Advanced Quantum ESPRESSO tutorial: Hubbard and Koopmans functionals from linear response Pavia (Italy), 28 August 2022

Hands-On Tutorial
Wannier interpolation of band structures
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#### Foreword

This document contains 2 exercises to help you learn wannier90 (W90) - the computer program that calculates maximally-localised Wannier functions (MLWFs).

# Required executables and tutorial files

- QE: pw.x, bands.x, plotband.x, pw2wannier90.x
- Wannier90: wannier90.x, kmesh.pl

#### The tutorial files are hosted on GitHub repo:

https://github.com/materialscloud-org/QuantumESPRESSO-school-2023/tree/main/Day1 You can download the files through git clone:

git clone https://github.com/materialscloud-org/QuantumESPRESSO-school-2023/

#### Software for visualization

- To plot the band structure:
  - gnuplot http://www.gnuplot.info/index.html
  - xmgrace https://plasma-gate.weizmann.ac.il/Grace/
- To plot the Wannier functions, Fermi surface:
  - xcrysden http://www.xcrysden.org/
  - VESTA https://jp-minerals.org/vesta/en/

These are already pre-installed in the Quantum Mobile.

### 1 Silicon valence bands

In this exercise you will learn how to obtain maximally-localised Wannier functions (MLWFs) for the valence bands of silicon.

- Go to the exercise\_wannier\_1 folder and inspect the input file O1\_scf.in. The first step is to perform a ground-state calculation for a silicon crystal (FCC) with two atoms per unit cell. Check if you understand all parameters (you can use the web page https://www.quantum-espresso.org/Doc/INPUT\_PW.html for keywords that you do not know or ask us). To visualize the crystal structure, there are two options, you can choose either one as you prefer:
  - Quantum ESPRESSO input generator and structure visualizer: open the following link in a browser,
    - https://www.materialscloud.org/work/tools/qeinputgenerator, click Choose File button, select the O1\_scf.in file, and click "Generate the PWscf input file" button. In the new webpage you find a 3D visualization of the structure.
  - xcrysden (if you have installed in locally in your computer): either opening the program and selecting from the menu File→Open PWscf...→Open PWscf Input File and selecting the input file, or directly from the command line with the command xcrysden --pwi O1\_scf.in.

```
&control
calculation
               = 'scf'
              = 'from_scratch'
restart_mode
              = 'si'
prefix
               = '../../files/pseudo/'
pseudo_dir
               = 'out/'
outdir
&system
                   0
ibrav
                   2
nat
                  1
ntyp
               = 25.0
ecutwfc
               = 200.0
ecutrho
&electrons
conv_thr
                 1.0d-10
ATOMIC_SPECIES
Si 28. Si.pbe-n-van.UPF
ATOMIC_POSITIONS crystal
Si -0.25
           0.75
                 -0.25
    0.00
           0.00
                   0.00
K_POINTS automatic
10 10 10 0 0 0
CELL_PARAMETERS bohr
-5.1
     0.0
            5.1
0.0
      5.1
             5.1
-5.1
       5.1
              0.0
```

- The Si pseudopotential that we will use for the calculation has  $Z_{val} = 4$  (this information can be obtained reading the first lines of the pseudopotential file, for instance with the command less ../../files/pseudo/Si.pbe-n-van.UPF). Using the information given above, and knowing that FCC Si is a semiconductor, how many occupied valence bands do you expect (and why)?
- Run the ground state calculation using the pw.x code of the Quantum ESPRESSO suite. The syntax for codes in the Quantum ESPRESSO suite is: command < inputfile > outputfile (i.e. pw.x < 01\_scf.in > scf.out).
- After the calculation finishes, inspect the output file scf.out to check if there are any errors/warnings. Compare your answer to the previous point (number of electrons and occupied valence bands) with the information provided in the output file.
- Now we want to plot the band structure of silicon (we will use this plot also for the next exercise, where we need also the conduction bands: therefore, we plot also a few of the lowest conduction bands). Copy the file O1\_scf.in to the file O2\_bands.in. Do the following modifications to the file O2\_bands.in (use the INPUT\_PW documentation from the link above for an explanation of the meaning of the flags, if needed):
  - In the CONTROL namelist, change the calculation keyword from 'scf' to 'bands' to perform a band structure calculation starting from the ground state density obtained from the scf run.
  - Ask the code to print 12 bands (flag nbnd=12 in the SYSTEM namelist).
  - Set diago\_full\_acc = .true. in the ELECTRONS namelist (see documentation for the meaning of this flag).
  - Change the k-point list to plot the band structure along the following path (coordinates are given in crystal units), using 50 points per segment:

```
* L(0.5, 0.5, 0.5) \rightarrow \Gamma(0, 0, 0)
* \Gamma(0, 0, 0) \rightarrow X(0.5, 0, 0.5)
```

You can do this simply using the following K\_POINTS card:

```
K_POINTS crystal_b
3
0.5 0.5 0.5 50
0. 0. 0. 50
0.5 0. 0.5 50
```

- Run the calculation using the pw.x code, as explained before.
- When the calculation has finished, run the file O3\_bandsx.in through the bands.x executable (make sure to read and understand the input file):

```
ibrun -np 2 bands.x < 03_bandsx.in > bandsx.out
```

```
&bands
prefix = 'si'
outdir = 'out/'
```

```
filband = 'bands.dat'
lsym = .false.
/
```

This will produce the bands.dat file.

- Finally, execute the plotband.x code (interactively) and answer to its questions. In particular, the input file is the bands.dat file created in the previous step; call the xmgrace file qebands.agr. When asked, call the ps file qebands.ps. You will be asked to provide the value of the Fermi level, which in this case can be put equal to the highest occupied energy level (see the output file scf.out). When asked for the deltaE and reference E for the energy axis, type 2 and Fermi level (use space to separate the 2 numbers), you can also tweak these 2 numbers to adjust the visual output of the figure. At the end, open the xmgrace file (or directly the postscript PS file) and inspect the band structure, identifying the valence and conduction bands.
- Now we are ready to calculate the wavefunctions on a complete grid of k-points. Copy the O2\_bands.in file that you created before to O5\_nscf.in, and modify the following:
  - Change the calculation type from 'bands' to 'nscf'.
  - Change the number of bands from 12 back to the number of valence bands that you expect (see your answer a few lines above: the answer should be 4), since for this exercise we need only the valence bands.
  - Change the k-point list to a full  $4 \times 4 \times 4$  Monkhorst-Pack mesh, that will be used to calculate the overlap matrices needed to obtain Wannier functions. To obtain the list of k-points, use the kmesh.pl utility in the utility folder of the Wannier90 code using the following command for a  $4 \times 4 \times 4$  mesh:

/home/max/koopmans/quantum\_espresso/q-e/wannier90-3.1.0/utility/kmesh.pl

To allow easier usage, let's add it to the \$PATH variable:

export PATH=/home/max/koopmans/quantum\_espresso/q-e/wannier90-3.1.0/utility:\$PATH (use the command without parameters to get an explanation of its usage).

- Run the nscf calculation using the pw.x code: mpirun -np 2 pw.x < 05\_nscf.in > nscf.out

- Now we have to prepare the input file for Wannier90. Open the file ex1.win, which is a template of the Wannier90 input file (note that Wannier90 input file must have the .win extension). Change the values marked with XXX inserting the correct values. In particular:
  - Insert the num\_bands value (this must be equal to the nbnd value set in the nscf calculation).
  - Insert the num\_wann value (this is the number of requested Wannier functions: in this case without disentanglement, this is equal to the num\_bands value).
  - Set the mp\_grid value to 4 4 4 (since we are using a  $4 \times 4 \times 4$  k-mesh).

- Insert, between the begin kpoints and end kpoints lines, the list of the 64 kpoints, one per line. Note that while pw.x requires four numbers per line (the three coordinates of each kpoint, and the weight), Wannier90 needs only three numbers (the three coordinates). To obtain these lines, use again the kmesh.pl utility, but this time specifying a fourth parameter to get the list in the Wannier90 format:

```
kmesh.pl 4 4 4 wan
```

Note Using the kmesh.pl utility, we are sure that we provide enough significant digits, and that the list of k-points given to pw.x and to Wannier90 is the same.

- Inspect the remaining part of the input file, using the Wannier90 user guide (that can be found on the https://github.com/wannier-developers/wannier90/raw/v3.1.0/doc/compiled\_docs/user\_guide.pdf page) for the input flags that you do not understand. Try to understand, in particular, the projections section. Can you say where the four s-like orbitals are located with respect to the Si atoms? \_\_\_\_\_\_

```
use_ws_distance = .true.
num_bands
                    XXX
num_wann
               = XXX
num_iter
               = 100
iprint
num_dump_cycles = 10
num_print_cycles =
!! To plot the WFs
wannier_plot
                       = plot
                   = true
wannier_plot_supercell = 3
! wannier_plot_list
                   = 1,5
!! To plot the WF interpolated bandstructure
bands_plot = true
begin kpoint_path
L 0.50000 0.50000 0.5000 G 0.00000 0.00000 0.0000
G 0.00000 0.00000 0.0000 X 0.50000 0.00000 0.5000
end kpoint_path
!! !! Bond-centred s-orbitals
begin projections
f=-0.125,-0.125, 0.375:s
f = 0.375, -0.125, -0.125:s
f=-0.125, 0.375,-0.125:s
f=-0.125,-0.125,-0.125:s
end projections
begin atoms_frac
Si -0.25 0.75 -0.25
Si 0.00 0.00
                0.00
end atoms_frac
```

```
begin unit_cell_cart
bohr
-5.10
        0.00
             5.10
0.00
       5.10
              5.10
-5.10
       5.10
               0.00
end unit_cell_cart
mp_grid = XXX XXX XXX
begin kpoints
XXX
end kpoints
```

- Finally, we are ready to perform a Wannier90 calculation. This is done in three steps:
  - 1. We first run a preprocessing step using the command

```
wannier90.x -pp ex1
```

which produce a ex1.wout file and ex1.nnkp file, that contains the relevant information from the Wannier90 input file in a format to be used in the next step.

2. Then we run the pw2wannier90.x code (of the Quantum ESPRESSO distribution). The input file for pw2wannier90.x is provided (file 06\_pw2wan.in). We are asking the code to calculate the overlap matrices  $M_{mn}$  (that will be written in the ex1.mmn file) and the  $A_{mn}$  matrices (file ex1.amn). Since we want to plot the Wannier functions in real space, we need also the  $u_{nk}(r)$  wavefunctions on a real-space grid. We thus also set the write\_unk flag in 06\_pw2wan.in, that will produce a set of files with names UNK00001.1, UNK00002.1, ... Finally, the code will also produce a ex1.eig file, with the eigenvalues on the initial  $4 \times 4 \times 4$  k-grid (Note: this is not needed to obtain the MLWFs of an insulator, but only for the interpolation and band plotting routines). Note that the pw2wannier90.x expects to find the ex1.nnkp file produced in the previous step. Run the code using

mpirun -np 2 pw2wannier90.x < 06\_pw2wan.in > pw2wan.out

```
%inputpp
outdir = 'out/'
prefix = 'si'
seedname = 'ex1'
write_amn = .true.
write_mmn = .true.
write_unk = .true.
/
```

3. Finally we can run Wannier90 to obtain MLWFs. Execute

#### wannier90.x ex1

and, when it finishes, inspect the output file, called ex1.wout.

- Check lines containing <-- DLTA to check for the convergence of the spread during the iterations.
- Check the lines after the string Final state: you find the centers and spreads of the maximally-localised Wannier functions.

- To check if the obtained MLWFs are correct, it is typically needed to:
  - \* Compare the Wannier-interpolated band structure with the ab initio one: the provided Wannier90 input file computes the interpolated band plot; you can try to compare the ab initio bandplot obtained in the steps before with the interpolated band structure (files ex1\_band.dat, and ex1\_band.gnu)
    - · To plot it with gnuplot: run gnuplot in terminal, and in gnuplot, type

```
set xtics nomirr
set x2tics
set x2range [0:0.21721815E+01]
set xrange [0:1.3195]
plot 'bands.dat.gnu' w p pt 7, 'ex1_band.dat' axes x2y1 w l
```

Note you can reuse the script in following exercises when comparing band structures, by replacing the file names qebands.agr and ex1\_band.dat.

· Or plot it with xmgrace: in terminal, type:

```
xmgrace qebands.agr ex1_band.dat
```

Note that you may need to rescale the x axis.

Note that the Wannier90 code also outputs in the ex1\_band.kpt file a list of the kpoints used for the interpolation, that could be used to plot the band structure on the same grid.

- \* Plot the real-space Wannier functions and check if they are real: if you ask Wannier90 to plot the Wannier functions, it will print also the ratio of the imaginary and real part of each of them at the end of the ex1.wout file: check that the value is small.
- Plot one of the Wannier functions, which are output in files ex1\_00001.xsf, .... To visualize the Wannier functions, you need to install VESTA or xcrysden in your computer, and download the xsf files:
  - \* using xcrysden: open the xsf file, then choose Tools—Data Grid—OK, and then choose a reasonable isovalue, activate the Render +/- isovalue flag, and press Submit.
  - \* using VESTA: open the xsf file, VESTA can automatically find a isovalue.

### 2 Silicon valence and conduction bands

xcrysden --pwi 01\_scf.in

In this exercise you will learn how to obtain maximally-localised Wannier functions (MLWFs) for the valence and conduction bands of silicon.

- Go to the exercise\_wannier\_2/ folder and inspect the input file O1\_scf.in. The first step is to perform a ground-state calculation for a silicon crystal (FCC) with two atoms per unit cell. Check if you understand all parameters (you can use the web page https://www.quantum-espresso.org/Doc/INPUT\_PW.html for keywords that you do not know or ask us). To visualize the crystal structure, there are two options, you can choose either one as you prefer:
  - Quantum ESPRESSO input generator and structure visualizer: open the following link in a browser,

```
https://www.materialscloud.org/work/tools/qeinputgenerator, click Choose File button, select the O1_scf.in file, and click "Generate the PWscf input file" button. In the new webpage you find a 3D visualization of the structure.
```

- xcrysden: either opening the program and selecting from the menu File  $\to$  Open PWscf... $\to$  Open PWscf Input File and selecting the input file, or directly from the command line with the command

```
&control
 calculation = 'scf'
 restart_mode = 'from_scratch'
 prefix
                = '../../files/pseudo/'
 pseudo_dir
                = 'out/'
 outdir
&system
  ibrav
                    0
 nat
                 = 1
 ntyp
                = 25.0
 ecutwfc
                = 200.0
  ecutrho
&electrons
  conv_thr
                    1.0d-10
ATOMIC_SPECIES
 Si 28. Si.pbe-n-van.UPF
ATOMIC_POSITIONS crystal
 Si -0.25 0.75 -0.25
      0.00
             0.00
                    0.00
K_POINTS automatic
  10 10 10 0 0 0
CELL PARAMETERS bohr
 -5.1
         0.0
               5.1
  0.0
         5.1
               5.1
  -5.1
         5.1
                0.0
```

• Run the ground state calculation using the pw.x code of the Quantum ESPRESSO suite.

```
mpirun -np 2 pw.x < 01_scf.in > scf.out
```

- After the calculation finishes, inspect the output file scf.out to check if there are any errors/warnings. Compare your answer to the previous point (number of electrons and occupied valence bands) with the information provided in the output file.
- Now we want to plot the band structure of silicon. If you have already calculated the DFT bands in the previous exercise you can skip this step. Copy the file O1\_scf.in to a new file O2\_bands.in. Do the following modifications to the file O2\_bands.in (use the INPUT\_PW documentation from the link above for an explanation of the meaning of the flags, if needed):
  - In the CONTROL namelist, change the calculation keyword from 'scf' to 'bands' to perform a band structure calculation starting from the ground state density obtained from the scf run.
  - Ask the code to print 12 bands (flag nbnd=12 in the SYSTEM namelist).
  - Set diago\_full\_acc = .true. in the ELECTRONS namelist (see documentation for the meaning of this flag).
  - Change the k-point list to plot the band structure along the following path (coordinates are given in crystal units), using 50 points per segment:

```
* L(0.5, 0.5, 0.5) \rightarrow \Gamma(0, 0, 0)
* \Gamma(0, 0, 0) \rightarrow X(0.5, 0, 0.5)
```

You can do this simply using the following K\_POINTS card:

```
K_POINTS crystal_b
3
0.5 0.5 0.5 50
0. 0. 0. 50
0.5 0. 0.5 50
```

- Run the calculation using the pw.x code, as explained before.
- When the calculation has finished, run the file O3\_bandsx.in through the bands.x executable (make sure to read and understand the input file):

mpirun -np 2 bands.x < 03\_bandsx.in > bandsx.out

```
&bands
  prefix = 'si'
  outdir = 'out/'
  filband = 'bands.dat'
  lsym = .false.
/
```

This will produce the bands.dat file.

- Finally, execute the plotband.x code (interactively) and answer to its questions. In particular, the input file is the bands.dat file created in the previous step; call the xmgrace file qebands.agr. When asked, call the ps file qebands.ps. You will be asked to provide the value of the Fermi level, which in this case can be put equal to the highest occupied energy level (see the output file scf.out). When asked for the deltaE and reference E for the energy axis, type 2 and Fermi level (use space to separate the 2 numbers), you can also tweak these 2 numbers to adjust the visual output of the figure. At the end, open the xmgrace file (or directly the postscript PS file) and inspect the band structure, identifying the valence and conduction bands.
- Now we are ready to calculate the wavefunctions on a complete grid of k-points. Copy the O2\_bands.in file that you created before to O5\_nscf.in, and modify the following:
  - Change the calculation type from 'bands' to 'nscf'.
  - Keep the number of bands as 12.
  - Change the k-point list to a full 4 × 4 × 4 Monkhorst-Pack mesh, that will be used to calculate the overlap matrices needed to obtain Wannier functions. To obtain the list of k-points, use the kmesh.pl utility which is bundled in the Wannier90 code, that you can find it here:

/home/max/koopmans/quantum\_espresso/q-e/wannier90-3.1.0/utility/kmesh.pl

To allow easier usage, let's add it to the \$PATH variable:

export PATH=/home/max/koopmans/quantum\_espresso/q-e/wannier90-3.1.0/utility:\$PATH

After this you can directly call the kmesh.pl utility from the command line.

Then generate the kpoint coordinates using the following command for a  $4 \times 4 \times 4$  mesh:

kmesh.pl 4 4 4

(use the command without parameters to get an explanation of its usage).

The output of the command will be a list of k-points, which you can copy and paste in the K\_POINTS card of the O5\_nscf.in file.

- Run the nscf calculation using the pw.x code:

```
mpirun -np 2 pw.x < 05_nscf.in > nscf.out
```

- Now we have to prepare the input file for Wannier90. Open the file si2.win, which is a template of the Wannier90 input file (note that Wannier90 input file must have the .win extension). Change the values marked with XXX inserting the correct values. In particular:
  - Insert the num\_bands value (this must be equal to the nbnd value set in the nscf calculation
    if the exclude\_bands parameter is not specified).
  - Insert the num\_wann value (this is the number of requested Wannier functions: in this case for valence and conduction bands, this is equal to 8, why? You may find some hints from the projections keyword as will be shown in later paragraph).
  - Set the mp\_grid value to 4 4 4 (since we are using a  $4 \times 4 \times 4$  k-mesh).

- Insert, between the begin kpoints and end kpoints lines, the list of the 64 kpoints, one per line. Note that while pw.x requires four numbers per line (the three coordinates of each kpoint, and the weight), Wannier90 needs only three numbers (the three coordinates). To obtain these lines, use again the kmesh.pl utility, but this time specifying a fourth parameter to get the list in the Wannier90 format:

```
kmesh.pl 4 4 4 wan
```

Note Using the kmesh.pl utility, we are sure that we provide enough significant digits, and that the list of k-points given to pw.x and to Wannier90 is the same.

- Set the maximum energy for the frozen window (flag dis\_froz\_max) inside the energy gap (use the band plot obtained in previous step to get a value for this flag).
- Set the maximum energy for the disentanglement (flag dis\_win\_max) to an energy large enough so as to contain enough bands for each k point; 17.0 eV should be a reasonable value (check where this value lies in the band plot).
- Inspect the remaining part of the input file, using the Wannier90 user guide (that can be found on the https://github.com/wannier-developers/wannier90/raw/v3.1.0/doc/compiled\_docs/user\_guide.pdf page) for the input flags that you do not understand. Try to understand, in particular, the projections section (project to 4 sp³ orbitals for each Si atom in the unit cell).

```
use_ws_distance = .true.
num_bands
                    XXX
num_wann
                    XXX
num_iter
                = 100
                    XXX
dis_win_max
dis_froz_max
                    XXX
dis_num_iter = 100
!! To plot the WFs
! restart
                           plot
wannier_plot
                      = true
wannier_plot_supercell = 3
!! To plot the WF interpolated band structure
bands_plot = true
begin kpoint_path
L 0.50000 0.50000 0.5000 G 0.00000 0.00000 0.0000
G 0.00000 0.00000 0.0000 X 0.50000 0.00000 0.5000
end kpoint_path
begin projections
Si: sp3
end projections
begin atoms_frac
Si -0.25 0.75 -0.25
Si 0.00 0.00
                0.00
```

```
end atoms_frac
begin unit_cell_cart
bohr
-5.10
        0.00
               5.10
 0.00
        5.10
               5.10
-5.10
        5.10
               0.00
end unit_cell_cart
mp_grid = XXX XXX XXX
begin kpoints
XXX
end kpoints
```

- Finally, we are ready to perform a Wannier90 calculation. This is done in three steps:
  - 1. We first run a preprocessing step using the command

```
wannier90.x -pp si2
```

which produce a si2.wout file and si2.nnkp file, that contains the relevant information from the Wannier90 input file in a format to be used in the next step.

2. Then we run the pw2wannier90.x code (of the Quantum ESPRESSO distribution). The input file for pw2wannier90.x is provided (file 06\_pw2wan.in). We are asking the code to calculate the overlap matrices  $M_{mn}$  (that will be written in the si2.mmn file) and the  $A_{mn}$  matrices (file si2.amn). Since we want to plot the Wannier functions in real space, we need also the  $u_{nk}(r)$  wavefunctions on a real-space grid. We thus also set the write\_unk flag in 06\_pw2wan.in, that will produce a set of files with names UNK00001.1, UNK00002.1, ... Finally, the code will also produce a si2.eig file, with the eigenvalues on the initial  $4 \times 4 \times 4$  k-grid. Note that the pw2wannier90.x expects to find the si2.nnkp file produced in the previous step. Run the code using

mpirun -np 2 pw2wannier90.x < 06\_pw2wan.in > pw2wan.out

```
&inputpp
  outdir = 'out/'
  prefix = 'si'
  seedname = 'si2'
  write_amn = .true.
  write_mmn = .true.
  write_unk = .true.
//
```

3. Finally we can run Wannier90 to obtain MLWFs. Execute

```
wannier90.x si2
```

and, when it finishes, inspect the output file, called si2.wout.

 Before the start of the maximal localisation iterations, there is a section (containing the string <--DIS) with the iterations of the disentanglement procedure. It is impor-</li> tant that at the end of this section the convergence is achieved (with a string <<< Disentanglement convergence criteria satisfied >>>).

- Check lines containing <-- DLTA to check for the convergence of the spread during the maximal localisation iterations.
- Check the lines after the string Final state: you find the centers and spreads of the maximally-localised Wannier functions.
- To check if the obtained MLWFs are correct, it is typically needed to:
  - \* Compare the Wannier-interpolated band structure with the ab initio one: the provided Wannier90 input file computes the interpolated band plot; you can try to compare the ab initio bandplot obtained in the steps before with the interpolated band structure (files si2\_band.dat, and si2\_band.gnu)
    - To plot it with gnuplot: run gnuplot in terminal, and in gnuplot, type
      {gnuplot}
      set xtics nomirr
      set x2tics
      set x2range [0:0.21721815E+01]
      set xrange [0:1.3195]
      plot 'bands.dat.gnu' w p pt 7, 'si2\_band.dat' axes x2y1 w l
      Note you can reuse the script in following exercises when comparing band
      structures, by replacing the file names qebands.agr and si2\_band.dat.
    - $\cdot$  Or plot it with xmgrace: in terminal, type:

```
xmgrace qebands.agr si2_band.dat
```

Note that you may need to rescale the x axis.

The Wannier90 code also outputs in the si2\_band.kpt file a list of the kpoints used for the interpolation, that could be used to plot the band structure on the same grid.

- \* Plot the real-space Wannier functions and check if they are real: if you ask Wannier90 to plot the Wannier functions, it will print also the ratio of the imaginary and real part of each of them at the end of the si2.wout file: check that the value is small.
- \* A practical note: Especially when using disentanglement, it is possible that the disentanglement convergence is not achieved, and/or that the obtained Wannier functions are not real, and/or that the interpolated band structure differs significantly from the ab initio one within the frozen window. Then, you need to change/tune the number of Wannier functions, the projections you chose and/or the energy values for the frozen and disentanglement windows, until you get "good" Wannier functions.
- Plot one of the Wannier functions, which are output in files si2\_00001.xsf, ....
   To visualize the Wannier functions, you need to install xcrysden or VESTA in your computer, and download the xsf files:
  - \* using xcrysden: open the xsf file, then choose Tools—Data Grid—OK, and then choose a reasonable isovalue, activate the Render +/- isovalue flag, and press Submit.
  - \* using VESTA: open the xsf file, VESTA can automatically find a isovalue.

• Optional (Do it only if you have enough time): Do the symmetry and the centers of the Wannier functions agree with your intuition? (We would like 4 sp<sup>3</sup>-like orbitals centered on each Si atom, with similar spreads). Try to rerun everything with a 6 × 6 × 6 kgrid for the nscf and Wannier90 step to check if the results improve, and how the spreads change with respect to the grid density? Also try to tune the dis\_froz\_max and see the difference of MLWF spreads and band interpolation?

# 3 (Optional) Lead Fermi surface and band structure

In this exercise we will see how to interpolate the band structure of lead, in particular around the Fermi energy. The first goal is to obtain the Fermi surface from Wannier interpolation. This will clearly show some of the advantages of using Wannier-interpolation schemes with respect to full k-points direct calculations. In order to build a MLWFs model that describes the band structure around the Fermi energy, we need to have an idea of the orbital character of the bands we are interested in. The crystal structure of pure lead has one atom per primitive cell, and since we also want to describe some states above the Fermi Energy, we include five d orbitals  $(d_{xy}, d_{xz}, d_{yx}, d_{z^2})$  and  $d_{x^2-y^2}$ , and four  $sp^3$  orbitals. Hence our guess for the projections is

- $\bullet$  5 d orbitals centered on the lead atom
- $4 sp^3$  orbitals centred on the lead atom

Now we are ready to obtain MLWFs and describe the states of lead around the Fermi-level.

- Directory: exercise\_wannier\_3/
- Input Files
  - O1\_scf.in The PWSCF input file for the ground state calculation

```
&control
    calculation='scf'
    restart_mode='from_scratch',
    pseudo_dir = '../../files/pseudo/',
    outdir='./out'
    prefix='pb'
&system
    ibrav = 2, celldm(1) = 9.3555, nat = 1, ntyp = 1,
    ecutwfc = 47.0, ecutrho = 189,
    occupations='smearing', smearing='cold', degauss=0.02
&electrons
    conv_thr = 1.0e-9
    mixing_beta = 0.7
ATOMIC_SPECIES
  Pb 207.2 Pb.pbe-dn-kjpaw_psl.0.2.2.UPF
ATOMIC_POSITIONS (crystal)
  Pb 0.0 0.0 0.0
K_POINTS (automatic)
888 000
```

- 04\_nscf.in The PWSCF input file to obtain Bloch states on a uniform grid

```
&control
calculation='nscf'
pseudo_dir = '../../files/pseudo/',
```

```
outdir='./out'
prefix='pb'
/
&system
ibrav = 2, celldm(1) = 9.3555, nat = 1, ntyp = 1,
ecutwfc = 47.0, ecutrho = 189,
occupations='smearing', smearing='cold', degauss=0.02
nosym=.true.,nbnd=13
/
&electrons
conv_thr = 1.0e-9
ATOMIC_SPECIES
Pb 207.2 Pb.pbe-dn-kjpaw_psl.0.2.2.UPF
ATOMIC_POSITIONS
Pb 0.0 0.0 0.0
K_POINTS crystal
0.00000000 0.00000000 0.00000000 1.953125e-03
0.00000000 0.00000000 0.12500000 1.953125e-03
```

- 05\_pw2wan.in Input file for pw2wannier90

```
&inputpp
outdir = './out'
prefix = 'pb'
seedname = 'lead'
write_amn = .true.
write_mmn = .true.
//
```

- lead.win The wannier90 input file

```
G 0.00 0.00 0.00
                     K 0.00 0.50 -0.50
End Kpoint_Path
! SYSTEM
begin unit_cell_cart
bohr
-4.67775 0.00000 4.67775
0.00000 4.67775 4.67775
-4.67775 4.67775 0.00000
end unit_cell_cart
begin atoms_frac
Pb 0.00 0.00 0.00
end atoms_frac
begin projections
Pb:d;sp3
end projections
! KPOINTS
mp_grid : 8 8 8
begin kpoints
0.0000000 0.0000000 0.00000000
0.00000000 0.00000000 0.12500000
```

 Run PWSCF to obtain the ground state of lead pw.x < 01\_scf.in > scf.out

```
2. Run PWSCF to obtain the Bloch states on a uniform k-point grid pw.x < 04_nscf.in > nscf.out
```

- 3. Run wannier90 to generate a list of the required overlaps (written into the lead.nnkp file). wannier90.x -pp lead
- 4. Run pw2wannier90 to compute the overlap between Bloch states and the projections for the starting guess (written in the lead.mmn and lead.amn files).
  pw2wannier90.x < 05\_pw2wan.in > pw2wan.out
- 5. Run wannier90 to compute the MLWFs. wannier90.x lead

Inspect the output file lead.wout.

1. Use Wannier interpolation to obtain the Fermi surface of lead. Rather than re-running the whole calculation we can use the unitary transformations obtained in the first calculation and restart from the plotting routine. Add the following keywords to the lead.win file:

```
restart = plot
fermi_energy = [insert your value here]
fermi_surface_plot = true
```

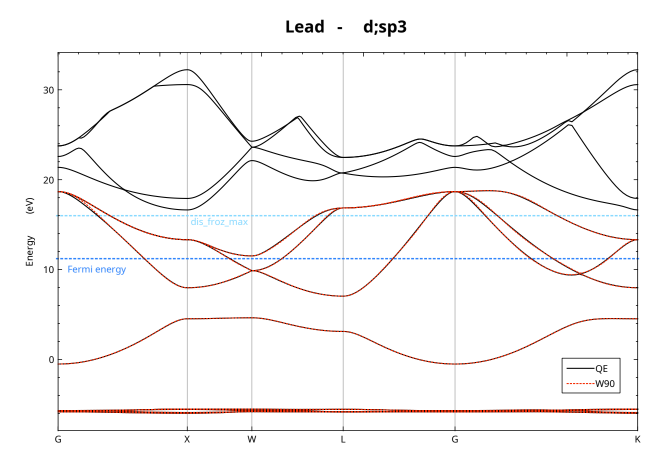
and re-run wannier90. The value of the Fermi energy can be obtained from the output of the initial first principles calculation. wannier90 calculates the band energies, through wannier interpolation, on a dense mesh of k-points in the Brillouin zone. The density of this grid is controlled by the keyword fermi\_surface\_num\_points. The default value is 50 (i.e., 50<sup>3</sup> points). The Fermi surface file lead.bxsf can be viewed using XCrySDen, e.g.,

```
xcrysden --bxsf lead.bxsf
```

2. Plot the interpolated band structure. A suitable path in k-space is

## Further ideas (if you have time)

- Compare the Wannier interpolated band structure with the full PWSCF band structure. Obtain MLWFs using a denser k-point grid. To plot the band structure you can use the PWSCF tool bands.x.
- Investigate the effects of the outer and inner energy windows on the interpolated bands.
- Instead of extracting a subspace of  $d+sp^3$  states, we could extract a different nine dimensional space (i.e., with s, p and d character). Examine this case and compare the interpolated band structures.
- Remove the low-energy d states from the wannierization (hint: use the exclude\_bands option in the Wannier90 input file) and compare both the spread and band structure you obtain.



Band Structure of lead showing the position of the Fermi energy and inner energy windows.