Correlated band theory implemented as a combination of density functional theory with exact diagonalization [DFT+U(ED)] of the Anderson impurity term with Coulomb repulsion $U$ in the open 14-orbital $5f$ shell is applied to UTe$_2$. The small gap for $U=0$, evidence of the half-filled $j = \frac{5}{2}$ subshell of $5f^3$ uranium, is converted for $U=3$ eV to a flat band semimetal with small heavy-carrier Fermi surfaces that will make properties sensitive to pressure, magnetic field, and off-stoichiometry, as observed experimentally. Two means of identification from the Green’s function give a mass enhancement of the order of 12 for already heavy (flat) bands, consistent with the common heavy fermion characterization of UTe$_2$. The predicted Kondo temperature around 100 K matches the experimental values from resistivity. The electric field gradients for the two Te sites are calculated by DFT+U(ED) to differ by a factor of seven, indicating a strong site distinction, while the anisotropy factor $\eta = 0.18$ is similar for all three sites. The calculated uranium moment $<M>_{1/2}$ of 3.5$\mu_B$ is roughly consistent with the published experimental Curie-Weiss values of 2.8$\mu_B$ and 3.3$\mu_B$ (which are field-direction dependent), and the calculated separate spin and orbital moments are remarkably similar to Hund’s rule values for an $f^3$ ion. The $U=3$ eV spectral density is compared with angle-integrated and angle-resolved photoemission spectra, with agreement that there is strong $5f$ character at, and for several hundred meV below, the Fermi energy. Our results support the picture that the underlying ground state of UTe$_2$ is that of a half-filled $j = \frac{5}{2}$ subshell with two half-filled $m_j = \pm \frac{5}{2}$ orbitals forming a narrow gap by hybridization, then driven to a conducting state by configuration mixing (spin-charge fluctuations). UTe$_2$ displays similarities to UPt$_3$ with its $5f$ dominated Fermi surfaces rather than a strongly localized Kondo lattice system.

I. INTRODUCTION

Recently discovered superconductivity (SC) in the heavy fermion material UTe$_2$ [1, 2] below 1.7K, earlier studied in single crystal form, shows a number of peculiar aspects. Unlike in several other U superconductors (UGe$_2$, UCoGe, URhGe, UCoAl) which display coexisting superconductivity and ferromagnetism, no long range magnetic order in the ground state has been observed. The magnetic susceptibility [1,2] has Curie-Weiss character, with magnetic moment in 2.8-3.3$\mu_B$/U range. Magnetic properties in a metal that do not order suggest UTe$_2$ is associated with a class of conducting spin liquids. The large and anisotropic Curie-Weiss magnetic susceptibility suggest antiferromagnetic coupling, whereas other probes [NMR Knight shift and spin-lattice relaxation rate ($1/T_1$)] suggest critical ferromagnetic fluctuations that could mediate SC in UTe$_2$ [1, 2]. The phase diagrams versus field and its direction, temperature, and pressure – including re-entrant superconductivity with application of field – are unusually complex, but those complications will not be addressed in this paper.

UTe$_2$ is however a heavy fermion metal, with a large Sommerfeld coefficient $\gamma \approx 120$ mJ/K$^2$, indicative of Kondo screening of local moments. The resistivity $\rho \sim 1\mu\Omega$ cm is slowly increasing from room temperature down to 75K, then it decreases rapidly over two orders of magnitude just above $T_c$, typical heavy-fermion (HF) behavior with a Kondo temperature of $\sim$100K. The susceptibility for field along the easy $a$ axis increases strongly below 100K, becoming a factor of five or more larger than in the other two directions. The interpretation has been that of critical magnetic fluctuations around incipient FM order along the $a$ axis. A considerable number of measurements of electrical and thermal conductivity, NMR spectra, and penetration depth in magnetic fields gives strong justification of a point-node, Weyl superconducting state.

Based on a non-vanishing specific heat coefficient $C_v/T$ below $T_c$, UTe$_2$ had originally been suggested by Ran and collaborators [5] to provide a new phase of superconducting matter with a ‘Bogoliubov Fermi surface,’ a Fermi surface that is gapless over not points, or lines, but an area. However, subsequent extension of the measurements to 50 mK revealed an upturn in $C_v/T$. The normal state $C_v$, i.e. above the critical magnetic field) was fit with the conventional form augmented by a residual contribution $C_x$ of unknown origin, modeled by a
FIG. 1: (Color online) (A) Projected densities of states/eV (per unit cell) for nonmagnetic UTe$_2$ from the DFT+U(OP) functional ($U=J=0.5$ eV), showing the projections for U 5f (blue) and Te 5p (magenta). Inset: the orthorhombic Immm UTe$_2$ crystal structure. (B) The $j=5/2$ and $7/2$ projected hybridization functions $\Delta(z) = \frac{1}{\pi n_f} \text{Im} \text{Tr}[G_0^{-1}(z)]$ (eV) obtained from LDA calculations. (C) Total densities of states/eV-unit cell, for nonmagnetic UTe$_2$ from the DFT+U(ED) functional. Note that $U=6$ eV opens a gap.

$C_x / T = A_{\text{div}} T^{-1/3}$ form in the region of 50 mK to $T_c$. $C_x$ would arise from thermal excitations that are not gapped by SC.

Supposing this extrapolation is reasonable up to around 2K (taken literally, it would give a divergent entropy), the excess entropy is (we estimate) of the order of $S_x \sim \alpha_x k_B \ln 2$, with $\alpha_x \sim 10^{-2}$, small enough to be extrinsic in origin.

U-based compounds provide a spectacular variety of behaviors, from ground states to unusual spectra. The ferromagnetic superconductors mentioned above have been reviewed and compared by Aoki, Ishida, and Floquet. These compounds seem to require correlation corrections beyond conventional density functional theory (DFT) in its local or semilocal approximations to account for their properties. Superconducting UPt$_3$ ($T_c < 1$ K), on the other hand, has six complicated Fermi surfaces that are described quite well by DFT calculations, with excellent agreement requiring only energy shifts of a few tens of meV. Isovalent and isostructural UPd$_3$, on the other hand, is reproduced only when the U 5f states are localized (removed from the valence states).

The electronic structure of UTe$_2$ has been studied from a first-principles itinerant (local density approximation [LDA]) viewpoint making use of density functional theory (DFT), initially by Aoki and collaborators and Fujimori et al. LDA however predicts a small gap for this heavy fermion metal, so these works were followed by others who have incorporated some correlation correction related to a (semi)localized nature of the 5f states. Two of the current authors applied a DFT+orbital polarization approach [DFT+U(OP), using a Coulomb repulsion $U$ and Hund’s exchange $J_H$] encouraging orbital polarization. Analysis indicated two roughly half-filled 5f orbitals, $m_j = \pm \frac{1}{2}$, at the Fermi level, suggesting multiband half-filled physics.

Some of the most basic theoretical questions are unsettled. Standard DFT calculations give a U 5f occupation near $\frac{3}{4}$. From the viewpoint of strongly localized 5f orbitals ($U=6$ eV), Miao et al. investigated the effect of removing the 5f-states from consideration, i.e. using ThTe$_2$ without 5f occupation as an underlying model of Fermi level bands. Applying dynamical mean field theory (DFT+DMFT) to UTe$_2$ with a large value of repulsion $U$ (with similar results obtained by Xu et al.), the 5f bands were shifted in Mott insulator fashion away from $E_F$, leaving a dispersive Te band and a large Fermi surface that was supported by their angle-resolved photoemission spectra (ARPES). A similar shift of 5f bands was observed by Ishizuka and coauthors applying the DFT+U method, even for moderate values of $U$ in the 1-2 eV range. ARPES data taken at higher energy with longer escape depth however give clear evidence of strong 5f character extending to $E_F$. Both the DFT+U and DFT+DMFT methods predict a predominant 5f$^2$ con-
configuration (our own DFT+$U$ calculations do so as well), whereas uranium core level spectroscopy\textsuperscript{16} indicates the $5f^3$ configuration is dominant, as in several other itinerant uranium intermetallics. We will present evidence that supports a dominant $f^3$ configuration. We address these differences within this paper, with some emphasis on angle-integrated PES (AIPES).

Less strongly correlated methods, such as we will present, leave flat $5f$ bands at $E_F$ in a hybridized band picture consistent with $5f$ dominated Fermi surfaces and with AIPES data, whereas transport and thermodynamic properties will be enhanced by residual (dynamic) interactions. Note that some U compounds, for example UPt$_3$ and UBe$_{13}$, are also heavy fermion metals that do not order magnetically but display exotic superconducting gap symmetry and $5f$ electron dominated Fermi surfaces.\textsuperscript{7}

This itinerant-localized dichotomy is itself not so unusual in metallic actinide compounds, reflecting the “dual” nature\textsuperscript{17} often exhibited by open $5f$ shells. Early on, Hill recognized the shortest U-U separation as a critical parameter in uranium intermetallics and provided the “Hill plot”\textsuperscript{18,19} of ordering temperature (magnetic or superconducting) versus U-U separation. If shorter than 3.5 Å, the $5f$ states are itinerant and sometimes superconducting, if longer they display a local moment and usually order. This critical separation is not absolute: the Hill criterion was found to be violated when the heavy fermion metals UPt$_3$ and UBe$_{13}$ were synthesized and found to be superconducting without magnetic order. Still, the Hill criterion is a very useful and physically motivated guide. In this picture, the nearest neighbor U-U separation in UTe$_2$ of 3.78 Å puts it well into the localized regime of the Hill plot: magnetic rather than superconducting. Just how UTe$_2$ violates the Hill criterion is a fundamental question in the understanding of this fascinating compound.

This dual nature of the $5f$ shell requires that both local and itinerant features of the $f$-electrons may need to be allowed in a description of the electronic structure of UTe$_2$. Ishizuka \textit{et. al.}\textsuperscript{15} performed DFT+$U$ calculations (related somewhat to the DFT+$U$(OP) results mentioned above) for empirically reasonable values of the effective Coulomb $U_{eff} = U - J = 1.1-2.0$ eV, $J=0$. Constraining their DFT+$U$ calculations to the nonmagnetic phase, they obtained a metallic band structure, with reconstruction occurring around $U \sim 1.1$ eV. Contrary to the results of Refs.\textsuperscript{13,14}, the electronic states near the Fermi energy ($E_F$) formed narrow bands with predominantly $5f$-character. The fundamental limitation of these calculations is that they rely on a single-determinant approximation for the $5f$-manifold of the U atom, and the DFT+$U$ method is more adept in describing occurrence and effects of long-range magnetic order. Our calculations parallel to those of Ishizuka \textit{et al.}, indicate that their choice of Hund’s exchange $J=0$, which neglects the anisotropy $U_{m,m'}$ of the repulsive interaction and misses Hund’s exchange, has a substantial effect of the resulting band structure.

In this work we study electron correlation effects in UTe$_2$ beyond the static mean-field DFT+$U$ approximation. We extend DFT method making use an Anderson impurity treatment of the $5f$ shell treated with exact diagonalization (ED) techniques and including the full self-consistency over the charge density. In Sec. II we describe the basic equations of the DFT+$U$(ED) method. In Sec. III the electronic structure results for a conventional value of Coulomb $U=3$ eV are presented and compared to large Coulomb $U=6$ eV results. Sec. IV is devoted to a comparison with photoemission results, and a Summary in Sec. V concludes the paper.

II. STRUCTURE AND METHODS

A. Structure

UTe$_2$ crystallizes in a body-centered orthorhombic $Immm$ structure\textsuperscript{3} (space group $\#71$) with $a=4.161$ Å, $b=6.122$ Å, $c=13.955$ Å, this volume containing two formula units (f.u.) (see Fig.\textsuperscript{1A}). The atomic sites and symmetries are: U, (0,0.13544, 0) mm2; Te1, (0.5, 0.2509, 0) 4j mm2; Te2, (0.0259, 0.5, 0) 4h m2m. The U-U separations are, in increasing order, 3.78 Å dimer separations directed along the $c$-axis, 4.16 Å along the $a$-axis (the lattice constant), 4.89 Å in the $b$-c plane, and 6.12 Å along the $b$-axis (the lattice constant). The normal to the cleavage plane lies $23.7^\circ$ from the $b$-axis in the $b$-c plane. The structure is sometimes pictured as U ladders lying in the cleavage plane, or even two sets of U ladders, but the large U-U separation makes any quasi-one-dimensional aspects difficult to identify. Note: much of our description will use the orthorhombic pseudo-zone with dimensions $\frac{2\pi}{a}$, $\frac{2\pi}{b}$, $\frac{2\pi}{c}$.

B. Formalism of the DFT+$U$(ED) method

Given the combination of itinerant behavior with localized physics in actinide materials and the Fermi liquid character of UTe$_2$, we aim for an effective (low energy) band structure that nevertheless includes essential effects of on-site repulsion and Hund’s exchange on the uranium site and related configurations, and does so self-consistently. We use an extension of the widely used DFT+$U$ method\textsuperscript{20} that makes use of a combination of DFT with the exact diagonalization (ED) of the multiconfigurational $5f$ shell of uranium, in the spirit of a generalized and orbital-occupation (charge, with spin) self-consistent generalization of the Anderson impurity model analogous that that done in dynamical mean field theory.

This DFT+$U$(ED) method takes advantage of the fact that electron interactions in the $s$, $p$, and $d$ shells are well described in DFT, whereas interelectronic interactions within the $5f$ shell are treated explicitly. We use the
fully anisotropic, rotationally invariant implementation of the $+U$ interaction in the DFT+U method in the full-potential linearized augmented plane wave (FP-LAPW) basis that includes both scalar-relativistic and spin-orbit coupling (SOC) effects. The calculations were carried out in the observed paramagnetic state, however both spin and orbital moment effects, such as rms values of the moments, are included in this $U$(ED) extension.

The effects of the interaction Hamiltonian $H_{\text{int}}$ on the electronic structure are described with the aid of an auxiliary impurity model describing the complete fourteen-orbital U 5$f$ shell. This multi-orbital impurity model includes the full Coulomb interaction, the SOC that is very strong in both U and Te, and the crystal field. The corresponding Hamiltonian can be written (see, for example, Hewson),

$$H_{\text{int}} = \sum_{qmm'} \epsilon_{\sigma\sigma'}^{\text{q}} f_{\sigma\sigma'}^{\text{q}} a_{\text{qmm}}^\dagger a_{\text{qmm'}} + \sum_{\sigma\sigma'} \epsilon_\sigma f_{\sigma\sigma}^\dagger f_{\sigma\sigma}$$

where $f_{\sigma\sigma}^\dagger$ creates an electron in orbital $m$ and spin $\sigma$ in the 5$f$ shell and $b_{\sigma\sigma}^\dagger$ creates an electron in state $m\sigma$ in the “bath” that consists of those host band states that hybridize with the impurity 5$f$ shell. $\epsilon_\sigma$ is the energy of the non-interacting 5$f$ ‘impurity’ level, and $\epsilon^q$ are the bath energies.

The parameter $\xi$ specifies the strength of SOC obtained from the atomic potential, and $\Delta_{\text{CF}}$ is the crystal-field (CF) potential at the impurity, as described below. The matrices $V^q$ describe the hybridization between the 5$f$ states and the bath orbitals at energy $\epsilon^q$.

In these calculations two sets of Slater integrals (in eV) $F_0 = 3.00, F_2 = 6.024, F_4 = 4.025, F_6 = 2.94$, and $F_0 = 6.00, F_2 = 6.024, F_4 = 4.025, F_6 = 2.94$ were chosen to specify the Coulomb interaction matrix $U_{\text{mmmm''}}$ in Eq. (1). They correspond to values of $U = 3$ eV and $U = 6$ eV respectively, with exchange $J = 0.51$ eV.

### C. Computational procedure

#### 1. Overview

The calculation uses the same DFT foundation and interaction matrix as implemented by Shick and collaborators and used in conventional DFT+U calculations, but it generalizes the Shick et al. ionic limit approximation (“Hubbard $\Gamma$”) to (a) include hybridization with the environment, and (b) applies exact diagonalization to evaluate the impurity Green’s function, much like some versions of dynamical mean field theory. A crystal field potential is included in the formalism, but for UTe$_2$ it is expected that the DFT treatment of CF is sufficient, as done for other actinides.

To specify the bath parameters, we use the previously reported LDA results for non-magnetic UTe$_2$ (the conventional basic underlying electronic structure before many-body interaction is considered), repeated for this study. Since the interaction $U$-terms in $H_{\text{int}}$ are couched in spin-orbital rather than $j,\gamma$ language, it is judicious to treat SOC in $H_{\text{int}}$. It is assumed that the first and fourth terms in Eq. (1) are diagonal in $\{j,\gamma\}$ representation.

Next, we obtain $V_j^{\text{q}}$ and $\epsilon_q^{\text{q}}$, for $j = \frac{1}{2}$ and $\frac{3}{2}$, from the hybridization functions

$$\Delta_j(z) = \frac{1}{\pi n_j} \text{Im} \text{Tr}_j[G_0^{-1}(z)]$$

where $\text{Tr}_j$ is the trace over the $j$-subspace, with $n_j = 6$ for $j = 5/2$, $n_j = 8$ for $j = 7/2$. $G_0(z)$ is the non-interacting DFT Green’s function extended to the complex energy $z$ plane, equal to the crystal Green’s function in Eq. (1) with the self-energy $\Sigma$ set to zero. The hybridization functions $\Delta_j(z)$ obtained from the LDA calculation without additional adjustment, are shown in Fig. 1. Since the Green’s function is the lowest-lying eigenstates of the many-body Hamiltonian $H_{\text{imp}}$ and to calculate the one-particle Green’s matrix $[G_{\text{imp}}(z)]_{\gamma\gamma'}$ in the subspace of the $f$ spin-orbitals $\{\phi_{\gamma} = \phi_{\gamma\gamma'}\}$ at low temperature ($k_B T = \beta^{-1} = (1/500)$ eV, $T = 40$ K). The expression for $G_{\text{imp}}$ is

$$[G_{\text{imp}}(z)]_{\gamma\gamma'} = \frac{1}{Z} \sum_{\alpha,\beta} \langle \alpha | c_{\gamma} | \beta \rangle \langle \beta | c_{\gamma'}^{\dagger} | \alpha \rangle \exp [-\beta E_{\alpha} + \beta E_{\alpha}']$$

where $Z$ is the partition function, $E_{\alpha}$ is the energy of the eigenstate $| \alpha \rangle$ of Eq. (1), and $z$ is the (complex) energy. The self-energy matrix $\Sigma_{\gamma,\gamma'}(z)$ is then obtained from the inverse of the Green’s function matrix $G_{\text{imp}}$.

The self-energy is then inserted into the local Green’s function $G(z)$

$$G_{\gamma\gamma'}(z) = \int_{\text{BZ}} \frac{d^3k}{V_{\text{BZ}}} \left[ z + \mu - H_{\text{DFT}}(k) - \Sigma(z) \right]_{\gamma\gamma'}^{-1},$$

where $H_{\text{DFT}}(k)$ is the DFT Hamiltonian and $\mu$ is the chemical potential.
calculated in a single-site approximation as described previously [24, 26] from which 5f orbital occupations are obtained. \( V_{BBZ} \) is the volume of the Brillouin zone (BZ). The self-energy is adjusted at each iteration until self-consistency is reached. Since this method has not been used previously, we provide additional details for purposes of clarity in Appendix A, which provides a flow chart and a step-by-step description of the procedure.

3. Density matrix self-consistency

In a single site approximation, the local Green’s function matrix \( G(z) \) for the 5f electrons in the manifold is

\[
G(z) = \left( G_0^{-1} + \Delta \mu - \Sigma(z) \right)^{-1},
\]

where \( G_0(z) \) is the non-interacting Green’s function \( \Delta \mu \) is a correction to the chemical potential chosen to ensure that \( n_f = -\pi^{-1} \text{Im} \int_{-\infty}^{E_F} dz G(z) \) is equal to the number of correlated \( f \)-electrons obtained from Eq. (7). Then, with the aid of this local Green’s function \( G(z) \), we evaluate the occupation matrix

\[
n_{\gamma\gamma'} = -\frac{1}{\pi} \text{Im} \int_{-\infty}^{E_F} dz G_{\gamma\gamma'}(z).
\]

For the energy integrations we use \( \text{Im} \Sigma(z)/\pi = 0.01 \) eV, and a grid along the real \( z \) axis of 0.01 eV.

The matrix \( n_{\gamma\gamma'} \) is used to construct an effective LDA+U potential \( V_U \), which is inserted into the Kohn–Sham-like equations [21]:

\[
\left[ -\nabla^2 + V_{\text{LDA}}(\mathbf{r}) + V_U + \xi (1 \cdot s) \right] \Phi_k(\mathbf{r}) = \xi_k \Phi_k(\mathbf{r}).
\]

For the spherically-symmetric LDA+U double-counting term (included in the potential \( V_U \)) we have adopted the fully localized limit (FLL) form \( V_{dc} = U (n_f - 1/2) - J (n_f - 1/2) / 2 \). We also note that the LDA potential \( V_{\text{LDA}} \) in Eq. (7) acting on the \( f \)-states is corrected to exclude the non-spherical double-counting with \( V_U \). [20]

The equations in Eq. (7) are iteratively solved until self-consistency over the charge density is reached. The DFT+U Green function matrix \( G_{+U} \) is calculated from Eq. (7) substituting the self-energy \( \Sigma(\epsilon) \) by the DFT+U potential \( V_U \).

The new value of the 5f-shell occupation is obtained from the solution of Eq. (7), and defines the new value of \( \epsilon_f = -V_{dc} \) in Eq. (11). The \( f \)-shell SOC parameter (an atomic quantity), and the CF matrix \( \Delta_{\text{CF}} \) in Eq. (1) are determined in each iteration. The CF matrix \( \Delta_{\text{CF}} \) in Eq. (1) is obtained by projecting the self-consistent solutions of Eq. (7) into the \( \{\phi_i\} \) local \( f \)-shell basis, giving the “local Hamiltonian”

\[
[H_{\text{loc}}]_{\gamma\gamma'} = \int_{\epsilon_0}^{\epsilon_F} d\epsilon \left[ N(\epsilon) \right]_{\gamma\gamma'} \approx \epsilon_0 \delta_{\gamma\gamma'} + \left[ \xi \cdot s + \Delta_{\text{CF}} \right]_{\gamma\gamma'} + \left[ V_U \right]_{\gamma\gamma'},
\]

where \( [N(\epsilon)]_{\gamma\gamma'} \) is the \( f \)-projected density of states (fDOS) matrix (whose integral to \( E_F \) gives the familiar occupation matrix), \( \epsilon_0 \) is the bottom of the valence band, \( \epsilon_F \) is the upper cut-off, and \( \epsilon_0 \) is the mean position of the non-interacting 5f level. The matrix \( \Delta_{\text{CF}} \) is then obtained by removing the interacting DFT+U potential and SOC \( [\xi \cdot s]_{\gamma\gamma'} \) from \( H_{\text{loc}} \). As mentioned, for UTe2 it is anticipated that the CF is represented sufficiently by DFT+U, and this step is neglected.

The self-consistency loop is closed by calculating the non-interacting Green’s function \( G_0 \),

\[
G_0(z) = \left( G_0^{-1} + \Delta \mu - \Sigma(z) \right)^{-1}
\]

and the next iteration is started by solving Eq. (1) for the updated \( \epsilon_f \), \( \xi \), and \( \Delta_{\text{CF}} \), and calculating the new self-energy \( \Sigma(\epsilon) \). The self-consistent procedure was repeated until the convergence of the 5f-manifold occupation matrix, with \( n_f \) converged to less than 0.01.

III. COMPUTATIONAL RESULTS

The U and Te atoms projected densities of states (DOS) for \( U=3 \) eV, and the total densities of states for \( U=3 \) eV and 6 eV, are shown in Fig. 1. The band structures for two different Coulomb \( U \) values are shown in Fig. 2. For all values of \( U \), the DOS near \( E_F \) is almost entirely due to U 5f states (see Table I). As several groups have noted, there is a small gap for \( U=0 \), the curious LDA result.

Increasing \( U \), the two flat bands near \( E_F \) become flatter, approach each other, and by \( U=3 \) eV become inverted, leaving a band crossing along \( \Gamma-X \) very near \( X \) almost exactly at the Fermi level. Increasing \( U \) to 6 eV, the bands separate leaving a small 10 meV bandgap, again separating disjoint valence and conduction bands. This moving away from \( E_F \) of the 5f bands is qualitatively consistent with DMFT results using \( U=6 \) eV. However, we do not obtain any highly dispersive Te 5p, U 5d band in the background, crossing \( E_F \), as in DMFT. In fact, for \( U=6 \) eV we obtain again a small gap, whereas (to repeat) UTe2 is observed to be a heavy fermion metal, not necessarily in conflict with a semimetal before dynamic correlations are included. Henceforward we focus on our \( U=3 \) eV results with flat 5f bands crossing \( E_F \).

A. Band structure and Fermi level quantities

Density functional theory with (semi)local exchange-correlation functional (\( U=0 \)) give an insulating band structure for UTe2, albeit with a very small 10–15 meV gap. Such a gap generally signals a bonding-antibonding separation of bands, but no such description has been forthcoming for UTe2. The gap reflects a different brand of band insulator. Every calculation reveals that the
large uranium SOC separates the $j = \frac{5}{2}$ and $j = \frac{7}{2}$ sub-shells separated by 1.5 eV (see Fig. 1), and with a $5f^3$ configuration the former subshell is half-filled. This SOC separation is much larger than any crystal field splitting. The previous work of some of the present authors\textsuperscript{12} established that, with ferromagnetic order imposed, two orbitals, $m_j = \pm \frac{5}{2}$, are half-filled, then hybridized, leading to the gap which is however very small. How physical UTe\textsubscript{2} obliterates this gap and in the process emerges as a nearly magnetic but superconducting material is the fundamental issue in the electronic structure of this compound.

Henceforward we focus on $U=3$ eV results unless otherwise stated, since this value is sufficient to restore a conducting band structure, and is typical of values in most uranium intermetallics. From Fig. 2, this “insulator-metal” transition arises from a hybridization reconfiguration of energy levels at the zone boundary points $U = (\frac{\pi}{2},0,\frac{\pi}{2})$ and $X = (\frac{\pi}{2},0,0)$. A distinctive feature is that all four bands shown in Fig. 2 are exceedingly flat along $X-U$ (the $k_z$ direction), unlike for $U=0$ or $U=6$ eV. Another feature is the band crossing along $\Gamma-X$ very near the point $X$. The unoccupied band for $U=6$ eV is nearly dispersionless along the three directions $X-U-R-S$, before mixing with dispersive bands in other regions of the zone.

The values of $N(E_F)$, $N_f(E_F)$, and Fermi velocities along the three crystal axes for $U=3$ eV are provided in Table I. The Fermi velocities are the r.m.s. FS averaged values $v_{F,x} = \sqrt{\langle v_{k,x}^2 \rangle_{FS}}$, and similarly for $yy$ and $zz$ components. The anisotropy is only 10-15%, indicative of three dimensional conduction; anisotropy is larger for the individual bands. The magnitudes for the separate bands, 0.3-1.4 $\times 10^5$ cm/s, indicate very heavy carriers even before renormalization by dynamical processes (electronic and phononic).

\begin{table}[h]
\centering
\begin{tabular}{|c|c|c|c|c|}
\hline
Atom & site & $\eta$ & $V_{zz}$ & $V_{yy}$ & $V_{xx}$ \\
\hline
U & 4i & 0.183 & 13.85 & -8.19 & -5.66 \\
Te1 & 4j & 0.173 & 4.56 & -2.67 & -1.89 \\
Te2 & 4h & 0.176 & 33.28 & -19.56 & -13.71 \\
\hline
\end{tabular}
\caption{Electric Field Gradient}
\end{table}

From Table I one sees that 90% of $N(E_F)$ is provided by the U $5f$ states. $N(E_F) = 11.0$ states/eV for $U = 3$ eV corresponds to a band Sommerfeld constant $\gamma = 13.0$ mJ/mol-K\textsuperscript{-2}. This implies a mass enhancement of nearly 9 from dynamic interactions compared to the experimental value. Note that there is a strong peak in the DOS (up to 59 states/eV) just 10 meV above $E_F$, corresponding to $\gamma = 70$ mJ/mol-K\textsuperscript{-2} which is within a factor of two of the experimental value of $\gamma = 120$ mJ K\textsuperscript{-2} mol\textsuperscript{-1}.\textsuperscript{11} The narrow peak just above $E_F$ implies a strong dependence of properties on stoichiometry.
FIG. 3: Upper panel: The Fermi surfaces of UTe$_2$ from DFT+U(ED), for Coulomb $U=3$ eV. The colors provide the relative Fermi velocities. The high symmetry k-points are $\Gamma[0,0,0]$, $X[\pi/a,0,0]$, $U[\pi/a,0,\pi/c]$, $R[\pi/a,\pi/b,\pi/c]$, $S[\pi/a,\pi/b,0]$, $Y[0,\pi/b,0]$. Lower panels: the 5$f$ fat-band structure of UTe$_2$, with the circle size indicating the amount of $m_l, m_s, m_j = m_l + m_s$ character in the bands, as labeled.
In materials with large SOC where the \( j = \frac{5}{2}, j = \frac{7}{2} \) splitting dominates site anisotropy and crystal field splitting but symmetry is low, state characters are not very transparent. The \( |m_l, m_s| > \) and \( |j, m_j| = m_l + m_s > \) decompositions of \( N_j(E_F) \) are provided in Appendix B, Table 11. The important bit of information is that the \( |\pm\rangle \) components (equal by symmetry) are three times larger than the \( |\pm\rangle \) components, and five times larger than \( |\pm\rangle \) components, with the \( m_j = \pm \frac{5}{2} \) components being negligible. This values reflect strong spin-orbital polarization at the Fermi level in UTe\(_2\), which also shows up in the strong anisotropy of the electric field gradients, below.

The Fermi surfaces (FS) are displayed in the upper panel of Fig. 3. The FS has three types of sheets: from the lower band is the hole sheet centered at \( \Gamma \) and a fluted hole column along \( X - U \), with masses varying by \( \sim 50\% \) over the sheets. The second band gives two symmetry related electron ellipsoids midway between \( U \) and \( R \), providing the required charge compensation. The corresponding Fermi velocities, with relative values shown by the colorbar in Fig. 3 and mean values provided in Table 1, have somewhat less than factor-of-two anisotropies.

The array of band graphics in the lower part of Fig. 3 presented with fat-band character and plotted along Cartesian directions, provides the relative amounts of the stated spin-orbital characters of bands near the Fermi level. The major contribution to FS-1 along the \( X - U \) line arises from \( |\pm\rangle \) > orbitals. The \( \Gamma \)-point centered hole spheroid is more \( |\pm\rangle \) > in character. The electron sheet along \( U - R \) arises from a mixture all three of these orbitals. As in our previous work\[12\], we find that the Fermi level states are dominated by \( |\pm\rangle \) > orbitals with some contribution from the \( |\pm\rangle \) orbital.

Our DFT+U(ED) results, obtained without restriction to a single-determinant reference state in determining the density, can be contrasted with previous beyond-DFT results. Admittedly, the groups that are involved are searching for a treatment that will provide a realistic platform for further considerations and experimental properties. Conventional DFT+U was applied by Ishizuka et al.\[14\], who chose the non-standard Hund’s \( J=0 \) approach, neglecting Hund’s exchange and anisotropy of the \( U_{m,m',m'',m'''} \) matrix. A crossover in behavior was found for values of \( U \) in the 1-1.5 eV range. Our earlier treatment of orbital polarization by the DFT+U(OP) method\[12\] focused on ferromagnetic alignment based on the observed large Curie-Weiss moment (which however does not order) so it is less directly comparable. The problem posed by the unphysical gap in LDA was addressed by Harima\[27\] with a hands-on shift of the 5\( f \) energy by 1.36 eV, giving small Fermi surfaces arising from flat bands much as we find, and by 2.72 eV, which led to a band structure with large Fermi surfaces. As mentioned, the work of Miao et al.\[13\] was strongly influenced by the apparent similarity of a dispersive band crossing \( E_F \) in ARPES that is like that in ThTe\(_2\), which has no 5\( f \) bands.

### B. Local U atom 5\( f \)-shell properties

The calculated 5\( f \) occupation within the uranium atomic sphere is \( n_f = 2.73 \), close to the value obtained in DFT+U(OP) calculations\[12\], and supporting the viewpoint of a reference U 5\( f^3 \) configuration. This value is a slight underestimate, since 5\( f \) orbitals extend somewhat beyond the atomic spheres used to obtain this number. The electronic structure results discussed above have been more consistent with an \( f^2 \) viewpoint, but without the comparison of calculated moments with the experimental Curie-Weiss moment that we have provide below.

#### 1. X-ray absorption

X-ray absorption spectroscopy (XAS) provides additional local information. XAS yields two intensities \( I_{5/2} : 4d_{5/2} \rightarrow 5f_{5/2,7/2} \) and \( I_{7/2} : 4d_{3/2} \rightarrow 5f_{5/2,7/2} \) and the branching ratio\[28\] \( B = I_{5/2}/(I_{5/2}+I_{3/2}) = 0.71 \). These are connected to the partial occupations \( n_{5/2} = 2.47 \) and \( n_{7/2} = 0.26 \). Our value of \( B \) can be directly compared to future experimental results for XAS and electron energy-loss spectroscopies. The \( |j, m_j| > \) decompositions of the U 5\( f \)-occupations are provided in Appendix B.

#### 2. Curie-Weiss moment

For the self-consistently determined impurity energy position \( \epsilon_f = -V_{da} \), we obtain a doubly (Kramers) degenerate ground state with spin, orbital, and total moments of \( S = 1.34, L = 5.80, J = 4.53 \) in \( \mu_B \). The calculated \( g \)-factor is 0.78, and \( \langle m_j \rangle = \pm 0.55 \). The Curie-Weiss magnetic moment \( \mu_{CW} = g/\langle J + 1 \rangle = 3.52 \mu_B \), calculated as the rms value over occupied uranium configurations, is in reasonable agreement with experimental values of 2.8\( \mu_B \)\[1\], and 3.3\( \mu_B \)\[2\]. These can be compared to the textbook values for an \( f^3 \) configuration: \( S = -\frac{3}{2}, L = 6, J = \frac{9}{2} \), resulting in \( g = 0.72 \). These Hund’s rule numbers, not normally reliable for 5\( f \) materials, are remarkably similar to the DFT+U(ED) results.

#### 3. Mass enhancement

We obtain an estimated average mass enhancement, without contributions from dynamical corrections, in two ways. From the mean spectral density,

\[
\frac{m^*}{m} = \frac{Tr[\hat{Z}^{-1}\hat{N}(E_F)]}{Tr[\hat{N}(E_F)]},
\]

where \( \hat{Z}^{-1} = [\hat{I} - dRc|\Sigma(e)|/de]^{-1} \) is the quasiparticle residue matrix, and \( \hat{N}(E_F) = -\frac{1}{2} \text{Im} \text{Tr}[\hat{G}(E_F)] \) is the spectral density matrix obtained from Eq. \[1\]. The quasiparticle weight \( Z(\frac{\mu}{m_F}) = 0.06 \) is obtained. This small value of \( Z \) indicates the strongly correlated character of \( f \)-electrons at \( E_F \). The renormalized perturbation theory \[23\] expression for the Kondo temperature is

\[
T_K = \frac{\pi^2}{4} Z \Delta(E_F). \tag{10}
\]

From the value of the hybridization function \( \Delta(E_F) \approx 50 \text{ meV} \), the predicted \( T_K \approx 100 \text{ K} \), very similar to experimental values from resistivity. \[1\]

4. Electric field gradients

Electric field gradients (EFG) provide a measure of the charge distribution (mostly from the local charge) that is available from all-electron calculations. Following the analysis of the electronic structure quantities given above, we have calculated the electric field gradients \[29\] in UTe\(_2\). Their values, together with the dimensionless asymmetry parameter for the \( x - y \) plane values,

\[
\eta = \frac{V_{xx} - V_{yy}}{V_{zz}} = \frac{V_{yy} - V_{xx}}{V_{yy} + V_{xx}}
\]

are provided in Table \[I\]. The second expression follows from the traceless nature of the EFG tensor. The EFG component \( V_{zz} \) is proportional to the nuclear quadrupolar resonance (NQR) frequency \( \nu_Q \). Experimental measurements of NQR require stable isotopes with nuclear spin \( I \geq 1 \). No such measurements have performed to date, so our calculations provide a prediction of this measure of the anisotropy of the charge density and resulting Hartree potential at the nuclei.

The most notable result is that the values for the Te2 site are roughly a factor of 7 larger than for the Te1 site, reflecting a substantially different charge distribution around the two sites. The values for uranium lie midway between, and the anisotropy factors \( \eta \approx 0.18 \) are the same for all three atomic sites. Measurement of some of these will provide useful information on the electronic density distribution, and orbital polarization, of UTe\(_2\).

IV. COMPARISON BETWEEN BAND STRUCTURE AND PHOTOEMISSION DATA

Comparison of calculated bands (or spectral density) with PES data is the most direct means of determining the basis of the electronic structure. Due to a number of experimental challenges – matrix element effects, energy and \( k \) resolution, band broadening due to dynamical effects, surface sensitivity – comparison can yet leave uncertainty, especially in quantum materials with strong dynamical processes. Angle-integrated averages over the momentum and matrix element dependence, giving the zone-averaged spectral density, provides the most unambiguous information – the spectral distribution of valence band states – on fundamental aspects of the electronic structure. We remind that a band picture provides an optimum set of single particle characteristics (orbitals and eigenvalues) to best describe ground state characteristics – energies, charge and spin densities and quantities derivable from them, for example, EFGs – and by continuity in metals, near ground state quantities. Single particle excitations involve self-energies that are minor in many metals but become central in describing heavy fermion metals. Occupied multiplets and configurations are sampled in the DFT+U(ED) method to determine spin-orbital occupations, while dynamic effects are not included in the band structure we present.

A. AIPES

Data taken at 20K with \( h\nu=800 \text{ eV} \) photon energy and 120 meV resolution are compared in Fig. \[4\] with the resolution-broadened DFT+U(ED) DOS; other details of the experiment and analysis are described by Fujimori and collaborators.\[11\]. The unbroadened DOS is shown in Fig. \[1\]. As mentioned above, the most significant region for comparison is at and close below the Fermi energy, where the dynamical self-energy is small. The leading edges at low energy in Fig. \[4\] are in extremely good experiment-theory agreement. This result is crucial, because DFT+U(ED) displays flat 5f bands at and

![Figure 4: Measured angle-integrated photoemission spectrum (green) plotted with the resolution broadened (120 meV Gaussian) and Fermi-Dirac cutoff (20 K) DFT+U(ED) DOS. The agreement of the onsets below \( E_F \) reflect flat and heavy bands that become the basis of the heavy fermion superconductor state. The broadening of the experimental spectrum down to \( \sim\text{4 eV} \) is the expected effect of dynamical fluctuations in the uranium 5f shell. The DFT+U(ED) DOSes are enhanced by a factor of five to reveal possible Te influence on structure in the -4 eV to -1 eV range, which is mostly due to single particle excitations from the 5f shell.](image-url)
FIG. 5: Upper right panel: Brillouin zone structure of UTe$_2$, panels A, B, C: ARPES spectra (left) together with the band structure (right) for the designated directions and with energies aligned. The size of the calculated points provide the relative amount of 5$f$ character of the wavefunction. Panels A and C provide new data, while panel B is a replot of earlier data of one of the authors to provide clear comparison with our theoretical results. The k-points notations of the orthorhombic pseudo-zone Γ[0,0,0], X[π/a,0,0], U [π/a,0,π/c], Z [0,0,π/c], Y[0,π/b,0], Γ′[0,2π/b,0] are used in the text.

immediately below $E_F$, whereas both DFT+$U$[15] and DFT+DMFT[13] displace 5$f$ weight away from $E_F$ by several hundred meV for DMFT, or more for DFT+$U$. This energy shift is much larger than the experimental resolution, so the corresponding “5$f$ edge” in those spectra should be displaced by this amount from $E_F$, which is not seen in the data. These AIPES data thus support the view that flat bands lie at $E_F$, consistent with giving small renormalized Fermi surfaces that provide the platform for the observed exotic superconducting states that are observed to be extraordinarily sensitive to magnetic field, pressure, and stoichiometry.

The AIPES intensity extending to 3-4 eV binding energy can be interpreted in terms of the excitations involving 5$f^3$ and 5$f^2$ multiplets, as discussed by Miao et al. [13] These considerations involve the relative participations of populations $f^2$ and $f^3$. The DFT+DMFT treatment of Miao et al. lead to a dominant $f^2$ ($^3H_4$, 84%) state; DFT+$U$ (with $U$ and $J$ values described above) gives exactly $f^2$ (two strongly bound 5$f$ bands). Our DFT+$U$(ED) method leads to a dominant $f^3$ description. Summing the spin-orbital occupations provided in Appendix B (2.73) gives a slight underestimate due to 5$f$ orbital tails extending beyond the sphere boundary. The disjoint valence and conduction bands discussed above argues for a half-filled $\frac{1}{2}$ subshell, which is $f^3$.

B. ARPES

ARPES data were obtained in the photon energy range 565-800 eV with energy resolution of 90-115 meV, with other aspects of the sample and setup described in Ref. [11]. In Fig. 5 we show ARPES results compared with the relevant band lines. For the U-Z-U line shown in Fig. 5A, agreement between the correlated bands and ARPES data near $E_F$ is apparent, with a dispersive band dropping down at $Z = [0,0,π/c]$ being resolved clearly. For the X-Γ-X direction in Fig. 5B, the calculations yield heavy $f$-bands located near and touching $E_F$. The bands lie at the same energy as intense emission in the experimental data, and are separated by less than the experimental resolution and sometimes crossing, so theory and experiment are consistent although incoherence in the data at this low energy cannot be ruled out. This intensity is at variance with the presence of a light band observed by Miao et al. [13] using photon energies in the 30-150 eV range.

Along the Γ-Y-Γ’ line in Fig. 5C, incoherence in the data lies in the energy region of a dispersive band, apparently reflecting weakly dispersing, largely incoherent, 5$f$ shell excitations. The strong intensity at [0,0,0] around -60 meV is in a region where our band structure predicts flat 5$f$ bands. Note also that Fig. 5C indicates a dispersive Te $p$ band passing through the 5$f$ bands and mixing
strongly in the calculations. Thus $f - p$ mixing is substantial, with a result that the Te $p$ character is strongly excluded from the $5f$ band regions, as is clear from the projected DOS in Fig. 1.

Both the calculated bands near $E_F$ as well as this data, and also the ARPES data of Ref. [11], differ from the results of Ref. [13]. We attribute the differences in the experimental ARPES data to the higher surface sensitivity of the spectra taken in the 30-150 eV [13] where the escape depth is $\sim$8-12 Å. The energy dependence of the electron escape depth [30] indicates that it is roughly twice as large [30] in our energy range. Thus our data are more bulk sensitive while those of Miao et al. are more surface impacted, where confinement imposed by the surface can lead to the U $5f$-electrons becoming more localized than in the bulk. The low energy ARPES data of Miao et al. were mostly interpreted by the band structure of ThTe$_2$, i.e., any without $5f$ bands whatsoever. Conversely, our ARPES data together with our correlated band results, along with earlier AIPES data [11] emphasize the presence of heavy $5f$ bands near $E_F$.

V. SUMMARY

While a great deal of experimental data has been collected that is relevant to the complex phase diagram of UTe$_2$, and there are several theoretical suggestions about the character and symmetry of its superconducting and magnetic phases, there is not yet any consensus emerging on its basic electronic structure. Given its heavy fermion properties this may not be so surprising, but relative to modestly correlated DFT-based calculations, the dynamical mass enhancement is substantial but not particularly large. The Hill criterion for the critical uranium atom separation [15, 19] is 3.5 Å, and clearly UTe$_2$ lies on the localized side of that limit, but it does not order magnetically at zero field. The Hill criterion is however sometimes violated in uranium compounds, so that guidance it provides is limited.

There is evidence that neither a fully localized nor simple itinerant picture holds for UTe$_2$. Our correlated band DFT+U(ED) calculations, motivated by the Anderson impurity model and taking into account the multiconfiguration aspect of the U $5f$ shell, suggest that both local and itinerant characteristics of the $f$-electrons appear near the Fermi level and need to be treated together. The uranium magnetic moment is given well compared to small Fermi surfaces that will be responsive to pressure and magnetic field, a sensitivity that is very clear in the emerging experimental phase diagram.

Finally, our low energy band structure shows evidence of Te $6p$ – U $5f$ mixing, which repels the Te $6p$ character from the low energy region rather than opening up to allow a dispersive Te band to cross the Fermi level. Still, the Te influence is important, and the large U-U separation placing it in magnetic ion regime suggests that exchange coupling in UTe$_2$ proceeds primarily of RKKY character mediated through the Te conduction bands. The overall picture is one of Fermi level bands dominated by $5f$ character as in UPd$_2$ versus a more strongly localized moment as in the Kondo lattice picture.

VI. ACKNOWLEDGMENTS

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Appendix A: Density matrix self-consistency in DFT+U(ED)

The calculation follows in several respects that of Shick and collaborators for their “DFT+Hubbard I approximation” study of elemental actinides. A few generalizations have been adopted here, with the main steps being illustrated in the flow diagram in Fig. 6. The various steps occur in the following order.

1. From a DFT/LDA calculation the crystal Green’s function $G_0(z)$ is constructed and initial parameters to solve Eq. (1) are chosen as reasonable estimates. For example, the repulsion $U$ and Hund’s rule $J$ constants (from the Slater parameters) must be specified.

2. From the (self-consistent) DFT calculation, the hybridization matrix is evaluated, from which the bath energies $\epsilon_{q=1}$ and hybridization strengths $V_q^j$ are chosen to represent mixing with the environment, as described in Sec. II.C. The input parameters to the DFT+U(ED) calculation are now determined.

3. Carry out an exact diagonalization of $H_{int}$ of Eq. (1). This step is represented by the top panel of the flow chart in Fig. 6. Use eigenvalues and eigenvectors to construct the impurity Green’s function $G_{imp}$ in Eq. (3) and the corresponding self-energy; adjust $\mu$ in Eq. (5) to fix the value of $n_f$ to the current value from the full calculation equation (7). Calculate the occupation matrix equation (6) as stated in Sec. II.C.3.

4. From the occupation matrix, set up the DFT+U potential, solve self-consistently the Kohn-Sham equation (7), to find the updated charge density according to some reliable prescription typically present in DFT codes. Obtain the DFT+U Green’s function. See the central panel in Fig. 6.

5. Obtain a new non-interacting Green’s function $G_0$, a new position $\epsilon_f$ of the impurity level, a new $\xi$ and (when used) $\Delta_{CF}$. This step is represented in the bottom panel in Fig. 6.

6. Now close the self-consistency loop. When output and input of the 5$f$-manifold occupation $n_f$ converge agree within a specified criterion, the self-consistency loop is exited, and analysis of the results follows.

Appendix B: Decomposition of the U 5$f$ DOS at $E_F$, and the U 5$f$ occupations

The essential aspects of the electronic structure of UTe$_2$ finally reduce to participation of various spin-orbitals in the 5$f$ occupation, and to their play at states at the Fermi level. Table II provides the Fermi level quantities in both $(\ell,s)$ and $(j,m_j)$ representations. The primary result to notice is the large and equal participation of the $m_j = \pm \frac{7}{2}$ orbitals.

| $m_j$ | -3 | -2 | -1 | 0 | 1 | 2 | 3 |
|-------|----|----|----|---|---|---|---|
| Spin↑| 0.48 | 0.23 | 0.85 | 0.81 | 0.10 | 0.07 | 0.01 |
| Spin↓| 0.01 | 0.07 | 0.10 | 0.81 | 0.85 | 0.23 | 0.48 |

Table III provides the strong spin-orbit decompositions for moment along the easy $\hat{a}$-axis, these values are for Coulomb $U=3$ eV.

Table III | $m_i, m_s >$ and $|j, m_j >$ decompositions of the U atom $f$-projected DOS at $E_F$ (in 1/eV) are provided for the unpolarized system. The magnetic quantization is along the easy $\hat{a}$ axis, these values are for Coulomb $U=3$ eV.

Table III

| $m_i$ | $m_s$ | $j$ | $m_j$ | $E_F$ (1/eV) |
|-------|-------|-----|-------|-------------|
| -3 | -2 | -1 | 0 | 1 | 2 |
| Spin↑ | 0.48 | 0.23 | 0.85 | 0.81 | 0.10 | 0.07 | 0.01 |
| Spin↓ | 0.01 | 0.07 | 0.10 | 0.81 | 0.85 | 0.23 | 0.48 |

Table III provides the strong spin-orbit decompositions for moment along the easy $\hat{a}$-axis. The $j = \frac{7}{2}$ contribution is minor.
TABLE III: The $|j,m_j>$ decompositions of the U atom $f$-occupation for the unpolarized system. The magnetic quantization is along the easy $\hat{a}$ axis. Coulomb $U=3$ eV and 6 eV values are provided.

| $j$ / $m_j$ | -7/2 | -5/2 | -3/2 | -1/2 | 1/2 | 3/2 | 5/2 | 7/2 |
|------------|------|------|------|------|-----|-----|-----|-----|
| 5/2        | 0.412 | 0.410 | 0.415 | 0.415 | 0.410 | 0.412 |
| 7/2        | 0.031 | 0.037 | 0.031 | 0.030 | 0.030 | 0.031 | 0.037 | 0.031 |

$U=6$ eV

| $j$ / $m_j$ | -7/2 | -5/2 | -3/2 | -1/2 | 1/2 | 3/2 | 5/2 | 7/2 |
|------------|------|------|------|------|-----|-----|-----|-----|
| 5/2        | 0.410 | 0.417 | 0.410 | 0.410 | 0.417 | 0.410 |
| 7/2        | 0.027 | 0.019 | 0.033 | 0.022 | 0.022 | 0.033 | 0.019 | 0.027 |

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