Roever, M.; Malindretos, J.; Bedoya-Pinto, A.; Rizzi, A.; Rauch, C.; Tuomisto, F.

Tracking defect-induced ferromagnetism in GaN:Gd

Published in:
Physical Review B

DOI:
10.1103/PhysRevB.84.081201

Published: 01/08/2011

Please cite the original version:
Tracking defect-induced ferromagnetism in GaN:Gd

Martin Roever, Joerg Malindretos,* Amilcar Bedoya-Pinto, and Angela Rizzi

IV. Physikalisches Institut, Georg-August-Universität Göttingen, Göttingen, Germany

Christian Rauch and Filip Tuomisto

Department of Applied Physics, Aalto University, Aalto, Finland

(Received 22 March 2011; published 10 August 2011)

We report on the magnetic properties of GaN:Gd layers grown by molecular beam epitaxy. A poor reproducibility with respect to the magnetic properties is found in these samples. Our results show strong indications that defects with a concentration of the order of $10^{19}$ cm$^{-3}$ might play an important role for the magnetic properties. Positron annihilation spectroscopy does not support a direct connection between the ferromagnetism and the Ga vacancy in GaN:Gd. Oxygen codoping of GaN:Gd promotes ferromagnetism at room temperature and points to a role of oxygen for mediating ferromagnetic interactions in Gd-doped GaN.

DOI: 10.1103/PhysRevB.84.081201 PACS number(s): 71.15.Eq, 76.30.Kg, 75.50.Pp, 78.70.Bj

The origin of the ferromagnetic order in GaN:Gd is still unknown. It was shown that cation vacancies can provide local moments which exchange directly by their overlapping wave functions and provide a long-ranged coupling mechanism. According to Neugebauer et al., the cation vacancy in GaN is a rather unlikely defect if the Fermi energy is not close to the conduction-band minimum. Furthermore, nitrogen or oxygen interstitials in GaN:Gd can also provide local moments with a long-ranged coupling and their formation is much more likely than the cation vacancy.

In this Rapid Communication we show experimental results about the reproducibility and long-term stability of the ferromagnetic phase in Gd-doped GaN layers. The theory of the Ga vacancy model is probed experimentally by positron annihilation spectroscopy (PAS). Oxygen codoping of GaN:Gd was carried out to introduce oxygen impurities on purpose. All GaN:Gd samples were grown by plasma-assisted molecular beam epitaxy on metalorganic chemical vapor deposition (MOCVD) GaN/Al$_2$O$_3$(0001) templates. If not stated otherwise, the MBE process was carried out with our optimized parameters for the growth of unintentionally doped GaN layers, namely, $760 \degree$C substrate temperature and slightly metal-rich conditions. Gd (4N) was supplied during growth by a high-temperature effusion cell. Oxygen codoping was carried out applying oxygen gas (4.8 sccm) with a flux ranging from 0.1 to 0.5 sccm into the growth chamber. The growth time was kept constant for all epitaxial layers, resulting in a thickness of $\sim 500$ nm for the standard GaN:Gd layers and $200$ nm for the oxygen codoped samples.

A superconducting quantum interference device (SQUID) MPMS 5 from Quantum Design was used to measure the magnetic properties. All data have been corrected for diamagnetic background and trapped field artifacts. The resolution limit is $\sim 3 \times 10^{-7}$ emu, which translates to $\sim 5 \times 10^{18}$ $\mu_B$/cm$^3$ for our sample geometry. Reference measurements on the substrate material were made on a regular basis and no ferromagnetic contributions were found.

The Gd concentration is measured by time of flight secondary ion mass spectroscopy. The resolution limit of Gd isotopes is $\sim 10^{16}$ cm$^{-3}$. All lower concentrations are extrapolated by effusion theory. The contamination with transition metals can be estimated to be at least a factor of $10^2$ lower than the Gd concentration.

The Doppler broadening of the positron annihilation radiation was measured by two germanium detectors with a...
Oxygen codoping of three samples (blue diamonds) was found to induce a ferromagnetic order at room temperature. On the other hand, several samples that have been grown later with the same nominal growth parameters (black circles) show a vanishing saturation magnetization at room temperature. The dependence of the effective saturation magnetization on the Gd concentration is shown in Fig. 2 and compares well with values in the literature. The observed trend could be described by a constant additional magnetic moment $M_{\text{eff}} = 2.5 \times 10^{19} \mu_B / \text{cm}^3$ independent of the Gd concentration, according to the expression $m_{\text{Gd,eff}} = m_{\text{Gd}} + M_{\text{def}} / c_{\text{Gd}}$. Assuming a defect-driven mechanism of the ferromagnetism, an effective defect concentration of this order would be required. In this scenario colossal magnetic moments could be explained with a high but not unrealistic concentration of magnetically active defects. However, it is unclear why these defects only form in the presence of Gd while their amount would be independent of the Gd concentration.

Electrical-transport properties of GaN:Gd grown on highly resistive 6H-SiC(0001) substrates show that even Gd concentrations lower than the background carrier concentration of typically $10^{17} \text{ cm}^{-3}$ increase the resistivity by several orders of magnitude. The growth conditions of these samples were optimized in the conventional line-shape parameters $S(\langle E_\gamma - 511 \text{ keV} \rangle < 0.75 \text{ keV})$ and $W(2.86 \text{ keV} < \langle E_\gamma - 511 \text{ keV} \rangle < 7.33 \text{ keV})$. All points have been normalized to the value of a suitable reference sample where positrons annihilate solely in the delocalized state in the GaN lattice.

The dependence of the effective saturation magnetization $M_{\text{eff}}$ on the Gd concentration was also pointed out in experiments with Gd-implanted GaN. A spectrum of positron annihilation radiation is analyzed using the conventional line-shape parameters $S(\langle E_\gamma - 511 \text{ keV} \rangle < 0.75 \text{ keV})$ and $W(2.86 \text{ keV} < \langle E_\gamma - 511 \text{ keV} \rangle < 7.33 \text{ keV})$. All points have been normalized to the value of a suitable reference sample where positrons annihilate solely in the delocalized state in the GaN lattice.

The structural quality of the Gd doped epitaxial layers below a Gd concentration of $10^{20} \text{ cm}^{-3}$, at which GdN clusters are formed, resembles those of unintentionally doped GaN, as deduced from x-ray diffraction analysis as well as transmission electron microscopy (not shown). An x-ray absorption near edge structure (XANES) analysis revealed that the Gd is incorporated substituional on Ga sites in a 3+ valence state and no significant bond length variations of the Ga-N bond were found.

In Fig. 1(a) the saturation magnetization $M_s$ at 300 K measured for many GaN:Gd epitaxial layers is plotted against the sample ID, i.e., in chronological order. In all samples the Gd concentration is in the range of $10^{14}$–$10^{19} \text{ cm}^{-3}$. The series around sample number J0180 clearly shows sizable $M_s$ values. On the other hand, several samples that have been grown later with the same nominal growth parameters (black circles) show a vanishing saturation magnetization at room temperature, with the exception of J0264 and J0265. The green triangles correspond to samples grown in a wide range of III-V ratios, slightly different substrate temperatures, and within the same range of Gd concentrations, but no influence of these parameters on the magnetic properties was found. Oxygen codoping of three samples (blue diamonds) was found to induce a ferromagnetic order at room temperature with $M_s$ values comparable to that of the first series. The saturation magnetization value increases with the oxygen supply during growth. It should be noted that GaN:Gd samples without oxygen codoping grown directly before and after the codoped samples exhibit no ferromagnetism. Selected samples have been repeatedly characterized by SQUID. The saturation magnetization is observed to decrease with time [Fig. 1(b)].
The observed variable-range hopping in an impurity band of magnitude as compared to unintentionally doped GaN.20

Secondary ion mass spectroscopy (not shown) reveals oxygen concentrations above \(5 \times 10^{16} \text{ cm}^{-3}\) in our GaN:Gd layers (J0175, J0188) and clearly higher values for the oxygen codoped samples (J0273–J0275). The high resistivity of the GaN:Gd layers indicates that the oxygen atoms are not mainly incorporated substitutionally on nitrogen sites, where they would form shallow donors.

With the aim of investigating possible correlations between ferromagnetism in GaN:Gd and the presence of cation vacancies, we performed positron annihilation spectroscopy (PAS). Depth-dependent Doppler broadening spectra have been recorded at room temperature (Fig. 3). From these, the \(S\) and \(W\) line-shape parameters representative for the GaN:Gd layers are determined and displayed in Fig. 4, together with characteristic values for the GaN lattice and the gallium vacancy.23 In order to ensure that the determined parameters at room temperature are representative for the vacancy concentration in the samples and not influenced by positrons detrapping from vacancies to negative ions,24 temperature-dependent measurements up to 550 K have been performed for selected samples (Fig. 4, small triangles). The reference value for the GaN lattice is determined by measuring a reference sample in which no positron trapping to vacancy defects is observed (Fig. 3).

The identity of dominant positron traps in a certain sample can be evaluated based on its position in the \(SW\) plot. The latter is given as the linear combination of the characteristic values of the involved annihilation states, weighted with the fraction of positrons annihilating in these states. The measured \(SW\) points for the GaN:Gd layers indicate a broad defect landscape in the different samples which is dominated by trapping to vacancy clusters. The strong deviation from the Ga vacancy line and high relative \(S\) parameters of up to 1.09 are characteristic for vacancy clusters with a considerably larger open volume than the isolated Ga vacancy. The relevant vacancy cluster point is not known exactly, but similar \(S\) parameters have been observed before in MBE-grown GaN.25 We can estimate the vacancy cluster density to lie in the range between \(1 \times 10^{16} \text{ cm}^{-3}\) and \(1 \times 10^{18} \text{ cm}^{-3}\). Since the accurate \(SW\) reference point for these defects is not known, the upper bound could even be lower by about one order of magnitude.

Considering only the ferromagnetic samples in Fig. 4 (red circles), the strongest cluster signal is found for the sample with the lowest Gd concentration (J0176, \(c_{\text{Gd}} = 1.5 \times 10^{14} \text{ cm}^{-3}\)) and the cluster signal decreases with increasing Gd doping. In the sample with the most stable ferromagnetism with time and highest absolute saturation magnetization (J0175, \(c_{\text{Gd}} = 9.6 \times 10^{17} \text{ cm}^{-3}\)), the measured \(SW\) point is located close to the GaN lattice and hence no positron trapping to vacancy defects is observed. At room temperature the positron diffusion length in this sample is considerably reduced compared to the GaN reference (Fig. 3). This hints at the presence of a high concentration of negatively charged ions,24 which can trap positrons through the formation of Rydberg states and possess annihilation characteristics close to the GaN lattice. It should be noted that detrapping from vacancy defects could be explicitly excluded in this sample, due to the temperature-dependent measurements. For the nonmagnetic samples the PAS results do not evidence any clear correlation between the Gd concentration and the vacancy type or concentration. Sample J0225, whose Gd content of \(5 \times 10^{16} \text{ cm}^{-3}\) is comparable to that of J0175, lies close to the saturation point for single Ga vacancy-related defects. Nevertheless, this sample never showed any ferromagnetic behavior. So, the defect landscape that has been probed by PAS, namely, single gallium vacancies and larger open...
volume defects, does not directly correlate with the magnetic properties of the GaN:Gd epitaxial layers. We can therefore conclude that the cation vacancy models proposed in literature are not supported by our experiments.1–3

Based on formation energy arguments, interstitial nitrogen as well as oxygen in octahedral sites next to Gd have been proposed to be a more likely source of defect-induced magnetism in GaN:Gd than cation vacancies.4 Both types of interstitials induce a spin split band of localized states originating from the nonbonding N-2p or O-2p orbitals, which lies well within the energy gap. Both configurations would be consistent with the observed compensation of the unintentional n-type dopants in GaN:Gd and with the electrical transport by hopping.20 Due to the large affinity of Gd to unintentional donors, which lies well within the energy gap. Both configurations between the native donors and the compensating defects. If our samples could be explained by an uncontrolled balance of magnetic moments. The efficient gettering of residual donors in GaN than cation vacancies.4 Both types of interstitials induce a spin split band of localized states originating from the nonbonding N-2p or O-2p orbitals, which lies well within the energy gap. Both configurations would be consistent with the observed compensation of the unintentional n-type dopants in GaN:Gd and with the electrical transport by hopping.20 Due to the large affinity of Gd to O, it is also possible that oxygen donors are attracted into interstitial positions by the Gd, thus further contributing to the compensation and providing extra localized magnetic moments. The efficient gettering of residual donors in GaN by Gd was also reported for bulk samples grown under high pressure26 and was theoretically confirmed by Mitra and Lambrecht for the case of substitutional oxygen.4

The nonreproducibility of the ferromagnetic properties in our samples could be explained by an uncontrolled balance between the native donors and the compensating defects. If the concentration of localized magnetic moments were near the percolation threshold, small variations of their amount could switch the macroscopic ferromagnetic state, as discussed in the Appendix of Osorio-Guillen et al.5 In this case the favorable effect of codoping with oxygen during growth, as seen for the samples in Fig. 1 (blue diamonds), might be explained by an increase of interstitial oxygen in presence of Gd. Therefore, the model of a ferromagnetic state induced by interstitial oxygen is at least qualitatively consistent with our data. It should be noted that attributing the favorable effect of oxygen codoping to carrier induced ferromagnetism due to ON donors seems rather unlikely considering the high resistivity of ferromagnetic GaN:Gd.9,20

In conclusion, room-temperature ferromagnetism and colossal magnetic moments of dilute GaN:Gd layers grown by MBE are confirmed. However, the reproducibility and the long-term stability of the magnetic properties are poor. Uncontrolled parameters seem to be responsible for the ferromagnetic properties. It is shown that a concentration of magnetically active defects of $\sim 10^{19} \text{cm}^{-3}$ might explain the observed magnetic properties. PAS measurements rule out single gallium vacancies as the origin of magnetic coupling in as-grown GaN:Gd and do not show any direct connection of gallium vacancy clusters to room-temperature ferromagnetism in this material either. Oxygen codoping is observed to promote ferromagnetism in GaN:Gd. However, more investigations are necessary to indubitably identify the role of oxygen and the magnetic coupling mechanism in this material.

We thank Andreas Laufer (Justus-Liebig-Universität Gießen) for the SIMS measurements and the Deutsche Forschungs Gesellschaft for funding within the SFB 602.

---