

Cray Scientific Libraries: Overview and Performance

**Cray XE6 Performance Workshop
University of Reading
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Contents

- **LibSci overview and usage**
- **BFRAME / CrayBLAS**
- **LAPACK**
- **ScaLAPACK**
- **FFTW / CRAFFT**
- **CASK**
- **LibSci for Accelerators**

What are libraries for?

- **Building blocks for writing scientific applications**
- **Historically – allowed the first forms of code re-use**
- **Later – became ways of running optimized code**
- **These days the complexity of the hardware is very high**
- **Cray PE insulates the user from that complexity**
 - Cray module environment
 - CCE
 - Performance tools
 - Tuned MPI libraries (+PGAS)
 - Optimized Scientific libraries

Cray scientific libraries are designed to give maximum possible performance from Cray systems with minimum effort

What makes Cray libraries special

1. Node performance

- Highly tune BLAS etc at the low-level

2. Network performance

- Optimize for network performance
- Overlap between communication and computation
- Use the best available low-level mechanism
- Use adaptive parallel algorithms

3. Highly adaptive software

- Using auto-tuning and adaptation, give the user the known best (or very good) codes at runtime

4. Productivity features

- Simpler interfaces into complex software

Scientific libraries – functional view

FFT

FFTW

CRAFFT

Sparse

Trilinos

PETSc

CASK

Dense

BLAS

LAPACK

ScaLAPACK

IRT

Libsci Usage all fits on one slide

- **LibSci**

- The drivers should do it all for you. Don't explicitly link.
- For threads, set OMP_NUM_THREADS
 - Threading is used within libsci.
 - If you call within parallel region, single thread used
 - -WI, -ydgemm_ reveals where the link was resolved

- **FFTW**

- Module load fftw (there are also wisdom files you can pick up)

- **PETSc**

- Module load petsc (or module load petsc-complex)
- Use as you would your normal petsc build

- **Trilinos**

- Module load trilinos

- **CASK – no need to do anything you get optimizations free**

Your friends

- module command (module --help)
- PrgEnv modules :
- Component modules
- csmlversion (tool)

TUNER/STUNER> module avail PrgEnv

PrgEnv-cray/3.1.35	PrgEnv-gnu/4.0.12A	PrgEnv-
pathscale/3.1.37G		
PrgEnv-cray/3.1.37AA	PrgEnv-gnu/4.0.26A	PrgEnv-
pathscale/3.1.49A		
PrgEnv-cray/3.1.37C	PrgEnv-gnu/4.0.36(default)	PrgEnv-
pathscale/3.1.61		
PrgEnv-cray/3.1.37E	PrgEnv-intel/3.1.35	PrgEnv-
pathscale/4.0.12A		
PrgEnv-cray/3.1.37G	PrgEnv-intel/3.1.37AA	PrgEnv-
pathscale/4.0.26A		
PrgEnv-cray/3.1.49A	PrgEnv-intel/3.1.37C	PrgEnv-
pathscale/4.0.36(default)		
PrgEnv-cray/3.1.61	PrgEnv-intel/3.1.37E	PrgEnv-pgi/3.1.35
PrgEnv-cray/4.0.12A	PrgEnv-intel/3.1.37G	PrgEnv-
pgi/3.1.37AA		
PrgEnv-cray/4.0.26A	PrgEnv-intel/3.1.49A	PrgEnv-pgi/3.1.37C
PrgEnv-cray/4.0.36(default)	PrgEnv-intel/3.1.61	PrgEnv-
pgi/3.1.37E		
PrgEnv-gnu/3.1.35	PrgEnv-intel/4.0.12A	PrgEnv-pgi/3.1.37G
PrgEnv-gnu/3.1.37AA	PrgEnv-intel/4.0.26A	PrgEnv-
pgi/3.1.49A		
PrgEnv-gnu/3.1.37C	PrgEnv-intel/4.0.36(default)	PrgEnv-
pgi/3.1.61		
PrgEnv-gnu/3.1.37E	PrgEnv-pathscale/3.1.35	PrgEnv-
pgi/4.0.12A		
PrgEnv-gnu/3.1.37G	PrgEnv-pathscale/3.1.37AA	PrgEnv-
pgi/4.0.26A		
PrgEnv-gnu/3.1.49A	PrgEnv-pathscale/3.1.37C	PrgEnv-
pgi/4.0.36(default)		
PrgEnv-gnu/3.1.61	PrgEnv-pathscale/3.1.37E	

- Cray driver scripts ftn, cc, CC

```
----- /opt/cray/modulefiles -----
-----
xt-libsci/10.5.02      xt-libsci/11.0.04      xt-libsci/11.0.05.1
xt-libsci/11.0.03      xt-libsci/11.0.04.8    xt-libsci/11.0.05.2(default)
```

```
xt-libsci/10.5.02      xt-libsci/11.0.04.8    xt-libsci/11.0.05.2(default)
```

Check you got the right library!

- Add options to the linker to make sure you have the correct library loaded.
- `-Wl` adds a command to the linker from the driver
- You can ask for the linker to tell you where an object was resolved from using the `-y` option.
 - E.g. `-Wl, -ydgemm_`

```
./main.o: reference to dgemm_  
/opt/xt-libsci/11.0.05.2/cray/73/mc12/lib/libsci_cray_mp.a(dgemm.o) :  
definition of dgemm_
```

Note : explicitly linking “-lsci” is bad! This won’t be found from libsci 11+ (and means single core library for 10.x!)

Threading

- **LibSci is compatible with OpenMP**
- Control the number of threads to be used in your program using **OMP_NUM_THREADS**
 - e.g. in job script

```
setenv OMP_NUM_THREADS 16
```
 - Then run with `aprun -n1 -d16`
- **What behavior you get from the library depends on your code**
 1. No threading in code
 - The BLAS call will use OMP_NUM_THREADS threads
 2. Threaded code, outside parallel region
 - The BLAS call will use OMP_NUM_THREADS threads
 3. Threaded code, inside parallel region
 - The BLAS call will use a single thread

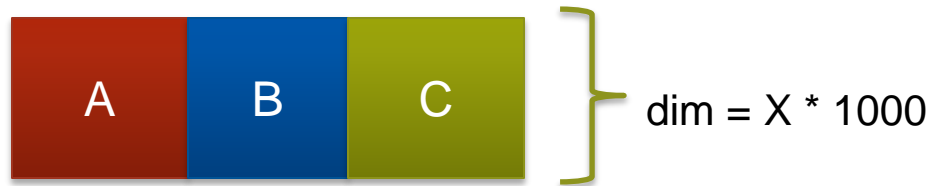
Emphasis

- A large subset of HPC customers care very deeply about each of the following
 - BLAS – explicit calls in their code
 - LAPACK – linear solvers
 - LAPACK – eigensolvers
 - ScaLAPACK
 - Serial FFT
- Our job is to make them work at extreme performance on Cray hardware
- A flaming-hot GEMM library can support wide usage

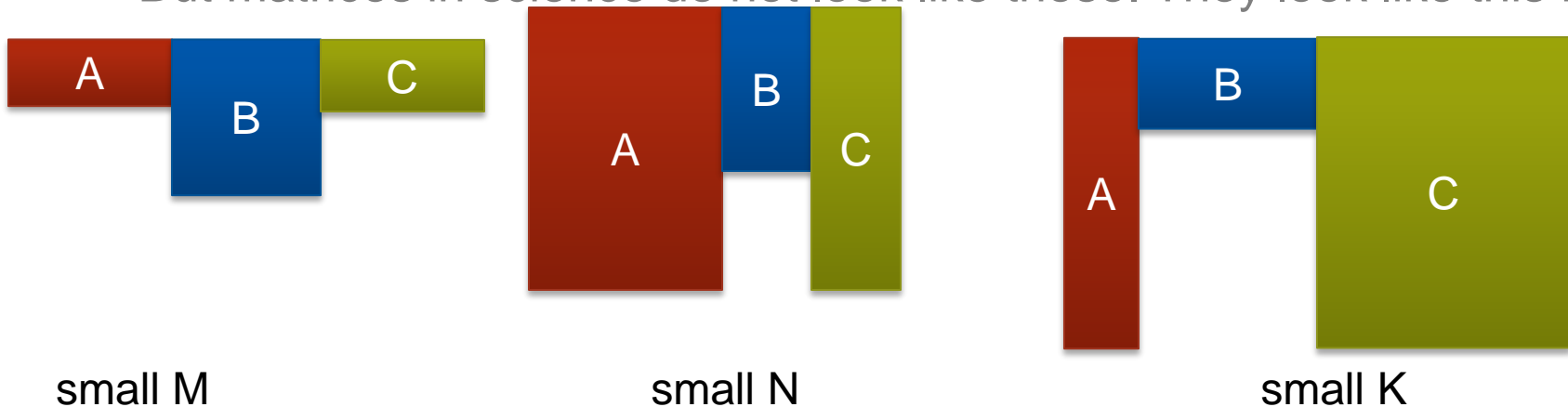


But that is very hard

- Vendor libraries can be very heavily tuned for a specific problem.
- Tunings are not general, unfortunately.
- Your library is probably tuned for this



- But matrices in science do not look like those. They look like this :



BFRAME / CrayBLAS

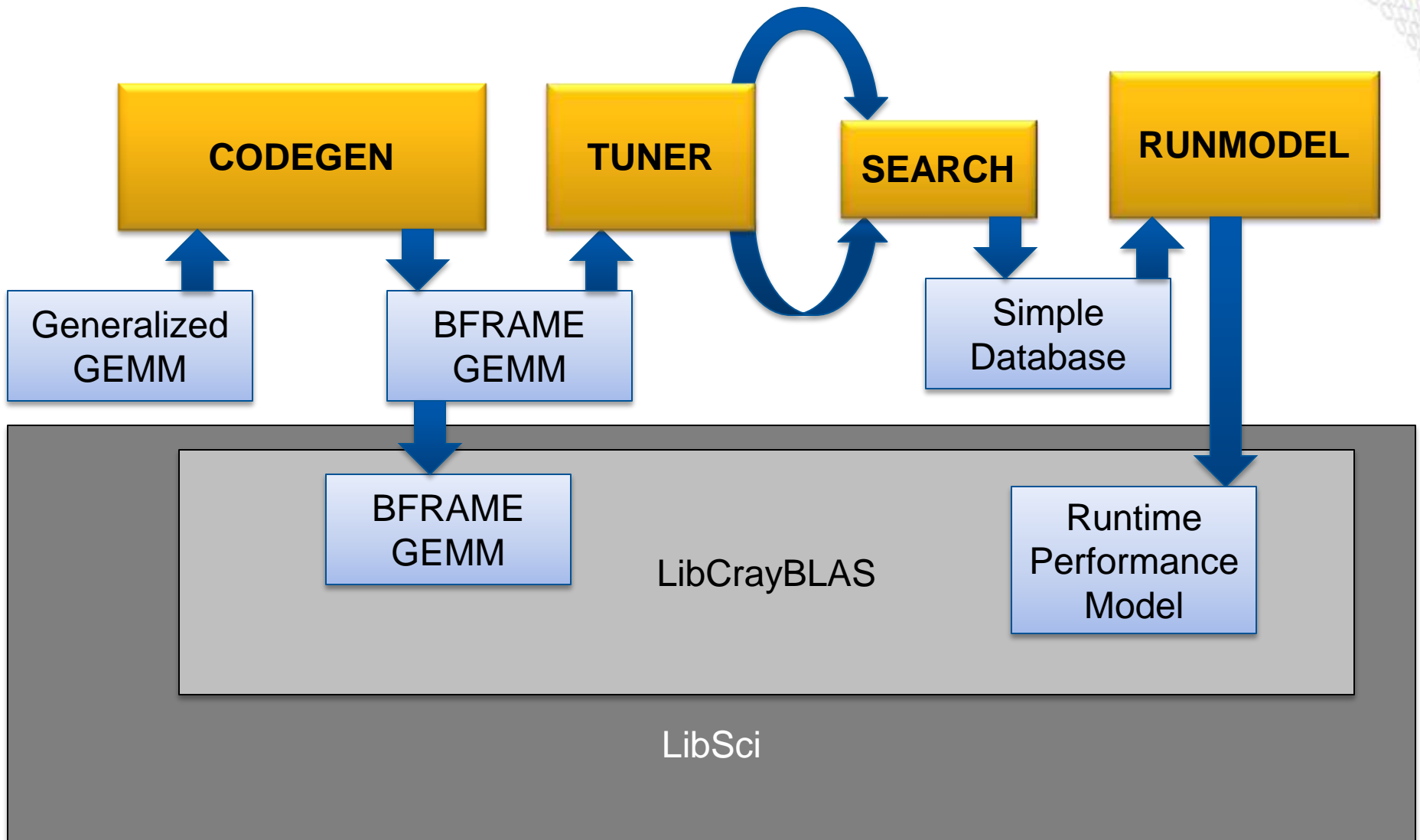
- **Goal**

- Entire auto-tuning framework for BLAS on CPUS & GPUS
- Must provide better performance than alternatives for “real” matrices
- Improve threaded performance for any problem size on any number of threads
- Encapsulate all tunings into a run-time performance model

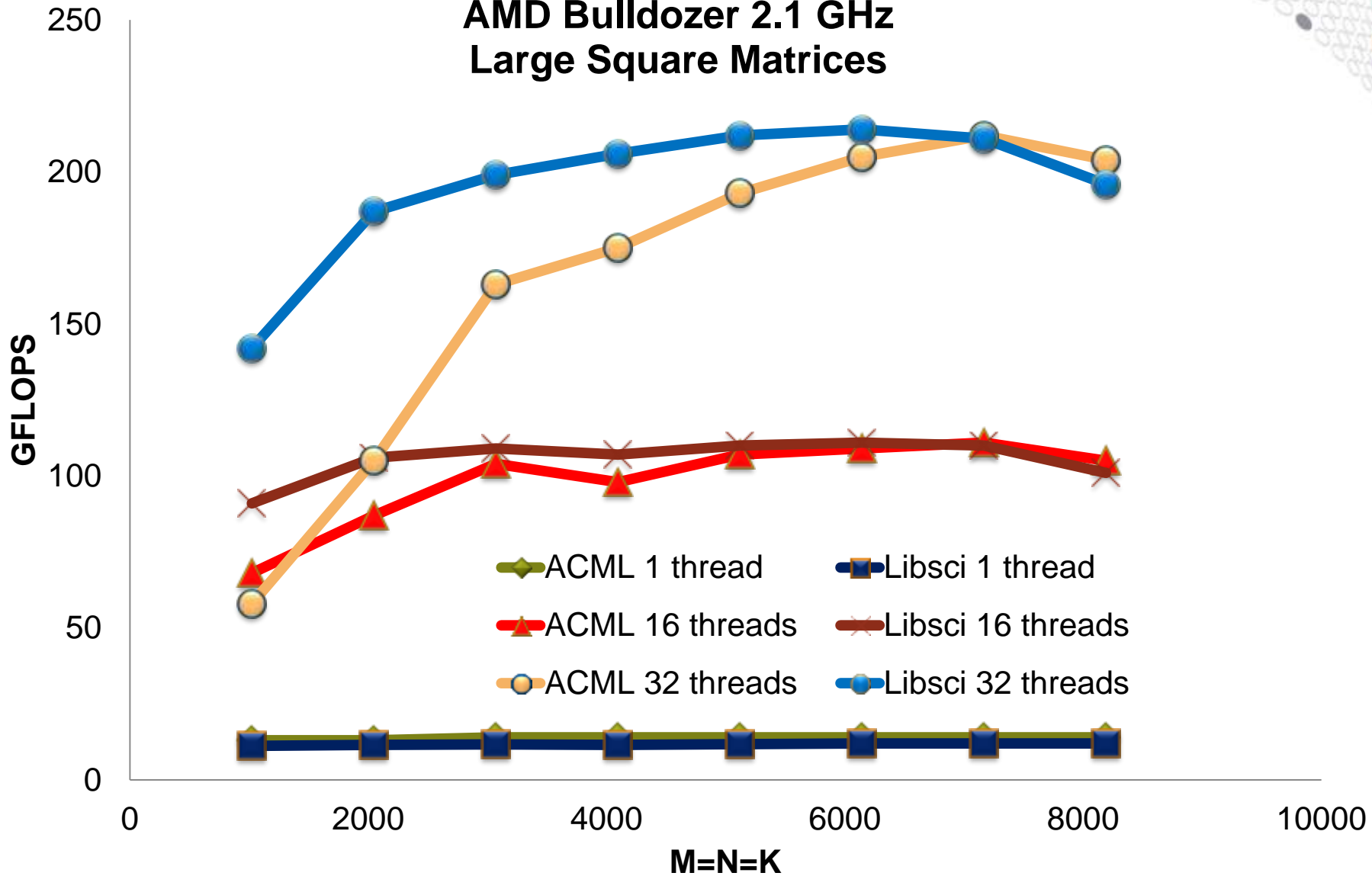
- **Design :**

1. Generalized GEMM implementation
2. Offline Auto tuning
3. Runtime performance modeling

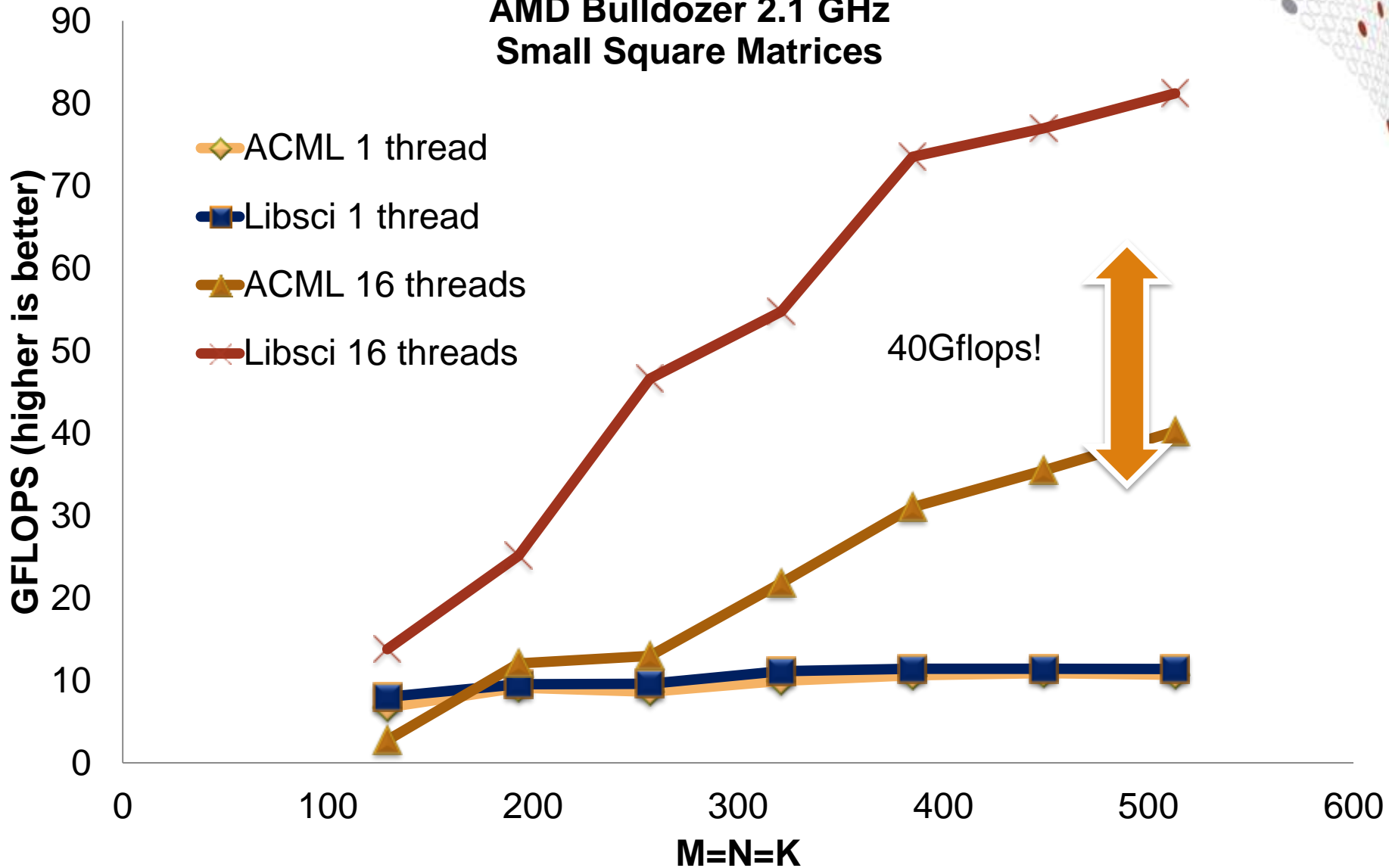
BFRAME infrastructure



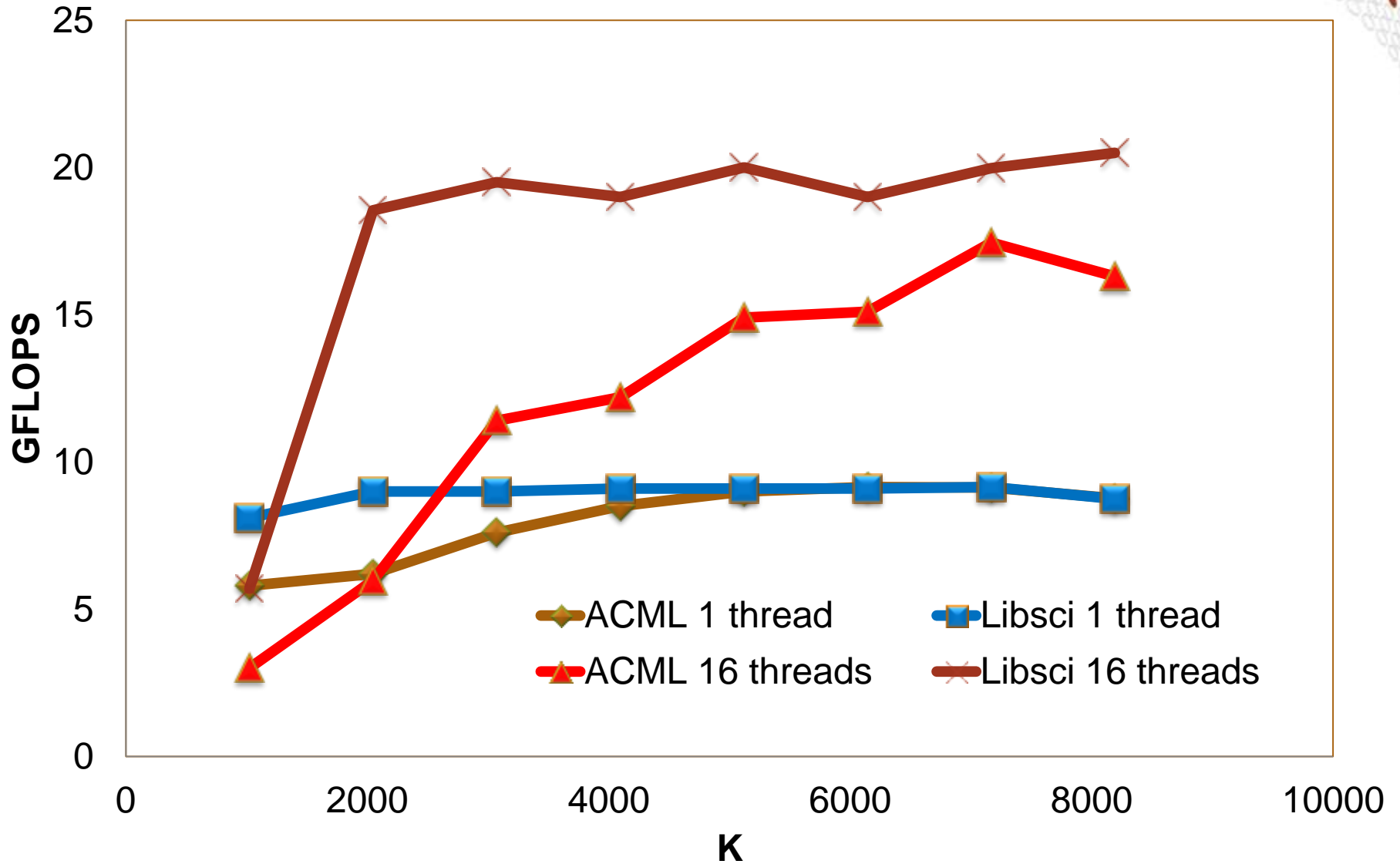
CrayBLAS DGEMM : AMD Bulldozer 2.1 GHz Large Square Matrices



CrayBLAS DGEMM : AMD Bulldozer 2.1 GHz Small Square Matrices



CrayBLAS DGEMM : AMD Bulldozer 2.1 GHzM : Varying K dimension



Tuning requests

- **CrayBLAS is an auto-tuned library**
 - Generally, excellent performance is possible for all shapes and sizes
- **However, even the adaptive CrayBLAS can be improved by tuning for exact sizes and shapes**
- **Send your specific tuning requirements to**

crayblas@cray.com

- **Just send the routine name, and the list of calling sequences**

Advanced optimizations for BLAS

- **For ZGEMM only**

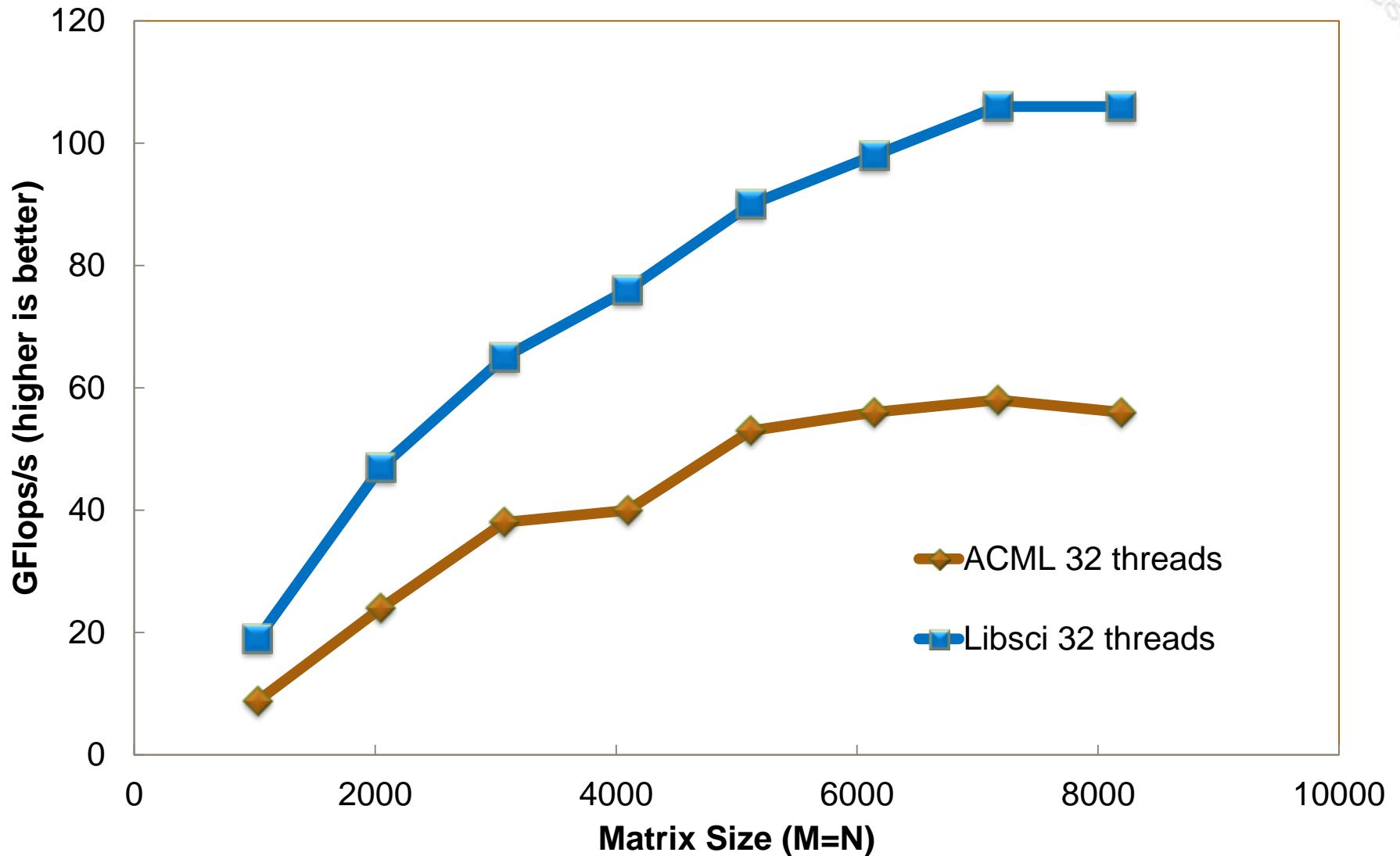
- Complex matrix multiplication can be performed using real matrix additions, for fewer flops
- You can turn on the 3M algorithm
- Set the environment variable
ZGEMM_USE_3M=1
- Note : there is an accuracy trade-off, though this should be safe most of the time

Threaded LAPACK

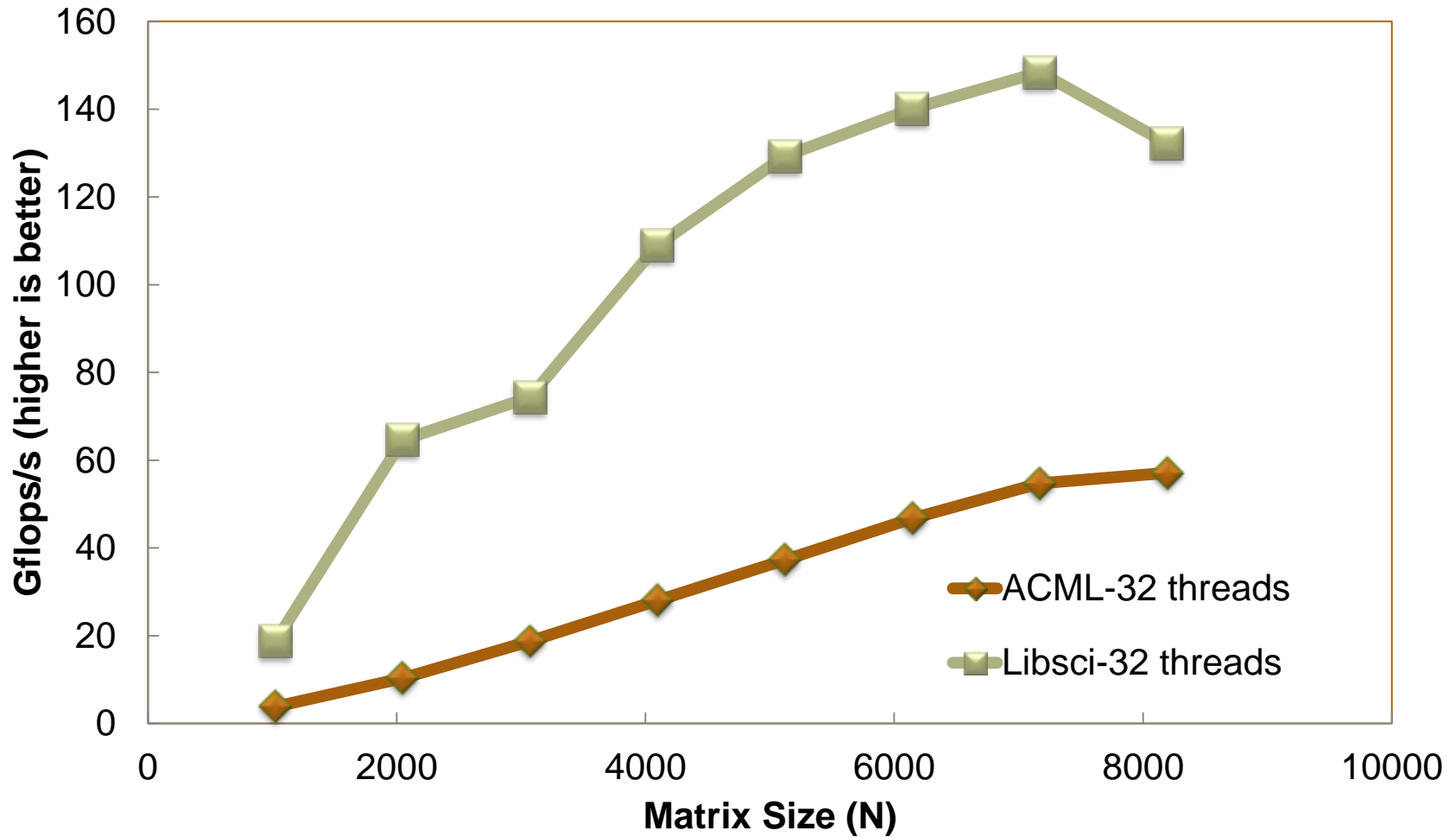
- Threaded LAPACK works exactly the same as threaded BLAS
- Anywhere LAPACK uses BLAS, those BLAS can be threaded
- Some LAPACK routines are threaded at the higher level
- No special instructions

LAPACK DGETRF (LU)

AMD Bulldozer 2.1Ghz :: July 2012



LAPACK DPOTRF (Cholesky) AMD Bulldozer 2.1Ghz ::Aug 2012

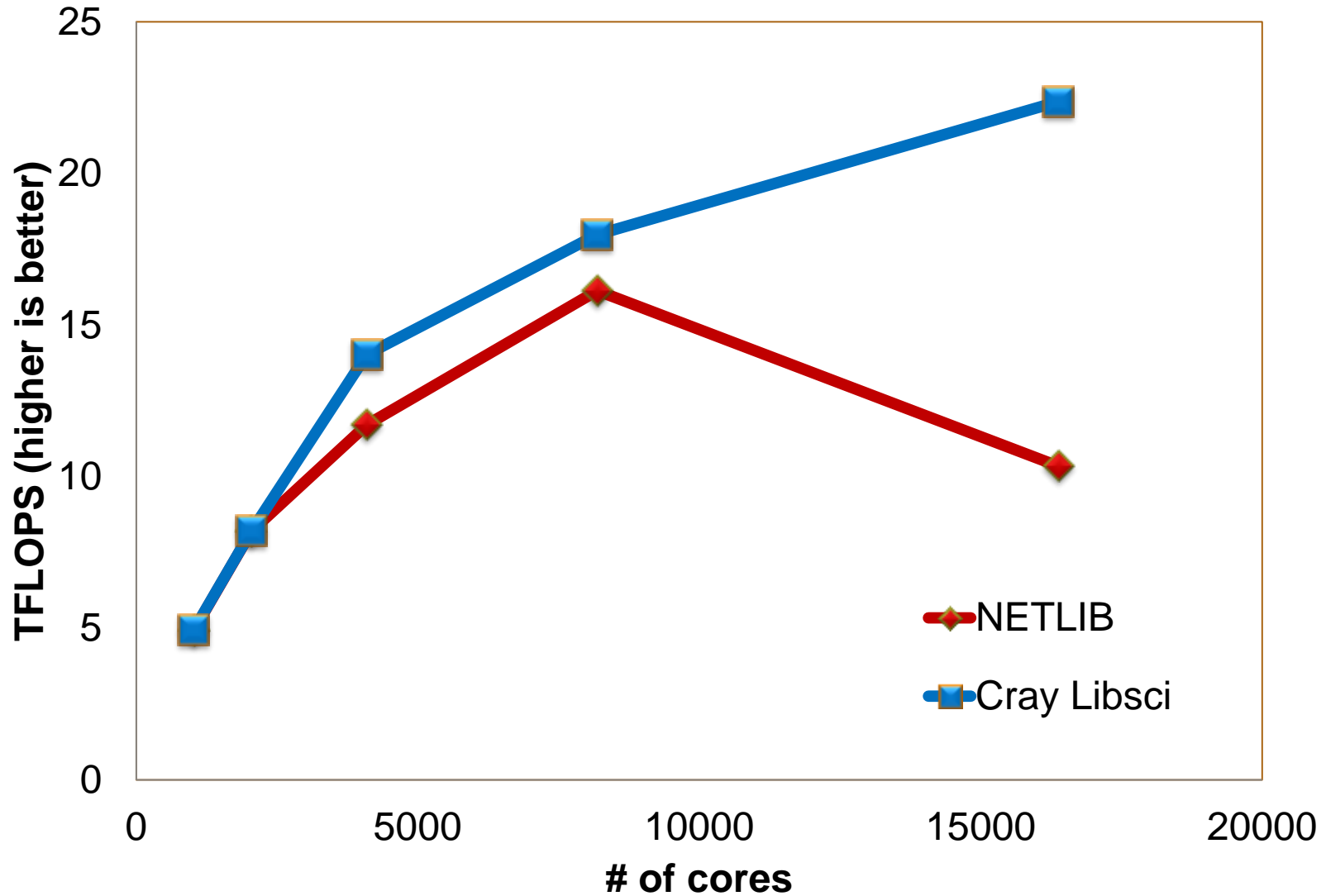


ScaLAPACK

- **ScaLAPACK in libsci is optimized for Gemini interconnect**
 - New collective communication procedures are added
 - Default topologies are changed to use the new optimizations
 - Much better strong scaling
- **It also benefits from the optimizations in CrayBLAS**
- **IRT can provide further improvements (see later)**

ScaLAPACK Double LU

Cray XE6 :: AMD Bulldozer 2.1 GHz
M = 131072 :: July 2012



Iterative Refinement Toolkit

- **Mixed precision can yield a big win on x86 machines.**
- **SSE (and AVX) units issue double the number of single precision operations per cycle.**
- **On CPU, single precision is always 2x as fast as double**
- **Accelerators sometimes have a bigger ratio**
 - Cell – 10x
 - Older NVIDIA cards – 7x
 - New NVIDIA cards (2x)
 - Newer AMD cards (> 2x)
- **IRT is a suite of tools to help exploit single precision**
 - A library for direct solvers
 - An automatic framework to use mixed precision under the covers

Iterative Refinement Toolkit - Library

- Various tools for solves linear systems in mixed precision
- Obtaining solutions accurate to double precision
 - For well conditioned problems
- Serial and Parallel versions of LU, Cholesky, and QR
- 2 usage methods
 - **IRT Benchmark routines**
 - Uses IRT 'under-the-covers' without changing your code
 - Simply set an environment variable
 - Useful when you cannot alter source code
 - **Advanced IRT API**
 - If greater control of the iterative refinement process is required
 - Allows
 - condition number estimation
 - error bounds return
 - minimization of either forward or backward error
 - 'fall back' to full precision if the condition number is too high
 - max number of iterations can be altered by users

IRT library usage

Decide if you want to use advanced API or benchmark API

benchmark API :

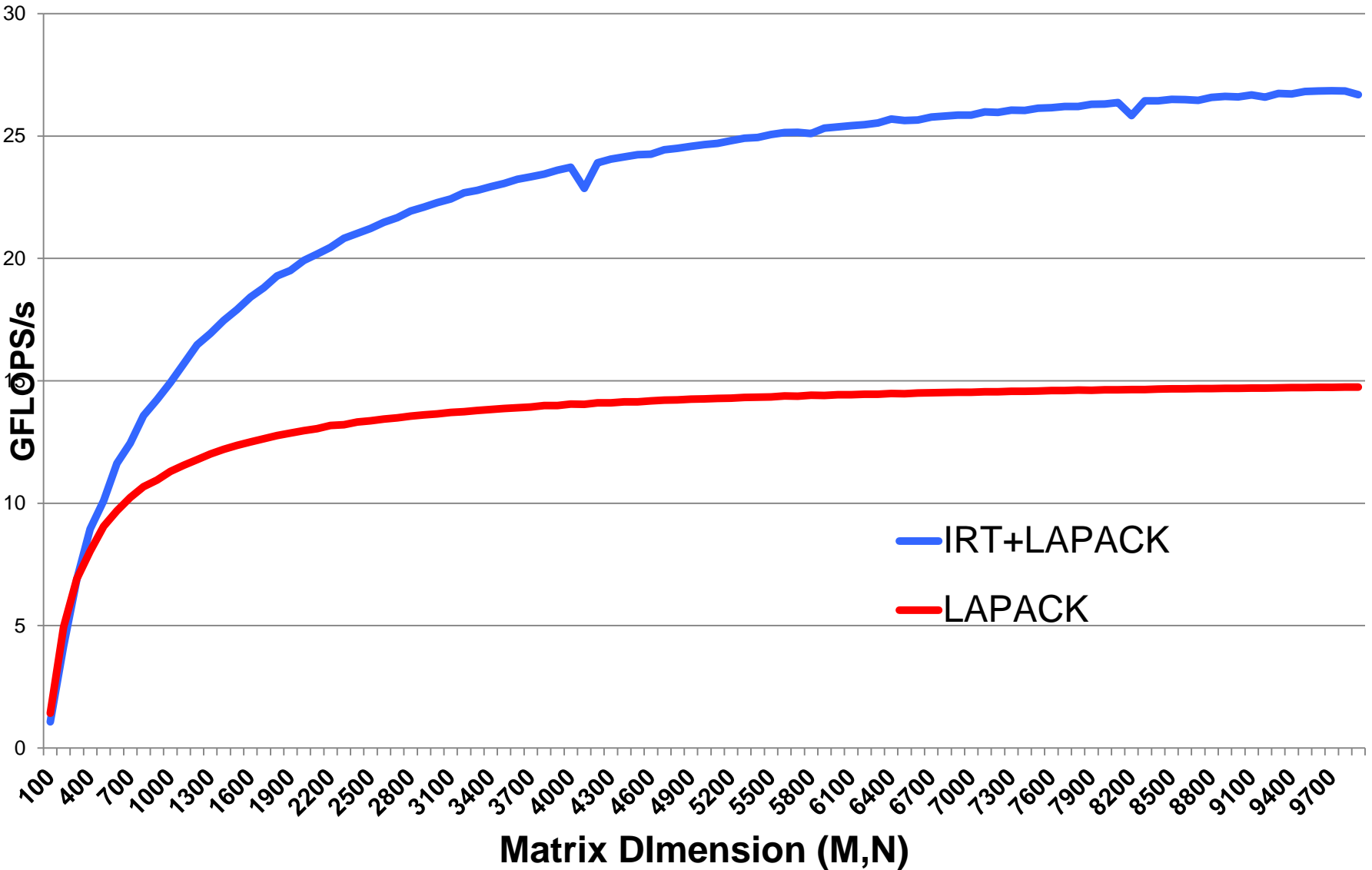
```
setenv IRT_USE_SOLVERS 1
```

advanced API :

1. locate the factor and solve in your code (LAPACK or ScaLAPACK)
2. Replace factor and solve with a call to IRT routine
 - e.g. dgesv -> irt_lu_real_serial
 - e.g. pzgesv -> irt_lu_complex_parallel
 - e.g. pzposv -> irt_po_complex_parallel
3. **Set advanced arguments**
 - Forward error convergence for most accurate solution
 - Condition number estimate
 - “fall-back” to full precision if condition number too high

Note : “info” does not return zero when using IRT !!

IRT with LAPACK LU DGETRF
AMD Bulldozer 2.1 GHz
2threads :: September 2012

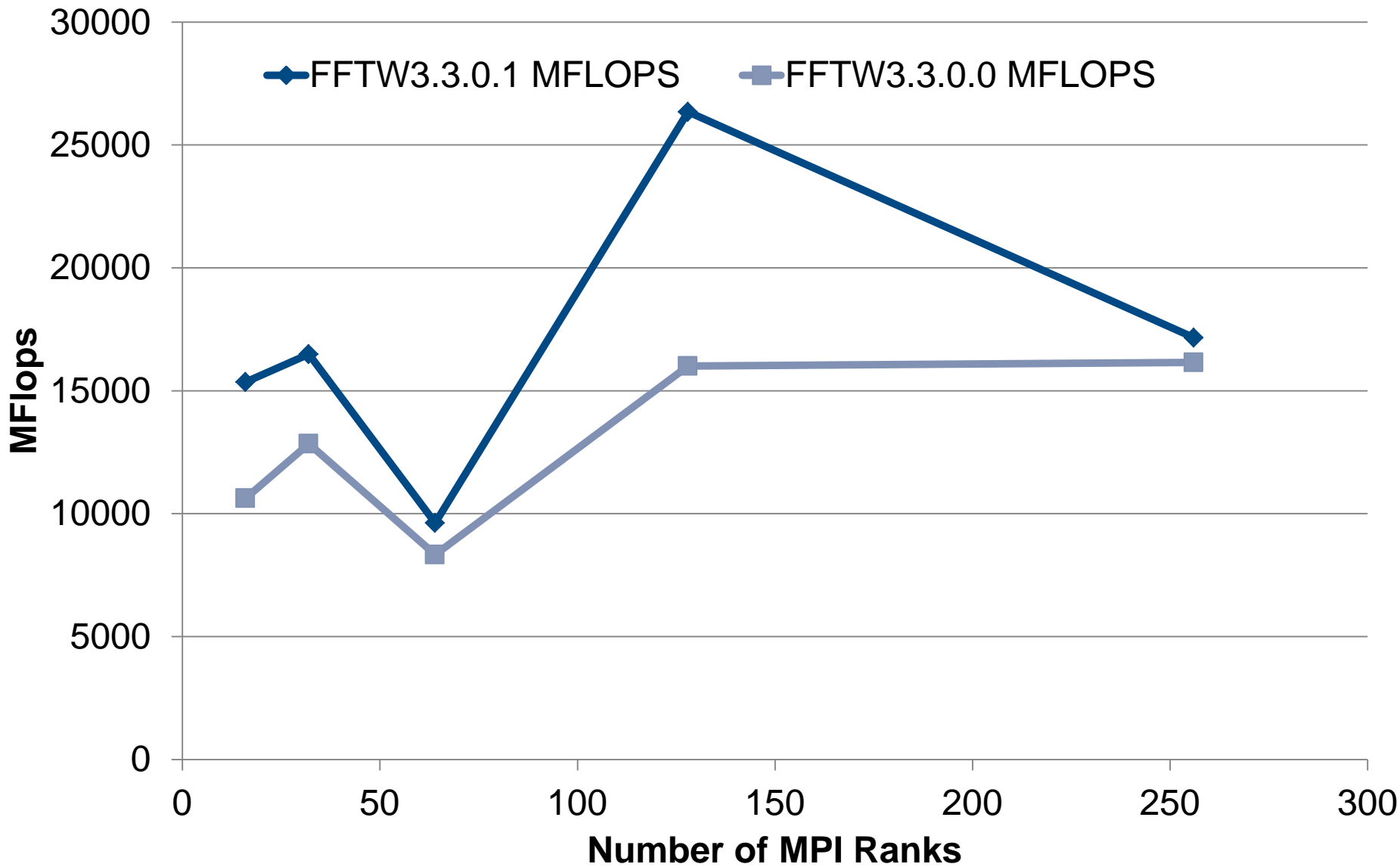


- **Cray's main FFT library is FFTW from MIT**
 - Some additional optimizations for Cray hardware
- **Usage is simple**
 - Load the module
 - In the code, call an FFTW plan
- **Cray's FFTW provides wisdom files for these systems**
- **You can use the wisdom files to skip the plan stage**
- **This can be a significant performance boost**
- **FFTW 3.1.0.1 includes Cray optimizations for IL processors**

FFTW 2d FFT 544x544

AMD Bulldozer : 2.1 GHz : 2 threads

July 2012



Cray Adaptive FFT (CRAFFT)

- **Serial CRAFFT is largely a productivity enhancer**
- **Also a performance boost due to “wisdom” usage**
- **Some FFT developers have problems such as**
 - Which library choice to use?
 - How to use complicated interfaces (e.g., FFTW)
- **Standard FFT practice**
 - Do a plan stage
 - Do an execute
- **CRAFFT is designed with simple-to-use interfaces**
 - Planning and execution stage can be combined into one function call
 - Underneath the interfaces, CRAFFT calls the appropriate FFT kernel

CRAFFT usage

1. Load module fftw/3.2.0 or higher.
2. Add a Fortran statement “use crafft”
3. call `crafft_init()`
4. Call `crafft transform` using none, some or all optional arguments (as shown in red)

In-place, implicit memory management :

```
call crafft_z2z3d(n1,n2,n3,input,ld_in,ld_in2,isign)
```

in-place, explicit memory management

```
call crafft_z2z3d(n1,n2,n3,input,ld_in,ld_in2,isign,work)
```

out-of-place, explicit memory management :

```
crafft_z2z3d(n1,n2,n3,input,ld_in,ld_in2,output,ld_out,ld_out2,isign,work)
```

Note : the user can also control the planning strategy of CRAFFT using the `CRAFFT_PLANNING` environment variable and the `do_exe` optional argument, please see the `intro_crafft` man page.

Parallel CRAFFT

- **Parallel CRAFFT is meant as a performance improvement to FFTW2 distributed transforms**
 - Uses FFTW3 for the serial transform
 - Uses ALLTOALL instead of ALLTOALLV where possible
 - Overlaps the local transpose with the parallel communications
 - Uses a more adaptive communication scheme based on input
 - Lots of more advanced research in one-sided messaging and active messages
- **Can provide impressive performance improvements over FFTW2**
- **Currently implemented**
 - complex-complex
 - Real-complex and complex-real
 - 3-d and 2-d
 - In-place and out-of-place
 - 1 data distribution scheme but looking to support more (please tell us)
 - C language support for serial and parallel
 - Generic interfaces for C users (use C++ compiler to get these)

parallel CRAFFT usage

1. Add “use crafft” to Fortran code
2. Initialize CRAFFT using `crafft_init`
3. Assume MPI initialized and data distributed (see manpage)
4. Call `crafft`, e.g. (optional arguments in red)

2-d complex-complex, in-place, internal mem management :

```
call crafft_pz2z2d(n1,n2,input,isign,flag,comm)
```

2-d complex-complex, in-place with no internal memory :

```
call crafft_pz2z2d(n1,n2,input,isign,flag,comm,work)
```

2-d complex-complex, out-of-place, internal mem manager :

```
call crafft_pz2z2d(n1,n2,input,output,isign,flag,comm)
```

2-d complex-complex, out-of-place, no internal memory :

```
crafft_pz2z2d(n1,n2,input,output,isign,flag,comm,work)
```

Each routine above has manpage. Also see 3d equivalent :

```
man crafft_pz2z3d
```

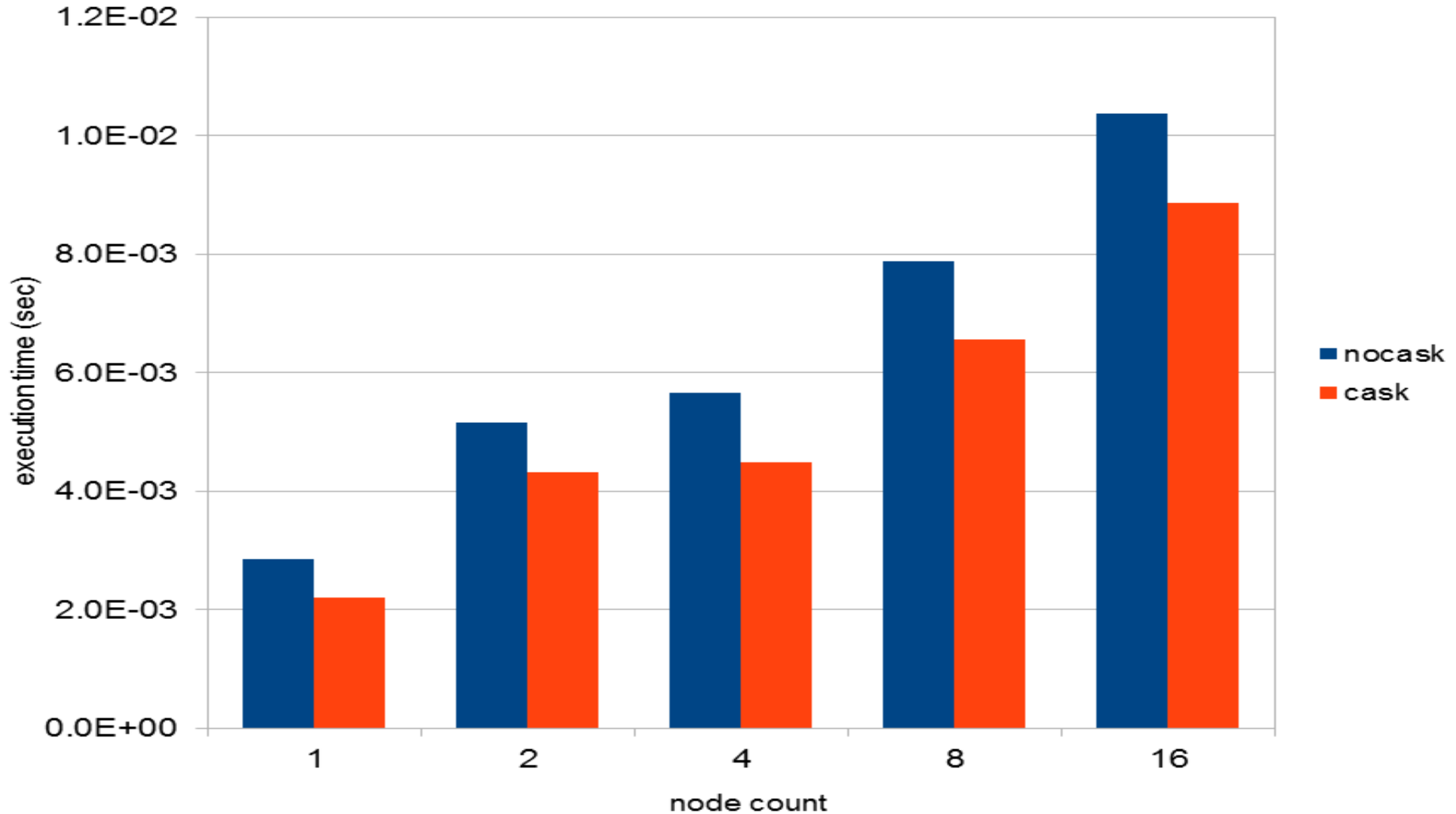
Cray Adaptive Sparse Kernel (CASK)



- Sparse matrix operations in PETSc and Trilinos on Cray systems are optimized via CASK
- CASK is a product developed at Cray using the Cray Auto-tuning Framework
- Offline :
 - ATF program builds many thousands of sparse kernel
 - Testing program defines matrix categories based on density, dimension etc
 - Each kernel variant is tested against each matrix class
 - Performance table is built and adaptive library constructed
- Runtime
 - Scan matrix at very low cost
 - Map user's calling sequence to nearest table match
 - Assign best kernel to the calling sequence
 - Optimized kernel used in iterative solver execution

CASK + PETSc AMD IL

PETSc ex50 Performance Summary
Driven cavity simulation



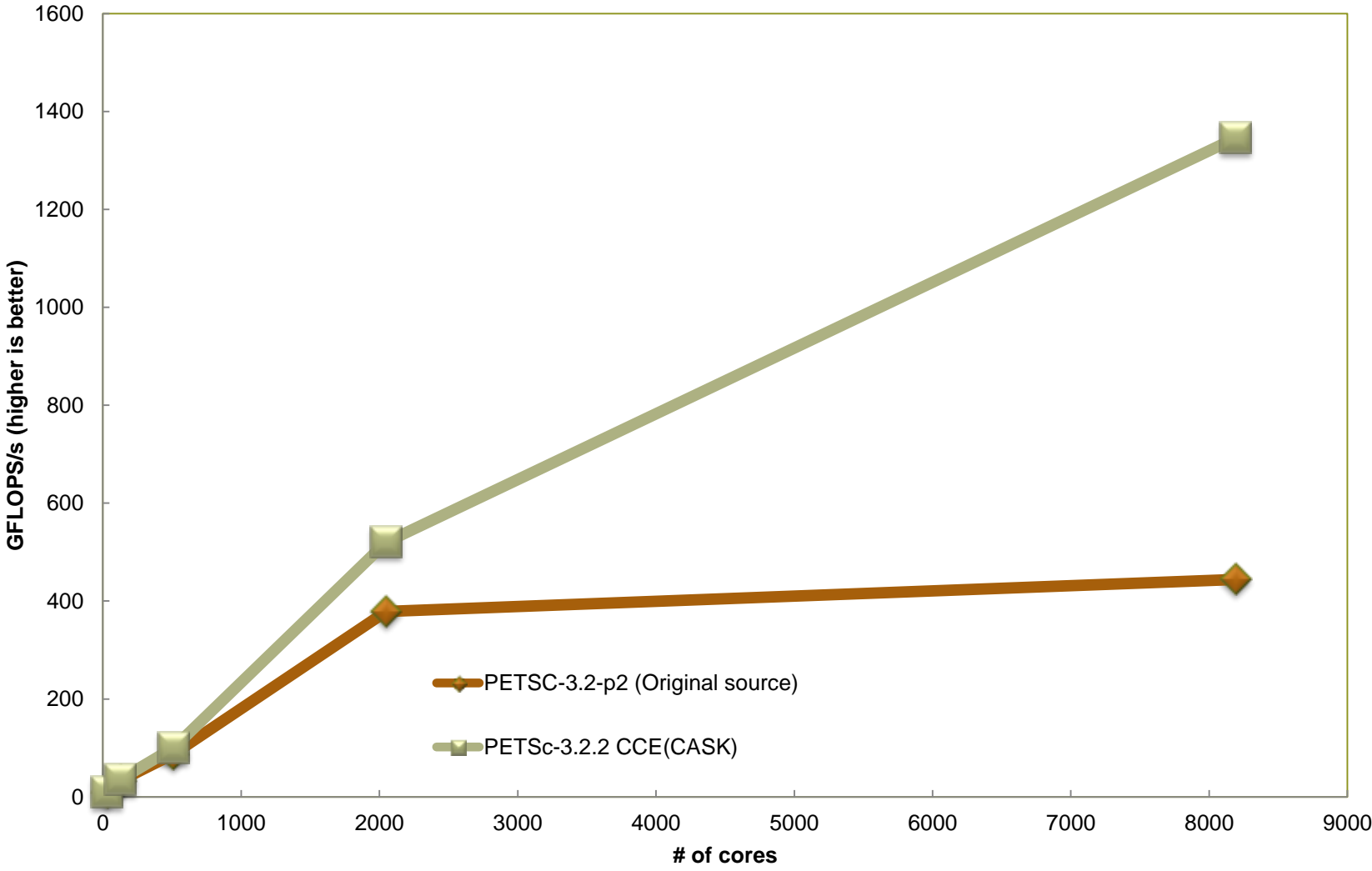
PETSc, Linear System Solution

2D Laplacian Problem

Weak Scalability

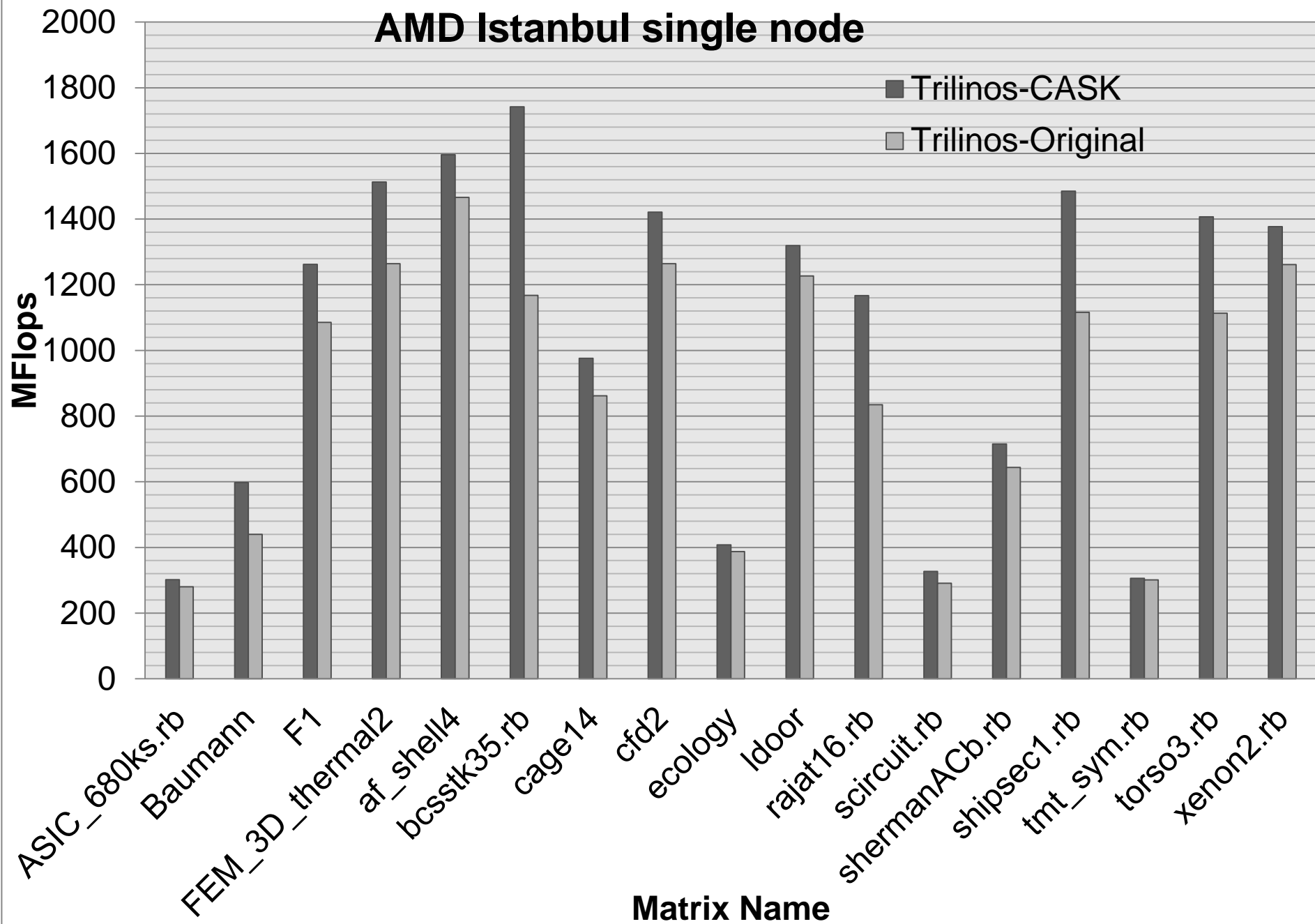
N=262,144 --- 268M

AMD Bulldozer 2.1G :: July 2012



Trilinos + CASK

AMD Istanbul single node



LibSci for Accelerators

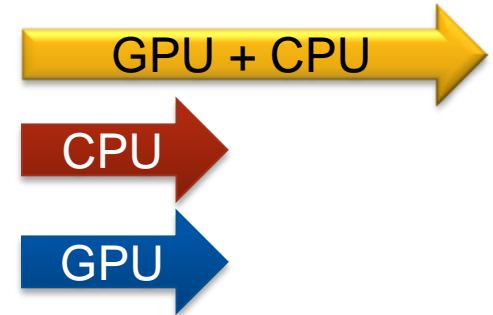
- **Provide basic libraries for accelerators, tuned for Cray**
- **Must be independent to openACC, but fully compatible**
- **Multiple use case support**
 - Get the base use of accelerators with no code change
 - Get extreme performance of GPU with or without code change
 - Extra tools for support of complex code
- **Incorporate the existing GPU libraries into libsci**
- **Provide additional performance and usability**
- **Maintain the Standard APIs where possible!**

Three interfaces for three use cases

- Simple interface

```
dgetrf(M, N, A, lda, ipiv, &info)
```

```
dgetrf(M, N, d_A, lda, ipiv, &info)
```



- Device interface

```
dgetrf_acc(M, N, d_A, lda, ipiv, &info)
```

- CPU interface

```
dgetrf_cpu(M, N, A, lda, ipiv, &info)
```


Usage - Basics

- Supports Cray and GNU compilers.
- Fortran and C interfaces (column-major assumed)
 - Load the module *craype-accel-nvidia20*.
 - Compile as normal (dynamic libraries used)
 - To enable threading in the CPU library, set OMP_NUM_THREADS
 - E.g. export OMP_NUM_THREADS=16
 - Assign 1 single MPI process per node
 - Multiple processes cannot share the single GPU
 - Execute your code as normal

Libsci_acc Example

- Starting with a code that relies on dgemm.
- The library will check the parameters at runtime.
- If the size of the matrix multiply is large enough, the library will run it on the GPU, handling all data movement behind the scenes.
- NOTE: Input and Output data are in CPU memory.

```
call dgemm('n','n',m,n,k,alpha,&  
          a,lda,b,ldb,beta,c,ldc)
```

Libsci_acc Example

- If the rest of the code uses OpenACC, it's possible to use the library with directives.
- All data management performed by OpenACC.
- Calls the device version of dgemm.
- All data is in CPU memory before and after data region.

```
!$acc data copy(a,b,c)
```

```
!$acc parallel
```

```
!Do Something
```

```
!$acc end parallel
```

```
!$acc host_data use_device(a,b,c)
```

```
call dgemm_acc('n','n',m,n,k,&  
              alpha,a,lda,&  
              b,ldb,beta,c,ldc)
```

```
!$acc end host_data
```

```
!$acc end data
```

Libsci_acc Example

- Libsci_acc is a bit smarter than this.
- Since 'a,' 'b', and 'c' are device arrays, the library knows it should run on the device.
- So just dgemm is sufficient.

```
!$acc data copy(a,b,c)
```

```
!$acc parallel
```

```
!Do Something
```

```
!$acc end parallel
```

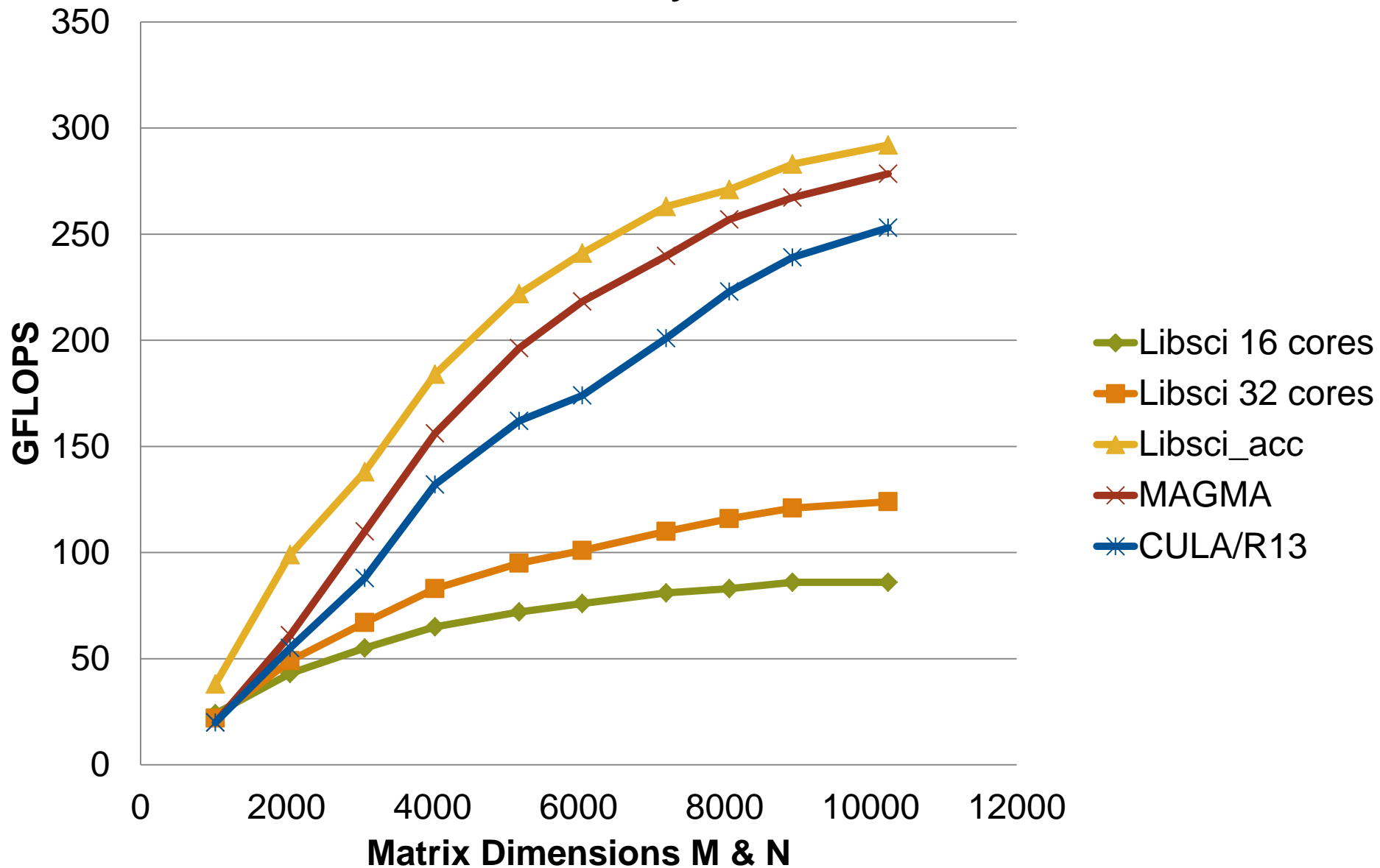
```
!$acc host_data use_device(a,b,c)
```

```
call dgemm      ('n','n',m,n,k,&  
                alpha,a,lda,&  
                b,ldb,beta,c,ldc)
```

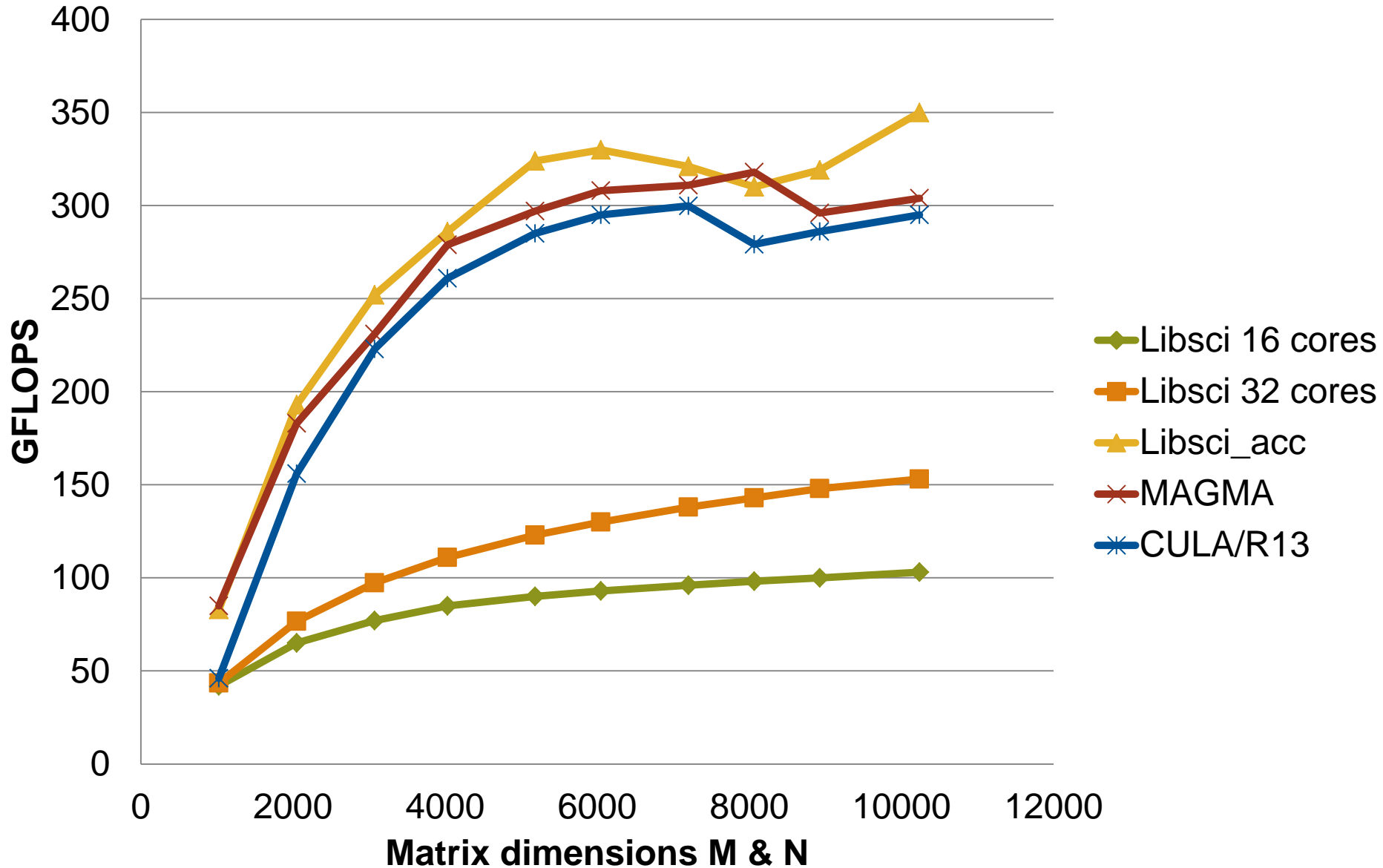
```
!$acc end host_data
```

```
!$acc end data
```

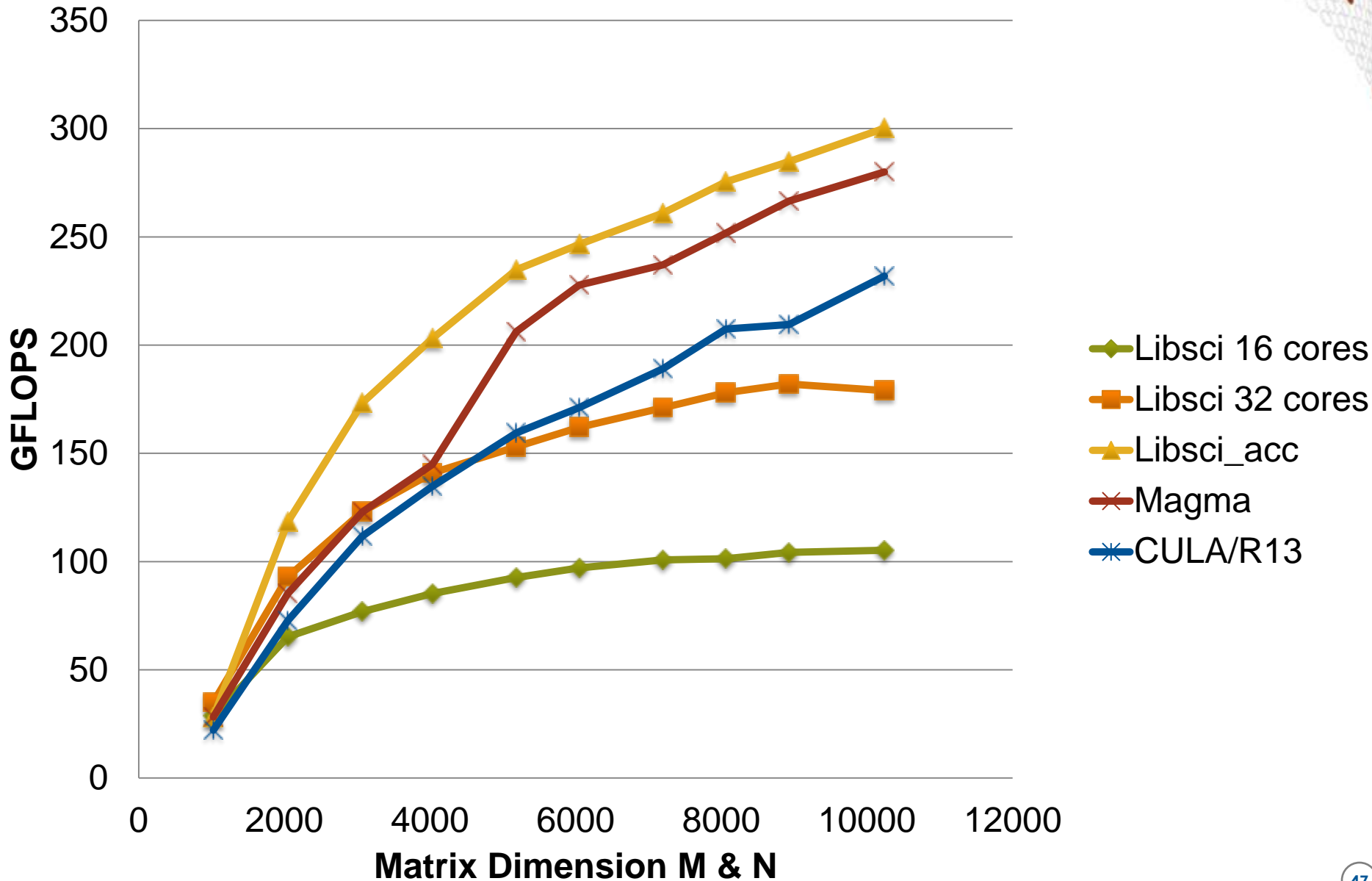
Libsci_acc : Double : LU (DGETRF) Cray XK6 (and XE6) July 2012



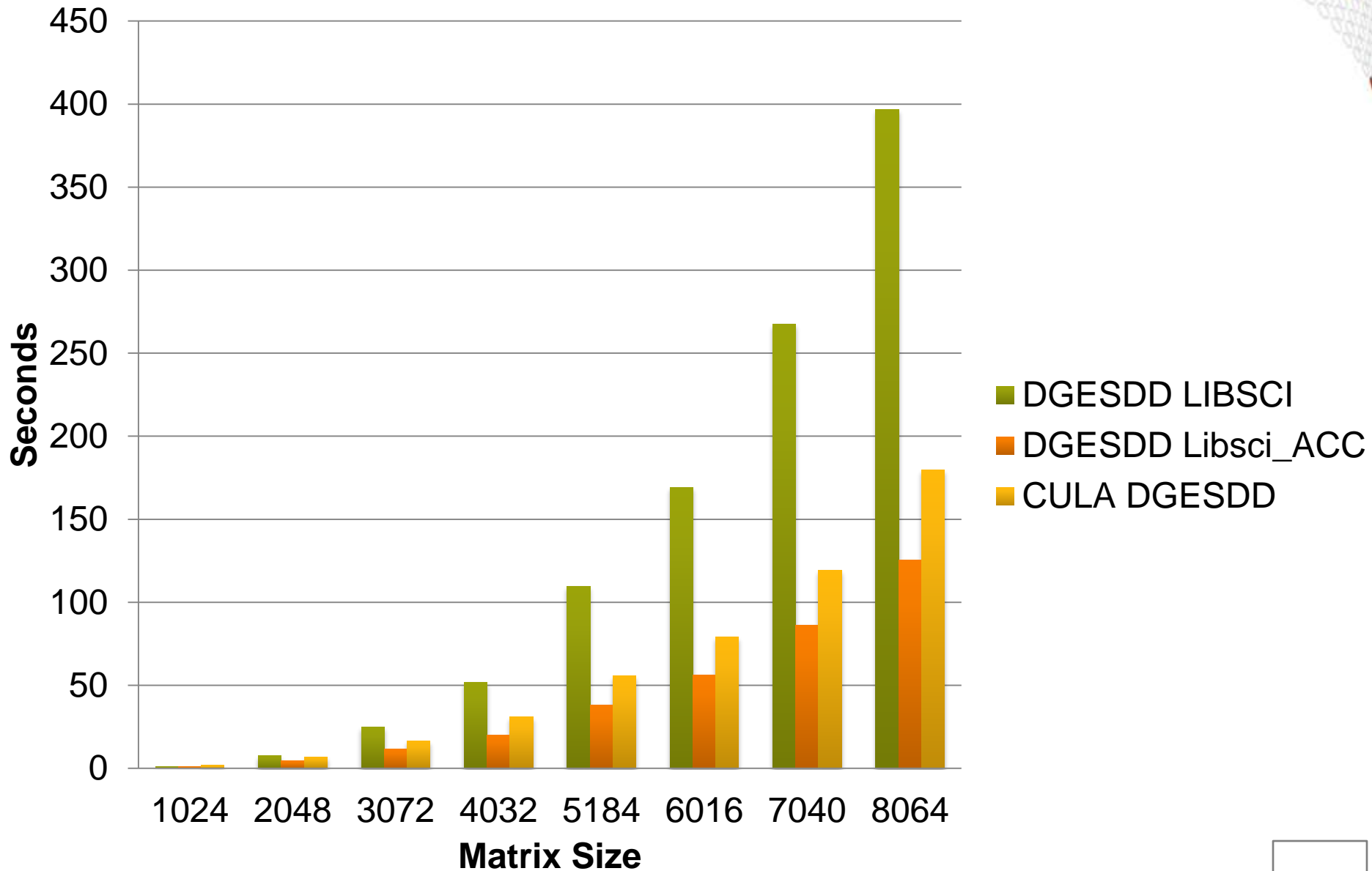
Libsci_acc : Double Complex LU (ZGETRF) Cray XK6 (and XE6) July 2012



Libsci_acc : Double Complex Cholesky (DPOTRF) Cray XK6 (and XE6) July 2012



libsci_acc : Double Complex LU (ZGETRF) Cray XK6 July 2012



Thanks

- **For specific CrayBLAS optimizations**
 - crayblas@cray.com
- **For more information on Cray libraries**
 - Cray Centre of Excellence
 - tedwards@cray.com