User guide of Wannier_tools

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1. Installaion

- (a) Check out the repository by "git clone https://github.com/quanshengwu/wannier_tools.git" or download the zip file.
- (b) cd wannier_tools/soc directory, Edit Makefile, Change the blas library " libs= " to your mkl library, you can use your own blas and lapack library. Then type make. The executive binary is copied to directory wannier_tools/bin
- (c) add the wannier_tools/bin path to your PATH by "export PATH=yourdirecory/wannier_tools/bin:\$PATH"

2. The introduction of input file

Before execute wann_tools, you should edit the input file in the directory wannier_tools/example by your own necessary. Here we take Bi2Se3 as an example.

At present, The input file is not very flexible. It is a normal fortran free format file.

```
1 wannier90 hr.dat!input file
2 output
                ! output file's filename
3| T
              ! BulkBand calc
41 F
              ! SlabBand calc
5| F
              ! WireBand calc
6| F
              ! SlabSS calc
7| F
              ! SlabArc calc
8 F
              ! SlabSpintexture_calc
9| F
              ! wanniercenter calc
10| F
               ! berry
11 40
               ! Nk
```

```
12 100
               ! omega number
 13 -2.00 2.00
                 ! omegamin omegamax
 14 0
              !E arc
 15 10
              ! nslab
 16| 1
              ! np
 17 18
             ! NumOccupied
 18 18
              ! ntotch
 19| 2
              ! soc
              ! eta
 20 0.001
 21 4.4195
               ! e-fermi
 22 0.0
               ! surf onsite
 23 000
              ! Bx By Bz
 24 -2.069 -3.583614 0.000000
                               ! crystal lattice information
 25 | 2.069 -3.583614 0.000000
 26 0.000 2.389075 9.546667
                       ! number of atoms in unit cell
 27 5
 28 Direct
                        ! Direct or Cartesian coordinate
 29 Bi 0.3990 0.3990 1.1970
 30 Bi 0.6010 0.6010 1.8030
 31 Se 0 0 0
 32 Se 0.2060 0.2060 0.6180
 33 Se 0.7940 0.7940 2.3820
 34 3 3 3 3 ! number of projectors
 35 Bi px py pz ! projectors
 36 Bi px py pz
 37 Se px py pz
 38| Se px py pz
 39 Se px py pz
 40
                ! number of k line only for bulk band
 41 G 0.00000 0.00000 0.0000 X 0.50000 0.00000 0.5000 !k
point path
 42 X 0.50000 0.00000 0.5000 W 0.75000 0.25000 0.5000
 43 W 0.75000 0.25000 0.5000 L 0.50000 0.50000 0.5000
 44 L 0.50000 0.50000 0.5000 G 0.00000 0.00000 0.0000
 45 G 0.00000 0.00000 0.0000 K 0.75000 0.37500 0.3750
 46 100
          ! Umatrix for new lattice 001
 47 010
```

line 1: wannier90_hr.dat ! input file

The file name of wannier90_hr.dat, which is generate from wannier90 package (http://www.wannier.org).

Currently, we support two formats, one is from wannier90. One is from Openmx, which rewrite wannier90_hr.dat.

line 2: output! output file

Filename for some outputs. You can check this file for running status.

From line 3-10: give some control parameters for this program. The value should be F or T relate t False and True respectively.

Flag option	Function	Output	Test
BulkBand_calc	Band structure for bulk	bulkek.dat, bulkek.gnu	yes
SlabBand_calc	Band structure for slab system	slabek.dat,slabek.gnu	yes
WireBand_calc	Band structure for ribbon system	ribbonek.dat,ribbonek.g nu	yes
SlabSS_calc	Surface spectrum A(k,E) along a k line and energy interval	dos.dat_l, dos.dat_r surfdos_l.gnu,surfdos_r. gnu	yes
SlabArc_calc	Surface spectrum A(k1,k2,E0) for a fixed	arc.dat_l, arc.dat_r arc_l.gnu,arc_r.gnu	yes

	energy E0		
SlabSpintexture_cal	calculate spin texture for slab system	spindos.dat	No
wanniercenter_calc	calculate wannier centers for slabs	wanniercenterky0.dat	No
berry_calc	calculate berry phase along one k line		No

For Bulkband calculation, you should specify number of k lines and the special klines in line

41-45 (these line number is relatively). The output for this subroutine are bulkek.dat and bulkek.gnu. You

can run "gnuplot bulkek.gnu", or "xmgrace bulkek.dat" to get bandstucture plot.

For Slab band calculation, you only need to change Nk in line 11 to get more points for k line.

The output for this subroutine are slabek.dat and slabek.gnu. You can run "gnuplot slabek.gnu", or "xmgrace slabek.dat" to get bandstucture plot. If you use gnuplot, you should check the png files.

The procedure for ribbon band calculation is similar to slab band calculation. the data is different by the names.

For surface state calculation, the output files are dos.dat_l, dos.dat_r, surfdos_l.gnu, surfdos_r.gnu. dos.dat_l and dos.dat_r are the spectrum weight for each k point and each energy, related to the top and bottom surface state respectively. You can run "gnuplot surfdos_l.gnu" to get surfdos_l.png.

For Fermi arc calculation, basically, the principle for this calculation is similar to surface state

calculation. The difference is to get spectrum A(k, \$\omega\$) for a fixed energy \$\omega\$ but (kx, ky) in

2D in this calculation, while get get spectrum A(k, \$\omega\$) for some special k lines and an energy interval for the surface state calculation.

SlabSpintexture_calc and wanniercenter_calc are not tested yet.

line 11: 40 ! Nk

Number of k points for any necessary subroutines, like bulk, slab, ribbon band structure calculation

line 12:500 ! omega number

Number of energies when calculate surface state, this value determines

the resolution of the surface state pictures. If it's large, the calculation will be slow linearly scaling.

You can set it to be 200 at first, if the resolution of the picture is not enough, then increase it.

line 13: -2.00 2.00 ! omegamin omegamax

Energy interval for surface state calculation.

line 14: 0 ! E_arc

Energy at which the fermi arc is calculated.

line 15: 10 ! nslab

Number of slabs for slab band. If you calculate the band structure for nanoribbon, it represent the thickness of one direction. The thickness for the other direction is Np.

see Np.

line 16: 1 ! Np

Number of principle layers for calculating surface state and fermi arc. If your wannier function is not enough localized. For instance, there are big hoppings beyond the home unit cell. Then we should set this value to 2 or 3 or 4. In principle, you should check the convergency of the surface state by increasing this value. The computation will be cubically grows as a functional of Np. at the beginning, you can set Np=1.

If you calculate band structure for nanoribbon, then this value is the thickness of another direction of nanoribbon. See slab.

line 17:18 ! NumOccupied

This value is not valid at present, leave it alone. However, don't delete this line.

line 18: 18 ! Ntotch

number of electrons, if you want to determine the fermi level, this value will work.

line 19: 1 ! soc

The flag for soc, if there is no spin-orbital included in your wannier90_hr.dat, then set it to be zero. else, set it to be 1.

line 20: 0.001 ! eta

Broadening of green's function when calculate the surface states. If you don't know what it is, leave it unchanged.

line 21: 4.4195 ! e-fermi in eV

Fermi level of your wannier90_hr.dat.

line 22-24:

```
-2.069 -3.583614 0.000000 ! crystal lattice information 2.069 -3.583614 0.000000 0.000 2.389075 9.546667
```

Three lattice vectors coordinates.

line 25-33:

```
      5
      ! number of atoms in unit cell

      Direct
      ! Direct or Cartesian coordinate

      Bi 0.3990
      0.3990
      1.1970

      Bi 0.6010
      0.6010
      1.8030

      Se 0
      0
      0

      Se 0.2060
      0.2060
      0.6180

      Se 0.7940
      0.7940
      2.3820
```

Set the number of atoms in your wannier90_hr.dat when you constructed it.

Set the coordinate system type, Direct or Cartesian Set the name and coordinate of each atoms. this coordinate is based on the three lattice vectors. It's easy to find it for through the wannier90.win.

line 34- 39:

```
3 3 3 3 3 ! number of projectors
Bi px py pz
Bi px py pz
Se px py pz
Se px py pz
Se px py pz
Se px py pz
```

Number of projectors for each atom when you construct your wannier functions. And the name of projectors.

line 40-45:

5 ! number of k line\\

G 0.00000 0.00000 0.0000 X 0.50000 0.00000 0.5000 ! k point path\\
X 0.50000 0.00000 0.5000 W 0.75000 0.25000 0.5000\\
W 0.75000 0.25000 0.5000 L 0.50000 0.50000 0.5000\\
L 0.50000 0.50000 0.5000 G 0.00000 0.00000 0.0000\\
G 0.00000 0.00000 0.00000 K 0.75000 0.3750

Number of high-symmetric k point lines. Look the format carefully. This k lines are in unit of the reciprocal lattice built by the lattice vector given in line 24-26

line 46-48:

1 0 0 ! Umatrix for new lattice a11, a12, a13\\

0 1 0 ! a21 a22 a23\\

0 0 1 ! a31 a32 a33

These three lines are very important for slabs and ribbon calculation. It defines the surface we needed. Basically, you should know of which surface you want to get the surface, and of which direction you want to simulate the ribbon. So we need to define the new lattice vector system like this, choose two vectors on the surface we want to study, and choose another vector which is not in this plane. The volume of the cell constructed by these three vectors should be the same of the previous one. And the orientation should be the same by checking \$R_1'\cdot(R_2'\times R_3')= R_1\cdot(R_2\times R_3)\$. The relations between old and new basis are

\begin{eqnarray}

$$R_1'=a_{11} R_1 + a_{12} R_2 + a_{13} R_3 \ R_2'=a_{21} R_1 + a_{22} R_2 + a_{23} R_3 \ R_3'=a_{31} R_1 + a_{32} R_2 + a_{33} R_3 \ \end{eqnarray}$$

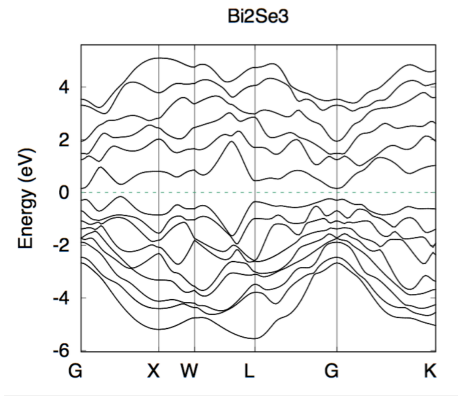
The slab calculations are base on the surface constructed by vector \$R_1', R_2'\$. The ribbon is along \$R_3'\$ direction, two surfaces are constructed by vector \$R_1', R_3'\$ and \$R_2', R_3'\$ respectively.

\end{enumerate}

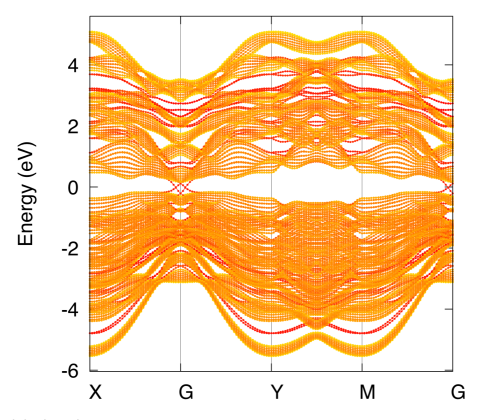
3. Examples (Bi2Se3)

The wannier90_hr.dat is compressed as Bi2Se3_hr.tar.gz in examples/ folder. You can decompressed by a command "tar xzvf Bi2Se3_hr.tar.gz". The input.dat file can also be found in that folder.

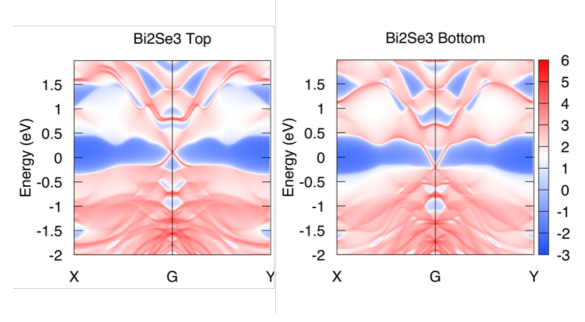
(1). Bulk band structure



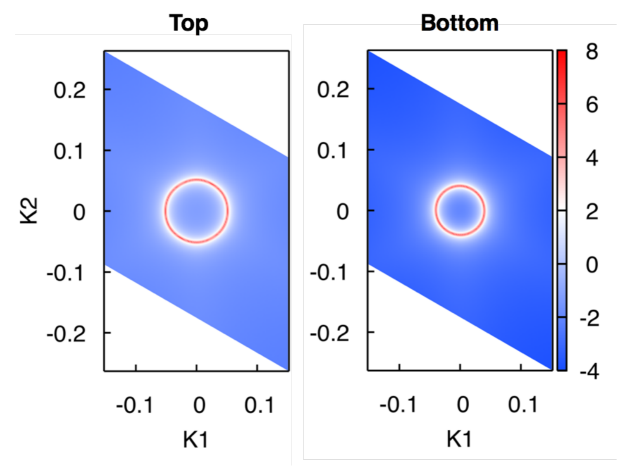
(2). (001) oriented slabs.



(3). (001) surface state



(4). (001) surface spectrum for a fixed energy E=0eV



Comment: The Wannier TB constructed here has not perfect symmetry, which will cause different chemical potential on both surfaces. So The top and bottom surface states are not exactly the same. However the topological properties will not be changed by change the surface chemical potential.

4. License and agreement.

If you use our code, please cite this website "Q.S.Wu, https://github.com/quanshengwu/wannier_tools" . If you have good ideas to improve this code, do not hesitate to contact me. Your contribution will be recorded.

5. Enjoy and good luck.