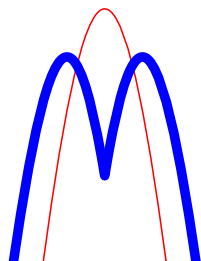


MOLPRO



Installation Guide Version 2015.1

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1 Obtaining the distribution materials

MOLPRO is distributed to licensees on a self-service basis using the world-wide web. Those entitled to the code should obtain it from <https://www.molpro.net/download> supplying the username and password given to them. The web pages contain both source code and binaries, although not everyone is entitled to source code, and binaries are not available for every platform.

Execution of MOLPRO, whether a supplied binary or built from source, requires a valid licence key. Note that the key consists of two components, namely a list of comma-separated key=value pairs, and a password string, and these are separated by '&'. In most cases the licence key will be automatically downloaded from the website when building or installing the software.

2 Installation of pre-built binaries

Binaries are given as self-extracting tar archives which are installed by running them on the command line. There are binaries tuned for several architectures. These also support parallel execution. The parallel binaries are built using GA with MPI.

The tar archives are fully relocatable, the location can be changed when running the script interactively, the default is `/usr/local`.

If the script finds a licence key which has been cached in `$HOME/.molpro/token` from a previous install then that key will be installed with the software. If the script cannot find a key or automatically download it from the molpro website then the script will prompt that this part of the install has failed. All files of Molpro are installed, but the user must then manually install the key with the library files in a file named `.token`, e.g.:

```
/usr/local/molpro-mpptype-arch/lib/.token
```

Other configuration options as described in section 3.5 may also be specified in the script file:

```
/usr/local/bin/molpro
```

3 Installation from source files

3.1 Overview

There are usually four distinct stages in installing MOLPRO from source files:

Configuration	A shell script that allows specification of configuration options is run, and creates a configuration file that drives subsequent installation steps.
Compilation	The program is compiled and linked, and other miscellaneous utilities and files, including the default options file, are built. The essential resulting components are <ol style="list-style-type: none">1. The <code>molpro</code> shell script which launches the main executable. In serial case one can directly run the main executable.2. The <code>molpro.exe</code> executable, which is the main program. For parallel computation, multiple copies of <code>molpro.exe</code> are started by a single instance of <code>molpro</code> shell script using the appropriate system utility, e.g. <code>mpirun</code>.

	3. Machine-ready basis-set, and other utility, libraries.
Validation	A suite of self-checking test jobs is run to provide assurance that the code as built will run correctly.
Installation	The program can be run directly from the source tree in which it is built, but it is usually recommended to run the procedure that installs the essential components in standard system directories.

3.2 Prerequisites

The following are required or strongly recommended for installation from source code.

1. A Fortran 2003 compiler. On HPC systems the latest vendor-supplied compiler should be used. The program is regularly tested with recent versions of GNU and Intel Fortran compilers.
2. GNU *make*, freely available from <http://www.fsf.org> and mirrors. GNU *make* must be used; most system-standard makes do not work. In order to avoid the use of a wrong *make*, it may be useful to set an alias, e.g., `alias make='gmake -s'`. A recent version of GNU *make* is required, 3.80 or above.
3. Libxml2 library and header files.
4. About 10GB disk space (strongly system-dependent; more with large-blocksize file systems, and where binary files are large) during compilation. Typically 100Mb is needed for the finally installed program. Large calculations will require larger amounts of disk space.
5. One or more large scratch file systems, each containing a directory that users may write on. There are parts of the program in which demanding I/O is performed simultaneously on two different files, and it is therefore helpful to provide at least two filesystems on different physical disks if other solutions, such as striping, are not available. The directory names should be stored in the environment variables `$TMPDIR`, `$TMPDIR2`, `$TMPDIR3`,... These variables should be set before the program is installed (preferably in `.profile` or `.cshrc`), since at some stages the installation procedures will check for them (cf. section 3.5).
6. If the program is to be built for parallel execution then the Global Arrays toolkit or the MPI-2 library is needed. For building MOLPRO with the Global Arrays toolkit, we recommend the latest stable version (although earlier versions may also work). This is available from <http://www.emsl.pnl.gov/docs/global> and should be installed prior to compiling MOLPRO. For building MOLPRO with the MPI-2 library, we recommend to use the built-in MPI-2 library, which may have advantages of optimization on some platforms. If there is no built-in one on the platform, a fresh MPI-2 library (e.g.: MPICH2, see <http://http://www.mpich.org/>) should be installed prior to compiling MOLPRO. Many MPI-2 libraries, including Intel MPI, Bull MPI, MPICH2, and Open MPI, have been tested, and others untested could also work.
7. The source distribution of MOLPRO, which consists of a compressed tar archive with a file name of the form `molpro.2015.1.tar.gz`. The archive can be unpacked using `gunzip` and `tar`.

3.2.1 Fedora packages

To build using GNU compilers one should ensure the following packages are installed (via dnf):

gcc-c++	provides GNU C and C++ compiler,
gcc-gfortran	provides GNU Fortran compiler,
libxml2-devel	provides libxml2 library & headers,
make	provides GNU make.

Optionally one can choose to install:

blas-devel	provides a BLAS library,
lapack-devel	provides a LAPACK library,

which will be used instead of compiling the equivalent MOLPRO routines.

3.2.2 openSUSE packages

To build using GNU compilers one should ensure the following packages are installed (via YaST):

gcc-c++	provides GNU C and C++ compiler,
gcc-fortran	provides GNU Fortran compiler,
libxml2-devel	provides libxml2 library & headers,
make	provides GNU make.

Optionally one can choose to install:

blas	provides a BLAS library,
lapack	provides a LAPACK library,

which will be used instead of compiling the equivalent MOLPRO routines.

3.2.3 Ubuntu packages

To build using GNU compilers one should ensure the following packages are installed via (apt-get):

build-essential	provides GNU C++ compiler,
curl	provides curl for downloading patches,
gfortran	provides GNU Fortran compiler,
libxml2-dev	provides libxml2 library & headers,
openssh-server	provides ssh access to localhost.

Optionally one can choose to install:

libblas-dev provides a BLAS library,
liblapack-dev provides a LAPACK library,

which will be used instead of compiling the equivalent MOLPRO routines. Set up password-less ssh by running the following commands and not entering a password when prompted:

```
ssh-keygen -t rsa
cat ~/.ssh/id_rsa.pub >> ~/.ssh/authorized_keys
```

This must be done for each user account which will be running MOLPRO.

3.2.4 macOS

It is recommended to install and manage the additional compilers and utilities needed for building and running MOLPRO using the Homebrew system. Visit <http://brew.sh>, and copy the installation command into a terminal window. On an empty system, you will be prompted to install command line tools; do this first. Then accept all prompts from the Homebrew installation script.

The following will install all utilities that might be needed for building and running MOLPRO for development in serial; for production calculations in parallel, a binary can be downloaded.

```
brew install Caskroom/cask/xquartz homebrew/x11/grace \
  open-babel doxygen automake wget \
  Caskroom/cask/java homebrew/science/jmol
```

3.3 Configuration

Once the distribution has been unpacked, change to the `Molpro` directory that has been created. Having changed to the `Molpro` directory, you should check that the directory containing the Fortran compiler you want to use is in your `PATH`. Then run the command

```
./configure
```

which creates the file `CONFIG`. This file contains machine-dependent parameters, such as compiler options. Normally `CONFIG` will not need changing, but you should at the least examine it, and change any configuration parameters which you deem necessary. Any changes made to `CONFIG` will be lost next time `./configure` is invoked, so it is best to supply as many of these as possible via the command line.

The `configure` procedure may be given command line options, and, some commonly used options include:

1. In the case of building for parallel execution, the option `--enable-mpp` must be given on the command line. This enables both `mpp` and `mppx` parallelism; for the distinction between these two parallelism modes, please refer to the user manual, section 2. The option `--enable-mpp` must be set to the location of the Global Arrays library directory or the MPI-2 library include directory.

For the case of using the Global Arrays toolkit, one example can be

```
./configure --enable-mpp=/usr/local/ga-install/lib
```

If using a Global Arrays build with an MPI library the appropriate MPI executable should appear first in PATH when more than one is available.

Queries regarding Global Arrays installations should be sent directly to the Global Arrays team, any Molpro related queries will assume a fully functional Global Arrays suite with all internal tests run successfully.

For the case of using the MPI-2 library, one example can be

```
./configure --enable-mpp=/usr/local/mpich2-install/include
```

and the directory should contain file `mpi.h`. Please ensure the built-in or freshly built MPI-2 library fully supports MPI-2 standard and works properly.

For desktop or single node installations, there are a series of options prefixed with `auto` which build any prerequisites, and can be passed to `--enable-mpp`, eg.

```
./configure --enable-mpp=auto-ga-mpich
```

2. For good performance it is important to use appropriate BLAS libraries; in particular, a fast implementation of the matrix multiplication `dgemm` is very important for MOLPRO. Therefore you should use a system tuned BLAS library whenever available.

MOLPRO will automatically detect the most appropriate BLAS library in many cases. In certain cases, in particular when the BLAS library is installed in a non-default location, `configure` should be directed to the appropriate directory with:

```
./configure --with-blas-path=/path/to/lib/dir
```

3. The Molpro installation directory can be given with `--prefix`.

3.4 Compilation and linking

After configuration, the remainder of the installation is accomplished using the GNU `make` command. Remember that the default `make` on many systems will not work, and that it is essential to use GNU `make` (cf. section 3.2). Everything needed to make a functioning program together with all ancillary files is carried out by default simply by issuing the command

```
make
```

in the MOLPRO base directory. Most of the standard options for GNU `make` can be used safely; in particular, `-j` can be used to speed up compilation on a parallel machine. The program can then be accessed by making sure the `bin/` directory is included in the PATH and issuing the command `molpro`. If MPI library is used for building Global Arrays or building MOLPRO directly, please be aware that some MPI libraries use `mpd` daemons to launch parallel jobs. In this case, `mpd` daemons must already be running before `make`.

3.5 Adjusting the default environment for MOLPRO

The default running options for MOLPRO are stored in the script `bin/molpro`. After program installation, either using binary or from source files, this file should be reviewed and adjusted, if necessary, to make system wide changes.

3.6 Tuning

MOLPRO can be tuned for a particular system by running in the root directory the command

```
make tuning
```

This job automatically determines a number of tuning parameters and appends these to the file `bin/molpro`. Using these parameters, MOLPRO will select the best BLAS routines depending on the problem size. This job should run on an empty system. It may typically take 10 minutes, depending on the processor speed, and you should wait for completion of this run before doing the next steps.

3.7 Testing

At this stage, it is essential to check that the program has compiled correctly. The makefile target `test` (i.e., command `make test`) will do this using the full suite of test jobs, and although this takes a significantly long time, it should always be done when porting for the first time. A much faster test, which checks the main routes through the program, can be done using `make quicktest`. For parallel installation, it is highly desirable to perform this validation with more than one running process. This can be done conveniently through the `make` command line as, for example,

```
make MOLPRO_OPTIONS=-n2 test
```

If any test jobs fail, the cause must be investigated. If, after due efforts to fix problems of a local origin, the problem cannot be resolved, the developers of MOLPRO would appreciate receiving a report. There is a web-based mechanism at <https://www.molpro.net/bugzilla> at which as many details as possible should be filled in. It may also be helpful to attach a copy of the `CONFIG` file along with the failing output. Please note that the purpose of such bug reports is to help the developers improve the code, and not for providing advice on installation or running.

3.8 Installing the program for production

Although the program can be used in situ, it is usually convenient to copy only those files needed at run time into appropriate installation directories as specified at configuration time (see section 3.3) and stored in the file `CONFIG`. To install the program in this way, do

```
make install
```

The complete source tree can then be archived and deleted. The overall effect of this is to create a shell script in the `INSTBIN` directory. The name should relate to the architecture, type of build, integer etc. Symbolic links relating to the type of build are then made, and finally providing that `INSTBIN/molpro` is not a file, a symbolic link is created to the new script. In some cases it is preferable to create a localized script in `INSTBIN/molpro` which will not be over written. The overall effect of this cascade of links is to provide, in the normal case, the commands `molpro` and one or both of `molpros` (serial) and `molprop` (parallel) for normal use, with the long names remaining available for explicit selection of particular variants.

For normal single-variant installations, none of the above has to be worried about, and the `molpro` command will be available from directory `INSTBIN`.

During the install process the key from `$HOME/.molpro/token` is copied to `PREFIX/.token` so that the key will work for all users of the installed version.

3.9 Installation of documentation

The documentation is available on the web at <http://www.molpro.net/info/users>. It is also included with the source code. The PDF user's manual is found in the directory `Molpro/doc/manual.pdf`, with the HTML version in the directory `Molpro/doc/manual/index.html`. After `make install` the documentation is installed in the `doc` subdirectory of `PREFIX` specified in `CONFIG` file generated by the `configure` command. Numerous example input files are included in the manual, and can alternatively be seen in the directory `Molpro/examples`.

3.10 Simple building for single workstations Linux or Mac OS X

The following instructions are quick instructions for installing MOLPRO on a single-workstation Linux or Mac OS X system. The instructions assume GNU compilers have been installed (details of getting GNU compilers for common Linux distributions are contained in the prerequisites section), but these can be substituted with alternative compilers. For serial MOLPRO:

```
./configure CXX=g++ FC=gfortran
make
```

For parallel MOLPRO one can use:

```
./configure CXX=g++ FC=gfortran --enable-mpp=/path/to/ga-install/lib
make
```

if Global Arrays has already been built. There is a simpler option, providing the `curl` utility is installed, and the machine is connected to the internet:

```
./configure CXX=g++ FC=gfortran --enable-mpp=auto-ga-mpich
make
```

which will automatically download and install MPICH and Global Arrays.

3.11 Installation on a Cygwin system

On a Windows machine Cygwin should be installed. In addition to the default package list one should also install the packages listed in table 1. If undertaking development work table 2

Package	Package Group	Reason
gcc-core	Devel	compiling C files
gcc-fortran	Devel	compiling Fortran files
gcc-g++	Devel	compiling C++ files
make	Devel	need GNU make
ca-certificates	Net	download boost
curl	Net	token download

Table 1: Cygwin requirements for user install

contains a list of potentially useful packages.

With the above steps, `configure` can be run and the Molpro built in the normal way.

Package	Package Group	Reason
bison	Devel	bison
gdb	Devel	gdb
git	Devel	git
libxslt	Libs	xsltproc
openssh	Net	ssh
vim	Editors	vi

Table 2: Cygwin packages for developers