
call mpi init thread (MPI THREAD MULTIPLE, iprovided, ierr)
call mpi_comm_rank(MPI_COMM_WORLD, irank, ierr)
call mpi_comm_size(MPI_COMM_WORLD, nranks, ierr)
if (iprovided >= MPI_THREAD_MULTIPLE) then ! All threads can call MPI
!$OMP parallel private(j, ithread, nthreads)
 nthreads=OMP GET NUM THREADS()
                                                Communicate between ranks.
 ithread =OMP_GET_THREAD_NUM()
 call pwork(ithread, irank, nthreads, nranks...)
                                                Threads use tags to differentiate.
 if(irank == 0) then
   call mpi_send(ithread,1,MPI_INTEGER, 1, ithread, MPI_COMM_WORLD, ierr)
 else
   call mpi_recv( j,1,MPI_INTEGER, 0, | ithread, MPI_COMM_WORLD, istat, ierr)
   print*, "Yep, this is ",irank," thread ", ithread," I received from ", j
 endif
!$OMP end parallel
endif
```

## NUMA Control in Code, at the Thread Level

- Within a code, Scheduling Affinity and Memory Policy (SCAMPer?)
  can be examined and changed using libnuma routines:
  - sched\_getaffinity, sched\_setaffinity
  - get\_mempolicy, set\_mempolicy

see man pages

- This is the only way to set affinities and policies that differ per thread
- To make scheduling assignments, set bits in a mask:



**Assignment to Core 0** 

**Assignment to Core 15** 

**Assignment to Core 0 or 15** 

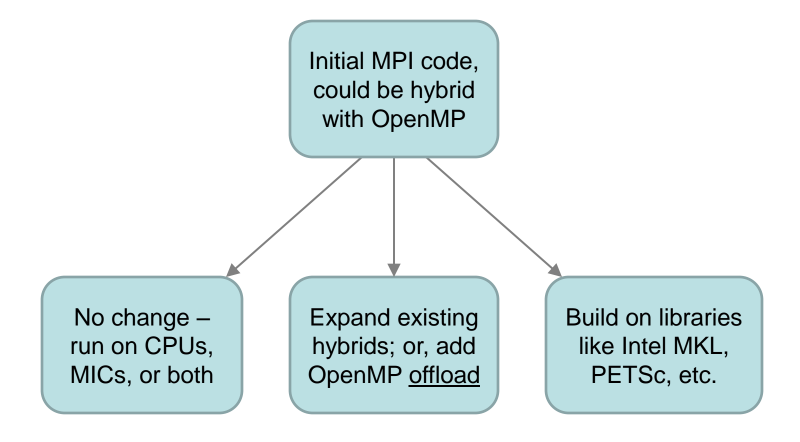
## **Code Example for Scheduling Affinity**

```
#include <spawn.h>
                       //C API parameters and prototypes
. . .
                       //Set core number
int icore=3;
                 //Allocate mask
cpu set t cpu mask;
CPU ZERO( &cpu mask); //Set mask to zero
CPU SET (icore, &cpu mask); //Set mask with core #
sizeof(cpu mask),
                   &cpu mask);
```

# Programming for MIC: Hybrid and Heterogeneous

- Each Stampede node currently has 2 processors + 1 MIC card
- MIC = Many Integrated Cores = a "coprocessor" on a PCIe card that features >60 cores; released as Xeon Phi™
  - Represents Intel's response to GPGPU, especially NVIDIA's CUDA
  - Answers the question: if 8 modern Xeon cores fit on a die, how many early Pentiums would fit?
- MIC answers CUDA's API problem: just compile like any normal code
  - Instruction set is x86 with support for 64-bit addressing
  - Recent x86 extensions may not be available
  - Developers use familiar Intel compilers, libraries, and tools
- However, MIC adds yet another level of programming complexity
  - Stampede is a multi-core machine where not all the cores are the same

# **MIC Strategies for HPC Codes**



# **OpenMP Offload Constructs: Base Program**

```
#include <omp.h>
#define N 10000
void foo(double *, double *, int );
int main(){
  int i; double a[N], b[N], c[N];
  for(i=0;i<N;i++){ a[i]=i; b[i]=N-1-i;}
  foo(a,b,c,N);
void foo(double *a, double *b, double *c, int n){
  int i;
  for(i=0;i<n;i++) { c[i]=a[i]*2.0e0 + b[i]; } }
```

- Objective: offload foo to a device
- Use OpenMP to do the offload

# **OpenMP Offload Constructs: Requirements**

```
#include <omp.h>
#define N 10000
#pragma <offload_function_spec>
void foo(double *, double *, int );
int main(){
  int i; double a[N], b[N], c[N];
  for(i=0;i<N;i++){ a[i]=i; b[i]=N-1-i;}
  #pragma <offload_this>
  foo(a,b,c,N);
#pragma <offload function spec>
void foo(double *a, double *b, double *c, int n){
  int i;
  #pragma omp parallel for
  for(i=0;i<n;i++) { c[i]=a[i]*2.0e0 + b[i]; } }
```

- Direct (Intel) compiler to offload function or block
- "Decorate" function and prototype
- Ideally, familiar-looking OpenMP directives work on device

# **Pros and Cons of MIC Programming Models**

- Offload engine: MIC serves as coprocessor for the host
  - Pros: distinct hardware gets distinct role; programmable via simple calls to a library such as MKL, or via directives (we'll go into depth on this)
  - Cons: PCle is the only path for most work; must try to retain data on card
- "Symmetric" #1: Everything is just an MPI core
  - Pros: MPI works for all cores (though 1 MIC core < 1 server core)</li>
  - Cons: memory may be insufficient to support a μOS plus lots of data;
     fails to take good advantage of shared memory; PCIe is a bottleneck
- "Symmetric" #2: MIC and host are just different SMPs
  - Pros: MPI/OpenMP works for both host and MIC; more efficient use of limited PCIe bandwidth and limited MIC memory
  - Cons: hybrid programming is already tough on homogeneous SMPs, but
     OpenMP-based hybrids may be the best path for scaling to 60 cores/MIC

## Quick Guide to KMP\_AFFINITY

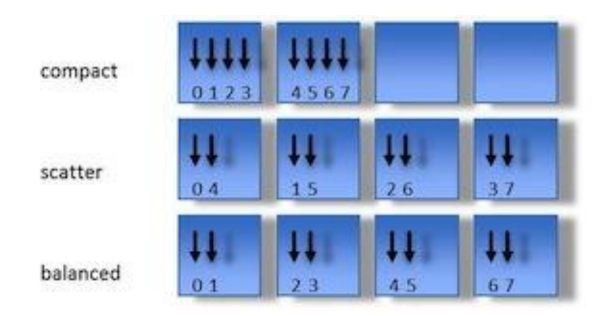
- Set this environment variable to influence thread affinity generally
- Useful for CPU and/or MIC models based on OpenMP (SMP, offload)
   export KMP AFFINITY=<type> (for SMP)

```
export MIC KMP AFFINITY=<type> (for offload)
```

Туре	Effect
compact	Pack threads close to each other.
explicit	Use the proclist modifier to pin threads.
none	Does not pin threads.
scatter	Round-robin threads to cores.
balanced (Phi only)	Use scatter, but keep OMP thread ids consecutive.

## KMP\_AFFINITY Types and Thread Placement

- Imagine a system with 4 cores and 4 hardware threads /core
- Placement of 8 threads is illustrated for the 3 types
- Compact type does not fully utilize all cores; not recommended



## **Roadmap: What Comes Next?**

- Expect many of the upcoming large systems to be accelerated
- MPI + OpenMP will be the main HPC programming model
  - If you are not using Intel TBBs or Cilk
  - If you are not spending all your time in libraries (MKL, etc.)
- Many HPC applications are pure-MPI codes
  - Start thinking about upgrading to a hybrid scheme
  - Adding OpenMP is a larger effort than adding the extra MIC directives
- Special MIC/OpenMP considerations
  - Many more threads will be needed:
     60+ cores on production Xeon Phi™ → 60+/120+/240+ threads
  - Good OpenMP scaling (and vectorization) are much more important

## Conclusions

- On heterogeneous NUMA systems like Stampede, placement and binding of processes and their associated memory are important performance considerations.
- Process Affinity and Memory Policy have a significant effect on pure MPI, pure OpenMP, and Hybrid codes—both on host and on MIC.
- Simple numactl commands and APIs allow users to control affinity of processes and threads and memory assignments.
- Future prospects for hybrid programming:
  - Core counts will increase on both processors and coprocessors.
  - Even more effort will be focused on process scheduling and data locality.
  - Expect to see more multi-threaded libraries; be alert for their potential interaction with your own multithreading strategy.

### References

 Yun (Helen) He and Chris Ding, Lawrence Berkeley National Laboratory, June 24, 2004: <u>Hybrid OpenMP and MPI Programming</u> and <u>Tuning (NUG2004)</u>.

www.nersc.gov/nusers/services/training/classes/NUG/Jun04/NUG2004\_yhe\_hybrid.ppt

- Texas Advanced Computing Center: <u>Stampede User Guide</u>, see
   numa section. www.tacc.utexas.edu/services/userguides/stampede
- Message Passing Interface Forum: MPI-2: MPI and Threads (specific section of the MPI-2 report).

http://www.mcs.anl.gov/research/projects/mpi/mpi-standard/mpi-report-2.0/node162.htm

 Intel Corp.: <u>Thread Affinity Interface (Linux and Windows)</u>, from the Intel Fortran Compiler User and Reference Guides.

http://www.intel.com/software/products/compilers/docs/fmac/doc\_files/source/extfile/optaps\_for/common/optaps\_openmp\_thread\_affinity.htm

# **Extra Slides: MPI-2 and Multithreading**

## **MPI-2** and Thread Safety

- Consider thread safety when calling MPI from threads
- Use MPI\_Init\_thread to select/determine the level of thread support
  - Supported in MPI-2, substitute for the usual MPI\_Init
- Thread safety is identified/controlled by MPI's provided types
  - Single means no multi-threading
  - Funneled means only the master thread can call MPI
  - Serialized means multiple threads can call MPI, but only 1 call can be in progress at a time
  - Multiple means MPI is thread safe
- Monotonic values are assigned to parameters

## MPI-2's MPI\_Init\_thread

#### Syntax:

- Input: rqd, or "required" (integer)
  - Indicates the desired level of thread support
- Output: pvd, or "provided" (integer)
  - Indicates the available level of thread support
- If thread level rqd is supported, the call returns pvd = rqd
- Otherwise, pvd returns the highest provided level of support

# **MPI-2 Thread Support Levels**

Support Levels	Description	
MPI_THREAD_SINGLE	Only one thread will execute.	
MPI_THREAD_FUNNELED	Process may be multi-threaded, but only the main thread will make MPI calls (calls are "funneled" to main thread). *Default*	
MPI_THREAD_SERIALIZE	Process may be multi-threaded, and any thread can make MPI calls, but threads cannot execute MPI calls concurrently; they must take turns (calls are "serialized").	
MPI_THREAD_MULTIPLE	Multiple threads may call MPI, with no restriction.	

## **Example: Single-Threaded MPI Calls**

```
Fortran
                                                 C
include 'mpif.h'
                                  #include <mpi.h>
program hybsimp
                                  int main(int argc,
                                    char **arqv) {
                                  int rank, size, ie, i;
                                  ie= MPI Init(&argc,&argv[]);
call MPI Init(ie)
call MPI Comm rank(...irk,ie)
                                  ie= MPI Comm rank(...&rank);
                                  ie= MPI Comm size(...&size);
call MPI Comm size(...isz,ie)
                                  //Setup shared mem, comp/comm
!Setup shared mem, comp/comm
!$OMP parallel do
                                  #pragma omp parallel for
  do i=1,n
                                    for(i=0; i<n; i++) {
    <work>
                                      <work>
  enddo
                                  // compute & communicate
!Compute & communicate
call MPI Finalize(ierr)
                                  ie= MPI Finalize();
end
```

#### Funneled MPI Calls via Master

- Must have support for MPI\_THREAD\_FUNNELED or higher
- Best to use OMP\_BARRIER
  - there is no implicit barrier in the master workshare construct, OMP\_MASTER
  - in the example, the master thread will execute a single MPI call within the OMP\_MASTER construct
  - all other threads will be sleeping

## **Example: Funneled MPI Calls via Master**

```
Fortran
include 'mpif.h'
                                  #include <mpi.h>
                                  int main(int argc,
program hybmas
                                     char **argv) {
                                  int rank, size, ie, i;
!$OMP parallel
                                  #pragma omp parallel
  !$OMP barrier
                                     #pragma omp barrier
  !$OMP master
                                     #pragma omp master
  call MPI <Whatever>(...,ie)
                                       ie= MPI <Whatever>(...);
  !$OMP end master
  !$OMP barrier
                                     #pragma omp barrier
!$OMP end parallel
end
```

## Serialized MPI Calls and OpenMP

- Must have support for MPI\_THREAD\_SERIALIZED or higher
- Best to use OMP\_BARRIER only at beginning, since there is an implicit barrier in the SINGLE workshare construct, OMP\_SINGLE
  - Example is the simplest one: any thread (not necessarily master)
     will execute a single MPI call within the OMP\_SINGLE construct
  - All other threads will be sleeping

## **Example: Serialized MPI Calls and OpenMP**

```
Fortran
include 'mpif.h'
                                   #include <mpi.h>
                                   int main(int argc,
program hybsing
                                     char **arqv) {
                                   int rank, size, ie, i;
call MPI Init thread( &
                                   ie= MPI Init thread(
MPI THREAD SERIALIZED, ipvd, ie)
                                  MPI THREAD SERIALIZED, ipvd);
!$OMP parallel
                                   #pragma omp parallel
  !$OMP barrier
                                     #pragma omp barrier
  !$OMP single
                                     #pragma omp single
  call MPI <Whatever>(...,ie)
                                       ie= MPI <Whatever>(...);
  !$OMP end single
  !Don't need OMP barrier
                                     //Don't need omp barrier
!$OMP end parallel
end
```

## Overlapping Work & MPI Calls

- One core is capable of saturating the lanes of the PCIe network link...
  - Why use all cores to communicate?
  - Instead, communicate using just one or several cores
  - Can do work with the rest during communication
- Must have support for MPI\_THREAD\_FUNNELED or higher to do this
- Can be difficult to manage and load-balance!

## **Example: Overlapping Work & MPI Calls**

```
Fortran
include 'mpif.h'
                                   #include <mpi.h>
program hybsing
                                   int main(int argc,
                                     char **arqv) {
                                   int rank, size, ie, i;
!$OMP parallel
                                   #pragma omp parallel
  if (ithread .eq. 0) then
                                     if (thread == 0) {
  call MPI <Whatever>(...,ie)
                                       ie= MPI <Whatever>(...);
  else
   <work>
                                     if(thread != 0){
  endif
                                       <work>
!$OMP end parallel
end
```