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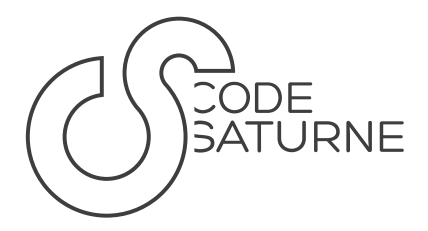
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Code_Saturne documentation

Code_Saturne version 4.0.5 installation guide

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1 Code_Saturne Automated or manual installation

Code_Saturne may be installed either directly through its GNU Autotools based scripts (the traditional configure, make, make install) sequence), or using an automated installer (install_saturne.py), which generates an initial setup file when run a first time, and builds and installs Code_Saturne and some optional libraries based on the edited setup when run a second time. The use of this automated script is briefly explained in the top-level README file of the Code_Saturne sources, as well as in the comments of setup file. It is not detailed further in this documentation, which details the manual installation, allowing a finer control over installation options.

Note that when the automatic installer is run, it generates a build directory, in which the build may be modified (re-running configure, possibly adapting the command logged at the beginning of the config.status file) and resumed.

2 Installation basics

The installation scripts of *Code_Saturne* are based on the GNU Autotools, (Autoconf, Automake, and Libtool), so it should be familiar for many administrators. A few remarks are given here:

- As with most software with modern build systems, it is recommended to build the code in a separate directory from the sources. This allows multiple builds (for example production and debug), and is considered good practice. Building directly in the source tree is not regularly tested, and is not guaranteed to work, in addition to "polluting" the source directory with build files.
- By default, optional libraries which may be used by <code>Code_Saturne</code> are enabled automatically if detected in default search paths (i.e. <code>/usr/</code> and <code>/usr/local</code>. To find libraries associated with a package installed in an alternate path, a <code>--with-<package>=...</code> option to the <code>configure</code> script must given. To disable the use of a library which would be detected automatically, a matching <code>--without-<package></code> option must be passed to <code>configure</code> instead.
- Most third-party libraries usable by *Code_Saturne* are considered optional, and are simply not used if not detected, but the libraries needed by the GUI are considered mandatory, unless the --disable-gui or --disable-frontend option is explicitly used.

When the prerequisites are available, and a build directory created, building and installing *Code_Saturne* may be as simple as running:

```
$ ../../code_saturne-4.0.5/configure
$ make
$ make install
```

The following chapters give more details on *Code_Saturne*'s recommended third-party libraries, configuration recommendations, troubleshooting, and post-installation options.

3 Compilers and interpreters

For a minimal build of Code_Saturne on a Linux or Posix system, the requirements are:

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- A C compiler, conforming at least to the C99 standard.
- A Fortran compiler, conforming at least to the Fortran 95 standard and supporting the ISO_C_BINDING Fortran 2003 module.
- A Python 2 interpreter, with Python version 2.6 or 2.7.

For parallel runs, an MPI library is also necessary (MPI-2 or MPI-3 conforming). To build and use the GUI, Libxml2 and PyQt4 (which in turn requires Qt4 and SIP) are required. Other libraries may be used for additional mesh format options, as well as to improve performance. A list of those libraries and their role is given in §4.2.

For some external libraries, such as Catalyst (see 4.2), a C++ compiler is also required.

In practice, the code is known to build and function properly at least with the GNU compilers 4.4 and above (up to 5.1 at this date), Intel compilers 11 and above (up to 15 at this date), IBM XL Fortran 14 and C 12 compilers.

Note also that while *Code_Saturne* makes heavy use of Python, this is for scripts and for the GUI only; The solver only uses compiled code, so we may for example use a 32-bit version of Python with 64-bit *Code_Saturne* libraries and executables. On supercomputers with separate front-end and compute node architectures, Python is not required on the compute nodes (except as a dependency for the optional Paraview/Catalyst support). Also, the version of Python used by ParaView/Catalyst may be independent from the one used for building *Code_Saturne*.

4 Third-Party libraries

For a minimal build of *Code_Saturne*, a Linux or Posix system with C and Fortran compilers (C99 and Fortran 95 with Fortran 2003 ISO C bindings conforming respectively), a Python (2.6 or later) interpreter and a make tool should be sufficient. For parallel runs, an MPI library is also necessary (MPI-2 or MPI-3 conforming). To build and use the GUI, Libxml2 and Qt4 with PyQt4 Python bindings (which in turn requires SIP) are required. Other libraries may be used for additional mesh format options, as well as to improve performance. A list of those libraries and their role is given in §4.2.

4.1 Installing third-party libraries for Code_Saturne

Third-Party libraries usable with Code_Saturne may be installed in several ways:

- On many Linux systems, most of libraries listed in §4.2 are available through the distribution's package manager. This requires administrator privileges, but is by far the easiest way to install third-party libraries for *Code_Saturne*.
 - Note that distributions usually split libraries or tools into runtime and development packages, and that although some packages are installed by default on many systems, this is generally not the case for the associated development headers. Development packages usually have the same name as the matching runtime package, with a <code>-dev</code> postfix added. For example, on a Debian or Ubuntu system <code>,libxml2</code> is usually installed by default, but <code>libxml2-dev</code> must also be installed for the <code>Code_Saturne</code> build to be able to use the former.
- On many large compute clusters, Environment Modules allow the administrators to provide multiple versions of many scientific libraries, as well us compilers or MPI libraries, using the module command. More details on Environment Modules may be found at http://modules.sourceforge.net. When being configured and installed Code_Saturne checks for modules loaded

 $^{^{1}}$ On Mac OS X systems, package managers such as Fink or MacPorts also provide package management, even though the base system does not.

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with the module command, and records the list of loaded modules. Whenever running that build of *Code_Saturne*, the modules detected at installation time will be used, rather than those defined by default in the user's environment. This allows using versions of *Code_Saturne* built with different modules safely and easily, even if the user may be experimenting with other modules for various purposes.

• If not otherwise available, third-party software may be compiled an installed by an administrator or a user. An administrator will choose where software may be installed, but for a user without administrator privileges or write access to usr/local, installation to a user account is often the only option. None of the third-party libraries usable by Code_Saturne require administrator privileges, so they may all be installed normally in a user account, provided the user has sufficient expertise to install them. This is usually not complicated (provided one reads the installation instructions, and is prepared to read error messages if something goes wrong), but even for an experienced user or administrator, compiling and installing 5 or 6 libraries as a prerequisite significantly increases the effort required to install Code_Saturne.

Even though it is more time-consuming, compiling and installing third-party software may be necessary when no matching packages or Environment Modules are available, or when a more recent version or a build with different options is desired.

4.2 List of third-party libraries usable by Code_Saturne

The list of third-party software usable with *Code_Saturne* is provided here:

- BLAS (Basic Linear Algebra Subroutines) may be used by the cs_blas_test unit test to compare the cost of operations such as vector sums and dot products with those provided by the code and compiler. If no third-party BLAS is provided, Code_Saturne reverts to its own implementation of BLAS routines, so no functionality is lost here. Optimized BLAS libraries such as Atlas, MKL, ESSL, or ACML may be very fast for BLAS3 (dense matrix/matrix operations), but the advantage is usually much less significant for BLAS 1 (vector/vector) operations, which are almost the only ones Code_Saturne has the opportunity of using. Code_Saturne uses its own dot product implementation (using a superblock algorithm, for better precision), and $y \leftarrow ax + y$ operations, so external BLAS1 are not used for computation, but only for unit testing (so as to be able to compare performance of built-in BLAS with external BLAS). The Intel MKL BLAS may also be used for matrix-vector products, so it is linked with the solver when available, but this is also currently only used in unit benchmark mode. Note that in some cases, threaded BLAS routines might oversubscribe processor cores in some MPI calculations, depending on the way both Code_Saturne and the BLAS were configured and interact, and this can actually lead to lower performance. Use of BLAS libraries is thus useful as a unit benchmarking feature, but has no influence on full calculations.
- PyQt4 is required by the Code_Saturne GUI. PyQt4 in turn requires Qt4, Python,and SIP. Without this library, the GUI may not be built, although XML files generated with another install of Code_Saturne may be used if Libxml2 is available.
- Libxml2 is required to read XML files edited with the GUI. If this library is not available, only user subroutines may be used to setup data.
- HDF5 is necessary for MED, and may also be used by CGNS.
- CGNSlib is necessary to read or write mesh and visualization files using the CGNS format, available as an export format with many third-party meshing tools. CGNS version 3.1 or above is required.
- MED is necessary to read or write mesh and visualization files using the MED format, mainly used by the SALOME platform (www.salome-platform.org).

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- libCCMIO is necessary to read or write mesh and visualization files generated or readable by STAR-CCM+ using its native format.
- SCOTCH or PT-SCOTCH may be used to optimize mesh partitioning. Depending on the mesh, parallel computations with meshes partitioned with these libraries may be from 10% to 50% faster than using the built-in space-filling curve based partitioning.
 - As SCOTCH and PT-SCOTCH use symbols with the same names, only one of the 2 may be used. If both are detected, PT-SCOTCH is used. Versions 6.0 and above are supported.
- Metis of Parmetis are alternative mesh partitioning libraries. These libraries have a separate source tree, but some of their functions have identical names, so only one of the 2 may be used. If both are available, Parmetis will be used. Partitioning quality is similar to that obtained with Scotch of PT-Scotch.
 - Though broadly available, the PARMETIS license is quite restrictive, so PT-SCOTCH may be preferred (*Code_Saturne* may be built with both METIS and SCOTCH libraries). Also, the METIS license was changed in March 2013 to the Apache 2 license, so it would not be surprising for future PARMETIS versions to follow. METIS 5.0 or above and PARMETIS 4.0 or above are supported.
- Catalyst (http://www.paraview.org/in-situ/) or full ParaView may be used for co-visualization or in-situ visualization. This requires ParaView 4.2 or above.
- eos-1.2 may be used for thermodynamic properties of fluids. it is not currently free, so usually available only to users at EDF, CEA, or organisms participating in projects with those entities.
- freesteam (http://freesteam.sourceforge.net) is a free software thermodynamic properties library, implementing the IAPWS-IF97 steam tables, from the International Association for the Properties of Water and Steam (IAPWS). Version 2.0 or above may be used.

For developers, the GNU Autotools (Autoconf, Automake, Libtool) as well as gettext will be necessary. To build the documentation, pdfLATFX and fig2dev (part of TransFig) will be necessary.

4.3 Notes on some third-party tools and libraries

4.3.1 Python and PyQt4

The GUI is written in PyQt4 (Python bindings for Qt4), so but Qt4 and the matching Python bindings must be available. On most modern Linux distributions, this is available through the package manager, which is by far the preferred solution. When running on a system which does not provide these libraries, there are several alternatives:

- build Code_Saturne without the GUI. If built with Libxml2, XML files produced with the GUI are still usable, so if an install of Code_Saturne with the GUI is available on an other machine, the XML files may be copied on the current machine. This is certainly not an optimal solution, but in the case where users have a mix of desktop or virtual machines with modern Linux distributions and PyQt4 installed, and a compute cluster with an older system, this may avoid requiring a build of Qt4 and PyQt4 on the cluster if users find this too daunting.
- Install a local Python interpreter, and add Qt4 bindings to this interpreter.

 Python (http://www.python.org) and Qt4 (http://qt.nokia.com/products) must be downloaded and installed first, in any order. The installation instructions of both of these tools are quite clear, and though the installation of these large packages (especially Qt4) may be a lengthy process in terms of compilation time, but is well automated and usually devoid of nasty surprises.².

²The only case in which the *Code_Saturne* developers have has issues with Qt4 was when trying to force an install into 64-bit mode with the GNU compilers (version 4.1.2) on a PowerPC 64 architecture running SLES 10 Linux, on which compilers default to building 32 bit code, although 64 bit is available. Using default options on the same machine led to a perfectly functional 32-bit Qt installation

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Once Python is installed, the SIP bindings generator (http://riverbankcomputing.co.uk/software/sip/intro) must also be installed. This is a small package, and configuring it simply requires running python configure.py in its source tree, using the Python interpreter just installed.

Finally, the PyQt4 bindings (http://riverbankcomputing.co.uk/software/pyqt/intro) may be installed, in a manner similar to SIP.

When this is finished, the local Python interpreter contains the PyQt4 bindings, and may be used by Code_Saturne's configure script by passing PYTHON=<path_to_python_executable.

• add Python Qt4 bindings as a Python extension module for an existing Python installation. This is a more elegant solution than the previous one, and avoids requiring rebuilding Python, but if the user does not have administrator privileges, the extensions will be placed in a directory that is not on the default Python extension search path, and that must be added to the PYTHONPATH environment variable. This works fine, but for all users using this build of Code_Saturne, the PYTHONPATH environment variable will need to be set.³

The process is similar to the previous one, but SIP and PyQt4 installation requires a few additional configuration options in this case. See the SIP and PyQt4 reference guides for detailed instructions, especially the *Building a Private Copy of the SIP Module* section of the SIP guide.

4.3.2 Scotch and PT-Scotch

Note that both SCOTCH and PT-SCOTCH may be built from the same source tree, and installed together with no name conflicts.

For better performance, PT-SCOTCH may be built to use threads with concurrent MPI calls. This requires initializing MPI with MPI_Init_thread with MPI_THREAD_MULTIPLE (instead of the more restrictive MPI_THREAD_SERIALIZED, MPI_THREAD_FUNNELED, or MPI_THREAD_SINGLE, or simply using MPI_Init). As Code_Saturne does not support thread models in which different threads may call MPI functions simultaneously, and the use of MPI_THREAD_MULTIPLE may carry a performance penalty, we prefer to sacrifice some of PT-SCOTCH's performance by requiring that it be compiled without the -DSCOTCH_PTHREAD flag. This is not detected at compilation time, but with recent MPI libraries, PT-SCOTCH will complain at run time if it notices that the MPI thread safety level in insufficient.

Detailed build instructions, including troubleshooting instructions, are given in the source tree's INSTALL.txt file. In case of trouble, note especially the explanation relative to the dummysizes executable, which is run to determine the sizes of structures. On machines with different front-end and compute node architectures, it may be necessary to start the build process, let it fail, run this executable manually using mpirun, then pursue the build process.

4.3.3 MED

The Autotools installation of MED is simple on most machines, but a few remarks may be useful for specific cases.

MED has a C API, is written in a mix of C and C++ code, and provides both a C (libmedC) and an Fortran API (libmed). Both libraries are always built, so a Fortran compiler is required, but Code_Saturne only links using the C API, so using a different Fortran compiler to build MED and Code_Saturne is possible.

MED does require a C++ runtime library, which is usually transparent when shared libraries are used. When built with static libraries only, this is not sufficient, so when testing for a MED library, the Code_Saturne configure script also tries linking with a C++ compiler if linking with a C compiler fails. This must be the same compiler that was used for MED, to ensure the runtime matches. The choice of this C++ compiler may be defined passing the standard CXX variable to configure.

³In the future, the *Code_Saturne* installation scripts could check the PYTHONPATH variable and save its state in the build so as to ensure all the requisite directories are searched for.

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Also, when building MED in a cross-compiling situation, --med-int=int or --med-int=int64_t (depending on whether 32 or 64 bit ids should be used) should be passed to its configure script to avoid a run-time test.

4.3.4 libCCMIO

Different versions of this library may use different build systems, and use different names for library directories, so using both the --with-ccm= or --with-ccm-include= and --with-ccm-lib= options to configure is usually necessary. Also, the include directory should be the toplevel library, as header files are searched under a libccmio subdirectory⁴

A libCCMIO distribution usually contains precompiled binaries, but recompiling the library is recommended. Note that at least for version 2.06.023, the build will fail building dump utilities, due to the -1 adf option being placed too early in the link command. To avoid this, add LDLIBS=-ladf to the makefile command, for example:

make -f Makefile.linux SHARED=1 LDLIBS=-ladf

(SHARED=1 and DEBUG=1 may be used to force shared library or debug builds respectively).

Finally, if using libCCMIO 2.6.1, remove the libcgns* files from the libCCMIO libraries directory if also building *Code_Saturne* with CGNS support, as those libraries are not required for CCMIO, and are are an obsolete version of CGNS, which may interfere with the version used by *Code_Saturne*.

Note that libCCMIO uses a modified version of CGNS's ADF library, which may not be compatible with that of CGNS. When building with shared libraries, the reader for libCCMIO uses a plugin architecture to load the library dynamically. For a static build with both libCCMIO and CGNS support, reading ADF-based CGNS files may fail. To work around this issue, CGNS files may be converted to HDF5 using the adf2hdf utility (from the CGNS tools). By default, CGNS post-processing output files use HDF5, so this issue is rarer on output.

4.3.5 freesteam

This library's build instructions mention bindings with ascend, but those are not necessary in the context of *Code_Saturne*, so building without them is simplest. Its build system is based on scons, and builds on relatively recent systems with Python 2.7 should be straightforward.

With Python versions lower than 2.6, the command-line arguments allowing to choose the installation prefix (so as to place it in a user directory) are ignored, and its SConstruct file is not complete enough to allow setting flags for linking with an alternative, user-installed Python library outside the default linker search path. In this case, editing the SConstruct file to change the default paths is an ugly, but simple solution.

4.3.6 libxml2

This library is usually available on all general purpose Linux distributions, so it does not generally need to be rebuilt, although the associated development package (and headers) must as usual be installed.

For HPC clusters, it might not be available by default on compute nodes, so it may need to be compiled. This library may be downloaded atftp://xmlsoft.org/libxml2/ (with a possible mirror at ftp://fr.rpmfind.net/pub/libxml/). All versions tested so far (at least versions 2.6 to 2.9) have worked well with Code_Saturne, but to avoid compilation issues (as well as generating a smaller library), it may be useful to add the --with-ftp=no --with-http=no options to libxml2's configure script.

⁴this is made necessary by libCCMIO version 2.6.1, in which this is hard-coded in headers including other headers. In more recent versions such as 2.06.023, this is not the case anymore, and an include subdirectory is present, but it does not contain the libccmioversion.h file, which is found only under the libccmio subdirectory, and is required by Code_Saturne to handle differences between versions, so that source directory is preferred to the installation include.

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4.3.7 Paraview or Catalyst

By default, this library is built with a GUI, but it may also be be built using OSMesa for offscreen rendering. The build documentation on the ParaView website and Wiki details this. For use with Code_Saturne, the recommended solution is to build or use a standard ParaView build for interactive visualization, and to use its CatalystScriptGeneratorPlugin to generate Python co-processing scripts. A second build, using OSMesa, may be used for in-situ visualization. This is the Version Code_Saturne will be linked to. A recommended cmake command for this build contains:

```
$ cmake \
-DCMAKE_INSTALL_PREFIX=$INSTALL_PATH/arch/$machine_name_osmesa \
-DPARAVIEW_BUILD_QT_GUI=OFF \
-DPARAVIEW_USE_MPI=ON \
-DPARAVIEW_ENABLE_PYTHON=ON \
-DPARAVIEW_INSTALL_DEVELOPMENT_FILES=ON \
-DVTK_USE_X=OFF \
-DOPENGL_INCLUDE_DIR=$MESA_INSTALL_PREFIX/include \
-DOPENGL_g1_LIBRARY=\"\" \
-DOPENGL_g1_LIBRARY=$MESA_INSTALL_PREFIX/lib/libGLU.so \
-DVTK_OPENGL_HAS_OSMESA=ON \
-DOSMESA_INCLUDE_DIR=$MESA_INSTALL_PREFIX/include \
-DOSMESA_LIBRARY=$MESA_INSTALL_PREFIX/include \
-DOSMESA_LIBRARY=$MESA_INSTALL_PREFIX/lib/libOSMesa.so \
$PARAVIEW_SRC_PATH
```

5 Preparing for build

If the code was obtained as an archive, it must be unpacked:

```
tar xvzf saturne.tar.gz
```

If for example you unpacked the directory in a directory named /home/user/Code_Saturne, you will now have a directory named /home/user/Code_Saturne/saturne.

It is recommended to build the code in a separate directory from the source. This also allows multiple builds, for example, building both an optimized and a debugging version. In this case, choose a consistent naming scheme, using an additional level of sub-directories, for example:

```
$ mkdir saturne_build
$ cd saturne_build
$ mkdir prod
$ cd prod
```

Some older system's make command may not support compilation in a directory different from the source directory (VPATH support). In this case, installing and using the GNU gmake tool instead of the native make is recommended.

5.1 Source trees obtained through a source code repository

For developers obtaining the code was obtained through a version control system such as Subversion, an additional step is required:

```
$ cd saturne
$ ./sbin/bootstrap
$ cd ..
```

In this case, additional tools need to be available:

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- GNU Autotools: Autoconf, Automake, Libtool (2.2 or 2.4), and Gettext.
- Bison (or Yacc) and Flex (or Lex)
- PdfLaTeX and TransFig

These tools are not necessary for builds from tarballs; they are called when building the tarball (using make dist), so as to reduce the number of prerequisites for regular users, while developers building the code from a repository can be expected to need a more complete development environment.

Also, to build and install the documentation when building the code from a repository instead of a tarball, the following stages are required:

```
$ make doc
$ make install-doc
```

6 Configuration

Code_Saturne uses a build system based on the GNU Autotools, which includes its own documentation.

To obtain the full list of available configuration options, run: configure --help.

Note that for all options starting with --enable-, there is a matching option with --disable-. Similarly, for every --with-, --without- is also possible.

Select configuration options, then run configure, for example:

```
$ /home/user/Code_Saturne/4.0.5/src/code_saturne-4.0.5/configure \
--prefix=/home/user/Code_Saturne/4.0/arch/prod \
--with-med=/home/user/opt/med-3.1 \
CC=/home/user/opt/mpich-3.0/bin/mpicc FC=gfortran
```

In the rest of this section, we will assume that we are in a build directory separate from sources, as described in §5. In different examples, we assume that third-party libraries used by *Code_Saturne* are either available as part of the base system (i.e. as packages in a Linux distribution), as Environment Modules, or are installed under a separate path.

6.1 Debug builds

It may be useful to install debug builds alongside production builds of *Code_Saturne*, especially when user subroutines are used and the risk of crashes due to user programming error is high. Running the code using a debug build is significantly slower, but more information may be available in the case of a crash, helping understand and fix the problem faster.

Here, having a consistent and practical naming scheme is useful. For a side-by-side debug build for the example above, we simply replace prod by dbg in the --prefix option, and add --enable-debug to the configure command:

```
$ cd ..
$ mkdir dbg
$ cd dbg
$ ../../code_saturne-4.0.5/configure \
--prefix=/home/user/Code_Saturne/4.0/arch/dbg \
--with-med=/home/user/opt/med-3.1 \
--enable-debug \
CC=/home/user/opt/mpich-3.0/bin/mpicc FC=gfortran
```

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6.2 Shared or static builds

By default, on most architectures, *Code_Saturne* will be built with shared libraries. Shared libraries may be disabled (in which case static libraries are automatically enabled) by adding --disable-shared to the options passed to configure. On some systems, such as BlueGene/Q, the build may default to static libraries instead.

It is possible to build both shared and static libraries by adding --disable-static to the configure options, but the executables will be linked with the shared version of the libraries, so this is rarely useful (the build process is also slower in this case, as each file is compiled twice).

In some cases, a shared build may fail due to some dependencies on static-only libraries. In this case, --disable-shared will be necessary. Disabling shared libraries has also been seen to avoid issues with linking on Mac OSX systems.

In any case, be careful if you switch from one option to the other: as linking will be done with shared libraries by default, a build with static libraries only will not completely overwrite a build using shared libraries, so uninstalling the previous build first is recommended.

6.3 Relocatable builds

By default, a build of *Code_Saturne* is not movable, as not only are library paths hard-coded using *rpath* type info, but the code's scripts also contain absolute paths.

To ensure a build is movable, pass the --enable-relocatable option to configure.

Movable builds assume a standard directory hierarchy, so when running configure, the <code>--prefix</code> option may be used, but fine tuning of installation directories using options such as <code>--bindir</code>, <code>--libdir</code>, or <code>--docdir</code> must not be used (these options are useful to install to strict directory hierarchies, such as when packaging the code for a Linux distribution, in which case making the build relocatable would be nonsense anyways, so this is not an issue. ⁵

6.4 Compiler flags and environment variables

As usual when using an Autoconf-based configure script, some environment variables may be used. configure --help will provide the list of recognized variables. CC and FC allow selecting the C and Fortran compiler respectively (possibly using an MPI compiler wrapper).

Compiler options are usually defined automatically, based on detection of the compiler (and depending on whether --enable-debug was used). This is handled in a config/cs_auto_flags.sh and libple/config/ple_auto_flags.sh scripts. These files are sourced when running configure, so any modification to it will be effective as soon as configure is run. When installing on an exotic machine, or with a new compiler, adapting this file is useful (and providing feedback to the Code_Saturne development team will enable support of a broader range of compilers and systems in the future.

The usual CPPFLAGS, CFLAGS, FCCFLAGS, LDFLAGS, and LIBS environment variables may also be used, an flags provided by the user are appended to the automatic flags. To completely disable automatic setting of flags, the --disable-auto-flags option may be used.

6.5 MPI compiler wrappers

MPI environments generally provide compiler wrappers, usually with names similar to mpicc for C, mpicxx for C++, and mpif90 for Fortran 90. Wrappers conforming to the MPI standard recommen-

⁵In the special case of packaging the code, which may require both fine-grained control of the installation directories and the possibility to support options such as dpgg's --instdir, it is assumed the packager has sufficient knowledge to update both *rpath* information and paths in scripts in the executables and python package directories of a non-relocatable build, and that the packaging mechanism includes the necessary tools and scripts to enable this.

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dations should provide a -show option, to show which flags are added to the compiler so as to enable MPI. Using wrappers is fine as long as several third-party tools do not provide their own wrappers, in which case either a priority must be established. For example, using HDF5's h5pcc compiler wrapper includes the options used by mpicc when building HDF5 with parallel IO, in addition to HDF5's own flags, so it could be used instead of mpicc. On the contrary, when using a serial build of HDF5 for a parallel build of Code_Saturne, the h5cc and mpicc wrappers contain different flags, so they are in conflict.

Also, some MPI compiler wrappers may include optimization options used to build MPI, which may be different from those we wish to use that were passed.

To avoid issues with MPI wrappers, it is possible to select an MPI library using the --with-mpi option to configure. For finer control, --with-mpi-include and --with-mpi-lib may be defined separately.

Still, this may not work in all cases, as a fixed list of libraries is tested for, so using MPI compiler wrappers is the simplest and safest solution. Simply use a CC=mpicc or similar option instead of --with-mpi.

There is no need to use an FC=mpif90 or equivalent option: in *Code_Saturne*, MPI is never called directly from Fortran code, so Fortran MPI bindings are not necessary.

6.6 Environment Modules

As noted in §4.1, on systems providing Environment Modules with the module command, Code_Saturne's configure script detects which modules are loaded and saves this list so that future runs of the code use that same environment, rather than the user's environment, so as to allows using versions of Code_Saturne built with different modules safely and easily.

Given this, it is recommended that when configuring and installing *Code_Saturne*, only the modules necessary for that build of for profiling or debugging be loaded. Note that as *Code_Saturne* uses the module environment detected and runtime instead of the user's current module settings, debuggers requiring a specific module may not work under a standard run script if they were not loaded when installing the code.

The detection of environment modules may be disabled using the --without-modules option, or the use of a specified (colon-separated) list of modules may be forced using the --with-modules= option.

6.7 Remarks for very large meshes

If *Code_Saturne* is to be run on large meshes, several precautions regarding its configuration and that of third-party software must be taken.

in addition to local connectivity arrays, $Code_Saturne$ uses global element ids for some operations, such as reading and writing meshes and restart files, parallel interface element matching, and post-processing output. For a hexahedral mesh with N cells, the number of faces is about 3N (6 faces per cell, shared by 2 cells each). With 4 cells per face, the $face \rightarrow vertices$ array is of size of the order of $4 \times 3N$, so global ids used in that array's index will reach 2^{31} for a mesh in the range of $2^{31}/12 \approx 178.10^6$. In practice, we have encountered a limit with slightly smaller meshes, around 150 million cells.

Above 150 million hexahedral cells or so, it is thus imperative to configure the build to use 64-bit global element ids. This is the default. Local indexes use the default int size. To slightly decrease memory consumption if meshes of this size are never expected (for example on a workstation or a small cluster), the --disable-long-gnum option may be used.

Recent versions of some third-party libraries may also optionally use 64-bit ids, independently of each other or of *Code_Saturne*. This is the case for the SCOTCH and METIS, MED and CGNS libraries. In the case of graph-based partitioning, only global cell ids are used, so 64-bit ids should not in theory

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be necessary for meshes under 2 billion cells. In a similar vein, for post-processing output using nodal connectivity, 64-bit global ids should only be an imperative when the number of cells or vertices approaches 2 billion. Practical limits may be lower, if some intermediate internal counts reach these limits earlier.

Partitioning a 158 million hexahedral mesh using serial METIS 5 or SCOTCH on a front-end node with 128 Gb memory is possible, but partitioning the same mesh on cluster nodes with "only" 24 Gb each may not, so using parallel partitioning PT-SCOTCH or PARMETIS should be preferred.

6.8 Installation with the SALOME platform

To enable SALOME platform (http://www.salome-platform.org) integration, the --with-salome configuration option should be used, so as to specify the directory of the SALOME installation (note that this should be the main installation directory, not the default application directory, also generated by SALOME's installers).

With SALOME support enabled, both the CFDSTUDY salome module (available by running code_saturne salome) after install) and the *Code_Aster* coupling adapter should be available.

Note that specifying a SALOME directory does not automatically allow the *Code_Saturne* configure script to find some libraries which may be available in the SALOME distribution, such as HDF5, MED, or CGNS. It is possible to use those libraries by specifying the appropriate paths in the matching options, but other versions and builds may also be used (as those file formats are portable, MED or CGNS files produced by either *Code_Saturne* or SALOME remain interoperable).

Also note that for SALOME builds containing their own Python interpreter and library, using that same interpreter for <code>Code_Saturne</code> may avoid some issues, but may then require sourcing the SALOME environment or at least its Python-related <code>LD_LIBRARY_PATH</code> for the main <code>Code_Saturne</code> script to be usable.

6.9 Example configuration commands

Most available prerequisites are auto-detected, so to install the code to the default /usr/local sub-directory, a command such as:

\$../../code_saturne-4.0.5/configure

should be sufficient.

For the following examples, Let us define environment variables respectively reflecting the *Code_Saturne* source path, installation path, and a path where optional libraries are installed:

- \$ SRC_PATH=/home/projects/Code_Saturne/4.0.5
- \$ INSTALL_PATH=/home/projects/Code_Saturne/4.0
- \$ CS_OPT=/home/projects/opt

For an install on which multiple versions and architectures of the code should be available, configure commands with all bells and whistles (except SALOME support) for a build on a cluster named athos, using the Intel compilers (made available through environment modules) may look like this:

```
$ module purge
$ module load intel_compilers/14.0.0.080
$ module load open_mpi/gcc/1.6.5
$ $SRC_PATH/code_saturne-4.0.5/configure \
--prefix=$INSTALL_PATH/arch/athos_ompi \
--with-blas=/opt/intel/l_ics2013.1.039/composerxe_xe_2013_sp1/mkl \
--with-libxml2=$CS_0PT/libxml2-2.8/arch/athos \
--with-hdf5=$CS_0PT/hdf5-1.8.10/arch/athos \
--with-med=$CS_0PT/med-3.1/arch/athos \
--with-cgns=$CS_0PT/cgns-3.2/arch/athos \
--with-ccm=$CS_0PT/libccmio-2.06.23/arch/athos \
--with-scotch=$CS_0PT/scotch-6.0/arch/athos_ompi \
--with-metis=$CS_0PT/parmetis-4.0/arch/athos_ompi \
--with-eos/$CS_0PT/eos-1.2.0/arch/athos_ompi \
CC=mpicc FC=ifort CXX=icpc
```

In the example above, we have appended the <code>_ompi</code> postfix to the architecture name for libraries using MPI, in case we intend to install 2 builds, with different MPI libraries (such as Open MPI and MPICH-based Intel MPI). Note that optional libraries using MPI must also use the same MPI library. This is the case for PT-SCOTCH or PARMETIS, but also HDF5, CGNS, and MED if they are built with MPI-IO support. Similarly, C++ and Fortran libraries, and even C libraries built with recent optimizing C compilers, may require runtime libraries associated to that compiler, so if versions using different compilers are to be installed, it is recommended to use a naming scheme which reflects this. In this example, HDF5, CGNS and MED were built without MPI-IO support, as <code>Code_Saturne</code> does not yet exploit MPI-IO for these libraries.

To avoid copying platform-independent data (such as the documentation) from different builds multiple times, we may use the same --datarootdir option for each build so as to install that data to the same location for each build.

6.10 Cross-compiling

On machines with different front-end and compute node architectures, such as IBM Blue Gene/Q, cross-compiling is necessary. To install and run *Code_Saturne*, 2 builds are required:

- a "front-end" build, based on front-end node's architecture. This is the build whose code_saturne command, GUI, and documentation will be used, and with which meshes may be imported (i.e. whose Preprocessor will be used). This build is not intended for calculations, though it could be used for mesh quality criteria checks. This build will thus usually not need MPI.
- a "compute" build, cross-compiled to run on the compute nodes. This build does not need to include the GUI, documentation, or the Preprocessor.

A debug variant of the compute build is also recommended, as always. Providing a debug variant of the front-end build is not generally useful.

A post-install step (see §8) will allow the scripts of the front-end build to access the compute build in a transparent manner, so it will appear to the users that they are simply working with that build.

Depending on their role, optional third-party libraries should be installed either for the front-end, for the compute nodes, or both:

- BLAS will be useful only for the compute nodes, and are generally always available on large compute facilities.
- Python and PyQt4 will run on the front-end node only.

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- Libxml2 must be available for the compute nodes if the GUI is used.
- HDF5, MED, CGNSlib, and libCCMIO may be used by the Preprocessor on the front-end node to import meshes, and by the main solver on the compute nodes to output visualization meshes and fields.
- Scotch or Metis may be used by a front-end node build of the solver, as serial partitioning of large meshes requires a lot of memory.
- PT-SCOTCH or PARMETIS may be used by the main solver on the compute nodes.

6.10.1 Cross-compiling configuration for Blue Gene/Q

In our example, the front-end node is based on an IBM Power architecture, running under Red Hat Enterprise Linux 6, on which the Python/Qt4 environment should be available as an RPM package, and installed by the administrators. If this is not possible, the Python/Qt4 aspects of the Blue Gene/P example may be adapted here.

On the compute nodes, the IBM XL compilers produce static object files by default, so specifying the --disable-shared option is not necessary for libraries using Autotools-based installs when using those compilers, though using --build=ppc64 --host=bluegeneq in this case ensures the cross-compiling environment is detected. This environment is the suggested default, and prevents shared library builds. To allow building with shared libraries while still ensuring the cross-compiling environment is detected, use --host=powerpc64-bgq-linux instead.

As the front-end nodes of Blue Gene/Q machines may be expected to run Red Hat EL 6.x linux variants, instead of 5.x for blue Gene/P, more up-to date compilers and libraries (such as Python/Qt4) should be available as packages, easily installable by the system administrators.

For the compute nodes, the following remarks may be mode for prerequisites:

- LibXml: to reduce the size and simplify the installation of this library, the --with-ftp=no, --with-http=no, and --without-modules options may be used with configure.
- HDF5: building this library with its configure script is a pain⁶, but installing HDF5 1.8.9 or above using CMake is as simple as on a workstation, and simply requires choosing the correct compilers and possibly a few other options (in the EDF *Code_Saturne* build, the GCC compilers were chosen to reduce risks, and the Fortran wrappers were not needed, so not built).
- CGNS: building CGNS 3.1 or 3.2 is based on CMake, and no specific problems have been observed.
- MED: building MED 3.0.5 or above for *Code_Saturne* is easier than previous versions, as a new --disable-fortan option is available for the configure script. Both the C and C++ compiler wrappers must be specified, and the link may fail with the GNU compilers, due to some shared library issue (trying to force --disable-shared). With the IBM XL compilers, the same build works fine, as long as the CXXFLAGS=-qlanglvl=redefmac) option is passed. Adding the HDF5 tools path to the \$PATH environment variable for the configuration stage may also be required.

For an example, let us start with the front-end build:

```
$ $$RC_PATH/code_saturne-4.0.5/configure \
--prefix=$INSTALL_PATH/arch/frontend \
--with-hdf5=$CS_OPT/hdf5-1.8.13/arch/frontend \
--with-med=$CS_OPT/med-3.1/arch/frontend \
--with-cgns=$CS_OPT/cgns-3.2/arch/frontend \
--with-scotch=$CS_OPT/scotch-6.0/arch/frontend
```

⁶It requires running a yodconfigure script and adapting other scripts (see documentation), then running this as a submitted job (or under a SLURM allocation if you are lucky enough to use this resource manager).

For the compute node, we use the same version of Python (which is used only for the GUI and scripts, which only run on the front-end or service nodes), but the compilers are cross-compilers for the compute nodes:

```
$ $SRC_PATH/code_saturne-4.0.5/configure \
--prefix=$INSTALL_PATH/arch/bgq \
--with-blas=/opt/ibmmath/essl/5.1 \
--with-blas-type=ESSL \
--with-libxml2=$CS_OPT/libxml2-2.8/arch/bgq \
--with-hdf5=$CS_OPT/hdf5-1.8.13/arch/bgq \
--with-med=$CS_OPT/med-3.1/arch/bgq \
--with-cgns=$CS_OPT/cgns-3.2/arch/bgq \
--with-scotch=$CS_OPT/scotch-6.0/arch/bgq \
--with-scotch=$CS_OPT/scotch-6.0/arch/bgq \
--disable-sockets --disable-dlloader -disable-nls \
--disable-frontend --enable-long-gnum \
--build=ppc64 --host=bluegeneq \
CC=/bgsys/drivers/ppcfloor/comm/xl/bin/mpixlc_r \
CXX=/bgsys/drivers/ppcfloor/comm/xl/bin/mpixlcxx_r \
FC=bgxlf95_r
```

The C++ compiler is specified, as it will be needed for the link stage due to C++ dependencies in the MED library, which is a static library in this example (see $\S4.3.3$).

Note that in the above examples, we specified an install of the SCOTCH partitioning library both for the front-end and for the compute nodes. The implies a serial build of SCOTCH on the front-end node, and a parallel build (PT-SCOTCH) on the compute nodes. Both are optional, and the serial partitioning on the front-end nodes should only be used as a backup or as a reference for parallel partitioning. Unless robustness or quality issues are encountered with parallel partitioning, it should supercede serial partitioning, as it allows for a simpler toolchain even for large meshes. Similarly, METIS could be used on the front-end node, and PARMETIS on the compute nodes.

6.10.2 Compiling for Cray X series

For Cray X series, when using the GNU compilers, installation should be similar to that on standard clusters. Using The Cray compilers, options such as in the following example are recommended:

```
$ $$RC_PATH/code_saturne-4.0.5/configure \
--prefix=$INSTALL_PATH/arch/xc30 \
--with-libxm12=$CS_OPT/libxm12-2.8/arch/xc30 \
--with-hdf5=$CS_OPT/hdf5-1.8.13/arch/xc30 \
--with-med=$CS_OPT/med-3.1/arch/xc30 \
--with-cgns=$CS_OPT/cgns-3.2/arch/xc30 \
--with-scotch=$CS_OPT/scotch-6.0/arch/xc30 \
--with-scotch=$CS_OPT/scotch-6.0/arch/xc30 \
--disable-sockets --disable-nls \
--disable-shared \
--host=x86_64-unknown-linux-gnu \
CC=cc \
CXX=CC \
FC=ftn
```

In case the automated environment modules handling causes issues, adding the --without-modules option may be necessary. In that case, caution must be exercised so that the user will load the same modules as those used for installation. This is not an issue if modules for *Code_Saturne* is also built, and the right dependencies handled at that level.

Note that to build without OpenMP with the Cray compilers, CFLAGS="h noomp" and FCFLAGS="h

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noomp" need to be added.

6.11 Troubleshooting

If configure fails and reports an error, the message should be sufficiently clear in most case to understand the cause of the error and fix it. Do not forget that for libraries installed using packages, the development versions of those packages are also necessary, so if configure fails to detect a package which you believe is installed, check the matching development package.

Also, whether it succeeds or fails, configure generates a file named config.log, which contains details on tests run by the script, and is very useful to troubleshoot configuration problems. When configure fails due to a given third-party library, details on tests relative to that library are found in the config.log file. The interesting information is usually in the middle of the file, so you will need to search for strings related to the library to find the test that failed and detailed reasons for its failure.

7 Compile and install

Once the code is configured, it may be compiled and installed; for example, to compile the code (using 4 parallel threads), then install it:

\$ make -j 4 && make install

To compile the documentation, add:

\$ make pdf && make install-pdf

To clean the build directory, keeping the configuration, use make clean; To uninstall an installed build, use make uninstall. To clear all configuration info, use make distclean (make uninstall will not work after this).

7.1 Installing to a system directory

When installing to a system directory, such as /usr or /usr/local, some Linux systems may require running ldconfig as root or sudoer for the code to work correctly.

8 Post-install

Once the code is installed, a post-install step may be necessary for computing environments using a batch system, for separate front-end and compute systems (such as Blue Gene systems), or for coupling with SYRTHES 4 or Code_Aster. The global default MPI execution commands and options may also be overridden.

Copy or rename the <install-prefix>/etc/code_saturne.cfg.template to <install-prefix>/etc/code_saturne.cfg, and uncomment and define the applicable sections.

If used, the name of the batch system should match one of the templates in <install-prefix>/share/code_saturne/batch, and those may also be edited if necessary to match the local batch configuration⁷

Also, the compute_versions section allows the administrator to define one or several alternate builds which will be used for compute stages. This is especially useful for installation on BlueGene type machines, where 2 separate builds are required (one for the front-end nodes and one for the compute nodes). The compute-node build may be configured using the --disable-frontend option so as only

⁷Some batch systems allow a wide range of alternate and sometimes incompatible options or keywords, and it is for all practical purposes impossible to determine which options are allowed for a given setup, so editing the batch template to match the local setup may be necessary.

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to build and install the components required to run on compute-nodes, while the front-end build may be configured without MPI support. The front-end build's post-install step allows definition of the associated compute build.

All default MPI execution commands and options may be overriden using the mpi section. Note that only the options defined in this section are overridden; defaults continue to apply for all others.

9 Installing for SYRTHES coupling

Coupling with SYRTHES 4 requires defining the path to SYRTHES 4 at the post-install stage.

When coupling with SYRTHES 4, both <code>Code_Saturne</code> and SYRTHES must use the same MPI library, and must use the same version of the PLE (Parallel Location and Exchange) library from <code>Code_Saturne</code>. By default, PLE is built as a sub-library of <code>Code_Saturne</code>, but a standalone version may be configured and built, using the <code>libple/configure</code> script from the <code>Code_Saturne</code> source tree, instead of the top-level <code>configure</code> script. <code>Code_Saturne</code> may then be configured to use the existing install of PLE using the <code>--with-ple</code> option. Similarly, SYRTHES must also be configured to use PLE.

Alternatively, SYRTHES 4 may simply be configured to use the PLE library from an existing Code_Saturne install.

10 Shell configuration

If *Code_Saturne* is installed in a non-default system directory (i.e. outside /usr or /usr/local, it is recommended to define an alias (in the user's .alias or .profile file, so as to avoid needing to type the full path when using the code:

alias code_saturne="\$prefix/code_saturne-\$version/bin/code_saturne"

Note that when multiple versions of the code are installed side by side, using a different alias for each will allow using them simultaneously, with no risk of confusion.

If using the bash shell, you may also source a bash completion file, so as to benefit from shell completion for *Code_Saturne* commands and options, either using

. <install-prefix>/etc/bash_completion.d/code_saturne

or

source <install-prefix>/etc/bash_completion.d/code_saturne

On some systems, only the latter syntax is effective. For greater comfort, you should save this setting in your .bashrc or .bash_profile file.

11 Caveats

11.0.1 Moving an existing installation

Never move an non-relocatable installation of Code_Saturne. Using LD_LIBRARY_PATH or LD_PRELOAD may allow the executable to run despite rpath info not being up-to-date, but in environments where different library, versions are available, there is a strong risk of not using the correct library. In addition, the scripts will not work unless paths in the installed scripts are updated.

To build a relocatable installation, see section 6.3.

If you are packaging the code and need both fine-grained control of the installation directories, and the possibility to support options such as dpgg's --instdir, it is assumed you have sufficient knowledge to update both *rpath* information and paths in scripts in the executables and python package directories,

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and that the packaging mechanism includes the necessary tools and scripts to enable this.

In any other case, you should not even think about moving a non-relocatable build.

If you need to test an installation in a test directory before installing it in a production directory, use the make install DESTDIR=<test_prefix> provided by the Autotools mechanism rather than configuring an install for a test directory and then moving it to a production directory. Another less elegant but safe solution is to configure the build for installation to a test directory, and once it is tested, re-configure the build for installation to the final production directory, and rebuild and install.