Installation guide for esys-Escript

Release - 5.7 (r1613964016)

Escript development team

February 22, 2021

Centre for Geoscience Computing (GeoComp)
The University of Queensland
Brisbane, Australia

Email: esys@esscc.uq.edu.au

Guide to Documentation

Documentation for esys.escript comes in a number of parts. Here is a rough guide to what goes where.

 $\textbf{install.pdf} \quad \text{``Installation guide for } \textit{esys-Escript''} \text{: Instructions for compiling } \textit{escript}$

for your system from its source code.

cookbook.pdf "The escript COOKBOOK": An introduction to escript for new users

from a geophysics perspective.

user.pdf "esys-Escript User's Guide: Solving Partial Differential Equations with

Escript and Finley": Covers main escript concepts.

inversion.pdf "esys.downunder: Inversion with escript": Explanation of the

inversion toolbox for escript.

sphinx_api directory Documentation for *escript Python* libraries.

escript_examples(.tar.gz)/(.zip) Full example scripts referred to by other parts of the documentation.

doxygen directory Documentation for C++ libraries (mostly only of interest for develop-

ers).

Contents

1	Intro	roduction	5
2	Insta	alling from Flatpak	7
3	Insta	alling inside Anaconda	9
4	Insta	alling from Docker	11
5	Insta	alling from Source	13
	5.1	Parallel Technologies	13
		5.1.1 What parallel technology do I need?	13
	5.2	MacOS	14
	5.3	Building	14
		5.3.1 Debian	14
		5.3.2 Ubuntu	15
		5.3.3 OpenSuse	15
		5.3.4 CentOS	16
		5.3.5 Fedora	16
		5.3.6 MacOS 10.10 and later (macports)	17
		5.3.7 MacOS 10.13 and later (homebrew)	17
		5.3.8 FreeBSD	17
		5.3.9 Windows	17
		5.3.10 Other Systems / Custom Builds	19
	5.4	Cleaning up	20
	5.5	Optional Extras	20
		5.5.1 Trilinos	20
	5.6	Testing esys.escript	21
A	Requ	uired compiler features	23
В	Trili	inos	25
		Debian "buster" configuration	25
		B.1.1 Required packages	25
		B.1.2 Example configuration file (without MPI)	25
		B.1.3 Example configuration file (with MPI)	26

Contents 3

4 Contents

Introduction

This document describes how to install esys- $Escript^1$ on to your computer. To learn how to use Escript please see the Cookbook, User's guide or the API documentation.

Escript is primarily developed on Linux desktop, SGI ICE and MacOS X systems. It can be installed in several ways:

- 1. Binary packages ready to run with no compilation required. These are available in Debian and Ubuntu repositories, so just use your normal package manager (so you don't need this guide). They are also available for Anaconda Python 3.
- 2. Using flatpak
- 3. From source that is, it must be compiled for your machine. This is the topic of this guide.

See the site https://answers.launchpad.net/escript-finley for online help. Chapter 5 covers installing from source. Appendix A lists some c++ features which your compiler must support in order to compile escript. This version of escript has the option of using Trilinos in addition to our regular solvers. Appendix B covers features of Trilinos which escript needs.

Chapter 1. Introduction

¹For the rest of the document we will drop the *esys*-

Installing from Flatpak

To install esys.escript on any linux distribution using flatpak¹, type

flatpak install flathub au.edu.uq.esys.escript

This wil download and install esys.escript on your machine. The esys.escript build installed utilises both the Trilinos and PASO solver libraries, with openMP but without OPENMPI.

After flatpak has finished downloading and installing esys.escript, you can launch an esys.escript window from the menu or run esys.escript in a terminal with the command:

flatpak run au.edu.uq.esys.escript [other arguments to pass to escript]

Finally, to uninstall esys.escript from your machine, type

flatpak uninstall escript

¹For most linux distributions this can be installed from the repository. Otherwise, flatpak is available at https://flathub.org/home

CHAPTER **THREE**

Installing inside Anaconda

There are precompiled binaries of esys.escript available for Anaconda python (for Linux), available in the conda-forge channel on Anaconda cloud. They can be installed using the command

conda install esys-escript -c conda-forge

Installing from Docker

To install an esys.escript Docker container on your machine, first install Docker and then type:

```
docker pull esysescript/esys-escript
```

Once installed, you can launch an escript session using the command:

```
docker run -ti esysescript/esys-escript run-escript
```

If you would also like to mount a folder, you can use the command:

```
docker run -ti -v $(pwd):/app/ esysescript/esys-escript run-escript
```

Additionally, if you wish to run an escript named test_program.py, you can use the command:

Installing from Source

This chapter describes installing esys.escript from source on unix/posix like systems (including MacOSX) and Windows 10.

5.1 Parallel Technologies

It is likely that the computer you run esys.escript on, will have more than one processor core. Escript can make use of multiple cores [in order to solve problems more quickly] if it is told to do so, but this functionality must be enabled at compile time. Section 5.1.1 gives some rough guidelines to help you determine what you need. There are two technologies which esys.escript can employ here.

- OpenMP the more efficient of the two [thread level parallelism].
- MPI Uses multiple processes (less efficient), needs less help from the compiler (not supported on Windows).

Escript is primarily tested on recent versions of the GNU and Intel suites ("g++"/"icpc"), and Microsoft Visual C++ (MSVC). However, it also passes our tests when compiled using "clang++". Escript now requires compiler support for some features of the C++11 standard. See Appendix A for a list.

Our current test compilers include:

- g++ 8
- clang++ 7
- intel icpc v17
- MSVC 2017 or 2019

Note that:

- OpenMP will not function correctly for $g++ \le 4.2.1$ (and is not currently supported by clang).
- icpc v11 has a subtle bug involving OpenMP and C++ exception handling, so this combination should not be used.

5.1.1 What parallel technology do I need?

If you are using any version of Linux released in the past few years, then your system compiler will support OpenMP with no extra work (and give better performance); so you should use it. MSVC 2017 and 2019 also have OpenMP support on Windows (OpenMP 2.0). You will not need MPI unless your computer is some form of cluster. MPI is not recommended on Windows as it will interfer with Jupyter.

If you are using BSD or MacOSX and you are just experimenting with esys.escript, then performance is probably not a major issue for you at the moment so you don't need to use either OpenMP or MPI. This also

applies if you write and polish your scripts on your computer and then send them to a cluster to execute. If in the future you find escript useful and your scripts take significant time to run, then you may want to recompile esys.escript with more options.

Note that even if your version of esys.escript has support for OpenMP or MPI, you will still need to tell the system to use it when you run your scripts. If you are using the run-escript launcher, then this is controlled by the -t, -p, and -n flags. If not, then consult the documentation for your MPI libraries (or the compiler documentation in the case of OpenMP 1).

If you are using MacOSX, then see the next section, if not, then skip to Section 5.3.

5.2 MacOS

This release of esys.escript has only been tested on OSX 10.13. For this section we assume you are using either homebrew or MacPorts as a package manager². You can of course install prerequisite software in other ways. For example, we have had *some* success changing the default compilers used by those systems. However this is more complicated and we do not provide a guide here.

Both of those systems require the XCode command line tools to be installed³.

5.3 Building

Escript is built using *SCons*. To simplify the installation process, we have prepared *SCons_options.py* files for a number of common systems⁴. The options files are located in the *scons/templates* directory. We suggest that the file most relevant to your OS be copied from the templates directory to the scons directory and renamed to the form XXXX_options.py where XXXX should be replaced with your computer's (host-)name. If your particular system is not in the list below, or if you want a more customised build, see Section 5.3.10 for instructions.

```
Debian - 5.3.1Ubuntu - 5.3.2
```

• OpenSuse - 5.3.3

• Centos - 5.3.4

• Fedora - 5.3.5

• MacOS (macports) - 5.3.6

• MacOS (homebrew) - 5.3.7

• FreeBSD - 5.3.8

• Windows - 5.3.9

Once these are done proceed to Section 5.4 for cleanup steps.

All of these instructions assume that you have obtained the <code>esys.escript</code> source (and uncompressed it if necessary).

5.3.1 Debian

These instructions were prepared on Debian 10 Buster.

As a preliminary step, you should install the dependencies that Escript requires from the repository. If you intend to use Python 2.7, then you should install the following

14 5.2. MacOS

¹It may be enough to set the OMP_NUM_THREADS environment variable.

²Note that package managers will make changes to your computer based on programs configured by other people from various places around the internet. It is important to satisfy yourself as to the security of those systems.

³As of OSX10.9, the command xcode-select --install will allow you to download and install the commandline tools.

⁴These are correct a time of writing but later versions of those systems may require tweaks. Also, these systems represent a cross section of possible platforms rather than meaning those systems get particular support.

```
sudo apt-get install python-dev python-numpy python-pyproj python-gdal sudo apt-get install python-sympy python-matplotlib python-scipy sudo apt-get install libboost-python-dev libboost-random-dev sudo apt-get install libnetcdf-dev libnetcdf-cxx-legacy-dev libnetcdf-c++4-dev sudo apt-get install scons lsb-release libsuitesparse-dev gmsh
```

If you intend to use Python 3.0+, then you should install the following

```
sudo apt-get install python3-dev python3-numpy python3-pyproj python3-gdal sudo apt-get install python3-sympy python3-matplotlib python3-scipy sudo apt-get install libboost-python-dev libboost-random-dev sudo apt-get install libnetcdf-dev libnetcdf-cxx-legacy-dev libnetcdf-c++4-dev sudo apt-get install libsuitesparse-dev scons lsb-release gmsh
```

In the source directory execute the following (substitute buster_py2 or buster_py3 for XXXX):

```
scons -j1 options_file=scons/templates/XXXX_options.py
```

If you wish to test your build, you can use the following:

```
scons -j1 py_tests options_file=scons/templates/XXXX_options.py
```

5.3.2 Ubuntu

These instructions were prepared on Ubuntu 20.04 LTS Focal Fossa.

As a preliminary step, you should install the dependencies that Escript requires from the repository. For Python 3.0+, you should install the following packages:

```
sudo apt-get install python3-dev python3-numpy python3-pyproj python3-gdal sudo apt-get install python3-sympy python3-matplotlib python3-scipy sudo apt-get install libnetcdf-cxx-legacy-dev libnetcdf-c++4-dev libnetcdf-dev sudo apt-get install libboost-random-dev libboost-python-dev libboost-iostreams-dev sudo apt-get install scons lsb-release libsuitesparse-dev
```

Then navigate to the source directory and execute the following

```
scons -j1 options_file=scons/templates/focus_options.py
```

5.3.3 OpenSuse

These instructions were prepared using OpenSUSE Leap 15.2.

As a preliminary step, you should install the dependencies that Escript requires from the repository. If you intend to use Python 2.7, then you should install the following packages

```
sudo zypper in python-devel python2-numpy python-gdal sudo zypper in python2-scipy python2-sympy python2-matplotlib sudo zypper in gcc gcc-c++ scons netcdf-devel libnetcdf_c++-devel sudo zypper in libboost_python-py2_7-1_66_0-devel libboost_numpy-py2_7-1_66_0-devel sudo zypper in libboost_iostreams1_66_0-devel suitesparse-devel
```

If you intend to use Python 3.0, then you should instead install the following packages

```
sudo zypper in python3-devel python3-numpy python3-GDAL sudo zypper in python3-scipy python3-sympy python3-matplotlib sudo zypper in gcc gcc-c++ scons netcdf-devel libnetcdf_c++-devel sudo zypper in libboost_python-py3-1_66_0-devel libboost_numpy-py3-1_66_0-devel sudo zypper in libboost_random1_66_0-devel libboost_iostreams1_66_0-devel sudo zypper in suitesparse-devel
```

Now to build escript itself. In the escript source directory execute the following (substitute *opensuse_py2* or *opensuse_py3* as appropriate for XXXX):

```
scons -j1 options_file=scons/templates/XXXX_options.py
```

If you wish to test your build, you can use the following:

```
scons -j1 py_tests options_file=scons/templates/XXXX_options.py
```

Now go to Section 5.4 for cleanup.

5.3.4 CentOS

It is possible to install esys.escript on both CentOS releases 7 and 8. We include separate instructions for each of these CentOS releases in this section.

CentOS release 7

The core of escript works, however some functionality is not available because the default packages for some dependencies in CentOS are too old. At present, it is not possible to compile <code>esys.escript</code> using Python 3.0+ on CentOS 7 as Python 3.0+ versions of many of the dependencies are not currently available in any of the CentOS repositories, but this may change in the future. In this section we only outline how to install a version of <code>esys.escript</code> that uses Python 2.7.

First, add the EPEL repository.

```
yum install epel-release.noarch
```

Install packages:

```
yum install netcdf-devel netcdf-cxx-devel gdal-python
yum install python-devel numpy scipy sympy python2-scons
yum install python-matplotlib gcc gcc-c++ boost-devel
yum install boost-python gdal-devel suitesparse-devel pyproj
```

Now to build escript itself. In the escript source directory:

```
scons -j1 options_file=scons/templates/centos7_0_options.py
```

Now go to Section 5.4 for cleanup.

CentOS release 8

The core of escript works in CentOS 8, however some functionality is not available because the default packages for some dependencies in CentOS are too old. This install is for Python 3.

First, add the EPEL, PowerTools and Okay repositories:

```
yum update
yum install epel-release.noarch dnf-plugins-core
yum config-manager --set-enabled PowerTools
rpm -ivh http://repo.okay.com.mx/centos/8/x86_64/release/okay-release-1-3.el8.noarch.rpm
yum update
```

Now, install the packages:

```
yum install python3-devel python3-numpy python3-scipy python3-pyproj
yum install boost-devel boost-python3 boost-python3-devel
yum install gcc gcc-c++ scons
yum install suitesparse suitesparse-devel
```

Finally, you can compile esys.escript with the command

```
scons -j1 options_file=scons/templates/centos8_0_options.py
```

5.3.5 Fedora

These instructions were prepared using Fedora 31 Workstation.

To build the a version of esys.escript that uses Python 2.7, install the following packages:

16 5.3. Building

```
yum install gcc-c++ scons suitesparse-devel
yum install python2-devel boost-python2-devel
yum install python2-scipy
yum install netcdf-devel netcdf-cxx-devel netcdf-cxx4-devel
```

To build the a version of esys.escript that uses Python 3.0+, install the following packages:

```
yum install gcc-c++ scons suitesparse-devel
yum install python3-devel boost-python3-devel
yum install python3-scipy python3-pyproj python3-matplotlib
yum install netcdf-devel netcdf-cxx-devel netcdf-cxx4-devel
```

In the source directory execute the following (substitute fedora_py2 or fedora_py3 for XXXX):

```
scons -j1 options_file=scons/templates/XXXX_options.py
```

Now go to Section 5.4 for cleanup.

5.3.6 MacOS 10.10 and later (macports)

The following will install the capabilities needed for the macports_10.10_options.py file (later versions can use the same options file).

```
sudo port install scons
sudo port select --set python python27
sudo port install boost
sudo port install py27-numpy
sudo port install py27-sympy
sudo port select --set py-sympy py27-sympy
sudo port install py27-scipy
sudo port install py27-pyproj
sudo port install py27-pyproj
sudo port install py27-gdal
sudo port install netcdf-cxx
sudo port install silo
```

```
scons -j1 options_file=scons/templates/macports_10.10options.py
```

5.3.7 MacOS 10.13 and later (homebrew)

The following will install the capabilities needed for the homebrew_10.13_options.py file.

```
brew install scons
brew install boost-python
brew install netcdf
```

There do not appear to be formulae for sympy or pyproj so if you wish to use those features, then you will need to install them separately.

```
scons -j1 options_file=scons/templates/homebrew_10.13_options.py
```

5.3.8 FreeBSD

At time of writing, numpy does not install correctly on FreeBSD. Since numpy is a critical dependency for esys.escript, we have been unable to test on FreeBSD.

5.3.9 Windows

These instructions were prepared for Microsoft Windows 10. Start by installing esys.escript dependencies.

- Microsoft Visual Studio
 - 1. Download the Microsoft Visual Studio Community 2017 (or VS 2019 if preferred) installer from

```
https://visualstudio.microsoft.com.
```

- 2. Launch the Visual Studio installer, selecting Individual components:
 - VS 2017: VC++ 2017 latest v141 tools
 VS 2019: MSVC v142 VS 2019 C++ build tools
 - Windows 10 SDK
 - MSBuild
 - Visual C++ tools for CMake
- Anaconda
 - 1. Download the Python 3.7 64-Bit Graphical Installer for Windows from

```
https://anaconda.org/.
```

2. Launch the Anaconda installer, selecting installation type: **Just Me** and destination folder: C:\Users\%USERNAME%\Anaconda3.

Next, open Windows Command Prompt (cmd.exe) and set-up the esys.escript dependencies.

- · Conda environment
 - 1. Create and activate a new environment

```
C:\Users\%USERNAME%\Anaconda3\Scripts\activate
conda create --name escript python=3.7
conda deactivate
C:\Users\%USERNAME%\Anaconda3\Scripts\activate escript
```

2. Install required conda modules

```
conda install numpy==1.15.4 matplotlib==2.2.2 sympy==1.1.1
  boost gdal git pyproj scons scipy m2-patch mumps gmsh
  -c defaults -c conda-forge
```

- Vcpkg
 - 1. Build vcpkg package manager

```
cd C:\Users\%USERNAME%
git clone https://github.com/Microsoft/vcpkg.git
cd vcpkg
bootstrap-vcpkg
```

2. Install the CppUnit vcpkg package

```
vcpkg install cppunit:x64-windows
```

Once the dependencies are installed and set-up, you can download and build esys.escript from source.

1. Activate the conda environment (if not active).

```
C:\Users\%USERNAME%\Anaconda3\Scripts\activate escript
```

2. Set-up the Command Prompt for the 64-bit MSVC command line build environment.

```
"C:\Program Files (x86)\Microsoft Visual Studio\2017\
    Community\VC\Auxiliary\Build\vcvars64.bat"
```

3. Add CppUnit to the Windows System Path.

```
set PATH=%PATH%;C:\Users\%USERNAME%\vcpkg\packages\cppunit_x64-windows\bin
```

4. Download the esys.escript source code tarball from

18 5.3. Building

```
https://launchpad.net/escript-finley
```

Extract the tarball to C:\Users\%USERNAME%\escript

5. Build and install the netCDF-4 C++ library before starting the esys.escript build. Download the netCDF-4 C++ v4.3.1 source code tarball from

```
https://github.com/Unidata/netcdf-cxx4/archive/v4.3.1.tar.gz
```

Extract the tarball to C:\Users\%USERNAME%\escript

6. Apply the provided patch.

```
cd C:\Users\%USERNAME%\escript\netcdf-cxx4-4.3.1
patch -p1 < ..\src\tools\anaconda\Anaconda3\netcdf-cxx4.patch</pre>
```

7. Configure, build, and install netcdf-cxx4.

```
mkdir build

cd build

cmake -G "Visual Studio 15 2017 Win64" -DBUILD_SHARED_LIBS=OFF

-DCMAKE_INSTALL_PREFIX="%CONDA_PREFIX%\Library"

-DCMAKE_LIBRARY_PATH="%CONDA_PREFIX%\Library\lib"

-DCMAKE_PREFIX_PATH="%CONDA_PREFIX%\Library"

-DNETCDF_LIB_NAME="netcdf" -DHDF5_LIB_NAME="hdf5" ..

cmake --build . --config Release

cmake --build . --config Release --target install
```

8. Kick-off the esys.escript build when the netCDF-4 C++ install is complete.

```
cd C:\Users\%USERNAME%\escript\src
scons -j1 build_full options_file=scons/templates/windows_msvc141_options.py
```

9. Once the build completes successfully, you can validate esys.escript using the provided test script.

```
python utest.py C:\Users\%USERNAME%\escript\src\build -t8
```

5.3.10 Other Systems / Custom Builds

esys.escript has support for a number of optional packages. Some, like netcdf need to be enabled at compile time, while others, such as sympy and the projection packages used in esys.downunder are checked at run time. For the second type, you can install them at any time (ensuring that python can find them) and they should work. For the first type, you need to modify the options file and recompile with scons. The rest of this section deals with this.

To avoid having to specify the options file each time you run scons, copy an existing _options.py file from the scons/ or scons/templates/ directories. Put the file in the scons directory and name it your-machinename_options.py. 5. For example: on a machine named toybox, the file would be scons/toybox_options.py.

Individual lines can be enabled/disabled, by removing or adding # (the python comment character) to the beginning of the line. For example, to enable OpenMP, change the line

```
#openmp = True
to
openmp = True
```

If you are using libraries which are not installed in the standard places (or have different names) you will need to change the relevant lines. A common need for this would be using a more recent version of the boost::python library. You can also change the compiler or the options passed to it by modifying the relevant lines.

⁵If the name has - or other non-alpha characters, they must be replaced with underscores in the filename

MPI

If you wish to enable or disable MPI, or if you wish to use a different implementation of MPI, you can use the mpi configuration variable. You will also need to ensure that the mpi_prefix and mpi_libs variables are uncommented and set correctly. To disable MPI use, mpi = 'none'.

Testing

As indicated earlier, you can test your build using scons py_tests. Note however, that some features like netCDF are optional for using esys.escript, the tests will report a failure if they are missing.

5.4 Cleaning up

Once the build (and optional testing) is complete, you can remove everything except:

- bin
- esys
- lib
- doc
- CREDITS
- LICENSE
- README

The last three aren't strictly required for operation. The doc directory is not required either but does contain examples of escript scripts.

You can run escript using path_to_escript_files/bin/run-escript.

Where path_to_escript_files is replaced with the real path.

Optional step

You can add the escript bin directory to your PATH variable. The launcher will then take care of the rest of the environment.

5.5 Optional Extras

Some other packages which might be useful include:

- Lapack and UMFPACK direct solvers (install the relevant libraries and enable them in the options file).
- support for the Silo file format (install the relevant libraries and enable them in the options file).
- VisIt visualisation package. Can be used independently but our weipa library can make a Visit plug-in
 to allow direct visualisation of escript simulations.
- gmsh meshing software used by our pycad library.
- Mayavi2 another visualisation tool.

5.5.1 Trilinos

esys.escript now has some support for Trilinos⁶ solvers and preconditioners. The most significant limitation is that the current Trilinos release does not support block matrices so esys.escript can only use Trilinos solvers for single PDEs (i.e. no PDE systems).

If your distribution does not provide Trilinos packages you can build a working version from source. (See Appendix B)

6https://trilinos.org/

5.4. Cleaning up

5.6 Testing esys.escript

esys.escript has extensive testing that can be used to confirm that the program is working correctly. To run the unit testing, compile esys.escript with the flag build_full. This will build esys.escript normally and then create a shell script named utest.sh. Once this file has been created, you can run unit testing using the command

```
sh utest.sh path_to_build_folder '-tT -nN -pP'
```

where T, N and P represent the number of threads, nodes and processes to run the testing on. Some of these terms can be omitted. For example, to run the testing in serial, you would run

```
sh utest.sh path_to_build_folder '-t1'
```

Note that a careless selection of these parameters may cause the testing program to skip many of the tests. For example, if you compile <code>esys.escript</code> with OpenMP enabled but then instruct the testing program to run on a single thread, many of the OpenMP tests will not be run.

Required compiler features

Building escript from source requires that your c++ compiler supports at least the following features:

- std::complex<>
- Variables declared with type auto
- Variables declared with type decltype (T)
- extern template class to prevent instantiation of templates.
- template class *classname*<*type*>; to force instantiation of templates
- isnan() is defined in the std:: namespace

The above is not exhaustive and only lists language features which are more recent that our previous baseline of c++99 (or which we have recently begun to rely on). Note that we test on up to date versions of g++, icpc & clang++ so they should be fine.

Note that in future we may use c++14 features as well.

В

Trilinos

In order to solve PDEs with complex coefficients, escript needs to be compiled with Trilinos support. This requires that your version of Trilinos has certain features enabled. Since some precompiled distributions of Trilinos are not built with these features, you may need to compile Trilinos yourself as well.

While we can't provide support for building Trilinos, we provide here two configuration files which seem to work for Debian 10 "buster". One of these cmake script builds Trilinos with MPI support and one builds Trilinos without MPI support.

B.1 Debian "buster" configuration

B.1.1 Required packages

The following packages should be installed to attempt this build:

```
cmake
g++
libsuitesparse-dev
libmumps-dev
libboost-dev
libscotchparmetis-dev
libmetis-dev
```

B.1.2 Example configuration file (without MPI)

Appendix B. Trilinos 25

```
-D CMAKE_Fortran_COMPILER=`which gfortran` \
     -D PYTHON_EXECUTABLE=/usr/bin/python3 \
     -D CMAKE_INSTALL_PREFIX=/usr/local/ \
     -D Trilinos_ENABLE_CXX11=ON \
     -D CMAKE_C_FLAGS=' -w -fopenmp' \
     -D CMAKE_CXX_FLAGS=' -w -fopenmp'\
     -D Trilinos_ENABLE_Fortran=ON \
     -D BUILD_SHARED_LIBS=ON \
     -D TPL_ENABLE_BLAS=ON \
     -D TPL ENABLE Boost=ON \
     -D TPL_ENABLE_Cholmod=ON \
     -D TPL_ENABLE_LAPACK=ON \
     -D TPL_ENABLE_METIS=ON \
     -D TPL ENABLE SuperLU=OFF \
     -D TPL ENABLE UMFPACK=ON \
     -D TPL_BLAS_INCLUDE_DIRS=/usr/include/suitesparse \
     -D TPL_Cholmod_INCLUDE_DIRS=/usr/include/suitesparse \
     -D TPL_Cholmod_LIBRARIES='libcholmod.so; libcolamd.so' \
     -D TPL_UMFPACK_INCLUDE_DIRS=/usr/include/suitesparse \
     -D TPL_Boost_INCLUDE_DIRS=/usr/local/boost/include \
      -D TPL_Boost_LIBRARY_DIRS=lib \
     -D Trilinos_ENABLE_Amesos=ON \
     -D Trilinos_ENABLE_Amesos2=ON \
     -D Trilinos_ENABLE_AztecOO=ON \
     -D Trilinos ENABLE Belos=ON \
     -D Trilinos ENABLE Ifpack=ON \
     -D Trilinos_ENABLE_Ifpack2=ON \
     -D Trilinos_ENABLE_Kokkos=ON \
     -D Trilinos_ENABLE_Komplex=ON \
     -D Trilinos_ENABLE_ML=ON \
     -D Trilinos_ENABLE_MueLu=ON \
     -D Trilinos_ENABLE_Teuchos=ON \
     -D Trilinos_ENABLE_Tpetra=ON \
     -D Trilinos_ENABLE_ALL_OPTIONAL_PACKAGES=ON \
     -D Kokkos_ENABLE_AGGRESSIVE_VECTORIZATION=ON \
     -D Tpetra_INST_COMPLEX_DOUBLE=ON \
     -D Trilinos_ENABLE_COMPLEX_DOUBLE=ON \
     -D Teuchos_ENABLE_COMPLEX=ON \
     -D Tpetra_INST_INT_INT=ON \
     -D Tpetra_ENABLE_DEPRECATED_CODE=ON \
     -D Trilinos_ENABLE_OpenMP=ON \
     -D Trilinos_ENABLE_MPI=OFF \
     -D Trilinos_ENABLE_EXPLICIT_INSTANTIATION=ON \
     -D Kokkos_ENABLE_COMPILER_WARNINGS=ON \
     -D Amesos2_ENABLE_Basker=ON \
     -D Tpetra_INST_SERIAL:BOOL=ON \
     -D Trilinos ENABLE TESTS=OFF \
$EXTRA ARGS \
$TRILINOS PATH
```

-D CMAKE_CXX_COMPILER=`which g++` \

B.1.3 Example configuration file (with MPI)

```
# Set this to the root of your Trilinos source directory.
```

#!/bin/bash

```
TRILINOS_PATH=../trilinos_source
# You can invoke this shell script with additional command-line
# arguments. They will be passed directly to CMake.
EXTRA_ARGS=$@
rm -f CMakeCache.txt
cmake \
      -D MPI_C_COMPILER=`which mpicc` \
      -D MPI_CXX_COMPILER=`which mpic++`
      -D MPI_Fortran_COMPILER=`which mpif90` \
      -D PYTHON EXECUTABLE=/usr/bin/python3 \
      -D CMAKE INSTALL PREFIX=/usr/local/ \
      -D Trilinos_ENABLE_CXX11=ON \
      -D CMAKE_C_FLAGS=' -w -fopenmp' \
      -D CMAKE_CXX_FLAGS=' -w -fopenmp'\
      -D Trilinos_ENABLE_Fortran=ON \
      -D BUILD_SHARED_LIBS=ON \
      -D TPL_ENABLE_BLAS=ON \
      -D TPL_ENABLE_Boost=ON \
      -D TPL_ENABLE_Cholmod=ON \
      -D TPL_ENABLE_LAPACK=ON \
      -D TPL ENABLE METIS=ON \
      -D TPL_ENABLE_SuperLU=OFF \
      -D TPL_ENABLE_UMFPACK=ON \
      -D TPL_ENABLE_PARMETIS=ON \
      -D TPL ENABLE SCALAPACK=ON \
      -D TPL_ENABLE_MUMPS=OFF \
      -D TPL_BLAS_INCLUDE_DIRS=/usr/include/suitesparse \
      -D TPL_Cholmod_INCLUDE_DIRS=/usr/include/suitesparse \
      -D TPL_Cholmod_LIBRARIES='libcholmod.so; libamd.so; libcolamd.so' \
      -D TPL_UMFPACK_INCLUDE_DIRS=/usr/include/suitesparse \
      -D TPL_Boost_INCLUDE_DIRS=/usr/local/boost/include \
      -D TPL_Boost_LIBRARY_DIRS=lib \
      -D TPL_MUMPS_INCLUDE_DIRS='/usr/include/mumps-seq-shared/' \
      -D Trilinos_ENABLE_Amesos=ON \
      -D Trilinos_ENABLE_Amesos2=ON \
      -D Trilinos_ENABLE_AztecOO=ON \
      -D Trilinos_ENABLE_Belos=ON \
      -D Trilinos_ENABLE_Ifpack=ON \
      -D Trilinos_ENABLE_Ifpack2=ON \
      -D Trilinos_ENABLE_Kokkos=ON \
      -D Trilinos_ENABLE_Komplex=ON \
      -D Trilinos_ENABLE_ML=ON \
      -D Trilinos_ENABLE_MueLu=ON \
      -D Trilinos ENABLE Teuchos=ON \
      -D Trilinos_ENABLE_Tpetra=ON \
      -D Trilinos_ENABLE_ALL_OPTIONAL_PACKAGES=ON \
      -D KOKKOS_ENABLE_AGGRESSIVE_VECTORIZATION=ON \
      -D Tpetra_INST_COMPLEX_DOUBLE=ON \
      -D Teuchos_ENABLE_COMPLEX=ON \
      -D Tpetra_INST_INT_INT=ON \
      -D Tpetra_ENABLE_DEPRECATED_CODE=ON \
```

Appendix B. Trilinos 27

```
-D Trilinos_ENABLE_OpenMP=ON \
-D Trilinos_ENABLE_MPI=ON \
-D Trilinos_ENABLE_EXPLICIT_INSTANTIATION=ON \
-D KOKKOS_ENABLE_COMPILER_WARNINGS=ON \
-D Amesos2_ENABLE_Basker=ON \
-D Tpetra_INST_SERIAL:BOOL=ON \
-D Trilinos_ENABLE_TESTS=OFF \
$EXTRA_ARGS \
$TRILINOS_PATH
```