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Hybrid Programming with OpenMP and MPI

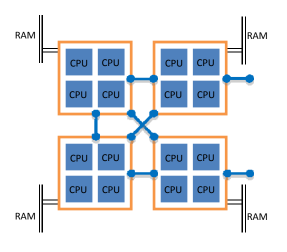
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Workshop: Introduction to Parallel Computing on Ranger, May 20, 2010 Based on materials developed by Kent Milfeld at TACC



RAM Arrangement on Ranger

- Distributed memory
 - each node has its own local memory
 - not directly addressable from other nodes
- Multichip nodes
 - each node has 4 chips (sockets)
- Multicore chips
 - each chip (socket) has 4 cores
- Memory spans entire nodes
 - node's full local memory is addressable from any core in any socket
- Memory is attached to sockets
 - 4 cores sharing the socket have fastest access to attached memory





Dealing with NUMA

How do we deal with NUMA (Non-Uniform Memory Access)? Two standard models for parallel programs -

- 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 at node
- Automatic memory coherency at node
- Lower (memory) latency and data movement within node
- Can synchronize on memory instead of barrier

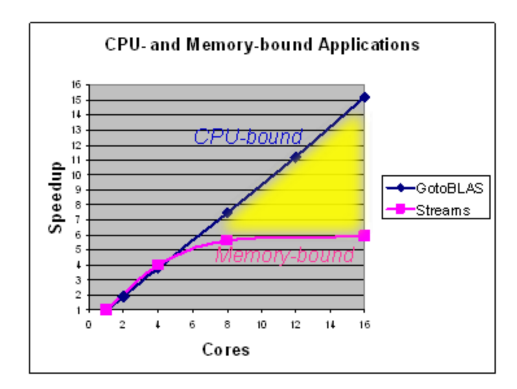
Why not hybrid?

- 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

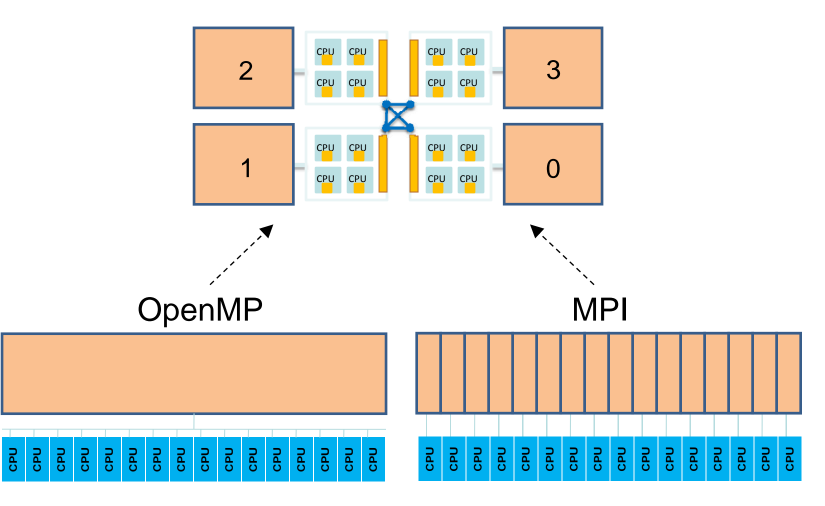
- Balance the computational load
- Reduce memory traffic, especially for memory-bound applications





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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

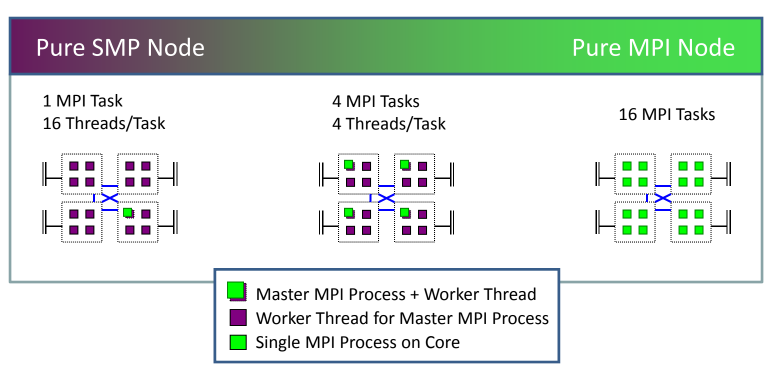


Some Possible MPI + Thread Configurations

- 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 Ranger, e.g.
- Treat each socket as an SMP
 - launch one MPI process on each socket
 - create parallel threads sharing same-socket memory
 - typically want 4 threads/socket on Ranger, 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.



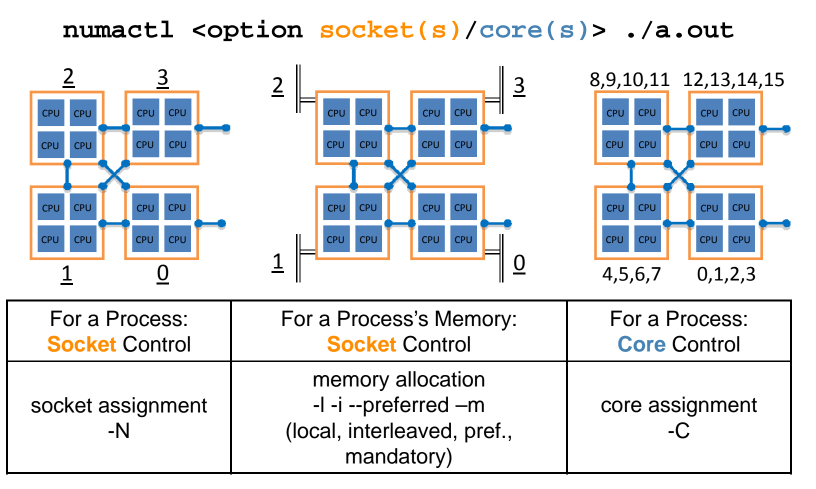
Process Affinity and Memory Policy

- 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:
 - dynamically on a running process (knowing process id)
 - at start of process execution (with wrapper command)
 - within program through F90/C API



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Using numactl





Quick Guide to numactl

Socket Affinity	-N	{0,1,2,3}	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	{0,1,2,3}	Allocate round robin (interleave) on these sockets. No fallback.
Memory Policy	preferred=	{0,1,2,3} select one	Allocate on this socket; fallback to any other if full.
Memory Policy	-m	{0,1,2,3}	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

- Make sure 1 process per node is created
- Specify total cores allocated by batch (nodes x 16)
- Set number of threads for each process
- PAMPering at job level
 - controls behavior for ALL tasks
 - no simple/standard way to control thread-core affinity

job script (Bourne shell)	job script (C shell)
•••	•••
#! -pe 1way 192	#! -pe 1way 192
•••	•••
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 4 tasks/node, 4 threads/task

Example script setup for a square (6x6 = 36) processor topology...

- Make sure 4 processes per node are created (one per socket)
- Specify total cores allocated by batch (nodes x 16)
- Specify actual cores used with MY_NSLOTS
- Set number of threads for each process
- PAMPering at process level, must create script to manage affinity

job script (Bourne shell)	job script (C shell)
• • •	•••
#! -pe 4way 48	#! -pe 4way 48
export MY_SLOTS=36	setenv MY_NSLOTS 36
export OMP_NUM_THREADS=4	setenv OMP_NUM_THREADS 4
ibrun numa.sh	ibrun numa.csh



Script for Socket Affinity

- Example script to extract rank for numactl options, a.out execution (TACC MPI systems always assign ranks sequentially on a node)
- No simple/standard way to control thread-core affinity

numa.sh	numa.csh
#!/bin/bash	#!/bin/csh
export MV2_USE_AFFINITY=0	setenv MV2_USE_AFFINITY 0
export MV2_ENABLE_AFFINITY=0	<pre>setenv MV2_ENABLE_AFFINITY 0</pre>
#TasksPerNode TPN=`echo \$PE sed 's/way//'` [! \$TPN] && echo TPN null! [! \$TPN] && exit 1	<pre>#TasksPerNode set TPN=`echo \$PE sed 's/way//'` if(! \${%TPN}) echo TPN null! if(! \${%TPN}) exit 0</pre>
sk=\$((\$PMI_RANK % \$TPN))	@ sk = \$PMI_RANK % \$TPN
numactl -N \$sk -m \$sk ./a.out	numactl -N \$sk -m \$sk ./a.out



Basic Hybrid Program Template

Start with MPI initialization

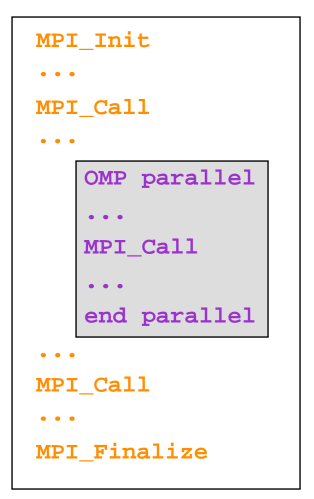
(Serial regions are executed by the master thread of the MPI process)

Create OMP parallel regions within each MPI process

- MPI calls may be allowed here too
- MPI rank is known to all threads

Call MPI in single-threaded regions

Finalize MPI

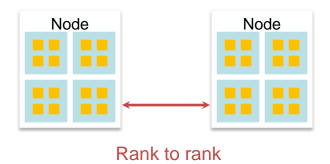




Types of MPI Calls Among Threads

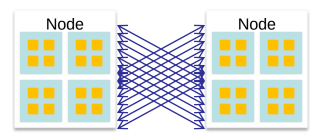
Single-threaded messaging

- 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



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_THREAD_SINGLE < MPI_THREAD_FUNNELED
 < MPI_THREAD_SERIALIZED < MPI_THREAD_MULTIPLE



MPI-2's MPI_Init_thread

Syntax:

call MPI_Init_thread(irqd, ipvd, ierr)
<pre>int MPI_Init_thread (int *argc, char ***argv,</pre>	<pre>int rqd, int *pvd)</pre>
<pre>int MPI::Init_thread(int& argc, char**& argv,</pre>	int rqd)

- 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 (MPI calls are "serialized").
MPI_THREAD_MULTIPLE	Multiple threads may call MPI, with no restriction.



Example: Single-Threaded MPI Calls

Fortran	С
include 'mpif.h'	<pre>#include <mpi.h></mpi.h></pre>
program hybsimp	int main(int argc,
	char **argv) {
	int rank, size, ie, i;
call MPI_Init(ie)	<pre>ie= MPI_Init(&argc,&argv[]);</pre>
<pre>call MPI_Comm_rank(irk,ie)</pre>	<pre>ie= MPI_Comm_rank(&rank);</pre>
<pre>call MPI_Comm_size(isz,ie)</pre>	<pre>ie= MPI_Comm_size(&size);</pre>
!Setup shared mem, comp/comm	//Setup shared mem, comp/comm
!\$OMP parallel do	<pre>#pragma omp parallel for</pre>
do i=1,n	<pre>for(i=0; i<n; i++){<="" pre=""></n;></pre>
<work></work>	<work></work>
enddo	}
!Compute & communicate	// compute & communicate
call MPI_Finalize(ierr)	<pre>ie= MPI_Finalize();</pre>
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	С
include 'mpif.h'	<pre>#include <mpi.h></mpi.h></pre>
program hybmas	int main(int argc,
	char **argv) {
	int rank, size, ie, i;
!\$OMP parallel	<pre>#pragma omp parallel {</pre>
!\$OMP barrier	#pragma omp barrier
!\$OMP master	<pre>#pragma omp master {</pre>
<pre>call MPI_<whatever>(,ie)</whatever></pre>	<pre>ie= MPI_<whatever>();</whatever></pre>
!\$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	С
include 'mpif.h'	<pre>#include <mpi.h></mpi.h></pre>
program hybsing	<pre>int main(int argc, char **argv) {</pre>
	int rank, size, ie, i;
call MPI_Init_thread(&	ie= MPI_Init_thread(
<pre>MPI_THREAD_SERIALIZED, ipvd, ie)</pre>	<pre>MPI_THREAD_SERIALIZED, ipvd);</pre>
!\$OMP parallel	<pre>#pragma omp parallel {</pre>
!\$OMP barrier	#pragma omp barrier
!\$OMP single	#pragma omp master {
<pre>call MPI_<whatever>(,ie)</whatever></pre>	<pre>ie= MPI_<whatever>();</whatever></pre>
!\$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	С
include 'mpif.h'	<pre>#include <mpi.h></mpi.h></pre>
program hybsing	<pre>int main(int argc, char **argv) {</pre>
	int rank, size, ie, i;
!\$OMP parallel	<pre>#pragma omp parallel {</pre>
if (ithread .eq. 0) then	if $(thread == 0)$ {
<pre>call MPI_<whatever>(,ie)</whatever></pre>	<pre>ie= MPI_<whatever>();</whatever></pre>
else	}
<work></work>	if(thread != 0){
endif	<work></work>
	}
!\$OMP end parallel	}
end	}



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



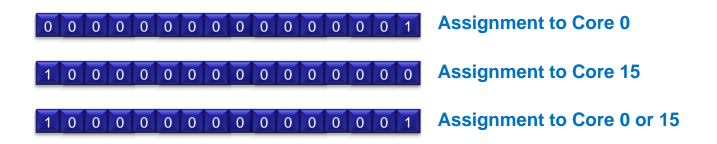
Example: Multiple Threads Calling MPI

```
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)
!$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
                                            ithread, MPI_COMM_WORLD, ierr)
  call mpi send(ithread,1,MPI INTEGER, 1,
 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
SOMP END PARALLEL
end
```



NUMA Control in Code

- Within a code, **Scheduling Affinity** and **Memory Policy** can be examined and changed through:
 - sched_getaffinity, sched_setaffinity
 - get_memorypolicy, set_memorypolicy
- To make scheduling assignments, set bits in a mask:





Code Example for Scheduling Affinity

```
. . .
#include <spawn.h>
                           //C API parameters and prototypes
. . .
int icore=3;
                             //Set core number
cpu set t cpu mask;
                             //Allocate mask
. . .
CPU ZERO( &cpu mask); //Set mask to zero
CPU SET(icore,&cpu mask); //Set mask with core #
err = sched_setaffinity( (pid_t)0 , //Set the affinity
                        sizeof(cpu mask),
                        &cpu mask);
```



Conclusions and Future Prospects

- On NUMA systems like Ranger, 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.
- Simple numactl commands and APIs allow users to control process and memory assignments.
- Future prospects for hybrid programming:
 - 8-core and 16-core socket systems are on the way, so 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

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