Performance Considerations: Hardware Architecture

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Parallel Optimization and Scientific Visualization for Ranger
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based on material developed by Kent Milfeld, TACC

www.cac.cornell.edu
HW challenges on Ranger?

- Distributed memory - each node has its own - not readily accessible from other nodes
- Multichip nodes - each node has four chips
- Multicore chips - each chip has four cores
- Memory is associated with chips - more accessible from cores on same chip
How do we deal with NUMA?

- NUMA = Non-Uniform Memory Access
- Distributed memory: MPI
- Shared memory: Threads
  - pthreads
  - OpenMP
- Both: Hybrid programming
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?

• Only profitable if on-node aggregation of MPI parallel components is faster as a single SMP algorithm (or a single SMP algorithm on each socket).
Hybrid - Motivation

• Load Balancing
• Reduce Memory Traffic
Node Views

OpenMP

MPI

Process-Affinity
Memory-Allocation

2

3

1

0
NUMA Operations

- Where do threads/processes and memory allocations go?
- If Memory were completely uniform there would be no need to worry about these two concerns. Only for NUMA (non-uniform memory access) is (re)placement of processes and allocated memory (NUMA Control) of importance.
- Default Control: Decided by policy when process exec’d or thread forked, and when memory allocated. Directed from within Kernel.

NUMA Control is managed by the Kernel.
NUMA Control can be changed with numactl.
NUMA Operations

• Ways Process Affinity and Memory Policy can be changed:
  – Dynamically on a running process (knowing process id)
  – At process execution (with wrapper command)
  – Within program through F90/C API

• Users can alter Kernel Policies
  (setting Process Affinity and Memory Policy == PAMPPer)
  – Users can PAMPPer their own processes.
  – Root can PAMPPer any process.
  – Careful, libraries may PAMPPer, too!
NUMA Operations

- Process Affinity and Memory Policy can be controlled at **socket** and **core** level with **numactl**.

**Command:**
```
numactl < options socket/core > ./a.out
```

**Process: Socket References**
- process assignment: `-N`

**Memory: Socket References**
- memory allocation: `–l` `–i` `--preferred` `–m` (local, interleaved, pref., mandatory)

**Process: Core References**
- core assignment: `–C`

**Process References:**

- **Socket References**
  - 0, 1, 2, 3
  - 4, 5, 6, 7
  - 8, 9, 10, 11
  - 12, 13, 14, 15

- **Core References**
  - 0
  - 1
  - 2
  - 3
# NUMA Quick Guide

<table>
<thead>
<tr>
<th>cmd</th>
<th>option</th>
<th>arguments</th>
<th>description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Socket Affinity</td>
<td>numactl</td>
<td>-N</td>
<td>Execute process on cores of this (these) socket(s) only.</td>
</tr>
<tr>
<td>Memory Policy</td>
<td>numactl</td>
<td>-l</td>
<td>Allocate on current socket; fallback to any other if full.</td>
</tr>
<tr>
<td>Memory Policy</td>
<td>numactl</td>
<td>-i</td>
<td>Allocate round robin (interleave) on these sockets. No fallback.</td>
</tr>
<tr>
<td>Memory Policy</td>
<td>numactl</td>
<td>--preferred=</td>
<td>Allocate on this socket; fallback to any other if full.</td>
</tr>
<tr>
<td>Memory Policy</td>
<td>numactl</td>
<td>-m</td>
<td>Allocate only on this (these) socket(s). No fallback</td>
</tr>
<tr>
<td>Core Affinity</td>
<td>numactl</td>
<td>-C</td>
<td>Execute process on this (these) core(s) only.</td>
</tr>
</tbody>
</table>
Modes of MPI/Thread Operation

- **SMP Nodes**
  - Single MPI task launched per node
  - Parallel Threads share all node memory, e.g. 16 threads/node on Ranger.

- **SMP Sockets**
  - Single MPI task launched on each socket
  - Parallel Thread set shares socket memory, e.g. 4 threads/socket on Ranger

- **No Shared Memory (all MPI)**
  - Each core on a node is assigned an MPI task.
  - (not really hybrid, but in master/worker paradigm master could use threads)
Modes of MPI/Thread Operation

Pure MPI Node

- 16 MPI Tasks
- 4 MPI Tasks
- 1 MPI Task

Pure SMP Node

- 4 Threads/Task
- 16 Threads/Task

Legend:
- Green: MPI Task on Core
- Purple: Master Thread of MPI Task
- Purple: Worker Thread of MPI Task
SMP Nodes

Hybrid Batch Script  16 threads/node

- Make sure 1 task is created on each node
- Set total number of cores (nodes x 16)
- Set number of threads for each node
- PAMPering at job level
- Controls behavior for ALL tasks
- No simple/standard way to control thread-core affinity

```
job script (Bourne shell)
...
#! -pe 1way 192
...
export OMP_NUM_THREADS=16
ibrun numactl –i all ./a.out
```

```
job script (C shell)
...
#! -pe 1way 192
...
setenv OMP_NUM_THREADS 16
ibrun numactl –i all ./a.out
```
SMP Sockets

Hybrid Batch Script  4 tasks/node, 4 threads/task

- Example script setup for a square (6x6 = 36) processor topology.
- Create a task for each socket (4 tasks per node).
- Set total number of cores allocated by batch (nodes x 16 cores/node).
- Set actual number of cores used with MY_NSLOTS.
- Set number of threads for each task
- PAMPering at task level
  - Create script to extract rank for numactl options, and a.out execution (TACC MPI systems always assign sequential ranks on a node).
  - No simple/standard way to control thread-core affinity

<table>
<thead>
<tr>
<th>job script (Bourne shell)</th>
<th>job script (C shell)</th>
</tr>
</thead>
<tbody>
<tr>
<td>...</td>
<td>...</td>
</tr>
<tr>
<td>#! -pe 4way 144</td>
<td>#! -pe 4way 144</td>
</tr>
<tr>
<td>...</td>
<td>...</td>
</tr>
<tr>
<td>export MY_NSLOTS =36</td>
<td>setenv MY_NSLOTS 36</td>
</tr>
<tr>
<td>export OMP_NUM_THREADS=4</td>
<td>setenv OMP_NUM_THREADS 4</td>
</tr>
<tr>
<td>ibrun numa.sh</td>
<td>ibrun numa.csh</td>
</tr>
</tbody>
</table>
SMP Sockets

Hybrid Batch Script  4 tasks/node, 4 threads/task

<table>
<thead>
<tr>
<th>numa.sh</th>
<th>numa.csh</th>
</tr>
</thead>
<tbody>
<tr>
<td>#!/bin/bash</td>
<td>#!/bin/tcsh</td>
</tr>
<tr>
<td>export MV2_USE_AFFINITY=0</td>
<td>setenv MV2_USE_AFFINITY 0</td>
</tr>
<tr>
<td>export MV2_ENABLE_AFFINITY=0</td>
<td>setenv MV2_ENABLE_AFFINITY 0</td>
</tr>
<tr>
<td>#TasksPerNode</td>
<td>#TasksPerNode</td>
</tr>
<tr>
<td>TPN = `echo $PE</td>
<td>sed 's/way//`</td>
</tr>
<tr>
<td>[ ! $TPN ] &amp;&amp; echo TPN NOT defined!</td>
<td>if(! ${%TPN}) echo TPN NOT defined!</td>
</tr>
<tr>
<td>[ ! $TPN ] &amp;&amp; exit 1</td>
<td>if(! ${%TPN}) exit 0</td>
</tr>
<tr>
<td>socket = $(( $PMI_RANK % $TPN ))</td>
<td>@ socket = $PMI_RANK % $TPN</td>
</tr>
<tr>
<td>exec numactl -N $socket -m $socket ./a.out</td>
<td>exec numactl -N $socket -m $socket ./a.out</td>
</tr>
</tbody>
</table>
Hybrid – Program Model

- Start with MPI initialization
- Create OMP parallel regions within MPI task (process).
  - Serial regions are the master thread or MPI task.
  - MPI rank is known to all threads
- Call MPI library in serial and parallel regions.
- Finalize MPI
MPI with OpenMP -- Messaging

**Single-threaded messaging**

MPI from serial region or a single thread within parallel region

**Multi-threaded messaging**

MPI from multiple threads within parallel region
Requires thread-safe implementation
Threads calling MPI

- Use MPI_Init_thread to select/determine MPI’s thread level of support (in lieu of MPI_Init). MPI_Init_thread is supported in MPI2.
- Thread safety is controlled by “provided” types: single, funneled, serialized and multiple:
  - Single means there is no multi-threading.
  - Funneled means only the master thread calls MPI.
  - Serialized means multiple threads can call MPI, but only 1 call can be in progress at a time (serialized).
  - Multiple means MPI is thread safe.
- Monotonic values are assigned to Parameters:
  - MPI_THREAD_SINGLE < MPI_THREAD_FUNNELED < MPI_THREAD_SERIALIZED < MPI_THREAD_MULTIPLE
MPI2 MPI_Init_thread

Syntax:

```c
void MPI_Init_thread(int irequired, int iprovided, int ierr);
int MPI_Init_thread(int *argc, char ***argv, int required, int *provided);
int MPI::Init_thread(int &argc, char **&argv, int required);
```

<table>
<thead>
<tr>
<th>Support Levels</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>MPI_THREAD_SINGLE</td>
<td>Only one thread will execute.</td>
</tr>
<tr>
<td>MPI_THREAD_FUNNELED</td>
<td>Process may be multi-threaded, but only main thread will make MPI calls (calls are &quot;funneled&quot; to main thread). “Default”</td>
</tr>
<tr>
<td>MPI_THREAD_SERIALIZE</td>
<td>Process may be multi-threaded, any thread can make MPI calls, but threads cannot execute MPI calls concurrently (MPI calls are &quot;serialized&quot;).</td>
</tr>
<tr>
<td>MPI_THREAD_MULTIPLE</td>
<td>Multiple threads may call MPI, no restrictions.</td>
</tr>
</tbody>
</table>

If supported, the call will return provided = required. Otherwise, the highest level of support will be provided.
Hybrid Coding

Fortran

```fortran
include ‘mpif.h’
program hybsimp

call MPI_Init(ierr)
call MPI_Comm_rank (...,irank,ierr)
call MPI_Comm_size (...,isize,ierr)
! Setup shared mem, comp. & Comm

 !$OMP parallel do
do i=1,n
<work>
  enddo
! compute & communicate

call MPI_Finalize(ierr)
end
```

C

```c
#include <mpi.h>
int main(int argc, char **argv){
  int rank, size, ierr, i;

  ierr= MPI_Init(&argc,&argv[]);
ierr= MPI_Comm_rank (...,&rank);
ierr= MPI_Comm_size (...,&size);
//Setup shared mem, compute & Comm

  #pragma omp parallel for
  for(i=0; i<n; i++){
    <work>
  }
// compute & communicate

  ierr= MPI_Finalize();
}
```
MPI Call through Master

- **MPI_THREAD_FUNNELED**
- **Use OMP_BARRIER** since there is no implicit barrier in master workshare construct (OMP_MASTER).
- All other threads will be sleeping.
include ‘mpif.h’
program hybmas

$OMP parallel

 $OMP barrier
 $OMP master

call MPI_<Whatever>(...,ierr)
 $OMP end master

 $OMP barrier

$OMP end parallel
end
MPI Call within Single

- **MPI_THREAD_SERIALIZED**

- **Only OMP_BARRIER** is at beginning, since there is an implicit barrier in SINGLE workshare construct (OMP_SINGLE).

- All other threads will be sleeping.

- (The simplest case is for any thread to execute a single mpi call, e.g. with the “single” omp construct. See next slide.)
Serialize through Single

Fortran

```fortran
include ‘mpif.h’
program hybsing

call mpi_init_thread(MPI_THREAD_SERIALIZED, iprovided,ierr)

 !$OMP parallel

 !$OMP barrier
 !$OMP single

 call MPI_<whatever>(...,ierr)

 !$OMP end single

 !$OMP barrier

 !$OMP end parallel
end
```

C

```c
#include <mpi.h>
int main(int argc, char **argv){
    int rank, size, ierr, i;
    mpi_init_thread(MPI_THREAD_SERIALIZED, iprovided)

    #pragma omp parallel

    {
        #pragma omp barrier
        #pragma omp single

        ierr=MPI_<Whatever>(...)

        //pragma omp barrier

    }

    //pragma omp barrier

}
```
Overlapping Communication and Work

• One core can saturate the PCI-e \( \leftrightarrow \) network bus. Why use all to communicate?

• Communicate with one or several cores.

• Work with others during communication.

• Need at least \texttt{MPI\_THREAD\_FUNNELED} support.

• Can be difficult to manage and load balance!
## Overlapping Communication and Work

### Fortran

```
include 'mpi.h'
program hybover

!$OMP parallel
   if (ithread.eq.0) then
      call MPI_<whatever>(...,ierr)
   else
      <work>
   endif

!$OMP end parallel
end
```

### C

```
#include <mpi.h>
int main(int argc, char **argv){
   int rank, size, ierr, i;

   #pragma omp parallel
   {
      if (thread == 0){
         ierr=MPI_<Whatever>(...)
      }
      if(thread != 0){
         work
      }
   }
```
Thread-rank Communication

• Can use thread id and rank in communication

• Example illustrates technique in multi-thread “ping” (send/receive).
Thread-rank Communication

... 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(i, ithread, nthreads)
...

  nthreads = OMP_GET_NUM_THREADS()
  ithread = OMP_GET_THREAD_NUM()
call pwork(ithread, irank, nthreads, nranks...)
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, istatus, ierr)
  print*, "Yep, this is ", irank, " thread ", ithread, " I received from ", j
endif

!$OMP END PARALLEL
end
NUMA in Code

- **Scheduling Affinity** and **Memory Policy** can be changed within code through:
  - `sched_get/setaffinity`
  - `get/set_memorypolicy`

- **Scheduling**: Bits in a mask are set for assignments.
**NUMA in Code**

- **Scheduling Affinity**

```c
#include <spawn.h>
...
int icore=3;
cpu_set_t cpu_mask;
...
CPU_ZERO(&cpu_mask);
CPU_SET(icore,&cpu_mask);
err = sched_setaffinity((pid_t)0,
                        sizeof(cpu_mask),
                        &cpu_mask);
```

<table>
<thead>
<tr>
<th>C API params/protos</th>
</tr>
</thead>
<tbody>
<tr>
<td>Set core number</td>
</tr>
<tr>
<td>Allocate mask</td>
</tr>
<tr>
<td>Set mask to zero</td>
</tr>
<tr>
<td>Set mask with core #</td>
</tr>
<tr>
<td>Set the affinity</td>
</tr>
</tbody>
</table>
Conclusion

- Placement and binding of processes, and allocation location of memory are important performance considerations in pure MPI/OpenMP and Hybrid codes.

- Simple `numactl` commands and APIs allow users to control process and memory assignments.

- 8-core and 16-core socket systems are on the way; even more effort will be focused on process scheduling and memory location.

- Expect to see more multi-threaded libraries.
References

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

- www-unix.mcs.anl.gov/mpi/mpi-standard/mpi-report-2.0/node162.htm#Node162

- services.tacc.utexas.edu/index.php/ranger-user-guide#Process%20Affinity%20and%20Memory%20Policy

- www.mpi-forum.org/docs/mpi2-report.pdf