# Hands-on Tutorial on Electronic Structure Computations: Transport with Wannier functions

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exercises by Giovanni Pizzi (2013)

The files for this tutorial can be found in the archive:

/home/users/tutorial/user/public/Lab4\_Wannier/lab4\_wannier.tar.gz

Copy the content of this folder to your personal folder (do not create extra sub-folders) on the cluster; when you are inside your personal folder, type:

cp /home/users/tutorial/user/public/Lab4\_Wannier/lab4\_wannier.tar.gz ./

In your personal folder on the cluster, untar the archive with the command

tar -xzvf lab4\_wannier.tar.gz

This will create a folder Lab4\_Wannier in your personal directory.

Remember you can always check the online input description for quantum espresso pw.x code:

http://www.quantum-espresso.org/wp-content/uploads/Doc/INPUT PW.html

And download the Wannier90 tutorial at:

www.wannier.org

# Exercise 0 (don't do this): Running instructions if you want to make your cluster crash

The codes (Quantum Espresso, Wannier90) should be already in your path, to check this, issue the command:

which wannier90.x which pw.x

Similarly for other executables needed in the exercises below. You will afterwards call these executables by simply typing:

pw.x wannier90.x

(+) You can also run in parallel, if you want, by using (if you want to use 6 cpus):

mpiexec -np 6 pw.x < inputfile.in > outputfile.out

Then, your cluster will crash, most probably: (.

#### Running instructions if you want to be kind to your cluster

Inside each exercise folder, you have a set of files starting with run\_ (run\_pw.sh, run\_pw2wannier.sh, run\_wannier90.sh, run\_bands.sh, run\_plotband.sh).

Wherever in this tutorial you are asked to run an executable as in (+), use the script instead (note the disappearence of the symbols < and >)

```
./run_pw.sh inputfile.in outputfile.out
```

For the wannier90.x executable, there are two scripts, one has to be used when the wannier90.x -pp command is issued (there is no explicit output file here):

```
./run_wannier90pp.sh prefix.win
```

And the other one when it is called without -pp flag (also here no output file)

```
./run_wannier90.sh prefix.win
```

The script will work only if you have your Lab4 Wannier directory immediately inside your personal folder.

### Exercise 1 - Silicon valence bands

In this exercise we will learn to obtain the Wannier functions for the valence bands of silicon.

- Go to the ex1 folder. Inspect the input file O1\_scf.in. It is a ground-state calculation for a silicon crystal with two atoms per unit cell, with a FCC cell. Check if you understand all parameters (you can use the web page
  - http://www.quantum-espresso.org/wp-content/uploads/Doc/INPUT\_PW.html for keywords that you do not know). Visualize the unit cell with xcrysden (either opening the program and selecting from the menu File  $\rightarrow$  Open PWscf... $\rightarrow$  Open PWscf Input File and selecting the input file, or directly from the command line with the command xcrysden --pwi 01\_scf.in
- 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 pseudo/Si.pbe-n-van.UPF). With 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 (pw.x < 01\_scf.in > scf.out). You may want to use the parallelization to run the simulation faster, using (to use 8 processors)

```
mpiexec -np 8 pw.x < 01_scf.in > scf.out [./run_pw.sh 01_scf.in scf.out]
```

- When the calculation finishes, inspect the output file to check if there were any errors/warnings. Check your answer to the previous point (number of electrons and occupied valence bands).
- 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\_band.in. Do the following modifications to the file O2\_band.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 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 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 crystalline units), using 50 points per segment:
    - \*  $L(0.5, 0.5, 0.5) \rightarrow \Gamma(0, 0, 0)$
    - \*  $\Gamma(0,0,0) \to X(0.5,0,0.5)$

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. [./run\_pw.sh 02\_bands.in bands.out]
- When the calculation has finished, run the file O3\_bandplot.in through the bands.x executable (be sure to read and understand the input file):

mpiexec -np 8 bands.x < 03\_bandplot.in > bandplot.out[./run\_bands.sh 03\_bandplot.in bandplot.out]

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 abinitiobands.agr. When asked, call the 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 02\_band.in file that you created before to 04\_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  $4 \times 4 \times 4$  mesh, that will be used to calculate the discrete derivatives needed for the calculation of the Wannier functions. To obtain the list, 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:

```
kmesh.pl 4 4 4 [./run_kmesh.sh 4 4 4] (use the command without parameters to get an explanation of its usage). ON THE CLUSTER, RUN WITH ./run_kmesh.sh 4 4 4
```

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

```
mpiexec -np 8 pw.x < 04_nscf.in > nscf.out [./run_pw.sh 04_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 k coordinates, and the weight), Wannier90 wants only three numbers (the three coordinates). To obtain the 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 [./run_kmesh.sh 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 http://www.wannier.org/user\_guide.html page) for the input flags that you do not understand. Try to understand, in particular, the projections section. Can you say which is the location of the four s-like orbitals sitting with respect to the Si atoms?
- Now, we are ready to perform the Wannier calculation. This is done in three steps:

- 1. We first run a preprocess step using the command (it **must** be run in serial)

  wannier90.x -pp ex1 [./run\_wannier90pp.sh ex1]

  to produce a ex1.nnkp file, that contains the relevant information from the Wannier90 input file in a format to be used in the next step.
- 2. We will now run the pw2wannier90.x code (of the Quantum Espresso suite). The input file for pw2wannier90.x is provided (file 05\_pw2wan.in). We are asking the code to calculate the overlap matrices M<sub>mn</sub> (that will be written on the ex1.mmn file) and the A<sub>mn</sub> matrices (file ex1.amn). Since we want to plot the Wannier functions in real space, we need also the u<sub>nk</sub>(r) wavefunctions on a real-space grid. We thus also set the write\_unk flag in 05\_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 × 4 × 4 k-grid (Note: this is not needed for the Wannierization, 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

mpiexec -np 8 pw2wannier90.x < 05\_pw2wan.in > pw2wan.out[./run\_pw2wannier90.sh 05\_pw2wan.in
pw2wan.out]

3. Now we can finally run the Wannierization. Execute

wannier90.x ex1 [./run wannier90.sh ex1]

(it must be run in serial) 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 Wannier functions.
- To check if the obtained Wannier functions 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.sh); otherwise, to use xmgrace, you can run

xmgrace abinitiobands.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 them toward 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, ... 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.####)

## Exercise 2 - Silicon valence and conduction bands

- Copy all the ex1 folder to a new folder named ex2. The first step (01\_scf.in) is identical: if you copied also the out directory, you don't need to rerun it.
- In the O4\_nscf.in file, change the value of nbnd to 12 to calculate the eigenenergies and wavefunctions for 12 bands, and run the nscf calculation (using pw.x as above).
- Rename ex1. win to ex2. win, and modify the following flags:
  - Change num\_bands to 12 to be consistent with the new nscf run
  - Change the projections to 4 sp<sup>3</sup> orbitals for each Si atom in the unit cell: to do this, the projections section should read

begin projections
Si:sp3
end projections

- Change the num\_wann flag to the right number of Wannier functions: how many do we want, according to the projections list given above?
- Set the maximum energy for the frozen window (flag dis\_froz\_max) inside the energy gap (use the band plot obtained in exercise 1 to get a value for this flag) [6.5]
- 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 OK (check where this value lies in the band plot).
- Inside the file 05\_pw2wan.in, change the seedname to 'ex2' to reflect the new name of the .win file.
- Run wannier -pp ex2 [./run\_wannier90pp.sh ex2]
- Run pw2wannier90 using 05\_pw2wan.in as input file
- Run wannier ex2 [./run\_wannier90.sh ex2]
- Check the output:
  - Before the start of the Wannierization iterations, there is a new section (containing the string <--DIS) with the iterations of the disentanglement part. It is important that at the end of this section the convergence is achieved (with a string <<< Disentanglement convergence criteria satisfied >>>).
  - 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.
- Check final WF centers and verify that WF are real; you may also want to plot the Wannier functions, or compare the interpolated band structure with the ab-initio one (obtained in exercise 1).
- Optional part, to be done only at the end, if you have enough time: Are the Wannier functions as expected? (We would like  $4 \text{ sp}^3$ -like orbitals centered on each atom, with similar spreads). Try to rerun everything with a  $6 \times 6 \times 6$  kgrid for the nscf and Wannier90 step to check if the results improve.

### Exercise 3 - Linear C chain

We will study now ballistic transport in a linear chain of equispaced carbon atoms. Go to the ex3 folder.

- Inspect the scf input file O1\_scf.in. It is a linear chain of C atoms along the x axis, with C-C distance of 1.3 Å. Run the scf calculation with pw.x. How many electrons are there in the system? \_\_\_\_\_ (####Note: there may be a bug in old versions of xcrysden when the input cell is given in angstrom in the pw.x input file. As a workaround, run first pw.x, and then open the output file with xcrysden --pwo scf.out####)
- Copy the O1\_scf.in file to O2\_band.in, and modify the following (similarly to what was done in exercise 1):
  - change the calculation type from 'scf' to 'bands'
  - set the number of bands (nbnd) to 16
  - set the diago\_full\_acc flag to .true.
  - Set the k-point list to a line in k space (using K\_POINTS crystal\_b) from  $\Gamma = (0,0,0)$  to X = (0.5,0,0) [in crystal coordinates] with 50 points inbetween.
- Run pw.x using 02 band.in as input file. Check the output to verify that the calculation was successful.
- Use the provided O3\_bandplot.in file as input for bands.x, and then run bandplot.x as described in exercise 1 to produce a band plot.
- Copy 02\_band.in to 04\_nscf.in, and modify the following:

- change the calculation type from 'bands' to 'nscf'	
<ul> <li>change the kpoint list to a 6x1x1 mesh (use kmesh.pl to generate the k-point list), that we will use f the Wannierization.</li> </ul>	or
• Run pw.x on the 04_nscf.in input file. Check the output. Get the value of the Fermi energy from to output: eV. Using this value and the band plot obtained before: Has this chain a metallic insulating behavior?	
• Open the ex3.win file, which is a template of the Wannier90 input, try to understand the meaning of input flags, and replace the values marked with XXX. In particular:	all
- num_bands (use the value set in the nscf input file)	
<ul> <li>num_wann (understand which and how many projections were used by inspecting the projections section of the input file, and then set the correct value here)</li> </ul>	on
<ul> <li>Set the dis_froz_max to the value you got for the Fermi energy, and explain why this is a reasonal choice:</li> </ul>	ole
<ul> <li>Using the band plot, set the window range for transport (tran_win_max and tran_win_min) large enouge to include the whole band plot.</li> </ul>	gh
- Set the $mp\_grid$ value to the correct grid size $(6 \times 1 \times 1)$ , and fill the kpoints section with the 6 kpoint (again, use the $kmesh.pl$ script)	$_{ m its}$
• Check the 05_pw2wan.in input file for pw2wannier90.x, then:	
- Run wannier90.x -pp ex3 (check the output)	
- Run pw2wannier90.x using 05_pw2wan.in as input (check the output)	
- Run wannier90.x ex3	
• Check the output of the wannierization (if the disentanglement convergence was achieved, if the Wannerizatic cycles reached convergence, if the Wannier functions are real). If everything went smoothly, you should get atom-centered p-like orbitals, and 2 mid-bond s-like orbitals, with smaller spread. (###Try to plot wixcrysden one WF for each type, to see how they look like.####)	4
• Compare the interpolated band structure with the ab-initio one.	
<ul> <li>Using the knowledge of the symmetry of the system and the degeneracy of the bands, can you say whi are the "p" bands and which are the "s" bands?</li> </ul>	$\mathrm{ch}$
<ul> <li>Do the interpolated band structure and the ab-initio band structure agree within the frozen window And outside? Were you expecting this?</li> </ul>	w?
Using the obtained band structures, can you give an estimate of the lower bound for dis_win_max th you expect to be needed to get converged Wannier functions?	at
• In the ex3.wout file, there is a new section called "Transport". Inspect the results and try to understand output.	its
• The activation of the transport flag also produces two new files: ex3_dos.dat (the Density Of States) as ex3_qc.dat (the Quantum Conductance). Plot them together using the command xmgrace ex3_dos.dat ex3_qc.dat (if you use this command, the black curve will be the DOS and the red one will be the QC).	nd

- Can you explain the values of the quantum conductance? \_\_\_\_\_

- Compare the DOS (black curve) with the band structure you plotted before. Were you expecting divergences at the band edges? Why? \_\_\_\_\_\_\_ Is the DOS behavior compatible with the band structure?

#### Exercise 3 bis - Dimerized C chain

(Do this exercise only if you have enough time, otherwise go to exercise 4)

- Copy the ex3 folder to ex3bis.
- Do all as you did for exercise 3, but changing the atomic coordinates of the C atoms in the files 01\_scf.in, 02\_band.in, 04\_nscf.in and ex3.win such that the x coordinates are the following: 0.6775 and 1.9225 (we are changing the positions of the C atoms such that now the interatomic C-C distances alternate with a shorter and a longer distance (a single and a triple bond).
- What do you expect that will happen to the band structure? Do you expect that you can use similar parameters for the Wannierization? \_\_\_\_\_
- Can you explain the difference in the DOS and in the quantum conductance with respect to exercise 3?
- You can compare the centers and the spreads of the Wannier functions of exercise 3 and 3 bis, and also visually inspect them to understand if there is a significant change or if they remain similar.

### Exercise 4 - Linear C chain with a defect

- Inspect the input file O1\_scf.in, understanding the input flags; (###use also xcrysden to visualize the structure (as already discussed before, due to a bug in xcrysden, you should open the *output* pw.x file, after running the scf step)###). It is a C chain with 22 atoms, where the distance of C atoms is the same of exercise 3, except for the two central atoms which are nearer. How many k-points are being used? Is it a resonable approximation? \_\_\_\_\_\_ (In any case, note, that for the lcr transport in Wannier90, Γ-sampling only is required).
- How many electrons are there in the system?
- Inspect the O2\_nscf.in file: the number of bands (nbnd) value is missing: put the correct value (you should deduce this value from the one used in exercise 3 and considering the number of atoms in the unit cell).
- Before opening the ex4.win file: which initial projections would you use, knowing the results of exercise 3 (and 3 bis)? \_\_\_\_\_
- Open the ex4.win file, inspect it, and replace the values marked with XXX. In particular:
  - num\_bands
  - num\_wann (deduce it with the same method you used for num\_bands, and then verify your result using the list of projections provided in the input file)
  - dis\_froz\_max and dis\_win\_max (hint: how are they related to the values used in example 3?)
  - tran\_win\_max and tran\_win\_min (hint: how are they related to the values used in example 3?)
- Inspect the O3\_pw2wan.in file. Note that for lcr transport calculations, we need a new flag: write\_unkg = .true. (the Wannier90 documentation explains why)
- run the scf calculation, followed by the nscf calculation, the wannier90 preprocess run, the pw2wannier90.x calculation, and finally the Wannierization.
- Check the output of the Wannierization, verifying the convergence and the results (WF centers, spreads). You may want to plot (some of) the Wannier functions: to do this modify 03\_pw2wan.in (adding the flag write\_unk = .true.) and ex4.win to plot also (some of the) Wannier functions (add the wannier\_plot and wannier\_plot\_list flags; see the Wannier90 manual for the documentation; take also into account the possibility of using the restart=plot flag.)
- Plot the DOS and the QC; compare the QC with the QC obtained in exercise 3.
- Check the decay of matrix elements inspecting the ex4\_hr.dat file (produced using to the plot\_hr flag) that contains the Hamiltonian in real space between Wannier functions (read the Wannier90 documentation for the explanation of the file format). Is the value dist\_cutoff=5Å that we used in the Wannier90 input flag sufficient, or too small? \_\_\_\_\_\_