Hybrid Programming

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This short talk is a conglomeration of larger presentations given by the above authors at a variety of tutorials including SC08, SC09 (upcoming) ParCFD 2009, SciDAC 2009 We are grateful for the use of these slides at the NUG User Group Meeting

Computer Centers and Vendors are Responding with New Designs

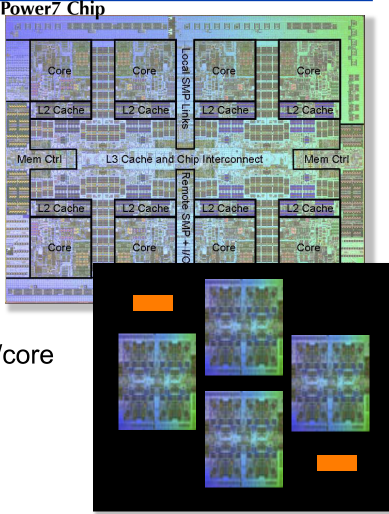
- Virtually all upcoming systems have various forms of heterogeneous parallelism
 - NERSC6 with its multicore design TBA
 - Blue Waters with its Power7 hardware threaded design

8 cores, 12 execution units/core, 4-way SMT/core

- ASC Sequoia (follow-on to BlueGene design) with anticipated support for transactional memory
- Experts everywhere are preparing for this architecture revolution with new languages, extensions to old languages, tools (and angst)
- Our goal at NERSC is to make this as painless as possible for application scientists
- We invite you to comment on our plans

Blue Waters Project Power7 Chip: Heart of Blue Waters

- Base Technology
 - 45 nm, 576 mm²
 - 1.2 B transistors
- Chip
 - 8 cores
 - 12 execution units/core
 - 4-way SMT/core
 - Caches
 - 32 KB I,D-cache, 256 KB L2/core
 - 32 MB L3 (private/shared)
 - Dual DDR3 memory controllers
 - 100 GB/s sustained memory bandwidth





Slide Courtesy Thom Dunning, NCSA

Blue Waters Project Blue Waters Computing System

System Attribute	Ranger	Blu	e Waters
Vendor Processor	Sun AMD Barcelona	IBM IBM Power7	
Peak Performance (PF) Sustained Performance (PF)	0.579 <0.05	>20	~1
Number of Cores/Chip Number of Processor Cores	4 62,976	2 >3	8 >200,000
Amount of Memory (TB)	123	>6	>800
Amount of Disk Storage (PE Amount of Archival Storage	· · · · · · · · · · · · · · · · · · ·	>5 >200	>10 >500
External Bandwidth (Gbps)	10	>10	100-400

Red numbers are ratios between Blue Waters and Ranger NERSC Users Group Meeting 2009

Blue Gene/P integrates a Double "Hummer" Floating Point Unit for added functionality

•Four PowerPC 450 (PPC450) cores, cache coherent

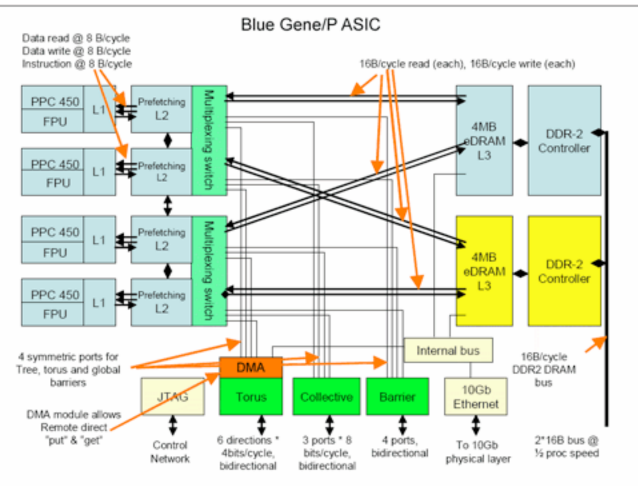
• Double "Hummer" floating point unit (FPU) per core

64-bit (double precision)

• Capable of SIMD floating-point multiply add (FMA) yielding 4 flops/cycle per core = 3.4 Gflop/s core

L2 Cache, 2 KB, private per core, 128-byte lines
L3 Cache, 8 MB, shared

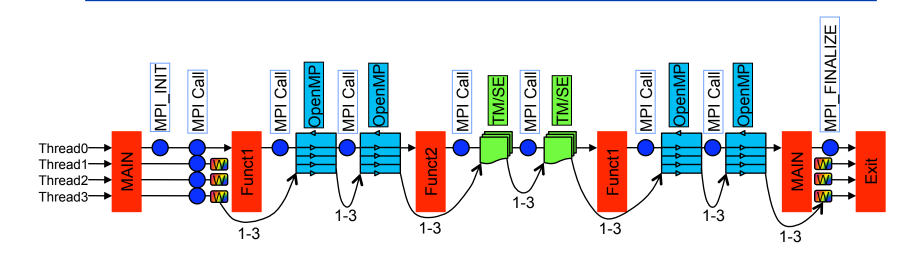
across all 4 cores, 128-byte lines



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Next generation Sequoia's programming model will likely be multilevel



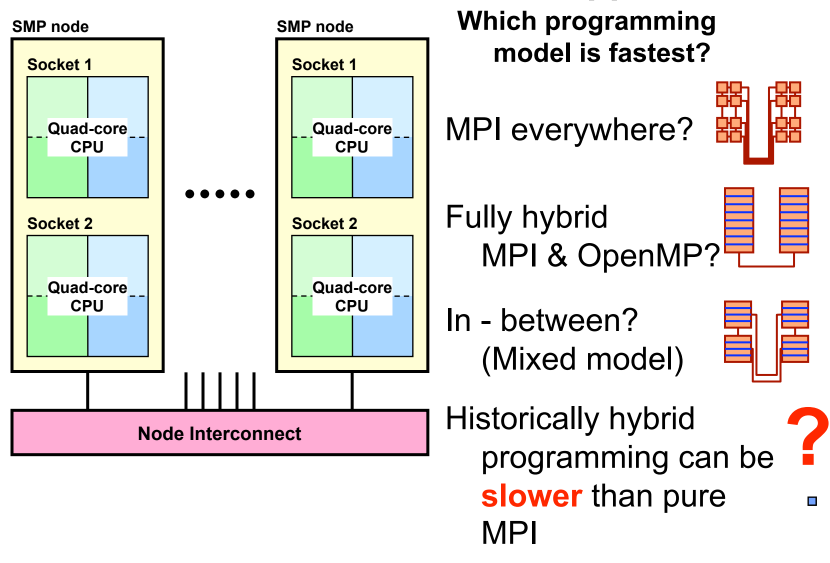
Possibly ~1.5 Million cores and even more threads!

Note: Speculative Execution (SE) and Transactional Memory (TM)

Diagram courtesy Mark Seager, LLNL

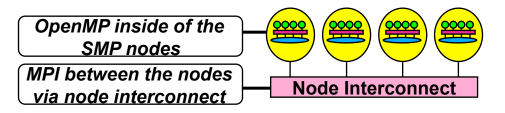
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Within the MPI-OpenMP hybrid model, there are variants depending on system and application

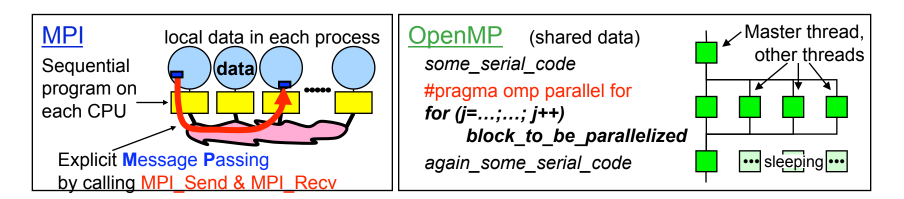


Programming models can be designed for hybrid systems

- Pure MPI (one MPI process on each CPU) "MPI-Everywhere"
- Hybrid MPI+OpenMP
 - shared memory OpenMP
 - distributed memory MPI



- Other: Virtual shared memory systems, PGAS, HPF, ...
- New Models combine MPI and UPC or CAF (see Lusk, et al. SC09, SIAM Feb. 2009.)



MPI and Threads

- MPI describes parallelism between processes (with separate address spaces)
- Thread parallelism provides a shared-memory model within a process, commonly Pthreads and OpenMP
- In the threads model of parallel programming, a single process can have multiple, concurrent execution paths
 - Pthreads (Posix Threads) is a standard library implementation that can be used for parallel programming
 - Pthreads generally provides more complicated and dynamic approaches
 - OpenMP is a set of compiler directives, callable runtime library routines, and environment variables that extend Fortran, C and C++
 - OpenMP provides convenient features for loop-level parallelism
 - OpenMP 3.0 adds task parallelism (released May 2008)

Hybrid Programming generally combines message passing with shared memory

- Other choices for a shared memory model include an implementations by Microsoft and others, as well as Pthreads
- In this talk, we concentrate on combining MPI with OpenMP
- OpenMP consists mostly of directives, where as MPI, related message passing libraries, and Pthreads methods consist of library routines
- Data movement in message passing libraries must be explicitly programmed, in contrast to OpenMP, where it happens automatically as data is read and written
- One goal of OpenMP is to make parallel programming easy
- It is also designed to be implemented incrementally
- We first explain some basic concepts of OpenMP, since we assume you are starting with an MPI program, or knowledge of MPI

Basic Concepts of OpenMP

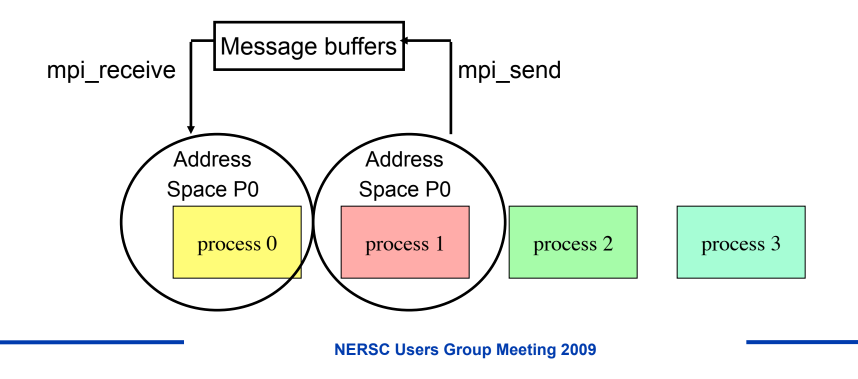
- OpenMP is an explicit programming model, namely the programmer specifies the parallelism. The compiler and run time system translate this into the parallel execution model.
- The task of the programmer is to correctly identify the parallelism and the dependencies
- OpenMP can have both implicit and explicit synchronization points
- Although OpenMP can be used beyond loop parallelism, we recommend studying loop parallelism first as a way to understand certain concepts such as private and shared variables, false sharing, race conditions, ...

MPI Memory Model

Message Passing Interface

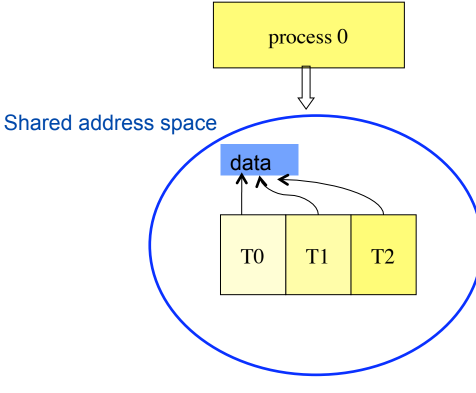
• Memory Model:

- MPI assumes a private address space
- Private address space for each MPI Process
- Data needs to be explicitly communicated
- Applies to distributed and shared memory computer architectures



OpenMP Memory Model

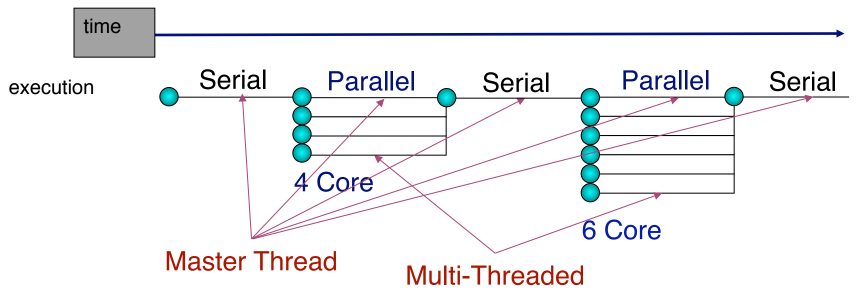
- OpenMP assumes a shared address space
- No communication is required between threads
- Thread Synchronization is required when accessing shared data



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OpenMP Code General Structure

- Fork-Join Model:
- Execution begins with a single "Master Thread"
- A team of threads is created at each parallel region
- Threads are joined at the end of parallel regions
- Execution is continued after parallel region by the Master Thread until the beginning of the next parallel region



The ParLab at Berkeley sponsored a Parallel Computing Bootcamp

- Note: All Bootcamp taped lectures, slides can be downloaded
 - See <u>http://parlab.eecs.berkeley.edu/bootcampagenda</u>
 - Including:
 - OpenMP
 - PGAS Languages
 - OpenCL
 - And several other interesting lectures
- From this website, you can also download the following:
 - makefile that should work on all NERSC clusters if you uncomment appropriate lines,
 - job-franklin-serial, job-franklin-pthreads4, job-franklin-openmp4, job-franklin-mpi4,
 - job-bassi-serial, job-bassi-pthreads8, job-bassi-openmp8, job-bassi-mpi8
 - sample batch files to launch jobs on Franklin and Bassi. Use qsub to submit on Franklin and Ilsubmit to submit on Bassi.
- We are also preparing sample codes to be available from the NERSC Website
- We recommend if you don't know OpenMP, you watch the ParLab BootCamp talk by Tim Mattson

Comments on Example from ParLAB Talk by Tim Mattson, Intel Corp.

Program to compute PI by quadrature, Tim starts with a simple serial program:

Next he shows Steps of Parallelization by OpenMP

• Identify concurrency in the loop, iterations may be executed concurrently:

- Steps: isolate data tha must be shared from data that will be local to a task
- Redefine x to remove loop carried dependence
- Look at how to rewrite the "reduction," where results from each iteration are accumulated into a single global sum
- Trick to promote scalar "sum" to an array indexed by the number of threads to create thread local copies of shared data

Examines differences between explicitly specifying the OpenMP other methods

• Final loop with explicit safe update of shared data

```
#include <omp.h>
static long num steps = 100000;
#define NUM 4
double step:
void main()
           double pi, sum = 0.0;
           step = 1.0/(double) num steps;
#pragma omp parallel num_threads(NUM)
           int i, ID;
                      double x, psum=0.0;
           ID = omp_get_thread_num();
           for (i=ID;i<=num steps;i+=nthreads){</pre>
                       x=(i+0.5)*step;
                       psum + = 4.0/(1.0 + x^*x);
           #pragma omp critical
                       sum += psum
}
           pi = step * sum;
```

 Along the way, example exposes concepts of private and shared data, false sharing, and other concepts. Refer to ParLab talk/slides.

Another version of the same PI program uses common OpenMP constructs

- Private clause for creating data local to a thread
- Reduction clause for managing data dependencies

 Most people would write the OpenMP in this fashion, but the example serves to illustrate some of the steps behind the directives and the thought process for shared/private variables

In order to do hybrid programming, models must be combined via Standards

- Hybrid programming (two programming models) requires that the standards make commitments to each other on semantics.
- OpenMP's commitment: if a thread is blocked by an operating system call (e.g. file or network I/O), the other threads remain runnable.
 - This is a major commitment; it involves the thread scheduler in the OpenMP compiler's runtime system and interaction with the OS.
 - What this means in the MPI context: An MPI call like MPI_Recv or MPI_Wait only blocks the calling thread.
- MPI's commitments are more complex...

The MPI Standard Defines 4 Levels of Thread Safety that affect Hybrid

- Note that these are not specific to Hybrid OpenMP Models
- The are in the form of commitments that the multithreaded application makes to the MPI implementation
 - MPI_THREAD_SINGLE: only one thread in the application
 - MPI_THREAD_FUNNELED: only one thread makes MPI calls, the Master Thread in the OpenMP context (next slide)
 - MPI_THREAD_SERIALIZED: Multiple threads make MPI calls, but only one at a time (not concurrently)
 - MPI_THREAD_MULTIPLE: Any thread may make MPI calls at any time, no restrictions
- MPI-2 defines an alternative to MPI_Init
 - MPI_Init_thread(requested, provided)
 - Allows applications to say what level it needs, and the MPI implementation to say what it provides

What This Means in the OpenMP Context

• MPI_THREAD_SINGLE

- There is no OpenMP multithreading in the program.
- MPI_THREAD_FUNNELED
 - All of the MPI calls are made by the master thread i.e., all MPI calls are
 - Outside OpenMP parallel regions, or
 - Inside OpenMP master regions, or
 - Guarded by call to MPI_Is_thread_main MPI call.
 - (same thread that called MPI_Init_thread)

• MPI_THREAD_SERIALIZED

#pragma omp parallel

#pragma omp atomic

...MPI calls allowed here...

```
• MPI_THREAD_MULTIPLE
```

– Anything goes; any thread may make an MPI call at any time

Threads and MPI in MPI-2

- An implementation is not required to support levels higher than MPI_THREAD_SINGLE; that is, an implementation is not required to be thread safe in order to be standard-conforming
- A fully thread-compliant implementation will support MPI_THREAD_MULTIPLE
- A portable program that does not call MPI_Init_thread should assume that only MPI_THREAD_SINGLE is supported

For MPI_THREAD_MULTIPLE

- When multiple threads make MPI calls concurrently, the outcome will be as if the calls executed sequentially in some (any) order
- Blocking MPI calls will block only the calling thread and will not prevent other threads from running or executing MPI functions
- It is the user's responsibility to prevent races when threads in the same process post conflicting MPI calls
- User must ensure that collective operations on the same communicator, window, or file handle are correctly ordered among threads

The Current Situation

- All MPI implementations support MPI_THREAD_SINGLE (duh).
- They probably support MPI_THREAD_FUNNELED even if they don't admit it.
 - Does require thread-safe malloc
 - Probably OK in OpenMP programs
- "Thread-safe" usually means MPI_THREAD_MULTIPLE.
- This is hard for MPI implementations that are sensitive to performance, like MPICH2.
 - Lock granularity issue
 - Working on lock-free MPICH2 implementation
- "Easy" OpenMP programs (loops parallelized with OpenMP, communication in between loops) only need FUNNELED.
 - So don't need "thread-safe" MPI for many hybrid programs
 - But watch out for Amdahl's Law!

How to determine thread support

MPI_Init_thread(&argc,&argv, MPI_THREAD_MULTIPLE,&provided);
printf("Supports level %d of %d %d %d %d %d\n",
 provided,
 MPI_THREAD_SINGLE,
 MPI_THREAD_FUNNELED,
 MPI_THREAD_SERIALIZED,
 MPI_THREAD_MULTIPLE);

Example output:

>Supports level 1 of 0 1 2 3

Support may vary depending on chosen compiler and MPI library.

Thread Support Levels on Franklin-PGI

PGI compiler

- with libmpich.a:
 - + MPICH_MAX_THREAD_SAFETY not set: Supports level 0 of 0 1 2 3
 - + MPICH_MAX_THREAD_SAFETY='single': Supports level 0 of 0 1 2 3
 - + MPICH_MAX_THREAD_SAFETY='funneled': Supports level 1 of 0 1 2 3
 - + MPICH_MAX_THREAD_SAFETY='serialized': Supports level 2 of 0 1 2 3

+ MPICH_MAX_THREAD_SAFETY='multiple': This library does not support multiple.

- with libmpich_threadm.a:
 - + MPICH_MAX_THREAD_SAFETY not set: Supports level 0 of 0 1 2 3
 - + MPICH_MAX_THREAD_SAFETY='single': Supports level 0 of 0 1 2 3
 - + MPICH_MAX_THREAD_SAFETY='funneled': Supports level 1 of 0 1 2 3
 - + MPICH_MAX_THREAD_SAFETY='serialized': Supports level 2 of 0 1 2 3
 - + MPICH_MAX_THREAD_SAFETY='multiple': Supports level 3 of 0 1 2 3

Thread Support Levels on Franklin-PathScale and gcc

PathScale and gcc compilers give same result

- with libmpich.a:
 - + MPICH_MAX_THREAD_SAFETY not set: Supports level 0 of 0 1 2 3
 - + MPICH_MAX_THREAD_SAFETY='single': Supports level 0 of 0 1 2 3
 - + MPICH_MAX_THREAD_SAFETY='funneled': Supports level 1 of 0 1 2 3
 - + MPICH_MAX_THREAD_SAFETY='serialized': Supports level 2 of 0 1 2 3
 - + MPICH_MAX_THREAD_SAFETY='multiple': This library does not support multiple.
 - with libmpich_threadm.a not available

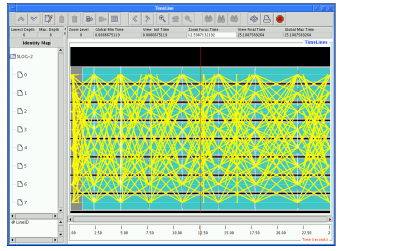
Thread Support Levels on Franklin-Summary

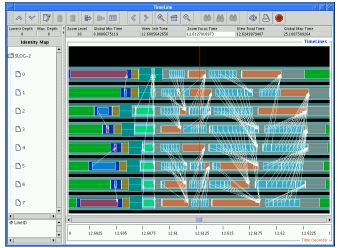
- Without setting the env variable, we always get '0'.
- To increase the supported level,
 - We need to explicitly set the environment variable.
- The MPI_THREAD_MULTIPLE level is currently supported only for PGI.
 - To get this level, one needs to link against libmpich_threadm.a.

Thanks to Woo-Sun Yang, NERSC User Services, for testing this on Franklin

Argonne has developed tools to Visualize the Behavior of Hybrid Programs

- Jumpshot is a logfile-based parallel program visualizer of the "standard" type. Uses MPI profiling interface.
- Recently it has been augmented in two ways to improve scalability.
 - Summary states and messages are shown as well as individual states and messages.
 - Provides a high-level view of a long run.
 - SLOG2 logfile structure allows fast interactive access (jumping, scrolling, and zooming) for large logfiles.

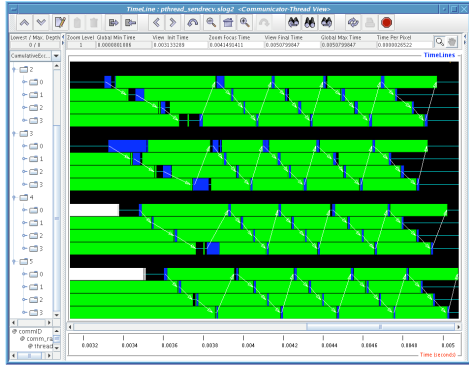




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Jumpshot and Multithreading

- Newest additions are for multithreaded and hybrid programs that use pthreads.
 - Separate timelines for each thread id
 - Support for grouping threads by communicator as well as by process



Using Jumpshot with Hybrid Programs

- SLOG2/Jumpshot needs two properties of the OpenMP implementation that are not guaranteed by the OpenMP standard
 - OpenMP threads must be pthreads
 - Otherwise, the locking in the logging library necessary to preserve exclusive access to the logging buffers would need to be modified.
 - These pthread ids must be reused (threads are "parked" when not in use)
 - Otherwise Jumpshot would need zillions of time lines.
- At NERSC, we are currently examining the use of Jumpshot and other tools for the analysis of hybrid programs
 - Part of the Cray Center of Excellence (COE) Program

Some Available Performance Tools

- All/Most Major HPC platforms
 - IPM Integrated Performance Manager, NERSC and others
 - Performance Visualization for Parallel Programs
 - <u>http://www-unix.mcs.anl.gov/perfvis</u> (jumpshot, etc)
 - TAU <u>http://www.cs.uoregon.edu/research/paracomp/tau/</u>
 - C++, Java, C, Fortran 90, HPF, and HPC++, and OpenMP
 - Vampir: <u>http://www.vampir-ng.com</u> for parallel programs (Tracing supported through TAU, KOJAK, and Adaptor)
 - Pablo: http://www-pablo.cs.uiuc.edu/ I/O and visualization
 - PAPI: Performance analysis using hardware counters
 - http://icl.cs.utk.edu/projects/papi/
 - Aims: http://www.nas.nasa.gov/Groups/Tools/Projects/AIMS/
 - Paradyn can operate on executable <u>http://www.cs.wisc.edu/~paradyn/</u>
 - PARAVER <u>http://www.cepba.upc.es/paraver</u>
 - MPI, OpenMP, Performance counters, system activity
 - GNU gprof
 - KOJAK http://www.fz-juelich.de/zam/kojak/
- Also most vendors support their own tools (e.g., Cray Apprentice2)
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Visualization and Analysis of Parallel Programs

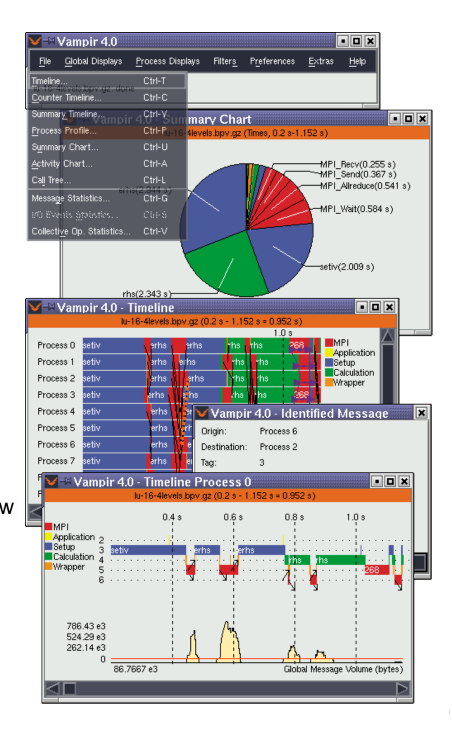


- Trace data based
- Global displays show all selected processes

 Timeline: application execution over time
 Counter Timeline: counter dev. over time
 Summary Timeline: shows parallelism
 Summary Chart: aggregated profiling
 Activity Chart: presents per–process profiling
 Call Tree: draws dynamic call tree
 Message Statistics: for each process pair
 I/O Statistics: MPI I/O operation statistics
 Collective Op. Statistics: collective operations

 Process displays show a single process per window
 - -Activity Chart
 - -Timeline
 - -Call Tree

Vampir Slides courtesy of Wolfgang E. Nagel, Holger Brunst TU Dresden. www.vampir-ng.com



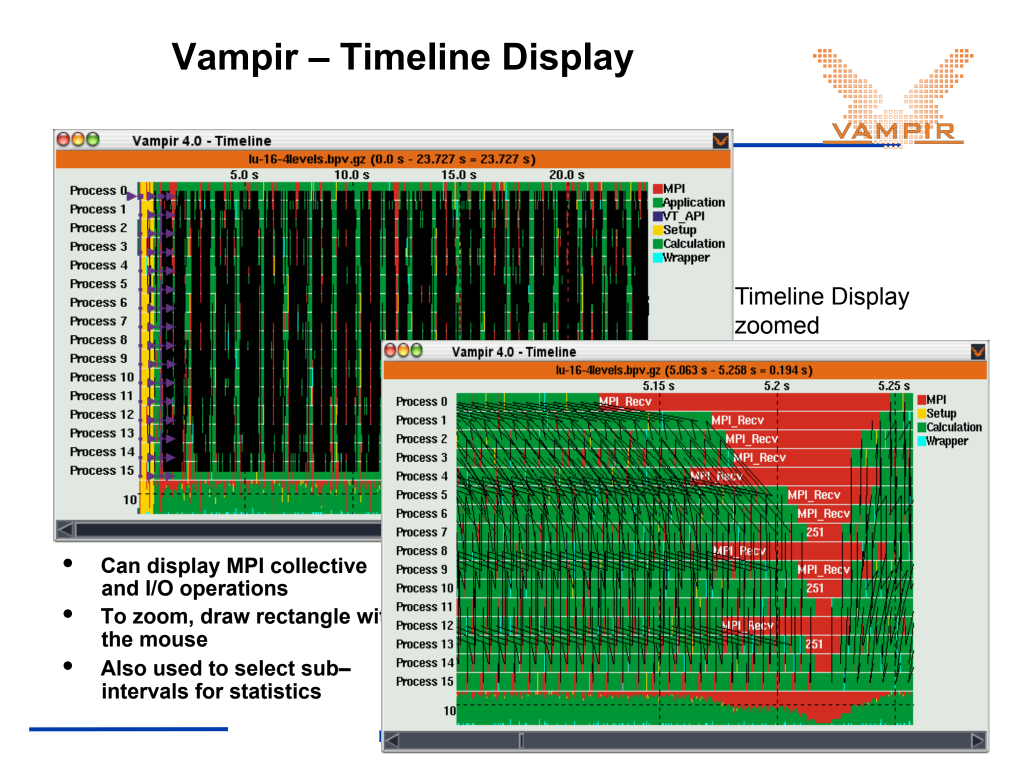
Vampir – Routine Profiles and Statistics



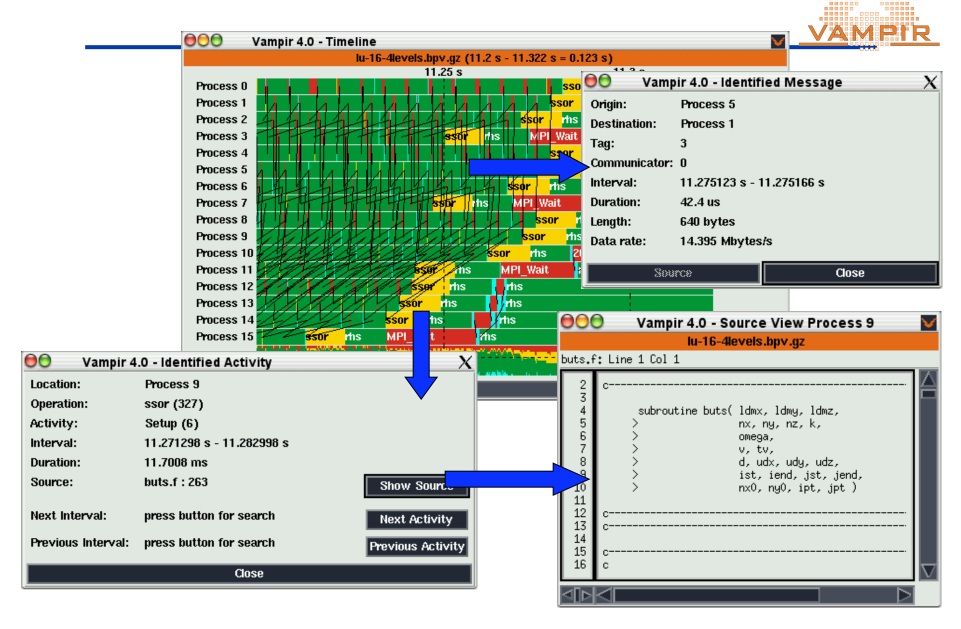
• Setup Summary Chart

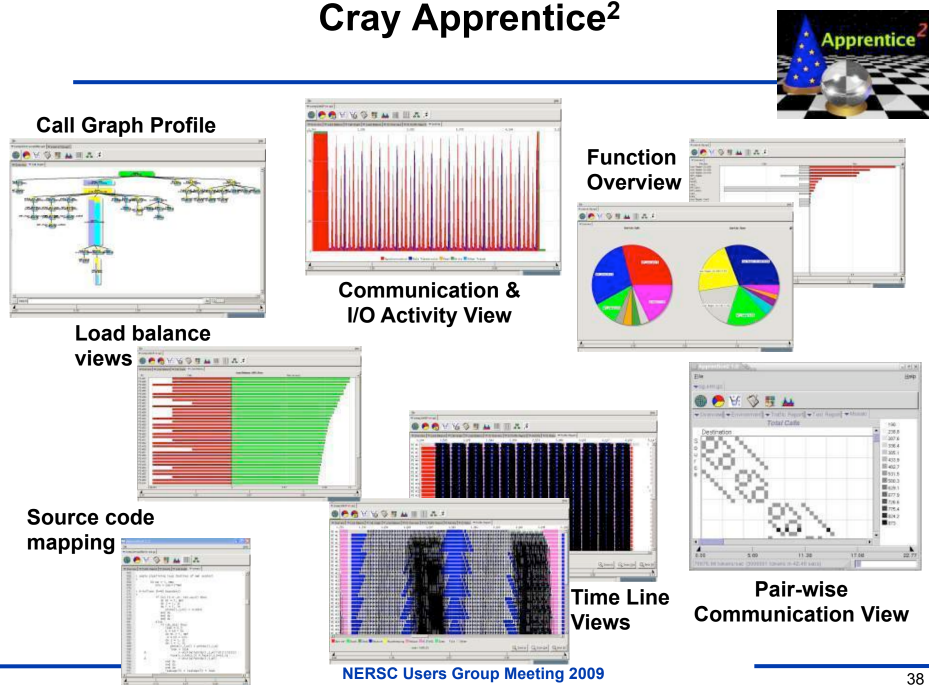
- Global Displays ⇒
 Summary Chart
- Context Menu \Rightarrow Select \Rightarrow All Symbols
- Context Menu ⇒
 Options ⇒Per Process
- Selected group: All Master Threads

r.gz (Times, 0.0 s-24.224 s) (Per Process 5.001 s 4.671 s 4.367 s 2.734 s 2.681 s
4.367 s 2.734 s
2.734 s
2.681 s
2.188 s
0.531 s
0.497 s
0.309 s
0.297 s
0.246 s
0.213 s
0.115 s
89.816 ms
76.059 ms
44.973 ms
43.142 ms
10.348 ms

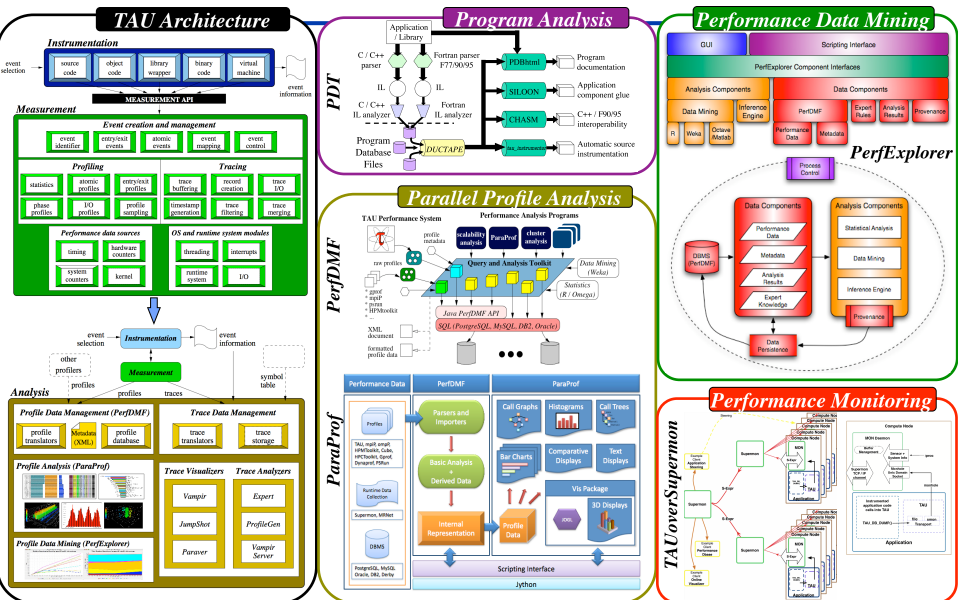


Vampir – Timeline Display– Info Dialogs





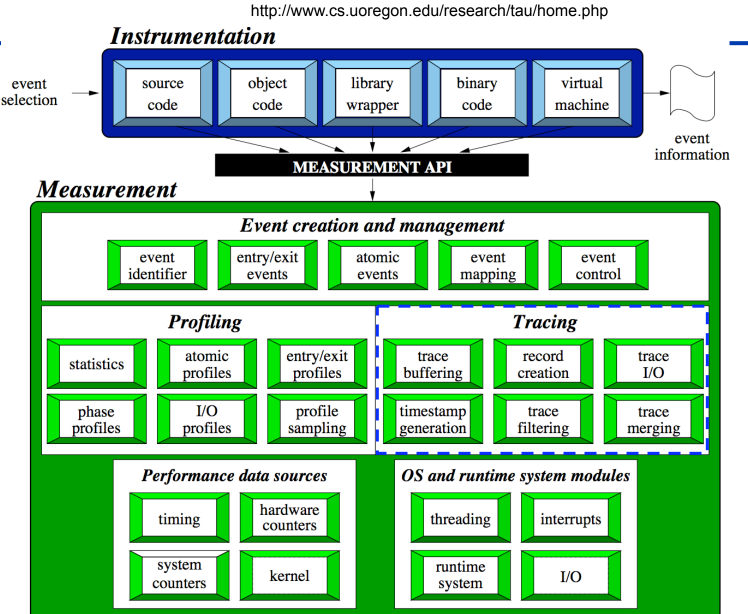
TAU Performance System Components



http://www.cs.uoregon.edu/research/tau/home.php

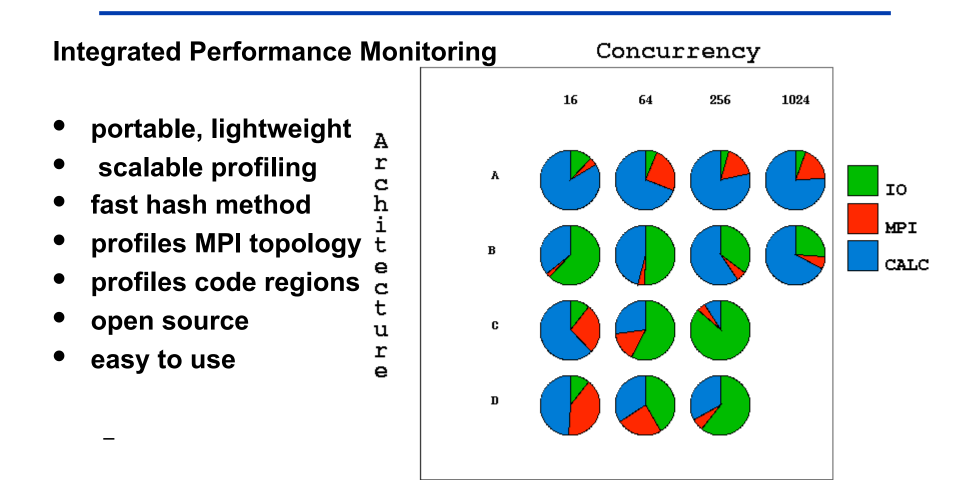
NERSC Users Group Meeting From Sameer Shende U. Oregon 39

TAU Performance System Architecture



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IPM Performance Tool



But, your code's "hot" spots may vary depending on architecture

Extremely difficult to use tools on very large numbers of processors: research area

- TotalView scalability
 - BlueGene/L port included significant scaling requirements
 - Collaborative scalability project targeting traditional OSs
 - Scalable subset debugging on BG/L
- Scalable, lightweight tools
 - "Focused tools": fewer features, but extreme scalability
 - Stack Trace Analysis Tool (STAT)
 - Scalable infrastructure: LaunchMON, MRNet
- Automatic error detection
 - Memory correctness tools
 - Umpire
 - Threading tools

Adapted from Scott Futral, Dong Ahn, and Greg Lee, LLNL

Advice for Debugging and Tuning MPI-OpenMP Hybrid Codes

- Debug and Tune MPI code and OpenMP code separately.
- Decide which loop to parallelize. Better to parallelize outer loop. Decide whether Loop permutation or loop exchange is needed.
- Choose between loop-based or SPMD.
- Use different OpenMP task scheduling options.
- Experiment with different combinations of MPI tasks and number of threads per MPI task.
- Adjust environment variables.
- Aggressively investigate different thread initialization options and the possibility of overlapping communication with computation.
- See afternoon section on hybrid programming

Adapted from Helen He LBL-NERSC

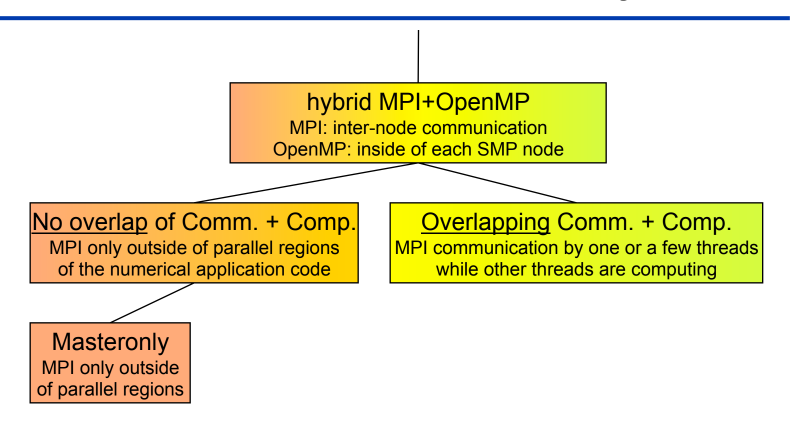
Performance Strategies

After isolating portions of code to optimize, what next?

Global Performance Strategies

- Degree of parallelism over entire code
- Number of parallel tasks as compared with available numbers of processors
- Uniformity of tasks / load balancing
- Synchronization requirements
- Processor-specific strategies
 - If cache-based architecture: data locality
 - Parallelism via a "super scalar" processor
 - Pipelining
 - Vector processors
- But, the way things are progressing, you may want to wait on optimization and instead consider
 - New languages
 - Auto-tuning options

Rabensiefners Models for Hybrid



45

Hybrid Masteronly

Masteronly MPI only outside of parallel regions Advantages

- No message passing inside of the SMP nodes
- No topology problem

for (iteration)

```
#pragma omp parallel
numerical code
/*end omp parallel */
```

/* on master thread only */
MPI_Send (original data
 to halo areas
 in other SMP nodes)
MPI_Recv (halo data
 from the neighbors)
} /*end for loop

Major Problems

- All other threads are sleeping while master thread communicates!
- What is inter-node bandwidth?
- MPI-lib must support at least MPI_THREAD_FUNNELED

Overlapping Communication and Computation

MPI communication by one or a few threads while other threads are computing

```
if (my_thread_rank < ...) {
    MPI_Send/Recv....
    i.e., communicate all halo data
} else {
    Execute those parts of the application
    that do not need halo data
    (on non-communicating threads)
}
Execute those parts of the application
    that need halo data
    (on all threads)</pre>
```

This can get very complicated. Looks a little bit more like MPI Programming.

Overlapping Communication and Computation

MPI communication by one or a few threads while other threads are computing

Three problems:

- the application problem:
 - one must separate application into:
 - code that can run before the halo data is received
 - code that needs halo data

→ very hard to do !!!

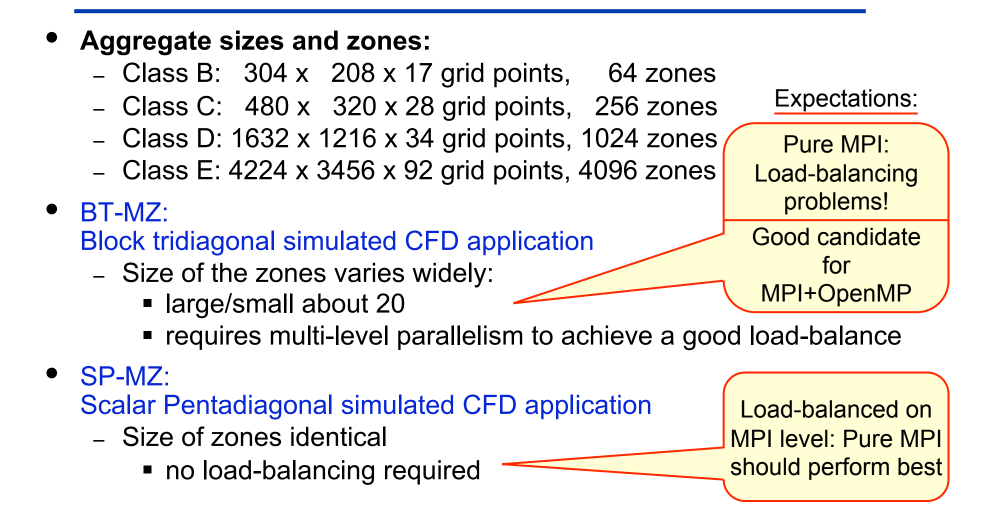
→ loss of ma OpenMP s (see next)	omp. via ik e ing directives ajor support slide)	<pre>if (my_thread_rank < 1) { MPI_Send/Recv } else { my_range = (high-low-1) / (num_threads-1) + 1; my_low = low + (my_thread_rank+1)*my_range; my_high=high+ (my_thread_rank+1+1)*my_range; my_high = max(high, my_high) for (i=my_low; i<my_high; <="" i++)="" pre="" {="" }=""></my_high;></pre>
• the load bala	ncing problem	}

Overlapping Communication and Computation

MPI communication by one or a few threads while other threads are computing

```
#pragma omp parallel
<u>Subteams</u>
                             {
   Important proposal
                             #pragma omp single onthreads( 0 )
   for OpenMP 3.x
   or OpenMP 4.x
                                MPI Send/Recv....
                             #pragma omp for onthreads( 1 : omp get numthreads()-1 )
                                for (.....)
                                { /* work without halo information */
   Barbara Chapman et al.:
   Toward Enhancing
                                } /* barrier at the end is only inside of the subteam */
   OpenMP's Work-Sharing
   Directives.
                             #pragma omp barrier
   In proceedings, W.E.
                             #pragma omp for
   Nagel et al. (Eds.): Euro-
                                for (.....)
   Par 2006, LNCS 4128, pp.
                                { /* work based on halo information */
   645-654, 2006.
                             /*end omp parallel */
```

Multi-zone NAS Parallel Benchmarks – Characteristics

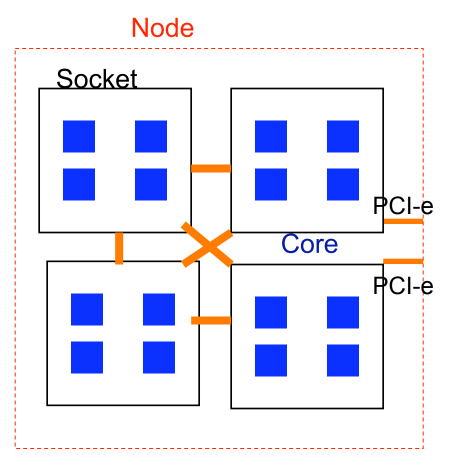


Sun Constellation Cluster Ranger (1)

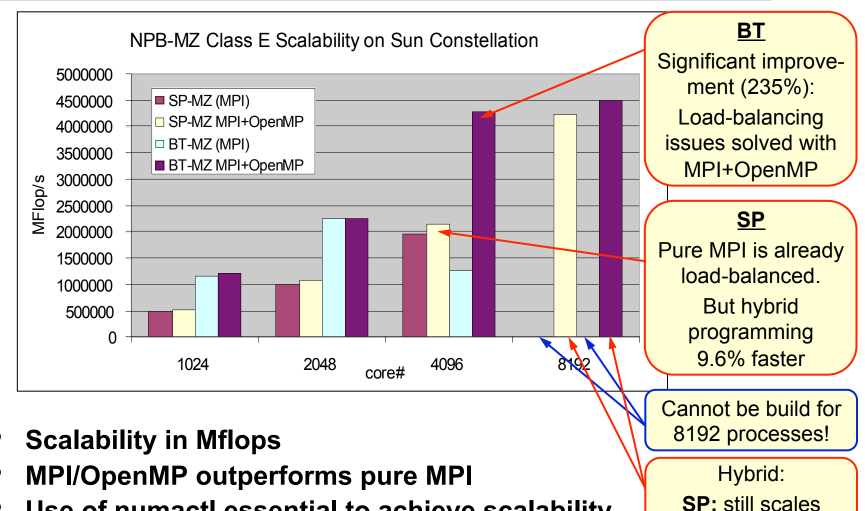
- Located at the Texas Advanced Computing Center (TACC), University of Texas at Austin (http://www.tacc.utexas.edu)
- 3936 Sun Blades, 4 AMD Quad-core 64bit 2.3GHz processors per node (blade), 62976 cores total
- 123TB aggregrate memory
- Peak Performance 579 Tflops
- InfiniBand Switch interconnect
- Sun Blade x6420 Compute Node:
 - 4 Sockets per node
 - 4 cores per socket
 - HyperTransport System Bus
 - 32GB memory

Sun Constellation Cluster Ranger (2)

- Compilation:
 - PGI pgf90 7.1
 - mpif90 -tp barcelona-64 -r8
- Cache optimized benchmarks
 Execution:
 - MPI MVAPICH
 - setenv
 OMP_NUM_THREAD
 NTHREAD
 - Ibrun numactl bt-mz.exe
- numactl controls
 - Socket affinity: select sockets to run
 - Core affinity: select cores within socket
 - Memory policy: where to allocate memory
 - http://www.halobates.de/numaapi3.pdf



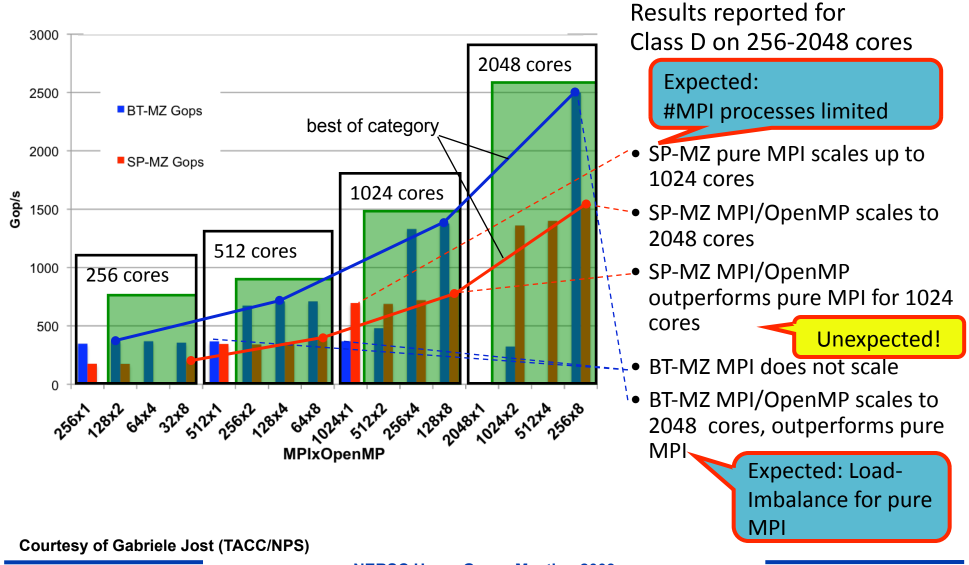
NPB-MZ Class E Scalability on Ranger



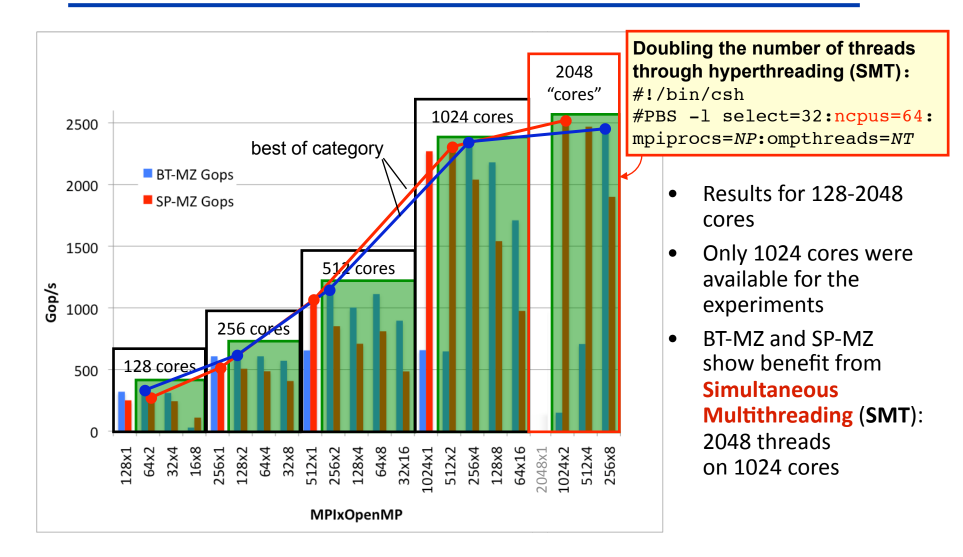
• Use of numactl essential to achieve scalability

BT: does not scale

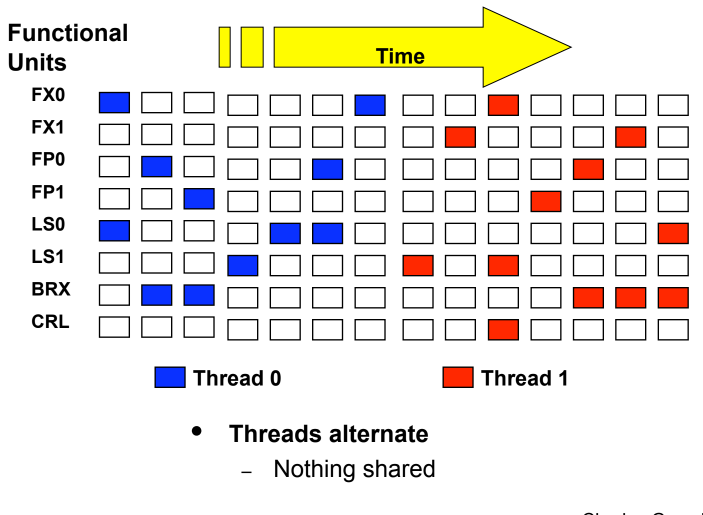
Cray XT5: NPB-MZ Class D Scalability



NPB-MZ Class D on IBM Power 6: Exploiting SMT for 2048 Core Results

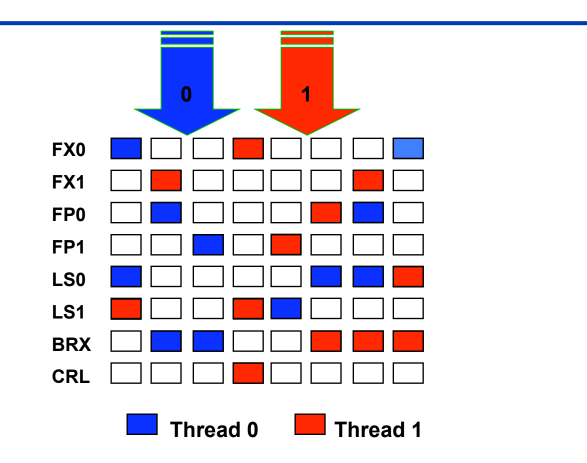


Conventional Multi-Threading



Charles Grassl, IBM

Simultaneous Multi-Threading

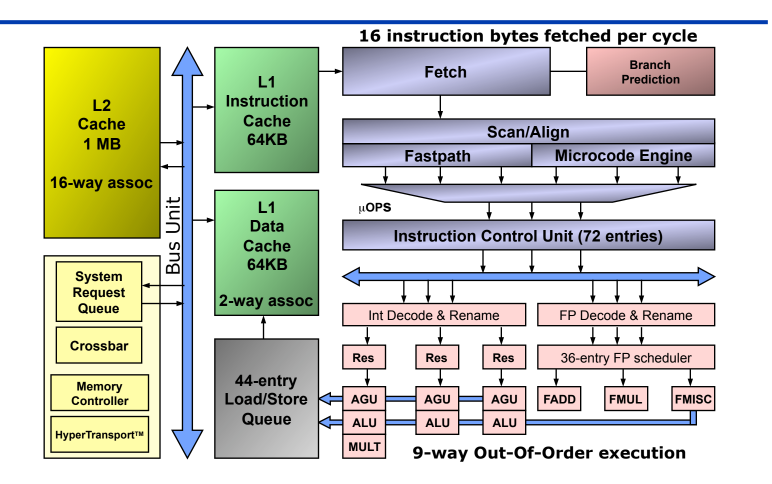


Simultaneous execution

- Shared registers
- Shared functional units

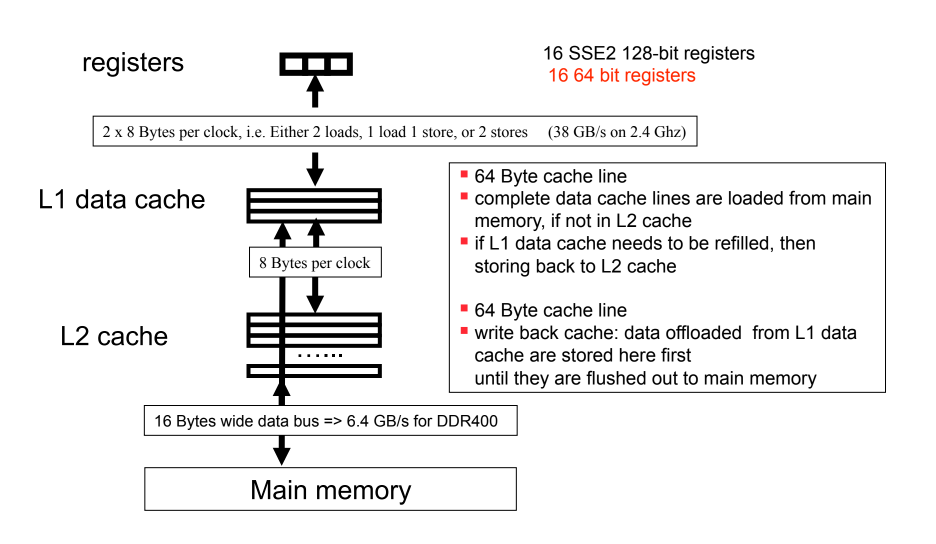
Charles Grassl, IBM

AMD Opteron Processor



- 36 entry FPU instruction scheduler
- 64-bit/80-bit FP Realized throughput (1 Mul + 1 Add)/cycle: 1.9 FLOPs/cycle
- 32-bit FP Realized throughput (2 Mul + 2 Add)/cycle: 3.4+ FLOPs/cycle

Simplified memory hierachy on the AMD Opteron



Hybrid parallelization affords new opportunities

- Nested Parallelism
- Load-Balancing
- Memory consumption
- Opportunities, if MPI speedup is limited due to "algorithmic" problem
- MPI scaling problems

Load-Balancing (on same or different level of parallelism)

• OpenMP enables

- Cheap *dynamic* and *guided* load-balancing
- Just a parallelization option (clause on omp for / do directive)
- Without additional software effort
- Without explicit data movement
- On MPI level
 - Dynamic load balancing requires moving of parts of the data structure through the network
 - Significant runtime overhead
 - Complicated software / therefore not implemented
- MPI & OpenMP
 - Simple static load-balancing on MPI level, medium quality dynamic or guided on OpenMP level cheap implementation

Memory consumption

• Shared nothing

- Heroic theory
- In practice: Some data is duplicated

MPI & OpenMP With n threads per MPI process:

- Duplicated data is reduced by factor n

Memory consumption (continued)

Future: With 100+ cores per chip the memory per core is limited.

- Data reduction through usage of shared memory may be a key issue
- Domain decomposition on each hardware level
 - Maximizes
 - Data locality
 - Cache reuse
 - Minimizes
 - CCnuma accesses
 - Message passing
- No halos between domains inside of SMP node
 - Minimizes
 - Memory consumption

How many multi-threaded MPI processes per SMP node

- SMP node = with **m sockets** and **n cores/socket**
- How many threads (i.e., cores) per MPI process?
 - Too many threads per MPI process
 - \rightarrow overlapping of MPI and computation may be necessary,
 - \rightarrow some NICs unused?
 - Too few threads
 - \rightarrow too much memory consumption (see previous slides)
- Optimum
 - somewhere between 1 and m x n

Opportunities, if MPI speedup is limited due to "algorithmic" problems

- Algorithmic opportunities due to larger physical domains inside of each MPI process
 - \rightarrow If multigrid algorithm only inside of MPI processes
 - → If separate preconditioning inside of MPI nodes and between MPI nodes
 - \rightarrow If MPI domain decomposition is based on physical zones

To overcome MPI scaling problems

 Reduced number of MPI messages, reduced aggregated message size

compared to pure MPI

- MPI has a few scaling problems
 - Handling of more than 10,000 processes
 - Irregular Collectives: MPI_....v(), e.g. MPI_Gatherv()
 - Scaling applications should not use MPI_....v() routines
 - MPI-2.1 Graph topology (MPI_Graph_create)
 - MPI-2.2 MPI_Dist_graph_create_adjacent
 - Creation of sub-communicators with MPI_Comm_create
 - MPI-2.2 introduces a new scaling meaning of MPI_Comm_create
- Hybrid programming reduces all these problems (due to a smaller number of processes)

Summary of Hybrid Programming

<u>MPI + OpenMP</u>

- Significant opportunity → higher performance on fixed number of cores
- NPB-MZ examples
 - BT-MZ \rightarrow strong improvement (as expected)
 - SP-MZ \rightarrow small improvement (none was expected)
- Often however, no speedup is obtained, especially for naïve implementations
- Hybrid MPI + OpenMP can solve certain problems with MPI-Everywhere
 - Load balancing
 - Memory consumption
 - Two levels of parallelism
 - Outer \rightarrow distributed memory \rightarrow halo data transfer \rightarrow MPI
 - Inner \rightarrow shared memory \rightarrow ease of SMP parallelization \rightarrow OpenMP

Summary (cont)

<u> MPI+OpenMP – Pitfalls:</u>

- Problems with OpenMP performance remain
 - On ccNUMA \rightarrow e.g., first touch
 - Pinning of threads on cores
- Problems with combination of MPI & OpenMP
 - topology and mapping problems
 - mismatch problems
- Most hybrid programs → Masteronly style
- Overlapping communication and computation with several threads
 - Requires thread-safety quality of MPI library
 - Loss of OpenMP support \rightarrow future OpenMP subteam concept

Hybrid Programming: more information

"Hybrid MPI and OpenMP Parallel Programming"

SC09 half-day tutorial M09, Monday, 8:30am – 12:00pm

Reminder, these slides are excerpted various tutorials given by Rolf Rabenseifner, Gabriele Jost, Rusty Lusk, Alice Koniges, et al. including SC08, SC09 (upcoming) ParCFD 2009, SciDAC 2009 We are grateful for the use of these slides at the NUG User Group Meeting