Measurement of Very Small
Zeeman Splittings in GaN:Mn,Mg
by Faraday Rotation

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In this work we demonstrate an application of Faraday rotation for measuring an extremely small Zeeman splitting of an Mn related absorption line placed at 1.417 eV in optical absorption spectrum of Mn and Mg doped gallium nitride. Analysis of the collected spectra allowed us to determine the value of the splitting as equal to $0.12 \pm 0.01$ meV at 6 T. This data should help in establishing the nature of the observed absorption band.

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1. Introduction

Faraday rotation can be used for a precise analysis of magnetic field induced variation of sharp absorption lines, both in energy position (Zeeman splitting) and in intensity [1]. In this work we demonstrate an application of the Faraday rotation for measuring an extremely small Zeeman splitting of a sharp Mn related infrared line in optical absorption spectrum of GaN:Mn,Mg, occurring in a moderate field of a few tesla.

2. Sample and experiment

Single GaN:Mn,Mg crystal was grown by equilibrium high pressure technique from nitrogen solution in liquid gallium. Manganese and magnesium were added
into gallium during growth. Secondary ion mass spectroscopy (SIMS) analysis determined content of magnesium as equal to about 0.1% and content of manganese below 0.001%.

Transmission and Faraday rotation were measured in a standard setup with the sample mounted strain-free in a superconducting magnet and immersed in superfluid helium. To obtain each Faraday rotation spectrum a series of transmission spectra was measured for various positions of the polarizer placed in front of the spectrograph (Fig. 1). Faraday rotation was determined by fitting parabolas (Fig. 2) to the dependence of the transmitted light intensity on the analyzer angle in the vicinity of the minimum transmission, as described in Ref. [1].

![Fig. 1. Idea of Faraday rotation determination by measuring a series of transmission spectra at different positions of the analyzer.](image1.png)

![Fig. 2. Determination of the Faraday rotation spectrum by finding transmission minima vs. rotation angle.](image2.png)

The analysis of the spectra was based on the relation between the logarithm of the transmission and the doubled phase of the transmitted light [1], representing respectively the real and imaginary part of the same analytic function.

3. Results and analysis

Near infrared absorption spectrum of the sample is presented in Fig. 3. A sharp zero-phonon line (ZPL) at 1.417 eV energy is followed by structures that
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reflect phonon density states of gallium nitride. Origin of the presented band is not fully established yet. It has been primarily attributed to photoionization transition from GaN valence band to bound Mn acceptor state [2]. Character of the band is however typical of intra-impurity transition of transition metal ions observed in GaN [3–5]. Thus an internal transition within $d$ shell of Mn ion cannot be excluded as the origin of the band.

In Fig. 4 transmission spectrum of GaN:Mn,Mg in energy range of ZPL together with a Lorentzian fit is presented. Lorentzian fit with a parabolic background reproduces well the spectrum, according to the equation

$$\ln(I(\omega)) = A\text{Re}\left(\frac{i}{\omega_0 - \omega + i\gamma}\right) + C\omega^2 + D.$$ 

The determined values of the amplitude $A$, resonance energy $\hbar\omega_0$ and HWHM width $h\gamma$ are given in the onset of Fig. 4. They were used subsequently for the analysis of the rotation spectrum, measured at 6 T (Fig. 5).
In principle, three possible sources of the Faraday rotation related to a given absorption line can be distinguished, namely differences between its two circularly polarized components in energy (Zeeman effect), in intensity, and/or in width. As pointed out in Ref. [1], for a symmetric line, the Zeeman mechanism produces also a symmetric rotation spectrum, whereas the two other mechanisms lead to asymmetric, dispersive type shapes. We found that the measured rotation spectra can be well described by the symmetric Zeeman type shape, therefore we exclude the other mechanisms from the interpretation. The experimental Faraday rotation spectrum was fitted with the expression

\[ \hat{\theta}(\omega) = \frac{1}{4} \Delta \omega A \frac{\partial}{\partial \omega_0} \text{Im} \left( \frac{i}{\omega_0 - \omega + i\gamma} \right) + F \omega^2 + G, \]

using the line parameters determined previously from the zero-field transmission spectrum (allowing a slight readjustment of \( \omega_0 \) for the shift in magnetic field). A parabolic background \( F \omega^2 + G \) was also allowed. The expression was obtained assuming a Zeeman splitting into two components of opposite circular polarizations, separated by \( \Delta \omega \) much smaller than the line width. The factor 1/4 results

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**Fig. 5.** Faraday rotation spectrum of the zero-phonon line. Lorentzian fit (solid) of the experimental data (points) produces Zeeman splitting value of 0.12 meV.

**Fig. 6.** Direct absorption experiment in magnetic field 6 T reveals Zeeman splitting of the zero-phonon line of about 0.13 meV.
from taking into account the fact that the rotation angle is equal to half of the phase shift between the two components. A good fit was obtained, yielding Zeeman splitting value $\hbar \Delta \omega = 0.12$ meV, indeed much smaller than the line width. This result is consistent with direct observation of the splitting between the two circular components (Fig. 6).

4. Conclusions

Faraday rotation experiment combined with direct absorption measurements performed on a GaN:Mn,Mg bulk crystal allowed us to reveal a very small influence of the magnetic field on a zero-phonon line at 1.417 eV. Assuming a doublet structure of the line we obtained a value of $0.12 \pm 0.01$ meV for its Zeeman splitting at 6 T. This data should help in establishing the nature of the observed absorption band.

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References