## **SANDIA REPORT**

SAND2015-XXXX Unlimited Release Printed January 19, 2015

# **xSDK User Manual**

#### Alicia Marie Klinvex

Prepared by Sandia National Laboratories Albuquerque, New Mexico 87185 and Livermore, California 94550

Sandia National Laboratories is a multi-program laboratory managed and operated by Sandia Corporation, a wholly owned subsidiary of Lockheed Martin Corporation, for the U.S. Department of Energy's National Nuclear Security Administration under contract DE-AC04-94AL85000.

MAN ROSE

Approved for public release; further dissemination unlimited.



Issued by Sandia National Laboratories, operated for the United States Department of Energy by Sandia Corporation.

**NOTICE:** This report was prepared as an account of work sponsored by an agency of the United States Government. Neither the United States Government, nor any agency thereof, nor any of their employees, nor any of their contractors, subcontractors, or their employees, make any warranty, express or implied, or assume any legal liability or responsibility for the accuracy, completeness, or usefulness of any information, apparatus, product, or process disclosed, or represent that its use would not infringe privately owned rights. Reference herein to any specific commercial product, process, or service by trade name, trademark, manufacturer, or otherwise, does not necessarily constitute or imply its endorsement, recommendation, or favoring by the United States Government, any agency thereof, or any of their contractors or subcontractors. The views and opinions expressed herein do not necessarily state or reflect those of the United States Government, any agency thereof, or any of their contractors.

Printed in the United States of America. This report has been reproduced directly from the best available copy.

Available to DOE and DOE contractors from

U.S. Department of Energy

Office of Scientific and Technical Information

P.O. Box 62

Oak Ridge, TN 37831

Telephone: (865) 576-8401 Facsimile: (865) 576-5728

E-Mail: reports@adonis.osti.gov
Online ordering: http://www.osti.gov/bridge

#### Available to the public from

U.S. Department of Commerce National Technical Information Service 5285 Port Royal Rd Springfield, VA 22161

Telephone: (800) 553-6847 Facsimile: (703) 605-6900

E-Mail: orders@ntis.fedworld.gov

Online ordering: http://www.ntis.gov/help/ordermethods.asp?loc=7-4-0#online



### SAND2015-XXXX Unlimited Release Printed January 19, 2015

## xSDK User Manual

Alicia Marie Klinvex

#### Abstract

Some application developers need to be able to use Trilinos together with other libraries, such as PETSc. This is nontrivial because these libraries all expect the data to be stored in different ways, and the way that you call a PETSc KSP linear solver, for instance, looks fundamentally different from the way you would call a Belos linear solver. The IDEAS software productivity project plans to address this problem with the Extreme-scale Scientific Software Development Kit (xSDK). The xSDK will provide an interoperability layer that enables easy installation and combined usage of the IDEAS libraries, including PETSc, Hypre, and SuperLU. This document describes the various interoperability layers and how to install and use the xSDK.

# Introduction

The following are the libraries included in the xSDK. TODO: Maybe each team should write a paragraph or so for this.

Two of the libraries, PETSc and Trilinos are comprehensive numerical software packages<sup>1</sup> that can solve a large variety of problems with both their own algorithmic implementations and wrappers to other numerical libraries. SuperLU and hypre are specialized libraries that provide sophisticated solvers for a restricted family of problems. SuperLU provides sparse direct linear solvers and ILU preconditioners while hypre provides iterative solvers for linear systems.

The xSDK provides a common system for installing and working with all of these packages so, in some sense, xSDK is meta-package.

## **Trilinos**

The Trilinos Project is an effort to develop algorithms and enabling technologies within an object-oriented software framework for the solution of large-scale, complex multi-physics engineering and scientific problems. Trilinos is organized into 66 different packages, each with a specific focus. These packages include linear and nonlinear solvers, preconditioners (including algebraic multigrid), graph partitioners, eigensolvers, and optimization algorithms, among other things. Users are only required to install the subset of packages related to the problems they are trying to solve.

Trilinos supports MPI+X, where X can be CUDA, OpenMP, etc.

## **PETSc**

PETSc is a suite of data structures and routines for the scalable solution of scientific applications modeled by partial differential equations. It includes linear solvers, preconditioners, nonlinear solvers, and ODE integrators. It also provides a variety of scalable constrained and unconstrained optimization solvers. PETSc utilizes the MPI program model and does not use threads. It does not support solving linear systems with multiple right-hand-sides. While it does not include eigensolvers, there is an eigensolver package called SLEPc built on

 $<sup>^{1}</sup>$ Note that the term "Trilinos package" has a specific meaning that, here we just mean package in the generic sense

top of PETSc with a very similar interface. The library libMesh and the framework MOOSE provide finite element solvers that utilize PETSc.

## hypre

Hypre is a library for solving large, sparse linear systems of equations on massively parallel computers. While it contains linear solvers and an eigensolver called LOBPCG, it is probably most well-known for its preconditioners.

Certain subsets of hypre are multithreaded.

## SuperLU

SuperLU is a general purpose library for the direct solution of large, sparse, nonsymmetric systems of linear equations on high performance machines. There are three separate versions of this code: SuperLU (for sequential machines), SuperLU\_MT (for shared memory parallel machines), and SuperLU\_DIST (for distributed memory).

# Chapter 1

# **xSDK** Installation

TODO: Ignore this. Barry is going to take care of the circular dependency issue; then we can revisit the installation procedure.

## Using PETSc configuration script

The easiest way to install the various xSDK libaries (Trilinos, PETSc, Hypre, and SuperLU) is the following three step process:

- 1. Download and unpack the PETSc tarball from http://www.mcs.anl.gov/petsc/, or pull from the repo via git clone https://bitbucket.org/petsc/petsc.
- 2. Run the PETSc configuration script.
- 3. Type make, then make install. Don't worry; the PETSc configuration script will remind you to do this.

The PETSc configuration script is very sophisticated, as you can tell by typing ./configure --help. It will produce a list for you of all the different configuration options (organized into different categories). The most interesting of these configuration options is --download-<PACKAGE>, where PACKAGE can be hypre, superlu, trilinos, etc. If you install PETSc via the following line:

./configure --prefix=<HOME>/petsc-install --download-xsdk
PETSc will download and install hypre, SuperLU, Trilinos, as well as commonly needed dependent package for you, and PETSc itself will be installed to the directory <HOME>/petsc-install.

If PETSc encounters any problems, such as being unable to find your MPI installation, it will output a helpful error message explaining what the problem is and how you can fix it.

If PETSc finds itself unable to download any packages you request (because you are behind a firewall, for instance), it will output the following message explaining how to fix the problem:

\_\_\_\_\_\_

```
Trying to download http://ftp.mcs.anl.gov/pub/petsc/externalpackages/hypre-2.10.0b-p1.tar.gz for HYPRE
______
 Trying to download ftp://ftp.mcs.anl.gov/pub/petsc/externalpackages/hypre-2.10.0b-p1.tar.gz for HYPRE
_____
Unable to download package hypre from: http://ftp.mcs.anl.gov/pub/petsc/externalpackages/hypre-2.10.0b-p1.tar.gz
* If URL specified manually - perhaps there is a typo?
* If your network is disconnected - please reconnect and rerun ./configure
* Or perhaps you have a firewall blocking the download
* Alternatively, you can download the above URL manually, to /yourselectedlocation/hypre-2.10.0b-p1.tar.gz
 and use the configure option:
 --download-hypre=/yourselectedlocation/hypre-2.10.0b-p1.tar.gz
Unable to download package hypre from: ftp://ftp.mcs.anl.gov/pub/petsc/externalpackages/hypre-2.10.0b-p1.tar.gz
* If URL specified manually - perhaps there is a typo?
* If your network is disconnected - please reconnect and rerun ./configure
st Or perhaps you have a firewall blocking the download
* Alternatively, you can download the above URL manually, to /yourselectedlocation/hypre-2.10.0b-p1.tar.gz
 and use the configure option:
 --download-hypre=/yourselectedlocation/hypre-2.10.0b-p1.tar.gz
```

## Without PETSc configuration script

If you would like to manually install Trilinos and enable the PETSc, hypre, and SuperLU interfaces, you may specify those options in the Trilinos configuration script (which I will call do-configure). Note that Trilinos requires all third party libraries to be installed before the configuration process. An example configuration script is below:

Program 1.1. do-configure

```
#!/bin/bash
                         TRILINOS_HOME=/home/amklinv/TrilinosDir/Trilinos
                          PETSC DIR=/home/amklinv/PETSc/petsc-3.6.0
                          PETSC_ARCH=arch-linux2-cxx-debug
                         PETSC_LIB="-Wl,-rpath,/home/amklinv/PETSc/petsc-3.6.0/arch-linux2-cxx-debug/lib -L/home/amklinv/PETSc/petsc-3.6.0/arch-linux2-cxx-debug/lib -lpetsc -Wl,-rpath,/home/amklinv/PETSc/petsc_install/lib -L/home/amklinv/PETSc/petsc_install/lib -lsuperlu_4.3 -lsuperlu_dist_4.0 -lHYPRE -Wl,-rpath,/home/amklinv/lapack-3.5.0 -L/home/amklinv/lapack-3.5.0 -llapack -lrefblas -lparmetis -lmetis -L/projects/install/rhel6-x86_64/sems/compiler/gcc/4.7.2/projects/install/rhel6-x86_64/sems/compiler/gcc/4.7.2/base/lib/gcc/x86_64-unknown-linux-gnu/4.7.2 -L/projects/install/rhel6-x86_64/sems/compiler/gcc/4.7.2/base/lib/gcc/x86_64-unknown-linux-gnu/4.7.2 -L/projects/install/rhel6-x86_64/sems/compiler/gcc/4.7.2/base/lib/gcc/x86_full-linux-gnu/4.7.2 -L/projects/install/rhel6-x86_full-linux-gnu/4.7.2 -L/projects/install/rhel6-x86_full-li
                                                         -Logic transfer of the compiler of the compile
                          PETSC INCLUDE PATH="${PETSC DIR}/${PETSC ARCH}/include:${PETSC DIR}/include:${PETSC DIR}
 11
                           \label{lineary_discrete} \begin{tabular}{ll} HYPRE\_LIBRARY\_DIRS="/home/amklinv/PETSc/petsc-3.6.0/arch-linux2-cxx-debug/external packages/hypre-2.10.0b-p1/src/hypre/lib" | https://scatter.accepts/lineary-linux2-cxx-debug/external packages/hypre-2.10.0b-p1/src/hypre-lib" | https://scatter.accepts/lineary-linux2-cxx-debug/external packages/hypre-2.10.0b-p1/src/hypre-lib" | https://scatter.accepts/lineary-linux2-cxx-debug/external packages/hypre-2.10.0b-p1/src/hypre-lib" | https://scatter.accepts/lineary-linux2-cxx-debug/external packages/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-2.10.0b-p1/src/hypre-
                          HYPRE_INCLUDE_DIRS="/home/amklinv/PETSc/petsc-3.6.0/arch-linux2-cxx-debug/externalpackages/hypre-2.10.0b-p1/src/hypre/
 12
13
                          rm -rf CMakeFiles CMakeCache.txt
14
16
                                      -D CMAKE_BUILD_TYPE=DEBUG \
                                    -D Trilinos_ENABLE_STRONG_C_COMPILE_WARNINGS:BOOL=OFF \
                                     -D CMAKE_INSTALL_PREFIX:PATH="/home/amklinv/TrilinosDir\trilinos-install" \
 19
                                     -D TPL_ENABLE_MPI:BOOL=ON \
                                     -D Trilinos_ENABLE_EXPLICIT_INSTANTIATION:BOOL=ON \
                                     -D Trilinos ENABLE ALL OPTIONAL PACKAGES: BOOL=OFF
                                     -D Trilinos_ENABLE_Amesos:BOOL=ON \
                                    -D Trilinos_ENABLE_Amesos2:BOOL=ON
-D Trilinos_ENABLE_Anasazi:BOOL=ON
                                      -D Trilinos_ENABLE_Aztec00:B00L=0N
                           -D Trilinos_ENABLE_Belos:BOOL=ON \
```

```
-D Trilinos_ENABLE_Epetra:BOOL=ON \
29
       -D Trilinos_ENABLE_EpetraExt:BOOL=ON
       -D Trilinos_ENABLE_Galeri:BOOL=ON \
       -D Trilinos_ENABLE_Ifpack:BOOL=ON \
-D Trilinos_ENABLE_Ifpack2:BOOL=ON
       -D Trilinos_ENABLE_Isorropia:BOOL=ON \
       -D Trilinos_ENABLE_Kokkos:BOOL=ON
       -D Trilinos_ENABLE_ML:BOOL=ON
       -D Trilinos_ENABLE_Tpetra:BOOL=ON \
       -D Trilinos_ENABLE_TrilinosCouplings:BOOL=ON \
37
       -D Trilinos_ENABLE_Triutils:BOOL=ON
39
       -D Trilinos_ENABLE_NOX:BOOL=ON \
40
       -D Trilinos_ENABLE_Zoltan:BOOL=ON
        -D Trilinos_ENABLE_Zoltan2:BOOL=ON
42
43
       -D EpetraExt_USING_PETSC:BOOL=ON \
       -D NOX_ENABLE_PETSC:BOOL=ON '
       -D NOX_ENABLE_ABSTRACT_IMPLEMENTATION_PETSC:BOOL=ON \
-D TPL_ENABLE_PETSC:BOOL=ON \
45
       -D PETSC_LIBRARY_DIRS:FILEPATH="${PETSC_LIB}" \
       -D PETSC_INCLUDE_DIRS:FILEPATH="${PETSC_INCLUDE_PATH}" \
48
        -D TPL_PETSC_LIBRARIES:STRING="${PETSC_LIB}"
       -D TPL_PETSC_INCLUDE_DIRS:STRING="${PETSC_INCLUDE_PATH}" \
51
52
       -D Amesos2_ENABLE_KLU2::BOOL=ON \
53
       -D EpetraExt_ENABLE_HYPRE:BOOL=ON \
54
        -D Ifpack_ENABLE_HYPRE:BOOL=ON
56
       -D TPL ENABLE HYPRE: BOOL=ON \
       D HYPRE_LIBRARY_DIRS:FILEPATH="${HYPRE_LIBRARY_DIRS}" \
-D HYPRE_INCLUDE_DIRS:FILEPATH="${HYPRE_INCLUDE_DIRS}" \
57
59
       -D Belos_ENABLE_TESTS:BOOL=ON \
60
       -D Belos_ENABLE_EXAMPLES:BOOL=ON \
62
       -D Ifpack2_ENABLE_TESTS:BOOL=ON \
        -D Ifpack2_ENABLE_EXAMPLES:BOOL=ON
       -D DART_TESTING_TIMEOUT:STRING=300 \
       -D BLAS LIBRARY NAMES:STRING="libf77blas.so.3" \
65
        -D BLAS_LIBRARY_DIRS:PATH="/usr/lib64/atlas"
       -D LAPACK_LIBRARY_NAMES:STRING="liblapack.so.3" \
-D LAPACK_LIBRARY_DIRS:PATH="/usr/lib64/atlas" \
67
68
\frac{70}{71}
       -D Trilinos_EXTRA_REPOSITORIES:STRING=preCopyrightTrilinos \
-D Trilinos_ENABLE_xSDKTrilinos:BOOL=ON \
        -D xSDKTrilinos_USING_PETSC:BOOL=ON
73
       -D xSDKTrilinos_USING_HYPRE:BOOL=ON \
      ${TRILINOS_HOME}
```

Let's examine this script in more detail. Since we are interested in the PETSc interface, we must first define PETSC\_DIR and PETSC\_ARCH. PETSC\_LIB looks intimidating at first, but you can generate it automatically by typing make getlinklibs in PETSC\_DIR. We then define the PETSc and hypre include paths. Everything else looks pretty normal until we reach lines 43–50, which enable the Epetra-based PETSc interface. Lines 54–58 enable the Epetra-based hypre interface. Then lines 70–73 enable the Tpetra-based interfaces living in the pre-Copyright package xSDKTrilinos.

# Chapter 2

# xSDK Interface Usage

This section describes the individual interfaces and their usage.

## Trilinos-PETSc

There is a two-way interface between PETSc and Trilinos which allows users to use PETSc datatypes with Trilinos and vice-versa.

### Using PETSc Mat and Vec with Trilinos solvers

TODO: Introduction and examples. We have two new interfaces in Trilinos to support using PETSc Mat anywhere a Tpetra::RowMatrix or Tpetra::CrsMatrix can be used. For packages requiring a Tpetra::RowMatrix or Tpetra::Operator, such as Anasazi and Belos, you may wrap a PETSc Mat in our Tpetra::PETScAIJMatrix; otherwise, you can copy it to a Tpetra::CrsMatrix. We will demonstrate each of those functions in the examples below.

Our first example (Program ??) shows how to compute the smallest eigenpairs of a PETSc Mat, specificially Poisson2D, using Trilinos' Anasazi package.

Program 2.1. PETSc\_AnasaziEx.cpp

```
#include "petscksp.h"
#include "AnasaziBasicEigenproblem.hpp"
#include "AnasaziConfigDefs.hpp"
#include "AnasaziTpetraAdapter.hpp"
#include "AnasaziRTRSolMgr.hpp"
#include "Teuchos_ParameterList.hpp"
      #include "Tpetra_PETScAIJMatrix.hpp
      int main(int argc, char **args)
10
         using Teuchos::RCP;
11
         using Teuchos::rcp;
         using std::cout;
14
         using std::endl;
         typedef Tpetra::PETScAIJMatrix <>
                                                                                   PETScAIJMatrix:
         typedef PETScAIJMatrix::scalar_type
typedef PETScAIJMatrix::local_ordinal_type
                                                                                   Scalar:
         typedef PETScAIJMatrix::global_ordinal_type
                                                                                   GO:
         typedef PETScAIJMatrix::node_type
                                                                                   Node;
         typedef Tpetra::Vector < Scalar, LO, GO, Node >
      typedef Tpetra::Map<LO,GO,Node>
```

```
typedef Tpetra::Operator <Scalar, LO, GO, Node>
24
        typedef Tpetra::MultiVector < Scalar, LO, GO, Node >
                                                                      MV:
        typedef Anasazi::RTRSolMgr <Scalar, MV, OP>
        typedef Anasazi::BasicEigenproblem SScalar,MV,OP>
typedef Anasazi::OperatorTraits<Scalar,MV,OP>
26
                                                                      Problem;
28
        typedef Anasazi::MultiVecTraits < Scalar, MV>
29
30
31
        PetscInt
                           m = 50, n = 50;
                          nev = 4;
32
        PetscInt
        PetscErrorCode ierr;
34
        MPI_Comm
                           comm;
35
        PetscInt
                          Istart, Iend, Ii, i, j, J, rank;
37
38
        PetscInitialize(&argc,&args,NULL,NULL);
        ierr = PetscOptionsGetInt(PETSC_NULL, "-mx",&m,PETSC_NULL);CHKERRQ(ierr);
ierr = PetscOptionsGetInt(PETSC_NULL, "-my",&n,PETSC_NULL);CHKERRQ(ierr);
ierr = PetscOptionsGetInt(PETSC_NULL, "-nev",&nev,PETSC_NULL);CHKERRQ(ierr);
40
        ierr = MatCreate(PETSC COMM WORLD.&A):CHKERRQ(ierr):
43
        ierr = MatSetSizes(A, PETSC_DECIDE, PETSC_DECIDE, m*n, m*n); CHKERRQ(ierr);
        ierr = MatSetType(A, MATAIJ); CHKERRQ(ierr)
        ierr = MatMPIAIJSetPreallocation(A,5,PETSC_NULL,5,PETSC_NULL); CHKERRQ(ierr);
46
        ierr = MatSetUp(A); CHKERRQ(ierr);
48
        ierr = PetscObjectGetComm( (PetscObject)A, &comm); CHKERRQ(ierr);
        ierr = MPI_Comm_rank(comm,&rank); CHKERRQ(ierr);
49
51
        ierr = MatGetOwnershipRange(A,&Istart,&Iend);CHKERRQ(ierr);
        52
54
         if (i<m-1) {J = Ii + n; ierr = MatSetValues(A,1,&Ii,1,&J,&v,INSERT_VALUES); CHKERRQ(ierr);}
if (j>0) {J = Ii - 1; ierr = MatSetValues(A,1,&Ii,1,&J,&v,INSERT_VALUES); CHKERRQ(ierr);}
if (j<n-1) {J = Ii + 1; ierr = MatSetValues(A,1,&Ii,1,&J,&v,INSERT_VALUES); CHKERRQ(ierr);}
55
57
          v = 4.0; ierr = MatSetValues(A,1,&Ii,1,&Ii,&v,INSERT_VALUES);CHKERRQ(ierr);
59
        }
60
        ierr = MatAssemblyBegin(A, MAT_FINAL_ASSEMBLY); CHKERRQ(ierr);
        ierr = MatAssemblyEnd(A,MAT_FINAL_ASSEMBLY); CHKERRQ(ierr);
63
        RCP < PETScAIJMatrix > tpetraA = rcp(new PETScAIJMatrix(A));
65
66
        RCP<MV> initGuess = rcp(new MV(tpetraA->getDomainMap(),4,false));
        initGuess->randomize();
68
69
        RCP < Problem > problem = rcp(new Problem(tpetraA,initGuess));
\frac{70}{71}
        problem -> setNEV(nev);
        problem ->setHermitian(true);
72
        problem -> setProblem();
73\\74
        Teuchos::ParameterList pl;
        Pol.set("Verbosity", Anasazi::IterationDetails + Anasazi::FinalSummary);
pl.set("Convergence Tolerance", 1e-6);
RCP<SolMgr> solver = rcp(new SolMgr(problem, pl));
79
        Anasazi::ReturnType returnCode = solver->solve();
        Anasazi::Eigensolution < Scalar, MV > sol = problem ->getSolution();
80
        std::vector<Anasazi::Value<Scalar> > evals = sol.Evals;
82
        RCP < MV > evecs = sol.Evecs;
        ierr = PetscFinalize(); CHKERRQ(ierr);
85
        return 0:
```

#### Lines 1–7 Include statements

Lines 11–28 Typedefs and using statements to make the code more readable

#### Lines 30–36 PETSc datatypes

Lines 38–41 Get the command line arguments using PETSc. This example has three of them: the number of mesh points in the x direction, number of mesh points in the y direction, and the number of desired eigenpairs.

#### Lines 43–62 Create the PETSc Mat and set its values.

**Line 64** Wrap the PETSc Mat in a Tpetra::PETScAIJMatrix. Since Anasazi only requires a Tpetra::Operator<sup>1</sup>, we do not have to deep copy the data to a Tpetra::CrsMatrix.

**Line 66** Create a random initial guess for the eigensolver. Note that we can treat tpetraA just like any other Tpetra::RowMatrix and obtain its domain map via getDomainMap().

#### Is the data copied or wrapped?

If you are using a part of Trilinos that requires Operator or RowMatrix, the data is wrapped. If you need a CrsMatrix specifically, the data is deep-copied.

### Using Trilinos datatypes with PETSc KSP solvers

If you would like to use Trilinos datatypes, such as Tpetra::Operator and Tpetra::MultiVector, with a PETSc KSP linear solver, you may use our new Belos interface: PETScSolMgr. This interface is very similar to that of the other native Belos linear solvers, which makes solving linear systems such as AX = B a simple process.

- 1. (Optional) Create a Tpetra::Operator for the preconditioner  $M \approx A$ . You may use the preexisting preconditioners of Ifpack2 and MueLu, or you may create your own custom preconditioner. Alternatively, you may choose not to use a preconditioner at all.
- 2. Create a Belos::LinearProblem containing the operator A, the initial guess X, the right-hand side B, and the preconditioner M (if you have one).
- 3. Create a Teuchos::ParameterList containing the parameters you wish to set. These parameters are summarized in Table 2.4.
- 4. Create a Belos::PETScSolMgr with the LinearProblem and ParameterList from the previous steps.
- 5. Call solve()

The following example (Program 2.2) illustrates this process in greater detail.

<sup>&</sup>lt;sup>1</sup>RowMatrix is a specific type of Operator, and CrsMatrix is a specific type of RowMatrix. Therefore, you can use a RowMatrix anywhere an Operator is accepted, but you can't necessarily use a RowMatrix anywhere a CrsMatrix is expected.

Parameter	Description	Default Value					
Maximum	integer defining the maximum number of itera-	1000					
Iterations	tions to be performed.						
Solver	string defining the linear solver to be used.	KSPGMRES					
	A list of all valid linear solver options can be						
	found at <pre>http://www.mcs.anl.gov/petsc/</pre>						
	<pre>petsc-current/docs/manualpages/KSP/</pre>						
	KSPType.html						
Verbosity	Belos::MsgType defining the amount of out-	Belos::Errors					
	put the program should produce. Op-						
	tions include Belos::Errors, Belos::Warnings, Be-						
	los::IterationDetails, Belos::TimingDetails, and						
	Belos::StatusTestDetails						
Convergence	double defining the tolerance of the linear solver	$10^{-8}$					
Tolerance							

Table 2.1. Belos::PETScSolMgr parameters

Program 2.2. Tpetra\_KSPEx.cpp

```
#include "BelosTpetraAdapter.hpp"
#include "BelosPETScSolMgr.hpp"
#include "Ifpack2_Factory.hpp"
#include "Teuchos_CommandLineProcessor.hpp"
 3
       #include "Teuchos_ParameterList.hpp"
       #include "Tpetra_CrsMatrix.hpp"
#include "Tpetra_DefaultPlatform.hpp"
       #include "Tpetra_MultiVector.hpp"
#include "MatrixMarket_Tpetra.hpp"
 9
10
       int main(int argc, char *argv[]) {
          typedef double
typedef Teuchos::ScalarTraits<ST>
12
                                                                             ST:
13
14
          typedef SCT::magnitudeType
          typedef Tpetra::MultiVector<>
typedef Tpetra::Operator<>
15
                                                                             MV:
          typedef Belos::MultiVecTraits<ST,MV>
17
                                                                           MVT;
          typedef Belos::OperatorTraits<ST,MV,OP> OPT;
18
19
          typedef Tpetra::CrsMatrix<>
20
          typedef Ifpack2::Preconditioner<>
                                                                         Prec;
^{-2}_{21}
22
          using Teuchos::ParameterList;
          using Teuchos::RCP;
using Teuchos::rcp;
23
25
26
          Teuchos::oblackholestream blackhole:
          Teuchos::GlobalMPISession mpiSession (&argc, &argv, &blackhole);
28
          RCP < const Teuchos::Comm < int > > comm = Tpetra::DefaultPlatform::getDefaultPlatform().getComm();
29
          const int myRank = comm->getRank();
          double tol = 1e-6;
31
         std::string filename("/home/amklinv/matrices/cage4.mtx");
std::string ksptype("gmres");
Teuchos::CommandLineProcessor cmdp(false,false);
cmdp.setOption("filename",&filename,"Filename for test matrix.");
cmdp.setOption("tol",&tol,"Relative residual tolerance.");
cmdp.setOption("ksptype",&ksptype,"Type of linear solver to be used.");
if (cmdp.parse(argc,argv) != Teuchos::CommandLineProcessor::PARSE_SUCCESSFUL) {
    return -1;
}
32
          std::string filename("/home/amklinv/matrices/cage4.mtx");
34
35
36
37
38
39
40
42
          RCP<CrsMatrix> A = Tpetra::MatrixMarket::Reader<CrsMatrix>::readSparseFile(filename,comm);
          RCP<MV> B = rcp(new MV(A->getRowMap(),1,false));
RCP<MV> X = rcp(new MV(A->getRowMap(),1,false));
43
          MVT::MvInit(*X);
MVT::MvInit(*B,1);
45
46
          Ifpack2::Factory factory;
```

```
RCP < Prec > M = factory.create("RELAXATION", A.getConst());
         ParameterList ifpackParams;
50
         ifpackParams.set("relaxation: type","Jacobi");
         M->setParameters(ifpackParams);
M->initialize();
         M->compute();
55
         ParameterList belosList;
         belosList.set( "Maximum Iterations", 100 );
belosList.set( "Convergence Tolerance", tol );
belosList.set( "Solver", ksptype );
58
         RCP<Belos::LinearProblem <double,MV,OP> > problem
= rcp( new Belos::LinearProblem <double,MV,OP>( A, X, B ) );
61
         problem -> setLeftPrec( M );
63
64
         problem -> setProblem();
         RCP< Belos::PETScSolMgr<double,MV,OP> > solver
= rcp( new Belos::PETScSolMgr<double,MV,OP>(problem, rcp(&belosList,false)) );
         solver -> solve();
```

#### Lines 1–9 Include statements

Lines 12–24 Typedefs and using statements to make the code more readable

Lines 26–29 Set up MPI

Lines 31-40 Parse command line arguments. This program allows the user to specify the filename for the matrix, the tolerance for the linear solve, and which linear solver is used. A list of all valid linear solver options can be found at http://www.mcs.anl.gov/petsc/petsc-current/docs/manualpages/KSP/KSPType.html.

**Lines 42–46** Set up the linear system by reading the matrix from a file, setting the initial guess for the solution as  $\vec{0}$  and setting the right hand side as  $\vec{1}$ .

Lines 48–54 Set up the Ifpack2 Jacobi preconditioner.

**Lines 56–59** Set the maximum number of iterations, convergence tolerance, and which PETSc KSP solver is being used.

Lines 61–64 Set up the linear problem for the Belos solver.

Lines 66-68 Solve the linear system.

### Can I use this to solve linear systems with multiple right-hand sides?

Yes. Unfortunately, PETSc has no support for multivectors at this time, so each of the right hand sides will be processed independently. If you want block or pseudo-block linear solvers, those are available within Trilinos.

#### Is the data copied or wrapped?

The raw Tpetra matrix (or operator) data is wrapped rather than deep copied.

# Trilinos-hypre

#### TODO: Introduction.

In this example (Program 2.3), we will examine how to use hypre solvers and preconditioners with Tpetra objects.

#### Program 2.3. Hypre\_SolveEx.cpp

```
#include "Ifpack2_Preconditioner.hpp"
      #include "Ifpack2_Hypre.hpp"
#include "Teuchos_CommandLineProcessor.hpp"
      #include "Tpetra_CrsMatrix.hpp"
#include "Tpetra_DefaultPlatform.hpp"
      int main(int argc, char *argv[]) {
         using Teuchos::Array;
         using Teuchos::RCP;
         using Teuchos::rcp;
11
         using Teuchos::ParameterList;
         using Ifpack2::FunctionParameter;
         using Ifpack2::Hypre::Prec;
         using Ifpack2::Hypre::Solver;
         typedef Tpetra::CrsMatrix<>::scalar_type Scalar;
         typedef Tpetra::CrsMatrix<>::local_ordinal_type LO;
17
         typedef Tpetra::CrsMatrix <>::global_ordinal_type GO;
         typedef Tpetra::CrsMatrix<>::node_type Node;
typedef Tpetra::DefaultPlatform::DefaultPlatformType Platform;
         typedef Tpetra::CrsMatrix<Scalar> CrsMatrix;
         typedef Tpetra::MultiVector<Scalar> MV;
typedef Ifpack2::Preconditioner<Scalar> Preconditioner;
24
25
         typedef Tpetra::Map<> Map;
         Teuchos::oblackholestream blackhole;
         Teuchos::GlobalMPISession mpiSession(&argc,&argv,&blackhole);
Platform &platform = Tpetra::DefaultPlatform::getDefaultPlatform();
         RCP < const Teuchos::Comm < int > > comm = platform.getComm();
31
         // Get parameters from command-line processor
int nx = 10;
Scalar tol = 1e-6;
Teuchos::CommandLineProcessor cmdp(false,true);
         cmdp.setOption("nx",&nx, "Number of mesh points in x direction.");
cmdp.setOption("tolerance",&tol, "Relative residual used for solver.");
         if(cmdp.parse(argc,argv) != Teuchos::CommandLineProcessor::PARSE_SUCCESSFUL) {
        return -1;
39
41
         // Create the 2D Laplace operator
         int n = nx*nx;
         RCP < Map > map = rcp(new Map(n,0,comm));
RCP < CrsMatrix > A = rcp(new CrsMatrix(map,5));
         for(LO i = 0; i<nx; i++) {
          for(LO j = 0; j<nx; j++) {
```

```
GO row = i*nx+j;
 49
                  if (!map->isNodeGlobalElement(row))
                     continue:
 51
                  Array <LO> indices;
 52
                  Array < Scalar > values;
 53
 54
 55
                  if(i > 0) {
 56
                     indices.push_back(row - nx);
 57
                     values.push_back(-1.0);
                  if(i < nx-1) {
 59
                     indices.push_back(row + nx);
 60
                     values.push_back(-1.0);
 62
 63
                  indices.push_back(row);
                  values.push_back(4.0);
 65
                  if(j > 0) {
 66
                     indices.push_back(row-1);
67
68
                     values.push_back(-1.0);
 69
                  if(j < nx-1) {
 70
                      indices.push_back(row+1);
 71
                     values.push_back(-1.0);
 73
74
                  A->insertGlobalValues(row,indices,values);
 75
76
77
           A->fillComplete();
 79
           \label{eq:rcp(new MV(A->getRowMap(),1));} $$ RCP < MV > X = rcp(new MV(A->getRowMap(),1));
            RCP<MV> B = rcp(new MV(A->getRowMap(),1,false));
 80
            B->randomize();
 82
            // Create the parameters for hypre
 84
            RCP < FunctionParameter > functs[10];
           functs[0] = rcp(new FunctionParameter(Prec,
functs[1] = rcp(new FunctionParameter(Prec,
                                                                                     &HYPRE_BoomerAMGSetPrintLevel, 1)); // print AMG solution info &HYPRE_BoomerAMGSetCoarsenType, 6)); // Falgout coarsening
 85
           functs[2] = rcp(new FunctionParameter(Prec, functs[3] = rcp(new FunctionParameter(Prec, functs[4] = rcp(new FunctionParameter(Prec,
                                                                                     &HYPRE_BoomerAMGSetRelaxType, 6)); // Sym GS/Jacobi hybrid &HYPRE_BoomerAMGSetNumSweeps, 1)); // Sweeps on each level
 87
 88
                                                                                      &HYPRE_BoomerAMGSetTol, 0.0));
           functs[5] = rcp(new FunctionParameter(Prec, &HTFRE_BOOMETANGSetMaxIter, 1));
functs[6] = rcp(new FunctionParameter(Solver, &HYPRE_PCGSetMaxIter, 1000));
functs[7] = rcp(new FunctionParameter(Solver, &HYPRE_PCGSetTol, tol));
functs[8] = rcp(new FunctionParameter(Solver, &HYPRE_PCGSetTwoNorm, 1));
 90
                                                                                                                                                  // Do only one iteration
// Maximum iterations
 91
                                                                                                                                                   // Convergence tolerance
                                                                                                                                                  // Use the two-norm as the stopping
            functs[9] = rcp(new FunctionParameter(Solver, &HYPRE_PCGSetPrintLevel, 2));
 94
                                                                                                                                                  // Print solve info
 95
           // Create the hypre solver and preconditioner
RCP
RCP
RCP
RCP
RCP
RCP
RCP
RCP
FrameterList hypreList;
hypreList.set("SolveOrPrecondition", Solver);
hypreList.set("Solver", Ifpack2::Hypre::PCG);
hypreList.set("Preconditioner", Ifpack2::Hypre::BoomerAMG);
hypreList.set("SetPreconditioner", true);
hypreList.set("NumFunctions", 10);
hypreList.set
RCP
FunctionParameter>
**("Functions", functs);
 96
 98
100
101
102
103
            hypreList.set < RCP < FunctionParameter > *> ("Functions", functs);
104
105
            prec -> setParameters(hypreList);
106
            prec -> compute();
108
           // Perform solve
prec -> apply (*B,*X);
109
           return 0;
112
```

#### Lines 1–5 Include statements

Lines 8–24 Typedefs and using statements to make the code more readable

Lines 26–30 Set up MPI

Lines 32–40 Parse command line arguments. This program allows the user to specify how large the problem should be and how accurately the linear system should be solved.

Lines 42–76 Set up the 2D Laplace operator.

Lines 78–81 Create a random right-hand-side and initialize the solution vector to 0.

Lines 83–94 Set hypre options (documented in the hypre user and reference manuals found at http://computation.llnl.gov/project/linear\_solvers/software.php). In this example, we have elected to use the conjugate gradient method with an algebraic multigrid preconditioner. We have specified a particular coarsening and relaxation type. The most important thing to note about BoomerAMG is that if you would like to use it as a preconditioner, you must set its maximum number of iterations to 1; otherwise, hypre will assume you meant to use it as a linear solver. We then set the tolerance and maximum number of iterations for hypre's conjugate gradient solver.

Lines 96–106 Create the hypre solver. Line 99 specifies that we will be using a hypre linear solver, and line 100 says it will be the conjugate gradient method. Line 101 says we would also like to use BoomerAMG. Remember that you must also set "SetPreconditioner" to true, or the preconditioner will not be used. Lines 103 and 104 specify the hypre parameters such as print level, tolerance, and maximum number of iterations.

**Lines 108–109** Solve the linear system. apply actually calls the hypre linear solve routine PCG with a BoomerAMG preconditioner, as we specified above.

In the next example (Program 2.4), we will examine how to use hypre preconditioners with Belos solvers.

Program 2.4. Hypre\_BelosEx.cpp

```
#include "BelosTpetraAdapter.hpp"
     #include "BelosPseudoBlocKCGSolMgr.hpp"

#include "Ifpack2_Preconditioner.hpp"

#include "Ifpack2_Hypre.hpp"

#include "Teuchos_CommandLineProcessor.hpp"

#include "Teuchos_CommandLineProcessor.hpp"
     #include "Tpetra_DefaultPlatform.hpp"
     int main(int argc, char *argv[]) {
10
       using Teuchos::Array;
        using Teuchos::RCP;
11
        using Teuchos::rcp;
13
        using Teuchos::ParameterList;
        using Ifpack2::FunctionParameter;
14
        using Ifpack2::Hypre::Prec;
        typedef Tpetra::CrsMatrix<>::scalar_type Scalar;
        typedef Tpetra::CrsMatrix<>::local_ordinal_type LO;
        typedef Tpetra::CrsMatrix<>::global_ordinal_type GO;
typedef Tpetra::CrsMatrix<>::node_type Node;
19
21
        typedef Tpetra::DefaultPlatform::DefaultPlatformType Platform;
        typedef Tpetra::CrsMatrix < Scalar > CrsMatrix;
        typedef Tpetra::MultiVector < Scalar > MV;
        typedef Tpetra::Operator < Scalar > OP;
25
        typedef Ifpack2::Preconditioner < Scalar > Preconditioner;
        typedef Tpetra::Map<> Map;
        typedef Belos::PseudoBlockCGSolMgr < Scalar, MV, OP > Solver;
        Teuchos::oblackholestream blackhole;
```

```
{\tt Teuchos::GlobalMPISession\ mpiSession\,(\&argc\,,\&argv\,,\&blackhole)\,;}
 31
 32
         Platform &platform = Tpetra::DefaultPlatform::getDefaultPlatform():
 33
         RCP < const Teuchos::Comm < int > > comm = platform.getComm();
 34
 35
          // Get parameters from command-line processor
         int nx = 10;
Scalar tol = 1e-6;
 36
 37
         bool verbose = false;
 38
 39
         Teuchos::CommandLineProcessor cmdp(false,true);
         40
 42
 43
           return -1;
 44
         }
 45
 46
 47
          // Create the 2D Laplace operator
 48
         int n = nx*nx;
RCP < Map > map = rcp(new Map(n,0,comm));
 49
         RCP<CreMatrix> A = rcp(new CrsMatrix(map,5));
for(LO i = 0; i<nx; i++) {
  for(LO j = 0; j<nx; j++) {
 50
 51
 52
 53
              GO row = i*nx+j;
              if(!map->isNodeGlobalElement(row))
 54
 55
                continue:
 56
              Array < LO > indices;
 57
              Array < Scalar > values;
 59
 60
              if(i > 0) {
 61
               indices.push_back(row - nx);
 62
                values.push_back(-1.0);
 63
              if(i < nx-1) {
 64
                indices.push_back(row + nx);
 65
                 values.push_back(-1.0);
 67
              indices.push back(row):
 68
 69
              values.push_back(4.0);
 70
71
              if(j > 0) {
                indices.push_back(row-1);
 72
                 values.push_back(-1.0);
 \frac{73}{74}
              if(j < nx-1) {
 75
                 indices.push_back(row+1);
 76
                values.push_back(-1.0);
 77
 78
              A->insertGlobalValues(row,indices,values);
           }
 79
 80
 81
         A->fillComplete();
 82
 83
          // Create the vectors
         RCP<MV> X = rcp(new MV(A->getRowMap(),1));
RCP<MV> B = rcp(new MV(A->getRowMap(),1,false));
 84
 85
 86
         B->randomize();
 87
 88
                      the parameters
 89
         RCP < FunctionParameter > functs[6];
         functs[0] = rcp(new FunctionParameter(Prec, &HYPRE_BoomerAMGSetPrintLevel, 1)); // print AMG solution info
functs[1] = rcp(new FunctionParameter(Prec, &HYPRE_BoomerAMGSetCoarsenType, 6)); // Falgout coarsening
 90
         functs[2] = rcp(new FunctionParameter(Prec, &HYPRE_BoomerAMGSetRelarType, 6)); // Sym GS/Jacobi hybrid functs[3] = rcp(new FunctionParameter(Prec, &HYPRE_BoomerAMGSetNumSweeps, 1)); // Sweeps on each level functs[4] = rcp(new FunctionParameter(Prec, &HYPRE_BoomerAMGSetTol, 0.0)); // Conv tolerance zero functs[5] = rcp(new FunctionParameter(Prec, &HYPRE_BoomerAMGSetMaxIter, 1)); // Do only one iteration
 92
 93
 95
                                                                                                                // Do only one iteration!
 96
 98
         RCP<Preconditioner> prec = rcp(new Ifpack2::Ifpack2_Hypre<Scalar,LO,GO,Node>(A));
ParameterList hypreList;
 99
         hypreList.set("SolveOrPrecondition", Prec);
hypreList.set("Preconditioner", Ifpack2::Hypre::BoomerAMG);
hypreList.set("NumFunctions", 6);
100
101
102
103
         hypreList.set<RCP<FunctionParameter>*>("Functions", functs);
104
         prec -> setParameters(hypreList);
105
         prec -> compute();
106
107
             Create the linear problem
         RCP< Belos::LinearProblem<Scalar,MV,OP> > problem = rcp(new Belos::LinearProblem<Scalar,MV,OP>(A,X,B));
108
         problem ->setHermitian();
109
         problem -> setLeftPrec(prec);
110
         problem -> setProblem();
112
113
          // Create the Belos linear solver
         RCP<ParameterList> belosList = rcp(new ParameterList());
114
115
         belosList->set("Convergence Tolerance", tol);
         if(verbose)
116
117
            belosList->set("Verbosity", Belos::Errors + Belos::Warnings + Belos::TimingDetails + Belos::StatusTestDetails);
118
            belosList->set("Verbosity", Belos::Errors + Belos::Warnings);
120
         RCP < Solver > newSolver = rcp(new Solver(problem, belosList));
121
```

```
123 newSolver->solve();
124
125 return 0;
126 }
```

Lines 1–86 These lines are not substantially different from the previous example. Again, we set up our operator, solution vector, and right hand side.

Lines 88–95 This time, we have elected not to use a hypre linear solver, so we only set the parameters related to the AMG preconditioner. Again, it is very important to set the maximum number of iterations if you wish to use AMG as a preconditioner.

Lines 97–105 Create the hypre preconditioner. This time, we specify that we would like to precondition rather than solve, since we will be using a Belos linear solver.

Lines 107–111 Create a Belos::LinearProblem that encapsulates the operator, solution vector, right-hand side, and preconditioner. We also specify that our operator is Hermitian so that Belos allows us to use PCG.

Lines 113–120 Create a Belos linear solver. The Belos solvers have many parameters, but we only specify the convergence tolerance and verbosity (what information will be printed).

Lines 122–123 Solve the linear system using a Belos pseudo-block conjugate gradient solver with hypre's BoomerAMG preconditioner.

## Is the data wrapped or copied?

The Tpetra matrix data is deep-copied to a hypre matrix.

# Trilinos-SuperLU

TODO.

Option	Arguments	Default	Description
$-mat\_superlu\_dist\_statprint$	Yes/No	No	Prints statistics on factorization
Someone needs to with -help and add the rest			

Table 2.2. PETSc SuperLU\_Dist Options

## PETSc to Other Packages

PETSc uses run-time binding of data structures and solver algorithms which means that in most causes you do not need to change your simulation code to switch between solvers in hypre, SuperLU, and Trilinos. You use the PETSc options database (for example command line arguments) to select your solvers; it is also possible to hardware particular solvers into the source code but we do not recommend this approach. The solvers in the other xSDK packages are interfaced to PETSc in the PETSc abstract PC preconditioners interface<sup>2</sup>. To select a particular preconditioner one uses

-pc\_type typename [-pc\_\* other potential options related to the solver]

### PETSc utilizing SuperLU

PETSc can use both SuperLU and SuperLU\_Dist solvers. Since the SuperLU\_Dist solvers are parallel and in general are faster and require less memory than SuperLU we will on discuss the details of utilizing SuperLU\_Dist. It is utilized with

-pc\_type lu -pc\_factor\_mat\_solver\_package superlu\_dist [-mat\_superlu\_dist\_\* SuperLU\_Dist

All the options will be printed if you run with -pc\_type lu -pc\_factor\_mat\_solver\_package superlu\_dist -help | grep superlu\_dist. We also display many of the options below

## PETSc utilizing hypre

PETSc utilizes several of the solvers/preconditioners in PETSc via the command

-pc\_type hypre -pc\_hypre\_type [boomeramg (default),pilut,parasails,ams,ads]

Since BoomerAMG is the crown jewel of hypre we recommend its use whenever possible and do not discuss here the use of the other solvers. Running with -pc\_type hypre -help | grep hypre for example will show the options for the pilut preconditioner.

<sup>&</sup>lt;sup>2</sup>This same interface is used for direct solvers

Option	Arguments	Default	Description
-pc_hypre_boomeramg_max_levels	l	Problem dependent	Sets maximum numb
Someone needs to with -help and add the rest			

Table 2.3. PETSc hypre BoomerAMG Options

Option	Arguments	Default	Description
-pc_ml_maxNlevels	l	10	Sets maximum number of levels
Someone needs to with -help and add the rest			

Table 2.4. PETSc ML Options

## PETSc utilizing Trilinos

PETSc can utilizes the ML solver in Trilinos.

-pc\_type ml [-pc\_ml\_\* options for ML]

Running with -pc\_type ml -help | grep pc\_ml will show the available options.

