MAGNETIC FIELD INFLUENCE ON AUGER EFFECT ON SHALLOW DONORS IN CdF$_2$:Mn$^{2+}$ LUMINESCENCE

A. KAMIŃSKA AND A. SUCHOCKI

Institute of Physics, Polish Academy of Sciences
Al. Lotników 32/46, 02-668 Warsaw, Poland

Direct observation of the suppression of the Auger effect on shallow donors by the magnetic field in the luminescence of manganese ions in semiconducting CdF$_2$:Mn crystals is presented. The magnetic field decreases the probability of the Auger effect, which is spin-dependent energy transfer from the manganese ions to the electrons occupying shallow donors. This results in the increase in the decay times of the luminescence.

PACS numbers: 73.20.Hb, 73.50.Gr

The mechanism of the nonradiative recombination of the excited states of impurities in insulating crystals is a topic of interest in solid-state physics. One of them is energy migration on the system of excited ions and energy transfer to different types of defects.

In semiconductors, the presence of carriers opens a new recombination channel — the Auger effect (AE). The excitation energy of the localized dopant can be nonradiatively transferred either to the free carrier or the carrier bound to some defect center. In any case the energy accepting carrier is promoted during the AE high into the respective band (the conduction band for the electrons and valence band for the holes). This effect has been found in CdF$_2$ crystals doped with Mn$^{2+}$ and Y$^{3+}$ ions [1]. The Coulomb interaction, responsible for the AE, conserves the spin and its projection. Since magnetic impurity is involved in this AE, its probability should be dependent on spin polarization of particles participating in the process.

Mn$^{2+}$ ($3d^5$) in CdF$_2$ is a very efficient activator of a blue-green luminescence (2.44 eV) corresponding to transitions from the first excited state $^4T_{1g}$ ($^G$) to the $^6A_{1g}$ ($^S$) ground state of the Mn$^{2+}$ ion in a centrosymmetric environment. The radiative decay time $\tau_r$ of the Mn$^{2+}$ luminescence in CdF$_2$ is equal to 158 ms at 4.2 K [1].

CdF$_2$ is one of the crystals with the fluorite structure. In opposite to the other fluorite crystals, which are good insulators, CdF$_2$ may be converted to a low-resistance $n$-type semiconductor by doping with transition metal or rare earth
ions (substituting Cd ion). The crystal used in our experiment was doped with 1 mol% of Mn and 0.05 mol% of Y. It was converted to the conductivity state by the annealing in H₂ atmosphere (to remove the interstitial compensating fluorine ions). Free electrons liberated at the surface were trapped by Y ions and neutral shallow donors were formed. Their activation energy is about 110 meV for low concentration of donors [2].

The sample was placed into the superconducting Oxford magnet system, in the range of magnetic field up to 7 teslas. The luminescence was excited by the 337.1-nm line of a pulsed nitrogen laser (excitation pulse duration of about 10 ns), which is resonant with the intrashell 3d⁵ transitions of Mn²⁺ ions. The decay kinetics were measured with the use of Multichannel Scaler/Averager SR 430. The measurements were performed at 2.2 K.

In the presence of shallow donors the Mn²⁺ luminescence becomes weaker and the decay time shorter because of ΑE — the excitation energy transfer from the Mn²⁺ ions to the electrons. At temperatures above 40 K the free electron ΑE dominates. Its probability is proportional to the free carriers concentration and this effect is governed by the dipole–dipole interaction. Below 40 K, when practically all electrons are frozen out at the shallow donors, the energy is transferred to the donors, this causes their ionization with the electrons excited high into the conduction band. It is a resonant energy transfer, since the luminescence spectrum of the sensitizer ion (Mn²⁺) overlaps the absorption spectrum of the activator ions (in our case — the photoionization spectrum of the shallow donors). Because of continuous character of the photoionization spectrum of the shallow Y donors this resonance condition is fulfilled, although the photoionization energy of shallow donor is much lower than the energy of the Mn²⁺ luminescence.

It has been found that the ΑE due to the electron bound on shallow donors in CdF₂:(Mn, Y) is dominated by the exchange interaction [1]. Therefore the energy transfer probability P is exponentially dependent on the distance R, between the interacting centers

\[ P = P₀ \exp \left( -\frac{2R}{a_B} \right) = \frac{1}{\tau_r} \exp \left( -\frac{2R}{a_B} \right), \]  

where τ_r is the radiative lifetime of sensitizer ions, a_B is the shallow donor Bohr radius (a_B = 7 Å for CdF₂ [2]). The γ parameter describes the strength of the exchange interaction and it is equal to γ = 2R₀/a_B, where R₀ is the critical distance, at which the ΑE rate equals the radiative probability.

The value of γ depends on the magnetic field and in its absence is equal to 14 [1]. This value can be found by the fit of Inokuti–Hirayama model [3], which describes the luminescence decay kinetics. Figure 1 shows the dependence of ΑE probability on the magnetic field, derived from the fit of the Inokuti–Hirayama model to the experimental luminescence decay kinetics.

In the external magnetic field the energy levels of Mn²⁺ ions split into Zee- man sublevels. The probability of Auger effect depends on the spin polarization of shallow donors and manganese ions. In the magnetic field all thermalized electrons have the same spin-down polarization at low temperatures. The Coulomb interaction responsible for the Auger effect conserves the total spin and its projection. Therefore only transitions between the Zeeman sublevels of the ⁴T₁g and
$^6A_{1g}$ states of Mn$^{2+}$ ion with the same spin projection are allowed. The transitions between the sublevels with different spin projection are forbidden in the strong magnetic field since that would require different polarization of the spin of electron on the shallow donor and in the conduction band. This is the reason for the suppression of the AE in the presence of magnetic field.

The dependence of the probability of the Auger effect on the spin polarization of the particles can be found in the general form if we neglect any spin–orbital coupling in the system. According to the theoretical model of Nawrocki et al. [4] it is given by

$$P_A(B) \propto \sum_{\sigma, \sigma' = \pm 1/2} P_{\sigma\sigma}(B)P_{\sigma\sigma'}(B) \sum_{\mu=3/2} C_{\sigma\sigma'}^{\mu}p_{\mu}(B), \quad (2)$$

where $P_{\sigma\sigma}(B)$ is the probability for the electron to be in the state with spin projection $\sigma$

$$P_{\sigma\sigma}(B) = Q_e^{-1}\exp\left(-\frac{g\mu_BB}{kT}\sigma\right), \quad Q_e = \sum_{\sigma=\pm1/2} \exp\left(-\frac{g\mu_BB}{kT}\sigma\right). \quad (3)$$

The relative probabilities of the Auger transition for different initial spin polarizations $C^{\mu\sigma\sigma'}$ can be calculated from the Clebsh–Gordan coefficients.

The probability for the Mn$^{2+}$ ion to be in the state with spin projection $\mu$, $p_{\mu}(B)$ is expressed by

$$p_{\mu}(B) = \frac{\exp(-\lambda\mu)}{Q_{\text{Mn}}}, \quad \lambda = \frac{g_{\text{Mn}}\mu_BB}{kT},$$

$$Q_{\text{Mn}} = \sum_{\mu=-3/2}^{3/2} \exp(-\lambda\mu) = \frac{\sinh(2\lambda)}{\sinh(\lambda/2)}. \quad (4)$$

The solid line in Fig. 1 represents the computer fit of Eq. (2) to the experimental data of the AE probability dependence on the external magnetic field $B$.

In conclusion, we have observed the suppression of the AE by the magnetic field applied to the CdF$_2$:(Mn,Y) crystal, which is explained by spin polarization.
of electrons involved in the AE and Mn$^{2+}$ ions. The presented theory is in very good agreement with the experimental data.

The authors are indebted to Dr. Marek Potemski of IMP-CNRS High Magnetic Field Laboratory in Grenoble, France, for valuable discussion.

References