Advanced Topics in MPI

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Outline

- What is MPI and where does it fit into HPC?
- Selected topics in MPI programming beyond the basics
- MPI profiling interface and tools
  - SLOG/Jumpshot: visualizing parallel performance
  - FPMPI: gathering summary statistics
  - Collchk: runtime checking of correct use of collective operations
- MPI and threads: hybrid programming
- One-sided communication
- MPI at Exascale
- Recent Activities of the MPI Forum
- ADLB: a scalable load-balancing library built on MPI
What is MPI?

- MPI (Message-Passing Interface) is a message-passing library interface standard.
  - Specification, not implementation
  - Library, not a language
  - Classical message-passing programming model

- MPI-1 was defined (1994) by a broadly-based group of parallel computer vendors, computer scientists, and applications developers.
  - 2-year intensive process

- MPI-2 was standardized in 1997

- Implementations appeared quickly and now MPI is taken for granted as vendor-supported software on any parallel machine.

- Portable, open-source implementations exist for virtually every system in the world; MPICH2 and OpenMPI are widely-used implementations
MPI in a Nutshell

- A parallel program consists of multiple processes, each with its own program counter, call stack, and address space.
  - A process may be multi-threaded, in which case each thread has its own program counter and call stack, and they share the address space.
  - A process’s address space is not accessible to other processes via the compiler-generated load and store instructions.

- Data is moved from one address space to another using MPI
  - Pair-wise exchange model (send-receive)
    • Useful for computations with data dependencies (I wait for data from another process using MPI receive).
  - Group communication model (collective operations)
    • Coordinated data exchange between multiple processes – takes advantage of group semantics for performance improvements.
    • Applications form groups of sizes appropriate for their computation.
  - One-sided communication operations (Put, Get)
    • Useful for asynchronous or uncoordinated computations.
  - I/O capabilities (MPI-IO)
    • Data movement from memory space to the file-system (pair-wise and
Timeline of the MPI Standard

- **MPI-1 (1994)**
  - Basic point-to-point communication, collectives, datatypes, etc.

- **MPI-2 (1997)**
  - Added parallel I/O, RMA, dynamic processes, C++ and Fortran90 bindings, semantics of interaction with threads, etc.

- **----- Stable for 10 years -----**

- **MPI-2.1 (2008)**
  - Minor clarifications and bug fixes to MPI-2

- **MPI-2.2 (2009)**
  - Today’s official standard
  - Small updates and additions to MPI 2.1. Backward compatible

- **MPI-3 (in progress, expected early 2012)**
  - Major new features and additions to extend MPI to exascale
  - Organized into several working groups
  - Draft 1 was released last November; Draft 2 will be released this November
Where does MPI fit into Exascale?

- Despite its record of success, there are concerns about whether we should abandon MPI and look for an entirely new way of programming massive parallelism.
- Most of these questions are derived from a misunderstanding of what MPI’s role is now and how actively it is evolving to meet these concerns.
- Is MPI too low-level to be a “productive” programming model?
  - Some call it the “assembly language of parallel programming”
  - Actually, since it is portable, it’s the C
  - It wasn’t designed for ease of use, but rather for capabilities needed to develop sophisticated portable parallel libraries.
- Can MPI scale to the numbers of address spaces and threads that will be needed/provided in the future?
  - Being addressed now by both standards (MPI-3 Forum) and implementations.
- Will MPI be able to interoperate with other programming models that we will need for parallelism within an address space?
Beyond Elementary MPI
Message Passing, Buffering, Deadlocks

- Message passing is a simple programming model, but there are some special issues
  - Buffering and deadlock
  - Deterministic execution
  - Performance
**Buffers**

- When you send data, where does it go? One possibility is:

  ![Diagram showing data flow from Process 0 to Process 1]

  - User data
  - Local buffer
  - the network
  - Local buffer
  - User data
Avoiding Buffering

- It is better to avoid copies:

This requires that `MPI_Send` wait on delivery, or that `MPI_Send` return before transfer is complete, and we wait later.
Sources of Deadlocks

- Send a large message from process 0 to process 1
  - If there is insufficient storage at the destination, the send must wait for the user to provide the memory space (through a receive)
- What happens with this code?

<table>
<thead>
<tr>
<th>Process 0</th>
<th>Process 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Send (1)</td>
<td>Send (0)</td>
</tr>
<tr>
<td>Recv (1)</td>
<td>Recv (0)</td>
</tr>
</tbody>
</table>

- This is called “unsafe” because it depends on the availability of system buffers in which to store the data sent until it can be received
Some Solutions to the “unsafe” Problem

- Order the operations more carefully:

<table>
<thead>
<tr>
<th>Process 0</th>
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</thead>
<tbody>
<tr>
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<td>Recv (0)</td>
</tr>
<tr>
<td>Recv (1)</td>
<td>Send (0)</td>
</tr>
</tbody>
</table>

- Supply receive buffer at same time as send:

<table>
<thead>
<tr>
<th>Process 0</th>
<th>Process 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sendrecv (1)</td>
<td>Sendrecv (0)</td>
</tr>
</tbody>
</table>
More Solutions to the “unsafe” Problem

- Supply own space as buffer for send

<table>
<thead>
<tr>
<th>Process 0</th>
<th>Process 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bsend(1)</td>
<td>Bsend(0)</td>
</tr>
<tr>
<td>Recv(1)</td>
<td>Recv(0)</td>
</tr>
</tbody>
</table>

- Use non-blocking operations:

<table>
<thead>
<tr>
<th>Process 0</th>
<th>Process 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isend(1)</td>
<td>Isend(0)</td>
</tr>
<tr>
<td>Irecv(1)</td>
<td>Irecv(0)</td>
</tr>
<tr>
<td>Waitall</td>
<td>Waitall</td>
</tr>
</tbody>
</table>
Communication Modes

- MPI provides multiple modes for sending messages:
  - Synchronous mode (MPI_Ssend): the send does not complete until a matching receive has begun. (Unsafe programs deadlock.)
  - Buffered mode (MPI_Bsend): the user supplies a buffer to the system for its use. (User allocates enough memory to make an unsafe program safe.
  - Ready mode (MPI_Rsend): user guarantees that a matching receive has been posted.
    • Allows access to fast protocols
    • undefined behavior if matching receive not posted

- Non-blocking versions (MPI_Issend, etc.)
- MPI_Recv receives messages sent in any mode.
Buffered Mode

- When MPI_Isend is awkward to use (e.g. lots of small messages), the user can provide a buffer for the system to store messages that cannot immediately be sent.

  ```c
  int bufsize;
  char *buf = malloc( bufsize );
  MPI_Buffer_attach( buf, bufsize );
  ...
  MPI_Bsend( ... same as MPI_Send ... )
  ...
  MPI_Buffer_detach( &buf, &bufsize );
  ```

- MPI_Buffer_detach waits for completion.
- Performance depends on MPI implementation and size of message.
MPI_Sendrecv

- Allows simultaneous send and receive
- Everything else is general.
  - Send and receive datatypes (even type signatures) may be different
  - Can use Sendrecv with plain Send or Recv (or Irecv or Ssend_init, ...)
  - More general than “send left”

<table>
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<th>Process 1</th>
</tr>
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<tr>
<td>SendRecv(1)</td>
<td>SendRecv(0)</td>
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</table>
Understanding Performance: Unexpected Hot Spots

- Basic performance analysis looks at two-party exchanges
- Real applications involve many simultaneous communications
- Performance problems can arise even in common grid exchange patterns
- Message passing illustrates problems present even in shared memory
  - Blocking operations may cause unavoidable memory stalls
Basic MPI: Looking Closely at a Simple Communication Pattern

- Many programs rely on “halo exchange” (ghost cells, ghost points, stencils) as the core communication pattern
  - Many variations, depending on dimensions, stencil shape
  - Here we look carefully at a simple 2-D case

- Unexpected performance behavior
  - Even simple operations can give surprising performance behavior.
  - Examples arise even in common grid exchange patterns
  - Message passing illustrates problems present even in shared memory
    - Blocking operations may cause unavoidable stalls
Processor Parallelism

- Decomposition of a mesh into 1 patch per process
  - Update formula typically $a(i,j) = f(a(i-1,j), a(i+1,j), a(i,j+1), a(i,j-1), ...)$
  - Requires access to “neighbors” in adjacent patches
Sample Code

- Do i=1,n_neighbors
  Call MPI_Send(edge, len, MPI_REAL, nbr(i), tag, comm, ierr)
Enddo
Do i=1,n_neighbors
  Call MPI_Recv(edge, len, MPI_REAL, nbr(i), tag, comm, status, ierr)
Enddo

- What is wrong with this code?
Deadlocks!

- All of the sends may block, waiting for a matching receive (will for large enough messages)
- The variation of
  ```
  if (has down nbr)
      Call MPI_Send( ... down ... )
  if (has up nbr)
      Call MPI_Recv( ... up ... )
  ...
  ```
  sequentializes (all except the bottom process blocks)
Sequentialization

Start  Start  Start  Start  Start  Start  Send  Recv  Send  Recv
Send  Send  Send  Send  Send  Send  Send  Recv  Send  Recv
Send  Recv
Send  Recv
Send  Recv
Send  Recv
Send  Recv
Fix 1: Use Irecv

- Do $i = 1, n_{neighbors}$
  - Call MPI_Irecv(edge, len, MPI_REAL, nbr(i), tag, comm, requests(i), ierr)
Enddo
- Do $i = 1, n_{neighbors}$
  - Call MPI_Send(edge, len, MPI_REAL, nbr(i), tag, comm, ierr)
Enddo
- Call MPI_Waitall(n_neighbors, requests, statuses, ierr)

- Does not perform well in practice. Why?
Timing Model

- Sends interleave
- Sends block (data larger than buffering will allow)
- Sends control timing
- Receives do not interfere with Sends
- Exchange can be done in 4 steps (down, right, up, left)
Mesh Exchange - Step 1

- Exchange data on a mesh
Mesh Exchange - Step 2

- Exchange data on a mesh
Mesh Exchange - Step 3

- Exchange data on a mesh
Mesh Exchange - Step 4

- Exchange data on a mesh
Mesh Exchange - Step 5

- Exchange data on a mesh
Mesh Exchange - Step 6

- Exchange data on a mesh
Timeline from IBM SP

- Note that process 1 finishes last, as predicted
Distribution of Sends

'SEND' state length distribution

(in seconds)
68 states of 96 (70%)
Why Six Steps?

- Ordering of Sends introduces delays when there is contention at the receiver
- Takes roughly twice as long as it should
- Bandwidth is being wasted
- Same thing would happen if using memcpy and shared memory
Fix 2: Use Isend and Irecv

- Do i=1,n_neighbors
  Call MPI_Irecv(edge,len,MPI_REAL,nbr(i),tag,
                comm,request(i),ierr)
Enddo
Do i=1,n_neighbors
  Call MPI_Isend(edge,len,MPI_REAL,nbr(i),tag,
                comm,request(n_neighbors+i),ierr)
Enddo
Call MPI_Waitall(2*n_neighbors, request, statuses,
                 ierr)
Mesh Exchange - Steps 1-4

- Four interleaved steps
Note processes 5 and 6 are the only interior processors; these perform more communication than the other processors.
Lesson: Defer Synchronization

- Send–receive accomplishes two things:
  - Data transfer
  - Synchronization

- In many cases, there is more synchronization than required

- Use nonblocking operations and MPI_Waitall to defer synchronization
**MPI Message Ordering**

- Multiple messages from one process to another will be **matched** in order, not necessarily completed in order

<table>
<thead>
<tr>
<th>Rank 0</th>
<th>Rank 1</th>
<th>Rank 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>MPI_Isend(dest=1)</td>
<td>MPI_Irecv(any_src, any_tag)</td>
<td>MPI_Isend(dest=1)</td>
</tr>
<tr>
<td>MPI_Isend(dest=1)</td>
<td>MPI_Irecv(any_src, any_tag)</td>
<td>MPI_Isend(dest=1)</td>
</tr>
<tr>
<td></td>
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<td></td>
<td>MPI_Irecv(any_src, any_tag)</td>
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</tbody>
</table>
MPI Profiling Interface
Tools Enabled by the MPI Profiling Interface

- The MPI profiling interface: how it works
- Some freely available tools
  - Those to be presented in other talks
  - A few that come with MPICH2
    - SLOG/Jumpshot: visualization of detailed timelines
    - FPMPI: summary statistics
    - Collcheck: runtime checking of consistency in use of collective operations
The MPI Profiling Interface

Call MPI_Send

Call MPI_Bcast

User Program

MPI_Send

Profiling Library

MPI_Send

PMPI_Send

MPI_Bcast

MPI_Send

MPI_Bcast

MPI Library
Performance Visualization with Jumpshot

- For detailed analysis of parallel program behavior, timestamped events are collected into a log file during the run.
- A separate display program (Jumpshot) aids the user in conducting a post-mortem analysis of program behavior.
- We use an indexed file format (SLOG-2) that uses a preview to select a time of interest and quickly display an interval, without ever needing to read much of the whole file.
Viewing Multiple Scales

Each line represents 1000’s of messages

Detailed view shows opportunities for optimization

1000x zoom
Pros and Cons of this Approach

- Cons:
  - Scalability limits
    - Screen resolution
    - Big log files, although
      - Jumpshot can read SLOG files fast
      - SLOG can be instructed to log few types of events
    - Use for debugging only indirect

- Pros:
  - Portable, since based on MPI profiling interface
  - Works with threads
  - Aids understanding of program behavior
    - Almost always see something unexpected
Looking at MILC in SPEC2007

- Curious amount of All_reduce in initialization – why?
MILC

- The answer, and how
The answer – why

Deep in innermost of quadruply nested loop, an innocent-looking line of code:

If ( i > myrank() ) …

And myrank is a function that calls MPI_Comm_rank

– It actually doesn’t cost that much here, but

– It illustrates that you might not know what your code is doing what you think it is
  – Not a scalability issue (found on small # of processes)
Detecting Consistency Errors in MPI Collective Operations

- The Problem: the specification of MPI_Bcast:
  \[
  \text{MPI\_Bcast(\ buf, count, datatype, root, comm )}
  \]
  requires that
  - \textbf{root} is an integer between 0 and the maximum rank.
  - \textbf{root} is the same on all processes.
  - The message specified by \textbf{buf, count, datatype} has the same \textit{signature} on all processes.

- The first of these is easy to check on each process at the entry to the MPI_Bcast routine.
- The second two are impossible to check locally; they are consistency requirements requiring communication to check.
- There are many varieties of consistency requirements in the MPI collective operations.
Datatype Signatures

- Consistency requirements for messages in MPI (buf, count, datatype) are not on the MPI datatypes themselves, but on the signature of the message:
  - \{type_1, type_2, \ldots\} where type_i is a basic MPI datatype

- So a message described by (buf1, 4, MPI_INT) matches a message described by (buf2, 1, vectype), where vectype was created to be a strided vector of 4 integers.

- For point-to-point operations, datatype signatures don’t have to match exactly (it is OK to receive a short message into a long
Approach

- Use the MPI profiling interface to intercept the collective calls, “borrow” the communicator passed in, and use it to check argument consistency among its processes.
  - For example, process 0 can broadcast its value of root, and each other process can compare with the value it was passed for root.
- For datatype consistency checks, we will communicate hash values of datatype signatures.
- Reference: Falzone, Chan, Lusk, Gropp, “Collective Error
Types of Consistency Checks

- **Call** – checks that all processes have made the same collective call (not MPI_Allreduce on some processes and MPI_Reduce on others).
  - Used in all collective functions
- **Root** – checks that the same value of root was passed on all processes
  - Used in Bcast, Reduce, Gather(v), Scatter(v), Spawn, Spawn_multiple, Connect
- **Datatype** – checks consistency of data arguments
More Types of Consistency Checks

- **MPI_IN_PLACE** – checks whether all process or none of the processes specified MPI_IN_PLACE instead of a buffer.
  - Used in Allgather(v), Allreduce, and Reduce_scatter
- **Local leader and tag** – checks consistency of these arguments
  - Used only in MPI_Intercomm_create
- **High/low** – checks consistency of these arguments
  - Used only in MPI_Intercomm_merge
- **Dims** – checks consistency of these arguments
Still More Types of Consistency Checks

- **Graph** – checks graph consistency
  - Used in Graph_create and Graph_map
- **Amode** – checks file mode argument consistency
  - Used in File_open
- **Size, datarep, flag** – checks consistency of these I/O arguments
  - Used in File_set_size, File_set_automicity, File_preallocate
- **Etype** – checks consistency of this argument
  - Used in File_set_view
- **Order** – checks that split-collective calls are properly ordered
Example Output

- We try to make error output instance specific:
  - Validate Bcast error (Rank 4) – root parameter (4) is inconsistent with rank 0’s (0)
  - Validate Bcast error (Rank 4) – datatype signature is inconsistent with Rank 0’s
  - Validate Barrier (rank 4) – collective call (Barrier) is inconsistent with Rank 0’s (Bcast)
Experiences

- Finding errors
  - Found error in MPICH2 test suite, in which a message with one MPI_INT was allowed to match sizeof(int) MPI_BYTEs.
  - MPICH2 allowed the match, but shouldn’t have.
  - Ran large astrophysics application (FLASH) containing many collective operations
    - Collective calls all in third-party AMR library (Paramesh), but could still be examined through MPI profiling library approach.
    - Found no errors 😊 (☹️)

- Portability, Performance
  - Linux cluster (MPICH2)
  - Blue Gene (IBM’s BG/L MPI)
  - Relative overhead decreases as size of message increases
    - The extra checking messages are much shorter than the real messages
  - Overhead can be relatively large for small messages
    - Opportunities for optimization remain
MPI and Threads
MPI and Threads

- MPI describes parallelism between processes (with separate address spaces)
- Thread parallelism provides a shared-memory model within a process
- OpenMP and Pthreads are common models
  - OpenMP provides convenient features for loop-level parallelism. Threads are created and managed by the compiler, based on user directives.
  - Pthreads provide more complex and dynamic approaches. Threads are created and managed explicitly by the user.
Programming for Multicore

- Almost all chips are multicore these days
- Today’s clusters often comprise multiple CPUs per node sharing memory, and the nodes themselves are connected by a network
- Common options for programming such clusters
  - All MPI
    - Use MPI to communicate between processes both within a node and across nodes
    - MPI implementation internally uses shared memory to communicate within a node
  - MPI + OpenMP
    - Use OpenMP within a node and MPI across nodes
  - MPI + Pthreads
    - Use Pthreads within a node and MPI across nodes
- The latter two approaches are known as “hybrid programming”
MPI’s Four Levels of Thread Safety

- MPI defines four levels of thread safety. These are in the form of commitments the application makes to the MPI implementation.
  - MPI_THREAD_SINGLE: only one thread exists in the application
  - MPI_THREAD_FUNNELED: multithreaded, but only the main thread makes MPI calls (the one that called MPI_Init or MPI_Init_thread)
  - MPI_THREAD_SERIALIZED: multithreaded, but only one thread at a time makes MPI calls
  - MPI_THREAD_MULTIPLE: multithreaded and any thread can make MPI calls at any time (with some restrictions to avoid races – see next slide)

- MPI defines an alternative to MPI_Init
  - MPI_Init_thread(requested, provided)
    - Application indicates what level it needs; MPI implementation returns
Specification of `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 application post conflicting MPI calls — e.g., accessing an info object from one thread and freeing it from another thread.
- User must ensure that collective operations on the same communicator, window, or file handle are correctly ordered among threads.
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
- A fully thread-safe implementation will support MPI_THREAD_MULTIPLE
- A program that calls MPI_Init (instead of MPI_Init_thread) should assume that only MPI_THREAD_SINGLE is supported
- A threaded MPI program that does not call MPI_Init_thread is an incorrect program (common user error we see)
### An Incorrect Program

<table>
<thead>
<tr>
<th>Process 0</th>
<th>Process 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thread 1</td>
<td>MPI_Bcast(comm)</td>
</tr>
<tr>
<td>Thread 2</td>
<td>MPI_Barrier(comm)</td>
</tr>
</tbody>
</table>

- 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.
An implementation must ensure that the above example 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.
The Current Situation

- All MPI implementations support MPITHREAD_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
- Many (but not all) implementations support THREAD_MULTIPLE
  - Hard to implement efficiently though (lock granularity issue)
- “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!
Performance with MPI_THREAD_MULTIPLE

- Thread safety does not come for free
- The implementation must protect certain data structures or parts of code with mutexes or critical sections
- To measure the performance impact, we ran tests to measure communication performance when using multiple threads versus multiple processes
  - Details in our Parallel Computing (journal) paper (2009)
Tests with Multiple Threads versus Processes

Diagram: Two sets of boxes connected by arrows. The left set contains labels 'T', 'T', 'T', 'T' and the right set contains labels 'T', 'T'. Below, the left set contains labels 'P', 'P', 'P', 'P' and the right set contains labels 'P', 'P'.
Concurrent Bandwidth Test on Linux Cluster

MPICH2 version 1.0.5
Open MPI version 1.2.1
Concurrent Bandwidth Test on a single SMP (Sun and IBM)
Concurrent Latency Test on Linux Cluster

MPICH2 version 1.0.5
Open MPI version 1.2.1
Concurrent Latency Test on a single SMP (Sun and IBM)
What MPI’s Thread Safety Means in the Hybrid MPI + 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**
  - Any thread may make an MPI call at any time
Visualizing Hybrid Programs with Jumpshot

- Recent additions to Jumpshot 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 MPI+OpenMP 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 (which uses Pthread locks) 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
Three Platforms for Hybrid Programming Experiments

- **Linux cluster**
  - 24 nodes, each with two Opteron dual-core processors, 2.8 Ghz each
  - Intel 9.1 Fortran compiler
  - MPICH2–1.0.6, which has MPI_THREAD_MULTIPLE
  - Multiple networks; we used GigE

- **IBM Blue Gene/P**
  - 40,960 nodes, each consisting of four PowerPC 850 MHz cores
  - XLF 11.1 Fortran cross-compiler
  - IBM’s MPI V1R1M2 (based on MPICH2), has MPI_THREAD_MULTIPLE
  - 3D Torus and tree networks

- **SiCortex SC5832**
  - 972 nodes, each consisting of six MIPS 500 MHz cores
  - Pathscale 3.0.99 Fortran cross-compiler
  - SiCortex MPI implementation based on MPICH2, has MPI_THREAD_FUNNELED
  - Kautz graph network
Experiments

- **Basic**
  - Proved that necessary assumptions for our tools hold
    - OpenMP threads are Pthreads
    - Thread id’s are reused

- **NAS Parallel Benchmarks**
  - NPB-MZ-MPI, version 3.1
  - Both BT and SP
  - Two different sizes (W and B)
  - Two different modes (“MPI everywhere” and OpenMP/MPI)
    - With four nodes on each machine

- Demonstrated satisfying level of portability of programs and tools across three quite different hardware/software environments
It Might Not Be Doing What You Think

- An early run:

- Nasty interaction between the environment variables OMP_NUM_THREADS and NPB_MAX_THREADS
More Like What You Expect

- BT class B on 4 BG/P nodes, using OpenMP on each node
MPI Everywhere

- BT class B on 4 BG/P nodes, using 16 MPI processes
## Observations on Experiments

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Cluster</th>
<th>BG/P</th>
<th>SiCortex</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bt-mz.W.16x1</td>
<td>1.84</td>
<td>9.46</td>
<td>20.60</td>
</tr>
<tr>
<td>Bt-mz-W.4x4</td>
<td>0.82</td>
<td>3.74</td>
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<td>0.78</td>
<td>3.00</td>
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<tr>
<td>Sp-mz.B.4x6</td>
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<td>211.78</td>
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- Time in seconds
- On the small version of BT (W), hybrid was better
- For SP and size B problems, MPI everywhere is better
- On SiCortex, more processes or threads are better than fewer
Observations

- This particular benchmark has been studied much more deeply elsewhere
  - Rolf Rabenseifner, “Hybrid parallel programming on HPC platforms,” Proceedings of EWOMP’03.
- Adding “hybridness” to a well-tuned MPI application is not going to speed it up. So this NPB study doesn’t tell us much.
- More work is needed to understand the behavior of hybrid programs and what is needed for future application development.
One-Sided Communication
One-Sided Communication

- A process can directly access another process’s memory (with a function call)
- Three data transfer functions
  - \texttt{MPI\_Put}, \texttt{MPI\_Get}, \texttt{MPI\_Accumulate}
  \begin{center}
  \begin{tikzpicture}
  \node (MPI_Put) {\texttt{MPI\_Put}};
  \node [right of=MPI_Put] (MPI_Get) {\texttt{MPI\_Get}};
  \draw [->] (MPI_Put) -- (MPI_Get);
  \end{tikzpicture}
  \end{center}
- Three synchronization methods
  - \texttt{MPI\_Win\_fence}
  - \texttt{MPI\_Win\_post/start/complete/wait}
  - \texttt{MPI\_Win\_lock/unlock}
Remote Memory Access Windows and Window Objects

= address spaces

= window object
Window Creation

- **MPI_Win_create** exposes local memory to RMA operation by other processes in a communicator
  - Collective operation
  - Creates window object

  ```
  MPI_Win_Create(base, size, disp_unit, info, comm, win)
  ```

- **MPI_Win_free** deallocates window object
Fence Synchronization

- `MPI_Win_fence(win)` is collective over the communicator associated with the window object
- (The numbers in parentheses refer to the target ranks)

Process 0

- `MPI_Win_fence(win)`
- `MPI_Put(1)`
- `MPI_Get(1)`
- `MPI_Win_fence(win)`

Process 1

- `MPI_Win_fence(win)`
- `MPI_Put(0)`
- `MPI_Get(0)`
- `MPI_Win_fence(win)`
Post-Start-Complete-Wait Synchronization

- Scalable: Only the communicating processes need to synchronize
- (The numbers in parentheses refer to the target ranks)

Process 0

MPI_Win_start(1)
MPI_Put(1)
MPI_Get(1)
MPI_Win_complete(1)

Process 1

MPI_Win_post(0,2)
MPI_Win_wait(0,2)

Process 2

MPI_Win_start(1)
MPI_Put(1)
MPI_Get(1)
MPI_Win_complete(1)
Lock-Unlock Synchronization

- “Passive” target: The target process does not make any synchronization call
- (The numbers in parentheses refer to the target ranks)

Process 0

- MPI_Win_create
- MPI_Win_lock(1, shared)
- MPI_Put(1)
- MPI_Get(1)
- MPI_Win_unlock(1)
- MPI_Win_free

Process 1

- MPI_Win_create
- MPI_Win_lock(1, shared)
- MPI_Put(1)
- MPI_Get(1)
- MPI_Win_unlock(1)
- MPI_Win_free

Process 2

- MPI_Win_create
- MPI_Win_lock(1, shared)
- MPI_Put(1)
- MPI_Get(1)
- MPI_Win_unlock(1)
- MPI_Win_free
Performance Tests

- “Halo” exchange or ghost-cell exchange operation
  - Each process exchanges data with its nearest neighbors
  - Part of mpptest benchmark
  - One-sided version uses all 3 synchronization methods

- Ran on
  - Sun Fire SMP at Univ. of Aachen, Germany
  - IBM p655+ SMP at San Diego Supercomputer Center
One-Sided Communication on Sun SMP with Sun MPI

Halo Performance on Sun

Bytes

uSec

sendrecv-8
psendrecv-8
putall-8
putpscwalloc-8
putlockshared-8
putlocksharednb-8
One-Sided Communication on IBM SMP with IBM MPI
MPI at Exascale

Rajeev Thakur
Mathematics and Computer Science Division
Argonne National Laboratory
MPI on the Largest Machines Today

- Systems with the largest core counts in June 2010 Top500 list
  - Juelich BG/P 294,912 cores
  - Oak Ridge Cray XT5 224,162 cores
  - LLNL BG/L 212,992 cores
  - Argonne BG/P 163,840 cores
  - LLNL BG/P (Dawn) 147,456 cores
  (All these systems run MPICH2-based MPI implementations)

- In a couple of years, we will have systems with more than a million cores

- For example, in 2012, the Sequoia machine at Livermore will
Future Extreme Scale Platforms

- Hundreds of thousands of “nodes”
- Each node has large numbers of cores, including
  - Regular CPUs and accelerators (e.g., GPUs)
Multiple Cores Per Node

- All Large Cores
- Mixed Large and Small Cores
- Many small cores
- All small cores

Memory Floating Point Cores

+ 3D Stacked Memory

SDRAM

Different Classes of Chips
- Home
- Games/Graphics
- Business
- Scientific
Scaling MPI to Exascale

- MPI already runs on the largest systems today at ~300,000 cores

- What would it take to scale MPI to exascale systems with millions of cores?

- On exascale, MPI is likely to be used as part of a “hybrid programming” model (MPI+X), much more so than it is today
  - MPI being used to communicate between “address spaces”
  - With some other “shared-memory” programming model (OpenMP, UPC, CUDA, OpenCL) for programming within an address space

- How can MPI support efficient “hybrid” programming on exascale systems?
Scaling MPI to Exascale

- Although the original designers of MPI were not thinking of exascale, MPI was always intended and designed with scalability in mind. For example:
  - A design goal was to enable implementations that maintain very little global state per process
  - Another design goal was to require very little memory management within MPI (all memory for communication can be in user space)
  - MPI defines many operations as collective (called by a group of processes), which enables them to be implemented scalably and efficiently

- Nonetheless, some parts of the MPI specification may need to be fixed for exascale
  - Being addressed by the MPI Forum in MPI-3
Factors Affecting MPI Scalability

- Performance and memory consumption
- A nonscalable MPI function is one whose time or memory consumption per process increase linearly (or worse) with the total number of processes (all else being equal)
- For example
  - If memory consumption of MPI_Comm_dup increases linearly with the no. of processes, it is not scalable
  - If time taken by MPI_Comm_spawn increases linearly or more with the no. of processes being spawned, it indicates a nonscalable implementation of the function
- Such examples need to be identified and fixed (in the specification and in implementations)
- The goal should be to use constructs that require only constant space per process
Requirements of a message-passing library at extreme scale

- No $O(n\text{procs})$ consumption of resources (memory, network connections) per process
- Resilient and fault tolerant
- Efficient support for hybrid programming (multithreaded communication)
- Good performance over the entire range of message sizes and all functions, not just latency and bandwidth benchmarks
- Fewer performance surprises (in implementations)

These issues are being addressed by the MPI Forum for MPI–3 and by MPI implementations
Scalability Issues in the MPI Specification

- Some functions take parameters that grow linearly with number of processes
- E.g., irregular (or “v”) version of collectives such as MPI_Gatherv
- Extreme case: MPI_Alltoallw takes six such arrays
  - On a million processes, that requires 24 MB on each process
- On low-frequency cores, even scanning through large arrays takes time (see next slide)

- Solution: The MPI Forum is considering a proposal to define sparse, neighborhood collectives that could be used instead of irregular collectives
Zero-byte MPI_Alltoallv time on BG/P

- This is just the time to scan the parameter array to determine it is all 0 bytes. No communication performed.
Scalability Issues in the MPI Specification

- **Graph Topology**
  - In MPI 2.1 and earlier, requires the entire graph to be specified on each process
  - Already fixed in MPI 2.2 – new distributed graph topology functions

- **One-sided communication**
  - Synchronization functions turn out to be expensive
  - Being addressed by RMA working group of MPI-3

- **Representation of process ranks**
  - Explicit representation of process ranks in some functions, such as MPI_Group_incl and MPI_Group_excl
  - Concise representations should be considered
Scalability Issues in the MPI Specification

- **All-to-all communication**
  - Not a scalable communication pattern
  - Applications may need to consider newer algorithms that do not require all-to-all

- **Fault tolerance**
  - Large component counts will result in frequent failures
  - Greater resilience needed from all components of the software stack
  - MPI can return error codes, but need more support than that
  - Being addressed in the fault tolerance group of MPI-3
MPI Implementation Scalability

- MPI implementations must pay attention to two aspects as the number of processes is increased:
  - memory consumption of any function, and
  - performance of all collective functions
    - Not just collective communication functions that are commonly optimized
    - Also functions such as MPI_Init and MPI_Comm_split
Process Mappings

- MPI communicators maintain mapping from ranks to processor ids
- This mapping is often a table of $O(nprocs)$ size in the communicator
- Need to explore more memory-efficient mappings, at least for common cases
- More systematic approaches to compact representations of permutations (research problem)
Communicator Memory Consumption

- NEK5000 is a well-known fluid dynamics code developed by Paul Fischer and colleagues at Argonne
- When they first tried to scale this code on the BG/P, it failed on as little as 8K processes because the MPI library ran out of communicator memory
- NEK5000 calls MPI_Comm_dup about 64 times (because it makes calls to libraries)
- 64 is not a large number, and, in any case, MPI_Comm_dup should not consume $O(nprocs)$ memory (it doesn’t in MPICH2)
- We ran an experiment to see what was going on...
Communicator Memory Consumption with original MPI on BG/P

- Run MPI_Comm_dup in a loop until it fails. Vary the number of processes.
What was going on --- and the fix

- The default MPI_Comm_dup in IBM’s MPI was allocating memory to store process mapping info for optimizing future calls to collective communication (Alltoall)
- Allocated memory was growing linearly with system size
- One could disable the memory allocation with an environment variable, but that would also disable the collective optimizations

- On further investigation we found that they really only needed one buffer per thread instead of one buffer per new communicator

- Since there are only four threads on the BG/P, we fixed the problem by allocating a fixed buffer pool within MPI
Communicator Memory Consumption Fixed

- NEK5000 code failed on BG/P at large scale because MPI ran out of communicator memory. We fixed the problem by using a fixed buffer pool within MPI and provided a patch to IBM.
Using a buffer pool enables all collective optimizations and takes up only a small amount of memory.
Scalability of MPI_Init

- Cluster with 8 cores per node. TCP/IP across nodes
- Setting up all connections at Init time is too expensive at large scale; must be done on demand as needed

![Graph showing eager versus lazy connection MPI_Init time](image)
Scalable Algorithms for Collective Communication

- MPI implementations typically use
  - $O(\log p)$ algorithms for short messages (binomial tree)
  - $O(m)$ algorithms, where $m$=message size, for large messages
    - E.g., bcast implemented as scatter + allgather
- $O(\log p)$ algorithms can still be used on a million processors for short messages
- However, $O(m)$ algorithms for large messages may not scale, as the message size in the allgather phase can get very small
  - E.g., for a 1 MB bcast on a million processes, the allgather phase involves 1 byte messages
- Hybrid algorithms that do logarithmic bcast to a subset of nodes, followed by scatter/allgather may be needed
- Topology-aware pipelined algorithms may be needed
- Use network hardware for broadcast/combine
Enabling Hybrid Programming

- MPI is good at moving data between address spaces
- Within an address space, MPI can interoperate with other “shared memory” programming models
- Useful on future machines that will have limited memory per core
- (MPI + X) Model: MPI across address spaces, X within an address space
- Examples:
  - MPI + OpenMP
  - MPI + UPC/CAF (here UPC/CAF address space could span multiple nodes)
  - MPI + CUDA/OpenCL on GPU-accelerated systems
- Precise thread-safety semantics of MPI enable such hybrid models
MPI-3 Hybrid Proposal on Endpoints

- In MPI today, each process has one communication endpoint (rank in MPI_COMM_WORLD)
- Multiple threads communicate through that one endpoint, requiring the implementation to do use locks etc., which are expensive
- This proposal (originally by Marc Snir) allows a process to have multiple endpoints
- Threads within a process attach to different endpoints and communicate through those endpoints as if they are separate ranks
- The MPI implementation can avoid using locks if each thread communicates on a separate endpoint
MPI-3 Hybrid Proposal on Endpoints

- Today, each MPI process has one communication endpoint (rank in MPI_COMM_WORLD)
- Multiple threads communicate through that one endpoint, requiring the implementation to do use locks etc. (expensive)

**Current MPI Design**

Separate address spaces for each endpoint
MPI-3 Hybrid Proposal on Endpoints

- The proposal is to allow a process to have multiple endpoints
- Threads within a process attach to different endpoints and communicate through those endpoints as if they are separate ranks
- The MPI implementation can avoid using locks if each thread

### Proposed MPI Design

![Diagram of MPI-3 Hybrid Proposal on Endpoints]

Multiple endpoints are mapped in the same address space
Recent Efforts of the MPI Forum
MPI Standard Timeline

- MPI-1 (1994)
  - Basic point-to-point communication, collectives, datatypes, etc

- MPI-2 (1997)
  - Added parallel I/O, RMA, dynamic processes, C++ bindings, etc

- ---- Stable for 10 years ----

- MPI-2.1 (2008)
  - Minor clarifications and bug fixes to MPI-2

- MPI-2.2 (2009)
  - Today’s official standard
  - Small updates and additions to MPI 2.1. Backward compatible

- MPI-3 (in progress, expected late 2011)
  - Major new features and additions to extend MPI to exascale
  - Organized into several working groups
MPI 2.2 (Today’s Official MPI Standard)

- Led by Bill Gropp
- Officially approved by the MPI Forum at the Sept 2009 meeting
- Small updates to the standard
  - Does not break backward compatibility
- Spec can be downloaded from the MPI Forum web site [www.mpi-forum.org](http://www.mpi-forum.org)
- Also available for purchase as a book from [https://fs.hlrs.de/projects/par/mpi/mpi22/](https://fs.hlrs.de/projects/par/mpi/mpi22/)
- Supported by MPICH2 1.2
New Features in MPI 2.2

- Scalable graph topology interface
  - Existing interface requires the entire graph to be specified on all processes, which requires too much memory on large numbers of processes
  - New functions allow the graph to be specified in a distributed fashion (MPI_Dist_graph_create, MPI_Dist_graph_create_adjacent)

- A local reduction function
  - MPI_Reduce_local(inbuf, inoutbuf, count, datatype, op)
  - Needed for libraries to implement user-defined reductions

- MPI_Comm_create extended to enable creation of multiple disjoint communicators

- Regular (non-vector) version of MPI_Reduce_scatter called MPI_Reduce_scatter_block
New Features in MPI 2.2

- MPI_IN_PLACE option added to MPI_Alltoall, Alltoally, Alltoallw, and Exscan
- The restriction on the user not being allowed to access the contents of the buffer passed to MPI_Isend before the send is completed by a test or wait has been lifted
- New C99 datatypes (MPI_INT32_T, MPI_C_DOUBLE_COMPLEX, etc) and MPI_AINT/ MPI_OFFSET
New Features being considered in MPI-3

- **Note:** All these are still under discussion in the Forum and not final

- Support for hybrid programming (Lead: Pavan Balaji, Argonne)
  - Extend MPI to allow multiple communication endpoints per process
  - Helper threads: application sharing threads with the implementation

- Improved RMA (Leads: Bill Gropp, UIUC, and Rajeev Thakur, Argonne)
  - Fix the limitations of MPI-2 RMA
  - New compare-and-swap, fetch-and-add functions
  - Collective window memory allocation
  - Test for completion of individual operations
  - Others...
New Features being considered in MPI-3

- New collectives (Lead: Torsten Hoefler, UIUC)
  - Nonblocking collectives already voted in (MPI_Ibcast, MPI_Ireduce, etc)
  - Sparse, neighborhood collectives being considered as alternatives to irregular collectives that take vector arguments

- Fault tolerance (Lead: Rich Graham, Oak Ridge)
  - Detecting when a process has failed; agreeing that a process has failed
  - Rebuilding communicator when a process fails or allowing it to continue in a degraded state
  - Timeouts for dynamic processes (connect–accept)
  - Piggybacking messages to enable application-level fault tolerance
New Features being considered in MPI-3

- **Fortran 2008 bindings (Lead: Craig Rasmussen, LANL)**
  - Full and better quality argument checking with individual handles
  - Support for choice arguments, similar to (void *) in C
  - Passing array subsections to nonblocking functions
  - Many other issues

- **Better support for Tools (Lead: Martin Schulz, LLNL)**
  - MPIT performance interface to query performance information internal to an implementation
  - Standardizing an interface for parallel debuggers
MPI Forum Mailing Lists and Archives

- Web site: http://lists.mpi-forum.org/

- Lists
  - mpi-forum
  - mpi-22, mpi-3
  - mpi3-coll
  - mpi3-rma
  - mpi3-ft
  - mpi3-fortran
  - mpi3-tools
  - mpi3-hybridpm

- Further info: http://meetings.mpi-forum.org/
- Wiki: https://svn.mpi-forum.org/trac/mpi-forum-web/wiki
What are we doing in MPICH2
Goals of the MPICH2 project

- Be the MPI implementation of choice for the highest-end parallel machines
  - 7 of the top 10 machines in the June 2010 Top500 list use MPICH2-based implementations

- Carry out the research and development needed to scale MPI to exascale
  - Optimizations to reduce memory consumption
  - Fault tolerance
  - Efficient multithreaded support for hybrid programming
  - Performance scalability

- Work with the MPI Forum on standardization and early prototyping of new features
MPICH2 collaboration with vendors

- Enable vendors to provide high-performance MPI implementations on the leading machines of the future

- Collaboration with IBM on MPI for the Blue Gene/Q
  - Aggressive multithreaded optimizations for high concurrent message rates
  - Recent publications in Cluster 2010 and EuroMPI 2010

- Collaboration with Cray for MPI on their next-generation interconnect (Gemini)

- Collaboration with UIUC on MPICH2 over LAPI for Blue Waters

- Continued collaboration with Intel, Microsoft, and Ohio State
Conclusions

- MPI has succeeded because
  - features are orthogonal (complexity is the product of the number of features, not routines)
  - complex programs are no harder than easy ones
  - open process for defining MPI led to a solid design
  - programmer can control memory motion and program for locality (critical in high-performance computing)
  - precise thread-safety specification has enabled hybrid programming

- MPI is ready for scaling to extreme scale systems with millions of cores barring a few issues that can be (and are being) fixed by the MPI Forum and by MPI implementations
The MPI Standard (1 & 2)
MPI 2.2 Standard
Tutorial Material on MPI, MPI-2

Using MPI
Portable Parallel Programming with the Message Passing Interface, Second Edition

William Gropp
Ewing Lusk
Anthony Skjellum

Using MPI 2
Advanced Features of the Message Passing Interface

William Gropp
Ewing Lusk
Rajeev Thakur

http://www.mcs.anl.gov/mpi/{usingmpi,usingmpi2}
ADLB: The Asynchronous Dynamic Load-Balancing Library

An approach to extreme scalability with an extremely simple programming model (for some applications)
Outline

- Introduction
  - Simple programming models
  - Load balancing
  - Scalability problems

- ADLB
  - What it is
  - How it works
  - The API

- Example applications
  - Fun – Sudoku solver
  - Serious – GFMC: complex Monte Carlo physics application
  - Useful – batcher: running independent jobs
Two Classes of Parallel Programming Models

- **Data Parallelism**
  - Parallelism arises from the fact that physics is largely local
  - Same operations carried out on different data representing different patches of space
  - Communication usually necessary between patches (local)
    - global (collective) communication sometimes also needed
  - Load balancing sometimes needed

- **Task Parallelism**
  - Work to be done consists of largely independent tasks, perhaps not all of the same type
  - Little or no communication between tasks
  - Usually needs a separate “master” task for scheduling
  - Load balancing essential
Load Balancing

- **Definition:** the assignment (scheduling) of tasks (code + data) to processes so as to minimize the total idle times of processes

- **Static load balancing**
  - all tasks are known in advance and pre-assigned to processes
  - works well if all tasks take the same amount of time
  - requires no coordination process

- **Dynamic load balancing**
  - tasks are assigned to processes by coordinating process when processes become available
  - Requires communication between manager and worker processes
  - Tasks may create additional tasks
  - Tasks may be quite different from one another
Generic Master/Slave Algorithm

- Easily implemented in MPI
- Solves some problems
  - implements dynamic load balancing
  - termination
  - dynamic task creation
  - can implement workflow structure of tasks
- Scalability problems
  - Master can become a communication bottleneck (granularity dependent)
  - Memory can become a bottleneck (depends on task description size)
The ADLB Vision

- No explicit master for load balancing; slaves make calls to ADLB library; those subroutines access local and remote data structures (remote ones via MPI).
- Simple Put/Get interface from application code to distributed work queue hides MPI calls
  - Advantage: multiple applications may benefit
  - Wrinkle: variable-size work units, in Fortran, introduce some complexity in memory management
- Proactive load balancing in background
  - Advantage: application never delayed by search for work from other slaves
  - Wrinkle: scalable work-stealing algorithms not obvious
The ADLB Model (no master)

- Doesn’t really change algorithms in slaves
- Not a new idea (e.g. Linda)
- But need scalable, portable, distributed implementation of shared work queue
  - MPI complexity hidden here
API for a Simple Programming Model

- Basic calls
  - ADLB_Init( num_servers, am_server, app_comm)
  - ADLB_Server()
  - ADLB_Put( type, priority, len, buf, answer_dest )
  - ADLB_Reserve( req_types, handle, len, type, prio, answer_dest)
  - ADLB_Ireserve( ... )
  - ADLB_Get_Reserved( handle, buffer )
  - ADLB_Set_Done()
  - ADLB_Finalize()

- A few others, for tuning and debugging
  - ADLB_{Begin,End}_Batch_Put()
  - Getting performance statistics with ADLB_Get_info(key)
API Notes

- Return codes (defined constants)
  - ADLB_SUCCESS
  - ADLB_NO_MORE_WORK
  - ADLB_DONE_BY_EXHAUSTION
  - ADLB_NO_CURRENT_WORK (for ADLB_Ireserve)

- Batch puts are for inserting work units that share a large proportion of their data

- Types, answer_rank, reserve_rank can be used to implement some common patterns
  - Sending a message
  - Decomposing a task into subtasks
  - Maybe should be built into API
How It Works

Application Processes

ADLB Servers

put/get
The ADLB Server Logic

- **Main loop:**
  - MPI_Iprobe for message in busy loop
  - MPI_Recv message
  - Process according to type
    - Update status vector of work stored on remote servers
    - Manage work queue and request queue
    - (may involve posting MPI_Isends to isend queue)
  - MPI_Test all requests in isend queue
  - Return to top of loop

- **The status vector replaces single master or shared memory**
  - Circulates every .1 second at high priority
  - Multiple ways to achieve priority
A Tutorial Example: Sudoku

1 2 3 4 5 6 7 8 9

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Parallel Sudoku Solver with ADLB

Program:
if (rank = 0)
    ADLB_Put initial board
    ADLB_Get board (Reserve+Get)
while success (else done)
    ooh
    find first blank square
    if failure (problem solved!)
        print solution
        ADLB_Set_Done
    else
        for each valid value
            set blank square to value
            ADLB_Put new board
            ADLB_Get board

Work unit = partially completed “board”
How it Works

- After initial Put, all processes execute same loop (no master)
Optimizing Within the ADLB Framework

- Can embed smarter strategies in this algorithm
  - ooh = “optional optimization here”, to fill in more squares
  - Even so, potentially a lot of work units for ADLB to manage

- Can use priorities to address this problem
  - On ADLB_Put, set priority to the number of filled squares
  - This will guide depth-first search while ensuring that there is enough work to go around
    - How one would do it sequentially

- Exhaustion automatically detected by ADLB (e.g., proof that there is only one solution, or the case of an invalid input board)
Green’s Function Monte Carlo - the defining application

- Green’s Function Monte Carlo -- the “gold standard” for ab initio calculations in nuclear physics at Argonne (Steve Pieper, PHY)
- A non-trivial master/slave algorithm, with assorted work types and priorities; multiple processes create work; large work units
- Has scaled to 2000 processors on BG/L a little over four years ago, then hit scalability wall.
- Need to get to 10’s of thousands of processors at least, in order to carry out calculations on $^{12}$C, an explicit goal of the UNEDF SciDAC project.
- The algorithm has had to become even more complex, with more types and dependencies among work units, together with smaller work units
- Wanted to maintain master/slave structure of physics code
Experiments with GFMC/ADLB on BG/P

- Using GFMC to compute the binding energy of 14 neutrons in an artificial well ("neutron drop" = teeny-weeny neutron star)
- A weak scaling experiment

<table>
<thead>
<tr>
<th>BG/P cores</th>
<th>ADLB Servers</th>
<th>Configs</th>
<th>Time (min.)</th>
<th>Efficiency (incl. serv.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4K</td>
<td>130</td>
<td>20</td>
<td>38.1</td>
<td>93.8%</td>
</tr>
<tr>
<td>8K</td>
<td>230</td>
<td>40</td>
<td>38.2</td>
<td>93.7%</td>
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<tr>
<td>16K</td>
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<td>39.6</td>
<td>89.8%</td>
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<tr>
<td>32K</td>
<td>905</td>
<td>160</td>
<td>44.2</td>
<td>80.4%</td>
</tr>
</tbody>
</table>

- Recent work: “micro-parallelization” needed for $^{12}$C, OpenMP in GFMC.
  - a successful example of hybrid programming, with ADLB + MPI + OpenMP
Progress with GFMC

Efficiency = compute_time/wall_time – 25 Feb 2010

Number of nodes (4 OpenMP cores per node)

Efficiency in %

12C ADLB+GFMC

Oct 2009
Jun 2009
Feb 2009
Another Physics Application - Parameter Sweep

- Luminescent solar concentrators
  - Stationary, no moving parts
  - Operate efficiently under diffuse light conditions (northern climates)
- Inexpensive collector, concentrate light on high-performance solar cell
- This application was parallelized by “non-parallel” programmers using ADLB without learning MPI
The “Batcher”

- Simple but useful
- Input is a file of Unix command lines
- ADLB worker processes execute each one with the Unix “system” call
  - therefore need this call available on each node
    - true for Unix clusters
    - problematic on machines with custom compute-node kernels
- 100-line program, mainly error-checking
ADLB Uses Multiple MPI Features

- ADLB_Init returns separate application communicator, so application can use MPI for its own purposes if it needs to.
- Servers are in MPI_Iprobe loop for responsiveness.
- MPI_Datatype for some complex, structured messages (status)
- Servers use nonblocking sends and receives, maintain queue of active MPI_Request objects.
- Queue is traversed and each request kicked with MPI_Test each time through loop; could use MPI_Testany. No MPI_Wait.
- Client side uses MPI_Ssend to implement ADLB_Put in order to conserve memory on servers, MPI_Send for other actions.
- Servers respond to requests with MPI_Rsend since MPI_Irecvs are known to be posted by clients before requests.
- MPI provides portability: laptop, Linux cluster, SiCortex, BG/P
- MPI profiling library is used to understand application/ADLB behavior.
Getting ADLB

- Web site is [http://www.cs.mtsu.edu/~rbutler/adlb](http://www.cs.mtsu.edu/~rbutler/adlb)
- To download adlb:
  - `svn co http://svn.cs.mtsu.edu/svn/adlbm/trunk adlbm`
- What you get:
  - source code
  - configure script and Makefile
  - README, with API documentation
  - Examples
    - Sudoku
    - Batcher
      - Batcher README
    - Traveling Salesman Problem
- To run your application
  - configure, make to build ADLB library
  - Compile your application with mpicc, use Makefile as example
  - Run with mpiexec
- Problems/complaints/kudos to {lusk,rbutler}@mcs.anl.gov
Future Directions

- **API design**
  - Some higher-level function calls might be useful
  - User community will generate these

- **Implementations**
  - The one-sided version
    - implemented
    - single server to coordinate matching of requests to work units
    - stores work units on client processes
    - Uses MPI_Put/Get (passive target) to move work
    - Hit scalability wall for GFMC at about 8000 processes
  - The thread version
    - uses separate thread on each client; no servers
    - the original plan
    - maybe for BG/Q, where there are more threads per node
    - not re-implemented (yet)
Where We Are Now

- ADLB is a research project working its way toward being useful general-purpose software.
- More users sought, especially those with more straightforward applications than GFMC!
- Its point is to explore whether extreme scalability in an application can be achieved without extreme complexity in application code.
Conclusions

- The Philosophical Accomplishment: Scalability need not come at the expense of complexity

- The Practical Accomplishment: Maybe this can accelerate the development of your application.
The End