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# Installation guide for *esys-Escript*

*Release - 3.4.2*  
*(r4925)*

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# Introduction

This document describes how to install *esys-Escript*<sup>1</sup> on to your computer. To learn how to use Escript please see the Cookbook, User's guide or the API documentation. If you use the Debian or Ubuntu and you have installed the `python-escript-doc` package then the documentation will be available in the directory `/usr/share/doc/python-escript-doc`, otherwise (if you haven't done so already) you can download the documentation bundle from launchpad.

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

1. Binary packages – ready to run with no compilation required. Bundles are available for:

- `.deb` files for Debian and Ubuntu Linux

Please see Appendix A for Windows instructions. The rest of this guide assumes you are using a posix like system.

2. From source – that is, it must be compiled for your machine. This will be required if there is no binary package for your machine or if extra functionality is required such as MPI parallelisation.

*The major change from the point of view of installation is that we no longer provide binary packages compiled against the “support bundles”. This means, there are no longer binary packages for MacOS or generic Linux. One can still build from source and we have endeavoured to make this process as straight forward as possible.*

See the site <https://answers.launchpad.net/escript-finley> for online help. Chapter 2 describes how to install binary packages on Debian/Ubuntu systems. Chapter 3 covers installing from source. Appendix A gives brief instructions for Windows.

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<sup>1</sup>For the rest of the document we will drop the *esys-*



# Debian/Ubuntu Binary Installation

We provide `.deb` files for the following distributions:

Debian (i386 or amd64):

- 6 — *Squeeze*
- 7 — *Wheezy*

Ubuntu (i386 or amd64):

- 12.04 — *Precise Pangolin (LTS)*
- 13.10 — *Saucy Salamander*
- 14.04 — *Trusty Tahr (LTS)*

Two packages make up the `esys.escript` system: The main package which contains all system itself and the (optional) documentation package. The main package will be named `python-escript-X-D_A.deb` where `X` is the version, `D` is the distribution codename (eg “wheezy” or “precise”) and `A` is the architecture. For example, `python-escript-3.4-1-precise_amd64.deb` would be the file for Ubuntu 12.04 for 64bit processors. There is a common documentation for all distributions called `python-escript-doc-X_all.deb`. To install Escript, download the appropriate `.deb` file(s) and execute the following commands as root (you need to be in the directory containing the file):

**(For Ubuntu users)**

You will need to either install `aptitude`<sup>1</sup> or substitute `apt-get` where this guide uses `aptitude`.

```
sudo apt-get install aptitude
```

```
dpkg --unpack python-escript*.deb
aptitude install python-escript python-escript-doc
```

Installing `escript` should not remove any (non-`escript`) packages from your system. If `aptitude` suggests removing `python-escript` then choose ‘N’. If it wants to remove `escript-noalias` or `escript`, then choose ‘Y’. It should then suggest installing some dependencies choose ‘Y’ here.

If you use `sudo` (for example on Ubuntu) enter the following instead:

```
sudo dpkg --unpack python-escript*.deb
sudo aptitude install python-escript python-escript-doc
```

There are a number of optional dependencies which you should also install unless you are sure you don’t need them:

```
aptitude install python-sympy python-matplotlib python-scipy
aptitude install python-pyproj python-gdal
```

This should install Escript and its dependencies on your system. Please notify the development team if something goes wrong.

---

<sup>1</sup>Unless you are short on disk space `aptitude` is recommended



# Installing from Source

This chapter assumes you are using a unix/posix like system (including MacOSX).

## 3.1 Parallel Technologies

It is likely that the computer you run `esys.escript` on, will have more than one processor core. `esys.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 3.1.1 gives some rough guidelines to help you determine what you need.

There are two technologies which `esys.escript` can employ here.

- OpenMP – more efficient of the two [thread level parallelism].
- MPI – Uses multiple processes (less efficient), needs less help from the compiler.

Esript is primarily tested on recent versions of the GNU and Intel suites (“g++” / “icpc”). However, it also passes our tests when compiled using “clang++”. The table below shows what methods are available with which compilers.

	Serial	OpenMP	MPI
$\leq$ g++-4.2.1	✓	<sup>1</sup>	✓
g++ (recent $\geq$ 4.3.2)	✓	✓	✓
icpc(10)	✓	✓	✓
icpc(11)	✓	<sup>2</sup>	✓
icpc(12)	✓	✓	✓
clang++	✓		✓

Where both OpenMP and MPI are marked, `esys.escript` can be compiled with either or both. A ✓ mark means that combination passes our tests.

### 3.1.1 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; so you should use it. You will not need MPI unless your computer is some form of cluster.

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 reinstall `esys.escript` with more options.

<sup>1</sup>The OpenMP support in g++-4.2.1 is buggy/non-functional.

<sup>2</sup>There is a subtle bug in icpc-11 when OpenMP and c++ exception handling are combined.

In general, for a single computer, OpenMP will give better performance (both time and memory) so use it if possible. If OpenMP is not an option, then use MPI.

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` and `-p` options. If not, then consult the documentation for your MPI libraries (or the compiler documentation in the case of OpenMP<sup>3</sup>).

If you are using MacOSX, then see the next section, if not, then skip to Section 3.3.

## 3.2 MacOS

To build `esys.escript` from source you need to choose and install the following (if you already have some parts installed, skip that section).

- XCode and the associated command line tools. [Section 3.2.1]
- A package manager. [Section 3.2.2]
- A compiler. [Section 3.2.3]

As noted above, different compilers have varying support for parallel processing technologies so your choice of compiler may limit your options of parallel technologies. Once you have the pieces in this section, go to Section 3.3 for build instructions.

### 3.2.1 Initial compiler

*This information is accurate as far as we can tell at time of writing but things may change.*

Even if you wish to install and use a different compiler later, the first step is generally to get some compiler onto your system.

As of OSX10.9, the command `xcodebuild` will allow you to download and install the commandline tools seemingly without an AppleID. For previous releases, it seems to be trickier to get the basic compiler without doing one of the following:

- purchasing an iTunes gift card.
- giving Apple access to your credit card.
- signing up as an Apple developer<sup>4</sup> and giving up personal information.

If you install XCode, you will need to download the “command line tools” optional package [see XCode documentation for details].

There are also a number of projects on the net which aim to deliver compilers for MacOS. Use at your own risk. For example:

- <http://hpc.sourceforge.net>
- <http://kennethreitz.org/experiments/xcode-gcc-and-homebrew>

### 3.2.2 OSX Package Managers

Once you have a working compiler, you will need to consider how you will install the components `esys.escript` needs. While it is certainly possible to compile and install these pieces manually, it can be easier to use a tool which handles some of the details for you. This is especially true if you will need to uninstall or upgrade them later.

On OSX, there are a number of options. The popular ones at time of writing seem to be `macports` and `homebrew`. (We have not experimented with `fink`). `esys.escript` does not assume that you are using a particular package manager<sup>5</sup>. The configuration instructions/files in this guide are just examples that we have found to work.

<sup>3</sup>It may be enough to set the `OMP_NUM_THREADS` environment variable.

<sup>4</sup>If you do this you can download a “command line tools” package which installs the relevant compilers without needing to install all of XCode.

<sup>5</sup>or even if you are using one at all.

Note 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.

Please consult the documentation for your chosen package manager to determine how to set it up.

### 3.2.3 A compiler

While the command line tools described in 3.2.1, do contain a c++ compiler, in recent versions of OSX, it is likely to be some version of clang++<sup>6</sup>. If you don't need OpenMP or genuine g++ or you have installed an OpenMP supporting compiler using some other method, then you can skip to Section 3.3.

If you are still reading this part, we assume that you wish to use g++ to build escript on your system and that you are using a package manager. The challenge here isn't installing the compiler itself. The issue is that the other dependencies that esys.escript needs have probably<sup>7</sup> been compiled by clang rather than the compiler you want to use. A lot of the time this is fine, but in the case of c++ libraries such as boost, the result can be two different standard c++ libraries<sup>8</sup> being used in the same program and confusing matters. Apart from giving up, there are two ways to try to solve this problem.

1. Changing the compiler that your package manager uses to compile its packages. *This is not for the faint hearted!* But we do provide some instructions in Appendix B.
2. Install most of the libraries using your package manager to install most things and then compile the c++ parts yourself. (The most important one is boost). We have tested this on Macports and it works well. The only trick is to make sure that your "replacement" components are in a separate directory which is ahead of you Macports directory in the various paths (PATH, LD\_LIBRARY\_PATH, DYLD\_LIBRARY\_PATH).

## 3.3 Building

To simplify things for people, we have prepared \_options.py files for a number of systems<sup>9</sup>. The \_options.py files are locate in the scon/os directory. We suggest that the file most relavent to your os be copied from the os directory to the scon directory and renamed to the form XXXX\_options.py where XXXX should be replaced with your computer's name. If your particular system is not in the list below, or if you want a more customised build<sup>10</sup>, see Section 3.3.9 for instructions.

- Debian - 3.3.1
- Ubuntu - 3.3.2
- OpenSuse - 3.3.3
- Centos - 3.3.4
- Fedora - 3.3.5
- MacOS (macports) - 3.3.6
- MacOS (homebrew) - 3.3.7
- FreeBSD - 3.3.8

Once these are done proceed to Section 3.4 for cleanup steps.

All of these instructions assume that you have obtained the source (uncompressed it if necessary).

---

<sup>6</sup>even if it is labelled as g++ it is likely clang++ under the hood.

<sup>7</sup>or "will probably be compiled" in the case of homebrew.

<sup>8</sup>libstdc++ and libc++ on OSX 10.9.

<sup>9</sup>These 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.

<sup>10</sup>for example, you want MPI functionality or you wish to use a different compiler

### 3.3.1 Debian

```
sudo aptitude install python-dev python-numpy libboost-python-dev libnetcdf-dev
sudo aptitude install sconslsb-release
sudo aptitude install python-sympy python-matplotlib python-scipy
sudo aptitude install python-pyproj python-gdal
```

#### *Optional step*

If for some reason, you wish to rebuild the documentation, you would also need the following:

```
sudo aptitude install python-sphinx doxygen python-docutils texlive
sudo aptitude install zip texlive-latex-extra latex-xcolor
```

In the source directory execute the following (substitute squeeze or wheezy as appropriate for XXXX):

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

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

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

### 3.3.2 Ubuntu

If you have not installed aptitude, then substitute apt-get in the following.

```
sudo aptitude install python-dev python-numpy libboost-python-dev libnetcdf-dev
sudo aptitude install sconslsb-release
sudo aptitude install python-sympy python-matplotlib python-scipy
sudo aptitude install python-pyproj python-gdal
```

#### *Optional step*

If for some reason, you wish to rebuild the documentation, you would also need the following:

```
sudo aptitude install python-sphinx doxygen python-docutils texlive
sudo aptitude install zip texlive-latex-extra latex-xcolor
```

In the source directory execute the following (substitute precise, quantal or raring as appropriate for XXXX):

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

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

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

### 3.3.3 OpenSuse

These instructions were prepared using release 13.1.

Install packages from the main distribution:

```
sudo zypper install libboost_python1_53_0 python-devel python-numpy
sudo zypper install python-scipy python-sympy python-matplotlib libnetcdf_c++-devel
sudo zypper install gcc-c++ sconslboost-devel netcdf-devel
```

These will allow you to use most features except some parts of the esys inversion library. If you wish to use those, you will need some additional packages [python-pyproj, python-gdal]. This can be done after Escript installation.

#### *Optional step*

Add [http://ftp.suse.de/pub/opensuse/repositories/Application:/Geo/openSUSE\\_13.1/](http://ftp.suse.de/pub/opensuse/repositories/Application:/Geo/openSUSE_13.1/) to your repositories in zypper.

```
sudo zypper install python-pyproj, python-gdal
```

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

```
scons -j1 options_file=scons/os/opensuse13.1_options.py
```

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

```
scons -j1 py_tests options_file=scons/os/opensuse13.1_options.py
```

Now go to Section 3.4 for cleanup.

### 3.3.4 Centos

These instructions were prepared using release 6.5. The core of escript works, however some functionality is not available because the default packages for some dependencies in Centos are too old.

Install packages from the main distribution:

```
yum install python-devel numpy scipy scons boost-devel
yum install python-matplotlib gcc-c++
yum install boost-python
```

The above packages will allow you to use most features except saving and loading files in netCDF format and the esys inversion library. If you wish to use those features, you will need to install some additional packages. NetCDF needs to be installed when you compile if you wish to use it.

#### *Optional step*

Add the EPEL repository.

```
rpm -U http://download.fedoraproject.org/pub/epel/6/x86_64/epel-release-6-8.noarch.rpm
```

```
yum install netcdf-devel gdal-python
```

For some coordinate transformations, esys can also make use of the python interface to a tool called proj. There does not seem to be an obvious centos repository for this though. If it turns out to be necessary for your particular application, the source can be downloaded.

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

```
scons -j1 options_file=scons/os/centos6.5_options.py
```

Now go to Section 3.4 for cleanup.

### 3.3.5 Fedora

These instructions were prepared using release 20.

Install packages

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

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

```
scons -j1 options_file=scons/os/fedora18_options.py
```

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

```
scons -j1 py_tests options_file=scons/os/fedora18_options.py
```

Now go to Section 3.4 for cleanup.

### 3.3.6 MacOS (macports)

```
port install python27
port select --set python python27
port install scons
port install openmpi
port install py27-numpy
port install boost
```

```
port install py27-sympy
port select --set py-sympy py27-sympy
install py27-scipy
install py27-pyproj
install py27-gdal
install py27-netcdf4
install netcdf-cxx
```

The options file we provide as an example below uses `clang` as its compiler. If you wish to use `g++`, then you will need to modify the

```
tools_names = ['clang']
```

to

```
tools_names = ['default']
```

```
scons -j1 options_file=scons/os/macports_options.py
```

### 3.3.7 MacOS (homebrew)

Note that these steps add “non-official” packages. You will also want to make sure that the homebrew Python is executed in preference to the system Python<sup>11</sup>.

```
brew install python
brew install scons
brew install boost
brew tap samueljohn/python
brew tap homebrew/science
pip install nose
brew install gfortran
brew install samueljohn/python/numpy
brew install scipy
brew install gdal
brew install openmpi
brew install matplotlib
brew install netcdf --enable-cxx-compat
```

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/os/homebrew_options.py
```

### 3.3.8 FreeBSD

#### Release 9.1

The following passes the majority of tests but there is an issue related to some features in the inversion library. The list of installations “works” but is not guaranteed to be minimal<sup>12</sup>.

Install the following packages:

- python
- scons
- boost-python-libs
- bash

Now install the following ports:

---

<sup>11</sup>Putting `/usr/local/bin` at the front of your `PATH` is one way to do this.

<sup>12</sup>Depending on your needs you might be able to get by with a smaller set of packages.

- science/py-scipy
- science/netcdf
- science/silo
- math/py-sympy
- graphics/py-pyproj
- graphics/py-gdal
- net/openmpi (optional)

You will need to add `/usr/local/mpi/openmpi/bin` to your path if you wish to build with MPI. Next choose (or create) your options file. In this case we have three prepared in the `scons/os` directory:

- `freebsd91_options.py`
- `freebsd91_mpi_options.py` If you would like to use MPI.
- `freebsd91_gcc46_options.py` Use this if you have managed to change compilers to gcc4.6 (and would like to use OpenMP).

In the escript source directory (where `ZZZ` is your options file):

```
scons -j1 options_file=ZZZ
```

## Release 10.0

Install the following packages:

- python
- scon
- boost-python-libs
- bash
- netcdf
- silo
- py27-scipy
- py27-gdal
- py27-matplotlib
- py27-pyproj
- py27-sympy

Next choose (or create) your options file. For the setup as above the escript source comes with a prepared file in `scons/os/freebsd10.0_options.py`. Finally to build escript issue the following in the escript source directory (replace the options file as required):

```
scons -j1 options_file=scons/os/freebsd10.0_options.py
```

*Note:* Some packages installed above are built with gcc 4.7. Somewhere in the toolchain a system-installed gcc library is pulled in which is incompatible with the one from version 4.7 and would prevent escript from executing successfully. As explained in the FreeBSD documentation<sup>13</sup> this can be fixed by adding a line to `/etc/libmap.conf`:

```
libgcc_s.so.1 gcc47/libgcc_s.so.1
```

<sup>13</sup>see <http://www.freebsd.org/doc/en/articles/custom-gcc/article.html>

### 3.3.9 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` 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/os/` directories. Put the file in the `scons` directory and name it `yourmachinename_options.py`.<sup>14</sup> 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.

#### 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. To disable MPI use, `mpi = 'none'`. You will also need to ensure that the `mpi_prefix` and `mpi_libs` variables are uncommented and set correctly.

#### Python3

`esys.escript` works with `python3` but until recently, many distributions have not distributed `python3` versions of their packages. You can try it out though by modifying the following variables:

```
pythoncmd='python3'  
  
usepython3=True  
  
pythonlibname='whateveryourpython3libraryiscalled'
```

#### 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.

## 3.4 Cleaning up

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

- `bin`
- `esys`
- `lib`
- `doc`
- `CREDITS.TXT`

---

<sup>14</sup>If the name has - or other non-alpha characters, they must be replaced with underscores in the filename

- README\_LICENSE

The last two 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.

## 3.5 Optional Extras

Some other packages which might be useful include:

- support for silo 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 files.
- gmsh — meshing software used by our `pycad` library.
- mayavi — another visualisation tool.



---

# Windows binary installation

There is no automated install/uninstall procedure for Escript on Windows at this time. Where builds are available, they will be built for Windows XP and has not been tested on newer versions. If you want to use MPI, make sure to download the MPI version [You will also need MPICH2 1.0.8 (<http://www.mcs.anl.gov/research/projects/mpich2/>)]. Other dependencies are:

- pythonxy (<http://www.pythonxy.com>) or
  - Python 2.7 (<http://python.org>)
  - Numpy 1.3.0 (<http://sourceforge.net/projects/numpy/files/NumPy>)
  - SymPy 0.7.1 (<http://sympy.org>)
- Optional:
  - gmsh 2.4.0 (required to use pycad, must be in your PATH) - <http://www.geuz.org/gmsh>
  - matplotlib 0.99 - <http://matplotlib.sourceforge.net>

Unpack the escript zip file, then:

- copy the `esys` directory to your Python 2.7 site-packages folder<sup>1</sup> (usually `C:\Python27\Lib\site-packages`).
- copy the `.dll` files from `esys_dlls` to a directory on your PATH. For example copy the directory to `C:\Python27\libs\esys_dlls` and add `C:\Python27\libs\esys_dlls` to your PATH.<sup>2</sup>

---

<sup>1</sup>Substitute the relevant Python version

<sup>2</sup>Failing to do so my result in the error message: “This application has failed to start because the boost\_python-vc71-mt-1.33.1.dll was not found.”



# Changing compilers for package managers

Here are some brief instructions / hints for changing the compilers used by package managers in OSX and FreeBSD. If you are intend to change the compiler or python versions, you should install the new compiler first, then the new version of python. After this, you can install the other dependencies. *Please note that neither homebrew, MacPorts nor BSD recommend changing the default C compiler so do so at your own risk. If you want OpenMP though, there does not seem to be a choice.*

The following steps seemed to work when we tested them but we cannot make guarantees.

## Changing compiler on FreeBSD(9.1)

The following sequence passes our unit tests under OpenMP.

- Install `gcc46` from ports.
- Modify `/etc/make.conf` to set the default compiler to be `gcc46`
- Install the remaining dependencies.
- Configure `esys.escript` to build with OpenMP.

We chose version 4.6 rather than a later one because one of the optional dependencies (scipy) will try to install it anyway.

## Changing compiler on MacPorts

The following sequence passes our unit tests under OpenMP.

- `port install gcc47`
- Set the default compiler for the command line with: `port select --set gcc mp-gcc47`
- Set the default compiler for macports by adding the following line to `/opt/local/etc/macports/macports.conf`:

```
#Added by user to coerce macports into using a newer compiler
default_compiler      macports-gcc-4.7
```

- Now install the remainder of the dependencies using `port -s install X`. This will build them from source rather than downloading precompiled versions. (This will unfortunately mean larger downloads.). In some cases, some dependencies will not build properly from source. In those cases(where XYZ is the dependency that fails to build), `port clean XYZ` then `port install XYZ`. Then *try to install the rest from source*.

Installing from source via macports is necessary to convince macports to link against the libraries provided by your new compiler rather than the default one.

### **Changing compiler on Homebrew**

There is no configuration file that needs to be changed in order to use a different compiler. You do need to do things in the correct order and make sure that you have made any required changes to your environment variables before moving on to the next step. In particular, the compiler must be installed before anything else, followed by Python.