Hund bands in spectra of multiorbital systems

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Spectroscopy experiments are routinely used to characterize the behavior of strongly correlated systems. An in-depth understanding of the different spectral features is thus essential. Here, we show that the spectrum of the multiorbital Hubbard model exhibits unique Hund bands that occur at energies given only by the Hund coupling $J_H$, as distinct from the Hubbard satellites following the interaction $U$. We focus on experimentally relevant single-particle and optical spectra that we calculate for a model related to iron chalcogenide ladders. The calculations are performed via the density-matrix renormalization group and Lanczos methods. The generality of the implications is verified by considering a generic multiorbital model within dynamical mean-field theory.

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Introduction. Strongly correlated systems are at the heart of modern condensed matter physics. The celebrated single-band Hubbard model, describing (doped) Mott insulators, is still extensively studied in the context of Cu-based high-temperature superconductivity [1–3]. An equally exciting case is that of iron-based superconductors where the presence of several active orbitals leads to novel effects beyond the “standard” Mott physics [4–6]. A nontrivial example is the orbital-selective Mott phase (OSMP) [5,7–10], where Mott-localized and itinerant electrons coexist.

A key probe of electronic excitations is the single-particle spectral function $A(k, \omega)$, characterizing the excitations’ dispersion. It is experimentally accessible by angle-resolved photoemission spectroscopy (ARPES) [11,12]. To understand the origin of different spectral features, it is convenient to consider idealized models that can be studied theoretically and monitor how the signatures of correlations (e.g., the Hubbard bands) evolve with increasing Coulomb interaction $U$. This is especially true for quantum systems of reduced dimensionality, for which quasiexact numerical methods [13,14], or even closed analytical solutions [15], provide unbiased information on the elementary excitations. However, even in reduced dimensionality obtaining accurate results for the multiorbital Hubbard model remains challenging. The difficulty lies in the exceptionally large Hilbert space. Because of that, the spectral functions are often calculated using the dynamical mean-field theory (DMFT) [16–19]. This approach, which strictly applies at large dimensionality, avoids the finite-size limitation, but often relies on solvers in Matsubara frequencies and hence the resulting spectral functions are blurred due to analytical continuation (see Ref. [20], which discusses this and introduces a method to alleviate the problem).

In this Letter, we numerically investigate the spectral functions of several multiorbital models. Our main result is summarized in Fig. 1(a). The electronic spectrum of a single-orbital model (without the Hund coupling $J_H \rightarrow 0$) consists of the usual upper and lower Hubbard bands (UHB and LHB, respectively) that develop with $U$. In multiorbital systems, the finite $J_H$ gives rise to additional excitations. Some of these states can appear at energies between UHB and LHB that depend exclusively on $J_H$ (i.e., are independent of $U$), paving the way to measure $J_H$ directly. Since such excitations occur due to the Hund coupling and have a robust dispersion [see Figs. 1(b) and 1(c) and Ref. [21] for the full spectrum of $A(k, \omega)$], we call them Hund bands. We recognize that the Hund bands arise whenever single-particle removal/addition processes yield a higher multiplet of the dominant valence subspace. This can occur provided (i) the higher multiplets exist, (ii) these multiplets are allowed by the selection rules upon adding/removing a particle, and (iii) the charge fluctuations are significant. All these requirements are met for Hund’s metals. Earlier work documented multiplet splittings in the Hubbard bands [20,31,32], in the fully occupied orbital [33], found additional “holon-doublon” peaks [34–39], and analyzed the energy-level structure, revealing multiplets that violate the Hund’s rules [40]. Here, we stress that charge excitations independent of $U$ are a generic consequence of the multiorbital systems.

To reach these conclusions, we use the density-matrix renormalization group method (DMRG) [41–46] and Lanczos diagonalization [2,47]. To show that our findings are generic, we study both the two- and three-orbital Hubbard model. Furthermore, we supplement our analysis with the effective model of the OSMP—the generalized Kondo-Heisenberg Hamiltonian. Finally, we confirm our findings with DMFT calculations. Our results apply to many experiments investigating the spectral properties of multiorbital materials, particularly iron-based compounds [48,49], ruthenates [33,50–52], iridates [53,54], and nickel oxides [55–59].
Model. We focus on the SU(2)-symmetric multiorbital Hubbard-Kanamori chain,

\[ H_H = -\sum_{\gamma \neq \ell} t_{\gamma \ell} c_{\gamma \ell \sigma}^\dagger c_{\gamma \ell + 1 \sigma} + H.c. + \sum_{\gamma \ell} \Delta_\gamma n_{\gamma \ell} \]

\[ + U \sum_{\gamma \ell} n_{\gamma \ell} n_{\gamma \ell} + (U - 5J_H/2) \sum_{\gamma < \ell} n_{\gamma \ell} n_{\gamma \ell} \]

\[ - 2J_H \sum_{\gamma < \ell} S_{\gamma \ell} \cdot S_{\gamma \ell} + J_0 \sum_{\gamma < \ell} [P_{\gamma \ell}^\dagger P_{\gamma \ell} + H.c.]. \]  

(1)

Here, \( c_{\gamma \ell \sigma}^\dagger \) creates an electron with spin \( \sigma \) at orbital \( \gamma \) of site \( \ell \), \( t_{\gamma \ell} \) is the symmetric hopping matrix in orbital space, \( \Delta_\gamma \) denotes the crystal-field splitting. \( n_{\gamma \ell} = \sum_{\sigma} n_{\gamma \ell \sigma} \) represents the total density of electrons. \( U \) is the standard repulsive Hubbard interaction. \( J_H \) is the Hund coupling between spins \( S_{\gamma \ell} \) at different orbitals \( \gamma \). The last term \( P_{\gamma \ell}^\dagger P_{\gamma \ell} \) denotes interorbital pair hopping, \( P_{\gamma \ell} = c_{\gamma \ell \uparrow} c_{\gamma \ell \downarrow} \). We assume open boundary conditions, as required by DMRG. For the two-orbital model, \( \gamma \in \{1, 2\} \), we used (in eV) \( t_{11} = -0.5, t_{12} = -0.15, t_{21} = 0, \Delta_1 = 0, \Delta_2 = 0.8 \); whereas for the three-orbital model: \( \gamma \in \{0, 1, 2\}, t_{00} = t_{11} = -0.5, t_{02} = -0.15, t_{02} = t_{12} = 0.1, t_{01} = 0, \Delta_0 = -0.1, \Delta_1 = 0, \Delta_2 = 0.8 \). These values were previously used to study the iron-based ladders of the 123 family [9, 10, 60–63]. The bandwidth of the two-orbital model, \( W = 2.1 \), is used as the energy unit [64]. All energy labels given throughout the text are independent of the \( J_H/U \) ratio.

We also study the minimal model of the OSMP: The generalized Kondo-Heisenberg model (gKH). This model was derived [10, 62, 63] to capture the static and dynamic properties of BaFe2Se3 iron-based ladder [65–67]. It describes interacting itinerant electrons (with spin \( s_i \)) coupled via Hund coupling to the localized spins \( S_i \).

\[ H_K = -t_i \sum_{\sigma} (c_{i \sigma}^\dagger c_{i+1 \sigma} + H.c.) + U \sum_{\ell} n_{\ell \uparrow} n_{\ell \downarrow} \]

\[ + K \sum_{\ell} S_{\ell \uparrow} \cdot S_{\ell \downarrow} - 2J_H \sum_{\ell} S_{\ell \uparrow} \cdot S_{\ell \uparrow}. \]  

(2)

For the gKH model, \( t_i = -0.5, K = 4t_i^2/U, \ t_i = -0.15 \), matching the OSMP of our two-orbital Hubbard model [10].

Huband bands. Let us study the orbital-resolved single-particle spectral function \( A_{\gamma}(k, \omega) \) and the density of states (DOS) \( A_{\gamma}(\omega) \) for the gKH model; see Fig. 2(b). The atom realizes the noninteger filling \( n = 1.5 \). In the \( U \rightarrow 0 \) limit, we recover the noninteracting behavior: A single metallic band. However, already at \( U/W \approx 0.8 \), i.e., close to the OSMP transition [9, 10], the three-peak structure is visible in \( A_1(\omega) \), and becomes clearer the larger the interaction \( U \) becomes. Since the three-peak structure is most pronounced for \( U \gg W \), it is instructive to examine the atomic limit \( U, J_H \rightarrow \infty \) of the gKH model; see Fig. 2(b). The atom realizes the noninteger filling \( n = 1.5 \). The structure is also visible in the itinerant orbital \( (\gamma = 1) \), Fig. 1(c), with an electron density equal to 1.5. Notice that the itinerant orbital’s spectrum is accurately reproduced by the effective gKH model.

Let us take a closer look at how the three-peak spectrum develops with the interaction \( U \). Figure 2(a) shows \( A_1(\omega) \) for the gKH model at noninteger filling \( n = 1.5 \). In the \( U \rightarrow 0 \) limit, we recover the noninteracting behavior: A single metallic band. However, already at \( U/W \approx 0.8 \), i.e., close to the OSMP transition [9, 10], the three-peak structure is visible in \( A_1(\omega) \), and becomes clearer the larger the interaction \( U \) becomes. Since the three-peak structure is most pronounced for \( U \gg W \), it is instructive to examine the atomic limit \( U, J_H \rightarrow \infty \) of the gKH model; see Fig. 2(b). The atom realizes the noninteger filling \( n = 1.5 \) provided the ground states (gs) of the 1- and 2-electron subspaces are degenerate, which is achieved at \( \mu = U + J_H/2 \). Then, the gs consists of a local interorbital triplet, denoted as \( |T_1 \rangle \), and a 3-electron singlet, denoted as \( |D \rangle \). By
removing an electron from the triplet, one creates a holon in the itinerant orbital (\( |T\rangle \rightarrow |H\rangle \)), with the cost of energy \( U + J_H \). Interestingly, from the doubly occupied state, one can remove an electron in two different ways. Depending on the spin projection of the removed electron, one can arrive at a local triplet or singlet, \(|D\rangle \rightarrow |T\rangle \) or \(|D\rangle \rightarrow |S\rangle \), respectively. The former is a zero-energy transition between degenerate states of the gs, while the latter costs an energy \( 2J_H \) as it breaks the Hund’s rule. In Fig. 2(a), we plot the relevant energy scales of the atomic limit (\( U + J_H \) and \( 2J_H \)) and find good agreement with the full many-body calculations of the gKH chain.

**Projections on the atomic configurations.** To make a stronger case for the atomic-limit interpretation of the three-particle peak spectrum, we decompose the spectral function of the full many-body calculation into individual transitions [38]. To this end, we use the projector \( P \) onto specific configurations of the on-site Ising basis \(|\gamma = 1, 2, \ldots\rangle \), i.e., \( \langle c_{\gamma,L/2,\sigma}\rangle \langle c_{\gamma,L/2,\sigma}\rangle \) \( \mathcal{P}_{\gamma,L/2,\sigma} \) [21]. For clarity, we discuss only the hole part (below \( \mu \)), as the electron part can be described analogously. Upon removing an electron from the itinerant orbital, we distinguish three contributions. (i) In

**Noninteger vs integer filling.** As shown above, for noninteger filling (doped system), the atomic limit is enough to explain the Hund bands. When the atomic gs of adjacent particle-number subspaces, say \( N \) and \( N - 1 \), are degenerate, there is no cost \( U \) for the transition from the gs of subspace \( N \) to the gs of subspace \( N - 1 \). The excitation cost is zero; it is compensated by \( \mu \) which is tuned to cause the degeneracy. However, if the \( N - 1 \) subspace contains not only the gs but also higher multiplets, these multiplets can be accessed in the photoemission process \( N \rightarrow N - 1 \) with just the energy \( \propto J_H \). Analogous reasoning applies to inverse transitions \( N - 1 \rightarrow N \). Thus, remarkably, this results in \( U \)-independent Hund bands.

Consider now this behavior in a more general system, hosting more atomic configurations with different \( n \). In Fig. 3 we present the three-orbital Hubbard model (3oH) results [69] for various electron fillings. For \( n = 4.5 \), the atomic limit of our setup [21] predicts one Hund excitation (between states with 5 and 4 electrons) with energy \( 2J_H \) [70], along with several \( U \)-dependent Hubbard excitations. We pinpoint the Hund band using the projector analysis, shown in Fig. 3(b). We differentiate transitions arriving at \( |↑↑, ↑, ↑\rangle \) and \( |↑↑, ↓, ↓\rangle \). Similarly, for the \( n = 3.5 \) filling, the atomic limit implies Hund bands in photoemission at 3\( J_H \) and 5\( J_H \). They are shown in Fig. 3(c). The 3\( J_H \) band is a transition to a low-spin \( S = 1/2 \) state \( |P \rangle \rightarrow |↑↑, ↓\rangle \); see Fig. 3(a). The 5\( J_H \) band originates in states of the form \( |↑↑, 0, ↑\rangle \) \( \propto |0, ↓, ↑\rangle \), where “-” is degenerate with the 3\( J_H \) excitation while “+” forms the 5\( J_H \) peak. The latter are the holon-doublon states [34–39]. Their origin was discussed in [34,36] but without realizing they are a particular example of the generic physics of Hund bands revealed here. Surprisingly, the 2\( J_H \) band persists even for \( n = 3.5 \) (as implied by the smaller but nonvanishing weight of \( |↑↑, ↓, ↓\rangle \)), inducing a third Hund peak, absent in the atomic spectrum. The intensity of this mode decreases with \( U \).

By contrast, for integer filling \( n \in \{1, 2, 3, \ldots\} \), the atomic limit alone does not predict the Hund bands. The atom lacks the necessary charge fluctuations as its gs does not span adjacent particle-number subspaces. Thus, only the “standard” Hubbard bands should be observed [5,71]. However, in the lattice, the charge fluctuations are possible provided the interaction \( U \) is not too large at a given filling \( n \). For half filling, the fluctuations vanish already for \( U \approx W \) and the Hubbard
bands are well developed [see, e.g., the inset of Fig. 2(a)]. Away from half filling, $U \sim W$ does not suppress the fluctuations. They are significant even at integer $n$, and vanish only at elevated $U \sim 10 W$ [5,61,72]. Consequently, the many-body gs has significant contribution of states with neighboring local occupations, $|n-1|$ and $|n+1|$. Adding/removing particles in these states allows reaching the higher multiplets of the atomic ground-state subspace $|n\rangle$, and the Hund bands emerge.

Consider the $n = 4$ case, i.e., one electron above half filling for 3oH. In the atom, the gs has only 4-electron configurations, but in the lattice we find significant on-site fluctuations to 5- and 3-electron states [4,73]. In Fig. 3(d), we project onto the same configurations as for $n = 4.5$ and again find the $2J_H$ Hund band (originating in the $n = 5 \rightarrow 4$ transitions). We should notice only half of the peak is exhausted by the projection onto $|\uparrow\downarrow, \downarrow, \uparrow\rangle$ and our results also indicate a weak $U$ dependence. We could not discern Hund bands corresponding to electron addition processes from 3-electron states: For a high-spin initial configuration $n = 3$, $S = 3/2$, the selection rules forbid reaching the low-spin $n = 4$, $S = 0$ state [74].

Conclusions. We showed that the charge fluctuations and finite Hund exchange present in the multiorbital Hubbard model cause the formation of unique bands of excitations. These Hund bands are formed by the energetically costly low-angular-momentum states (i.e., on-site configurations which break the Hund’s rules) and they do not depend on Hubbard $U$. The latter makes them distinct from the Hubbard-band multiplet splittings. Among the Hund bands the canonical spin-singlet mode ($\omega \simeq 2J_H$) is especially prevalent.

Our results are a generic consequence of multiorbital systems. They originate in the existence of higher multiplets; hence they do not depend on the presence of the orbital-selective Mott phase (see [21] for additional discussion) or on the system’s dimensionality. To confirm this, in Fig. 4(a), we present DMFT calculations in infinite dimensions. We focus on generic rather than material-specific features and consider a semicircular density of states and orbital degeneracy [21]. The DMFT results clearly show the $2J_H$ mode. In Supplemental Material [21], we repeat the calculations for a typical $t_{2g}$ DOS and also find the Hund band.

Our findings are relevant for ARPES, resonant inelastic x-ray scattering [75], Raman spectroscopy [76,77], nonequilibrium investigations [78–80], and reflectivity/transmission measurements [81–84]. Figure 4(b) demonstrates the last: It presents how the optical conductivity [21] evolves with $U$ for the gKH model at $n = 1.5$. Crucially, we observe the Hund band at $\omega \simeq 2J_H$. Often, such additional spectral features are attributed to the interband transitions. Here, we showed that additional modes can also originate in the Hund exchange and, consequently, can be used to estimate the value of $J_H$.

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