Direct evidence of ferromagnetism without net magnetization observed by x-ray magnetic circular dichroism


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We have performed x-ray magnetic circular dichroism experiments to study the cancellation of spin and orbital magnetic moments in (Sm,Gd)Al₂, a ferromagnet without net magnetization at a certain compensation temperature, Tcomp. We verified the existence of long-range order for both spin and orbital magnetic moments at Tcomp. The spin and orbital magnetic moments of the Sm ion are found always antiparallel coupled and the magnitude of its orbital magnetic moment is always larger than that of spin one, so the cancellation of magnetic moments cannot be achieved by only Sm 4f electrons. We show that the addition of spin magnetic moments of Gd ions and conduction electrons, which are ferromagnetically coupled with the spin magnetic moment of Sm ions, cancels out the surplus orbital magnetic moments in Sm ions completely and results in the zero magnetization at Tcomp. All our experimental results can be reproduced well by atomic multiplet calculations.

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I. INTRODUCTION

Samarium atoms in solids usually contribute three electrons to the conduction band and the atoms become Sm³⁺ with the 4f⁵ configuration. The spin and orbital magnetic moments (Mₕ and Mₑ) of the Sm³⁺ ground state, which can be decided from Hund’s rules as ⁹/₂ Hₕₙ/₂, tend to cancel each other out because of their similar magnitudes and antiparallel orientation owing to the spin-orbital interaction, and only a small surplus orbital magnetic moment is left. As a result, the contributions from conduction electrons and doped ions to the magnetism of samarium compounds become important. The tuning of the magnetic properties was demonstrated by Adachi and Ino such that the magnetization became zero at a certain compensation temperature, Tcomp, when some of the samarium atoms in SmAl₂ were substituted with the gadolinium ones. They suggest that (Sm,Gd)Al₂ is a ferromagnet (with long-range order for both Mₕ and Mₑ) without net magnetization, as the total spin and orbital magnetic moments, including the contributions from Gd ions and conduction electrons, cancel each other out at Tcomp. Their consideration is very direct and simple in that the electron configuration of Gd³⁺ ion is ⁴f⁷, that is to say half-filled, its total orbital angular momentum is zero and only a spin magnetic moment is left, leading to the possibility of canceling out the surplus orbital magnetic moment in Sm³⁺. In this work, we verify this mechanism experimentally.

Although the existence of ferromagnetic ordering of Mₕ at Tcomp (Ref. 6) and both Mₕ and Mₑ below and above Tcomp (Ref. 7) was proved, the ferromagnetic ordering of Mₑ at Tcomp has not been clearly observed, and no experimental measurement on the magnetic moments of dilute Gd ions has been reported. Therefore, further quantitative studies for the individual behavior of Mₕ and Mₑ inside Sm and Gd ions are indispensable for confirming the ferromagnetic property completely and for understanding the mechanism of the zero net magnetization. X-ray magnetic circular dichroism (XMCD) is most suitable for this task because we can evaluate Mₕ and Mₑ separately by sum rules in an element-selective (Sm or Gd) and electronic-state-selective (the state of ⁴f or conduction electrons) manner through the tuning of the photon energy. In this work, the Mₕ absorption edges are used, which give us information about the ⁴f electrons in Sm and Gd.

II. EXPERIMENT

The measurements were carried out at the beam line BL25SU of SPring-8 synchrotron radiation facility. A twin-helical undulator was adopted to generate left or right circularly polarized soft x rays with high (>99%) circular polarization. X-ray absorption spectra were measured in the total photoelectron yield mode. A magnetic field of 1.4 T was applied by permanent magnets normal to the sample surface. The XMCD spectrum is often obtained by taking the difference of absorption when reversing the sample magnetization. However for (Sm,Gd)Al₂, we have shown that this is not applicable near Tcomp because of the insensitivity of the sample to external magnetic field due to its tiny magnetization. To resolve this problem, the XMCD needs to be observed by the difference of absorption caused by the reversing of helicity of incident photons. Our previous work shows that the 1.4 T magnetic field is not enough to saturate the sample magnetization and the degree of magnetization is not the same when magnetized at different temperatures be-
cause the sample is a polycrystal and its net magnetization is small.9 We note that even for nonferromagnetic materials, small XMCD can still be observed under an external magnetic field. To eliminate this possibility and to maintain the magnetization of the sample unchanged,10 which is important for the studies on temperature dependence of magnetic moments, in this work, the sample was magnetized in advance at a certain temperature $T_m$ and the external magnetic field was removed before the measurements. We obtained the XMCD by taking the difference of two spectra in two successive scans. After the first scan the helicity of incident photons was reversed for the second scan. With this method, the accuracy of experimental data was not high enough to provide us with reliable information for understanding the mechanism of the cancellation of magnetic moments at $T_{\text{comp}}$11 The error is mainly caused by the time-dependent instability of the measurement system. If we can reverse the helicity of incident photons at every data point, this problem will be resolved. The twin helical undulator of SPring-8 BL25SU beam line has the capability of reversing the helicity of synchrotron radiation by the adjustment of local orbits in the straight section of the storage ring as fast as 10 Hz.12 In this work, the XMCD measurements were done by this helicity-switching method with 1 Hz frequency. The helicity was reversed at every photon energy and the XMCD was measured as $\mu^+ - \mu^-$, where $\mu^+$ and $\mu^-$ are the x-ray absorption coefficients with the helicity of the circular polarization parallel and antiparallel to the direction of the fixed magnetization orientation. The sample studied here is a Sm$_{0.982}$Gd$_{0.018}$Al$_2$ polycrystal ($T_{\text{comp}}=81$ K), the same as that used in Ref. 5. Its surface was carefully scraped before every measurement to remove contamination.

III. DATA ANALYSIS

Although the linear dichroism of Sm$^{3+}$ and Gd$^{3+}$ is not zero, our calculation results have shown that the difference between $\int_{M^*} M^0 (\mu^+ + \mu^-)$ and $\int_{M^*} M^0 (\mu^+ - \mu^-)$ is less than 10% for Sm$^{3+}$ and negligibly small for Gd$^{3+}$, where $M^0$ is the absorption coefficient for linear polarized light with the polarization parallel to the quantization axis. Then the expectation values of $M_L$ and $M_S$ of the 4f electrons can be extracted from XMCD spectra by the sum rules13-15

$$\langle M_L \rangle = -\langle L_Z \rangle \mu_B = 2n\frac{\int_{M^*} M^0 (\mu^+ + \mu^-)}{\int_{M^*} M^0 (\mu^+ - \mu^-)} \mu_B,$$

$$\langle M_S \rangle = -2\langle S_Z \rangle \mu_B = \left(2n\frac{\int_{M^*} M^0 (\mu^+ + \mu^-) - \frac{3}{2}\int_{M^*} M^0 (\mu^+ - \mu^-)}{\int_{M^*} M^0 (\mu^+ - \mu^-) + 6\langle T_Z \rangle} \right) \mu_B,$$

where $L_Z$, $S_Z$, $T_Z$, and $n$ are the orbital and spin angular momenta, magnetic dipole operator, and the number of holes in the 4f shell, respectively. The replacement of the integration of $\mu^0$ by that of $\frac{1}{2}(\mu^+ + \mu^-)$ results in the same small relative error ($<3\%$) for both spin and orbital magnetic moments (please note that $\langle T_Z \rangle$ is almost proportional to $\langle S_Z \rangle$ as shown in Table III), and this error can be absorbed by the multiplying factor related to the unsaturated magnetization effect as discussed later. For rare-earth elements with less-than-half filled 4f shells, the off-diagonal term of the 3d-4f exchange interaction causes a large amount of $3d_{5/2}-3d_{3/2}$ mixing and the sum rule for $M_T$ becomes invalid.16 To define a correction factor $X_l/X_E$ to describe the effect of this mixing, where

$$X_l = -\frac{\int_{M^*} (\mu^+ - \mu^-) - \frac{3}{2}\int_{M^*} (\mu^+ - \mu^-)}{\frac{3}{2}\int_{M^*} (\mu^+ + \mu^-)},$$

related to the absorption, and

$$X_E = \frac{2}{3n} \langle S_Z \rangle + \frac{2}{n} \langle T_Z \rangle.$$

For 3d electrons, $\langle T_Z \rangle$ is zero for polycrystal samples because of the decoupling between the quadrupolar charge distribution and magnetic spin orientation due to the small spin-orbital interaction17 and $\langle T_Z \rangle$ need not be considered. However, for 4f electrons, the crystal field is weaker than the spin-orbital interaction and $\langle T_Z \rangle$ is almost free of the crystal field18 and isotropic, so its value for polycrystal samples is about the same as that for free ions.

IV. RESULTS AND DISCUSSIONS

The Sm $M_{4.5}$ absorption and XMCD spectra obtained at different temperatures are shown in Fig. 1. Panel (c) shows the raw XMCD spectra with the sample being magnetized parallel and antiparallel to the sample surface normal. The near-zero constant background dichroism away from the absorption edges and the symmetry of these two spectra confirm that the systematic errors can be removed easily by background subtraction and the measurements can be done with the sample being magnetized along only one direction. At 81 K, the compensation temperature, both the spectra with $T_m$ higher and lower than $T_{\text{comp}}$ [see Fig. 1(d) and its caption] were taken, and the polarities of their dichroism are opposite to each other.

The XMCD measurements for Gd are somewhat difficult due to its small concentration. To remove the very high background dichroism, the measurement at every temperature was done by two scans with different magnetization directions, parallel and antiparallel to the sample normal. The average absorption and related XMCD spectra after background subtraction are shown in Fig. 2.

For atomic Sm$^{3+}$ and Gd$^{3+}$ ions, theoretical XMCD spectra and the expectation values of $L_Z$, $S_Z$, and $T_Z$ of their 4f electrons were obtained from full atomic multiplet calculations. In our calculations, the electrostatic Coulomb and exchange parameters were taken from Ref. 19. The magnetic effect was introduced by a molecular field and its intensity at temperature $T$ can be evaluated by solving the equation

$$B_m \mu_B = J_{ff} S_B (2S \mu_B B_S/kT)$$

where $B_m$, $S$, $J_{ff}$, $B_S(x)$ are the molecular field intensity, the spin angular momentum of the Sm$^{3+}$ ground state (is 5/2 here), the Heisenberg exchange parameter, and the Brillouin function, respectively. Our calculation results show that the slopes of the magnetic moment...
versus temperature [see Fig. 3(a)] are mainly determined by the molecular field intensity, whereas those of Sm$^{3+}$ are a little bit reduced at low temperatures due to the crystal field. The best fitting between theoretical and experimental slopes is achieved when the crystal-field splitting between $G_7$ and $G_8$ levels of Sm$^{3+}$ ground state is set as 2 meV, and the same $J_{ff}$ of 0.96 meV, which is the half of that estimated from the sample Curie temperature $T_C$ (about 130 K) as $J_{ff} = 3kT_C/[2S(S+1)]$, is adopted for both Sm$^{3+}$ and Gd$^{3+}$ ions because of the dilute concentration of Gd$^{3+}$. We can see that the experimental results can be reproduced very well by the theoretical calculations as shown in Fig. 3(a). The calculation results of relative energy level, the expectation values of spin and orbital angular momentum and magnetic dipole operator for the lowest multiplets of 4f electrons in atomic Sm$^{3+}$ and Gd$^{3+}$ ions when the molecular field is 2.25 meV (corresponding to the 48 K temperature; see Table III) are shown in Tables I and II, respectively. The strength of the molecular field estimated from above equations, the correction factor

FIG. 1. (a) Sm $M_{4,5}$ x-ray absorption spectra observed at temperatures of 48, 65, 81, 97, and 119 K with $T_m = 110$ K are compared with the theoretical spectra of atomic Sm$^{3+}$. Solid (broken) lines represent the absorptions with the helicity of the incident photons parallel (antiparallel) to the sample magnetization. (b) XMCD spectra deduced from the spectra in panel (a). (c) The experimental XMCD spectra taken at 119 K with the sample being magnetized parallel (solid line, $\mu_+ - \mu_-$) and antiparallel (broken line, $\mu_- - \mu_+$) to the sample normal with $T_m = 110$ K. (d) XMCD spectra taken at 81 K with $T_m = 110$ K (solid lines, deduced from the spectra in panel (a)) and $T_m = 43$ K (broken line; the corresponding absorption spectra are not shown).

FIG. 2. (a) The average experimental spectra of Gd $M_{4,5}$ absorption with $T_m = 110$ K are compared with the theoretical ones of atomic Gd$^{3+}$. The definition of solid and broken lines is the same as that in Fig. 1(a). (b) The XMCD spectra deduced from the spectra in panel (a).

FIG. 3. (a) The experimental expectation values of $m_{S_s}^{Sm}$ (circle), $m_{S_s}^{Gd}$ (cross), $m_{L_n}^{Sm}$ (triangle), and $m_{L_3}^{Gd}$ (times) (all enlarged by a factor of 3.05 to eliminate the unsaturated magnetization effect) when the sample was magnetized at 110 K (open marks) and 43 K (filled marks) are compared with theoretical estimations for atomic Sm (broken lines) and Gd (dashed lines) trivalent ions. (b) The distributions to the net magnetization (+, right axes) from the Sm ion (circle), Gd ion (square), and conduction electrons (times) are shown with the net magnetization of SmAl$_2$ (triangle), which is the gross magnetic moments of both Sm ions and conduction electrons.
TABLE I. The calculation results of relative energy level, the expectation values of spin angular momentum, orbital angular momentum, and magnetic dipole operator for the lowest multiplets of 4f electrons in atomic Sm and Gd.

<table>
<thead>
<tr>
<th>Energy (meV)</th>
<th>$S_Z$</th>
<th>$L_Z$</th>
<th>$T_Z$</th>
</tr>
</thead>
<tbody>
<tr>
<td>113</td>
<td>−1.829</td>
<td>2.494</td>
<td>0.348</td>
</tr>
<tr>
<td>118</td>
<td>−1.782</td>
<td>2.515</td>
<td>0.350</td>
</tr>
<tr>
<td>119</td>
<td>−1.653</td>
<td>2.476</td>
<td>0.341</td>
</tr>
<tr>
<td>120</td>
<td>−1.041</td>
<td>2.676</td>
<td>0.342</td>
</tr>
<tr>
<td>121</td>
<td>−0.575</td>
<td>2.756</td>
<td>0.343</td>
</tr>
<tr>
<td>122</td>
<td>−0.201</td>
<td>2.866</td>
<td>0.344</td>
</tr>
<tr>
<td>123</td>
<td>0.124</td>
<td>2.996</td>
<td>0.345</td>
</tr>
<tr>
<td>124</td>
<td>0.757</td>
<td>3.126</td>
<td>0.346</td>
</tr>
<tr>
<td>125</td>
<td>1.483</td>
<td>3.246</td>
<td>0.347</td>
</tr>
</tbody>
</table>

The experimental spectra and magnetic moments of 4f electrons in (Sm, Gd)Al$_2$ can be described as being caused by the exchange interaction between the 4f electrons and conduction electrons, and thus its magnitude should be proportional to the spin magnetic moment of 4f electrons. The coefficient $C$ can be determined from the experimental $m_{con} = C m_{Sm}$, where the sizes of magnetic moments are deduced from the experimental $m_{Sm}$ as shown in Fig. 3(b).

The contributions from different sources to the magnetism of Sm$_{0.98}$Gd$_{0.018}$Al$_2$ are shown in Fig. 3(b). When temperature decreases, the circular dichroism [see Figs. 1(b) and 2(b)], the sizes of magnetic moments [see Fig. 3(a)], and the net magnetic moments (vector sum of spin and magnetic moments) of every part [conduction electrons, Sm or Gd]...
ions; see Fig. 3(b)] all increase. The net magnetic moment of Sm$^{3+}$ ions ($\Theta$, $m_{\text{Sm}}=0.982(m_S^{\text{Sm}}+m_L^{\text{Sm}})$), where 0.982 is the Sm concentration) is always orbital dominant. The magnitude of $m_{\text{con}}$ ($\times$) is not enough to cancel out the surplus orbital magnetic moment in Sm$^{3+}$ and the net magnetic moment of SmAl$_2$ ($\Delta$, $m_{\text{SmAl}_2}=m_{\text{Sm}}+m_{\text{con}}$) cannot reach zero. The net magnetic moment of Gd ions ($\Box$, $m_{\text{Gd}}=0.018(m_S^{\text{Gd}}+m_L^{\text{Gd}})$) is almost of spin kind only [the orbital one is negligibly small as shown in Fig. 3(a)] and its participation cancels out the surplus orbital magnetic moment completely. The total, or to say, the net magnetic moment of (Sm,Gd)Al$_2$ ($+$, $M_T=m_{\text{Sm}}+m_{\text{con}}+m_{\text{Gd}}$) becomes zero at $T_{\text{comp}}$.

V. CONCLUSIONS

In conclusion, the existence of ferromagnetic ordering for both spin and orbital magnetic moments in Sm$_{0.982}$Gd$_{0.018}$Al$_2$ at the compensation temperature is proved perfectly by our XMCD study. The cancellation of spin and orbital magnetic moments cannot be achieved by Sm ions only. The spin magnetic moment of conduction electrons is not enough to cancel out the surplus orbital magnetic moment in Sm ions and the addition of spin magnetic moment from Gd ions cancels out the orbital magnetic moment thoroughly at $T_{\text{comp}}$ and results in the zero net magnetization. The magnetic moments of Sm and Gd ions in (Sm,Gd)Al$_2$ can be well described in an atomic picture.

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1 The contributions of $M_S$ and $M_L$ to the paramagnetic moment of the free Sm$^{3+}$ ion can be estimated to be $25\mu_B/\sqrt{35}$ and $-30\mu_B/\sqrt{35}$, respectively.


7 J. W. Taylor, J. A. Duffy, A. M. Bebb, M. R. Lees, L. Bouchenoire, S. D. Brown, and M. J. Cooper, Phys. Rev. B 66, 161319(R) (2002). It seems, however, that the direction of the sample magnetization was not well controlled in their experiment of x-ray diffraction, though this condition is crucial.


10 It seems the observed almost zero form factors near $T_{\text{comp}}$ in Ref. 7 is the result of the unsuccesful control of the magnetization of their sample.


18 The electronic states of Gd$^{3+}$ is not influenced by the crystal field because its zero total orbital angular momentum. For Sm$^{3+}$, our calculations show that when the fourth- and sixth-order crystal-field parameters $A_4(r^4)$ and $A_6(r^6)$ change from 0 to 30 meV, $(T_2)$ varies less than 0.8%.


20 The Hamiltonian of the spin magnetic moment under the molecular field $B_m$ can be expressed as $2\mu B_2 B_m=2(g-1)\mu_B/B_2$ (see Ref. 2).