



THERMO_PW User's Guide (v.2.1.1)

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Introduction

This guide covers the installation and usage of the THERMO_PW package. It assumes some familiarity with QUANTUM ESPRESSO. For this please consult the web site: <http://www.quantum-espresso.org>.

THERMO_PW computes material properties. At low level, it calls QUANTUM ESPRESSO routines and, at high level, it has pre-processing tools to reduce the information provided by the user and post-processing tools that use the output of QUANTUM ESPRESSO to produce plots of material properties directly comparable with experiment.

THERMO_PW has the following directory structure, contained in a directory `thermo_pw/` that should be put in the root directory of QUANTUM ESPRESSO:

<code>Doc/</code>	: contains this user's guide and other documentation
<code>examples/</code>	: some examples
<code>examples_qe/</code>	: QUANTUM ESPRESSO examples run using THERMO_PW
<code>inputs/</code>	: a collection of useful inputs
<code>pseudo_test/</code>	: a collection of inputs to test a pseudopotential library
<code>space_groups/</code>	: a collection of structures for many space groups
<code>lib/</code>	: source files for modules used by THERMO_PW
<code>qe/</code>	: routines of QUANTUM ESPRESSO that require some change
<code>fft</code>	: some fft routines that can run on the GPU.
<code>lapack</code>	: some lapack routines that can run on the GPU.
<code>src/</code>	: source files for THERMO_PW
<code>tools/</code>	: source files for auxiliary tools
<code>tools_input/</code>	: examples of inputs for the auxiliary tools

The THERMO_PW package can calculate the following quantities:

- Plot of the Brillouin zone (the structure can be seen by reading the input of THERMO_PW by the XCrySDen program).
- Plot of the X-rays powder diffraction pattern of the input crystal.
- Total energy at fixed geometry.
- Total energy as a function of the kinetic energy cut-off.

- Total energy as a function of $\mathbf{k}$ -points and smearing.
- Electronic band structure at fixed geometry.
- Electronic density of states at fixed geometry. Electronic thermodynamic properties: energy, free energy, entropy, and heat capacity.
- Electronic heat capacity as a function of temperature (for metals only).
- Complex dielectric constant as a function of the complex frequency ω at fixed geometry. Complex index of refraction for all systems except monoclinic and triclinic. Reflectivity at normal incidence and adsorption coefficient for cubic solids.
- Inverse dielectric constant at a given wavevector $\mathbf{q}$ as a function of the complex frequency ω at fixed geometry.
- Phonon frequencies at fixed geometry.
- Phonon dispersions at fixed geometry and harmonic thermodynamic properties: vibrational energy, vibrational free energy, vibrational entropy, and constant volume heat capacity as a function of temperature. Atomic Debye-Waller factors as a function of temperature.
- Frozen ions and relaxed ions elastic constants at fixed geometry.
- Relaxed ions temperature dependent elastic constants at fixed unperturbed geometry.
- Fit of the total energy as a function of the lattice parameters with a quadratic or quartic polynomial and determination of equilibrium lattice parameters. Murnaghan or (third or fourth order) Birch-Murnaghan fit. Enthalpy as a function of pressure. Crystal parameters and volume as a function of pressure.
- Electronic band structure at the minimum of the total energy.
- Electronic density of states at the minimum of the total energy. Electronic thermodynamic properties.
- Complex dielectric constant as a function of the complex frequency ω at the minimum of the total energy. Complex index of refraction for all systems except monoclinic and triclinic. Reflectivity at normal incidence and adsorption coefficient for cubic solids.
- Inverse dielectric constant at a given wavevector $\mathbf{q}$ as a function of the complex frequency ω at the minimum of the total energy.
- Phonon frequencies at the minimum of the total energy.
- Phonon dispersions and harmonic thermodynamic quantities at the minimum of the total energy.

- Frozen ions and relaxed ions elastic constants at the minimum of the total energy.
- Anharmonic properties within the quasi-harmonic approximation: lattice parameters, thermal expansion tensor, volume, volume thermal expansion, and constant strain heat capacity as a function of temperature; phonon frequencies and mode Grüneisen parameters interpolated at a given geometry or at the equilibrium geometry at a given temperature (limited to cubic, tetragonal, orthorhombic, and hexagonal systems). Bulk modulus and pressure derivative of the bulk modulus, isobaric heat capacity, isoentropic bulk modulus, and average Grüneisen parameter as a function of temperature (limited to cubic systems). Minimum Helmholtz (or Gibbs at finite pressures) free energy as a function of temperature.
- Isothermal and isoentropic elastic constants and elastic compliances as a function of temperature within the “quasi-static” approximation.
- Isothermal and isoentropic elastic constants and elastic compliances as a function of temperature within the “quasi-harmonic” approximation.
- Surface band structure identification and plot of the projected bulk band structure.

THERMO_PW can run on both serial and parallel machines using all the parallelization options of QUANTUM ESPRESSO. Moreover, THERMO_PW can run using several images. When possible, the image parallelization is used in an asynchronous way. One image takes the role of master and distributes the work to all the images that carry it out independently. Presently the total energies of several geometries for the determination of the equilibrium geometry are calculated in parallel when there are several images. Stresses or total energies at different strained geometries needed for the calculation of the elastic constants are calculated in parallel. The phonon calculations are carried out in parallel, each image doing one irreducible representation of one $\mathbf{q}$ point. For frequency dependent calculation, each frequency, or group of frequencies, can be calculated in parallel by different images. The phonon dispersions of several geometries needed for the quasi-harmonic calculation of the thermodynamic properties or of the elastic constants can be calculated in parallel (one geometry at a time or all geometries together).

1.1 People

The THERMO_PW code is primarily designed, written, and maintained by Andrea Dal Corso (SISSA - Trieste).

Some routines have been contributed by SISSA PhD students and post-docs. Among them I mention M. Palumbo, O. Motornyi, A. Urru, C. Malica, X. Gong, B. Thakur, and A. Ahmed.

I would like also to thank all the people that contributed with comments, requests of improvements, and bug reports.

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Installing, Compiling, and Running

2.1 Installing

The THERMO_PW package is tightly bound to QUANTUM ESPRESSO. It cannot be compiled without it. To download and compile QUANTUM ESPRESSO, please refer to the general User's Guide, available in the file `Doc/user_guide.pdf` in the root QUANTUM ESPRESSO directory, or on the web site

<http://www.quantum-espresso.org>.

The main distribution page of Thermo_pw is

https://dalcorso.github.io/thermo_pw/

where you can download one of the `.tar.gz` files. Please match carefully THERMO_PW and QUANTUM ESPRESSO versions as illustrated in Table 1. For the versions of QUANTUM ESPRESSO not listed here, there is no THERMO_PW package. The source files of THERMO_PW can be obtained by unpacking the `.tar.gz` file in the root QUANTUM ESPRESSO directory, for instance with the command

```
tar -xzf thermo_pw.2.1.1.tar.gz
```

You can also download the `git` version of THERMO_PW as described at the web page: https://dalcorso.github.io/thermo_pw/thermo_pw_help.html. The `git` version of THERMO_PW contains the most recent features and bug fixes but it might work only with the `git` version of QUANTUM ESPRESSO and its use for production is not recommended.

Please read the web page:

http://dalcorso.github.io/thermo_pw/thermo_pw_help.html

for updated information about the compatibility between the `git` version of THERMO_PW and QUANTUM ESPRESSO. This web page contains also information on critical bugs found in the THERMO_PW package and should be consulted before using THERMO_PW.

THERMO_PW	QUANTUM ESPRESSO	release date
2.1.1	7.5	13/10/2025
2.1.0	7.4.1	24/09/2025
2.0.3	7.4.1	23/06/2025
2.0.2	7.4	20/06/2025
2.0.1	7.3.1	15/04/2025
2.0.0	7.3	24/09/2024
1.9.1	7.3	14/02/2024
1.9.0	7.2	26/01/2024
1.8.1	7.2	05/05/2023
1.8.0	7.1	26/04/2023
1.7.1	7.1	05/07/2022
1.7.0	7.0	05/07/2022
1.6.1	7.0	10/01/2022
1.6.0	6.8	27/12/2021
1.5.1	6.8	22/07/2021
1.5.0	6.7	19/07/2021
1.4.1	6.7	29/12/2020
1.4.0	6.6	22/12/2020
1.3.2	6.6	17/8/2020
1.3.1	6.6	13/8/2020
1.3.0	6.5	12/8/2020
1.2.1	6.5	23/1/2020
1.2.0	6.4.1	28/12/2019
1.1.1	6.4.1	16/04/2019
1.1.0	6.4	16/04/2019
1.0.9	6.3	6/3/2019
1.0.0	6.3	17/7/2018
0.9.9	6.2.1	5/7/2018
0.9.0	6.2.1	20/12/2017
0.8.0	6.2	24/10/2017
0.8.0-beta	6.2-beta	31/8/2017
0.7.9	6.1	06/7/2017
0.7.0	6.1	18/3/2017
0.6.0	6.0.0	5/10/2016
0.5.0	5.4.0	26/4/2016
0.4.0	5.3.0	23/1/2016
0.3.0	5.2.1, 5.2.0	23/6/2015
0.2.0	5.1.2	13/3/2015
0.1.0	5.1.1	28/11/2014

Table 2.1: Compatibility between the versions of THERMO_PW and of QE.

2.2 Compiling

In order to compile THERMO_PW, the main Makefile and the files `install/makedeps.sh` and `install/plugins_makefile` of QUANTUM ESPRESSO must be changed. This is done by giving the command `make join_qe` inside the `thermo_pw` directory. This command exchanges the `thermo_pw` files with those of the QUANTUM ESPRESSO package.

Typing `make thermo_pw` inside the main QUANTUM ESPRESSO directory, or `make` inside the `thermo_pw` directory, produces the executable `thermo_pw/src/thermo_pw.x` that appears in the QUANTUM ESPRESSO `bin/` directory. A few other tool codes are produced as well and linked in the QUANTUM ESPRESSO `bin/` directory.

With some versions of QE the sequence of commands which is known to compile `thermo_pw` is the following:

```
Untar qe (with the version you want to run)
tar -xzf qe.tgz
cd main qe directory
tar -xzf thermo_pw.tgz
cd thermo_pw
make join_qe
cd ..
./configure
make thermo_pw
```

Running `./configure` before `make join_qe` might not work properly. To run `thermo_pw` with GPU you need a `pgi fortran` compiler (for instance the one contained in the `Nvidia hpc sdk`) and should run `configure` with a command similar to:

```
./configure --with-cuda=$CUDA_ROOT --with-cuda-runtime=11.8
--with-cuda-cc=80 --with-scalapack=no --with-cuda-mpi=yes
```

Please check on your system which is the correct version of `cuda-runtime` and of `cuda-cc`. The numbers given here are indicative.

Starting from version 1.5.1 `thermo_pw` can be compiled also with `cmake`. The command `make join_qe` substitutes the `CMakeLists.txt` file of QE with the one contained in `thermo_pw`. After writing `make join_qe` follow the QE instructions to compile using `cmake`. Two new commands are available: `make thermo_pw` produces `thermo_pw.x` and `make tpw_tools` produces the tools codes.

THERMO_PW has been written on a PC with the Linux operating system using a `gfortran` compiler and `openMPI` parallelization. It has been run in parallel on a Linux cluster with several hundreds processors. It has not been tested with other combinations of computer/operating system, but it is supposed to run on the same systems where QUANTUM ESPRESSO runs. If you have a machine in which you can compile and run QUANTUM ESPRESSO but not `thermo_pw`, please report the problem.

2.3 Searching help and reporting bugs

For problems installing `thermo_pw` or for help with some of its features, please subscribe to the `thermo_pw-forum` mailing list (https://lists.quantum-espresso.org/mailman/listinfo/thermo_pw-forum). Requests of new features are also welcome. If you think you have found a bug in `thermo_pw`, you can report it to the mailing list or write me: `dalcorso.at.sissa.it`.

2.4 Uninstalling

In order to remove THERMO_PW, give the command `make leave_qe` in the `thermo_pw` directory. Then just remove the directory. Note that the command `make leave_qe` is needed to restore the original QUANTUM ESPRESSO files.

2.5 Running THERMO_PW

In order to run THERMO_PW, you need an input for `pw.x`, a file called `thermo_control`, and an input for `ph.x` (which must be called `ph_control`) if required by the task. These files must be in your working directory. The input of `pw.x` can have any name and is given as input to the THERMO_PW code. It is better not to specify an `outdir` directory in the `ph.x` input. Specifying an `outdir` directory is not forbidden, but for some tasks `thermo_pw.x` might add a geometry number to `outdir` and the `outdir` written in the `ph.x` input must be consistent.

A typical command for running THERMO_PW is:

```
mpirun -n np thermo_pw.x -ni ni ... < input_pw > output_thermo_pw
```

where `np` is the number of processors and `ni` is the number of images. The dots indicate the other QUANTUM ESPRESSO parallelization options that you can find in its manual.

Note that it is very easy to waste resources using too many images. Unused images wait for the working images to complete their tasks wasting cpu-time in an endless loop. Some options do not use the image feature, so you have to know how the calculation is divided and the number of images must not be larger than the number of tasks (below I give this number for each option). If you have doubts on this point use one image (`ni=1`).

The outputs of the THERMO_PW code are one or more `postscript` or `pdf` files with plots of the material properties. THERMO_PW produces also files with the data of the plot and scripts for the `gnuplot` program. Usually, the user does not need to modify these files, but they allow the improvement of the figures when needed. The plot of the Brillouin zone (BZ) is made with the help of the `asymptote` code. `Thermo_pw` produces a script for the `asymptote` code and can also run it to produce the `pdf` file of the BZ.

Input variables

The `pw.x` and `ph.x` input files are described in the **QUANTUM ESPRESSO** documentation. In this section we discuss only the creation of the file `thermo_control`. This file contains a namelist:

```
&INPUT_THERMO
  what=' ',
  ...
/
```

The `what` variable controls the sequence of calculations made by `THERMO_PW`. For each possible value of `what`, we discuss briefly the input variables that control the output plots. Usually default values of the input variables are sufficient to carry out the basic `THERMO_PW` tasks and you are not supposed to set any variable except `what`, but in some cases these input variables give more control on the calculation and on its accuracy.

`THERMO_PW` writes on files the data to plot and a script to plot these data. The output `postscript` or `pdf` files are produced by invoking the `gnuplot` program. Usually any modern Linux distribution provides a package to install this code, or has it already installed. You can also download the package from <http://www.gnuplot.info/>.

The following input variables control the use of the `gnuplot` code:

```
lgnuplot      : if .TRUE. gnuplot is called by the
                program and the postscript or pdf files are
                immediately available. Otherwise give the
                command gnuplot gnuplot_files/*.
                Default: logical .TRUE.
gnuplot_command : the command used to call gnuplot.
                Default: character(len=*) 'gnuplot'
flgnuplot     : initial part of the name of the files where
                gnuplot scripts are written.
                Default: character(len=*) 'gnuplot.tmp'
flect        : extension of the output files. Presently .ps
                and .pdf are supported for postscript or pdf
```

output. The latter is available only if gnuplot supports the pdfcairo terminal.
Default: `character(len=*) '.ps'`

If your system has not gnuplot you can disable the production of the postscript or pdf files and use other graphical tools to plot the output data.

3.1 Temperature and pressure

Several quantities in THERMO_PW can be calculated as a function of temperature at zero pressure or at selected pressures. Moreover the equilibrium geometry can be searched at fixed pressure minimizing the enthalpy instead of the energy. In some cases it is also possible to plot some quantities as a function of pressure at zero temperature or at selected temperatures. There is therefore the necessity to specify uniform meshes of temperatures (pressures) or to choose selected temperatures (pressures). The meshes are specified giving the minimum and maximum temperatures (pressures) and the interval between temperature (pressure) points. Selected temperatures (pressures) are instead defined giving their number and their values. The code will choose these temperatures (pressures) on the uniform mesh taking the point closest to the specified temperature (or pressure). For the options where these features are active, the values of temperature and pressure are controlled by the following variables:

```
tmin      : minimum temperature for the mesh of temperatures.  
           Default: real 1 K  
tmax      : maximum temperature for the mesh of temperatures.  
           Default: real 800 K  
deltat    : interval between two temperatures. Be careful  
           with this value because it is used also to  
           compute temperature derivatives numerically.  
           Too small or too large values could give  
           inaccurate anharmonic properties.  
           Default: real 3 K  
ntemp     : number of temperatures.  
           Default: integer determined from previous data  
pressure   : The external pressure. The crystal parameters are  
           calculated minimizing the enthalpy at this  
           pressure. Given in kbar units.  
           Default: real 0.0 kbar  
pmin      : minimum pressure for the mesh of pressures.  
           Default: real -50.0 kbar  
pmax      : maximum pressure for the mesh of pressures.  
           Default: real 100.0 kbar  
deltap    : Interval between two pressures in the mesh of pressures.  
           Default: real 1.0 kbar  
ntemp_plot : number of temperatures in the plots where the  
           temperature is a parameter. When 0 these plots are  
           not produced.  
           Default: integer 0  
temp_plot(ntemp_plot) : A real array of dimension ntemp_plot with  
           the values of the temperature.  
           No Default value, must be given (in K) when  
           ntemp_plot is not zero.
```

`npres_plot` : number of pressures in the plots where the pressure is a parameter. When 0 these plots are not produced.
Default: integer 0

`press_plot(npres_plot)` : A real array of dimension `npres_plot` with the values of the pressure.
No Default value, must be given (in kbar) when `npres_plot` is not zero.

`nvol_plot` : number of volumes in the plots where the volume is a parameter. When 0 these plots are not produced.
Default: integer 0

`ivol_plot(nvol_plot)` : an integer array with the volumes to consider in the plots where the volume is a parameter. The chosen volumes can be only among one of the `ngeo(1)` geometries.
No Default value, must be given when `nvol_plot` is not zero.

Note that when you fix the external pressure, the geometries chosen to fit the enthalpy must be about the minimum geometry at that pressure. Similarly, when using pressure ranges, the number of simulated geometries must be large enough to cover the selected range of pressures, otherwise the plotted quantities might be inaccurate.

3.2 Coordinates and structure

The `thermo_pw` code requires the Bravais lattice of the solid. Moreover for computing some quantities it assumes that the direct lattice vectors are those provided by the routine `latgen.f90` of the `QUANTUM ESPRESSO` distribution. For this reason it is not recommended to use `ibrav=0` in the `pw.x` input. The preferred method is to give the value of `ibrav` and use the primitive vectors provided by `QUANTUM ESPRESSO`. It is also possible to specify the `space_group` number and the coordinates of the nonequivalent atoms. When the `pw.x` input contains the `ibrav=0` option, `thermo_pw` writes on output the values of `ibrav`, `celldm`, and of the atomic coordinates that should be used in the input of `pw.x` to simulate the same solid and stops. There are however two input variables of `thermo_pw` that can modify this behavior:

`continue_zero_ibrav` : when `ibrav=0` in the input of `pw.x` and this variable is set to `.TRUE.` `thermo_pw` runs with `ibrav=0` (not recommended except when you deal with a supercell). When this variable is `.FALSE.` and `ibrav=0` the behavior depends on `find_ibrav`.
Default: logical `.FALSE.`

`find_ibrav` : This variable is active only when `continue_zero_ibrav=.FALSE.` When this variable is set to `.TRUE.` and the input of `pw.x` has `ibrav=0`, `thermo_pw` finds the values of `ibrav`, `celldm`, and of the atomic positions that produce the same crystal and continue the calculation. The geometry used by `thermo_pw` might be rotated with respect to the input and have different primitive vectors. When this variable is `.FALSE.` the code stops after writing in output `ibrav`, `celldm`, and the atomic positions. These variables can be copied in the `pw.x` input. Note that the automatic identification of the lattice does not work for supercells.
Default: logical `.FALSE.`

3.3 **what='scf'**

With this option the code computes only the total energy. This is a single calculation as if running `pw.x` with the given input. No other input variable is necessary. An example for this option can be found in `example01`.

Number of tasks for this option: 1.

3.4 **what='scf_ke'**

With this option the code makes several self-consistent calculations, in parallel on several images, varying the kinetic energy cut-off for the wavefunctions and for the charge density. In the input of `pw.x` one specifies the minimum values for these two cut-offs. These values are then increased in fixed intervals controlled by the following variables. The energy is then plotted as a function of the wavefunctions kinetic energy cut-off, a different curve for each value of the charge density cut-off.

The variables that control this option are:

```
nke          : number of kinetic energies tested for the
                wavefunctions cut-off.
                Default: integer 5
deltake      : delta of wavefunctions kinetic energy cut-off
                in Ry (can be either positive or negative).
                Default: real 10 Ry
nkeden       : number of kinetic energies tested for the
                charge density cut-off.
                Default: integer 1
deltakeden   : delta of charge density kinetic energy
                cut-off in Ry (can be either positive or
                negative).
                Default: real 100 Ry.
flkeconv     : name of the file where the data with the
                total energy as a function of the kinetic
                energy is written.
                Default: character(len=*) 'output_keconv.dat'
flpskeconv   : name of the postscript file with the plot
                of the total energy as a function of the
                kinetic energy cut-off.
                Default: character(len=*) 'output_keconv'
```

An example for this option can be found in `example10`.

Number of tasks for this option: `nke * nkedens`.

3.5 **what='scf_nk'**

With this option the code makes several self-consistent calculations, in parallel on several images, varying the size of the **k**-point grid, and optionally for metals the smearing parameter `degauss`. In the input of `pw.x` the minimum value of these parameters is given and these values are increased in fixed intervals controlled by the following variables. On output the energy is plotted as a function of the mesh size, one curve for each smearing parameter.

The variables that control this option are:

```
nnk          : the number of different values of nk to test.
                Default: integer 5
deltank(3)    : the interval between nk values. All three values
                of nk1, nk2, and nk3 are updated simultaneously.
                Default: integer 2 2 2
nsigma        : the number of smearing intervals.
                Default: integer 1
deltasigma    : the distance between different smearing values
                (can be either positive or negative).
                Default: 0.005 Ry
flnkconv      : file where the data with the k point convergence
                is written.
                Default: character(len=*) 'output_nkconv.dat'
flpsnkconv    : name of the postscript file with the k points
                convergence plot.
                Default: character(len=*) 'output_nkconv'
```

An example for this option can be found in `example11`.

Number of tasks for this option: `nnk * nsigma`.

3.6 `what='scf_bands'`

With this option the code makes a self-consistent calculation followed by a band structure calculation. This option is not parallelized over images and should be used with one image. The output of the band structure calculation is further processed in order to produce a plot of the band structure. The zero of the energy is the highest valence band of the first **k** point in insulators and the Fermi energy in metals.

The energy bands plot can be modified by the following variables:

`emin_input` : minimum energy for the band dispersion plot (in eV).
Default: real minimum of the bands

`emax_input` : maximum energy for the band dispersion plot (in eV).
Default: real maximum of the bands

`nbnd_bands` : the number of bands in the band calculation.
Default: integer $2 \times \text{nbnd}$, where `nbnd` is the number of bands given in `pw.x` input or calculated by `pw.x`.

`only_bands_plot`: if the files with the bands and the representations are already on files, this option allows to change the parameters of the plot (such as the maximum or minimum energy) and do another plot without additional calculation. If the files are missing and this variable is `.TRUE.` an error occurs. Note that using this option you cannot change the path.
Default: logical `.FALSE.`

`lsym` : if `.TRUE.` does the symmetry analysis of the bands.
Default: `.TRUE.`

`enhance_plot`: if `.TRUE.` writes on the band plot the point group labels, and colors with different background colors lines at the zone border.
Default: `.FALSE.`

`long_path` : if `.TRUE.` plots the bands in all the Brillouin zone path. Otherwise makes a faster calculation on a short path. The short path is indicated also for two-dimensional layers perpendicular to the `z` direction.
Default: `.TRUE.`

`old_path` : if `.TRUE.` use an alternative path, usually more similar to the one used in experimental papers (available only for a few lattices).
Default: `.FALSE.`

`path_fact` : A factor that multiply the number of points along each line of the default path. Note that this is a real number so you can also decrease the default number of points along each line.
Default: real 1.0

`filband` : file where the bands are written in the QE format.
Default: `character(len=*) 'output_band.dat'`

`flpband` : file(s) where the bands are written in gnuplot format.
Default: `character(len=*) 'output_pband.dat'`

`flpsband` : postscript file with the electronic band structure.
Default: `character(len=*) 'output_band'`

Number of tasks for this option: 1.

By default, the bands are plotted along a fixed path in the Brillouin zone, but the user can modify this behavior giving the path at the end of the `INPUT_THERMO` namelist with the same format used for the `pw.x` input. The automatic path generation is not available for base-centered monoclinic and for triclinic Bravais lattices. For these lattices the path must be given explicitly. The following variables control the path:

`q_in_band_form` : only the first and last point of each k path are given. The weight of each k point is an integer, the number of points in the line that starts at this k point.
Default: logical `.TRUE.`

`q_in_cryst_coord` : the k - points are given in crystal coordinates. For centered lattices the crystal coordinates refer to the primitive cell (not the conventional one). Same convention as in QE.
Default: logical `.FALSE.`

`point_label_type` : the label definition (see the BZ manual).
Default: `SC`

`q2d` : the q points define a rectangle in reciprocal space. See the QE guide for more details.
Default: logical `.FALSE.`

`is_a_path` : if `.TRUE.` the q points are in a path in reciprocal space. This is usually the case except when `q2d=.TRUE.` or when the input points are in an arbitrary order. Set this to `.FALSE.` only if you want to skip the plot of the bands.
Default: logical `.TRUE.`

Note that the path is not given in the input of `pw.x` that contain instead the information to generate the mesh of k points for the self-consistent calculation. An example for this option can be found in `example02`. If you give explicitly the path, be careful with options that require geometry changes (see below). Only automatic paths, or path given through letter labels are easily recalculated. The other paths could turn out to be correct only for one geometry.

It is also possible to separate the self-consistent and the band calculation, by running first `thermo_pw.x` using `what='scf'` and then running, on the same directory, `thermo_pw.x` using `what='scf_bands'`. The same input can be used in the two calculations, only the `thermo_control` file need to be changed. The number of processors and pools can be changed in the same cases in which this is possible in `pw.x`. You cannot however run twice `thermo_pw.x` on the same directory using `what='scf_bands'` and two different paths, you must use two different working directories.

3.7 `what='scf_2d_bands'`

With this option the code makes a self-consistent calculation followed by a band structure calculation as with the option `what='scf_bands'`, but it assumes that the cell contains a slab with surfaces perpendicular to the z direction. Therefore the two-dimensional Bravais lattice of the surface is identified and the default path is chosen on the two-dimensional Brillouin zone. There are three options: Plot of the projected band structure (PBS); plot of the bands of the slab; plot of the bands of the slab above the projected band structure (the Fermi energies are aligned). In the first case the code computes several paths of $\mathbf{k}$ -points parallel to the surface (at different k_z) and does not plot the individual bands but selects the energy regions in which there are bulk states. The second case is similar to a standard band plot. The default path contains only $\mathbf{k}$ -points parallel to the surface (with $k_z = 0$). The third case assumes that the projected band structure has been already calculated and the information to plot it can be found on the file `flpbs`. For the rest it is similar to case two. For each direction, bands belonging to different irreducible representations of the point co-group of $\mathbf{k}$ can be plotted in the same panel or on different panels. This option is controlled by the following variables:

`lprojpbs`: When `.TRUE.` the projected band structure (PBS) is calculated if `nkz > 1` otherwise it is read from file. Usually this variable is `.TRUE.`. Set it to `.FALSE.` if you do not want to see the PBS, or if you want to see the bands of a bulk projected on the surface Brillouin zone without the PBS.
Default: logical `.TRUE.` (forced to `.FALSE.` if `what` is not `'scf_2d_bands'`)

`nkz`: The number of `kz` values used for the PBS plot. If `lprojpbs` is `.FALSE.` a plot of the bulk bands projected on the surface Brillouin zone is produced.
Default: integer 4 (forced to 1 if `what` is not `'scf_2d_bands'`).

`gap_thr`: minimum size (in eV) of the gaps in the PBS.
Default: real 0.1 eV

`sym_divide`: When `.TRUE.` the bands belonging to different irreducible representations are plotted in different panels. This option can be controlled by variables specified in the path (see below).
Default: logical `.FALSE.`

`identify_sur`: When `.TRUE.` the surface bands are searched and identified on the surface band structure.
Default: logical `.FALSE.`

`dump_states`: If `.TRUE.` and `identify_sur` is `.TRUE.` dump on the file `'dump/state_k_#'` the planar averages of the density (and in the noncollinear case also of the magnetization density) of each state. One file for

each k point is produced and # is the number of the k points. (Use with a small number of k points or it might create quite large files).
Default: logical .FALSE.

sur_layers: The number of surface layers on which we add the charge density of each state to check if it is a surface state.
Default: integer 2

sur_thr: the threshold (in percentage) of the charge density that must be on the surface layers to identify a state as a surface state.
Default: calculated from the actual charge density values of the states.

sp_min : minimum distance between layers. Two atoms form different layers only if their distance along z is larger than this number. Should be smaller than the interplanar distance (in a.u.) written by the tool gener_3d_slab.
Default: real 2.0 a.u.

subtract_vacuum: if .TRUE. the charge density of each state on vacuum is subtracted (to remove the vacuum states that are confused with surface states)
Default: .TRUE.

force_bands: when .TRUE. the bands are plotted in any case. Used to plot the bulk bands on top of the PBS, mainly for debugging.
Default: logical .FALSE.

only_bands_plot: if the files with the bands, the representations, the pbs and the projections are already on files, this option allows to change the parameters of the plot (such as the maximum energy or sur_thr) and do another plot without any additional calculation. If the files are missing and this variable is .TRUE. an error occurs.
Default: logical .FALSE.

flpbs: the name of the file that contains the information on the projected band structure.
Default: character(len=*) 'output_pbs'

flprojlayer: the name of the file that contains the information of the projection of the charge density of each state on each layer. Calculated only when identify_sur is .TRUE..
Default: character(len=*) 'output_projlayer'

The bands and the gnuplot scripts are saved on the same files that would be used with the option `what='scf_bands'`.

This option is not yet available for `lsda` calculations. Number of tasks for this option: 1. Image parallelization is not useful with this option.

By default the symmetry separation is not carried out. The code plots the bands of the slab on the same panel with a different color for each representation as in the bulk band structure plot (color refer to the representations of the slab point co-group of $\mathbf{k}$). In order to plot in different panels the different representations the user can specify `sym_divide= .TRUE..` By default this option is disabled and its use is rather tricky. In order to use it you must indicate explicitly the path on the two dimensional Brillouin zone using the option `q_in_band_form=.TRUE..` Close to the starting point of a given line you indicate the number of representations for that line (0 means all representations) and which ones. For instance for a (111) surface of an fcc metal in the direction $\bar{\Gamma} - \bar{M}$ you may want to plot separately the states even or odd with respect to the mirror plane perpendicular to the surface that contains the $\bar{\Gamma} - \bar{M}$ line. In order to do so you can specify the path as follows:

```
5
gG    30  0
K      30  0
M      30  1  1
gG      30  1  2
M        1  0
```

The representations to plot are indicated by their numbers (in this case 1 or 2). The number of the representation and the point co-group of each $\mathbf{k}$ -point can be found in the output of `thermo_pw`. These representation numbers refer to the point co-group of each $\mathbf{k}$ -point in the slab when you plot the slab band structures and to the point co-group of each $\mathbf{k}$ -point in the bulk when you plot a PBS. Some particular values of k_z , such as $k_z = 0$ might have a point co-group in the bulk different from the point co-group of a point with a generic k_z but in this case the representations are transformed into those of the smaller group using the group-subgroup relationships and the symmetry descent of the irreducible representations (only when `sym_divide=.TRUE.`). The representations of the smaller point co-group have to be used in the PBS input.

In general, the point co-group of a $\mathbf{k}$ -point $\mathbf{k} = (\mathbf{k}_{\parallel}, k_z)$ with component $\mathbf{k}_{\parallel}$ parallel to the surface and a generic k_z in the bulk is different from the point co-group of a $\mathbf{k}$ -point $\mathbf{k} = (\mathbf{k}_{\parallel}, 0)$ in the slab. Moreover, experimentally one cannot consider symmetries of the slab that exchange the two surfaces, and therefore the point co-group a $\mathbf{k}$ -point $\mathbf{k} = (\mathbf{k}_{\parallel}, 0)$ on the surface is a subgroup of the point co-group of $\mathbf{k} = (\mathbf{k}_{\parallel}, 0)$ on the slab. The point co-group $\mathbf{k} = (\mathbf{k}_{\parallel}, k_z)$ in the bulk does not contain operations that exchange k_z with $-k_z$ but it might be larger than the point co-group of $\mathbf{k} = (\mathbf{k}_{\parallel}, 0)$ on the surface because it might contain symmetries of the bulk that require fractional translations perpendicular to the surface that are not symmetries neither of the slab nor of the surface.

The point co-group of a given $\mathbf{k}$ -point $\mathbf{k} = (\mathbf{k}_{\parallel}, 0)$ on the surface can be found by removing from the corresponding slab point co-group the operations that exchange the two surfaces. It is also the group formed from the intersection

of the point co-group of $\mathbf{k} = (\mathbf{k}_{\parallel}, 0)$ in the slab and of the point $\mathbf{k} = (\mathbf{k}_{\parallel}, k_z)$ in the bulk. It is the user responsibility to specify the same number of panels for the PBS and for the slab calculation and to assure that the representations plotted in each panel correspond to each other. Returning to the example of the (111) surface of an fcc, in the direction $\bar{\Gamma} - \bar{K}$ the slab has C_2 symmetry about the x -axis, a symmetry that the surface has not. Therefore you can plot with two different colors the bands that belong to the A or B representations of the slab, (states even or odd with respect to a 180° rotation about the x axis, an operation that exchanges the two surfaces) but you cannot separate the PBS into even or odd states with respect to the C_2 symmetry. You might specify two different panels with the A or B bands in each, but the PBS in the two panels will be the same. On the contrary, for a $\mathbf{k}$ -point along the $\bar{\Gamma} - \bar{M}$ direction, the point co-group has the C_s symmetry both for the slab and for the surface, so you can separate both the PBS and the surface states in two different panels.

There is no input variable to control or change the colors or style of the plot. To change the defaults you can modify directly the `gnuplot` script, it is written in such a way that a change of a few variables can control the entire plot.

3.8 what='scf_dos'

With this option the code makes a self-consistent calculation followed by a band structure calculation on a uniform mesh of **k**-points and computes and plots the electronic density of states.

This option does not use the image parallelization and should be used with one image.

This option is controlled by the following variables:

```
deltae      : energy interval for electron dos plot (in Ry).
              Default: real 0.01 Ry.
ndose       : number of energy points in the dos plot.
              Default: determined from previous data
nk1_d, nk2_d, nk3_d : thick mesh for dos calculation.
              Default: integer 16, 16, 16
k1_d, k2_d, k3_d : the shift of the k point mesh.
              Default: integer 1, 1, 1
sigmae      : the smearing used for dos calculation (in eV).
              If 0.0 uses the degauss of the electronic
              structure calculation in metals and 0.01 Ry
              in insulators.
              Default: real 0.0
legauss     : When .TRUE. computes the electronic dos using
              a gaussian smearing. When .false. uses the same
              smearing of the electronic structure calculation
              in metals or gaussian smearing in insulators.
              Default: logical .false.
fleldos     : name of the file that contains the electron dos
              data.
              Default: character output_eldos.dat
flpseldos   : name of the postscript file that contains the
              electron dos picture.
              Default: character output_eldos
fleltherm   : name of the file that contains the electron
              thermodynamic data.
              Default: character output_eltherm.dat
flpseltherm : name of the postscript file that contains the
              plot of the electron thermodynamic quantities.
              Default: character 'output_eltherm'
```

The minimum and maximum energy, as well as the number of bands, are specified as with the option `what='scf_bands'`. However with the present option no energy shift is applied to the bands and the minimum and maximum energies refer to the unshifted eigenvalues. Note that after a calculation with `what='scf_dos'` you can run the tool code `epsilon_tpw.x` to evaluate the frequency dependent dielectric constant (for insulators only).

With this option, in the metallic case, the code computes the electronic thermodynamic quantities of a gas of independent electrons whose energy levels

give the calculated density of states and produces a postscript file with the electronic excitation energy, free energy, entropy, and constant strain heat capacity as a function of temperature. The zero of the electron energy is the energy at the smallest temperature required in input when it is lower than 4 K or 4 K.

Number of tasks for this option: 1.

3.9 `what='plot_bz'`

With this option the code writes a script to make a plot of the Brillouin zone (BZ) and of the path (the default one or the one given in input). The script must be read by the `asymptote` code, available at <http://asymptote.sourceforge.net/>. In many Linux distributions this code is available as a separate package, but it is not installed by default.

The following variables control the plot:

```
lasymptote : if .TRUE. asymptote is called by the program and
              the pdf file with the plot of the BZ is produced.
              Default: logical .FALSE.

flasy       : initial part of the name of the file where the
              asymptote script is written and of the name of the
              pdf file.
              Default: character(len=*) 'asy_tmp'

asymptote_command : the command that invokes asymptote and
                    produces the pdf file of the BZ.
                    Default: character(len=*) 'asy -f pdf -noprc
                    flasy.asy'

npx         : used only in the monoclinic cell, this parameter
              is needed to determine the shape of the Brillouin
              zone. The default value is usually large enough,
              but for particular shapes of the monoclinic
              Brillouin zone it could be small. If the code
              stops with an error asking to increase npx, double
              it until the error disappears.
              Default: integer 8
```

The structure of the solid can be seen using the `XCrySDen` code reading the input file of `pw.x`. You can find the code at <http://www.xcrysden.org/>. `THERMO_PW` produces also a file in the `xsf` format called `prefix.xsf`, where the variable `prefix` is given in the input of `pw.x`. This can be useful when the nonequivalent atomic positions and the space group are given in the input of `pw.x`. To see an `xsf` file, give the command `xcrysden -xsf file.xsf`.

With this option the code produces also a file with the X-ray powder diffraction intensities for the solid. A plot shows the scattering angles and the relative intensity of each peak. Note that this plot is made using a superposition of atomic charges, not the self-consistent charge. By setting the flag `lformf=.TRUE.` the atomic form factors of all the atomic types used to calculate the intensities are plotted. By setting the flag `lxrdp=.TRUE.` the intensities plot is done also after the cell optimization and after a self-consistent calculation for the options that support it. The variables that control these plots are:

```
lambda      : The X-ray wavelength (in A) used to calculate the
              scattering angles.
              Default: Cu alpha line 1.541838 A if lambda_element
```

```

                                is empty
lambda_elem : The anode element, used to set the X-ray
                                wavelength. Supported elements 'Cr', 'Fe', 'Co',
                                'Cu', 'Mo'. NB: lambda must be zero to use
                                lambda_elem, otherwise the value of lambda given
                                in input is used.
                                Default: character(len=2) ' '
flxrdp      : name of the file where the scattering angles and
                                intensities are written.
                                Default: character 'output_xrdp.dat'
flpsxrdp    : name of the postscript file with the X-ray
                                diffraction spectrum.
                                Default: character 'output_xrdp'
lxdp        : if .TRUE. compute the xrdp also after the cell
                                optimization with all the options mur_lc_...
                                with the uniformly strained atomic positions
                                and after the scf calculation if supported by
                                the option.
                                Default: logical .FALSE.
lformf      : if .TRUE. plot also the form factor of each
                                atom type present in the solid. Note that the
                                atom type is recognized from the atom name in
                                the thermo_pw input. The name must coincide
                                with the symbols in the periodic table. (Cu, H,
                                Li, Lil, ... are correct, CU, LI, H1 ... are
                                wrong).
                                Default: logical .FALSE.
smin        : minimum value of s used in the atomic form
                                factor plot.
                                Default: real 0.0
smax        : maximum value of s used in the atomic form
                                factor plot.
                                Default: real 1.0
nspoint     : number of points in which the atomic form
                                factor is calculated.
                                Default: integer 200
lcm         : when .TRUE. the code uses the Cromer-Mann
                                coefficients from the International Tables of
                                Crystallography to compute the atomic form
                                factors, otherwise uses the Doyle-Turner or
                                Smith-Burge parameters.
                                Default: logical .FALSE.
flformf     : name of the file in which the atomic form
                                factor is written. The code adds a number to
                                each file name and creates a file per atom type.
                                Default: character 'output_formf.dat'

```

`flpsformf` : name of the postscript file with the atomic form factor. The code adds a number to each file name and creates a file per atom type.
Default: character `'output_formf'`

3.10 `what='scf_ph'`

With this option the code makes a self-consistent calculation followed by a phonon calculation. The phonon calculation is controlled by the file `ph_control` and can be at a single $\mathbf{q}$ point or on a mesh of $\mathbf{q}$ points. The different representations are calculated in parallel when several images are available. No other input variable is necessary. The outputs of this calculation are the dynamical matrices files.

`thermo_pw` adds to the `ph.x` code the ability to compute the complex dielectric constant tensor of insulators as a function of a complex frequency for the study of optical properties within time-dependent density functional perturbation theory (TD-DFPT). The code produces also the complex index of refraction for all systems except monoclinic and triclinic. For cubic solids it makes also a plot of the reflectivity for normal incidence and of the adsorption coefficient. As a default the TD-DFPT algorithm uses the Sternheimer equation and a self-consistent loop, but it is also possible to use a Lanczos chain. The option is activated in the `ph.x` input by setting `epsil=.TRUE.` and `fpol=.TRUE.`, but at variance with the `ph.x` code, the frequencies must be specified as complex numbers. The following additional variables can be put in the input of the `ph.x` code, to select the frequency range and the number of frequencies to compute:

`freq_line` : if this variable is `.TRUE.`, after the `FREQUENCY` keyword the code expects the number of frequency points and the starting and final frequencies. If `.FALSE.` the number of frequencies and a list of frequencies are given. The frequencies are complex numbers and are given with a real and an imaginary part (in Ry), without parenthesis.
Default: `.FALSE.`

`delta_freq` : When `freq_line` is `.TRUE.` instead of giving the last frequency of the line one can give the distance between two frequency points `delta_freq` as a complex number. The last point of the line is calculated using the number of frequencies `nfs` and the first frequency. When `delta_freq` is not zero the last frequency is not used and can be omitted.
Default: `complex, (0.0, 0.0).`

`start_freq` : Number of the initial frequency calculated in the job in the sequence of frequencies.
Default: `integer 1`

`last_freq` : Number of the final frequency calculated in the job in the sequence of frequencies.
Default: `integer nfs (total number of frequencies)`

`lfreq_ev` : If `.TRUE.` the units of the frequencies are eV instead of the default Ry units.
Default: `logical .FALSE.`

`linear_im_freq`: This option is used only when `freq_line=.TRUE.`. When `linear_freq_im` is `.TRUE.`, the imaginary part of each frequency is calculated as $\eta * \text{freq}$ where η is the imaginary part of the first frequency on the frequency line.
Default: logical `.FALSE.`

`llanczos`: When this flag is `.TRUE.` at finite frequencies a Lanczos algorithm is used to solve the linear system. Can be very fast but might require much more memory than the standard algorithm. Presently it is incompatible with images.
Default: `.FALSE.`

`lanczos_steps`: steps of the Lanczos chain.
Default: integer 2000

`lanczos_steps_ext`: steps of the extrapolated Lanczos chain
Default: integer 10000

`lanczos_restart_steps`: number of steps between saving of the Lanczos status. If 0 the status is saved only at the end of the run. Use `recover=.TRUE.` to resume an interrupted Lanczos chain or to increase the number of steps.
Default: integer 0

`extrapolation` : extrapolation type. Presently only 'no' or 'average' are available. In the first case no extrapolation is applied, in the second the average of the beta and gamma is used.
Default: character 'average'

`pseudo_hermitian` : when `.TRUE.` a pseudo-hermitian algorithm is used to make the Lanczos steps. Should be twice faster than the default non hermitian algorithm.
Default: `.TRUE.`

`only_spectrum` : Computes only the spectrum assuming that the Lanczos chain coefficients are in a file. It gives error if the number of requested Lanczos steps is larger than those available on file.
Default: logical `.FALSE.`

`lcg`: When this flag is `.TRUE.` a global conjugate gradient algorithm is used to compute the dielectric constant and the phonon frequencies. It will not require mixing, but will use more memory than the standard algorithm (for insulators only). It is not available for the frequency dependent case.
Default: `.FALSE.`

When a non zero wave-vector $\mathbf{q}$ is specified in the input of the phonon code, the previous options produce the inverse of the dielectric constant as a

function of the frequency at the wave-vector $\mathbf{q}$ (this option can be used both for insulators and metals).

Additional variables can be specified in the THERMO_PW input to control where the frequency dependent dielectric constant is written and plotted and how the work is divided among images:

```
flepsilon : beginning of the name of the file where the
            frequency dependent dielectric constant is
            written at finite  $q$  (the code adds the
            extensions _re and _im).
            Default: character(len=*) 'epsilon'
flpsepsilon : name of the postscript file where the frequency
            dependent dielectric constant is plotted at finite  $q$ .
            Default: character(len=*) 'output_epsilon'
floptical : beginning of the name of the file where the
            frequency dependent dielectric constant and the
            complex index of refraction are written (the code
            adds the extensions _xx, _yy, and _zz for the solids
            that need to distinguish the different directions).
            Default: character(len=*) 'optical'
flpsoptical : name of the postscript file where the frequency
            dependent dielectric constant, the complex index of
            refraction and for cubic solid also the reflectivity
            and the absorption coefficient are plotted.
            Default: character(len=*) 'output_optical'
omega_group : number of frequencies calculated together by
            each image. This variable is used only with images.
            Default: integer 1.
```

An example for this option can be found in example03, example16, example17, example20, and example21.

Number of tasks for this option: for a phonon calculation the number of parallelizable tasks of the phonon code (smaller but of the order of the number of $\mathbf{q}$ points times $3N_{at}$, where N_{at} is the number of atoms in the unit cell), for a dielectric constant calculation using Sternheimer equation `nfs/omega_group`, number of frequencies divided by the number of frequencies in each group, for a dielectric constant calculation using Lanczos 1 (images not allowed).

It is also possible to separate the self-consistent and the phonon calculation, by running first `thermo_pw.x` using `what='scf'` and then running, on the same directory, `thermo_pw.x` using `what='scf_ph'`. The same input can be used in the two calculations, only the `thermo_control` file need to be changed. The number or processors/pools/images can be changed in the same cases in which this is possible in Quantum ESPRESSO.

Using images in a phonon calculation with the master/slave approach has an overhead because each image must recalculate the initialization and the

band structure at each task, or check if the bands are already on disk, calculated previously by the same image. On some systems with slow disks it could be faster to recalculate the bands instead of reading them from disk. It is also possible to use the image breaking suggested by the `ph.x` code that keeps, as much as possible, on the same image the tasks that require the same initialization without recomputing it. The input variables that control this part of the calculation are:

```
force_band_calculation : if .TRUE. the bands are never read
                        from disk but recalculated when needed.
                        Default: logical .FALSE.
use_ph_images          : if .TRUE. each image makes a set of tasks so
                        as to minimize the number of band calculations
                        and phonon initialization.
                        Default: logical .FALSE. if nimage>1 .TRUE.
                        nimage=1.
sym_for_diago          : When .TRUE. use symmetry to calculate the
                        bands and the unperturbed wavefunctions instead
                        of diagonalizing the Hamiltonian.
                        Default: logical .FALSE.
```

3.11 `what='scf_disp'`

With this option the code makes a self-consistent calculation followed by a phonon dispersion calculation at a fixed geometry. The geometry is given in the input of `pw.x`. The dynamical matrices are used to calculate the interatomic force constants that are then used to calculate the dynamical matrices and hence the phonon frequencies along a path in the Brillouin zone and on a much thicker mesh of $\mathbf{q}$ points. The path can be generated automatically or given in input as in a band structure calculation (see above `what='scf_bands'`). The uniform mesh of $\mathbf{q}$ points is controlled by `thermo_control`. The code uses the phonon frequencies calculated on the thick mesh of $\mathbf{q}$ points to get the phonon density of states using a smearing approach. The density of states is used to calculate the harmonic thermodynamic properties: vibrational energy, free energy, entropy, and constant strain heat capacity. The same thermodynamic quantities are calculated also by direct integration over the Brillouin zone and compared in the plots. When `with_eigen=.TRUE.`, the atomic B-factors are calculated as a function of temperature by the generalized vibrational density of states or by a direct integration over the Brillouin zone. Note that presently no interpolation formula is used at low temperatures so `thermo_pw` can not be used to obtain thermodynamic properties at very low temperatures. The extensive quantities plotted in the figures refer to an Avogadro number of unit cells. If you need them per mole you have to divide by the number of formula units in a unit cell.

The input variables that control this option are:

```
freqmin_input : minimum frequency for phonon dos plot.
                  Default: real determined from phonon frequencies
freqmax_input : maximum frequency for phonon dos plot.
                  Default: real determined from phonon frequencies
deltafreq      : frequency interval for phonon dos plot.
                  Default: real 1 cm-1
ndos_input     : number of frequency points in the dos plot.
                  Default: determined from previous data
nq1_d, nq2_d, nq3_d : thick mesh for phonon dos calculation.
                  Default: integer 192, 192, 192
phdos_sigma    : the smearing used for phonon dos calculation
                  (in cm-1).
                  Default: real 2. cm-1
idebye         : if 1, 2, or 3 the code computes the Debye
                  temperature as a function of temperature from
                  the free energy, the vibrational energy, or the
                  heat capacity respectively.
                  Default: integer 0
after_disp     : if .TRUE. the dynamical matrices are supposed
                  to be already available in files in the current
                  directory. This option is needed to restart when
                  the outdir directory has been erased and ph.x
```

cannot be run without redoing the scf calculation.
The exact restart point depends on the files
already available on the current directory.
Default: logical .FALSE.

fildyn : the name of the dynamical matrix file, as
would be specified in the input of ph. To be used
when after_disp is .TRUE..
Default: character ' '

zasr : type of acoustic sum rule applied to the ifc.
Default: character(len=*) 'Simple'

ltherm_dos : if .TRUE. the thermal properties are calculated
from the phonon dos.
Default: logical .TRUE.

ltherm_freq : if .TRUE. the thermal properties are calculated
from the direct integration using the phonon
frequencies.
Default: logical .TRUE.

lgruneisen_gen: If .TRUE. phonons are computed only in a reduced set of
geometries and first (and second) derivatives of the
free energy with respect to crystal parameters are
evaluated by finite differences.
Default: .FALSE.

ggrun_recipe: the degree of the polynomial used to fit the free energy
for the general Gruneisen theory
1 linear polynomial (requires 2 nvar + 1 phonons)
2 quadratic polynomial (requires 3^nvar phonons)
3 quadratic polynomial (requires nvar (nvar+3) /2 +1
phonons)
For energy the polynomial degree is 2 or 4
controlled by the lquartic variable.
Default: 2

icenter_grun The reference geometry to compute phonons. The 3*nvar,
nvar*(nvar+3)/2+1 or the 2 * nvar + 1 geometries are
calculated starting from this.
The user must check that this point is not on the border
of the grid.
Default: Integer, the point of coordinates (ngeo(i)+1)/2
in each direction (transformed into an integer).

flfrc : file where the interatomic force constants are
written.
Default: character(len=*) 'output_frc.dat.gl'

flfrq : file where matdyn writes the interpolated
frequencies.
Default: character(len=*) 'output_frq.dat.gl'

flvec : file where the eigenvectors of the dynamical
matrix are written.

```

                                Default: character(len=*) 'matdyn.modes'
fldosfrq      : file where the frequencies used to calculate
                the phonon density of states are saved.
                                Default: character(len=*) 'save_frequencies.dat'
fldos         : file where the phonon dos is written.
                                Default: character(len=*) 'output_dos.dat.gl'
fltherm       : file where the harmonic thermodynamic
                quantities are written.
                                Default: character(len=*) 'output_therm.dat.gl'
flpsdisp      : postscript file of the phonon dispersions.
                                Default: character(len=*) 'output_disp'
flpsdos       : postscript file of the phonon dos.
                                Default: character(len=*) 'output_dos'
flpstherm     : postscript file of the harmonic thermodynamic
                quantities.
                                Default: character(len=*) 'output_therm'

```

This option requires `ldisp=.TRUE.` in the phonon input.

An example for this option can be found in `example04`.

Number of tasks for this option: number of parallelizable tasks of the phonon code (smaller but of the order of number of **q** points times $3N_{at}$, where N_{at} is the number of atoms in the unit cell).

3.12 `what='scf_elastic_constants'`

With this option the code calculates the elastic constants of the solid at the geometry given in the `pw.x` input. There are four different algorithms that at convergence should give the same results. In two of them, depending on the Laue class, the code calculates the nonzero components of the stress tensor for a set of strains and obtains the elastic constants from the numerical first derivative of the stress with respect to strain. The two algorithms `standard` and `advanced` differ only for the choice of the unit cell. In the `standard` method the code applies the strain to the primitive vectors of the unstrained solid and uses `ibrav=0` and the strained vectors to compute the stress tensor. The `advanced` method, available only for selected Bravais lattices, tries to optimize the calculation by choosing strains for which the number of needed **k**-points is reduced. Moreover it identifies the Bravais lattice of the strained solid and recalculates the primitive vectors with the conventions of QUANTUM ESPRESSO. When available this should be the most efficient method. The other two algorithms are called `energy_std` and `energy`. Using the `energy_std` or `energy` algorithm the elastic constants are calculated from a polynomial fit of the total energy as a function of strain without computing stress. This option usually requires more independent strains. It can be used when stress calculation is not implemented in QUANTUM ESPRESSO. The difference between `energy_std` and `energy` is the same between the algorithms that use the stress. With `energy_std` the code applies the strain using `ibrav=0`, while with `energy` an optimized cell is used. As the `advanced` algorithm the `energy` algorithm is available only for selected Bravais lattices. For all methods the number of strains is `ngeo_strain` for each independent strain. For each strain, the code relaxes the ions to their equilibrium positions if `frozen_ions=.FALSE.` or keeps them in the strained positions if `frozen_ions=.TRUE.`. Note that elastic constant calculations with `frozen_ions=.FALSE.` might require smaller force convergence threshold than standard calculations. The default value of `forc_conv_thr` must be changed in the `pw.x` input.

If you know in which direction the atoms relax and if there is only one atom that moves in each strain type, one can use `frozen_ions=.TRUE.`, and compute the energy at several atomic positions choosing the one that minimizes the energy. This approach computes more energy configurations but does not relax them and should provide results identical to `frozen_ions=.FALSE.`. It can be used only with `energy` and `energy_std` algorithms, but should work also with the free energy in `what='elastic_constants_geo'`.

At finite pressure all methods give the elastic constants that relate linearly stress and strain.

The input variables that control this option are:

```
frozen_ions: if .TRUE. the elastic constants are calculated
              keeping the ions frozen in the strained positions.
              Default: logical .FALSE.
ngeo_strain: the number of strained configurations used to
```

calculate each derivative.
Default: integer 4 ('standard' and 'advanced'),
6 ('energy')

elastic_algorithm: 'standard', 'advanced', 'energy_std' or
'energy'. See discussion above.
Default: character 'standard'

delta_epsilon: the interval of strain values between two
geometries. To avoid a zero strain geometry
that might have a different symmetry ngeo_strain
must be even.
Default: real 0.005

epsilon_0: a minimum strain. For small strains the ionic
relaxation routine requires a very small threshold
to give the correct internal relaxations and
sometimes fail to converge. In this case you
can increase delta_epsilon, but if delta_epsilon
becomes too large you can reach the nonlinear
regime. In this case you can use a small
delta_epsilon and a minimum strain (To be used
only for difficult systems).
Default: real 0.0

poly_degree: degree of the polynomial used to interpolate
stress or energy. ngeo_strain must be larger
than poly_degree+1.
Default: 3 ('standard', 'advanced', 2
if ngeo_strain < 6), 4 ('energy', 3 if
ngeo_strain < 6).

stype(1:strain_types): For each strain type for which this
variable is .TRUE. it is possible to compute
the energy by moving one atom in one direction
and minimizing the energy with respect to the
ionic position. This option requires frozen_ions
=.TRUE. and should give the same result of
frozen_ions=.FALSE. if the atom that moves and
its direction is chosen correctly.
Default: logical .FALSE..

move_at(1:strain_types): The atom that moves in each strain type.
Default: integer none (it must be set
if stype(istrain)=.TRUE.).

atom_dir(3,1:strain_types): The versor of the atomic movement in
Cartesian coordinates (the vector can be given with
any module. It is normalized by the code).
Default: real none (it must be set if
stype(istrain)=.TRUE.).

atom_step(1:strain_types): the step for each atomic displacement.
Default : real 0.01 a.u..

<code>nmove</code>	Number of atomic positions sampled for minimizing the energy at each strain type. Default: integer 5.
<code>lcm_ec</code>	If <code>.TRUE.</code> the center of mass of the cell is conserved in the displacement introduced with previous variables. Default: logical <code>.TRUE.</code> .
<code>fl_el_cons:</code>	the name of the file that contains the elastic constants. Default: <code>character(len=*) 'output_el_con.dat'</code>

The three algorithms are equivalent only at convergence both with **k**-point sampling and with the kinetic energy cut-off, but large differences between the elastic constants obtained with the standard and advanced algorithms might point to insufficient **k**-point sampling. Large differences between the elastic constants obtained with the `energy_std` or `energy` algorithms with respect to the other two might point to insufficient kinetic energy cut-off.

Number of tasks for this option: `ngeo_strain` times the number of independent strains.

Using the elastic constants tensor the code can calculate and print a few auxiliary quantities: the bulk modulus, the poly-crystalline averages of the Young modulus, of the shear modulus, and of the Poisson ratio. Both the Voigt and the Reuss averages are printed together with the Hill average. The Voigt-Reuss-Hill average of the shear modulus and of the bulk modulus are used to compute average sound velocities. The average of the Poisson ratio and the bulk modulus allow the estimation of the Debye temperature. The Debye temperature is calculated also with the exact formula evaluating the average sound velocity from the angular average of the sound velocities calculated for each propagation direction solving the Christoffel wave equation. The exact Debye temperature is used within the Debye model to calculate the Debye's vibrational energy, free energy, entropy, and constant strain heat capacity. These quantities are plotted in a postscript file as a function of temperature.

3.13 `what='mur_lc'`

With this option the code runs several self-consistent calculations at different geometries. The runs can be done in parallel when several images are available. This option has two working modes controlled by the logical variable `lmurn`. When `lmurn=.TRUE.` the total energy as a function of the volume is interpolated by an equation of state (Murnaghan or Birch-Murnaghan) and a plot of the energy as a function of the volume, of the pressure as a function of the volume and of the enthalpy as a function of pressure are produced. The volume is changed by changing only `celldm(1)`. `celldm(2)...``celldm(6)` remain fixed at the values given as input of `pw.x` or can be read from file using the option `lgeo_from_file=.TRUE.`. When `lmurn=.FALSE.` the energy is calculated in a uniform grid of parameters composed of `ngeo(1) × ngeo(2) ... × ngeo(6)` points. The energies are fitted with a quadratic or quartic polynomial of N_k variables, where N_k is the number of independent crystal parameters for the given crystal system. A plot of the energy as a function of the lattice constant is produced for cubic systems. For solids of the hexagonal, tetragonal, and trigonal systems contour plots of the energy as a function of the two crystal parameters (a and c/a or a and $\cos \alpha$) are plotted. For orthorhombic systems contour plots of the energy as a function of a and b/a are plotted for each value of c/a . Presently no graphical tool is implemented to plot the energy for monoclinic and triclinic crystal systems. However in all cases the enthalpy as a function of pressure is shown. Moreover separate plots show the crystal parameters as well as the volume as a function of pressure. When in the directory `elastic_constants` there are the elastic constants for each geometry (calculated by the option `what=elastic_constants_geo`), these are interpolated at the pressure dependent crystal parameters and shown on output. Using the input variable `lgeo_to_file=.TRUE.` the code writes on file the crystal parameters that for each `celldm(1)` of the grid of crystal structures give a uniform pressure. When `lmurn=.FALSE.` the bulk modulus is not calculated. To obtain it, you can calculate the elastic constants at the minimum geometry (see the option `what='mur_lc_elastic_constants'`). With this option the pressure control is active. You can specify a finite pressure and the enthalpy is minimized instead of the energy. Note however that if the minimum is distant from the starting configuration its associated error can be large, larger for the quadratic than for the Murnaghan interpolation. For this reason the present option should be used starting from the minimum found by `pw.x` using the `vc-relax` option and the pressure should not be too different from the pressure used for `vc-relax`. Note that with this option the atomic coordinates are relaxed at each geometry even if you specified `calculation='scf'` in the `pw.x` input. Use `frozen_ion=.TRUE.` if you want to keep them fixed. To increase the maximum number of ionic iterations use `calculation='relax'` and give `nstep` (otherwise the default is 20). Only the `bfgs` relaxation is supported by this option. When `lel_free_energy=.TRUE.` the code makes also an electronic bands dos calculation at each geometry, computes the electronic thermodynamic quantities as a function of tempera-

ture and writes them in separate files. These files can be used to add the electronic contribution to the anharmonic properties with the option `mur_lc_t`. This option can be controlled by the following variables:

`ngeo(1), ..., ngeo(6)` : the number of geometries to use for each `celldm` parameter. The lattice constant of these geometries is calculated from the input of `pw.x`. `celldm(1), ..., celldm(6)` of this input is used for the central geometry. For the others `celldm(1), ..., celldm(6)`, are changed in steps of `step_ngeo(1), ..., step_ngeo(6)`. `ngeo(1)` must be odd. Only the values of `celldm` relevant for each Bravais lattice are actually changed.
Default: integer 1,1,1,1,1,1 for `what=scf_*`, 9,1,1,1,1,1 for `what=mur_lc_*` and `lmurn=.TRUE.` or for cubic systems, 5 on all the relevant `celldm` parameters when `lmurn=.FALSE.` and the system is not cubic.

`step_ngeo(1), ..., step_ngeo(6)` : The step between the lattice constants at different geometries. `step_ngeo(1)` is, in atomic units, the change of `a`, `step_ngeo(2)`, `step_ngeo(3)` are dimensionless and are the changes of the ratios `b/a`, `c/a`, `step_ngeo(4)`, `step_ngeo(5)`, `step_ngeo(6)` are the changes in degree of the angles `alpha`, `beta`, and `gamma`. The cosine of the angle is calculated by the program.
Default: real 0.05 a.u., 0.02, 0.02, 0.5, 0.5, 0.5

`lmurn` : if `.TRUE.` the fit with an equation of state is done. Only `ngeo(1)` values of the energy are fitted, the other values of `ngeo` are not used. if `.FALSE.` use a quadratic or quartic function to interpolate the energy as a function of all `celldm` parameters. The number of self-consistent calculations is `ngeo(1) x ngeo(2) x ngeo(3) x ngeo(4) x ngeo(5) x ngeo(6)`. In this case only the minimum energy and the optimal `celldm` are given in output.
Default: `.TRUE.`

`ieos` : choose the equation of state to use (only when `lmurn=.TRUE.`):
1 - Birch-Murnaghan third order
2 - Birch-Murnaghan fourth order
4 - Murnaghan
Default: integer 4

`show_fit` : if `.TRUE.` show the contour plot of the fitted energy instead of the energy. Used by default when `reduced_grid` is `.TRUE.`.

Default: logical .FALSE.

frozen_ions: if .TRUE. the atomic coordinates are obtained by applying the strain to the coordinates given in the pw.x input to the new cell parameters (equivalent to keep the crystal coordinates fixed) and kept fixed. If .FALSE. the atomic coordinates are relaxed at each geometry.
Default: logical .FALSE.

vmin_input : minimum volume for the plot of the energy as a function of volume.
Default: real 0.98 times the volume of the first geometry.

vmax_input : maximum volume for the plot of the energy as a function of volume.
Default: real 1.02 times the volume of the last geometry.

deltav : distance between two volumes in the plot of the energy as a function of the volume.
Default: real calculated from nvol.

nvol : number of volumes in equation of state plot.
Default: integer 51

lquartic : if .TRUE. fit the energy with a quartic polynomial.
Default: logical .TRUE.

lsolve : choose the algorithm used to fit the quartic polynomial parameters.
Allowed values:
1 explicitly minimize χ^2 (usually less accurate than the other two. Should be used only for tests).
2 Use the QR algorithm to minimize χ^2 (lapack routine dgels)
3 Use the SVD algorithm to minimize χ^2 (lapack routine dgels).
Default: integer 2

flevdat : file where the equation of state is written. The results of the fit are then written in flevdat.ev.out.
Default: character(len=*) 'output_ev.dat'

flpsmur : postscript file of the equation of state plot.
Default: character(len=*) 'output_mur'

lel_free_energy : if .TRUE. computes the electronic thermodynamic properties (energy, free energy, entropy, and constant strain heat capacity) at each temperature and plots them. See the scf_dos option for the parameters that control the calculation.
Default: .FALSE.

ncontours : the number of contours in the energy plot. These levels can be determined automatically by the code

or defined by the user. The energy levels can be defined after the INPUT_THERMO namelist but before the path, as a list:

```
energy_level(1)      color(1)
```

```
...
```

```
energy_level(ncontours)  color(ncontours)
```

Color is a string of the type color_red, color_green, etc.

The list of available colors is at the beginning of each gnuplot script. energy_level is in Ry units.

Default: integer 9

do_scf_relax : if .TRUE. the code makes a self-consistent relax calculation at the equilibrium geometry to find the optimized atomic coordinates. This step is needed only for solids that have internal degrees of freedom in the unstrained configuration. If .FALSE. the coordinates of the input geometry are strained uniformly to the equilibrium geometry. Default: logical .FALSE.

lgeo_from_file : if .TRUE. the input geometries are read from file. ngeo(1) must have the total number of geometries and lmurn must be .TRUE..
Default : .FALSE.

lgeo_to_file : if .TRUE. at the end of the calculation the code writes in a file the geometries that correspond to the optimized crystal parameters for each value of celldm(1) of the grid of geometries.
Default : .FALSE.

flenergy : name of the file that contains the energy in a form that can be used by gnuplot to make contour plots.
Default: character(len=*) 'output_energy'

flgeom : name of the file that contains the geometries requested with the flags lgeo_to_file or lgeo_from_file. The file is in the directory energy files.
Default: character(len=*) 'output_geometry'

flpsenergy : file with the contour plots of the energy as a function of the crystal parameters.
Default: character(len=*) 'output_energy'

An example for this option can be found in example05.

Number of tasks for this option:

ngeo(1) **when** lmurn=.TRUE.,

ngeo(1) × ngeo(2) × ngeo(3) × ngeo(4) × ngeo(5) × ngeo(6) **when** lmurn=.FALSE..

3.14 **what='mur_lc_bands'**

With this option the code computes the band structure at the geometry that minimizes the energy (or the enthalpy at finite pressure). See `what='scf_bands'` and `what='mur_lc'` for a list of the variables that control these two options. An example for this option can be found in `example06`.
Number of tasks for this option: see `what='mur_lc'` and `what='scf_bands'`.

3.15 **what='mur_lc_dos'**

With this option the code computes the electronic dos at the geometry that minimizes the energy (or the enthalpy at finite pressure). See `what='scf_dos'` and `what='mur_lc'` for a list of the variables that control these two options. Number of tasks for this option: see `what='mur_lc'` and `what='scf_dos'`.

3.16 **what='mur_lc_ph'**

This option is similar to `what='scf_ph'` but the phonon calculation is made at the geometry that minimizes the energy (or the enthalpy at finite pressure). See `what='scf_ph'` and `what='mur_lc'` for a list of the variables that control these options. An example for this option can be found in `example07`. Number of tasks for this option: Maximum between the number of tasks needed by the `what='mur_lc'` option and the number of tasks of the phonon code (see above the option `what='scf_ph'`).

3.17 **what='mur_lc_disp'**

This option is similar to `what='scf_disp'` but the phonon calculation is made at the geometry that minimizes the energy (or the enthalpy at finite pressure). See `what='scf_disp'` and `what='mur_lc'` for a list of the variables that control these options. An example for this option can be found in `example08`.

Number of tasks for this option: Maximum between the number of tasks needed by the `what='mur_lc'` option and the number of tasks of the phonon code (see above the option `what='scf_ph'`).

3.18 **what='mur_lc_elastic_constants'**

As `what='scf_elastic_constants'` but the calculation is made at the geometry that minimizes the energy (or the enthalpy at finite pressure). See `what='scf_elastic_constants'` and `what='mur_lc'` for a list of the variables that control these two options. An example for this option can be found in `example13`.

Number of tasks for this option: Maximum between the number of tasks needed by the `what='mur_lc'` option and the number of tasks needed for the `what='scf_elastic_constants'` option.

3.19 `what='mur_lc_t'`

With this option the code calculates the anharmonic properties within the quasi-harmonic approximation. The outputs of the code are the values of crystal parameters (`celldm`) as a function of temperature. This calculation is done by computing the phonon dispersions on all the geometries specified as in `what='mur_lc'` (or on a subset of these geometries) and minimizing the Helmholtz free energy. Separate plots of the phonon dispersions are obtained for all the calculated geometries. For each geometry the code produces also plots of the phonon density of states and of the harmonic thermodynamic quantities. From `celldm` as a function of temperature the code computes the thermal expansion tensor, the volume, and the volume thermal expansion as a function of temperature. The frequencies at all the calculated geometries are interpolated by quadratic or quartic polynomials of the crystal parameters and can be shown at crystal parameters given in input or at those that minimize the free energy at a temperature given in input. The interpolated frequencies are shown on the same path used for the phonon dispersions. In addition to the frequencies the code produces also several plots of the derivatives of the frequencies with respect to the crystal parameters multiplied by the crystal parameters.

When an equation of state is used to interpolate the Helmholtz free energy (`lmurn=.TRUE.`), in addition to the volume, the bulk modulus and the pressure derivative of the bulk modulus are plotted as a function of temperature. Moreover the isobaric heat capacity, the isoentropic bulk modulus, and the average Grüneisen parameter are calculated as a function of temperature. The mode Grüneisen parameters are calculated with cubic interpolations of the phonon frequencies. Using the variable `with_eigen` one can calculate these parameters as derivatives of the phonon frequencies (default) or as expectation values of the derivatives of the dynamical matrix on the central geometry eigenvectors (might require a lot of RAM). The mode Grüneisen parameters are used to calculate the volume thermal expansion and the result is compared with the volume thermal expansion derived from the numerical derivative of the equilibrium volume obtained from the minimization of the Helmholtz free energy. When the Helmholtz free energy is interpolated with a quadratic or cubic polynomial (`lmurn=.FALSE.`), by default, the code computes only the temperature dependence of the lattice parameters and of the volume, the volume thermal expansion, and the thermal expansion tensor. However if a file with the elastic constants is found in the `elastic_constants` directory and `lb0_t=.FALSE.` the bulk modulus is calculated and assumed independent from the temperature so that also the isobaric specific heat, the isoentropic bulk modulus, and the average Grüneisen parameter are calculated as a function of temperature. The derivatives of the frequencies with respect to the crystal parameters are used to calculate the thermal expansion tensor which is compared with that obtained from the numerical derivatives of the crystal parameters obtained from the minimization of the Helmholtz free energy. If many files with the elastic constants, one for each geometry, as produced with

the option `elastic_constants_geo`, are found in the `elastic_constants` directory and `lb0_t=.TRUE.`, the bulk modulus and the elastic constants are computed as a function of temperature within the “quasi-static approximation” and are used to calculate the other thermodynamic properties. If one or many elastic constants files are found in the `anhar_files` directory and `lb0_t=.TRUE.` the bulk modulus and elastic constants are computed as a function of temperature within the “quasi-harmonic approximation” and are used to calculate the other thermodynamic properties (see a more detailed discussion in the option `what=elastic_constants_geo`). In addition to the quantities plotted for cubic solids, the code plots also the elastic constants and the bulk modulus as a function of the temperature and the elastic compliances and the compressibility as a function of temperature. The elastic constants are interpolated with a quadratic (`lquartic=.FALSE.`) or quartic (`lquartic=.TRUE.`) polynomial of the crystal parameters. Moreover the thermal stresses and the generalized average Grüneisen parameters are plotted. These possibilities are implemented only for cubic, tetragonal, hexagonal, trigonal, and orthorhombic systems.

With this option the pressure control is active. You can specify a finite pressure and the Gibbs energy is minimized instead of the Helmholtz free energy. Note however that if the minimum is distant from the starting configuration its associated error can be large, larger for the quadratic than for the equation of state interpolation.

By using the variables `ntemp_plot` and `temp_plot` (see the Section Temperature and pressure) the code produces also plots of the thermal expansion, of the bulk modulus, of the average Grüneisen parameter, and of the product of the thermal expansion and the bulk modulus as a function of pressure. These plots contain several lines one for each chosen temperature. By using the variables `npress_plot` and `press_plot` (see the Section Temperature and pressure) in addition to the plots of the volume, the bulk modulus, the thermal expansion, the isobaric heat capacity, the isoentropic bulk modulus, and the average Grüneisen parameter as a function of temperature calculated at the input pressure, the code produces also a plot of the same quantities for all the `npress_plot` pressures. Note that these plots may be inaccurate if the chosen pressures are at geometries distant from the set of geometries chosen by the code for the anharmonic calculation, so these plots might require values of `ngeo` larger than the default. Presently the use of these variables is limited to cubic solids. By using the variables `nvol_plot` and `ivol_plot` the code produces a plot of the thermal pressure as a function of temperature for several volumes.

The Helmholtz (or Gibbs at finite pressure) free-energy can be interpolated in two ways depending on the variable `ltherm_glob`. When `ltherm_glob=.FALSE.` (default) the vibrational (plus the electronic) free energy is fitted separately by a polynomial, while only the static energy is fitted by an equation of state. When `ltherm_glob=.TRUE.` a different equation of state is used at each temperature and no polynomial interpolation is needed. The electronic free energy is added if the flag `lel_free_energy=.TRUE.`. This electronic free energy can

be calculated in two ways. When `hot_electrons=.FALSE.` the code expects to find in the directory `therm_files` the file with the electronic free energy computed with the option `what=mur_lc`. When `hot_electrons=.TRUE.` the free energy due to excited electrons is calculated from the total energy evaluated as a function of `sigma`. The total energy for several values of `sigma` must be put in directories called `restart2`, `restart3` ... `restart#nsigma`.

The input variables that control these plots are those described in the option `what='mur_lc'` and `what='mur_lc_disp'` in addition to the following:

```
grunmin_input : minimum y coordinate for the Gruneisen
                parameter plot.
                Default: real, calculated from the Gruneisen
                parameters.
grunmax_input : maximum y coordinate for the Gruneisen
                parameter plot.
                Default: real, calculated from the Gruneisen
                parameters.
volume_ph      : The frequencies and Gruneisen parameters inter-
                  plotted at this volume are plotted on a postscript
                  file. When volume_ph=0.0 the volume is calculated
                  from temp_ph. This option is available only for
                  cubic solids. Otherwise use celldm_ph.
                  Default: 0.0 (in (a.u.)**3)
celldm_ph      : The frequencies and Gruneisen parameters
                  interpolated at this crystal parameters are
                  plotted on a postscript file.
                  If this is 0.0 the celldm are calculated from
                  temp_ph. To have accurate Gruneisen parameters
                  and interpolated frequencies set the central
                  geometry as close as possible to celldm_ph. When
                  all nstep are odd, the central geometry is the one
                  given in the input of pw.x.
                  Default: 0.0 (celldm(1) in a.u., celldm(2-6)
                  dimensionless)
temp_ph        : The frequencies and Gruneisen parameters inter-
                  plotted at the volume (cubic systems) or at
                  celldm (anisotropic systems) that minimize the
                  free energy at this temperature are plotted on a
                  postscript file (only when volume_ph=0.0 or
                  celldm_ph(1)=0.0).
                  Default: real tmin (in K)
with_eigen     : if .TRUE. use the eigenvectors of the
                  dynamical matrix to calculate the Gruneisen
                  parameters used for anharmonic properties.
                  Could require a lot of RAM. Note however that
                  eigenvectors are always used to calculate
                  the plotted Gruneisen bands (both in cubic
```

and anisotropic solids).
Default: logical .FALSE.

ltherm_glob : when .FALSE. the vibrational (plus electronic) free energy is fitted by a polynomial, while only the static energy is fitted by an equation of state. When .TRUE. a different equation of state is fitted at each temperature.
Default: logical .FALSE.

poly_degree_ph : degree of the polynomial used to interpolate the vibrational free energy. Presently only the values 1, 2, 3, or 4 are available for anisotropic solids.
Default: integer 4

poly_degree_cv : degree of the polynomial used to interpolate the heat capacity. Presently only the values 1, 2, 3, or 4 are available for anisotropic solids.
Default: integer 4

poly_degree_bfact : degree of the polynomial used to interpolate the b factor. Presently only the values 1, 2, 3, or 4 are available for anisotropic solids.
Default: integer 4

poly_degree_elc : degree of the polynomial used to interpolate the elastic constants. Presently only the values 1, 2, 3, or 4 are available for anisotropic solids.
Default: integer 4

poly_degree_grun : degree of the polynomial used to interpolate the frequencies (Used only when lmurn=.TRUE. otherwise it is 2).
Default: integer 4

lv0_t : if .TRUE. the calculation of the thermal expansion with Gruneisen parameters uses the equilibrium geometry as a function of temperature computed from the free energy minimization, otherwise the equilibrium geometry at T=0 K. If reduced_grid=.TRUE. or both ltherm_freq=.FALSE. and ltherm_dos=.FALSE. the input geometry is used when lv0_t=.FALSE.
Default: logical .TRUE.

lb0_t : if .TRUE. the calculation of the thermal expansion with Gruneisen parameters uses the bulk modulus as a function of temperature computed from the free energy minimization, otherwise the bulk modulus computed at T=0 K (lmurn=.TRUE.). For lmurn=.FALSE. the code expects a single elastic constant file when lb0_t=.FALSE. and an

elastic constants file for each geometry when `lb0_t=.TRUE.`. Note that if `lb0_t=.FALSE.` and there are many elastic constants files the code use a constant bulk modulus computed with the elastic constants found in the file of the central geometry. If `lb0_t=.TRUE.` and there is a single elastic constants file all the quantities that depend on elastic properties are not computed. Default: logical `.TRUE.`.

`add_empirical` : If `.TRUE.` adds to the free energy an empirical term that can represent the anharmonic contribution or the electronic contribution. Default: `.FALSE.`.

`efe` : The type of empirical free energy
 1) $(\alpha_1 + \alpha_2 * V) T^2$
 2) $-2/3 k_B \ln \alpha_1 (v/v_0)^{\alpha_2} T^2$
 Default: 0 (must be explicitly given)

`alpha1` : parameter of the empirical free energy (see above) in eV/K² in 1), in 1/K in 2). Default: 0.0

`alpha2` : parameter of the empirical free energy (see above) in eV / K² / A³ in 1), adimensional in 2) Default: 0.0

`v0p` : parameter of the empirical free energy (equilibrium volume in (a.u.)³ Default: 0.0

`hot_electrons` : If `.TRUE.` the electronic free energy is computed from the energy as a function of smearing, otherwise it is read from files computed from electron dos. Must be used together with `lel_free_energy=.TRUE.`. Default: `.FALSE.`.

`nsigma` : The number of smearing values for which there are restart files. Note that `nsigma` includes the present restart, so the code expects to find `restart2`, `restart3`, ..., `restart#nsigma`. Default: 0 (option not used). Minimum value 3 to make a quadratic fit of the energy.

`sigma_ry(nsigma)`: the value of the smearing for each directory restart. In Ry units. Default: Must be set for each `nsigma` by the user.

`lhugoniot` : If `.TRUE.` the code plots `T(p)` and `V(p)` along the Hugoniot curve. Default: `.FALSE.`.

`flgrun` : file where the Gruneisen parameters are written. Default: character(len=*) 'output_pgrun.dat'

`flpgrun` : file where the Gruneisen parameters in a plotable

form are written.
Default: character(len=*) 'output_grun.dat'

flpsgrun : name of the postscript file with the Gruneisen parameters plot. The frequencies are written in a file with the same name plus the string _freq.
Default: character(len=*) 'output_grun'

flanhar : file where the anharmonic thermodynamic quantities are written.
Default: character(len=*) 'output_anhar.dat'

flpsanhar : postscript file of the anharmonic quantities.
Default: character(len=*) 'output_anhar'

fact_ngeo(1)...fact_ngeo(6) : With these factors the vibrational free energy is interpolated using a smaller number of geometries with respect to the total energy. The phonons are always calculated at geometry 1, then fact_ngeo(i)-1 geometries are not calculated and so on. The last calculated geometry must be ngeo(i). This happens when fact_ngeo(i) divides ngeo(i)-1. For even ngeo(i), fact_ngeo(i) must be 1. For odd ngeo(i) the following table gives a few examples

ngeo	fact_ngeo	calculated geometries
3	2	1,3
5	2	1,3,5
7	2	1,3,5,7
7	3	1,4,7
9	2	1,3,5,7,9
9	4	1,5,9
11	2	1,3,5,7,9,11
11	5	1,6,11

Default: integer 1,1,1,1,1,1
This option is not active when one of the ngeo_ph(i) is different from ngeo(i).

ngeo_ph(1),...,ngeo_ph(6) These variables are set to compute the phonon dispersions in a subset of the geometries used to compute the total energy. All values must be smaller than the corresponding ngeo and even or odd as the corresponding ngeo. step_ngeo remains the same for the two meshes. The following table gives a few examples:

ngeo	ngeo_ph	phonon calculated in geometries
5	3	2,3,4
6	2	3,4
6	4	2,3,4,5
7	3	3,4,5
7	5	2,3,4,5,6
9	3	4,5,6

```

          9          5          3,4,5,6,7
      Default: integer ngeo(1),...,ngeo(6)
reduced_grid: if .TRUE. the computed geometries are only along
one dimensional lines. So each parameter is varied
independently keeping the others fixed at the input
values. This option sets ltherm_freq=.FALSE.,
ltherm_dos=.FALSE., lv0_t=.FALSE. and lb0_t=.FALSE..
With this option the thermal expansion is
calculated only using the Gruneisen parameters
at the input geometry. The multidimensional fit
of the free energy is not done so this method
should be faster than the default one, but it
is less precise. This option is used only with
lmurn=.FALSE. and requires a file with the
elastic constants at the input geometry.
Default: logical .FALSE.
all_geometries_together: if .TRUE. all the phonon calculation
for all the geometries are used for the image
parallelization. To be used only if you have many
images (and CPUs) available.
Default: logical .FALSE.

```

The output files corresponding to different geometries can be identified by the presence of the letters `g1`, `g2`, ... in the filename. To exploit all the features of this option please write the dynamical matrices in `.xml` format (using a `fildyn` with the `.xml` extension). An example for this option can be found in `example09`.

Number of tasks for this option: Maximum between the number of tasks needed by the `what='mur_lc'` option and the number of tasks of the phonon code (see above the option `what='scf_ph'`).

When `all_geometries_together=.TRUE.`: number of tasks of the phonon code times the number of geometries.

3.20 `what='elastic_constants_geo'`

With this option the code can compute the elastic constants and elastic compliances as a function of temperature using the quasi-harmonic approximation. For the same geometries that are used to compute the elastic constants with the `elastic_algorithm='energy_std'` or `elastic_algorithm='energy'`, the code can compute the phonon dispersions and compute the elastic constants at each temperature as the second derivatives of the Helmholtz free energy with respect to strain. The second derivatives are corrected so that the stress-strain elastic constants are shown in the plots and in output. The temperature dependent elastic constants are calculated on a regular grid of unperturbed geometries, the same geometries chosen by the option `what='mur_lc'`, and written on separate files, one for each unperturbed geometry, inside the directory `anhar_files`. In order to plot the elastic constants as a function of temperature within the 'quasi-harmonic' approximation, it is necessary to make another calculation with `what='mur_lc_t'` having on files the elastic constants calculated for each geometry with the present option. In this case `thermo_pw` will be able to calculate the anharmonic properties using temperature dependent elastic constants and bulk moduli obtained by interpolating the "fixed-geometry quasi-harmonic" elastic constants computed by this option at the crystal parameters found at each temperature from the minimization of the free energy. The variables `fact_ngeo` and `ngeo_ph` are not available with this option. Using `start_geometry_qha` and `last_geometry_qha` it is possible to compute the temperature dependent elastic constants for selected or for a single unperturbed configuration. The use of `start_geometry` and `last_geometry` is also allowed but it refers to the global number of geometries necessary to compute the elastic constants in all the grid.

Since the calculation of the Helmholtz free energy derivatives is quite heavy, it has to be requested explicitly using the flag `use_free_energy=.TRUE..` By default, the code computes only the elastic constants at $T = 0$ K as second derivatives of the energy and writes them on files in the directory `elastic_constants`. A run of `thermo_pw` using `what='mur_lc_t'` having on files the $T = 0$ K elastic constants in the directory `elastic_constants` allows the calculation of the anharmonic properties using temperature dependent elastic constants and bulk moduli obtained by interpolating (within the "quasi-static approximation") the elastic constants computed by this option at the crystal parameters that, at each temperature, minimize the free energy. When both the $T = 0$ K and the temperature dependent elastic constants are on file in the directory `elastic_constants` and `anhar_files` respectively, the latter are used. When both `use_free_energy=.TRUE.` and `lel_free_energy=.TRUE.`, the electronic free energy is added to the free energy before computing the elastic constants. In this case the code expects to find on file (in `therm_files`) the electronic thermodynamic properties for each perturbed geometry. These files are produced with this same option and the flags `use_free_energy=.FALSE.` and `lel_free_energy=.TRUE..` In this case the code computes the electronic

thermodynamic properties at each perturbed geometry and writes them on file, without calculating the elastic constants. Note that the user must be careful to use the same value for the `lel_free_energy` flag with this option and in the following `mur_lc_t` calculation that interpolates the elastic constants. After computing the elastic constants at $T = 0$ K with the present option one can also run another `thermo_pw` calculation with the option `what='mur_lc'` computing the crystal parameters for a range of pressures (using `pmax` and `pmin` input variables). If the elastic constants are found in the directory `elastic_constants` they are interpolated at the pressure dependent crystal parameters and plotted on output. The variables that control this run are:

`use_free_energy` : when `.TRUE.` computes the elastic constants as second derivatives of the Helmholtz free energy with respect to strain. When `.FALSE.` the elastic constants are computed as second derivatives of the energy or using the stress-strain algorithms.
Default: `.FALSE.`

`start_geometry_qha` : Among the geometries considered by the option `mur_lc_t` the calculations of elastic constants are done starting from this geometry.
Default: integer 1

`last_geometry_qha` : Among the geometries considered by the option `mur_lc_t` the calculations of elastic constants are done only up to this geometry.
Default: integer total number of geometries.

An example for this option with `use_free_energy=.FALSE.` can be found in `example22` while an example with `use_free_energy=.TRUE.` can be found in `example23`.

Number of tasks for this option: The product of the number of tasks needed by the `what='scf_elastic_constants'` option and the number of geometries used with `what=mur_lc_t` when `use_free_energy=.FALSE.`.. When `use_free_energy=.TRUE.` and `all_geometries_together=.TRUE.` the number of tasks of the previous case is further multiplied by the number of tasks needed to compute a phonon dispersion (see above the option `what='scf_ph'`). When `use_free_energy=.TRUE.` and `all_geometries_together=.FALSE.` the number of tasks of this option is equal to the number of tasks needed to compute a phonon dispersion.

Restarting an interrupted run

There are several situations that might require the restart of the THERMO_PW code. We must distinguish two different cases: THERMO_PW stopped while running QUANTUM ESPRESSO routines, because the code reached the maximum cpu time or because some external event stopped the run, or THERMO_PW stopped after doing some post-processing task. This second case comprises also the normal termination of THERMO_PW and the necessity to change some details of the plot rerunning the post-processing tools without redoing the QUANTUM ESPRESSO calculations.

Support for the first case is based on the recover features provided by QUANTUM ESPRESSO routines and usually works when images are not used. This restarting method needs files in the `outdir` directory. In this case THERMO_PW behaves as QUANTUM ESPRESSO except for the fact that `max_seconds` in the input of `pw.x` or of `ph.x` is not active. To run `thermo_pw` for a fixed number of seconds `max_seconds` must be set in the THERMO_CONTROL namelist. If the code stopped inside `pw.x`, `restart_mode` must be set to 'restart' in the input of `pw.x` while if the code stopped inside `ph.x` routines `recover` must be set to `.TRUE.` in the input of `ph.x`.

When running `thermo_pw` with several images and calculating a phonon dispersion or using the `what='mur_lc_t'` option, `max_seconds` is controlled by the image driver of `thermo_pw`. Presently, after `max_seconds` a signal is sent to the asynchronous driver and the master stops sending new works to the images. It stops the code when all the images have terminated their current task. Recovering from this point is possible without losing any previous work by keeping the `outdir` directory and by setting `recover=.TRUE.` in the `ph.x` input. Note however that when `thermo_pw` is stopped by the operating system in an unclean way this restart method could not work.

As a last resource you can remove the `outdir` directory, and THERMO_PW will not recalculate the quantities contained in files that are already in the working directory. Completed phonon calculations at a given geometry for which the dynamical matrices are available are not redone if you put the `fildyn` name in the `thermo_control` namelist. It is possible to stop `thermo_pw` after the calculation of the phonon dispersions for a fixed number of geometries by setting the input variable `max_geometries` in the THERMO_CONTROL

namelist or also specify exactly which geometries to do in a given run using the variables `start_geometry` and `last_geometry` or `start_geometry_qha` and `last_geometry_qha`.

In general the restart of `thermo_pw` from a post-processing task is much easier. Each routine checks if a file with the same name as the file that it would produce is already in the working directory, and if this happens, it reads its content and returns. This feature cannot be disabled from input. In order to recalculate a given quantity, just remove the file that contains it from the working directory. For instance in an anharmonic calculation, if you have already all the dynamical matrices for all the geometries and you do not have any more the `outdir` directory, it is possible to skip entirely the phonon calculations and the reading of the files produced by `pw.x` by setting the variable `after_disp=.TRUE.` and giving the name of the dynamical matrices file using the variable `fildyn` in the `THERMO_CONTROL` namelist. In this case `THERMO_PW` can compute the anharmonic properties with a different set of temperatures, or with a different sampling on the phonon frequencies, etc.. You need to erase the output files that contain the phonon dos, or the thermal properties from a previous calculation, keeping the dynamical matrices files and the `restart` directory and rerun `THERMO_PW`. Similarly if the files containing the bands energy eigenvalues are already in the working directory, it is possible to set the input variable `only_bands_plot` to change the bands plot without redoing the bands calculation. Note however that in this case it is not possible to change the Brillouin zone path.

The following variables can be used to stop `THERMO_PW` before it concludes all the calculations:

`max_seconds`: the code stops after `max_seconds` have elapsed.
Note that the check is not done continuously so the clean stop might occur a few minutes after `max_seconds`.
Default: `real 10E8`

`max_geometries`: the code stops after computing the dispersions in `max_geometries`.
Default: `integer 1000000`

`start_geometry`: the code starts doing the phonons for `start_geometry`.
Default: `integer 1`

`last_geometry`: the code does only the phonons for geometries with index lower than `last_geometry`.
Default: `integer total number of geometries`.

Note that the first time that you use `start_geometry` and `last_geometry` the codes makes a self-consistent `pw.x` run for all the geometries and saves the results in the `outdir` directory. If for any reason the `outdir` directory is removed after computing some geometries, just remove also the `restart`

directory and the code will recreate the information in `outdir` but will not recalculate the dynamical matrices already available.

Finally we consider some typical runs of the most time consuming option `what='mur_lc_t'`, but what we say is valid also for the computation of the phonon dispersions at a single geometry. With a single processor or with a personal computer with a small number of processors you can run the code without interruption. In general in these cases it is not useful to use the image parallelization before exploiting all the parallelization levels of QUANTUM ESPRESSO, since images have an overhead due to the necessity of reinitialize the phonon calculation and recalculate the bands. If you cannot complete the calculation in a single run you need to send several times the THERMO_PW run. In this case you can set `start_geometry=last_geometry`, `max_seconds` in the `thermo_pw` input, and `recover=.TRUE.` in the `ph.x` input. You need to repeat this for all the geometries and finally you can collect the results and compute the anharmonic properties. Note that when `start_geometry` or `last_geometry` are set by the user the anharmonic calculation is skipped.

Thermo_pw on the GPU

Thermo_pw uses the QUANTUM ESPRESSO routines which are GPU aware, so if you compile with the flag `-D__CUDA`, GPU is active in thermo_pw. Starting from version 1.9.0 THERMO_PW contains also an experimental GPU version which improves on the QUANTUM ESPRESSO routines when the system is small and requires many **k**-points. This version which is activated by setting `many_k=.TRUE.` in the INPUT_THERMO namelist uses several experimental routines presently available only in thermo_pw. It has many limitations so it has to be used only for particular cases. It is implemented with Davidson diagonalization for LDA and GGA functionals. It is working for norm conserving, ultrasoft, and PAW pseudopotentials, both scalar and fully relativistic, but not for hybrid functionals or for LDA+U schemes. When the response to a phonon perturbation is calculated by thermo_pw with the flag `many_k=.TRUE.` the new routines are used, while the old ones are used for the other perturbations. The method is found to be useful only when FFT sizes are smaller than approximately $48 \times 48 \times 48$. It is not compatible with **G**-vectors parallelization, but it can be used with pools parallelization. It must be used with a number of MPI processes equal to the number of GPUs and with a number of pools equal to the number of MPI processes (one pool per GPU). It uses CUDA fortran so it is active only on NVIDIA GPUs. It has been tested on marconi100 at CINECA loading modules `hpc-sdk/2022-binary` or `hpc-sdk/2023-binary`. The `many_k` routines are active also on the CPU version of the code for testing purposes, but there is no advantage to use them with CPU.

`many_k`: If `.TRUE.`, the part of the code that loads on the GPU several wavefunctions and Hamiltonians is used.
Default: logical `.FALSE.`

`memgpu`: The available free memory in one GPU in GByte units.
(If the code crashes due to memory allocation problems, decrease this value).
Default: real 10.

Tools

The directory `tools` contains a few tools that can be useful to build structures of solids, surfaces, ribbons, and nanowires. Moreover it contains some miscellaneous codes that give additional information on the internal conventions of `THERMO_PW` or further process its output. Currently it contains the following programs:

- `average_grun.x` reads the thermal expansion, the isothermal bulk modulus, the volume, the isochoric heat capacity, and the temperature and computes the average Grüneisen parameter, the isobaric heat capacity, and the isoentropic bulk modulus.
- `bravais_lattices.x` tests the module `lattices.f90` of the library. Presently it can read three primitive lattice vectors of a Bravais lattice and find the `ibrav` code and the `celldm` parameters of the input lattice. It can also read two sets of primitive vectors and decide if they describe the same Bravais lattice. In the positive case it gives the orientation of one lattice with respect to the other. For an example of its use see `tools_input/bravais_lattice.in`.
- `change_dynmat_name.x` A simple tool to change the number of the geometry to a set of dynamical matrix files. For an example of its use see `tools_input/change_name.in`.
- `change_e_name.x` A simple tool to change the number of the geometry to a set of energy files inside the `restart` directory.
- `change_tel_name.x` A simple tool to change the number of the geometry to the electronic thermodynamic files inside the `therm_file` directory. It can be used also to change the names of the elastic constant files, both those in `elastic_constants` and in `anhar_files` directories.
- `crystal_point_group.x` is a crystal point group calculator. It can give several information about the crystallographic point groups, such as the list of symmetry operations, the product of two rotations, the product table, the class structure, the character tables of the irreducible representations of the point group and of the double point group and the projective

representations. It gives the list of subgroups and supergroups of a given group and the compatibility tables of a given group with its subgroups. It can also decompose the Kronecker product representations. Finally it can list the conjugate groups and calculate the intersection of two point groups. For an example of its use see `tools_input/crystal_point_group.in`.

- `debye.x` reads from input the Debye temperature and the number of atoms per unit cell and writes in a file the thermodynamic quantities (vibrational energy, free energy, entropy, and heat capacity) as a function of temperature computed using the Debye model. When the system has only one atomic type it writes the atomic B factor as a function of temperature computed using the Debye model. For an example of its use see `tools_input/debye.in`.
- `density.x` reads the volume of a unit cell of a solid and its mass (in a.m.u.) and computes the density, or reads the density of a solid and the mass (in a.m.u.) of a unit cell and computes the volume.
- `elastic.x` reads the elastic constants of a solid and computes the elastic compliances, the bulk modulus, and a few poly-crystalline averages. It uses the THERMO_PW library so the output is the same. For an example of its use see `tools_input/elastic.in`.
- `emp_f.x` reads a set of parameters for the construction of the Helmholtz free energy of a solid as a function of volume and temperature and prints the thermodynamic quantities. Presently it supports the models of Dorogokupets, Litasov, Mie-Gruneisen-Debye, and high temperature Birch-Murnaghan equations (still in experimental form).
- `emp_g.x` reads a set of parameters for the construction of the Gibbs free energy of a solid as a function of pressure and temperature and prints the thermodynamic quantities. Presently it supports only the model of Gustafson for tungsten.
- `epsilon_tpw.x` generalizes the routine `epsilon.f90` of the QUANTUM ESPRESSO distribution. It calculates the complex dielectric constant of a solid as a function of the frequency for independent electrons using the LDA or GGA eigenvalues. It is limited to insulators, but supports norm-conserving, ultrasoft, and PAW pseudopotentials. It supports both scalar relativistic and fully relativistic pseudopotentials and it uses the point group symmetry of the solid to reduce the number of $\mathbf{k}$ -points. For an example of its use see `example14`.
- `gener_nanowire.x` reads a two dimensional (2D) Bravais lattice index and atomic coordinates and generates a sheet of type (m, n) . A sheet contained between the two vectors $\mathbf{C} = m \mathbf{a}_1 + n \mathbf{a}_2$ and $\mathbf{T} = p \mathbf{a}_1 + q \mathbf{a}_2$ can be also generated and wrapped about $\mathbf{C}$ in a nanotube form ($\mathbf{a}_1$ and $\mathbf{a}_2$ are the primitive lattices of the 2D Bravais lattice). For lattices that allow it,

p and q can be determined automatically so that $\mathbf{T}$ is perpendicular to $\mathbf{C}$. For an example of its use see `tools_input/gener_nanowire.in`.

- `gener_2d_slab.x` reads a two dimensional Bravais lattice index and atomic coordinates and generates an infinite ribbon perpendicular to $\mathbf{G} = m\mathbf{b}_1 + n\mathbf{b}_2$, where $\mathbf{b}_1$ and $\mathbf{b}_2$ are the primitive reciprocal lattice vectors of the 2D Bravais lattice. The number of rows of the ribbon, and the number of atoms per row are given as input variables. For an example of its use see `tools_input/gener_2d_slab.in`.
- `gener_3d_slab.x` reads a three dimensional Bravais lattice index and atomic coordinates and generates an infinite slab perpendicular to $\mathbf{G} = m\mathbf{b}_1 + n\mathbf{b}_2 + o\mathbf{b}_3$, where $\mathbf{b}_1$, $\mathbf{b}_2$ and $\mathbf{b}_3$ are the primitive reciprocal lattice vectors of the Bravais lattice. The number of layers of each slab, and the size of the surface unit cell are given as input parameters. For an example of its use see `tools_input/gener_3d_slab.in`.
- `hex_trig.x` reads the values of a and c of the conventional hexagonal cell of a rhombohedral lattice (in Ångstrom), and gives as output the size a_r (in a.u.) and the cosine of the angle α of the rhombohedral cell. This information can be written in the input of `pw.x` for this type of cells. It is used to convert the structural information contained in a CIF file to the `pw.x` input.
- `kovalev.x` writes the correspondence between point group symmetry operations defined in the Kovalev tables and those used by QUANTUM ESPRESSO.
- `mag_point_group.x` gives a few information on the magnetic point groups.
- `merge_interp.x` This is a driver of the spline interpolation routines of QE. It can read the meshes and the functions to interpolate from two different files and provide the first function on the mesh of the second. In the data files lines that start with `#` are considered comments.
- `optical.x` contains a few utilities for optical properties calculations. It transforms a complex dielectric constant into a complex index of refraction and computes the reflectivity or the absorption coefficient for cubic system. It converts also from energy of the photon in eV to the frequency in Hz or the wavelength in nm.
- `pdec.x` reads the temperature dependent elastic constants files calculated for several pressures and makes a plot of the elastic constants as a function of pressure at several temperatures.
- `plot_sur_states.x` reads the dump file produced by THERMO_PW in a `what='scf_2d_bands'` calculation that contains the planar averages of all the states, and plots the states with the $\mathbf{k}$ point and the band numbers requested in input. For an example of its use see `tools_input/plot_sur_states.in`.

- `rotate_tensors.x` applies a rotation to a tensor of rank 1, 2, 3, or 4 defined in a coordinate system 1 and finds the form of the tensor in a new coordinate system 2.
- `space_groups.x` gives several information on space groups. It can give the names of the space group given the number reported in the International Tables for Crystallography (ITA), or the number given one of the names, translate the names between different editions of the ITA tables or the Shönflies name. It gives the list of coset representatives of each space group and the list of symmorphic space groups. For an example of its use see `tools_input/space_groups.in`.
- `supercell.x` reads a three dimensional Bravais lattice index and the atomic coordinates of the atoms inside a unit cell and produces a supercell with $n1 \times n2 \times n3$ cells of the original unit cell or a supercell delimited by three arbitrary Bravais lattice vectors given in crystal or cartesian coordinates. For centered cells there is the option to consider $n1$, $n2$, and $n3$ for the primitive or for the centered Bravais lattices. The input unit cell can be specified also by giving the space group and the coordinates of the nonequivalents atoms. It can be useful to study defects or to calculate all the atomic positions starting from the space-group and the nonequivalent positions. It is also possible to give the input Bravais lattice by using `ibrav=0` and the three principal vectors. In this case, before generating the supercell, the code rotates the Bravais lattice so that it has the same orientation of the vectors described in the `thermo.pdf` guide. For an example of its use see `tools_input/supercell.in`.
- `test_colors.x` produces a postscript file with the `gnuplot` colors that can be used in the plots.
- `test_eos.x` reads the parameters of an equation of state, and optionally the coefficients of a polynomial. Produces in output a file with the energy, the pressure, the bulk modulus, and its first and possibly second derivative with respect to pressure. It also checks the analytic results with those obtained by numerical finite differences writing on file the relative errors.
- `template.x` This is an example on how to write a tool code that can interface with the `QE` routines and all the library routines of `thermo_pw`. The template activates the parallelization options of `QE` and can use images.
- `translate.x` reads a set of atomic positions and a translation vector and translates the atomic positions. It can read also a rotation matrix and roto-translate the atomic positions.
- `units.x` writes on output the numerical constants used to write the guide `units.pdf` and `equilibrium.pdf`. It computes also the error associated to each conversion factor.

For a detailed description of the input variables please look at the beginning of the `fortran` sources of each code.

Examples, examples_qe, inputs, pseudo_test, space_groups, tools_inputs

The directories `examples`, `examples_qe`, `inputs`, `pseudo_tests`, `space_groups` and `tools_inputs` contain a set of examples that can be studied in order to learn how to use the THERMO_PW package. The `examples` directory contains inputs that run quickly but do not give converged results. These examples can be studied to see how the THERMO_PW code works in the different cases. The reference directory of each example contains all the output files produced by the run. A one-to-one comparison with the output produced by running the example script is however not possible due to the asynchronous nature of the runs. Only the plotted physical quantities should be the same.

The directory `examples_qe` is used by developers. It contains examples mostly imported from QUANTUM ESPRESSO that are used to check that QUANTUM ESPRESSO functionalities are not spoiled by `thermo_pw`.

The directory `inputs` contains a set of realistic inputs and reasonably converged results. Not all output files are reported in the reference directory of each run. The `inputs` examples are divided according to the structure type and many material properties are calculated for each structure. This directory can be seen as a gallery of the results that can be obtained by the THERMO_PW code, or as a source of information for the construction of a particular input geometry.

The directory `pseudo_test` contains a set of inputs that can be used to test a pseudopotential library. It illustrates how to use THERMO_PW for high-through-put calculations.

The directory `space_groups` contains a collection of structures ordered by the space group number. They are used to test the space groups routines. These inputs are also examples for the keyword `space_group` and for the use of Wyckoff positions to give the atomic coordinates in the `pw.x` input. You can also use these structures for your calculations, but note that the cut-off energies and the **k**-point meshes are not converged.

The directory `tools_inputs` give some examples of the inputs of the auxiliary

tools programs.

Color codes

In this section we briefly summarize the color codes of some of the figures that can be obtained from `thermo_pw`.

- Total energy versus kinetic energy. This is a figure of the total energy versus wave-functions kinetic energy cut-offs. When the test requires several charge density cut-offs there is a different curve for each charge density cut-off. The curve corresponding to the lowest charge density cut-off is `red`, the one corresponding to the highest is `blue`, all the others are `green`. Note that the total energy of the last configuration (highest wave function and charge density cut offs) is subtracted from all energies.
- Total energy versus size of the **k**-point mesh. This is a figure of the total energy as a function of the size of the **k**-point mesh. When the test requires several values of `degauss`, there is a different curve for each `degauss`. The curve corresponding to the first `degauss` is `red`, the one corresponding to the last is `blue`, all the others are `green`. Note that the total energy of the last configuration (highest number of points and lowest `degauss`) is subtracted from all energies.
- Total energy as a function of volume (`lmurn=.TRUE.`). This plot is composed of three figures: the total energy as a function of the volume, the pressure as a function of the volume and the enthalpy as a function of pressure. All curves are `red`. The points on the first curve are the energies calculated by `pw.x`, the continuous curve is the fit.
- Total energy as a function of one or two crystallographic parameters (`lmurn=.FALSE.`). When there is a single parameter the curve is `red` as in the case `lmurn=.TRUE.`. When there are two parameters a contour plot of the energy as a function of two parameters is shown. The contour levels, their number and their colors can be given in input. By default the code shows nine levels with three colors. From the lowest to the highest levels, the colors are `red`, `green`, and `blue`. The energy value of each level is written on output. When the user requests more levels without specifying their colors, the code continues with three `yellow` levels, then `pink`, `cyan`, `orange`, `black`, and when more than 24 levels are requested the

sequence of colors is repeated. When `lgeo_to_file=.TRUE.` the path written on file is shown in this plot with an `orange` points connected by a line. For orthorhombic solids the code produces many postscript figures, one for each value of c/a on the grid. In each figure there is a contour plot of the energy as a function of a and b/a . The colors of the levels follow the same conventions of the previous case. When the levels are chosen by the code the entire energy range (for all c/a) is divided into nine levels so each figure might have less than nine curves. For crystal systems with more crystallographic parameters, this figure is not available.

- Elastic constants (elastic compliances) as a function of pressure. The elastic constants (elastic compliances) are shown in different plots in red. In a final plot all the elastic constants (elastic compliances) are shown on the same figure in red, green, blue, yellow, pink, cyan, orange and black with same order of the previous plots. When there are more than eight elastic constants the colors are repeated.
- The crystal parameters and the volume as a function of pressure. When (`lmurn=.FALSE.`) the code plots the lattice parameters as a function of pressure, as well as the volume as a function of pressure (so far tested only for cubic cases). All plots are red.
- Energy bands. In this figure the bands have the color of their irreducible representation. Each line of the path can have a different point group and set of representations. See the `point_groups.pdf` file for the list of representations and their color code. When the symmetry analysis is not done all the bands are red.
- Energy bands with `enhance_plot=.TRUE.`. In this case the background color of the panels with lines at the zone border are gray, yellow, or pink. Gray means that the point group (or double point group) representations are used, yellow or pink means that a gauge transformation was applied and projective representations might have been used. A yellow background indicates that no switch from the point group to the double point group or viceversa was made, while a pink background means that such a switch was necessary.
- Electron density of states. This is a plot composed of two figures, the first contains the electron density of states, the second the integral of the density of states up to that energy. The dos is red. In the local spin density case, the dos for spin up is red the one for spin down is blue and with a negative sign. The integrated density of states is blue. In the spin polarized case, the curve shows the integral of the sum of the up and down density of states.
- Electronic energy, free energy, entropy, and isochoric heat capacity (metals only). This plot is composed of four pictures one for each quantity. There is a single blue curve per plot.

- Dielectric constant as a function of frequency ($q = 0$). There are two plots, one for the real part and one for the imaginary part. Other two plots contain the real and imaginary part of the complex index of refraction. For cubic solids other two plots show the reflectivity for normal incidence and the absorption coefficient. All curves are in red. For hexagonal, trigonal, and tetragonal systems the xx component is in red, while the zz component is in green. For orthorhombic systems the xx component is in red, the yy component in green and the zz component in blue. For monoclinic and triclinic systems the plot is not available.
- Inverse of the dielectric constant as a function of frequency ($q \neq 0$). There are four plots: the real and imaginary part of $\varepsilon(q, \omega)$ and the real and imaginary part of $1/\varepsilon(q, \omega)$. They are all in red. Note that the latter is really calculated, while the first is just its inverse.
- Phonon dispersions. In this figure the phonon dispersions have the color of their irreducible representations. The same comments made for the plot of the band structure apply here.
- Phonon dos. There is one picture with a single red curve.
- Vibrational energy, free energy, entropy, and isochoric heat capacity. This plot is composed of four figures each one showing one quantity. In red the quantities obtained using the phonon density of states, in blue those obtained from integration over the Brillouin zone. In some cases the red curve is not visible because it is exactly below the blue one.
- Atomic B factors as a function of temperature. This plot is composed of one figure for each atom for cubic solids and of two figures for each atom in the other cases. One figure contains B_{xx} (red, pink), B_{yy} (blue, light_blue) and B_{zz} (dark_green, green) as a function of temperature. The first color refers to quantities calculated from generalized phonon density of states while the second refers to quantities calculated by Brillouin zone integration. If the curves coincide, only the last one (green) will be visible. The second figure, when plotted shows B_{xy} (red, pink), B_{xz} (blue, light_blue), and B_{yz} (dark_green, green).
- Debye vibrational energy, free energy, entropy, and isochoric heat capacity. This plot is composed of four figures, one for each quantity. The curves are in blue and the word Debye appears in the y axis label.
- Crystal parameters as a function of pressure at several temperatures. Volume as a function of pressure at several temperatures. The number of plots depends on the crystal system. In these plots the first temperature is red, the others follow in the order green, blue, yellow, pink, cyan, orange, black. If there are more temperatures the sequence is repeated.
- Helmholtz free energy as a function of volume (`lmurn=.TRUE.`). The free energy calculated using the phonon dos (integral over the Brillouin zone)

is red (blue). When required in input this figure contains also the free energy as a function of volume for several temperatures. The color sequence red, green, blue, yellow, pink, cyan, orange, black indicates the different temperatures. In this case the same figure contains also the Gibbs energy as a function of pressure for several temperatures, the vibrational (plus electronic if available) free energy as a function of volume for several temperatures and the electronic free energy as a function of volume for several temperatures.

- The equilibrium volume as a function of temperature (`lmurn=.TRUE.`). The equilibrium volume obtained from the free energy calculated using the phonon dos (integral over the Brillouin zone) is red (blue). When required in input this figure contains also the volume as a function of temperature at several pressures. The color sequence red, green, blue, yellow, pink, cyan, orange, black indicates the different pressures. In this figure there is also the equilibrium volume divided by the equilibrium volume at $T = 300$ K (and the same pressure) is plotted as a function of temperature. When required in input this figure contains also the equilibrium volume as a function of pressure at several temperatures with the same color sequence. In this case also the equilibrium volume divided by the equilibrium volume at $T = 300$ K and zero pressure is plotted as a function of pressure for several temperatures.
- When requested in input, the pressure as a function of volume at several temperatures (`lmurn=.TRUE.`). The color sequence red, green, blue, yellow, pink, cyan, orange, black, indicates the different temperatures. In the same figure there is also the thermal pressure as a function of volume for several temperatures with the same color sequence and the thermal pressure as a function of temperature for several volumes with the same color sequence.
- The isothermal bulk modulus as a function of temperature (`lmurn=.TRUE.`). The isothermal bulk modulus obtained interpolating the free energy calculated using the phonon dos (integral over the Brillouin zone) is red (blue). When required in input this figure contains also the isothermal bulk modulus as a function of temperature at several pressures. The color sequence red, green, blue, yellow, pink, cyan, orange, black, indicates the different pressures. When required in input this figure contains also the isothermal bulk modulus as a function of pressure at several temperatures with the same color sequence. The same figure contains also the isoentropic bulk modulus as a function of temperature and the difference between isothermal and isoentropic bulk moduli. When required in input this figure contains also the isoentropic bulk modulus and the difference isoentropic-isothermal bulk moduli as a function of temperature for several pressures or as a function of pressure for several temperatures.
- Volume thermal expansion as a function of temperature (`lmurn=.TRUE.`).

The thermal expansion obtained from the free energy calculated using the phonon dos (integral over the Brillouin zone) is `red` (`blue`). The one obtained from the mode Grüneisen parameters is `green`. When required in input this figure contains also the thermal expansion as a function of temperature at several pressures. The color sequence `red`, `green`, `blue`, `yellow`, `pink`, `cyan`, `orange`, `black`, indicates the different pressures. When required in input this figure contains also the thermal expansion as a function of pressure at several temperatures with the same color sequence.

- The isochoric heat capacity as a function of temperature (`lmurn=.TRUE.`). The isochoric heat capacity calculated using the phonon dos (integral over the Brillouin zone) is `red` (`blue`). When required in input this figure contains also the isochoric heat capacity as a function of temperature for several pressures. The color sequence `red`, `green`, `blue`, `yellow`, `pink`, `cyan`, `orange`, `black`, indicates the different pressures. When required in input this figure contains also the isochoric heat capacity as a function of pressure at several temperatures with the same color sequence. The same figure contains also the isobaric heat capacity as a function of temperature and the difference isobaric-isochoric heat capacity. When required in input this figure contains also the isobaric heat capacity and the difference isobaric-isochoric heat capacity as a function of temperature for several pressures or as a function of pressure for several temperatures.
- Average Grüneisen parameter as a function of temperature (`lmurn=.TRUE.`). The parameter obtained from phonon dos (integral over the Brillouin zone) is `red` (`blue`). The one obtained from the mode Grüneisen parameters is `green`. When required in input this figure contains also the average Grüneisen parameter as a function of temperature at several pressures. The color sequence `red`, `green`, `blue`, `yellow`, `pink`, `cyan`, `orange`, `black`, indicates the different pressures. When required in input this figure contains also the average Grüneisen parameter as a function of pressure at several temperatures with the same color sequence.
- Crystallographic parameters, volume, Helmholtz (or Gibbs at finite pressure) free energy, thermal expansion tensor, volume thermal expansion, constant strain heat capacity (C_ϵ), isobaric heat capacity (C_P), difference $C_P - C_V$ of isobaric and isochoric heat capacities, difference $C_\sigma - C_\epsilon$ of constant stress and constant strain heat capacities (note that $C_\sigma = C_P$), difference $C_V - C_\epsilon$ of isochoric and constant strain heat capacities, difference $B_S - B_T$ of the isoentropic and isothermal bulk modulus, and average Grüneisen parameter as a function of temperature (`lmurn=.FALSE.`). The number of figures in this plot depends on the crystal system and on the presence of one or more files with the elastic constants. It shows a as a function of temperature for cubic solids, a , c/a , and c for tetragonal and hexagonal solids. For orthorhombic solids it shows also b/a and

b while for trigonal solids it shows a and $\cos \alpha$. For monoclinic solids it shows a , b/a , b , c/a , c , and $\cos \alpha$ (c-unique) or $\cos \beta$ (b-unique). All the six crystallographic parameters as a function of temperature are shown for triclinic solids. All quantities calculated using the phonon density of states are in red, those calculated integrating over the Brillouin zone are in blue with the exception of the thermal expansion tensor. When this tensor is diagonal with all identical components it follows the above rules while the tensor computed from mode Grüneisen parameters is in green. For hexagonal, tetragonal and trigonal solids α_{xx} follows the above rules while α_{zz} is pink, cyan, and orange when computed from phonon density of states, Brillouin zone integration, or mode Grüneisen parameters, respectively. In the orthorhombic case α_{xx} and α_{zz} have the same colors, while α_{yy} is gold, olive, and light-blue in the three cases, respectively. For the other crystal systems the thermal expansion tensor is not given. The thermal expansion tensor from the mode Grüneisen parameters is calculated only when the `elastic_constants` directory contains at least one file with the elastic constants. In this case also $C_P - C_V$, $C_\sigma - C_\epsilon$, $C_V - C_\epsilon$, and the average Grüneisen parameters are calculated using this thermal expansion tensor and plotted in green. The volume used in these calculations is the blue curve if `ltherm_freq=.TRUE.` or as in the red curve if `ltherm_freq=.FALSE.` and `ltherm_dos=.TRUE.`. When both `ltherm_freq=.FALSE.` and `ltherm_dos=.FALSE.` the volume is kept fixed at the equilibrium volume at $T = 0$ K. The same applies for the bulk modulus calculated from a single elastic constant file when the flag `lb0_t=.FALSE.` or computed within the “quasi-static” approximation when `lb0_t=.TRUE.`. The C_P , $C_\sigma - C_\epsilon$, $C_V - C_\epsilon$, and the average Grüneisen parameter are plotted only in presence of one or more elastic constants file.

- Thermal stresses as a function of temperature. This plot is composed of one figure in cubic solids and of two figures in the other cases. One figure contains b_{xx} (red, pink), b_{yy} (blue, light_blue) and b_{zz} (dark_green, green) as a function of temperature. The first color refers to quantities calculated from phonon density of states while the second refers to quantities calculated by Brillouin zone integration. If the curves coincide, only the last one (green) will be visible. The second figure, when plotted, shows b_{xy} (red, pink), b_{xz} (blue, light_blue), and b_{yz} (dark_green, green).
- Mode Grüneisen parameters. In this plot the mode Grüneisen parameters have the color of the irreducible representation of the phonon dispersion curve of which they are the derivative. The same comments made for the band structure plot apply here.
- Generalized average Grüneisen parameters as a function of temperature. This plot is composed of one figure in cubic solids and of two figures in the other cases. One figure contains γ_{xx} (red, pink), γ_{yy} (blue, light_blue) and γ_{zz} (dark_green, green) as a function of temperature. The first color

refers to quantities calculated from phonon density of states while the second color refers to quantities calculated by Brillouin zone integration. If the curves coincide, only the last one (green) will be visible. The second figure, when plotted shows γ_{xy} (red, pink), γ_{xz} (blue, light_blue), and γ_{yz} (dark_green, green).

- Phonon dispersions at the geometry that corresponds to a given temperature. The colors are assigned on the basis of the irreducible representation of each mode. The same comments made for the band structure plot apply here.
- Temperature dependence of the isothermal and isoentropic elastic constants within the “quasi-static”, “fixed geometry quasi-harmonic” or “quasi-harmonic” approximation. There is a plot for each non-zero elastic constant and a plot of the bulk modulus. The number of plots depends on the Laue class. Elastic constants interpolated at the geometry computed using the phonon density of states are in `red` (isothermal) and `green` (isoentropic), those calculated from integration over the Brillouin zone are in `blue` (isothermal) and `orange` (isoentropic).
- Temperature dependence of the isothermal and isoentropic elastic compliances within the “quasi-static”, “fixed geometry quasi-harmonic” or “quasi-harmonic” approximation. There is a plot for each non-zero elastic compliance and a plot of the compressibility. The number of plots depends on the Laue class. Elastic compliances interpolated at the geometry computed using the phonon density of states are in `red` (isothermal) and `green` (isoentropic), those calculated from integration over the Brillouin zone are in `blue` (isothermal) and `orange` (isoentropic).
- Temperature dependence of the isothermal elastic constants within the “fixed geometry quasi-harmonic” approximation for all the geometries of the mesh. There is a plot for each non-zero elastic constant and a plot of the bulk modulus. The number of plots depends on the Laue class. Elastic constants of the different geometries are in the sequence `red`, `green`, `blue`, `yellow`, `pink`, `cyan`, `orange`, `black`. When there are more than eight geometries the sequence is repeated. The same colors are used for the elastic constants obtained with the phonon density of states or from the integration over the Brillouin zone.
- Temperature dependence of the isothermal elastic compliances within the “fixed geometry quasi-harmonic approximation” for all the geometries of the mesh. There is a plot for each non-zero elastic compliance and a plot of the compressibility. The number of plots depends on the Laue class. Elastic compliances of the different geometries are in the sequence `red`, `green`, `blue`, `yellow`, `pink`, `cyan`, `orange`, `black`. When there are more than eight geometries the sequence is repeated. The same colors are used for the elastic compliances obtained with the phonon density of states or from the integration over the Brillouin zone.

- Anharmonic quantities as a function of temperature (pressure) plotted for several pressures (temperatures) chosen using `npres_plot` (`ntemp_plot`). Each pressure (temperature) is plotted with a different color, in the sequence red, green, blue, yellow, pink, cyan, orange, black from `press_plot(1)` to `press_plot(npres_plot)`. If there are more than eight pressures (temperatures) the sequence of colors is repeated.

Documentation

In addition to this user's guide, this directory contains the following documents:

- `tutorial.pdf`: a short guide that indicates where to find the information needed to compute a given quantity.
- `point_groups.pdf`: a description of the crystallographic point groups, character tables of the irreducible representations of point groups and of the double point groups and tables of the projective representations, for the interpretation of the color codes in the band and phonon dispersion plots.
- `thermo.pdf`: some notes on the thermodynamic expressions implemented in THERMO_PW.
- `developer_guide.pdf`: some notes on the internal logic of THERMO_PW.
- `unit.pdf`: some notes on the atomic units used in THERMO_PW.
- `equilibrium.pdf`: some notes on the atomic units used in THERMO_PW for the equilibrium thermodynamic tensors.