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Trilinos 3.1 Tutorial

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Abstract

This document introduces the use of Trilinos, version 3.1. Trilinos has been written to support, in a rigorous manner, the solver needs of the engineering and scientific applications at Sandia National Laboratories.

Aim of this manuscript is to present the basic features of some of the Trilinos packages. The presented material includes the definition of distributed matrices and vectors with Epetra, the iterative solution of linear system with AztecOO, incomplete factorizations with IFPACK, multilevel methods with ML, direct solution of linear system with Amesos, and iterative solution of nonlinear systems with NOX. With the help of several examples, some of the most important classes and methods are detailed to the unexperienced user. For the most majority, each example is largely commented throughout the text. Other comments can be found in the source of each example.

This document is a companion to the Trilinos User's Guide [10] and Trilinos Development Guides [11, 12]. Also, the documentation included in each of the Trilinos' packages is of fundamental importance.
Acknowledgments

The authors would like to acknowledge the support of the ASCI and LDRD programs that funded development of Trilinos.
Trilinos 3.1 Tutorial

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1 Introduction

The Trilinos Project is an effort to facilitate the design, development, integration and ongoing support of mathematical software libraries. Goal of the Trilinos Project is develop parallel solver algorithms and libraries within an object-oriented software framework for the solution of large-scale, complex multiphysics engineering and scientific applications. The emphasis is 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 abstract interfaces.

1.1 Getting Started with Trilinos

The Trilinos Project uses a two-level software structure designed around collections of packages. A Trilinos package is an integral unit, usually developed to solve a specific task, by a (relatively) small group of expert of the field. Packages exist underneath the Trilinos top level, which provides a common look-and-feel. Each package has its own structure, documentation and set of examples. In principle, Trilinos packages can live independently. However, each package is even more valuable when combined with other Trilinos packages.

Trilinos is a large software project, and currently about twenty packages are included. Fully understanding all the functionalities of the Trilinos packages requires time. The entire set of packages covers a wide range of numerical methods for large scale computing. Some packages are focused on the development of computational schemes, like for instance the solution of linear and nonlinear systems, to the definition of parallel preconditioners for Krylov methods, eigenvalue computation. Other packages are more focused on implementation issues (like definition of matrices and vectors, abstract classes for linear operators). The first Chapters of this tutorial will be focused on implementation issues, while the last Chapters will have a more "mathematical" taste.

Each package offers sophisticated features, that cannot be "unleashed" at a very first usage. For each package, we will outline only the basic features, and we refer to the documentation of each package for a more involved usage. Our goal is to present enough material so that the reader can successfully use the described packages. In fact, for new users, it is neither easy, nor necessary, to manage all the Trilinos functionalities. At the beginning, it is more important for them to understand how to manage the basic classes, such as vector, matrix and linear system classes. However, it is clear that for a fine-tuning, the reader will have to look through each package's documentation and examples.
Although all packages have the same importance in the Trilinos structure, a typical user will probably — at least at the beginning — make use of the following packages:

- **Epetra.** This 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 of them can “speak” other languages). Each Trilinos package is able to accept in input Epetra objects. This allows powerful combinations among the various Trilinos functionalities.

- **AztecOO.** This is a linear solve package based on preconditioned Krylov methods. It supports all the Aztec interfaces and functionality, but also provides significant new functionality.

- **IFPACK.** This is a package to perform various incomplete factorizations, and it is here used in conjunction with AztecOO.

- **ML.** This is an algebraic multilevel preconditioner package, which provided scalable preconditioning capabilities for a variety of problem classes. It is here used in conjunction with AztecOO.

- **Amesos.** This package provides a common interface to various direct solvers (generally available outside the Trilinos framework), both sequential and parallel.

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

- **Triutils.** This is a collection of various utilities, that can be extremely useful in some phases of software development.

Table 1 gives a partial overview of what can be accomplished using Trilinos.

This tutorial is divided into 10 chapters:

- Chapter 2 describes the Epetra_Vector class;
- Chapter 3 introduces the Epetra_Matrix class;
- Chapter 4 briefly describes some other Epetra classes;
- Chapter 5 shows how to solve linear systems with AztecOO;
- Chapter 6 presents the basic usage of IFPACK;
<table>
<thead>
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<tr>
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<td>solve a linear system with preconditioned Krylov accelerators, like CG, GMRES, Bi-CGSTAB, TFQMR:</td>
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<td>interface with various direct solvers, as UMFPACK, MUMPS, SuperLU and others:</td>
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<td>Computation of eigenvalue of large, sparse matrices:</td>
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<td>Templated interface to BLAS and LAPACK, arbitrary-precision arithmetic, parameter lists:</td>
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</tr>
<tr>
<td>Definition of abstract interfaces to vectors, linear operators, and solvers:</td>
<td>TSF*, TSFCore*, TSFExtended*</td>
</tr>
</tbody>
</table>

**Table 1.** Partial overview of what can be done with Trilinos. *: not covered in this tutorial.
• Chapter 7 introduces multilevel preconditioners based on ML;
• Chapter 8 introduces the Amesos package;
• Chapter 9 outlines the main features of the Trilinos nonlinear solver package, NOX.
• Chapter 10 presents some tools provided with the Triutils package.

Remark 1. As already pointed out, Epetra objects are meant to be the “common language” spoken by all the Trilinos packages, and therefore the new user must become familiar with those objects. Therefore we suggest to read Chapters 2-4 before considering other Trilinos packages. Also, Chapter 5 should be read before Chapters 6 and 7 (even if both IFPACK and ML can be compiled and run without AztecOO).

This tutorial assume a basic background in numerical methods for PDEs, and in iterative linear and nonlinear solvers. Although not strictly necessary, the reader is suppose to have a certain familiarity with distributed memory computing and, to a minor extent, with MPI.

Note that this tutorial is not a substitute of 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 [7]. Developers should also consider the Trilinos Developers’ Guide, which addresses many topics, including the development tools used by Trilinos’ developers, and how to include a new package.

1.2 Installing Trilinos

To obtain Trilinos, please refer to the instructions reported at the following web site:

http://software.sandia.gov/Trilinos

Trilinos has been compiled on a variety of architectures, including Linux, Sun Solaris, SGI Irix, DEC, and many others. Trilinos has been designed to support parallel applications. However, it can be compiled and run on serial computer. Detailed comments on the installation, and an exhaustive list of FAQs, can be found at the web pages:

1Trilinos provides a variety of services to a developer wanting to integrate a package into Trilinos. They include Autoconf [1], Automake [2] and Libtool [3]. Those tools provide a robust, full-featured set of tools for building software across a broad set of platforms. Although these tools are not official standards, they are widely used. All existing Trilinos packages use Autoconf and Automake. Libtool support will be added in future releases.
Before using Trilinos, users might decide to set the environmental variables `TRILINOS_HOME`, indicating the full path of the Trilinos directory, `TRILINOS_LIB`, indicating the location of the compiled Trilinos library, and `TRILINOS_ARCH`, containing the architecture and the communicator currently used. For example, using the BASH shell, command lines of the form

```
export TRILINOS_HOME=/home/msala/Trilinos
export TRILINOS_ARCH=LINUX.MPI
export TRILINOS_LIB=${TRILINOS_HOME}/${TRILINOS_ARCH}
```

can be places in the users' `.bashrc` file.

Here, we briefly report the procedure one should follow in order to compile Trilinos as required by the examples reported in the following chapters 2-10. Suppose we want to compile under LINUX with MPI. The installation procedure can be are reported below. ($ indicates the shell prompt.)

```
$ cd ${TRILINOS_HOME}
$ mkdir ${TRILINOS_ARCH}
$ cd ${TRILINOS_ARCH}
$ ../configure --prefix="${TRILINOS_HOME}/${TRILINOS_ARCH}" \
  --enable-mpi --with-mpi-compilers \ 
  --enable-triutils --enable-aztecoo \ 
  --enable-ifpack \ 
  --enable-ml --enable-nox | tee configure_${TRILINOS_ARCH}.log
$ make | tee make_${TRILINOS_ARCH}.log
$ make install | tee make_install_${TRILINOS_ARCH}.log
```

**Remark 2.** All Trilinos packages can be build to run with or without MPI. If MPI is enabled (using `--enable-mpi`), the users must know the procedure for beginning MPI jobs on their computer system(s). In some cases, options must be set on the configure line to specify the location of MPI include files and libraries.

---

2Amesos can be more difficult to compile for the unexperienced user, as it required some information about the packages to interface. Suggestions about the configuration of Amesos are reported in Chapter 8. More details about the installation of Trilinos can be found in [10].
1.3 Compiling and Linking a program using Trilinos

In order to compile and link (part of) the Trilinos library, the user can decide to use a Makefile as reported below. This Makefile refers to one of the examples, reported in the NOX subdirectory of this tutorial.

```bash
1: TRILINOS_HOME = /home/msala/Trilinos/
2: TRILINOS_ARCH = LINUX_MPI
3: TRILINOS_LIB = $(TRILINOS_HOME)$(TRILINOS_ARCH)
4:
5: include $(TRILINOS_HOME)/build/makefile.$(TRILINOS_ARCH)
6:
7: MY_COMPILER_FLAGS = -DHAVE_CONFIG_H $(CXXFLAGS) -c -g\n8: -I$(TRILINOS_LIB)/include/
9:
10: MY_LINKER_FLAGS = $(LDFLAGS) $(TEST C OBJ) \n11: -L$(TRILINOS_LIB)/lib/ \n12: -lnoxepetra -lnox -lifpack \n13: -laiztecoo -lepeta -llapack -lblas $(ARCH_LIBS)
14:
15: exl: ex1.cpp
16: $(CXX) ex1.cpp $(MY_COMPILER_FLAGS)
17: $(LINKER) ex1.o $(MY_LINKER_FLAGS) -o ex1.exe
```

Line number have been reported for reader's convenience.

The lines 1-3 can be omitted, see Section 1.2. Line 5 includes basic definitions of Trilinos. (Note that, on some architectures, one may need to use gmake instead of make.) In line 7, the variable HAVE_CONFIG_H is defined. Linker flags of lines 10-13 defines the library to link (location of BLAS and LAPACK can change on different platforms). The variable ARCH_LIBS is defined in line 5.

To run the compiled example in a sequential environment, simply type

```
$ ./ex1.exe
```

In a MPI environment, the user might have to use an instruction of type

```
$ mpirun -np 2 ./ex1.exe
```
Please check the local MPI documentation for more details.

### 1.4 Copyright and Licensing of Trilinos

Trilinos is released under the Lesser GPL GNU License.

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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.

### 1.5 Programming Language Used in this Tutorial

Trilinos is written in C++ (for most packages), and in C. Some interfaces are provided to FORTRAN code (mainly BLAS and LAPACK routines). Even if a limited support is included for C programs (and a more limited for FORTRAN code), to unleash the full power of Trilinos we suggest to use C++. All the example programs contained in this tutorial will be in C++; some packages contains examples in C.
1.6 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,
title = "{Trilinos Developers Guide Part II: ASCI Software Quality Engineering Practices Version 1.0}"",
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}

These BiBTeX information can be downloaded from the web page

http://software.sandia.gov/Trilinos/citing.html
1.7 A Note on Directory Structure

Each Trilinos package is contained in the subdirectory

\${TRILINOS_HOME}/packages

The structure of all packages is quite similar (although not exactly equal). As a general line, source files are in

\${TRILINOS_HOME}/packages/<package-name>/src

Example files are reported in

\${TRILINOS_HOME}/packages/<package-name>/examples

and test files in

\${TRILINOS_HOME}/packages/<package-name>/test

The documentation is reported

\${TRILINOS_HOME}/packages/<package-name>/doc

Often, Trilinos developers use Doxygen\(^3\). For instance, to create the documentation for Epetra, we use can type

$ cd \${TRILINOS_HOME}/packages/epetra/doc
$ doxygen Doxyfile

and then browse it using an HTML reader, or compiling the \LaTeX{} file using

$ cd \${TRILINOS_HOME}/packages/epetra/doc/latex
$ make

\(^3\)Copyright ©1997-2003 by Dimitri van Heesch. More information can be found at the web address http://www.stack.nl/ dimitri/doxygen/.
1.8 List of Trilinos Developers

A list of the Trilinos’ developers, updated to December 2003, would include the following names (in alphabetical order):

2 Working with Epetra Vectors

Probably, the first mathematical entities defined by a numerical method is a vector. Within the Trilinos framework, vectors are usually constructed starting from Epetra Classes.

Epetra vectors can be used to store double values (like the solution of a PDE problem, the right-hand side of a linear system, or the nodal coordinates), as well as integer data values (such as a set of indexes).

Epetra vectors can be 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.

This Chapter will show some of the Trilinos capabilities to work with vectors. Vector classes can be used to perform common vector operations, as dot products, vector scalings and norms, or fill with constant or random values.

During the Chapter, the user be introduced to:

- The Epetra_Comm object (in Section 2.1);
- The Epetra_Map object (in Section 2.2);
- Creating and assembling Epetra vectors (in Sections 2.3 and 2.4);
- Redistributing vectors (in Section 2.5).

2.1 Epetra Communicator Objects

The Epetra_Comm class is an interface that encapsulates the general information and services needed for the other Epetra classes to run on a 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 implementations:
- Epetra_SerialComm (for serial executions);
- Epetra_MpiComm (for MPI distributed memory executions).

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

```c
#include "Epetra_config.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 argc, 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

```c
#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 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 as

```c
int NumProc = Comm.NumProc();
int MyPID = Comm.MyPID();
```
File \{TRILINOS\_HOME\}/doc/tutorial/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

Very often, various distributed objects such as matrices or vectors, have identical distribution of elements among the processes. This distribution of elements (or points) is here called a map, and its actual implementation within the Trilinos project is given by the Epetra\_Map class (or, more generally, by an Epetra\_BlockMap). Basically, the class handles the definition of:

- global number of elements (called NumGlobalPoints);
- the local number of elements (called NumMyPoints);
- the global numbering of all local nodes (an integer vector of size NumMyPoints, called MyGlobalElements).

There are essentially three ways to define an map. The easiest way is to specify the global number of elements:

Epetra\_Map Map(NumGlobalPoints, 0, Comm);

In this case, the constructor takes the global dimension of the vector (here indicated as NumGlobalPoints), the base index (0 for C or C++ arrays, 1 for FORTRAN arrays, but it can be any number), and an Epetra\_Comm object (introduced in Section 2.1). As a result, each process will be assigned a contiguous list of elements.

Another way to build the Epetra\_Comm object is to furnish the local number of elements:

Epetra\_Map Map(-1, NumMyPoints, 0, Comm);

This will create a vector of size $\sum_{i=0}^{NumProc-1} NumMyPoints$. Each process will get a contiguous set of elements. These two approached are coded in file \{TRILINOS\_HOME\}/doc/tutorial/epetra/ex2.cpp.
Another, more involved way, to create an Epetra_Map, is to specify on each process both the number of local elements, and the global numbering of each local element. To better explain this, let us consider the following code, in which a vector, of global dimension 5, is split among 2 processes p0 and p1. p0 owns nodes 0 and 4, while p1 nodes 1, 2, and 3.

```cpp
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);

The complete code is reported in `${TRILINOS_HOME}/doc/tutorial/epetra/ex3.cpp`.

A Map object can be queried for the global and local number of elements, using

```cpp
int NumGlobalElements = Map.NumGlobalElements();
int NumMyElements = Map.NumMyElements();
```

and for the global ID of local elements, using

```cpp
int * MyGlobalElements = Map.MyGlobalElements();
```

or, equivalently,

```cpp
int MyGlobalElements[NumMyElements];
Map.MyGlobalElements(MyGlobalElements);
```
The class Epetrahlap is derived from Epetra_BlockMap. This 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 this class. An example of use of block maps is reported in `${TRILINOS_HOME}/doc/tutorial/epetra/ex23.cpp`.

Note that different maps can coexist in the same part of the code. This allows the user to easily define vectors with different distributions (even for vectors of the same size). Two classes are provided to transfer data from one map to another. Those classes (Epetra_Import and Epetra_Export) are discussed in Section 2.5.

**Remark 3.** Most Epetra objects overload the `<` operator. For example, to visualize information about the Map, one can simply write

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

This Section has presented the construction of very basic map objects. However, map objects of very general form can be constructed. 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 `${TRILINOS_HOME}/doc/tutorial/epetra/ex20.cpp` and `${TRILINOS_HOME}/doc/tutorial/epetra/ex21.cpp` can be used to better understand the potentiality of Epetra Maps.

**Remark 4.** 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 or for parallel runs. 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).

To create a sequential vector containing `Length` elements, one can use the following command:
Epetra_SerialDenseVector x(Length);

Other constructors are available; check the Epetra Class Documentation.

The class Epetra_SerialDenseVector enables the construction and use of real-valued, double-precision dense vectors. The Epetra_SerialDenseVector class is intended to provide convenient vector notation but derives all significant functionality from Epetra_SerialDenseMatrix class. The vector can be filled using the [ ] or ( ) operators. Both methods return the specified element of the vector. However, using ( ), bounds, checking is enforced. Using using [ ], no bounds checking is done unless Epetra is compiled with EPETRA_ARRAY_BOUNDS_CHECK.

Remark 5. To construct replicated Epetra objects on distributed memory machines, the user may consider the class Epetra_LocalMap. This class allows the constructions of those replicated local objects and keeps information that describe the distribution.

File ${TRILINOS_HOME}/doc/tutorial/epetra/ex4.cpp shows some basic operations on dense vectors.

2.4 Creating and Assembling a Distributed Vector

To create a distributed vector, the first step is to define a map. (Actually, this is true for all distributed Epetra objects.) After that, an Epetra_Vector object can be constructed with an instruction of type

Epetra_Vector x(Map);

This constructor allocates space for the vector and set all the elements to zero. A copy constructor can be used as well:

Epetra_Vector y(x);

Alternatively, the user can pass a pointer to an array of double precision values:

Epetra_Vector x(Copy,Map,LocalValues);

Note the word Copy is input to the constructor. Epetra allows two data access modes:
1. **Copy mode**: Allocates memory and makes a copy of the user-provided data. In this case, the user data is not needed after construction;

2. **View mode**: Creates a “view” of the user’s data. In this case, the user data is required to remain untouched for the life of the vector (or modified carefully). It is worth noting that the View mode is very dangerous from a data hiding perspective. Therefore, users are strongly encouraged to develop code using Copy mode first and only use View mode in a secondary 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

   \[
   \text{Epetra\_Vector } z(\text{View}, \text{Map}, \text{double\_vector});
   \]

   where \text{double\_vector} is a pointer to the vector of doubles.

Regardless of how a vector has been created, one can use the \texttt{[]} operator to access a vector element:

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

where \texttt{i} is in the local index space.

Epetra also defines some functions to set vector elements in local or global index space. \texttt{ReplaceMyValues} or \texttt{SumIntoMyValues} will replace or sum values into a vector with a given indexed list of values, with indexes in the \textit{local} index space; \texttt{ReplaceGlobalValues} or \texttt{SumIntoGlobalValues} will replace or sum values into a vector with a given indexed list of values in the \textit{global} index space. It is important to note that a process cannot set a vector entries locally owner by another process. In other words, both global and local insert and replace functions refers to the part of a vector assigned to the calling process. Intra-process communications can be performed using Import and Export objects, covered in Section 2.5.

Another way is to put vector values in a user-provided array. For instance, one may have:

\[
\begin{align*}
\text{double } & *x\_values; \\
& x\_values = \text{new double}[\text{MyLength}]; \\
& x.\text{ExtractCopy}( x\_values ); \\
& \text{for( int } i=0 \text{; } i<\text{MyLength} \text{; ++i } ) x\_values[i] *= 10; \\
& \text{for( int } i=0 \text{; } i<\text{MyLength} \text{; ++i } ) \\
& \phantom{=}x.\text{ReplaceMyValues}( 1, 0, x\_values+i, \&i );
\end{align*}
\]
(File \${TRILINOS_HOME}/doc/tutorial/epetra/ex5.cpp reported the complete source.) It is important to note that 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. To that aim, one has to call

```cpp
double * pointer;
x.ExtractView( &pointer );
```

Now, modifying the values of pointer will affect the internal data of the Epetra_Vector x. An example of the use of ExtractView is reported in file \${TRILINOS_HOME}/doc/tutorial/epetra/ex5.cpp.

**Remark 6.** 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. The reader may look to the file \${TRILINOS_HOME}/doc/tutorial/epetra/ex7.cpp for an example of use of multi-vectors.

The user can also consider the function ResetView, which allows a (very) lightweight 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 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 \${TRILINOS_HOME}/doc/tutorial/epetra/ex8.cpp

It is possible to perform a certain number of operations on vector objects. Some of them are reported in Table 2. Example \${TRILINOS_HOME}/doc/tutorial/epetra/ex18.cpp works with some of the functions reported in the table.

### 2.5 Epetra_Import and Epetra_Export

Epetra_Import and Epetra_Export are two classes meant for efficient importing of off-processors elements. Epetra_Import and Epetra_Export are used to construct a communication plan that can be called repeatedly by computational classes such the Epetra multi-vectors of the Epetra matrices.

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```
int NumMyElement()  
    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 2. Some methods of the class Epetra_Vector
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 reserve 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 functionalities of these two classes, we consider the following example. Suppose that vector $x$ of global length 4, is distributed over two processes. Process 0 owns nodes 0,1,2, while process 1 owns nodes 1,2,3. This means that nodes 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 node 1 and 2 from the 2 processes. This is done in the following example (the complete code is reported in ${TRILINOS_HOME}/doc/tutorial/epetra/ex9.cpp).

```cpp
int NumGlobalElements = 4; // global dimension of the problem
int NumMyElements; // local nodes
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 map
Epetra_Map Map(-1,MyGlobalElements.Length(),
                MyGlobalElements.Values(),0, Comm);
```

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// 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:

[malsa:epetra] > mpirun -np 2 ./ex31.exe
Epetra::Vector
    MyPID  GID  Value
    0     0    10
    1     1    20
    1     2    20
    1     3    20
Epetra::Vector
    MyPID  GID  Value
3 Working with Epetra Matrices

Epetra contains several matrix classes. Epetra matrices can be defined to be *serial* or *parallel*:

- Examples of serial matrices are, for instance, the matrix corresponding to a given element in a finite-element discretization, or the Hessemerg matrix in the GMRES method. Those matrices are of small size, and therefore they are not distributed among the processors (but they can be replicated).

- For distributed sparse matrices, the basic class is Epetra_RowMatrix. This class is meant for double-precision matrices with row access (as required in a matrix-vector product), and it is a pure virtual class. Various classes are derived from Epetra_RowMatrix. Among them, here we recall:
  - 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.

This Chapter will show some of the Trilinos capabilities to work with matrices. During the Chapter, the user be introduced to:

- Create (serial) dense matrices (in Section 3.1);
- Create sparse point matrices (in Section 3.2);
- Create sparse block matrices (in Section 3.3);
- Insert non-local elements using finite-element matrices (in Section 3.4).

### 3.1 Serial Dense Matrices

Epetra provides functionalities for sequential dense matrices with the class Epetra_SerialDenseMatrix. A possible way to create a serial dense matrix $D$ of dimension $n \times m$ is
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().)

Epetra_SerialDenseMatrix are 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. Epetra_SerialDenseMatrix is intended to provide a very basic support for 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). 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.

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 );
AtimesB.Multiply('N','N',l.O, A, B, 0.0);
cout << AtimesB;
```

The complete code is reported in file `${TRILINOS_HOME}/doc/tutorial/epetra/ex10.cpp.`
To solve a linear system with a dense matrix, one has to create an Epetra_SerialDenseSolver. This class uses the most sophisticated techniques available in the LAPACK library. The class is built on the top of BLAS and LAPACK, and thus has excellent performances and numerical capabilities.

Given an Epetra_SerialDenseMatrix and two Epetra_DSerialDenseVectors x and b, the general approach is as follows:

```cpp
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,

```cpp
double rcond = Solve.RCOND();
```

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

File `${TRILINOS_HOME}/doc/tutorial/epetra/exll.cpp` outlines some of the capabilities of the Epetra_SerialDenseSolver class.

The Epetra_LAPACK class provides access to most of the same functionality as Epetra_SerialDenseSolver. The primary difference is that Epetra_LAPACK is a "thin" layer on the top of LAPACK, while Epetra_SerialDenseSolver attempts to provide easy access to the more sophisticated aspects of solving dense linear systems.

As a general rule, we can say that Epetra_LAPACK should be preferred when the user is looking for 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 want to work with a more object-oriented interface, he/she will probably use Epetra_SerialDenseMatrix.

### 3.2 Distributed Sparse Matrices

Epetra provided an extensive set of methods 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, as scaling, norm, matrix-vector multiplication and matrix-multivector multiplication.

Application do not need to know about the particular storage format, and other implementation details such as data layout, 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, generally much more efficient for problems with multiple degree of freedom per node, is introduced in Section 3.3. If required, other matrix formats can be supported via the Epetra.Operator, described in Section 4.3.

Remark 7. Some numerical algorithms require the application of the linear operator only. For this reason, some applications find convenient to not store a given matrix. Epetra can handle this situation using with the Epetra.Operator class; see Section 4.3.

The process of creating a sparse matrix is more involved with respect to that of dense matrices. This is because, in order to obtain excellent numerical performances, one has to provide an estimation of the nonzero elements on each row of the sparse matrix. (Recall that dynamic allocation of new memory and copying the old storage into the new one is an expensive operation.)

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

- allocate an integer array 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 Nnz;
- fill the sparse matrix.

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:

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

As the present stage of development, no functions are provided to perform a matrix-matrix product between to distributed objects. However, the interested user can convert the Epetra matrix into an ML matrix (called ML.Operator), perform the matrix-matrix multiplication with ML functions, and convert back the resulting ML.Operator into an Epetra matrix.
The example illustrates how to construct the matrix, and how to perform matrix-vector operations. The code can be found in `{TRILINOS_HOME}/doc/tutorial/epetra/ex12.cpp.

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

```c
int NumGlobalElements = 5;
```

We create a map, and define the local number of rows and the global numbering for each local row:

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

In particular, we have that `j = 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:

- 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 second approach can be coded 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). So we need 2 off-diagonal terms (except for the first and last equation). Here `NumNz[i]` is the Number of nonzero terms in the i-th global equation on this process.

Now, we create an `Epetra_CsrMatrix` as

```c
Epetra_CsrMatrix M(NumGlobalElements, NumMyElements, NumNz);
```
Epetra_CrsMatrix A(Copy, Map, NumNz);

and we add rows one-at-a-time. A has been created in Copy mode, and relies on the specified map. To fill its values, we need some additional variables: Indexes and Values. Those will contain the global column number and the values of the nonzeros for each row.

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;
        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 indexes have been inserted using global indexes. As a final operation, we can transform the matrix into local indexes. This phase in required in order to perform efficient parallel matrix-vector products and other matrix operations.

A.FillComplete();

The above presentation refers to a rather common case: In a parallel matrix-vector product

\[ AX = B, \]
the map used to define the parallel distribution of the matrix, is the same of the (multi-
vectors $X$ and $B$. This means that the rows of $A$ are distributed among the processes in
the same way of the elements of $X$ and $B$. However, Epetra allows the user to handle the
more general case of a matrix defined using a Map, is different from that of $X$ and that of
$B$. In fact, each Epetra matrix is defined by four maps:

- Two maps, called RowMap and ColumnMap, are used to determine the set of rows
and the columns of the elements assigned to a given processor. In general, one pro-
cessor cannot set elements assigned to other processors. (However, some classes,
derived from the Epetra_RowMatrix, can perform data exchange; see for instance
Epetra_FECCrsMatrix or Epetra_FEVbrMatrix.) RowMap and ColumnMap determine
the pattern of the matrix, as it is used during the construction. They can be obtained
using the methods RowMap() and ColMap() of the Epetra_RowMatrix class. Usually,
the user does not specify a ColumnMap, which is automatically created by Epetra.
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 completely different from RowMap and
ColumnMap, meaning that a matrix can be constructed using a certain data distribu-
tion, then used on vectors with another data distribution. DomainMap and RangeMap
can differ. Those two maps can be obtained using the methods DomainMap() and
RangeMap().

The potentialities of this approach are better explained using an example, reported in the
example file `{TRILINOS_HOME}/doc/tutorial/epetra/ex24.cpp. In this ex-
ample, to be run using two processors, we build up two maps: MapA will be used to con-
struct the matrix, while MapB to define the parallel layout of the vectors $X$ and $B$. For the
sake of simplicity, $A$ is diagonal.

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):

```cpp
for( int i=0 ; i<NumElementsA ; ++i ) {
    double one = 2.0;
    int indices = MyGlobalElementsA[i];
    A.InsertGlobalValues(MyGlobalElementsA[i], 1, &one, &indices );
}
```

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

```
A.FillComplete(MapB, MapB);
```

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

```
Epetra_Vector VecB(MapB);  Epetra_Vector VecB2(MapB);
A.Multiply(false, VecB, VecB2);
```

**Remark 8.** Although presented for Epetra.CrsMatrix objects, the distinction between RowMap, ColMap, DomainMap, and RangeMap is valid for all classes derived from Epetra.RowMatrix.

Example `${TRILINOS_HOME}/doc/tutorial/epetra/ex14.cpp` shows the use of some of the methods of the Epetra.CrsMatrix class. The code prints out several information about the structure of the matrix, and some of its properties. The output will be approximatively as here reported:

```
[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 are reported for Epetra_CrsMatrix:

- Example ${TRILINOS_HOME}/doc/tutorial/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 are hardwired. The interested user can easily modify those lines in order to read the grid information from a file.

- Example ${TRILINOS_HOME}/doc/tutorial/epetra/ex15.cpp explains how to export an Epetra_CrsMatrix to file in a MATLAB format. The output of this example will be as follows:

```
[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;
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 ${TRILINOS_HOME}/doc/tutorial/epetra/ex16.cpp, which exports an Epetra_Vector to MATLAB format.
```
3.3 Creating VBR Matrices

The following code shows how to work with VBR matrices. 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 it is done in the following example.

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

\[ \text{Epetra\_Map Map(NumGlobalElements,0,Comm);} \]

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

\[ \text{// local number of elements} \]
\[ \text{int NumMyElements = Map.NumMyElements();} \]
\[ \text{// global numbering of local elements} \]
\[ \text{int * MyGlobalElements = new int [NumMyElements];} \]
\[ \text{Map.MyGlobalElements( MyGlobalElements );} \]

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 \( \text{ElementSizeList} \) will contain the block size of local element \( i \).

\[ \text{Epetra\_IntSerialDenseVector ElementSizeList(NumMyElements);} \]
\[ \text{for( int i=0 ; i<NumMyElements ; ++i )} \]
\[ \text{ElementSizeList[i] = 1+MyGlobalElements[i];} \]

Here \( \text{ElementSizeList} \) is declared as Epetra\_IntSerialDenseVector, but an int array is fine as well.

Now we can create a map for the block distribution:

\[ \text{Epetra\_BlockMap BlockMap(NumGlobalElements,NumMyElements, MyGlobalElements, ElementSizeList.Values(),0,Comm);} \]
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.

```cpp
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 )
            Values[k] = 1.0*i;
        B.SubmitBlockEntry(Values,BlockRows,BlockCols,BlockRows,BlockCols);
    }
    B.EndSubmitEntries();
}
```

Note that, with VBR matrices, we have to insert one block at time. This required two more
instructions, one to start this process (BeginInsertGlobalValues), and another one
to commit the end of submissions (EndSubmitEntries).

Please refer to `${TRILINOS_HOME}/doc/tutorial/epetra/ex17.cpp` for
the entire source.
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 of setting non-local matrix elements. We will illustrate this using the following example, reported in `${TRILINOS_HOME}/doc/tutorial/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

```cpp
Epetra_FECrsMatrix A(Copy,Map,1);
```

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

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

The Function `SumIntoGlobalValues` adds the coefficients specified in `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).

At this point, we need to exchange data, to that each matrix element not owned by process 0 could be send to the owner, as specified by `Map`. This is accomplished by calling, on all processes,

```cpp
A.GlobalAssemble();
```

A simple

```cpp
cout << A;
```

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

Epetra includes a certain number of classes that can greatly help to develop 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

To retrieve elapsed and wall-clock time can be problematic because of several 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.

A Epetra_Time object is defined as

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

To compute the elapsed time required by a piece of code, then user should put the instruction

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

before the code to the timed. Then, the methods ElapsedTime() and WallTime() will return the elapsed time and wall-clock time, respectively. ElapsedTime() returns the elapsed time from the creation of this object.
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:

```cpp
Epetra_Flops counter();
```

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

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

Then, we perform the desired computations on Epetra objects (in this case, the vector-vector problem):

```cpp
x.Dot(y,&dotProduct);
```

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

```cpp
total_flops = counter.Flops();
```

which are the total number of *serial* flops. This will also reset the flop counter.

*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:

```cpp
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 << endl;

This code is reported in ${TRILINOS_HOME}/doc/tutorial/epetra/ex20.cpp. The output will be approximately as follows:

[mwala: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 9. Operation count are serial count, and therefore keep trace of local operations only.

Remark 10. Each computational class has a Flops() method, that can queried for the flop count of that object.

4.3 Epetra.Operator and Epetra_RowMatrix Classes

Matrix-free methods can be easily introduced in the Epetra framework using one of the following two classes:

- Epetra.Operator;
Technically, both classes are pure virtual classes (that is, they specify 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_FE_CrsMatrix, and Epetra_FE_VbrMatrix.

In the following, we consider for instance how to apply a matrix to a vector without explicitly constructing the matrix. The matrix is the classical finite-difference discretization of a Laplace on a 1D grid with constant grid-size. 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 TriDiagonalOperator, and derived from the Epetra_Operator class.

class TriDiagonalOperator : public Epetra_Operator {
public:
    // ... definitions here, constructors and methods
private:
    Epetra_Map Map_;  // value in the sub-diagonal
    double diag_minus_one_;  // value in the diagonal
    double diag_;  // value in the super-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 Apply method, which may be coded as follows:

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 ) {

            // code here
        }
    }
}

int main() {
    // code here
    return 0;
}

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\[
Y[\text{vec}][0] = \text{diag} \ast X[\text{vec}][0];
\]
break;
\]
// more general case (Length >= 2)
// first row
\[
Y[\text{vec}][0] = \text{diag} \ast X[\text{vec}][0] + \text{diag}_{\text{plus one}} \ast X[\text{vec}][1];
\]
// intermediate rows
for( int i=1 ; i<Length-1 ; ++i ) {
\[
Y[\text{vec}][i] = \text{diag} \ast X[\text{vec}][i] + \text{diag}_{\text{plus one}} \ast X[\text{vec}][i+1]
+ \text{diag}_{\text{minus one}} \ast X[\text{vec}][i-1];
\]
}
// final row
\[
Y[\text{vec}][\text{Length-1}] = \text{diag} \ast X[\text{vec}][\text{Length-1}]
+ \text{diag}_{\text{minus one}} \ast X[\text{vec}][\text{Length-2}];
\]
return true;
}\]

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

\[
\text{TriDiagonalOperator TriDiagOp(-1.0,2.0,-1.0,Map);}
\]

and we can apply this operator to a vector as:

\[
\text{DiagOp.Apply(x,y);}
\]

\[
\text{${TRILINOS_HOME}/doc/tutorial/epetra/ex21.cpp$ reports the entire source code.}
\]

**Remark 11.** 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 patterns for intra-process data exchange. For this purpose, Epetra\_Import classes can be used. File \$\{TRILINOS_HOME\}/doc/tutorial/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's 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 monoprocess case. Extensions to multi-processes case requires to handle ghost-nodes updates.

As a first step, we create a class, derived from the Epetra::Operator class,

class MSRMatrix : public Epetra::Operator
{

public:
  // constructor
  MSRMatrix(Epetra::Map Map, int * bindx, double * val) :
    Map_(Map), bindx_(bindx), val_(val)
  {}

  ~MSRMatrix() // destructor
  {}

  // Apply the RowMatrix to a MultiVector
  int Apply(const Epetra::MultiVector & X, Epetra::MultiVector & Y ) const
  {
    int Nrows = bindx_[0]-1;

    for( int i=0 ; i<Nrows ; i++ ) {
      // diagonal element
      for( int vec=0 ; vec<X.NumVectors() ; ++vec ) {
        Y[vec][i] = val_[i]*X[vec][i];
      }
      // off-diagonal elements
      for( int j=bindx_[i] ; j<bindx_[i+1] ; j++ ) {
        for( int vec=0 ; vec<X.NumVectors() ; ++vec ) {
          Y[vec][bindx_[j]] += val_[j]*X[vec][bindx_[j]];
        }
      }
    }
    return 0;
  }
}
} /* Apply */
... other functions ...

private:
int * bindx_; double * val_;
}

As stated by the fragment of code above, the constructor takes the two MSR vectors, and an Epetra_Map. The complete code is reported in ${TRILINOS_HOME}/doc/tutorial/epetra/ex2!

4.4 Epetra_LinearProblem

A linear problem of type $AX = B$ is defined by an Epetra_LinearProblem class. This 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 problems can be used to solve linear systems with iterative methods (typically, using AztecOO, covered in Chapter 5), or with direct solvers (typically, using Amesos, described in Chapter 8.)

Once the linear problem has been defined, the user can:

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

4.5 Concluding Remarks

More details about the Epetra project, and a technical description of classes and methods, can be found in [5, 9].
5 Iterative Solution of Linear Systems with AztecOO

AztecOO is a package that extends the Aztec library [20]. Aztec is the legacy iterative solver at the Sandia National Laboratories. It has been extracted from the MPSalsa reacting flow code [17, 15], and it is currently installed in dozens of Sandia's applications. AztecOO extends this package, using C++ classes to enable more sophisticated use.

AztecOO is intended for the iterative solution of linear systems of the form

\[ A X = B, \]  

(1)

when \( A \in \mathbb{R}^{n \times n} \) is the linear system matrix, \( X \) the solution, and \( B \) the right-hand side. Both \( X \) and \( B \) are Epetra_Vector objects.

In this Chapter, we will:

- Outline the basic issues of the iterative solution of linear systems (in Section 5.1);
- Present the basic usage of AztecOO (in Section 5.2);
- Define one-level domain decomposition preconditioners (in Section 5.3);
- Use of AztecOO problems as preconditioners to other AztecOO problems (in Section 5.4).

5.1 Theoretical Background

Aim of this Section is to briefly present some aspects of the iterative solution of linear 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.

One can distinguish between two different aspects of the iterative solution of a linear system. The first one is 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 from previous approximations. This leads to specific iteration methods, like conjugate gradient or GMRES. The second aspect is the transformation of the given system to one that can be more efficiently solved by a particular iteration method. This is called 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 (1) by

\[ P^{-1}AX = P^{-1}B \]

(left-preconditioning), or by

\[ AP^{-1}PB = B \]

(right-preconditioning), using a 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.

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

1. \( P \) is a good approximation of \( A \) in some sense. Although no general theory is available, we can say that \( P \) should act so that \( P^{-1}A \) is near to being the identity matrix and its eigenvalues are clustered within a sufficiently small region of the complex plane;

2. \( P \) is efficient, in the sense that the iteration method converges much faster, in terms of CPU time, for the preconditioned system. In other words, preconditioners must be selected in such a way that the cost of constructing and using them is offset by the improved convergence properties they permit to achieve;

3. \( P \) or \( P^{-1} \) can take advantage of the architecture of modern supercomputers, that is, can be constructed and applied in parallel environments.

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

The choice of \( P \) varies from "black-box" algebraic techniques which can be applied to general matrices to "problem dependent" preconditioners which exploit special features of a particular class of problems. Although problem dependent preconditioners can be very powerful, there is still a practical need for efficient preconditioning techniques for large classes of problems. 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, since the user has to supply few information about the matrix and the problem to be solved.
AztecOO itself implements a variety of preconditioners, from “classical” methods such as Jacobi and Gauss-Seidel, to polynomial and domain-decomposition based preconditioners. More preconditioners can be given to an AztecOO Krylov accelerator, by using the Trilinos packages IFPACK and ML, covered in Chapter 6 and 7, respectively.

5.2 Basic Usage of AztecOO

To solve a linear system with AztecOO, one must create an `Epetra_LinearProblem` object with the command

```c
Epetra_LinearProblem Problem(&A, &x, &b);
```

where `A` is an Epetra matrix, and `x`, `b` two Epetra vectors. Then, the user must create an AztecOO object,

```c
AztecOO Solver (Problem);
```

and specify how to solve the linear system. All AztecOO options are set using two vectors, `options` (integer) and `params` (double), as detailed in the Aztec’s User Guide.

To choose among the different AztecOO parameters, the user can create two vectors, usually called `options` and `params`, set them to the default values, and then override with the desired parameters: Default values can be set by

```c
int options[AZ_OPTIONS_SIZE];
double params[AZ_PARAMS_SIZE];
AZ_defaults(options, params);
```

followed by, for instance,

```c
Solver.SetAllAztecOptions(options);
Solver.SetAllAztecParams(params);
```

---

5At the current stage of development, AztecOO does not handle Epetra.MultiVectors. It accepts Multi.Vectors, but it will solve the linear system corresponding to the first multivector only.

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Those two functions will copy the values of options and params in internal variables of the AztecOO object.

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

```c
Solver.SetAztecOption( AZ_precond, AZ_Jacobi );
Solver.SetAztecParams( AZ_tol, 1e-12 );
```

to specify a point Jacobi preconditioner, and a tolerance of $10^{-12}$. (We refer to the Aztec documentation for more details about the various Aztec settings.)

To solve the linear system the user may call

```c
Solver.Iterate(1000, 1E-9);
```

The complete code is in `${TRILINOS_HOME}/doc/tutorial/aztec/ex1.cpp`.

Note that the matrix must be in local coordinates (that is, the command `A.FillComplete()` has been called before attempting to solve the linear system). Note also that the procedure to solve a linear system with AztecOO is identical for sequential and parallel runs. However (for certain choices of the preconditioners), the convergence rate can change as the number of processes used in the computation varies.

When this function returns, one can retrieve the number of iterations performed by the linear solver using `Solver.NumIters()`, while `Solver.TrueResidual()` gives the (nonscaled) norm the residual.

### 5.3 One-level Domain Decomposition Preconditioners with AztecOO

In this Section, we will consider preconditioners based on one-level overlapping domain decomposition preconditioners, of the form

$$ P^{-1} = \sum_{i=1}^{M} R_i^T \bar{A}_i^{-1} R_i, $$

where $P$ is the preconditioning operator, $M$ the number of subdomains. $R_i$ is a rectangular matrix, composed by 0's and 1's, which restricts a global vector to the subspace defined by
the interior of each subdomain, and $\tilde{A}_i$ is an approximation of

$$A_i = R_i A R_i^T.$$  

($\tilde{A}_i$ can be equal to $A_i$). Typically, $\tilde{A}_i$ differs from $A_i$ when incomplete factorizations are used in (2) to apply $\tilde{A}_i^{-1}$, or when a matrix different from $A$ is used in (3).

In order to use a preconditioner of the form (3), the user has to specify

Solver.SetAztecOption( AZ_precond, AZ_dom_decomp );

followed by the choice of incomplete factorization (and possibly with that of corresponding parameters, for instance the level-of-fill),

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

By default, AztecOO will consider zero-overlap among the rows of $A^6$. However, this value of overlap can be changed by, for instance,

Solver.SetAztecOption( AZ_overlap, 1 );

**Remark 12.** By using AztecOO in conjunction with ML, one can easily implement a two-level domain decomposition schemes. The reader is referred to Section 7.3.

**Remark 13.** Another Trilinos package can be used to compute incomplete factorizations, IFPACK. It is covered in Chapter 6.

### 5.4 Use of AztecOO Problems as a Preconditioner for AztecOO

One may wish to use an AztecOO solver in the preconditioning phase, as done in ${TRILINOS_HOME}/doc/tb.$ The main steps are here reported.

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:

---

Footnote 6: For point matrices arising from the FE discretization of the PDE problem with local functions, this is equivalent to one mesh element of overlap.
Epetra_LinearProblem A_Problem(&A, &x, &b);
AztecOO A_Solver(A_Problem);

Now, we have to define the preconditioner. For the sake of simplicity, we here suppose to use the same Epetra_Matrix A in the preconditioning phase. However, the two matrices can in principle be different (although of the same size).

Epetra_CrsMatrix P(A);

(This operation is in general expensive as involves the copy constructor.) Then, we create the linear problem which will be used as preconditioner. This requires several steps. (Note that all the P prefix identifies preconditioner' objects.)

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

   Epetra_LinearProblem P_Problem;
P_Problem.SetOperator(&P);

2. As we wish to use AztecOO to solve the prec step (in a recursive way), we have to define an AztecOO object:

   AztecOO P_Solver(P_Problem);

3. Now, we customize certain parameters:

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

4. The last step is to create an AztecOO.Operator, so that we can set the Aztec's preconditioner with, and we set the user's 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. Finally, we solve the linear system:

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

5.5 Concluding Remarks

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 unscaled residual;
- `SetAztecDefaults()` can be used to restore default values in the options and params vectors.

The official documentation of AztecOO can be found in [8].
6 Incomplete Factorizations with IFPACK

IFPACK provides a suite of object-oriented algebraic preconditioners for the solution of preconditioned iterative solvers. IFPACK offers a variety of overlapping (one-level) Schwarz preconditioners, The packages uses Epetra for basic matrix-vector calculations, and accepts user matrices via abstract matrix interface. A concrete implementation for Epetra matrices is provided. The package separates graph construction for factorization, improving performances in a substantial manner with respect to other factorization packages.

In this Chapter we present how to use IFPACK objects as a preconditioner for an AztecOO solver.

In this Chapter, we will

- Set the notation (in Section 6.1);
- Show how to compute incomplete Cholesky factorizations (in Section 6.2);
- Present IFPACK’s RILU-type factorizations (in Section 6.3).

6.1 Theoretical Background

Aim of this Section is to briefly present some aspects on incomplete factorization methods, 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.

A broad class of effective preconditioners is based on incomplete factorization of the linear system matrix, and it is usually indicated as ILU. The ILU-type preconditioning techniques lie between direct and iterative methods and provide a balance between reliability and numerical efficiency.

The preconditioner is given 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.

The incomplete LU factorization of a matrix \( A \) can be described as follows. Let \( A_0 = A \). Then, for \( k = 2, \ldots, n \), we have

\[
A_{k-1} = \begin{pmatrix} B_k & F_k \\ E_k & C_k \end{pmatrix}.
\]
Thus, we can write the $k$-step of the Gaussian elimination in a block form as

$$A_{k-1} = \begin{pmatrix} I & 0 \\ E_k B_k^{-1} & I \end{pmatrix} \begin{pmatrix} B_k & F_k \\ 0 & A_k \end{pmatrix},$$

where $A_k = C_k - E_k B_k^{-1} F_k$. If $B_k$ is a scalar, then we have the typical point-wise factorization, otherwise we have a block factorization. Pivoting, if it is necessary, can be accomplished by reordering $A_k$ at every step.

To make the factorization incomplete, entries as dropped in $A_k$, i.e. the factorization proceeds with

$$\tilde{A}_k = A_k - R_k,$$

where $R_k$ is the matrix of dropped entries.

Dropping can be performed by position, for example, dropping those entries in the update matrix $E_k B_k^{-1} F_k$ that are not in the pattern of $C_k$. This simple ILU factorization is known as ILU(0). 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($f$), where $f$ 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 than level-of-fill methods. The general strategy is to compute an entire row of the $\tilde{L}$ and $\tilde{U}$ matrices, and then keep only the biggest entries in a certain number. 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.

### 6.2 Incomplete Cholesky Factorizations

Ifpack_CrsLct is a class for constructing and using incomplete Cholesky factorizations of an Epetra_CrsMatrix. The factorization is produced based 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 very easy to 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

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

where $$e = (1, 1, \ldots, 1)^T$$. (More details can be found in the IFPACK's documentation.) This estimation can be computed as follows:

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

Please refer to file `${TRILINOS_HOME}/doc/tutorial/ifpack/exl.cpp` for a complete example of incomplete Cholesky factorization.

### 6.3 RILU 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

```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 14.** 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
ILUT = 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 = ILUT->InitValues(A);
```

The ILUT object can be used with AztecOO simply setting

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

where `solver` is an AztecOO object. Example `{$TRILINOS_HOME}/doc/tutorial/ifpack/ex2.cpp` shows the use of 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.

6.4 Concluding Remarks

More documentation on the IFPACK package can be found in [6, 4].
7 Multilevel Methods with ML

The ML package defines a class of preconditioners based on multilevel methods [18]. While technically any linear system can be considered, ML should be used on linear systems, like elliptic PDEs, that are known to work well with multilevel methods.

ML is a large package, that can be used to a variety of purposes. ML provides multilevel solvers, as well as multilevel preconditioners, and it can handle geometric as well as algebraic methods.

In this Chapter we present:

- Outline the basic issues of multilevel schemes (in Section 7.1);
- Present the use of ML objects as a preconditioner for an AztecOO solver objects (in Section 7.2);
- Outline the steps required to implement two-level domain decomposition methods, with a coarse grid defined using aggregation procedures (in Section 7.3).

As other Trilinos packages, ML can be compiled and run independently from Epetra, that is, it can accept input matrix in formats different from the Epetra_RowMatrix or Epetra_Operator. Should the reader be interested in running ML without Epetra, or using a C code (and not a C++ code), then we refer to the ML guide, contained in the $\{TRILINOS_HOME\}/packages/ml/doc/.

7.1 Theoretical Background

Aim of this Section is to briefly present some aspects on multilevel methods. 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.

Multilevel methods require the operator to be defined on a sequence of coarser spaces, an iterative method that evolves the solution (called a smoother) and interpolation operators that transfer information between the spaces. The principle behind the algorithm is that the high-frequency errors can be efficiently solved on the fine space, while the low-frequency are treated on the coarser one, where there frequencies manifest themselves as
high-frequencies. A very popular multilevel methods are multigrid methods. Geometric
multigrid (GMG) methods cannot be applied without the existence of an underlying grid
(this is their major limitation). This led to the development of algebraic multigrid method
(AMG), initiated by Ruge and Stüben. In AMG, both the matrix hierarchy and the pro-
longation operators are constructed just from the stiffness matrix. Since the automatic
generation of a grid-hierarchy for GMG and especially the proper assembly of all com-
ponents would be a very difficult task for unstructured problems, the automatic algebraic
construction of a virtual grid is a big advantage.

A function to solve (1) using a multilevel method can be defined as follows:

\[
\text{MGM}(X, B, k) \begin{cases} 
  \text{if}(k == 0) & X = A_k \backslash B; \\
  \text{else} & \\
    X = S_{k^1}(X, B); \\
    D = R_{k-1,k}(B - A_k X); \\
    V = 0; \\
    \text{MGM}(V, D, k-1); \\
    X = X + P_{k,k-1} V; \\
    X = S_{k^2}(U, B); \\
\end{cases}
\]

In the above method, \(S_{k}^{1}\) and \(S_{k}^{2}\) are two smoothers, \(R_{k-1,k}\) is a restriction operator from
level \(k\) to \(k - 1\), and \(P_{k,k-1}\) is a prolongator from \(k - 1\) to \(k\).

In a variational setting, the matrices \(A_k\) can be constructed as

\[
A_k = R_{k-1,k} A_k P_{k,k-1}.
\]

Alternatively, when a grid is available at level \(k - 1\), one can discretize the PDE operator
on grid \(k - 1\).

**Remark 15.** In this tutorial, we will consider multilevel methods based on aggregation
schemes only.

### 7.2 ML as a Preconditioner for AztecOO

In order to use ML as a preconditioner, we need to define an AztecOO Solver, as outlined
in Chapter 5.
ML requires the user to define a structure, to store internal data. This structure is usually called `ml_handle`:

```c
ML *ml_handle;
```

We intend to use ML as a “black-box” (or gray-box) multilevel preconditioner, using aggregation procedures to define the multilevel hierarchy. The variable

```c
int N_levels = 10;
```

defines the maximum number of levels, while

```c
ML_Set_PrintLevel(3);
```

toggle the output level (from 0 to 10, 10 being verbose mode and 0 silent mode).

The ML handle is created using

```c
ML_Create(&ml_handle, N_levels);
```

ML can accept in input very general matrices. Basically, the user has to specify the number of local rows, and provide a function to update the ghost nodes (that is, nodes requires in the matrix-vector product, but assigned to another process). For Epetra matrices, this is done by the following function

```c
EpetraMatrix2MLMatrix(ml_handle, 0, &A);
```

Note that `A` is not converted to ML format. Instead, proper wrappers are defined. (Here, `A` is the Epetra matrix for which we aim to construct a multilevel preconditioner.)

ML requires another structure, called `ML_Aggregate`, to store the information about the aggregates at various levels:

```c
ML_Aggregate *agg_object;
ML_Aggregate_Create(&agg_object);
```

The multilevel hierarchy is constructed with the instruction
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. (We refer to the ML manual for more details about the input parameters.)

We still need to define the smoother, for instance a symmetric Gauss-Seidel:

ML_Gen_Smoother_SymGaussSeidel(ml_handle, ML_ALL_LEVELS,
        ML_BOTH, 1, ML_DEFAULT);

and to generate the solver as

ML_Gen_Solver (ml_handle, ML_MGV, 0, N_levels-1);

Finally, we can create an Epetra_Operator, based on the previously defined ML hierarchy

Epetra_ML_Operator MLop/ml_handle,comm,map,map);

and set the preconditioning operator of our AztecOO solver,

solver.SetPrecOperator(&MLop);

At this point, we can call Iterate() as usual,

solver.Iterate(Niters, le-12);

The entire code is reported in ${TRILINOS_HOME}/doc/tutorial/ml/exl.cpp. The output will be approximatively as reported below.

[msala:ml]> mpirun -np 2 ./exl.exe

******************************************************************************
* ML Aggregation information                                                 *
63
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: max coarse size = 1
ML_Aggregate: max no. of levels = 10

ML_Aggregate-Coarsen begins
ML_Aggregate-CoarsenUncoupled: current level = 0
Aggregation(UVB): Total nonzeros = 128 (Nrows=30)
Aggregation(UC): Phase 0 - no. of bdry pts = 0
Aggregation(UC): Phase 1 - nodes aggregated = 28 (30)
Aggregation(UC): Phase 1 - total aggregates = 8
Aggregation(UC_Phase2_3): Phase 1 - nodes aggregated = 28
Aggregation(UC_Phase2_3): Phase 2a - additional aggregates = 0
Aggregation(UC_Phase2_3): Phase 2 - total aggregates = 8
Aggregation(UC_Phase2_3): Phase 3 - boundary nodes = 0
Aggregation(UC_Phase2_3): 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
ML_Aggregate-CoarsenUncoupled: current eps = 0.000000e+00
Aggregation(UVB): Total nonzeros = 46 (Nrows=8)
Aggregation(UC): Phase 0 - no. of bdry pts = 0
Aggregation(UC) : Phase 1 - nodes aggregated = 6 (8)
Aggregation(UC) : Phase 1 - total aggregates = 2
Aggregation(UC_Phase2_3) : Phase 1 - nodes aggregated = 6
Aggregation(UC_Phase2_3) : Phase 1 - total aggregates = 2
Aggregation(UC_Phase2_3) : Phase 2a- additional aggregates = 0
Aggregation(UC_Phase2_3) : Phase 2 - total aggregates = 2
Aggregation(UC_Phase2_3) : Phase 2 - boundary nodes = 0
Aggregation(UC_Phase2_3) : 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
7.3 Two-level Domain Decomposition Preconditioners with ML

In order to use the example reported in this Section, one should compile ML with the configure flag `--with-ml-metis`. In this way, ML will use the graph decomposition library METIS to create the coarse-level matrix\(^7\).

Two-level domain decomposition methods have been proved to be very effective for the iterative solution of many different kind of linear systems. For some classes of problems, a very convenient way to define the coarse grid operator is to use aggregation procedure. This is very close to what presented in Section 7.2. 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)\(^8\). This can be done as follows.

First, we need to define an AztecOO problem, an ML structure, and an MLAggregate structure. Then, we limit ourselves to 2-level scheme.

```cpp
int N_levels = 2;
```

Then, we specify the aggregation scheme as

```cpp
ML_Aggregate_Set_CoarsenScheme_METIS(agg_object);
```

and define the number of aggregates (here, 4) to be defined on each process as

```cpp
ML_Aggregate_Set_LocalNumber( ml_handle, agg_object, 0, 4 );
```

As smoother, we can adopt a subdomain-based Gauss-Seidel smoother.

The creation of the multilevel hierarchy and the solution of the linear system will be as reported in Section 7.2.

The entire code is reported in `${TRILINOS_HOME}/doc/tutorial/ml/ex2.cpp`.

\(^7\)Note that ML has to be aware of the location of the METIS include files and the METIS library. The user can use the configure flags `--with-incdirs` and `--with-ldflags`. Please type `configure --help` for more information. If you don’t have METIS, or you don’t want to re-configure ML, you will be able to run the example of this Section. However, you will be limited to use only one aggregate per process.

\(^8\)Aggregation schemes based on ParMETIS are also available. Please refer to the help of the ML configure for more details.
7.4 Concluding Remarks

More documentation about ML can be found in [21, 19, 19].
8 Interfacing Direct Solvers with Amesos

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

\[ AX = B \]  

(4)

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.

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

```c
AmesosProblem.Solver();
```

Amesos will take care of redistributing data among the processors, if necessary.

This Chapter starts with few notes on the installation of the third-part packages required by Amesos. Then, the Chapter will present the use of Amesos objects, to interface with the following packages:

- UMFPACK, version 4.1 (in Section 8.2);
- SuperLUdist, version 2.0 (in Section 8.3);
- A generic interface to various direct solvers is presented (in Section 8.4).

8.1 Installation of Trilinos third-part Packages

Amesos is an interface to other packages, mainly developed outside the Trilinos framework\(^9\). In order to use those packages, the user should carefully check copyright and licensing of those third party codes. Please refer to the web page or the documentation of each particular package for details.

\(^9\)Currently, SuperLU is included in the Trilinos framework.
Amesos supports a variety of direct solvers for linear systems of equations, and you are likely to use Amesos with only few of them. We suggest to define the shell variable TRILINOS_3PL to define the directory used to stored third-part packages. For instance, under BASH, you may have a line of type

```bash
export TRILINOS_3PL=/home/msala/Trilinos3PL
```

in your .bashrc file. Then, you may decide to create a directory to hold include files and libraries. For instance, to compile under LINUX with MPI:

```bash
$ mkdir ${TRILINOS_3PL}/LINUX_MPI
$ mkdir ${TRILINOS_3PL}/LINUX_MPI/include
$ mkdir ${TRILINOS_3PL}/LINUX_MPI/lib
```

(Note that this will reflect the directory structure used by Trilinos, see Section 1.2.) While installing a package, you can now copy all include files and libraries in these directories.

Using this setting, you can configure Amesos with a command of type

```bash
$ cd ${TRILINOS_HOME}/packages/amesos
$ ./configure --prefix=${TRILINOS_HOME}/LINUX_MPI \
   --enable-mpi --with-mpi-compilers \
   --enable-amesos-umfpack \
   --enable-amesos-superludist \
   --with-amesos-superludistlib="
   "$ {TRILINOS_3PL}/SuperLU_DIST_2.0/libsuperlu_LINUX.a"
```

(This command is followed by make and make install, as usual.) This will enable UMFPACK and SuperLUdist, which are the two packages covered in this Chapter.

For more details about the configuration options of Amesos, please refer to Amesos documentation.

### 8.2 UMFPACK

File ${TRILINOS_HOME}/doc/tutorial/amesos/ex1.cpp shows how to use Amesos to solve a linear system with UMFPACK\(^\text{10}\).

\(^\text{10}\)UMFPACK is a set of routines solving sparse linear systems via LU factorization. It is copyrighted by Timothy A. Davis. More information can be obtained at the web page
Suppose that $A$, $x$ and $b$ are an Epetra_RowMatrix and two Epetra_MultiVector, respectively, or compatible dimensions. Amesos objects for the solution of linear systems requires an Epetra_LinearProblem object, plus another object, AMESOS::Parameter::List, used to specify the parameters.

```cpp
Epetra_LinearProblem Problem(&A,&x,&b);
AMESOS::Parameter::List params;
```

Then, only few lines are required: We can define an Amesos object and solve the problem,

```cpp
Amesos_Umfpack UmfpackProblem(Problem,params);
UmfpackProblem.Solve();
```

or, alternatively, it is possible to specify when symbolical factorization, numerical factorization and solution occur,

```cpp
Amesos_Umfpack UmfpackProblem(Problem,params);
UmfpackProblem.SymbolicFactorization();
UmfpackProblem.NumericFactorization();
UmfpackProblem.Solve();
```

Note that exactly the same code can be run with more than one processor. In this case, being UMFPACK a serial solver, Amesos will take care to gather all required data on a processor, solve sequentially the linear system, and then broadcast the solution.

### 8.3 SuperLUdist

Solving using SuperLUdist\[^{11}\] is not much different from what presented in Section 8.2. Instead of declaring an Amesos_Umfpack object, one can proceed as follows:

```cpp
Amesos_Superludist * SuperludistProblem =
    new Amesos_Superludist(Problem,params);
```

[^11]: SuperLU DIST is a parallel extension to the serial SuperLU library. It is targeted for the distributed memory parallel machines. Copyright (c) 2003, The Regents of the University of California, through Lawrence Berkeley National Laboratory. Please refer to the web site [http://www-users.cs.umn.edu/~xjia/ SuperLU](http://www-users.cs.umn.edu/~xjia/SuperLU) for more information.
followed by a call to Solve(), possibly preceded by SymbolicFactorization() and NumericFactorization().

Remark 16. We have declared a pointer to an Amesos_Superludist object because the destructor of this object contains some MPI calls. As in example

${}\text{TRILINOS.HOME}/\text{doc/tutorial/amesos/ex2.cpp}$ the destructor is called at the end of the main function (after a call to MPI_Finalize()), we have to delete this object using the C++ statement

```cpp
delete SuperludisProblem;
```

before the call to MPI_Finalize().

8.4 A Generic Interface to Various Direct Solvers

All Amesos objects are derived from the Amesos_BaseClass object. Using the capabilities of C++, one may decide to code a generic interface to a direct solver as follows:

```cpp
// parameter vector for Amesos
AMESOS::Parameter::List ParamList;

// prepare the linear solver
Amesos_BaseSolver * AmesonSolver;

switch( choice ) {  
    case ML_SOLVE_WITH_AMESOS_UMFPACK:  
        AmesonSolver =  
            new Amesos_Umfpack( *LinearProblem, ParamList );  
        break;
    case ML_SOLVE_WITH_AMESOS_SUPERLUDIST:  
        AmesonSolver =  
            new Amesos_Superludist( *LinearProblem, ParamList );  
        break;
    default:  
        cerr << "Error" << endl;
}
```

Now, factorization and solution are the same for all the packages:
AmesosProblem->SymbolicFactorization();
AmesosProblem->NumericFactorization();
AmesosProblem = (void *) AmesosProblem ;
9 Solving Nonlinear Systems with NOX

NOX is a suite of solution methods for the solution of nonlinear systems of type

\[ F(x) = 0, \]

with

\[ F(x) = \begin{pmatrix} f_1(x_1, \ldots, x_n) \\ \vdots \\ f_n(x_1, \ldots, x_n) \end{pmatrix} \]

is a nonlinear vector function. The Jacobian matrix of \( F \), \( J \), is defined by

\[ J_{i,j} = \frac{\partial F_i}{\partial x_j}(x). \]

NOX aims to solve (5) 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 issues of the solution of nonlinear systems (in Section 9.1);
- Introduce the NOX package (in Section 9.2);
- Describe how to introduce a NOX solver in an existing code (in Section 9.3);
- Present Jacobian-free methods (in Section 9.6).

9.1 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 (5). The algorithm is as follows: given $x_0$, for $k = 1, \ldots$ until convergence, solve

$$J_k(x_{k-1}) (x_k - x_{k-1}) = -F(x_{k-1}), \quad J_k(x_{k-1}) = \left[ \frac{\partial F}{\partial x}(x_{k-1}) \right].$$

(6)

Newton method introduces a local full linearization of the equation. 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 (6) 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 (6) 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 (5), 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.

### 9.2 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 NOX::Abstract::Vector (which provides support for basic vector operations as dot products), and 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 --enable-nox-epetra has to be specified (see Section 1.2)\(^\text{12}\).

### 9.3 Introducing NOX in an Existing Code

Two basic steps are required to implement a NOX::Epetra interface. First, a concrete class derived from 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

   ```
   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 numerical 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. 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 Epetra_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)
   ```

\(^{12}\)Other two concrete implementation are provided, for LAPACK and PETSc. The user may wish to configure NOX with --enable-nox-lapack or --enable-nox-petsc. Examples can be compiled with the options --enable-nox-lapack-examples, --enable-nox-petsc-examples, and --enable-nox-epetra-examples.
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

\[
\text{computePreconditioner(const Epetra\_Vector & x, Epetra\_Operator & M)}
\]

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 \texttt{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 \texttt{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,

\[
\text{Epetra\_Operator} \text{ } \& \text{ } J = \text{UserProblem.getJacobian()},
\]

created as a NOX matrix-free operator,

\[
\text{NOX::Epetra::MatrixFree} \text{ } \& \text{ } J = \text{MatrixFree(userDefinedInterface, solutionVec)},
\]

or created by NOX using a finite differencing,

\[
\text{NOX::Epetra::FiniteDifference} \text{ } \& \text{ } J = \text{FIXME}...
\]

At this point, users have to create an instantiation of the \texttt{NOX::Epetra::Interface} derived object,

\[
\text{UserInterface interface(}...\text{)},
\]

and finally construct the group,

\[
\text{NOX::Epetra::Group group(printParams, lsParams, interface, FIXME)}.
\]
9.4 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 (5) 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

$$\alpha = \left(\sqrt{\frac{5 - 1}{2}}, \sqrt{\frac{5 - 1}{2}}\right),$$

the other being the reflection of $\alpha$ among the $x_2$ axis.

Code \$\{TRILINOS_HOME\}/doc/tutorial/nox/exl.cpp applies the Newton method to this problem, with $x_0 = (0.5, 0.5)$ as a starting solution. The output is approximately as follows:

```
[mmsala:nox] > mpirun -np 1 ./exl.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
**************************

**************************
-- 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 Iter = 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"
9.5 A 2D Nonlinear PDE Problem

In this Section, we consider the solution of the following nonlinear PDE problem:

\[
\begin{align*}
-\Delta u + \lambda e^u &= 0 \quad \text{in } \Omega = (0, 1) \times (0, 1) \\
\end{align*}
\]

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 $\{\text{TRILINOS_HOME}/doc/tutorial/nox/ex2.cpp$, we build the Jacobian matrix as an Epetra_CrsMatrix, and we use NOX to solve problem (7) 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,

\[
\text{SimpleProblemInterface Interface(&Problem);}
\]

Once a NOX::Epetra:Interface object has been defined, the procedure is almost identical to that of the previous Section.

### 9.6 Jacobian-free Methods

In Section 9.5, the entries of the Jacobian matrix have been explicitly coded. For more complex discretization schemes, 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.

### 9.7 Concluding Remarks

The documentation of NOX can be found in [13].

A library of continuation classes, called LOCA [14, 16], 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.
Triutils is a collection of various utilities, that can help development and testing. Mainly, triutils contains functions or classes to generate matrices in various formats (MSR, VBR, Epetra), to read matrices (in HB or COO format), to convert matrices from one format to another, and to process the command line. Programs using triutils should include the file Trilinos_Util.h.

In this Chapter, we will present:

- How to read a matrix (and possibly right-hand side and solution vectors) from an Harwell/Boeing file format (in Section 10.1);
- How to retrieve a parameter specified on the command line (in Section 10.2).

10.1 Reading a HB problem

It is possible to read matrix, solution and right-hand side, from a file written in the Harwell/Boeing format. This is done in ${TRILINOS_HOME}/doc/tutorial/triutils/ex1.cpp. The key instructions are the following.

First, we define pointers to Epetra_Vector and Epetra_Matrix objects:

```cpp
// Pointers because of Trilinos_Util_ReadHb2Epetra
Epetra_Map * readMap;
Epetra_CrsMatrix * readA;
Epetra_Vector * readx;
Epetra_Vector * readb;
Epetra_Vector * readxexact;
```

The HB problem is read with the instruction

```cpp
Trilinos_Util_ReadHb2Epetra(fileName, Comm, readMap, readA, readx, readb, readxexact);
```

Here, Comm is an Epetra_SerialComm or Epetra_MpiComm object, and FileName an array of character containing the name of the HB file.
This creates an Epetra_Matrix and two Epetra_Vectors, with all the elements assigned to processor zero. This is because the HB file does not contain any information about the distribution of the elements to the processors. Should the user need to solve the linear problem in parallel, thus he has to redistributed readA. In this case, the first step is to specify a map. For instance, we can use a linear map:

```cpp
int NumGlobalElements = readMap->NumGlobalElements();
Epetra_Map map(NumGlobalElements, 0, Comm);
```

and create and exporter to distribute read-in matrix and vectors:

```cpp
Epetra_Export exporter(*readMap, map);
Epetra_CrsMatrix A(Copy, map, 0);
Epetra_Vector x(map);
Epetra_Vector b(map);
Epetra_Vector xexact(map);
// this is the data distribution phase
x.Export(*readx, exporter, Add);
b.Export(*readb, exporter, Add);
xexact.Export(*readxexact, exporter, Add);
A.Export(*readA, exporter, Add);
```

Finally, we can destroy the objects used to store the non-distributed HB problem:

```cpp
delete readA;
delete readx;
delete readb;
delete readxexact;
delete readMap;
```

and solve the distributed linear system with the method of choice.

### 10.2 ShellOptions

ShellOptions is a class to manage the input arguments and shell variables. With this class, it is easy to handle input line arguments and shell variables. For instance, the user can write
$ ./ex2.exe -nx 10 -tol 1e-6 -solver=cg

and then easily retrieve the value of nx, tol, and solver.

A simple code using this class is as follows:

```cpp
int main(int argc, char *argv[]) {
    ShellOptions Args(argc, argv);
    int nx = Args.GetIntOption("-nx", 123);
    int ny = Args.GetIntOption("-ny", 145);
    double tol = Args.GetDoubleOption("-tol", 1e-12);
    string solver = Args.GetStringOption("-solver");

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

    return 0;
}
```

Each line option can have a value or not. For options with a value, the user can specify this values as follows. Let -tolerance be the name of the option and 1e-12 its value. Both choices are valid:

- `-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 (`-`). Each option cannot have more than one value.

To use this class, the user has to build the database using the `argc`, `argv` input arguments. Then, to retrieve the option value, the user has to use one of the following functions: `GetIntOption`, `GetDoubleOption`, and `GetStringOption`.

If option name is not found in the database, a value of 0, 0.0 or an empty string is returned. If needed, the user can also specify a default value to return when the option name...
is not found in the database. Method HaveOption can be used to query the database for an option.

File `${TRILINOS_HOME}/doc/tutorial/triutils/ex2.cpp` gives an example of the usage of this class.
References


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