Field-Induced Magnetic Transitions in the One-Dimensional Compound Ca₃Co₂O₆

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Magnetization measurements on the one-dimensional oxide Ca₃Co₂O₆ having a triangular net of Co₂O₆ chains have been carried out both in static and pulsed high magnetic fields. The M/H vs. T curve obeys the Curie-Weiss law at high temperatures. Below 25 K, however, M/H increases abruptly, and a plateau is observed at 1/3 of the full moment in the M-H curve, suggesting a ferrimagnetic state of the ferromagnetic chains due to the antiferromagnetic interchain interaction. At low H, this system is considered to be in a partially disordered antiferromagnetic state for 10 K < T < 25 K and in a ferrimagnetic state below 10 K. The observed plateau in the M-H curve for T ≤ 5 K is broader than for 10 K ≤ T ≤ 20 K, indicating spin freezing at lower temperatures. The results can be discussed in terms of the triangular Ising spin systems.

KEYWORDS: ferrimagnetic transition, triangular lattice, frustration, Ca₃Co₂O₆, one-dimensional magnetic chain, Ising spin system, partially disordered antiferromagnetic phase

The cooperative phenomena associated with magnetic linear-chain systems are of great interest. Experimentally, many studies have been carried out on CsCoCl₃, RbFeCl₃ and other ABO₃-type compounds.¹⁻⁵ At low temperatures, these nearly one-dimensional systems exhibit transitions to a variety of three-dimensional ordered states. Such transition is induced by relatively weak coupling between chains of strongly coupled spins. The sign and magnitude of interchain and intrachain interactions are expected to determine the type of ordered state. Theoretically, the Ising spin model with antiferromagnetic nearest-neighbor interactions on the triangular lattice is investigated most extensively because of its simplicity. The unique magnetic properties of CsCoCl₃ associated with the triangular lattice are known to be well explained by this model.⁶⁻⁷

Recently, Fjellvåg et al. have determined the crystal structure of Ca₃Co₂O₆,⁸ which is related to those of Sr₃ABO₆ (where A=Ni, Cu, Zn, B=Ir, Pt).⁹⁻¹⁰ Ca₃Co₂O₆ belongs to the space group R̅3c. The structure of Ca₃Co₂O₆ is shown in Fig. 1. Figure 1(a) shows the perspective view along the hexagonal [110] direction. Sr₃ABO₆ consists of infinite chains of alternating face-sharing AO₆ trigonal prisms and BO₆ octahedra (anti-trigonal prisms), whereas Co atoms occupy both A and B sites in Ca₃Co₂O₆. Consequently, the Co₂O₆ infinite chains are built by successive alternating CoO₆ trigonal prisms (with Co1) and CoO₆ octahedra (with Co2) along the hexagonal c-axis. One-dimensional character along the c-axis can be expected because the Co₂O₆ chains are widely spaced; the interchain separation is 5.24 Å which is about twice as large as the intrachain Co distance of 2.59 Å.⁸ Each Co₂O₆ chain is surrounded by six chains forming a triangular net as shown in Fig. 1(b). Thus, we can expect cooperative phenomena caused by the triangular arrangement of the Co₂O₆ chains. It should be noted that the phase of each chain is different from the phase of three nearest-neighbor chains by 1/6 along the c-axis (the chain direction) and from the phases of the other three by 2/6 along the same axis (see Fig. 1(a)).

To elucidate the magnetic behavior of Ca₃Co₂O₆, we have performed magnetization measurements at various temperatures under both static and pulsed fields up to 5 T and 40 T, respectively. The main finding is that the M-H curves below 25 K show stepwise magnetization; the first step is the ferrimagnetic plateau at 1/3 of the
saturation magnetization and the second step over 10 T is full magnetization (1.8 \mu_B/Co).

The Ca$_3$Co$_2$O$_6$ sample was prepared by a solid state reaction method from CaCO$_3$ and CoO with 99.999% purities. Powders were ground, pelletized and calcined in air at 1173 K for one day. Then, the sample was reheated at 1273 K in air for one week. Using the powder X-ray diffraction, the sample was confirmed to be in a single phase of Ca$_3$Co$_2$O$_6$ with the space group R3c and with the lattice parameters a = 9.071 Å and c = 10.375 Å in the hexagonal expression which was almost in agreement with the data of Fjellvåg et al. The magnetization was measured by using a SQUID magnetometer (Quantum Design MPMS) as a function of temperature (2 K \leq T \leq 300 K) and field (0 T \leq H \leq 5 T). The magnetic susceptibility \chi (=M/H) for 300 K \leq T \leq 770 K was measured at H=0.5 T using a torsion magnetic balance. High field magnetization measurements were performed up to 40 T for powdered Ca$_3$Co$_2$O$_6$ by an induction method with well balanced pick-up coils. The duration time of the pulsed magnetic field was 12 msec, corresponding to (\Delta H/\Delta t) = 6.7 T/msec. We also performed magnetization measurements up to 6.2 T under the condition of (\Delta H/\Delta t) = 2.0 T/msec.

First, we show, in Fig. 2, the temperature dependence of M/H measured using a SQUID magnetometer and a torsion magnetic balance. As seen in the inset of this figure, \chi is fitted well to the Curie-Weiss law over the range of 100 to 800 K with the Weiss temperature \theta = 40.4 K and the paramagnetic effective Bohr-magneton number \mu_B/Co, accompanied with a constant susceptibility \chi_0 = 1.7 \times 10^{-4} emu/Co mol. Below 100 K, the deviation from the Curie-Weiss law becomes appreciable. When the temperature is decreased to 25 K (=T_c), the values of M/H start to increase suddenly, and the magnetization curve shows non linear field dependence.

A ferromagnetic intrachain interaction is deduced from the abrupt increase in the M/H vs. T curve and the positive value of \theta. The irreversibility in M/H vs. T curves appears at 10 K between zero-field-cooled (ZFC) and field-cooled (FC) processes.

Figure 3 shows the representative magnetization curves in static fields up to 5 T measured using a SQUID magnetometer. In the paramagnetic region above T_c, the magnetization is proportional to H, as seen in the M-H curve at 30 K, whereas, below T_c, non linearities appear in M-H curves. One can see a clear plateau in the M-H curves in the temperature range from 8 K to 16 K for H < 3 T (see Fig. 3(a)), implying a ferrimagnetic ordering of ferromagnetic chains.

The magnetizations measured using pulsed fields are shown in Fig. 4. The magnetization at 35 K, where the spin system is in a paramagnetic state, is saturated toward 1.8 \mu_B/Co. It should be noted that the M-H curve is almost temperature independent for H > 10 T. This suggests that the magnetic transition at T_c is not due to the change in the amplitude of Co moment which would be associated with a transition between the high spin state and the low spin one nor that of charge ordering between Co1 and Co2.

As seen in Fig. 4(b), one can observe a stepwise change in the magnetization at 15 K. The observed plateau is not clear and hysteresis is observed, in contrast to the M-H curve at 14 K in static fields. Hysteresis becomes clearer for T \leq 5 K and will be discussed later. The plateau at 1/3 of the full moment strongly suggests a ferrimagnetic alignment of ferromagnetic chains. In the paramagnetic state, the short range ferromagnetic order develops predominantly along the magnetic chains. When the short range order is well developed as temperature
decreases, each ferromagnetic chain would behave like a magnetic moment. Therefore, the resultant magnetic structure should have two-dimensional character, resulting in a triangular-lattice antiferromagnet. The ferromagnetic state indicates an antiferromagnetic interchain interaction. In the triangular lattice Heisenberg spin system, a 120° structure, where spins rotate successively by 120°, is stable. Therefore, Co spins would have Ising-like character. The molecular field theory for an antiferromagnetic triangular Ising lattice gives phase transition from a paramagnetic to a ferrimagnetic state. Some compounds with a triangular net have been reported to exhibit ferrimagnetic ordering. In the case of NaNi acac₃-benzene, however, there is no onedimensional chain. As for Ce₂Eu, the ground state is a 120° state and a ferrimagnetic state is induced by a field, which is explained by an energy crossing between these two states. In the case of CsCoCl₃, intrachain interaction is antiferromagnetic, in contrast to Ca₃Co₂O₅. To the authors' knowledge, this is the first observation of the ferrimagnetic ordering of well-developed ferromagnetic chains among triangular lattice systems.

From the Arrott plots for the magnetization data in static fields, Ca₃Co₂O₅ is in a ferrimagnetic state below T_c=10 K at zero field. Therefore, the observed metamagnetic transition below 10 K is from a ferrimagnetic to a ferromagnetic state. For T_c< T< T_c, on the other hand, a partially disordered antiferromagnetic (PDA) phase is possible in low fields, in which 2/3 of the chains exhibit antiferromagnetic order, leaving the remaining 1/3 incoherent with respect to the other chains. According to ref. 6, a PDA phase between a paramagnetic and a ferrimagnetic states is predicted at zero magnetic field. From neutron diffraction measurements at T=12 K (> T_c2) and under zero magnetic field, we successfully observed magnetic reflections, which are likely to be those from the PDA phase. In a PDA state only incoherent chains would contribute to the increase in magnetization as seen in the M-H curve at 15 K. Since the propagation of a domain wall in the ferromagnetic chain is considered to be more difficult than in the case of the antiferromagnetic chain, thermal fluctuations and the interaction between two or three solitons may be important for facilitating the soliton motion in the ferromagnetic chain. Further study is needed in order to confirm the existence of the PDA phase.

Finally, let us discuss the magnetic behavior for T ≤ 10 K. As seen in Fig. 3(b), prominent hysteresis is observed and the metamagnetic transition is broadened, compared with that for T ≥ 10 K. The hysteresis at 2 K is much larger than that at 5 K. As mentioned above, the M/H vs. T curve shows a spin freezing below 10 K; the irreversibility between (M/H)_FC and (M/H)_FC was observed around 10 K. In the case of a pulsed field with (ΔH/Δt) = 6.7 T/msec, on the other hand, the magnetization at 4 K is completely different from that obtained in the static field measurements (cf. Fig. 3(b) and Fig. 4(c)). In the case of (ΔH/Δt) = 2.0 T/msec, a plateau in the M-H curves was observed, as shown in the inset of Figs. 4(b) and 4(c). This result indicates that the ferromagnetic chains (or Co spins) respond to an external field at a rate of the order of 10⁻³ sec. For NaNi acac₃-benzene, similar behavior is observed under the condition of (ΔH/Δt)=300 T/msec, which is explained as the energy level crossing of the d-orbitals of Ni²⁺. However, the energy level crossing is not likely to occur for Ca₃Co₂O₅ because the field sweeping rate is much smaller than that in the case of ref. 13. We cannot give a reasonable explanation for the change from the ferromagnetic to the spin freezing phase. The difference in the configuration of nearest-neighbor chains and/or an interchain interaction between separated chains such as the third and forth nearest-neighbor chains may play a part.

In summary, we propose a rough magnetic phase diagram of Ca₃Co₂O₆, as shown in Fig. 5.

Although we have clarified the magnetic properties of Ca₃Co₂O₅ in terms of the one-dimensional ferromagnetic chain, the moment of each cobalt ion is not clear. Since there are two different Co sites, two possibilities can be considered regarding the Co valence: (1) Co ions at both sites are trivalent. (2) one Co ion is divalent and the other tetravalent. It is also unknown whether Co spins are in the low spin state or the high spin state in connection with the energy level splitting of d-orbitals by the crystalline field of an octahedron and a trigonal prism. Furthermore, assuming that the g factor is 2, the saturation moment corresponds to about two thirds of the value expected from the experimental μ_B=3.88 μ_B/Co. This small value should be interpreted using the fictitious spin S' of Co²⁺ ions as in the case of Ising spin

![Fig. 4. Magnetization curves of Ca₃Co₂O₅ under high pulsed magnetic fields up to 40 T with (ΔH/Δt) = 6.7 T/msec at 35 K (a), 15 K (b), and 4 K (c). The arrows along the curves indicate the directions of the field sweep. The inset represents magnetization curves up to 6.2 T with (ΔH/Δt) = 2.0 T/msec at 15 K (b) and 4 K (c) plotted in the same units as the main plot.](image-url)
magnetic interaction between ferromagnetic chains on a triangular lattice. Ferrimagnetic transition temperature $T_{c2}$ at zero field is 10 K. A spin freezing behavior is observed at lower temperatures. A PDA state is possibly realized for $T_{c2} < T < T_{c1}$.


Fig. 5. Schematic phase diagram as a function of $H$ and $T$. FR, a ferrimagnetic state; F, a ferromagnetic state; SF, a spin freezing state; P, a paramagnetic state; PDA, a partially disordered antiferromagnetic state.

For the further understanding of the magnetic properties of $\text{Ca}_3\text{Co}_2\text{O}_6$, specific heat, neutron diffraction, NMR and ESR measurements are now in progress. A single crystal will be required although all attempts have been unsuccessful to date.

In conclusion, we have measured magnetization up to 40 T for $\text{Ca}_3\text{Co}_2\text{O}_6$ and observed a plateau at 1/3 of the full moment. This indicates ferrimagnetic alignment between the ferromagnetic chains, due to the antiferromagnetic systems.$^5,17$