MagneticTB: A package for tight-binding model of magnetic and non-magnetic materials

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Abstract
We present a Mathematica program package MagneticTB, which can generate the tight-binding model for arbitrary magnetic space group. The only input parameters in MagneticTB are the (magnetic) space group number and the orbital information in each Wyckoff positions. Some useful functions including getting the matrix expression for symmetry operators, manipulating the energy band structure by parameters and interfacing with other software are also developed. MagneticTB can help to investigate the physical properties in both magnetic and non-magnetic system, especially for topological properties.

Program summary
Program title: MagneticTB
Licensing provisions: GNU General Public Licence 3.0
Programming language: Mathematica
External routines/libraries used: ISOTROPY (iso.byu.edu)
Developer’s repository link: https://github.com/zhangzeyingvv/MagneticTB
Nature of problem: Construct the symmetry adopted tight-binding model for the system with arbitrary magnetic space group.

Keywords: Tight-binding method, Representation theory, Magnetic space group, Mathematica

1. Introduction
Tight-binding method is a powerful tool to investigate the novel properties in condensed mater physics [1–5]. Compared with first-principles method, tight-binding method can greatly simplify calculations. Moreover, after considering (magnetic) space group symmetry, the tight-binding model can give more reliable results. For example, in topological materials, symmetry plays an important role to protect the topological properties, such as $Z_2$ topological insulator protected by time reversal symmetry [6], topological crystalline insulators and topological nodal semimetals protected by space group symmetries [7, 8] and magnetic topological crystalline insulator protected by magnetic space group symmetries, i.e. the combination of space group operations and time reversal [9, 10].

At present, a lot of researchers using symmetry adopted tight-binding model to investigate physical properties of electronic system [11–21]. However, most of the software packages are mainly focused on the tight-binding model for non-magnetic materials [22–26] and only a few of them can be used to construct the symmetry adopted tight-binding model automatically. For the first-principles level, Wannier90 can generate the Wannier-tight-binding model by interfacing with first-principles software, but the symmetry

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Preprint submitted to Elsevier
adopted Wannier function can not be applied to other first-principles software (such as VASP, ABINIT) except Quantum-Espresso [27–30]. FPLO can generate symmetry adopted tight-binding model with proper parameters for given structures [31]. Meanwhile, both Wannier90 and FPLO do not support magnetic symmetry. For the model level, GTPack, Qsymm and MathemaTB can generate the tight-binding model with space group symmetry but does not support magnetic symmetry directly [32–34]. WannierTools can do the symmetrization of non-magnetic tight-binding model but cannot generate the tight-binding model by itself [35].

Therefore, it is necessary to develop a package which can construct the tight-binding model with magnetic space group symmetry automatically. Here we introduce a software package: MagneticTB, a tool for generating the tight-binding model for the system with arbitrary magnetic space group. The required input information of this package is only the magnetic space group number and the orbital information in each Wyckoff positions. It can help to investigate the physical properties of the given symmetry. We also present some useful functions including get the matrix expression for symmetry operators, manipulate the band structure by parameters and interface with other software.

This paper is organized as follows. In Sec. 2, we give an introduction of symmetry adopted tight-binding methods. In Sec. 3, The usage of MagneticTB are given, including how to install and run MagneticTB. In Sec. 4, we give three concrete examples, such examples show the specific capabilities of the MagneticTB. Finally, conclusions are given.

2. Symmetry adopted tight-binding method

In periodic system, the bases of tight-binding model can be written as Bloch sum [11]

$$\psi_{lmk}(r) = \frac{1}{\sqrt{N}} \sum_{R_j} e^{ik(R_j + d^m_j)} \varphi_m(r - R_j - d^n_l)$$

where \(N\) is the number of unit cells in the crystal, \(R_j\) is the translation vector of the Bravais lattice, \(d^n_l\) is the position of \(n\)-th Wyckoff position’s \(l\)-th atom in the unit cell (for each \(Q\) and \(d^n_l\) there exists one and only one pair of \(d^n_l\) and \(R\) which satisfies \(Qd^n_l = d^n_{l'} + R\) where \(Q\) is arbitrary group element in the (magnetic) space group and \(R\) is a lattice vector), \(\varphi_m(r)\) is the \(m\)-th atomic orbital basis for position \(d^n_l\) but located at coordinate origin. Then Eq. (1) satisfies the Bloch theorem \(\psi_{lmk}(r + R_j) = e^{ikR_j} \psi_{lmk}(r)\). The tight-binding Hamiltonian can be written as:

$$H_{lm'k'}^{n'n'}(k) = \sum_{R_j} e^{ik(R_j + d^n_l - d'^m_{l'})} E_{nm'm'}(d^n_l, R_j + d'^m_{l'})$$

(2)

$$E_{nm'm'}(d^n_l, R_j + d'^m_{l'}) = \langle \varphi_m(r - d^n_l) | \hat{H} | \varphi_{m'}(r - d'^m_{l'} - R_j) \rangle$$

for simplicity, we rewrite the atomic orbitals in vector form: \(\Phi^m(r - R_j - d^n_l) = \{ \varphi_m(r - R_j - d^n_l) \}, (m = 1, ..., M_n). Then E_{nm'm'}(d_j, R_j + d_{l'}) (m = 1, ..., M_n; m' = 1, ..., M_{n'}) form an \(M_n \times M_{n'}\) matrix:

$$E(d^n_l, R_j + d'^m_{l'}) = \langle \Phi^m(r - d^n_l) | \hat{H} | \Phi^{m'}(r - R_j - d'^m_{l'}) \rangle$$

(3)

Then the Hamiltonian can be rewritten as:

$$H_{lm'k'}^{n'n'}(k) = \sum_{R_j} e^{ik(R_j + d^n_l - d'^m_{l'})} E(d^n_l, R_j + d'^m_{l'})$$

(4)

\(E(d^n_l, R_j + d'^m_{l'})\) is the hopping matrix between \(n\)-th Wyckoff position’s \(l\)-th atom to \(n'\)-th Wyckoff position’s \(l'\)-th atom. When the lattice is in invariant under some symmetry \(E(d^n_l, R_j + d'^m_{l'})\) may not independent for arbitrary \(d^n_l\) and \(d'^m_{l'}\). Fortunately, for symmetry operation \(Q\) group representation theory gives us explicit expression for the relationship between \(E(Qd^n_l, Q(R_j + d'^m_{l'}))\) and \(E(d^n_l, R_j + d'^m_{l'})\).
For the case that symmetry operation does not contain the time reversal $\mathcal{T}$, i.e. $Q = \{R|t\}$, where $R$ and $t$ are the rotation and translation part of $Q$ respectively, we have

$$E(Qd_{l}^n, Q(R_j + d_{l}^n)) = D^n(R)E(d_{l}^n, R_j + d_{l}^n)D^{n\dagger}(R)$$  \hspace{1cm} (5)

For the case that operation $Q$ contains time reversal symmetry $\mathcal{T}$, i.e. $Q = \{R|t\mathcal{T}\}$, we have

$$E(Qd_{l}^n, Q(R_j + d_{l}^n)) = D^n(R\mathcal{T})E^*(d_{l}^n, R_j + d_{l}^n)D^{n\dagger}(RT)$$  \hspace{1cm} (6)

Where $D^n(R)(D^n(R\mathcal{T}))$ are the $M_n \times M_n$ representation matrices of $R(R\mathcal{T})$ under atomic orbital bases $\Phi^n(r)$ (not necessarily irreducible representations), $E^*(d_{l}^n, R_j + d_{l}^n)$ is complex conjugate of $E(d_{l}^n, R_j + d_{l}^n)$ (see Fig. 1 for example). It is clear that for spinless cases with time reversal symmetry $\mathcal{T}$, when the basis functions are real, $D(\mathcal{T})$ is equal to identity matrix, indicating that $E(d_{l}^n, R_j + d_{l}^n)$ are real matrices.

The next step is to get the analytical expressions of $D^n(R)(D^n(R\mathcal{T}))$. For a fixed $n$, we don’t have to worry about mixing superscripts of $n$ in $D^n(R)$ because transformations can only occur under the same $n$. So we temporarily use $D(R)$ rather than $D^n(R)$ in this step. Consider the following four cases:

i. Spinless system and $Q$ does not contain $\mathcal{T}$.

ii. Spinless system and $Q$ contain $\mathcal{T}$.

iii. Spinful system and $Q$ does not contain $\mathcal{T}$.

iv. Spinful system and $Q$ contain $\mathcal{T}$.

In case i, $D(R)$ can be obtained simply by solve the linear equation [36]

$$\hat{R}\Phi(r) = \Phi(R^{-1}r) = \Phi(r)D(R)$$  \hspace{1cm} (7)

in which $\hat{R}$ is the function operator for the rotation $R$. In case. ii, we define $\mathcal{F}(r) = \hat{T}\Phi(r)$, for spinless system $\mathcal{T} = \mathcal{K}$, hence, $\mathcal{F}(r) = \Phi^s(r)$ and then solve the linear equation

$$\hat{R}\mathcal{F}(r) = \mathcal{F}(R^{-1}r) = \Phi(r)D(R\mathcal{T})$$  \hspace{1cm} (8)

In case. iii, since the spin matrix is orbital independent we define the basis function as

$$\Phi^s(r) = \{\Phi(r)^\uparrow, \Phi(r)^\downarrow\}$$  \hspace{1cm} (9)
The two spinors $\uparrow = (1, 0)^T$ and $\downarrow = (0, 1)^T$, and under the rotation $\hat{R}$ they are transformed according to

$$\hat{R}(\uparrow, \downarrow) = (\uparrow, \downarrow)D^{\downarrow}(R)$$

For proper rotation $R$, $D^{\downarrow}(R) = \exp\left(-\frac{1}{2}i\alpha \mathbf{n} \cdot \mathbf{\hat{\sigma}}\right)$, where $\alpha$ is the rotation angle of $R$, $\mathbf{n}$ is the unit vector along rotation axis, for improper rotation $S$, $R = IS$, $I$ is the inversion symmetry, $D^{\downarrow}(S) = D^{\downarrow}(R)$ [37, 38].

Then $D(R)$ can be obtained by solving the linear equation

$$\hat{R}\Phi_s(r) = \Phi_s(r)D(R)$$

Case. (iv) is similar to case. (ii), the only difference is replace the time reversal operator $\mathcal{T} = \mathcal{K}$ by $\mathcal{T} = i\mathbf{\hat{\sigma}}_y \mathcal{K}$ and consider the spin rotation matrices. The above four cases cover all the possibilities of $D(R)$ and $D(R^T)$.

Then the operator (or representation matrix) for $Q$ can be defined as

$$P^{n' \leftarrow n}(Q) = \begin{cases} 
\delta_{nn'} & \text{Q does not contain } \mathcal{T} \\
\tilde{\delta}_{nn'} D^n(R) & \text{Q contains } \mathcal{T}
\end{cases}$$

where $\tilde{\delta}_{nn', Qd^n}$ is equal to 1 only when $d^n$ and $Qd^n$ differ by a lattice vector and to 0 otherwise (it can also be written as $\tilde{\delta}_{nn', Qd^n} = \delta_{nn', Qd^n} + R_s$ if a suitable lattice vector $R_s$ is chosen). The Hamiltonian under constraint of $Q$ can be written as

$$P(Q)^{-1}H(k)P(Q) = H(R^{-1}k)$$

for $Q = \{R|t\}$, and

$$P(Q)^{-1}H(k)P(Q) = H^*(-R^{-1}k)$$

for $Q = \{R|t\}^T$ [See Appendix A for proof of Eqs.(5–6) and Eqs.(13–14)]. Eqs.(13–14) are key point to generate the symmetry adopted tight-binding model. In MagneticTB we first generate the Hamiltonian with only translation symmetry and then use Eqs.(13–14) to the simplify the Hamiltonian (see Fig. 1(b)) for the workflow of MagneticTB. The database for tight-binding model of 1651 magnetic space group can be found in our later work [39].

3. Capabilities of MagneticTB

3.1. Installation

To install the MagneticTB, unzip the "MagneticTB.zip" file and copy the MagneticTB directory to any of the following four paths:

- FileNameJoin[{$UserBaseDirectory", "Applications$}]
- FileNameJoin[{$BaseDirectory", "Applications$}]
- FileNameJoin[{$InstallationDirectory", "AddOns", "Packages$}]
- FileNameJoin[{$InstallationDirectory", "AddOns", "Applications$}]

Then one can use the package after running Needs["MagneticTB"] . The version of Mathematica should higher or equal to 11.0.
3.2. Running
3.2.1. Core module

To initialize the program one should identify the magnetic space group and the orbital information in each Wyckoff positions. Here we provide a function `msgop` to show the symmetry information of an arbitrary magnetic space group. The only input of the `msgop` is the magnetic space group number, one can get the symmetry information by the following code:

```
msgop[gray[191]]
msgop[bnsdict[[191, 236]]]
msgop[ogdict[[191, 8, 1470]]]
```

Magnetic space group (BNS): {191.236,P6'/mm'm}

```
Primitive Lattice vector: {{a,0,0},{-a/2,(Sqrt[3] a)/2,0},{0,0,c}}
Conventional Lattice vector: {{a,0,0},{-a/2,(Sqrt[3] a)/2,0},{0,0,c}}
```

```
{{1, ,{{1,0,0},{0,1,0},{0,0,1}},{0,0,0}, F},
{3z,{{0,1,0},{1,1,0},{0,0,1}},{0,0,0},F},
{2x,{{1,1,0},{-1,0,0},{0,0,1}},{0,0,0},F},
...
```

doesn't return the magnetic space group code of gray space group 191, `bnsdict[[191,236]]` return the magnetic space group code of BNS No. 191.236, `ogdict[[191,8,1470]]` return the magnetic space group code of OG No. 191.8.1470. Then the `msgop` will print the standard lattice vector and the symmetry operations (for primitive cell) of the corresponding magnetic space group, which can be the input of the `init` function. Notice the magnetic space group code is build-in constant in MagneticTB, users should use `gray`, `bnsdict`, `ogdict` functions rather than inputting the magnetic space group code directly.

Then one can feed the above information to `init` function then the basic results of input structure can be generated. The `init` function have five mandatory options neamly, `lattice`, `lattpar`, `wyckoffposition`, `symminformation` and `basisFunctions`. The `lattice` is the lattice vector of magnetic system which can contain parameters, `lattpar` is the parameters in lattice vector to determine the bond length of magnetic system, and `wyckoffposition` is a list to designate atomic position and magnetization-direction for each Wyckoff positions in the magnetic system. The format of `wyckoffposition` is:

```
{{a, m1}, {a2, m2}, ...}
```

where `a` and `m`, represent one of the atomic positions and its magnetization-directions of the `i`-th Wyckoff position, respectively. The `symminformation` contain the elements of the coset of magnetic space group with respect to translation group, which can direct use the output of `msgop` (notice that the output of `msgop` is standard symmetry operation from ISOTROPY [40, 41]. However, users can also use the non-standard structure as input, not limit to the output of `msgop`). The format of `symminformation` is:

```
{{n1, R1, t1, A1}, {n2, R2, t2, A2}, ...}
```

where `ni` is the name of symmetry operation, `Ri` and `ti` are the rotation and translation part of symmetry operation, and `Ai` represents whether the symmetry operation is combined with time reversal symmetry ("T" for true and "F" for false). Finally, the `basisFunctions` is the basis function for each Wyckoff position. The format of `basisFunctions` is:

```
{b1, b2, ..., }
```

where `bi` is the list of basis functions of the `i`-th Wyckoff position. The build-in basis functions for spinless case in MagneticTB is shown in Table 1. When spin is considered, for build-in basis functions, add "up" or "dn" after basis functions string, e.g. for |p↑⟩, the basis function for spin-up case is "pup". However, users may use other basis functions such as f, |3/2, 1/2⟩ orbitals. In such cases, users can directly input the analytical expression of basis functions. For example, if only consider fxyz orbital, one should input `basisFunctions -> {{x*y*z}}`. The analytical expressions of basis functions can be simply obtained from quantum mechanics or group theory books [32, 42, 43].
Table 1: String codes representing basis functions and available values for \texttt{basisFunctions}

| Basis function String | Basis function String |
|-----------------------|-----------------------|
| s                     | "s"                   |
| p                     | "py"                  |
| z                     | "pz"                  |
| p x                   | "px"                  |
| p y                   | "py"                  |
| p z                   | "pz"                  |
| p x + ip y            | "px+ipy"              |
| p x − ip y            | "px−ipy"              |
| d z<sup>2</sup>        | "dz2"                 |
| d x y                  | "dxy"                 |
| d y z                  | "dyz"                 |
| d x z                  | "dxz"                 |

\texttt{sgop=sgop[gray[191]];}

\texttt{init [}
\texttt{lattice -> \{a,0,0\},\{(−a/2),(Sqrt[3] a)/2,0\},\{0,0,c\}},
\texttt{latpar -> \{a -> 1, c -> 3\},}
\texttt{wyckoffposition -> \{\{(1/3, 2/3, 0), \{0, 0, 0\}\},}
\texttt{symminformation -> \texttt{sgop},
\texttt{basisFunctions -> \{"pz"\}}];

Table 2: Basic results of \texttt{init}

| properties | illustrate of properties |
|------------|--------------------------|
| atompos    | atomic position and magnetization-direction for each atom |
| wcc        | \(d_l\) for each basis function |
| reclatt    | reciprocal lattice vector for given structure |
| symmetryops| \(P_Q\) for each symmetry operation |
| unsymham   | generate the Hamiltonian with only translation symmetry |
| symmcompile | summary of \(P(Q)\), see main text for detail |
| bondclassify| summary of bonds information, see main text for detail |

After inputting the above five options appropriately, one can run \texttt{init} and obtain the basic results. Here we introduce two important basic results: \texttt{symmcompile} and \texttt{bondclassify}, the other properties are given in Table 2. The format of \texttt{symmcompile} is

\[
\texttt{symmcompile} \{\{N_1, \{n_1, R_1, t_1, A_1\}, P_1, R^k_1\}, \{N_2, \{n_2, R_2, t_2, A_2\}, P_2, R^k_2\}, ...\}
\]

where \(N_i, P_i, R^k_i\) are the label, the symmetry operator (Eq. (12)) and the rotation acting on \(k\) space of the \(i\)-th symmetry operation, respectively. For example

\texttt{symmcompile}

\[
\{1,\{1\},\{1,0,0\},\{0,1,0\},\{0,0,1\}\},\{0,0,0\}, \text{ F.},\{1,0,0\},\{0,1,0\},\{0,0,1\}\},
\{1,\{1\},\{1,0,0\},\{0,1,0\},\{0,0,1\}\},\{0,0,0\}, \text{ F.},\{1,0,0\},\{0,1,0\},\{0,0,1\}\},
\{0,0,0\}, \text{ F.},\{1,0,0\},\{1,0,0\},\{0,0,1\}\},
\{0,0,0\}, \text{ F.},\{0,1,0\},\{1,0,0\},\{0,0,1\}\},
\{0,0,0\}, \text{ F.},\{1,0,0\},\{1,0,0\},\{0,0,1\}\},...\}
\]

The format of \texttt{bondclassify} is

\[
\{l_i, n_i, \{\{p_{i,j}, p_{i,k}\}, ...\}\}, ...
\]

where \(l_i\) is the bond length of the \((i-1)\)-th neighbour hopping \((i = 1 \text{ for on-site hopping})\), \(n_i\) is the number of the \((i-1)\)-th neighbour’s bonds, \(\{p_{i,j}, p_{i,k}\}\) is the atomic position of \(n_i\) corresponding bonds.

Till this moment, symmetry adopted tight-binding model for magnetic system is ready to be generated. By using the \texttt{symham[n]} function, one can obtain the symmetry adopted tight-binding model. When \(n = 1\), \texttt{symham[1]} return the Hamiltonian with only on-site hopping, \(n = 2\) return the Hamiltonian with only nearest-neighbour hopping and so on. By default, MagneticTB will check all the input symmetry operations to ensure the Hamiltonian is correct. However, it may last long time when the structure is complex. In principle, only
the generators of the (magnetic) space group are enough to get the Hamiltonian. Therefore, one can specify
the symmetry operations by \texttt{symmetryset-list} in \texttt{symham} where \texttt{list} is the list of indexes of the symmetry
operations. For example, \texttt{symham[2,symmetryset->{2}]} will generate the Hamiltonian with only \(C_{6z}\) symmetry
for nearest-neighbour hopping. \texttt{symmetryset} can not only save computing resource but can also investigate
the Hamiltonian for symmetry breaking cases. The parameters for each neighbour in MagneticTB are given
in Table. 3.

| On-site energy | Nearest | Second-nearest | Third-nearest | \((k-1)\)-th nearest |
|----------------|---------|----------------|--------------|----------------------|
| \(e_1,e_2,..\) | \(t_1,t_2,..\) | \(r_1,r_2,..\) | \(s_1,s_2,..\) | \(pkn1,pkn2,..\) |

3.2.2. Plot module

After tight-binding model being generated, there may exist many parameters, one can use \texttt{bandManipulate}
function to manipulate the band structure to investigate the relationship between band structure and
parameters. The format of \texttt{bandManipulate} is

\[
\texttt{bandManipulate[{{{k1,k2},{name of k1, name of k2}},...},np,Hamiltonian]}
\]

where \(np\) is the number of \(k\) points per line. Then one can easily check the band structure of different
parameters. When the proper parameters are obtained, one can use \texttt{bandplot} to plot the band structure

\[
\texttt{bandplot[{{{k1,k2},{name of k1, name of k2 }},...}, np,Hamiltonian,parameters]}
\]

see section. 4 for concrete example.

3.2.3. IO module

In MagnetTB, one can get the tight-binding model for magnetic system. However, MagneticTB do not
calculate the other properties (such as surface states, finding the gap-less point and so on) directly, since
it will generally cost too much computing resources. It is better to do such heavy calculations by Fortran,
Python or C. Therefore we develop \texttt{hopp} function to convert the symmetry adopted tight-binding model to
"wannier90_hr.dat" format, which is convenient to interface with WannierTools [35], Z2Pack [44], PythTB
[45] and our home-made package Wannflow [46, 47]. The "wannier90_hr.dat" in Wannier90 use the following
convention [27], namely conventions II:

\[
\begin{aligned}
\tilde{\psi}_{nlm}^m(r) &= \frac{1}{\sqrt{N}} \sum_{R_j} e^{ik \cdot R_j} \psi_{nlm}^m(r - R_j - d_n^m) \\
\tilde{H}_{nn'}^{ll'}(k) &= \sum_{R_j} e^{ik \cdot R_j} E_{mnv}(d_n^m, R_j + d_{n'}^{l'})
\end{aligned}
\] (15)

which is different form MagneticTB in Eq. (2), the relationship between two conventions is

\[
\begin{aligned}
\tilde{H}(k) &= V(k)H(k)V^\dagger(k) \\
V_{nn'}^{ll'}(k) &= e^{ik \cdot d_{n'}^{l'}} \delta_{nn'}
\end{aligned}
\] (16)

In MagneticTB (convention I) the operation matrix defined in Eq. (12) is \(k\) independent while the Hamiltonian
is non-periodic by shifting the reciprocal vector \(G\)

\[
H(k + G) = V^\dagger(G)H(k)V(G)
\] (17)

By contrast, in conventions II the Hamiltonian is periodic. i.e \(\tilde{H}(k + G) = \tilde{H}(k)\). The format of \texttt{hopp}
function is

\[
\]
See section 4 for concrete example. One can also use symmham[],ham] to generate the Wolfram expression for Hamiltonian in convention II. Notice that hopp function (but not symmham[]) is only applied to the output of symham function, and that expressions explicitly including Sin or Cos may not work well. Be careful to use it.

4. Examples

4.1. Three-band tight-binding model for MoS$_2$

MoS$_2$ monolayer has direct bandgap in the visible range, strong spin-orbit coupling, and rich valley related physics, which make it an candidate for nanoelectronic, optoelectronic, and valleytronic applications [48, 49]. The space group of MoS$_2$ is $P6/m2$ (space group No. 187). Considering the Mo atom at 1$a$ Wyckoff position and using the $d_{z^2}$, $d_{xy}$, and $d_{x^2−y^2}$ orbitals, the model can be obtained by

\[
\text{sgop} = \text{msgop}[\text{gray}[187]]; \\
\text{tran} = \{(1, 1, 0), (0, 1, 0), (0, 0, 1)\}; \\
\text{sgoptr} = \text{MapAt}[\text{FullSimplify}[\text{tran}.#\cdot\text{Inverse}[\text{tran}] &, \text{sgop}, \{\ldots, 2\}]; \\
\text{init} \{ \\
\text{lattice} -> \{(1, 0, 0), (1/2, \text{Sqrt}[3]/2, 0), (0, 0, 10)\}, \\
\text{lattpar} -> \{\}, \\
\text{wyckoffposition} -> \{\{0, 0, 0\}, \{0, 0, 0\}\}, \\
\text{symminformation} -> \text{sgoptr}, \\
\text{basisFunctions} -> \{\text{”dz2”, ”dxy”, ”dx2−y2”}\}; \}
\]

\[
\text{mos2} = \text{Sum}[\text{symham}[\text{i}, \text{symmetryset} = \{9, 11, 13\}], \{\text{i}, \{1, 2\}\}]; \\
\text{mos2Liu} = \text{mos2} /. \text{Thread}[\{\text{kx, ky, kz}\} -> (\{\text{kx, ky, kz}\} (2\pi))\cdot\text{Inverse}[\text{reclatt}]]; \\
\]

This gives exactly the same results as in Ref. [14]. The relationship of parameters between Ref. [14] and MagneticTB are

| Ref. [14] | $\epsilon_1$ | $\epsilon_2$ | $t_0$ | $t_1$ | $t_2$ | $t_{11}$ | $t_{12}$ | $t_{22}$ |
|----------|-------------|-------------|------|------|------|--------|--------|--------|
| MagneticTB | $e_1$ | $e_2$ | $t_1$ | $t_2$ | $t_3$ | $t_4$ | $t_5$ | $t_6$ |

4.2. Graphene

Graphene with linear dispersion around Fermi level is one of the most important materials in spintronics [50]. The magnetic space group of graphene is $P6/mmm1'$ (BNS No. 191.234). There are two C atoms at 2$c$ Wyckoff position, and the bands near Fermi energy are mainly from $p_z$ orbital. The above information is enough to establish the tight-binding model near Fermi energy of graphene. The model can be obtained as follow

\[
\text{Needs}["\text{MagneticTB’"]} \\
\text{sgop} = \text{msgop}[\text{gray}[191]]; \\
\text{init} \{ \\
\text{lattice} -> \{(a, 0, 0), (-a/2, \text{Sqrt}[3] a, 0), (0, 0, c)\}, \\
\text{lattpar} -> \{a -> 1, c -> 3\}, \\
\text{wyckoffposition} -> \{\{1/3, 2/3, 0\}, \{0, 0, 0\}\}, \\
\text{symminformation} -> \text{sgop}, \\
\text{basisFunctions} -> \{\text{”pz”}\}; \\
\text{ham} = \text{Sum}[\text{symham}[\text{i}], \{\text{i}, \{3\}\}]; \text{MatrixForm}[\text{ham}]
\]

output:

\[
\begin{bmatrix}
    e_1 + 2r_1(\cos(k_x + k_y) + \cos k_x + \cos k_y) \\
    t_1e^{i(-\frac{\sqrt{3}a}{2} - \frac{k_x}{2})} + t_1e^{i(\frac{\sqrt{3}a}{2} - \frac{k_y}{2})} + t_1e^{i(\frac{\sqrt{3}a}{2} + \frac{k_y}{2})} \\
    e_1 + 2r_1(\cos(k_x + k_y) + \cos k_x + \cos k_y)
\end{bmatrix}
\]

For spin-orbital coupling case, the only thing which needs to change is the basis functions

basisFunctions -> {"pzup", "pzdn"}
and the corresponding Hamiltonian reads
\[
\begin{pmatrix}
e_1 + h^- & 0 & h' & 0 \\
0 & e_1 + h^+ & 0 & h' \\
0 & 0 & e_1 + h^+ & 0 \\
0 & 0 & 0 & e_1 + h^-
\end{pmatrix}
\]
where
\[h^\pm = \pm 2r_1 (\sin k_x + \sin k_y - \sin k_x + k_y) + 2r_2 (\cos k_x + k_y) + \cos k_x + \cos k_y),
\]
\[h' = t_1 e^{i\left(-\frac{2\pi x}{a} - \frac{\pi y}{b}\right)} + t_1 e^{i\left(\frac{2\pi x}{a} + \frac{2\pi y}{b}\right)}.\]

Such model is corresponding to the first $Z_2$ topological insulator \[6\]. After get the Hamiltonian, we can use \texttt{bandManipulate} and \texttt{bandplot} to plot the band structure, and click the "ExportData" button to print the value of parameters.

\[
\text{path}=\{
\{\{0,0,0\},\{0,1/2,0\}\},\{\text{\textquotedbl}CapitalGamma\text{\textquotedbl},\text{\textquotedbl}M\text{\textquotedbl}\}\},
\{\{0.1/2.0\},\{1/3.1/3.0\}\},\{\text{\textquotedbl}M\text{\textquotedbl},\text{\textquotedbl}K\text{\textquotedbl}\}\},
\{\{1/3.1/3.0\},\{0,0,0\}\},\{\text{\textquotedbl}K\text{\textquotedbl},\{\text{\textquotedbl}CapitalGamma\text{\textquotedbl}\}\}\};
\text{bandManipulate}[\text{path}, 20, \text{ham}]
\text{bandManipulate}[\text{path}, 20, \text{hamsoc}]
\text{bandplot}[\text{path}, 200, \text{ham}, \{e_1 \to 0, t_1 \to 0.5, r_1 \to 0\}]
\text{bandplot}[\text{path}, 200, \text{hamsoc}, \{e_1 \to 0, t_1 \to 0.5, r_1 \to 0.02, r_2 \to 0\}]
\]

Moreover we can get the wannier90_hr.dat by \texttt{hop} function for this model.

\[
\text{hop}[\text{hamsoc}, \{e_1 \to 0, r_1 \to 0.02, r_2 \to 0, t_1 \to 0.5\}]
\text{Generated by MagneticTB}
\]

![Figure 2](image-url)

Figure 2: Output of \texttt{bandManipulate} and \texttt{bandplot} for graphene without considering spin (a-b) and with spin (c-d).

Moreover we can get the wannier90_hr.dat by \texttt{hop} function for this model.
1 1 1 1 1 1 1
-1 -1 0 1 1 0.0000000 0.0200000
-1 -1 0 2 1 0.0000000 0.0000000
-1 -1 0 3 1 0.0000000 0.0000000
-1 -1 0 4 1 0.0000000 0.0000000
-1 -1 0 1 2 0.0000000 0.0000000
-1 -1 0 2 2 0.0000000 0.0200000

4.3. Magnetic C-3 Weyl point

The charge-3 (C-3) Weyl point is a 0D two-fold band degeneracy with Chern number $|C| = 3$. Encyclopedia of emergent particles tell us that the C-3 Weyl point always appear at least in a pair or coexist with nodal surface in nonmagnetic systems [51]. Here we confirm that in magnetic system, due to the breaking of time reversal symmetry $T$, the C-3 Weyl point can uniquely coexist with conventional Weyl points. Consider the type IV magnetic space group $P_{c3}$ (BNS No. 143.3). The generator of the group is $C_3z$ and $\{E|001\}^T$. Put $|p_x + ip_y \uparrow\rangle, |p_x - ip_y \downarrow\rangle$ basis functions at Wyckoff position 2a, and then the symmetry operator for $C_3z$ and $E\{001\}^T$ are

$$C_3z = -\sigma_z$$
$$\{E\{001\}^T = i\sigma_y$$

Under this bases the effective Hamiltonian at $\Gamma$ point can be written as

$$H_{C-3 WP} = \epsilon + \alpha k_z \sigma_x + ck_\parallel^2 + \beta(k_x + e^{-i\frac{2\pi}{3}}k_y)^3 \sigma_z + h.c. \quad (18)$$

where $\epsilon, c$ are real parameters and $\alpha, \beta$ are complex parameters. Besides, there are another three essential Weyl points locate at $(\pi, 0, 0), (0, \pi, 0), (\pi, \pi, 0)$. Since the $C_3z$ symmetry does not change the Chern number of Weyl points, the Chern number at $M$ has to be $\pm 1$. According to no-go theorem, the Chern number of $\Gamma$ is $\mp 3$. One can easily check that the Chern number of Eq. (18) is $\pm 3$. The degeneracies of $\Gamma$ and $M$ are because $(\{E\{001\}^T)^2 = -1$ at $(0/\pi, 0/\pi, 0)$. The model can be obtained as follow

\begin{verbatim}
sgop = msgop[bnscdict[{143, 3}]]; init[lattice -> {{Sqrt[3]/2, -1/2, 0}, {0, 1, 0}, {0, 0, 2}}, latpar -> {}]; wyckoffposition -> {{0, 0, 0}, {0, 0, 1}}]; symminformation -> sgop; basisFunctions -> {{x + I y, 0}, {x - I y}}]; c3w = Sum[symham[i], {i, {2, 4}}]; c3w2band = Table[c3w[i, j], {i, {1, 4}}, {j, {1, 4}}];
\end{verbatim}

The band structure of $c3w2band$ is shown in Fig. 3(a).

4.4. Magnetic cubic nodal-line

Topological high order nodal line is that the energy difference between the bands are non-linear, and the order of energy dispersion around the degeneracy points plays an important role in different physical properties, such as density of states, Berry phase and Landau-level [52]. Recently, Zhang et. al. proposed high order nodal-line in magnetic system [53]. In this example, we use MagneticTB to generate magnetic cubic nodal-line. Generally, the magnetic cubic nodal-line is protected by $C_{6z}$ and $M_x$ symmetries, there are many magnetic space groups which contains the above two symmetries. Consider the type IV magnetic space group $P_{6cc}$ (BNS No. 184.196), and put $|p_x + ip_y \uparrow\rangle, |p_x - ip_y \downarrow\rangle$ basis functions at Wyckoff position 2a. Then the tight-binding model can be generated by

\begin{verbatim}
sgop = msgop[bnscdict[{184, 196}]]; init[lattice -> {{Sqrt[3]/2, -1/2, 0}, {0, 1, 0}, {0, 0, 2}}, latpar -> {}];
\end{verbatim}

wyckoffposition -> {{0, 0, 0}, {0, 0, 1}}.
syminformation -> sgop.
basisFunctions -> {{{x + I y, 0}, {0, x - I y}}};
cnl = Sum[symham[i, symmetryset -> {2, 7, 13}], {i, 1, 5}];
cnl2band = Table[cnl[[i, j]], {i, {1, 4}}, {j, {1, 4}}];
MatrixForm[cnl2band]
path = {{
  {0, 1/10, 1/4}, {0, 0, 1/4}, {"Q", "P"},
  {{0, 0, 1/4}, {0, 0, 1/4}, {"P", "Q"}}};
bandManipulate[path, 20, cnl2band]

One can check that no matter how the parameters change, the dispersion of arbitrary point along \( \Gamma \)-\( A \) on \( k_x-k_y \) plane are non-linear, see Fig. 3(b) which is consistent with Ref. [53].

5. Conclusion

In conclusion, we have developed a software package to generate the symmetry-adopted tight-binding model for arbitrary magnetic space group. The input parameters for MagneticTB are clear and easy to set and both spinless and spinful Hamiltonian can be generated automatically. Besides, some useful functions such as manipulating the band structure, interfacing with other software are implemented, which can be used for further study on the magnetic systems. Moreover, MagneticTB can not only be used to investigate physical properties of electronic systems, but also be used to study photonics, ultracold, acoustic and mechanical systems [54–56]. Finally, an exciting direction for future is to apply the magnetic field for the tight-binding model in MagneticTB [57].

Acknowledgments

ZZ acknowledges the support by the NSF of China (Grant No. 12004028), the China Postdoctoral Science Foundation (Grant No. 2020M670106), the Fundamental Research Funds for the Central Universities (ZY2018). GBL acknowledges the support by the National Key R&D Program of China (Grant No. 2017YFB0701600). YY acknowledges the support by the National Key R&D Program of China (Grant No. 2020YFA0308800), the NSF of China (Grants Nos. 11734003, 12061131002), the Strategic Priority Research Program of Chinese Academy of Sciences (Grant No. XDB30000000).
Appendix A.

In this appendix, we use the translation operation $T(d)$. For $Q = \{ R|v \}$, we have $QT(d) = \{ R|v \} \{ E|d \} = \{ R|Rd + v \} = T(Qd)R$. Thus

\[
E(d_i^n, (R_j + d_i^n')) = \langle \hat{Q}(d_i^n)| \hat{R}(Q(d_i^n + R_j)) | \hat{Q}(d_i^n + R_j) \rangle \phi'(r)
\]

which completes the proof of Eq.(5).

For $Q = \{ R|v \} T$, notice time reversal symmetry does not change the real space coordinates, i.e. $Td = d$, we have $QT(d) = \{ R|v \} T \{ E|d \} = \{ R|Rd + v \} T = T(Qd)RT$. Use the fact that for anti-unitary operator $\hat{A}$, $\langle \hat{A}\psi, \hat{A}\phi \rangle = \langle \psi|\phi \rangle^*$, then

\[
E^+(d_i^n, (R_j + d_i^n')) = \langle \hat{Q}*\hat{T}(d_i^n)| \hat{R}\hat{T}(Q(d_i^n + R_j)) | \hat{Q}(d_i^n + R_j) \rangle \phi'(r)
\]

which completes the proof of Eq.(6).

Proof of Eq.(13),

\[
[H(k)P(Q)]^{nn'}_{ll'} = \sum_{\mu \nu} H(k)^{ll'}_{\mu \nu} P^{\mu n'}_{\mu l'}(Q)
\]

\[
= \sum_{\mu \nu} \sum_{R_j} e^{ik \cdot (R_j + d_i^n - d_i^n)} E(d_i^n, R_j + d_i^n) \delta_{\nu l'} Q_{d_i^n + R_j} D^{n'}(R)
\]

\[
= \sum_{R_j} e^{ik \cdot (R_j + Qd_i^n + R_j - d_i^n)} E(d_i^n, R_j + Qd_i^n + R_j) D^{n'}(R)
\]

\[
\text{use Eq.(5) } \sum_{R_j} e^{i R^{-1} k \cdot (R_j + d_i^n - Q^{-1} d_i^n)} E(d_i^n, Q(R_j + d_i^n')) D^{n} (R) E(Q^{-1} d_i^n, R_j + d_i^n')
\]
\[ [P(Q)H(R^{-1}k)]_{\text{li}}^{\mu\nu'} = \sum_{\mu'\nu'} P^{\mu\nu}_{\mu'}(Q) H(R^{-1}k)_{\mu'\nu'} \]
\[ = \sum_{\mu'\nu'} \sum_{R_j} \delta_{\nu\nu'} \delta_{\mu\mu'} \delta_{\mu'} Q_d R_j D^n(R) e^{iR^{-1}k(R_j + d^{\mu'}_n - d^{\nu'}_n)} E(d^{\mu'}_n, R_j + d^{\nu'}_n) \]
\[ = \sum_{R_j} D^n(R) e^{iR^{-1}k(R_j + d^{\mu'}_n - d^{\nu'}_n)} E(Q^{-1}(d^{\mu'}_n - R_s), R_j + d^{\nu'}_n) \]
\[ = \sum_{R_j} e^{iR^{-1}k(R_j + d^{\mu'}_n - d^{\nu'}_n + R^{-1}R_s)} D^n(R) E(Q^{-1}d^{\mu'}_n - R^{-1}R_s, R_j + d^{\nu'}_n) \]
\[ = \sum_{R_j} e^{iR^{-1}k(R_j + d^{\mu'}_n - d^{\nu'}_n + R^{-1}R_s)} D^n(R) E(Q^{-1}d^{\mu'}_n, R_j + d^{\nu'}_n + R^{-1}R_s) \]
\[ \Rightarrow H(k) P(Q) = P(Q) H(R^{-1}k) \quad (A.4) \]

In the above derivation we use the relation \( Q^{-1}(d^{\mu'}_n - R_s) = Q^{-1}d^{\mu'}_n - R^{-1}R_s \) \((Q^{-1} \text{ is not linear})\). Compare the last lines of the above two equations and we can find they are equal to each other, i.e.
\[ [H(k) P(Q)]_{\text{li}}^{\mu\nu'} = [P(Q) H(R^{-1}k)]_{\text{li}}^{\mu\nu'} \Rightarrow H(k) P(Q) = P(Q) H(R^{-1}k) \quad (A.5) \]

Proof of Eq.\((14)\): similar to Eq.\((A.3)\) and Eq.\((A.4)\) we have
\[ [H(k) P(Q)]_{\text{li}}^{\mu\nu'} = \sum_{\mu'\nu'} H(k)_{\mu\mu'} P^{\mu\nu'}_{\mu'} (Q) \]
\[ = \sum_{\mu'\nu'} \sum_{R_j} e^{i\mu} (R_j + d^{\mu'}_n - d^{\nu'}_n) E(d^{\mu'}_n, R_j + d^{\nu'}_n) \delta_{\mu\mu'} \delta_{d^{\mu'}_n, Q_d R_j} D^n(R T) \]
\[ = \sum_{R_j} D^n(R T) e^{iR^{-1}k(R_j + d^{\mu'}_n - d^{\nu'}_n + R^{-1}R_s)} E(Q^{-1}(d^{\mu'}_n, R_j + d^{\nu'}_n) D^n(R T) \]
\[ \Rightarrow \text{use Eq.\((6)\)} \sum_{R_j} e^{iR^{-1}k(R_j + d^{\mu'}_n - d^{\nu'}_n + R^{-1}R_s)} E(Q^{-1}d^{\mu'}_n, R_j + d^{\nu'}_n) \]
\[ [P(Q) H^*(-R^{-1}k)]_{\text{li}}^{\mu\nu'} = \sum_{\mu'\nu'} P^{\mu\nu}_{\mu'}(Q) H^*(-R^{-1}k)_{\mu'\nu'} \]
\[ = \sum_{\mu'\nu'} \sum_{R_j} \delta_{\nu\nu'} \delta_{\mu\mu'} \delta_{\mu'} Q_d R_j D^n(R T) e^{iR^{-1}k(R_j + d^{\mu'}_n - d^{\nu'}_n)} E^*(d^{\mu'}_n, R_j + d^{\nu'}_n) \]
\[ = \sum_{R_j} D^n(R T) e^{iR^{-1}k(R_j + d^{\mu'}_n - d^{\nu'}_n - R^{-1}R_s)} E^*(Q^{-1}(d^{\mu'}_n - R_s), R_j + d^{\nu'}_n) \]
\[ = \sum_{R_j} e^{iR^{-1}k(R_j + d^{\mu'}_n - d^{\nu'}_n + R^{-1}R_s)} D^n(R T) E^*(Q^{-1}d^{\mu'}_n - R^{-1}R_s, R_j + d^{\nu'}_n) \]
\[ = \sum_{R_j} e^{iR^{-1}k(R_j + d^{\mu'}_n - d^{\nu'}_n + R^{-1}R_s)} D^n(R T) E^*(Q^{-1}d^{\mu'}_n, R_j + d^{\nu'}_n + R^{-1}R_s) \]
\[ \Rightarrow H(k) P(Q) = P(Q) H^*(-R^{-1}k) \quad (A.8) \]
It is easy to verify that for $A, B$ not containing $T$ and $C, D$ containing $T$, $P(Q)$ obey the following corepresentation algebra:

$$P(A)P(B) = P(AB)$$
$$P(A)P(C) = P(AC)$$
$$P(C)P^*(A) = P(CA)$$
$$P(C)P^*(D) = P(CD)$$

(A.9)

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