Moving from the:
Cray XE6 to the Cray XC30
Continuity
The High Productivity Vision

● Cray systems are designed to be High Productivity as well as High Performance Computers

● The Cray Programming Environment (PE) continues to provide a simple and consistent interface to users and developers.
  ● Renewed focus on improving scalability and reducing complexity

● The default Programming Environment provides:
  ● the highest levels of application performance
  ● a rich variety of commonly used tools and libraries
  ● a consistent interface to multiple compilers and libraries
  ● an increased automation of routine tasks

● Cray is committed to extending, developing and refining the PE.
  ● Frequent communication and feedback to/from users
  ● Strong collaborations with third-party developers
Cray Software Ecosystem

- CrayPAT
- Cray Apprentice2
- Reveal
- Cray PETSc, CASK
- Cray Iterative Refinement Toolkit

Compilers
- GNU
- Intel

Applications
- LSTC
- accelrys
- CD-adapco
- MathWorks
- Dassault Systemes
- Metacomp Technologies
- Altair

Debugging
- PGI
- TotalView Technologies
- allinea
- TotalView Technologies

Performance Tools
- ParaTools
- Lustre
- nag
- IMSL

I/O & Libraries
- Lustre
- nag
- IMSL

DVS

Cray Linux Environment
An Adaptive Linux OS optimized specifically for HPC

ESM – Extreme Scalability Mode
• No compromise scalability
• Low-Noise Kernel for scalability
• Native Comm. & Optimized MPI
• Application-specific performance tuning and scaling

CCM – Cluster Compatibility Mode
• No compromise compatibility
• Fully standard x86/Linux
• Standardized Communication Layer
• Out-of-the-box ISV Installation
• ISV applications simply install and run

CLE run mode is set by the user on a job-by-job basis to provide full flexibility
Cray Integrated Programming Environment

1. Applications

2. Compiler

3. Static Analysis

4. Runtime Information

5. Program Analysis Tools

6. Source-to-Source optimizations

Performance Analysis Overview

High Level Profile/Tracing

Performance Problem Analyzer

Export/Import Program Analyses

Performance Feedback

Queries for Application Optimization

Database containing application information

Executing Application

Runtime Information
### Cray Programming Environment Distribution

**Focus on Performance and Productivity**

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<td>Cray PETSc (with CASK)</td>
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<td>Cray Trilinos (with CASK)</td>
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- **Cray developed**
- **Licensed ISV SW**
- **3rd party packaging**
- **Cray added value to 3rd party**
XE6 and XC30 Software Components
Generational Commonality

**XE6 Software Stack**

- **MPI Applications**
- **PGAS/SHMEM Applications**

- Gemini-aware optimized PE components
  - MPI
  - SHMEM
  - UPC & CAF PGAS Languages

- User Applications
- Programming Models

- Generic APIs
  - User-level
  - Distributed Memory API (DMAPP)
  - Generic Network Interface (uGNI)

- Kernel
  - CLE Linux Core
  - Kernel-level Generic Network Interface (kGNI)
  - Gemini - Hardware Abstraction Layer (HAL)

- Network

**XC30**

- **MPI Applications**
- **PGAS/SHMEM Applications**

- Aries-aware optimized PE components
  - MPI
  - SHMEM
  - UPC & CAF PGAS Languages

- User Applications
- Programming Models

- Generic APIs
  - User-level
  - Distributed Memory API (DMAPP)
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- Kernel
  - CLE Linux Core
  - Kernel-level Generic Network Interface (kGNI)
  - Aries - Hardware Abstraction Layer (HAL)

- Network

**Common Base**
The Cray System
Transition
Compilers

Like HECToR will be three compilers installed, however Intel replaces the PGI compiler.

Only the most recent versions will be available (e.g. those released over the last few months)

- **Cray Compilation Environment (CCE)**
  - Coincides with new 8.2 release

- **Intel Composer Suite**
  - New compiler over HECToR

- **GNU Compiler Collection**
  - The standard set of expected compilers and tools.
CCE Overview

- **Cray technology focused on scientific applications**
  - Takes advantage of automatic vectorization
  - Takes advantage of automatic shared memory parallelization

- ** Standards conforming languages and programming models**
  - ANSI/ISO C99 and C++2003 compliant
  - OpenMP 3.1 compliant, working on OpenMP 4.0
  - OpenACC 1.0

- **OpenMP and automatic multithreading fully integrated**
  - Share the same runtime and resource pool
  - Aggressive loop restructuring and scalar optimization done in the presence of OpenMP
  - Consistent interface for managing OpenMP and automatic multithreading
Improvements for CCE/8.2.0 on XC30

● Full support for optimising for “ivybridge” processors.

● Improved performance of performance-critical maths intrinsics.

● New Coarray C++ template library that implements coarray concepts to C++.

● GNU and CCE OpenMP libraries are now compatible. Linking must be performed by CCE with PrgEnv-cray.
CCE – GNU – Intel compilers

● More or less all optimizations and features provided by CCE are available in Intel and GNU compilers
  ● GNU compiler serves a wide range of users & needs
    ● Default compiler with Linux, some people only test with GNU
    ● Defaults are conservative (e.g. -O1)
      ● -O3 includes vectorization and most inlining
    ● Performance users set additional options
  ● Intel compiler is typically more aggressive in the optimizations
    ● Defaults are more aggressive (e.g. -O2), to give better performance “out-of-the-box”
      ● Includes vectorization; some loop transformations such as unrolling; inlining within source file
    ● Options to scale back optimizations for better floating-point reproducibility, easier debugging, etc.
    ● Additional options for optimizations less sure to benefit all applications
  ● CCE is even more aggressive in the optimizations by default
    ● Better inlining and vectorization
    ● Aggressive floating-point optimizations
    ● OpenMP enabled by default
## Cray, Intel and GNU compiler flags

<table>
<thead>
<tr>
<th>Feature</th>
<th>Cray</th>
<th>Intel</th>
<th>GNU</th>
</tr>
</thead>
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<tr>
<td>Listing</td>
<td>-hlist=a</td>
<td>-opt-report3</td>
<td>-fdump-tree-all</td>
</tr>
<tr>
<td>Free format (ftn)</td>
<td>-f free</td>
<td>-free</td>
<td>-ffree-form</td>
</tr>
<tr>
<td>Vectorization</td>
<td>By default at -O1 and above</td>
<td>By default at -O2 and above</td>
<td>By default at -O3 or using -ftree-vectorize</td>
</tr>
<tr>
<td>Inter-Procedural Optimization</td>
<td>-hwp</td>
<td>-ipo</td>
<td>-flto (note: link-time optimization)</td>
</tr>
<tr>
<td>Floating-point optimizations</td>
<td>-hfpN, N=0...4</td>
<td>-fp-model [fast</td>
<td>fast=2</td>
</tr>
<tr>
<td>Suggested Optimization</td>
<td>(default)</td>
<td>-O2 -xAVX</td>
<td>-O2 -mavx -ftree-vectorize -ffast-math -funroll-loops</td>
</tr>
<tr>
<td>Aggressive Optimization</td>
<td>-O3 -hfp3</td>
<td>-fast</td>
<td>-Ofast -mavx -funroll-loops</td>
</tr>
<tr>
<td>OpenMP recognition</td>
<td>(default)</td>
<td>-fopenmp</td>
<td>-fopenmp</td>
</tr>
<tr>
<td>Variables size (ftn)</td>
<td>-s real64</td>
<td>-real-size 64</td>
<td>-freal-4-real-8</td>
</tr>
<tr>
<td></td>
<td>-s integer64</td>
<td>-integer-size 64</td>
<td>-finteger-4-integer-8</td>
</tr>
</tbody>
</table>
Compiler man pages and documentation

- For more information on individual compilers

<table>
<thead>
<tr>
<th>PrgEnv</th>
<th>C</th>
<th>C++</th>
<th>Fortran</th>
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<tr>
<td>PrgEnv-cray</td>
<td>man craycc</td>
<td>man crayCC</td>
<td>man crayftn</td>
</tr>
<tr>
<td>PrgEnv-intel</td>
<td>man icc</td>
<td>man icpc</td>
<td>man ifort</td>
</tr>
<tr>
<td>PrgEnv-gnu</td>
<td>man gcc</td>
<td>man g++</td>
<td>man gfortran</td>
</tr>
<tr>
<td>Wrappers</td>
<td>man cc</td>
<td>man CC</td>
<td>man ftn</td>
</tr>
</tbody>
</table>

- To verify that you are using the correct version of a compiler, use:
  - `-V` option on a cc, CC, or ftn command with CCE and Intel
  - `--version` option on a cc, CC, or ftn command with GNU

- Cray Reference Manuals:
  - C and C++: [http://docs.cray.com/books/S-2179-81/](http://docs.cray.com/books/S-2179-81/)
  - Fortran: [http://docs.cray.com/books/S-3901-81/](http://docs.cray.com/books/S-3901-81/)
## Interlagos/Ivybridge Comparison

<table>
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<th>AMD Opteron “Interlagos”</th>
<th>Intel Xeon “Ivybridge”</th>
</tr>
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<tbody>
<tr>
<td><strong>Base Clock Speed</strong></td>
<td>2.3 GHz</td>
<td>2.7 GHz</td>
</tr>
<tr>
<td><strong>Cores per die</strong></td>
<td>6</td>
<td>12</td>
</tr>
<tr>
<td><strong>Dies per node</strong></td>
<td>4</td>
<td>2</td>
</tr>
<tr>
<td><strong>Each cores has:</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>User threads</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Function group</td>
<td>1 SSE (vector)</td>
<td>1 AVX (vector)</td>
</tr>
<tr>
<td>bits wide</td>
<td>128 bits wide</td>
<td>256 bits wide</td>
</tr>
<tr>
<td>functional units</td>
<td>1 add and 1 multiply</td>
<td>1 add and 1 multiply</td>
</tr>
<tr>
<td>Cache: L1</td>
<td>32KB</td>
<td>32KB</td>
</tr>
<tr>
<td>Cache: L2</td>
<td>512KB</td>
<td>256KB</td>
</tr>
<tr>
<td>L3 Cache (per die)</td>
<td>6 MB</td>
<td>30 MB</td>
</tr>
<tr>
<td>Total Cache per core</td>
<td>1.5 MB</td>
<td>2.75 MB</td>
</tr>
<tr>
<td><strong>Cache BW Per core (GB/s)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>L1/L2/L3</td>
<td>35 / 3.2 / 3.2</td>
<td>100 / 40 / 23</td>
</tr>
<tr>
<td>Stream TRIAD BW/node</td>
<td>52 Gbytes/s</td>
<td>100 Gbytes/s</td>
</tr>
<tr>
<td>Peak DP FLOPs per core</td>
<td>4 flops/clk</td>
<td>8 flops/clk</td>
</tr>
<tr>
<td>Peak DP FLOPs per node</td>
<td>294 GFlops</td>
<td>518 GFlops</td>
</tr>
<tr>
<td>Main memory latency</td>
<td>110ns</td>
<td>82ns</td>
</tr>
</tbody>
</table>
The XC30 Compute node features:

- **2 x Intel® Xeon® Sockets/die**
  - 12 core Ivybridge
  - QPI interconnect
  - Forms 2 NUMA nodes
- **8 x 1833MHz DDR3**
  - 8 GB per Channel
  - 64 GB total
- **1 x Aries NIC**
  - Connects to shared Aries router and wider network
  - PCI-e 3.0
Intel® Xeon® Ivybridge 12-core socket/die

- 4 x 1866 MHz DDR3 Channels
- Ring bus
- Quick Path Interconnect (inter die)
- DDR3 Memory Controller
- Shared L3 Cache
- Core
- Core
- Core
- Core
- Core
- Core
- Core
- Core
- Core
- Core
- Core
- Core
- QPI
- System
- PCIe-3.0
- External I/O (Aries)
- Socket/die
Intel Xeon Ivybridge Core Structure

- Manufactured on a 22nm Process
- 256 bit AVX Instructions (4 double precision floating point)
  - 1 x Add
  - 1 x Multiply
  - 1 x Other
- 2 Hardware threads (Hyperthreads)
- Peak DP FP per node 8FLOPS/clock
Placement and Scheduling
There are always two stages to launching a job on a Cray XC30 with a batch scheduler.

1. Requesting the right amount of resource from the batch scheduler

2. Launching the parallel application on the allocated compute nodes

Both of these stages have been affected by upgrades to software and hardware when compared to Cray XE6
Glossary of terms

PE/Processing Element
● A discrete software process with an individual address space. One PE is equivalent to:
  1 MPI Rank, 1 Coarray Image, 1 UPC Thread, or 1 SHMEM PE

Threads
● A logically separate stream of execution inside a parent PE that shares the same address space

CPU
● The minimum piece of hardware capable of running a PE. It may share some or all of its hardware resources with other CPUs
  Equivalent to a single “Intel Hyperthread”

Compute Unit
● The individual unit of hardware for processing, may be seen described as a “core”. Poite may provide multiple CPUs.
Hyperthreads

- Hyper-threads are a feature of modern Intel processors
  - Essentially they are a form of hardware multithreading that avoids costly context switches.
  - Designed to mask long pauses in execution, e.g. network communication, memory accesses.
- Allows OS to schedule two processes (PEs) simultaneously on the same hardware core (Compute unit)
  - Each thread context is held in hardware and scheduled automatically (virtually no overhead of a context switch)
- The Cray XC30 software and hardware stacks provide full support for using Hyperthreads.
  - 48 CPUs visible to the OS
    - CPUs 0-11 & 24-35 on Socket 0
    - CPUs 12-23 & 36-47 on Socket 1
    - Shared compute unit pairs of CPUs are 0&24, 7&31, 8&32 and 15&39 etc.
An Ivybridge XC30 blade has two sockets, each with 12 Intel Cores. Each Intel core is termed a “Compute Unit”.

In “Single stream” mode we elect to use only one Hyperthread (CPU) from each of these “Compute Units”.

This means there is a total of 24 “CPUs” per node.

Therefore, without oversubscribing, there can only be a maximum of 24 PEs and threads assigned to one node.
Users can choose whether to use single or dual stream mode at runtime using aprun’s “-j1” or “-j2” options. “-j1” is single stream mode where, aprun binds PEs and ranks to the 24 Compute Units (e.g. only use CPUs 0-23)
Default Binding - CPU

- By default `aprun` will bind each PE to a single CPU for the duration of the run.
- This prevents PEs moving between CPUs.
- All child processes of the PE are bound to the same CPU.
- PEs are assigned to CPUs on the node in increasing order from 0. e.g.

```
aprun -n 48 -N 24 -j1 a.out
```
Default Thread Binding (pt 1)

- You can inform `aprun` how many threads will be created by each PE by passing arguments to the `–d` (depth) flag.
- `aprun` does not create threads, just the master PE.
- PEs are bound to a single CPU and reserve space according to the depth argument, e.g.

```
aprun -n 24 -N 12 -d2 -j1 a.out
```
Each subsequently created child processes/thread is bound by the OS to the next CPU \textit{(modulo by the depth argument)}. e.g.

\begin{verbatim}
OMP_NUM_THREADS=2
aprun -n 24 -N 12 -d2 -j1 a.out
\end{verbatim}

Each PE becomes the master thread and spawns a new child thread. The OS binds this child thread to the next CPU.
BEWARE – Intel Helper Threads

● The Intel OpenMP runtime is different to GNU and CCE.
  ● It creates an extra thread as a shepherd … (n+1 threads spawned)
  ● It also has its own method of binding to CPUs (KMP_AFFINITY)

● Unfortunately both of these options can make things more complicated on Cray XC30 with aprun binding features.

● Cray’s default advice…
  ● Don’t use KMP_AFFINITY to bind threads:
    ● export KMP_AFFINITY=disabled
    ● aprun -cc [numa_node|none] <exe>
  ● Study man aprun in detail
  ● Contact the helpdesk/Centre of Excellence
The alternative mode is “Dual stream” mode each of these “Compute Units” can host two Hyperthreads, or “CPUs”.

This means there is a total of 48 CPUs per node.

And so the maximum number of PEs and threads that can be assigned without oversubscription is 48 PEs.
Include Hyperthreads “-j2” Dual Stream Mode

Specifying “-j2” in aprun assigns PEs to all of the 48 CPUs available. However CPUs that share a common Compute Unit are assigned consecutively.

This means threads will share Compute Units with default binding.
Some Hyperthread advice

- Pure MPI, Mix-mode/hybrid and MPMD all continue to function as they did on Cray XE6 by default.
  - Just remember that number of NUMA nodes drops to 2.

- By default, aprun will use only 1 CPU per compute unit.
  - We expect this to the optimal way to run most HPC codes.

- The default mode leaves the Hyperthreads open for the Operating System to use
  - Can be used for CPU specialisation and MPI Progress engines.

- We expect only codes with low computational intensity to benefit from actively using hyperthreads
  - Others may transparently benefit from the additional OS and library features
The version of PBS has changed between HECToR and Archer (now running PBS 12)

This is a requirement, but it also allows for several improvements that were previously unavailable

- The numbers of queues will be significantly reduced (perhaps just 4)
- One of these queues will potentially support quick job turn-around
  create a “debug” queue

However, the upgrade has changed the way users ask for resources from the PBS batch scheduler.

- Users will now request resources in quanta of nodes rather than MPI ranks
- Older mpp* style notation will be rejected by the scheduler as it is no longer accepted.
PBS Select notation

PBS now asks users to select “chunks” of resources for their jobs.

This replaces the older notation so users must remove all mppwidth, mppnpn and mppdepth statements from batch scripts.

The simplest way to think of a chunk is as one entire node of the XC30. Users can submit jobs requesting numbers of nodes using:

```
#PBS -l select=<num_nodes>
```

(or via the command line qsub options)
Writing job scripts

ARCHER jobs will have access to new environment variables, e.g. $NUM\_NODES which contains the number of nodes allocated to the job.

Scripts are then free to launch jobs via aprun using any layout as long as it does not exceed the number of nodes allocated. E.g. for generic aprun

```
aprun -n n -N N -d d -j j a.out
```

\[
(d \times N) \leq (24 \times j) \leq \text{ceiling}(n / N) \leq $NUM\_NODES
\]

**NB.** OMP\_NUM\_THREADS will be set by default to 1.
All job scripts should set or unset this variable as necessary.
PBS – Fine tuning

PBS allows to users to specify more details about how many PEs and OpenMP threads will run in a chunk.

Users can specify:
-1 select=<#nodes>:mpiprocs=<#ppn>:ompthreads=<#threads>

On ARCHER jobs default to mpiprocs=24 and ompthreads=1.

Each job will be launched with environment variables:

$NUM_NODES=<#nodes>
$NUM_PPN=<#ppn>          # copied from mpiprocs
$NUMDEPTH=<#threads>     # copied from ompthreads
Some simple examples

```
#!/bin/bash
#PBS -l select=64:mpiprocs=24
#PBS -l walltime=6:00:00

NUM_WIDTH=${[ ${NUM_NODES} * ${NUM_PPN} ]}
cd ${PBS_O_WORKDIR}

export OMP_NUM_THREADS=1 # Added for safety
aprun -n ${NUM_WIDTH} -N ${NUM_PPN} mpiapp.exe
```

```
#!/bin/bash
#PBS -l select=32:mpiprocs=24:ompthreads=2
#PBS -l walltime=12:00:00

NUM_WIDTH=${[ ${NUM_NODES} * ${NUM_PPN} ]}
cd ${PBS_O_WORKDIR}

export OMP_NUM_THREADS=${NUM_DEPTH}
aprun -n ${NUM_WIDTH} -N ${NUM_PPN} -d ${NUM_DEPTH} -j2 app.exe
```
The simplest example

```bash
#!/bin/bash
#PBS -l select=64
#PBS -l walltime=6:00:00

cd ${PBS_O_WORKDIR}
aprun -B mpiapp.exe
```

**ARCHER queues default to** `mpiprocs=24:ompthreads=1`

Therefore this `aprun` will use:

```bash
aprun -n 1536 -N 24 -d 1 -j1 mpiapp.exe
```
Using the Higher Memory Nodes

ARCHER has one group of nodes that has 128 GB of main memory per node.

To tell the scheduler that you require these nodes, add `bigmem=true` to the select request.

e.g.

```
-l select=64:mpiprocs=24:ompthreads=1:bigmem=true
```
Libraries
Intel provides the Intel Math Kernel Library (MKL) with the composer suite.

Like cray-libsci, MKL provides sets of routines for scientific and engineering applications (also includes financial applications).

Routines are highly optimised for Intel’s own processors and can offer extremely good performance.

Distributed as mutually exclusive libraries, it is not a modle. Instead advice on linking with Intel and GNU compilers can be found here:

Linking with MKL and PrgEnv-crav

● PrgEnv-crav compatible with sequential, not threaded, MKL

● Assume you have loaded the intel module as well as cce (this defines the $INTEL_PATH)
  ● Typical case: You want to use MKL BLAS and/or LAPACK
    -L ${INTEL_PATH}/mkl/lib/intel64/ \\ 
    -Wl,--start-group \\ 
    -lmkl_intel_lp64 -lmkl_sequential -lmkl_core \\ 
    -Wl,--end-group
  ● Another typical case: You want to use MKL serial FFTs/DFTs
    Same as above (need more for FFTW interface)
  ● A less typical case: You want to use MKL distributed FFTs
    -L ${INTEL_PATH}/mkl/lib/intel64/ \\ 
    -Wl,--start-group \\ 
    -lmkl_cdft_core -lmkl_intel_lp64 -lmkl_sequential \ 
    -lmkl_core -lmkl_blacs_intelmpi_lp64 \ 
    -Wl,--end-group

● The Intel MKL Link Line Advisor can tell you what to add to your link line
  ● http://software.intel.com/sites/products/microkt/
MPI Features / Functionality for XC

● MPI on XC behaves essentially the same as MPI on XE
  ● uGNI interface is the same for XE and XC
  ● MPICH3 code base is nearly the same
    ● Messaging Paths (VSHORT, EAGER, RENDEZVOUS) are Identical

● Enhanced Features for XC
  ● Modified MPI Asynchronous Progress Engine Threads
    ● Threads can be placed on unused Intel hyper thread cores

● XC Hardware Collective Engine (CE)
  ● XC supports hardware-offload of Barrier & Allreduce collectives
  ● Invoke these via MPICH_USE_DMAPP_COLL env variable
  ● Must also link libdmapp into your application directly.
MPI - Async Progress Engine Support

- Used to improve communication/computation overlap
  - Each MPI rank starts a “helper thread” during MPI_Init
- Helper threads progress MPI engine while application computes
- Only inter-node messages that use Rendezvous Path are progressed (relies on BTE for data motion)

To enable on XC when using 1 stream per core:
- export MPICH_NEMESIS_ASYNC_PROGRESS=1
- export MPICH_MAX_THREAD_SAFETY=multiple
- export MPICH_GNI_USE_UNASSIGNED_CPUS=enabled
- Run application: aprun -n XX a.out

To enable on XC when using 2 streams per core recommend running with the corespec option:
- export MPICH_NEMESIS_ASYNC_PROGRESS=1
- export MPICH_MAX_THREAD_SAFETY=multiple
- Run application with corespec: aprun -n XX -r [1-2] a.out

- 10% or more performance improvements with some apps
Non Blocking MPI (inc MPI-3)

Collective computation, 4096 PE’s

Average completion time [µs]

Reduction size [B]

Overlapped availability [%]

Reduction size [B]

Data courtesy of P. Manninen Cray Finland
CPU Specialisation

- Despite the low-noise nature of the XC30’s CNL Linux OS it occasionally is necessary to run OS/kernel/daemon processes on CPUs.
- If all CPUs are in use then the OS must swap a user process out to execute the OS/kernel/deammon process.
- Normally this introduces only a small amount of noise to the application which evens out over the length of the run.
- However, there are certain pathological cases which amplify these delays if there are frequent synchronisations between nodes (e.g. collectives) preventing scaling.
- CPU specialisation reserves some CPUs for the OS/system/daemon tasks (like OS, MPI progress engines, daemons). This improves overall performance.
CPU Specialisation (pt 2)

- On the XC30 the reserved CPU’s are automatically chosen to be from any unused CPUs on Compute Units (e.g. spare Hyperthreads), even if “-j1” has been selected.

- Users specify how many CPUs to reserve by adding a “-r <CPUs>” flag to the aprun command. The sum total of “-r” and “-N” must not exceed 48 (the total number of CPUs on the node). E.g

  aprun -n 1024 -N 24 -r 8 -j 1 a.out

- Required for use with MPI environment variables, MPICH_GNI_USE_UNASSIGNED_CPUS and MPICH_NEMESIS_ASYNC_PROGRESS flags.
Hyperthreading optimization chart

Single Stream Mode – No MPI Async:
Collect performance baseline here
Maximize per cpu performance
Little MPI communication overlap for medium size messages

Dual Stream Mode – Without MPI Async
Goals:
Optimizing per node performance or
Maximizing performance by using many PEs
Is this “better”?

Single Stream Mode – With MPI Async
Goals:
Maximize per cpu performance
Improve communication performance
Does overall performance improve?

Dual Stream Mode – With MPI Async
Goals:
Optimizing per node perf. or
Maximizing perf. using many PEs and
Improve communication performance…
But give up using 1 or 2 hyperthreads
Is this “better”? 
How to get more help

- Support website: [http://www.archer.ac.uk](http://www.archer.ac.uk)
- Cray XC30 Advanced Tools Workshop
  Edinburgh @ EPCC
  28th – 29th January 2014