PYATB v1.0.0

PYATB

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CHAPTER

ONE

INTRODUCTION



PYATB (Python ab initio tight binding simulation package) is an open-source software package designed for computing electronic structures and related properties based on the ab initio tight binding Hamiltonian. The Hamiltonian can be directly obtained after conducting self-consistent calculations with first-principles packages using numerical atomic orbital (NAO) bases, such as ABACUS. The package comprises three modules - Bands, Geometric, and Optical, each providing a comprehensive set of tools for analyzing different aspects of a material's electronic structure.

The Bands module enables users to calculate essential properties of band structures, including the partial density of states (PDOS), fat bands, Fermi surfaces, Wyel/Dirac points, and more. Additionally, the band unfolding method is utilized to obtain the energy band spectra of a supercell by projecting the electronic structure of the supercell onto the Brillouin zone of the primitive cell.

In the Geometric module, users have access to tools for computing Berry phase and Berry curvature-related quantities, such as electric polarization, Wilson loops, Chern numbers, anomalous Hall conductivities, and more.

The Optical module offers a range of optical property calculations, including optical conductivity and nonlinear optical responses, such as shift current and Berry curvature dipole.

PYATB is licensed under GPLv3.0, making it freely available for use and modification by the scientific community. The main developers of PYATB are Gan Jin, Hongsheng Pang, Yuyang Ji, Zujian Dai, under the supervision of Prof. Lixin

He at the University of Science and Technology of China.

1.1 Capabilities

PYATB provides three major modules: *Bands module*, *Geometric module*, *Optical module*, each with its own set of functions:

· Bands module

Band structre

Allows users to calculate the energy bands and wave functions using three different k-point modes: k-point, k-line, and k-mesh.

- Band unfolding

Calculates the spectra weight by unfolding the energy bands of a supercell into the Brillouin zone (BZ) of the primitive cell.

- Fermi energy

Calculates the Fermi energy at a given temperature.

- Fermi surface

Plots the Fermi surface.

- Find nodes

Allows users to search for degenerate points of the energy bands in the BZ within a specified energy window. This function can be used to find the Weyl/Dirac points in Weyl/Dirac semimetals.

- DOS and PDOS

Calculates the density of states (DOS) and partial density of states (PDOS) of particular orbitals.

- Fat band

Provides the contribution of each atomic orbital to the electronic wave functions at each k-point in the BZ.

- Spin texture

Plots the spin polarization vector as a function of momentum in the BZ.

• Geometric module

- Wilson loop

Enables users to calculate the \mathbb{Z}_2 number by tracking the Wannier centers along the Wilson loop.

- Electric polarization

Evaluates the electric polarization in various directions for non-centrosymmetric materials based on the Berry phase theory.

- Berry curvature

Computes the Berry curvature in the BZ.

- Anomalous Hall conductivity

Calculates the anomalous Hall conductivity using Berry curvature.

- Chern number

Calculates the Chern number of a system for any given k-plane.

Chirality

Examines the chirality of Weyl points by calculating the Berry curvature on a sphere around the k point.

· Optical module

1.1. Capabilities 2

- JDOS

Calculates the joint density of states (JDOS), which characterizes both electronic states and optical transitions.

- Optical conductivity and dielectric function

Calculates the frequency-dependent optical conductivity and dielectric function.

- Shift current

Calculates the shift current conductivity tensor for the bulk photovoltaic effect.

- Berry curvature dipole

Calculates the Berry curvature dipole which leads to the nonlinear anomalous Hall effects.

1.2 Methodology

PYATB is based on the ab initio tight binding model, where the parameters of the Hamiltonian are generated directly from the self-consistent calculations using first-principles software based on numerical atomic orbitals (NAO) bases, such as ABACUS.

NOTE: ABACUS (Atomic-orbital Based Ab-initio Computation at UStc) is an open-source package based on density functional theory (DFT). For a detailed introduction, please refer to https://abacus.ustc.edu.cn/.

In a periodic system, the Kohn-Sham equation at a given k point can be written as,

$$H|\Psi_{n\mathbf{k}}\rangle = E_{n\mathbf{k}}|\Psi_{n\mathbf{k}}\rangle.$$

Here $\Psi_{n\mathbf{k}}$ is the Bloch wave function of the n-th band, and can be expressed under NAO as,

$$|\Psi_{n\mathbf{k}}\rangle = \frac{1}{\sqrt{N}} \sum_{\mu} C_{n\mu}(\mathbf{k}) \sum_{\mathbf{R}} \mathrm{e}^{i\mathbf{k}\cdot\mathbf{R}} |\mathbf{R}\mu\rangle,$$

where $|\mathbf{R}\mu\rangle \equiv \phi_{\mu} (\mathbf{r} - \tau_{\mu} - \mathbf{R})$ is the μ -th atomic orbital, in the \mathbf{R} -th unit cell, and τ_{μ} denotes the center position of this orbital. The composite index $\mu = (\alpha, i, \zeta, l, m)$, where α is the element type, i is the index of the atom of each element type, ζ is the multiplicity of the radial functions for the angular momentum l, and m is the magnetic quantum number. The coefficient of the NAO is given by $C_{n\mu}(\mathbf{k})$.

Under the NAO base, the Kohn-Sham equation becames a eigenvalue problem,

$$H(\mathbf{k})C_n(\mathbf{k}) = E_{n\mathbf{k}}S(\mathbf{k})C_n$$
.

where $H(\mathbf{k})$, $S(\mathbf{k})$ and $C_n(\mathbf{k})$ are the Hamiltonian matrix, overlap matrix and eigenvectors of the n-th band, respectively.

To obtain the $H(\mathbf{k})$ and $S(\mathbf{k})$, we first calculate tight binding Hamiltonian in real space via first-principles softwares based on NAOs, such as ABACUS,

$$H_{\nu\mu}(\mathbf{R}) = \langle \mathbf{0}\nu | H | \mathbf{R}\mu \rangle ,$$

$$S_{\nu\mu}(\mathbf{R}) = \langle \mathbf{0}\nu | \mathbf{R}\mu \rangle .$$

Once we have the $H_{\nu\mu}(\mathbf{R})$ and $S_{\nu\mu}(\mathbf{R})$, we can obtain the Hamiltonian matrix and the overlap matrix at arbitrary \mathbf{k} points using the following relation,

$$\begin{split} H_{\nu\mu}(\mathbf{k}) &= \sum_{\mathbf{R}} \mathrm{e}^{i\mathbf{k}\cdot\mathbf{R}} H_{\nu\mu}(\mathbf{R})\,, \\ S_{\nu\mu}(\mathbf{k}) &= \sum_{\mathbf{R}} \mathrm{e}^{i\mathbf{k}\cdot\mathbf{R}} S_{\nu\mu}(\mathbf{R})\,, \end{split}$$

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When solving for the geometric properties of the bands, we also need the dipole matrix between the NAOs, namely:

$$r_{\nu\mu,a}(\mathbf{R}) = \langle \mathbf{0}\nu | r_a | \mathbf{R}\mu \rangle \quad a = x, y, z.$$

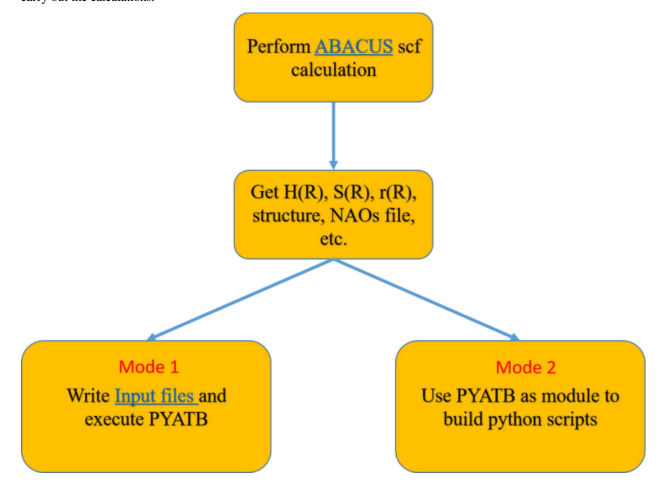
The dipole matrix $\mathbf{A}_{\nu\mu}^{R}(\mathbf{k})$ can then be obtained by Fourier transform,

$$A^R_{\nu\mu,a}(\mathbf{k}) = \sum_{\mathbf{R}} \mathrm{e}^{i\mathbf{k}\cdot\mathbf{R}} r_{\nu\mu,a}(\mathbf{R}).$$

After obtaining the tight binding parameters $H_{\nu\mu}(\mathbf{R})$, $S_{\nu\mu}(\mathbf{R})$, and $r_{\nu\mu,a}(\mathbf{R})$ from the first principles software, the electronic structures and related physical propertie can be calculated using PYATB. More description of the methods used in PYATB can be found in Ref. ???

1.3 Workflow

The workflow of PYATB involves several key steps. First,one need to perform self-consistent calculations using ABAUCS to generate tight binding Hamiltonan, including $H_{\nu\mu}(\mathbf{R})$, $S_{\nu\mu}(\mathbf{R})$, and $r_{\nu\mu,a}(\mathbf{R})$. Some of the functions may require crystal structur file and atomic orbital data. Once all the necessary files are obtained, the user can either write an Input file to perform the corresponding function computation using PYATB or write a script utilizing the PYATB module to carry out the calculations.



1.3. Workflow 4

CHAPTER

TWO

INSTALLATION

2.1 Download

The latest version of the PYATB package can be obtained from the GitHub repository:

git clone https://github.com/pyatb/pyatb.git

2.2 Prerequisites

Currently, PYATB is only supported on Linux systems and has not been tested on Windows or Mac systems. To use PYATB, the following prerequisites must be installed:

- Python 3.7 or newer
- C++ compiler
- Intel MKL
- pybind11
- Eigen3
- NumPy
- · SciPy
- mpi4py
- Matplotlib

2.3 Install

1. Before installing PYATB, you need to install the pybind11 module and mpi4py module:

```
pip install pybind11
```

and

pip install mpi4py

2. You can set the addresses of the C++ compiler (non-MPI version), MKL library, and Eigen library through the siteconfig.py file, and modify the relevant variables to point to the correct paths for your system.

- To set the C++ compiler, you can modify the variable "compiler" in the siteconfig.py. For example, you can set compiler = 'icpc'.
- To set the MKL library, you can modify the variable "mkl_library_dir" and "mkl_include_dir" in the siteconfig.py.
- To set the Eigen library, you can modify the variable "eigen_include_dir" in the siteconfig.py. Eigen is a C++ template library for linear algebra. You can find and download it on the official website Eigen.
- 3. After completing the above preparatory work, you can install PYATB with the following command:

```
pip install ./
```

4. After completing the installation process, you can access the pyatb executable and corresponding module, which can be imported using the import pyatb command.

2.4 Related to the Intel MKL library

If you encounter the following problem when running pyatb using the intel oneapi MKL library under Anaconda:

```
undefined symbol: mkl_sparse_optimize_bsr_trsm_i8
```

This problem may be caused by a conflict between the mkl library in Anaconda and the mkl library in intel oneapi. You can solve it by using the following method:

```
export LD_PRELOAD=/opt/intel/oneapi/mkl/2022.0.2/lib/intel64/libmkl_def.so.2:\
/opt/intel/oneapi/mkl/2022.0.2/lib/intel64/libmkl_avx2.so.2:\
/opt/intel/oneapi/mkl/2022.0.2/lib/intel64/libmkl_core.so:\
/opt/intel/oneapi/mkl/2022.0.2/lib/intel64/libmkl_intel_lp64.so:\
/opt/intel/oneapi/mkl/2022.0.2/lib/intel64/libmkl_intel_thread.so:\
/opt/intel/oneapi/compiler/2022.0.2/linux/compiler/lib/intel64_lin/libiomp5.so
```

Note that you need to replace the MKL path with your own environment. For libiomp5.so, its location may sometimes be in the directory where the intel compiler is located.

Another solution is to use the MKL library in the Anaconda virtual environment. If you cannot find it, you can install the MKL library and its include files using the following commands:

```
conda install -c conda-forge mkl conda install -c conda-forge mkl-include
```

CHAPTER

THREE

RUN

PYATB supports mixed parallelism of OpenMP and MPI, and you need to determine the number of threads and processes to run depending on the actual configuration of your computer.

For example, set the number of threads to 2,

export OMP_NUM_THREADS=2

and then use 6 processes to run PYATB,

mpirun -n 6 pyatb

BRIEF INTRODUCTION OF THE INPUT FILES

4.1 Input

The Input file describes the basic information about the system and the parameters required to calculate the function. For a complete list of the input parameters, please consult this *input list*.

The following is an example of a Input file:

```
INPUT_PARAMETERS
   nspin
                                   4
   package
                                   ABACUS
   fermi_energy
                                   9.481194886038594
   fermi_energy_unit
   HR_route
                                   data-HR-sparse_SPIN0.csr
   SR_route
                                   data-SR-sparse_SPIN0.csr
                                   new-data-rR-tr_SPIN4
   rR_route
   HR_unit
                                  Ry
                                   Bohr
   rR_unit
                                   8000
   max_kpoint_num
}
LATTICE
   lattice_constant
                                  1.8897162
   lattice_constant_unit
                                  Bohr
   lattice_vector
    -2.069 -3.583614 0.000000
    2.069 -3.583614 0.000000
    0.000 2.389075 9.546667
}
BAND_STRUCTURE
                                   0
   wf_collect
   kpoint_mode
                                   line
   kpoint_num
   high_symmetry_kpoint
   0.00000 0.00000 0.0000 100 # G
   0.00000 0.00000 0.5000 100 # Z
   0.50000 0.50000 0.0000 100 # F
   0.00000 0.00000 0.0000 100
    0.50000 0.00000 0.0000 1
```

The Input file consists of three main sections (INPUT_PARAMETERS, LATTICE, FUNCTION), each of which is represented by the keyword + {}, namely:

```
KEYWORD
{
   parameter xx
}
```

The **INPUT_PARAMETERS** section is utilized to specify public parameters and indicate the HR, SR, and rR files required for the calculation.

The **LATTICE** section outlines lattice information and includes three parameters: lattice_constant, lattice_constant_unit, and lattice_vector.

The **FUNCTION** section lists the parameters necessary for each calculation function. These parameters differ depending on the specific function being used.

NOTE:

- The keyword of the section must be capitalized, and the parameters in the section must be lowercase.
- Each parameter value is provided by specifying the name of the input variable and then putting the value after the name, separated by one or more blank characters(space or tab).
- Any line starting with # or // will be ignored. To ignore a function section, add # in front of the section name. This will cause all parameters within that function section to be ignored.
- Proper case is required to avoid reading errors. For example, eV must not be written as ev, and Bohr must not be written as bohr.

4.2 HR, SR, rR

HR, SR, and rR are important input files for PYATB and contain information about the tight binding model, which can be generated by ABACUS.

In the PYATB program, if only the *Bands module* is used, it is enough to provide HR and SR, and specifying rR is not required. However, when the functions of the calculations are related to the *Geometric module* or *Optical module*, the rR file must be specified.

4.2.1 How to generate HR, SR, rR files using ABACUS

Run the scf calculation of ABACUS normally, specifies the parameters out_mat_hs2 and out_mat_r in the IN-PUT file, and three files of sparse format data-HR-sparse_SPIN0.csr, data-SR-sparse_SPIN0.csr, data-rR-sparse_csr (and data-HR-sparse_SPIN1.csr when nspin is 2), will be generated in OUT* folder.

NOTE:

• nspin=4 and nspin=1, nspin=2 output different files. nspin=4 files contain imaginary numbers, so the correct nspin parameter needs to be strictly specified when executing pyatb programs.

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4.3 Other file

For the calculation of some functions, structure files and NAOs (numerical atomic orbital bases) files are also required.

• The structure file

This file describes the structural information about the system, e.g., lattice constant, lattice vectors, and positions of the atoms within a unit cell. The structure file format of ABACUS is currently used.

• The NAOs files

This file describes the numerical atomic orbitals used to expand the Hamiltonian. The NAOs file format of ABACUS is currently used.

4.3. Other file 10

FUNCTIONS

5.1 BAND_STRUCTURE

An example of calculating the energy band structure of the topological insulator Bi_2Se_3 is provided in the folder examples/Bi2Se3.

The Input file is:

```
INPUT_PARAMETERS
   nspin
   package
                                   ABACUS
   fermi_energy
                                   9.557219691497478
   fermi_energy_unit
   HR_route
                                   data-HR-sparse_SPIN0.csr
   SR_route
                                   data-SR-sparse_SPIN0.csr
                                   data-rR-sparse.csr
   rR_route
   HR_unit
                                   Ry
                                   Bohr
   rR_unit
                                   8000
   max_kpoint_num
}
LATTICE
   lattice_constant
                                  1.8897162
   lattice_constant_unit
                                  Bohr
   lattice_vector
    -2.069 -3.583614 0.000000
    2.069 -3.583614 0.000000
    0.000 2.389075 9.546667
}
BAND_STRUCTURE
                                   0
   wf_collect
   kpoint_mode
                                   line
   kpoint_num
   high_symmetry_kpoint
   0.00000 0.00000 0.0000 100 # G
   0.00000 0.00000 0.5000 100 # Z
   0.50000 0.50000 0.0000 100 # F
    0.00000 0.00000 0.0000 100
    0.50000 0.00000 0.0000 1
```

wf_collect: Whether to output wave function matrix information. The wave function file stores the expansion coefficients of NAOs.

There are three ways to set k-points: k-point, k-line, and k-mesh, with the keyword kpoint_mode used to define the mode. The setting parameters for each mode differ, so please refer to the INPUT for detailed instructions. In this example, the line mode is used. In this mode, kpoint_num specifies the number of high-symmetry points, and high_symmetry_kpoint records the direct coordinates of these points and the number of k-points between each pair of high-symmetry points. Each row in the setting consists of four numbers, where the first three indicate the coordinates, and the last number specifies the number of k-points between the given k-point and the next high symmetry point.

After the calculation is done, a number of files will be generated in the Out/Band_Structure folder, including kpt.dat and band.dat. If kpoint_mode is set to line, the output will also include the energy band diagram and its corresponding drawing script.

5.2 BANDUNFOLDING

5.2.1 introduction

Band unfolding is a method of projecting the wave function of the supercell to the coupled k points in the original unit cell to obtain the spectral function.

The relationships between the lattice vectors of the large cell (A) and primitive cell (a) are given by

$$\begin{bmatrix} A_1 \\ A_1 \\ A_3 \end{bmatrix} = \begin{bmatrix} m_{11} & m_{12} & m_{13} \\ m_{21} & m_{22} & m_{23} \\ m_{31} & m_{32} & m_{33} \end{bmatrix} \begin{bmatrix} a_1 \\ a_2 \\ a_3 \end{bmatrix}$$

Detailed descriptions can be found in First-principles calculations of the surface states of doped and alloyed topological materials via band unfolding method.

5.2.2 example

Here is an example located in the examples/NV folder that showcases how to calculate the spectral function of the nitrogen-vacancy (NV) center in diamond.

The Input file is:

```
INPUT_PARAMETERS
    nspin
                                     1
   package
                                     ABACUS
                                     15.46412271260700
    fermi_energy
   fermi_energy_unit
   HR_route
                                     data-HR-sparse_SPIN0.csr
   SR_route
                                     data-SR-sparse_SPIN0.csr
   rR_route
                                     data-rR-sparse.csr
   HR_unit
                                     Ry
    rR_unit
                                     Bohr
}
LATTICE
    lattice_constant
                                     1.8897162
```

(continues on next page)

```
lattice_constant_unit
                                    Bohr
   lattice vector
    7.13366 0 0 #latvec1
    0 7.13366 0 #latvec2
    0 0 7.13366 #latvec3
BANDUNFOLDING
                                    STRII
    stru_file
                                    60
   ecut
   band_range
                                    10 250
   m_matrix
                                    -2 2 2 2 -2 2 2 2 -2
   kpoint_mode
                                    line
   kpoint_num
   high_symmetry_kpoint
   0.500000 0.000000 0.500000 300 # X
    0.500000 0.250000 0.750000 300
                                     # W
    0.500000 0.500000 0.500000 300
    0.000000 0.000000 0.000000 300
                                     # Gamma
    0.500000 0.000000 0.500000 1
```

stru_file: The structure file name of the supercell. This file indicates the crystal structure of the supercell and the corresponding orbital file. Make sure that both the structure file and the orbital file exist.

ecut: Determine the number of projections to the plane wave basis group, the energy unit is Ry.

band_range: Specify the energy band range of band unfolding.

m_matrix: Transformation matrix of supercell and primitive cell lattice vector

The k-point setting is determined according to the structure of the original cell, not the k-point in the supercell. For the k point setting of this function, please refer to the kpoint_mode module.

Once the task calculation is finished, three files will be generated in the Out/Bandunfolding folder. These files include kpt.dat and spectral_weight.dat, representing the k-point and spectral function of the original cell, respectively. Additionally, a drawing script called plot_unfold.py will also be included.

NOTE: When calculating a slab material, it is necessary to remove the thickness of the vacuum layer and make necessary modifications to the crystal vector and atomic positions specified in the LATTICE and structure files. Failing to make these modifications accurately can result in inaccurate results.

5.3 FERMI_ENERGY

5.3.1 introduction

This function is to calculate the Fermi energy of solid materials given temperature and electronic occupation number. If the system is an insulator, fermi energy is given by the valence band maximum (VBM).

For each \mathbf{k} point, the probability of finding an electron in any energy state should obey the Fermi-Dirac distribution. The integration of occupied electrons over the entire Brillouin zone should be the occupation number. Though which, the exact Fermi energy could be attained following Newton interpolation.

$$f(E, E_f, T) = \frac{1}{1 + e^{\left(\frac{E - E_f}{k_B T}\right)}}$$

$$N[E_f] = \int_{BZ} [d\mathbf{k}] \sum_n f(E_n, E_f, T)$$

5.3.2 example

You can find an example of how to calculate the Fermi energy of Copper in the examples/Cu folder.

The Input file is:

```
INPUT_PARAMETERS
   nspin
                        ABACUS
   package
   fermi_energy
                        Aut.o
   fermi_energy_unit
                        eV
   HR_route
                        data-HR-sparse_SPIN0.csr
                        data-SR-sparse_SPIN0.csr
   SR_route
   HR_unit
                        Bohr
   rR_unit
   max_kpoint_num
                         8000
}
LATTICE
   lattice_constant 6.91640
   lattice_constant_unit Bohr
   lattice_vector
   0.50 0.50
                        0.00
   0.50
             0.00
                        0.50
          0.50
   0.00
                        0.50
FERMI_ENERGY
   temperature
   electron_num
                        11
   grid
                        50 50 50
   epsilon
                        1e-4
```

electron_num: The number of the electrons in the system. The number of valence electrons, which is the sum of all atomic valence electrons of the system, can be obtained from the self-consistent output of the first-principles software.

epsilon: The max tolerable error of Newton interpolation. If two steps of Newton interpolation differs less than this epsilon, the calculation would stop and output the result.

temperature: The temperature of the system, unit in K.

After the task calculation is completed, there will be one file in the Out/Fermi_Energy folder, namely fermi_energy.dat, showing the calculated Fermi energy.

5.4 FERMI_SURFACE

5.4.1 introduction

The Fermi surface is a function that calculates the iso-energy surface of a given energy level. If the given energy level is the Fermi energy, then it will plot the Fermi surface.

5.4.2 example

You can find an example of how to calculate the Fermi surface of Copper in the examples/Cu folder.

The Input file is:

```
INPUT_PARAMETERS
   nspin
                         1
  package
                        ABACUS
   fermi_energy
                       Auto
   fermi_energy_unit
                       eV
   HR_route
                       data-HR-sparse_SPIN0.csr
   SR_route
                       data-SR-sparse_SPIN0.csr
   HR_unit
                        Bohr
   rR_unit
                        8000
   max_kpoint_num
}
LATTICE
   lattice_constant 6.91640
   lattice_constant_unit Bohr
   lattice_vector
   0.50 0.50
0.50 0.00
                        0.00
                        0.50
   0.00
             0.50
                        0.50
}
FERMI_ENERGY
                         0
   temperature
   electron_num
                        11
   grid
                        50 50 50
   epsilon
                        1e-4
FERMI_SURFACE
   bar
                         1e-5
   kpoint_mode
                        mp
   k_start
                         0 0 0
                         1 0 0
   k_vect1
                        0 1 0
   k_vect2
                         0 0 1
   k_vect3
                         50 50 50
   mp_grid
```

bar: The max tolerable error bar for the Fermi surface.

nbands: If you know the energy band range where the Fermi energy is located, you can set this parameter to speed up the calculation. There are two numbers in total, indicating the range of the energy band. The default value is 0 0, that is, all energy bands are considered.

For the k point setting of this function, please refer to the kpoint_mode module.

After the task calculation is completed, there will be two files in the Out/Fermi_Surface folder, namely fermi_surface_kpt.dat and plot_fermi_surface.py, corresponding to the k-point found on the Fermi surface and a plotting script.

NOTE: To visualize iso-energy surface at specific energy levels, the fermi_energy parameter in IN-PUT_PARAMETERS can be customized.

5.5 FIND_NODES

5.5.1 introduction

Find nodes is a method of finding k points with degenerate energy bands in a given energy range. This function can be used to find the Weyl/Dirac points in Weyl/Dirac semimetals.

5.5.2 example

An example of how to locate the Weyl point of $MnSb_2Te_4$ is provided in the examples/MnSb2Te4 folder.

The Input file is:

```
INPUT PARAMETERS
{
   nspin
                                4
                                ABACUS
   package
   fermi_energy
                                9.9700823666762375
   fermi_energy_unit
   HR_route
                                data-HR-sparse_SPIN0.csr
                                data-SR-sparse_SPIN0.csr
   SR_route
                                data-rR-sparse.csr
   rR_route
   HR_unit
                                Ry
   rR_unit
                                Bohr
}
LATTICE
                                1.8897162
   lattice_constant
   lattice_constant_unit
                                Bohr
   lattice_vector
    4.2885731815169859 \\ -0.0001203117360283 \\ -0.0000592245216563
    2.1441823977387200 3.7140734770500492 -0.0000592245216895
    }
FIND_NODES
   energy_range
                                9.870 10.070
   k_start
                                0.0 \ 0.0 \ -0.2
                                0.0 0.0 0.0
   k_vect1
                                0.0 0.0 0.0
   k_vect2
```

(continues on next page)

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energy range: The energy range in which the program searches for degenerate points, the energy unit is eV.

Set the search space of k points: Selecting a parallel hexahedron in k-space requires an origin (k_start) and three vectors ($k_vect1, k_vect2, k_vect3$) that are not parallel to each other in general. In this example, k_vect2 and k_vect3 are zero vectors, so the chosen search space is k-line from (0.0, 0.0, -0.2) to (0.0, 0.0, 0.4).

After the task calculation is completed, there will be three files in the Out/Find_Nodes folder, namely nodes.dat, plot_nodes.py and nodes.pdf, corresponding to the degenerate k-point(s) in direct coordinate and a plotting script with its plot.

5.6 PDOS

5.6.1 introduction

The distribution of electronic states at various energies is characterized by the density of states (DOS), while the partial density of states (PDOS) is a useful tool for analyzing the contribution of individual atomic orbitals to the DOS.

The implementation of PDOS is given by

$$g_{\mu}(E) = \frac{1}{N_{\mathbf{k}}} \sum_{\mathbf{k}} \sum_{n} \sum_{\nu} C_{n\nu}^{*}(\mathbf{k}) S_{\nu\mu}(\mathbf{k}) C_{n\mu}(\mathbf{k}) \delta(E - E_{n\mathbf{k}}),$$

where $C_{n\mu}(\mathbf{k})$ is the coefficient of the NAO.

5.6.2 example

An example (refer to folder examples/Si2) of calculating the PDOS of the diamond Si is given here.

The Input file is:

```
INPUT_PARAMETERS
   nspin
   package
                      ABACUS
   fermi_energy
                      6.389728305291531
   fermi_energy_unit eV
   HR_route
                      data-HR-sparse_SPIN0.csr
   SR_route
                      data-SR-sparse_SPIN0.csr
   rR_route
                      data-rR-sparse.csr
   HR_unit
                      Ry
   rR_unit
                      Bohr
}
LATTICE
                          1.8897162
   lattice_constant
```

(continues on next page)

5.6. PDOS 17

```
lattice_constant_unit
                          Bohr
   lattice vector
   0.00000000000 2.71500000000 2.71500000000
   2.71500000000 0.00000000000 2.715000000000
   2.71500000000 2.71500000000 0.00000000000
}
PDOS
{
   stru_file STRU
e_range -5.0 17.0
   de
               0.01
   sigma
            0.07
   kpoint_mode mp
   mp_grid
               12 12 12
}
```

stru_file: The structure file name of the supercell. This file indicates the crystal structure of the supercell and the corresponding orbital file. Make sure that both the structure file and the orbital file exist.

e_range: Specify the energy range of dos, the unit is eV.

de: specifies the energy interval.

sigma: Specify the parameters of Gaussian smearing.

For the k point setting of this function, please refer to the kpoint_mode module.

After the task calculation is completed, there will be three files in the Out/PDOS folder, namely TDOS.dat and PDOS.xml, $plot_dos.py$. Specify the projected atomic orbital index in the plot script, and then draw the PDOS plot.

5.7 FAT_BAND

5.7.1 introduction

A fat band can provide information about the contributions of specific atomic orbitals or groups of orbitals to the electronic bands of a material at given k points.

5.7.2 example

In the examples/Si2 folder, there is an example of how to calculate the fat band of the diamond Si.

The Input file is:

(continues on next page)

5.7. FAT_BAND 18

```
rR_unit
                       Bohr
LATTICE
    lattice_constant
                           1.8897162
    lattice_constant_unit
    lattice_vector
   0.00000000000 2.71500000000 2.71500000000
    2.71500000000 0.00000000000 2.715000000000
    2.71500000000 2.71500000000 0.00000000000
}
FAT_BAND
   band_range
                                  1 8
   stru_file
                                  STRU
   kpoint_mode
                                  line
   kpoint_num
   high_symmetry_kpoint
    0.50000 0.50000 0.5000 100
    0.00000 0.00000 0.0000 100
    0.50000 0.00000 0.5000 100
    0.37500 -0.37500 0.0000 100
    0.00000 0.00000 0.0000 1
}
```

band_range: There are two numbers (separated by spaces) to indicate which bands are selected for projection, counting from 1.

stru_file: The structure file name. This file indicates the crystal structure and the corresponding orbital file. Make sure that both the structure file and the orbital file exist.

For the k point setting of this function, please refer to the kpoint_mode module.

Once the task calculation is finished, you will find four files in the Out/Fat_Band folder. These files include band. dat and pband.dat, fatband.xml, and plot_fatband.py. They contain valuable information about the original bands, the coefficients of the bands projected onto each atomic orbital (the number of atomic orbitals is equal to the number of basis sets), an XML formatted file of the projected bands, and a script to visualize the fat band.

5.8 SPIN_TEXTURE

5.8.1 introduction

Spin texture refers to the spatial distribution of electron spins in momentum space. In PYATB, the spin texture is calculated as follows,

$$\langle \Psi_{n\mathbf{k}}|\hat{\sigma}_i|\Psi_{n\mathbf{k}}\rangle = \sum_{\mu,\nu,s,s'} C^*_{n,\mu s}(\mathbf{k}) S_{\mu\nu,ss'}(\mathbf{k}) \hat{\sigma}_{i,ss'} C_{n,\nu s'}(\mathbf{k}),$$

where $\hat{\sigma}_i$ are the Pauli matrices, with i=x, y, z, and $s=\uparrow, \downarrow$ is the spin index.

5.8.2 example

Here, we provide an example of calculating the spin texture of Bi₂Se₃ (refer to folder examples/Bi₂Se₃).

The Input file is:

```
INPUT_PARAMETERS
{
   nspin
   package
                                   ABACUS
   fermi_energy
                                   9.557219691497478
   fermi_energy_unit
                                   eV
   HR_route
                                   data-HR-sparse_SPIN0.csr
   SR_route
                                   data-SR-sparse_SPIN0.csr
   rR_route
                                   data-rR-sparse.csr
   HR_unit
   rR_unit
                                   Bohr
                                   8000
   max_kpoint_num
}
LATTICE
{
                                   1.8897162
   lattice_constant
   lattice_constant_unit
                                   Bohr
   lattice_vector
   -2.069 -3.583614 0.000000
    2.069 -3.583614 0.000000
    0.000 2.389075 9.546667
}
SPIN_TEXTURE
    nband
                       78
   kpoint_mode
                       direct
   kpoint_num
                       20
   kpoint_direct_coor
   0.010000 0.000000 0.000000
   0.009511 0.003090 0.000000
   0.008090 0.005878 0.000000
   0.005878 0.008090 0.000000
   0.003090 0.009511 0.000000
   0.000000 0.010000 0.000000
  -0.003090 0.009511 0.000000
  -0.005878 0.008090 0.000000
  -0.008090 0.005878 0.000000
  -0.009511 0.003090 0.000000
  -0.010000 0.000000 0.000000
  -0.009511 -0.003090 0.000000
  -0.008090 -0.005878 0.000000
  -0.005878 -0.008090 0.000000
  -0.003090 -0.009511 0.000000
  -0.000000 -0.010000 0.000000
   0.003090 -0.009511 0.000000
   0.005878 -0.008090 0.000000
   0.008090 -0.005878 0.000000
    0.009511 -0.003090 0.000000
}
```

nband: Denote the band number of which spin texture is calculated.

For the k point setting of this function, please refer to the kpoint_mode module.

After the task calculation is completed, there will be three files in the Out/Spin_Texture folder, namely kpt. dat and spin_texture.dat,plot_spin_texture.py, corresponding to the k-point and the spin texture, the drawing script.

5.9 WILSON LOOP

5.9.1 introduction

We can determine the Z2 topology number of the topological insulator by Wilson loop. Computing the six time reversal invariant planes, we can obtain the Z2 topology metrics (ν_0 , ν_1 ν_2 ν_3). These six planes are k1=0.0, k1=0.5, k2=0.0, k2=0.5, k3=0.0, k3=0.5, respectively. When selecting a plane, only half of the plane needs to be selected.

$$\nu_0 = Z2(ki = 0) + Z2(ki = 0.5) \quad mod \quad 2$$
 $\nu_i = Z2(ki = 0.5)$

where i = 1, 2, 3 refers to the x, y and z directions.

The Wilson loop is implemented as follows: $\$W_n(\mathbf{k_2}) = \frac{i}{2\pi} \int_0^{2\pi} d\mathbf{k_1} \langle u_{n,\mathbf{k_1},\mathbf{k_2}} | \partial_{\mathbf{k_1}} | u_{n,\mathbf{k_1},\mathbf{k_2}} \rangle. \$$

5.9.2 example

Here is an example of how to calculate the Wilson loop of the topological insulator Bi_2Se_3 located in the examples/Bi2Se3 folder.

The Input file is:

```
INPUT_PARAMETERS
   nspin
                                   4
                                   ABACUS
   package
   fermi_energy
                                   9.557219691497478
   fermi_energy_unit
                                   data-HR-sparse_SPIN0.csr
   HR_route
   SR_route
                                   data-SR-sparse_SPIN0.csr
   rR_route
                                   data-rR-sparse.csr
   HR_unit
                                   Bohr
   rR_unit
   max_kpoint_num
                                   8000
}
LATTICE
   lattice_constant
                                   1.8897162
   lattice_constant_unit
                                   Bohr
   lattice_vector
    -2.069 -3.583614 0.000000
    2.069 -3.583614 0.000000
    0.000 2.389075 9.546667
WILSON_LOOP
```

(continues on next page)

```
      occ_band
      78

      k_start
      0.0 0.0 0.5

      k_vect1
      1.0 0.0 0.0

      k_vect2
      0.0 0.5 0.0

      nk1
      101

      nk2
      101
```

occ_band: The number of occupied energy bands of an insulator.

k_start: The origin point coordinates used to describe a Brillouin zone plane.

k_vect1: The expansion vector is a vector used to define a Brillouin zone plane, and it is also the direction of integration for calculations.

k_vect2: The expansion vector is a vector used to define a Brillouin zone plane, and it is also the direction of Wilson loop evolution for calculations.

nk1: k_vect1 is divided into nk1 k-points.

nk2: k_vect2 is divided into nk2 k-points.

After the task calculation is completed, there will be two files in the Out/Wilson_Loop folder, namely wilson_loop.dat and plot_wl.py, corresponding to the Wilson loop of each k-point in the k_vect2 direction and drawing script.

5.10 POLARIZATION

5.10.1 introduction

We calculate the spontaneous polarization of periodic solids by so-called Modern Theory of Polarization, namely Berry phase theory. The electric polarization $\bf P$ is a modulo a quantum $e{\bf R}/V_c$ multi-valued function, corresponding to the following implementation equation:

$$\mathbf{P} = \frac{-e}{(2\pi)^3} \sum_{n}^{occ} \int_{BZ} \mathbf{A}_n(\mathbf{k}) d^3k,$$

where $A_n(\mathbf{k})$ is the Berry connection of a single band.

5.10.2 example

Here, we present an example (located in the examples/PbTiO3 folder) showcasing the calculation of the polarization of PbTiO3.

The Input file is:

(continues on next page)

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```
rR_route
                      data-rR-sparse.csr
   HR_unit
                      Ry
                      Bohr
   rR_unit
LATTICE
{
                          7.3699
   lattice_constant
   lattice_constant_unit Bohr
   lattice_vector
   1.0000000000
                      0.0000000000
                                           0.0000000000
   0.000000000
                      1.0000000000
                                           0.0000000000
   0.000000000
                      0.0000000000
                                           1.0000000000
}
POLARIZATION
                22
   occ_band
   nk1
                1.0
   nk2
                10
   nk3
                3
   atom_type
               STRU
   stru_file
   valence_e
               14 12 6
}
```

occ_band: The number of occupied energy bands of an insulator.

nk1: This refers to the number of samples taken in the x direction of the reciprocal lattice vector \mathbf{G} .

nk2: This refers to the number of samples taken in the y direction of the reciprocal lattice vector \mathbf{G} .

nk3: This refers to the number of samples taken in the z direction of the reciprocal lattice vector \mathbf{G} .

stru_file: Specify the strucutre file. NAOs files are not required.

atom_type: The number of element types in the system.

valence_e: The number of valence electrons per element.

The parameters nk1, nk2, and nk3 correspond to the number of k-points used for the integration in the three lattice directions, and increasing their values leads to a more accurate and convergent result.

After completing the task, polarization.dat appears in the Out/Polarization folder which contains the electric polarization of the three lattice directions.

5.11 BERRY_CURVATURE

5.11.1 introduction

Berry curvature is of fundamental importance for understanding some basic properties of solid materials and is essential for the description of the dynamics of Bloch electrons.

The Berry curvature of a single energy band is defined as follows:

$$\Omega_n(\mathbf{k}) = \nabla \times \mathbf{A}_n(\mathbf{k}),$$

where Berry phase $\mathbf{A}_n(\mathbf{k}) = i \langle u_{n\mathbf{k}} | \nabla_{\mathbf{k}} | u_{n\mathbf{k}} \rangle$, $|u_{n\mathbf{k}}\rangle$ is the periodic part of the Bloch wave function.

We calculated the Berry curvature:

$$\Omega_{\alpha\beta}(\mathbf{k}) = \sum_{n} f_n(\mathbf{k}) \Omega_{n,\alpha\beta}(\mathbf{k}),$$

where f_n is the Fermi occupation function.

Detailed descriptions can be found in Calculation of Berry curvature using non-orthogonal atomic orbitals.

5.11.2 example

An example (refer to folder example/Fe) of calculating the Berry curvature of the fcc-Fe is given here.

The Input file is:

```
INPUT_PARAMETERS
{
                      4
   nspin
                      ABACUS
   package
                     18.18839115931923
   fermi_energy
   fermi_energy_unit eV
                data-HR-sparse_SPIN0.csr
   HR_route
   SR_route
                    data-SR-sparse_SPIN0.csr
   rR_route
                    data-rR-sparse.csr
                    Ry
   HR_unit
                      Bohr
   rR_unit
}
LATTICE
   lattice_constant
                         5.4235
   lattice_constant_unit
                         Bohr
   lattice_vector
   0.5 0.5 0.5
   -0.5 0.5 0.5
   -0.5 -0.5 0.5
}
BERRY_CURVATURE
   method
   kpoint_mode
                          line
   kpoint_num
   high_symmetry_kpoint
   0.0 0.0 0.0 100 # G
   0.5 -0.5
             -0.5
                     100 # H
   0.75 0.25 -0.25 100 \# P
   0.5 0.0 -0.5
                     100 # N
   0.0 0.0
             0.0
                     100 # G
   0.5 0.5
               0.5
                     100 # H
   0.5 0.0
               0.0
                     100 # N
   0.0 0.0
               0.0
                     100 # G
   0.75 0.25 -0.25 100 # P
   0.5
         0.0
               0.0
                     1
                         # N
```

method: Method for calculating Berry curvature. 0 means direct calculation, 1 means calculation by Kubo formula.

occ_band: The number of occupied energy bands of an insulator. When this value is not set, it will be determined according to the Fermi energy.

For the k point setting of this function, please refer to the kpoint_mode module.

Upon completion of the task calculation, two files will be generated in the Out/Berry_Curvature directory: kpt. dat and berry_curvature.dat. These files respectively contain the k-point coordinates and the total Berry curvature for each k-point.

5.12 AHC

5.12.1 introduction

The dc anomalous Hall conductivity (AHC) is simply given as the Brillouin zone integral of the Berry curvature of occupying energy bands,

$$\sigma_{xy} = -\frac{e^2}{\hbar} \sum_{n}^{occ} \int_{\rm BZ} \frac{d\mathbf{k}}{(2\pi)^3} f_n(\mathbf{k}) \Omega_{n,z}(\mathbf{k}).$$

5.12.2 example

An example (refer to folder example/Fe) of calculating the AHC of the fcc-Fe is given here.

The Input file is:

```
INPUT_PARAMETERS
   nspin
                     ABACUS
   package
   fermi_energy 18.18839115931923
   fermi_energy_unit eV
                      data-HR-sparse_SPIN0.csr
   HR_route
   SR_route
                      data-SR-sparse_SPIN0.csr
   rR_route
                     data-rR-sparse.csr
   HR_unit
                      Ry
   rR_unit
                     Bohr
}
LATTICE
                         5.4235
   lattice_constant
   lattice_constant_unit Bohr
   lattice_vector
    0.5 0.5 0.5
   -0.5 0.5 0.5
   -0.5 -0.5 0.5
}
AHC
   integrate_mode
                          Grid
   integrate_grid
                          100 100 100
   adaptive_grid
                          20 20 20
   adaptive_grid_threshold 100
```

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integrate_mode: Specifies the mode of integration, which can be grid integration and adaptive integration.

integrate_grid: Specifies a uniform grid for grid integration.

adaptive_grid: Specifies the grid for adaptive densification.

adaptive_grid_threshold: Specifies the cut-off value of adaptive densification, the unit is ².

Once the calculation is complete, the resulting file ahc.dat will be generated in the Out/AHC directory.

5.13 CHERN_NUMBER

5.13.1 introduction

Chern number is a topological invariant used to explain the quantized Hall conductivity. The Chern invariant is the total Berry flux in the 2D Brillouin zone,

$$n = \frac{1}{2\pi} \oint_{\mathbf{S}} \mathbf{\Omega} \cdot d\mathbf{S}$$

where n is an integer, **S** is any closed 2D manifold, Ω is total Berry curvature (flux).

To calculate the Chern number, you must first select a closed 2D surface in the Brillouin.

5.13.2 example

An example (refer to folder examples/MnBi2Te4-wey1) of calculating Chern number of the Weyl semimetal $MnBi_2Te_4$ is given here.

The Input file is:

```
INPUT_PARAMETERS
   nspin
                                ABACUS
   package
                                9.2309138700265265
   fermi_energy
   fermi_energy_unit
   HR_route
                                data-HR-sparse_SPIN0.csr
   SR_route
                                data-SR-sparse_SPIN0.csr
   rR_route
                                data-rR-sparse.csr
   HR_unit
                                Ry
   rR_unit
                                Bohr
}
LATTICE
{
                                1.8897162
   lattice_constant
   lattice_constant_unit
                                Bohr
   lattice_vector
   2.188670000000001 3.7908876409999999 0.0000000000000000
   2.188670000000001 1.263629209999999 13.7730333333000008
CHERN_NUMBER
```

(continues on next page)

```
0
method
occ band
                                 109
                                 Grid
integrate_mode
                                100 100 1
integrate_grid
                                 20 20
adaptive_grid
adaptive_grid_threshold
                                100
                                 0 0 0
k_start
k_vect1
                                 1 0 0
                                 0 1 0
k_vect2
```

method: Method for calculating berry curvature. 0 means direct calculation, 1 means calculation by Kubo formula.

occ_band: The number of occupied energy bands of an insulator. When this value is not set, it will be determined according to the Fermi energy.

To determine a plane of k-space requires an origin (k_start) and two vectors that are not parallel to each other (k_vect1, k_vect2) .

k_start: The origin point coordinates used to describe a Brillouin zone plane.

k_vect1: The expansion vector used to describe a Brillouin zone plane.

k_vect2: The expansion vector used to describe a Brillouin zone plane.

For k-point integration, please refer to the [Setting of integration] section.

After the task calculation is completed, the <code>chern_number.dat</code> file will appear in the <code>Out/Chern_Num</code> folder which contains the Chern number specific results.

5.14 CHIRALITY

5.14.1 introduction

Examines the chirality of Weyl points by calculating the Berry curvature on a sphere around the k point.

5.14.2 example

An example is provided in the examples/MnSb2Te4 directory to investigate the chirality of the Weyl point in $MnSb_2Te_4$.

The Input file is:

```
INPUT_PARAMETERS
{
                                     4
   nspin
                                     ABACUS
   package
                                     9.9700823666762375
   fermi_energy
   fermi_energy_unit
   HR_route
                                     data-HR-sparse_SPIN0.csr
   SR_route
                                     data-SR-sparse_SPIN0.csr
   rR_route
                                     data-rR-sparse.csr
   HR_unit
                                     Ry
   rR_unit
                                     Bohr
```

(continues on next page)

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```
LATTICE
                                    1.8897162
    lattice_constant
    lattice_constant_unit
                                    Bohr
    lattice_vector
    4.2885731815169859 -0.0001203117360283
                                               -0.0000592245216563
    2.1441823977387200 3.7140734770500492
                                               -0.0000592245216895
    2.1439936829776851
                          1.2378353300160352
                                               13.4147375436875418
}
CHIRALITY
   k_vect
                                    0.0000 0.0000 -0.0538
   radius
   point_num
                                    100
```

k_vect: The k-point direct coordinates need to be calculated.

radius: The radius of the integrating sphere. The unit is $^{-1}$.

point_num: The number of k-points that are uniformly sampled on a spherical surface.

The chirality.dat file that record the results are in the Out/Chirality folder.

5.15 JDOS

5.15.1 introduction

Joint density of states (JDOS) is used to describe the density of states of electrons excited from the valence band to the conduction band, which relates to the absorption spectrum and the dielectric function of system.

The implementation of JDOS per crystal cell is given by

$$D_{joint}(\omega) = \frac{V_c}{\hbar} \int \frac{d^3k}{(2\pi)^3} \sum_{n,m} f_{nm} \delta(\omega_{mn} - \omega).$$

where V_c is the cell volume, $f_{nm} = f_n - f_m$ and $\hbar \omega_{mn} = E_m - E_n$ are differences between occupation factors and band energies, respectively.

Currently JDOS is only used to calculate insulators and semiconductors.

5.15.2 example

An example of calculating the JDOS of a perovskite $CsPbI_3$ is presented here (refer to folder examples/CsPbI3).

The Input file is:

(continues on next page)

5.15. JDOS 28

```
fermi_energy_unit
   HR_route
                   data-HR-sparse_SPIN0.csr
   SR_route
                  data-SR-sparse_SPIN0.csr
                   data-rR-sparse.csr
   rR_route
   HR_unit
   rR_unit
                   Bohr
}
LATTICE
   lattice_constant 1.8897261258369282
   lattice_constant_unit Bohr
   lattice_vector
      }
JDOS
   occ_band
              0.5 10
   omega
             0.01
   domega
              0.2
   eta
   grid
             30 30 30
```

occ_band: Specifies the occupied energy band of the system. Currently, only insulator or semiconductor materials can be calculated.

omega: Specifies the photon energy, the unit is eV.

domega: Specifies the energy interval of the omega.

eta: Specify the parameters of Gaussian smearing.

grid: Specifies the uniform k-point grid used to calculate the JDOS.

Once the task calculation is finished, two files will be generated in the Out/JDOS folder: JDOS. dat and $plot_jdos$. py. The former contains the JDOS data, while the latter is a script used for plotting the JDOS.

5.16 OPTICAL_CONDUCTIVITY

5.16.1 introduction

The frequency-dependent optical conductivity expressed by the Kubo-Greenwood formula can be formulated as

$$\sigma_{\alpha\beta}(\hbar\omega) = -\frac{ie^2\hbar}{NV_{\rm cell}} \sum_{\mathbf{k}} \sum_{n,m} \left(\frac{f_{n\mathbf{k}} - f_{m\mathbf{k}}}{E_{n\mathbf{k}} - E_{m\mathbf{k}}} \right) \frac{\langle \psi_{n\mathbf{k}} | v_\alpha | \psi_{m\mathbf{k}} \rangle \langle \psi_{m\mathbf{k}} | v_\beta | \psi_{n\mathbf{k}} \rangle}{\hbar\omega + E_{n\mathbf{k}} - E_{m\mathbf{k}} + i\eta} \,.$$

The imaginary part of the dielectric function is

$$\epsilon_i^{\alpha\beta}(\omega) = -\frac{e^2\pi}{\epsilon_0\hbar} \int \frac{d\mathbf{k}}{\left(2\pi\right)^3} \sum_{nm} f_{nm} r_{nm}^\alpha r_{mn}^\beta \delta\left(\omega_{mn} - \omega\right) \,,$$

The real part of the dielectric function is obtained by the Kramer-Kronig transformation,

$$\epsilon_r^{\alpha\beta}(\omega) = \delta_{\alpha\beta} + \frac{2}{\pi} \mathbf{P} \int_0^\infty d\omega' \frac{\omega' \epsilon_i^{\alpha\beta} (\omega')}{\omega'^2 - \omega^2} .$$

The linear optical spectrum can be calculated through the dielectric function, such as refractive index $n(\omega)$, extinction coefficient $\kappa(\omega)$, absorption coefficient $\kappa(\omega)$, energy-loss function $L(\omega)$, reflectivity $R(\omega)$:

$$\begin{split} n(\omega) &= \left[\frac{\sqrt{\varepsilon_1^2 + \varepsilon_2^2} + \varepsilon_1}{2}\right]^{\frac{1}{2}} \\ \kappa(\omega) &= \left[\frac{\sqrt{\varepsilon_1^2 + \varepsilon_2^2} - \varepsilon_1}{2}\right]^{\frac{1}{2}} \\ \alpha(\omega) &= \frac{\sqrt{2}\omega}{c} \left[\sqrt{\varepsilon_1^2 + \varepsilon_2^2} - \varepsilon_1\right]^{\frac{1}{2}} \\ L(\omega) &= \operatorname{Im}\left(\frac{-1}{\varepsilon(\omega)}\right) = \frac{\varepsilon_2}{\varepsilon_1^2 + \varepsilon_2^2} \\ R(\omega) &= \frac{(n-1)^2 + k^2}{(n+1)^2 + k^2} \end{split}$$

5.16.2 example

Here, we provide an example (located in the examples/Si folder) demonstrating the calculation of the optical conductivity and dielectric function for diamond Si.

The Input file is:

```
INPUT_PARAMETERS
   nspin
   package ABACUS fermi_energy 6.389728305291531
    fermi_energy_unit eV
   HR_route data-HR-sparse_SPINO.csr
SR_route data-SR-sparse_SPINO.csr
    rR_route
                        data-rR-sparse.csr
    HR_unit
    rR_unit
                        Bohr
}
LATTICE
    lattice_constant 1.8897162
    lattice_constant_unit Bohr
   lattice_vector
   0.00000000000 2.71500000000 2.71500000000
    2.71500000000 0.00000000000 2.71500000000
    2.71500000000 2.71500000000 0.00000000000
OPTICAL_CONDUCTIVITY
                  4
    occ_band
                  0
                      10
    omega
```

(continues on next page)

```
domega 0.01
eta 0.1
grid 50 50 50
}
```

occ_band: Used to specify the occupied energy band of an insulator or semiconductor. Currently this function can only calculate insulators or semiconductors.

omega: Specifies the photon energy, the unit is eV.

domega: Specifies the energy interval of the omega.

eta: Specify the parameters of Gaussian smearing.

grid: Specifies the uniform k-point grid used to calculate the optical conductivity.

After completing the task, five main files are generated in the Out/Optical_Conductivity folder, namely optical_conductivity_real_part.dat, optical_conductivity_imag_part.dat, dielectric_function_real_part.dat, dielectric_function_imag_part.dat and plot_optical.py.

5.17 SHIFT CURRENT

5.17.1 introduction

The shift current is an intrinsic contribution to the bulk photovoltaic effect (BPVE). It describes the photocurrent generated by light illumination on homogeneous non-centrosymmetric crystals. The shift current is a second-order optical response. It can be expressed as a DC current, generated by a monochromatic photoelectric field $\mathbf{E}(t) = \mathbf{E}(\omega) \mathrm{e}^{i\omega t} + \mathbf{E}(-\omega) \mathrm{e}^{-i\omega t}$, where

$$J^a = 2\sigma^{abc}(0; \omega, -\omega)E_b(\omega)E_c(-\omega).$$

Here, a, b, c = x, y, z, and $\sigma^{abc}(0; \omega, -\omega)$ is the shift current tensor,

$$\sigma^{abc}(0;\omega,-\omega) = \frac{\pi e^3}{\hbar^2} \int \frac{d\mathbf{k}}{8\pi^3} \sum_{n,m} f_{nm} \text{Im} \left[I_{mn}^{abc} + I_{mn}^{acb} \right] \delta \left(\omega_{mn} - \omega \right)$$

where $I_{mn}^{abc} = r_{mn}^b r_{nm;a}^c$, r_{nm}^a is the inter-band dipole matrix, and $r_{nm;a}^b$ is the generalized derivative of the dipole matrix.

Detailed descriptions can be found in Second-order optical response in semiconductors.

5.17.2 example

Here, we provide an example of calculating the shift current conductivity of the monolayer WS_2 (refer to folder examples/WS2).

The Input file is:

(continues on next page)

```
0.3484302262859574
    fermi_energy
    fermi_energy_unit
   HR_route
                            data-HR-sparse_SPIN0.csr
                            data-SR-sparse_SPIN0.csr
   SR_route
    rR_route
                            data-rR-sparse.csr
    HR_unit
                            Bohr
    rR_unit
LATTICE
                           1.8897162
   lattice_constant
   lattice_constant_unit Bohr
   lattice_vector
   3.183820900165 0.0
                                      0.0
   -1.591910450082 2.757269780643 0.0
                                      20.086904001384
    0.0
                    0.0
}
SHIFT_CURRENT
   occ_band
                            13
                            0
   omega
                            0.01
   domega
   smearing_method
                           1
                            0.1
    grid
                            1000 1000 1
```

occ_band: Used to specify the occupied energy band of an insulator or semiconductor. Currently this function can only calculate insulators or semiconductors.

omega: Specifies the photon energy, the unit is eV.

domega: Specifies the energy interval of the omega.

eta: Specify the parameters of Gaussian smearing.

grid: Specifies the uniform k-point grid used to calculate the shift current.

Upon completion of the task, two essential files are generated in the Out/Shift_Current folder. These files include shift_current.dat and plot_shift_current.py.

5.18 BERRY_CURVATURE_DIPOLE

5.18.1 introduction

In a system with time-reversal symmetry, the Berry curvature is an odd function of \mathbf{k} , i.e., $\Omega_a(\mathbf{k}) = -\Omega_a(-\mathbf{k})$. As a result, the integration of the Berry curvature over the BZ is zero. However, if the system lacks a inversion symmetry, a higher-order nonlinear AHC can arise. More specifically, $j_a^0 = \chi_{abc} E_b(\omega) E_c(-\omega)$ and $j_a^{2\omega} = \chi_{abc} E_b(\omega) E_c(\omega)$, describe a rectified current and the second harmonic, respectively, whereas ω is the driving frequency. The coefficient χ_{abc} is given by

$$\chi_{abc} = -\varepsilon_{adc} \frac{e^3 \tau}{2(1 + i\omega \tau)} D_{bd}.$$

where

$$D_{ab} = \int_{k} f_0 \left(\frac{\partial \Omega_b}{\partial k_a} \right)$$

is called the Berry curvature dipole.

In PYATB, the Berry curvature dipole at a given temperature T and chemical potential μ is calculated using the following formula:

$$D_{ab}(\mu,T) = -\int \frac{\partial f_0(E,\mu,T)}{\partial E} D_{ab}(E) dE.$$

5.18.2 example

Here, we provide an example of calculating the Berry curvature dipole of the trigonal Te (refer to folder examples/Te).

The Input file is:

```
INPUT_PARAMETERS
                           4
   nspin
   package
                           ABACUS
   fermi_energy
                           9.574476774876963
   fermi_energy_unit
                          eV
   HR_route
                           data-HR-sparse_SPIN0.csr
                          data-SR-sparse_SPIN0.csr
   SR_route
                          data-rR-sparse.csr
   rR_route
   HR_unit
                          Ry
   rR_unit
                          Bohr
   max_kpoint_num
                          28000
}
LATTICE
                   1.8897162
   lattice_constant
   lattice_constant_unit
                           Bohr
   lattice_vector
   2.22 -3.84515 0.000
   2.22
          3.84515 0.000
   0.00
          0.00000 5.910
}
BERRY_CURVATURE_DIPOLE
                          9.474 10.074
   omega
                          0.001
   domega
   integrate_mode
                          Grid
   integrate_grid
                           400 400 400
   adaptive_grid
                           20 20 20
   adaptive_grid_threshold 20000
```

omega: To set the energy range for the Berry curvature dipole, you can adjust it based on the Fermi energy level. the unit is eV.

domega: Specifies the energy interval of the omega.

For k-point integration, please refer to the [Setting of integration] section.

Once the task has been finished, three crucial files are produced in the Out/Berry_Curvature_Dipole directory. These files consist of bcd_step2.dat, kpoint_list, and plot_bcd.py. The first file stores the Berry curvature dipole's magnitude, while the second file records the refined k-point coordinates. The third file contains the script used for generating the visualization of the dipole.

DETAILED INTRODUCTION OF THE INPUT FILES

6.1 Full List of INPUT Keywords

• INPUT_PARAMETERS

nspin | package | fermi_energy | fermi_energy_unit | HR_route | SR_route | rR_route | binary | HR_unit | rR_unit | max_kpoint_num | sparse_format

• LATTICE

lattice_constant | lattice_constant_unit | lattice_vector

• BAND_STRUCTURE

wf_collect | kpoint_mode

• BANDUNFOLDING

stru_file | ecut | band_range | m_matrix | kpoint_mode

• FERMI_ENERGY

temperature | electron_num | grid | epsilon |

• FERMI_SURFACE

bar | nbands | kpoint_mode

• FIND_NODES

energy_range | k_start | k_vect1 | k_vect2 | k_vect3 | initial_grid | initial_threshold | adaptive_gridadaptive_threshold | kpoint_mode

• PDOS

stru_file | e_range | de | sigma | kpoint_mode

• FAT_BAND

band_range | stru_file | kpoint_mode

• SPIN_TEXTURE

nband | kpoint_mode

• WILSON_LOOP

occ_band | k_start | k_vect1 | k_vect2 | nk1 | nk2

• POLARIZATION

occ_band | nk1 | nk2 | nk3 | atom_type | stru_file | valence_e

• BERRY_CURVATURE

method | occ_band | kpoint_mode

AHC

method | integrate_mode

• CHERN NUMBER

method | occ_band | k_start | k_vect1 | k_vect2 | integrate_mode

• CHIRALITY

method | k_vect | radius | point_num

JDOS

occ_band | omega | domega | eta | grid

• OPTICAL_CONDUCTIVITY

occ_band | omega | domega | eta | grid

• SHIFT_CURRENT

occ_band | omega | domega | smearing_method | eta | grid | method

• BERRY_CURVATURE_DIPOLE

omega | domega | integrate_mode

Setting of k points

• When kpoint_mode is 'mp'

mp_grid | k_start | k_vect1 | k_vect2 | k_vect3

• When kpoint_mode is 'line'

kpoint_num | high_symmetry_kpoint

• When kpoint_mode is 'direct'

kpoint_num | kpoint_direct_coor

Setting of integration

• When integrate mode is 'Grid'

integrate_grid | adaptive_grid | adaptive_grid_threshold

• When integrate_mode is 'Adaptive'

relative_error | absolute_error | initial_grid

6.1.1 INPUT_PARAMETERS

nspin {#input_nspin}

- Type: Integer
- **Description**: Indicates the spin component of the wave function, related to the structure of the HR file.
 - 1: regardless of spin.
 - 2: the wave function is divided into two groups, one group is all up and one group is all down.

- 4: the wave function has both up and down components.
- Default: No default value

package {#input_package}

- Type: String
- Description: Indicates data sources for HR, SR, rR.
- Default: ABACUS

fermi_energy {#input_fermi_energy}

- Type: Real
- **Description**: Indicates the Fermi energy of the system. When set to Auto, the FERMI_ENERGY function needs to be added.
- Default: Auto

fermi_energy_unit {#input_fermi_energy_unit}

- Type: String
- Description: The unit of Fermi. Can be set to Ry, eV.
- Default: eV

HR_route {#input_HR_route}

- Type: String
- **Description**: Path to HR matrix file. When nspin=2, two sets of paths need to be provided.
- Default: No default value

SR_route {#input_SR_route}

- Type: String
- **Description**: Path to the SR matrix file.
- Default: No default value

rR_route {#input_rR_route}

- Type: String
- **Description**: Path to the rR matrix file.
- Default: No default value

binary {#input_binary}

• Type: Boolean

• **Description**: Whether HR, SR, and rR files are binary files.

• Default: 0

HR_unit {#input_HR_unit}

• Type: String

• Description: The unit of HR. Can be set to Ry, eV.

• **Default**: Ry

rR_unit {#input_rR_unit}

• Type: String

• **Description**: The unit of rR. Can be set to Bohr, Angstrom.

• Default: Bohr

max_kpoint_num {#input_max_kpoint_num}

• Type: Integer

• **Description**: The upper limit of the number of k points stored in the memory during program calculation, which is used to control the memory consumption during calculation.

• **Default**: 8000

sparse_format {#input_sparse_format}

• **Type**: Boolean

• Description: Whether HR, SR, rR matrices are stored in memory is sparse storage.

• Default: 0

6.1.2 LATTICE

lattice_constant {#lattice_lattice_constant}

• Type: Real

• **Description**: The lattice constant of the system.

lattice_constant_unit {#lattice_lattice_constant_unit}

• Type: String

• **Description**: The unit of the lattice constant. Can be set to Bohr, Angstrom.

• Default: Bohr

lattice vector {#lattice lattice vector}

• Type: Real

• **Description**: The 3 lattice vectors of the system. Each lattice vector is a row, with a total of 3 rows and 9 parameters.

• Default: No default value

6.1.3 BAND STRUCTURE

wf_collect {#bandstructure_wf_collect}

• Type: Boolean

• **Description**: Whether to output wave function matrix information.

• Default: No default value

kpoint_mode {#bandstructure_kpoint_mode}

• Type: String

• **Description**: Used to set the k point. See *Setting of k points*

• Default: No default value

6.1.4 BANDUNFOLDING

stru_file {#bandunfolding_stru_file}

• Type: String

• Description: Specify the strucutre file path.

• Default: No default value

ecut {#bandunfolding_ecut}

• Type: Real

• Description: Used to determine the number of plane wave basis sets. Unit is Ry.

• **Default**: 10

band_range {#bandunfolding_band_range}

- Type: Integer
- **Description**: Specifies the range of supercell energy band index within which the energy bands will be calculated. There are two parameters, representing the starting band index and the end band index, the index counts from 1.
- Default: No default value

m_matrix {#bandunfolding_m_matrix}

- Type: Real
- **Description**: The lattice vector transformation matrix between the supercell and the primitive cell, with 9 parameters, is written on the same line.
- Default: No default value

kpoint_mode {#bandunfolding_kpoint_mode}

- Type: String
- **Description**: Used to set the k point of unitcell. See *Setting of k points*
- Default: No default value

6.1.5 FERMI_ENERGY

temperature {#fermienergy_temperature}

- Type: Real
- **Description**: temperature. The unit is K
- Default: 0

electron_num {#fermienergy_electron_num}

- Type: Integer
- **Description**: The total number of electrons in the system.
- Default: No default value

grid {#fermienergy_grid}

- Type: Integer
- **Description**: The grid to use for Newton interpolation. There are three parameters.
- **Default**: 10 10 10

epsilon {#fermienergy_epsilon}

• **Type**: Real

• **Description**: Newton interpolation parameters, absolute accuracy.

• **Default**: 0.001

6.1.6 FERMI_SURFACE

bar {#fermisurface_bar}

• Type: Real

• **Description**: The max tolerable error bar for the Fermi surface

• Default: No default value

nbands {#fermisurface_nbands}

• Type: Integer

• **Description**: If you know the energy band range where the Fermi energy is located, you can set this parameter to speed up the calculation. There are two numbers in total, indicating the range of the energy band. The default value is 0 0, that is, all energy bands are considered.

• **Default**: 0 0

kpoint_mode {#fermisurface_kpoint_mode}

• Type: String

• **Description**: Used to set the k point. See *Setting of k points*

• Default: No default value

6.1.7 FIND_NODES

energy_range {#findnodes_energy_range}

• Type: Integer

• Description: The energy range in which the program searches for degenerate points, the energy unit is eV.

k_start {#findnodes_k_start}

• **Type**: Real

• **Description**: The origin point coordinates used to describe a Brillouin zone plane.

• **Default**: 0.0 0.0 0.0

k vect1 {#findnodes k vect1}

• Type: Real

• **Description**: The expansion vector used to describe a Brillouin zone plane.

• **Default**: 1.0 0.0 0.0

k vect2 {#findnodes k vect2}

• Type: Real

• **Description**: The expansion vector used to describe a Brillouin zone plane.

• **Default**: 0.0 1.0 0.0

k vect3 {#findnodes k vect3}

• Type: Real

• **Description**: The expansion vector used to describe a Brillouin zone plane.

• **Default**: 0.0 0.0 1.0

initial_grid {#findnodes_initial_grid}

• Type: Integer

• Description: Set the initial grid for searching degenerate k points. There are three parameters.

• **Default**: 10 10 10

initial_threshold {#findnodes_initial_threshold}

• Type: Real

• **Description**: The energy unit is eV. In the initial grid, only the k-points whose band differences are less than the threshold can enter the search for the next round of degenerate points.

• **Default**: 0.1

adaptive_grid {#findnodes_adaptive_grid}

• Type: Integer

• **Description**: The refined grid will refine the k-points that reach the initial_threshold in the initial grid. There are three parameters.

• Default: 20 20 20

adaptive threshold {#findnodes adaptive threshold}

• Type: Real

• **Description**: The minimum difference considered in independent bands, the energy unit is eV. This means if the band gap is below this bar, it will be recognized as degenerate bands.

• **Default**: 0.001

kpoint_mode {#findnodes_kpoint_mode}

• Type: String

• **Description**: Used to set the k point. See *Setting of k points*

• Default: No default value

6.1.8 PDOS

stru_file {#pdos_stru_file}

• Type: String

• **Description**: The structure file name. This file records the structure of the lattice, the types of elements, and the atomic orbitals used. Make sure that both the structure file and the orbital file exist.

• **Default**: No default value

e_range {#pdos_e_range}

• Type: Real

• Description: The range of energy E. There are two parameters, indicating the starting point and the ending point.

• Default: No default value

de {#pdos_de}

• Type: Real

• **Description**: The interval dE for the energy E.

• **Default**: 0.01

sigma {#pdos_sigma}

• **Type**: Real

• **Description**: Parameters for gauss smearing.

• **Default**: 0.001

kpoint mode {#pdos kpoint mode}

• Type: String

• Description: Used to set the k point. See Setting of k points

• Default: No default value

6.1.9 FAT_BAND

band_range {#fatband_band_range}

• Type: Integer

• **Description**: There are two numbers (separated by spaces) to indicate which bands are selected for projection, counting from 1.

• Default: No default value

stru_file {#fatband_stru_file}

• Type: String

• **Description**: The structure file name. This file indicates the crystal structure and the corresponding orbital file. Make sure that both the structure file and the orbital file exist.

• Default: No default value

kpoint_mode {#fatband_kpoint_mode}

• Type: String

• **Description**: Used to set the k point of unitcell. See *Setting of k points*

• Default: No default value

6.1.10 SPIN_TEXTURE

nband {#spintexture_nband}

• Type: Integer

• **Description**: A band index. (Band index counts from 1)

kpoint_mode {#spintexture_kpoint_mode}

• Type: String

• **Description**: Used to set the k point. See *Setting of k points*

• Default: No default value

6.1.11 WILSON_LOOP

occ_band {#wilsonloop_occ_band}

• Type: Integer

• **Description**: The number of occupied energy bands of an insulator.

• Default: No default value

k_start {#wilsonloop_k_start}

• Type: Real

• Description: The origin point coordinates used to describe a Brillouin zone plane.

• **Default**: 0.0 0.0 0.0

k_vect1 {#wilsonloop_k_vect1}

• Type: Real

• **Description**: The expansion vector is a vector used to define a Brillouin zone plane, and it is also the direction of integration for calculations.

• **Default**: 1.0 0.0 0.0

k_vect2 {#wilsonloop_k_vect2}

• Type: Real

• **Description**: The expansion vector is a vector used to define a Brillouin zone plane, and it is also the direction of Wilson loop evolution for calculations.

• **Default**: 0.0 1.0 0.0

nk1 {#wilsonloop_nk1}

• Type: Integer

• **Description**: k_vect1 is divided into nk1 k-points.

• **Default**: 100

nk2 {#wilsonloop_nk2}

• Type: Integer

• **Description**: k_vect2 is divided into nk2 k-points.

• **Default**: 100

6.1.12 POLARIZATION

occ_band {#polarization_occ_band}

• Type: Integer

• Description: The number of occupied energy bands of an insulator.

• Default: No default value

nk1 {#polarization_nk1}

• Type: Integer

• **Description**: The number of samples in the x direction of reciprocal lattice vector **G**.

• Default: No default value

nk2 {#polarization_nk2}

• Type: Integer

• Description: The number of samples in the y direction of reciprocal lattice vector G.

• Default: No default value

nk3 {#polarization_nk3}

• **Type**: Integer

• Description: The number of samples in the z direction of reciprocal lattice vector G.

• Default: No default value

atom_type {#polarization_atom_type}

• Type: Integer

• **Description**: The number of element types in the system.

stru_file {#polarization_stru_file}

• Type: String

• **Description**: Specify the strucutre file. NAOs files are not required.

• Default: No default value

valence_e {#polarization_valence_e}

• Type: Integer

• **Description**: The number of valence electrons per element.

• Default: No default value

6.1.13 BERRY_CURVATURE

method {#berrycurvature_method}

• Type: Integer

• **Description**: Method for calculating berry curvature. 0 means direct calculation, 1 means calculation by Kubo formula.

• Default: 0

occ_band {#berrycurvature_occ_band}

• Type: Integer

• **Description**: The number of occupied energy bands of an insulator. When this value is not set, it will be determined according to the Fermi energy.

• Default: -1

kpoint_mode {#berrycurvature_kpoint_mode}

• Type: String

• **Description**: Used to set the k point. See *Setting of k points*

• Default: No default value

6.1.14 AHC

method {#ahc_method}

• Type: Integer

• **Description**: Method for calculating berry curvature. 0 means direct calculation, 1 means calculation by Kubo formula.

• Default: 0

integrate_mode {#ahc_integrate_mode}

• Type: String

• **Description**: Used for integration settings. See *Setting of integration*.

• Default: No default value

6.1.15 CHERN_NUMBER

method {#chernnumber_method}

• Type: Integer

• **Description**: Method for calculating berry curvature. 0 means direct calculation, 1 means calculation by Kubo formula.

• Default: 0

occ_band {#chernnumber_occ_band}

• Type: Integer

• **Description**: The number of occupied energy bands of an insulator. When this value is not set, it will be determined according to the Fermi energy.

• Default: -1

k_start {#chernnumber_k_start}

• Type: Real

• Description: The origin point coordinates used to describe a Brillouin zone plane.

• **Default**: 0.0 0.0 0.0

k_vect1 {#chernnumber_k_vect1}

• Type: Real

• **Description**: The expansion vector used to describe a Brillouin zone plane.

• **Default**: 1.0 0.0 0.0

k_vect2 {#chernnumber_k_vect2}

• Type: Real

• **Description**: The expansion vector used to describe a Brillouin zone plane.

• **Default**: 0.0 1.0 0.0

integrate_mode {#chernnumber_integrate_mode}

• Type: String

• **Description**: Used for integration settings. See *Setting of integration*.

• Default: No default value

6.1.16 CHIRALITY

method {#chirality_method}

• Type: Integer

• **Description**: Method for calculating berry curvature. 0 means direct calculation, 1 means calculation by Kubo formula.

• Default: 0

k_vect {#chirality_k_vect}

• Type: Real

Description: The k-point coordinates need to be calculated. There are three parameters to represent the coordinates.

• Default: No default value

radius {#chirality_radius}

• Type: Real

• **Description**: The radius of the integrating sphere. The unit is $^{-1}$.

• Default: No default value

point_num {#chirality_point_num}

• Type: Integer

• **Description**: The number of k-points that are uniformly sampled on a spherical surface.

• Default: No default value

6.1.17 JDOS

occ_band {#jdos_occ_band}

• Type: Integer

Description: Specifies the occupied energy band of the system. Currently, only insulator or semiconductor materials can be calculated.

omega {#jdos_omega}

• Type: Real

• **Description**: Specifies the photon energy, the unit is eV. There are two parameters, indicating the starting point and the ending point.

• Default: No default value

domega {#jdos_domega}

• Type: Real

• **Description**: The energy interval of ω .

• Default: No default value

eta {#jdos_eta}

• Type: Real

• **Description**: Specify the parameters of Gaussian smearing.

• **Default**: 0.01

grid {#jdos_grid}

• Type: Integer

• **Description**: The grid for integration. There are 3 parameters in total.

• Default: No default value

6.1.18 OPTICAL CONDUCTIVITY

occ_band {#opticalconductivity_occ_band}

• Type: Integer

• **Description**: Used to specify the occupied energy band of an insulator or semiconductor. Currently this function can only calculate insulators or semiconductors.

• Default: No default value

omega {#opticalconductivity_omega}

• Type: Real

• **Description**: The range of ω . There are two parameters, indicating the starting point and the ending point. Unit is eV.

domega {#opticalconductivity_domega}

• **Type**: Real

• **Description**: The energy interval of ω .

• Default: No default value

eta {#opticalconductivity_eta}

• Type: Real

• Description: Parameters for triangular smearing.

• **Default**: 0.01

grid {#opticalconductivity_grid}

• Type: Integer

• **Description**: The grid for integration. There are 3 parameters in total.

• Default: No default value

6.1.19 SHIFT_CURRENT

occ_band {#shiftcurrent_occ_band}

• Type: Integer

• **Description**: Used to specify the occupied energy band of an insulator or semiconductor. Currently this function can only calculate insulators or semiconductors.

• Default: No default value

omega {#shiftcurrent_omega}

• Type: Real

• **Description**: The range of ω . There are two parameters, indicating the starting point and the ending point. Unit is eV.

• Default: No default value

domega {#shiftcurrent_domega}

• Type: Real

• **Description**: The energy interval of ω .

smearing_method {#shiftcurrent_smearing_method}

• Type: Integer

• **Description**: The method of smearing. 0: no smearing. 1: Gaussian smearing. 2: adaptive smearing.

• Default: 1

eta {#shiftcurrent eta}

• Type: Real

• Description: Specify the parameters of Gaussian smearing.

• **Default**: 0.01

grid {#shiftcurrent grid}

• Type: Integer

• **Description**: The grid for integration. There are 3 parameters in total.

• Default: No default value

method {#shiftcurrent method}

• Type: Integer

• **Description**: Specify the method to calculate the shift current. 0 represents calculation using the Sternheimer equation, 1 represents the first order partial derivative calculation.

• Default: 1

6.1.20 BERRY CURVATURE DIPOLE

omega {#berrycurvaturedipole_omega}

• Type: Real

• **Description**: To set the energy range for the Berry curvature dipole, you can adjust it based on the Fermi energy level. The unit is eV. There are two parameters.

• Default: No default value

domega {#berrycurvaturedipole_domega}

• Type: Real

• **Description**: Specifies the energy interval of the omega.

integrate_mode {#berrycurvaturedipole_integrate_mode}

• Type: String

• **Description**: Used for integration settings. See *Setting of integration*. Since the integration is of a tensor, only 'Grid'integrate_mode is available.

• Default: No default value

6.2 Setting of k points

As long as the kpoint_mode parameter exists in FUNCTIONS, the following setting methods are to be followed.

6.2.1 When kpoint_mode is 'mp'

mp_grid

• Type: Integer

• **Description**: The grid dividing the Brillouin zone. There are three parameters to divide the three-dimensional Brillouin zone.

• Default: No default value

k_start

• Type: Real

• Description: The origin point coordinates of the Brillouin zone.

• **Default**: 0.0 0.0 0.0

k_vect1

• **Type**: Real

• **Description**: Expanded vector of the Brillouin zone.

• **Default**: 1.0 0.0 0.0

k_vect2

• **Type**: Real

• **Description**: Expanded vector of the Brillouin zone.

• **Default**: 0.0 1.0 0.0

k_vect3

• Type: Real

• Description: Expanded vector of the Brillouin zone.

• **Default**: 0.0 0.0 1.0

6.2.2 When kpoint mode is 'line'

kpoint_num

• Type: Integer

• **Description**: The number of high symmetry points.

• Default: No default value

high_symmetry_kpoint

• Type: Real

• **Description**: Fractional coordinates of high symmetry points and line densities of corresponding k-lines. The first three parameters are the fractional coordinates of the high symmetry points, and the fourth parameter is the line density.

• Default: No default value

6.2.3 When kpoint_mode is 'direct'

kpoint_num

• Type: Integer

• **Description**: the number of k points.

• Default: No default value

kpoint_direct_coor

• Type: Real

• **Description**: Fractional coordinates of the k point.

6.3 Setting of integration

As long as the integrate_mode parameter exists in FUNCTIONS, the following setting methods are followed.

6.3.1 When integrate_mode is 'Grid'

integrate_grid

• Type: Integer

• Description: Low precision grid for integration. There are three parameters.

• Default: 4 4 4

adaptive_grid

• Type: Integer

• Description: High precision grid for integration. There are three parameters.

• **Default**: 4 4 4

adaptive_grid_threshold

• Type: Real

• Description: If the value of a k point is greater than this value, then the k point will be adapted.

• **Default**: 50.0

6.3.2 When integrate mode is 'Adaptive'

relative_error

• Type: Real

• **Description**: The relative error of the adaptive integral.

• Default: 1e-6

absolute_error

• Type: Real

• **Description**: The absolute error of the adaptive integral.

• **Default**: 0.1

initial_grid

• Type: Real

• **Description**: The initial grid for adaptive integration. There are three parameters.

• **Default**: 1 1 1