Abstract

Theoretical condensed matter research is plagued by a fundamental issue of complexity. The sheer amount of degrees of freedom in a material on any technologically relevant scale is overwhelming (e.g. $\sim 10^{23}$ electrons per cm³), and makes it impossible to describe the quantum mechanical wavefunction exactly.

The Hamiltonian plays a central role in the description of crystals, the subject of this thesis. It can be decomposed into various parts, and their interactions. Depending on the physics under scrutiny it then often suffices to solve only one of those parts. This can be either because the energy scales and associated timescales that govern the constituents are very different, or because the interactions between them are small. One example, often put into practice, is the separation of electronic and phononic (lattice) degrees of freedom, leading to the well-known Born-Oppenheimer approximation, decoupling their respective motion. Another is the often neglected spin-orbit coupling, due to the tiny prefactor associated with its relativistic origin.

Solving these subproblems then allows for progress to be made in understanding the physics that govern them. However, there will inevitably be systems for which this interaction is not small and leads to fascinating new physics that manifestly depends on both subsystems combined. In this thesis we focus on these cases and how they arise in functional materials, with the occasional eye towards applications in technology.

The reason why these cross-order couplings can be interesting for technological applications, is that often one of the orders is more robust with respect to perturbations, and therefore more long-lived, but also harder to control efficiently. By exploiting the cross-order coupling in certain materials, one could potentially control the long lived order by applying perturbations to the more easily controllable order.

In giant Rashba effect systems, the coupling between spin and ferroelectric order leads to a linear spin-splitting of the band structure, whose sign depends on the orientation of the ferroelectric polarization. We show that, rather than the relativistic Rashba effect, a combination of electrostatics and atomic spin-orbit coupling lies at the origin of the large splitting.

The coupling between magnetism and ferroelectricity in multiferroic $GdMn_2O_5$ leads to a never before observed four-state hysteresis loop for the ferroelectric polarization, which depends on the magnitude, angle and history of the applied magnetic field. As we will show, this four-state hysteresis loop is accompanied by a full 360° rotation of spins in the material, which resembles the crankshaft of a car, converting the linear back-and-forth motion of the magnetic field into a rotational motion of the spins.

In a thin film of elemental Chromium, the ultrafast dynamics of a spin density wave, coupled to a slower varying charge density wave, allows for a high degree of control of the latter through excitations of the former. This allows us to predict the sequence of optical pulses to be applied to the material in order to follow closely an enveloping signal function.

And finally, the coupling between ferroelectricity and strain in BaTiO₃ leads to a softening at purely ferroelectric domain walls, allowing for some mechanical control of the position of this wall.

We utilize both theoretical and computational tools to understand the nature of these interactions, how they lead to cross-order coupling in these materials, and how this then translates into the experimentally observed behavior.

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1.1 Spin-orbit coupling

1.2 Wannier Functions

In a lot of the work presented in this Thesis we construct simplified model Hamiltonians to try and describe the physics that are manifested real materials. For topics involving the electronic structure, either directly in describing the Rashba-splitting of energy bands in Chapter ??, or indirectly to parametrize the magnetic exchanges between localized spins as in Chapter ??, tight-binding models are often the most appropriate tool. These are constructed by defining a localized, often atomic-like, set of orbitals which is repeated inside each unit cell, and the so-called hopping terms between them. The first benefit of this simplification is that it makes the numerics of the problem more tractable, since a limited set of orbitals leads to small dimensions of the matrices that represent various operators. Secondly, the representation in terms of real-space localized wavefunctions, as compared with the extended Bloch functions (BFs) that diagonalize the Hamiltonian, often also leads to a more intuitive picture. There are two main methods that are used to construct the tightbinding Hamiltonian. The first is to use a semi-empirical approach, where first a set of localized orbitals is chosen, in terms of which the elements of the Hamiltonian can be written down as a combination of certain model parameters, the amount of which is reduced by using symmetry arguments. In the case of real materials one then uses experimental measurements to fit these model parameters.

The second is to extract these orbitals, and parameters from a first-principles based simulation such as density functional theory. In thesis we favor the latter since it is more flexible, not depending on experiments while still providing quantitative results. This also allows for an easy comparison of multiple materials in order to better understand how certain physics manifest themselves in different cases.

The main question to answer then becomes how to define a such a set of localized orbitals, when most simulations for periodic systems handle extended BFs $|\psi_{\mathbf{k}}(\mathbf{r})\rangle = e^{i\mathbf{k}\cdot\mathbf{r}}u_{\mathbf{k}}(\mathbf{r})$, where \mathbf{k} denotes a wavevector in the first Brillouin zone (BZ), $e^{i\mathbf{k}\cdot\mathbf{r}}$ the plane wave envelope function, and $u_{\mathbf{k}}(\mathbf{r})$ the cell-periodic part (see the left panel of Fig. 1.1(a)). There are many ways to do this, but the most natural one in this case is that of the Wannier functions (WFs) [Wannier1937].

We will here give a short recap of the excellent review done by Marzari et. al.

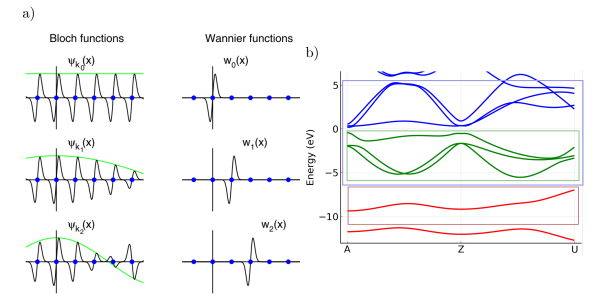


Figure 1.1: a) A comparison between Bloch functions (left) and Wannier functions (right). The green line in the left panel denotes the envelope function $e^{i\mathbf{k}\cdot\mathbf{r}}$. b) Bandstructure of semiconductor GeTe, demonstrating three cases for Wannierization: red is a single disconnected band, green shows a disconnected composite manifold of connected bands, blue shows the situation when bands need to be disentangled.

[Marzari2012], starting with the most clear case of a single isolated band n with BFs $|\psi_{n\mathbf{k}}(\mathbf{r})\rangle$ (see red window in Fig. 1.1(b)), where the BF can be written as a straightforward discrete Fourier transform of the WF:

$$|\psi_{n\mathbf{k}}(\mathbf{r})\rangle = e^{i\mathbf{k}\cdot\mathbf{r}} |u_{n\mathbf{k}}(\mathbf{r})\rangle = \sum_{\mathbf{R}} e^{i\mathbf{k}\cdot\mathbf{R}} |w_{n\mathbf{R}}(\mathbf{r})\rangle,$$
 (1.1)

with $|u_{n\mathbf{k}}(\mathbf{r})\rangle$ the periodic part of the BF, $|w_{n\mathbf{R}}(\mathbf{r})\rangle$ the localized WF centered in the unit cell defined by lattice vector \mathbf{R} , and n the band index. The comparison between BFs and WFs is made in Fig. 1.1. When k=0 we can see that the periodic part of the BF $u_{n\mathbf{k}}$ is simply the sum over the WFs centered at different unit cells. In the case of $k \neq 0$, similar to how the envelope part modulates the periodic part of the BF, the contribution of each WF to the total sum needs to be modulated in a discrete way through $e^{i\mathbf{k}\cdot\mathbf{R}}$.

The inverse Fourier transform over the BF in the first Brillouin Zone (BZ) can then be performed in order to generate the localized WFs,

$$|w_{n\mathbf{R}}(\mathbf{r})\rangle = \frac{V}{(2\pi)^3} \int_{BZ} d\mathbf{k} e^{-i\mathbf{k}\cdot\mathbf{R}} |\psi_{n\mathbf{k}}(\mathbf{r})\rangle,$$
 (1.2)

where V denotes the real-space volume of the unit cell. In these and following equations, the normalization convention is used such that $\int_V d\mathbf{r} \langle \psi_{n\mathbf{k}}(\mathbf{r}) | \psi_{n\mathbf{k}}(\mathbf{r}) \rangle = 1$. As can be seen from the right panel in Fig. 1.1, WFs centered in different unit cells are shifted copies of one another.

One of the most useful properties of the WFs lies in the realization that, through the gauge freedom at each k of the BF¹, the shape of the WFs is not unique:

$$|\psi_{n\mathbf{k}}(\mathbf{r})\rangle \Rightarrow \left|\tilde{\psi}_{n\mathbf{k}}(\mathbf{r})\right\rangle = e^{i\phi_n(\mathbf{k})}\left|\psi_{n\mathbf{k}}(\mathbf{r})\right\rangle$$
 (1.3)

$$|w_{n\mathbf{R}}(\mathbf{r})\rangle \Rightarrow |\tilde{w}_{n\mathbf{R}}(\mathbf{r})\rangle = \frac{V}{(2\pi)^3} \int_{BZ} d\mathbf{k} e^{-i(\mathbf{k}\cdot\mathbf{R} + \phi_n(\mathbf{k}))} |\psi_{n\mathbf{k}}(\mathbf{r})\rangle,$$
 (1.4)

This means that by varying $\phi_n(\mathbf{k})$ we can adapt the WF basis to the needs of the particular problem under investigation. In the case of a single isolated band this is a bonafide luxury exploitable to simplify the task at hand. However, in pretty much all practical problems, it are groups of bands that we are interested in. The simplest case is when these form a composite manifold that is detached from other bands, such as the green valence bands in Fig. 1.1(b).

It is natural to generalize the notion of the single band gauge transformation (in that case amounting to a single free phase), to this manifold of states:

$$\left| \tilde{\psi}_{m\mathbf{k}}(\mathbf{r}) \right\rangle = \sum_{m} U_{\mathbf{k}}^{nm} \left| \psi_{n\mathbf{k}}(\mathbf{r}) \right\rangle.$$
 (1.5)

Here n, m are band indices enumerating the bands inside the composite manifold. These transformed states are no longer eigenstates of the Hamiltonian if $U_{\mathbf{k}}^{nm}$ has offdiagonal elements. Traces performed over such an isolated set of bands are invariant w.r.t. to these gauge transformations, leading to the invariance of observables such as the Hamiltonian which lies at the core of the ability to isolate and focus on this subproblem. Choosing right gauge is a necessity in order to extract well localized WFs, however, because composite band manifolds, in general, will harbor crossings and degeneracies where the BFs become non-analytic and thus the variation of the periodic parts u_{nk} will not be smooth in the variation over k. This is an issue, since it is a well known fact that only smooth functions in reciprocal space lead to well localized ones in real space when the inverse Fourier transform is performed. Indeed, in Eq. 1.2, only then will there be a cancellation of the terms from the fast varying exponent $e^{i\mathbf{k}\cdot\mathbf{R}}$ when \mathbf{R} becomes large. It is thus important to choose a gauge at each k-point such that the rotated wavefunctions $|\tilde{\psi}_{nk}(r)\rangle$ vary as smoothly as possible, which will ultimately lead to the WFs that are as localized as possible². The localization functional can be written down as

$$\Omega = \sum_{n} \langle w_{n\mathbf{0}} | r^2 | w_{n\mathbf{0}} \rangle - |\langle w_{n\mathbf{0}} | \mathbf{r} | w_{n\mathbf{0}} \rangle|^2.$$
 (1.6)

¹The solution to Schrödinger equation does not determine uniquely its phase.

²It can be proved that these WFs will be exponentially localized in the case of normal insulators.

This can be split up into a gauge-invariant and gauge-dependent part

$$\Omega = \Omega_I + \tilde{\Omega},\tag{1.7}$$

$$\Omega_{I} = \sum_{n} \left[\langle w_{n\mathbf{0}} | r^{2} | w_{n\mathbf{0}} \rangle - \sum_{m,\mathbf{R}} \left| \langle w_{m\mathbf{R}} | \mathbf{r} | w_{n\mathbf{0}} \rangle \right|^{2} \right]$$
(1.8)

$$\tilde{\Omega} = \sum_{n} \sum_{m, \mathbf{R} \neq n, \mathbf{0}} \left| \left\langle w_{m\mathbf{R}} \right| \mathbf{r} \left| w_{n\mathbf{0}} \right\rangle \right|^{2}. \tag{1.9}$$

For a set of isolated bands it thus suffices to minimize the gauge-dependent part in order to find the unique set of maximally localized Wannier functions (MLFW) that consitute these bands [Kunes2004, Marzari2012]. We can write down the position dependent operators in reciprocal space as [Blount1962]

$$\langle w_{n\mathbf{R}} | \mathbf{r} | w_{m\mathbf{0}} \rangle = i \frac{V}{(2\pi)^3} \int d\mathbf{k} e^{i\mathbf{k}\cdot\mathbf{R}} \langle u_{n\mathbf{k}} | \mathbf{\nabla}_{\mathbf{k}} | u_{m\mathbf{k}} \rangle$$
 (1.10)

and

$$\langle w_{n\mathbf{R}} | r^2 | w_{m\mathbf{0}} \rangle = -\frac{V}{(2\pi)^3} \int d\mathbf{k} e^{i\mathbf{k}\cdot\mathbf{R}} \langle u_{n\mathbf{k}} | \mathbf{\nabla}_{\mathbf{k}}^2 | u_{m\mathbf{k}} \rangle.$$
 (1.11)

By then minimizing $\tilde{\Omega}$ in terms of different gauge transformations $U_{\mathbf{k}}^{nm}$ applied to the $|u_{n\mathbf{k}}\rangle$, we can find a set of MLFW.

If maximal localization is not the goal, tweaking the U_k^{nm} allows for the generation of WFs with other desirable characteristics in a way that, e.g., they obey certain ionic-site symmetries. This can be achieved by projecting onto atomic-like orbitals, which can be very useful in gaining a further understanding in terms of orbitals with definite and known properties. More details on the different construction methods and can be found in Ref. [Marzari2012].

Up to now, only isolated sets of bands were discussed. However, in many cases such a set does not exist in the region of interest. This leads to sets of band which are termed *entangled*, being connected to bands that lie outside this energy range (see the blue bands in Fig. 1.1(b)). If we are seeking J WFs, we thus need to somehow select at each k-point J states $\left|\tilde{\psi}_{nk}\right\rangle$ from a bigger set J_k , that can then be used in the above described localization procedure:

$$\left|\tilde{\psi}_{n\mathbf{k}}\right\rangle = \sum_{m=1}^{J_{\mathbf{k}}} V_{\mathbf{k}}^{nm} \left|\psi_{m\mathbf{k}}\right\rangle. \tag{1.12}$$

 $V_{\mathbf{k}}^{nm}$ are rectangular matrices of dimension $J \times J_{\mathbf{k}}$. This leads to a two-step process where first a subspace is selected for each k-point, followed by the final gauge selection to arrive at the final localized WFs.

$$|w_{n\mathbf{R}}(\mathbf{r})\rangle = \frac{V}{(2\pi)^3} \int_{BZ} d\mathbf{k} e^{-i\mathbf{k}\cdot\mathbf{R}} \sum_{m=1}^{J} U_{\mathbf{k}}^{nm} \sum_{l=1}^{J_{\mathbf{k}}} V_{\mathbf{k}}^{ml} |\psi_{l\mathbf{k}}(\mathbf{r})\rangle$$
(1.13)

There are again many ways to do this subspace selection (or *disentanglement*). The most prevalent two are to use another projection based method, or one that again

focuses on minimizing the real space spread of the WFs. We utilize the latter in this work. It is based on finding the V_k^{nm} that lead to the maximal overlap of the selected subspaces with their neighbors at each k-point. The matrices will be generally rectangular in this case, since they transform a set of J_k wavefunctions into a smaller one with only J wavefunctions. As discussed before, this smoothness in reciprocal space translates into the subspace as a whole being more localized in real space. Indeed, it can be shown [Marzari2012] that this intrinsic smoothness is exactly what Ω_I measures in Eq. 1.8. Following this disentanglement procedure, we can apply the procedures described for the isolated manifold of bands to ultimately lead to the WFs which are very well localized. In many cases, it is also desirable to adapt the algorithm in such a way that the BFs inside a "frozen" energy window are exactly interpolated by the resulting WF base and tight-binding Hamiltonian.

In the previous equations a continuous integration over the BZ was performed for the fourier transform from BFs to WFs, however, in reality a discrete k-grid and Fourier transform is used. To keep close to the continuous case, the Fourier transform pair is defined as

$$\left|\tilde{\psi}_{n\mathbf{k}}(\mathbf{r})\right\rangle = \sum_{\mathbf{R}} e^{i\mathbf{k}\cdot\mathbf{R}} \left|w_{n\mathbf{R}}(\mathbf{r})\right\rangle,$$
 (1.14)

$$|w_{n\mathbf{R}}(\mathbf{r})\rangle = \frac{1}{N} \sum_{\mathbf{k}} e^{-i\mathbf{k}\cdot\mathbf{R}} \left| \tilde{\psi}_{n\mathbf{k}}(\mathbf{r}) \right\rangle,$$
 (1.15)

where N denotes the number of unit cells in the periodic supercell in real space, or the number of k-points in the discrete mesh over the BZ. This discretization enforces periodic boundary conditions on the BFs over this supercell, meaning that the WFs in this definition also have this supercell periodicity. The localization then means that inside the supercell the WFs are localized. If the interpolation is continuous, the supercell is infinite, restoring the earlier definitions and original notion of WFs.

The final process to extract a tight-binding Hamiltonian in a WF basis, starting from an ab-initio DFT simulation, can thus be summarized as follows:

- 1. perform a self-consistent DFT calculation in order to find the ground state density and BFs over a discrete k-mesh to be used in the Wannierization
- 2. find which trial orbitals (e.g. atomic-like) are most suitable for the bands or problem under investigation. This can be done for example by perforing a projected density of states calculation.
- 3. select an inner "frozen" window with the most important bands, and, if entangled, and an outer window from which to disentangle J-dimensional subspaces for each k-point
- 4. use a projection on the trial orbitals as the initial guess and then optimize the smoothness of the subspaces as a whole to find V_{k}^{ml} matrices to identify the J-dimensional subspaces.

5. find the final gauges U_{k}^{nm} in order to minimize the spread of each of the J WFs.

Further details on the Wannierization process can be found in Ref. [Marzari2012], and specific details on the implementation in the Wannier90 package used throughout this work can be found in Ref. [Mostofi2014AnFunctions].

When the $U_{\mathbf{k}}$ and $V_{\mathbf{k}}$ matrices are found, any property $f(\mathbf{k})$ defined on the k-mesh in terms of the BFs used in the first-principles calculation can be transformed into $F(\mathbf{R})$ in the Wannier representation. Since the WFs are well localized, $F(\mathbf{R})$ tends to decay rapidly with $|\mathbf{R}|$. It is important to realize that, depending on the coarseness of the k-grid used in the first-principles calculation, $F(\mathbf{R})$ can only be calculated for limited values of $|\mathbf{R}|$, spanning the supercell defined by the discretization of the BZ. Using these short-ranged real-space $F(\mathbf{R})$, it is then possible to interpolate the values of $f(\mathbf{k})$ by performing the inverse procedure. This allows to efficiently calculate $f(\mathbf{k})$ on a finer k-mesh compared with the one used in the first-principles calculations, provided the WFs are well localized inside the supercell defined by the original coarse k-mesh. This efficient interpolation is one of the great advantages of using this technique, since the basis, and thus dimensions of the matrices, is much lower than that used in the first-principles calculation (e.g. many plane waves).

The main quantity we will use this interpolation for is the Hamiltonian:

$$\tilde{H}_{\mathbf{k}}^{nm} = \left\langle \tilde{\psi}_{n\mathbf{k}} \middle| H \middle| \tilde{\psi}_{n\mathbf{k}} \right\rangle = \sum_{\mathbf{R}} e^{i\mathbf{k}\cdot\mathbf{R}} \left\langle w_{n\mathbf{0}} \middle| H \middle| w_{n\mathbf{R}} \right\rangle, \tag{1.16}$$

where the tilde is used to distinguish the k-space wavefunctions from BFs, and $|w_{nR}\rangle$ denote the WFs centered at unit cell identified by lattice translation R. To recover the bandstructure from this Hamiltonian can then be diagonalized by unitary transformations W_k such that

$$H_{\mathbf{k}}^{nm} = [W_{\mathbf{k}}^{\dagger} \tilde{H}_{\mathbf{k}}^{nm} W_{\mathbf{k}}]^{nm} = \delta_{nm} \varepsilon_{n\mathbf{k}}. \tag{1.17}$$

This concludes this very condensed introduction and overview of the use and construction of Wannier functions as a tool to aid in the theoretical understanding of the behavior of electrons in extended systems.