Outline

Part 1
- Introduction
  - Review of hardware & parallel programming models
  - NERSC NECAP
- Principles of High Performance Parallel Programming (HPPP)
- EMGeo: basic
- EMGeo: intermediate

Part 2
- Know MIC and programming model
- Multi-level parallelism: Nested OpenMP

Part 3
- PARSEC
- EMGeo: advanced
- Conclusions
About the Presenter: Jeongnim Kim, PhD

- Sr. HPC application Engineer at Joe Curley's MICRO (MIC Ramp Organization) group; working for code modernization and optimization on Xeon and Xeon Phi™

- Has been active in computational materials science and HPC since 1993
  - Used most of parallel computing platforms at DOE and NSF HPC centers: Intel Paragon, Cray T3D/T3E, SGI Origin 2000, Intel Itanium, IBM Power 3-7, Cray XT/XE/XK/XC, and IBM Blue Gene Q
  - Distributed programming on Intel Paragon (1994); OpenMP programming on SGI Origin (1998)

- Prior to joining Intel in April 2014
  - Worked for Oak Ridge National Laboratory (ORNL) and National Center for Supercomputing Applications and Materials Computation Center, University of Illinois, Urbana-Champaign
  - Developed QMCPACK and led Quantum Monte Carlo collaboration between ORNL, ANL, LLNL, Sandia and UI

- PhD in condensed matter theory from the Ohio State University, USA, and a BS in Physics from Korea Advanced Institute of Science and Technology, Korea
How to exploit OpenMP* for high-performance parallel applications

Someone said

- Shared-memory programming models on multi- and many-core processors are critical. You must hybridize your application!

- OpenMP* is so easy. All you have to do is to find loops and put `OMP parallel do` over the loops.

- OpenMP* 4.x let you express your intention of vectorization of the loops and compilers can vectorize them.

- MKL comes with threaded numerical libraries. Use threads with GEMM or FFT.

Then, you are thinking

“I tried OpenMP but the performance is much worse than MPI. Where is the performance?”
Distributed-shared-memory programming (a.k.a., hybrid programming)

“I tried OpenMP but the performance is much worse than MPI. Why bother?”

A good question! But,

- The laws of physics say otherwise: finite electron velocity, limited parallel channels, multiple hops, ....

- Just ask how many instructions are needed to execute a put or get. E.g., a simple send/recv = memory→[MPI buffer]→ memory.

- Moving data with MPI must be more expensive than memory to cache.

So, what is going on?
This workshop aims to

- Refresh your knowledge of hardware, software and parallel programming
- Remind you of Parallel computing 101
- Use NESAP codes to discuss processes to exploit modern hardware
- Introduce advanced OpenMP* concepts and techniques
- Promote code design and thinking out of box
Disclaimers

- OpenMP* (MPI) is selected as the de-facto standard for shared (distributed) parallel programming model.
- Processes based on the experiences with numerous HPC applications.
- Materials using MPI/Fortran applications chosen by NERSC
- Each process will be marked by the target developers

<table>
<thead>
<tr>
<th>Dev0</th>
<th>New member of the team; cannot find code documentations (or hidden) and everyone is busy.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dev1</td>
<td>Computer scientist or engineer; know nothing about the application (science); have to work with the “domain scientists”.</td>
</tr>
<tr>
<td>Dev2</td>
<td>Designed and wrote the code and “invented” the algorithms.</td>
</tr>
<tr>
<td>Dev3</td>
<td>Jeongnim Kim (instructor)</td>
</tr>
<tr>
<td>Dev4</td>
<td>Balint Joo or work at MICRO and PCL</td>
</tr>
</tbody>
</table>
Cray XC30: a distributed shared-memory cluster
Trends in Parallel Machines: clusters of SMPs

Top 10 systems in November 2014: clusters of SMPs using specialized interconnects

- Tianhe-2: Xeon + Xeon Phi
- Titan: Opteron + Tesla
- Sequoia: Blue Gene Q
- K Computer: MIPS
- XC30: Xeon

 Canonical HPC systems: clusters of SMPs using commodity interconnects

Your desktops and laptops: a SMP node with multi/many cores

Each system is an optimized solution of high performance and low cost (manufacturing, building, power, support)
Why Parallel Computing?

We have parallel computers. Need to use them well!

Parallel computing uses multiple computing units in parallel to

- solve problems more quickly than a single processor ("strong scaling")
- solve larger problems in the same time as a single processor ("weak scaling")
- solve problems with higher fidelity

Enables computational simulations for breakthrough discovery and prediction.

High-performance parallel computing is hard and requires

- Finding enough parallelism
- Deciding the optimal granularity, locality and load balance
- Coordination and synchronization

Real-world applications/algorithms are complex and often hierarchical; monolithic programming model is limited; no silver bullets
Parallel Programming for Performance

- Numerical and system libraries
- Distributed-memory parallel programming: MPI, PGAS
  - Map on to a set of memory domains, e.g. nodes, sockets, cores
  - Explicit and implicit data exchanges and synchronization
- Shared-memory parallel programming: OpenMP*, Pthreads, TBB, Cilk™ Plus, OpenCL*
- Vector programming: auto-vectorization, OpenMP* 4.0
Cori Applications: **NERSC 44 NESAP**

http://www.hpcwire.com/2014/09/03/nersc-reveals-44-nesap-code-teams/

<table>
<thead>
<tr>
<th>Benchmark</th>
<th>Parallelism</th>
<th>Language</th>
<th>2014</th>
<th>2017</th>
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<tbody>
<tr>
<td></td>
<td>MPI</td>
<td>THREADS</td>
<td>Fortran</td>
<td>C</td>
</tr>
<tr>
<td>MILC/CHROMA +</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>Nuclear QMC</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>BerkeleyGW/NWCHEM(PW)/QE/ VASP +</td>
<td>x</td>
<td>*</td>
<td>x</td>
<td>P</td>
</tr>
<tr>
<td>NWCHEM/CP2K +</td>
<td>x</td>
<td>*</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>GTC-P/GTCP-C</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td>x</td>
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<tr>
<td>QBOX</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td>x</td>
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<tr>
<td>LAMMPS/NAMD +</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td>O</td>
</tr>
<tr>
<td>HACC</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td>O</td>
</tr>
<tr>
<td>AMG-2013</td>
<td>x</td>
<td>x</td>
<td>x</td>
<td>P</td>
</tr>
</tbody>
</table>

**BGQ & KNC: Optimized and Ported**

**Red:** non-DOE applications
Why can’t we just stick to MPI*?

- We have clusters of SMPs.
  - Each node has 10-100 of cores and multiple threads per core.
  - Some hardware claims to support millions and soon billions of concurrency.
  - Multiple memory & cache levels with various sharing modes: L1 shared by 4 HT on KNC
- Cannot wait for a magic MPI implementation which does all.

- Applications can use the large memory available per SMP node
  - Eliminate/reduce data replications: only one copy of shared constant data is needed.
  - No extra data copies with put/get

- Consider MPI* time and resource use at scale
  - Scaling of collectives: $O(C \log C)$ vs $O(N \log N)$, $C=(1-1000)N$
  - Serialization of point-to-point communications
  - Data for MPI abstractions and communications
Evolution in computation, memory and communication

<table>
<thead>
<tr>
<th></th>
<th>Cray T3E-1350 [1]</th>
<th>Cray XC30 (Edison@NERSC)</th>
<th>XC30/T3E</th>
</tr>
</thead>
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<tr>
<td></td>
<td>processor clock</td>
<td>2.4GHz</td>
<td>3.64</td>
</tr>
<tr>
<td>SMP</td>
<td>1 CPU</td>
<td>2x12 cores</td>
<td>24</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>1</td>
</tr>
<tr>
<td>peak GF/s</td>
<td>1.350 /CPU</td>
<td>460.8 /SMP 19.2 /core</td>
<td>341</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>14.2</td>
</tr>
<tr>
<td>peak memory BW</td>
<td>1.2 GB/s/CPU</td>
<td>89 GB/s/SMP*</td>
<td>74</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>3</td>
</tr>
<tr>
<td>memory</td>
<td>256 MB/CPU</td>
<td>64 GB/SMP 2.67 GB/core</td>
<td>256</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>10.4</td>
</tr>
<tr>
<td>peak bisection BW</td>
<td>166 GB/s (512 CPUs)</td>
<td>11 GB/s/node</td>
<td>34</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>1.4</td>
</tr>
<tr>
<td>MPI latency (μsec)</td>
<td>6</td>
<td>0.25-3.7</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>0.125*</td>
</tr>
</tbody>
</table>

Cori/T3E Cori/Edison

<table>
<thead>
<tr>
<th></th>
<th>&gt; 2000</th>
<th>&gt; 6</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>&gt; 370**</td>
<td>&gt;5**</td>
</tr>
<tr>
<td></td>
<td>34</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>1</td>
</tr>
</tbody>
</table>

- Assume serialized MPI zero-message point-to-point communications.
- ** Depend on DDR4 or On-package Memory

High-performance parallel computing

Moving data is expensive!
- Node-node
- Socket-Socket; Processor-(co)processor
- Core-core
- SIMD lanes

At each parallel level
- Find enough parallelism
- Decide the optimal granularity
- Optimize locality and data movement
- Ensure load balance
- Reduce the impact of coordination and synchronization

All the parallel units have to be coordinated with maximum overlap of data movement and computing.
Set the goals and priorities (Dev*)

- Define performance and your performance goal
  - Strong scaling: reduced time-to-solution at any cost
  - Weak scaling: “constant” time-to-solution with increasing resources
  - Both at a sustained high performance

- Set your priorities
  - Performance, Performance, Performance
  - Optimize (performance, portability, maintenance, ....)

- Know your type, your team and ecosystem
  - Incremental development from the bottom (evolutionary)
  - Transformative development (revolutionary)
  - Iterative process of using both
High-performance Hybrid Programming 101 (Dev*)

- Apply computing 101: const, restrict, C99, alignment, remove branching ....
- Map the data and algorithms to the hierarchical memory and communication hardware and the parallel programming models
- Maximize the shared memory use: eliminate/reduce data replications.
  - Remember only one copy of shared constant data per task is needed!
- Maximize the distributed memory use: localize the data and do not share
  - Think what is needed for high-performance MPI applications
  - Use private data and thread-local storage
- Consider cost of OpenMP* or any thread-based (parallel programming) methods
  - Creating/destroying a team of threads is not FREE!
  - Implicit synchronization and barriers
  - Cache coherency
  - False sharing and write/read conflicts.
EMGeo: Part 1 for Dev0/Dev1

Know your application
Design experiments
Bottom-up transformation
Know EMGeo

Excerpts from README.md

- “EMGeo is a Fortran 90 pure MPI code“
- While the code is somewhat complex, the good news is that the 220 line `qmr` subroutine found in `krysolver.f90` takes up over 90% of the wall-clock run time under typical configurations. Further, this QMR solver routine spends a significant portion of time in **ELLPACK-format** sparse matrix-vector multiply operation appearing within the main loop (lines 243-255 of `krysolver.f90`).”
- “a finite difference (FD) code for electromagnetic imaging in geophysical exploration”
- “uses two levels of parallelism: FD method and multiple FD problems”
- “The FD problem domains are decomposed on an I x J x K grid of MPI ranks (inner level)”
- “**Please** refer questions to Scott before attempting to contact Michael.”
Set the goal(s) and design experiments

Goal: Transform EMGeo to attain *sustained performance* with any combination of MPI tasks and OpenMP threads

- Workload in run directory: p##_IxJxKxd1 where I*J*K=*MPI tasks*
  - p01_1x1x1xd1 p04_2x2x1xd1 p08_2x2x2xd1 p64_4x4x4xd1

- Establishing the baseline
  - Strong scaling with respect to MPI task
  - Hotspots analysis on a quad-socket HWS: p04_2x2x1xd1 and p08_2x2x2xd1

- Bottom-up transformation

- Results
Baseline performance on HSW-EX (quad 18-core)

- `I_MPI_PIN_DOMAIN`
  - 4 = socket; 8 = auto:9; 64 = core
- Just confirmed README.md
- QMR is *the* hotspot
- Domain-decomposition with boundary exchanges: constant total memory footprint
- Super-scaling from 1 to 4 task!
- Excellent strong scaling and all the parts scale well.
- 10% in MPI at 64 tasks: allreduce, send/recv

![Elapsed time (sec)](chart)

<table>
<thead>
<tr>
<th>MPI</th>
<th>1</th>
<th>4</th>
<th>8</th>
<th>64</th>
</tr>
</thead>
<tbody>
<tr>
<td>qmr</td>
<td>400</td>
<td>200</td>
<td>100</td>
<td>0</td>
</tr>
<tr>
<td>solve_yavg_coeff</td>
<td>4.05x</td>
<td>7.7x</td>
<td>24.7x</td>
<td></td>
</tr>
</tbody>
</table>
Hotspot analysis: Loops and functions

- 92.6% in `qmr` in `krysolver.f90`

```plaintext
do i=1,n
  MPI_ALLREDUCE
  do i=1, n
    enddo
  MPI_ALLREDUCE
  ...
```

- All the loops are the same size and LHS (lvalue) is linearly accessed.
- Go ahead and put OMP pragma
- Just careful about `allreduce` and make local variables private
QMR* in krysolver

* quasi-minimum-residue, a Krylov space solver

```c
!$OMP PARALLEL DO REDUCTION(+:ay) PRIVATE(csum,mcole,j)
do i=1,n
    ay = ay + cpnt(i)*dconjg(cpnt(i))
    csum = czero
    mcole=lshift(rshift(mcol(i),1),1)
do j=1,mcole,2
    csum = csum + mtx(j,i)*cgs(index(j,i))
    csum = csum + mtx(j+1,i)*cgs(index(j+1,i))
enddo
if(mcole.ne.mcol(i)) csum=csum+mtx(mcol(i),i)*cgs(index(mcol(i),i))
cvk1(i) = csum
enddo
!$OMP END PARALLEL DO
call MPI_ALLREDUCE(ay,dsum,1,MPI_DOUBLE_PRECISION,MPI_SUM, ..)
enddo
beta = czero

do i=1,n
    cvk1(i) = cpnt(i) - cvk1(i)
    crk(i) = cvk1(i)
    beta = beta + cvk1(i)*cvk1(i)
enddo

call MPI_ALLREDUCE(beta,csum,1,MPI_DOUBLE_COMPLEX, ..)
```

SpMV: sparse matrix-vector multiplication
QMR* in krysolver

```
do i=1,n
    ay = ay + cpnt(i)*dconjg(cpnt(i))
    csum = czero
    mcol = lshift(rshift(mcol(i),1),1)
    do j=1,mcol,2
        csum = csum + mtx(j,i)*cgs(index(j),cgs(index(j)))
    enddo
    if(mcol.ne.mcol(i)) csum = csum + mtx(mcol(i),cvs)
    cvk1(i) = csum
enddo
```

```
call MPI_ALLREDUCE(ay,dsum,1,MPI_DOUBLE_COMPLEX)
beta = czero
$OMP PARALLEL DO REDUCTION(+:beta)
```

```
do i=1,n
    cvk1(i) = cpnt(i) - cvk1(i)
    crk(i) = cvk1(i)
    beta = beta + cvk1(i)*cvk1(i)
enddo
```

```
call MPI_ALLREDUCE(beta,csum,1,MPI_DOUBLE_COMPLEX, ...)
```

```
Results: p64 vs p04 with 8 threads

Summary of p64 (using 64 cores)

- Elapsed Time: 12.385s
- Total Thread Count: 1
- Paused Time: 0s
- CPU Time: 12.050s
  - Spin Time: 0.409s
  - Overhead Time: 0s
  - Effective Time: 11.641s

Top Hotspots
This section lists the most active functions in your application.

<table>
<thead>
<tr>
<th>Function</th>
<th>CPU Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>[Loop at line 274 in qmr]</td>
<td>4.408s</td>
</tr>
<tr>
<td>[Loop at line 356 in qmr]</td>
<td>3.252s</td>
</tr>
<tr>
<td>[Loop at line 291 in qmr]</td>
<td>1.091s</td>
</tr>
<tr>
<td>[Loop at line 272 in qmr]</td>
<td>0.890s</td>
</tr>
<tr>
<td>[Loop at line 307 in qmr]</td>
<td>0.760s</td>
</tr>
<tr>
<td>[Others]</td>
<td>1.649s</td>
</tr>
</tbody>
</table>

Summary of p04 (using 32 cores)

- Elapsed Time: 16.250s
- Total Thread Count: 8
- Paused Time: 0s
- CPU Time: 105.419s
  - Effective Time: 92.985s
  - Spin Time: 12.202s
  - Overhead Time: 0.232s
- OpenMP Analysis. Collection Time: 16.250
  - Serial Time (outside any parallel region): 4.491s (27.6%)
  - Parallel Region Time: 11.759s (72.4%)
    - Estimated Ideal Time: 11.258s (69.3%)
    - Potential Gain: 0.501s (3.1%)

Top OpenMP Regions by Potential Gain
This section lists OpenMP regions with the highest potential for performance improvement.

<table>
<thead>
<tr>
<th>OpenMP Region</th>
<th>Potential Gain (%)</th>
<th>Elapsed Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>gmr_Somp$paralleltk@unknown:190:209</td>
<td>0.0%</td>
<td>0s</td>
</tr>
<tr>
<td>gmr_Somp$paralleltk@unknown:275:284</td>
<td>0.0%</td>
<td>0s</td>
</tr>
<tr>
<td>gmr_Somp$paralleltk@unknown:359:377</td>
<td>0.0%</td>
<td>0s</td>
</tr>
<tr>
<td>gmr_Somp$paralleltk@unknown:294:209</td>
<td>0.0%</td>
<td>0s</td>
</tr>
<tr>
<td>gmr_Somp$paralleltk@unknown:310:315</td>
<td>0.0%</td>
<td>0s</td>
</tr>
</tbody>
</table>

Amdahl's law?
Conversation with Dev2

- It should be straightforward to parallelize the other parts
  - MPI can do it. Then, why not OpenMP?
- Prediction: 11.48 sec = 1.21 (p64_2x2x2x2) + 10.27 (p04_2x2x1/16 OMP)
  - 8% gain as implied by the MPI time with 64 tasks

If Dev2 says, “What is the point? All these work for few % gain?”, then stop.

If Dev2 says, “That looks interesting. But, it just shows that the physics is not violated. Show me performance.”
EMGeo: Part 2
OpenMP Analysis: p04_2x2x1xd1 using 16 threads

- Overall no obvious load imbalance.
- Serial section: gen (solve_gen.f90) and solve_yavg_coeff.f90
Get rid of the “serial” bottleneck: Dev3

- Apply OpenMP in `solve_yang_coeff` at L108 and similar loops in `solve_gen.f90`
- It looks like all the temporary variables within the loop can be made private.

```
nrow = 0
nrow2= 0 ! rows on node space
do kndx=dim3_sim(5),dim3_sim(6)
    do jndx=dim3_sim(3),dim3_sim(4)
        do indx=dim3_sim(1),dim3_sim(2)
            nrow2 = nrow2 + 1
            do icomp=1,3
                nrow = nrow + 1
            enddo !icomp
        enddo !indx
    enddo !jndx
enddo !kndx
```

```
 !$OMP PARALLEL DO COLLAPSE(3) PRIVATE(nrow,nrow2,...)
do kndx=dim3_sim(5),dim3_sim(6)
    do jndx=dim3_sim(3),dim3_sim(4)
        do indx=dim3_sim(1),dim3_sim(2)
            nrow2 = compute_row(indx,jndx,kndx)
            nrow = nrow2*3;nrow2=nrow2+1
            do icomp=1,3
                nrow = nrow + 1
            enddo !icomp
        enddo !indx
    enddo !jndx
enddo !kndx
```

Results: disaster – NAN
What went wrong and how to proceed

- All the advertised gotchas exist: common block, hidden dependency ....
  ⇒ There are tools for that and Fortran users can fix them.

- Initialization determines the sparse-matrix storage ordering in ELLPACK-format and SpMV, need a critical look at
  - How data are ordered, allocated and initialized
  - How to facilitate SIMD optimization: collapse(2) vs collapse(3)
  - How auxiliary data structures are used; how many of them are used; why they are needed.

- Many solutions exist and time for serious discussion with Dev2 for transformative code design.
EMGeo on Cori

It will work *fine* on Xeon™ Phi

- Can use multiple MPIs on a node: no problem with memory use.
- Performance improvement through MPI/OpenMP on Xeon is real.
- Enough parallelisms to exploit; load-balancing is not difficult.
- Most of the critical loops are amenable to vectorizations.
- No hard serial bottlenecks exist. Just a matter of using OpenMP correctly.

Can it work *great* on Cori and future MICs?

- All these point to *Probably* but it is time to have serious conversation with the developers for code design and reset our goals.
Code design following best practices of today

- A Core is a new Node but threads are not MPI processes.
- Similar hierarchical architectures of CPUs: socket-core-SIMD
- Microarchitectures matter
  - Xeon™ HSW != KNL
  - Memory bandwidth, NUMAness, process vs thread, cache modes, SIMD, ...
  - Improved serial performance on KNL does not mean serial bottlenecks become magically uncritical.

Focus on

- Adaptive data partition and load balancing algorithms with MPI/OpenMP/SIMD
- Code pruning to facilitate compiler optimization
- Portable and performance portable code: encapsulate targeted optimization
Co-Design vs. Code Design

- HPC Myths
  - The magic compiler
  - The magic programming model/language (DSL)
  - Special-purpose hardware
  - Co-Design?

- HPC Reality
  - Follow the architecture
  - Know the boundary conditions
  - There is no such thing as a ‘code port’
  - Think out of the box
  - Get the best team
  - Work together

Average performance speed-up on ~10 applications codes on Titan is ~2 (ranging from 1..few to 7), but of Titan’s 27 PFlops, only 2.5 PFlops are in the CPU! What is wrong with this picture?

Blasting Through the 10 Petaflops Barrier: HACC on the BG/Q

Salman Habib
NPP and MCS Divisions,
Argonne National Laboratory

Vitali Morozov

HPC Myths

Argonne National Laboratory

HPC Reality

Blaclarity NP and MCS Divisions,
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HPC Myths

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HPC Reality

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Average performance speed-up on ~10 applications codes on Titan is ~2 (ranging from 1..few to 7), but of Titan’s 27 PFlops, only 2.5 PFlops are in the CPU! What is wrong with this picture?

Blasting Through the 10 Petaflops Barrier: HACC on the BG/Q

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