# cpw2000 User Guide

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José Luís Martins Departamento de Física, Instituto Superior Técnico, Lisboa, Portugal (Retired)

INESC MN, Lisboa, Portugal

jlmartins@inesc-mn.pt
jose.l.martins@tecnico.ulisboa.pt

# Contents

## 1 Introduction

This is the current version of a old code that calcultes the electronic structure of crystals within the pseudopotential approximation with a plane-wave basis.

The very original code was written by Sverre Froyen at the University of California Berkeley. I do not know what was the first time his version of the code, as there was already an earlier pseudopotential code at Berkeley, I would guess that Physical Review B, 26, 3258 (1983) https://journals.aps.org/prb/abstract/10.1103/PhysRevB.28.3258 already used that version.

After a long conversation with Roberto Car about the seminal Car-Parrinello work, I wrote at Berkeley in Marvin Cohen's group the code for the iterative diagonalization of the plane-wave matrices Physical Review B 37, 6134 (1988).

Since then many people contributed to the code. At Minnesota Norm Troullier and the group of Jim Chelikowsky contributed to the development. Renata Wentzcovitch implemented the original molecular dynamics and the variational cell shape molecular dynamis. Nadia Binggeli implemented the Langevin molecular dynamics. At Lisbon the largest contribution was from Carlos Loia Reis.

## 2 Installation

#### 2.1 Step 1: Downloading and extracting the archive

The code is available from GitHub

https://github.com/jlm785/cpw2000

If you downloaded the cpw2000-5.x.y.tar.gz file (where x and y are the minor version numbers) just extract it,

```
$ tar xzf cpw2000-5.x.y.tar.gz
```

and you will have a cpw2000-5.x.y directory

If you cloned the git you already have the relevant code in the main directory.

## 2.2 Step 2: Generating the documentation

The documentation is in the cpw2000-5.x.y/Doc directory. It include this file, and the means to generate a detailed description of the code for developers.

For that detailed description you need doxygen (https://www.doxygen.nl) and graphviz (https://graphviz.org/), both are available on most distributions. To check that your computer has doxygen and graphviz installed run the commands

```
cpw2000-5.x.y/Doc$ dpkg -s doxygen
```

cpw2000-5.x.y/Doc\$ dpkg -s graphviz

on a Debian based distribution (Ubuntu et al), or use equivalent tools (yum, rpm, dnf zypper,...) or the relevant GUI.

If they are not available just install them

```
cpw2000-5.x.y/Doc$ sudo apt install doxygen
```

cpw2000-5.x.y/Doc\$ sudo apt install graphviz

again for Debian based distros, or use equivalent tools (yum, rpm, dnf zypper,...) or the relevant GUI.

Finally run doxyfile in the doxy directory

cpw2000-5.x.y/Doc/doxy\$ doxygen Doxyfile

If you know what you are doing you can edit cpw2000-5.x.y/Doc/doxy/Doxyfile.

Finally it is useful to create a link to the file cpw2000-5.x.y/Doc/html/index.html

cpw2000-5.x.y/Doc\$ ln -s html/index.html documentation.html

Opening that link in your browser will allow you to see the documentation for (almost) every file in the code.

## 2.3 Step 3: Compiling the code

The code has been tested with several Fortran compilers, ifort (from Intel one API), gfortran (from gnu), pgfortran (from Portland), and even LLVM (experimental Intel compiler).

The choice of compiler is in the cpw2000-5.x.y/Src/make.inc file. You may edit it for your convenience, but in principle you have to make just two choices. The first is to identify which CPU you are using, as it narrows the choice of possible compilers. This is done by commenting/uncommenting the lines below # compiler for the job.

The second is the compiler you want, and is the key decision. Again comment/uncomment the lines below # Suggestions for compilers.

For any other modification of the make.inc file it is assumed you know what you are doing... The make.inc file was adapted from the Wannier90 code https://wannier.org/. As compilers may evolve with time, if you run into problems check the documentation and corresponding textttmake.inc file from that distribution. Another code that uses a make.inc file is ELK https://elk.sourceforge.io/. There you can find suggestions for compiler options.

To compile just go to cpw2000-5.x.y/Src and type make

```
cpw2000-5.x.y$ cd Src
```

cpw2000-5.x.y/Src\$ make

You should get the main library libcpw\_compiler.a and a few executables cpw\_compiler.exe, etc... where compiler is the name of the chosen compiler. As you probably are not interested in comparing compilers, it is advisable to create links without the compiler name. For example, after compiling with ifort, I usually create the links

```
cpw2000-5.x.y$ ln -s Src/cpw_ifort.exe cpw.exe
cpw2000-5.x.y$ ln -s Src/cpw_post_process_ifort.exe cpw_post_process.exe
cpw2000-5.x.y$ ln -s Src/libpw_ifort.a libpw.a
```

In the examples it will be assumed that those links are created, but obviously the links could be anywhere (in a bin folder for example).

## 3 First time run of the code

I assume you have a working directory separate from the source. In the following I will call it cpw2000-5.x.y/WORK

#### 3.0.1 Files required to run the code

You will need a file with the crystal description and a pseudopotential file for each type of atom in the crystal.

The file with the crystal description is called cpw.in. You must have such a file in your working directory. You can find such a file for almost all elements in cpw2000-5.x.y/Structures/Elements and files for other structures in the other subdirectories of textttcpw2000-5.x.y/Structures. The format of the crystal description is the same as in the SIESTA code, texttthttps://siesta-project.org/siesta/index.html so you may find descriptions for other crystals in the "net". Beware that presently the code only accepts fractional lattice coordinates.

There is also a tool to obtain a cpw.in file by answering a few questions.

You will have to generate the pseudopotentials for each atom, or find someone who has done that for you and have a file in the relevant format. The name of the file is Xy\_POTKB\_F.DAT where Xy is the one or two character chemical symbol. This file should be (or at least a link) on your working directory.

The good news is that the code to generate pseudopotentials is available from GitHub

```
https://github.com/jlm785/pseudopotential
```

and if you run the test on Validation after you followed the relevant instructions to install the code you will get pseudopotential files for all elements.

The bad news is that some of those pseudopotentials where not extensively tested, and even those that are reported as tested may not be what you want. Just test them in a simple case before you proceed. As pseudopotentials calculations should reproduce first principles calculations in cpw2000-5.x.y/Structures/Elements you can find elk.in files that you can use to run LAPW all-electron calculations with the ELK code, https://elk.sourceforge.io/. Testing the pseudopotential will spare you a lot of grief later.

If you want just a "quick and dirty" calculation than you can use the pseudopotentials from Validation of the pseudopotential code.

#### 3.1 Running the code

```
Once you have the required files you run the code with cpw2000-5.x.y/WORK$ ../cpw.exe if you created the appropriate links, or cpw2000-5.x.y/WORK$ ../Src/cpw_compiler.exe otherwise. The code will write to default output, so redirect it with cpw2000-5.x.y/WORK$ ../cpw.exe > output.dat or cpw2000-5.x.y/WORK$ ../cpw.exe | tee output.dat.

You will get a PW_RHO_V.DAT file with the self-consistent potential. By running cpw2000-5.x.y/WORK$ ../cpw_post_process.exe you will start an interactive analysis of the results.
```

If both executables ran without errors you are set.

## 4 The cpw.in file

The cpw.in syntax is similar to the input file from the SIESTA code. The parser is different, esdf instead of fdf. The keywords are even the same when possible. Therefore part of the text of this section was pilfered from the SIESTA documentation.

The cpw.in contains all the physical data of the system and the parameters of the simulation to be performed.

This file is written in a special format called ESDF, developed by Chris J. Pickard. This format allows data to be given in any order, or to be omitted in favor of default values. It is used in other electronic structure codes such as CASINO and PARSEC. However to be consistent with SIESTA (which uses the fdf format) the special character indicating a block structure has been changed.

Here we offer a glimpse of the rules:

- The syntax is a 'data label' followed by its value. Values that are not specified in the datafile are assigned a default value.
- The labels are case insensitive, and characters \_ . in a data label are ignored. Thus, LatticeConstant and lattice\_constant represent the same label.
- All text following the # character is taken as comment.
- Logical values can be specified as T, true, .true., yes, F, false, .false., no.
- Character strings should **not** be in apostrophes.
- Real values which represent a physical magnitude must be followed by its units. It is important to include a decimal point in a real number to distinguish it from an integer, in order to prevent ambiguities when mixing the types on the same input line.
- Complex data structures are called blocks and are placed between '%block label' and a '%endblock label' (without the quotes).
- If the same label is specified twice, the first one takes precedence.
- If a label is misspelled it will not be recognized (there is no internal list of "accepted" tags in the program). You can check the actual value used by CPW2000 by looking for the label in the output (by default standard output).

This is an example for silicon:

LatticeConstant	10.2629	bohr	
%block LatticeVectors			
0.0000000	0.50000000		0.50000000
0.5000000	0.00000000		0.50000000
0.5000000	0.50000000		0.00000000
%endblock LatticeVectors			
NumberOfSpecies	1		

NumberOfAtoms

%block Chemical\_Species\_Label

1 14 Si

2

%endblock Chemical\_Species\_Label

AtomicCoordinatesFormat Fractional

%block AtomicCoordinatesAndAtomicSpecies

0.12500000 0.12500000 0.12500000 1 # Si 1 -0.12500000 -0.12500000 -0.12500000 1 # Si 1

%endblock AtomicCoordinatesAndAtomicSpecies

StructureSource Experiment

#-----

# Energy cutoff, bands, and Brillouin mesh

#-----

PWEnergyCutoff 12.0000 hartree

NumberOfEigenStates 10

%block kgrid\_Monkhorst\_Pack

 4
 0
 0
 0.500000

 0
 4
 0
 0.500000

 0
 0
 4
 0.500000

%endblock kgrid\_Monkhorst\_Pack

#### 4.1 General stuff

#### • SystemLabel

A *single* word (max. 20 characters *without blanks*) containing a nickname of the system. Reserved to be used to name output files in the future.

#### • PrintingLevel (integer)

Defines the detail of the printout. May take the values 1,2, or 3.

Default value: 1 (for molecular dynamics) or 3 (single geometry).

The higher the value the more details will be printed. Remember that too much information is noise. 1 is recommended for molecular dynamics. 3 for single calculation, or when things seem to go wrong.

#### • MD.TypeOfRun (string)

Choice on how the atoms move. Molecular dynamics or geometry optimization.

Default value: ONE

With the default it will run a single geometry. See below for the other options to run molecular dynamics calculations.

The value in cpw.in may be overridden by the first argument of the executable.

cpw2000-5.x.y/WORK\$ ../cpw.exe MICRO > output.dat

will perform a microcanonic molecular dynamics calculation irrespective of the value in cpw.in.

## 4.2 Crystal descriptors

These are the lines that describe the crytal structure.

## • LatticeConstant (length)

A physical value, requiring a real number followed by the units. Accepted units are: bohr, ang, nm, m. This is just to define the scale of the lattice vectors.

Default value: 1.0 bohr.

The code stops if the units are not present or not recognized. Internally the code uses atomic units (bohr).

#### • LatticeVectors (block)

The cell vectors are read in units of the lattice constant defined above. They are read as a matrix, each vector being one line.

Default value: Unit matrix.

The internal representation of the lattice vectors in the code is by their metric tensor. So the original orientation is space is lost! When the code needs to read or print information with orientation content it will use some *canonic* vectors based on the lattice symmetry. Those are printed at the start of the SCF code, and before questions to the user in the post-processing code. Just pay attention, they may be **different** from what is written in cpw.in! See the section on output for more details.

The code checks if the entered value is near a rational number or the square root of a rational number (with low denominators). If it is the case it will use that modified value and print a warning. This allows to have an exact symmetry, and the modifications are always small.

### • NumberOfSpecies (integer)

Number of different atomic species in the simulation. It must be the number of lines in the **ChemicalSpeciesLabel** block.

Default value: 1

If not present or nin positive it will use the number of lines in **ChemicalSpeciesLabel**. In case of inconsistency the execution terminates.

#### • NumberOfAtoms (integer)

Number of atoms in the simulation. It is the number of lines in the block **AtomicCoordinatesAndAtomicSpecies**.

Default value: 1

In case of inconsistency the execution terminates.

#### • ChemicalSpeciesLabel (block)

It specifies the different chemical species that are present, assigning them a number for further identification. cpw2000 recognizes the different atoms by the given atomic number.

One line for each species. The first number in a line is the species number, it is followed by the atomic number, and then by the chemical symbol. From H to Og all chemical symbols are recognized. There is an extra chemical symbol, ZZ with a number of protons (atomic number) that can be zero to allow adding special pseudopotentials to the crystal.

In case of inconsistency the execution terminates.

## • AtomicCoordinatesFormat (string)

Character string to specify the format of the atomic positions in input. It is not used!!! It is here for future use and compatibility with SIESTA.

#### • AtomicCoordinatesAndAtomicSpecies (block)

Block specifying the position and species of each atom. One line per atom, with three reals followed by one integer. In total the number of lines indicated by **NumberOfAtoms** must be present.

The three reals indicate the atomic positions in **FRACTIONAL lattice coordinates** followed by the species of atom on that position, as identified in the **Chemical\_Species\_Label** block.

In case of inconsistency the execution terminates.

#### • StructureSource (string)

Information on the source of the crytal structure. Not used by the code,

## 4.3 Major Self Consistent Field parameters

This are the parameters whose values you would mention in a paper to allow reproducibility.

#### • PWEnergyCutoff (energy)

Energy cutoff of the plane wave basis set expansion. Real value followed by the energy unit. Accepted units are: eV, Ry, Hartree, and a few other.

Default value: 5 Hartree

Internally the code uses Hartrees.

#### • NumberOfEigenStates (integer)

This parameter indicates the number of eigenstates to be calculated.

Default value: 10

In some future the default should be slightly larger than half the number of electrons.

By choosing a low value the cost of the diagonalization may be reduced by finding fewer eigenstates. However choosing a slightly larger number of active eigenstates than the bare minimum may help converge faster the occupied eigenstates and therefore the overall calculation. Note, that if the electronic temperature is greater than zero then the number of partially occupied states increases, depending on the band gap. The value specified must be greater than the number of occupied states (at least the number of electrons divided by two for a non-spin-polarized calculation) and less than the number of basis functions (which is extremly large for plane waves).

#### • kgridMonkhorstPack (block)

Specifies the Fourier integration grid, known in the literature as the Monkhorst-Pack grid, for the Brillouin zone integration. It is just the good old Gauss quadrature method with sine/cosine functions chosen for periodic functions.

Default value: The  $4 \times 4 \times 4$  sampling with 1/2 shift.

Specified as an integer  $3 \times 3$  matrix and a real vector. At present only the diagonal elements of the matrix are relevant, but this format will allow an extension of the method in the future and better compatibility with SIESTA.

It has three lines, each with three integers and a real.

$$m_{11}$$
  $m_{12}$   $m_{13}$   $d_1$   $m_{21}$   $m_{22}$   $m_{23}$   $d_2$   $m_{31}$   $m_{32}$   $m_{33}$   $d_3$ 

In the direction j the Brillouin zone will be divided in  $m_{jj}$  sections and a point with a shift of  $d_j$  will be selected.

To use only the  $\Gamma$  point (molecule in a supercell) use

It is usual to have  $m_{jj}$  an even number and  $d_j = 0.5$ . For hexagonal crystals and the conventional axis, it is also usual/convenient to have  $m_{11} = m_{22}$  a multiple of three. These are recipes that minimize the number of irreducible points and avoid breaking symmetry.

If the diagonal elements are non-positive it will default to  $m_{ii} = 1$ , printing a warning. It also warns the presence of unused non-zero off-diagonal values.

#### • XC.Authors (string)

Particular parametrization of the exchange-correlation functional.

Default value: CA

#### - CA

Local density approximation (LDA). Quantum Monte Carlo calculation of the homogeneous electron gas by D. M. Ceperley and B. J. Alder, Phys. Rev. Lett. **45**,566 (1980), as parametrized by J. P. Perdew and A. Zunger, Phys. Rev B **23**, 5075 (1981)

- PBE
  - GGA of J. P. Perdew, K. Burke and M. Ernzerhof, Phys. Rev. Lett. 77, 3865 (1996)
- TBL
   Meta-GGA of Tran and Blaha. F. Tran and P. Blaha, Phys. Rev. Lett. 102, 226401 (2009)

#### • Xc.TBL.C (real)

Sets Tran-Blaha constant if a positive value. If it is negative, the original constant is used. A value around 1.09 is usually a good choice. It can be used to "fine-tune" the band gap in simulations (second-priciples calculations).

Default value: 1.0

#### • TypeOfScfDiag (string)

Indicates how the SCF was performed.

#### - PW

The full plane-wave basis set. It is the "usual" or traditional choice.

#### - AO

Uses the atomic orbitals included in the pseudopotential file. Not available in very old files. It corresponds to a Linear Combination of Atomic Orbitals (LCAO) calculation. It is very fast, but with limited accuracy. Use for *exploratory* runs on complex structures.

#### - AOJC

Uses the atomic orbitals included in the pseudopotential file, but improves the LCAO wave-functions with a single Jacobian Correction. Slower than the AO option but still quite fast. It is very convenient to explore the Born-Oppenheimer energy surface in optimization and molecular dynamics. However one may want to check the *final* results with a full PW calculation.

#### - AOJCPW

Follows an AOJC calculation with a full PW calculation. Final results are the practically the same as the normal PW calculation, but may be faster. Check if that is the case for your crystals before using..

#### • DualApproximation (boolean)

Use the dual approximation.

Default value: .TRUE.

It uses a smaller grid for the calculation of the effective potential. Speeds up the calculations with a compromise on precision. For molecular dynamics it is very safe. For geometry optimization with small energy cutoffs may not be accurate enough.

## • ElectronicTemperature (temperature)

A real value followed by unit. The normal unit is the Kelvin, k, but will accept other energy units, for example meV

Default value: 0 K

Occupy orbitals with a Fermi-Dirac distribution with that temperature.

#### 4.4 Minor Self Consistent Field parameters

Other parameters of self-consistency that have less impact, unless "very wrong" choices are made. Probable safe to leave at default values.

#### • MaxSCFIterations (integer)

Maximum number of self-consistent iterations.

Default value: 30

If very large, computing time can be wasted in the cases convergence is not achieved. If very small the code can terminate without a result, also wasting computer time.

#### • ScfTolerance (real)

Convergence parameter (atomic units/Hartree) for self-consistency

Default value: 0.00001

The SCF iterations are deemed converged if the difference between the input and output values of all the components of the Fourier transform of the effective potential are smaller than this treshold.

For exploratory runs decrease this parameter.

## • MaxDiagIterations (integer)

Maximum number of Ritz steps in the iterative diagonalization subroutine.

Default value: 40

Increase if in the last few SCF iterations you still see warnings about noisy diagonalization. In early SCF iterations, those warnings disappear if you increase this parameter, but computing time will also increase without changing final results.

## • DiagTolerance (real)

Criteria for iterative diagonalization convergence.

Default value: 0.00001

The iterative diagonalization is deemed converged if the module of the error vector is smaller than that value. For exploratory runs you can use the value of 0.001.

#### 4.5 Unused Self Consistent Field parameters

Over the years only one option was left in the code.

## • TypeOfPseudoMixing (string)

Default value: BFGS

Type of effective potential mixing used to accelerate convergence. Broyden-Fletcher-Goldfarb-Shanno method.

Broyden 1st (BROYD1) is available in older codes.

#### • TypeOfPseudopotential

Type of pseudopotential that is used.

Default value: PSEUKB

Kleinman-Bylander separable pseudos are the only available option. In older versions, the non-separable pseudo was available (PSEUDO), as well as a gaussian integration separable pseudopotential (PSEUGA). Some subroutines and variables still keep the ga characters from those days.

#### 4.6 Molecular dynamics parameters

## • MD.TypeOfRun (string)

Choice on how the atoms move. Molecular dynamics or geometry optimization.

Default value: ONE

#### - ONE

Just one SCF calculation. Atoms do not move.

#### - RSTRT

Restart the molecular dynamics from the last atomic configuration. Use in case it was interrupted by some external reason (power failure). Just keep all the other parameters the same. It will reproduce what a non-interrupted calculation would find in the case of molecular dynamics. For optimization the minimization restarts, so it will not reproduce a non-interrupted calculation, but probably will converge to the same minimum.

#### MICRO

Microcanonic molecular dynamics. The total/free energy is conserved, depending whether the temperature is zero and the system is an insulator or temperature is finite and not too low in the case of a metal. How low the temperature can go depends on the density of integration k-points.

#### - LANG

Langevin molecular dynamics, in contact with a thermostat with temperature indicated by MD.TargetTemperature.

#### - VCSLNG

Langevin molecular dynamics but with a variational cell shape, that is the lattice vectors change with simulation time.

#### - LBFSYM

Minimization of the energy with respect to atomic positions with the LBF algorithm.

#### VCSLBF

Minimization of the energy with respect to atomic positions and cell shape with the LBF algorithm.

#### - EPILBF

Minimization of the energy with respect to atomic positions and "vertical" dimension of the cell with the LBF algorithm. It models an epitaxial situation. The initial  $\vec{a}_1$  and  $\vec{a}_2$  of **LatticeVectors** define the epitaxial surface. The cell only contracts and expands in the direction perpendicular to those vectors  $(\vec{b}_3)$ .

#### • MD.InitialTemperature (temperature)

A real value followed by unit. The normal unit is the Kelvin, k, but will accept other energy units, for example  $\mathtt{meV}$ 

Default value: 1000 K

The initial kinetic energy is set according to the temperature. If the initial potential energy is high, the temperature of the system may be quite higher. If unsure, first do a thermalization with Langevin, otherwise the system may break apart. Simulations with **TypeOfScfDiag** as AO or AOJC are great for fast thermalization.

#### • MD.NumberOfSteps (integer)

Number of steps of the molecular dynamics run or maximum number of optimization steps.

Default value: 100

#### • MD.LengthTimeStep (time)

A real value followed by unit. The usual unit are femtoseconds, fs, but will accept other time units s, ns, ps, autime.

Default value: 2.418884 fs = 100.0 autime

Time step for molecular dynamics. If ut is too small atoms barely move, if too large the integrator of the molecular dynamics (Verlet) becomes unstable. Instability appears in the non-conservation of energy. Few femtoseconds should be OK, but remember that light atoms move faster (square root of mass scaling). Simulations with **TypeOfScfDiag** as AO or AOJC are great for initial checks.

#### • MD.TargetTemperature (temperature)

A real value followed by unit. The normal unit is the Kelvin, k, but will accept other energy units, for example meV

Default value: 300 K

For molecular dynamics with a Langevin thermostat, it is the temperature of the thermal bath.

## • MD.FrictionFracInvTimeStep (real)

The friction coefficient for simulations with a Langevin thermostat.

Default value: 20.0

It is set with respect to the time step. It indicates how many time steps occur until some thermalization is achieved. If small the thermalization is of bad quality, if large it will take a long time to thermalize.

#### • MD.TargetPressure (pressure)

A physical value, requiring a real value followed by unit. Accepted units are: GPa, Pa, atm, Mbar, bar, MPa.

Default value: 0 GPa

In a molecular dynamics with variational cell shape, or cell optimization, it is the applied pressure.

## • MD.TargetStress (block)

Real  $3 \times 3$  matrix of an applied stress in addition to the applied pressure. The units are in GPa only.

Default value: 0

The matrix should be symmetric, and is in lattice coordinates, so some effort is needed to understand the orientation of the crystal.

#### • MD.CellMass (real)

Fictitious cell mass for variational cell shape molecular dynamics. It is in units of electron mass.

Default value: 10.0

Should be fairly larger than one for the cell dynamics to be slow, but not so large that the cell dynamics is too slow. Simulations with **TypeOfScfDiag** as AO or AOJC are great for initial checks.

## • MD.Seed (integer)

Seed for the pseudo-random generator for the thermostat and initial velocity.

Default value: 87697

Same seed will give the same trajectory. Different seeds will give different trajectories allowing trivial parallelization of simulations.

## • MD.UseKeatingCorrections (boolean)

Default value: .false.

For some tetrahedral semiconductors, it provides a correction to LDA that reproduces experimental bond lengths. Do not use with heavy elements (Sn). If the structure is not tetrahedral the code stops.

#### • MD.UseFixedkplusG (boolean)

Default value: .false.

Keep a fixed  $\vec{k} + \vec{G}$  expansion in a variational cell shape simulation to avoid the noise of the basis set changing during the simulation. Should only be used near equilibrium. abinit has a neater way of dealing with this noise, the code is prepared to use that trick, but it has not been implemented. Use **MD.CG.FixedkplusGTol** to define when it kicks in.

#### • MD.CG.Tolerance (force)

A physical value, requiring a real value followed by unit. The normal and internal unit is har/bohr, but will accept N, eV/ang, Ry/bohr.

Default value: 0.0001 har/bohr

A geometry optimization run stops when the components of the forces on all atoms are smaller than this value.

#### • MD.CG.StepMax (length)

A physical value, requiring a real value followed by unit. Accepted units are: bohr, ang, nm, m.

Default value: 0.01 bohr

Maximum displacement of atoms to avoid instabilities in the optimization.

#### • MD.CG.FixedkplusGTol (force)

A physical value, requiring a real value followed by unit. The normal and internal unit is har/bohr, but will accept N, eV/ang, Ry/bohr.

Default value: 0.01 har/bohr

The fixed  $\vec{k} + \vec{G}$  is used after the components of the forces (har/bohr) on all atoms are smaller than this value.

## 4.7 Symmetry

Treatment of symmetry during a molecular dynamics or optimization run.

## • UseSymmetry (boolean)

Default value: .TRUE. for optimization and single runs, .FALSE. for molecular dynamics runs.

Try to maintain the initial symmetry of the system. It is close to 100% fiable (as close as real number logic allows).

## • SymmTolerance (real)

Tolerance for symmetry identification.

Default value: 0.00001

If after a symmetry operation the difference between the positions (in lattice coordinates) is smaller than this value, the atoms are considered superposed.

## 4.8 Unfolding

If the structure is a supercell of some lattice it may be useful to plot the band structure in the "unfolded" Brillouin zone of the parent structure. The unfolding procedure was developed within an industrial collaboration (rede project), it was used in semiconductors with a parent fcc lattice, so unfolding was only extensively tested in that lattice.

#### • Rede.Superlattice (block)

The block has three lines with three integers. Each line indicates how each of the supercell lattice vectors relate to the lattice vectors of the parent structure.

Taking as an example the supercell described in Structures/Supercells/cpw\_Si6Ge6\_Si(001).in, which has 6 Si atomic layers and 6 germanium atomic layers on a (001) surface, the block is

%block Rede.Superlattice

%endblock Rede.Superlattice

the underlying parent fcc structure has lattice vectors  $\vec{A}_1 = (0, a/2, a/2)$ ,  $\vec{A}_2 = (a/2, 0, a/2)$  and  $\vec{A}_3 = (a/2, a/2, 0)$ . The supercell vectors are  $\vec{a}_1 = -\vec{A}_1 + \vec{A}_2 = (-a/2, a/2, 0)$ ,  $\vec{a}_2 = \vec{A}_3 = (a/2, a/2, 0)$  and  $\vec{a}_3 = 3\vec{A}_1 + 3\vec{A}_2 - 3\vec{A}_3 = (0, 0, 3a)$ . Notice that the in-plane vectors have been rotated by 45°, their length is  $\sqrt{2}a$  as the lattice constant is that of Si, it is consistent with epitaxy on Si (001). The third vector has a length slightly larger than 3a because it was allowed to relax in the presence of the larger Ge atoms.

#### • Rede.NumberOfLatticePlanes

In a supercell grown on a surface indicates the number of lattice planes. It is used to help in plots to determine band alignments. In the above example you have six lattice planes (but twelve atomic planes). It must be present for unfolding to work. That is an "hack" to deal with back compatibility of some files, but do not influence the unfolding. This "requirement" should be removed in future releases.

# 5 Default output

The code writes to the default output, which can be redirected to a file. Depending on the choice of **PrintingLevel** the output will be different. We will use as a first example the file Validation/Output/pw\_ref\_01.out.

#### 5.1 Initial information

On the top of the file we have the information about the version of code, and the date and time it was run.

```
density-functional pseudopotential plane-wave program version 4.99 run on the 19-Mar-21 at 13:30:54
```

notice that care was taken to code back the year 2000 bug in the 1990s.

Continuing with that example we find next some information about how many chemical species, and maximum number of atoms per chemical species the code is using.

The values set by size\_mxdtyp\_mxdatm\_esdf are:

```
The value of mxdtyp is: 2
The value of mxdatm is: 8
input read from cpw.in
```

notice that the information about which subroutine printed the information is included in this case.

Next information are the lattice vectors,

#### Primitive Translation Vectors

		in a.u.	in	lattice u	nits	
a1=	0.125697E+02	0.000000E+00	0.000000E+00	1.225	0.000	0.000
a2=	-0.418990E+01	0.118508E+02	0.000000E+00	-0.408	1.155	0.000
a3=	0.00000E+00	0.00000E+00	0.725711E+01	0.000	0.000	0.707

preceded by a warning that they were changed,

```
WARNING: avec(3, 3) changed from 0.707106780000 to 0.707106781187
```

informing that the code recognized a value very close to  $\sqrt{2}/2$  and changed it.

The line

```
Single geometry calculation
```

informs that MD.TypeOfRun was ONE.

To be reproducible the information needed to generate the pseudopotentials is printed

## potentials :

-----

Si ca nrl nc 19-SEP-18 atom 5.804 Improved Troullier - Martinskb-loc= 2 3s( 2.00) rc= 1.993p( 2.00) rc= 1.993d( 0.00) rc= 1.99 nql=4000 delql= 0.0100

see the user guide of the pseudopotential generation code for details.

Next the other chosen or default parameters are printed

Local Density Approximation (LDA) using Ceperley and Alder as parametrized by Perdew and Zunger

dual aproximation is used

The energy cutoff for wave-function kinetic energy is 10.000 Hartree

SCF is converged when the difference in potentials is less then 0.00001000 Iterative diagonalization converged when the error in |h| psi - e psi| is less then

The temperature for electron Fermi distribution is 0.000Kelvin

Plane-wave basis calculation

see previous section for the keywords associated.

## 5.2 Crystal data

In the output the initial crystal data is printed,

CRYSTAL DATA

1081.02690815 volume

real-space metric

157.99703		2.66567731 7.99703193	0.000000 0.000000 52.665677	00	metric metric	g22,g2	:3	
12.56968703	12.56	968703	7.25711219	length	1,2,3	(a.u.)		
109.47122063	109.47122063 90.00000000		90.00000000 angle 12,13,23 (degrees)					
position	(lattice c	oord.)	position (	cartesia	an cooi	rd. a.u.)	no.	type
0.0000	0.50000 0.62500	0.00000	0.72571E+01 0.36286E+01	0.0000		0.00000E+00 0.00000E+00	1 2	Si Si

posi posi

posi	Si	3	0.00000E+00	0.00000E+00	0.0000E+00	0.00000	0.00000	0.00000
posi	Si	4	0.00000E+00	-0.25658E+01	0.36286E+01	0.00000	0.12500	0.37500
posi	Si	5	0.36286E+01	0.51316E+01	0.36286E+01	0.50000	0.50000	0.00000
posi	Ge	6	0.36286E+01	-0.51316E+01	0.36286E+01	0.50000	0.00000	0.50000
posi	Ge	7	0.36286E+01	0.25658E+01	0.72571E+01	0.50000	0.62500	0.37500
posi	Ge	8	0.36286E+01	0.25658E+01	0.00000E+00	0.50000	0.12500	-0.12500

in what should be self-explanatory.

Here we encounter a characteristic of the output, in that we have a value 1081.02690815 followed by a "key" volume. This allows a quick search for that key in the output. For example cpw2000-5.x.y/WORK\$ grep volume ../Validation/Output/pw\_ref\_05.out will return the volumes encoutered in Langevin a molecular dynamics with variational cell shape

843.03621271	volume
844.20060174	volume
847.56015618	volume
852.95790991	volume
860.21678359	volume
869.14551064	volume
879.54271598	volume
891.20293979	volume
1092.59744351	volume
1091.08256914	volume
1089.72581021	volume
1088.52562153	volume
1087.47977919	volume
1086.58556009	volume
1085.84052430	volume
1085.24216798	volume

in a form that would be trivial to plot (for example with gnuplot).

#### 5.3 Crystal symmetry and reciprocal space info

The code recognizes the space group operations, although does not find its conventional name (we will later describe how to find it), it lists the symmetry operations, in lattice coordinates, first the matrix and than the eventual fractional coordinate.

#### rotation matrices and fractional translations in lattice coordinates

1	1 0	0	0	1	0	0	0	1	0.000000000 0.0000	0.00000
2	0 -1	0	-1	0	0	0	0	-1	-0.500000000 -0.5000	0.00000
3	0 -1	0	-1	0	0	0	0	1	-0.500000000 -0.5000	0.00000
4	1 0	Ω	0	1	Ο	0	Ω	-1	0 000000000 0 0000	000000 0 000000

we have 4 symmetry operations, the first is the identity and the last is the reflection with respect to the "z" axis. Remember that the code only uses the metric, and therefore the z-axis, which

in this case is "obvious" because angles 13 and 23 are 90 degrees. In the general case it may be more complicated.

Next the output informs that there are 13117 G-vectors, but only 1868 are unrelated by spatial or temporal symmetry, and some information about the maximum values of  $g_i$  encountered in the expansion  $\vec{G} = \sum_{i=1,3} g_i \vec{b}_i$  for the potential and charge densities.

```
13117 G-vectors are set up in 1868 stars, kmax = 17 17 10
```

It also informs that of the  $4 \times 4 \times 4$  integration grid in reciprocal space, there are 12 unrelated by symmetry, and that it will calculate 24 orbitals

```
12 k-points generated by program from parameters : 
 n = 4 4 4 s = 0.50 0.50 0.50 nb = 24
```

Computing time for starting (s): 0.01

## 5.4 The SCF cycle

The first information about the SCF cycle is the FFT grid size and the maximum and minimum values of the local self-consistent potential.

```
in fft for local potential n = 24 24 16 max and min of potential (Hartree) 0.1915 -4.8485 0.0000
```

In the first iterations one may encounter warnings of the type

```
WARNING WARNING: After h_kb_dia

The estimated error in energy has an accuracy
of 7.3 digits
```

First, 7 digit accuracy is already quite good. To be fast the code does not try to be very accurate in the early SCF iterations, saving computing time. As long as the warnings do not persist into the final iteration, these warnings can be ignored, they are here because they can be helpful if things go wrong...

The information for iteration 8 is

```
in fft for local potential n = 24 24 16

max and min of potential (Hartree) 0.1819 -4.7630 0.0000

the fermi level is at 5.9538 [eV]

total energy = -31.8310932988
  computing time for iteration 8 1.890
```

again grep may be used to find out what is happening along the SCF iterations.

## 5.5 Potential, energies and forces

Once self-consistency is achieved, the information about the reciprocal space expansion of the density and potential are printed,

iteration number 9

i	k-prot		t	Ek	den	V(out)	V(in)	delta V	Vionic
1	0	0	0	0.00000	32.00000	-0.33659			
2	-1	0	0	0.14055	0.01194	0.00052	0.00051	0.00001	-0.01183
					0.00024	0.00001	0.00001	-0.00000	-0.00000
3	-1	1	0	0.18740	0.01175	0.00039	0.00039	0.00000	-0.01138
					0.02455	0.00082	0.00081	0.00001	-0.02276
4	-1 -	-1	0	0.37480	0.00032	-0.00005	-0.00005	-0.00000	0.00971
					0.00000	0.00000	0.00000	-0.00000	-0.00000
5	0	0	1	0.37480	0.00298	0.00018	0.00018	0.00000	-0.02912
					0.00000	0.00000	-0.00000	0.00000	0.00000
6	-2	1	0	0.51535	0.01475	0.00007	0.00007	0.00000	0.00858
					0.00565	0.00006	0.00005	0.00000	0.00000
7	0 -	-1	-1	0.51535	-0.01402	-0.00007	-0.00007	-0.00000	-0.00858
					-0.00344	-0.00003	-0.00003	-0.00000	0.00000
8	-2	0	0	0.56220	4.98977	0.03271	0.03271	-0.00000	-0.16014
					-5.03324	-0.03301	-0.03301	-0.00000	0.15190
9	-1	1	1	0.56220	4.98124	0.03270	0.03270	-0.00000	-0.16014
					5.02352	0.03299	0.03299	0.00000	-0.15190

after the index, one has the expansion of the G-vector, the kinetic energy associated (half the length squared) the Fourier coefficient of the density, the Fourier coefficients of the input and output XC potential, followed by the difference delta V and the coefficients of the ionic pseudopotential. The self-consistency is done by potential mixing (many codes do density mixing) and the self-consistency condition is on delta V.

Next we have the details of the total energy

Iteration number	9	Energies (Ha)	Changes
Alaba Tarm	_	0.017057	
Alpha Term	=	0.917257	
Kinetic Energy	=	11.949083	0.000022
Local PP Energy	=	-8.515343	-0.000104
Nonlocal Energy	=	5.722961	0.000071
Harris-Foulkes	=	-31.831093	-0.000000
Eigenvalue Sum	=	1.020901	-0.000013
HXC Correction	=	8.135799	0.00001
Hartree Energy	=	2.166313	0.000017
XC Energy	=	-9.562418	-0.000005
Ewald Energy	=	-33.591688	

Total Energy = -31.831093 0.000000

The last column has the change with respect to the previous iteration, it is not zero in the contributions, but has six decimals in the variational total energy, meaning nice convergence. it should be noticed that the Harris-Foulkes energy is identical (within the six decimals) with the total energy, another sign of good convergence.

Th total energy is the sum of the five last contributions, the Harris-Foulkes energy is the sum of the four first contributions plus the Ewald energy.

The stress tensor is given in both lattice and cartesian coordinates

Contravaria	nt stress	tensor (a.u.)	Cartesian stress (GPa)						
0.000642	0.000195	0.000000	0.239973E+01	0.929093E-12	0.267779E-13				
0.000195	0.000642	0.000000	0.914063E-12	0.256751E+01	-0.116381E-13				
0.000000	0.000000	0.000836	0.254685E-13	-0.104815E-13	0.119894E+01				
0 0000698	6	2 05539463	nressure (au and (	Pa)	Total				

its trace (divided by three) is the pressure.

Final information is the force on the atoms,

	Force	(Lattice	coord.)	Force (Cartesian coord. a.u)	10.	type	
0.0	0053	-0.00062	0.00000	-0.69115E-03 -0.11838E-01 0.74502E-17	1	Si	forc
-0.00	0001	0.00001	-0.00000	-0.80366E-16 0.15823E-03 -0.33131E-15	2	Si	ford
0.0	0062	-0.00053	-0.00000	0.69115E-03 -0.11838E-01 -0.30332E-15	3	Si	ford
-0.00	0058	0.00058	0.00000	-0.57519E-15 0.11843E-01 0.15964E-15	4	Si	ford
-0.00	0053	0.00053	0.00000	-0.14862E-15 0.10863E-01 0.88501E-16	5	Si	ford
-0.00	0063	0.00063	-0.00000	-0.51521E-15 0.12831E-01 -0.18928E-15	6	Ge	ford
-0.00	0032	-0.00091	-0.00000	-0.89184E-02 -0.60653E-02 -0.27596E-15	7	Ge	ford
0.0	0091	0.00032	-0.00000	0.89184E-02 -0.60653E-02 -0.51362E-16	8	Ge	ford

indicating that we are close but not at equilibrium.

Computing time is given at the very end

```
Total computing time (s): 47.74 Elapsed time (s): 12.01
```

with the ratio between computing and elapsed time indicating that the system had four cores.

# 6 Other files from cpw.exe

By design the code tries to write the minimum number of files. With grep one should be able to produce relevant information.

For example

```
cpw2000-5.x.y/WORK$ grep "4 Si position" ../Validation/Output/pw_ref_04.out will produce the trajectory of the 4th atom in the simulation cell
```

0.37456	0.12804	0.00365	0.36474E+01 -0.25300E+01 0.26210E-01 4 Si	posi
0.37435	0.13059	0.01401	0.36644E+01 -0.25017E+01 0.10068E+00 4 Si	posi
0.37460	0.13249	0.02300	0.36800E+01 -0.24848E+01 0.16520E+00 4 Si	posi
0.37504	0.13449	0.02959	0.36977E+01 -0.24688E+01 0.21256E+00 4 Si	posi
0.37525	0.13653	0.03398	0.37141E+01 -0.24500E+01 0.24414E+00 4 Si	posi
0.37561	0.13857	0.03628	0.37314E+01 -0.24328E+01 0.26066E+00 4 Si	posi
0.37419	0.12132	-0.02379	0.35960E+01 -0.25952E+01 -0.17094E+00 4 Si	posi
0.37595	0.11977	-0.02458	0.35975E+01 -0.26292E+01 -0.17655E+00 4 Si	posi
0.37730	0.11958	-0.02441	0.36060E+01 -0.26451E+01 -0.17536E+00 4 Si	posi
0.37842	0.11900	-0.02299	0.36098E+01 -0.26624E+01 -0.16519E+00 4 Si	posi
0.37909	0.11856	-0.02009	0.36116E+01 -0.26738E+01 -0.14429E+00 4 Si	posi
0.37946	0.11954	-0.01604	0.36213E+01 -0.26676E+01 -0.11524E+00 4 Si	posi

allowing the user to plot it.

If the atoms move a cpw.out file will be written. With eventual modifications it can be used to continue the calculations with the final geometry. Notice that the default choices are different.

If the atoms move a RESTART.DAT file will be present. It may be used to continue the simulation for more steps, or restart if the calculation was interrupted by a power failure.

The most important file will be PW\_RHO\_V.DAT. It is a binary file, so it cannot be modified by accident but that may not allow portability between different operating systems. It contains the information about the crystal structure, charge density and potential.

# 7 Post processing

Once the self-consistent potential for some structure has been saved to the PW\_RHO\_V.DAT file, one can proceed to calculate many properties witn a post-processing code,

```
cpw2000-5.x.y/WORK$ ../cpw_post_process.exe
```

it is mainly an interactive code, asking questions and proceeding according to the answers. All the answers are printed back to a file replay\_post.dat. This way the user will have a record of the answers, and more important, may rerun the post processing, with eventually slight changes.

```
cpw2000-5.x.y/WORK$ cp replay_post.dat replay.dat
```

and then, with optional minor editing of the file,

```
cpw2000-5.x.y/WORK$ ../cpw_post_process.exe < replay.dat</pre>
```

The post processing code is in fact several independent subroutines (sub-programs) called from the same top level program.

That program loops at most 100 times (just edit if you want more!) asking what the user wants and reads an integer entered by the user.

If the integer is zero, the program stops. At lower levels of the program a choice of zero would return the execution to the calling level.

If the integer is not in the allowed range, the question is repeated. In lower levels of the program an out of range choice gives the user a second chance. If the user gives again a wrong choice the code returns to the calling level.

Looking at the examples

# 7.1 Band structures, density of states, optical response, quantum geometric,...

This is the part of the post-processing that has the most options. The code gives a summary of the main parameters of the SCF calculation (geometry, pseudopotentials, etc,...) and then informs the user what cutoff was used for the plane-wave expansion in the SCF calculation and asks the user what cutoff (s)he wants to use. Choosing the same cutoff is a safe option. In this case only Hartree units are allowed. The code then asks if it can use the dual approximation, again answering "yes" is safe. It then asks if the user wants to modify other parameters. those are

## • Eigenvalue precision

You can change the eigenvalue precision of the iterative diagonalization if you are not confortable with the default. You can manage with a low precision in most cases.

## • Fermi energy

The code will read from the file a reasonable guess of the Fermi energy. In several cases it will shift the calculated eigenvalues by that value. The estimate is not precise, and if you have a better estimate you can enter it.

In most cases you can answer no.

The code than gives a choice of several types of calculations.

#### 7.1.1 Band structure

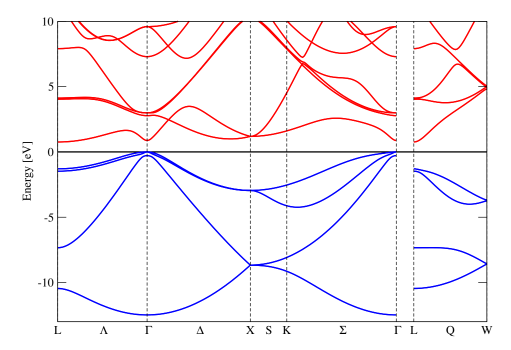
The first option is to plot a band structure. For that you need a path in the Brillouin zone. That information must in a file called BAND\_LINES.DAT in the working directory. In the documention the user can find information from QUANTUM ESPRESSO and ELK that can help write or modify the BAND\_LINES.DAT file. In Structures/Band\_Lines the user can find some examples.

An example of a BAND\_LINES.DAT file for an fcc lattice is

5	11	0.01							
0.5	0.5	0.5	0.0	0.0	0.0	80	L :	Lambda	${\tt Gamma}$
0.0	0.0	0.0	0.0	0.5	0.5	80	Gamm	a Delta	X
0.0	0.5	0.5	0.25	0.625	0.625	20	X	S	K
0.25	0.625	0.625	1.0	1.0	1.0	60	K	Sigma	${\tt Gamma}$
0.5	0.5	0.5	0.25	0.75	0.5	60	L	Q	W

In the first line one must specify the number of lines, 5 in this case, and how many bands (without spin) one wants to plot. A third optional value gives the spacing between calcualted points.

Each subsequent line indicates a pannel in the band structure plot. The initial and final point are indicated in reciprocal lattice coordinates. In the example the conventional choice of primitive lattice vectors was used. After the coordinates of the final point one has the number of points in that pannel, it is **not** used if in the first line the spacing between points is specified. Finally the labels of the first point, line and last point are specified. The code understands the meaning of the greek letters Gamma, Lambda, Delta, Sigma.



A band structure for Ge is shown in the figure. Notice that if the end of one pannel is the beginning of the next the pannels are joined. If not a small space is introduced.

The band structure can be calculted by several methods.

## • Full plane wave basis diagonalization

This is the "normal" case, slower but most precise. You will produced files with and without spin-orbit, in the GRACE and gnuplot formats.

The higher quality is the GRACE file, band\_so.agr and band.agr for the case with and without spin-orbit. But you have to have installed the graphical software, https://plasma-gate.weizmann.ac.il/Grace/, it is available on most linux distributions. However development seems to have stopped. It is a good start for publication quality figures.

Available in all distributions is gnuplot, just type

cpw2000-5.x.y/WORK\$ gnuplot band\_so.gp

and you will have a window with the band structure.

#### • Diagonalization in a Luttinger-Kohn basis

This is very fast but only accurate around the  $\Gamma$  point of the Brillouin zone. It will produce the similar files as the previous case but with an added  $_{\mathtt{lk}}$  in the file name.

#### • k.p method

It is a second order expansion of the hamiltonian in a Luttinger-Kohn basis around the  $\Gamma$  point. So it is very similar to the previous method, but now he files have a  $\_kp$  added to the file name.

## • 2-k-point Luttinger-Kohn interpolation

This is a very good compromise between speed and accuracy, but needs some additional information. It calculates the bands on a few points and then interpolates with the GLK method between them.

The code first asks how many interpolation points you want between calculated points. Notice that the values in BAND\_LINES.DAT that control the number of k-points is after interpolation. Besides the starting and end points on each pannel, you need to calculate a few (1 to 3) intermediate points to have a good interpolation, so the answer to that quaetion depends on how many final points you are going to have. The speedup in computing time is almost the number of intermediate points. Accuracy and computing time are again the compromises you have to take.

The code than asks if you want to accept the defaults of the calculation. If unsure accept the defaults. Otherwise read the code and the relevant article.

This option only produces the grace figures. They have an added \_lk\_int to the file names. There is also a band\_lk\_ref.agr file with the reference (fully calculated) bands. It should have visible straight lines if you are using the method efficiently.

## 7.2 Modified tight binding

For a fast dense sampling of the Brillouin zone, a modified tight-binding scheme can be used. It has many similarities with Wannier interpolation, and in fact can be used to write the files required to call WANNIER90.

### 7.3 Plot of charge densities and potentials

Plots or prepares the plots of real functions electron density,  $\rho(\vec{r})$ , effective potential,  $v_{\text{eff}}(\vec{r})$  and some potential contributions.

It can generate a three dimensional plot in the ".xsf format of XCRYSDEN compatible with VESTA, or a two dimensional level plot in a plane chosen by the user.

An interesting capability is to plot along a line these quantities averaged over the perpendicular plane. It is very useful to find band alignments.

#### 7.4 Plot densities of states

The data for the calculation of the density of states and respective plot can be generated with different approaches. This section of post processing reads the file with the data and generates the plot.

Choices of graphics options are separated from the sometimes very heavy calculations of the bands throughout the Brillouin zone.

#### 7.5 Exploration of a k.p model

The first option may store a k.p model in a file. This section allows the exploration of that model.

#### 7.6 Plot dielectric function

Like the case of density of states, the data for the dielectric function is first calculated throughout the Brillouin zone, the choices of graphics options are chosen in this section of the code.

#### 7.7 Plot of the wave-functions

It is similar to the plots of the charge density, but now the function to be plotted is complex, adding an extra choice of real part, imaginary part, or module, plus obviously the choice of which state to plot. Both 2D and 3D options are available.

#### 7.8 Crystalography

This section allows a reveryough examination of the crystal structure.

#### 7.9 Interface to Quantum Espresso

Quantum Espresso has many more developers than the present code. This section writes a QE input file. The pseudopotential code can write the required pseudopotentials, including Troullier-Martins pseudopotentials with spin-orbit (a feature that is not available in the QE toolchain). The consistency between the two codes are at the 4th or 5th decimal place in the total energy.

## 8 Where to find crystal structures

Many example of cpw.in files can be found in the Structures directory. Besides (almost) all the elements, there are many files collected over the years.

There is an auxiliary program in the toolbox, Tools/gen\_PW.f90 that can help the user to create a cpw.in file by answering some questions.

As mentioned before, the input file was chosen to be very similar to the SIESTA input file, so the user adapt it easily.

Another option is to convert any crystal file in the ".xyz format which is sufficiently simple to be adapted.

## 9 Toolbox

In Tools directory the user can find a few useful tools.

Most are sufficiently short that the user can easily read and modify the code.

It also has programs that call one of the options of post-processing. They were used to develop those options.

#### 9.1 Equation of state

From a few values of energy as a function of volume (or lattice constant for cubic systems), a least square fit is applied to the Murnaghan or Birch-Murnaghan equations of state. The output is the equilibrium volume, bulk modulus and pressure derivative of the Bulk modulus. If the user has several such data sets, the code will find the transition pressures between those structures.

#### 9.2 Remove gnuplot commands from 2D plot files

The 2D plot files are in the gnuplot format, with plotting directives. To remove the directives one can run Tools/convert\_2Dplot.f90

## 9.3 Crystalography from cpw.in

Doing a crystalographic analysis after a self-consistent calculation may be too late. With Tools/cpwin\_geom.f90 the analysis may be done from the cpw.in file. Read the crystallography sub-section of the post-processing section to see how it works.

## 9.4 Coulomb potential

The Coulomb potential for hydrogen can be generate by Tools/h\_pot\_kb.f90

## 9.5 Graphical interface for analysis of band structure

This is a very useful tool. From a file with the relevant data it generates a very powerful GUI. There is an earlier version BandInfoUi based on the grace plotting package, which is similar to the most recent package and therefore there is no need to describe in detail.

The most recent package is the QtBandViewer. It is a python code that uses the Qt user interface framework (https://www.qt.io).

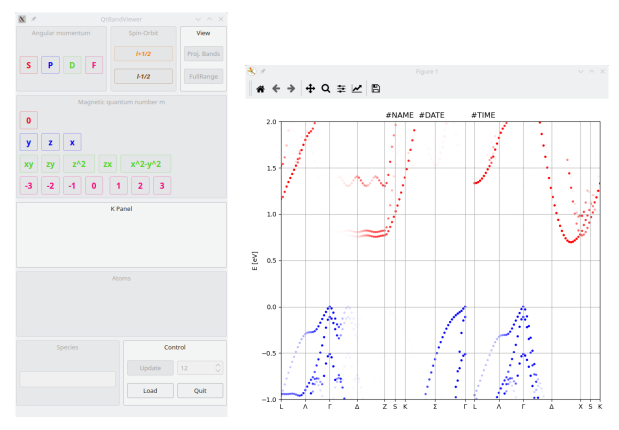
To install follow the instructions on Tools/QtBandViewer/AAAREADME. It is strongly recommended to create a python virtual environment under a working directory, and copy files there. By some obscure reasons (the writer of the guide does not use python) the python files should be copied. To use the tool you will need access to a .bv file format, by default named BAND\_SO.DAT.bv or BAND.DAT.bv with the band energies and the BAND\_LINES.DAT file used to generate those files. Those data files can be huge so avoid copying them.

As an example we will use the files generated from the structure file Structures/Supercells/cpw\_Si6Ge6\_Si(001).in with unfolding. The BAND\_LINES.DAT was copied from Structures/Supercells/band\_lines\_Si6Ge6\_Si(001).dat

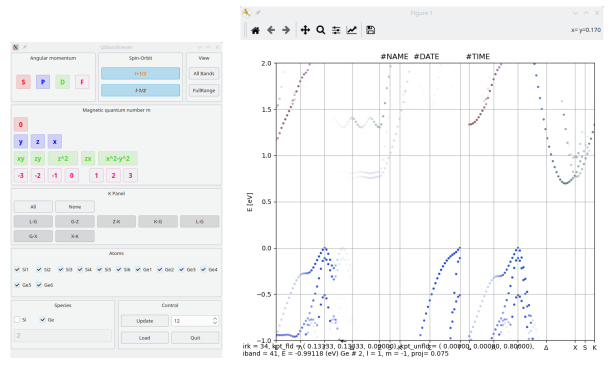
once the code is installed, run it

## cpw2000-5.x.y/WORK/QtBV\$ python3 QtBandViewer.py

The initial window is shown in the left of the figure. Press the Load button, and a new window shown on the right will display the unfolded band structure. Notice how the bottom of the conduction band appear to be in the  $\Gamma$ -X direction (parallel to the surface) and there are flat bands in the epitaxial growth direction  $\Gamma$ -Z.



If now we deselect Si leaving only the Ge contributions, and select the s, p and d angular momenta, we see that "dots" at the bottom of the band are still dark, but that the flat band almost disappears. This indicates that the bottom of the conduction band has a contribution from the Ge layers and there is a localized band in the Si layer.



Reselecting Si and deselecting Ge and keeping the s, p and d angular momenta, we confirm that the flat band is indeed mostly Si, the bottom of the conduction band has a Si contribution that is weaker than the Ge contribution. It is also apparent that for the valence band at the unfolded

 $\Gamma$  there is a band below the top and split-off bands that has a stronger Si content. For larger supercell periodicities it would converge to the band alignment between Si and strained Ge.

