HPC Made Easy: Using Docker to Distribute and Test Trilinos

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HPC Made Easy: Using Docker to Distribute and Test Trilinos

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by Sean Deal

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Abstract

Virtualization is an enticing option for computer science research given its ability to provide repeatable, standardized environments, but traditional virtual machines have too much overhead cost to be practical. Docker, a Linux-based tool for operating-system level virtualization, has been quickly gaining popularity throughout the computer science field by touting a virtualization solution that is easily distributable and more lightweight than virtual machines. This thesis aims to explore if Docker is a viable option for conducting virtualized research by evaluating the results of parallel performance tests using the Trilinos project.
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1 Introduction

Virtualization - the isomorphic mapping of a virtual guest system to a real host system - is an enticing option for many computing tasks. In particular, virtual machines (VMs) are commonly used for several purposes. VMs can be summarized as virtual systems that replicate an entire machine - for instance, VM software such as Oracle VM VirtualBox can be used to run a virtual instance of Ubuntu Linux on a Windows machine or vice versa. This level of virtualization is achieved by placing virtualizing software between the software and hardware of the host machine. When creating an instance of a system virtual machine, virtualizing software provides a guest operating system and facilitates access to the native hardware of the host machine. In the case of differing system architectures between the guest and host machines, the virtualization software also translates machine instructions [24].

Virtual machines are sufficient for most casual computing needs, such as wanting to run Windows software on a Mac. Due to their standardized environments, VMs have also been used for more serious computing endeavors, such as software development and research. However, VMs tend to be too impractical for repeatable and reliable research, as they are not easily scalable and can complicate the pipeline of studies using different combinations of tools [3].

Docker is a new technology that has been making waves in the computer science field, touting a method of virtualization that is both easily distributable and more lightweight than virtual machines. As Docker has been rising in popularity, it has caught the attention of researchers as a possible method of simplifying repeatable research [3]. This thesis aims to test the performance of Docker containers compared to native hardware. By running tests from the Trilinos project in parallel, we will evaluate various performance metrics in both environments. We will then attempt to conclude whether or not Docker truly can be of use to computer science researchers and provide an easier way to conduct repeatable research without sacrificing performance.
2 Background

2.1 Docker

Docker is a tool for operating system-level virtualization, which is a server virtualization method where the operating system’s kernel allows for multiple isolated userspace instances, referred to as containers. This allows multiple users to run operations as if they are working on their own dedicated server, while these containers are being run off of a single server. In addition, the server administrator has power to regulate workloads across these isolated containers. Because these containers are completely isolated, operations executed in one container will not affect other containers, even if they are running simultaneously [17].

Docker started its life as a component of the ‘Platform as a Service’ provider dotCloud. In March 2013, dotCloud released Docker as an open source project. Docker was originally built using Linux Containers (LXC), a userspace interface for the Linux kernel that allows users to create and manage Linux containers. LXC’s primary goal “is to create an environment as close as possible to a standard Linux installation but without the need for a separate kernel” [5].

Docker was touted as a repeatable and lightweight virtualization solution due to its heavy focus on isolation of both resources and file systems [1]. The benefits of Docker were embraced immediately by developers. One of the most welcomed benefits was the use of Docker for environment standardization in development. Previously, testing environments varied at each step along the development cycle, but by using Docker, developers could ensure that the environments used to develop and test the software would be consistent [16].

Just about one year later, in March 2014, Docker was updated to version 0.9, which included a major change to Docker’s infrastructure. Instead of exclusively using LXC to access Linux container functionalities, the Docker group developed their own execution environment called libcontainer [15]. This environment allows Docker to have
direct access to container APIs instead of relying on outside technology, though Docker still supports LXC as well as other execution environments (Figure 1). This meant that Docker was now one complete package and also opened the door for Docker to run on non-Linux platforms [25]. By allowing Docker to become a self-contained complete package, libcontainer was monumental in Docker’s rise to the top of the Linux container community.

Figure 1: A diagram showing the various execution environments compatible with Docker as of Docker 0.9. libvirt, lxc, and systemd-nspawn are separate from the Docker engine, while libcontainer is part of Docker. This diagram also shows Linux container APIs used by Docker [25].

Docker has quickly become the de facto standard for operating system-level virtualization. dotCloud, Inc. officially changed its name to Docker, Inc. in October 2013 to reflect its change in focus from the dotCloud service to Docker [10]. Docker, Inc. proceeded to sell dotCloud to the German company cloudControl in 2014; cloudControl filed for bankruptcy in February 2016, shutting down the dotCloud service that originally spawned Docker [20]. Meanwhile, Docker has teamed up with high-profile companies such as Google, Microsoft, Amazon, and IBM to create the Open Container Initiative.
This project is an effort to make Docker the true standard for Linux containers. Ideally, the Open Container Initiative will make the Docker container format and runtime the basis of this new standard, meaning that developers will be able to run their containerized applications in any runtime \cite{18}.

2.2 Trilinos

Trilinos is an open-source project consisting of several packages used for scalable science and engineering applications. The initial goals of the Trilinos project regarded the development of production-quality mathematical solvers. The project garnered success and recognition early, receiving an R&D 100 award in 2004 \cite{19}. Presently, there are more than fifty packages in Trilinos covering a broad range of algorithms in the areas of computational science and engineering (Figure 2).

*Trilinos strategic capability areas:*

- User experience
- Parallel programming environments
- Framework and tools
- Software engineering technologies and integration
- I/O support
- Meshes, geometry, and load balancing
- Discretizations
- Scalable linear algebra
- Linear and eigensolvers
- Embedded nonlinear analysis tools

Figure 2: Trilinos strategic capability areas. The primary package used in this thesis, Epetra, falls under the area of scalable linear algebra \cite{26}.

Trilinos packages are self-contained software components, each with their own requirements and dependencies. Trilinos is predominantly a community-driven project, so keeping packages mostly isolated from each other allows Trilinos developers to focus mainly on their own package. However, packages can also be built in combination with each other. Many packages are built in close relation with others, providing expanded
functionality. In addition, Trilinos has support for more than eighty third-party libraries which can be used in tandem with packages in Trilinos \cite{13}.

2.2.1 Epetra

Epetra is a package that implements serial and parallel linear algebra and provides the foundation for Trilinos solvers. Epetra’s uses include construction and use of sparse graphs, sparse matrices, and dense vectors. The package also includes wrappers that provide simplified interaction with BLAS and LAPACK, two common linear algebra packages outside of Trilinos \cite{8}. Epetra was the primary package used for performance testing in this thesis, the details of which will be explained later.

3 Benefits of Docker

3.1 Development

Because Docker containers are isolated userspace instances, they provide standardized environments that could be beneficial for both development and bug reproduction. Instead of several developers working on the same project from different machines, creating one or more standardized Docker images would provide a standard environment for all the developers to work from. This would help fix errors during development that may arise due to different developers having different versions of tools used to build and run Trilinos. In addition, standardized Docker containers can reduce costs needed for developers to maintain their own development environments.

Any image can be run on any operating system that supports Docker. For instance, a developer running Ubuntu can pull and work from a container based on Fedora. This allows developers to test their software in several environments and also allows several developers to work in the same environment regardless of their host machines’ operating system (Figure \ref{fig:docker}). By having standardized images for issue handling, bugs can be reproduced in several different environments regardless of the host operating system.
of the issue handler.

One of the key goals of Trilinos is universal interoperability, meaning that any combination of packages and third-party libraries that makes algorithmic sense can be built into a specific installation of Trilinos \[13\]. However, a problem arises when attempting to use an installation of Trilinos that was built for a different purpose. If the current installation does not include necessary packages or third-party libraries, Trilinos must be completely re-built and re-installed with the new packages included. Sometimes this re-building process can be as simple as changing the configuration file, but considering the large number of packages and third-party libraries compatible with Trilinos, this is not always an easy process \[2\].

One of the most intriguing areas of potential benefit is the use of Dockerfiles for creating new builds of Trilinos. Dockerfiles are short scripts that are used to automatically create containers and run specified commands in them. This means that a simple Dockerfile can be used to configure, build, and install a new installation of Trilinos and provide an image with this new installation included. By providing different configuration files to the same Dockerfile, it is possible to create many different images containing different builds of Trilinos.

### 3.2 Distribution

A key function of Docker is the use of images in conjunction with containers. Docker images are essentially snapshots of containers that are used as bases from which other containers are created. At any time, a container can be committed to either the host image or a new image, functionally saving the changes made inside the container. These images can be shared through the Docker Hub Registry, a hosting service integrated into Docker which acts as a repository for Docker images (Figure \[3\]). Any user can pull any public images and, if they are registered with Docker Hub, push their own images to the Registry through simple Docker commands.

As mentioned previously, standardized environments are a benefit to develop-
Figure 3: A visualization of multiple Trilinos developers with different host operating systems working from the same image. Changes made by any developer can be pushed to the Docker Hub and then pulled by other developers.

Docker can provide an easy pathway to distributing Trilinos to end users. Currently, users of Trilinos have to download the Trilinos source, configure it for their purposes, build it, and install it before they are able to start using it for their own applications. By providing images that have Trilinos already installed, users can start building their own applications right away without having to go through the Trilinos build process. These users could then create an image of their own application that uses the Trilinos image as a base and distribute that image to users of their application.

3.3 Comparison to Virtual Machines

Docker containers are commonly compared to virtual machines. Both containers and VMs are isolated instances, and both are built and run from a base image. The main
One drawback to Docker's approach to virtualization is that it is entirely Linux-based, meaning Docker does not run natively on Mac or Windows machines. Instead, Mac and Windows users must run a custom VM through VirtualBox that allows access to all the same Docker functionalities. For these users, Docker provides Docker Toolbox, which includes everything needed to run the VM and start using Docker [6]. However, not all Windows and Mac machines are capable of virtualization, and even if they are, enabling virtualization can be a tedious process.

On March 24, 2016, Docker announced a new beta for Docker for Mac and Win-

---

**Figure 4:** A visualization comparing virtual machines and Docker containers. VMs include the entire guest OS, whereas Docker builds containers directly from the operating system through the Docker Engine [7].

---

difference is that virtual machine instances include the entire guest operating system, whereas Docker containers are run using the Linux kernel directly through the Docker engine. By using built-in Linux functions such as *cgroups* and *namespaces*, Docker containers create an isolated workspace on the same kernel that is significantly more lightweight than a virtual machine instance (Figure 4). In addition, Docker images are much smaller in size when compared to VM images; for instance, an Ubuntu VM image is roughly 943 MB [21], while an Ubuntu Docker image is only about 188 MB [23].
dows which eliminates the need to run a VM through VirtualBox. This new beta directly
utilizes xhyve and Hyper-V, the built-in virtualization tools on Mac and Windows respec-
tively, to run an Alpine Linux distribution which in turn runs the Docker application.
Instead of running a VM, users on Windows and Mac now simply have to run the Docker
application. This is an intriguing development that will likely make Docker much
easier to use on non-Linux platforms. It does not remove all problems, though, as users
still need to enable virtualization on their machines before being able to use this new
Docker application.

4 Performance Testing

With Docker providing a simplified way to distribute applications, its appeal has spread
to the area of computational research, including the field of high-performance computing
(HPC). Trilinos itself is not an application, rather a collection of libraries, but its packages
can be used for a wide range of algorithms and technologies in the areas of computational
science and engineering. If Docker containers allow for performance at a similar level
as a native installation, the process of conducting repeatable computing research could
be greatly simplified.

4.1 Methods

For this thesis, performance testing was conducted in two environments. The first was
an eight node cluster named Melchior at CSB/SJU. Each node of Melchior uses an Intel
Xeon processor with 12 cores (Appendix 6.1). The second environment was a series of
Docker containers running on each node of Melchior (Appendix 6.2), with each container
being built off of the same base image. The installations of Trilinos were identical in both
environments (Appendix 6.3).

The Message Passing Interface (MPI) was used to conduct performance tests
in parallel; specifically, OpenMPI was used. MPI is a realization of the message-passing
model of parallel computation, which consists of a set of processes that only have local
memory but can communicate with other processes by sending and receiving messages [11]. By programming with MPI, programs are able to split the workload between a number of separate processes. Understandably, MPI is used frequently by computer researchers, as it allows complex or computation-intensive tasks to be done much quicker.

As previously mentioned, Epetra is a package within Trilinos that implements serial and parallel linear algebra. While Trilinos does not provide standalone software, its packages have executable tests that can be used to evaluate performance. One of these is the Epetra BasicPerfTest. This test takes parameters for the size of a mesh grid and, if running in parallel, a matrix of processors. It then sets up a grid of the type Epetra Map of the specified size on each processor and performs the following operations for each element of each matrix:

- MatVec - A simple solve of the equation \( y = Ax \). The MatVec is performed with new and old implementations, with and without optimized storage, and with a Trans variable set to 0 and 1, indicating whether to solve for the transpose of \( A \). All combinations are performed ten times each, resulting in eighty operations total.

- Lower/Upper Solve - An LU factorization. Both lower and upper triangular solves are performed, varying optimized storage and transpose similarly to the MatVec for a total of eighty operations.

The test then creates a vector of the type Epetra MultiVector of the same length as the matrix used above and performs these operations:

- Norm2 - The Euclidean norm of the vector. This operation is performed ten times.

- Dot - The dot product of the vector with itself. This operation is performed ten times.

- Update - A linear combination of the vector with itself, following the equation \( w = \alpha x + \beta y \) with \( \alpha = 1.0 \) and \( \beta = 1.0 \). This operation is performed ten times.
For all operations, the BasicPerfTest returns a result in millions of floating-point operations per second (Mega FLOPs, or MFLOPs), defined as

\[
\frac{\text{Number of floating-point operations in a program}}{\text{Execution time} \times 10^6}
\]

This serves as a more reliable indicator of performance than simply recording the time spent to complete an operation, as MFLOPs values are solely dependent upon the machine and the program [22].

The test was run several times, varying both the number of processes and the problem size. Grid sizes of 1000, 2000, and 4000 square were used, and each grid size was tested using 1, 8, 16, and 48 processes. The total number of equations evaluated for a given test is equal to \( g^2 \times p \) where \( g \) is the grid size and \( p \) is the number of processes. For each case, the test was performed five times, and results were recorded for the new MatVec with optimized storage and Trans=0, the lower triangular solve with optimized storage and Trans=0, the 10 Norm2’s, the 10 Dot products, and the 10 Updates. The harmonic mean and median of each operation were then calculated [14]. This was repeated for both the native Trilinos installation on the Melchior cluster and the installation in Docker containers. For the Docker installations, an equal number of containers was used on each node of Melchior to match the number of processes; for example, with 16 processes, 2 Docker containers were used on each of the 8 nodes of Melchior.
4.2 Results

4.2.1 Problem Size - 1000x1000

Figure 5: Performance results for the Epetra BasicPerfTest with a problem size of 1000x1000 and 1 process.
Figure 6: Performance results for the Epetra BasicPerfTest with a problem size of 1000x1000 and 8 processes. One Docker container was used on each node of Melchior.

Figure 7: Performance results for the Epetra BasicPerfTest with a problem size of 1000x1000 and 16 processes. Two Docker containers were used on each node of Melchior.
Figure 8: Performance results for the Epetra BasicPerfTest with a problem size of 1000x1000 and 48 processes. Six Docker containers were used on each node of Melchior.

4.2.2 Problem Size - 2000x2000

Figure 9: Performance results for the Epetra BasicPerfTest with a problem size of 2000x2000 and 1 process.
Figure 10: Performance results for the Epetra BasicPerfTest with a problem size of 2000x2000 and 8 processes. One Docker container was used on each node of Melchior.

Figure 11: Performance results for the Epetra BasicPerfTest with a problem size of 2000x2000 and 16 processes. Two Docker containers were used on each node of Melchior.
Figure 12: Performance results for the Epetra BasicPerfTest with a problem size of 2000x2000 and 48 processes. Six Docker containers were used on each node of Melchior.

4.2.3 Problem Size - 4000x4000

Figure 13: Performance results for the Epetra BasicPerfTest with a problem size of 4000x4000 and 1 process.
Figure 14: Performance results for the Epetra BasicPerfTest with a problem size of 4000x4000 and 8 processes. One Docker container was used on each node of Melchior.

Figure 15: Performance results for the Epetra BasicPerfTest with a problem size of 4000x4000 and 16 processes. Two Docker containers were used on each node of Melchior.
When attempting to test a problem size of 4000x4000 with 48 processes on both Melchior and Docker, the test process was killed with a signal indicating that the process ran out of memory.

5 Conclusion

The results of running the Epetra BasicPerfTest both on the Melchior cluster and in Docker containers have shown that there is little to no drop in performance when performing tests using Docker containers. On serial performance tests, Melchior and Docker performed virtually the same regardless of problem size (Figures 5, 9, 13). When running in parallel with MPI, the results became slightly more erratic, though Docker consistently performed very similarly to the native installation. Larger problem sizes seemed to be more conclusive, as the results were more consistent with fewer outliers. Interestingly, Docker seemed to perform best with a larger number of processes, as the Docker containers outperformed the native installation more consistently on tests with 48 processes (Figures 8, 12). One possible explanation is the use of multiple containers on each node of Melchior for these tests. Using separate containers may have allowed each process to remain tied to a single processor, cutting down on performance lost by switching processors in the middle of the test. This certainly suggests that Docker handles scalable applications very well, though further testing may be required to definitively conclude if Docker actually improves scalability.

These results are very exciting, especially when combined with the other benefits of Docker explained earlier. Since Docker provides an easy way to distribute applications, the prospect of distributing research-related programs or packages is very promising. Specifically with Trilinos, this means that a pre-installed copy of Trilinos could be distributed through the Docker Hub, and users could begin developing and distributing their applications much quicker, and with no drop in performance.

Docker does not seem to be a complete silver bullet to virtual machines, though. The lack of native Docker support on Windows and Mac is notable and possibly unavoid-
able due to Docker’s reliance on native Linux commands to create containers. Still, Docker has shown that they are doing all they can to improve the experience of Windows and Mac users through the Docker Toolbox and the recently announced Docker for Mac and Windows beta, which eliminates the need for VirtualBox. For Linux users, Docker indeed appears to be a favorable alternative to virtual machines, and these results show that the HPC community can also find benefit in using Docker for scalable applications.

5.1 Future Research

Moving forward, similar performance testing will be done using different Trilinos packages, such as AztecOO, which works closely with Epetra to provide an object-oriented interface to the Aztec linear solver library [12]. This will serve to further explore the performance capabilities of Docker containers. It would also be useful to vary not only the number of processes but also the number of Docker containers set up on each node of Melchior. Doing so would demonstrate whether or not the number of containers has an effect on performance and would shed more light on the scalability of Docker containers.

In addition, we will likely move to create official Trilinos images in the Docker Hub Registry. This will realize the prospect of providing Trilinos users with an installed version of Trilinos that they can develop their applications against and distribute their applications on top of to their users. It may also lead to the use of a consistent development environment for Trilinos developers. In addition, with the recent announcement of Docker for Mac and Windows, the pathway to using Docker on non-Linux platforms is becoming easier. It would be beneficial to do more in-depth exploration into using Docker on these platforms.
References


[5] Linux Containers. What’s lxc?


[20] Jordan Novet. Dotcloud, the cloud service that gave birth to docker, is shutting down february 29, 2016.


[23] Docker Hub Registry. Ubuntu official repository.


6 Appendix

6.1 Computing Architecture

The computing architecture of the Melchior cluster is as follows:

```plaintext
processor : 0
vendor_id : GenuineIntel
cpu family : 6
model : 45
model name : Intel(R) Xeon(R) CPU E5-2420 0 @ 1.90GHz
stepping : 7
microcode : 0x710
cpu MHz : 1199.968
cache size : 15360 KB
physical id : 0
siblings : 12
core id : 0
cpu cores : 6
apicid : 0
initial apicid : 0
fpu : yes
fpu_exception : yes
cpuid level : 13
wp : yes
flags : fpu vme de pse tsc msr pae mce cx8 apic sep mtrr pge
        mca cmov pat pse36 clflush dts acpi mmx fxsr sse sse2 ss ht tm pbe
        syscall nx pdpe1gb rdtscp lm constant_tsc arch_perfmon pebs bts
        rep_good nopl xtopology nonstop_tsc apicperf eagerfpu pni
        pclmulqdq dtes64 monitor ds_cpl vmx smx est tm2 ssse3 cx16 xtpr pdcm
        pccid dca sse4_1 sse4_2 x2apic popcnt tsc_deadline_timer aes xsave
        avx lahf_lm ida arat epb pni pts dtherm tpr_shadow vnmi flexpriority
        ept vpid xsave opt
bogomips : 3790.61
clfush size : 64
```
6.2 Docker Setup Notes

The following notes come from CSB/SJU Linux administrator Josh Trutwin, who was invaluable in helping to get Docker working properly on Melchior. IP addresses and MAC addresses have been obscured for security.

Firstly install docker, which requires a custom CentOS repository to run on our Red Hat Enterprise Linux 7 Workstation environment:

```bash
# cat /etc/yum.repos.d/docker.repo
[dockerrepo]
name=Docker Repository
baseurl=https://yum.dockerproject.org/repo/main/centos/7
enabled=0
gpgcheck=1
gpgkey=https://yum.dockerproject.org/gpg
```

Install docker:

```bash
# yum -y --disablerepo="*" --enablerepo=dockerrepo install docker-engine
```

Setup a private network on 10.0.x.y on the second NIC on each node of the cluster:

```bash
# cat /etc/sysconfig/network-scripts/ifcfg-eth1
DEVICE=eth1
TYPE=Ethernet
HWADDR=--:--:--:--:--:--
BOOTPROTO=none
ONBOOT=yes
BRIDGE=br0
```
# cat /etc/sysconfig/network-scripts/ifcfg-br0
DEVICE=br0
TYPE=Bridge
IPADDR=10.0.0.4  <---- This is different for each HPC, node 0 is 10.0.0.1, 2 is 10.0.0.2, etc
NETMASK=255.255.0.0
BOOTPROTO=None
ONBOOT=yes
DELAY=0

# /sbin/sysctl -w net.ipv4.ip_forward=1

# service network restart

Edit the docker service ExecStart configuration to assign a portion of the 10.1.x.y network to each docker instance - for example, hpc3 below:

# cat /usr/lib/systemd/system/docker.service
[Unit]
Description=Docker Application Container Engine
Documentation=https://docs.docker.com
After=network.target docker.socket
Requires=docker.socket

[Service]
Type=notify
ExecStart=/usr/bin/docker daemon --bridge=br0 --fixed-cidr=10.1.4.0/24
-MountFlags=slave
-LimitNOFILE=1048576
-LimitNPROC=1048576
-LimitCORE=infinity

[Install]
WantedBy=multi-user.target
Start Docker, set to run on boot:

```
# systemctl enable docker.service
# systemctl start docker.service
```

Verify:

```
[root@hpc3 ~]# docker run -it centos /bin/bash
[...]
[root@e11e04cb0b6b /]# ip addr show
1: lo: <LOOPBACK,UP,LOWER_UP> mtu 65536 qdisc noqueue state UNKNOWN
    link/loopback :00:00:00:00:00 brd 00:00:00:00:00:00
    inet ---.---.---.---/8 scope host lo
       valid_lft forever preferred_lft forever
    inet6 ::1/128 scope host
       valid_lft forever preferred_lft forever
35: eth0@if36: <BROADCAST,MULTICAST,UP,LOWER_UP> mtu 1500 qdisc noqueue state UP
    link/ether --:--:--:--:--:-- brd ff:ff:ff:ff:ff:ff link-netnsid 0
    inet 10.1.4.12/16 scope global eth0
       valid_lft forever preferred_lft forever
    inet6 ----::---:---:----:---/64 scope link
       valid_lft forever preferred_lft forever
```

### 6.3 Trilinos Configuration Script

The same configuration script was used for the Trilinos installation on Melchior and the installation in Docker.

```
rm -rf CMakeCache.txt CMakeFiles/
EXTRA_ARGS=$@
cmake \
-DCMAKE_BUILD_TYPE:STRING=RELEASE \
-DCMAKE_INSTALL_PREFIX=../install \
```
-DTPL_ENABLE_MPI:BOOL=ON \
-DMPI_BASE_DIR:PATH=/usr/lib64/openmpi \
-DTrilinos_ENABLE_OpenMP:BOOL=ON \
-D Trilinos_ENABLE_TESTS:BOOL=ON \
-D Trilinos_ENABLE_ALL_PACKAGES:BOOL=OFF \
-D Trilinos_ENABLE_Epetra:BOOL=ON \
-DTrilinos_ENABLE_CXX11=ON \
-DTrilinos_ASSERT_MISSING_PACKAGES=OFF \
-DBUILD_SHARED_LIBS:BOOL=OFF \
-D CMAKE_VERBOSE_MAKEFILE:BOOL=OFF \
-D Trilinos_VERBOSE_CONFIGURE:BOOL=OFF \
$EXTRA_ARGS \
../publicTrilinos

6.4 Epetra BasicPerfTest Source Code

//@HEADER
//
******************************************************************************
//
// Epetra: Linear Algebra Services Package
//
Copyright 2011 Sandia Corporation
//
// Under the terms of Contract DE-AC04-94AL85000 with Sandia
// Corporation,
// the U.S. Government retains certain rights in this software.
//
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// PROFITS; OR BUSINESS INTERRUPTION) HOWEVER CAUSED AND ON ANY THEORY
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//
// Questions? Contact Michael A. Heroux (maherou@sandia.gov)
//
//*************************************************************************

//@HEADER
// prototypes

void GenerateCrsProblem(int numNodesX, int numNodesY, int numProcsX, int numProcsY, int numPoints,
int *xoff, int *yoff,
const Epetra_Comm &comm, bool verbose, bool summary,
Epetra_Map *&map,
Epetra_CrsMatrix *&A,
Epetra_Vector *&b,
Epetra_Vector *&bt,
Epetra_Vector *&xexact, bool StaticProfile, bool MakeLocalOnly);

void GenerateCrsProblem(int numNodesX, int numNodesY, int numProcsX,
int numProcsY, int numPoints,
    int * xoff, int * yoff, int nrhs,
    const Epetra_Comm &comm, bool verbose, bool summary,
    Epetra_Map *& map,
    Epetra_CrsMatrix *& A,
    Epetra_MultiVector *& b,
    Epetra_MultiVector *& bt,
    Epetra_MultiVector *&xexact, bool StaticProfile,
        bool MakeLocalOnly);

void GenerateVbrProblem(int numNodesX, int numNodesY, int numProcsX,
int numProcsY, int numPoints,
    int * xoff, int * yoff,
    int nsizes, int * sizes,
    const Epetra_Comm &comm, bool verbose, bool summary,
    Epetra_BlockMap *& map,
    Epetra_VbrMatrix *& A,
    Epetra_Vector *& b,
    Epetra_Vector *& bt,
    Epetra_Vector *&xexact, bool StaticProfile,
        bool MakeLocalOnly);

void GenerateVbrProblem(int numNodesX, int numNodesY, int numProcsX,
int numProcsY, int numPoints,
    int * xoff, int * yoff,
    int nsizes, int * sizes, int nrhs,
    const Epetra_Comm &comm, bool verbose, bool summary,
Epetra_BlockMap * & map,
Epetra_VbrMatrix * & A,
Epetra_MultiVector * & b,
Epetra_MultiVector * & bt,
Epetra_MultiVector * & xexact, bool StaticProfile,
  bool MakeLocalOnly);

void GenerateMyGlobalElements(int numNodesX, int numNodesY, int
numProcsX, int numProcs,
  int myPID, int * & myGlobalElements);

void runMatrixTests(Epetra_CrsMatrix * A, Epetra_MultiVector * b,
Epetra_MultiVector * bt,
  Epetra_MultiVector * xexact, bool StaticProfile,
  bool verbose, bool summary);

void runLUMatrixTests(Epetra_CrsMatrix * L, Epetra_MultiVector * bL,
Epetra_MultiVector * btL, Epetra_MultiVector * xexactL,
  Epetra_CrsMatrix * U, Epetra_MultiVector * bU,
  Epetra_MultiVector * btU, Epetra_MultiVector *
  xexactU,
  bool StaticProfile, bool verbose, bool summary);

int main(int argc, char * argv[])
{
  int ierr = 0;
  double elapsed_time;
  double total_flops;
  double MFLOPs;

  #ifdef EPETRA_MPI

  // Initialize MPI
  MPI_Init(&argc, &argv);
  Epetra_MpiComm comm( MPI_COMM_WORLD );

  #endif
else
        Epetra_SerialComm comm;
#endif

bool verbose = false;
bool summary = false;

// Check if we should print verbose results to standard out
if (argc>6) if (argv[6][0]=='-' && argv[6][1]=='v') verbose = true;

// Check if we should print verbose results to standard out
if (argc>6) if (argv[6][0]=='-' && argv[6][1]=='s') summary = true;

if(argc < 6) {
    cerr << "Usage:
         " << argv[0] << " NumNodesX NumNodesY NumProcX NumProcY NumPoints [-v|-s]"
         << endl
    << "where:
         " << NumNodesX << " Number of mesh nodes in X direction per processor" << endl
    << NumNodesY << " Number of mesh nodes in Y direction per processor" << endl
    << NumProcX << " Number of processors to use in X direction" << endl
    << NumProcY << " Number of processors to use in Y direction" << endl
    << NumPoints << " Number of points to use in stencil (5, 9 or 25 only)" << endl
    << "-v|-s (Optional) Run in verbose mode if -v present or summary mode if -s present" << endl
    << "NOTES: NumProcX*NumProcY must equal the number of processors used to run the problem."
    << "Serial example:
         " << argv[0] << " 16 12 1 1 25 -v" << endl
    << argv[0] << " 16 12 1 1 25 -s" << endl
Run this program in verbose mode on 1 processor using a 16x12 grid with a 25 point stencil."

MPI example:

mpirun -np 32 "${argv[0]} 10 12 4 8 9 -v"

Run this program in verbose mode on 32 processors putting a 10x12 subgrid on each processor using 4 processors in the X direction and 8 in the Y direction. Total grid size is 40 points in X and 96 in Y with a 9 point stencil.

return(1);

//char tmp;

//if (comm.MyPID()==0) cout << "Press any key to continue..." << endl;

//if (comm.MyPID()==0) cin >> tmp;

//comm.Barrier();

comm.SetTracebackMode(0); // This should shut down any error traceback reporting

if (verbose && comm.MyPID()==0)
    cout << Epetra_Version() << endl << endl;

if (summary && comm.MyPID()==0) {
    if (comm.NumProc()==1)
        cout << Epetra_Version() << endl << endl;
    else
        cout << endl << endl; // Print two blank line to keep output columns lined up
}

if (verbose) cout << comm <<endl;
// Redefine verbose to only print on PE 0

if (verbose && comm.MyPID()!=0) verbose = false;
if (summary && comm.MyPID()!=0) summary = false;

int numNodesX = atoi(argv[1]);
int numNodesY = atoi(argv[2]);
int numProcsX = atoi(argv[3]);
int numProcsY = atoi(argv[4]);
int numPoints = atoi(argv[5]);

if (verbose || (summary && comm.NumProc()==1)) {
  cout << "Number of local nodes in X direction = " << numNodesX << endl
       << "Number of local nodes in Y direction = " << numNodesY << endl
       << "Number of global nodes in X direction = " << numNodesX*numProcsX << endl
       << "Number of global nodes in Y direction = " << numNodesY*numProcsY << endl
       << "Number of local nonzero entries = " << numNodesX*numNodesY*numPoints << endl
       << "Number of global nonzero entries = " << numNodesX*numNodesY*numPoints*numProcsX*numProcsY << endl
       << "Number of Processors in X direction = " << numProcsX << endl
       << "Number of Processors in Y direction = " << numProcsY << endl
       << "Number of Points in stencil = " << numPoints << endl;
}

// Print blank line to keep output columns lined up
if (summary && comm.NumProc()>1)
cout << endl << endl << endl << endl << endl << endl << endl <<
endl<< endl << endl;

if (numProcsX*numProcsY!=comm.NumProc()) {
  cerr << "Number of processors = " << comm.NumProc() << endl
  << "is not the product of " << numProcsX << " and " <<
  numProcsY << endl << endl;
  return(1);
}

if (numPoints!=5 && numPoints!=9 && numPoints!=25) {
  cerr << "Number of points specified = " << numPoints << endl
  << "is not 5, 9, 25" << endl << endl;
  return(1);
}

if (numNodesX*numNodesY <=0) {
  cerr << "Product of number of nodes is <= zero" << endl << endl;
  return(1);
}

Epetra_IntSerialDenseVector Xoff, XLoff, XUoff;
Epetra_IntSerialDenseVector Yoff, YLoff, YUoff;
if (numPoints==5) {
  // Generate a 5-point 2D Finite Difference matrix
  Xoff.Size(5);
  Yoff.Size(5);

  // Generate a 2-point 2D Lower triangular Finite Difference matrix
  XLoff.Size(2);
  YLoff.Size(2);
XLoff[0] = -1; XLoff[1] = 0;
YLoff[0] = 0; YLoff[1] = -1;

// Generate a 3-point 2D upper triangular Finite Difference matrix
XUoff.Size(3);
YUoff.Size(3);
XUoff[0] = 0; XUoff[1] = 1; XUoff[2] = 0;
YUoff[0] = 0; YUoff[1] = 0; YUoff[2] = 1;
}
else if (numPoints==9) {

// Generate a 9-point 2D Finite Difference matrix
Xoff.Size(9);
Yoff.Size(9);
Xoff[0] = -1; Xoff[1] = 0; Xoff[2] = 1;
Yoff[0] = -1; Yoff[1] = -1; Yoff[2] = -1;

// Generate a 5-point lower triangular 2D Finite Difference matrix
XLoff.Size(5);
YLoff.Size(5);
XLoff[0] = -1; XLoff[1] = 0; XLoff[2] = 1;
YLoff[0] = -1; YLoff[1] = -1; YLoff[2] = -1;
YLoff[3] = 0; YLoff[4] = 0;

// Generate a 4-point upper triangular 2D Finite Difference matrix
XUoff.Size(4);
YUoff.Size(4);
XUoff[0] = 1;
YUoff[0] = 0;
}

else {
    // Generate a 25-point 2D Finite Difference matrix
    Xoff.Size(25);
    Yoff.Size(25);
    int xi = 0, yi = 0;
    int xo = -2, yo = -2;
    Xoff[xi++] = xo++; Xoff[xi++] = xo++; Xoff[xi++] = xo++; Xoff[xi++] = xo++; Xoff[xi++] = xo++;
    Yoff[yi++] = yo ; Yoff[yi++] = yo ; Yoff[yi++] = yo ; Yoff[yi++] = yo ; Yoff[yi++] = yo ;
    xo = -2, yo++;
    Xoff[xi++] = xo++; Xoff[xi++] = xo++; Xoff[xi++] = xo++; Xoff[xi++] = xo++; Xoff[xi++] = xo++;
    Yoff[yi++] = yo ; Yoff[yi++] = yo ; Yoff[yi++] = yo ; Yoff[yi++] = yo ; Yoff[yi++] = yo ;
    xo = -2, yo++;
    Xoff[xi++] = xo++; Xoff[xi++] = xo++; Xoff[xi++] = xo++; Xoff[xi++] = xo++; Xoff[xi++] = xo++;
    Yoff[yi++] = yo ; Yoff[yi++] = yo ; Yoff[yi++] = yo ; Yoff[yi++] = yo ; Yoff[yi++] = yo ;
    xo = -2, yo++;
    Xoff[xi++] = xo++; Xoff[xi++] = xo++; Xoff[xi++] = xo++; Xoff[xi++] = xo++; Xoff[xi++] = xo++;
    Yoff[yi++] = yo ; Yoff[yi++] = yo ; Yoff[yi++] = yo ; Yoff[yi++] = yo ; Yoff[yi++] = yo ;
    xo = -2, yo++;
    Xoff[xi++] = xo++; Xoff[xi++] = xo++; Xoff[xi++] = xo++; Xoff[xi++] = xo++; Xoff[xi++] = xo++;
    Yoff[yi++] = yo ; Yoff[yi++] = yo ; Yoff[yi++] = yo ; Yoff[yi++] = yo ; Yoff[yi++] = yo ;
    xo = -2, yo++;
    Xoff[xi++] = xo++; Xoff[xi++] = xo++; Xoff[xi++] = xo++; Xoff[xi++] = xo++; Xoff[xi++] = xo++;
    Yoff[yi++] = yo ; Yoff[yi++] = yo ; Yoff[yi++] = yo ; Yoff[yi++] = yo ; Yoff[yi++] = yo ;
// Generate a 13-point lower triangular 2D Finite Difference matrix
XLoff.Size(13);
YLoff.Size(13);
xi = 0, yi = 0;
oxo = -2, yo = -2;
oxo = -2, yo++;
oxo = -2, yo++;

// Generate a 13-point upper triangular 2D Finite Difference matrix
XUoff.Size(13);
YUoff.Size(13);
xi = 0, yi = 0;
oxo = 0, yo = 0;
XUoff[xi++] = xo++; XUoff[xi++] = xo++; XUoff[xi++] = xo++; YUoff[yi++] = yo; YUoff[yi++] = yo; YUoff[yi++] = yo;
oxo = -2, yo++;
oxo = -2, yo++;
YUoff[yi++] = yo ; YUoff[yi++] = yo ; YUoff[yi++] = yo ; YUoff[yi++] = yo ; YUoff[yi++] = yo ;

Epetra_Map * map;
Epetra_Map * mapL;
Epetra_Map * mapU;
Epetra_CrsMatrix * A;
Epetra_CrsMatrix * L;
Epetra_CrsMatrix * U;
Epetra_MultiVector * b;
Epetra_MultiVector * bt;
Epetra_MultiVector * xexact;
Epetra_MultiVector * bL;
Epetra_MultiVector * btL;
Epetra_MultiVector * xexactL;
Epetra_MultiVector * bU;
Epetra_MultiVector * btU;
Epetra_MultiVector * xexactU;
Epetra_SerialDenseVector resvec(0);

//Timings
Epetra_Flops flopcounter;
Epetra_Time timer(comm);

int jstop = 1;
for (int j=0; j<jstop; j++) {
    for (int k=1; k<2; k++) {
        int nrhs=k;
        if (verbose) cout << "\n***************␣Results␣for␣" << nrhs << "␣RHS␣with␣";
        bool StaticProfile = (j!=0);
if (verbose) {
    if (StaticProfile) cout << "static profile\n";
    else cout << "dynamic profile\n";
}

GenerateCrsProblem(numNodesX, numNodesY, numProcsX, numProcsY, numPoints, Xoff.Values(), Yoff.Values(), nrhs, comm, verbose, summary, map, A, b, bt, xexact, StaticProfile, false);

runMatrixTests(A, b, bt, xexact, StaticProfile, verbose, summary);

delete A;
delete b;
delete bt;
delete xexact;

GenerateCrsProblem(numNodesX, numNodesY, numProcsX, numProcsY, Xoff.Length(), Xoff.Values(), Yoff.Values(), nrhs, comm, verbose, summary, mapL, L, bL, btL, xexactL, StaticProfile, true);

GenerateCrsProblem(numNodesX, numNodesY, numProcsX, numProcsY, Xoff.Length(), Xoff.Values(), Yoff.Values(), nrhs, comm, verbose, summary, mapU, U, bU, btU, xexactU, StaticProfile, true);
runLUMatrixTests(L, bL, btL, xexactL, U, bU, btU, xexactU,
              StaticProfile, verbose, summary);

delete L;
delete bL;
delete btL;
delete xexactL;
delete mapL;

delete U;
delete bU;
delete btU;
delete xexactU;
delete mapU;

Epetra_MultiVector q(*map, nrhs);
Epetra_MultiVector z(q);
Epetra_MultiVector r(q);

delete map;
q.SetFlopCounter(flopcounter);
z.SetFlopCounter(q);
r.SetFlopCounter(q);

resvec.Resize(nrhs);

flopcounter.ResetFlops();
timer.ResetStartTime();

// 10 norms
for( int i = 0; i < 10; ++i )
q.Norm2( resvec.Values() );

elapsed_time = timer.ElapsedTime();
total_flops = q.Flops();
MFOOPS = total_flops/elapsed_time/1000000.0;
if (verbose) cout << "\nTotal MFOOPS for 10 Norm2's =" << MFOOPS
 << endl;

if (summary) {
    if (comm.NumProc()==1) cout << "Norm2" << 't';
    cout << MFOOPS << endl;
}
flopcounter.ResetFlops();
timer.ResetStartTime();

// 10 dot's
for( int i = 0; i < 10; ++i )
    q.Dot(z, resvec.Values());

elapsed_time = timer.ElapsedTime();
total_flops = q.Flops();
MFOOPS = total_flops/elapsed_time/1000000.0;
if (verbose) cout << "Total MFOOPS for 10 Dot's =" << MFOOPS
 << endl;

if (summary) {
    if (comm.NumProc()==1) cout << "DotProd" << 't';
    cout << MFOOPS << endl;
}
flopcounter.ResetFlops();
timer.ResetStartTime();
// 10 dot's
for( int i = 0; i < 10; ++i )
q.Update(1.0, z, 1.0, r, 0.0);

elapsed_time = timer.ElapsedTime();
total_flops = q.Flops();
MFLOPs = total_flops/elapsed_time/1000000.0;
if (verbose) cout << "Total MFLOPs for 10 Updates = " << MFLOPs << endl;

if (summary) {
    if (comm.NumProc()==1) cout << "Update" << 't';
    cout << MFLOPs << endl;
}

#ifdef EPETRA_MPI
MPI_Finalize();
#endif

return ierr;

// Constructs a 2D PDE finite difference matrix using the list of x and y offsets.
// nx  (In) - number of grid points in x direction
// ny  (In) - number of grid points in y direction
// The total number of equations will be nx*ny ordered such that the x direction changes
// most rapidly:
// First equation is at point (0,0)
// Second at (1,0)
// ...
// nx equation at (nx-1,0)
// nx+1st equation at (0,1)

// numPoints (In) - number of points in finite difference stencil
// xoff  (In) - stencil offsets in x direction (of length numPoints)
// yoff  (In) - stencil offsets in y direction (of length numPoints)
// A standard 5-point finite difference stencil would be described as :

// numPoints = 5
// xoff = [-1, 1, 0, 0, 0]
// yoff = [ 0, 0, 0, -1, 1]

// nrhs - Number of rhs to generate. (First interface produces vectors,
// so nrhs is not needed

// comm  (In) - an Epetra_Comm object describing the parallel machine
// (numProcs and my proc ID)
// map   (Out) - Epetra_Map describing distribution of matrix and
// vectors/multivectors
// A    (Out) - Epetra_CrsMatrix constructed for nx by ny grid using
// prescribed stencil
// Off-diagonal values are random between 0 and 1. If
// diagonal is part of stencil,
// diagonal will be slightly diag dominant.
// b     (Out) - Generated RHS. Values satisfy b = A*xexact
// bt    (Out) - Generated RHS. Values satisfy b = A’*xexact
// xexact (Out) - Generated exact solution to Ax = b and b’ = A’*xexact

// Note: Caller of this function is responsible for deleting all output
// objects.

void GenerateCrsProblem(int numNodesX, int numNodesY, int numProcsX,
                        int numProcsY, int numPoints,
                        int * xoff, int * yoff,
const Epetra_Comm &comm, bool verbose, bool summary,
    Epetra_Map *map,
    Epetra_CrsMatrix *A,
    Epetra_Vector *b,
    Epetra_Vector *bt,
    Epetra_Vector *xexact, bool StaticProfile,
    bool MakeLocalOnly) {

    Epetra_MultiVector * b1, * bt1, * xexact1;

    GenerateCrsProblem(numNodesX, numNodesY, numProcsX, numProcsY,
                        numPoints,
                        xoff, yoff, 1, comm, verbose, summary,
                        map, A, b1, bt1, xexact1, StaticProfile,
                        MakeLocalOnly);

    b = dynamic_cast<Epetra_Vector *>(b1);
    bt = dynamic_cast<Epetra_Vector *>(bt1);
    xexact = dynamic_cast<Epetra_Vector *>(xexact1);

    return;
}

void GenerateCrsProblem(int numNodesX, int numNodesY, int numProcsX,
                          int numProcsY, int numPoints,
                          int * xoff, int * yoff, int nrhs,
                          const Epetra_Comm &comm, bool verbose, bool summary,
                          Epetra_Map *map,
                          Epetra_CrsMatrix *A,
                          Epetra_MultiVector *b,
                          Epetra_MultiVector *bt,
Epetra_MultiVector *&xexact, bool StaticProfile, bool MakeLocalOnly) {

Epetra_Time timer(comm);
// Determine my global IDs

int * myGlobalElements;
GenerateMyGlobalElements(numNodesX, numNodesY, numProcsX, numProcsY, comm.MyPID(), myGlobalElements);

int numMyEquations = numNodesX*numNodesY;

map = new Epetra_Map(-1, numMyEquations, myGlobalElements, 0, comm);
// Create map with 2D block partitioning.
delete [] myGlobalElements;

int numGlobalEquations = map->NumGlobalElements();

int profile = 0; if (StaticProfile) profile = numPoints;

if (MakeLocalOnly)
    A = new Epetra_CrsMatrix(Copy, *map, *map, profile, StaticProfile);
    // Construct matrix with rowmap=colmap
else
    A = new Epetra_CrsMatrix(Copy, *map, profile, StaticProfile); // Construct matrix

int * indices = new int[numPoints];
double * values = new double[numPoints];

double dnumPoints = (double) numPoints;
int nx = numNodesX*numProcsX;

for (int i=0; i<numMyEquations; i++) {

int rowID = map->GID(i);
int numIndices = 0;

for (int j=0; j<numPoints; j++) {
    int colID = rowID + xoff[j] + nx*yoff[j]; // Compute column ID based on stencil offsets
    if (colID>-1 && colID<numGlobalEquations) {
        indices[numIndices] = colID;
        double value = - ((double) rand())/ ((double) RAND_MAX);
        if (colID==rowID)
            values[numIndices++] = dnumPoints - value; // Make diagonal dominant
        else
            values[numIndices++] = value;
    }
    //cout << "Building row " << rowID << endl;
    A->InsertGlobalValues(rowID, numIndices, values, indices);
}

delete [] indices;
delete [] values;
double insertTime = timer.ElapsedTime();
timer.ResetStartTime();
A->FillComplete(false);
double fillCompleteTime = timer.ElapsedTime();

if (verbose)
    cout << "Time to insert matrix = " << insertTime << endl
        << "Time to complete fill = " << fillCompleteTime << endl;
if (summary) {
    if (comm.NumProc()==1) cout << "InsertTime" << 't';
    cout << insertTime << endl;
}
if (comm.NumProc()==1) cout << "FillCompleteTime" << ' \t';
cout << fillCompleteTime << endl;
}

if (nrhs<=1) {
    b = new Epetra_Vector(*map);
    bt = new Epetra_Vector(*map);
    xexact = new Epetra_Vector(*map);
}
else {
    b = new Epetra_MultiVector(*map, nrhs);
    bt = new Epetra_MultiVector(*map, nrhs);
    xexact = new Epetra_MultiVector(*map, nrhs);
}
xexact->Random(); // Fill xexact with random values
A->Multiply(false, *xexact, *b);
A->Multiply(true, *xexact, *bt);
return;

// Constructs a 2D PDE finite difference matrix using the list of x and y offsets.

// nx (In) - number of grid points in x direction
// ny (In) - number of grid points in y direction
// The total number of equations will be nx*ny ordered such that the x direction changes most rapidly:
// First equation is at point (0,0)
// Second at (1,0)
// ...
// nx equation at (nx-1,0)
// nx+1st equation at (0,1)

// numPoints (In) - number of points in finite difference stencil
// xoff (In) - stencil offsets in x direction (of length numPoints)
// yoff (In) - stencil offsets in y direction (of length numPoints)
// A standard 5-point finite difference stencil would be described as :
// numPoints = 5
// xoff = [-1, 1, 0, 0, 0]
// yoff = [0, 0, 0, -1, 1]

// nsizes (In) - Length of element size list used to create variable
// block map and matrix
// sizes (In) - integer list of element sizes of length nsizes
// The map associated with this VbrMatrix will be created by cycling
// through the sizes list.
// For example, if nsizes = 3 and sizes = [2, 4, 3], the block map
// will have elementsizes
// of 2, 4, 3, 2, 4, 3, ...

// nrhs - Number of rhs to generate. (First interface produces vectors,
// so nrhs is not needed

// comm (In) - an Epetra_Comm object describing the parallel machine
// (numProcs and my proc ID)
// map (Out) - Epetra_Map describing distribution of matrix and
// vectors/multivectors
// A (Out) - Epetra_VbrMatrix constructed for nx by ny grid using
// prescribed stencil
// Off-diagonal values are random between 0 and 1. If
diagonal is part of stencil,
// diagonal will be slightly diag dominant.
void GenerateVbrProblem(int numNodesX, int numNodesY, int numProcsX, int numProcsY, int numPoints, int * xoff, int * yoff, int nsizes, int * sizes, const Epetra_Comm &comm, bool verbose, bool summary, Epetra_BlockMap *& map, Epetra_VbrMatrix *& A, Epetra_Vector *& b, Epetra_Vector *& bt, Epetra_Vector *& xexact, bool StaticProfile, bool MakeLocalOnly) {

    Epetra_MultiVector * b1, * bt1, * xexact1;
    GenerateVbrProblem(numNodesX, numNodesY, numProcsX, numProcsY, numPoints, xoff, yoff, nsizes, sizes, 1, comm, verbose, summary, map, A, b1, bt1, xexact1, StaticProfile, MakeLocalOnly);

    b = dynamic_cast<Epetra_Vector *>(b1);
    bt = dynamic_cast<Epetra_Vector *>(bt1);
    xexact = dynamic_cast<Epetra_Vector *>(xexact1);

    return;
}
```c
void GenerateVbrProblem(int numNodesX, int numNodesY, int numProcsX,
    int numProcsY, int numPoints,
    int * xoff, int * yoff,
    int nsizes, int * sizes, int nrhs,
    const Epetra_Comm &comm, bool verbose, bool summary,
    Epetra_BlockMap *& map,
    Epetra_VbrMatrix *& A,
    Epetra_MultiVector *& b,
    Epetra_MultiVector *& bt,
    Epetra_MultiVector *& xexact, bool StaticProfile
    , bool MakeLocalOnly) {

    int i, j;

    // Determine my global IDs
    int * myGlobalElements;
    GenerateMyGlobalElements(numNodesX, numNodesY, numProcsX, numProcsY,
        comm.MyPID(), myGlobalElements);

    int numMyElements = numNodesX*numNodesY;

    Epetra_Map ptMap(~1, numMyElements, myGlobalElements, 0, comm); //
        Create map with 2D block partitioning.
    delete [] myGlobalElements;

    int numGlobalEquations = ptMap.NumGlobalElements();

    Epetra_IntVector elementSizes(ptMap); // This vector will have the
        list of element sizes
    for (i=0; i<numMyElements; i++)
        elementSizes[i] = sizes[ptMap.GID(i)%nsizes]; // cycle through
            sizes array
```
map = new Epetra_BlockMap(-1, numMyElements, ptMap.MyGlobalElements(),
    elementSizes.Values(),
    ptMap.IndexBase(), ptMap.Comm());

int profile = 0; if (StaticProfile) profile = numPoints;

if (MakeLocalOnly)
  A = new Epetra_VbrMatrix(Copy, *map, *map, profile); // Construct
  matrix rowmap=colmmap
else
  A = new Epetra_VbrMatrix(Copy, *map, profile); // Construct matrix

int * indices = new int[numPoints];

// This section of code creates a vector of random values that will
be used to create
// light-weight dense matrices to pass into the VbrMatrix
// construction process.

int maxElementSize = 0;
for (i=0; i< nsizes; i++) maxElementSize = EPETRA_MAX(maxElementSize, sizes[i]);

Epetra_LocalMap lmap(maxElementSize*maxElementSize, ptMap.IndexBase(),
    ptMap.Comm());
Epetra_Vector randvec(lmap);
randvec.Random();
randvec.Scale(-1.0); // Make value negative
int nx = numNodesX*numProcsX;

for (i=0; i<numMyElements; i++) {
  int rowID = map->GID(i);
int numIndices = 0;
int rowDim = sizes[rowID%nsizes];
for (j=0; j<numPoints; j++) {
    int colID = rowID + xoff[j] + nx*yoff[j]; // Compute column ID based on stencil offsets
    if (colID>-1 && colID<numGlobalEquations)
        indices[numIndices++] = colID;
}
A->BeginInsertGlobalValues(rowID, numIndices, indices);
for (j=0; j < numIndices; j++) {
    int colDim = sizes[indices[j]%nsizes];
    A->SubmitBlockEntry(&(randvec[0]), rowDim, rowDim, colDim);
}
A->EndSubmitEntries();
delete [] indices;
A->FillComplete();

// Compute the InvRowSums of the matrix rows
Epetra_Vector invRowSums(A->RowMap());
Epetra_Vector rowSums(A->RowMap());
A->InvRowSums(invRowSums);
rowSums.Reciprocal(invRowSums);

// Jam the row sum values into the diagonal of the Vbr matrix (to make it diag dominant)
int numBlockDiagonalEntries;
int * rowColDims;
int * diagoffsets = map->FirstPointInElementList();
A->BeginExtractBlockDiagonalView(numBlockDiagonalEntries, rowColDims);

for (i=0; i< numBlockDiagonalEntries; i++) {
    double * diagVals;
    int diagLDA;
    A->ExtractBlockDiagonalEntryView(diagVals, diagLDA);
    int rowDim = map->ElementSize(i);
    for (j=0; j<rowDim; j++) diagVals[j+j*diagLDA] = rowSums[diagoffsets[i]+j];
}

if (nrhs<=1) {
    b = new Epetra_Vector(*map);
    bt = new Epetra_Vector(*map);
    xexact = new Epetra_Vector(*map);
}
else {
    b = new Epetra_MultiVector(*map, nrhs);
    bt = new Epetra_MultiVector(*map, nrhs);
    xexact = new Epetra_MultiVector(*map, nrhs);
}

xexact->Random(); // Fill xexact with random values
A->Multiply(false, *xexact, *b);
A->Multiply(true, *xexact, *bt);
return;
}

void GenerateMyGlobalElements(int numNodesX, int numNodesY, int numProcsX, int numProcs,
                               int myPID, int * & myGlobalElements) {

myGlobalElements = new int[numNodesX*numNodesY];

int myProcX = myPID%numProcsX;
int myProcY = myPID/numProcsX;

int curGID = myProcY*(numProcsX*numNodesX)*numNodesY+myProcX*numNodesX;

for (int j=0; j<numNodesY; j++) {
    for (int i=0; i<numNodesX; i++) {
        myGlobalElements[j*numNodesX+i] = curGID+i;
    }
    curGID+=numNodesX*numProcsX;
}

void runMatrixTests(Epetra_CrsMatrix * A, Epetra_MultiVector * b,
                    Epetra_MultiVector * bt,
                    Epetra_MultiVector * xexact, bool StaticProfile,
                    bool verbose, bool summary) {

    Epetra_MultiVector z(*b);
    Epetra_MultiVector r(*b);
    Epetra_SerialDenseVector resvec(b->NumVectors());

    //Timings
    Epetra_Flops flopcounter;
    A->SetFlopCounter(flopcounter);
    Epetra_Time timer(A->Comm());
    std::string statdyn = "dynamic";
    if (StaticProfile) statdyn = "static";

    for (int j=0; j<4; j++) { // j = 0/2 is notrans, j = 1/3 is trans

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bool TransA = (j==1 || j==3);
std::string contig = "without";
if (j>1) contig = "with\(UUU\)";

#ifdef EPETRA_SHORT_PERFTEST
int kstart = 1;
#else
int kstart = 0;
#endif

for (int k=kstart; k<2; k++) {
  // Loop over old multiply vs. new multiply

  std::string oldnew = "old";
  if (k>0) oldnew = "new";

  if (j==2) A->OptimizeStorage();

  flopcounter.ResetFlops();
timer.ResetStartTime();

  if (k==0) {
    //10 matvecs
    #ifndef EPETRA_SHORT_PERFTEST
    for( int i = 0; i < 10; ++i )
      A->Multiply1(TransA, *xexact, z); // Compute z = A*xexact or z = A'*xexact using old Multiply method
    #endif
  }
  else {
    //10 matvecs
    for( int i = 0; i < 10; ++i )
      A->Multiply(TransA, *xexact, z); // Compute z = A*xexact or z = A'*xexact
double elapsed_time = timer.ElapsedTime();
double total_flops = A->Flops();

// Compute residual
if (TransA)
    r.Update(-1.0, z, 1.0, *bt, 0.0); // r = bt - z
else
    r.Update(-1.0, z, 1.0, *b, 0.0); // r = b - z
r.Norm2(resvec.Values());

if (verbose) cout << "ResNorm = " << resvec.NormInf() << ": ";
double MFLOPs = total_flops/elapsed_time/1000000.0;
if (verbose) cout << "Total MFLOPs for 10\'s with " << statdyn << " Profile (Trans = " << TransA
    << ") and " << contig << " optimized storage = " << MFLOPs << " \(s\) (" << elapsed_time << " s)"
    << endl;
if (summary) {
    if (A->Comm().NumProc()==1) {
        if (TransA) cout << "TransMv " << statdyn << " Prof " << contig
            << " OptStor " << \\
        else cout << "NoTransMv " << statdyn << " Prof " << contig
            << " OptStor " << \\
    }
    cout << MFLOPs << endl;
}
return;
void runLUMatrixTests(Epetra_CrsMatrix * L, Epetra_MultiVector * bL, 
Epetra_MultiVector * btL, Epetra_MultiVector * xexactL, 
Epetra_CrsMatrix * U, Epetra_MultiVector * bU, 
Epetra_MultiVector * btU, Epetra_MultiVector * xexactU, 
bool StaticProfile, bool verbose, bool summary) {

if (L->NoDiagonal()) {
    bL->Update(1.0, *xexactL, 1.0); // Add contribution of a unit 
    btL->Update(1.0, *xexactL, 1.0); // Add contribution of a unit 
}
if (U->NoDiagonal()) {
    bU->Update(1.0, *xexactU, 1.0); // Add contribution of a unit 
    btU->Update(1.0, *xexactU, 1.0); // Add contribution of a unit 
}

Epetra_MultiVector z(*bL);
Epetra_MultiVector r(*bL);
Epetra_SerialDenseVector resvec(bL->NumVectors());

//Timings
Epetra_Flops flopcounter;
L->SetFlopCounter(flopcounter);
U->SetFlopCounter(flopcounter);
Epetra_Time timer(L->Comm());
std::string statdyn = "dynamic";
if (StaticProfile) statdyn = "static\";
for (int j=0; j<4; j++) { // j = 0/2 is notrans, j = 1/3 is trans

    bool TransA = (j==1 || j==3);
    std::string contig = "without";
    if (j>1) contig = "withUuu";

    if (j==2) {
        L->OptimizeStorage();
        U->OptimizeStorage();
    }

    flopcounter.ResetFlops();
    timer.ResetStartTime();

    //10 lower solves
    bool Upper = false;
    bool UnitDiagonal = L->NoDiagonal(); // If no diagonal, then unit
        must be used
    Epetra_MultiVector * b = TransA ? btL : bL; // solve with the
        appropriate b vector
        for( int i = 0; i < 10; ++i )
            L->Solve(Upper, TransA, UnitDiagonal, *b, z); // Solve Lz = bL or
                    L'z = bLt

    double elapsed_time = timer.ElapsedTime();
    double total_flops = L->Flops();

    // Compute residual
    r.Update(-1.0, z, 1.0, *xexactL, 0.0); // r = bt - z
    r.Norm2(resvec.Values());

    if (resvec.NormInf()>0.000001) {
        cout << "resvec_U=U" << resvec << endl;
cout << "z_U" << z << endl;
cout << "xexactL_U" << *xexactL << endl;
cout << "r_U" << r << endl;
}

if (verbose) cout << "ResNorm_U" << resvec.NormInf() << ":\;
double MFLOPs = total_flops/elapsed_time/1000000.0;
if (verbose) cout << "Total MFLOPs for 10 Lower solves U" << statdyn << "Profile_U(TransU=U)" << TransA
  << "UandU" << contig << "OptUstorageU=U" << MFLOPs << "U(" << elapsed_time << "Us)" << endl;

if (summary) {
  if (L->Comm().NumProc()==1) {
    if (TransA) cout << "TransLSv" << statdyn<< "Prof" << contig << "OptStor" << '\t';
    else cout << "NoTransLSv" << statdyn << "Prof" << contig << "OptStor" << '\t';
  }
  cout << MFLOPs << endl;
}
flopcounter.ResetFlops();
timer.ResetStartTime();

//10 upper solves
Upper = true;
UnitDiagonal = U->NoDiagonal(); // If no diagonal, then unit must be used
b = TransA ? btU : bU; // solve with the appropriate b vector
for( int i = 0; i < 10; ++i )
  U->Solve(Upper, TransA, UnitDiagonal, *b, z); // Solve Lz = bL or L^Tz = bLt
elapsed_time = timer.ElapsedTime();
total_flops = U->Flops();

// Compute residual
r.Update(-1.0, z, 1.0, *xexactU, 0.0); // r = bt - z
r.Norm2(resvec.Values());

if (resvec.NormInf()>0.001) {
    cout << "U = " << *U << endl;
    //cout << "resvec = " << resvec << endl;
    cout << "z = " << z << endl;
    cout << "xexactU = " << *xexactU << endl;
    //cout << "r = " << r << endl;
    cout << "b = " << *b << endl;
}

if (verbose) cout << "ResNorm = " << resvec.NormInf() << ";");
MFLOPs = total_flops/elapsed_time/1000000.0;
if (verbose) cout << "Total MFLOPs for 10 Upper solves = " << MFLOPs << endl;
if (summary) {
    if (L->Comm().NumProc()==1) {
        if (TransA) cout << "TransUSv " << statdyn<< "Prof " << contig << "OptStor " << '	';
        else cout << "NoTransUSv " << statdyn << "Prof " << contig << " OptStor " << '	';
    }
    cout << MFLOPs << endl;
}
return;