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A MECHANISM FOR THE METAL INSULATOR TRANSITION AND VARIOUS PROPERTIES IN SAMARIUM COMPOUNDS

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Abstract. — The 4f states for the insulator side EuB₆ and the metallic side SmB₆ are investigated. Transport properties and ESR line widths of Gd and Eu in SmB₆ are evaluated satisfactorily on this model. Metal insulator transitions in Y and As doped SmS are studied on the bases of the molecular magnetic exciton and molecular magnetic impurity state in excellent agreement with experiments.

1. Introduction. — In various Sm compounds, Sm ions change the valency from divalence to trivalence, which causes the complicated metal-insulator transition accompanied with various anomalous properties. In this sense, these compounds have been studied very intensively in the recent several years [1]. Among them, Sm-chalcogenides, SmₓX, in which X means S, Se or Te, and Sm-hexaboride, SmB₆, have attracted much attention. In particular, SmB₆ is the most convenient material to study the mechanism for the above transition because the rigid lattice is formed mainly by boron and thus only a small change of the volume is observed with the above transition. At first, in the present paper, we treat the mechanism for the metal-insulator transition in SmB₆ based on the recent experimental results of our group and then treat that for SmS mostly doped with Y and As worked by IBM group.

2. Physical picture for SmB₆. — In rare earth borides, RB₆, R is the molecule Bₓ, which is similar to the chalcogen X in RX and thus exists as B₆⁻, forms the CsCl type lattice. Therefore, EuB₆, in which Eu⁺⁺⁺ is the stable state, should be insulator similar to EuX while other RB₆ should be metallic with one conduction electron per a rare earth atom. Recent remarkable improvement for the sample preparation now makes possible to produce very high purity samples so as to determine the Fermi surface of LaB₆ very clearly [2]. Thus we now know that the bottoms of the conduction band are at X-points of the εₓ character with the band width of a couple of eV. The situation of EuB₆ is, however, not simple because the existing samples are all metallic. This is due to the following facts. Recent experiment by Mercurio et al. [3] shows that the 4f levels are situated very closely to the bottom of the conduction band, may be several tenths of eV, much closer than these in EuX. While in RB₆, because of the rigid boron net work, some amount of the rare earth vacancies is always produced. In usual situation, this makes the sample p-type, such as Eu vacancies in EuX. However, in EuB₆, due to the small energy gap, this makes the sample n-type as shown in figure 1. The vacancy has effectively two negative charge and thus is compensated by producing two Eu⁺⁺⁺ ions as shown by sites 1 and 2 in figure 1. Then, the potential at site 3 is raised by

$$\Delta V = (2 - \sqrt{2}) e^2/4a,$$

in which s is the effective dielectric constant with the value of the order one. By substituting, for example, e = 4 and a = 4.2 Å, we have $$\Delta V = 0.59 \text{ eV},$$ which is nearly enough to push the 4f level at site 3, or any other equivalent site, up to the bottom of the conduc-
tion band. Thus, we expect that one Eu vacancy creates three Eu$^{+++}$ states and one conduction electron, which should be bound weakly at the above mentioned defect complex but delocalized very easily for a small concentration of the vacancies, may be the order of $10^{-3}$ to $10^{-4}$. In this way, EuB$_6$ can exist nearly always as the n-type degenerate semiconductor.

EuB$_6$ is ferromagnetic below about 15 K. Our best samples show the residual resistivities $10^{-3}$ $\Omega$-cm [4]. The resistivities increase sharply at the Curie temperature $T_C$ and become nearly constant, $5 \times 10^{-3}$ $\Omega$-cm, to the room temperature due to the strong scattering by the defect complexes and the d-f exchange interaction. The Hall constant measurement at 4.2 K shows large contribution of the anomalous Hall constant. From the normal part, the carrier number is estimated to be $10^{-2}$ per Eu atom and the mobility 45 cm$^2$/V s. In the paramagnetic region, it is difficult to separate the anomalous term from the normal term. However, the assumption of no anomalous term leads to a too small value of the carrier concentration. It is rather natural to assume that the carrier concentration is nearly temperature independent at least up to the room temperature and a relatively large amount of the anomalous Hall effect exists even in the paramagnetic region. Then the mobility in the paramagnetic region is estimated to be 10 cm$^2$/V s. Note that the mobility of LaB$_6$ is 80 cm$^2$/V s at the room temperature. This is important because in SmB$_6$ Nickerson et al. [5] proposed a very unusual band scheme based on their Hall constant measurement assuming no anomalous Hall effect. This is discussed later.

In SmB$_6$, Sm-vacancies can be created as far as the one third of the Sm sites with very small change of lattice constant, that is Sm$_{1-x}$Sm$_x$B$_6$ for $0 < x < 1/3$ [6]. For $x = 1/3$, all Sm should be trivalent. Detailed experimental study has been done in our group for various stoichiometries of Sm$_{1-x}$Sm$_x$B$_6$ [7]. All anomalous properties are explained consistently by the following model.

It is already well established that in nearly stoichiometric SmB$_6$, the ratio of Sm$^{++}$ and Sm$^{+++}$ is about $4:6$ to $3:7$ and is temperature independent [8]. Therefore, the 4f levels are now pushed up to the middle of the conduction band and the Fermi level is just in the 4f levels. Note that in SmX the 4f levels are pushed up about 1.5 eV from these of EuX. It is obviously impossible to treat the 4f levels by the usual one electron picture. The following picture may be one of the approximations for such states. The ground state of Sm$^{++}$ is the singlet, $S = 3$, $L = 3$ and $J = 0$. Therefore we choose it the bases state, or the effective vacuum, and introduce the Fermion creation and annihilation operators $\beta^+_v$ and $\beta^-_v$, respectively, in which the suffix $v$ means a state $J \neq 0$ for Sm$^{++}$ or any state for Sm$^{+++}$ but hereafter, for simplicity, $v$ is restricted to the six states for $S = 5/2$, $L = 5$ and $J = 5/2$ of Sm$^{+++}$. Note that in the cubic environment the $J = 5/2$ states are split into the doublet and the quartet with the energy gap of the order of 100 $\kappa$, in which $\kappa$ is the Boltzmann constant. Then the main term of the Hamiltonian may be written as

$$\mathcal{H}_0 = \sum_{ik} \epsilon_{ik} a^+_i a_k + \sum_{vn} \epsilon_v \beta^+_{vn} \beta^-_{vn} +$$

$$+ \sum_{vn, v'n} \beta^+_{vn} \beta^-_{v'n} V_{nn}(v,v') \quad (1)$$

in which $a^+_i$ and $a_i$ are the usual creation and annihilation operators for the conduction electron and the last term is an infinitely large correlation energy to avoid the state $\beta^+_{vn} \beta^-_{vn}$. Therefore, $\mathcal{H}_0$ corresponds to the extended Hubbard Hamiltonian. Note that to create a Sm$^{+++}$, one electron is freed which we assumed to be put at the bottom of the conduction band so as to fix the energy scale. Therefore, $\epsilon_v$ is negative and $\epsilon_{ik}$ is measured from the bottom of the conduction band. Note also that the third term, the effective transfer of the 4f electron, mostly comes from the mixing with the valence bands and is important in EuX, an order of 0.1 eV, for the 4f band width [9], but should be much smaller in SmB$_6$. The main terms of the interaction term may be written in the following simple form

$$\mathcal{H}_1 = \sum_{ik} \lambda_{ik}(ik, k', vv') a^+_i a_{k', v} \beta^+_{vn} \beta^-_{v'n} +$$

$$+ \sum V_{nn} \beta^+_{vn} \beta^-_{v'n} a^+_n + \text{c.c.} \quad (2)$$

in which the first term includes both the Coulomb type and the d-f exchange type interactions and the second term means the main term for the d-f mixing, which is the order of a few hundreds $\kappa$ both for SmB$_6$ and SmX$^9$.

Because the 4f band width is very small, in the zeroth approximation, we can pick up the first two terms of $\mathcal{H}_0$ and the Coulomb term of $\mathcal{H}_1$, which may be approximated by $\nu(n_d) n_d n_d$, in which $n_d$ and $n_d$ are the number of the conduction electrons and that of Sm$^{+++}$, respectively. This is the Hamiltonian treated by Falicov and Kimball [10] and, as the conduction band width is large, gives a nearly temperature independent values for $n_d$ and $n_d$. In the next step, we treat the third and fourth terms of $\mathcal{H}_0$ following the spirit of Hubbard approximation [11]. We further simplify the approximation as follows. For the $\nu$-th state of Sm$^{+++}$, the available sites to move around are $(c_3 + c_3) N$, in which $c_3$ and $c_3$ are the fractions of Sm$^{++}$ and the $\nu$-th state of Sm$^{+++}$, respectively, and $N$ is the total Sm sites. Then the $\nu$-th band width is reduced by the factor $(c_3 + c_3)$ while the density of state is kept constant. For the paramagnetic state, we have such six fold degenerate bands. When the d-f mixing term is taken into account, the conduction electrons near to the Fermi level mix strongly with the above mentioned 4f bands. From the symmetry consideration, no 5d conduction electrons at the Fermi level are free from the d-f mixing, and as the density of state of the 4f bands is two order of magnitude larger than
that of the conduction band, no conduction electron like states can exist near to the Fermi level. The situation is shown in figure 2, in which the energy scale for the 4f bands are shown by the reversed sign for convenience. If we assume \( c_2 = 0.4 \), then, \( n \) the paramagnetic region, \( C_v = 0.1 \), \( c_2 + c_v = 0.5 \) and the Fermi level is expected to be situated at the sharply falling part of the 4f bands. Then we immediately arrive the following conclusion that, similar to Pd [12], the susceptibility due to the Sm\(^{+++}\) bands should increases gradually with increasing temperature and makes a broad peak at about a few 10 K, while the coefficient for the linear \( T \) dependent term of the specific heat also increases gradually and makes a broad peak at about a few 10 K, while the coefficient for the linear \( T \) dependent term of the specific heat also increases gradually and makes a broad peak similar to the susceptibility. These behaviours are really observed experimentally [5, 7, 13]. The absolute values for the susceptibility and the specific heat are also consistent with the present picture of the narrow 4f bands with the band width of a few hundreds \( \kappa \).

\[
\sigma_v = \frac{N \rho_\nu(E_f)}{\kappa T} \sum \int_{-\infty}^{\infty} d(E) \langle \xi(E) \rangle \zeta^2(E, E) \tag{3}
\]

in which \( D_v(E) \) is the density of state for the \( \nu \)-th band, \( f_\nu = f(1 - f) \), where \( f \) is the Fermi distribution function, and \( \zeta^2(E, E) \) is the average of the square of the velocity matrix, \( \langle v_1 | v_x | v_2 \rangle \), on the equi-energy surface \( E \), where \( v_1 \) and \( v_2 \) are the eigenstates of the \( \nu \)-th band. The velocity operator \( v_\nu \) was defined here as

\[
v_\nu = i \hbar \sum_{m \neq \nu} (X_m - X_\nu) \xi_{\nu,m} \beta^+_m \beta_m \tag{4}
\]

and, in the narrow band limit, \( \zeta^2(E, E) \) may be evaluated to

\[
\zeta^2(E, E) = \frac{a^2 A^2}{3 \hbar^2 N_\nu} \tag{5}
\]

where \( N_\nu = N(c_2 + c_v) \), \( A_\nu = \varepsilon Z(c_2 + c_v) \), in which \( \varepsilon \) is the nearest neighbour transfer energy and \( Z \) is the number of the nearest neighbour sites, therefore six in the present case, and thus \( A_\nu \) means the effective band width for the \( \nu \)-th band. When the density of state is rewritten in the form \( D_v(E) = \rho(E)N_c/A_\nu \), where \( \rho(E) \) should be about unity at the center of the band, the conductivity \( \sigma_\nu \) is written finally in a simple form

\[
\sigma_\nu = \frac{\pi e^2 \hbar^2}{18 \hbar} \frac{N \rho_\nu(E_f)}{\kappa T} \tag{6}
\]

or, by substituting \( a = 4.15 \times 10^{-8} \) cm,

\[
N = 1.4 \times 10^{22} \text{cm}^{-3}, \quad \sigma_\nu = 10^9 \rho^2(E_f) \text{ (ohm-cm)}^{-1} \tag{7}
\]

and \( \sigma \) is six times \( \sigma_\nu \). Therefore, \( \sigma \) can be near \( 10^4 \) (\( \Omega\)-cm\(^{-1} \)) in maximum and this is really observed in Y doped SmS by Penney and Holtzberg [15]. However, in stoichiometric SmB\(_6\), \( \sigma \) is about 5 (\( \Omega\)-cm\(^{-1} \)) at 4.2 K [7, 13] and is still decreasing gradually with decreasing temperature. This should be reckoned as follows. Similar to the usual impurity band, when the potential fluctuation exceeds the band width due to the translational energy, the 4f band states are localized. In SmB\(_6\), we expect that the translational 4f band width is only a few hundreds \( \kappa \), much smaller than that in SmX. While, we always expect some Sm defects which cause the potential fluctuation. Therefore, the hopping type conduction is rather expected for the 4f band in SmB\(_6\). For the samples with more Sm vacancies or \( R^{+++} \) doping, the potential fluctuation is rather smeared out and the conductivity increases more than one order of magnitude [7]. In the high pressure phase of SmS [16], the potential fluctuation is expected to be relatively higher than that of Y doped sample and thus the conductivity decreases to the order of \( 10^3 \) (\( \Omega\)-cm\(^{-1} \)). These are in the boundary between the metallic and the hopping 4f band conduction.

As temperature increases, the population on the conduction band increases and thus the conduction band conduction \( \sigma_c \) given by

\[
\sigma_c = 2e^2 \int D(E) \mu(E) f(E) dE \tag{8}
\]
is observed, in which \( \mu \) is the mobility. If we use the simplified model that \( D(E) \) is a constant, \( D_1 \), and \( \mu(E) = 0 \) for \( E < E_i \), and \( \mu(E) = \mu_s \) for \( E > E_i \), the experimental curve above 4 K in SmB\(_6\) is well fitted by the following value; \( E_i - E_{f_1} = 20 \) \( \kappa \), \( \mu_s = 100 \) cm\(^2\)/V s. and \( D_1 = 10^{34} \) erg\(^{-1}\) s\(^{-1}\) = 1.1 state/eV Sm site.

\[
\Delta H_I = \frac{\hbar}{g \mu_B T_1} = 4 \pi \int |I(E) D(E)|^2 f_1(E) dE \quad (9)
\]

in which \( \mu_B \) is the Bohr magneton number and \( I(E) \) the \( d-f \) exchange constant depending on the character of the conduction electron. Again by using a simplified model that \( D(E) = D_1 \), \( I(E) = 0 \) for \( E < E_i \), and \( I(E) = I \) for \( E > E_i \), we obtain good agreement with experiment with the values; \( I = 0.024 \) eV and \( E_i = 8 \) \( \kappa \).

Note that \( E_i \) is smaller than \( E_f \) because for the resonance relaxation mechanism the small mobility part has equal contribution and thus it pushes down the effective boundary of the conduction band.

The hyperfine splitting is large in Eu and thus the total line width \( \Delta H \) is given by \( (\Delta H_0^2 + \Delta H_1^2)^{1/2} \), in which \( \Delta H_0 \) is due to the hyperfine splitting and also the dipole field and is temperature independent while \( \Delta H_1 \) comes from \( T_1 \). Because Eu exists as the divalent ion, the amplitude of the conduction electron on Eu site is small and thus the effective \( d-f \) exchange constant \( I \) should be much smaller than that in Gd. Therefore, in Eu, we rather expect the relaxation process through the 4f band to be more important. For this purpose, eq. (9) should be modified as follows.

\[
|D(E)|^2 \rightarrow |D(E)|^2 = 3|D_1(E)|^2 \quad (10a)
\]
\[
I(E)^2 \rightarrow (g - 1)^2 J(J + 1) |I_s(0)|^2 / 3 \quad (10b)
\]

in which the \( g \)-factor for \( J = 2.5 \) is \( 2/7 \) and \( I_s(0) \) is the sum of the exchange interaction between a Eu spin and the neighbour Sm\(^{+++}\) spins. Because \( I_s(0) \) is connected with the \( g \)-shift and the susceptibility, we can estimate \( I_s(0) \) from these experimental values to be about 4 \( \kappa \). Then, to get best fit with experimental \( \Delta H \), the 4f band width \( D_1 \) is determined to be 400 \( \rho_s(E_f) \) \( \kappa \), which is very reasonable. In this way, the life times of Gd and Eu spins in SmB\(_6\) are explained satisfactorily both qualitatively and quantitatively on our model.

3. Metal-Insulator Transition in Doped SmS. — As the space is limited, we describe the mechanisms for the metal-insulator transitions in Y and Sm doped SmS very briefly.

In pure SmS, the energy gap between the 4f levels and the bottoms of the conduction band at \( X \) points is expected to be about 1.2 eV for the Frank-Condon condition, no lattice distortion. Taking into account the lattice distortion effect, 0.5 eV for optical phonon, 0.16 eV for acoustic phonon and 0.07 eV for the polaron effect on the conduction electron, the real gap is evaluated to be about 0.3 eV. Furthermore, by considering the single exciton formation, the gap, or the zero phonon line in the optical absorption, is reduced to be about 0.3 eV. This contradicts with Wachters conjecture [17], but is consistent with experiments by IBM group [1]. However, as shown by Kasuya [18], the single exciton is not the most stable state but the molecular magnetic exciton, MME, probably three pairs of single exciton, is more stable. This means that once an exciton is created, the bound electron polarizes the near neighbour 4f spins through the \( d-f \) exchange interaction and thus digs a potential of \( IS \sim 0.3 \) eV for the second exciton electron, but, because of the three fold degeneracy of the conduction band, the kinetic energy loss of the fourth electron is more than the energy gain due to the \( d-f \) exchange potential. Thus, the formation energy for such a MME is estimated to be about 0.1 eV/single pair.

When the pressure is applied, the 4f-\( t_{2g} \) energy gap shrinks in the ratio \( -8 \) meV/kbar and thus the instability of the MME occurs at about 12 kbar. Actually, however, the first order transition occurs before it, at about 7 kbar.

When \( Y \), or any \( R^{+++} \), is doped, one conduction electron is trapped to each \( Y \) atom and forms the magnetic impurity state, MIS, which is essentially equal to the stable single ME. Therefore, around \( Y \),

(1) T. Penney and F. Holtzberg. Private communication.
the total formation energy of MME is reduced by nearly 0.3 eV. However, single Y is still not enough to make the formation energy negative and the pair Y atoms at least sitting on the next nearest neighbour sites is enough to make the MME, or in the present case the molecular MIS, to be instable, in which three neighbour Sm change to Sm^{+++} thus five electrons are trapped in total. However, among them, two electrons are trapped so loosely that they are delocalized very easily and form free conduction electrons. The above picture explains the experimental facts [1] excellently.

When As is doped, the valence bands on the neighbour S are pushed up which causes the 4f levels on the neighbouring Sm to be pushed up through the p-f mixing effect, or the antibonding effect [9]. This is enough to push the 4f levels on the nearest neighbour Sm sites up to more than 0.1 eV so that the MME instable. Now, around an As atom, six Sm^{+++} and six electrons are formed among which three electrons are delocalized easily. This picture is again in excellent agreement with experiment [19]. The detail and more topics will be published in separate papers.

4. **Conclusion.** — The 4f states in EuB_{6} and SmB_{6} have been studied based on the extended Hubbard model. Transport properties and the EPR line widths of Gd and Eu in SmB_{6} are explained satisfactorily on the above model.

Metal-insulator transition mechanisms in SmS, as well as Y and As doped SmS, are studied based on the instabilities of the molecular magnetic exciton and the molecular magnetic impurity state in excellent agreement with experiments.

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References


[2] Private communication by ISHIZAWA, Y.


[8] See, for example, references 1 and 5.


