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Development of the Gd(0001) band structure with film thickness

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By growing gadolinium films on W(110), it has been possible to observe the development of the Gd(0001) band structure with increasing film thickness. The changes in the 5d bands as a result of exposure to oxygen or the development of the valence-band structure with increasing film thickness lead to changes in the 5p multiplet oscillator strengths. These changes are a result of the influence of the band structure upon the final-state effects that give rise to such multiplet structures. This study provides conclusive evidence that there is a 5d contribution to the shallow 5p multiplet photoemission core-level structure.

I. INTRODUCTION

Both theoretical and experimental studies of the gadolinium band structure have been undertaken. Recently, a Gd(0001) magnetic surface state has been predicted and observed. In this paper we show that this surface state and the bulk band structure can only be distinguished for the relatively thicker films as a result of the development of the Gd(0001) band structure. Because the 5d bands play a significant role in photoemission from the 5p shallow core levels as a result of the final-state coupling, the development of the 5d band structure is expected to affect the 5p multiplets. Changes in the 5p multiplet structure with increasing film thickness can indeed be observed as are expected.

By depositing gadolinium on W(110), the development of the gadolinium band structure with increasing film thickness can be characterized. Films deposited on W(110) have been studied by a number of groups and several studies have established that the films grow and adopt the crystal structure of bulk gadolinium with the (0001) surface parallel with the W(110) surface (i.e., the c axis grows along the surface normal). This property of the gadolinium films grown on W(110) makes this rare-earth overlayer system particularly useful for investigating the development of the electronic band structure with increasing film thickness for crystalline gadolinium without the contamination problems common to the rare earths.

II. EXPERIMENT

The gadolinium was deposited on clean W(110) as has been described previously for gadolinium deposition onto other substrates. The crystallinity and quality of the films deposited on W(110) was carefully characterized by low-energy electron diffraction (LEED). The thickness of the gadolinium films was ascertained from a quartz-crystal oscillator thin-film monitor (Leybold-Inficon) but the film thicknesses cited in this work are approximate, as noted elsewhere. All binding energies cited in this work are referenced to the Fermi energy of the clean W(110) surface. Throughout this work, the light incidence angle was 70° so that the vector potential is largely normal to the surface (p-polarized light) unless otherwise noted. In this geometry, the surface state is greatly enhanced as discussed in detail elsewhere. The photoelectrons were collected normal to the surface (i.e., k∥ = 0 or Γγ) into hemispherical electron energy analyzers described elsewhere, with angular resolutions of ±1.5°. The light sources for this photoemission study were the NSLS vuv ring and the 1-GeV (Alladin) ring at the Synchrotron Radiation Center with the light dispersed in each case by a toroidal grating monochromator. The combined energy resolution (analyzer and monochromator) of the photoemission spectra varied from 0.15 to 0.2 eV full width at half-maximum for the photoenergies taken between 30 and 50 eV.

III. RESULTS AND DISCUSSION

With increasing amounts of gadolinium deposited upon W(110), the gadolinium features quickly become the most prominent features in the energy distribution curves as seen in Fig. 1. For the very thin gadolinium films, there are two prominent features: the 4f level at a binding energy of 8.6 eV and the 5d bands near the Fermi energy. For photoemission in the normal direction (k∥ = 0 or Γγ), the gadolinium bands become resolved into at least two distinct features with increasing coverage as seen in Fig. 1. With a 30-Å-thick gadolinium film, the feature near the Fermi energy has become quite narrow with a half-width of 300-400 meV. There is another 5d band feature at a somewhat larger binding energy of 1–2 eV binding energy. This feature becomes increasingly intense with increasing gadolinium film thickness. This higher-binding-energy feature has a binding energy of 1.5 eV by 30 Å, but has a half-width of some 800 meV or more.
FIG. 1. Photoemission spectra of gadolinium overlayers on W(110). The photoelectrons were collected normal to the surface and the light is $p$ polarized so as to increase the relative signal of the surface state relative to the other bands.

The state at 0.2 eV falls into a gap in the projected band structure at $\Gamma$, based upon calculated\textsuperscript{1-6,8-9} and experimentally derived\textsuperscript{11} band structures for bulk gadolinium. This state shows conservation of two dimensionality of state, is independent of film thickness, and is sensitive to the adsorption of oxygen and other contamination, as seen in Fig. 2. Since this state also has a very narrow full width at half-maximum, this state has been identified as a surface state.\textsuperscript{10} This state is predicted by a recently calculated band structure,\textsuperscript{9} which characterizes this state as a minority-spin magnetic surface state. This surface state does not fall into a gap of the projected tungsten density of states at $\Gamma$ and hybridization with the W(110) surface is possible, thus one cannot anticipate that this state will be as narrow for the thin films as one would expect for the thicker films, as is observed.

Another gadolinium band is increasingly distinct as a result of an increasing binding energy with increasing gadolinium coverage as seen in Fig. 1. There is increasing similarity of the bulk band-binding energies and oscillator strengths for the gadolinium overlayers on W(110) and the gadolinium 5$d$ bulk bands\textsuperscript{11} with increasing film thickness. These gadolinium-induced states at 1–2 eV below the Fermi energy have a binding energy that is sensitive to changes in photon energy, even for gadolinium films of 10 Å. Sensitivity to changes in $k_z$ (changes in photon energy) is characteristic of bulk bands, and the increasing resemblance of these states 1–2 eV below $E_F$ to the bulk band structure at $\Gamma$ (Ref. 11) with increasing film thickness is strongly indicative that these features are representative of the bulk band structure. It is important to realize that for the thinnest of gadolinium films, the lattice constant differs slightly from that of bulk gadolinium.\textsuperscript{16-18} This together with the additional complications of the two dimensionality of the film and hybridization of the gadolinium overlayer with the W(110) substrate results in the differences between the bulk band structure observed for the thicker films or by Himpsel and Reihl,\textsuperscript{11} and the band structure observed for the very thin gadolinium films.

With strong spin-orbit coupling the 5$p$ level separates into two $5p_{1/2}$- and $5p_{3/2}$-based levels as has been observed at 28 and 21 eV, respectively (Refs. 12, 13, 27, and the...
FIG. 3. Photoemission spectra across the gadolinium 5p envelope for a 10-, 15-, and 30-Å-thick gadolinium film on W(110). Note the change in the relative multiplet oscillator strengths. This is particularly apparent after the photoemission background is removed. The inset shows the effect of one 1-L oxygen exposure on the 15-Å-thick gadolinium film. The photon energy was 50 eV.

references therein). Fine structure can be observed within the 5p levels, particularly within the 5p_{3/2} envelope. We have recently identified this fine structure as a multiplet structure which is a result of J-f coupling in the final state.\(^{27}\) The relatively weaker exchange interaction of the 5p hole with the 4f and 5d electrons of gadolinium will lift the J degeneracy. Neglecting the 5d electrons for simplicity, the \(f = \frac{3}{2}\) term of the hole (in the 5p_{3/2} envelope) couples with the \(5s^{1/2} 4f\) levels (again neglecting the 5d electrons) to form four terms of \(\left(\frac{5}{3}\right)_{5}, \left(\frac{5}{3}\right)_{4}, \left(\frac{5}{3}\right)_{3}, \) and \(\left(\frac{5}{3}\right)_{2}\) (Ref. 27) as is observed at binding energies of 20.3 \(\pm 0.1\), 20.9 \(\pm 0.1\), 22.0 \(\pm 0.2\), and 23.4 \(\pm 0.3\) eV, respectively. Similar results, though not as pronounced, have also been obtained for terbium.

In Fig. 3, we present the 5p_{3/2} photoemission feature as a function of gadolinium film thickness. There is a clear indication that the \(\left(\frac{5}{3}\right)_{3}\) multiplet increases in oscillator strength relative to the other multiplets within the 5p_{3/2} envelope with increasing gadolinium film thickness. Furthermore, with oxygen adsorption there is a pronounced change in the relative 5p multiplet oscillator strengths as seen in the inset to Fig. 3. This can be correlated with the changes in the 5d band structure as seen in Fig. 2.

The results in Fig. 3 indicate that the 5p 5d \(\left(\frac{5}{3}\right)_{3}\) multiplet member, where \(j\) for the 5p_{3/2} hole is parallel to the 4f-derived J, is dramatically affected by oxidation and Gd film thickness. Oxidation destroys the gadolinium surface state (as seen in Fig. 2), while increasing film thickness leads to formation of a distinct surface state at \(E_{\text{F}}. \)\(^{10}\) We do not have conclusive data to establish whether the Gd \(\left(\frac{5}{3}\right)_{2}\) multiplet couples magnetically to the surface or bulk Gd 5d. We nonetheless postulate that the Gd 5p to Gd 5d final-state coupling (leading to the multiplet fine structure) is strongly influenced by the Gd 5p surface-state magnetic ordering. Clearly there is a change in the 5d band structure with increasing gadolinium film thickness while the occupied 4f levels at 8.6 eV binding energy are highly localized and exhibit little dependence upon film thickness as seen in Fig. 1. These results strongly support the postulate proposed elsewhere\(^{12-15}\) that photoemission from the 5p levels is strongly influenced by final-state interactions with the 5d levels, instead of solely a result of final-state interactions of the 4f\(^{2}\) electrons with the 5p hole.

In summary, we find that there are dramatic changes in the 5d band structure with increasing film thickness. Some of these changes are associated with the development of a bulk band structure and a well-defined gadolinium surface state. These changes in the electronic band structure of gadolinium with increasing film thickness on W(110) also result in surprising changes to the photoemission features from the gadolinium core levels as a result of final-state coupling effects with the 5d bands.

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