Trilinos Tutorial

For Trilinos Release 10.4
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Marzio Sala
Michael A. Heroux
David M. Day
James M. Willenbring
(Editors)

Sandia National Laboratories
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Preface

The Trilinos Project is an effort to facilitate the design, development, integration and ongoing support of mathematical software libraries. The goal of the Trilinos Project is to develop algorithms and enabling technologies within an object-oriented software framework for the solution of large-scale, complex multiphysics engineering and scientific applications. There is an emphasis on developing robust, scalable algorithms in a software framework, using abstract interfaces for flexible interoperability of components while providing a full-featured set of concrete classes that implement all the abstract interfaces.

This document introduces the use of Trilinos, Release 10.4.0 and subsequent minor releases 10.4.X. The presented material includes, among others, the definition of distributed matrices and vectors with Epetra, the iterative solution of linear systems with AztecOO, incomplete factorizations with IFPACK, multilevel and domain decomposition preconditioners with ML, direct solution of linear system with Amesos, eigenvalues and eigenvectors computations with Anasazi, and iterative solution of nonlinear systems with NOX.

This tutorial is an introduction to Trilinos, intended to help computational scientists effectively apply the appropriate Trilinos package to their applications. Only a small subset of all Trilinos packages are covered in this tutorial. Basic examples are presented that are fit to be imitated.

The developers of each Trilinos package are acknowledged for providing excellent documentation and examples (and the code itself!), without which this document would have never been possible. We also acknowledge the support of the ASCI and LDRD programs that funded development of Trilinos, and the support of D-INFK department of the ETH Zürich.

Oscar Chinellato (ETHZ/ICoS) and Jens Walter (ETHZ/ICOS) are acknowledged for the \LaTeX{} macros and styles used to format this document.

Marzio Sala
Micheal Heroux
David Day
James Willenbring
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1

Introduction

Marzio Sala, Michael Heroux, David Day, James Willenbring

1.1 Getting Started

The Trilinos framework uses a two level software structure that connects a system of packages. A Trilinos package is an integral unit, usually developed to solve a specific task, by a (relatively) small group of experts. Packages exist beneath the Trilinos top level, which provides a common look-and-feel. Each package has its own structure, documentation and set of examples, and it is possibly available independently of Trilinos. However, each package is even more valuable when combined with other Trilinos packages.

Trilinos is a large software project, and currently about fifty packages are included. The entire set of packages covers a wide range of algorithms and enabling technologies for the solution of large-scale, complex multi-physics engineering and scientific problems, as well as a large set of utilities to improve the development of software for scientific computing.

Clearly, a full understanding all the functionalities of the Trilinos packages requires time. Each package offers sophisticated features, difficult to “unleash” at a first sight. Besides that, a detailed description of each Trilinos package is beyond the scope of this document. For these reasons, the goal of this tutorial is to ensure that users have the background to make good use of the extensive documentation contained in each package.

We will describe the following subset of the Trilinos packages.

- **Epetra.** The package defines the basic classes for distributed matrices and vectors, linear operators and linear problems. Epetra classes are the common language spoken by all the Trilinos packages (even if some packages can “speak” other languages). Each Trilinos package accepts as input Epetra objects. This allows powerful combinations among the various Trilinos functionalities.

- **Triutils.** This is a collection of utilities that are useful in software development. Here, we present a command line parser and a matrix generator, that are used throughout this document to define example matrices.

- **AztecOO.** This is a linear solver package based on preconditioned Krylov methods. Aztec users will find that AztecOO supports all the Aztec interfaces and functionality, and also provides significant new functionality.

- **Belos.** Provides next-generation iterative linear solvers and a powerful linear solver developer framework.
• **IFPACK.** The package performs various incomplete factorizations, and is here used with AztecOO.

• **Teuchos.** This is a collection of classes that can be essential for advanced code development.

• **ML.** The algebraic multilevel and domain decomposition preconditioner package provides scalable preconditioning capabilities for a variety of problems. It is here used as a preconditioner for AztecOO solvers.

• **Anesos.** The package provides a common interface to certain sparse direct linear solvers (generally available outside the Trilinos framework), both sequential and parallel.

• **Anasazi.** The package provides a common interface to parallel eigenvalue and eigenvector solvers, for both symmetric and non-symmetric linear problems.

• **NOX.** This is a collection of nonlinear solvers, designed to be easily integrated into an application and used with many different linear solvers.

• **Zoltan.** A toolkit of parallel services for dynamic, unstructured, and/or adaptive simulations. Zoltan provides parallel dynamic load balancing and related services for a wide variety of applications, including finite element methods, matrix operations, particle methods, and crash simulations.

• **Tpetra.** Next-generation, templated version of Petra, taking advantage of the newer advanced features of C++.

• **Didasko.** This package contains all the examples reported in this tutorial. The sources of the examples can be found in the subdirectory `<your-trilinos-home>/packages/didasko/examples`.

Table 1.1 gives a partial overview of what can be accomplished using Trilinos.

**Remark 1.** As already pointed out, Epetra objects are meant to be the “common language” spoken by all the Trilinos packages, and are a natural starting point. For new users, Chapters 2-4 are a prerequisite to the later chapters. Chapters 6 is not essential to understand Trilinos, but the functionalities there presented are used in this document as a starting point for many examples. One of the classes described in Chapter 10, the Teuchos::ParameterList, is later used in Chapters 11 and 12. Chapter 7 should be read before Chapters 9 and 11 (even if both IFPACK and ML can be compiled and run without AztecOO).

The only prerequisites assumed in this tutorial are some familiarities with numerical methods for PDEs, and with iterative linear and nonlinear solvers. Although not strictly necessary, the reader is assumed to have some familiarity with distributed memory computing and, to a lesser extent, with MPI\(^1\).

Note that this tutorial is not a substitute for individual packages’ documentation. Also, for an overview of all the Trilinos packages, the Trilinos philosophy, and a description of the packages provided by Trilinos, the reader is referred to [HBH+03]. Developers should

\(^1\)Although almost no explicit MPI instructions are required in a Trilinos code, the reader should be aware of the basic concepts of message passing, like the definition of a communicator.
also consider the Trilinos Developers’ Guide, which addresses many topics, including the development tools used by Trilinos’ developers, and a description of how to include a new package.

1.2 Installation

To obtain Trilinos, please follow the instructions at the Trilinos download page:

http://trilinos.sandia.gov/download

Trilinos has been compiled on a variety of architectures, including various flavors of Linux, Sun Solaris, SGI Irix, DEC, Mac OSX, IBM AIX, ASC Red Storm, and others. Trilinos has been designed to support parallel applications. However, it also compiles and runs on serial computers. An introduction to Trilinos and a list of FAQs may be found at the web pages:

http://trilinos.sandia.gov/getting_started.html
http://trilinos.sandia.gov/faq.html

After obtaining Trilinos, the next step is its compilation. Instructions for building Trilinos are available online:

http://trilinos.sandia.gov/build_instructions.html

1.3 Copyright and Licensing of Trilinos

Trilinos is released under the Lesser GPL GNU Licence.

Trilinos is copyrighted by Sandia Corporation. Under the terms of Contract DE-AC04-94AL85000, there is a non-exclusive license for use of this work by or on behalf of the U.S. Government. Export of this program may require a license from the United States Government.

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Some parts of Trilinos are dependent on a third party code. Each third party code comes with its own copyright and/or licensing requirements. It is responsibility of the user to understand these requirements.
1.4 Programming Language Used in this Tutorial

Trilinos is written in C++ (for most packages), and in C. Some interfaces are provided to FORTRAN codes (mainly BLAS and LAPACK routines). Even if limited support is included for C programs (and a more limited for FORTRAN code), to unleash the full power of Trilinos we recommend C++. All the example programs contained in this tutorial are in C++; some packages (like ML) contain examples in C.

1.5 Referencing Trilinos

The Trilinos project can be referenced by using the following BiBTeX citation information:

@techreport{Trilinos-Overview,  
  title = "An Overview of Trilinos",  
  author = "Michael Heroux and Roscoe Bartlett and Vicki Howle Robert Hoekstra and Jonathan Hu and Tamara Kolda and Richard Lehoucq and Kevin Long and Roger Pawlowski and Eric Phipps and Andrew Salinger and Heidi Thornquist and Ray Tuminaro and James Willenbring and Alan Williams ",  
  institution = "Sandia National Laboratories",  
  number = "SAND2003-2927",  
  year = 2003}

@techreport{Trilinos-Dev-Guide,  
  title = "Trilinos Developers Guide",  
  author = "Michael A. Heroux and James M. Willenbring and Robert Heaphy",  
  institution = "Sandia National Laboratories",  
  number = "SAND2003-1898",  
  year = 2003}

@techreport{Trilinos-Dev-Guide-II,  
  author = "Michael A. Heroux and James M. Willenbring and Robert Heaphy",  
  institution = "Sandia National Laboratories",  
  number = "SAND2003-1899",  
  year = 2003}

@techreport{Trilinos-Users-Guide,  
  title = "Trilinos Users Guide",  
  author = "Michael A. Heroux and James M. Willenbring",  
  institution = "Sandia National Laboratories",  
  number = "SAND2003-2952",  
  year = 2003}

@techreport{Trilinos-Tutorial-5.0,  
  title = "Trilinos Tutorial",  
  author = "Michael Heroux and Roscoe Bartlett and Vicki Howle Robert Hoekstra and Jonathan Hu and Tamara Kolda and Richard Lehoucq and Kevin Long and Roger Pawlowski and Eric Phipps and Andrew Salinger and Heidi Thornquist and Ray Tuminaro and James Willenbring and Alan Williams ",  
  institution = "Sandia National Laboratories",  
  number = "SAND2003-2927",  
  year = 2003}
1.6 A Note on the Directory Structure

Each Trilinos package is contained in the subdirectory

\texttt{<your-trilinos-directory>/packages}

Each package generally contains sources, examples, tests and documentation subdirectories:

\texttt{<your-trilinos-directory>/packages/<package-name>/src}
\texttt{<your-trilinos-directory>/packages/<package-name>/examples}
\texttt{<your-trilinos-directory>/packages/<package-name>/test}
\texttt{<your-trilinos-directory>/packages/<package-name>/doc}

Developers’ documentation is written using Doxygen\footnote{Copyright \copyright 1997-2003 by Dimitri van Heesch. More information can by found at the web address \url{http://www.stack.nl/~dimitri/doxygen/}.}. The Doxygen documentation, and other available documentation, for each package is available online via

\url{http://trilinos.sandia.gov/packages/<package-name>/documentation.html}

The Doxygen documentation can also be generated from the source code directly when accessing a version-controlled copy of the Trilinos source code. For some packages, Doxygen documentation can also be generated from the distribution tarball.

For example, to create the documentation for Epetra, use the following commands:

$ cd <your-trilinos-home>/packages/epetra/doc
$ doxygen

Generally, both HTML and \LaTeX documentation are created by Doxygen. The browser of choice can be used to walk through the HTML documentation. To compile the \LaTeX sources, the commands are:

$ cd <your-trilinos-home>/packages/epetra/doc/latex
$ make

1.7 List of Trilinos Developers

A list of past and present Trilinos developers is available online at:

\url{http://trilinos.sandia.gov/team.html}
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<tr>
<th>Service provided/Task performed</th>
<th>Package</th>
<th>Tutorial</th>
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<td>Chapter 3</td>
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<td>Solve a linear system with preconditioned Krylov accelerators, CG, GMRES, Bi-CGSTAB, TFQMR:</td>
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<tr>
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Table 1.1: Partial overview of intended uses of Trilinos. ⋆: not covered in this tutorial.
Working with Epetra Vectors

Marzio Sala, Michael Heroux, and David Day.

A vector is a fundamental data structure required by almost all numerical methods. Within the Trilinos framework, vectors are usually constructed starting from Epetra classes.

An Epetra vector may store either double-precision values (like the solution of a PDE problem, the right-hand side of a linear system, or nodal coordinates), or integer data values (such as a set of indexes or global IDs).

An Epetra vector may be either serial or distributed. Serial vectors are usually small, so that it is not convenient to distribute them across the processes. Possibly, serial vectors are replicated across the processes. On the other hand, distributed vectors tend to be significantly larger, and therefore their elements are distributed across the processors. In this latter case, users must specify the partition they intend to use. In Epetra, this is done by specifying a communicator (introduced in Section 2.1) and an Epetra object called map (introduced in Section 2.2). A map is basically a partitioning of a list of global IDs.

During the Chapter, the user will be introduced to:

- The fundamental Epetra communicator object, Epetra_Comm (in Section 2.1);

- The Epetra_Map object (in Section 2.2);

- The creation and assembly of Epetra vectors (in Sections 2.3 and 2.4). The sections also present common vector operations, such as dot products, fill with constant or random values, vector scalings and norms;

- A tool to redistributing vectors across processes (in Section 2.5).

2.1 Epetra Communicator Objects

The Epetra_Comm virtual class is an interface that encapsulates the general information and services needed for the other Epetra classes to run on serial or parallel computer. An Epetra_Comm object is required for building all Epetra_Map objects, which in turn are required for all other Epetra classes.

Epetra_Comm has two basic concrete implementations:

- Epetra_SerialComm (for serial executions);
Chapter 2. Working with Epetra Vectors

- Epetra_MpiComm (for MPI distributed memory executions).

For most basic applications, the user can create an Epetra_Comm object using the following code fragment:

```c++
#include "Epetra_ConfigDefs.h"
#ifdef HAVE_MPI
#include "mpi.h"
#include "Epetra_MpiComm.h"
#else
#include "Epetra_SerialComm.h"
#endif
// .. other include files and others ...
int main( int argv, char *argv[]) {
  // .. some declarations here ...
  #ifdef HAVE_MPI
    MPI_Init(&argc, &argv);
    Epetra_MpiComm Comm(MPI_COMM_WORLD);
  #else
    Epetra_SerialComm Comm;
  #endif
  // ... other code follows ...
  Note that the MPI_Init() call and the
  #ifdef HAVE_MPI
    MPI_Finalize();
  #endif
  call, are likely to be the only MPI calls users have to explicitly introduce in their code.

  Most of Epetra_Comm methods are similar to MPI functions. The class provides methods
  such as MyPID(), NumProc(), Barrier(), Broadcast(), SumAll(), GatherAll(), MaxAll(),
  MinAll(), ScanSum(). For instance, the number of processes in the communicator, NumProc,
  and the ID of the calling process, MyPID, can be obtained by

  int NumProc = Comm.NumProc();
  int MyPID = Comm.MyPID();

  The file didasko/examples/epetra/ex1.cpp presents the use of some of the above introduced functions. For a description of the syntax, please refer to the Epetra Class Documentation.

2.2 Defining a Map

The distribution of a set of integer labels (or elements) across the processes is here called a map, and its actual implementation is given by the Epetra_Map class (or, more precisely, by an Epetra_BlockMap, from which Epetra_Map is derived). Basically, the class handles the definition of the:

- global number of elements in the set (called NumGlobalElements);
2.2. Defining a Map

### Table 2.1: Some methods of the class **Epetra_Map**

- local number of elements (called `NumMyElements`);

- global numbering of all local elements (an integer vector of size `NumMyElements`, called `MyGlobalElements`).

There are three ways to define an map. The easiest way is to specify the global number of elements, and let Epetra decide:

```c
Epetra_Map Map(NumGlobalElements, 0, Comm);
```
In this case, the constructor takes the global dimension of the vector, the base index\(^1\), and an \textit{Epetra\_Comm} object (introduced in Section 2.1). As a result, each process will be assigned a contiguous set of elements.

A second way to build the \textit{Epetra\_Comm} object is to furnish the local number of elements:

\begin{verbatim}
Epetra_Map Map(-1,NumMyElements,0,Comm);
\end{verbatim}

This will create a vector of size \(\sum_{i=0}^{NumProc-1} NumMyElements\). Each process will get a contiguous set of elements. These two approaches are coded in file \texttt{didasko/examples/epetra/ex2.cpp}.

A third more involved way to create an \textit{Epetra\_Map}, is to specify on each process both the number of local elements, and the global indexing of each local element. To understand this, consider the following code. A vector of global dimension 5 is split among processes \(p_0\) and \(p_1\). Process \(p_0\) owns elements 0 an 4, and process \(p_1\) elements 1, 2, and 3.

\begin{verbatim}
#include "Epetra_Map.h"
// ...
MyPID = Comm.MyPID();
switch( MyPID ) {
    case 0:
        MyElements = 2;
        MyGlobalElements = new int[MyElements];
        MyGlobalElements[0] = 0;
        MyGlobalElements[1] = 4;
        break;
    case 1:
        MyElements = 3;
        MyGlobalElements = new int[MyElements];
        MyGlobalElements[0] = 1;
        MyGlobalElements[1] = 2;
        MyGlobalElements[2] = 3;
        break;
}
Epetra_Map Map(-1,MyElements,MyGlobalElements,0,Comm);
\end{verbatim}

The complete code is reported in \texttt{didasko/examples/epetra/ex3.cpp}.

Once created, a Map object can be queried for the global and local number of elements, using

\begin{verbatim}
int NumGlobalElements = Map.NumGlobalElements();
int NumMyElements = Map.NumMyElements();
\end{verbatim}

and for the global ID of local elements, using

\begin{verbatim}
int * MyGlobalElements = Map.MyGlobalElements();
\end{verbatim}

\(^1\)The index base is the index of the lowest order element, and is usually, 0 for C or C++ arrays, and 1 for FORTRAN arrays. Epetra can indeed accept any number as index base. However, some other Trilinos package may require a C-style index base.
2.3. Creating and Assembling Serial Vectors

that returns a pointer to the internally stored global indexing vector, or, equivalently,

```cpp
int MyGlobalElements[NumMyElements];
Map.MyGlobalElements(MyGlobalElements);
```

that copies in the user’s provided array the global indexing.

The class Epetra::Map is derived from Epetra::BlockMap. The class keeps information that describes the distribution of objects that have block elements (for example, one or more contiguous entries of a vector). This situation is common in applications like multiple-unknown PDE problems. A variety of constructors are available for the class. An example of the use of block maps is reported in `didasko/examples/epetra/ex23.cpp`.

Note that different maps may coexist in the same part of the code. The user may define vectors with different distributions (even for vectors of the same size). Two classes are provided to transfer data from one map to another: Epetra::Import and Epetra::Export (see Section 2.5).

**Remark 2.** Most Epetra objects overload the `<<` operator. For example, to visualize information about the Map, one can simply write

```cpp
cout << Map;
```

We have constructed very basic map objects. More general objects can be constructed as well. First, element numbers are only labels, and they do not have to be consecutive. This means that we can define a map with elements 1, 100 and 10000 on process 0, and elements 2, 200 and 20000 on process 1. This map, composed by 6 elements, is perfectly legal. Second, each element can be assigned to more than one process. Examples `didasko/examples/epetra/ex20.cpp` and `didasko/examples/epetra/ex21.cpp` can be used to better understand the potential of Epetra::Maps.

**Remark 3.** The use of “distributed directory” technology facilitates arbitrary global ID support.

2.3 Creating and Assembling Serial Vectors

Within Epetra, it is possible to define sequential vectors for serial and parallel platforms. A sequential vector is a vector which, in the opinion of the programmer, does not need to be partitioned among the processes. Note that each process defines its own sequential vectors, and that changing an element of this vector on this process will not directly affect the vectors stored on other processes (if any have been defined).

The class Epetra::SerialDenseVector enables the construction and use of real-valued, double precision dense vectors. The Epetra::SerialDenseVector class provides convenient vector notation but derives all significant functionality from Epetra::SerialDenseMatrix class (see Section 3.1). The following instruction creates a sequential double-precision vector containing `Length` elements:

```cpp
#include "Epetra_SerialDenseVector.h"
Epetra_SerialDenseVector DoubleVector(Length);
```
Other constructors are available, as described in the Epetra Class Documentation. Integer vectors can be created as:

```cpp
#include "Epetra_IntSerialDenseVector.h"
Epetra_SerialIntDenseVector IntVector(Length);
```

We recommend Epetra_SerialDenseVector and Epetra_SerialIntDenseVector instead of more common C++ allocations (using `new`), because Epetra serial vectors automatically delete the allocated memory when destructed, avoiding possible memory leaks.

The vector can be filled using the `[]` or `()` operators. Both methods return a reference to the specified element of the vector. However, using `()`, bound checking is enforced. Using using `[]`, no bounds checking is done unless Epetra is compiled with `EPETRA_ARRAY_BOUNDS_CHECK`.

**Remark 4.** To construct replicated Epetra objects on distributed memory machines, the user may consider the class Epetra_LocalMap. The class constructs the replicated local objects and keeps information that describe the distribution.

The file `didasko/examples/epetra/ex4.cpp` illustrates basic operations on dense vectors.

### 2.4 Creating and Assembling a Distributed Vector

A distributed object is an entity whose elements are partitioned across more than one process. Epetra’s distributed objects (derived from the Epetra_DistObject class) are created from a Map. For example, a distributed vector can be constructed starting from an Epetra_Map (or Epetra_BlockMap) with an instruction of type

```cpp
Epetra_Vector x(Map);
```

(We shall see that this dependency on Map objects holds for all distributed Epetra objects.) This constructor allocates space for the vector and sets all the elements to zero. A copy constructor may be used as well:

```cpp
Epetra_Vector y(x);
```

A variety of sophisticated constructors are indeed available. For instance, the user can pass a pointer to an array of double precision values,

```cpp
Epetra_Vector x(Copy,Map,LocalValues);
```

Note the word `Copy` is input to the constructor. It specifies the Epetra_CopyMode, and refers to many Epetra objects. In fact, Epetra allows two data access modes:

1. **Copy**: allocate memory and copy the user-provided data. In this mode, the user data is not needed be the new Epetra_Vector after construction;

2. **View**: create a “view” of the user’s data. The user data is assumed to remain untouched for the life of the vector (or modified carefully). From a data hiding perspective, View mode is very dangerous. But is is often the only way to get the required performance. Therefore, users are strongly encouraged to develop code using the Copy mode. Only use View mode as needed in a secondary performance optimization phase. To use the View mode, the user has to define the vector entries using a (double) vector (of appropriate size), than construct an Epetra_Vector with an instruction of type.
Epetra_Vector z(View,Map,z_values);

where $z_{values}$ is a pointer a double array containing the values for $z$.

To set a locally owned element of a vector, one can use the [] operator, regardless of how a vector has been created. For example,

$$x[i] = 1.0*i;$$

where $i$ is in the local index space.

Epetra also defines some functions to set vector elements in local or global index space. ReplaceMyValues or SumIntoMyValues will replace or sum values into a vector with a given indexed list of values, with indexes in the local index space; ReplaceGlobalValues or SumIntoGlobalValues will replace or sum values into a vector with a given indexed list of values in the global index space (but locally owned). It is important to note that no process may set vector entries locally owned by another process. In other words, both global and local insert and replace functions refer to the part of a vector assigned to the calling process. Intra-process communications can be (easily) performed using Import and Export objects, covered in Section 2.5.

The user might need (for example, for reasons of computational efficiency) to work on Epetra_Vectors as if they were double * pointers. File didasko/examples/epetra/ex5.cpp shows the use of ExtractCopy(). ExtractCopy does not give access to the vector elements, but only copies them into the user-provided array. The user must commit those changes to the vector object, using, for instance, ReplaceMyValues.

A further computationally efficient way, is to extract a “view” of the (multi-)vector internal data. This can be done as follows, using method ExtractView(). Let $z$ be an Epetra_Vector.

```c++
double * z_values;
z.ExtractView( &z_values );
for( int i=0 ; i<MyLength ; ++i ) z_values[i] *= 10;
```

In this way, modifying the values of $z_{values}$ will affect the internal data of the Epetra_Vector $z$. An example of the use of ExtractView is reported in file didasko/examples/epetra/ex6.cpp.

**Remark 5.** The class Epetra_Vector is derived from Epetra_MultiVector. Roughly speaking, a multi-vector is a collection of one or more vectors, all having the same length and distribution. File didasko/examples/epetra/ex7.cpp illustrates use of multi-vectors.

The user can also consider the function ResetView, which allows a (very) light-weight replacement of multi-vector values, created using the Epetra_DataMode View. Note that no checking is performed to see if the values passed in contain valid data. This method can be extremely useful in the situation where a vector is needed for use with an Epetra operator or matrix, and the user is not passing in a multi-vector. Use this method with caution as it could be extremely dangerous. A simple example is reported in didasko/examples/epetra/ex8.cpp.

It is possible to perform a certain number of operations on vector objects. Some of them are reported in Table 2.2. Example didasko/examples/epetra/ex18.cpp works with some of the functions reported in the table.
int NumMyElements()
returns the local vector length on the calling processor

int NumGlobalElements()
returns the global length

int Norm1(double *Result) const
returns the 1-norm (defined as $\sum_i |x_i|$) (see also Norm2 and NormInf)

int NormWeighted(double *Result) const
returns the 2-norm, defined as $\sqrt{\frac{1}{n} \sum_{j=1}^n (w_j x_j)^2}$

int Dot(const Epetra MultiVector A, double *Result) const
computes the dot product of each corresponding pair of vectors

int Scale(double ScalarA, const Epetra MultiVector &A
Replace multi-vector values with scaled values of A, this=$ScalarA*$A

int MinValue(double *Result) const
compute minimum value of each vector in multi-vector (see also MaxValue and MeanValue)

int PutScalar(double Scalar)
Initialize all values in a multi-vector with constant value

int Random()
set multi-vector values to random numbers

<table>
<thead>
<tr>
<th>Table 2.2: Some methods of the class Epetra_Vector</th>
</tr>
</thead>
</table>

### 2.5 Epetra_Import and Epetra_Export classes

The Epetra_Import and Epetra_Export classes apply off-processor communication. Epetra_Import and Epetra_Export are used to construct a communication plan that can be called repeatedly by computational classes such as the Epetra multi-vectors of the Epetra matrices.

Currently, those classes have one constructor, taking two Epetra_Map (or Epetra_BlockMap) objects. The first map specifies the global IDs that are owned by the calling processor. The second map specifies the global IDs of elements that we want to import later.

Using an Epetra_Import object means that the calling process knows what it wants to receive, while an Epetra_Export object means that it knows what it wants to send. An Epetra_Import object can be used to do an Export as a reverse operation (and equivalently an Epetra_Export can be used to do an Import). In the particular case of bijective maps, either Epetra_Import or Epetra_Export is appropriate.

To better illustrate the use of these two classes, we present the following example. Suppose that the double-precision distributed vector $x$ of global length 4, is distributed over two processes. Process 0 own elements 0,1,2, while process 1 owns elements 1,2,3. This means
that elements 1 and 2 are replicated over the two processes. Suppose that we want to bring all the components of \( x \) to process 0, summing up the contributions of elements 1 and 2 from the 2 processes. This is done in the following example (c.f. didasko/examples/epetra/ex9.cpp).

```cpp
int NumGlobalElements = 4; // global dimension of the problem

int NumMyElements; // local elements
Epetra_IntSerialDenseVector MyGlobalElements;

if( Comm.MyPID() == 0 ) {
    NumMyElements = 3;
    MyGlobalElements.Size(NumMyElements);
    MyGlobalElements[0] = 0;
    MyGlobalElements[1] = 1;
    MyGlobalElements[2] = 2;
} else {
    NumMyElements = 3;
    MyGlobalElements.Size(NumMyElements);
    MyGlobalElements[0] = 1;
    MyGlobalElements[1] = 2;
    MyGlobalElements[2] = 3;
}

// create a double-precision map
Epetra_Map Map(-1,MyGlobalElements.Length(),
               MyGlobalElements.Values(),0, Comm);

// create a vector based on map
Epetra_Vector x(Map);
for( int i=0 ; i<NumMyElements ; ++i )
    x[i] = 10*( Comm.MyPID()+1 );
cout << x;

// create a target map, in which all the elements are on proc 0
int NumMyElements_target;

if( Comm.MyPID() == 0 )
    NumMyElements_target = NumGlobalElements;
else
    NumMyElements_target = 0;

Epetra_Map TargetMap(-1,NumMyElements_target,0,Comm);

Epetra_Export Exporter(Map,TargetMap);

// work on vectors
Epetra_Vector y(TargetMap);
```
y.Export(x,Exporter,Add);
cout << y;

Running this code with 2 processors, the output will be approximately the following:

```
[msala:epetra]> mpirun -np 2 ./ex31.exe
Epetra::Vector
<table>
<thead>
<tr>
<th>MyPID</th>
<th>GID</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>10</td>
</tr>
<tr>
<td>0</td>
<td>1</td>
<td>10</td>
</tr>
<tr>
<td>0</td>
<td>2</td>
<td>10</td>
</tr>
</tbody>
</table>
Epetra::Vector
<table>
<thead>
<tr>
<th>MyPID</th>
<th>GID</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>20</td>
</tr>
<tr>
<td>1</td>
<td>2</td>
<td>20</td>
</tr>
<tr>
<td>1</td>
<td>3</td>
<td>20</td>
</tr>
</tbody>
</table>
Epetra::Vector
Epetra::Vector
<table>
<thead>
<tr>
<th>MyPID</th>
<th>GID</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>10</td>
</tr>
<tr>
<td>0</td>
<td>1</td>
<td>30</td>
</tr>
<tr>
<td>0</td>
<td>2</td>
<td>30</td>
</tr>
<tr>
<td>0</td>
<td>3</td>
<td>20</td>
</tr>
</tbody>
</table>
```
Working with Epetra Matrices

Marzio Sala, Michael Heroux, David Day

Epetra contains several matrix classes. Epetra matrices can be defined to be either serial or distributed. A serial matrix could be the matrix corresponding to a given element in a finite-element discretization, or the Hessemberg matrix in the GMRES method. Those matrices are of (relatively) small size, so that it is not convenient to distribute them across the processes.

Other matrices, e.g. the linear system matrices, must be distributed to obtain scalability. For distributed sparse matrices, the basic Epetra class is Epetra_RowMatrix, meant for double-precision matrices with row access. Epetra_RowMatrix is a pure virtual class. The classes that are derived from Epetra_RowMatrix include:

- Epetra_CrsMatrix for point matrices;
- Epetra_VbrMatrix for block matrices (that is, for matrices which have a block structure, for example the ones deriving from the discretization of a PDE problem with multiple unknowns for node);
- Epetra_FECrsMatrix and Epetra_FEVbrMatrix for matrices arising from FE discretizations.

The purpose of the Chapter is to review the allocation and assembling of different types of matrices as follows:

- The creation of (serial) dense matrices (in Section 3.1);
- The creation of sparse point matrices (in Section 3.2);
- The creation of sparse block matrices (in Section 3.3);
- The insertion of non-local elements using finite-element matrices (in Section 3.4).

3.1 Serial Dense Matrices

Epetra supports sequential dense matrices with the class Epetra_SerialDenseMatrix. A possible way to create a serial dense matrix $D$ of dimension $n$ by $m$ is

```cpp
#include "Epetra_SerialDenseMatrix.h"
Epetra_SerialDenseMatrix D(n,m);
```
One could also create a zero-size object,

```
Epetra_SerialDenseMatrix D();
```

and then shape this object:

```
D.Shape(n,m);
```

(D could be reshaped using `ReShape()`.)

An Epetra_SerialDenseMatrix is stored in a column-major order in the usual FORTRAN style. This class is built on the top of the BLAS library, and is derived from Epetra_Blas (not covered in this tutorial). Epetra_SerialDenseMatrix supports dense rectangular matrices.

To access the matrix element at the i-th row and the j-th column, it is possible to use the parenthesis operator \( A(i,j) \), or the bracket operator \( A[j][i] \), note that i and j are reversed\(^1\).

As an example of the use of this class, in the following code we consider a matrix-matrix product between two rectangular matrices \( A \) and \( B \).

```cpp
int NumRowsA = 2, NumColsA = 2;
int NumRowsB = 2, NumColsB = 1;
Epetra_SerialDenseMatrix A, B;
A.Shape(NumRowsA, NumColsA);
B.Shape(NumRowsB, NumColsB);
// ... here set the elements of A and B
Epetra_SerialDenseMatrix AtimesB;
AtimesB.Shape(NumRowsA, NumColsB);
double alpha = 1.0, beta = 1.0;
AtimesB.Multiply('N','N',alpha, A, B, beta);
cout << AtimesB;
```

Multiply() performs the operation \( C = \alpha A + \beta B \), where \( A \) replaced by \( A^T \) if the first input parameter is \( T \), and \( B \) replaced by \( B^T \) if the second input parameter is \( T \). The corresponding source code file is `didasko/examples/epetra/ex10.cpp`.

To solve a linear system with a dense matrix, one has to create an Epetra_SerialDenseSolver. This class uses the most robust techniques available in the LAPACK library. The class is built on the top of BLAS and LAPACK, and thus has excellent performance and numerical stability\(^2\).

Another class, Epetra_LAPACK, provides a “thin” layer on top of LAPACK, while Epetra_SerialDenseSolver attempts to provide easy access to the more robust dense linear solvers.

Epetra_LAPACK is preferable if the user seeks a convenient wrapper around the FORTRAN LAPACK routines, and the problem at hand is well-conditioned. Instead, when the user wants (or potentially wants to) solve ill-conditioned problems or favors a more object-oriented interface, then we suggest Epetra_SerialDenseMatrix.

Given an Epetra_SerialDenseMatrix and two Epetra_SerialDenseVectors \( x \) and \( b \), the general approach is as follows:

\(^1\) The bracket approach is in general faster, as the compiler can inline the corresponding function. Instead, some compiler have problems to inline the parenthesis operator.

\(^2\) Another package, Teuchos, covered in Chapter 10, allows a templated access to LAPACK. ScalAPACK is supported through Amesos, see Chapter 12.
Epetra_SerialDenseSolver Solver();
Solver.SetMatrix(D);
Solver.SetVectors(x,b);

Then, it is possible to invert the matrix with Invert(), solve the linear system with Solve(), apply iterative refinement with ApplyRefinement(). Other methods are available; for instance,

double rcond=Solve.RCOND();

returns the reciprocal of the condition number of matrix D (or -1 if not computed).

didasko/examples/epetra/ex11.cpp outlines some of the capabilities of the Epetra_SerialDenseSolver class.

### 3.2 Distributed Sparse Matrices

Epetra provides an extensive set of classes to create and fill distributed sparse matrices. These classes allow row-by-row or element-by-element constructions. Support is provided for common matrix operations, including scaling, norm, matrix-vector multiplication and matrix-multivector multiplication\(^3\).

Using Epetra objects, applications do not need to know about the particular storage format, and other implementation details such as data layout, the number and location of ghost nodes. Epetra furnishes two basic formats, one suited for point matrices, the other for block matrices. The former is presented in this Section; the latter is introduced in Section 3.3. Other matrix formats can be introduced by deriving the Epetra_RowMatrix virtual class as needed.

**Remark 6.** Some numerical algorithms require the application of the linear operator only. For this reason, some applications choose not to store a given matrix. Epetra can handle this situation using with the Epetra_Operator class; see Section 4.3.

Creating a sparse matrix may be more complicated than creating a dense matrix. It is worthwhile to take steps to avoid unnecessary dynamic memory activities due to uncertainty in the number of elements in each row.

As a general rule, the process of constructing a (distributed) sparse matrix is as follows:

- allocate an integer array $\text{Nnz}$, whose length equals the number of local rows;
- loop over the local rows, and estimate the number of nonzero elements of that row;
- create the sparse matrix using $\text{Nnz}$;
- fill the sparse matrix.

---

\(^3\)Methods for matrix-matrix products are available through the EpetraExt package. Another alternative is to use the efficient matrix-matrix product of ML, which requires ML_Operator objects. One may use lightweight conversions to ML_Operator, perform the ML matrix-matrix product, then convert the result to Epetra Matrix.
virtual int Multiply (bool TransA, const Epetra_MultiVector &X, Epetra_MultiVector &Y) const=0
Returns the result of a Epetra_RowMatrix multiplied by a Epetra_MultiVector X in Y.

virtual int Solve (bool Upper, bool Trans, bool UnitDiagonal, const Epetra_MultiVector &X, Epetra_MultiVector &Y) const=0
Returns result of a local-only solve using a triangular Epetra_RowMatrix with Epetra_MultiVectors X and Y.

virtual int InvRowSums (Epetra_Vector &x) const=0
Computes the sum of absolute values of the rows of the Epetra_RowMatrix, results returned in x.

virtual int LeftScale (const Epetra_Vector &x)=0
Scales the Epetra_RowMatrix on the left with a Epetra_Vector x.

virtual int InvColSums (Epetra_Vector &x) const=0
Computes the sum of absolute values of the cols of the Epetra_RowMatrix, results returned in x.

virtual int RightScale (const Epetra_Vector &x)=0
Scales the Epetra_RowMatrix on the right with a Epetra_Vector x.

Table 3.1: Mathematical methods of Epetra_RowMatrix

As an example, in this Section we will present how to construct a distributed (sparse) matrix, arising from a finite-difference solution of a one-dimensional Laplace problem. This matrix looks like:

\[
A = \begin{pmatrix}
2 & -1 \\
-1 & 2 & -1 \\
& & \ddots & \ddots & \ddots & -1 \\
& & & & & -1 \\
& & & & & 2
\end{pmatrix}.
\]

The example illustrates how to construct the matrix, and how to perform matrix-vector operations. The code can be found in didasko/examples/epetra/ex12.cpp.

We start by specifying the global dimension (here is 5, but can be any number):

```cpp
int NumGlobalElements = 5;
```

We create a map (for the sake of simplicity linear), and define the local number of rows and the global numbering for each local row:

```cpp
Epetra_Map Map(NumGlobalElements,0,Comm);
int NumMyElements = Map.NumMyElements();
int * MyGlobalElements = Map.MyGlobalElements();
```

In particular, we have that \(j=\text{MyGlobalElements}[i]\) is the global numbering for local node \(i\). Then, we have to specify the number of nonzeros per row. In general, this can be done in two ways:
virtual bool Filled () const=0
If FillComplete() has been called, this query returns true, otherwise it returns false.

virtual double NormInf () const=0
Returns the infinity norm of the global matrix.

virtual double NormOne () const=0
Returns the one norm of the global matrix.

virtual int NumGlobalNonzeros () const=0
Returns the number of nonzero entries in the global matrix.

virtual int NumGlobalRows () const=0
Returns the number of global matrix rows.

virtual int NumGlobalCols () const=0
Returns the number of global matrix columns.

virtual int NumGlobalDiagonals () const=0
Returns the number of global nonzero diagonal entries, based on global row/column index comparisons.

virtual int NumMyNonzeros () const=0
Returns the number of nonzero entries in the calling processor's portion of the matrix.

virtual int NumMyRows () const=0
Returns the number of matrix rows owned by the calling processor.

virtual int NumMyCols () const=0
Returns the number of matrix columns owned by the calling processor.

virtual int NumMyDiagonals () const=0
Returns the number of local nonzero diagonal entries, based on global row/column index comparisons.

virtual bool LowerTriangular () const=0
If matrix is lower triangular in local index space, this query returns true, otherwise it returns false.

virtual bool UpperTriangular () const=0
If matrix is upper triangular in local index space, this query returns true, otherwise it returns false.

virtual const Epetra::Map & RowMatrixRowMap () const=0
Returns the Epetra::Map object associated with the rows of this matrix.

virtual const Epetra::Map & RowMatrixColMap () const=0
Returns the Epetra::Map object associated with the columns of this matrix.

virtual const Epetra::Import * RowMatrixImporter () const=0
Returns the Epetra::Import object that contains the import operations for distributed operations.

Table 3.2: Attribute access methods of Epetra::RowMatrix
• Furnish an integer value, representing the number of nonzero element on each row (the same value for all the rows);

• Furnish an integer vector NumNz, of length NumMyElements(), containing the nonzero elements of each row.

The first approach is trivial: the matrix is created with the simple instruction

\[ \text{Epetra\_CrsMatrix } A(\text{Copy,Map,3}); \]

(The Copy keyword is explained in Section 2.4.) In this case, Epetra considers the number 3 as a “suggestion,” in the sense that the user can still add more than 3 elements per row (at the price of a possible performance decay). The second approach is as follows:

```c++
int * NumNz = new int[NumMyElements];
for( int i=0 ; i<NumMyElements ; i++ )
if( MyGlobalElements[i]==0 || MyGlobalElements[i] == NumGlobalElements-1)
   NumNz[i] = 2;
else
   NumNz[i] = 3;
```

We are building a tridiagonal matrix where each row has (-1 2 -1). Here NumNz[i] is the number of nonzero terms in the i-th global equation on this process (2 off-diagonal terms, except for the first and last equation).

Now, the command to create an Epetra_CsrMatrix is

\[ \text{Epetra\_CsrMatrix } A(\text{Copy,Map,NumNz}); \]

We add rows one at a time. The matrix \( A \) has been created in Copy mode, in a way that relies on the specified map. To fill its values, we need some additional variables: let us call them Indexes and Values. For each row, Indexes contains global column indices, and Values the correspondingly values.

```c++
double * Values = new double[2];
Values[0] = -1.0; Values[1] = -1.0;
int * Indices = new int[2];
double two = 2.0;
int NumEntries;
for( int i=0 ; i<NumMyElements; ++i ) {
   if (MyGlobalElements[i]==0) {
      Indices[0] = 1;
      NumEntries = 1;
   } else if (MyGlobalElements[i] == NumGlobalElements-1) {
      Indices[0] = NumGlobalElements-2;
      NumEntries = 1;
   } else {
      Indices[0] = MyGlobalElements[i]-1;
```
3.2. Distributed Sparse Matrices

```c
NumEntries = 2;
}
A.InsertGlobalValues(MyGlobalElements[i], NumEntries, Values, Indices);
// Put in the diagonal entry
A.InsertGlobalValues(MyGlobalElements[i], 1, &two, MyGlobalElements+i);
}
```

Note that column indices have been inserted using global indices (but a method called `InsertMyValues` can be used as well). Finally, we transform the matrix representation into one based on local indexes. The transformation in required in order to perform efficient parallel matrix-vector products and other matrix operations.

```c
A.FillComplete();
```

This call to `FillComplete()` will reorganize the internally stored data so that each process knows the set of internal, border and external elements for a matrix-vector product of the form $B = AX$. Also, the communication pattern is established. As we have specified just one map, Epetra considers that the the rows of $A$ are distributed among the processes in the same way of the elements of $X$ and $B$. Although standard, this approach is only a particular case. Epetra allows the user to handle the more general case of a matrix whose Map differs from that of $X$ and that of $B$. In fact, each Epetra matrix is defined by four maps:

- Two maps, called RowMap and ColumnMap, define the sets of rows and columns of the elements assigned to a given processor. In general, one processor cannot set elements assigned to other processors. RowMap and ColumnMap define the pattern of the matrix, as it is used during the construction. They can be obtained using the methods `RowMatrixRowMap()` and `RowMatrixColMap()` of the Epetra_RowMatrix class. Usually, as a ColumnMap is not specified, it is automatically created by Epetra. In general RowMap and ColumnMap can differ.

- DomainMap and RangeMap define, instead, the parallel data layout of $X$ and $B$, respectively. Note that those two maps can be completely different from RowMap and ColumnMap, meaning that a matrix can be constructed using a certain data distribution, then used on vectors with another data distribution. DomainMap and RangeMap can differ. Maps can be obtained using the methods `DomainMap()` and `RangeMap()`.

The potential of the approach are illustrated by the example file `didasko/examples/epetra/ex24.cpp`. In this example, to be run using two processors, we build two maps: `MapA` will be used to construct the matrix, while `MapB` to define the parallel layout of the vectors $X$ and $B$. For the sake of simplicity, $A$ is diagonal.

```c
Epetra_CrsMatrix A(Copy, MapA, MapA, 1);
```

As usual in this Tutorial, the integer vector `MyGlobalElementsA` contains the global ID of local nodes. To assemble $A$, we cycle over all the local rows (defined by `MapA`):

---

4Some classes, derived from the Epetra_RowMatrix, can perform data exchange; see for instance Epetra_FECCrsMatrix or Epetra_FEVbrMatrix.
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for( int i=0 ; i<NumElementsA ; ++i ) {
  double one = 2.0;
  int indices = MyGlobalElementsA[i];
  A.InsertGlobalValues(MyGlobalElementsA[i], 1, &one, &indices);
}

Now, as both $X$ and $B$ are defined using MapB, instead of calling FillComplete(), invoke

A.FillComplete(MapB,MapB);

Now, we can create $X$ and $B$ as vectors based on MapB, and perform the matrix-vector product:

Epetra_Vector X(MapB); Epetra_Vector B(MapB);
A.Multiply(false,X,B);

Remark 7. Although presented for Epetra_CrsMatrix objects, the distinction between RowMap, ColMap, DomainMap, and RangeMap holds for all classes derived from Epetra_RowMatrix.

Example didasko/examples/epetra/ex14.cpp shows the use of some of the methods of the Epetra_CrsMatrix class. The code prints out information about the structure of the matrix and its properties. The output will be approximatively as reported here:

[msala:epetra]> mpirun -np 2 ./ex14
*** general Information about the matrix
Number of Global Rows = 5
Number of Global Cols = 5
is the matrix square = yes
$||A||_{\infty}$ = 4
$||A||_1$ = 4
$||A||_F$ = 5.2915
Number of nonzero diagonal entries = 5( 100 %)
Nonzero per row : min = 2 average = 2.6 max = 3
Maximum number of nonzero elements/row = 3
min( a_{i,j} ) = -1
max( a_{i,j} ) = 2
min( abs(a_{i,j}) ) = 1
max( abs(a_{i,j}) ) = 2
Number of diagonal dominant rows = 2 (40 % of total)
Number of weakly diagonal dominant rows = 3 (60 % of total)
*** Information about the Trilinos storage
Base Index = 0
is storage optimized = no
are indices global = no
is matrix lower triangular = no
is matrix upper triangular = no
are there diagonal entries = yes
is matrix sorted = yes

Other examples for Epetra_CrsMatrix include:
• Example didasko/examples/epetra/ex13.cpp implements a simple distributed finite-element solver. The code solves a 2D Laplace problem with unstructured triangular grids. In this example, the information about the grid is hardwired. The interested user can easily modify those lines in order to read the grid information from a file.

• Example didasko/examples/epetra/ex15.cpp explains how to export an Epetra_CrsMatrix to file in a MATLAB format. The output of this example will be approximately as follows:

```matlab
[msala:epetra]> mpirun -np 2 ./ex15
A = spalloc(5,5,13);
% On proc 0: 3 rows and 8 nonzeros
A(1,1) = 2;
A(1,2) = -1;
A(2,1) = -1;
A(2,2) = 2;
A(2,3) = -1;
A(3,2) = -1;
A(3,3) = 2;
A(3,4) = -1;
% On proc 1: 2 rows and 5 nonzeros
A(4,4) = 2;
A(4,5) = -1;
A(4,3) = -1;
A(5,4) = -1;
A(5,5) = 2;
```

A companion to this example is didasko/examples/epetra/ex16.cpp, which exports an Epetra_Vector to MATLAB format. Note also that the package EpetraExt contains several purpose tools to read and write matrices in various formats.

### 3.3 Creating Block Matrices

This section reviews how to work with block matrices (where each block is a dense matrix). This format has been designed for PDE problems with more than one unknown per grid node. The resulting matrix has a sparse block structure, and the size of each dense block equals the number of PDE equations defined on that block. This format is quite general, and can handle matrices with variable block size, as is done in the following example.

First, we create a map, containing the distribution of the blocks:

```cpp
Epetra_Map Map(NumGlobalElements,0,Comm);
```

Here, a linear decomposition is used for the sake of simplicity, but any map may be used as well. Now, we obtain some information about the map:

5Trilinos offers capabilities to deal with matrices composed by few sparse blocks, like for instance matrices arising from the discretization of the incompressible Navier-Stokes equations, through the Meros package (not covered in this tutorial).
// local number of elements
int NumMyElements = Map.NumMyElements();

// global numbering of local elements
int * MyGlobalElements = new int [NumMyElements];
Map.MyGlobalElements( MyGlobalElements );

A block matrix can have blocks of different size. Here, we suppose that the dimension of
diagonal block row $i$ is $i + 1$. The integer vector ElementSizeList will contain the block size
of local element $i$.

Epetra_IntSerialDenseVector ElementSizeList(NumMyElements);
for( int i=0 ; i<NumMyElements ; ++i )
   ElementSizeList[i] = 1+MyGlobalElements[i];

Here ElementSizeList is declared as Epetra_IntSerialDenseVector, but an int array is fine
as well.

Now we can create a map for the block distribution:

Epetra_BlockMap BlockMap(NumGlobalElements,NumMyElements,
               MyGlobalElements,
               ElementSizeList.Values(),0,Comm);

and finally we can create the VBR matrix based on BlockMap. In this case, nonzero elements
are located in the diagonal and the sub-diagonal above the diagonal.

Epetra_VbrMatrix A(Copy, BlockMap, 2);

int Indices[2];
double Values[MaxBlockSize];

for( int i=0 ; i<NumMyElements ; ++i ) {
    int GlobalNode = MyGlobalElements[i];
    Indices[0] = GlobalNode;
    int NumEntries = 1;
    if( GlobalNode != NumGlobalElements-1 ) {
        Indices[1] = GlobalNode+1;
        NumEntries++;
    }
    A.BeginInsertGlobalValues(GlobalNode, NumEntries, Indices);
    // insert diagonal
    int BlockRows = ElementSizeList[i];
    for( int k=0 ; k<BlockRows * BlockRows ; ++k )
        Values[k] = 1.0*i;
    B.SubmitBlockEntry(Values,BlockRows,BlockRows,BlockRows);
    // insert off diagonal if any
    if( GlobalNode != NumGlobalElements-1 ) {
        int BlockCols = ElementSizeList[i+1];
        for( int k=0 ; k<BlockRows * BlockCols ; ++k )
            Indexes[1] = GlobalNode+1;
3.4. Insert non-local Elements Using FE Matrices

The most important additional feature provided by the Epetra_FECrsMatrix with respect to Epetra_CrsMatrix, is the capability to set non-local matrix elements. We will illustrate this using the following example, reported in didasko/examples/epetra/ex23.cpp. In the example, we will set all the entries of a distributed matrix from process 0. For the sake of simplicity, this matrix is diagonal, but more complex cases can be handled as well.

First, we define the Epetra_FECrsMatrix in Copy mode as

\[
\text{Epetra_FECrsMatrix } A(\text{Copy,Map,1});
\]

Now, we will set all the diagonal entries from process 0:

\[
\text{if( Comm.MyPID() == 0 ) } \{
\text{ for( int } i=0 \text{ ; } i<\text{NumGlobalElements} \text{ ; ++i } ) \{ \\
\text{ int } \text{indices}[2]; \\
\text{ indices[0]} = i; \text{indices[1]} = i; \\
\text{ double } \text{value} = 1.0*i; \\
\text{ A.SumIntoGlobalValues(1,indices,&value); }
\text{ } \}
\text{ } \}
\]

The Function \text{SumIntoGlobalValues} adds the coefficients specified in \text{indices} (as pair row-column) to the matrix, adding them to any coefficient that may exist at the specified location. In a finite element code, the user will probably insert more than one coefficient at time (typically, all the matrix entries corresponding to an elemental matrix).

Next, we need to exchange data, to that each matrix element not owned by process 0 could be send to the owner, as specified by \text{Map}. This is accomplished by calling, on all processes,

\[
\text{A.Globalization();}
\]

A simple

\[
\text{cout } \ll \text{ A;}
\]

can be used to verify the data exchange.
Other Epetra Classes

Marzio Sala, Michael Heroux, David Day

Epetra includes classes that facilitate the development of parallel codes. In this Chapter we will recall the main usage of some of those classes:

- Epetra_Time (in Section 4.1);
- Epetra_Flops (in Section 4.2);
- Epetra_Operator and Epetra_RowMatrix (in Section 4.3);
- Epetra_LinearProblem (in Section 4.4).

4.1 Epetra_Time

Retrieving elapsed and wall-clock time is problematic due to platform-dependent and language-dependent issues. To avoid those problems, Epetra furnishes the Epetra_Time class. Epetra_Time is meant to insulate the user from the specifics of timing among a variety of platforms. Using Epetra_Time, it is possible to measure the elapsed time. This is the time elapsed between two phases of a program.

An Epetra_Time object is defined as

\[ \text{Epetra\_Time } \text{time}(\text{Comm}); \]

(Comm being an Epetra_Comm object, see Section 2.1.) To compute the elapsed time required by a given piece of code, then user should put the instruction

\[ \text{time.ResetStartTime();} \]

before the code to be timed. ElapsedTime() returns the elapsed time from the creation of this object or from the last call to ResetStartTime().

4.2 Epetra_Flops

The Epetra_Flops class provides basic support and consistent interfaces for counting and reporting floating point operations performed in the Epetra computational classes. All classes based on the Epetra_CompObject can count flops by the user creating an Epetra_Flops object and calling the SetFlopCounter() method for an Epetra_CompObject.

As an example, suppose you are interested in counting the flops required by a vector-vector product (between, say, \( x \) and \( y \)). The first step is to create an instance of the class:
Epetra_Flops counter();

Then, it is necessary to "hook" the counter object to the desired computational object, in the following way:

x.SetFlopCounter(counter);
y.SetFlopCounter(counter);

Then, perform the desired computations on Epetra objects, like

x.Dot(y,&dotProduct);

Finally, extract the number of performed operations and stored it in the double variable total_flops as

total_flops = counter.Flops();

This returns the total number of serial flops, and then resets the flop counter to zero.

Epetra_Time objects can be used in conjunction with Epetra_Flops objects to estimate the number of floating point operations per second of a given code (or a part of it). One can proceed as here reported:

Epetra_Flops counter;
x.SetFlopCounter(counter);
Epetra_Time timer(Comm);
x.Dot(y,&dotProduct);

double elapsed_time = timer.ElapsedTime();
double total_flops =counter.Flops();
cout << "Total ops: " << total_flops << endl;
double MFLOPs = total_flops/elapsed_time/1000000.0;
cout << "Total MFLOPs for mat-vec = " << MFLOPs << endl;

This code is reported in didasko/examples/epetra/ex20.cpp. The output will be approximately as follows:

[mmsala:epetra] > mpirun -np 2 ./ex20
Total ops: 734
Total MFLOPs for mat-vec = 6.92688

Total ops: 734
Total MFLOPs for mat-vec = 2.48021

Total ops: 246
Total MFLOPs for vec-vec = 0.500985

q dot z = 2
Total ops: 246
Total MFLOPs for vec-vec = 0.592825

q dot z = 2
Remark 8. Operation count are serial count, and therefore keep track of local operations only.

Remark 9. Each computational class has a \texttt{Flops()} method, that may be queried for the flop count of that object.

### 4.3 Epetra\_Operator and Epetra\_RowMatrix Classes

Matrix-free methods are introduced in the Epetra framework using either of the following two classes:

- Epetra\_Operator;
- Epetra\_RowMatrix.

Each class is a pure virtual class (specifying interfaces only), that enable the use of real-valued double-precision sparse matrices. Epetra\_RowMatrix, derived from Epetra\_Operator, is meant for matrices where the matrix entries are intended for row access, and it is currently implemented by Epetra\_CrsMatrix, Epetra\_VbrMatrix, Epetra\_FECrsMatrix, and Epetra\_FEVbrMatrix.

Consider for example the 3-point centered difference discretization of a one dimensional Laplacian on a regular grid. For the sake of simplicity, we avoid the issues related to intra-process communication (hence this code can be run with one process only).

The first step is the definition of a class, here called \texttt{TriDiagonalOperator}, and derived from the Epetra\_Operator class.

```cpp
class TriDiagonalOperator : public Epetra\_Operator {
public:
   // .. definitions here, constructors and methods
private:
   Epetra\_Map Map_;  
   double diag\_minus\_one_; // value in the sub-diagonal
   double diag_;       // value in the diagonal 
   double diag\_plus\_one_; // value in the super-diagonal 
}
```

As the class Epetra\_Operator implements several virtual methods, we have to specify all those methods in our class. Among them, we are interested in the \texttt{Apply} method, which may be coded as follows:

```cpp
int Apply( const Epetra\_MultiVector & X, 
          Epetra\_MultiVector & Y ) const { 
   int Length = X.MyLength();

   // need to handle multi-vectors and not only vectors
   for( int vec=0 ; vec<X\_NumVectors() ; ++vec ) {

      // one-dimensional problems here
      if( Length == 1 ) {
```
Y[vec][0] = diag_ * X[vec][0];
break;
}

// more general case (Length >= 2)
// first row
Y[vec][0] = diag_ * X[vec][0] + diag_plus_one_ * X[vec][1];

// intermediate rows
for( int i=1 ; i<Length-1 ; ++i ) {
    Y[vec][i] = diag_ * X[vec][i] + diag_plus_one_ * X[vec][i+1]
             + diag_minus_one_ * X[vec][i-1];
}
// final row
Y[vec][Length-1] = diag_ * X[vec][Length-1]
                   + diag_minus_one_ * X[vec][Length-2];
return true;
}

Now, in the main function, we can define a TriDiagonalOperator object using the specified constructor:

    TriDiagonalOperator TriDiagOp(-1.0,2.0,-1.0,Map);

and

    DiagOp.Apply(x,y);

computes the discrete Laplacian on x and returns the product in y.

    didasko/examples/epetra/ex21.cpp reportes the complete source code.

**Remark 10.** The clear disadvantage of deriving Epetra_Operator or Epetra_RowMatrix with respect to use Epetra_CrsMatrix or Epetra_VbrMatrix, is that users must specify their communication pattern for intra-process data exchange. For this purpose, Epetra_Import classes can be used. File didasko/examples/epetra/ex22.cpp shows how to extend ex21.cpp to the multi-process case. This example makes use of the Epetra_Import class to exchange data.

Another use of Epetra_Operator and Epetra_RowMatrix is to allow support for user defined matrix format. For instance, suppose that your code generates matrices in MSR format (detailed in the Aztec documentation). You can easily create an Epetra_Operator, that applies the MSR format to Epetra_MultiVectors. For the sake of simplicity, we will limit ourselves to the serial case. In the distributed, we must also handle ghost-node updates.

As a first step, we create a class, derived from the Epetra_Operator class,

    class MSRMatrix : public Epetra_Operator
{

public:
    // constructor

4.4. Epetra LinearProblem

A linear system \( AX = B \) is defined by an Epetra_LinearProblem class. The class requires an Epetra_RowMatrix or an Epetra_OPERATOR object (often an Epetra_CrsMatrix or Epetra_VbrMatrix), and two (multi-)vectors \( X \) and \( B \). \( X \) must have been defined using a map equivalent to the DomainMap of \( A \), while \( B \) using a map equivalent to the RangeMap of \( A \) (see Section 3.2).

Linear systems may be solved either by iterative methods (typically, using AztecOO, covered in Chapter 7), or by direct solvers (typically, using Amesos, described in Chapter 12).

Once the linear problem has been defined, the user can:
void SetPDL (ProblemDifficultyLevel PDL)
Set problem difficulty level.

void SetOperator (Epetra_RowMatrix *A)

void SetOperator (Epetra_Operator *A)

void SetLHS (Epetra_MultiVector *X)
Set left-hand side X of linear problem AX = B.

void SetRHS (Epetra_MultiVector *B)
Set right-hand side B of linear problem AX = B.

int CheckInput () const
Check input parameters for existence and size consistency.

int LeftScale (const Epetra_Vector &D)
Perform left scaling of a linear problem.

int RightScale (const Epetra_Vector &D)
Perform right scaling of a linear problem.

Epetra_Operator * GetOperator () const
Get a pointer to the operator A.

Epetra_RowMatrix * GetMatrix () const
Get a pointer to the matrix A.

Epetra_MultiVector * GetLHS () const
Get a pointer to the left-hand-side X.

Epetra_MultiVector * GetRHS () const
Get a pointer to the right-hand-side B.

ProblemDifficultyLevel GetPDL () const
Get problem difficulty level.

bool IsOperatorSymmetric () const
Get operator symmetry bool.

Table 4.1: Methods of Epetra_LinearProblem

- scale the problem, using LeftScale(D) or RightScale(D), D being an Epetra_Vector of compatible size;
- change X and B, using SetRHS(&B) and SetLHS(&X);
- change A, using SetOperator(&A).

Please refer to Table 4.1 for a summary of the methods.

4.5 Concluding Remarks on Epetra
More details about the Epetra project, and a technical description of classes and methods, can be found in [Her02].
MATLAB I/O with EpetraExt

Marzio Sala

EpetraExt is a collection of utilities for the handling of Epetra objects. In this Chapter we will explain the usage of the input/output of Epetra objects to/from Matrix Market format. We will consider the following classes:

- EpetraExt::VectorToMatrixMarketFile (in Section 5.1);
- EpetraExt::RowMatrixToMatrixMarketFile (in Section 5.2);

By using these classes, some Epetra objects can be easily exported to MATLAB.

5.1 EpetraExt::VectorToMatrixMarketFile

The Matrix Market format is a simple, portable and human-readable format. The specifications to create the corresponding ASCII files can be found at

http://math.nist.gov/MatrixMarket/

where MATLAB files to perform I/O can be downloaded. Let us consider that X is an Epetra_Vector object. This object can be saved on file in both serial and parallel computations by using the command

EpetraExt::VectorToMatrixMarketFile(fileName, X, descr1, descr2);

where FileName is the output file name (for example, X.mm), and descr1 and descr2 are two description strings. By using the mmread.m MATLAB script available at the Matrix Market web site, one can type within MATLAB

>> X = mmread('X.mm');

then use X for any MATLAB operation. For multivectors, one should use the class EpetraExt::MultiVectorToMatrixMarketFile instead.
5.2  EpetraExt::RowMatrixToMatrixMarketFile

Equivalently to what explained in Section 5.1, one can export an Epetra_RowMatrix derived class to a sparse MATLAB array. Let us consider for example the following code, that creates a distributed diagonal matrix:

```c++
int NumMyElements = 5;
Epetra_Map  Map(-1, NumMyElements, 0, Comm);
Epetra_CrsMatrix A(Copy, Map, 0); // create a simple diagonal matrix
for (int i = 0 ; i < NumMyElements ; ++i)
{
    int j = Map.GID(i);
    double value = 1.0;
    EPETRA_CHK_ERR(A.InsertGlobalValues(j, 1, &value, &j));
}
A.FillComplete();
```

The matrix can be exported as follows:

```c++
EpetraExt::RowMatrixToMatrixMarketFile("A.mm", A, "test matrix",
                                    "This is a test matrix");
```

and then read in MATLAB as

```matlab
>> A = mmread('A.mm')
```

```
A =

    (1,1)    1
    (2,2)    1
    (3,3)    1
    (4,4)    1
    (5,5)    1
```

This example is contained in didasko/examples/epetraext/ex3.cpp.

5.3  Concluding Remarks

EpetraExt contains other several classes not described here. Among them, we note that one can read Epetra_Map’s, Epetra_MultiVector’s and Epetra_CrsMatrix’s from file when previously saved using EpetraExt output functions. Also, EpetraExt offers reordering and coloring. The ZOLTAN interface is later described in Chapter 15.
Generating Linear Systems with Triutils

Marzio Sala

This Chapter presents two functionalities of Triutils, that will be extensively used in the examples of the later chapters:

- the Triutils command line parser (in Section 6.1);
- the Triutils matrix generator (in Section 6.2).

Some readers may choose to skip this Chapter because their application is their example. However, it does help to find the test matrices closest to the ones in their code for several reasons. Using well-chosen matrices, simple but sufficiently close to the final application, one can quickly test the performances of a given set of algorithms on the problem of interest, using a serial or a parallel environment. Several test matrices exhibit a known behavior (e.g., theory predicts the behavior of the condition number), and can be used to validate algorithms. Besides, as they can be quickly generate with few code lines, experts may use them to optimize or fix their code. Therefore, a short code using a gallery matrix may be used to communicate with developers.

6.1 Trilinos_Util::CommandLineParser

It is possible to use the Trilinos_Util::CommandLineParser class to parse the command line. With this class, it is easy to handle input line arguments and shell-defined variables. For instance, the user can write

[msala:triutils]$ex2.exe -nx 10 -tol 1e-6 -solver=cg -output

and, in the code, easily obtain the value of \(nx\), \(tol\), and \(solver\), using a simple code as follows:

```c
int main(int argc, char *argv[])
{

    Trilinos_Util::CommandLineParser CLP(argc,argv);
    int nx = CLP.Get("-nx", 123);
```
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```cpp
int ny = CLP.Get("-ny", 145);
double tol = CLP.Get("-tol", 1e-12);
string solver = CLP.Get("-solver","gmres");

bool Output = CLP.Has("-output");

cout << "nx = " << nx << endl;
cout << "ny = " << ny << " (default value)" << endl;
cout << "tol = " << tol << endl;
cout << "solver = " << solver << endl;

return 0;
}
```

In the command line, the user can specify a value for a given option in the following ways:

- `-tolerance 1e-12` (with one or more spaces)
- `-tolerance=1e-12` (with = sign and no spaces)

Option names must begin with one or more dashes (`-`). Options may have at most one value.

If option name is not found in the database, the default value is returned. If needed, the user can also specify a default value to return when the option name is not found in the database. The method `HaveOption` will query the database for an option.

File `didasko/examples/triutils/ex2.cpp` gives an example of the usage of this class.

### 6.2 Trilinos_Util::CrsMatrixGallery

The class `Trilinos_Util::CrsMatrixGallery` provides functions similar to the MATLAB’s `gallery` function\(^1\).

A typical constructor requires the problem type and an `Epetra_Comm`, and is followed by a set of instructions to specify the problem. The following example creates a matrix corresponding to the discretization of a 2D Laplacian on a Cartesian grid with 100 points:

```cpp
Trilinos_Util::CrsMatrixGallery Gallery("laplace_2d", Comm);
Gallery.Set("problem_size",100);
Gallery.Set("map_type","linear");
Gallery.Set("exact_solution","random");
```

The nodes are decomposed linearly, and the exact solution is a random vector.

The next example will read a matrix stored in Harwell/Boeing format:

```cpp
Trilinos_Util::CrsMatrixGallery Gallery("hb", Comm);
Gallery.Set("matrix_name","bcsstk14.rsa");
Gallery.Set("map_type","greedy");
```

\(^1\)Many of the matrices that can be created using `Trilinos_Util::CrsMatrixGallery` are equivalent or similar to those provided by the MATLAB® function `gallery`. In these cases, the reader is referred to the MATLAB documentation for more details about the matrices’ properties.
The example reads the matrix (and, if available, solution and right-hand side) from the file `bcsstk14.rsa`, and partitions the matrix across the processes using a simple greedy algorithm.

Once all the required parameters have been specified, the user can get a pointer to the constructed Epetra_CrsMatrix, to the exact and starting solution, and to the right-hand side (both Epetra_Vector’s):

```cpp
A = Gallery.GetMatrix();
ExactSolution = Gallery.GetExactSolution();
RHS = Gallery.GetRHS();
StartingSolution = Gallery.GetStartingSolution();
```

An Epetra_LinearProblem is defined by

```cpp
Epetra_LinearProblem Problem(A,StartingSolution,RHS);
```

Next one may use AztecOO to solve the linear system:

```cpp
AztecOO Solver(Problem);
Solver.SetAztecOption( AZ_precond, AZ_dom_decomp );
Solver.Iterate(1000,1E-9);
```

Using Trilinos_Util::MatrixGallery, one computes the true residual and the difference between computed and exact solution:

```cpp
double residual;
Gallery.ComputeResidual(&residual);
Gallery.ComputeDiffBetweenStartingAndExactSolutions(&residual);
```

A list of methods implemented by Trilinos_Util::CrsMatrixGallery is reported in Table 6.1.

The matrix can be written on a file in MATLAB format, using

```cpp
string FileName = "matrix.m";
bool UseSparse = false;
Gallery.WriteMatrix(FileName,UseSparse);
```

If `UseSparse` is true, the matrix is created in sparse format (using the MATLAB command `spalloc`).

To sum up, the main options reviewed here are:

- **problem_type** [string] Specifies the problem type. A list of currently available problems is reported later in this section.

- **problem_size** [int] Size of the problem. Note that some problems, defined on structured meshes, allow the specification of the number of nodes on the x-, y- and z-axis. Please refer to each problem’s description for more details.

- **nx** [int] Number of nodes in the x-direction (if supported by the specific problem).
GetMatrix()  
Returns a pointer to the internally stored Epetra_CrsMatrix.

GetExactSolution()  
Returns a pointer to the internally stored exact solution vector (as an Epetra_Vector).

GetStartingSolution()  
Returns a pointer to the internally stored starting solution vector (as an Epetra_Vector).

GetRhs()  
Returns a pointer to the internally stored right-hand side (as an Epetra_Vector).

GetLinearProblem()  
Returns a pointer to the internally stored Epetra_LinearProblem for the VBR matrix).

GetMap()  
Returns a pointer to the internally stored Epetra_Map.

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ny</td>
<td>[int] Number of nodes in the y-direction (if supported by the specific problem).</td>
</tr>
<tr>
<td>nz</td>
<td>[int] Number of nodes in the z-direction (if supported by the specific problem).</td>
</tr>
<tr>
<td>mx</td>
<td>[int] Number of processes in the x-direction (if supported by the specific problem).</td>
</tr>
<tr>
<td>my</td>
<td>[int] Number of processes in the y-direction (if supported by the specific problem).</td>
</tr>
<tr>
<td>mz</td>
<td>[int] Number of processes in the z-direction (if supported by the specific problem).</td>
</tr>
<tr>
<td>map_type</td>
<td>[string] Defines the data layout across the processes. See Table 6.2.</td>
</tr>
<tr>
<td>exact_solution</td>
<td>[string] Defines the exact solution. See Table 6.3.</td>
</tr>
<tr>
<td>starting_solution</td>
<td>[string] Defines the starting solution vector. It can be: zero or random.</td>
</tr>
</tbody>
</table>

Table 6.1: Methods of class Trilinos_Util::CrsMatrixGallery.
6.2. Trilinos_Util::CrsMatrixGallery

<table>
<thead>
<tr>
<th>map_type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>linear</td>
<td>Create a linear map. Elements are divided into continuous chunks among the processors. This is the default value.</td>
</tr>
<tr>
<td>box</td>
<td>Used for problems defined on Cartesian grids over a square. The domain is subdivided into ( mx \times my ) subdomains. ( mx ) and ( my ) are automatically computed if the total number of processes is a perfect square. Alternatively, ( mx ) and ( my ) are specified via <code>Set(&quot;mx&quot;,IntValue)</code> and <code>Set(&quot;my&quot;,IntValue)</code>.</td>
</tr>
<tr>
<td>interlaced</td>
<td>Elements are subdivided so that element ( i ) is assigned to process ( i % \text{NumProcs} ).</td>
</tr>
<tr>
<td>random</td>
<td>Assign each node to a random process</td>
</tr>
<tr>
<td>greedy</td>
<td>(only for HB matrices) implements a greedy algorithm to decompose the graph of the HB matrix among the processes</td>
</tr>
</tbody>
</table>

**Table 6.2:** Available `map_type` options.

<table>
<thead>
<tr>
<th>exact_solution</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>random</td>
<td>Random values</td>
</tr>
<tr>
<td>constant</td>
<td>All elements set to 1.0.</td>
</tr>
<tr>
<td>quad_x</td>
<td>Nodes are supposed to be distributed over the 1D segment ((0,1)), with equal spacing, and the solution is computed as (x(1-x)).</td>
</tr>
<tr>
<td>quad_xy</td>
<td>Nodes are supposed to be distributed over the square ((0,1)\times(0,1)), with equal spacing, and the solution is computed as (x(1-x)y(1-y)).</td>
</tr>
</tbody>
</table>

**Table 6.3:** Available `exact_solution` options.

A list of currently available problems is reported below. We use the following notation. `IntValue` always refer to a generic positive integer. The following symbols \( a,b,c,d,e,f,g \) always refer to double-precision values. Note that some matrices are dense, but still stored as Epetra_CrsMatrix, a sparse matrix format. The generic \((i,j)\) element of a given matrix is \(A_{i,j}\) (for simplicity, we suppose that indices start from 1). \( n \) represents the matrix size.

<table>
<thead>
<tr>
<th>problem</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>eye</td>
<td>Creates an identity matrix. The size of the problem is set using <code>Set(&quot;problem size&quot;, IntValue)</code>, or, alternatively, by <code>Set(&quot;nx&quot;, IntValue)</code>.</td>
</tr>
<tr>
<td>cauchy</td>
<td>Creates a particular instance of a Cauchy matrix with elements (A_{i,j} = 1/(i+j)). Explicit formulas are known for the inverse and determinant of a Cauchy matrix. For this particular Cauchy matrix, the determinant is nonzero and the matrix is totally positive.</td>
</tr>
</tbody>
</table>

\(^2\)It is understood that, in the actual implementation, indices start from 0.
cross_stencil_2d  Creates a matrix with the same stencil of laplace_2d, but with arbitrary values. The stencil is

\[
A = \begin{bmatrix}
e & b & a & c \\
b & a & c & d \\
a & c & d & e \\
d & e & a & d \\
\end{bmatrix}.
\]

The default values are \(a=5, \ b=c=d=e=1\). The problem size is specified as in laplace_2d.

cross_stencil_3d  Similar to the 2D case. The matrix stencil correspond to that of a 3D Laplace operator on a structured grid. On a given x-y plane, the stencil is as in laplace_2d. The value on the plane below is set using Set("f",F), and in the plane above with Set("g",G). The default values are \(a=7, b=c=d=e=f=g=1\). The problem size is specified as in laplace3d.

diag  Creates a diagonal matrix. The elements on the diagonal can be set using Set("a",value). Default value is \(a = 1\). The problem size is set as for eye.

fiedler  Creates a matrix whose element are \(A_{i,j} = |i - j|\). The matrix is symmetric, and has a dominant positive eigenvalue, and all the other eigenvalues are negative.

hanowa  Creates a matrix whose eigenvalues lie on a vertical line in the complex plane. The matrix has the 2x2 block structure (in MATLAB’s notation)

\[
A = \begin{bmatrix}
a * eye(n/2) & -diag(1:m) \\
diag(1:m) & a * eye(n/2) \\
\end{bmatrix}.
\]

The complex eigenvalues are of the form \(a k \sqrt{-1}\) and \(-k \sqrt{-1}\), for \(1 \leq k \leq n/2\). The default value for \(a\) is -1.

hb  The matrix is read from file. File name is specified by Set("file name", FileName). FileName is a C++ string. The problem size is automatically determined.

hilbert  This is a famous example of a badly conditioned matrix. The elements are defined as \(A_{i,j} = 1/(i+j)\).

jordblock  Creates a Jordan block with eigenvalue set via Set("a",DoubleVal). The default value is 0.1. The problem size is specified as for eye.

kms  Create the \(n \times n\) Kac-Murdock-Szegő Toeplitz matrix such that \(A_{i,j} = \rho^{|i-j|}\) (for real \(\rho\) only). Default value is \(\rho = 0.5\), or can be set as Set("a",value). The inverse of this matrix is tridiagonal, and the matrix is positive definite if and only if \(0 < |\rho| < 1\).
6.2. Trilinos_Util::CrsMatrixGallery

- **laplace_1d**: Creates the classical tridiagonal matrix with stencil [-1, 2, -1]. The problem size is specified as for eye.

- **laplace_1d_n**: As for laplace_1d, but with Neumann boundary condition. The matrix is singular.

- **laplace_2d**: Creates a matrix corresponding to the stencil of a 2D Laplacian operator on a structured Cartesian grid. The problem size is specified using `Set("problem size", IntValue)`. In this case, IntValue must be a perfect square. Alternatively, one can set the number of nodes along the x-axis and y-axis, using `Set("nx",IntValue)` and `Set("ny",IntValue)`.

- **laplace_2d_n**: As for laplace_2d, but with Neumann boundary condition. The matrix is singular.

- **laplace_3d**: Creates a matrix corresponding to the stencil of a 3D Laplacian operator on a structured Cartesian grid. The problem size is specified using `Set("problem size",IntValue)`. In this case, IntValue must be a cube. Alternatively, one can specify the number of nodes along the axis, using `Set("nx",IntValue)`, `Set("ny",IntValue)`, and `Set("nz",IntValue)`.

- **lehmer**: Returns a symmetric positive definite matrix, such that
  \[
  A_{i,j} = \begin{cases} 
  \frac{i}{j} & \text{if } j \geq i \\
  \frac{j}{i} & \text{otherwise} 
  \end{cases} 
  \]
  This matrix has three properties: is totally nonnegative, the inverse is tridiagonal and explicitly known, The condition number is bounded as \( n \leq \text{cond}(A) \leq 4 \times n \). The problem size is set as for eye.

- **minij**: Returns the symmetric positive definite matrix defined as \( A_{i,j} = \min(i,j) \). The problem size is set as for eye.

- **ones**: Creates a matrix with equal elements. The default value is 1, and can be changed using `Set("a",a)`.

- **parter**: Creates a matrix \( A_{i,j} = 1/(i - j + 0.5) \). This matrix is a Cauchy and a Toepliz matrix. Most of the singular values of A are very close to \( \pi \). The problem size is set as for eye.
pei  Creates the matrix
\[
A_{i,j} = \begin{cases} 
\alpha + 1 & \text{if } i \neq j \\
1 & \text{if } i = j.
\end{cases}
\]
The value of \( \alpha \) can be set as \text{Set("a",value)}, and it defaults to 1. This matrix is singular for \( \alpha = 0 \) or \(-n\).

recirc\_2d  Returns a matrix corresponding to the finite-difference discretization of the problem
\[-\mu \Delta u + (v_x, v_y) \cdot \nabla u = f\]
on the unit square, with homogeneous Dirichlet boundary conditions. A standard 5-pt stencil is used to discretize the diffusive term, and a simple upwind stencil is used for the convective term. Here,
\[
v_x = (y - 1/2)V, \quad v_y = (1/2 - x)V
\]
The value of \( \mu \) can be specified using \text{Set("diff", DoubleValue)}, and that of \( V \) using \text{Set("conv", DoubleValues)}. The default values are \( \mu = 10^{-5}, V = 1 \). The problem size is specified as in \text{laplace\_3d}.

ris  Returns a symmetric Hankel matrix with elements \( A_{i,j} = 0.5/(n - i - j + 1.5) \), where \( n \) is problem size. The eigenvalues of \( A \) cluster around \(-\pi/2\) and \(\pi/2\).

tridiag  Creates a tridiagonal matrix. The diagonal element is set using \text{Set("a", a)}, the subdiagonal using \text{Set("b", b)}, and the super-diagonal using \text{Set("c", c)}. The default values are \( a=2, b=c=1 \). The problem size is specified as for \text{eye}.

uni\_flow\_2d  Returns a matrix corresponding to the finite-difference discretization of the problem
\[-\mu \Delta u + (v_x, v_y) \cdot \nabla u = f\]
on the unit square, with homogeneous Dirichlet boundary conditions. A standard 5-pt stencil is used to discretize the diffusive term, and a simple upwind stencil is used for the convective term. Here,
\[
v_x = \cos(\alpha)V, \quad v_y = \sin(\alpha)V
\]
that corresponds to an unidirectional 2D flow. The value of \( \mu \) can be specified using \text{Set("diff", DoubleValue)}, and that of \( V \) using \text{Set("conv", DoubleValue)}, and that of \( \alpha \) using \text{Set("alpha", DoubleValue)}. The default values are \( V = 1, \mu = 10^{-5}, \alpha = 0 \). The problem size is specified as in \text{laplace3d}.

Class Trilinos\_Util::VrbMatrixGallery, derived from Trilinos\_Util::CrsMatrixGallery, can
be used to generate VBR matrices. The class creates an Epetra_CrsMatrix (following user’s defined parameters, as previously specified), then “expands” this matrix into a VBR matrix. This VBR matrix is based on an Epetra_BlockMap, based on the Epetra_Map used to define the Epetra_CrsMatrix. The number of PDE equations per node is set with parameter num_pde_eqns. The Epetra_CrsMatrix is expanded into a VBR matrix by replicating the matrix num_pde_eqns times for each equation.

A list of methods implemented by Trilinos_Util::VbrMatrixGallery is reported in Table 6.4.

<table>
<thead>
<tr>
<th>Method Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>GetVrbMatrix()</td>
<td>Returns a pointer to the internally stored VBR matrix.</td>
</tr>
<tr>
<td>GetVrbExactSolution()</td>
<td>Returns a pointer to the internally stored exact solution vector (as an Epetra_Vector).</td>
</tr>
<tr>
<td>GetVrbStartingSolution()</td>
<td>Returns a pointer to the internally stored starting solution vector (as an Epetra_Vector).</td>
</tr>
<tr>
<td>GetVrbRhs()</td>
<td>Returns a pointer to the internally stored right-hand side (as an Epetra_Vector).</td>
</tr>
<tr>
<td>GetVrbLinearProblem()</td>
<td>Returns a pointer to the internally stored Epetra_LinearProblem.</td>
</tr>
<tr>
<td>GetBlockMap()</td>
<td>Returns a pointer to the internally stored Epetra_BlockMap.</td>
</tr>
</tbody>
</table>

**Table 6.4:** Methods of class Trilinos_Util::VbrMatrixGallery.

Trilinos_Util::CrsMatrixGallery can be used in conjuction with Trilinos_Util::CommandLineParser as in the following code:

```c++
int main(int argc, char *argv[])
{
  #ifdef HAVE_MPI
    MPI_Init(&argc,&argv);
    Epetra_MpiComm Comm(MPI_COMM_WORLD);
  #else
    Epetra_SerialComm Comm;
  #endif

  Epetra_Time Time(Comm);

  Trilinos_Util::CommandLineParser CLP(argc,argv);
  Trilinos_Util::CrsMatrixGallery Gallery("", Comm);
  Gallery.Set(CLP);

  // get matrix
  Epetra_CrsMatrix * Matrix = Gallery.GetMatrix();
  Epetra_Vector * LHS = Gallery.GetLHS();
  Epetra_Vector * StartingSolution = Gallery.GetStartingSolution();
}
```
```cpp
Epetra_Vector * ExactSolution = Gallery.GetExactSolution();
Epetra_LinearProblem * Problem = Gallery.GetLinearProblem();

// various computatons...

// check that computed solution (in StartingSolution)
// is close enough to ExactSolution

double residual, diff;

Gallery.ComputeResidual(&residual);
Gallery.ComputeDiffBetweenStartingAndExactSolutions(&diff);

if( Comm.MyPID()==0 )
  cout << "\|b-Ax\|_2 = " << residual << endl;

if( Comm.MyPID()==0 )
  cout << "\|x_{exact} - x\|_2 = " << diff << endl;

#ifdef HAVE_MPI
  MPI_Finalize();
#endif

return 0 ;
```

This program can be executed with the following command line:

```
[msala:triutils]> mpirun -np 4 ex1.exe -problem_type=laplace_2d \
  -problem_size=10000
```

Matrix gallery option names shall be specified with an additional leading dash ("-"). Options values will be specified as usual.

**Remark 11.** Most of the examples reported in the following chapters use both `Trilinos_Util::CommandLineParser` and `Trilinos_Util::CrsMatrixGallery` to define the distributed matrix. The user is encouraged to test a given method using matrices with different numerical properties.
Iterative Solution of Linear Systems with AztecOO

Marzio Sala, Michael Heroux, David Day

The AztecOO [Her04] package extends the Aztec library [THHS99]. Aztec is the legacy iterative solver at the Sandia National Laboratories. It has been extracted from the MPSalsa reacting flow code [SMH+95, SDH+96], and it is currently installed in dozens of Sandia’s applications. AztecOO extends this package, using C++ classes to enable more sophisticated uses. AztecOO is intended for the iterative solution of linear systems of the form

\[ A X = B, \]  

(7.1)

when \( A \in \mathbb{R}^{n \times n} \) is the linear system matrix, \( X \) the solution, and \( B \) the right-hand side. Although AztecOO can live independently of Epetra, in this tutorial it is supposed that \( A \) is an Epetra_RowMatrix, and both \( X \) and \( B \) are Epetra_Vector or Epetra_MultiVector objects.

The Chapter reviews the following topics:

- Outline the basic issue of the iterative solution of linear systems (in § 7.1);
- Present the basic usage of AztecOO (in § 7.2);
- Define one-level domain decomposition preconditioners (in § 7.3);
- Use of AztecOO problems as preconditioners to other AztecOO problems (in § 7.4).

7.1 Theoretical Background

Our aim is to briefly present the vocabulary and notation required to define the main features available to users. The Section is neither exhaustive, nor complete. Readers are referred to the existing literature for a comprehensive presentation (see, for instance, [BBC+94, Axe94, Saa96]).

One can distinguish between two different aspects of the iterative solution of a linear system. The first one in the particular acceleration technique for a sequence of iterations vectors, that is a technique used to construct a new approximation for the solution, with information
Chapter 7. Iterative Solution of Linear Systems with AztecOO

from previous approximations. This leads to specific iteration methods, generally of Krylov type, such as conjugate gradient or GMRES. The second aspect is the transformation of the given system to one that is solved more efficiently by a particular iteration method. This is called \textit{preconditioning}. A good preconditioner improves the convergence of the iterative method, sufficiently to overcome the extra cost of its construction and application. Indeed, without a preconditioner the iterative method may even fail to converge in practice.

The convergence of iterative methods depends on the spectral properties of the linear system matrix. The basic idea is to replace the original system (7.1) by the left preconditioned system,

\[
P^{-1} A X = P^{-1} B
\]

or the right preconditioned system

\[
A P^{-1} P X = B
\]

using the linear transformation \(P^{-1}\), called preconditioner, in order to improve the spectral properties of the linear system matrix. In general terms, a preconditioner is any kind of transformation applied to the original system which makes it easier to solve.

From a modern perspective, the general problem of finding an efficient preconditioner is to identify a linear operator \(P\) with the following properties:

1. \(P^{-1} A\) is near to the identity matrix

2. The cost of applying the preconditioner is relatively low.

3. Preconditioning is scalable.

The role of the \(P\) in the iterative method is simple. At each iteration, it is necessary to solve an auxiliary linear system \(P z_m = r_m\) for \(z_m\) given \(r_m\). It is unnecessary to explicitly invert \(P\).

It should be stressed that computing the inverse of \(P\) is not mandatory; actually, the role of \(P\) is to “preconditioning” the residual at step \(m\), \(r_m\), through the solution of the additional system \(P z_m = r_m\). This system \(P z_m = r_m\) should be much easier to solve than the original system.

The choice of \(P\) varies from “black-box” algebraic techniques for general matrices to “problem dependent” preconditioners that exploit special features of a particular class of problems. Although problem dependent preconditioners may be very powerful, there is still a practical need for efficient preconditioning techniques for large classes of problems. Even the “black-box” preconditioners require parameters, and suitable parameter settings do vary. Between these two extrema, there is a class of preconditioners which are “general-purpose” for a particular – although large – class of problems. These preconditioners are sometimes called “gray-box” preconditioners; users supply a little information about the matrix and the problem to be solved.

The AztecOO preconditioners include Jacobi, Gauss-Seidel, polynomial and domain decomposition based methods [SBG96]. Furthermore preconditioners can be given to an AztecOO Krylov accelerator, by using the Trilinos packages IFPACK and ML, covered in §9 and 11, respectively.
7.2 Basic Usage

First we delineate the steps required to apply AztecOO to a linear system. To solve a linear system with AztecOO, one must create an \texttt{Epetra\_Linear\_Problem} object (see § 4.4) with the command

\begin{verbatim}
Epetra_LinearProblem Problem(&A,&x,&b);
\end{verbatim}

\(A\) is an Epetra matrix, and both \(x\) and \(b\) are Epetra vectors\(^1\). Second create an AztecOO object,

\begin{verbatim}
AztecOO Solver(Problem);
\end{verbatim}

Next specify how to solve the linear system. All AztecOO options are set using two vectors, one of integers and the other of doubles, as detailed in the Aztec’s User Guide [THHS99]. To choose among the different AztecOO parameters, the user can create two vectors, usually called \texttt{options} and \texttt{params}, set them to the default values, and then override with the desired parameters. Here’s how to set default values:

\begin{verbatim}
int options[AZ_OPTIONS_SIZE];
double params[AZ_PARAMS_SIZE];
AZ_defaults(options, params);
Solver.SetAllAztecOptions( options );
Solver.SetAllAztecParams( params );
\end{verbatim}

The latter two functions copy the values of \texttt{options} and \texttt{params} to variables internal to the AztecOO object.

Alternatively, it is possible to set specific parameters without creating \texttt{options} and \texttt{params}, using the AztecOO methods \texttt{SetAztecOption()} and \texttt{SetAztecParams()}. For instance,

\begin{verbatim}
Solver.SetAztecOption( AZ_precond, AZ_Jacobi );
\end{verbatim}

to specify a point Jacobi preconditioner. (Please see to the Aztec documentation [Her04] for more details about the various Aztec settings.)

Finally solve the linear system, say with a maximum of 1550 iterations and the residual error norm threshold \(10^{-9}\):

\begin{verbatim}
Solver.Iterate(1550,1E-9);
\end{verbatim}

The complete code is in \texttt{didasko/examples/aztec/ex1.cpp}.

Note that the matrix must be in local form (that is, the command \texttt{A.FillComplete()} must have been invoked before solving the linear system). The same AztecOO linear system solution procedure applies in serial and in parallel. However for some preconditioners, the convergence rate (and the number of iterations) depends on the number of processor.

When \texttt{Iterate()} returns, one can query for the number of iterations performed by the linear solver using \texttt{Solver.NumIters()}, while \texttt{Solver.TrueResidual()} gives the (unscaled) norm of the residual.

\(^1\)At the current stage of development, AztecOO does not take advantage of the Epetra\_Multi\_Vectors. It accepts Multi\_Vectors, but it will solve the linear system corresponding to the first multivector only. The Belos package implements block Krylov subspace methods that are significantly more efficient for linear systems with multiple simultaneous right-hand sides.
7.3 Overlapping Domain Decomposition Preconditioners

A one level overlapping domain decomposition preconditioner $P$ takes the form

$$P^{-1} = \sum_{i=1}^{M} R_i^T \tilde{A}_i^{-1} R_i,$$  \hspace{1cm} (7.2)

The number of subdomains is $M$, $R_i$ is a rectangular Boolean matrix that restricts a global vector to the subspace defined by the interior of the $i$th subdomain, and $\tilde{A}_i$ approximates $A_i = R_i A R_i^T$.

(\tilde{A}_i may equal $A_i$). Typically, $\tilde{A}_i$ differs from $A_i$ when incomplete factorizations are used in (7.2) to apply $\tilde{A}_i^{-1}$, or when a matrix different from $A$ is used in (7.3).

(7.3), The specification starts with

```
Solver.SetAztecOption( AZ_precond, AZ_dom_decomp );
```

Next if an incomplete factorization of $A_i$ will be used, then specify its parameters:

```
Solver.SetAztecOption( AZ_subdomain_solve, AZ_ilu );
Solver.SetAztecOption( AZ_graph_fill, 1 );
```

On the other hand, exact subdomain solves \(^2\) are specified like this:

```
Solver.SetAztecOption( AZ_subdomain_solve, AZ_lu );
```

The default amount of overlap is 0; this is equivalent to “block” Jacobi preconditioning with block size equal to the size of the subdomain. If amount of overlap is one ($k$),

```
Solver.SetAztecOption( AZ_overlap, 1 );
```

then the “block” is augmented by all distance one ($k$), neighbors in the sparse matrix graph.

**Remark 12.** Two-level domain decomposition schemes [SBG96] are available through AztecOO in conjunction with ML. Please see § 11.5.

**Remark 13.** The IFPACK Trilinos package (see § 9) computes different incomplete factorizations.

Consider for example a Laplace equation discretized with the 5-point stencil over a regular Cartesian grid on a square. If $h$ is the mesh size, then the condition number of the unpreconditioned system is $O(h^{-2})$. In theory [SBG96] if $H$ is the size of each (square) subdomain, a one level Schwarz preconditioner with minimal overlap yields a preconditioned linear system with condition number $O(hH)^{-1}$. The AztecOO preconditioner delivers the Table 7.1 gives the estimated condition numbers for the corresponding AztecOO preconditioner.

The corresponding source code is contained in `didasko/examples/aztecoo/ex3.cpp`

The code uses the Trilinos_Util_CrsMatrixGallery class to create the matrix. The command to estimate the condition number with $h = 1/60$ and $H = 1/3$ is

```
mpirun -np 9 ex3.exe -problem_type=laplace_2d \
    -problem_size=900 -map_type=box
```

\(^2\)AztecOO must be configured with the option `--enable-aztecoo-azlu`, and the package Y12M is required.
7.4 AztecOO Problems as Preconditioners

Preconditioners fall into two categories. In the first category are preconditioners that are a function of the entries of the coefficient matrix. Examples include Jacobi, Gauss-Seidel, incomplete factorizations, and domain decomposition preconditioners. The second category contains preconditioners such as polynomial preconditioners that are defined by the action of the matrix on some set of vectors. Note that only category two preconditioners apply in matrix free mode. The topic of this section is another type of category two preconditioner, preconditioning one Krylov subspace method by another Krylov subspace method.

AztecOO accepts Epetra Operator objects as preconditioners. That is any class, derived from an Epetra Operator implementing the method `ApplyInverse()` may also be used as a preconditioner, using `SetPreOperator()`. AztecOO itself can be used to define a preconditioner for AztecOO; the class AztecOO_Operator (which takes an AztecOO object in the construction phase) is derived from Epetra Operator.

File `didasko/examples/aztecoo/ex2.cpp` shows how to use an AztecOO solver in the preconditioning phase. The main steps are sketched here.

First, we have to specify the linear problem to be solved (set the linear operator, the solution and the right-hand side), and create an AztecOO object:

```cpp
Epetra_LinearProblem A_Problem(&A, &x, &b);
AztecOO A_Solver(A_Problem);
```

Now, we have to define the preconditioner. For the sake of clarity, we use the same Epetra Matrix A in the preconditioning phase. However, the two matrices may differ.

```cpp
Epetra_CrsMatrix P(A);
```

(This operation is in general expensive as it involves the copy constructor. It is used here for the sake of clarity.) Then, we create the linear problem that will be used as a preconditioner. This takes a few steps to explain. Note that all the P prefix identifies preconditioner’ objects.

1. Create the linear system solve at each preconditioning step, and and we assign the linear operator (in this case, the matrix A itself)

```cpp
Epetra_LinearProblem P_Problem;
P_Problem.SetOperator(&P);
```

2. As we wish to use AztecOO to solve the preconditioner step recursively, we have to define an AztecOO object:

```cpp
AztecOO P_Solver(P_Problem);
```

### Table 7.1: Condition number for one-level domain decomposition preconditioner on a square.

<table>
<thead>
<tr>
<th>h</th>
<th>$h = 1/30$</th>
<th>$h = 1/60$</th>
<th>$h = 1/120$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$H = 1/3$</td>
<td>40.01</td>
<td>83.80</td>
<td>183.41</td>
</tr>
<tr>
<td>$H = 1/4$</td>
<td>51.46</td>
<td>106.01</td>
<td>223.22</td>
</tr>
<tr>
<td>$H = 1/6$</td>
<td>79.19</td>
<td>150.40</td>
<td>311.26</td>
</tr>
<tr>
<td>$H = 1/8$</td>
<td>-</td>
<td>191.06</td>
<td>403.29</td>
</tr>
<tr>
<td>AZ_cg</td>
<td>Conjugate Gradient method (intended for symmetric positive definite matrices).</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AZ_cg_condnum</td>
<td>AZ_cg that also estimates the condition number of the preconditioned linear system (returned in params[AZ_condnum])</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AZ_gmres</td>
<td>restarted Generalized Minimal Residual method</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AZ_gmres_condnum</td>
<td>AZ_gmres that also estimates the condition number of the preconditioned linear operator (returned in params[AZ_condnum])</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AZ_cgs</td>
<td>Conjugate Gradient Squared method</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AZ_tfqmr</td>
<td>Transpose-Free Quasi Minimal Residual method</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AZ_bicgstab</td>
<td>Bi-Conjugate Gradient with Stabilization method</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AZ_lu</td>
<td>Serial sparse direct linear solver available if AztecOO is configured with --enable-aztecoo-azlu</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 7.2: options[AZ_solver] Choices

3. Specify a particular preconditioner:

   P_Solver.SetAztecOption(AZ_precond, AZ_Jacobi);
   P_Solver.SetAztecOption(AZ_output, AZ_none);
   P_Solver.SetAztecOption(AZ_solver, AZ_cg);

4. Create an AztecOO_Operator, to set the Aztec’s preconditioner with and set the users defined preconditioners:

   AztecOO_Operator
   P_Operator(&P_Solver, 10);
   A_Solver.SetPrecOperator(&P_Operator);

   (Here 10 is the maximum number of iterations of the AztecOO solver in the preconditioning phase.)

5. Solve the linear system:

   int Niters=100;
   A_Solver.SetAztecOption(AZ_kspace, Niters);
   A_Solver.SetAztecOption(AZ_solver, AZ_gmres);
   A_Solver.Iterate(Niters, 1.0E-12);

### 7.5 Concluding Remarks on AztecOO

The following methods are often used:

- **NumIters()** returns the total number of iterations performed on this problem;
• TrueResidual() returns the true unscaled residual;

• ScaledResidual() returns the true scaled residual;

• SetAztecDefaults() can be used to restore default values in the options and params vectors.

NumIters() is useful in performance optimization. A less costly preconditioner is viable if NumIters() is “small”, and a more costly preconditioner may be worthwhile if NumIters() is “large”. Many iterative methods with right preconditioning apply the residual error norm threshold to the preconditioned residual, $P^{-1}r$, instead of $r = b - Ax$. TrueResidual() will always return the unscaled residual norm.
Several goals motivated the development of the Belos linear solver package: interoperability, extensibility, and the capability to use and develop next-generation linear solvers. The intention of interoperability is to ease the use of Belos in a wide range of application environments. To this end, the algorithms written in Belos utilize an abstract interface for operators and vectors, allowing the user to leverage existing linear algebra libraries. The concept of extensibility drives development of Belos to allow users to make efficient use of available solvers while simultaneously enabling them to easily develop their own code in the Belos framework. This is encouraged by promoting code modularization and multiple levels of access to solvers and their data. Another motivation for Belos was the capability to use and develop next-generation linear solvers. These next-generation linear solvers should enable the efficient solution of

- Single right-hand side systems: $Ax = b$
- Simultaneously solved, multiple right-hand side systems: $AX = B$
- Sequentially solved, multiple right-hand side systems: $AX_i = B_i$, for $i = 1, \ldots, t$
- Sequences of multiple right-hand side systems: $A_iX_i = B_i$, for $i = 1, \ldots, t$

where $A \in \mathbb{R}^{n \times n}$, $x, b \in \mathbb{R}^n$, and $X, B \in \mathbb{R}^{n \times k}$.

In this Chapter, we outline the Belos linear solver framework and motivate the design. In particular, we present

- the Belos operator/vector interface (Section 8.1);
- the Belos linear solver framework (Section 8.2);
- a description of Belos classes (Section 8.3);
- the interface to the Epetra linear algebra package (Section 8.4);
- an example using Belos for the solution of a linear problem (Section 8.5).
Chapter 8. Iterative Solution of Linear Systems with Belos

8.1 The Belos Operator/Vector Interface

The Belos linear solver package utilizes abstract interfaces for operators and multivectors. This allows users to leverage existing linear algebra libraries and to protect previous software investment. Algorithms in Belos are developed at a high-level, where the underlying linear algebra objects are opaque. The choice in linear algebra is made through templating, and access to the functionality of the underlying objects is provided via the traits classes Belos::MultiVecTraits and Belos::OperatorTraits. These classes define opaque interfaces, specifying the operations that multivector and operator classes must support in order to be used in Belos without exposing low-level details of the underlying data structures.

The benefit of using a templated traits class over inheritance is that the latter requires the user to derive multivectors and operators from Belos-defined abstract base classes. The former, however, defines only local requirements: Belos-defined traits classes implemented as user-developed adapters for the chosen multivector and operator classes.

Belos::MultiVecTraits provides routines for the creation of multivectors, as well as their manipulation. In order to use a specific scalar type and multivector type with Belos, there must exist a template specialization of Belos::MultiVecTraits for this pair of classes. A full list of methods required by Belos::MultiVecTraits is given in Table 8.1.

Just as Belos::MultiVecTraits defined the interface required to use a multivector class with Belos, Belos::OperatorTraits defines the interface required to use the combination of a specific operator class with a specific multivector class. This interface defines a single method:

\[
\text{OperatorTraits<} \text{ScalarType, MV, OP>::Apply(const OP &Op, const MV &X, MV &Y)}
\]

This method performs the operation \( Y = Op(X) \), where \( Op \) is an operator of type \( \text{OP} \) and \( X \) and \( Y \) are multivectors of type \( \text{MV} \). In order to use the combination of \( \text{OP} \) and \( \text{MV} \), there must be a specialization of Belos::OperatorTraits for \( \text{ScalarType, OP} \) and \( \text{MV} \).

Calling methods of MultiVecTraits and OperatorTraits requires that specializations of these traits classes have been implemented for given template arguments. Belos provides the following specializations of these traits classes:

- Epetra_MultiVector and Epetra_Operator (with scalar type double)
- Thyra::MultiVectorBase and Thyra::LinearOpBase (with arbitrary scalar type)
  This allows Belos to be used with any classes that implement the abstract interfaces provided by the Thyra package.
- Belos::MultiVec and Belos::Operator (with arbitrary scalar type)
  This allows Belos to be used with any classes that implement the abstract base classes Belos::MultiVec and Belos::Operator.

For user-specified classes that do not match one of the above, specializations of MultiVecTraits and OperatorTraits will need to be created by the user for use by Belos. Test routines Belos::TestMultiVecTraits() and Belos::TestOperatorTraits() are provided by Belos to help in testing user-developed adapters.

8.2 The Belos Linear Solver Framework

The goals of flexibility and efficiency can interfere with the goals of simplicity and ease of use. For example, efficient memory use and low-level data access required in scientific codes
8.2. The Belos Linear Solver Framework

<table>
<thead>
<tr>
<th>Method name</th>
<th>Args</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clone</td>
<td>X, n</td>
<td>Creates a new empty multivector from X containing n columns.</td>
</tr>
<tr>
<td>CloneCopy</td>
<td>X</td>
<td>Creates a new multivector from X with a copy of all the vectors in X (deep copy).</td>
</tr>
<tr>
<td>CloneCopy</td>
<td>X, index</td>
<td>Creates a new multivector from X with only a copy of the index vectors (deep copy).</td>
</tr>
<tr>
<td>CloneView</td>
<td>X, index</td>
<td>Creates a new multivector from X that shares the index vectors from X (shallow copy).</td>
</tr>
<tr>
<td>GetVecLength</td>
<td>X</td>
<td>Returns the vector length of the multivector X.</td>
</tr>
<tr>
<td>GetNumberVecs</td>
<td>X</td>
<td>Returns the number of vectors in the multivector X.</td>
</tr>
<tr>
<td>MvTimesMatAddMv</td>
<td>X, Y, α, β</td>
<td>Apply a SerialDenseMatrix M to another multivector X, Y ← αXM + βY.</td>
</tr>
<tr>
<td>MvAddMv</td>
<td>X, Y, Z, α, β</td>
<td>Perform Z ← αX + βY.</td>
</tr>
<tr>
<td>MvTransMv</td>
<td>X, Y, α, C</td>
<td>Compute the matrix C ← αXH Y.</td>
</tr>
<tr>
<td>MvDot</td>
<td>X, Y, d</td>
<td>Compute the vector d where the components are the individual dot-products of the i-th columns of X and Y, i.e. d[i] = X[i]H Y[i].</td>
</tr>
<tr>
<td>MvScale</td>
<td>X, α</td>
<td>Scale the columns of the multivector X by α.</td>
</tr>
<tr>
<td>MvNorm</td>
<td>X, norm</td>
<td>Compute the 2-norm of each individual vector of X, norm[i] =</td>
</tr>
<tr>
<td>SetBlock</td>
<td>X, Y, index</td>
<td>Copy the vectors in X to the subset of vectors in Y indicated by index.</td>
</tr>
<tr>
<td>MvRandom</td>
<td>X</td>
<td>Replace the vectors in X with random vectors.</td>
</tr>
<tr>
<td>MvInit</td>
<td>X, α</td>
<td>Replace each element of X with α.</td>
</tr>
<tr>
<td>MvPrint</td>
<td>X, os</td>
<td>Print the multivector X to an output stream os.</td>
</tr>
</tbody>
</table>

Table 8.1: Methods required by Belos::MultiVecTraits interface.

can lead to complicated interfaces and violations of standard object-oriented development practices. In Belos, this problem is addressed by providing a multi-tiered access strategy for linear solver algorithms. Belos users have the choice of interfacing at one of two levels: either working at a high-level with a linear solver manager or working at a low-level directly with an iteration.

Consider as an example the conjugate gradient (CG) iteration. The essence of this iteration can be distilled into the following steps:

1. apply operator A to current direction vector \( p_i \): \( Ap_i = A * p_i \)
2. use \( Ap_i \) to compute step length \( \alpha_i \): \( \alpha_i = \frac{<r_i, r_i>}{<p_i, Ap_i>} \)
3. use \( \alpha_i \) and \( p_i \) to compute approximate solution \( x_{i+1} \): \( x_{i+1} = x_i + \alpha_i * p_i \)
4. compute new residual \( r_{i+1} \) using step length and \( Ap_i \): \( r_{i+1} = r_i - \alpha_i * Ap_i \)
5. compute improvement in the step $\beta_i$ using $r_i$ and $r_{i+1}$: 
\[ \beta_i = \frac{<r_{i+1}, r_{i+1}>}{<r_i, r_i>} \]

6. use $\beta_i$ and $r_{i+1}$ to compute the new direction vector $p_{i+1}$: 
\[ p_i = r_{i+1} + \beta_i \ast p_i \]

In implementing a CG solver, this iteration repeats until some stopping criterion has been satisfied. Many valid stopping criteria exist, however, they are distinct from the essential iteration as described above. A user wanting to perform CG iterations could ask the solver to perform these iterations until a user-specified stopping criterion, like maximum number of iterations or residual norm, was satisfied. This allows the user complete control over the stopping criteria and leaves the iteration responsible for a relatively simple bit of state and behavior.

This is the way that Belos has been designed. The iterations (encapsulating an iteration kernel and the associated state) are derived classes of the abstract base class Belos::Iteration. The goals of this class are three-fold:

- to define an interface used for checking the status of an iteration by a status test;
- to contain the iteration kernel associated with a particular linear solver;
- to contain the state associated with that iteration.

The status tests, assembled to describe a specific stopping criterion and queried by the iteration, are represented as subclasses of Belos::StatusTest. The communication between status test and iteration occurs inside of the iterate() method provided by each Belos::Iteration. This code generally takes the form:

```cpp
SomeIteration::iterate() {
    while ( statustest.checkStatus(this) != Passed ) {
        // perform iteration kernel
        //
    }
    return; // return back to caller
}
```

Each Belos::StatusTest provides a method, checkStatus(), which through queries to the methods provided by Belos::Iteration, determines whether the solver meets the criteria defined by that particular status test. After an iteration returns from iterate(), the caller has the option of accessing the state associated with the iteration and re-initializing it with a new state.

While this method of interfacing with the iteration is powerful, it can be tedious. This method requires that a user construct a number of support classes, in addition to managing calls to iterate(). The Belos::SolverManager class was developed to address this need. A solver manager is a class that wraps around an iteration, providing additional functionality while also handling lower-level interaction with the iteration that a user may not wish to handle. Solver managers are intended to be easy to use, while still providing the features and flexibility needed to solve real-world linear problems. For example, the Belos::BlockCGSolMgr takes only two arguments in its constructor: a Belos::LinearProblem specifying the linear problem to be solved and a Teuchos::ParameterList of options specific to this solver manager. The solver manager instantiates an iteration, along with the status
tests and other support classes needed by the iteration. To solve the linear problem, the user simply calls the \texttt{solve()} method of the solver manager. The solver manager performs repeated calls to the \texttt{iterate()} method, performs restarts or recycling, and places the final solution into the linear problem.

Users therefore have a number of options for computing solutions to linear problems with Belos:

- use an existing solver manager;
  In this case, the user is limited to the functionality provided by the current linear solvers.

- develop a new solver manager around an existing iteration;
  The user can extend the functionality provided by the iteration, specifying custom configurations for status tests, orthogonalization, restarting, recycling, etc.

- develop a new iteration/solver manager;
  The user can write an iteration that is not represented in Belos. The user still has the benefit of the support classes provided by Belos, and the knowledge that the new iteration / solver manager can be easily used by anyone already familiar with Belos.

\section{Belos Classes}

Belos is designed with extensibility in mind, so that users can augment the package with any special functionality that they need. However, the released version of Belos provides all functionality necessary for solving a wide variety of problems. This section lists and briefly describes the current abstract classes found in Belos. Each subsection also includes information on the derived classes provided by Belos.

\textbf{Remark 14.} \textit{Belos makes extensive use of the Teuchos utility classes, especially Teuchos::RCP (Section 10.7) and Teuchos::ParameterList (Section 10.6). Users are encouraged to become familiar with these classes and their correct usage.}

\subsection{Belos::LinearProblem}

Belos::LinearProblem is a templated container for the components of a linear problem, as well as the solutions. Both the linear problem and the linear solver in Belos are templated on the scalar type, the multivector type and the operator type. Before declaring a linear problem, users must choose classes to represent these entities. Having done so, they can begin to specify the parameters of the linear problem using the Belos::LinearProblem \texttt{set} methods:

- \texttt{setOperator} - set the operator $A$ for which the solutions will be computed
- \texttt{setLHS} - set the left-hand side (solution) vector $X$ for the linear problem $AX = B$
- \texttt{setRHS} - set the right-hand side vector $B$ for the linear problem $AX = B$
- \texttt{setLeftPrec} - set the left preconditioner for the linear problem
- \texttt{setRightPrec} - set the right preconditioner for the linear problem
- \texttt{setHermitian} - specify whether the problem is Hermitian
• setLabel - specify the label prefix used by the timers in this object

In addition to these set methods, Belos::LinearProblem defines a method setProblem() that gives the class the opportunity to perform any initialization that may be necessary before the problem is handed off to a linear solver, in addition to verifying that the problem has been adequately defined.

For each of the set methods listed above, there is a corresponding get function. These are the functions used the iterations and solver managers to get necessary information from the linear problem.

8.3.2 Belos::Iteration

The Belos::Iteration class defines the basic interface that must be met by any iteration class in Belos. The specific iterations are implemented as derived classes of Belos::Iteration. Table 8.2 lists the linear solver currently implemented in Belos.

<table>
<thead>
<tr>
<th>Solver</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>CG</td>
<td>A basic single-vector CG iteration for SPD linear problems.</td>
</tr>
<tr>
<td>BlockCG</td>
<td>A block CG iteration for SPD linear problems.</td>
</tr>
<tr>
<td>BlockGmres</td>
<td>A block GMRES iteration for non-Hermitian linear problems.</td>
</tr>
<tr>
<td>BlockFGmres</td>
<td>A block flexible GMRES iteration for non-Hermitian linear problems.</td>
</tr>
<tr>
<td>PseudoBlockGmres</td>
<td>A simultaneous single-vector GMRES iteration for non-Hermitian linear problems.</td>
</tr>
<tr>
<td>GCRODR</td>
<td>A deflation, restarted GCRO iteration for non-Hermitian linear problems.</td>
</tr>
</tbody>
</table>

Table 8.2: Iterations currently implemented in Belos.

The iteration interface provides two significant types of methods: status methods and solver-specific state methods. The status methods are defined by the Belos::Iteration abstract base class and represent the information that a generic status test can request from any linear solver. A list of these methods is given in Table 8.3.

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>getNumIters</td>
<td>Get the current number of iterations.</td>
</tr>
<tr>
<td>resetNumIters</td>
<td>Reset the number of iterations.</td>
</tr>
<tr>
<td>getNativeResiduals</td>
<td>Get the most recent residual norms native to the iteration.</td>
</tr>
<tr>
<td>getCurrentUpdate</td>
<td>Get the most recent solution update computed by the iteration.</td>
</tr>
</tbody>
</table>

Table 8.3: A list of generic status methods provided by Belos::Iteration.

The class Belos::Iteration, like Belos::LinearProblem, is templated on the scalar type, multivector type and operator type. The options for the iteration are passed in through the constructor, defined by Belos::Iteration to have the following form:
These classes are used as follows:

- **problem** - the linear problem to be solved; the solver will get the operator and vectors from here; see Section 8.3.1.

- **printer** - the output manager dictates verbosity level in addition to processing output streams; see Section 8.3.6.

- **tester** - the status tester dictates when the solver should quit `iterate()` and return to the caller; see Section 8.3.4.

- **ortho** - the orthogonalization manager defines the inner product and other concepts related to orthogonality, in addition to performing these computations for the solver; see Section 8.3.5.

- **params** - the parameter list specifies linear solver-specific options; see the documentation for a list of options support by individual solvers.

An iteration class also specifies a concept of state, i.e. the current data associated with the iteration. After declaring an iteration object, it is in an uninitialized state. For most iterations, to be initialized means to be in a valid state, containing all of the information necessary for performing an iteration step. Belos::Iteration provides two methods concerning initialization: `isInitialized()` indicates whether the iteration is initialized or not, and `initialize()` (with no arguments) instructs the iteration to initialize itself using the linear problem.

To ensure that iterations can be used as efficiently as possible, the user needs access to their state. To this end, each iteration provides low-level methods for getting and setting their state:

- **getState()** - returns an iteration-specific structure with read-only pointers to the current state of the iteration.

- **initialize(...)** - accepts an iteration-specific structure enabling the user to initialize the iteration object with a particular state.

The combination of these two methods, along with the flexibility provided by status tests, allows the user almost total control over linear solver iterations.

### 8.3.3 Belos::SolverManager

Using Belos by interfacing directly with the linear solver iterations is extremely powerful, but can be more entailed than necessary. Solver managers provide a way for users to encapsulate
specific solving strategies inside of an easy-to-use class. Novice users may prefer to use existing solver managers, while advanced users may prefer to write custom solver managers.

The Belos::SolverManager class provides a parameter list driven interface for solving linear systems and, as such, has very few methods. Only two constructors are supported: a default constructor and a constructor accepting a Belos::LinearProblem and a parameter list of solver manager-specific options. If the default constructor is used, the Belos::LinearProblem and parameter list can be passed in, post-construction, using the methods setProblem(...) and setParameters(...), respectively. It is also possible to get the valid parameters and current parameters from the solver manager by using the methods getValidParameters() and getCurrentParameters(), respectively. Most importantly, there is a solve() method that takes no arguments and returns either Belos::Converged or Belos::Unconverged. Consider the following simple example code:

```c++
// create a linear problem
Teuchos::RCP<Belos::LinearProblem<ScalarType,MV,OP> > problem = ...;
// create a parameter list
Teuchos::RCP<Teuchos::ParameterList> params;
params->set(...);
// create a solver manager
Belos::BlockCGSolMgr<ScalarType,MV,OP> CGsolver( problem, params );
// solve the linear problem
Belos::ReturnType ret = CGsolver.solve();
// get the solution from the problem
Teuchos::RCP< MV > sol = problem->getLHS();
```

**Remark 15.** Errors in Belos are communicated via exceptions. This is outside the scope of this tutorial; see the Belos documentation for more information.

As has been stated before, the goal of the solver manager is to create a linear solver iteration object, along with the necessary support objects. Another purpose of many solver managers is to manage and initiate the repeated calls to the underlying iteration’s iterate() method. For linear solver iterations that build a Krylov subspace to some maximum dimension (e.g., BlockGmres, BlockFGmres, etc.), the solver manager will also employ a strategy for restarting the solver when the subspace is full. This is something for which multiple approaches are possible. Also, there may be substantial flexibility in creating the support classes (e.g., sort manager, status tests) for the solver. An aggressive solver manager could even go so far as to construct a preconditioner for the linear problem, or switch its solution technique based on convergence behavior.

These examples are meant to illustrate the flexibility that specific solver managers may have in implementing the solve() routine. Some of these options might best be incorporated into a single solver manager, which takes orders from the user via the parameter list. Some of these options may better be contained in multiple solver managers, for the sake of code simplicity. It is even possible to write solver managers that contain other solvers managers; motivation for something like this would be to select the optimal solver manager at runtime based on some expert knowledge, or to create a hybrid method which uses the output from one solver manager to initialize another one.
8.3.4 Belos::StatusTest

By this point in the tutorial, the purpose of the Belos::StatusTest should be clear: to give the user or solver manager flexibility in stopping the linear solver iterations in order to interact directly with the iteration.

Many reasons exist for why a user would want to stop the iteration from continuing:

- some convergence criterion has been satisfied and it is time to quit;
- some part of the current solution has reached a sufficient accuracy to removed from the iteration;
- the solver has performed a sufficient or excessive number of iterations.

These are just some commonly seen reasons for ceasing the iteration, and each of these can be so varied in implementation/parametrization as to require some abstract mechanism controlling the iteration.

The following is a list of Belos-provided status tests:

- Belos::StatusTestCombo - this status test allows for the boolean combination of other status tests, creating near unlimited potential for complex status tests.
- Belos::StatusTestOutput - this status test acts as a wrapper around another status test, allowing for printing of status information on a call to checkStatus()
- Belos::StatusTestMaxIters - this status test monitors the number of iterations performed by the solver; it can be used to halt the solver at some maximum number of iterations or even to require some minimum number of iterations.
- Belos::StatusTestGenResNorm - this status test allows the user to construct one of a family of residual tests to monitor the residual norms of the current iterate.
- Belos::StatusTestImpResNorm - this status test monitors the implicit residual norm (e.g., native residual available through GMRES) and checks for loss of accuracy.

8.3.5 Belos::OrthoManager

Orthogonalization and orthonormalization are commonly performed computations in iterative linear solvers; in fact, for some linear solvers, they represent the dominant cost. Different scenarios may require different approaches (e.g., Euclidean inner product versus a weighted inner product, full versus partial orthogonalization). Combined with the plethora of available methods for performing these computations, Belos has left as much leeway to the users as possible.

Orthogonalization of multivectors in Belos is performed by derived classes of the abstract class Belos::OrthoManager. This class provides five methods:

- innerProd(X,Y,Z) - performs the inner product defined by the manager.
- norm(X) - computes the norm induced by innerProd().
- project(X,C,Q) - given an orthonormal basis Q, projects X onto the space perpendicular to colspan(Q), optionally returning the coefficients of X in Q.
Chapter 8. Iterative Solution of Linear Systems with Belos

- **normalize**(X,B) - returns an orthonormal basis for \(\text{colspan}(X)\), optionally returning the coefficients of \(X\) in the computed basis.

- **projectAndNormalize**(X,C,B,Q) - computes an orthonormal basis for subspace \(\text{colspan}(X) - \text{colspan}(Q)\), optionally returning the coefficients of \(X\) in \(Q\) and the new basis.

It should be noted that a call to **projectAndNormalize()** is not necessarily equivalent to a call to **project()** followed by **normalize()**. This follows from the fact that, for some orthogonalization managers, a call to **normalize()** may augment the column span of a rank-deficient multivector in order to create an orthonormal basis with the same number of columns as the input multivector. In this case, the code

```cpp
orthoMgr.project(X,C,Q);
orthoMgr.normalize(X,B);
```

could result in an orthonormal basis \(X\) that is not orthogonal to the basis in \(Q\).

Belos provides three orthogonalization managers:

- Belos::DGKSOrthoManager - performs orthogonalization using classical Gram-Schmidt with a possible correction step.

- Belos::ICGSOrthoManager - performs orthogonalization using iterated classical Gram-Schmidt.

- Belos::IMGSOrthoManager - performs orthogonalization using iterated modified Gram-Schmidt.

More information on these orthogonalization managers is available in the Belos documentation.

### 8.3.6 Belos::OutputManager

The output manager is a concrete class in Belos and exists to provide flexibility with regard to the verbosity of the linear solver. The output manager has two primary concerns: what output is printed and where the output is printed to. When working with the output manager, output is classified into one of the message types from Table 8.4.

The output manager in Belos provides the following output-related methods:

- **bool setVerbosity** (MsgType type) - Set the type of messages we need to print out information for.

- **bool setOStream** (const Teuchos::RCP<std::ostream> &os) - Set the output stream where information should be sent.

- **bool isVerbosity** (MsgType type) - Find out whether we need to print out information for this message type.

- **void print** (MsgType type, const string output) - Send output to the output manager.

- **ostream & stream** (MsgType type) - Create a stream for outputting to.
8.4 Using the Belos adapter to Epetra

The output manager is meant to ease some of the difficulty associated with I/O in a distributed programming environment. For example, consider some debugging output requiring optional computation. For reasons of efficiency, we may want to perform the computation only if debugging is requested; i.e., `isVerbosity(Belos::Debug) == true`. However, while we need all nodes to enter the code block to perform the computation, we probably want only one of them to print the output. For the Belos::OutputManager, the output corresponding to the verbosity level of the manager is sent to the output stream only on the master node; the output for other nodes is neglected.

### Table 8.4: Message types used by Belos::OutputManager.

<table>
<thead>
<tr>
<th>Message type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Errors</td>
<td>Errors (always printed)</td>
</tr>
<tr>
<td>Warnings</td>
<td>Warning messages</td>
</tr>
<tr>
<td>IterationDetails</td>
<td>Approximate eigenvalues, errors</td>
</tr>
<tr>
<td>OrthoDetails</td>
<td>Orthogonalization/orthonormalization checking</td>
</tr>
<tr>
<td>FinalSummary</td>
<td>Final computational summary (usually from SolverManager::solve())</td>
</tr>
<tr>
<td>TimingDetails</td>
<td>Timing details</td>
</tr>
<tr>
<td>StatusTestDetails</td>
<td>Status test details</td>
</tr>
<tr>
<td>Debug</td>
<td>Debugging information</td>
</tr>
</tbody>
</table>

8.4 Using the Belos adapter to Epetra

The Epetra package provides the underlying linear algebra foundation for many Trilinos solvers. By using the Belos adapter to Epetra, users not only avoid the trouble of implementing their own multivector and operator classes, but they also gain the ability to utilize any other Trilinos package which recognizes Epetra classes (such as AztecOO, IFPACK, ML, and others).

In order to use the Belos adapter to Epetra, users must include the following file:

```cpp
#include "BelosEpetraAdapter.hpp"
```

This file simply defines specializations of the Belos::MultiVecTraits and Belos::OperatorTraits classes, while also including the Epetra header files defining the multivector and operator classes.

Because Epetra makes exclusive use of double precision arithmetic, `Epetra_Operator` and `Epetra_MultiVector` are used only with scalar type `double`. For brevity, it is useful to declare type definitions for these classes:

```cpp
typedef double ST;
typedef Epetra_MultiVector MV;
typedef Epetra_Operator OP;
```

Multivectors will be of type `MV`:

```cpp
Teuchos::RCP<MV> X = Teuchos::rcp( new MV(...) );
```
Operators can be any subclass of `OP`, for example, an Epetra_CrsMatrix:

```cpp
Teuchos::RCP<OP> A
    = Teuchos::rcp( new Epetra_CrsMatrix(...) );
```

The Belos interface to Epetra defines a specialization of Belos::MultiVecTraits for Epetra_MultiVector and a specialization of Belos::OperatorTraits for Epetra_Operator applied to Epetra_MultiVector. Therefore, we can now specify a linear solver and preconditioner utilizing these computational classes. An example of defining and solving a linear problem using a Belos linear solver is given in the next section.

### 8.5 Defining and Solving a Linear Problem

This section gives sample code for solving a symmetric positive definite (SPD) linear problem using the block CG solver manager, Belos::BlockCGSolMgr. The example code in this section comes from the Didasko example `didasko/examples/belos/ex1.cpp`.

The first step in solving a linear problem is to define the problem using the Belos::LinearProblem class. Assume we have chosen classes to represent our scalars, multivectors and operators as `ST`, `MV` and `OP`, respectively. Given an operator `A`, a multivector `X` containing the initial guess, and a multivector `B` containing the right-hand side, all wrapped in Teuchos::RCP, we would define the linear problem as follows:

```cpp
Teuchos::RCP< LinearProblem<ST,MV,OP> > myProblem
    = Teuchos::rcp( new LinearProblem<ST,MV,OP>(A,X,B) );
myProblem->setHermitian();
bool ret = myProblem->setProblem();
if (ret != true) {
    // there should be no error in this example :)
}
```

The first line creates a Belos::LinearProblem object and wraps it in a Teuchos::RCP (Section 10.7). The second line specifies the symmetry of the linear problem. The third line signals that we have finished setting up the linear problem. This step must be completed before attempting to solve the problem; failure to do so will result in the solver manager throwing an exception.

Next, we create a parameter list wrapped in a Teuchos::RCP to specify the parameters for the solver manager:

```cpp
int verb = Belos::Warnings + Belos::Errors
    + Belos::FinalSummary + Belos::TimingDetails;
Teuchos::RCP<Teuchos::ParameterList> myPL;
myPL->set( "Verbosity", verb );
myPL->set( "Block Size", 4 );
myPL->set( "Maximum Iterations", 100 );
myPL->set( "Convergence Tolerance", 1.0e-8 );
```

Here, we have asked for the linear solver to output information regarding errors and warnings, as well as to provide a final summary after completing all iterations and print the timing information collected during the solve. We have also specified the tolerance for
convergence testing (used to construct a status test); the block size; and the maximum number of iterations (used to construct a status test). This solver manager permits other options as well, affecting the block size as well as the output; see the Belos documentation.

We now have all of the information needed to declare the solver manager and solve the problem:

```cpp
Belos::BlockCGSolMgr<ST,MV,OP> mySolver( myProblem, myPL );
```

The linear problem is solved with the instruction

```cpp
Belos::ReturnType solverRet = mySolver.solve();
```

The return value of the solver indicates whether the algorithm succeeded or not; i.e., whether the requested residual tolerance was achieved in the allotted number of iterations. Output from `solve()` routine in this example might look as follows:

Passed.......OR Combination ->

OK............Number of Iterations = 21 < 100
Converged....(2-Norm Imp Res Vec) / (2-Norm Res0)
  residual [ 0 ] = 6.38815e-09 < 1e-08
  residual [ 1 ] = 9.08579e-09 < 1e-08
  residual [ 2 ] = 9.26932e-09 < 1e-08
  residual [ 3 ] = 4.43278e-09 < 1e-08

================================================================================
TimeMonitor Results
Timer Name Local time (num calls)
================================================================================
Belos: Operation Op*x 0.000583 (22)
Belos: Operation Prec*x 0 (0)
Epetra_CrsMatrix::Multiply(TransA,X,Y) 0.000453 (22)
Belos: Orthogonalization 0.006996 (23)
Belos: BlockCGSolMgr total solve time 0.01208 (1)
================================================================================

The solution vector can be retrieved from the linear problem (where they were stored by the solver manager) as follows:

```cpp
Teuchos::RCP<MV> sol = myProblem->getLHS();
```

Then the residual can be checked using the computed solution, resulting in output that should look like:

```
***********************************************************
Results (outside of linear solver)
***********************************************************
Linear System Relative Residual
```
<table>
<thead>
<tr>
<th></th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>6.388147e-09</td>
</tr>
<tr>
<td>2</td>
<td>9.085794e-09</td>
</tr>
<tr>
<td>3</td>
<td>9.269324e-09</td>
</tr>
<tr>
<td>4</td>
<td>4.432775e-09</td>
</tr>
</tbody>
</table>

******************************************************
In this Chapter we discuss the use of IFPACK objects as preconditioners for AztecOO solvers. Specifically we present:

- Parallel distributed memory issues (in Section 9.2).
- Incomplete factorizations and notation (in Section 9.1).
- How to compute incomplete Cholesky factorizations (in Section 9.3).
- IFPACK’s RILU-type factorizations (in Section 9.4).

### 9.1 Theoretical Background

The aim of this section is to define concepts associated with incomplete factorization methods and establish our notation. This section is not supposed to be exhaustive, nor complete on this subject. The reader is referred to the existing literature for a comprehensive presentation.

A broad class of effective preconditioners is based on incomplete factorization of the linear system matrix. Such preconditioners are often referred to as incomplete lower/upper (ILU) preconditioners. ILU preconditioning techniques lie between direct and iterative methods and provide a balance between reliability and numerical efficiency. ILU preconditioners are constructed in the factored form $P = \tilde{L}\tilde{U}$, with $\tilde{L}$ and $\tilde{U}$ being lower and upper triangular matrices. Solving with $P$ involves two triangular solutions.

ILU preconditioners are based on the observation that, although most matrices $A$ admit an LU factorization $A = LU$, where $L$ is (unit) lower triangular and $U$ is upper triangular, the factors $L$ and $U$ often contain too many nonzero terms, making the cost of factorization too expensive in time or memory use, or both. One type of ILU preconditioner is ILU(0),
which is defined as proceeding through the standard LU decomposition computations, but
keeping only those terms in $L$ that correspond to nonzero terms in the lower triangle of $A$
and similarly keeping only those terms in $U$ that correspond to nonzero terms in the upper
triangle of $A$. Although effective, in some cases the accuracy of the ILU(0) may be insufficient
to yield an adequate rate of convergence. More accurate factorizations will differ from ILU(0)
by allowing some fill-in. The resulting class of methods is called ILU($k$), where $k$ is the level-
of-fill. A level-of-fill is attributed to each element that is processed by Gaussian elimination,
and dropping will be based on the level-of-fill. The level-of-fill should be indicative of the size
of the element: the higher the level-of-fill, the smaller the elements.

Other strategies consider dropping by value – for example, dropping entries smaller than
a prescribed threshold. Alternative dropping techniques can be based on the numerical size
of the element to be discarded. Numerical dropping strategies generally yield more accurate
factorizations with the same amount of fill-in as level-of-fill methods. The general strategy is
to compute an entire row of the $L$ and $U$ matrices, and then keep only a certain number of
the largest entries. In this way, the amount of fill-in is controlled; however, the structure of
the resulting matrices is undefined. These factorizations are usually referred to as ILUT, and
a variant which performs pivoting is called ILUTP.

When solving a single linear system, ILUT methods can be more effective than ILU($k$). However, in many situations a sequence of linear systems must be solved where the pattern
of the matrix $A$ in each system is identical but the values of changed. In these situations,
ILU($k$) is typically much more effective because the pattern of ILU($k$) will also be the same
for each linear system and the overhead of computing the pattern is amortized.

### 9.2 Parallel Incomplete Factorizations

Parallel direct sparse solvers that compute the complete factorization $A = LU$ are effective
on parallel computers. However, the effective scalability of these solvers is typically limited
to a speedup of order ten, regardless of the number of processors used. Also, it is typically
the factorization (constructing $L$ and $U$) that exhibits the best parallel speedup. The forward
and back triangular solves typically exhibit very poor parallel speedup.

The situation for ILU preconditioners is even worse. Complete factorizations can scale
well because of very important graph properties that can be determined at low cost. ILU
factorizations do not have the same properties, so predicting fill-in across the parallel machine
is not practically possible. Also, because ILU preconditioners require repeated forward and
back solves, they are more affected by the poor scalability of these operations.

Because ILU preconditioners do not scale well on parallel computers, a common practice
is to perform local ILU factorizations. In this situation, each processor computes a factoriza-
tion of a subset of matrix rows and columns independently from all other processors. This
additional layer of approximation leads to a block Jacobi type of preconditioner across pro-
cessors, where each block is solved using an ILU preconditioner. The difficulty with this type
of preconditioner is that it tends to become less robust and require more iterations as the
number of processors used increases. This effect can be offset to some extent by allowing
overlap. Overlap refers to having processors redundantly own certain rows of the matrix for
the ILU factorization. Level-1 overlap is defined so that a processor will include rows that
are part of its original set. In addition, if row $i$ is part of its original set and row $i$ of $A$ has a
nonzero entry in column $j$, then row $j$ will also be included in the factorization on that pro-
9.3. Incomplete Cholesky Factorizations

Recall that if a matrix is symmetric positive definite, it admits a Cholesky factorization of the form $A = LL^T$, where $L$ is lower triangular. Ifpack_CrsIct is a class for constructing and using incomplete Cholesky factorizations of an Epetra_CrsMatrix. It is built in part on top of the ICT preconditioner developed by Edmond Chow at Lawrence Livermore National Laboratory [LSC04]. Specific factorizations depend on several parameters:

- Maximum number of entries per row/column. The factorization will contain at most this number of nonzero elements in each row/column;
- Diagonal perturbation. By default, the factorization will be computed on the input matrix. However, it is possible to modify the diagonal entries of the matrix to be factorized, via functions SetAbsoluteThreshold() and SetRelativeThreshold(). Refer to the IFPACK’s documentation for more details.

It is easy to have IFPACK compute the incomplete factorization. First, define an Ifpack_CrsIct object,

```cpp
Ifpack_CrsIct * ICT = NULL;
ICT = Ifpack_CrsIct(A, DropTol, LevelFill);
```

where $A$ is an Epetra_CrsMatrix (already FillComplete’d), and DropTol and LevelFill are the drop tolerance and the level-of-fill, respectively. Then, we can set the values and compute the factors,

```cpp
ICT->InitValues(A);
ICT->Factor();
```

IFPACK can compute the estimation of the condition number

$$\text{cond}(L_iU_i) \approx \| (LU)^{-1} e \|_\infty,$$

where $e = (1, 1, \ldots, 1)^T$. (More details can be found in the IFPACK documentation.) This estimation can be computed as follows:

```cpp
double Condest;
ICT->Condest(false, Condest);
```

Please refer to file didasko/examples/ifpack/ex1.cpp for a complete example of incomplete Cholesky factorization.

9.4 RILUK Factorizations

IFPACK implements various incomplete factorization for non-symmetric matrices. In this Section, we will consider the Epetra_CrsRiluk class, that can be used to produce RILU factorization of a Epetra_CrsMatrix. The class required an Ifpack_OverlapGraph in the construction phase. This means that the factorization is split into two parts:
1. Definition of the level filled graph;
2. Computation of the factors.

This approach can significantly improve the performances of code, when an ILU preconditioner has to be computed for several matrices, with different entries but with the same sparsity pattern. An Ifpack_IlukGraph object of an Epetra matrix $A$ can be constructed as follows:

```cpp
Ifpack_IlukGraph Graph =
    Ifpack_IlukGraph(A.Graph(),LevelFill,LevelOverlap);
```

Here, `LevelOverlap` is the required overlap among the subdomains.

A call to `ConstructFilledGraph()` completes the process.

**Remark 16.** An Ifpack_IlukGraph object has two Epetra_CrsGraph objects, containing the $L_i$ and $U_i$ graphs. Thus, it is possible to manually insert and delete graph entries in $L_i$ and $U_i$ via the Epetra_CrsGraphInsertIndices and RemoveIndices functions. However, in this case FillComplete must be called before the graph is used for subsequent operations.

At this point, we can create an Ifpack_CrsRiluk object,

```cpp
ILUK = Ifpack_CrsRiluk(Graph);
```

This phase defined the graph for the incomplete factorization, without computing the actual values of the $L_i$ and $U_i$ factors. Instead, this operation is accomplished with

```cpp
int initerr = ILUK->InitValues(A);
```

The ILUK object can be used with AztecOO simply setting

```cpp
solver.SetPrecOperator(ILUK);
```

where `solver` is an AztecOO object. The example in `didasko/examples/ifpack/ex2.cpp` shows the use of the Ifpack_CrsRiluk class.

The application of the incomplete factors to a global vector, $z = (L_i U_i^{-1}) r$, results in redundant approximation for any element of $z$ that correspond to rows that are part of more than one local ILU factor. The OverlapMode defines how those redundant values are managed. OverlapMode is an Epetra_CombinedMode enum, that can assume the following values: Add, Zero, Insert, Average, AbxMax. The default is to zero out all the values of $z$ for rows that were not part of the original matrix row distribution.

### 9.5 Concluding Remarks on IFPACK

More documentation on the IFPACK package can be found in [SH05a]
The Teuchos Utility Classes

Heidi Thornquist

Teuchos (pronounced “te-fos”) is a collection of portable C++ tools that facilitate the development of scientific codes. Only a few of the many tools in Teuchos are mentioned in this section. For more details on all of the capabilities provided by Teuchos, please refer to the online documentation (http://trilinos.sandia.gov/packages/teuchos). Teuchos classes have been divided between a “standard” build and an “extended” build. The “standard” build contains the general purpose templated tools like BLAS/LAPACK wrappers, parameter lists, a command-line parser, serial dense matrices, timers, flop counters, and a reference-counted pointer class. These tools are built by default when Teuchos is enabled using the configure option --enable-teuchos. The “extended” build contains more special purpose tools like XML parsing and MPI communicators, which can be included in the Teuchos library by using the configure option --enable-teuchos-extended.

10.1 Introduction

In this Chapter, we will present the following “standard” build classes:

- **Teuchos::ScalarTraits** class (Section 10.2): The ScalarTraits class provides a basic interface to scalar types (float, double, complex<float>, complex<double>) that is used by the templated computational classes within Teuchos. It is the mechanism by which Teuchos’ capabilities can be extended to support arbitrary precisions.

- **Teuchos::SerialDenseMatrix** class (Section 10.3): The SerialDenseMatrix is a templated version of the Epetra_SerialDenseMatrix class that is most often used to interface with the templated BLAS/LAPACK wrappers.

- **Teuchos::BLAS** class (Section 10.4): The BLAS class provides templated wrappers for the native BLAS library and can be extended to support arbitrary precision computations.

- **Teuchos::LAPACK** class (Section 10.5): The LAPACK class provides templated wrappers for the native LAPACK library.

- **Teuchos::ParameterList** class (Section 10.6): ParameterList is a container that can be used to group all the parameters required by a given piece of code.
• **Teuchos::RCP** class (Section 10.7): RCP is a smart reference-counted pointer class, which provides a functionality similar to the garbage collector of Java.

• **Teuchos::TimeMonitor** class (Section 10.8): TimeMonitor is a timing class that starts a timer when it is initialized and stops it when the destructor is called on the class.

• **Teuchos::CommandLineProcessor** class (Section 10.9): CommandLineProcessor is a class that helps parse command line input arguments from (argc, argv[ ]).

## 10.2 **Teuchos::ScalarTraits**

The ScalarTraits class provides a basic interface to scalar types (float, double, complex<float>, complex<double>) that is used by the templated computational classes within Teuchos. This interface includes a definition of the magnitude type and methods for obtaining random numbers, representations of zero and one, the square root, and machine-specific parameters. The Teuchos classes that utilize this scalar traits mechanism are **Teuchos::SerialDenseMatrix**, **Teuchos::BLAS**, and **Teuchos::LAPACK**.

ScalarTraits enables the extension of Teuchos’ computational capabilities to any scalar type that can support its basic interface. In particular, this interface can be used for arbitrary precision scalar types. An interface to the arbitrary precision library ARPREC [BHLT02] is available if Teuchos is configured with `--enable-teuchos-arprec`. Teuchos must also be configured with the local ARPREC library paths (`--with-libs`, `--with-incdirs`, and `--with-libdirs`). To obtain more information on ARPREC or download the source code, see [http://crd.lbl.gov/~dhbailey/mpdist/](http://crd.lbl.gov/~dhbailey/mpdist/).

**Remark 17.** To enable complex arithmetic (complex<float> or complex<double>) support in ScalarTraits or any dependent classes, configure Teuchos with `--enable-teuchos-complex`.

## 10.3 **Teuchos::SerialDenseMatrix**

**Teuchos::SerialDenseMatrix** is a templated version of the SerialDenseMatrix class in Epetra (Chapter 3). It is most useful for interfacing with the templated BLAS and LAPACK wrappers, which will be discussed in Sections 10.4 and 10.5. However, by enabling the simple construction and manipulation of small dense matrices, the SerialDenseMatrix class has also been used as an independent tool in many Trilinos packages.

**Teuchos::SerialDenseMatrix** provides a serial interface to a small dense matrix of templated scalar type. This means a SerialDenseMatrix object can be created for any scalar type supported by Teuchos::ScalarTraits (Section 10.2). Boundschecking can be enabled for this class by configuring Teuchos with `--enable-teuchos-abc`. An exception will be thrown every time a matrix bound is violated by any method. This incurs a lot of overhead for this class, so boundschecking is only recommended as a debugging tool.

To use the **Teuchos::SerialDenseMatrix** class, include the header:

```cpp
#include "Teuchos_SerialDenseMatrix.hpp"
```

Creating a double-precision matrix can be done in several ways:
// Create an empty matrix with no dimension
Teuchos::SerialDenseMatrix<int,double> Empty_Matrix;
// Create an empty 3x4 matrix
Teuchos::SerialDenseMatrix<int,double> My_Matrix( 3, 4 );
// Basic copy of My_Matrix
Teuchos::SerialDenseMatrix<int,double> My_Copy1( My_Matrix ),
// (Deep) Copy of principle 3x3 sub-matrix of My_Matrix
My_Copy2( Teuchos::Copy, My_Matrix, 3, 3 ),
// (Shallow) Copy of 2x3 sub-matrix of My_Matrix
My_Copy3( Teuchos::View, My_Matrix, 2, 3, 1, 1 );

The matrix dimensions and strided storage information can be obtained:

int rows = My_Copy3.numRows(); // number of rows
int cols = My_Copy3.numCols(); // number of columns
int stride = My_Copy3.stride(); // storage stride

Matrices can change dimension:

Empty_Matrix.shape( 3, 3 ); // size non-dimensional matrices
My_Matrix.reshape( 3, 3 ); // resize matrices and save values

Filling matrices with numbers can be done in several ways:

My_Matrix.random(); // random numbers
My_Copy1.putScalar( 1.0 ); // every entry is 1.0
My_Copy2(1,1) = 10.0; // individual element access
Empty_Matrix = My_Matrix; // copy My_Matrix to Empty_Matrix

Basic matrix arithmetic can be performed:

Teuchos::SerialDenseMatrix<int,double> My_Prod( 3, 2 );
// Matrix multiplication ( My_Prod = 1.0*My_Matrix*My_Copy^T )
My_Prod.multiply( Teuchos::NO_TRANS, Teuchos::TRANS,
1.0, My_Matrix, My_Copy3, 0.0 );
My_Copy2 += My_Matrix; // Matrix addition
My_Copy2.scale( 0.5 ); // Matrix scaling

The pointer to the array of matrix values can be obtained:

double* My_Array = My_Matrix.values(); // pointer to matrix values
double* My_Column = My_Matrix[2]; // pointer to third column values

The norm of a matrix can be computed:

double norm_one = My_Matrix.normOne(); // one norm
double norm_inf = My_Matrix.normInf(); // infinity norm
double norm_fro = My_Matrix.normFrobenius(); // frobenius norm

Matrices can be compared:

// Check if the matrices are equal in dimension and values
if (Empty_Matrix == My_Matrix) {
    cout<< "The matrices are the same!" <<endl;
}

// Check if the matrices are different in dimension or values
if (My_Copy2 != My_Matrix) {
    cout<< "The matrices are different!" <<endl;
}
Chapter 10. The Teuchos Utility Classes

A matrix can be sent to the output stream:

```
cout << My_Matrix << endl;
```

This section presents examples of all the methods in the `Teuchos::SerialDenseMatrix` class and can be found in `didasko/examples/teuchos/ex1.cpp`. There is also a specialization of this class for serial dense vectors that includes additional creation, accessor, arithmetic, and norm methods (`Teuchos::SerialDenseVector`).

### 10.4 Teuchos::BLAS

The `Teuchos::BLAS` class provides templated wrappers for the native BLAS library. This class has been written to facilitate the interface between C++ codes and BLAS, which are written in Fortran. Unfortunately, the interface between C++ and Fortran function calls is not standard across all computer platforms. The `Teuchos::BLAS` class provides C++ wrappers for BLAS kernels that are specialized during the Teuchos configuration. This insulates the rest of Teuchos and its users from the details of the Fortran to C++ translation.

The `Teuchos::BLAS` class provides C++ wrappers for a substantial subset of the BLAS kernels (Figure 10.1). The native BLAS library implementations of those kernels will be used for the standard scalar types (float, double, complex<float>, complex<double>). However, `Teuchos::BLAS` also has a templated version of each of these kernels. Paired with `Teuchos::ScalarTraits` (Section 10.2), the `Teuchos::BLAS` class can be extended to provide arbitrary precision computations. To use the `Teuchos::BLAS` class, include the header:

```
#include "Teuchos_BLAS.hpp"
```

Creating an instance of the BLAS class for double-precision kernels looks like:

```
Teuchos::BLAS<int, double> blas;
```

This instance provides the access to all the BLAS kernels listed in Figure 10.1:

```
const int n = 10;
double alpha = 2.0;
double x[n];
for ( int i=0; i<n; i++ ) { x[i] = i; }
blas.SCAL(n, alpha, x, 1);
int max_idx = blas.IAMAX(n, x, 1);
cout << "The index of the maximum magnitude entry of x[] is the "
    << max_idx << "-th and x[ " << max_idx-1 << " ] = " << x[max_idx-1]
    << endl;
```

This is a small usage example, but its purpose is to illustrate that any of the supported BLAS kernels is a method of the `Teuchos::BLAS` class. This example can be found in `didasko/examples/teuchos/ex2.cpp`.

### 10.5 Teuchos::LAPACK

The `Teuchos::LAPACK` class provides templated wrappers for the native LAPACK library. This class has been written to facilitate the interface between C++ codes and BLAS, which are written in Fortran. Unfortunately, the interface between C++ and Fortran function calls
<table>
<thead>
<tr>
<th>BLAS Kernel</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ROTG</td>
<td>Computes a Givens plane rotation</td>
</tr>
<tr>
<td>SCAL</td>
<td>Scale a vector by a constant</td>
</tr>
<tr>
<td>COPY</td>
<td>Copy one vector to another</td>
</tr>
<tr>
<td>AXPY</td>
<td>Add one scaled vector to another</td>
</tr>
<tr>
<td>ASUM</td>
<td>Sum the absolute values of the vector entries</td>
</tr>
<tr>
<td>DOT</td>
<td>Compute the dot product of two vectors</td>
</tr>
<tr>
<td>NRM2</td>
<td>Compute the 2-norm of a vector</td>
</tr>
<tr>
<td>IAMAX</td>
<td>Determine the index of the largest magnitude entry of a vector</td>
</tr>
<tr>
<td>GEMV</td>
<td>Add a scaled matrix-vector product to another scaled vector</td>
</tr>
<tr>
<td>TRMV</td>
<td>Replaces a vector with its upper/lower-triangular matrix-vector product</td>
</tr>
<tr>
<td>GER</td>
<td>Updates a matrix with a scaled, rank-one outer product</td>
</tr>
<tr>
<td>GEMM</td>
<td>Add a scaled matrix-matrix product to another scaled matrix</td>
</tr>
<tr>
<td>SYMM</td>
<td>Add a scaled symmetric matrix-matrix product to another scaled matrix</td>
</tr>
<tr>
<td>TRMM</td>
<td>Add a scaled upper/lower-triangular matrix-matrix product to another scaled matrix</td>
</tr>
<tr>
<td>TRSM</td>
<td>Solves an upper/lower-triangular linear system with multiple right-hand sides</td>
</tr>
</tbody>
</table>

**Figure 10.1:** BLAS kernels supported by Teuchos::BLAS

is not standard across all computer platforms. The **Teuchos::LAPACK** class provides C++ wrappers for LAPACK routines that are specialized during the Teuchos configuration. This insulates the rest of Teuchos and its users from the details of the Fortran to C++ translation.

**Teuchos::LAPACK** is a serial interface only, as LAPACK functions are. Users interested in the parallel counterpart of LAPACK, ScaLAPACK, can use the Amesos package; see Chapter 12.

The **Teuchos::LAPACK** class provides C++ wrappers for a substantial subset of the LAPACK routines (Figure 10.2). The native LAPACK library implementations of those kernels will be used for the standard scalar types (float, double, complex<float>, complex<double>). Unlike **Teuchos::BLAS**, the **Teuchos::LAPACK** class does not have a templated version of these routines at this time, so it cannot offer arbitrary precision computations.

To use the **Teuchos::LAPACK** class, include the header:

```cpp
#include "Teuchos_LAPACK.hpp"
```

Creating an instance of the LAPACK class for double-precision routines looks like:

```cpp
Teuchos::LAPACK<int, double> lapack;
```

This instance provides the access to all the LAPACK routines listed in Figure 10.2:

```cpp
Teuchos::SerialDenseMatrix<int,double> My_Matrix(4,4);
Teuchos::SerialDenseVector<int,double> My_Vector(4);
My_Matrix.random();
My_Vector.random();

// Perform an LU factorization of this matrix.
int ipiv[4], info;
char TRANS = 'N';
lapack.GETRF( 4, 4, My_Matrix.values(), My_Matrix.stride(), ipiv, &info );

// Solve the linear system.
lapack.GETRS( TRANS, 4, 1, My_Matrix.values(), My_Matrix.stride(),
               ipiv, My_Vector.values(), My_Vector.stride(), &info );
```
This small example illustrates how easy it is to use the `Teuchos::LAPACK` class. Furthermore, it also exhibits the compatibility of the `Teuchos::SerialDenseMatrix` and `Teuchos::SerialDenseVector` classes with the `Teuchos::LAPACK` class. This example can be found in `didasko/examples/teuchos/ex3.cpp`. 
<table>
<thead>
<tr>
<th>LAPACK Routine</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>POTRF</td>
<td>Computes Cholesky factorization of a real symmetric positive definite (SPD) matrix.</td>
</tr>
<tr>
<td>POTRS</td>
<td>Solves a system of linear equations where the matrix has been factored by POTRF.</td>
</tr>
<tr>
<td>POTRI</td>
<td>Computes the inverse of a real SPD matrix after its been factored by POTRF.</td>
</tr>
<tr>
<td>POTCON</td>
<td>Estimates the reciprocal of the condition number (1-norm) of a real SPD matrix after its been factored by POTRF.</td>
</tr>
<tr>
<td>POSV</td>
<td>Computes the solution to a real system of linear equations where the matrix is SPD.</td>
</tr>
<tr>
<td>POEQU</td>
<td>Computes row and column scaling or equilibrating a SPD matrix and reduce its condition number.</td>
</tr>
<tr>
<td>PORFS</td>
<td>Improves the computed solution to a system of linear equations where the matrix is SPD.</td>
</tr>
<tr>
<td>POSVX</td>
<td>Expert SPD driver: Uses POTRF/POTRS to compute the solution to a real system of linear equations where the matrix is SPD. The system can be equilibrated (POEQU) or iteratively refined (PORFS) also.</td>
</tr>
<tr>
<td>GELS</td>
<td>Solves and over/underdetermined real linear system.</td>
</tr>
<tr>
<td>GETRF</td>
<td>Computes an LU factorization of a general matrix using partial pivoting.</td>
</tr>
<tr>
<td>GETRS</td>
<td>Solves a system of linear equations using the LU factorization computed by GETRF.</td>
</tr>
<tr>
<td>GETRI</td>
<td>Computes the inverse of a matrix using the LU factorization computed by GETRF.</td>
</tr>
<tr>
<td>GECON</td>
<td>Estimates the reciprocal of the condition number of a general matrix in either the 1-norm or ∞-norm using the LU factorization computed by GETRF.</td>
</tr>
<tr>
<td>GESV</td>
<td>Computes the solution of a linear system of equations.</td>
</tr>
<tr>
<td>GEEQU</td>
<td>Computes row and column scaling for equilibrating a linear system, reducing its condition number.</td>
</tr>
<tr>
<td>GERFS</td>
<td>Improves the computed solution to a system of linear equations and provides error bounds and backward error estimates for the solution [Use after GETRF/GETRS].</td>
</tr>
<tr>
<td>GESVX</td>
<td>Expert driver: Uses GETRF/GETRS to compute the solution to a real system of linear equations, returning error bounds on the solution and a condition estimate.</td>
</tr>
<tr>
<td>GEHRD</td>
<td>Reduces a real general matrix to upper Hessenberg form by orthogonal similarity transformations.</td>
</tr>
<tr>
<td>HSEQR</td>
<td>Compute the eigenvalues of a real upper Hessenberg matrix and, optionally, the Schur decomposition.</td>
</tr>
<tr>
<td>GEES</td>
<td>Computes the real Schur form, eigenvalues, and Schur vectors of a real nonsymmetric matrix.</td>
</tr>
<tr>
<td>GEEV</td>
<td>Computes the eigenvalues and, optionally, the left and/or right eigenvectors of a real nonsymmetric matrix.</td>
</tr>
<tr>
<td>ORGHR</td>
<td>Generates a real orthogonal matrix which is the product of the elementary reflectors computed by GEHRD.</td>
</tr>
<tr>
<td>ORMHR</td>
<td>Overwrites the general real matrix with the product of itself and the elementary reflectors computed by GEHRD.</td>
</tr>
<tr>
<td>TREVC</td>
<td>Computes some or all of the right and/or left eigenvectors of a real upper quasi-triangular matrix.</td>
</tr>
<tr>
<td>TREXC</td>
<td>Reorders the real Schur factorization of a real matrix via orthogonal similarity transformations.</td>
</tr>
<tr>
<td>LARND</td>
<td>Returns a random number from a uniform or normal distribution.</td>
</tr>
<tr>
<td>LARNV</td>
<td>Returns a vector of random numbers from a chosen distribution.</td>
</tr>
<tr>
<td>LAMCH</td>
<td>Determines machine parameters for floating point characteristics.</td>
</tr>
<tr>
<td>LAPY2</td>
<td>Computes sqrt (x^2 + y^2) safely, to avoid overflow.</td>
</tr>
</tbody>
</table>

Figure 10.2: LAPACK routines supported by Teuchos::LAPACK
10.6 Teuchos::ParameterList

The Teuchos::ParameterList class is a C++ container of \(<\text{key}, \text{value}\)> pairs, where the \text{key}\ is a character string (std::string) and the \text{value}\ can be almost any type of C++ object. The ability to hold almost any type of C++ object as a \text{value}\ in the same list makes this class very useful for storing parameters. This parameter list can then be passed to an object, like an iterative linear solver, which will use the information to define its behavior.

The Teuchos::ParameterList is currently being used by several Trilinos packages. For instance, all Amesos objects (see Chapter 12) and the smoothed aggregation preconditioning object ML_Epetra::MultiLevelPreconditioner (see Section 11.4) are configured through a Teuchos::ParameterList.

Remark 18. The parameter list stores a copy of the input object if it passed by reference. If the list is passed a pointer to an object, only the pointer is copied and not the object that it points to.

To use the Teuchos::ParameterList class, include the header:

```
#include "Teuchos_ParameterList.hpp"
```

Creating an empty parameter list looks like:

```
Teuchos::ParameterList My_List;
```

Setting parameters in this list can be easily done:

```
My_List.set("Max Iters", 1550);
My_List.set("Tolerance", 1e-10);
My_List.set("Solver", "GMRES");
```

The templated “set” method should cast the input \text{value}\ to the correct data type. However, in the case where the compiler is not casting the input \text{value}\ to the expected data type, an explicit cast can be used with the “set” method:

```
My_List.set("Tolerance", (float)(1e-10));
```

A hierarchy of parameter lists can be constructed using Teuchos::ParameterList. This means another parameter list is a valid \text{value}\ in any parameter list. To create a sublist in a parameter list and obtain a reference to it:

```
Teuchos::ParameterList& Prec_List = My_List.sublist("Preconditioner");
```

Now this parameter list can be filled with values:

```
Prec_List.set("Type", "ILU");
Prec_List.set("Drop Tolerance", 1e-3);
```

The parameter list can be queried about the existence of a parameter, sublist, or type:

```
// Has a solver been chosen?
bool solver_defined = My_List.isParameter("Solver");
// Has a preconditioner been chosen?
bool prec_defined = My_List.isSublist("Preconditioner");
// Has a tolerance been chosen and is it a double-precision number?
bool tol_double = My_List.template isType<double>("Tolerance");
// Has a drop tolerance been chosen and is it a double-precision number?
bool dtol_double = Teuchos::isParameterType<double>(Prec_List, "Drop Tolerance");
```
The last two methods for checking the parameter type are equivalent. There is some question as to whether the syntax of the first type-checking method (isType) is acceptable to older compilers. Thus, the second type-checking method (isParameterType) is offered as a portable alternative.

Parameters can be retrieved from the parameter list in quite a few ways:

// Get method that creates and sets the parameter if it doesn't exist.
int its = My_List.get("Max Iters", 1200);

// Get method that retrieves a parameter of a particular type.
float tol = My_List.template get<float>("Tolerance");

In the above example, the first “get” method is a safe way of obtaining a parameter when its existence is indefinite but required. The second “get” method should be used when the existence of the parameter is definite. This method will throw an exception if the parameter doesn’t exist. The safest way to use the second “get” method is in a try/catch block:

try {
    tol = My_List.template get<float>("Tolerance");
} catch (exception& e) {
    tol = 1e-6;
}

The second “get” method uses a syntax that may not be acceptable to older compilers. Optionally, there is another portable templated “get” function that can be used in the place of the second “get” method:

try {
    tol = Teuchos::getParameter<float>(My_List, "Tolerance");
} catch (exception& e) {
    tol = 1e-6;
}

A parameter list can be sent to the output stream:

cout<< My_List << endl;

For this parameter list, the output would look like:

Max Iters = 1550
Preconditioner ->
    Drop Tolerance = 0.001 [unused]
    Type = ILU [unused]
Solver = GMRES [unused]
Tolerance = 1e-10

It is important to note that misspelled parameters (with additional space characters, capitalizations, etc.) may be ignored. Therefore, it is important to be aware that a given parameter has not been used. Unused parameters can be printed with method:

My_List.unused( cout );

This section presents examples of all the methods in the Teuchos::ParameterList class and can be found in didasko/examples/teuchos/ex4.cpp.
10.7 Teuchos::RCP

The Teuchos::RCP class is a templated class for implementing automatic garbage collection in C++ using smart, reference-counted pointers. Using this class allows one client to dynamically create an object and pass the object around to other clients without fear of memory leaks. No client is required to explicitly call delete because object will be deleted when all the clients remove their references to it. The type of garbage collection performed by Teuchos::RCP is similar to those found in Perl and Java.

To use the Teuchos::RCP class, include the header:

```cpp
#include "Teuchos_RCP.hpp"
```

The data type used with Teuchos::RCP should not be a built-in data type (like int or double), this creates unnecessary overhead. Instead, it should be used to manage dynamic objects and data members that need to be shared with many clients. This means that the data type will most likely be a C++ class. Consider the class hierarchy:

```cpp
class A {
  public:
    A() {}
    virtual ~A(){}
    virtual void f(){}
  
};
class B1 : virtual public A {}; 
class B2 : virtual public A {}; 
class C : public B1, public B2 {}; 
```

Creating a reference-counted pointer to a dynamically allocated object (of type A) can be done several ways:

```cpp
// Create a reference-counted NULL pointer of type A.
RCP<A> a_null_ptr;
// Create a reference-counted pointer of non-const type A.
RCP<A> a_ptr = rcp(new A);
// Create a reference-counted pointer of const type A.
RCP<const A> ca_ptr = rcp(new A);
// Create a const reference-counted pointer of non-const type A.
const RCP<A> a_cptr = rcp(new A);
// Create a const reference-counted pointer of const type A.
const RCP<const A> ca_cptr = rcp(new A);
```

The Teuchos::RCP class can also perform implicit conversions between a derived class (B1) and its base class (A):

```cpp
RCP<B1> b1_ptr = rcp(new B1);
RCP<A> a_ptr1 = b1_ptr;
```

Other non-implicit type conversions like static, dynamic, or const casts can be taken care of by non-member template functions:

```cpp
RCP<const C> c_ptr = rcp(new C);
// Implicit cast from C to B2.
RCP<const B2> b2_ptr = c_ptr;
```
Using a reference-counted pointer is very similar to using a raw C++ pointer. Some of the operations that are common to both are:

```cpp
RCP<A> a_ptr2 = rcp(new A), // Initialize reference-counted pointers.
      a_ptr3 = rcp(new A); // "
      a_ptr2 = rcp(a_ptr3); // Assign from a raw pointer (only do this once!)
      a_ptr2 = a_ptr2; // Assign one smart pointer to another.
      ra_ptr2->f(); // Access a member of A using ->
      *ra_ptr2 = *a_ptr3; // Dereference the objects and assign.
```

However, a reference-counted pointer cannot be used everywhere a raw C++ pointer can. For instance, these statements will not compile:

```cpp
// Pointer arithmetic ++, --, +, - etc. not defined!
ra_ptr2++; // error
// Comparison operators ==, !=, <=, >= etc. not defined!
ra_ptr2 == ra_ptr1; // error
```

Because the two are not equivalent, the `Teuchos::RCP` class provides a way of getting the raw C++ pointer held by any `RCP<A>` object:

```cpp
A* true_ptr = a_ptr1.get();
```

These are just some of the basic features found in the `Teuchos::RCP` class. A more extensive tutorial of this powerful tool is “in the works” and will be made available to Teuchos users as soon as it is finished. The examples presented in this section can be found in `didasko/examples/teuchos/ex5.cpp`.

## 10.8 Teuchos::TimeMonitor

The `Teuchos::TimeMonitor` class is a container that manages a group of timers. In this way, it can be used to keep track of timings for various phases of the code. Internally, this class holds an array of `Teuchos::Time` objects. The `Teuchos::Time` class defines a basic wall-clock timer that can `start()`, `stop()`, and return the `totalElapsedTime()`.

To use the `Teuchos::TimeMonitor` class, include the header:

```cpp
#include "Teuchos_TimeMonitor.hpp"
```

To create a timer for the TimeMonitor to manage, call:

```cpp
RCP<Time> FactTime = TimeMonitor::getNewTimer("Factorial Time");
```
The `getNewTimer` method creates a new reference-counted `Teuchos::Time` object and adds it to the internal array. To avoid passing this timer into each method that needs timing, consider putting it in the global scope (declare it outside of `main(argc, argv[])`). Now, when we want to time a part of the code, the appropriate timer should be used to construct a local `TimeMonitor`:

```cpp
Teuchos::TimeMonitor LocalTimer(*FactTime);
```

This timer will be started during the construction of `LocalTimer` and stopped when the destructor is called on `LocalTimer`.

To obtain a summary from all the timers in the global `TimeMonitor`, use:

```cpp
TimeMonitor::summarize();
```

Information from each timer can also be obtained using the methods from `Teuchos::Time`. This section presents examples of all the methods in the `Teuchos::TimeMonitor` class and can be found in `didasko/examples/teuchos/ex6.cpp`.

### 10.9 Teuchos::CommandLineProcessor

`Teuchos::CommandLineProcessor` is a class that helps to parse command line input arguments and set runtime options. Additionally, a `CommandLineProcessor` object can provide the user with a list of acceptable command line arguments, and their default values.

To use the `Teuchos::CommandLineProcessor` class, include the header:

```cpp
#include "Teuchos_CommandLineProcessor.hpp"
```

Creating an empty command line processor looks like:

```cpp
Teuchos::CommandLineProcessor My_CLP;
```

To set an option, it must be given a name and default value. Additionally, each option can be given a help string. Although it is not necessary, a help string aids a users comprehension of the acceptable command line arguments. Some examples of setting command line options are:

```cpp
// Set an integer command line option.
int NumIters = 1550;
My_CLP.setOption("iterations", &NumIters, "Number of iterations");
// Set a double-precision command line option.
double Tolerance = 1e-10;
My_CLP.setOption("tolerance", &Tolerance, "Tolerance");
// Set a string command line option.
string Solver = "GMRES";
My_CLP.setOption("solver", &Solver, "Linear solver");
// Set a boolean command line option.
bool Precondition;
My_CLP.setOption("precondition", "no-precondition", &Precondition, "Preconditioning flag");
```

There are also two methods that control the strictness of the command line processor. For a command line processor to be sensitive to any bad command line option that it does not recognize, use:

```cpp
My_CLP.recogniseAllOptions(false);
```
Then, if the parser finds a command line option it doesn’t recognize, it will throw an exception. To prevent a command line processor from throwing an exception when it encounters an unrecognized option or when the help string is printed, use:

```java
My_CLP.throwExceptions(false);
```

Finally, to parse the command line, `argc` and `argv` are passed to the `parse` method:

```java
My_CLP.parse( argc, argv );
```

The --help output for this command line processor is:

```
Usage: ./ex7.exe [options]
options:
  --help        Prints this help message
  --pause-for-debugging  Pauses for user input to allow attaching a debugger
  --iterations int  Number of iterations
                   (default: --iterations=1550)
  --tolerance double  Tolerance
                       (default: --tolerance=1e-10)
  --solver string  Linear solver
                   (default: --solver="GMRES")
  --precondition bool  Preconditioning flag
                       (default: --precondition)
```

This section presents examples of all the methods in the `Teuchos::CommandLineProcessor` class and can be found in `didasko/examples/teuchos/ex7.cpp`. 
Multilevel Preconditioners with ML

Marzio Sala, Michael Heroux, David Day

The ML package defines a class of preconditioners based on multilevel methods [BHM00, TT00]. While theoretically ML preconditioners apply to any linear system, the range of applicability of the methods is limited at this time, primarily to certain linear elliptic partial differential equations discretized with linear shape functions. The ML package provides multilevel solvers and preconditioners based on geometric and algebraic coarsening schemes. Please contact the developers for information on the status of special purpose methods, such as those for the incompressible Navier-Stokes equations and Maxwell’s equations.

This Chapter will present:

• A multilevel preconditioning framework (in Section 11.1);
• How to use ml objects as AztecOO preconditioners (in Section 11.2);
• The ML_Epetra::MultiLevelOperator class (in Section 11.3);
• How to define black-box preconditioners using the ML_Epetra::MultiLevelPreconditioner Class (in Section 11.4);
• How to implement two level domain decomposition methods with aggregation based coarse matrix (in Section 11.5).

11.1 A Multilevel Preconditioning Framework

For certain combinations of iterative methods and linear systems, the error at each iteration projected onto the eigenfunctions has components that decay at a rate proportional to the corresponding eigenvalue (or frequency). Multilevel methods exploit this property [BHM00] by projecting the linear system onto a hierarchy of increasingly coarsened “meshes” so that each error component rapidly decays on at least one coarse “mesh.” The linear system on the coarsest “mesh”, called the coarse grid problem, is solved exactly. The iterative method is called the smoother, as a reflection of its diminished role as a way to damp out the high frequency error. The grid transfer (or interpolation) operators are called restriction (R) and prolongation (P) operators.

Multilevel methods are characterized by the sequence of coarse spaces, the definition of the operator each coarse space, the specification of the smoother, and the restriction and
prolongation operators. Geometric multigrid (GMG) methods are multilevel methods that require the user to specify the underlying grid, and in most cases a hierarchy of (not necessarily nested) coarsens grids. Both the automatic generation of a grid-hierarchy for GMG and the specification of the ML, designed for unstructured problems, are beyond the scope of the tutorial.

Algebraic multigrid (AMG) (see [BHM00, Section 8]) method development has been motivated by the demand for multilevel methods that are easier to use. In AMG, both the matrix hierarchy and the prolongation operators are constructed just from the stiffness matrix. Recall that to use Aztec00 or IFPACK, a user must supply a linear system, and select a preconditioning strategy. In AMG, the only additional information required from the user is to specify a coarsening strategy.

Readers that are unfamiliar with multigrid methods are strongly advised to review [BHM00] before using ML.

A multilevel method for (7.1) depends on the number of coarsen grid levels, and operators for each level. Levels are numbered from the coarsest level, 0, to the finest. The pre- and post-smoothers are denoted $S_k^{(1)}$ and $S_k^{(2)}$ respectively. $R_{k-1,k}$ is the restriction operator from level $k$ to $k-1$, and $P_{k,k-1}$ is a prolongator from $k-1$ to $k$. In AMG, the operators on the coarse meshes $A_k$ are defined by

$$A_{k-1} = R_{k-1,k} A_k P_{k,k-1}. $$

The AMG coarse grid operators may be less sparse than the corresponding GMG coarse grid operators, defined by assembling the coefficient matrix on the coarse grid. These pieces combine into a multigrid solver.

**Recursive Definition of the V-cycle Scheme**

\[
\text{MGM}(x, b, \text{Number of Levels})
\]

\[
\text{if } k > 0
\]

\[
x = S_k^{(1)}(x, b)
\]

\[
d = R_{k-1,k}(b - A_k x)
\]

\[
v = 0
\]

\[
\text{MGM}(v, d, k-1)
\]

\[
x = x + P_{k,k-1} v
\]

\[
x = S_k^{(2)}(x, b)
\]

\[
\text{else}
\]

\[
x = A_k^{-1} b
\]

**Remark 19.** The tutorial only discussed AMG methods. The interested reader is referred to [TH04] for information on GMG methods in ML.

## 11.2 ML Objects as AztecOO Preconditioners

ML may be used as a “black-box” multilevel preconditioner, using aggregation procedures to define the multilevel hierarchy. In order to use ML as a preconditioner, we need to define an AztecOO Solver, as outlined in Section 7. ML requires the user to define a structure that stores internal data. The convention in ML is to call the structure `ml_handle`. Next define
the maximum number of levels, the amount of diagnostic information written to the screen, which varies from none (0) to extremely verbose (10), and create the ML handle structure:

```c
ML *ml_handle;
int N_levels = 10;
ML_Set_PrintLevel(3);
ML_Create(&ml_handle, N_levels);
```

Next construct an ML preconditioner for an Epetra matrix. Additionally, ML requires a structure that stores information about the aggregates at each level called ML_Aggregate:

```c
EpetraMatrix2MLMatrix(ml_handle, 0, &A);
ML_Aggregate *agg_object;
ML_Aggregate_Create(&agg_object);
```

The multilevel hierarchy is constructed with the instruction

```c
N_levels = ML_Gen_MGHierarchy_UsingAggregation(ml_handle,
	0,
	ML_INCREASING,
	agg_object);
```

Here, 0 is the index of the finest level, and the index of coarser levels will be obtained by incrementing this value. Please see [SHT04] for more information on the input parameters.

Next define the smoother, such as symmetric Gauss-Seidel, and initialize the solver:

```c
ML_Gen_Smoother_SymGaussSeidel(ml_handle, ML_ALL_LEVELS,
	ML_BOTH, 1, ML_DEFAULT);
ML_Gen_Solver (ml_handle, ML_MGV, 0, N_levels-1);
```

Finally, use this ML hierarchy to create an Epetra_Operator, set the preconditioning operator of our AztecOO solver, and then call Iterate() as usual:

```c
ML_Epetra::MultiLevelOperator MLop(ml_handle,comm,map,map);
solver.SetPrecOperator(&MLop);
solver.Iterate(Niters, 1e-12);
```

The entire code is reported in didasko/examples/ml/ex1.cpp. The output is reported below.

```bash
[msala:ml]> mpirun -np 2 ./ex1.exe
************************************************************
* ML Aggregation information *
==============================================================
ML_Aggregate : ordering = natural.
ML_Aggregate : min nodes/aggr = 2
ML_Aggregate : max neigh selected = 0
ML_Aggregate : attach scheme = MAXLINK
ML_Aggregate : coarsen scheme = UNCOUPLED
ML_Aggregate : strong threshold = 0.000000e+00
ML_Aggregate : P damping factor = 1.333333e+00
```
ML_Aggregate : number of PDEs = 1
ML_Aggregate : number of null vec = 1
ML_Aggregate : smoother drop tol = 0.000000e+00
ML_Aggregate : max coarse size = 1
ML_Aggregate : max no. of levels = 10

*****************************************************************
ML_Gen_MGHierarchy : applying coarsening
ML_Aggregate_Coarsen begins
ML_Aggregate_CoarsenUncoupled : current level = 0
Aggregation(UVB) : Total nonzeros = 128 (Nrows=30)
Aggregation(UV) : Phase 0 - no. of bdry pts = 0
Aggregation(UV) : Phase 1 - nodes aggregated = 28 (30)
Aggregation(UV) : Phase 2 - total aggregates = 8
Aggregation(UV) : Phase 2 - boundary nodes = 0
Aggregation(UV) : Phase 3 - leftovers = 0 and singletons = 0
Aggregation time = 1.854551e-03
Gen_Prolongator : max eigen = 1.883496e+00
ML_Gen_MGHierarchy : applying coarsening
ML_Gen_MGHierarchy : Gen_RAP
RAP time for level 0 = 5.319577e-04
ML_Gen_MGHierarchy : Gen_RAP done
ML_Gen_MGHierarchy : applying coarsening
ML_Aggregate_Coarsen begins
ML_Aggregate_CoarsenUncoupled : current level = 1
Aggregation(UVB) : Total nonzeros = 46 (Nrows=8)
Aggregation(UV) : Phase 0 - no. of bdry pts = 0
Aggregation(UV) : Phase 1 - nodes aggregated = 6 (8)
Aggregation(UV) : Phase 2 - total aggregates = 2
Aggregation(UV) : Phase 3 - leftovers = 0 and singletons = 0
Aggregation time = 1.679042e-03
Gen_Prolongator : max eigen = 1.246751e+00
ML_Gen_MGHierarchy : applying coarsening
ML_Gen_MGHierarchy : Gen_RAP
RAP time for level 1 = 4.489557e-04
ML_Gen_MGHierarchy : Gen_RAP done
ML_Gen_MGHierarchy : applying coarsening
ML_Aggregate_Coarsen begins
Aggregation total setup time = 8.903003e-02 seconds
Smoothed Aggregation : operator complexity = 1.390625e+00.

************************************************************
***** Preconditioned CG solution
***** Epetra ML Operator
***** No scaling
************************************************************

iter:  0    residual = 1.000000e+00
iter:  1    residual = 1.289136e-01
iter:  2    residual = 4.710371e-03
iter:  3    residual = 7.119470e-05
iter:  4    residual = 1.386302e-06
iter:  5    residual = 2.477133e-08
iter:  6    residual = 6.141025e-10
iter:  7    residual = 6.222216e-12
iter:  8    residual = 1.277534e-13

Solution time: 0.005845 (sec.)
total iterations: 8
Residual = 6.99704e-13

11.3 The ML_Epetra::MultiLevelOperator Class

As with other Trilinos packages, ML can be compiled and run independently from Epetra. It accepts input matrix in formats different from the Epetra_RowMatrix or Epetra_Operator. However, as part of the Trilinos project, ML can be used to define a preconditioner operator for Epetra_LinearProblem objects (see for instance [Her02]). This means that, in a C++ framework, ML can be defined as an Epetra_Operator object, applied to an Epetra_MultiVector object, and used as a preconditioner for AztecOO. This can be done in two ways:

- By defining an ML_Epetra::MultiLevelOperator object, derived from the Epetra_Operator class. The constructor of this object requires already filled ML_Struct and ML_Aggregate structures. ML must have been configure with the option --enable-epetra.

- By defining an ML_Epetra::MultiLevelPreconditioner object, derived from the Epetra_RowMatrix class. Basically, the constructor of this object demands for an Epetra_RowMatrix pointer and a Teuchos parameter list, that contains all the user’s defined parameters. ML must have been configure with options --enable-epetra --enable-teuchos.

The first approach, described in Section 11.3, is more general, and can be applied to geometric and algebraic multilevel preconditioner, but it requires a deeper knowledge of the ML package. This is because the user has to explicitly construct the ML hierarchy, define
Chapter 11. Multilevel Preconditioners with ML

the aggregation strategies, the smoothers, and the coarse grid solver. The second approach, presented in Section 11.4, instead, although limited to algebraic multilevel preconditioners, allows the use of ML as a black-box preconditioner. This class automatically constructs all the components of the preconditioner, following the parameters specified in a Teuchos parameters’ list.

Next we walk through how to write some code to construct an ML preconditioner for an Epetra_RowMatrix \( A \). The ML_Epetra::MultiLevelOperator class is defined in the header file ml_MultiLevelOperator.h. Users must include it, and along with some subset of ml_config.h, AztecOO.h, Epetra_Operator.h, Epetra_MultiVector.h and Epetra_LinearProblem.h. Check the Epetra and AztecOO documentation for more information on this topic.

The next steps proceed exactly as in §11.2:

```cpp
ML *ml_handle;
int N_levels = 10;
ML_Set_PrintLevel(3);
ML_Create(&ml_handle, N_levels);
EpetraMatrix2MLMatrix(ml_handle, 0, &A);
ML_Aggregate *agg_object;
ML_Aggregate_Create(&agg_object);
N_levels = ML_Gen_MGHierarchy_UsingAggregation(ml_handle, 0,
ML_INCREASING,
agg_object);
ML_Gen_Smoother_SymGaussSeidel(ml_handle, ML_ALL_LEVELS,
ML_BOTH, 1, ML_DEFAULT);
ML_Gen_Solver (ml_handle, ML_MGV, 0, N_levels-1);
ML_Epetra::MultiLevelOperator MLop(ml_handle, comm, map, map);
```

At this point, our example diverges from §11.2. Instead of using ML to solve the linear system \( Ax = b \), where \( x \) and \( b \) are Epetra_MultiVector, use the ML operator as the precondition for an AztecOO linear system, and solve it:

```cpp
Epetra_LinearProblem Problem(A,&x,&b);
AztecOO Solver(Problem);
Solver.SetPrecOperator(&MLop);
Solver.SetAztecOption( AZ_solver, AZ_gmres );
Solver.Iterate(Niters, 1e-12);
```

### 11.4 The ML_Epetra::MultiLevelPreconditioner Class

An alternative to the ML_Epetra::MultiLevelOperator (that is also in the namespace ML_Epetra) is the MultiLevelPreconditioner class. Replace the header file ml_MultiLevelOperator.h discussed in section 11.3 by ml_MultiLevelPreconditioner.h. Table 11.1 reports the aggregation schemes currently available in ML.

A very simple fragment of code using this class is reported below. The reader may refer to file $ML_HOME/examples/ml_example_MultiLevelPreconditioner.cpp for a more complex example. To run example, first configure ML --enable-triutils.
11.4. The ML_Epetra::MultiLevelPreconditioner Class

Uncoupled

For a 1, 2, or 3 dimensional structured Cartesian grid with a 3, 9 or 27 point stencil respectively, construct aggregates of optimal size such that each aggregate resides on one processor.

MIS

Maximal independent set based coarsening with aggregates allowed to reside on multiple processes. The scheme minimizes the number of iterations, but the cost per iteration is high.

METIS

Use a serial graph partitioner to create aggregates residing on one processor. The number of nodes in each aggregate is specified with the option `aggregation: nodes per aggregate`. ML must be configured with `--with-ml-metis`.

ParMETIS

Use a parallel graph partitioner to create aggregates that may reside on multiple processors. ML must be configured with `--with-ml-parmetis3x`. The number of aggregates is specified by option `aggregation: global number`.

<table>
<thead>
<tr>
<th>Coarsening Scheme</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Uncoupled</td>
<td>For a 1, 2, or 3 dimensional structured Cartesian grid with a 3, 9 or 27 point stencil respectively, construct aggregates of optimal size such that each aggregate resides on one processor.</td>
</tr>
<tr>
<td>MIS</td>
<td>Maximal independent set based coarsening with aggregates allowed to reside on multiple processes. The scheme minimizes the number of iterations, but the cost per iteration is high.</td>
</tr>
<tr>
<td>METIS</td>
<td>Use a serial graph partitioner to create aggregates residing on one processor. The number of nodes in each aggregate is specified with the option <code>aggregation: nodes per aggregate</code>. ML must be configured with <code>--with-ml-metis</code>.</td>
</tr>
<tr>
<td>ParMETIS</td>
<td>Use a parallel graph partitioner to create aggregates that may reside on multiple processors. ML must be configured with <code>--with-ml-parmetis3x</code>. The number of aggregates is specified by option <code>aggregation: global number</code>.</td>
</tr>
</tbody>
</table>

Table 11.1: ML_Epetra::MultiLevelPreconditioner Coarsening Schemes

Jacobi

Point-Jacobi. Damping factor is specified using `smoother: damping factor`, and the number of sweeps with `smoother: sweeps`.

Gauss-Seidel

Point Gauss-Seidel.

Aztec

Use AztecOO’s built-in preconditioning functions as smoothers. Or, use approximate solutions with AztecOO as smoothers. The AztecOO vectors `options` and `params` can be set using `smoother: Aztec options` and `smoother: Aztec params`.

<table>
<thead>
<tr>
<th>Smoothers</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jacobi</td>
<td>Point-Jacobi. Damping factor is specified using <code>smoother: damping factor</code>, and the number of sweeps with <code>smoother: sweeps</code>.</td>
</tr>
<tr>
<td>Gauss-Seidel</td>
<td>Point Gauss-Seidel.</td>
</tr>
<tr>
<td>Aztec</td>
<td>Use AztecOO’s built-in preconditioning functions as smoothers. Or, use approximate solutions with AztecOO as smoothers. The AztecOO vectors <code>options</code> and <code>params</code> can be set using <code>smoother: Aztec options</code> and <code>smoother: Aztec params</code>.</td>
</tr>
</tbody>
</table>

Table 11.2: ML_Epetra::MultiLevelPreconditioner Smoothers

Jacobi

Use Jacobi as a solver.

Gauss-Seidel

Use Gauss-Seidel as a solver.

Amesos_KLU

Use Amesos’s KLU sequential solver.

Amesos_UMFPACK

Use UMFPACK.

Amesos_Superludist

Use SuperLU_DIST.

Amesos_MUMPS

Use MUMPS.

Table 11.3: ML_Epetra::MultiLevelPreconditioner Coarsest Grid Exact Solvers To use Amesos, ML must be configured with `--enable-amesos` and Amesos also be configured as needed.
#include "ml_include.h"
#include "ml_MultiLevelPreconditioner.h"
#include "Teuchos_ParameterList.hpp"

... 

// A is an Epetra_RowMatrix derived class object
// solver is an AztecOO object

Teuchos::ParameterList MList;

// default values for smoothed aggregation
ML_Epetra::SetDefaults("SA", MList);
MList.set("max levels", 6);
MList.set("increasing or decreasing", "decreasing");
MList.set("aggregation: type", "MIS");
MList.set("coarse: type", "Amesos_KLU");

ML_Epetra::MultiLevelPreconditioner * MLPrec =
    new ML_Epetra::MultiLevelPreconditioner(A, MList, true);

solver.SetPrecOperator(MLPrec);
solver.SetAztecOption(AZ_solver, AZ_gmres);
solver.Iterate(Niters, 1e-12);

...

delete MLPrec;

The general procedure is as follows. First, the user defines a Teuchos parameters' list. Second input parameters are set via method set(ParameterName, ParameterValue), where ParameterName is a string defining the parameter, and ParameterValue is the specified parameter, that can be any C++ object or pointer. This list is passed to the constructor, together with a pointer to the matrix, and a boolean flag. If this flag is set to false, the constructor will not compute the multilevel hierarchy until when MLPrec->ComputePreconditioner() is called. The hierarchy can be destroyed using MLPrec->Destroy(). For instance, the user may define a code like:

// A is still not filled with numerical values
ML_Epetra::MultiLevelPreconditioner * MLPrec =
    new ML_Epetra::MultiLevelPreconditioner(A, MList, false);

// compute the elements of A
...

// now compute the preconditioner
MLPrec->ComputePreconditioner();

// solve the linear system, and refill A
11.5 Two-Level Domain Decomposition Preconditioners with ML

In this fragment of code, the user defines the ML preconditioner, but does not create the preconditioner in the construction phase. This is of particular interest, for example, when ML is used in conjunction with nonlinear solvers (like NOX [KP04]).

We point out that the input parameter list is copied in the construction phase, hence later changes to MLList will not affect the preconditioner. Should one need to modify parameters in the MLPrec's internally stored parameter list, proceed as follows:

 ParameterList & List = MLprec->GetList();

and then directly modify List.

All ML options can have a common prefix, specified by the user in the construction phase. For example, suppose that we require ML: to be the prefix. The constructor will be

 MLList.set("ML: aggregation: type", "METIS");
 ML_Epetra::MultiLevelPreconditioner * MLprec =
 new ML_Epetra::MultiLevelPreconditioner(*A,
 MLList, 
 true, 
 Prefix);

where Prefix is a char array containing ML:.

Note that spaces are important: Do not include leading or trailing spaces, and separate words by just one space! Misspelled parameters will not be used, and can be detected calling method PrintUnused() after the construction of the multilevel hierarchy.

For a detailed list of all the parameters, we refer to the ML user's guide. Here, we report the most used parameters in Tables 11.1, 11.2 and 11.3.

11.5 Two-Level Domain Decomposition Preconditioners with ML

The idea of two level domain decomposition based on aggregation is to use a graph partitioner to partition the local or global graph into subgraphs, and then treat each subgraph as a large aggregate.

The example contained herein uses the graph decomposition library METIS to create the coarse-level matrix. If you don't have METIS, or just do not want to re-configure ML, you may run the example you will be limited to use only one aggregate per process. There are three changes to the Trilinos configuration. One flag tells the package (ML) to look for an external library, and the other two flag tells the compiler where to find the include directories and external library. Configure ML with the flags --with-mlmetis and --with-incdirs and --with-ldflags set to the locations of the METIS include files and library. Please type configure --help in the ML subdirectory for more information.

Two-level domain decomposition methods are effective for the iterative solution of many different kinds of linear systems. For some classes of problems, a very convenient way to define the coarse grid operator is to use an aggregation procedure. This is very close to what presented in Section 11.2. The main difference is that only two level methods are considered,
and that the coarse grid remains of (relatively) small size. The idea is to define a small number of aggregates on each process, using a graph decomposition algorithm (as implemented in the library METIS, for instance)\(^1\). This can be done as follows.

The linear system matrix \(A\), here coded as an Epetra_CrsMatrix\(^2\), corresponds to the discretization of a 2D Laplacian on a Cartesian grid. \(x\) and \(b\) are the solution vector and the right-hand side, respectively.

The AztecOO linear problem is defined as

\[
\text{Epetra\_LinearProblem problem(&A, &x, &b);}
\]

\[
\text{AztecOO solver(problem);}
\]

At this point, we can create the Teuchos parameters’ list, with the following parameters:

\[
\text{ParameterList MLList;}
\]

\[
\text{ML\_Epetra::SetDefaults("DD",MLList);}\]

\[
\text{MLList.set("max levels",2);}\]

\[
\text{MLList.set("increasing or decreasing","increasing");}
\]

\[
\text{MLList.set("aggregation: type", "METIS");}
\]

\[
\text{MLList.set("aggregation: nodes per aggregate", 16);}\]

\[
\text{MLList.set("smoother: pre or post", "both");}
\]

\[
\text{MLList.set("coarse: type","Amesos\_KLU");}
\]

\[
\text{MLList.set("smoother: type", "Aztec");}
\]

The last option tells ML to use the Aztec preconditioning function as a smoother. Aztec requires an integer vector \texttt{options} and a double vector \texttt{params}. Those can be defined as follows:

\[
\text{Teuchos::RCP<vector<int>> options = rcp(new vector<int>(AZ\_OPTIONS\_SIZE));}
\]

\[
\text{Teuchos::RCP<vector<double>> params = rcp(new vector<double>(AZ\_PARAMS\_SIZE));}
\]

\[
\text{AZ\_defaults(options,params);}\]

\[
\text{(*options)[AZ\_precond] = AZ\_dom\_decomp;}\]

\[
\text{(*options)[AZ\_subdomain\_solve] = AZ\_icc;}\]

\[
\text{MLList.set("smoother: Aztec options", options);}\]

\[
\text{MLList.set("smoother: Aztec params", params);}\]

The last two commands set the Teuchos reference-counted pointers, \texttt{options} and \texttt{params}, in the parameter list. Note that \textit{all} Aztec preconditioners can be used as smoothers for ML. At this point we can create the ML preconditioner as

\[
\text{ML\_Epetra::MultiLevelPreconditioner * MLPrec =}
\]

\[
\text{new ML\_Epetra::MultiLevelPreconditioner(A, MLList, true);}\]

\(^1\)Aggregation schemes based on ParMETIS are also available. Please refer to the help of the ML configure for more details.

\(^2\)Epetra\_VbrMatrix and Epetra\_RowMatrix can be used as well.
and check that no options have been misspelled, using

    MLPrec->PrintUnused();

AztecOO solver is called using

    solver.SetPrecOperator(MLPrec);

    solver.SetAztecOption(AZ_solver, AZ_cg_condnum);

    int Niters = 500;
    solver.SetAztecOption(AZ_kspace, 160);

    solver.Iterate(Niters, 1e-12);

Finally, some (limited) information about the preconditioning phase are obtained using

    cout << MLPrec->GetOutputList();

The entire code is reported in
didasko/examples/ml/ex2.cpp.
Interfacing Direct Solvers with Amesos

Marzio Sala

The Amesos package provides an object-oriented interface to several direct sparse solvers. Amesos will solve (using a direct factorization method) the linear systems of equations (7.1) where \( A \) is stored as an Epetra_RowMatrix object, and \( X \) and \( B \) are Epetra_MultiVector objects.

The Amesos package has been designed to face some of the challenges of direct solution of linear systems. In fact, many solvers have been proposed in the last years, and often each of them requires different input formats for the linear system matrix. Moreover, it is not uncommon that the interface changes between revisions. Amesos aims to solve those problems, furnishing a clean, consistent interface to many direct solvers.

12.1 Introduction to the Amesos Design

Using Amesos, users can interface their codes with a (large) variety of direct linear solvers, sequential or parallel, simply by a code instruction of type

\[
\text{AmesosProblem.Solver();}
\]

Amesos will take care of redistributing data among the processors, if necessary.

All the Amesos classes are derived from a base class mode, \texttt{Amesos_BaseSolver}. This abstract interface provides the basic functionalities for all Amesos solvers, and allows users to choose different direct solvers very easily – by changing an input scalar parameter. See Section 12.2 for more details.

In this Chapter, we will suppose that matrix \( A \) in equation (7.1) is defined as an Epetra_RowMatrix, in principle with nonzero entries on all the processes defined in the Epetra_Comm communicator in use. \( X \) and \( B \), instead, are Epetra_MultiVector, defined on the same communicator.

Amesos contains several classes:

- \texttt{Amesos_Lapack}: Interface to serial dense solver LAPACK.
- \texttt{Amesos_KLU}: Interface to Amesos’s internal solver KLU. KLU is a serial, unblocked code ideal for getting started, and for very sparse matrices, such as circuit matrices.
- \texttt{Amesos_Umfpack}: Interface to Tim Davis’s UMFPACK [Dav03]. UMFPACK is a serial solver.
Chapter 12. Interfacing Direct Solvers with Amesos

- **Amesos_Superludist**: Interface to Xiaoye S. Li's SuperLU solver suite, including SuperLU, SuperLU_DIST 1.0 and SuperLU_DIST 2.0 [LD03]. SuperLU is a serial solver, while SuperLU_DIST is a parallel solver.

- **Amesos_Mumps**: Interface to MUMPS 4.3.1 [ADLK03]. MUMPS is a parallel direct solver;

- **Amesos_Scalapack**: Interface to ScaLAPACK [BCC+97], the parallel version of LAPACK [92].

If the supported packages is serial, and one is solving with more than one process, matrix and right-hand side are shipped to process 0, solved, then the solution is broadcasted to the distributed solution vector $X$. For parallel solvers, instead, various options are supported, depending on the package at hand:

- The Amesos_Superludist interface can be used over all the processes, as well as on a subset of them. The matrix is kept in distributed form over the processes of interest;

- Amesos_Mumps can keep the matrix in a distributed form over all the processes, or the matrix can be shipped to processor 0. In both cases, all the processes in the MPI communicator will be used.

This Chapter, we will cover:

- The Amesos_BaseSolver interface to various direct solvers, presented (in Section 12.2).

### 12.2 Amesos_BaseSolver: A Generic Interface to Direct Solvers

All Amesos objects are constructed from the function class **Amesos**. Amesos allows a code to delay the decision about which concrete class to use to implement the Amesos_BaseSolver interface. The main goal of this class is to allow the user to select any supported (and enabled at configuration time) direct solver, simply changing an input parameter. Another remarkable advantage of Amesos_BaseSolver is that, using this class, users does not have to include the header files of the 3-part libraries in their code.

An example of use of this class is as follows. First, the following header files must be included:

```c
#include "Amesos.h"
#include "AmesosClassType.h"
```

Then, let A be an Epetra_RowMatrix object (for instance, and Epetra_CrsMatrix). We need to define a linear problem,

```c
Epetra_LinearProblem * Amesos_LinearProblem =
    new Epetra_LinearProblem;
Amesos_LinearProblem->SetOperator( A );
```

1 At present, MUMPS is the only Amesos class that can take advantage of the symmetry of the linear system matrix.

2 Note that Amesos does not contain interfaces to LAPACK routines. Other Trilinos packages already offer those routines (Epetra and Teuchos).

3 Using Amesos_BaseSolver, 3-part libraries header files are required in the compilation of Amesos only.
and to create an Amesos parameter list (which can be empty):

```cpp
Teuchos::ParameterList ParamList;
```

Now, let `Choice` be an string, with one of the following values:
- `Amesos_Klu`;
- `Amesos_Umfpack`;
- `Amesos_Mumps`;
- `Amesos_Superludist`;
- `Amesos_Scalapack`.

We can construct an `Amesos_BaseSolver` object as follows:

```cpp
Amesos_BaseSolver * A_Base;
Amesos A_Factory;
A_Base = A_Factory.Create(Choice, *Amesos_LinearProblem, ParamList);
assert(A_Base!=0);
```

Symbolic and numeric factorizations are computed using methods

```cpp
A_Base->SymbolicFactorization();
A_Base->NumericFactorization();
```

The numeric factorization phase will check whether a symbolic factorization exists or not. If not, method `SymbolicFactorization()` is invoked. Solution is computed (after setting of LHS and RHS in the linear problem), using

```cpp
A_Base->Solve();
```

The solution phase will check whether a numeric factorization exists or not. If not, method `SymbolicFactorization()` is called.

Users must provide the nonzero structure of the matrix for the symbolic phase, and the actual nonzero values for the numeric factorization. Right-hand side and solution vectors must be set before the solution phase, for instance using

```cpp
Amesos_LinearProblem->SetLHS(x);
Amesos_LinearProblem->SetRHS(b);
```

A common ingredient to all the Amesos classes is the Teuchos parameters’ list. This object, whose definition requires the input file `Teuchos_ParameterList.hpp`, is used to specify the parameters that affect the 3-part libraries. Here, we simply recall that the parameters’ list can be created as

```cpp
Teuchos::ParameterList AmesosList;
```

and parameters can be set as

```cpp
AmesosList.set(ParameterName,ParameterValue);
```

Here, `ParameterName` is a string containing the parameter name, and `ParameterValue` is any valid C++ object that specifies the parameter value (for instance, an integer, a pointer to an array or to an object).

For a detailed list of parameters we refer to [SS04].
Two goals motivated the development of the Anasazi eigensolver package: interoperability and extensibility. The intention of interoperability is to ease the use of Anasazi in a wide range of application environments. To this end, the algorithms written in Anasazi utilize an abstract interface for operators and vectors, allowing the user to leverage existing linear algebra libraries. The concept of extensibility drives development of Anasazi to allow users to make efficient use of Anasazi codes while simultaneously enabling them to easily develop their own code in the Anasazi framework. This is encouraged by promoting code modularization and multiple levels of access to solvers and their data. In this Chapter, we outline the Anasazi eigensolver framework and motivate the design. In particular, we present

- the Anasazi operator/vector interface (Section 13.1);
- the Anasazi eigensolver framework (Section 13.2);
- a description of Anasazi classes (Section 13.3);
- the interface to the Epetra linear algebra package (Section 13.4);
- an example using Anasazi for the solution of an eigenvalue problem (Section 13.5).

### 13.1 The Anasazi Operator/Vector Interface

The Anasazi eigensolver package utilizes abstract interfaces for operators and multivectors. The goals of this are to leverage existing linear algebra libraries and to protect previous software investment. Algorithms in Anasazi are developed at a high-level, where the underlying linear algebra objects are opaque. The choice in linear algebra is made through templating, and access to the functionality of the underlying objects is provided via the traits classes Anasazi::MultiVecTraits and Anasazi::OperatorTraits.

These classes define opaque interfaces, specifying the operations that multivector and operator classes must support in order to be used in Anasazi without exposing low-level details of the underlying data structures.
### Table 13.1: Methods required by MultiVecTraits interface.

<table>
<thead>
<tr>
<th>Method name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clone</td>
<td>Creates a new empty multivector containing a specified number of columns.</td>
</tr>
<tr>
<td>CloneCopy</td>
<td>Creates a new multivector with a copy of the contents of an existing multivector.</td>
</tr>
<tr>
<td>CloneView</td>
<td>Creates a new multivector that shares the selected contents of an existing multivector.</td>
</tr>
<tr>
<td>GetVecLength</td>
<td>Returns the vector length of a multivector.</td>
</tr>
<tr>
<td>GetNumberVecs</td>
<td>Returns the number of vectors in a multivector.</td>
</tr>
<tr>
<td>MvTimesMatAddMv</td>
<td>Apply a SerialDenseMatrix $M$ to another multivector $A$, $B \leftarrow \alpha AM + \beta B$.</td>
</tr>
<tr>
<td>MvAddMv</td>
<td>Perform $mv \leftarrow \alpha A + \beta B$.</td>
</tr>
<tr>
<td>MvTransMv</td>
<td>Compute the matrix $C \leftarrow \alpha A^H B$.</td>
</tr>
<tr>
<td>MvDot</td>
<td>Compute the vector $b$ where the components are the individual dot-products of the $i$-th columns of $A$ and $B$, i.e. $b[i] = A[i]^H B[i]$.</td>
</tr>
<tr>
<td>MvScale</td>
<td>Scale the columns of a multivector.</td>
</tr>
<tr>
<td>MvNorm</td>
<td>Compute the 2-norm of each individual vector of $A$.</td>
</tr>
<tr>
<td>SetBlock</td>
<td>Copy the vectors in $A$ to a subset of vectors in $B$.</td>
</tr>
<tr>
<td>MvRandom</td>
<td>Replace the vectors in $A$ with random vectors.</td>
</tr>
<tr>
<td>MvInit</td>
<td>Replace each element of the vectors in $A$ with $\alpha$.</td>
</tr>
<tr>
<td>MvPrint</td>
<td>Print the multi-vector to an output stream.</td>
</tr>
</tbody>
</table>

The benefit of using a templated traits class over inheritance is that the latter requires the user to derive multivectors and operators from Anasazi-defined abstract base classes. The former, however, defines only local requirements: Anasazi-defined traits classes implemented as user-developed adapters for the chosen multivector and operator classes.

Anasazi::MultiVecTraits provides routines for the creation of multivectors, as well as their manipulation. In order to use a specific scalar type and multivector type with Anasazi, there must exist a template specialization of Anasazi::MultiVecTraits for this pair of classes. A full list of methods required by Anasazi::MultiVecTraits is given in Table 13.1.

Just as Anasazi::MultiVecTraits defined the interface required to use a multivector class with Anasazi, Anasazi::OperatorTraits defines the interface required to use the combination of a specific operator class with a specific multivector class. This interface defines a single method:

\[
\text{OperatorTraits<ScalarType,MV,OP>::Apply(const OP &Op, const MV &x, MV &y)}
\]

This method performs the operation $y = Op(x)$, where $Op$ is an operator of type OP and $x$ and $y$ are multivectors of type MV. In order to use the combination of OP and MV, there must
be a specialization of Anasazi::OperatorTraits for ScalarType, OP and MV.

Calling methods of MultiVecTraits and OperatorTraits requires that specializations of these traits classes have been implemented for given template arguments. Anasazi provides the following specializations of these traits classes:

- Epetra::MultiVector and Epetra::Operator (with scalar type double)
- Thyra::MultiVectorBase and Thyra::LinearOpBase (with arbitrary scalar type)
  This allows Anasazi to be used with any classes that implement the abstract interfaces provided by the Thyra package.
- Anasazi::MultiVec and Anasazi::Operator (with arbitrary scalar type)
  This allows Anasazi to be used with any classes that implement the abstract base classes Anasazi::MultiVec and Anasazi::Operator.

For user-specified classes that don’t match one of the above, specializations of MultiVecTraits and OperatorTraits will need to be created by the user for use by Anasazi. Test routines Anasazi::TestMultiVecTraits() and Anasazi::TestOperatorTraits() are provided by Anasazi to help in testing user-developed adapters.

13.2 The Anasazi Eigensolver Framework

The goals of flexibility and efficiency can interfere with the goals of simplicity and ease of use. For example, efficient memory use and low-level data access required in scientific codes can lead to complicated interfaces and violations of standard object-oriented development practices.

In Anasazi, this problem is addressed by providing a multi-tiered access strategy for eigensolver algorithms. Anasazi users have the choice of interfacing at one of two levels: either working at a high-level with an eigensolver manager or working at a low-level directly with an eigensolver.

Consider as an example the block Davidson iteration. The essence of this iteration can be distilled into the following algorithm:

1. apply preconditioner $N$ to the current residuals: $H = NR$
2. use $H$ to expand current basis $V$
3. use new $V$ to compute a projected eigenproblem
4. solve the projected eigenproblem and form the Ritz vectors $X$ and Ritz values $\Theta$
5. compute the new residuals $R$

In implementing a block Davidson method, this iteration repeats until the basis $V$ is full (in which case it is time to restart) or some stopping criterion has been satisfied. Many valid stopping criteria exist, as well as many different methods for restarting the basis. Both of these, however, are distinct from the essential iteration as described above. A user wanting to perform block Davidson iterations could ask the solver to perform these iterations until a user-specified stopping criterion was satisfied or the basis was full, at which time the user would perform a restart. This allows the user complete control over the stopping criteria and
the restarting mechanism, and leaves the eigensolver responsible for a relatively simple bit of state and behavior.

This is the way that Anasazi has been designed. The eigensolvers (encapsulating an iteration and the associated state) are derived classes of the abstract base class Anasazi::Eigensolver. The goals of this class are three-fold:

• to define an interface used for checking the status of a solver by a status test;
• to contain the essential iteration associated with a particular eigensolver iteration;
• to contain the state associated with that iteration.

The status tests, assembled to describe a specific stopping criterion and queried by the eigensolver during the iteration, are represented as subclasses of Anasazi::StatusTest. The communication between status test and eigensolver occurs inside of the iterate() method provided by each Anasazi::Eigensolver. This code generally takes the form:

```cpp
SomeEigensolver::iterate() {
    while ( statustest.checkStatus(this) != Passed ) {
        // perform eigensolver iterations
    }
    return; // return back to caller
}
```

Each Anasazi::StatusTest provides a method, checkStatus(), which through queries to the methods provided by Anasazi::Eigensolver, determines whether the solver meets the criteria defined by that particular status test. After a solver returns from iterate(), the caller has the option of accessing the state associated with the solver and re-initializing the solver with a new state.

While this method of interfacing with the solver is powerful, it can be tedious. This method requires that user construct a number of support classes, in addition to managing call to Eigensolver::iterate(). The Anasazi::SolverManager class was developed to address this complaint. A solver manager is a class that wraps around an eigensolver, providing additional functionality while also handling lower-level interaction with the eigensolver that a user may not wish to handle. Solver managers are intended to be easy to use, while still providing the features and flexibility needed to solve real-world eigenvalue problems. For example, the Anasazi::BlockDavidsonSolMgr takes only two arguments in its constructor: an Anasazi::Eigenproblem specifying the eigenvalue problem to be solved and a Teuchos::ParameterList of options specific to this solver manager. The solver manager instantiates an eigensolver, along with the status tests and other support classes needed by the eigensolver. To solve the eigenvalue problem, the user simply calls the solve() method of the solver manager. The solver manager performs repeated calls to the eigensolver iterate() method, performs restarts and locking, and places the final solution into the eigenproblem.

Users therefore have a number of options for performing eigenvalue computations with Anasazi:

• use an existing solver manager;
  In this case, the user is limited to the functionality provided by the current eigensolvers.
• develop a new solver manager around an existing eigensolver;
The user can extend the functionality provided by the eigensolver, specifying custom configurations for status tests, orthogonalization, restarting, locking, etc.

• develop a new eigensolver/solver manager;
The user can write an eigensolver for an iteration that is not represented in Anasazi. The user still has the benefit of the support classes provided by Anasazi, and the knowledge that the new solver/solver manager can be easily used by anyone already familiar with Anasazi.

13.3 Anasazi Classes

Anasazi is designed with extensibility in mind, so that users can augment the package with any special functionality that they need. However, the released version of Anasazi provides all functionality necessary for solving a wide variety of problems. This section lists and briefly describes the current classes found in Anasazi.

The solution of an eigenvalue problem requires a minimum subset of Anasazi classes. The following is a list of classes that any Anasazi user must be familiar with to use the package.

Remark 20. Anasazi makes extensive use of the Teuchos utility classes, especially Teuchos::RCP (Section 10.7) and Teuchos::ParameterList (Section 10.6). Users are encouraged to become familiar with these classes and their correct usage.

13.3.1 Anasazi::Eigenproblem

Anasazi::Eigenproblem is a container for the components of an eigenvalue problem, as well as the solutions. By requiring eigenproblems to derive from Anasazi::Eigenproblem, Anasazi defines a minimum interface that can be expected of all eigenvalue problems by the classes that will work with the problems (e.g., eigensolvers and status testers).

Both the eigenproblem and the eigensolver in Anasazi are templated on the scalar type, the multivector type and the operator type. Before declaring an eigenproblem, users must choose classes to represent these entities. Having done so, they can begin to specify the parameters of the eigenvalue problem. The Anasazi::Eigenproblem defines set methods for the parameters of the eigenproblem. These methods are:

• setOperator - set the operator for which the eigenvalues will be computed
• setA - set the $A$ operator for the eigenvalue problem $Ax = Mx\lambda$
• setM - set the $M$ operator for the eigenvalue problem $Ax = Mx\lambda$
• setPrec - set the preconditioner for the eigenvalue problem
• setInitVec - set the initial iterate
• setAuxVecs - set the auxiliary vectors, a subspace used to constrain the search space for the solution
• setNEV - set the number of eigenvalues to be computed
• **setHermitian** - specify whether the problem is Hermitian

In addition to these **set** methods, Anasazi::Eigenproblem defines a method **setProblem()** that gives the class the opportunity to perform any initialization that may be necessary before the problem is handed off to an eigensolver, in addition to verifying that the problem has been adequately defined.

For each of the **set** methods listed above, there is a corresponding **get** function. These are the functions used by eigensolvers and solver managers to get the necessary information from the eigenvalue problem. In addition, there are two methods for storing and retrieving the results of the eigenvalue computation:

```cpp
const Eigensolution & getSolution();
void setSolution(const Eigensolution & sol);
```

The Anasazi::Eigensolution structure is described in Section 13.3.2.

Anasazi provides users with a basic implementation of Anasazi::Eigenproblem, called Anasazi::BasicEigenproblem (Section 13.5). This formulation provides all the functionality necessary to describe both generalized and standard linear eigenvalue problems.

### 13.3.2 Anasazi::Eigensolution

The Anasazi::Eigensolution structure was developed in order to facilitate setting and retrieving of solution data. The class contains the following information:

- **Teuchos::RCP< MV > Evecs**
  The computed eigenvectors.

- **Teuchos::RCP< MV > Espace**
  An orthonormal basis for the computed eigenspace.

- **std::vector< Value< ScalarType > > Evals**
  The computed eigenvalues.

- **std::vector< int > index**
  An index into **Evecs** to enable compressed storage of eigenvectors for real, non-Hermitian problems.

- **int numVecs**
  The number of computed eigenpairs.

All Anasazi solver managers place the results of the computation in the Anasazi::Eigenproblem class using an Anasazi::Eigensolution structure. The number of eigensolutions computed is given by field **numVecs**. The eigenvalues are always stored as two real values, even when templated on a complex data type or when the eigenvalues are real. Similarly, the eigenspace can always be represented by a multivector of width **numVecs**, even for non-symmetric eigenproblems over the real field. The storage scheme for eigenvectors requires more finesse.

When solving real symmetric eigenproblems, the eigenvectors can always be chosen to be real, and therefore can be stored in a single column of a real multivector. When solving eigenproblems over a complex field, whether Hermitian or non-Hermitian, the eigenvectors
may be complex, but the multivector is defined over the complex field, so that this poses no problem. However, real non-symmetric problems can have complex eigenvectors, which prohibits a one-for-one storage scheme using a real multivector. Fortunately, the eigenvectors in this scenario occur as complex conjugate pairs, so the pair can be stored in two real vectors. This permits a compressed storage scheme, which uses an index vector stored in the Eigensolution, allowing conjugate pair eigenvectors to be easily retrieved from \texttt{Evecs}.

The integers in \texttt{Anasazi::Eigensolution::index} take one of three values: \{0, +1, −1\}. These values allow the eigenvectors to be retrieved as follows:

- \texttt{index[i]} = 0: the \texttt{i}-th eigenvector is stored uncompressed in column \texttt{i} of \texttt{Evecs}.
- \texttt{index[i]} = +1: the \texttt{i}-th eigenvector is stored compressed, with the real component in column \texttt{i} of \texttt{Evecs} and the \textit{positive} complex component stored in column \texttt{i} + 1 of \texttt{Evecs}.
- \texttt{index[i]} = −1: the \texttt{i}-th eigenvector is stored compressed, with the real component in column \texttt{i} − 1 of \texttt{Evecs} and the \textit{negative} complex component stored in column \texttt{i} of \texttt{Evecs}.

Because this storage scheme is only required for non-symmetric problems over the real field, all other eigenproblems will result in an index vector composed entirely of zeroes. For the real non-symmetric case, the +1 index will always immediately precede the corresponding −1 index.

\textbf{Remark 21.} Solver managers all put the computed eigensolution into the eigenproblem class before returning from \texttt{solve()}. Eigensolvers do not; a user working directly with an eigensolver will need to recover the solution directly from the eigensolver state.

\subsection*{13.3.3 \texttt{Anasazi::Eigensolver}}

The \texttt{Anasazi::Eigensolver} class defines the basic interface that must be met by any eigensolver class in \texttt{Anasazi}. The specific eigensolvers are implemented as derived classes of \texttt{Anasazi::Eigensolver}. Table 13.2 lists the eigensolver currently implemented in \texttt{Anasazi}.

<table>
<thead>
<tr>
<th>Solver</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>BlockDavidson</td>
<td>A block Davidson solver for Hermitian eigenvalue problems.</td>
</tr>
<tr>
<td>BlockKrylovSchur</td>
<td>A block Krylov Schur solver for Hermitian or non-Hermitian eigenvalue problems.</td>
</tr>
<tr>
<td>LOBPCG</td>
<td>The locally optimal block preconditioned conjugate gradient method for Hermitian eigenproblems.</td>
</tr>
</tbody>
</table>

\textbf{Table 13.2:} Eigensolvers currently implemented in \texttt{Anasazi}.

Each eigensolver provides two significant types of methods: status methods and solver-specific state methods. The status methods are defined by the \texttt{Anasazi::Eigensolver} abstract base class and represent the information that a generic status test can request from any eigensolver. A list of these methods is given in Table 13.3.

The class \texttt{Anasazi::Eigensolver}, like \texttt{Anasazi::Eigenproblem}, is templated on the scalar type, multivector type and operator type. The options for the eigensolver are passed through the constructor, defined by \texttt{Anasazi::Eigensolver} to have the following form:
Chapter 13. Eigenvalue and Eigenvector Computations with Anasazi

Method | Description
--- | ---
getNumIters | Get the current number of iterations.
getRitzValues | Get the most recent Ritz values.
getRitzVectors | Get the most recent Ritz vectors.
getRitzIndex | Get the Ritz index needed for indexing compressed Ritz vectors.
getResNorms | Get the most recent residual norms, with respect to the OrthoManager.
getRes2Norms | Get the most recent residual 2-norms.
getRitzRes2Norms | Get the most recent Ritz residual 2-norms.
getCurSubspaceDim | Get the current subspace dimension.
getMaxSubspaceDim | Get the maximum subspace dimension.
getBlockSize | Get the block size.

Table 13.3: A list of generic status methods provided by Anasazi::Eigensolver.

Eigensolver(
    const Teuchos::RCP< Eigenproblem<ST,MV,OP> > &problem,
    const Teuchos::RCP< SortManager<ST,MV,OP> > &sorter,
    const Teuchos::RCP< OutputManager<ST> > &printer,
    const Teuchos::RCP< StatusTest<ST,MV,OP> > &tester,
    const Teuchos::RCP< OrthoManager<ST,OP> > &ortho,
    ParameterList &params)
);

These classes are used as follows:

- **problem** - the eigenproblem to be solved; the solver will get the problem operators from here.
- **sorter** - the sort manager selects the significant eigenvalues; see Section 13.3.6.
- **printer** - the output manager dictates verbosity level in addition to processing output streams; see Section 13.3.8.
- **tester** - the status tester dictates when the solver should quit iterate() and return to the caller; see Section 13.3.5.
- **ortho** - the orthogonalization manager defines the inner product and other concepts related to orthogonality, in addition to performing these computations for the solver; see Section 13.3.7.
- **params** - the parameter list specifies eigensolver-specific options; see the documentation for a list of options support by individual solvers.

In addition to specifying an iteration, an eigensolver also specifies a concept of state, i.e. the current data associated with the iteration. After declaring an Eigensolver object, the state of the solver is in an uninitialized state. For most solver, to be initialized mean to be in a valid state, containing all of the information necessary for performing eigensolver iterations.
Anasazi::Eigensolver provides two methods concerning initialization: `isInitialized()` indicates whether the solver is initialized or not, and `initialize()` (with no arguments) instructs the solver to initialize itself using random data or the initial vectors stored in the eigenvalue problem.

To ensure that solvers can be used as efficiently as possible, the user needs access to the state of the solver. To this end, each eigensolver provides low-level methods for getting and setting the state of the solver:

- `getState()` - returns a solver-specific structure with read-only pointers to the current state of the solver.
- `initialize(...)` - accepts a solver-specific structure enabling the user to initialize the solver with a particular state.

The combination of these two methods, along with the flexibility provided by status tests, allows the user almost total control over eigensolver iterations.

### 13.3.4 Anasazi::SolverManager

Using Anasazi by interfacing directly with eigensolvers is extremely powerful, but can be tedious. Solver managers provide a way for users to encapsulate specific solving strategies inside of an easy-to-use class. Novice users may prefer to use existing solver managers, while advanced users may prefer to write custom solver managers.

Anasazi::SolverManager defines only two methods: a constructor accepting an Anasazi::Eigenproblem and a parameter list of solver manager-specific options; and a `solve()` method, taking no arguments and returning either Anasazi::Converged or Anasazi::Unconverged. Consider the following example code:

```cpp
// create an eigenproblem
Teuchos::RCP< Anasazi::Eigenproblem<ScalarType,MV,OP> > problem = ...;
// create a parameter list
Teuchos::ParameterList params;
params.set(...);
// create a solver manager
Anasazi::BlockDavidsonSolMgr<ScalarType,MV,OP> solman(problem,params);
// solve the eigenvalue problem
Anasazi::ReturnType ret = solman.solve();
// get the solution from the problem
Anasazi::Eigensolution sol = problem->getSolution();
```

**Remark 22.** Errors in Anasazi are communicated via exceptions. This is outside the scope of this tutorial; see the Anasazi documentation for more information.

As has been stated before, the goal of the solver manager is to create an eigensolver object, along with the support objects needed by the eigensolver. Another purpose of many solver managers is to manage and initiate the repeated calls to the underlying solver's `iterate()` method. For solvers that build a Krylov subspace to some maximum dimension (e.g., Block-KrylovSchur and BlockDavidson), the solver manager will also assume the task of restarting the solver when the subspace is full. This is something for which multiple approaches are
possible. Also, there may be substantial flexibility in creating the support classes (e.g., sort manager, status tests) for the solver. An aggressive solver manager could even go so far as to construct a preconditioner for the eigenvalue problem.

These examples are meant to illustrate the flexibility that specific solver managers may have in implementing the `solve()` routine. Some of these options might best be incorporated into a single solver manager, which takes orders from the user via the parameter list given in the constructor. Some of these options may better be contained in multiple solver managers, for the sake of code simplicity. It is even possible to write solver managers that contain other solvers managers; motivation for something like this would be to select the optimal solver manager at runtime based on some expert knowledge, or to create a hybrid method which uses the output from one solver manager to initialize another one.

### 13.3.5 Anasazi::StatusTest

By this point in the tutorial, the purpose of the Anasazi::StatusTest should be clear: to give the user or solver manager flexibility in stopping the eigensolver iterations in order to interact directly with the solver.

Many reasons exist for why a user would want to stop the solver from iterating:

- some convergence criterion has been satisfied and it is time to quit;
- some part of the current solution has reached a sufficient accuracy to removed from the iteration ("locking");
- the solver has performed a sufficient or excessive number of iterations.

These are just some commonly seen reasons for ceasing the iteration, and each of these can be so varied in implementation/parametrization as to require some abstract mechanism controlling the iteration.

The following is a list of Anasazi-provided status tests:

- **Anasazi::StatusTestCombo** - this status test allows for the boolean combination of other status tests, creating near unlimited potential for complex status tests.
- **Anasazi::StatusTestOutput** - this status test acts as a wrapper around another status test, allowing for printing of status information on a call to `checkStatus()`
- **Anasazi::StatusTestMaxIters** - this status test monitors the number of iterations performed by the solver; it can be used to halt the solver at some maximum number of iterations or even to require some minimum number of iterations.
- **Anasazi::StatusTestResNorm** - this status test monitors the residual norms of the current iterate.
- **Anasazi::StatusTestOrderedResNorm** - this status test also monitors the residual norms of the current iterate, but only considers the residuals associated with the most significant part of the current iterate.
### 13.3. Anasazi Classes

#### Table 13.4: Options for Anasazi::BasicSort.

<table>
<thead>
<tr>
<th>Option</th>
<th>Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>SM</td>
<td>Sort eigenvalues in increasing order of magnitude</td>
</tr>
<tr>
<td>SR</td>
<td>Sort eigenvalues in increasing order of real part</td>
</tr>
<tr>
<td>SI</td>
<td>Sort eigenvalues in increasing order of imaginary part</td>
</tr>
<tr>
<td>LM</td>
<td>Sort eigenvalues in decreasing order of magnitude</td>
</tr>
<tr>
<td>LR</td>
<td>Sort eigenvalues in decreasing order of real part</td>
</tr>
<tr>
<td>LI</td>
<td>Sort eigenvalues in decreasing order of imaginary part</td>
</tr>
</tbody>
</table>

#### 13.3.6 Anasazi::SortManager

The purpose of a sort manager is to separate the eigensolver classes from the sorting functionality required by those classes. This satisfies the flexibility principle sought by Anasazi, by giving users the opportunity to perform the sorting in whatever manner is deemed to be most appropriate. Anasazi defines an abstract class Anasazi::SortManager with two methods, one for sorting real values and one for sorting complex values:

```cpp
ReturnType sort (... , std::vector<MagnitudeType> &evals,
                 std::vector<int> *perm)
ReturnType sort (... , std::vector<MagnitudeType> &r_evals,
                 std::vector<MagnitudeType> &i_evals,
                 std::vector<int> *perm)
```

Each of these sort routines will sort the eigenvalues according to some implementation and optionally return the permutation vector as well (useful for sorting associated vectors).

Anasazi provides a derived class Anasazi::BasicSort. This class provides basic sorting functionality, described in Table 13.4.

#### 13.3.7 Anasazi::OrthoManager

Orthogonalization and orthonormalization are commonly performed computations in iterative eigensolvers; in fact, for some eigensolvers, they represent the dominant cost. Different scenarios may require different approaches (e.g., Euclidean inner product versus \(M\) inner product, orthogonal projections versus oblique projections). Combined with the plethora of available methods for performing these computations, Anasazi has left as much leeway to the users as possible.

Orthogonalization of multivectors in Anasazi is performed by derived classes of the abstract class Anasazi::OrthoManager. This class provides five methods:

- `innerProd(X,Y,Z)` - performs the inner product defined by the manager.
- `norm(X)` - computes the norm induced by `innerProd()`.
- `project(X,C,Q)` - given an orthonormal basis \(Q\), projects \(X\) onto the space perpendicular to \(\text{colspan}(Q)\), optionally returning the coefficients of \(X\) in \(Q\).
- `normalize(X,B)` - returns an orthonormal basis for \(\text{colspan}(X)\), optionally returning the coefficients of \(X\) in the computed basis.
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<table>
<thead>
<tr>
<th>Message type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Errors</td>
<td>Errors (always printed)</td>
</tr>
<tr>
<td>Warnings</td>
<td>Warning messages</td>
</tr>
<tr>
<td>IterationDetails</td>
<td>Approximate eigenvalues, errors</td>
</tr>
<tr>
<td>OrthoDetails</td>
<td>Orthogonalization/orthonormalization checking</td>
</tr>
<tr>
<td>FinalSummary</td>
<td>Final computational summary (usually from SolverManager::solve())</td>
</tr>
<tr>
<td>TimingDetails</td>
<td>Timing details</td>
</tr>
<tr>
<td>StatusTestDetails</td>
<td>Status test details</td>
</tr>
<tr>
<td>Debug</td>
<td>Debugging information</td>
</tr>
</tbody>
</table>

Table 13.5: Message types used by Anasazi::OutputManager.

- **projectAndNormalize(X,C,B,Q)** - computes an orthonormal basis for subspace \( \text{colspan}(X) - \text{colspan}(Q) \), optionally returning the coefficients of \( X \) in \( Q \) and the new basis.

It should be noted that a call to **projectAndNormalize()** is not necessarily equivalent to a call to **project()** followed by **normalize()**. This follows from the fact that, for some orthogonalization managers, a call to **normalize()** may augment the column span of a rank-deficient multivector in order to create an orthonormal basis with the same number of columns as the input multivector. In this case, the code

```cpp
orthoman.project(X,C,Q);
orthoman.normalize(X,B);
```

could result in an orthonormal basis \( X \) that is not orthogonal to the basis in \( Q \).

Anasazi provides two orthogonalization managers:

- **Anasazi::BasicOrthoManager** - performs orthogonalization using multiple steps of classical Gram-Schmidt.

- **Anasazi::SVQBOorthoManager** - performs orthogonalization using the SVQB orthogonalization technique described by Stathopoulos and Wu.

More information on these orthogonalization managers is available in the Anasazi documentation.

### 13.3.8 Anasazi::OutputManager

The output manager in Anasazi exists to provide flexibility with regard to the verbosity of the eigensolver. Each output manager has two primary concerns: what output is printed and where the output is printed to. When working with the output manager, output is classified into one of the message types from Table 13.5.

Output manager in Anasazi are subclasses of the abstract base class Anasazi::OutputManager. This class provides the following output-related methods:

- **bool isVerbosity (MsgType type)** - Find out whether we need to print out information for this message type.
13.4 Using the Anasazi adapter to Epetra

- void print (MsgType type, const string output) - Send output to the output manager.
- ostream & stream (MsgType type) - Create a stream for outputting to.

The output manager is meant to ease some of the difficulty associated with I/O in a distributed programming environment. For example, consider some debugging output requiring optional computation. For reasons of efficiency, we may want to perform the computation only if debugging is requested; i.e., isVerbosity(Anasazi::Debug) == true. However, while we need all nodes to enter the code block to perform the computation, we probably want only one of them to print the output. Furthermore, the user may want different types of output treated in a different manner. The abstraction of the printing mechanism allows both of these goals to be met.

Anasazi provides a single output manager, Anasazi::BasicOutputManager. This class accepts an output stream from the user. The output corresponding to the verbosity level of the manager is sent to this stream only on the master node; the output for other nodes is neglected.

13.4 Using the Anasazi adapter to Epetra

The Epetra package provides the underlying linear algebra foundation for many Trilinos solvers. By using the Anasazi adapter to Epetra, users not only avoid the trouble of designing their own multivector and operator classes, but they also gain the ability to utilize any other Trilinos package which recognizes Epetra classes (such as AztecOO, IFPACK, and others).

In order to use the Anasazi adapter to Epetra, users must include the following file:

```cpp
#include "AnasaziEpetraAdapter.hpp"
```

This file simply defines specializations of the Anasazi::MultiVecTraits and Anasazi::OperatorTraits classes, while also including the Epetra header files defining the multivector and operator classes.

Because Epetra makes exclusive use of double precision arithmetic, Epetra_Operator and Epetra_MultiVector are used only with scalar type double. For brevity, it is useful to declare type definitions for these classes:

```cpp
typedef double ST;
typedef Epetra_MultiVector MV;
typedef Epetra_Operator OP;
```

Multivectors will be of type MV:

```cpp
Teuchos::RCP<MV> X
    = Teuchos::rcp( new MV(...) );
```

Operators can be any subclass of OP, for example, an Epetra_CrsMatrix:

```cpp
Teuchos::RCP<OP> A
    = Teuchos::rcp( new Epetra_CrsMatrix(...) );
```
The Anasazi interface to Epetra defines a specialization of Anasazi::MultiVecTraits for Epetra::MultiVector and a specialization of Anasazi::OperatorTraits for Epetra::Operator applied to Epetra::MultiVector. Therefore, we can now specify an eigenproblem and eigensolver utilizing these computational classes. An example defining an eigenvalue problem and solving the problem using an Anasazi eigensolver is given in the next section.

13.5 Defining and Solving an Eigenvalue Problem

This section gives sample code for solving a symmetric eigenvalue problem using the Block Krylov Schur solver manager, Anasazi::BlockKrylovSchurSolMgr. The example code in this section comes from the Didasko example didasko/examples/anasazi/ex1.cpp.

The first step in solving an eigenvalue problem is to define the eigenvalue problem. Assume we have chosen classes to represent our scalars, multivectors and operators as ST, MV and OP, respectively. Given an operator A and a multivector X containing initial vectors, both wrapped in Teuchos::RCP, we might define the eigenproblem as follows:

```cpp
Teuchos::RCP<BasicEigenproblem<ST,MV,OP> > MyProblem = Teuchos::rcp( new BasicEigenproblem<ST,MV,OP>(A,X) );
MyProblem->setHermitian( true );
MyProblem->setNEV( 4 );
bool ret = MyProblem->SetProblem();
if (ret != true) {
    // there should be no error in this example :)
}
```

The first line creates a Anasazi::BasicEigenproblem object and wraps it in a Teuchos::RCP (Section 10.7). The second line specifies the symmetry of the eigenproblem. The third line specifies the desired number of eigenvalues and eigenvectors. Lastly, the fourth signals that we have finished setting up the eigenproblem. This step must be completed before attempting to solve the problem.

If we were directly using the Anasazi::BlockKrylovSchur eigensolver, we would proceed by creating all of the support objects needed by the solver: a sort manager, an output manager, an orthogonalization manager, and a status test.

Instead, we will utilize a solver manager for solving the problem. First, we create a parameter list to specify the parameters for the solver manager:

```cpp
int verb = Anasazi::Warnings + Anasazi::Errors + Anasazi::FinalSummary + Anasazi::TimingDetails;
Teuchos::ParameterList MyPL;
MyPL.set( "Verbosity", verb );
MyPL.set( "Which", "SM" );
MyPL.set( "Block Size", 4 );
MyPL.set( "Num Blocks", 20 );
MyPL.set( "Maximum Restarts", 100 );
MyPL.set( "Convergence Tolerance", 1.0e-8 );
```

Here, we have asked for the eigensolver to output information regarding errors and warnings, as well as to provide a final summary after completing all iterations and to print the
timing information collected during the solve. We have also specified the tolerance for convergence testing (used to construct a status test); the block size and number of blocks (passed on to the solver); the desired eigenvalues (given to the sort manager); and the maximum number of restarts (used by the solver manager, the agent performing the restarts). This solver manager permits other options as well, affecting the step size as well as the convergence criteria; see the Anasazi documentation.

We now have all of the information needed to declare the solver manager and solve the problem:

\[
\text{Anasazi::BlockKrylovSchurSolMgr<ST,MV,OP> MyBlockKrylovSchur(MyProblem, MyPL );}
\]

The eigenproblem is solved with the instruction

\[
\text{Anasazi::ReturnType solverreturn = MyBlockKrylovSchur.solve();}
\]

The return value of the solver indicates whether the algorithm succeeded or not; i.e., whether the requested number of eigenpairs were found to a sufficient accuracy (as defined by the solver manager). Output from \text{solve()} routine in this example might look as follows:

```
================================================================================
BlockKrylovSchur Solver Status
The solver is initialized.
The number of iterations performed is 39
The block size is 4
The number of blocks is 20
The current basis size is 4
The number of auxiliary vectors is 0
The number of operations \text{Op*x} is 156
CURRENT RITZ VALUES
<table>
<thead>
<tr>
<th>Ritz Value</th>
<th>Ritz Residual</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.620281e-01</td>
<td>9.482040e-15</td>
</tr>
<tr>
<td>3.985070e-01</td>
<td>5.712179e-14</td>
</tr>
<tr>
<td>3.985070e-01</td>
<td>2.536671e-14</td>
</tr>
<tr>
<td>6.349859e-01</td>
<td>4.846649e-11</td>
</tr>
</tbody>
</table>
================================================================================
```

```
================================================================================
TimeMonitor Results
<table>
<thead>
<tr>
<th>Timer Name</th>
<th>Local time (num calls)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Operation \text{Op*x}</td>
<td>0.001701 (39)</td>
</tr>
</tbody>
</table>
```
Eigenvalues and eigenvectors can be retrieved from the eigenproblem (where they were stored by the solver manager) as follows:

```cpp
Anasazi::Eigensolution<ST,MV> sol = MyProblem->getSolution();
```

Four examples are provided with the tutorial:

- `didasko/examples/anasazi/ex1.cpp`: compute the eigenvectors corresponding to the smallest eigenvalues for a 2D Laplace problem using the block Krylov Schur solver

- `didasko/examples/anasazi/ex2.cpp`: solves the problem from `ex1` using instead the block Davidson eigensolver

- `didasko/examples/anasazi/ex3.cpp`: uses the block Krylov Schur solver to solve a non-Hermitian convection-diffusion problem

- `didasko/examples/anasazi/ex4.cpp`: uses the LOBPCG solver to solve the 2D Laplacian problem from `ex1`
Solving Nonlinear Systems with NOX

Marzio Sala, Michael Heroux, David Day

NOX is a suite of solution methods for the solution of nonlinear systems of type

\[ F(x) = 0, \] (14.1)

where

\[ F(x) = \begin{pmatrix} f_1(x_1, \ldots, x_n) \\ \vdots \\ f_n(x_1, \ldots, x_n) \end{pmatrix} \]

is a nonlinear vector function, and the Jacobian matrix of \( F \), \( J \), is defined by

\[ J_{i,j} = \frac{\partial F_i}{\partial x_j}(x). \]

NOX aims to solve \((14.1)\) using Newton-type methods. NOX uses an abstract vector and "group" interface. Current implementation are provided for Epetra/AztecOO objects, but also for LAPACK and PETSc. It provides various strategies for the solution of nonlinear systems, and it has been designed to be easily integrated into existing applications.

In this Chapter, we will

- Outline the basic issue of the solution of nonlinear systems (in Section 14.1);
- Introduce the NOX package (in Section 14.2);
- Describe how to introduce a NOX solver in an existing code (in Section 14.3);
- Present Jacobian-free methods (in Section 14.5).

14.1 Theoretical Background

Aim of this Section is to briefly present some aspects of the solution of nonlinear systems, to establish a notation. The Section is not supposed to be exhaustive, nor complete on this subject. The reader is referred to the existing literature for a rigorous presentation.
To solve system of nonlinear equations, NOX makes use of Newton-like methods. The Newton method defines a suite \( \{x_k\} \) that, under some conditions, converges to \( x \), solution of (14.1). The algorithm is as follows: given \( x_0 \), for \( k = 1, \ldots \) until convergence, solve

\[
J_k(x_{k-1}) (x_k - x_{k-1}) = -F(x_{k-1}), \quad J_k(x_{k-1}) = \left[ \frac{\partial F}{\partial x}(x_{k-1}) \right].
\]

(14.2)

Newton method introduces a local full linearization of the equations. Solving a system of linear equations at each Newton step can be very expensive if the number of unknowns is large, and may not be justified if the current iterate is far from the solution. Therefore, a departure from the Newton framework consists of considering inexact Newton methods, which solve system (14.2) only approximatively.

In fact, in practical implementation of the Newton method, one or more of the following approximations are used:

1. The Fréchet derivative \( J_k \) for the Newton step is not recomputed at every Newton step;
2. The equation for the Newton step (14.2) is solved only inexactly;
3. Defect-correction methods are employed, that is, \( J_k \) is numerically computed using low-order (in space) schemes, while the right-hand side is built up using high-order methods.

For a given initial guess, “close enough” to the solution of (14.1), the Newton method with exact linear solves converges quadratically. In practice, the radius of convergence is often extended via various methods. NOX provides, among others, line search techniques and trust region strategies.

### 14.2 Creating NOX Vectors and Group

NOX is not based on any particular linear algebra package. Users are required to supply methods that derive from the abstract classes \texttt{NOX::Abstract::Vector} (which provides support for basic vector operations as dot products), and \texttt{NOX::Abstract::Group} (which supports the linear algebra functionalities, evaluation of the function \( G \) and, optionally, of the Jacobian \( J \)).

In order to link their code with NOX, users have to write their own instantiation of those two abstract classes. In this tutorial, we will consider the concrete implementations provided for Epetra matrices and vectors. As this implementation is separate from the NOX algorithms, the configure option \texttt{--enable-nox-epetra} has to be specified (see Section 1.2)

### 14.3 Introducing NOX in an Existing Code

Two basic steps are required to implement a \texttt{NOX::Epetra} interface. First, a concrete class derived from \texttt{NOX::Epetra::Interface} has to be written. This class must define the following methods:

1. A method to compute \( y = F(X) \) for a given \( x \). The syntax is

\[ Other two concrete implementation are provided, for LAPACK and PETSc. The user may wish to configure NOX with \texttt{--enable-nox-lapack} or \texttt{--enable-nox-petsc}. Examples can be compiled with the options \texttt{--enable-nox-lapack-examples}, \texttt{--enable-nox-petsc-examples}, and \texttt{--enable-nox-epetra-examples}. \]
computeF(const Epetra_Vector & x, Epetra_Vector & y, 
   FillType flag)

with x and y two Epetra_Vectors, and flag an enumerated type that tells why this 
method was called. In fact, NOX has the ability to generate Jacobians based on nu-
merical differencing. In this case, users may want to compute an inexact (and hopefully 
cheaper) $F$, since it is only used in the Jacobian (or preconditioner).

2. A function to compute the Jacobian, whose syntax is

computeJacobian(const Epetra_Vector & x, 
   Epetra_Operator * Jac)

This method is optional method. It should be implemented when users wish 
to supply their own evaluation of the Jacobian. If the user does not wish to supply 
their own Jacobian, they should implement this method so that it throws an error if 
it is called. This method should update the Jac operator so that subsequent Epe-
tra_Operator::Apply() calls on that operator correspond to the Jacobian at the current 
solution vector $x$.

3. A method which fills a preconditioner matrix, whose syntax is

computePrecMatrix(const Epetra_Vector & x, 
   Epetra_RowMatrix & M)

It should only contain an estimate of the Jacobian. If users do not wish to supply their 
own Preconditioning matrix, they should implement this method such that if called, it 
will throw an error.

4. A method to apply the user’s defined preconditioner. The syntax is

computePreconditioner(const Epetra_Vector & x, Epetra_Operator & M)

The method should compute a preconditioner based upon the solution vector $x$ and store 
it in the Epetra_Operator $M$. Subsequent calls to the Epetra_Operator::Apply method 
will apply this user supplied preconditioner to epetra vectors.

Then, the user can construct a NOX::Epetra::Group, which contains information about 
the solution technique. All constructors require:

- A parameter list for printing output and for input options, defined as NOX::Parameter::List.
- An initial guess for the solution (stored in an Epetra_Vector object);
- an operator for the Jacobian and (optionally) and operator for the preconditioning 
  phase. Users can write their own operators. In particular, the Jacobian can be defined 
  by the user as an Epetra_Operator,

Epetra_Operator & J = UserProblem.getJacobian(),
created as a NOX matrix-free operator,

\[
\text{NOX::Epetra::MatrixFree} \& J = \text{MatrixFree(userDefinedInterface, solutionVec)},
\]

or created by NOX using a finite differencing,

\[
\text{NOX::Epetra::FiniteDifference} \& J = \text{FIXME...}
\]

At this point, users have to create an instantiation of the \text{NOX::Epetra::Interface} derived object,

\text{UserInterface interface(...)},

and finally construct the group,

\text{NOX::Epetra::Group} group(printParams, lsParams, interface).

### 14.3.1 A Simple Nonlinear Problem

As an example, define \( F : \mathbb{R}^2 \to \mathbb{R}^2 \) by

\[
F(x) = \left( \frac{x_1^2 + x_2^2 - 1}{x_2 - x_1^2} \right).
\]

With this choice of \( F \), the exact solutions of (14.1) are the intersections of the unity circle and the parabola \( x_2 - x_1^2 \). Simple algebra shows that one solution lies in the first quadrant, and has coordinates

\[
\alpha = \left( \frac{\sqrt{5} - 1}{2}, \frac{\sqrt{5} - 1}{2} \right),
\]

the other being the reflection of \( \alpha \) among the \( x_2 \) axis.

Code didasko/examples/nox/ex1.cpp applies the Newton method to this problem, with \( x_0 = (0.5, 0.5) \) as a starting solution. The output is approximatively as follows:

```
[msala:nox]> mpirun -np 1 ./ex1.exe
*************************************************
-- Nonlinear Solver Step 0 --
 f = 5.590e-01 step = 0.000e+00 dx = 0.000e+00
*************************************************

*************************************************
-- Nonlinear Solver Step 1 --
 f = 2.102e-01 step = 1.000e+00 dx = 3.953e-01
*************************************************

*************************************************
-- Nonlinear Solver Step 2 --
 f = 1.009e-02 step = 1.000e+00 dx = 8.461e-02
```
14.3. Introducing NOX in an Existing Code

**********************************************************************
Nonlinear Solver Step 3 --
f = 2.877e-05 step = 1.000e+00 dx = 4.510e-03 (Converged!)
**********************************************************************

-- Final Status Test Results --
Converged....OR Combination ->
  Converged....F-Norm = 2.034e-05 < 2.530e-04
  (Length-Scaled Two-Norm, Relative Tolerance)
??...........Number of Iterations = -1 < 20
**********************************************************************

-- Parameter List From Solver --
Direction ->
Method = "Newton" [default]
Newton ->
  Linear Solver ->
    Max Iterations = 400 [default]
  Output ->
    Achieved Tolerance = 8.6e-17 [unused]
    Number of Linear Iterations = 2 [unused]
    Total Number of Linear Iterations = 6 [unused]
    Tolerance = 1e-10 [default]
    Rescue Bad Newton Solve = true [default]
Line Search ->
Method = "More’-Thuente"
More’-Thuente ->
  Curvature Condition = 1 [default]
  Default Step = 1 [default]
  Interval Width = 1e-15 [default]
  Max Iters = 20 [default]
  Maximum Step = 1e+06 [default]
  Minimum Step = 1e-12 [default]
  Optimize Slope Calculation = false [default]
  Recovery Step = 1 [default]
  Recovery Step Type = "Constant" [default]
  Sufficient Decrease = 0.0001 [default]
  Sufficient Decrease Condition = "Armijo-Goldstein" [default]
Output ->
  Total Number of Failed Line Searches = 0 [unused]
  Total Number of Line Search Calls = 3 [unused]
  Total Number of Line Search Inner Iterations = 0 [unused]
  Total Number of Non-trivial Line Searches = 0 [unused]
Nonlinear Solver = "Line Search Based"
Output ->
    2-Norm of Residual = 2.88e-05  [unused]
    Nonlinear Iterations = 3  [unused]
Printing ->
    MyPID = 0  [default]
    Output Information = 2
    Output Precision = 3  [default]
    Output Processor = 0  [default]
Computed solution :
Epetra::Vector
    MyPID   GID   Value
    0       0     0.786
    0       1     0.618
Exact solution :
Epetra::Vector
    MyPID   GID   Value
    0       0     0.786
    0       1     0.618

14.4  A 2D Nonlinear PDE

In this Section, we consider the solution of the following nonlinear PDE problem:

\[
\begin{align*}
-\Delta u + \lambda e^u &= 0 \quad \text{in } \Omega = (0, 1) \times (0, 1) \\
u &= 0 \quad \text{on } \partial \Omega.
\end{align*}
\]  

(14.3)

For the sake of simplicity, we use a finite difference scheme on a Cartesian grid, with constant mesh sizes $h_x$ and $h_y$. Using standard procedures, the discrete equation at node $(i,j)$ reads

\[
-\frac{u_{i-1,j} + 2u_{i,j} - u_{i+1,j}}{h_x^2} + \frac{-u_{i,j-1} + 2u_{i,j} - u_{i,j+1}}{h_y^2} - \lambda e^{u_{i,j}} = 0.
\]

In example didasko/examples/nox/ex2.cpp, we build the Jacobian matrix as an Epetra_CrsMatrix, and we use NOX to solve problem (14.3) for a given value of $\lambda$. The example shows how to use NOX for more complex cases. The code defines a class, here called PDEProblem, which contains two main methods: One to compute $F(x)$ for a given $x$, and the other to update the entries of the Jacobian matrix. The class contains all the problem definitions (here, the number of nodes along the x-axis and the y-axis and the value of $\lambda$). In more complex cases, a similar class may have enough information to compute, for instance, the entries of $J$ using a finite-element approximation of the PDE problem.

The interface to NOX, here called SimpleProblemInterface, accepts a PDEProblem as a constructor,

SimpleProblemInterface Interface(&Problem);

Once a NOX::Epetra::Interface object has been defined, the procedure is almost identical to that of the previous Section.
14.5 Jacobian-free Methods

In Section 14.4, the entries of the Jacobian matrix have been explicitly coded. Sometimes, it is not always possible nor convenient to compute the exact entries of $J$. For those cases, NOX can automatically build Jacobian matrices based on finite difference approximation, that is,

$$J_{i,j} = \frac{F_i(u + h_j e_j) - F_i(x)}{h_j},$$

where $e_j$ is the $j$-unity vector. Sophisticated schemes are provided by NOX, to reduce the number of function evaluations.

14.6 Concluding Remarks on NOX

The documentation of NOX can be found in [KP04].

A library of continuation classes, called LOCA [SBRP+01, SLPS02], is included in the NOX distribution. LOCA is a generic continuation and bifurcation analysis package, designed for large-scalr applications. The algorithms are designed with minimal interface requirements over that needed for a Newton method to read an equilibrium solution. LOCA is built upon the NOX package. LOCA provided functionalities for single parameter continuation and multiple continuation. Also, LOCA provides a stepper class that repeatedly class the NOX nonlinear solver to compute points along a continuation curve. We will not cover LOCAL in this tutorial. The interested reader is referred to the LOCA documentation.
In order to get good parallel performance, the data distribution (map) is important. Poor data distribution can both lead to high communication among processes and also load imbalance, that is, some processes have more work than others.

Trilinos provides two packages for partitioning and load balancing: Zoltan and Isorropia. While Zoltan has a generic, data-structure-neutral interface, Isorropia provides Epetra interfaces to Zoltan that are more convenient for many Trilinos users.

15.1 Background

In parallel linear algebra applications, a critical part is to distribute the matrices among processes (processors). The vectors are often distributed to conform with the appropriate matrices, though not always. Matrices can be partitioned either along rows, columns, or by a 2-dimensional block decomposition. We limit our discussion to 1-dimensional data distributions, in particular, row distributions (which are best supported in Epetra). In this case, partitioning dense matrices is easy. For a matrix with \( n \) rows and with \( p \) processes, simply give each process \( n/p \) rows. For sparse matrices, the situation is more complicated. To achieve load-balance, one may wish that each process obtains approximately the same number of rows, or alternatively, similar number of nonzero entries. Additionally, the communication cost when applying the matrix should be small. Specifically, for iterative solvers, the communication cost in a matrix-vector product should be minimized.

A common abstraction of this problem is graph partitioning. This model assumes the matrix is symmetric, so the sparsity pattern of the matrix can be represented by an undirected graph. The graph partitioning problem is to partition the vertices into \( p \) sets such that the number of edges between sets are minimized. The number of cut edges approximates the communication cost in the parallel computation. Although graph partitioning is NP-hard to solve exactly, there are several fast algorithms that work well in practice. Zoltan provides a common interface to graph partitioners (and other algorithms). At present, the most widely used software for graph partitioning, are the METIS and ParMETIS [Kar, KK99] packages from University of Minnesota.

Recently, it has been shown [CA99] that hypergraph partitioning is a more accurate model for parallel matrix-vector communication cost. A parallel hypergraph partitioner is available in Zoltan [DBH+06, CBD+07] An advantage of hypergraph partitioning is that it also applies to rectangular matrices, not just square matrices.
15.2 Partitioning Methods

Zoltan and Isorropia currently support two types of partitioning methods: geometric and graph/hypergraph. The geometric methods are convenient if you have a vector of points in space, for example particles, or mesh points. The partitioner will then partition these points such that points that are close in Euclidean space will be assigned to the same or a nearby process.

The graph/hypergraph methods do not use geometry but rather rely on connectivity information, e.g., the sparsity pattern of a matrix. Multilevel algorithms for (hyper-)graph partitioning give high-quality partitionings, but take longer to compute than geometric methods.

15.3 Isorropia

Isorropia is the preferred partitioning package for most Trilinos users since it supports Epetra. The actual partitioning is done by calling Zoltan, so Zoltan is a required dependency.

Isorropia provides a simple interface for new users: createBalancedCopy. The input is an Epetra distributed data object (matrix, vector), and the output is a copy of the object (matrix, vector) but with a different (better) map (distribution):

```cpp
Epetra_CrsMatrix *A; // Original matrix
Epetra_CrsMatrix *bal_A; // Balanced matrix
bal_A = Isorropia::createBalancedCopy(A);
```

Note that the user is responsible for deallocating balA after use. The createBalancedCopy interface is rather limited and mainly intended for beginners. The full-fledged (primary) API contains the following three classes:

1. Partitioner
2. Redistributor
3. CostDescriber

The Partitioner performs the partitioning, but does not move any data. The Redistributor takes user data and a Partitioner as input, and redistributes the user data. The CostDescriber is optional and can be used to provide costs (weights) to the Partitioner for problem-specific partitioning.

Note that the primary API uses Teuchos::RCP reference-counted pointers for safer memory management.

```cpp
Epetra_CrsMatrix A;
Partitioner part(Teuchos::rcp(A));
Redistributor rd(Teuchos::rcp(part));
Teuchos::RCP<Epetra_CrsMatrix> bal_A = rd.redistribute(A);
```

The default partitioning method in Isorropia is hypergraph partitioning for a sparse matrix, and RCB for a vector.

Isorropia only supports a small number of parameters. The parameters are case insensitive. Advanced users who wish to specify detailed options should use Zoltan parameters. Isorropia supports a parameter sublist Zoltan, and these parameters are passed on to Zoltan.
15.4 Zoltan

Zoltan is a general-purpose package for parallel data management, including partitioning and load balancing [DBH+02]. Zoltan was developed independently of Trilinos, and does not depend on any other Trilinos packages. Thus, it can be built and used as a stand-alone library, if desired. Zoltan was written in C but also provides C++ and Fortran90 wrapper interfaces.

Zoltan currently provides all the partitioning methods in Isorropia. Zoltan also supports several third-party libraries, including the popular graph partitioners ParMetis and Scotch. To build Zoltan with ParMetis, you need to turn on `TPL_ENABLE_parmetis` in Cmake. Zoltan TPLs can also be used via Isorropia.
Templated Distributed Linear Algebra Objects with Tpetra

Marzio Sala

Tpetra is a package of classes for the construction and use of serial and distributed templated linear algebra objects, such as vectors, multi-vectors, and matrices. Tpetra is a direct descendant of Epetra, and as such it implements the Petra model. It provides capabilities that are very close to that of Epetra, but in a fully templated fashion. This Chapter will:

• Briefly introduce the concepts behind the design of Tpetra and templated programming (in Section 16.1);
• Presents the basic classes (in Section 16.2);
• Shows how to define and use vectors (in Section 16.4);
• Describes the matrix format (in Section 16.5).

This chapter only gives a broad overview of the Tpetra project; more details can be found in [SH05b] and on the Tpetra web page.

16.1 Introduction

The Epetra package, described in the previous Chapters, has proven to be very successful, robust, portable, and efficient. Its only drawback is that it only works with double-precision real values and integers. It cannot be used with complex or arbitrary-precision data, for example. Therefore, Epetra developers decided to create a new package, with most of Epetra’s capabilities, and a full support for templated programming as well. Two major types are used in Tpetra: the ordinal type and the scalar type\(^1\) As the name suggests, the OrdinalType is used to store information on how many of something is available. For example, this is the datatype for element IDs, or as a counting type. In Epetra, the OrdinalType is always int. In Tpetra, it will most likely be an int or long. However, it could be any type that is mathematically countable. ScalarType is the type of the actual data. In Epetra, this is always double; in Tpetra, it can for example be complex<double>, or almost any other type. For example, a small $3 \times 3$ matrix can be a valid type.

\(^1\)Actually Tpetra is based on two other types, but most users will only make use of ordinal and scalar types. Please consult the Tpetra manual for more details.
**Remark 23.** Changing Epetra to support programming involved quite more than simply replace all int and double instances with OrdinalType and ScalarType; in fact, almost all the Tpetra code is entirely new. However, since Tpetra developers reused many design and patterns from Epetra, it will be relatively easy for Epetra users to switch to Tpetra.

To use Tpetra, one has to know something about Teuchos; see Chapter 10 for an overview of this package. Although internally used for several tasks (the smart pointers, the BLAS kernels, and the flops-counting of all objects), Tpetra users mostly have to care only about the traits mechanism.

Traits are an important component of any templated code. Using arbitrary data types requires some care in code writing. Since the actual type is not known, it is very important not to make assumptions like someVar = 5.0, which may not works for a given data type. (For example, this operation may not be defined for a type representing a $3 \times 3$ matrix.) To solve this problem, a design pattern known as traits is adopted. Tpetra takes advantage of two Teuchos classes, ScalarTraits and OrdinalTraits, to define traits. Two traits are of particular importance: one and zero. Zero is the mathematical zero, that is, the value such that $x \times 0 = 0 \forall x$. One is the unity or identity, that is the value such that $x \times 1 = x \forall x$. As long as a type as these two traits defined, and defines the basic operations such as $=, +, -, \times, /$, then this type can be used as OrdinalType or ScalarType.

### 16.2 A Basic Tpetra Code

We now introduce the basic components of a basic Tpetra code. First, we need to include the header files:

```cpp
#include "Tpetra_ConfigDefs.hpp"
#include "Tpetra_MpiPlatform.hpp"
#include "Tpetra_MpiComm.hpp"
#endif
#include "Teuchos_ScalarTraits.hpp"
```

Then, we define (using a typedef statement) the ordinal type and scalar type:

```cpp
typedef int OrdinalType;
typedef double ScalarType;
```

Since the generic assignment `myVar = 0` or `myVar = 1` may not work with a given ordinal or scalar type, we must define four variables, containing the value zero and one for both ordinal and scalar type; Teuchos::ScalarTraits is used for this operation:

```cpp
OrdinalType const OrdinalZero = Teuchos::ScalarTraits<OrdinalType>::zero();
OrdinalType const OrdinalOne = Teuchos::ScalarTraits<OrdinalType>::one();
ScalarType const ScalarZero = Teuchos::ScalarTraits<ScalarType>::zero();
ScalarType const ScalarOne = Teuchos::ScalarTraits<ScalarType>::one();
```
At this point, exactly as it was done with Epetra, one has to define a communicator, either serial or parallel:

```cpp
#ifdef HAVE_MPI
MPI_Init(&argc,&argv);
Tpetra::MpiComm<OrdinalType, ScalarType> Comm(MPI_COMM_WORLD);
#else
Tpetra::SerialComm<OrdinalType, ScalarType> Comm;
#endif
```

Parallel communication is done using class Tpetra::Comm, whose functionalities are very close to that of the Epetra_Comm class. Both classes provide an insulating layer between the actual communications library used and the rest of Tpetra; this makes Epetra and Tpetra codes accessible in serial, MPI or shared memory environments.

The most important difference between Epetra and Tpetra communicator class is in the nomenclature. Epetra uses the term processor, while Tpetra adopts the more general image. This is because often the same processor runs multiple MPI jobs, or vice-versa, several processors may be running a single MPI job. Therefore, the term memory image defines a running copy of the code.

Tpetra::Comm provides the same collective communication operations that Epetra::Comm does. In addition, it provides several point-to-point operations that were not part of Epetra::Comm. Among these new operations, one has for example blocking sends and blocking receives.

An example of usage is as follows. First, let us define two working arrays, of type ScalarType and size 2:

```cpp
OrdinalType size = 2;
vector<ScalarType> V(size), W(size);
```

The values of V can be broadcasted from image 0 with instruction

```cpp
int RootImage = 0;
Comm.broadcast(&V[0], size, RootImage);
```

or use `sumAll()` as

```cpp
Comm.maxAll(&V[0], &W[0], size);
```

Example `didasko/examples/tpetra/ex1.cpp` shows the usage of Tpetra::Comm objects.

**Remark 24.** Note that using MPI with Tpetra is not as simple as it is with Epetra. Since Epetra is based on `int` and `double` data types, one simply has to transmit data using `MPI_INT` or `MPI_DOUBLE`. In Tpetra, instead, the data type is in principle not known: a data type may not be contiguous in memory, or be a user-defined class containing pointers or subclasses. The solution adopted by Tpetra developers involves traits. In this tutorial we suppose that the scalar type is one among `float`, `double` or `complex<double>`; for more involved types please contact the Tpetra developers.

Tpetra needs an additional class that does have no equivalence in the Epetra world: the Tpetra::Platform class. This class is responsible for creating Comm instances. This class
is required by the templated approach of Tpetra. In fact, virtual member functions and
templates are mutually exclusive: virtual member functions cannot be templated. Or, more
precisely, they can be templated at the class level, but not at the function level. More details
on this subject can be found in [SH05b].

Generally, two platforms must be created: one for OrdinalType’s, and the other for Scalar-
Type’s. The general syntax is:

```c++
#ifdef HAVE_MPI
const Tpetra::MpiPlatform <OrdinalType, OrdinalType>
    platformE(MPI_COMM_WORLD);
const Tpetra::MpiPlatform <OrdinalType, ScalarType>
    platformV(MPI_COMM_WORLD);
#else
const Tpetra::SerialPlatform <OrdinalType, OrdinalType> platformE;
const Tpetra::SerialPlatform <OrdinalType, ScalarType> platformV;
#endif
```

At this point, one can insert any of the snippets later presented. Before quitting `main()`,
one has to close MPI:

```c++
#ifdef HAVE_MPI
MPI_Finalize() ;
#endif
return(EXIT_SUCCESS);
```

### 16.3 Spaces

In the Tpetra lingo, a space is a set of elements, distributed across the available images. The
Epetra equivalence are the Epetra_Map and Epetra_BlockMap.

Tpetra has three space classes: Tpetra::ElementSpace, Tpetra::ElementBlockSpace, and
the Tpetra::VectorSpace.

Tpetra::ElementSpace objects are defined to have elements of size one, while variable
element sizes are supported by the Tpetra::BlockElementSpace class. Tpetra::VectorSpace,
instead, serves two purposes. In addition to creating Tpetra::Vectors, it acts as an “insulat-
ing” class between Tpetra::Vector’s and ElementSpace and BlockElementSpace. Through this
mechanism, Tpetra::Vector’s can be created and manipulated using one nonambiguous set of
vector indices, regardless of if it uses an Tpetra::ElementSpace or a Tpetra::BlockElementSpace
for distribution.

The distribution of elements in a Tpetra::ElementSpace or Tpetra::BlockElementSpace
can be arbitrary. Perhaps the simplest way to create a space is to specify the global element
of elements:

```c++
OrdinalType length = 10;
OrdinalType indexBase = OrdinalZero;
Tpetra::ElementSpace<OrdinalType>
    eElementSpace(length, indexBase, platformE);
Tpetra::VectorSpace<OrdinalType, ScalarType>
    vectorSpace(elementSpace, platformV);
```
More involved constructors exist as well, and they are not covered here because their usage is basically equivalent to that of Epetra::Map’s.

Some methods of Tpetra::ElementSpace are:

- `getNumGlobalElements()` returns the number of elements in this calling object;
- `getNumMyElements()` returns the number of elements belonging to the calling image;
- `getLID(OrdinalType GID)` returns the local ID of the global ID passed in, or throws exception 1 if not found on the calling image;
- `getGID(OrdinalType LID)` returns the global ID of the local ID passed in, or throws exception 2 if not found on the calling image.
- operators `==` and `!=` can be used to compare two spaces.

Tpetra::VectorSpace’s have similar query methods:

- `getNumGlobalEntries()` returns the number of entries in the calling object;
- `getNumMyEntries()` return the number of entries belonging to the calling image.
- `getLocalIndex(OrdinalType globalIndex)` returns the local index for a given global index.
- `OrdinalType getGlobalIndex(OrdinalType localIndex)` returns the global index for a given local index.
- `isMyLocalIndex (OrdinalType localIndex)` returns true if the local index value passed in is found on the calling image, returns false if it doesn’t.
- `isMyGlobalIndex(OrdinalType globalIndex)` returns true if the global index value passed in is found the calling image, returns false if it doesn’t.

## 16.4 Creating and Using Vectors

The most basic unit of linear algebra is the vector, implemented by class Tpetra::Vector. Vectors can be used to perform many mathematical operations, such as scaling, norms, dot products, and element-wise multiplies. It is also be used in conjunction with anothe Tpetra object, Tpetra::CisMatrix, described in the Section 16.5.

Tpetra::Vector is templated on ScalarType for the vector entries, and on OrdinalType for the vector indices. A VectorSpace object (described in Section 16.2) is needed for all Vector objects.

Note that for most of the mathematical methods that set this to the result of an operation on vectors passed as parameters, the this vector can be used as one of the parameters (unless otherwise specified).

Given a vectorSpace, a Teaptra::Vector is created as

```
Tpetra::Vector<OrdinalType, ScalarType> v1(vectorSpace);
```

Several methods are available to define the elements of a vector. To set all the elements to the same value, one can do:

```
v1.setAllToScalar(ScalarOne);
```
Otherwise, by using the [] operator,

\[ \text{OrdinalType MyLength = elementSpace.getNumMyElements();} \]

for (OrdinalType i = OrdinalZero ; i < MyLength ; ++i)
\[ v1[i] = (ScalarType)(10 + i); \]

Note that all Tpetra::Vector entries can only be accessed through their local index values. Global index values can be converted to local indices by using the VectorSpace::getLocalIndex method.

Another way is to extract a view of the internally stored data array (again, or all locally owned values):

\[ \text{OrdinalType NumMyEntries = v1.getNumMyEntries();} \]

\[ \text{vector<ScalarType*> data(NumMyEntries);} \]
\[ v1.extractView(&data[0]); \]

for (OrdinalType i = OrdinalZero ; i < MyLength ; ++i)
\[ *(data[i]) = (ScalarType)(100 + i); \]

Several methods are associated with a Tpetra::Vector object. For example, 
\[ \text{scale(ScalarType scalarThis)} \] scales the current values of the vector, \[ \text{norm1() returns the 1-norm, norm2() returns the 2-norm, normInf() the } \infty \text{ norm, minValue() computes the minimum value of the vector, maxValue() the maximum value, and meanValue() returns the mean (average) value. Update operations can be performed using method update(); for example, } \]
\[ x = \alpha \times x + \beta \times y \] is computed by
\[ x.update(beta, y, alpha); \]
while \[ x = \alpha \times x + \beta \times y + \gamma z \] by
\[ x.update(beta, y, gamma, y, alpha); \]
Finally, vectors can be printed as \texttt{cout} \texttt{<<} \texttt{v1}.

### 16.5 Creating and Using Matrices

In Tpetra, matrices are defined by the Tpetra::CisMatrix class. This class is meant for matrices that are either row- or column-oriented; this property is set in the constructor.

Constructing Tpetra::CisMatrix objects is a multi-step process. The basic steps are as follows:

1. Create a Tpetra::CisMatrix instance, using one of the constructors.
2. Enter values using the submitEntries methods.
3. Complete construction by calling fillComplete.

We now show these steps for the creation of a diagonal, distributed matrix. We suppose that elementSpace and vectorSpace are valid Tpetra::ElementSpace and Tpetra::VectorSpace, respectively.

First, if not already available, we need to extract the list of locally owned D’s from elementSpace,
16.5. Creating and Using Matrices

OrdinalType NumMyElements = elementSpace.getNumMyElements();
vector<OrdinalType> MyGlobalElements = elementSpace.getMyGlobalElements();

Then, we instantiate a Tpetra::CisMatrix object and we insert one element at-a-time:

Tpetra::CisMatrix<OrdinalType,ScalarType> matrix(vectorSpace);

for (OrdinalType LID = OrdinalZero ; LID < NumMyElements ; ++LID)
{
    OrdinalType GID = MyGlobalElements[LID];
    // add the diagonal element of value 'GID'
    matrix.submitEntry(Tpetra::Insert, GID, (ScalarType)GID, GID);
}

Method submitEntry requires the CombineMode (Tpetra::Insert, Tpetra::Add, Tpetra::Replace), the row (or column) ID, the value, and the column (or row) ID of the element to be inserted. Method submitEntries() can also be used to submit multiple entries with just one function call. The last step is performed by calling matrix.fillComplete().

Prior to calling fillComplete, data is stored in a format optimized for easy and efficient insertions/deletions/modifications. It is not a very efficient form for doing matvec operations though. After calling fillComplete, the data is transformed into a format optimized for matvec operations. It is not very efficient at modifying data though. So after fillComplete() has been called, the matrix should be viewed as const, and cannot be modified. Trying to do so will result in an exception being thrown.

Once freezed, a matrix can be queried for global number of nonzeros, rows, columns, diagonals, and maximum number of entries per row/column are returned with methods getNumGlobalNonzeros(), getNumGlobalRows(), getNumGlobalCols(), getNumGlobalDiagonals(), getGlobalMaxNumEntries(), respectively. The corresponding local information are returned by similar methods with My instead of Global. Other queries only have a global meaning: isRowOriented(), isFillCompleted(), normOne(), normInf().

To apply the matrix to a vector x and get the result in y, one can simply write matrix.apply(x, y, UseHermitian), where UseHermitian is a boolean variable that can be used to multiply with $A$ or $A^H$.

CisMatrix stores data using Compressed Index Space Storage. This is a generalization of Compressed Row Storage, but allows the matrix to be either row-oriented or column oriented. Accordingly, CisMatrix refers to data using a generalized vocabulary. The two main terms are the primary distribution and the secondary distribution:

In a row-oriented matrix, the primary distribution refers to rows, and the secondary distribution refers to columns. In a column-oriented matrix, the primary distribution refers to columns, and the secondary distribution refers to rows.

These distributions are specified by Tpetra::VectorSpace objects. If both the primary and secondary distributions are specified at construction, information about the secondary distribution is available from then on. But if only the primary distribution is specified at construction, then CisMatrix will analyze the structure of the matrix and generate a VectorSpace that matches the distribution found in the matrix. Note that this is done when fillComplete is called, and so information about the secondary distribution is not available prior to that.
Other two distributions are indeed used. The domain distribution specifies the distribution of the vector to which the matrix will be applied; while the range distribution specifies the distribution of the result vector \( y = Ax \).

### 16.6 Concluding Remarks

All Tpetra classes that represent a linear algebra object inherit from `Teuchos::CompObject`. The flop count of a CompObject represents the number of floating-point operations that have occurred on the calling image – it is not a global counter.

Tpetra also depends on Kokkos for serial kernels, which are now separated from the surrounding code, so that more optimized kernels can be developed and used without affecting Tpetra. Kokkos routines as purely serial; therefore any redistributions needed before or after the computation must be handled by Tpetra. Kokkos is a collection of the handful of sparse and dense kernels that determine the much of the performance for preconditioned Krylov methods. In particular, it contains function class for sparse matrix vector multiplication and triangular solves, and also for dense kernels that are not part of the standard BLAS.

The classes in this package are written in a way that they can be easily customized via inheritance, replacing only the small sections of code that are unique for a given platform or architectures. In this way we hope to provide optimal implementations of these algorithms for a broad set of platforms from scalar microprocessors to vector multiprocessors.

Kokkos is not intended as a user package, but to be incorporated into other packages that need high performance kernels. As such, it is not covered in this Tutorial.
Bibliography


