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Understanding mixed valent materials: Effects of dynamical core-hole screening in high-pressure x-ray spectroscopy

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Changes in the electronic structure of Yb, a material whose valence is modified under pressure, are observed with remarkable detail in x-ray absorption and emission data measured between ambient conditions and 20 GPa. These changes are reproduced by a theory that essentially does not rely on experimental parameters, and includes dynamical core-hole screening. From the combined experimental and theoretical data we can firmly establish on a quantitative level how the valency of an intermediate valence material is modified by pressure. In metallic Yb it increases from 2 to 2.55±0.05 between 0 and 20 GPa.

A crucial aspect of materials with incomplete 4f or 5f shells is the valency. It is the central parameter in understanding heavy-fermion materials, anomalous superconductors, and obviously intermediate valent (IV) materials. In the series of the rare-earth metals, Yb and Eu show a number of anomalous properties (like large equilibrium volume) demonstrating their divalent ground state, in contrast to the prevailing trivalent behavior of the rest of this group of elements. The valency in, e.g., metallic Yb reflects a balance between the energy required to excite a 4f electron into the valence band from the 4f14(sd)2 (Yb2+) ground state, and the larger cohesive energy of the 4f13(sd)3 (Yb3+) configuration. This balance may be altered by applying external pressure.

The relative stability of the Yb2+ and Yb3+ ionic configurations, and the possibility of a valence transition under pressure, were already discussed in Ref. 3. More recently, it has been shown that the equation of state of Yb under high pressure can be described very accurately by an ab initio calculation for the IV state, the Coulomb interaction between 4f electrons and 5d holes being of crucial importance. From the experimental side, the anomalous compressibility of Yb, suggesting a continuous valence transition to a fully trivalent state above 100 GPa, has turned out to be difficult to experimentally verify this on a microscopic scale.

The Yb valence is more directly probed by x-ray absorption spectroscopy (XAS) at the Yb L3 edge (2p63d94fN ed +hvin→2p64fN ed+hvout) is fixed, corresponding to “high-resolution” partial fluorescence yield XAS (PFY XAS) or Lα1 resonant inelastic x-ray scattering (RIXS) spectra, respectively. RIXS is especially powerful because the contribution of each electronic configuration in the ground state (4f13 or 4f14 for Yb) can be selectively enhanced by tuning hvout to specific energies along the absorption edge. We performed high-pressure experiments at beamline ID16 of the European Synchrotron Radiation Facility (Grenoble, France), which provides a focused (130×50 μm2) monochromatic x-ray beam at the sample position. The sample, a 25-μm-thick ytterbium foil (from Goodfellow), loaded in a diamond anvil cell, using high-strength Be as gasketing material and ethanol as pressure-transmitting medium. Pressure could be measured in situ by the conventional ruby fluorescence technique. Both the incident and the emitted radiation passed through the Be gasket in the horizontal scattering plane, to avoid strong absorption by the diamonds. X rays emitted at 90° from the incident beam were analyzed using the (620) Bragg reflection of a 1-mm-radius spherically bent silicon crystal, and then detected by a Peltier-cooled Si diode. We measured PFY XAS spectra by scanning the incident energy through the Yb L3 edge (hvin≈8.95 keV) while recording the intensity of the emitted characteristic Yb Lα1 fluorescence (hvp=7.415 keV). RIXS spectra were recorded for selected values of hvin across the L3 edge. The overall energy resolution was ~1.5 eV. Depending on the incident photon energy the acquisition time per RIXS spectrum varied between ~15 and 45 mins.

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FIG. 1. \( L_{\alpha 1} \) PFY XAS spectra of Yb at selected pressures.

PFY XAS data measured at pressures between 0 and 20 GPa are shown in Fig. 1. The broad line shape at ambient pressure is indicative of underlying structure that could not be appreciated in lower-resolution data. With increasing pressure, spectral weight is transferred from threshold to a new peak \( B \) at \( \sim 10 \) eV higher energy. Moreover, an intermediate peak \( A \) emerges from the broad threshold, and progressively drifts to higher energy. Changes are already quite visible at 4 GPa, the boundary between the Yb fcc and bcc phases. For a pure Yb\( ^{2+} \) or Yb\( ^{3+} \) ground state, due to dipole selection rules, the Yb L\( _{3} \) XAS spectrum is proportional to an unoccupied density of states (DOS) of \( d \) symmetry, modulated by matrix elements and distorted by the attractive core-hole potential. In IV Yb compounds, the line shape is interpreted as the superposition of two replicas of the \( d \) DOS, weighted by the ground state weights \( w(2+) \) and \( w(3+) \) of the Yb\( ^{2+} \) and Yb\( ^{3+} \) configurations, and offset by several eV’s. The replicas correspond to \( 2p^{5}4f^{2}3\text{ed} \) and \( 2p^{5}4f^{3}3\text{ed} \) final states, and their energy separation reflects the large Coulomb repulsion between the \( 2p \) and \( 4f \) holes. Therefore the observed spectral weight transfer indicates an increasing Yb\( ^{3+} \) weight in the hybrid ground state, in agreement with Ref. 10. The added element in the data of Fig. 1 is feature \( A \), which changes with pressure and suggests that the \( d \) DOS is modified as a function of this parameter. A closer analysis (see below) substantiates this interpretation.

Figure 2(a) shows the series of RIXS spectra measured through the L\( _{3} \) edge at 20 GPa. They are plotted as a function of the energy transfer \( E_T = h\nu_{in} - h\nu_{out} \). Spectral features whose energy tracks with \( h\nu_{in} \) in the Raman regime, appear at constant energy transfer. The Raman feature final states at \( \sim 1.526 \) and \( \sim 1.536 \) keV (indicated by the arrows) originate from the Yb\( ^{2+} \) and Yb\( ^{3+} \) ground state configurations. The dispersive peak (marked by the asterisk) observed at larger incident energies is the characteristic \( L_{\alpha 1} \) fluorescence, whose intensity follows the XAS profile. The 20 GPa RIXS spectra for all incident energies can be consistently decomposed, following the guidelines described in Ref. 12, to yield the intensity profiles of Yb\( ^{2+} \) and Yb\( ^{3+} \) and of the \( L_{\alpha 1} \) fluorescence [Fig. 2(b)]. Their energy dependence enables a qualitative interpretation of the PFY XAS spectra of Fig. 1: the Yb\( ^{2+} \) signal dominates near threshold (and peaks at \( h\nu_{in} = 8.942 \) keV), while the Yb\( ^{3+} \) intensity peaks at \( \sim 10 \) eV higher energy. The decomposition of the 4 and 10 GPa RIXS data leads to the same conclusion. At 0 GPa the spectra were fitted only with \( 2+ \) and fluorescence signals. Changes in the RIXS line shape with pressure are illustrated in Fig. 3(a) for the representative value \( h\nu_{in} = 8.942 \) keV, at the Yb\( ^{2+} \) resonance. We decomposed the 20 GPa spectrum into separate Yb\( ^{2+} \) and Yb\( ^{3+} \) contributions. The former is a rescaled replica of the ambient pressure line shape.

The Yb valence can be deduced at all pressures from the relative intensities of the Yb\( ^{2+} \) and Yb\( ^{3+} \) RIXS components, as \( I(\nu_{in}) = w(2+) + w(3+) \) (Fig. 2(b)) shows that the Yb valence increases from 2 to 2.55±0.05 between ambient pressure and 20 GPa. The error bars account for statistical errors, for corrections associated with the energy dependence of the penetration length, and for uncertainties in the precise resonance energy.

A theoretical description of the observed spectroscopic changes must go beyond standard ground state electronic structure theory, which is based on the local density approximation (or gradient corrected functionals): these calculations do not always reproduce well experimental data, which arise from a combination of the initial and final densities of states. The method presented here accounts for the superposition of initial and final electronic states and applies to many complex systems. In general, dynamical screening of the core hole created in the excitation process is expected to...
play an important role. A solution to this arduous theoretical problem may be obtained by the first-principles multiband version\textsuperscript{16} of the Mahan–Nozières–De Dominicis (MND) theory.\textsuperscript{17–19} A further complication arises for mixed valent materials, namely, that the spectral weights of the divalent and trivalent contributions must be evaluated. We have solved this part of the problem by calculating, for each pressure, the ground state weights $w(2+)$ and $w(3+)$ using the mean-field description of the Kimball–Falicov model, where the $d$-$f$ Coulomb interaction plays an important role.\textsuperscript{20} These weights are compared to experiment in Fig. 3(b), and we note that the agreement between theory and experiment is very good. We remark that all parameters in the Kimball–Falicov model (the energy difference between di- and trivalent configurations, the Falicov interaction parameter, the renormalized Fermi energy, and the $spd$-$f$ hybridization matrix element) were calculated from first principles. A relative offset of 10 eV was applied to the Green’s functions to reproduce the energy separation of the Yb$^{2+}$ and Yb$^{3+}$ core level binding energies (the calculated value for this energy splitting was 8.2 eV).

We then computed a complete solution of the dynamical screening problem\textsuperscript{16} for Yb$^{2+}$ and Yb$^{3+}$, where the initial and final state Green’s functions play a crucial role. The XA spectrum of the mixed valence compound was obtained by multiplying the Yb$^{2+}$ and Yb$^{3+}$ spectra with the theoretical weights $w(2+)$ and $w(3+)$. The first-principles calculations were done using the plane-wave VASP code\textsuperscript{23,24} within the projector augmented wave\textsuperscript{25} method. The exchange-correlation energy was described by the Ceperley and Alder functional\textsuperscript{16} parametrized by Perdew and Zunger.\textsuperscript{27} The mixed valency of Yb was modeled using supercells containing Yb atoms with three and two valence electrons. For the calculation at 10 GPa a cell with 10 Yb$^{2+}$ and 6 Yb$^{3+}$ atoms was used, to simulate a valency close to that determined by the Kimball–Falicov model.\textsuperscript{20} For the same reason, in the calculation at 20 GPa a cell with seven Yb$^{2+}$ and nine Yb$^{3+}$ was used. The Green’s functions were calculated for the Yb$^{2+}$ and Yb$^{3+}$ atoms with a set of neighboring atoms representing the mixed valent alloy.

FIG. 3. (a) $\sigma_\alpha$ RIXS spectra ($h\nu_{\text{ex}}$=8.942 keV) of Yb measured at ambient pressure (filled connected dots) and 20 GPa (hollow dots). The thick continuous line at 20 GPa is the sum of a scaled replica of the 0 GPa line shape representing the Yb$^{2+}$ component (light gray area) and of an Yb$^{3+}$ component (dark gray area). The latter is obtained as the difference between the measured spectrum and the Yb$^{2+}$ component. (b) Values of the Yb valency determined from the RIXS data (filled dots) and from our first-principles calculation (white dots). Experimental and theoretical values coincide at 0 GPa.

FIG. 4. Comparison of experimental PFY XAS spectra of metallic Yb at selected pressures (dotted lines) with calculation: shaded curves are the unbroadened Yb$^{2+}$ (light gray) and Yb$^{3+}$ (dark gray) XAS spectra calculated by the MND theory as described in the text. Thick lines represent the broadened sum of the latter curves. Also shown for comparison are the initial state local density approximation density of states (thin lines).
The Monkhorst-Pack scheme\textsuperscript{28} was applied with $5 \times 5 \times 5$ and $3 \times 3 \times 3$ $k$-point meshes, and a kinetic energy cutoff at $-200$ eV was used for the plane-wave basis set. In order to simulate a system with a core excitation we adopted the $Z+1$ approximation for the excited atom of our supercells. Finally, we added a Lorentzian broadening, whose full width at half maximum (FWHM) was taken to increase linearly above threshold to account for the final state lifetime:\textsuperscript{24} \( \Delta E(\text{FWHM}) = 0.6 + 0.15(E - E_{\text{threshold}}) \) eV. In order to compare the theoretical spectra with experiment, we introduced a Gaussian experimental broadening \( \Delta E(\text{FWHM}) = 1.5 \) eV. No phenomenological background was added to the calculated spectra.

A comparison of experimental and calculated spectra for 0, 10, and 20 GPa (Fig. 4) reveals an overall good agreement. The calculated features above 10 eV from the Fermi edge are partially obscured by the large spectral broadening and by the experimental background (not included in the calculation). The comparison provides an interpretation of the observed energy shift of feature A: it reflects the increasing separation with pressure between the center of mass of the unoccupied $d$ DOS of the $2^+ \rangle$ configuration and the Fermi energy. The reason for this is an increasing overlap between the wave functions of the $d$ orbitals with increasing pressure and decreasing lattice constant. The result is a broadening of the $d$ band and a corresponding shift of the spectral features. Also shown in Fig. 4 is the initial state local density approximation DOS, showing modestly good agreement with the data. The final state DOS (not shown here) is quite similar to the MND data. It should be noted that this finding is far from being a standard result of MND theory since the core-hole valence-electron interaction is known in many cases to produce an “edge singularity,” with an associated increase of the XAS intensity at the threshold energy: in a recent work\textsuperscript{16} it was shown that the relative intensities between various features of the XAS spectrum of graphite were significantly different in the MND calculation compared to a final state calculation.

The development in experimental capabilities in combination with advancements in the theoretical description of mixed valent materials is shown here to yield a remarkably detailed description of ground and excited state properties. The most noticeable property is the valency, recently discussed, e.g., by Strange \textit{et al.}\textsuperscript{22} that we now can monitor at a very detailed level as a function of pressure. Our theory reproduces the observations quite accurately. The excited state properties are revealed by the PFY XAS signal and also here theory and experiment agree well. The combined PFY XAS and RIXS data, and the comparison with theory, yield very detailed information on how the valency of a material changes with applied pressure. The method is an important development for our understanding of mixed valent materials in general.

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