Asynchronous Parallel Methods

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- What's the problem?
- What is an asynchronous method?
- Reducing synchronisation in existing models

The Problem

- Synchronisations often essential for program correctness
 - waiting for an MPI receive to complete before reading from buffer
 - barriers at the end of an OpenMP parallel loop
 - Dest the second
 - But they cost time
 - and slow down the calculation
- Cost is usually not the synchronisation operation itself
 - it is waiting for other tasks to catch up with each other
 - all calculations have some load imbalance from random fluctuations
 - a real problem as we increase the number of cores
- Try to reduce synchronisation
 - and let things happen in their "natural" order

Reference



- See:
 - "The Case of the Missing Supercomputer Performance: Achieving Optimal Performance on the 8,192 Processors of ASCI Q"
 - Fabrizio Petrini, Darren J. Kerbyson, Scott Pakin
 - http://dx.doi.org/10.1145/1048935.1050204
 - "[W]hen you have eliminated the impossible, whatever remains, however improbable, must be the truth."
 - Sherlock Holmes, Sign of Four, Sir Arthur Conan Doyle



An example

 "Although SAGE [the application] spends half of its time in allreduce (at 4,096 processors), making allreduce seven times faster leads to a negligible performance improvement."







- Simulation parameters (simplified)
 - total number of pixels: L x L
 - number of processors: P
 - decomposition: $P \ge 1$ (1D) or $\sqrt{P} \ge \sqrt{P}$ (2D)
- System properties (simplified)
 - floating-point operations per second: f
 - message-passing latency: T_I
 - message-passing bandwidth: B

Update

$$new_{i,j} = 0.25 * \left(old_{i-1,j} + old_{i+1,j} + old_{i,j-1} + old_{i,j+1} - edge_{i,j}\right)$$

- 5 floating-point operations per pixel

Delta calculation

$$delta = delta + \left(new_{i,j} - old_{i,j}\right) * \left(new_{i,j} - old_{i,j}\right)$$

- 3 flops per pixel

• Time taken: Time = flops _ per _ pixel *
$$\frac{N_{pixel}}{f}$$

 $N_{pixel} = \frac{L^2}{P}$

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Collectives



- Reduce frequency of calculation by a factor X
 - e.g. trade more calculation for fewer synchronisations

loop over iterations: update arrays; compute local delta; compute global delta using allreduce; stop if less than tolerance value; end loop loop over iterations: update arrays; every X iterations: local delta; global delta; can we stop?;

end loop

Possible because array updates independent of global values

- may not be true for, e.g., Conjugate Gradient
- can use different algorithms, e.g. Chebyshev iteration
- again, more iterations but less synchronisation

Barriers



• (Almost) never required for MPI program correctness

- Why?
 - because collectives do the appropriate synchronisation
 - because MPI_Recv is synchronous

Halo swapping







• Do not impose unnecessary ordering of messages

loop over sources:
 receive value from
 particular source;
end loop

loop over sources:
 receive value from
 any source;
end loop

- loop now just counts the correct number of messages
- Alternative
 - first issue a separate non-blocking receive for each source
 - then issue a single Waitall



• Do not impose unnecessary ordering of messages

loop over directions: send up; recv down; send down; recv up; end loop loop over directions:
 isend up; irecv down;
 isend down; irecv up;
end loop
wait on all requests;

• Extensions

- can now overlap communications with core calculation
- only need to wait for receives before non-core calculation
- wait for sends to complete before starting next core calculation

Overlapping





wait for completion of non-blocking sends/recvs complete calculation at the four edges

Halos of Depth D

- Use less frequent communication
 - smaller number of larger messages; increased computation



loop d=D:1:-1

Swap depth D every D iterations



- Need diagonal communications
 - and must swap halos of depth D-1 on edge(i,j)

• Standard method: run this code every iteration

```
MPI_Irecv(..., procup, ..., &reqs[0]);
MPI_Irecv(..., procdn, ..., &reqs[1]);
MPI_Isend(..., procdn, ..., &reqs[2]);
MPI_Isend(..., procup, ..., &reqs[3]);
MPI_Waitall(4, reqs, statuses);
```

• Persistent comms: setup once

MPI_Recv_init(..., procup, ..., &reqs[0]); MPI_Recv_init(..., procdn, ..., &reqs[1]); MPI_Send_init(..., procdn, &reqs[2]); MPI_Send_init(..., procup, ..., &reqs[3]);

• Every iteration:

```
MPI_Startall(4, reqs);
```

- Warning
 - message ordering not guaranteed to be preserved