The Magnetic Structure of GdAgSb$_2$ Determined by X-ray Resonant Exchange Scattering

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Introduction

The recent availability of high-quality single-crystal samples of RA$g$Sb$_2$ ($R$ = rare earth ion) compounds has stimulated new interest in the highly anisotropic electronic and magnetic properties of these materials and their manifestation in various physical phenomena [1, 2].

Most of the RA$g$Sb$_2$ compounds display long-range antiferromagnetic ordering at low temperature [3]. The reported transition temperatures for the heavy rare earth compounds, however, there are significant deviations from de Gennes scaling that are claimed to be free of CEF effects, a determination of the magnetic structure is needed to resolve issues regarding the role of CEF effects on magnetic ordering in this family of compounds. The extensive neutron scattering experiments on powder samples of RA$g$Sb$_2$ compounds reported by André et al. [4], however, did not include the Gd-compound, presumably because of the neutron opacity of the naturally abundant Gd ion.

Methods and Materials

Single crystals of GdAgSb$_2$ were grown by using a high-temperature flux technique at Ames Laboratory [5]. The resulting crystals are shaped as platelets, with the flat surface perpendicular to the crystallographic $c$ direction (the [001] unique axis of the tetragonal structure). Further polishing of the sample surface was not possible without degrading the sample quality (increasing the sample mosaicity) because of its softness. For the magnetic x-ray diffraction experiment, the cleanest sample, without noticeable flux inclusions from the growth process, was chosen. For this sample, the mosaic was measured to be $0.015^\circ$ at the (008) reflection at 17 keV.

The resonant x-ray scattering experiments were carried out on in the Midwest Universities Collaborative Access Team (MU-CAT) 6-ID undulator beamline at sector 6 of the APS by using a double-crystal Si(111) monochromator. The energy profile of the primary beam (predominantly $\pi$-polarized component to pass) was measured to be $\pi$-polarized component to pass as well.

At low temperature, in the purely $\pi$-polarized channel, a resonant scattering signal was observed at positions corresponding to a magnetic wave vector of $\tau = (0\bar{1}0)$, whereas no signal was observed on the $\tau = (\bar{1}00)$ positions. However, after changing the scattering angle of the analyzer crystal to make use of the PG (002) reflection, allowing both $\sigma$ and $\pi$ scattering to pass, magnetic peaks were observed at positions corresponding to $\tau = (\bar{1}\bar{1}0)$. These results indicate that the ordered moment direction in the antiferromagnetic state is oriented perpendicular to the magnetic wave vector. The energy profiles depicted in Figure 1 show both the dipolar origin of the scattering at the (0-128) reflection, arising from the strong interaction between the core hole in the 2p$_{3/2}$ orbital and the 5d band, and the quadrupolar origin of the scattering at the (1-128) reflection, arising from the weaker interaction between the 2p$_{3/2}$ core hole and the strongly localized 4f band [6, 7].

To confirm that the resonant enhancement at the (1-128) magnetic peak arises from electric quadrupole transitions with the moment transverse to the magnetic wave vector, Q-dependent integrated intensity measurements were performed by using those magnetic peaks (1-128) that could
FIG. 1. Integrated intensity of the (½08) (upper panel) and (0-½8) (middle panel) reflections of GdAgSb$_2$ as a function of energy across the Gd L$_3$ edge in the low-temperature magnetic phase. The background fluorescence from the sample is shown in the lower panel.

be accessed in the symmetric scattering geometry. The integrated intensity was measured with open detector slits by using the PG(002) reflection from the analyzer crystal. The rather broad mosaic of the analyzer, ~0.5°, is significantly larger than the full width at half maximum (FWHM) of magnetic satellites of ~0.1°, ensuring adequate integration of the scattered intensity. Proper integration over 2θ and χ was achieved by opening the detector slits until a flat top was observed in 2θ and χ scans through the magnetic peaks. The integrated intensities of several peaks that could be accessed in this geometry were calculated from rocking scans of the sample crystal through the magnetic reflections. The observed integrated intensities of the magnetic satellite are plotted in Fig. 2. The solid line represents a fit to the theoretical quadrupole resonance scattering cross section with the moment direction along [010], showing reasonable agreement with the data.

The temperature dependence of the integrated intensity (I ≈ μ$^2$) of the (½08) magnetic satellite peak at the L$_3$ resonance energy is displayed in Fig. 3. The solid line is a fit to the power law, I = A(T$_N$ - T)$^β$. A transition temperature T$_N$ = 13.0K obtained from the fit is close to the value T$_N$ = 12.8K obtained from the bulk susceptibility measurement [1].

**Discussion**

The principal result of this work is the ab initio determination of the magnetic structure of GdAgSb$_2$ by x-ray resonance exchange scattering. Below 13K, the local magnetic moments order antiferromagnetically, doubling the unit cell along the a direction (or equivalently in the tetragonal structure b direction) in the basal plane, with the moments ordered transverse to this direction also being within the basal plane. These results indicate that the commensurate antiferromagnetic ordering observed in several of the RAgSb$_2$ compounds is not necessarily driven by the strong CEF effects as proposed by André et al. Indeed, it should be noted that in the related RNi$_2$B$_2$C family, the Gd compound, as well as many of the strongly anisotropic member compounds (R = Er, Tb, Ho), order in an incommensurate structure determined largely by Fermi surface nesting [8, 9].

The interactions that conspire to produce magnetic ordering in many of these systems are quite complex, involving RKKY-type indirect exchange, CEF anisotropies, and, especially for the light rare earths, hybridization effects and possible changes in the indirect exchange coupling. A great deal of further study is clearly necessary to unravel the contributions of all of these mechanisms.
FIG. 3. Temperature dependence of the integrated intensity of the (ν08) magnetic satellite of GdAgSb₂. The solid line is a fit to a power law as described in the text.

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References