

MAGNETIC PROPERTIES OF NEW COMPOUNDS RMg₂Cu₉ WITH TWO-DIMENSIONAL ALIGNMENT OF R ATOMS*

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Structural and magnetic properties of new compounds RMg₂Cu₉ were examined. The results indicated that RMg₂Cu₉ crystallize in the CeNi₃ type hexagonal structure for R = La, Ce, Pr, Nd, Sm and show magnetic order for R = Ce and Sm, that is in contrast with RMg₂Ni₉ system. Some contrastive physical properties in RMg₂T₉ (T = Ni, Cu) might be originated in both the two-dimensionality of R atoms and the number of carrier.

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1. Introduction

CeMg₂Ni₉ crystallizes in the hexagonal PuNi₃ type structure, while CeMg₂Cu₉ crystallizes in the CeNi₃ type structures [1], both of which are built up from the stacking of single layers of CeT₅ (T = Ni, Cu) and double layers of MgT₂ along the *c*-axis. It have been clarified that CeMg₂T₉ system reveal some novel physical properties, for example, no magnetic long range order in RMg₂Ni₉ with R = Ce, Pr, Nd and Gd and almost pressure-independence of T_N up to 0.9 GPa for CeMg₂Cu₉, that might be reflected two-dimensionality of R atoms [2]. To clarify the influence of two-dimensional arrangement of R atoms, we studied physical properties of light-rare-earth RMg₂Cu₉ (R = La, Pr, Nd and Sm) compounds. In this paper, we report the crystal structure and magnetic properties of new compounds RMg₂Cu₉ (R = La, Ce, Pr, Nd and Sm). Details of the sample preparation and the other experimental procedures are given in Ref. [3].

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2. Results and discussion

2.1. Crystal structure

Figure 1(a) shows powder X-ray diffraction pattern for RMg_2Cu_9 ($\text{R} = \text{La}, \text{Ce}, \text{Pr}, \text{Nd}$ and Sm) together with the simulation pattern assumed CeNi_3 type hexagonal structure. Comparing our experimental pattern with the simulation one, we can conclude that RMg_2Cu_9 ($\text{R} = \text{La}, \text{Ce}, \text{Pr}, \text{Nd}$ and Sm) is in a single phase of CeNi_3 type hexagonal structure. The refined lattice constants are listed in Table I. The lattice constants of RMg_2Cu_9 obey the lanthanide contraction, indicating that R is in a tri-valent state, being in contrast with RMg_2Ni_9 system that Ce is in an intermediate valence state [4].

From the above simulation result, the crystal structure of RMg_2Cu_9 can be drawn in figure 1(b), which is built up of alternating single layers of RCu_5 and double layers of MgCu_2 along the c -axis. In this structure, the R-R nearest neighbor distance along the c -axis are almost two-times larger than that in the c -plane (see Table I). Thus, these compounds are characterized by two-dimensional alignment of R atoms.

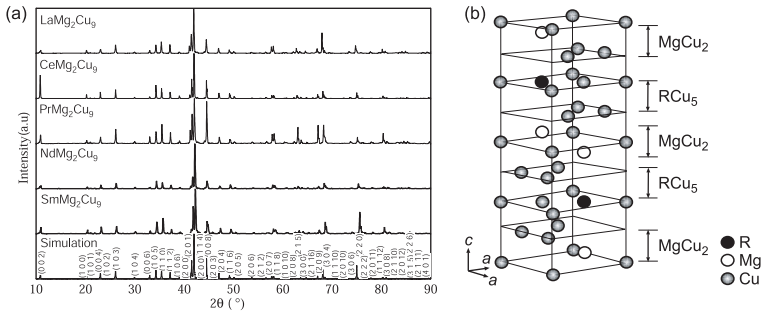


Fig. 1. (a) Powder X-ray diffraction profiles of RMg_2Cu_9 ($\text{R} = \text{La}, \text{Ce}, \text{Pr}, \text{Nd}$ and Sm) and the simulation pattern assumed CeNi_3 type hexagonal structure. (b) The crystal structure of RMg_2Cu_9 with a hexagonal CeNi_3 type.

TABLE I

Structural and magnetic properties of RMg_2Cu_9 ($\text{R} = \text{La}, \text{Ce}, \text{Pr}, \text{Nd}$ and Sm).

RMg_2Cu_9	Lattice constance			R-R nearest neighbor distance		μ_{eff} (μ_B)		θ_p (K)	T_N (K)
	a (Å)	c (Å)	c/a	a (Å)	c (Å)	Observed	Theoretical		
LaMg_2Cu_9	5.073	16.270	3.207	5.073	8.646	2.4	2.54	-15.7	2.7
CeMg_2Cu_9	5.061	16.260	3.213	5.061	8.639	3.3	3.58	-21.7	
PrMg_2Cu_9	5.052	16.251	3.217	5.052	8.633	3.6	3.62	-1.56	
NdMg_2Cu_9	5.041	16.236	3.221	5.041	8.624				10
SmMg_2Cu_9	5.027	16.213	3.225	5.027	8.610				

2.2. Magnetic properties

The magnetization curves for RMg_2Cu_9 at 2 K are shown in figure 2. For LaMg_2Cu_9 , the magnetization decrease linearly with increasing magnetic field, indicating LaMg_2Cu_9 is diamagnetism. On the other hand, the magnetization for $\text{R} = \text{Ce}, \text{Pr}$ and Sm , linearly increase with increasing magnetic field, indicating that these compounds are antiferromagnetism or paramagnetism. For NdMg_2Cu_9 on the contrary, the magnetization gradually increases with increasing magnetic field.

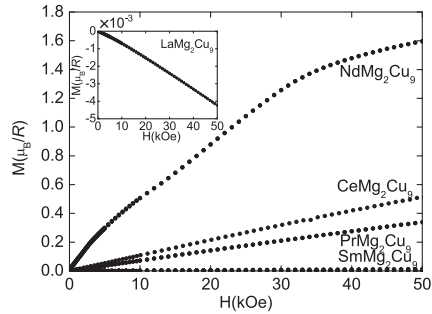


Fig. 2. Magnetization curve for RMg_2Cu_9 at 2 K. The inset shows the magnetization curve for LaMg_2Cu_9 .

Figure 3 shows the temperature dependence of the magnetic susceptibility χ for RMg_2Cu_9 with (a) $\text{R} = \text{Ce}, \text{Pr}$ and Nd , (b) La , and Sm . The χ of

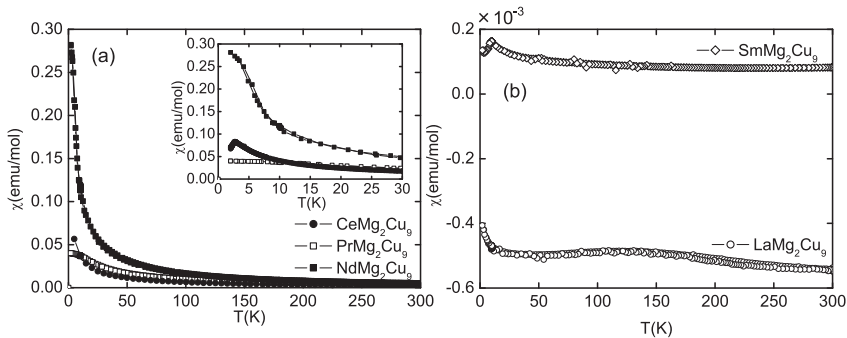


Fig. 3. Temperature dependence of the magnetic susceptibility at 10 kOe for RMg_2Cu_9 (a) $\text{R} = \text{Ce}, \text{Pr}, \text{Nd}$ and (b) La, Sm .

LaMg_2Cu_9 and SmMg_2Cu_9 are almost independent of temperature, reflecting diamagnetism and Van Vleck paramagnetism. The χ of CeMg_2Cu_9 and SmMg_2Cu_9 exhibit a small peak at 2.7 K and 10 K respectively, suggesting

the existence of antiferromagnetic order below these temperatures. On the other hand, the χ of PrMg_2Cu_9 was observed no magnetic anomaly at 2 K, which is due to singlet ground state caused by a crystal electric field splitting. The χ for NdMg_2Cu_9 shows induced-ferromagnetic behavior below about 4 K. The $1/\chi$ for $\text{R} = \text{Ce}, \text{Pr}, \text{Nd}$ follows the Curie–Weiss Law with effective magnetic moments that are close to theoretical values expected for the R^{3+} free ion. The effective magnetic moment μ_{eff} , Curie–Weiss temperature θ_p and magnetic ordering temperature are summarized in Table I. Since the θ_p for NdMg_2Cu_9 is near 0 K, it seems likely that the ground state of NdMg_2Cu_9 is in a paramagnetic state and shows induced-ferromagnetism in high field at $T < 4$ K.

It should be noted that RMg_2Cu_9 for $\text{R} = \text{Ce}$ and Sm show antiferromagnetic order at low temperature, while RMg_2Ni_9 show no-magnetic order even for Gd-system with large spin components. The reason of no magnetic order for RMg_2Ni_9 was originally thought to be the two-dimensionality of rare earth arrangement [3]. However, there is antiferromagnetic order in the RMg_2Cu_9 with $\text{R} = \text{Ce}$ and Sm , although nearest neighbor R–R distances along c -axis are larger than that in the RMg_2Ni_9 . Therefore, it seems likely that the reason of no magnetic order for RMg_2Ni_9 is not only in a two-dimensionality of R atoms, but also in a low carrier number of conduction electrons which could be observed as a high electrical resistivity compared to that in the Cu system [2].

3. Summary

We have clarified that new compounds of RMg_2Cu_9 crystallize in the hexagonal CeNi_3 type structure for $\text{R} = \text{La}, \text{Ce}, \text{Pr}, \text{Nd}, \text{Sm}$ and show magnetic order for $\text{R} = \text{Ce}$ and Sm , that is in contrast with the RMg_2Ni_9 system. The physical properties of RMg_2T_9 for $\text{T} = \text{Ni}, \text{Cu}$ therefore might be originated in both the two-dimensionality of R atoms and the number of carrier.

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