



Developer's Manual for QUANTUM ESPRESSO(v. 4.1cvs)

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1 Introduction

1.1 Who should read (and who should *write*) this guide

The intended audience of this guide is everybody who wants to:

- know how Quantum ESPRESSO works, including its internals
- modify/customize/add/extend/improve/clean up Quantum ESPRESSO
- know how to read data produced by Quantum ESPRESSO

The same category of people should also "write" this guide, of course.

1.2 Who may read this guide but will not necessarily profit from it

People who want to know about the capabilities of Quantum ESPRESSO, or who want just to use Quantum ESPRESSO, should read the User Guide.

People who want to know about the methods or the physics behind Quantum ESPRESSO should read first the relevant literature.

1.3 How to Contribute to Quantum ESPRESSO

You can contribute to a better Quantum ESPRESSO by

- answering other people's questions on the mailing list (correct answers are strongly preferred to wrong ones).
- suggesting changes: note however that suggestions requiring a significant amount of work are welcome only if accompanied by implementation or by a promise of future implementation (fulfilled promises are strongly preferred to forgotten ones).
- porting to new/unsupported architectures or configurations: see the Installation mechanism section. You shouldn't need new preprocessing flags: those already existing should be sufficient. If you really need a new one, look into file `include/defs.h`.`README` to learn what is there and how it is used.
- pointing out bugs in the software and in the documentation (reports of real bugs are strongly preferred to reports of nonexistent bugs). A bug report should include enough information to be reproduced: typically, version number, hardware/software combination(s) for which the problem arises, whether it is reproducible or erratic, whether it happens in serial or parallel execution or both, and, most important, an input and output exhibiting such behavior (fast to execute if possible). The error message alone is usually not a sufficient piece of information.
- adding new features to the code. If you like to have something added to Quantum ESPRESSO, contact the developers. Unless there are technical reasons not to include your changes, we will try to make you happy (no warranty that we will actually succeed).

For extensive changes, the ideal procedure is as follows:

- download the current CVS version (see Using CVS) and work on that version
- when you are happy with your modified version, make a copy of it, then update your copy with `cvcs update`
- if you get no conflicts and everything is still working, you have won. Send the modified files to the developers.
- if you get conflicts, or if the updated code doesn't work any longer, you haven't yet won. Look into the conflicting section: in most cases conflicts are trivial (format changes, white spaces) or easily solved (the part of the code you were modifying has been moved to another place, for instance). Sometimes, somebody else has done changes that are incompatible with yours during the same period. Look into the ChangeLog (use the script `cvcs2cl.pl` to get an updated ChangeLog) to understand what may have happened. Use `cvcs update -D 2006-oct-12` to update to the CVS version of 12 oct 2006: this may be useful if you suspect that the incompatible change happened after that date. CVS versions can be *tagged*; tags are visible with `cvcs status -v`. Use `cvcs update -r TAG` to update to the CVS version with tag TAG. In all cases, use `cvcs update -A` to revert to the last version.

Developers can be contacted via the `pw_forum` mailing list.

2 Structure of the distribution

2.1 Contents of the various directories

2.1.1 Modules

2.1.2 Sources

2.1.3 Utilities

2.1.4 Libraries

Subdirectory `flib/` contains libraries written in fortran77 (*.f) and in fortran-90 (*.f90). The latter should not depend on any module, except for "kind" and "constants".

Subdirectory `clib/` contains libraries written in C (*.c). Functions that are called by fortran should be preprocessed using the macros:

1. `F77_FUNC(func,FUNC)` for function "func", not containing underscore(s) in name
2. `F77_FUNC(f_nc,F_NC)` for function "f_nc", containing underscore(s) in name

These macros are defined in file `include/c_defs.h`. This file must be included by all *.c files. The macros are automatically generated by "configure" and choose the correct case (lowercase or uppercase) and the correct number of final underscores. See file `include/defs.h`.README for more info.

2.2 Installation mechanism

The code contains C-style preprocessing directives. There are two ways to do preprocessing of fortran files:

- directly with the fortran compiler, if supported;
- by first pre-compiling with the C preprocessor `cpp`.

In the first case, one needs to specify in the `make.sys` file the fortran compiler option that tells the compiler to pre-process first. In the second case, one needs to specify the C precompiler and options (if needed) in `make.sys`. Normally, "configure" should take care of this.

2.2.1 How to edit the configure script

The "configure" script is generated from its source file "configure.ac" by the GNU Autoconf utility (<http://www.gnu.org/software/autoconf/>). Don't edit "configure" directly: whenever it gets regenerated, your changes will be lost. Instead, go to the "install/" directory, edit "configure.ac", then run "autoconf" to regenerate "configure". If you want to keep the old "configure", make a copy first.

GNU Autoconf is installed by default on most Unix/Linux systems. If you don't have it on your system, you'll have to install it. Also, if it bumps an error message saying "configure.in not found", you'll have to install a more recent version. Version 2.53 at least is known to work.

"configure.ac" is a regular Bourne shell script (i.e., "sh" – not csh!), except that:

- capitalized names starting with "AC_" are Autoconf macros. Normally you shouldn't have to touch them.

- square brackets are normally removed by the macro processor. If you need a square bracket (that should be very rare), you'll have to write two.

You may refer to the GNU Autoconf Manual for more info.

"make.sys.in" is the source file for "make.sys", that "configure" generates: you might want to edit that file as well. The generation procedure is as follows: if "configure.ac" contains the macro "AC_SUBST(name)", then every occurrence of "@name@" in the source file will be substituted with the value of the shell variable "name" at the point where AC_SUBST was called.

Similarly, "configure.msg" is generated from "configure.msg.in": this file is only used by "configure" to print its final report, and isn't needed for the compilation. We did it this way so that our "configure" may also be used by other projects, just by replacing the Quantum ESPRESSO-specific "configure.msg.in" by your own.

"configure" writes a detailed log of its operation to "config.log". When any configuration step fails, you may look there for the relevant error messages. Note that it is normal for some checks to fail.

2.2.2 How to add support for a new architecture

In order to support a previously unsupported architecture, first you have to figure out which compilers, compilation flags, libraries etc. should be used on that architecture. In other words, you have to write a "make.sys" that works: you may use the manual configuration procedure for that (see the Quantum ESPRESSO User Guide). Then, you have to modify "configure" so that it can generate that "make.sys" automatically.

To do that, you have to add the case for your architecture in several places throughout "configure.ac":

1. Detect architecture

Look for these lines:

```
if test "$arch" = ""
then
    case $host in
        ia64-*-linux-gnu )      arch=ia64    ;;
        x86_64-*-linux-gnu )    arch=x86_64  ;;
        *-pc-linux-gnu )       arch=ia32    ;;
        etc.
```

Here you must add an entry corresponding to your architecture and operating system. Run "config.guess" to obtain the string identifying your system. For instance on a PC it may be "i686-pc-linux-gnu", while on IBM SP4 "powerpc-ibm-aix5.1.0.0". It is convenient to put some asterisks to account for small variations of the string for different machines of the same family. For instance, it could be "aix4.3" instead of "aix5.1", or "athlon" instead of "i686"...

2. Select compilers

Look for these lines:

```

# candidate compilers and flags based on architecture
case $arch in
ia64 | x86_64 )
    ...
ia32 )
    ...
aix )
    ...
etc.

```

Add an entry for your value of \$arch, and set there the appropriate values for several variables, if needed (all variables are assigned some reasonable default value, defined before the "case" block):

- "try_f90" should contain the list of candidate Fortran 90 compilers, in order of decreasing preference (i.e. configure will use the first it finds). If your system has parallel compilers, you should list them in "try_mpif90".

- "try_ar", "try_arflags": for these, the values "ar" and "ruv" should be always fine, unless some special flag is required (e.g., -X64 With sp4).

- you should define "try_dflags" if the Quantum ESPRESSO codes contain any "#ifdef" specific to your machine: for instance, on IBM machines, "try_dflags=-D__AIX" . A list of such flags can be found in file include/defs.h.README

You shouldn't need to define the following two:

- "try_arflags_dynamics" should be set to the same as "try_arflags" except in special cases in which dynamically libraries are to be built (presently only on PowerPC Mac with OS-X and xlf compiler)

- "try_iflags" should be set to the appropriate "-I" option(s) needed by the preprocessor or by the compiler to locate *.h files to be included; try_iflags="-I../include" should be good for most cases

For example, here's the entry for IBM machines running AIX:

```

aix )
    try_mpif90="mpxlf90_r mpxlf90"
    try_f90="xlf90_r xlf90 $try_f90"
    try_arflags="-X64 ruv"
    try_arflags_dynamic="-X64 ruv"
    try_dflags="-D__AIX -D__XLF"
    ;;

```

The following step is to look for both serial and parallel fortran compilers:

```

# check serial Fortran 90 compiler...
...
AC_PROG_F77($f90)
...

```

```

        # check parallel Fortran 90 compiler
...
        AC_PROG_F77($mpif90)
...
echo setting F90... $f90
echo setting MPIF90... $mpif90

```

A few compilers require some extra work here: for instance, if the Intel Fortran compiler was selected, you need to know which version because different versions need different flags.

At the end of the test,

- \$mpif90 is the parallel compiler, if any; if no parallel compiler is found or if `-disable-parallel` was specified, \$mpif90 is the serial compiler
- \$f90 is the serial compiler

Next step: the choice of (serial) C and Fortran 77 compilers. Look for these lines:

```

# candidate C and f77 compilers good for all cases
try_cc="cc gcc"
try_f77="$f90"

case "$arch:$f90" in
*:f90 )
    ....
etc.

```

Here you have to add an entry for your architecture, and since the correct choice of C and f77 compilers may depend on the fortran-90 compiler, you may need to specify the f90 compiler as well. Again, specify the compilers in `try_cc` and `try_f77` in order of decreasing preference. At the end of the test,

- \$cc is the C compiler
- \$f77 is the Fortran 77 compiler, used to compile *.f files (may coincide with \$f90)

3. Specify compilation flags.

Look for these lines:

```

# check Fortran compiler flags
...
case "$arch:$f90" in
ia64:ifort* | x86_64:ifort* )
    ...
ia64:ifc* )
    ...
etc.

```

Add an entry for your case and define:

- "try_fflags": flags for Fortran 77 compiler.
- "try_f90flags": flags for Fortran 90 compiler. In most cases they will be the same as in Fortran 77 plus some others. In that case, define them as "\$(FFLAGS) -something_else".
- "try_fflags_noopt": flags for Fortran 77 with all optimizations turned off: this is usually "-O0". These flags must be used for compiling flib/dlamch.f (part of our version of Lapack): it won't work properly with optimization.
- "try_ldflags": flags for the linking phase (not including the list of libraries: this is decided later).
- "try_ldflags_static": additional flags to select static compilation (i.e., don't use shared libraries).
- "try_dflags": must be defined if there is in the code any `#ifdef` specific to your compiler (for instance, `-D__INTEL` for Intel compilers). Define it as "\$try_dflags -D..." so that pre-existing flags, if any, are preserved.
- if the Fortran 90 compiler is not able to invoke the C preprocessor automatically before compiling, set "have_cpp=0" (the opposite case is the default). The appropriate compilation rules will be generated accordingly. If the compiler requires that any flags be specified in order to invoke the preprocessor (for example, "-fpp " – note the space), specify them in "pre_fdflags".

For example, here's the entry for ifort on Linux PC:

```
ia32:ifort* )
    try_fflags="-O2 -tpp6 -assume byterecl"
    try_f90flags="\$(FFLAGS) -nomodule"
    try_fflags_noopt="-O0 -assume byterecl"
    try_ldflags=""
    try_ldflags_static="-static"
    try_dflags="$try_dflags -D__INTEL"
    pre_fdflags="-fpp "
    ;;
```

Next step: flags for the C compiler. Look for these lines:

```
case "$arch:$cc" in
*:icc )
    ...
*:pgcc )
    ...
etc.
```

Add an entry for your case and define:

- "try_cflags": flags for C compiler.
- "c_ldflags": flags for linking, when using the C compiler as linker. This is needed to check for libraries written in C, such as FFTW.

- if you need a different preprocessor from the standard one (\$CC -E), define it in "try_cpp".

For example for XLC on AIX:

```
aix:mpcc* | aix:xlcc* | aix:cc )
    try_cflags="-q64 -O2"
    c_ldflags="-q64"
    ;;
```

Finally, if you have to use a nonstandard preprocessor, look for these lines:

```
echo $ECHO_N "setting CPPFLAGS... $ECHO_C"
case $cpp in
    cpp) try_cppflags="-P -traditional" ;;
    fpp) try_cppflags="-P" ;;
    ...
```

and set "try_cppflags" as appropriate.

4. Search for libraries

To instruct "configure" to search for libraries, you must tell it two things: the names of libraries it should search for, and where it should search.

The following libraries are searched for:

- BLAS or equivalent. Some vendor replacements for BLAS that are supported by Quantum ESPRESSO are:

```
MKL on Linux, 32- and 64-bit Intel CPUs
ACML on Linux, 64-bit AMD CPUs
essl on AIX
complib.sgimath on sgi origin
SCSL on sgi altix
SUNperf on sparc
cxml on alpha
```

Moreover, ATLAS is used over BLAS if available.

- LAPACK or equivalent. Some vendor replacements for LAPACK that are supported by Quantum ESPRESSO are:

```
mkl on linux SUNperf on sparc
```

- FFTW (version 3) or another supported FFT library. The latter include:

```
essl on aix ACML on Linux, 64-bit AMD CPUs SUNperf on sparc
```

- the MASS vector math library on aix
- an MPI library. This is often automatically linked by the compiler

If you have another replacement for the above libraries, you'll have to insert a new entry in the appropriate place.

This is unfortunately a little bit too complex to explain. Basic info: "AC.SEARCH.LIBS(function, name, ...)" looks for symbol "function" in library "libname". If that is found, "-lname" is appended to the LIBS environment variable (initially empty). The real thing is more complicated than just that because the "-Ldirectory" option must be added to search in a nonstandard directory, and because a given library may require other libraries as prerequisites (for example, Lapack requires BLAS).

2.3 Adding new directories or routines

3 Algorithms

3.1 Diagonalization

3.2 Self-consistency

3.3 Structural optimization

3.4 Symmetrization

3.5 Gamma tricks

In calculations using only the Γ point ($k=0$), the Kohn-Sham orbitals can be chosen to be real functions in real space, so that $\psi(G) = \psi^*(-G)$. This allows us to store only half of the Fourier components. Moreover, two real FFTs can be performed as a single complex FFT. The auxiliary complex function Φ is introduced: $\Phi(r) = \psi_j(r) + i\psi_{j+1}(r)$ whose Fourier transform $\Phi(G)$ yields

$$\psi_j(G) = \frac{\Phi(G) + \Phi^*(G)}{2}, \psi_{j+1}(G) = \frac{\Phi(G) - \Phi^*(G)}{2i}.$$

A side effect on parallelization is that G and $-G$ must reside on the same processor. As a consequence, pairs of columns with $G_{n'_1, n'_2, n'_3}$ and $G_{-n'_1, -n'_2, n'_3}$ (with the exception of the case $n'_1 = n'_2 = 0$), must be assigned to the same processor.

4 Structure of the code

4.1 Modules and global variables

4.2 Meaning of the most important variables

4.3 Conventions for indices

4.4 Preprocessing

The code contains C-style preprocessing directives. Most fortran compilers directly support them; some don't, and preprocessing is "hand-made" by the makefile using the C preprocessor `cpp`. The C preprocessor may:

- assign a value to a given expression. For instance, command `#define THIS that`, or the option in the command line: `-DTHIS=that`, will replace all occurrence of `THIS` with `that`.
- include file (command `#include`)
- expand macros (command `#define`)
- execute conditional expressions such as

```
#ifdef __expression
...code A...
#else
...code B...
#endif
```

If "expression" is defined (with a `#define` command or from the command line with option `D__expression`), then `...code A...` is sent to output; otherwise `...code B...` is sent to output.

The file `include/defs.h.README` contains a list of definitions that are used in the code. In order to make preprocessing options easy to see, preprocessing variables should start with two underscores, as `__expression` in the above example. Traditionally "preprocessed" variables are also written in uppercase.

4.5 Performance issues

4.6 Portability issues

5 Parallelization

In parallel execution, PW starts N independent processes (do not start more than one per processor!) that communicate via calls to MPI libraries. Each process has its own set of variables and knows nothing about other processes' variables. Variables that take little memory are replicated, those that take a lot of memory (wavefunctions, G-vectors, R-space grid) are distributed.

Beware: replicated calculations may either be performed independently on each processor, or performed on one processor and broadcast to all others. The first approach requires less programming, but it is unsafe: in principle all processors should yield exactly the same results, if they work on the same data, but sometimes they don't (depending on the machine, compiler, and libraries). Even a tiny difference in the last significant digit can eventually cause serious trouble if allowed to build up, especially when a replicated check is performed (in which case the code may "hang" if the check yields different results on different processors). Never assume that the value of a variable produced by replicated calculations is exactly the same on all processors: when in doubt, broadcast the value calculated on a specific processor (the "root" processor) to all others.

5.1 Paradigms

5.2 Implementation

5.2.1 Data distribution

Quantum ESPRESSO employ arrays whose memory requirements fall into three categories.

- *Fully Scalable*: Arrays that are distributed across processors of a pool. Fully scalable arrays are typically large to very large and contain one of the following dimensions:
 - number of plane waves, npw (or max number, npwx)
 - number of Gvectors, ngm
 - number of grid points in the R space, nrxx

Their size decreases linearly with the number of processors in a pool.

- *Partially Scalable*: Arrays that are distributed across processors of the ortho or diag group. Typically they are much smaller than fully scalable array, and small in absolute terms for moderate-size system. Their size however increases quadratically with the number of atoms in the system, so they have to be distributed for large systems (hundreds to thousands atoms). Partially scalable arrays contain none of the dimensions listed above, two of the following dimensions:
 - number of states, nbnd
 - number of projectors, nkb

Their size decreases linearly with the number of processors in a ortho or diag group.

- *Nonscalable*: All the remaining arrays, that are not distributed across processors. These are typically small arrays, having dimensions like for instance:
 - number of atoms, nat
 - number of species of atoms, nsp

The size of these arrays is independent on the number of processors.

5.2.2 Parallel fft

6 File Formats

6.1 Data file(s)

Quantum ESPRESSO restart file specifications: Paolo Giannozzi scripsit AD 2005-11-11, Last modified by Andrea Ferretti 2006-10-29

6.1.1 Rationale

Requirements: the data file should be

- efficient (quick to read and write)
- easy to read, parse and write without special libraries
- easy to understand (self-documented)
- portable across different software packages
- portable across different computer architectures

Solutions:

- use binary I/O for large records
- exploit the file system for organizing data
- use XML
- use a small specialized library (iotk) to read, parse, write
- ensure the possibility to convert to a portable formatted file

Integration with other packages:

- provide a self-standing (code-independent) library to read/write this format
- the use of this library is intended to be at high level, hiding low-level details

6.1.2 General structure

Format name: `''QEXML''`

Format version: `''1.4.0''`

The "restart file" is actually a "restart directory", containing several files and sub-directories. For CP/FPMD, the restart directory is created as `''$prefix.$ndw/''`, where `$prefix` is the value of the variable "prefix". `$ndw` the value of variable `ndw`, both read in input; it is read from `''$prefix.$ndr/''`, where `$ndr` the value of variable `ndr`, read from input. For PWscf, both input and output directories are called `''$prefix.save/''`.

The content of the restart directory is as follows:

`''data-file.xml''`

which contains:

- general information that doesn't require large data
- atomic structure, lattice, symmetries, parameters of
- pointers to other files or directories containing bu
- such as grids, wavefunctions, charge density, potent

`''charge_density.dat''`

contains the charge density

`''spin_polarization.dat''`

contains the spin polarization (rhoup-rhodw) (LSDA cal

`''magnetization.x.dat''`

<code>''magnetization.y.dat''</code>	contain the spin polarization along x,y,z (noncollinear)
<code>''magnetization.z.dat''</code>	
<code>''lambda.dat''</code>	contains occupations (Car-Parrinello dynamics only)
<code>''mat_z.1''</code>	contains occupations (ensemble-dynamics only)

`<pseudopotentials>` A copy of all pseudopotential files given in input

`<k-point dirs>` One or more subdirectories `''K00001/''`, `''K00002/''`, etc,

Each k-point directory contains:

<code>''evc.dat''</code>	containing the wavefunctions for spin-unpolarized calculations
<code>''evc1.dat''</code>	
<code>''evc2.dat''</code>	containing the spin-up and spin-down wavefunctions, respectively for spin polarized (LSDA) calculations;
	in a molecular dynamics run, also wavefunctions at the previous step
<code>''evcm.dat''</code>	for spin-unpolarized calculations OR
<code>''evcm1.dat''</code>	
<code>''evcm2.dat''</code>	for spin polarized calculations;
<code>''gkectors.dat''</code>	with the details of specific k+G grid;
<code>''eigenval.xml''</code>	containing the eigenvalues for the corresponding k-points
<code>''eigenval1.xml''</code>	
<code>''eigenval2.xml''</code>	for spin-polarized calculations;

- All files `''*.xml''` are XML-compliant, formatted file;
- Files `''mat_z.1''`, `''lambda.dat''` are unformatted files, containing a single record;
- All other files `''*.dat''`, are XML-compliant files, but they contain an unformatted record.

6.1.3 Structure of file `''data-file.xml''`

* `''XML Header''`: whatever is needed to have a well-formed XML file

* `''Body''`: introduced by `<Root>`, terminated by `</Root>`. Contains first-level tags on

* `''First-level tags''`: contain either

** second-level tags

** "data tags": tags containing data (values for a given variable)

** "file tags": tags pointing to a file

`''data tags syntax''` ([...] = optional) :

`<TAG type="vartype" size="n" [UNIT="units"] [LEN="k"]>`

values (in appropriate units) for variable corresponding to TAG:

n elements of type vartype (if character, of length k)

`</TAG>`

where TAG describes the variable into which data must be read;

"vartype" may be "integer", "real", "character", "logical";

 if type="logical", LEN=k" must be used to specify the length
 of the variable character; size="n" is the dimension.

 Acceptable values for "units" depend on the specific tag.

''Short syntax'', used only in a few cases:

```
<TAG attribute="something"/> .
```

For instance:

```
<FFT_GRID nr1="NR1" nr2="NR2" nr3="NR3"/>
```

defines the value of the FFT grid parameters nr1, nr2, nr3
 for the charge density

6.1.4 Sample

* Header:

```
<?xml version="1.0"?>
<?iotk version="1.0.0test"?>
<?iotk file_version="1.0"?>
<?iotk binary="F"?>
```

These are meant to be used only by iotk (actually they aren't)

* First-level tags:

- <HEADER> (global information about fmt version)
- <CONTROL> (miscellanea of internal information)
- <STATUS> (information about the status of the CP simulation)
- <CELL> (lattice vector, unit cell, etc)
- <IONS> (type and positions of atoms in the unit cell etc)
- <SYMMETRIES> (symmetry operations)
- <ELECTRIC_FIELD> (details for an eventual applied electric field)
- <PLANE_WAVES> (basis set, cutoffs etc)
- <SPIN> (info on spin polarizaztion)
- <MAGNETIZATION_INIT> (info about starting or constrained magnetization)
- <EXCHANGE_CORRELATION>
- <OCCUPATIONS> (occupancy of the states)
- <BRILLOUIN_ZONE> (k-points etc)
- <PHONON> (info for phonon calculations)
- <PARALLELISM> (specialized info for parallel runs)
- <CHARGE-DENSITY>
- <TIMESTEPS> (positions, velocities, nose' thermostats)
- <BAND_STRUCTURE_INFO> (dimensions and basic data about band structure)
- <EIGENVALUES> (eigenvalues and related data)
- <EIGENVECTORS> (eigenvectors and related data)

* Tag description

```
<HEADER>
  <FORMAT>      (name and version of the format)
  <CREATOR>      (name and version of the code generating the file)
</HEADER>

<CONTROL>
  <PP_CHECK_FLAG>  (whether the file can be used for post-processing)
  <LKPOINT_DIR>    (whether kpt-data are written in sub-directories)
  <Q_REAL_SPACE>   (whether augmentation terms are used in real space)
</CONTROL>

<STATUS> (optional)
  <STEP>    (number $n of steps performed, i.e. we are at step $n)
  <TIME>    (total simulation time)
  <TITLE>   (a job descriptor)
  <ekin>    (kinetic energy)
  <eht>     (hartree energy)
  <esr>     (Ewald term, real-space contribution)
  <eself>   (self-interaction of the Gaussians)
  <epseu>   (pseudopotential energy, local)
  <enl>     (pseudopotential energy, nonlocal)
  <exc>     (exchange-correlation energy)
  <vave>    (average of the potential)
  <enthal>  (enthalpy: E+PV)
</STATUS>

<CELL>
  <BRAVAIS_LATTICE>
  <LATTICE_PARAMETER>
  <CELL_DIMENSIONS> (cell parameters)
  <DIRECT_LATTICE_VECTORS>
    <UNITS_FOR_DIRECT_LATTICE_VECTORS>
    <a1>
    <a2>
    <a3>
  <RECIPROCAL_LATTICE_VECTORS>
    <UNITS_FOR_RECIPROCAL_LATTICE_VECTORS>
    <b1>
    <b2>
    <b3>
</CELL>

<IONS>
  <NUMBER_OF_ATOMS>
  <NUMBER_OF_SPECIES>
```



```

<UNITS_FOR_ATOMIC_MASSES>
For each $n-th species $X:
  <SPECIE.$n>
    <ATOM_TYPE>
    <MASS>
    <PSEUDO>
  </SPECIE.$n>
<PSEUDO_DIR>
<UNITS_FOR_ATOMIC_POSITIONS>
For each atom $n of species $X:
  <ATOM.$n SPECIES="$X">
</IONS>

<SYMMETRIES>
  <NUMBER_OF_SYMMETRIES>
  <INVERSION_SYMMETRY>
  <NUMBER_OF_ATOMS>
  <UNITS_FOR_SYMMETRIES>
For each symmetry $n:
  <SYMM.$n>
    <INFO>
    <ROTATION>
    <FRACTIONAL_TRANSLATION>
    <EQUIVALENT_IONS>
  </SYMM.$n>
</SYMMETRIES>

<ELECTRIC_FIELD> (optional)
  <HAS_ELECTRIC_FIELD>
  <HAS_DIPOLE_CORRECTION>
  <FIELD_DIRECTION>
  <MAXIMUM_POSITION>
  <INVERSE_REGION>
  <FIELD_AMPLITUDE>
</ELECTRIC_FIELD>

<PLANE_WAVES>
  <UNITS_FOR_CUTOFF>
  <WFC_CUTOFF>
  <RHO_CUTOFF>
  <MAX_NUMBER_OF_GK-VECTORS>
  <GAMMA_ONLY>
  <FFT_GRID>
  <GVECT_NUMBER>
  <SMOOTH_FFT_GRID>
  <SMOOTH_GVECT_NUMBER>
  <G-VECTORS_FILE>      link to file "gvectors.dat"

```

```

    <SMALLBOX_FFT_GRID>
</PLANE_WAVES>

<SPIN>
    <LSDA>
    <NON-COLINEAR_CALCULATION>
    <SPIN-ORBIT_CALCULATION>
    <SPIN-ORBIT_DOMAG>
</SPIN>

<EXCHANGE_CORRELATION>
    <DFT>
    <LDA_PLUS_U_CALCULATION>
    if LDA_PLUS_U_CALCULATION
        <NUMBER_OF_SPECIES>
        <HUBBARD_LMAX>
        <HUBBARD_L>
        <HUBBARD_U>
        <HUBBARD_ALPHA>
    endif
</EXCHANGE_CORRELATION>

if hybrid functional
    <EXACT_EXCHANGE>
        <x_gamma_extrapolation>
        <nqx1>
        <nqx2>
        <nqx3>
        <exxdiv_treatment>
        <yukawa>
        <ecutvcut>
        <exx_fraction>
        <screening_parameter>
    </EXACT_EXCHANGE>
endif

<OCCUPATIONS>
    <SMEARING_METHOD>
    if gaussian smearing
        <SMEARING_TYPE>
        <SMEARING_PARAMETER>
    endif
    <TETRAHEDRON_METHOD>
    if use tetrahedra
        <NUMBER_OF_TETRAHEDRA>
        for each tetrahedron $t
            <TETRAHEDRON.$t>

```

```

endif
<FIXED_OCCUPATIONS>
if using fixed occupations
  <INFO>
  <INPUT_OCC_UP>
  if lsda
    <INPUT_OCC_DOWN>
  endif
endif
</OCCUPATIONS>

<BRILLOUIN_ZONE>
  <NUMBER_OF_K-POINTS>
  <UNITS_FOR_K-POINTS>
  <MONKHORST_PACK_GRID>
  <MONKHORST_PACK_OFFSET>
  For each k-point $n:
    <K-POINT.$n>
  </BRILLOUIN_ZONE>

<PHONON>
  <NUMBER_OF_MODES>
  <UNITS_FOR_Q-POINT>
  <Q-POINT>
</PHONON>

<PARALLELISM>
  <GRANULARITY_OF_K-POINTS_DISTRIBUTION>
</PARALLELISM>

<CHARGE-DENSITY>
  link to file "charge_density.rho"
</CHARGE-DENSITY>

<TIMESTEPS> (optional)
  For each time step $n=0,M
    <STEP$n>
      <ACCUMULATORS>
      <IONS_POSITIONS>
        <stau>
        <svel>
        <taui>
        <cdmi>
        <force>
      <IONS_NOSE>
        <nhpcl>
        <nhpdim>

```

```

        <xnhp>
        <vnhp>
    <ekincm>
    <ELECTRONS_NOSE>
        <xnhe>
        <vnhe>
    <CELL_PARAMETERS>
        <ht>
        <htve>
        <gvel>
    <CELL_NOSE>
        <xnhh>
        <vnhh>
    </CELL_NOSE>
</TIMESTEPS>

<BAND_STRUCTURE_INFO>
    <NUMBER_OF_BANDS>
    <NUMBER_OF_K-POINTS>
    <NUMBER_OF_SPIN_COMPONENTS>
    <NON-COLINEAR_CALCULATION>
    <NUMBER_OF_ATOMIC_WFC>
    <NUMBER_OF_ELECTRONS>
    <UNITS_FOR_K-POINTS>
    <UNITS_FOR_ENERGIES>
    <FERMI_ENERGY>
</BAND_STRUCTURE_INFO>

<EIGENVALUES>
    For all kpoint $n:
        <K-POINT.$n>
            <K-POINT_COORDS>
            <WEIGHT>
            <DATAFILE>
            </K-POINT.$n>
</EIGENVALUES>

<EIGENVECTORS>
    <MAX_NUMBER_OF_GK-VECTORS>
    For all kpoint $n:
        <K-POINT.$n>
            <NUMBER_OF_GK-VECTORS>
            <GK-VECTORS>
            for all spin $s
                <WFC.$s>
                <WFCM.$s>

```

link to file `"/K$n/eigenval.xml"`

link to file `"/K$n/gkvecvectors.dat"`

link to file `"/K$n/evc.dat"`

link to file `"/K$n/evcm.dat"` (optional)
containing wavefunctions at preceding step

```
</K-POINT.$n>
</EIGENVECTORS>
```

6.2 Restart files

7 Modifying/adding/extending Quantum ESPRESSO

7.1 Hints, Caveats, Do's and Dont's

- Before doing anything, inquire whether it is already there, or under development.
- Before starting writing code, inquire whether you can reuse code that is already available in the distribution. Avoid redundancy: the only bug-free software line is the one that doesn't exist.
- When you make some change:
 - Check that are not spoiling other people's work. In particular, search the distribution for codes using the routine or module you are modifying and change its usage or its calling sequence everywhere.
 - Do not forget to add/update documentation and examples as well.
 - Do not forget that your change must work on many different combinations of hardware and software, in both serial and parallel execution.
- Please do not include files with DOS \hat{M} characters or tabulators \hat{I} .

Important: when you modify the program sources, run the `install/makedeps.sh` script or type `make depend` to update files `make.depend` in the various subdirectories.

7.2 Programming style (or lack of it)

Guidelines for developers:

- preprocessing options should be capitalized and start with two underscores. Examples: `__AIX`, `__LINUX`, ...
- fortran commands should be capitalized: `CALL something()`
- variable names should be lowercase: `foo = bar/2`
- indent DO's and IF's with three white spaces (editors like emacs will do this automatically for you)
- do not write crammed code: leave spaces, insert empty separation lines
- comments (introduced by a `!`) should be used to explain what is not obvious from the code, not to repeat what is already evident. Obscure comments serve no purpose.

- do not use machine-dependent extensions or sloppy syntax. Standard f90 requires that a & is needed both at end of line AND at the beginning of continuation line if there is a ' or " " spanning two lines. Some compilers do not complain if the latter & is missing, others do.
- use DP (defined in module "kinds") to define the type of real and complex variables
- all constants should be defined to be of kind DP. Preferred syntax: 0.0_dp.
- use "generic" intrinsic functions: SIN, COS, etc.
- conversions should be explicitly indicated. For conversions to real, use DBLE, or else REAL(...,KIND=dp). For conversions to complex, use CMPLX(...,KIND=dp). For complex conjugate, use CONJG. For imaginary part, use AIMAG. IMPORTANT: Do not use REAL or CMPLX without KIND=dp, or else you will lose precision (except when you take the real part of a double precision complex number).

7.3 Adding or modifying input variables

New input variables should be added to "Modules/input_parameters.f90", then copied to the code internal variables in the "input.f90" subroutine. The namelists and cards parsers are in : "Modules/read_namelists.f90" and "Modules/read_cards.f90". Files "input_parameters.f90", "read_namelists.f90", "read_cards.f90" are shared by all codes, while each code has its own version of "input.f90" used to copy input values into internal variables

EXAMPLE: suppose you need to add a new input variable called "pippo" to the namelist control, then:

1. add pippo to the input_parameters.f90 file containing the namelist control

```
INTEGER :: pippo = 0
NAMELIST / control / ....., pippo
```

Remember: always set an initial value!

2. add pippo to the control_default subroutine (contained in module read_namelists.f90)

```
subroutine control_default( prog )
...
IF( prog == 'PW' ) pippo = 10
...
end subroutine
```

This routine sets the default value for pippo (can be different in different codes)

3. add pippo to the control_bcast subroutine (contained in module read_namelists.f90)

```
subroutine control_bcast( )
...
call mp_bcast( pippo )
...
end subroutine
```

8 Using CVS

The package is available read-only using anonymous CVS. Developers may have read-write access if needed. Note that the latest (development) version may not work properly, and sometimes not even compile properly. Use at your own risk.

CVS (Concurrent Version System) is a software that allows many developers to work and maintain a single copy of a software in a central location (repository). It is installed by default on most Unix machines, or otherwise it can be very easily installed: see <http://www.cvshome.org>. For a tutorial, see: http://www.loria.fr/molli/cvs/cvs-tut/cvs_tutorial_toc.html. You will also need a working installation of ssh (secure shell) to use CVS.

8.1 Anonymous CVS

You have to define the environment variables CVS_RSH and CVSROOT. For csh/tcsh, use

```
setenv CVS_RSH ssh
setenv CVSROOT :pserver:anonymous@scm.qe-forge.org:/cvsroot/q-e
```

For sh/ksh/bash, use

```
export CVS_RSH=ssh
export CVSROOT=:pserver:anonymous@scm.qe-forge.org:/cvsroot/q-e
```

Then:

```
cvs login
```

Do not specify any password, just press "Enter".

8.2 Read/Write CVS

The environment variable CVS_RSH is defined as above, but CVSROOT is set to a different value. For csh/tcsh, use

```
setenv CVS_RSH ssh
setenv CVSROOT :ext:your-account@scm.qe-forge.org:/cvsroot/q-e
```

For sh/ksh/bash use

```
export CVS_RSH=ssh
export CVSROOT=:ext:your-account@scm.qe-forge.org:/cvsroot/q-e
```

You need to have an account on <http://www.qe-forge.org> ("your-account" above) with privileges as Quantum ESPRESSO developer. You will be prompted for your password at each cvs operation.

8.3 CVS operations

For the first code download:

```
cvcs checkout -P espresso
```

The code appears in directory "espresso/". To update the code to the current version:

```
cvcs update -dP
```

in the directory containing the distribution. Option "-d" ensures that newly added directories are downloaded, "-P" that empty directories are removed. It is possible to download the version at a given date, or corresponding to a given "tag" (set by the developers, usually just before extensive changes or at public releases).

When updating, you should get lines looking like

```
cvcs server: Updating Modules
P Modules/Makefile
U Modules/control_flags.f90
```

where P means "patched", U means "updated" (a new file is added); or

```
M PW/Makefile
```

where M means "locally modified" (i.e., wrt the CVS repository version); or

```
cvcs server: FPMD/control.f90 is no longer in the repository
```

if a file has been meanwhile deleted, or moved, or renamed; but no lines like

```
C Somedir/SomeFile
cvcs server: conflict while updating Somedir/SomeFile
```

This means that somebody else has modified the same parts of the code that you have locally modified; conflicting files will contain lines like

```
>>>>>>>>>>>
    something
=====
    something else
<<<<<<<<<<<<
```

If this happen, you must edit the file manually to remove conflicts.

You can compare your local copy with any version of the repository using

```
cvcs diff -r tag
```

or with the repository at a given date using

```
cvcs diff -D date
```


You can also compare two different versions/snapshots of the repository, by specifying two different tags or dates. Options "-i" (ignore case), "-w" (ignore all white spaces), "-b" (ignore changes in the number of white spaces) may be useful to distinguish true changes from esthetic ones (such as changes in indentation).

READ-WRITE ACCESS ONLY:

In order to save your changes to the repository, use "cvs commit". In order to add a file, first use "cvs add", then "cvs commit". In order to delete a file, first use "cvs delete", then "cvs commit". In order to rename a file, delete the file with the old name, add the file with the new name.

In order to add a new directory (let us say "dir/", and if you have the permission to do so:

- create directory "dir/"; do "cvs add dir" (this will create the CVS subdirectory in the new directory "dir/")
- copy all files into "dir/", then from inside "dir/" add files: "cvs add *.f90 Makefile" (for instance)
- "cvs commit" will save the new directory

8.4 CVS hints

It is convenient to create a `.cvsrc` file in your home directory containing the following lines:

```
cvs -z3
diff -wib
update -dP
checkout -P
```

These options will be automatically used.

You can use the same repository read-only with anonymous cvs, read-write with your qe-forge account. Download your repository using anonymous cvs as explained above; then re-define the `$CVSROOT` environment variable to point to the appropriate value for read-write access. You will keep using anonymous cvs because cvs will look for the contents of file `CVS/Root`, but if you use syntax:

```
cvs -d$CVSROOT commit ...
```

you can perform write operations (you will be prompted for your qe-forge password). In this way you will need to type your password only once in a while.

You will find very convenient to use the Web-CVS interface: <http://qe-forge.org/cgi-bin/cvstrac/qe/index>, in particular the "Timeline" and "Browse" options, useful to follow what has been done and what is going on in the development version.