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Spectroscopic Evidence for the 5f Coulomb Interaction in UAl2 and UPt3

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The 5f spectral weight in the heavy-fermion materials UAl2 and UPt3 has been measured by electron spectroscopy, and compared to that in density-functional calculations. The one-electron 5f widths are both too large to account for the enhanced values of the specific-heat coefficient γ, and too small to account for the measured widths. In the Anderson Hamiltonian, both discrepancies arise from the 5f Coulomb interaction. It is pointed out that an enhanced γ value does not imply enhanced magnitude of the 5f spectral weight at E_f.

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A number of materials containing Ce and U have very large low-temperature specific heats, leading to the descriptive term "heavy fermion." The ground states vary from magnetic to superconducting, and it is generally agreed that the Ce and U f electrons give rise to these novel and variable properties. A central assumption of current many-body approaches to these materials is the existence of an f Coulomb repulsion U_ff which is larger than the one-electron f width. For cerium materials this is well established (2-5) by the observation of separate peaks split by U_ff in combined photoemission (PES) and bremsstrahlung isochromat (BIS) spectra, which measure the f spectral weight of the single-particle Green’s function. For metallic uranium materials the situation is much different. Typically, separated peaks are not observed and in uranium metal, for example, the total observed f width is comparable to the one-electron width obtained in band calculations. For only one metallic uranium material has spectroscopic evidence for U_ff been reported, and this material is not of the heavy-fermion type (6) Thus far UBe13 is the only heavy-fermion uranium material for which a complete BIS-PES 5f spectrum has been published (7). The total 5f width was found to be about the same as in uranium metal, i.e., much too large to relate to the large specific heat. No conclusion about the relative roles of one-electron and Coulomb interactions was reached.

In this paper we report combined BIS-PES spectra for the heavy-fermion uranium materials UAl2 and UPt3. The specific heats of both materials (9,10) have a large T-linear coefficient γ, and a T^2 ln T term often taken (9-13) to imply spin fluctuations. Additionally, UPt3 is a superconductor for which triplet Cooper pairing has been speculated (10). We find BIS spectra which are more structured than found for UBe13, and can compare our 5f spectral weights to density-functional calculations (14,15) very recently available for these two materials. These calculations provide a one-electron fiduciary for the interpreting of the observed 5f widths, leading to an interesting conclusion not given previously, that the one-electron 5f bandwidths are, on the one hand, too large to account for the large γ values, but are, on the other hand, too small to account for the measured 5f widths. Appealing to existing theoretical results for the Anderson Hamiltonian, we argue below that both these apparently contradictory effects arise simultaneously from the Coulomb interaction U_ff.

BIS and PES spectra were obtained at a photon energy of 1486.6 eV with use of a Vacuum Generators ESCALAB, operated under conditions to yield resolutions of 0.5 and 0.8 eV, respectively. Clean sample surfaces were obtained by the fracturing of polycrystalline ingots in a vacuum of 5×10^-11 Torr, after which the sample was transferred under the same vacuum to the measurement position and cooled to ~100 K. Valence-band and core-level PES were then measured, followed by BIS. During data taking the chamber pressure was always below 7×10^-11 Torr for x-ray pho-
toelectron spectroscopy and $1 \times 10^{-10}$ Torr for BIS. Subsequently the PES and BIS measurements were repeated at room temperature, with no significant differences in the data.

Room-temperature PES measurements with variable photon energy were performed at the Stanford Synchrotron Radiation Laboratory, using beam line I-1 for UPt$_3$ and beam line III-1 for UA$_2$. The UPt$_3$ sample was the same as that used for the PES-BIS measurements, while the UA$_2$ sample was a small unoriented single crystal. Both samples were fractured and measured in a vacuum of $8 \times 10^{-11}$ Torr, in a vacuum chamber equipped with a commercial cylindrical-mirror analyzer to measure electron energies. The resolution, which is determined almost entirely by the electron analyzer for the photon energies used, was 0.4 eV.

Figures 1 and 2 show resonant photoemission and BIS spectra for UA$_2$ and UPt$_3$, respectively, with their inelastic backgrounds removed by a standard method. The PES spectra for photon energy $h\nu = 92$ eV have the U 5$f$ emission resonantly suppressed and thereby reveal Al 3$s$-3$p$ or Pt 5$d$ emission, plus some U 5$d$ emission which is only partly suppressed by the resonance. The PES spectra labeled U 5$f$ result from subtraction of the $h\nu = 92$-eV spectrum from that for $h\nu = 98$ eV, a photon energy for which the U 5$f$ emission is resonantly enhanced. For UPt$_3$ the two spectra were scaled to match below $-5$ eV before subtracting. The results are generally consistent with previous resonant and x-ray PES on UPt$_3$ and uv PES on UA$_2$. The portions above $E_F$ are BIS data taken with $h\nu = 1486.6$ eV, for which the U 5$f$ spectral weight is known to dominate. The BIS and U 5$f$ PES spectra have been scaled to reproduce the $n_f$ values, $\approx 2.5$ for both materials, obtained from the density-functional calculations, which are expected to be accurate for this average ground-state property. The PES spectra are similar to one another and to those of UIr$_3$B$_2$ and UBe$_{13}$ while the two BIS spectra differ from one another in how many peaks occur above $E_F$. They also differ from that reported for UBe$_{13}$ in that both show very clearly, in addition to the slope change marking the onset of $E_F$, a second slope change which occurs after a distinct plateau at $\approx 0.5$ eV, and is followed by a very sharp rise to the peak at $\approx 1$ eV.

For comparison with our data, Figs. 1 and 2 also show the results of recent density-functional calculations of the electronic structure of UA$_2$ and UPt$_3$. From partial densities of states, one finds for UA$_2$ that the U $f$ weight is almost entirely confined between $-1$ and 2 eV, the U $d$ weight occurs in this region and extends further below $E_F$ to $-3$ eV, and the weight between $-3$ and $-10$ eV is of Al $s$-$p$ origin. For UPt$_3$, the U-site weight from $-0.5$ to 1.5 eV is dominated by $f$ states, the weight from $-0.5$ to $-4.5$ is

![FIG. 1. Top: PES and BIS spectra for UA$_2$, as described in the text. Bottom: local density-functional density of states from Ref. 14.](image1)

![FIG. 2. Top: PES and BIS spectra for UPt$_3$, as described in the text. Bottom: local density-functional density of states from Ref. 15.](image2)
about equally $d$ and $f$ states, and the weight between $-4.5$ and $-7$ eV is dominantly $d$ states. The Pt-site weight is dominantly $d$ everywhere, and relative to Pt metal, the empty and filled Pt $5d$ states are split off above and below $E_F$ to reduce greatly the Pt weight at $E_F$. There is substantial U-Pt and U-Al hybridization. A sharp contrast with the calculated $f$ bands of uranium metal is provided by the sharpness of the various subbands, the clear spin-orbit separation into spin-$\pm \frac{7}{2}$ and $\pm \frac{5}{2}$ groups, and the sharp cutoffs above $E_F$ at 1.5 and 2 eV for the two materials. It is an important point that the theoretical curves extend above $E_F$ far enough to show all the $5f$ structure.

Many features of the density-functional results are seen in the experimental spectra. For UPt$_3$ the $h\nu = 92$-eV spectrum shows a small Pt weight at $E_F$, and the calculated width, if not the detailed shape, of the Pt states. U $5f$ weight appears at the top of the Pt $5d$ band, a predicted consequence of U-Pt hybridization. For UAl$_2$ the $h\nu = 92$-eV spectrum shows the calculated Al $5p$ states, including the dip at $-5$ eV, and the U $d$ states nearer $E_F$. The spin-orbit splitting of the U $5f$ bands suggests a spin-orbit origin of the 0.5-eV shoulder and 1-eV peak found in the BIS spectra of both materials.

The most striking and informative feature of Figs. 1 and 2 is that the measured widths of the $5f$ spectral weights greatly exceed the one-electron ones. Above $E_F$ the BIS spectra show very much weight over an energy range as great as 6 eV above the $5f$ cutoffs, and below the $E_F$ the PES spectra show much more weight in the $-1$ to $-2$-eV region than is predicted or can be explained by the experimental resolution. Indeed, recent high-resolution PES spectra$^{21}$ of UBe$_{13}$ and UPt$_3$, which show much sharpening of the $5f$ structure, nonetheless have much $5f$ weight between $-1$ and $-2$ eV.

Although direct comparison to our spectra implies that the one-electron $5f$ widths are too small, comparison of calculated and experimental $\gamma$ values suggests$^{14,15}$ that the one-electron $5f$ widths are instead too large, in that the experimental $\gamma$ values and the relation $\gamma = \pi^2 k_F^2 N(E_F)/3$ imply values of $N(E_F)$, the density of states at $E_F$, which are much larger than calculated, in spite of the sharpness of the subband structure. As described below, this situation occurs for the Anderson Hamiltonian because of $U_{ff}$, which provides the basic mechanism for enhancing the $\gamma$ values so dramatically and at the same time increases the total spreading of the $5f$ weight by displacing energetically unfavorable $5f$ valence states away from $E_F$. From Figs. 1 and 2 a conservative lower limit on $U_{ff}$ is $\sim 1.5$ eV, consistent with previous theoretical$^{22}$ estimates, $U_{ff} \sim 2$ eV, for metallic uranium.

Given the large U-U separations (3.38 and 4.1 Å for UAl$_2$ and UPt$_3$, respectively) the $N_f$-fold degenerate Anderson Hamiltonian is most widely used to model the interplay of one-electron and Coulomb interactions in these materials. Theories for the lattice case are still in progress,$^{3,23}$ but for an impurity$^{24,25}$ with the local orbital occupation $n_f < 1$, and in the limit of large degeneracy and Coulomb interaction, the situation described qualitatively above is well understood quantitatively, as follows. For $U_{ff} = 0$ the Green's-function spectral weight $\zeta(E)$ is a single Lorentzian of degeneracy $N_f$ and one-electron hybridization width $\Delta$, centered above the Fermi level such that $\zeta(E_F) = \pi n_f^2/\Delta$. This is the impurity analog, with neglect of spin-orbit coupling, of the density-functional results. For the same value of $n_f$, but with $U_{ff}$ large, $\zeta(E)$ spreads over a range $\sim U_{ff}$, leaving around $E_F$ only the Kondo resonance, having a reduced width approximating the Kondo temperature $\delta << \Delta$. A remarkable consequence$^{26}$ of the Friedel sum rule is that $\zeta(E_F)$ has the same value, for fixed $n_f$, as for $U_{ff} = 0$. The usual expression above for $\gamma$ continues to hold, if $N(E_F)$ is taken as the quasiparticle density of states, given by $N(E_F) = Z^{-1} \zeta(E_F)$.

This width $\Delta$ is too large to account for the enhanced specific heat, but is too small to account for the total $f$ width. The enhanced $\gamma$ value is reflected in the PES-BIS spectrum only by the small width $\delta$ of features near $E_F$, but not by a large magnitude of $\zeta(E_F)$, contrary to commonly stated$^{17,18,28}$ expectations. Typically the BIS-PES experimental resolution is inadequate to measure $\delta$ directly so that $N(E_F)$ can be obtained from BIS-PES spectra only by use of theoretical relations$^{24}$ between $\delta$ and the large-energy-scale features of the spectrum, such as $U_{ff}$.

For cerium materials, the theoretical results for the impurity Anderson Hamiltonian have led to a unified interpretation$^{5,24,29,30}$ of much of the large-energy-scale spectroscopic and low-energy-scale transport data, apart from lattice-coherence effects. A major barrier, that$^5,18$ measured BIS-PES $4f$ spectra often have more weight near $E_F$ than expected from the Kondo resonance, has been removed by very recent theories which find that this weight arises from spin-orbit sidebands$^{30,31}$ of the Kondo resonance, and from $f^2$ valence states present for finite $U_{ff}$. These results for cerium are encouraging that the substantial weight near $E_F$ in our uranium spectra can also be accounted for with the Anderson Hamiltonian. For example, the expectation of spin-orbit sidebands provides an explanation of the spectral weight occurring above and below $E_F$ at the uranium spin-orbit energy. More generally, the sideband spectrum around $E_F$, as well as the relation between high and low energy scales, should reflect the details of the quantum struc-
ture of the relevant valence states, and so a final judgment must await explicit calculations in the regime $n_f \sim 2.5$, for which the relevant valence states, $n_f = 2, 3$, and 4, have a quantum structure richer than just spin-orbit splitting.

To summarize, the one-electron $5f$ bandwidths for UA$_2$ and UP$_3$ are too large to account for the large $\gamma$ values and too small to account for the measured BIS-PES widths, precisely the situation that occurs in the Anderson Hamiltonian as a result of $U_{jj}$. Success with cerium hopes that the situation can be modeled quantitatively by the Anderson Hamiltonian when the theory is extended to treat larger numbers of $f$ electrons.

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15P. Strange and B. L. Gyorgy, in Proceedings of the Conference on Electronic Structure and Properties of Rare Earth and Actinide Intermetallics, St. Pölten, Austria, 3–6 September 1984 (unpublished). The authors of Ref. 8 have also calculated for UP$_3$.


23The existence of a Kondo resonance [R. M. Martin, Phys. Rev. Lett. 48, 362 (1982)] and the form of the relation between $\gamma$, $N(E_F)$, and $\xi(E_F)$ discussed in the text are required also for the lattice case by general results of J. M. Luttinger [Phys. Rev. 119, 1153 (1961)].


27It can even occur that material $A$ having a larger $\gamma$ than that of material $B$ may have a smaller $\xi(E_F)$ with a suitably larger $Z^{-1}$.


