QE-2019: Hands-on session - Day-1



Topics of Day-1 hands-on session:

- 1. Installation/compilation of Quantum ESPRESSO (example0.QE-compilation)
- 2. How to run basic PWscf (pw.x) calculations
- 3. How to run post-processing calculations to plot molecular-orbitals and charge-density (pp.x), DOS (dos.x), and band-structure (bands.x)
- 4. How to calculate low-dimensional systems (example1.benzene and example2.graphene)
- 5. How to make basic convergence tests (example3.Si)
- 6. How to deal with metals (example4.Al)
- 7. How to deal with ultrasoft-pseudopotentials and with spin polarization (example5.Fe)

About Quantum ESPRESSO



More info about Quantum ESPRESSO can be found in:

- https://www.quantum-espresso.org/
- Quantum ESPRESSO (QE) documentation:
 - on-line manuals at www.quantum-espresso.org/resources/users-manual
 - Doc/ sub-directories in the QUANTUM ESPRESSO distribution
 - input data description: most programs contained in QE have their own input file description in the form of hyperlinked INPUT_***.html files (where *** stands for the name of the program)

Hands-on material



Hands-on material for each day is contained within its own directory:

- Day-1/ hands-on exercises for Day-1
- Day-2/, Day-3/, Day-4/, Day-5/ hands-on exercises for Day-2 to Day-5

Please go to the Day-1/ directory and execute: git pull this will update the hands-on exercises to the latest version from the GitLab.

- All directories contain a README.md file with instructions how to run exercise(s)
- Naming of files is described in README-filenames.md (in Day-1/).
- To help recognizing for which program a given input file is intended, the filename starts with the name of the program, i.e.:
 - pw.*.in input file for pw.x program
 - pp.*.in input file for pp.x program
 - etc.

Disclaimer: many examples use lousy convergence thresholds to speed-up calculations

0. Compilation of Quantum ESPRESSO



Please go to the Day-1/ directory.

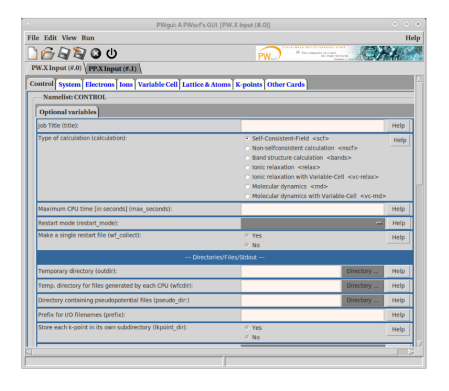
• cd exampleO.QE-compilation/

TODO: please complete the slide ...

Preparation of Quantum ESPRESSO input files

A few tools are available that aid at editing Quantum ESPRESSO input files:

- PWgui a QE input file builder GUI (pwgui)
- QE-emacs-modes makes editing of input files easier with emacs editor. It provides syntax highlighting, basic auto-indentation, and several utility commands; its manual is available in the QE sub-directory GUI/QE-modes/Doc/user_guide.pdf



```
~ A X
                emacs@catalyst.ijs.si
File Edit Options Buffers Tools Help
   calculation = 'scf'
              = 'silicon'
   outdir
               = '/temp/tone/pw/Si'
   tstress
   tprnfor
               = .true.
   ibrv = 2 ! mistyped variable is not highlighted
   A = 5.43
   nat = 2
   ntyp = 1
   ecutwfc = 18
&ELECTRONS
  conv_thr = 1.0d-8
ATOMIC SPECIES
  Si 28.086 Si.pz-vbc.UPF
ATOMIC POSITIONS alat
          0.00 0.00
0.25 0.25
K POINTS automatic
   4 4 4 1 1 1
-:--- scf.Si.in
```

About QE-emacs-modes

QE-emacs-modes package provides syntax highlighting, auto-indentation, and several utility commands, in particular:

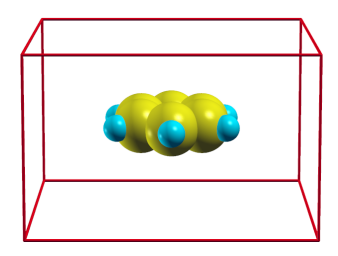
- Alt-x prog-insert-template inserts a respective input file template
- Alt-x prog-NAMELIST inserts a blank namelist named NAMELIST
- Alt-x prog-CARD inserts a blank namelist named CARD
- Alt-x prog-variable inserts a namelist variable named variable
- Alt-x prog-mode toggles the respective mode
- Alt-x indent-region indents region

where

- prog is one of qe, pw, neb, cp, ph, ld1, or pp (these stands for pw.x, neb.x, ... Quantum ESPRESSO (QE) programs)
- NAMELIST is the uppercase name for a given QE namelist
- CARD is the uppercase name for a given QE card
- variable is the lowercase name for a given namelist variable

1. How to describe a molecule with Quantum ESPRESSO

With Quantum ESPRESSO we can describe a molecule by putting it in a big box.



- move to Day-1/example1.benzene/ directory
- look at the input file pw.benzene.scf.in. It is composed of three "namelists" &CONTROL (note that calculation ='scf' is the default value), &SYSTEM, &ELECTRONS, followed by three "cards" ATOMIC_SPECIES, ATOMIC_POSITIONS, K_POINTS
- instructions for how to run the example are in README.md

Disclaimer: the box used in this example is very small as to speed-up calculations

1. How to calculate and plot molecular orbitals

Here are the two needed input files for calculation of molecular orbitals of benzene (actually, $sign(\psi(\mathbf{r}))|\psi(\mathbf{r})|^2$), opened with emacs using QE-emacs-modes:

```
emacs@cl
File Edit Options Buffers Tools Help
&CONTROL
   prefix='benzene',
 &SYSTEM
   ibrav = 6
   A = 11.0
   C = 7.0
   nat = 12,
   ntyp = 2,
   nbnd = 16
   ecutwfc = 20.0.
   ecutrho = 200.0,
   assume isolated = 'martyna-tuckerman',
 &ELECTRONS
ATOMIC SPECIES
   C 1.0 C.pbe-rrkjus.UPF
  H 1.0 H.pbe-rrkjus.UPF
ATOMIC POSITIONS angstrom
  H 5.5000000 7.98563953
      5.5000000
                6.89520922
      6.7089386 6.19812524
     7.6529918
                6.74309454
                4.80187470
     6.7089386
                4.25690561
      7.6529918
     5.5000000
                4.10479062
  H 5.5000000 3.01436043
      4.2910612 4.80187468
  H 3.3470081 4.25690556
    4.2910613 6.19812528
  H 3.3470082
                6.74309458
K_POINTS gamma
-:--- pw.benzene.scf.in All (18,2)
                                        (QE-pw.x)
Beginning of buffer
```

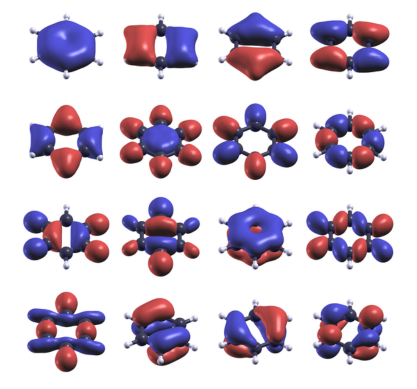
```
(v) (A) (X
                    emacs@cl
File Edit Options Buffers Tools Help
&INPUTPP
    prefix
             = 'benzene',
   filplot = 'psi2.benzene',
   plot num = 7,
    kpoint = 1,
   kband(1) = 1,
   kband(2) = 16,
   lsign = .true.,
&PL0T
   fileout
                 = '.xsf',
   iflag
   nfile
   output format = 5,
   weight(1) = 1.0,
-:--- pp.benzene.psi2.in All (1,0)
                                         (QE-pp.x)
```

1. How to calculate and plot molecular orbitals

To plot molecular orbitals (actually the signed square modulus, $sign(\psi(\mathbf{r}))|\psi(\mathbf{r})|^2$), we need to:

- calculate Kohn-Sham states with pw.x (i.e. make an SCF calculation)
- instruct pp.x to write them in a suitable format to specified files
- plot orbitals with xcrysden

See README.md for detailed instructions.



1. How to plot molecular orbitals with xcrysden

Execute in the terminal:

```
$ pw.x < pw.benzene.scf.in > pw.benzene.scf.out
$ pp.x < pp.benzene.scf.in > pp.benzene.scf.out
```

The resulting molecular orbitals (i.e., $sign(\psi(r))|\psi(r)|^2$) are written to psi2.benzene_*.xsf

• Plot one of the generated XSF files with xcrysden, e.g.:

```
$ xcrysden --xsf psi2.benzene_K001_B006.xsf
```

and follow these instructions:

- use the menu Tools-->Data Grid; a new window opens, press [OK]
- an isosurface-control window appear; specify the Isovalue, say 0.005 and press [Submit]
- click the Render +/- isovalue radiobutton and again press [Submit]
- rotate and zoom the structure according to your preference
- save the displayed state via the menu File-->Save Current State (e.g., save to my-display.xcrysden)
- try this with other orbitals, e.g.:

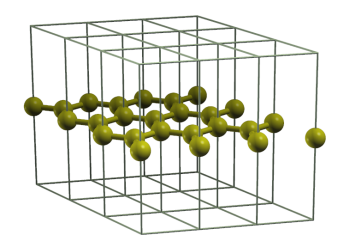
```
$ xcrysden --xsf psi2.benzene_K001_B005.xsf --script my-display.xcrysden
```

To plot all orbitals, execute: ./plot-psi2.sh

2. How to calculate a 2D-periodic system: graphene

A 2D-periodic system (e.g., a graphene sheet) is modelled by adding a vacuum layer in the 3rd direction.

- move to Day-1/example2.graphene/ directory
- look at the input file pw.graphene.scf.in; graphene has a 2-atom hexagonal unit cell in the xy plane: ibrav=4, celldm(1)=4.654, celldm(3)= $some\ suitably\ large\ value,\ e.g.\ 3.0;$



(remember: celldm(1) in Bohr radii, celldm(3)=c/a; alternatively: A=2.463, C=7.388 in Å)

• atomic positions:

ATOMIC_POSITIONS (alat)
C 0.000000 0.000000 0.000000
C 0.000000 0.5773503 0.000000

or, equivalently:

ATOMIC_POSITIONS (crystal)

C 0.000000 0.000000 0.000000

C 0.333333 0.666667 0.000000

 \bullet k-points: use a dense grid in the xy plane only, e.g.

K_POINTS (automatic)

9 9 1 0 0 0

(a uniform $9\times9\times1$ grid, centered on $\mathbf{k}=(0,0,0)$)

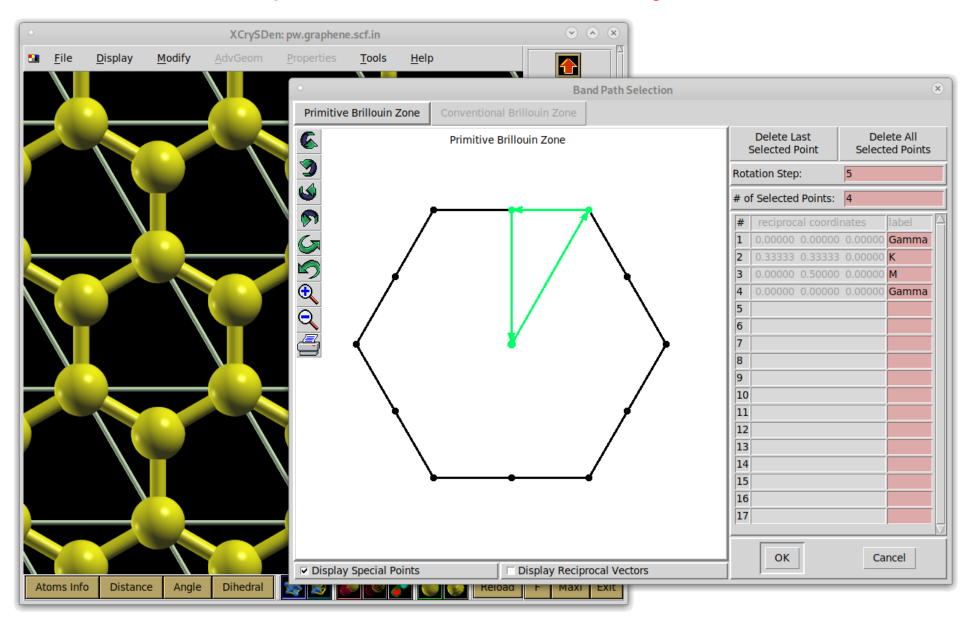
2. Graphene: DOS and bands (spaghetti)

- DOS is typically calculated by a pw.x SCF calculation followed by a pw.x non-SCF calculation (calculation = 'nscf') with a denser k-point grid, and finally using dos.x post-processing code.
- to calculate the bands (spaghetti plot), the pw.x SCF calculation is followed by a pw.x "bands"-type non-SCF calculation (calculation = 'bands'), for which we need a suitable path of k-points. The most difficult (?) part is to figure out a suitable path of k-points.

You may either use the "k-path selection" tool of xcrysden or the SeeK-path web site at http://materialscloud.org/tools/seekpath.

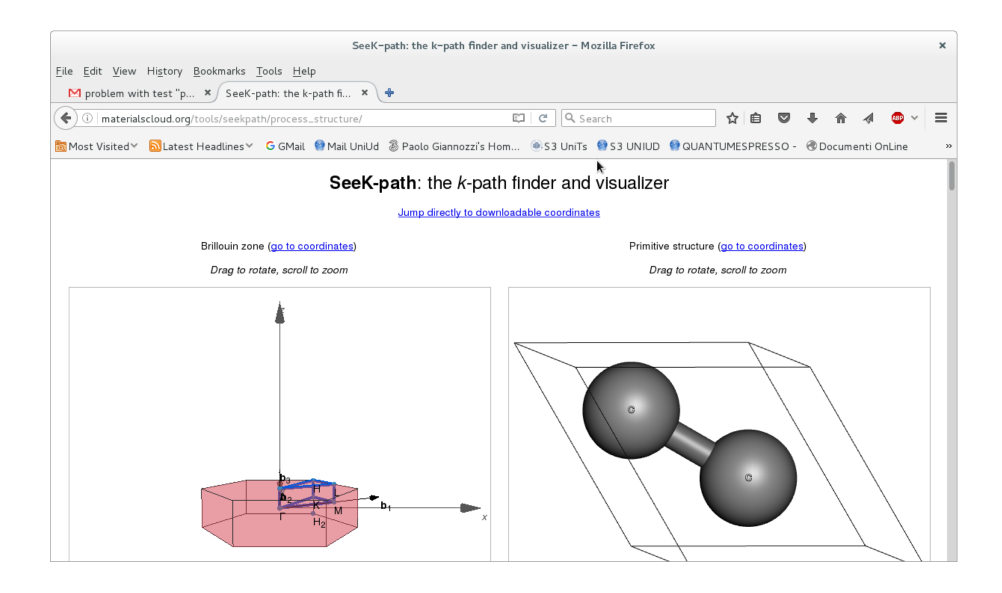
• instructions for how to calculate DOS and bands are in README.md

K-path selection tool of xcrysden



(important: to save k-path in Quantum ESPRESSO format, explicitly specify the *.pwscf extension)

SeeK-path @ http://materialscloud.org/tools/seekpath



3. Bulk system: Silicon

Self-consistent calculation (and a series of tests) for Silicon in the diamond structure:

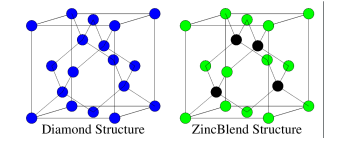
- move to Day-1/example3.Si directory
- look at the input file pw.si.scf.in. It is composed of three "namelists" &CONTROL (note that calculation = 'scf' is the default value), &SYSTEM, &ELECTRONS, followed by three "cards" ATOMIC_SPECIES, ATOMIC_POSITIONS, K_POINTS
- in the &CONTROL namelist notice the following two variables (they are commented):
 - outdir: temporary directory for large files. Must be writable, will be created if not existent. You may set environment variable ESPRESSO_TMPDIR instead.
 - pseudo_dir: directory where pseudopotential (PP) files are kept. It must exist, be readable, and contain the required PP file (in this example, Si.pz-vbc.UPF for Silicon). You may set environment variable ESPRESSO_PSEUDO instead.

(note that for the hands-on exercises we rely on ESPRESSO_TMPDIR and ESPRESSO_PSEUDO environmental variables, hence we don't need to set outdir and pseudo_dir variables)

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Providing atomic structure in input

How is the crystal structure defined? This is a very simple case: the diamond lattice is an fcc (face-centered cubic) lattice with two atoms per unit cell. You need to specify:



- What is the Bravais lattice?ibrav=2, meaning fcc lattice
- How many and which parameters are needed to completely define Bravais lattice? just one: celldm(1)=10.2, lattice parameter a in a.u.
- How many atoms there are in the unit cell?
 nat=2: two atoms
- How many different atomic species are present?ntyp=1: one species
- Which ones, described by which pseudopotential?
 See card ATOMIC_SPECIES
- Where the atoms are located in the unit cell? See card ATOMIC_POSITIONS: here, in Cartesian axes, in units of a ("alat")

Notice that there are several alternative methods to specify an atomic structure!

Brillouin zone (BZ) sampling

k-points are described in the **K_POINTS** card. One has to choose

Whether to provide a list of k-points or a uniform grid

• If a list is chosen: provide a list of k-points in the Irreducible BZ and corresponding symmetry weights; the latter do not need to add up to 1, they are normalized by the code

Frequently Asked Question: where do I find special k-points and their weights? Answer: 1) in papers, 2) use an auxiliary code kpoints.x, 3) use uniform grids

• If a uniform grid is chosen: Monkhorst-Pack parameters (H.J. Monkhorst and J.D. Pack, *Phys. Rev. B* **13**, 5188 (1976)), and offsets along the three directions

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Running the pw.x code

For serial (single processor) execution you can use

```
$ pw.x -in pw.si.scf.in > pw.si.scf.out
```

(note: input redirection pw.x < pw.si.scf.in works but it is not recommended on parallel machines)

Look at the directory specified by **outdir** (in our case **\$ESPRESSO_TMPDIR**) and its content:

```
$ ls $ESPRESSO_TMPDIR
silicon.save silicon.xml
```

(to see only these files, you may need to use ls \$ESPRESSO_TMPDIR/silicon.*)

The directory contains a data directory with binary data files for further processing and an XML file with general information on the run. The name of the various files is determined by the value of the prefix variable and by their content.

Do not run two instances of pw.x that access the same outdir with the same prefix! Unpredictable behavior may follow (the directory is used for temporary files as well). In case of trouble, clean outdir.

Running the pw.x code (II)

Examine output file pw.si.scf.out, look how self-consistency proceeds:

```
$ grep -e 'total energy' -e estimated pw.si.scf.out}
                                  -15.79103344 Ry
    total energy
    estimated scf accuracy
                                    0.06376674 Ry
    total energy
                             = -15.79409289 Ry
    estimated scf accuracy
                                    0.00230109 Rv
    total energy
                             = -15.79447822 Ry
                                    0.00006291 Ry
    estimated scf accuracy
                             = -15.79449510 Ry
    total energy
                                    0.00000448 Ry
    estimated scf accuracy <
                             = -15.79449593 Ry
    total energy
    estimated scf accuracy <
                                    0.0000005 Ry
    The total energy is the sum of the following terms:
```

Notice that there are 8 electrons in the cell: 2 (pseudo-)atoms/cell with 4 electrons. The system is a non-magnetic insulator, so just the lowest 4=8/2 valence bands (Kohn-Sham states) are computed.

Convergence w.r.t. the kinetic energy cutoff

The kinetic energy cutoff ecutwfc (in Ry) determines the size of the Plane-Wave (PW) basis set used to expand wave-functions (i.e. Kohn-Sham orbitals) (for Norm-Conserving PP, the PW set for charge density ecutrho=4*ecutwfc, do not specify it)

- A manual test of convergence w.r.t. kinetic energy cutoff entails the following tasks (BEWARE: we will not do it manually)
 - 1. change value of ecutwfc in pw.si.scf.in input to, e.g., 16, 20, 24, 28, 32 Ry
 - 2. for each value of ecutwfc, run pw.x and collect the final energy
 - 3. collect the data in a file, say si.etot_vs_ecut (i.e. each line should contain two values: ecutwfc and "total-energy")
 - 4. plot the energies collected in si.etot_vs_ecut using your preferred plotting program, for instance:

```
$ gnuplot
gnuplot> plot 'si.etot_vs_ecut' with lines
```

because such a manual procedure is very cumbersome we use scripts instead

Convergence w.r.t. kinetic energy cutoff (II)

To make convergence tests easier and faster, scripts are commonly used. To this end, Unix shell-scripts have been traditionally used, yet there are other more "fancy" alternatives, e.g., PWTK scripts.

- A Unix shell-script is located in ex1.ecutwfc.classic/sub-directory (file: ecutwfc.sh)
- A PWTK script is located in ex1.ecutwfc/ sub-directory (file: ecutwfc.pwtk)

```
Unix shell-script
#!/bin/sh
rm -f si.etot_vs_ecut.dat
for ecut in 12 16 20 24 28 32
   cat > pw.si.scf.$ecut.in << EOF
   prefix='silicon',
 &SYSTEM
   ibrav = 2,
   celldm(1) = 10.2,
   nat = 2,
   ntyp = 1,
   ecutwfc = $ecut,
&ELECTRONS
ATOMIC_SPECIES
  Si 28.086 Si.pz-vbc.UPF
ATOMIC POSITIONS
  Si 0.00 0.00 0.00
  Si 0.25 0.25 0.25
K_POINTS automatic
 4 4 4 1 1 1
EOF
   pw.x -in pw.si.scf.$ecut.in > pw.si.scf.$ecut.out
   grep -e 'kinetic-energy cutoff' -e ! pw.si.scf.\ecut.out \
       awk '/kinetic-energy/ {ecut=$(NF-1)}
                            {print ecut, $(NF-1)}' >> si.etot_vs_ecut.dat
done
```

```
load_fromPWI ../pw.si.scf.in

set fid [open si.etot_vs_ecut.dat w]

foreach ecut {12 16 20 24 28 32} {
    SYSTEM "ecutwfc = $ecut"
    runPW pw.Si.scf.$ecut.in

    puts $fid "$ecut [::pwtk::pwo::totene pw.Si.scf.$ecut.out]"
} close $fid
```

PWTK scripting: basics



The basic philosophy is to keep the syntax close to original input syntax!

```
pw.x input file
```

```
&CONTROL
   calculation = 'scf'
&SYSTEM
   ecutwfc = 25.0
   ecutrho = 200.0
ATOMIC POSITIONS alat
   Si
         0.00 0.00
                     0.00
   Si
         0.25 0.25 0.25
K POINTS automatic
           1 1 1
```

pw.Si.in

Run from terminal as:

```
pw.x -in pw.Si.in > pw.Si.out
```

script

```
CONTROL {
   calculation = 'scf'
SYSTEM {
   ecutwfc = 25.0
   ecutrho = 8*25.0
ATOMIC POSITIONS alat {
   Si
         0.0
              0.0
                   0.0
         1/4 1/4
                   1/4
   Si
K POINTS automatic {
runPW pw.Si.in
```

Si bulk.pwtk

Run from terminal as:

```
pwtk Si bulk.pwtk
```

PWTK scripting: basics



- PWTK scripts are basically Tcl-scripts, hence they use Tcl-syntax
- namelists and cards have the same names as in QE (with a few exceptions), but they are all written in **upper-case**. Their content is encapsulated in curly braces:

```
CONTROL { calculation = 'scf', outdir = '/tmp/pwscf/'}
ATOMIC_POSITIONS { ... }
```

instead of curly braces, one can also use double-quotes ("..."), e.g.:
 SYSTEM " celldm(1) = \$a "

- the difference between curly braces {...} and double-quotes "..." is that inside double-quotes the variable \$a is substituted by its value, whereas inside curly braces the \$a is treated literally (i.e. no substitution)
- real numbers can be specified as mathematical expressions (e.g. ecutrho = 8*25.0)
- indices of ntyp-type array variables, such as starting_magnetization(i), can be specified with atomic labels, e.g.:

```
SYSTEM { starting_magnetization(Fe) = -0.8 } where Fe is one among atomic species defined in the ATOMIC_SPECIES card.
```





- namelist variables can be set on-the fly (we will use this heavily)
- the order of namelists and cards is not important; PWTK knows how to construct proper input files
- namelist and cards can be called many times, but there is a big difference how multiple calls are handled for namelists and cards: cards are handled in overwrite mode, whereas namelists are handled in a kind of append mode. For example, the following is OK:

```
CONTROL { calculation = 'scf' }
CONTROL { outdir = '/tmp/qe' }
and is equivalent to:
CONTROL { calculation = 'scf' , outdir = '/tmp/qe' }
```

• to unset a namelist variable, set it to an empty-value; for example to unset the outdir variable, use:

```
CONTROL { outdir = }
```

• today we will use the following PWTK functions:

- PW TK
- load_fromPWI loads input data from an existing pw.x input file
- ::pwtk::pwo::totene returns the converged total energy from pw.x output
- seq like the Unix seq command (returns a sequence of numbers)
- runPW constructs pw.x input file and runs a calculation
- runPP similar as runPW but for the pp.x program
- runDOS ... for the dos.x program
- runPROJWFC ... for the projwfc.x program
- the PWTK web-site is:

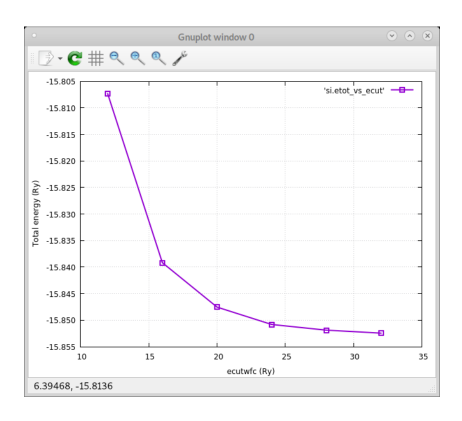
```
http://pwtk.ijs.si/ or http://pwtk.quantum-espresso.org/
```

PWTK documentation is available at:

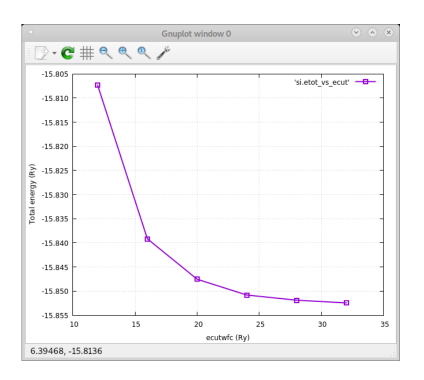
http://pwtk.ijs.si/toc_index.html

Convergence w.r.t. the kinetic energy cutoff (III)

- To run the convergence test via the Unix shell-script move to ex1.ecutwfc.classic sub-directory (read the README.md file) and execute
 - \$./ecutwfc.sh
- To run the convergence test via the PWTK script move to ex1.ecutwfc subdirectory (read the README.md file) and execute
 - \$ pwtk ecutwfc.pwtk



Convergence w.r.t. the kinetic energy cutoff (IV)



Reminder:

- Convergence w.r.t the cutoff is a property of the *pseudopotential(s)* used.
- Convergence of the *absolute energy* is typically slower than convergence of *interesting physical properties*, e.g. structure.
- Absolute values of total energy do not have any physical meaning (and depend upon the specific PP): only energy differences do

Convergence w.r.t. k-points

A sufficiently dense grid of k-points is needed in order to account for *periodicity*.

To test the convergence w.r.t. k-points, you need to edit the K_POINTS card and request *automatic* Monkhorst-Pack grids:

```
K_POINTS automatic
nk1 nk2 nk3     k1 k2 k3
```

then step-wise increase nk1=nk2=nk3 to, e.g., 2, 4, 6, 8 (keep k1=k2=k3=1) and a run pw.x calculation for each value of nk1=nk2=nk3.

For example, with PWTK this can be achieved with the following snippet:

```
load_fromPWI pw.si.scf.in

foreach k {2 4 6 8} {
    K_POINTS automatic "$k $k $k 1 1 1"
    runPW pw.si.scf.$k.in
}
```

Convergence w.r.t. k-points (II)

Description of the K_POINTS card for *automatic* mode:

```
K_POINTS automatic
nk1 nk2 nk3     k1 k2 k3
```

The first three nk1 nk2 nk3 numbers mean "there are nk1,nk2,nk3 grid points along crystal axis 1,2,3"; the second three k1 k2 k3 numbers, either 0 or 1, mean "grid starts from 0" or "displaced by half a step" along crystal axis 1,2,3

Also note that:

- Convergence is not necessarily monotonic: there is no variational principle w.r.t.
 number of k-points
- The 2 2 2 1 1 1 Monkhorst-Pack grid is the same as the "two Chadi-Cohen points" (see: D.J. Chadi and M.L. Cohen, Phys. Rev. B 8, 5747 (1973))

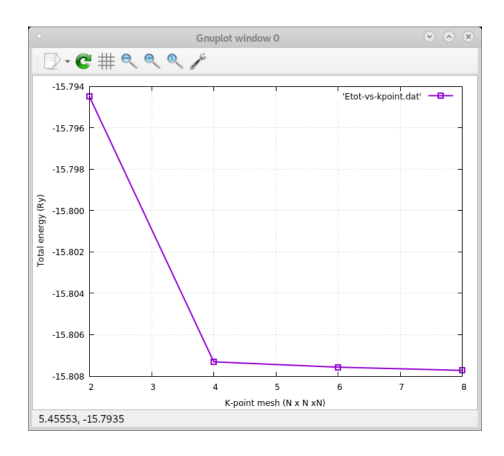
Convergence w.r.t. k-points (III)

The PWTK script for testing the convergence with respect to k-points is located in Day-1/example3.Si/ex2.kpoints directory (see README.md for detailed instructions).

Within this directory execute:

\$ pwtk kpoints.pwtk

You should get a plot like this one:



Equation of State: silicon

Equilibrium in Si is determined by the minimum-energy lattice parameter alone: there are no forces on atoms, by symmetry (you can verify this by setting tprnfor=.true. in namelist &CONTROL and looking for forces reprinted at the end).

To find the lattice parameter:

- Choose suitable values for ecutwfc and the k-point grid (e.g. 30 Ry and 4 4 4 1 1 1)
- Run pw.x for values of celldm(1) ranging from 9.7 to 10.7 in steps of 0.1 a.u.

With PWTK this can be achieved with the following snippet:

```
load_fromPWI pw.si.scf.in

foreach alat [seq 9.7 0.1 10.7] {
    SYSTEM "celldm(1) = $alat"
    runPW pw.si.scf.$alat.in
}
```

Equation of State: silicon (II)

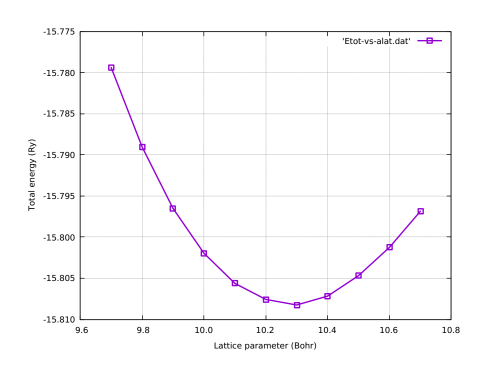
The corresponding PWTK script is located in Day-1/example3.Si/ex3.alat directory (see README.md for detailed instructions).

Within this directory execute:

\$ pwtk alat.pwtk

The experimental lattice parameter for Si is 5.47 Å or 10.26 a.u.. This is a case where plain simple LDA yields remarkable results. You may experiment changing cutoff, k-points, pseudopotential, ...

You should find that:



- The energy vs lattice parameter E(a) curves are shifted down rather uniformly with increasing cutoff and are not strongly dependent on k-points.
- Structural properties and energy differences converge faster than total energies.

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Equation of State: silicon (III)

Use the code ev.x to fit your results to a phenomenological equation-of-state (EOS, e.g. Murnaghan) and to get accurate values for the lattice parameter and for the bulk modulus.

The ev.x code prompts for some data and reads a data-file like the one produced by the alat.pwtk script (the data-file is Etot-vs-alat.dat). For cubic systems a data-file should contain the following rows:

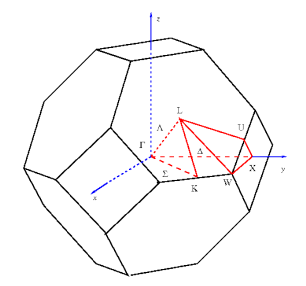
```
a_1 E(a_1)
a_2 E(a_2)
a_3 E(a_3)
```

. . .

Band Structure of Silicon

The scheme to calculate the bands (spaghetti plot) is the following:

- 1. SCF pw.x calculation (calculation = 'scf')
- 2. "bands"-type non-SCF pw.x calculation (fixed-potential) with:
 - calculation = 'bands'
 - the number of Kohn-Sham states explicitly set (variable nbnd)
 - a suitable path of k-points specified in K_POINTS (in this example we use the $L-\Gamma-X-W-K-L$ path)



3. bands.x calculation, which, among others, produces data-files for the spaghetti plot

Important: outdir and prefix must be the same for "bands" and "scf" pw.x calculations and for the bands.x calculation
Important: the k-point path must be continuous in k-space

Band Structure of Silicon (II)

The input for the bands.x program is the following:

```
&BANDS

prefix='...', outdir='...', filband = 'Si.bands.dat', lsym=.true.
/
```

Two files are produced: Si.bands.dat.gnu, directly plottable with gnuplot, and Si.bands.dat, for further processing by the auxiliary command plotband.x.

If option lsym=.true., bands.x performs a symmetry analysis. An additional file Si.bands.dat.rep is generated, containing information on symmetry labels of the various bands.

The snippet for running all three calculations manually is:

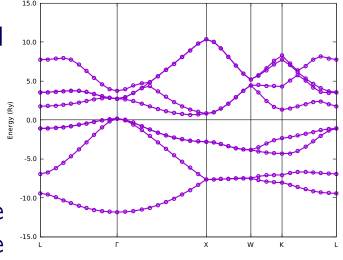
```
$ pw.x -in pw.Si.scf.in > pw.Si.scf.out
$ pw.x -in pw.Si.bands.in > pw.Si.bands.out
$ pw.x -in bands.Si.in > bands.Si.scf.out
```

but we will use a PWTK script instead.

Band Structure of Silicon (III)

To execute the PWTK script that will perform all the needed calculations to plot the bands, proceed as follows:

- move to directory Day-1/example3.Si/ex4.bands and read the README.md file
- set suitable values for celldm(1), ecutwfc, and K_POINTS
- and execute: pwtk bands.pwtk
- you may set the **Efermi** value to the top of the occupied bands in the gnuplot file plot.gp (see the instructions in README.md); then re-plot spaghetti with: gnuplot plot.gp



Remark: in PWTK, once outdir and prefix are set, they are automatically inherited for subsequent calculations.

Auxiliary program plotbands.x

plotband.x prompts for terminal input:

```
$ plotband.x
Input file > Si.bands.dat
Reading 8 bands at 39 k-points
Range: -5.6940 16.4680eV Emin, Emax > -5.6940 16.4680
high-symmetry point: -0.5000 0.5000 0.5000 x coordinate
                                                         0.0000
high-symmetry point: 0.0000 0.0000 0.0000 x coordinate 0.8660
high-symmetry point: 0.0000 0.0000 1.0000 x coordinate 1.8660
high-symmetry point: 0.0000 0.5000 1.0000 x coordinate
                                                        2.3660
high-symmetry point: 0.0000 0.7500 0.7500 x coordinate 2.7196
high-symmetry point: -0.5000 0.5000 0.5000
                                                        3.3320
                                         x coordinate
output file (gnuplot/xmgr) > Si.bands.plot
bands in gnuplot/xmgr format written to file Si.bands.plot
output file (ps) > (press Return)
```

If symmetry analysis was performed in the previous step, the output is written to several plottable files Si.bands.plot.N.M, where N labels the high-symmetry lines, M labels irreducible representations.

4. A metallic example: Aluminum

Aluminum is even simpler than Silicon: one atom per unit cell in a fcc lattice. **BUT:** it is a metal, only valence bands and a few k-points will not suffice.

- move to the Day-1/example4.Al directory
- read the pw.x input file pw.al.scf.in
- notice the presence of new variables: occupations, smearing, degauss;
- run pw.x as:

```
$ pw.x -in pw.al.scf.in > pw.al.scf.out
```

- in the output file notice that
 - the number of bands (Kohn-Sham states) is automatically set to a value larger than the number of electrons divided by 2
 - the Fermi energy is computed.

Convergence with respect to k-points, degauss, and smearing

This is a "three-dimensional" convergence test, where we will vary the number of k-points and values of degauss and smearing variables. In particular, we will vary:

- smearing variable, possible values: 'gauss' (or 'g'), 'marzari-vanderbilt'
 (or 'm-v'), 'methfessel-paxton' (or 'm-p')
- degauss variable, in range from 0.01 to 0.15
- k-points using the automatic grids of 6 6, 12 12, and 16 16 16

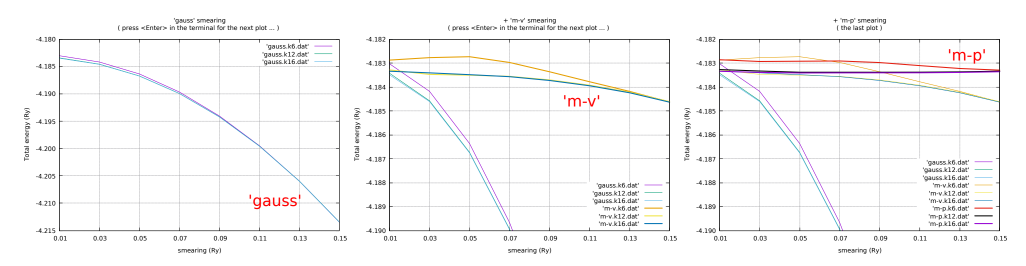
With PWTK this can be achieved with the following snippet:

Convergence with respect to k-points, degauss, and smearing (II)

- move to Day-1/example4.Al/ex1.degauss directory
- execute: pwtk degauss.pwtk

Notice how much slower the convergence is for metals than for insulators!

Both m-v and m-p depend much less upon degauss and allow for faster and safer convergence than simple gaussian broadening. For Al and m-v or m-p smearing, good convergence is achieved for a 12 12 k-point grid and degauss ~ 0.01 to 0.05 Ry.



Notice that you cannot reduce the broadening too much: the energy levels must have some overlap, or else the advantage of broadening is lost!

How to plot charge-density

Example Day-1/example4.Al/ex2.chdens shows how to calculate the valence and the all electron charge density (the latter requires PAW a potential and a very large cutoff energy)

- move to Day-1/example4.Al/ex2.chdens directory (chdens is an acronym for charge-density)
- execute: pwtk 1-chdens.pwtk this script calculates and "plots" the valence charge density; notice that the electron charge is located mainly in interstitial regions (due to the use of a pseudo-potential, there is almost no charge in close vicinity of nuclei; see next page);
- the scheme to calculate and plot the charge-density is:
 - 1. make an SCF pw.x calculation
 - 2. make a post-processing pp.x calculation (plot_num=0 for charge density) and instruct the program to write charge density in a suitable format
 - 3. plot the charge density by xcrysden (let's plot density in contour/colorplane style; follow the instructions of tutor and select density range from 0.0 to 0.05)

How to plot charge-density (II)

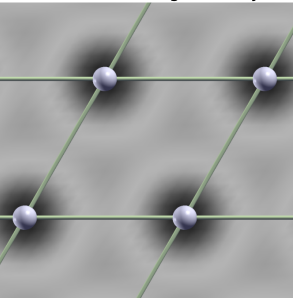
• to calculate all-electron **valence** and **total** charge densities, execute:

\$ pwtk 2-chdens-paw.pwtk

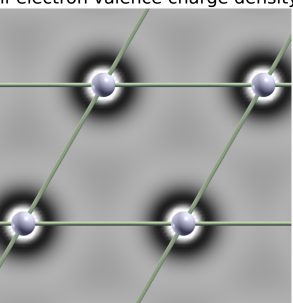
(note that plot_num=17 for all-electron valence density and plot_num=21 for all-electron total density)

 comparison between PP valence-density vs. all-electron densities (valence and total):

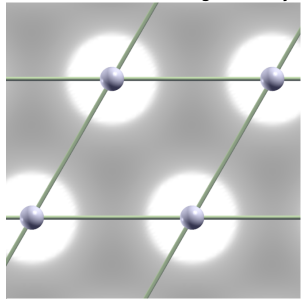
PP valence charge density



all-electron valence charge density



all-electron total charge density



5. A magnetic example: Iron

Iron has two remarkable features: it is magnetic and it requires an Ultrasoft PP (USPP) since its localized 3d atomic states are very hard.

- move to the Day-1/example5.Fe directory and read the pw.x input file pw.fe_fm.scf.in
- the structure is bcc (ibrav=3) with one atom per unit cell
- notice the presence of variables nspin and of starting_magnetization, indicating LSDA (nspin=2) with unconstrained total magnetization and initial symmetry broken; plus, variables for metallic calculations
- notice that this calculation uses GGA (PBE): it is specified inside the PP file (can be guessed from the PP file name), reprinted on output as "Exchange-correlation"
- also notice that with USPP, it is typically needed to set ecutrho $> 4 \times$ ecutwfc (it should be at least 8 to 12 times larger)

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Magnetic structures

Run pw.x in the usual way (pw.x -in $pw.fe_fm.scf.in > pw.fe_fm.scf.out$). In the output, notice:

- the number of k-points is doubled w.r.t the non-magnetic case: the first set of k-points contains spin-up states, the second set spin-down states

 (use verbosity='high' in namelist &CONTROL if there are more than 100 k-points)
- in the output notice such lines:

```
total magnetization = 2.41 Bohr mag/cell absolute magnetization = 2.60 Bohr mag/cell
```

Since there is a single (magnetic) atom per unit cell, the only possible magnetic structure is ferromagnetic.

Magnetic structures: going antiferomagnetic

- in order to reach antiferromagnetic states, you need to:
 - introduce a supercell with two sublattices of different species of atoms (even if they are the same, it is important that they are labeled as different)
 - start with opposite initial magnetization for the two sublattices
- Can you write input data for an AFM structure? (hint: split bcc into two simple cubic sublattices, **ibrav=1**, with two atoms at (0,0,0) and $(\frac{1}{2},\frac{1}{2},\frac{1}{2})$).

As a convenience, an antiferromagnetic file is provided (pw.fe_afm.scf.in)

You can compare the ferromagnetic and antiferromagnetic files by:

\$ diff pw.fe_fm.scf.in pw.fe_afm.scf.in

Convergence check for USPP

For computational efficiency, it is convenient to keep ecutwfc as low as possible, while ecutrho is less critical (look at the CPU time report at the end of an output: there are very many fftw, depending upon ecutwfc, while a much smaller number of fft depends upon ecutrho)

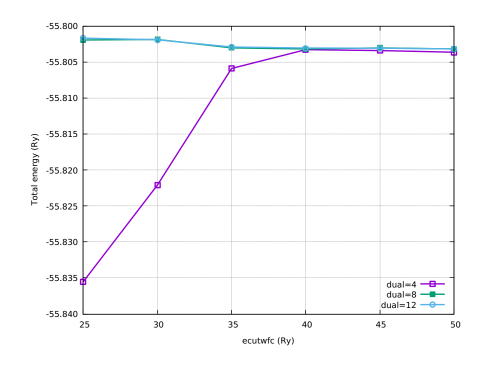
Set the ecutrho/ecutwfc ratio (dual) to 4, 8, 12 and compute the energy vs ecutwfc curve. For dual = 4 it will look funny: energy *increases* with increasing cutoff (see next page), but for a higher dual (i.e. better description of augmentation charge) the normal behavior is observed.

The corresponding PWTK snippet is:

Convergence check for USPP (II)

A PWTK script to make convergence check for USPP is provided, i.e.:

- move to directory:
 Day-1/example5.Fe/ex1.ecut
- execute: pwtk ecut.pwtk
- you should obtain such a plot (notice that curves for dual=8 and 12 almost coincide)

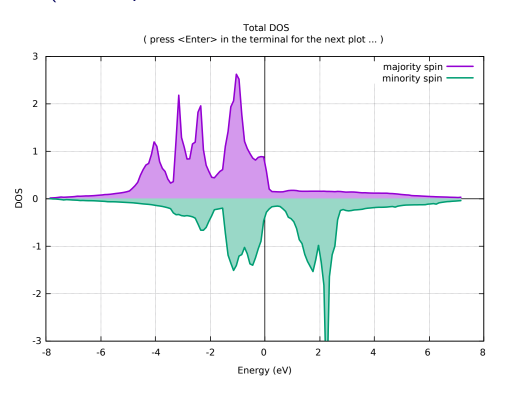


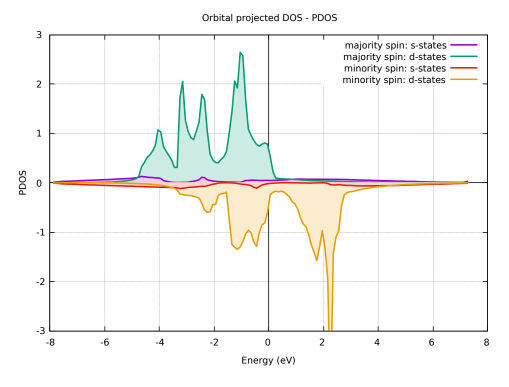
Homework: for converged values of both cutoffs and k-points, you may compare the stability of iron in the bcc, fcc, hcp phases (the latter being a slightly more complicated structure)

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Density of States (DOS) and Projected DOS (PDOS)

- 1. move to Day-1/example5.Fe/ex2.dos directory and
- 2. edit the dos.pwtk script and set ecutwfc and ecutrho to appropriate values
- 3. execute: pwtk dos.pwtk (this script calculates both total DOS and DOS projected (PDOS) to atomic orbitals)





Density of States (DOS) and Projected DOS (PDOS)

The scheme to calculate DOS and PDOS consists of:

- 1. an SCF pw.x calculation (calculation='scf')
- 2. a non-SCF pw.x calculation (calculation='nscf'), where:
 - the same prefix and outdir are used as in the preceding SCF calculation
 - a denser k-point mesh is specified
 - in this example the *linear tetrahedron method* is used (variable occupations='tetrahedra')
- 3. a dos.x calculation to calculate DOS (DOS is written to a file as specified by the fildos variable in the dos.x input; also here the values prefix and outdir are the same as for SCF pw.x calculation)
- 4. a projwfc.x calculation to calculate PDOS projected to atomic states (this calculation is analogous to dos.x one, inputs are also very similar; you can compare them as:

```
$ diff dos.Fe.in projwfc.Fe.in
```

the difference is that dos.x uses the fildos whereas projwfc.x uses the filpdos variable)

Explore the content of fildos and filpdos files!