Unfolding first-principles band structures

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A general method is presented to unfold band structures of first-principles super-cell calculations with proper spectral weight, allowing easier visualization of the electronic structure and the degree of broken translational symmetry. The resulting unfolded band structures contain additional rich information from the Kohn-Sham orbitals, and absorb the structure factor that makes them ideal for a direct comparison with angular resolved photoemission spectroscopy experiments. With negligible computational expense via the use of Wannier functions, this simple method has great practical value in the studies of a wide range of materials containing impurities, vacancies, lattice distortions, or spontaneous long-range orders.

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The electronic band structure is no doubt one of the most widely applied analysis tools in the first-principles electronic structure calculations of crystals, especially within the Kohn-Sham framework [1] of density functional theory [2]. It contains the basic ingredients to almost all the textbook descriptions of crystal properties (e.g. transport, optical and magnetic properties, and the semiclassical treatment [3]). Furthermore, the theoretical band structure, when formulated within the quasiparticle picture of the one-particle Green function, has a direct experimental connection with angular-resolved photoemission spectroscopy (ARPES).

However, the usefulness of the band structure, as well as the agreement with ARPES spectra, diminishes rapidly when a large “super cell” is involved. The use of super cells is a common practice in modern first-principles studies when the original periodicity of the system is modified via the introduction of “external” influences from impurities or lattice distortions. They are also widely applied in the presence of spontaneous translational symmetry breaking, say by a charge density wave, a spin density wave, or an orbital ordering. As illustrated in Fig. 1 when the period of the super cell grows longer, the corresponding first Brillouin zone of the super cell (SBZ) shrinks its size. In turn, bands in the first Brillouin zone of the normal cell (NBZ) get “folded” into the SBZ. For a very large super cell, the resulting SBZ can be tiny in size but contain a large number of “horizontal” looking bands that no longer resemble the original band structure or the experimental ARPES spectra, and cease to be informative besides giving a rough visualization of the density of states (DOS). The information is now hidden in the Kohn-Sham orbitals, instead of the dispersion of the bands.

In this Letter, by explicitely utilizing these Kohn-Sham orbitals, we present a method to unfold the band structure of the SBZ back to the larger NBZ with proper spectral weight. Making use of the corresponding Wannier functions, the method can be greatly simplified to negligible computational cost. The resulting unfolded band structure incorporates explicitly the structure factor and thus facilitates significantly a direct comparison with ARPES experiments. Furthermore, the unfolded band structure illustrates very clearly the influence of the symmetry breaker (e.g.: impurities, vacancies, dopants, lattice distortions) via direct comparison with the nominal normal-cell band structure. In the case of spontaneous symmetry breaking, it gives a direct visualization of the strength of each band’s coupling to the order parameters. In light of the amazingly rich information, we expect countless applications of this simple method to a wide range of studies employing super cells, including systems with charge density wave, spin density wave, or orbital ordering, and in the studies of impurities and lattice distortions, to name a few.

Theoretically, the folding of the bands results from the introduction of additional coupling, $V_{k,j,k',j'}$, between the
originally uncoupled Kohn-Sham orbitals $|kj⟩$ and $|krtj⟩$ in the NBZ. (Here $k$ and $j$ denote the crystal momentum and the band index.) This coupling extends the period of Kohn-Sham orbitals to a longer one compatible with the size of the super cell. Equivalently, this coupling, no matter how small it is, mixes the original orbitals of different normal-cell crystal momentum $k$ and forces us to label them with a supercell crystal momentum $K$ as the new quantum number in the SBZ. (In the following, upper-/lower-case symbols refer to variables corresponding to the super/norm cell, respectively.) Our method is based on the simple idea that unless $V$ is extremely strong, it is much more convenient and informative to represent the band structure or more precisely the spectral function $A = −\text{Im}G/\pi$ of the retarded one-particle Green function, $G$, not in the new eigen-orbital $|KJ⟩$ basis, but in the $|kj⟩$ basis of the normal cell instead:

$$G_{kj,k′j′}(ω) = G_{0kj,k′j′}(ω)δ_{k,k′}δ_{j,j′} - V_{kj,k′j′},$$

where $G_0$ represents a conceptual system with the period of the normal cell before $V$ is applied. Clearly, $G$ smoothly recovers the original period of $G_0$ as $V$ approaches zero. Thus,

$$A_{kj,k′j′}(ω) = \sum_{KJ} |⟨kj|KJ⟩|^2 A_{KJ,KJ}(ω)$$

(2)

should resemble the band structure of the normal cell with deviations in both the dispersion and in the spectral weight that reflect the effects of $V$. Note that while the coupling $V$ introduces non-diagonal elements of $A_{kj,k′j′}(ω)$, we focus only on the diagonal elements here for simplicity, without loss of generality. It is straightforward to show that in the case of $V = 0$, the weight of the bands that follow the bands of the normal cell is exactly one, and that of the rest of the folded bands vanishes. One thus recovers exactly the original band structure of the normal cell as expected. That is, the unfolded band structure is invariant against any arbitrary choice of super cell.

In addition, it is often desirable to also measure in each band the contribution of local orbitals with well-defined characters (e.g.: $p_x$, $d_{xy}$, $e_g$, or bonding/antibonding). This can be achieved rigorously via the use of local Wannier orbitals $|rn⟩$: 

$$A_{kn,kn}(ω) = \sum_{KJ} |⟨kn|KJ⟩|^2 A_{KJ,KJ}(ω),$$

(3)

where $|kn⟩ = \sum_r |rn⟩⟨rn|kn⟩ = \sum_r |rn⟩e^{ikr}/\sqrt{l}$ are the Fourier transform of the Wannier orbitals $|rn⟩$ of orbital index $n$ and associated with the lattice vector $r$. (Here $l$ denotes the number of $k$-points in the NBZ.) Given a consistent definition of the Wannier functions of the super-cell calculation that maps $|RN⟩$ of the super cell to $|R+r, n⟩$ of the normal cell, where $r = r(N)$ is a normal-cell lattice vector within the first super cell, and $nt = nτ(N)$ is the corresponding normal-cell orbital index, the use of Wannier function also reduces dramatically the computational expense by turning the factor

$$⟨kn|KJ⟩ = \sum_{RN} ⟨kn|RN⟩⟨RN|KN⟩⟨KN|KJ⟩$$

$$= \sum_{RN} ⟨kn|R + r(N), nτ(N)⟩⟨RN|KN⟩⟨KN|KJ⟩$$

$$= \sqrt{l/l'} \sum_{RN} e^{i(K−k)·R} e^{−ik·r(N)} δ_{n,nτ(N)}⟨KN|KJ⟩$$

$$= \sqrt{l/l'} \sum_{N} e^{−ik·r(N)} δ_{n,nτ(N)}δ_{|k⟩,K}⟨KN|KJ⟩$$

(4)

into merely a structure factor that is a sum of coefficients of the eigen-orbital $|KJ⟩$ of the super cell in the Wannier function basis, modulated by the proper phase that encapsulates the internal position in the super cell. Here $|k⟩$ denotes the $k$-point folded into the SBZ from $k$. Since $A_{KJ,KJ}$ is just a delta function at the eigenvalue $δ(ω − ϵ_{kJ})$, this final expression in essence requires only a simple coding to plot all the eigenvalues of the super cell in the larger NBZ with a proper weight.

Of course, the above definition only makes sense when the Wannier functions $|RN⟩ ↔ |rn⟩$ and $|RNn⟩ ↔ |rnn⟩$ (that are translational symmetric in the normal cell unit: same $n$ different $r$) are approximately identical. Therefore, the “gauge” [4] of constructing $|RN⟩$ and $|RNn⟩$ with the same $n$ must be controlled accordingly. In the presence of a potential that breaks the translational symmetry of the normal cell, for example, coming from a CDW, lattice distortions, impurities, etc., the commonly employed [6–10] maximally localized Wannier function [4] and other minimization-based methods [8, 9] risk defining the gauge differently in the super cell in favor of better localization, and thus should be used with extreme caution. We found that a maximum projection method [10, 12] with consistent projection between the normal and the super cells works well to satisfy this requirement. Equations (3) and (4) should in principle also be applicable in many existing codes employing atomic center local orbitals as basis [13, 14], as long as the non-orthogonal nature of those bases is taken into account. Of course, these methods do not benefit from the energy resolution of the Wannier functions that allows unfolding only the bands within the physically relevant energy range.

The unfolded band structure also has an important direct connection to the ARPES measurement. For systems with enlarged unit cells due to weak symmetry breaking, the ARPES spectra typically shows different band structures in different Brillouin zones of the super cell, distinctly different from the results of first-principles calculations, which have all the bands in the SBZ. In some cases, the observed ARPES spectra might even appear ignorant about the SBZ [15]. This significant mismatch is typically regarded as the effect of the “matrix element” and left unaddressed by both theorists and ex-
perimentalists, making a direct comparison very difficult. Within the “sudden approximation”, the ARPES intensity is proportional to \[16, 17\]

\[
\sum_{KJ} \left| |e| \cdot (f |p| KJ) \right|^2 A_{KJ,KJ}(\omega) \\sim \sum_{KJkn} \left| |e| \cdot \langle f |p| kn \rangle \right|^2 \langle kn |KJ \rangle |^2 A_{KJ,KJ}(\omega)
\]

\[= \sum_{kn} \left| |e| \cdot (f |p| kn) \right|^2 A_{kn,kn}(\omega), \tag{5}\]

where \(e\) denotes the polarization vector of light, and \(|f\) the “final state” of the photoelectron. Clearly, except the polarization dependent dipole matrix element, \(|e| \cdot (f |p| kn)\), the unfolded spectral function, \(A_{kn,kn}(\omega)\), contains almost the full information of the experimental spectrum by absorbing the additional structure factor \(\langle kn |KJ \rangle\), absent in the typical super-cell solution, \(A_{KJ,KJ}(\omega)\). Obviously, the inclusion of this additional matrix element would facilitate significantly the comparison between the theory and the ARPES experiment.

As an example, let’s consider the effect of Na impurities in Na-doped cobaltates, Na\(_x\)CoO\(_2\) at \(x = 1/3\). In typical first-principles studies\[18, 19\], the impurity is incorporated via a super cell as demonstrated in Fig. 2(b) in comparison with the undoped normal cell shown in Fig. 2(a). Fig. 2(d) and (c) show the corresponding band structures obtained with standard DFT calculations. Since in this example the super cell is three times larger than the normal cell, the corresponding SBZ is three times smaller and contains three times more bands. Clearly, even for such a small super cell, the change of the size/orientation of the SBZ and more importantly the large number of folded bands, make it practically impossible to cleanly compare with the band structure in the NBZ of the undoped parent compound. In fact, to many untrained eyes, these two band structures may appear entirely unrelated.

By contrast, the unfolded band structure shown in Fig. 2(e), demonstrates a strong resemblance to the band structure of the undoped compound. This allows a clear visualization of the effects of the (periodic) Na impurities on the original Co and O bands. Specifically, besides the introduction of additional Na-s bands, one observes shifts in band energies, gap openings and the nearby “shadow bands”, all of which reflects the influence of the Na impurity on these bands. What is really nice here is the cleanness of the unfolded band structure in general, owing to the weak intensity of the shadow bands. As expected, the influence of the Na impurity is only minor on most Co-d and O-p bands, while the Na-s bands themselves show sizable effects of broken translational symmetry. The size of the gap opening and the intensity of the shadow bands actually reflect directly the strength of each band’s coupling to the broken translational symmetry of the normal cell (in this specific case, to the charge-density-wave order parameter introduced by the periodic presence of Na atoms.) Of course, for a simulation of randomly positioned impurities, these CDW-related features are entirely artificial, and the unfolded band structure makes apparent the alarming limitation of common practice of using small super cells in the study of impurities. On the other hand, in many other cases, for example the super modulation of the lattice, these features would actually correspond to a physical order parameter and provide valuable information.

As another example, let’s consider a spontaneous orbital ordering in A-type anti-ferromagnetic LaMnO\(_3\). Figure 3 (c) and (d) show the similar comparison of band structures without and with the long-range staggered orbital order, corresponding to unit cells shown in Fig. 3 (a) (normal cell) and Fig. 3 (b) (super cell), respectively. Both results are obtained via the LSDA+\(U\) (\(U=8eV\), \(J_H=0.88eV\)) approximation without lattice relaxation for simplicity, without loss of generality. By comparing the band structures with and without the orbital order on the equal footing, the detailed information of the spontaneous orbital order should be visualized explicitly.

Just like in the Na\(_x\)CoO\(_2\) case, the straightforward results of the orbital ordered (OO) band structure (Fig. 3 (d)) of the super cell calculation \[20\] can hardly be compared with the non-OO one (Fig. 3 (c)). By contrast,
the unfolded band structure (Fig. 3 (e)) of the OO case resembles strongly the non-OO case. In fact, one finds that only those bands of Mn-$e_g$ character (red, blue, and green) show strong coupling to the OO order parameter with large gap openings and intensive shadow bands, while the rest of the bands are basically uncoupled to the orbital order. In addition, from the significant energy gain corresponding to the large OO gap ($\Delta_{OO}$) near the Fermi level of the red and blue bands, it is apparent that essentially the orbital order is driven only by the spin-majority $e_g$-orbitals (red and blue). All these effects are of course entirely consistent with the existing “electronic interaction assisted Jahn-Teller picture” [21–23], in which the degenerate Mn-$e_g$ orbitals split to gain energy and stabilize the system at low-temperature. However, this unfolded band structure represents probably the best visualization of such physics in real materials with details of first-principles calculations.

In conclusion, a simple method for unfolding first-principles band structures of super cell calculations is presented. Proper spectral weights are obtained with negligible computational cost by making use of the Kohn-Sham orbitals with the help of carefully chosen Wannier functions. The inclusion of the structure factor in the resulting unfolded band structure makes it ideal for direct comparison with the ARPES measurement. The resulting unfolded band structures allow an easy visualization of each band’s coupling to the order parameter of spontaneous broken translational symmetry, as well as their couplings to the external symmetry breakers like the impurities and lattice distortions. Our method should prove valuable in the study of a wide range of problems requiring the use of super cells, including systems with impurities, vacancies, and lattice distortions, and broken symmetry phases of strongly correlated materials, to name a few.

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