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Spatially Correlated Distributions of Local Metallic Properties in Bulk and Nanocrystalline GaN

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We compare local electronic structure at different atom types of a metallic semiconductor in bulk and nanocrystalline form. Multinuclear magic-angle-spinning nuclear magnetic resonance (MAS NMR) establishes that GaN synthesized as an intentionally-doped bulk powder or as annealed nanocrystalline particles exhibits metallic behavior and a wide distribution of differing electronic environments in both forms. Bulk polycrystalline wurtzite GaN doped with 0.13% Ge as a shallow donor exhibits a temperature-independent distribution of ⁷¹Ga Knight shifts over the temperature range 123-473 K. Each Knight shift frequency in the inhomogeneously-broadened spectrum is characterized by a 71 Ga spin-lattice relaxation time T_1 that is in good agreement with the value predicted by the Knight-Korringa relation across the broad range of temperatures. The ¹⁴N spectrum shows a slightly smaller Knight shift distribution with spin-lattice relaxation time T_1 values at 295 K across the distribution also in good agreement with the Knight-Korringa relation. Similarly, annealed nanocrystalline wurtzite GaN (50-100 nm, and without Ge) exhibits a ⁷¹Ga Knight shift distribution and T_1 values (at 295 K) that follow the same Knight-Korringa behavior. Thus, both bulk and nanocrystalline forms of GaN are n-type and well above the metal-insulator transition (MIT), the nanocrystals most likely as a result of incorporation of shallow donor oxygen atoms during synthesis. Carriers in both forms of sample exhibit the near-ideal characteristics of a degenerate Fermi gas of non-interacting spins. The observation of NMR signals from both atom types, Ga and N, allows for the direct spatial correlation of the local electronic structure at the two sites in the lattice, specifically the s-orbital character of the electronic wavefunction of conduction band electrons at the Fermi edge. The relative values of these carrier wavefunction probabilities (nearly twice as great for the N atom as for the Ga) are in line with theoretical predictions. Analyses of ⁷¹Ga, ¹⁴N, and ¹⁵N NMR results, including double-resonance 2D ¹⁵N{⁷¹Ga} measurements, reveal electronic disorder in the form of broad distributions of local metallic properties (Knight shifts) that are shown to be spatially correlated on a sub-nanometer scale.

I. INTRODUCTION

Semiconductors heavily doped with shallow donors become metallic, acquiring useful electronic and photonic properties. Mott provided a quantitative estimate of the donor concentration required to induce such a transition [1], known as the Metal-Insulator (MIT) or Mott transition. Physically, this corresponds to a transition between two extrema: 1) dilute isolated and independent shallow donors, that at low temperatures are un-ionized and have weakly-bound electrons whose wavefunctions [2] typically extend over several lattice constants; and 2) concentrated and interacting shallow donors, whose overlapping wavefunctions form an impurity band that can lie below the conduction band or, at higher concentrations, enter into it. Magnetic resonance techniques based upon observing either the electron spin, e.g., electron paramagnetic resonance (EPR) or the nuclear spin via the electron spin, e.g., electron-nuclear double-resonance (ENDOR) have provided very detailed insights on local electronic structures in the first regime [3–6], but not the second. Detailed experimental and theoretical understanding of the

spatially-varying electronic structures of heavily-doped semiconductors is challenging because of the electronic disorder arising from the assumed random substitutions of shallow donors in the lattice [7].

One of the few means of observing such electronic disorder over the atomic scale is provided by magnetic resonance techniques that exploit well-understood interactions between electron and nuclear spins. Extensive magnetic resonance studies on doped Si around the MIT, including nuclear magnetic resonance (NMR) studies [7– 14] have yielded important insights into local electronic structure. However, an unambiguous interpretation of the NMR results has proven difficult because of complications due to formation of impurity bands distinct from the conduction band [12, 13]. Furthermore, no direct experimental evidence has yet been obtained for spatial variations in the electronic structure about individual silicon atoms in metallic doped samples.

By working with the compound semiconductor hexagonal (or wurtzite) gallium nitride (h-GaN), which has technologically-important applications in solid-state lighting and power electronics [15, 16], we have for the first time directly correlated spatial variations in the electronic structures of constituent atoms in semiconductor samples well above the MIT. The insights come from two

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main sources. There are frequency displacements, called Knight shifts (K), of the ⁷¹Ga, ¹⁴N and ¹⁵N NMR signals that arise here from nuclear hyperfine interactions with electrons in the conduction band having *s*-orbital character [17]. There are also associated contributions to the overall spin-lattice relaxation rate T_1^{-1} from the Korringa relaxation term $T_{1,K}^{-1}$ [17]. In this work we show that both bulk polycrystalline

GaN prepared from an alkali metal flux containing Ga and Ge under N_2 pressure and containing 0.13 wt% Ge as a substitutional n-type dopant (GaN:Ge) [18],[19], as well as annealed nanocrystalline GaN powders, are highly metallic, i.e. are well above the MIT. Our results resolve widely differing interpretations of the electronic structure of nanocrystalline GaN powders manifested by their ⁶⁹Ga, ⁷¹Ga and ¹⁵N NMR spectra [14, 20-26]. The results also represent a detailed electronic structural characterization of a degenerately doped semiconductor in bulk and nanocrystalline forms. While both GaN:Ge and nanocrystalline GaN materials exhibit properties that are characteristic of degenerate Fermi gases of non-interacting spins, the local metallic properties, as reflected by the Knight shifts and Korringa relaxation rates, vary widely over sub-nanometer length scales. We determine the ratio of s-orbital carrier wavefunction probabilities at the Fermi edge, with the N atoms having significantly more s-character than the Ga atoms regardless of the local metallic property. Furthermore, the spatial correlation of local metallic properties at different neighboring spin pairs (in this case 71 Ga and 14 N or ¹⁵N) provides a strong indication that the Knight shift distributions are due to electronic disorder related to the proximities to randomly located shallow donors.

II. RESULTS AND DISCUSSION

A. Bulk GaN:Ge Powder

1. ⁷¹Ga and ¹⁴N Knight shifts

Both ⁷¹Ga and ¹⁴N magic-angle spinning (MAS) NMR spectra of GaN:Ge show broad Knight-shifted signals arising from distributions of ⁷¹Ga and ¹⁴N electronic environments. Figure 1(a) shows the central transitions (+1/2 to -1/2) of six overlaid ⁷¹Ga spectra acquired at temperatures from 123 to 473 K. The small narrow peak at ca. 332 ppm corresponds to undoped h-GaN [27] and is assigned to regions of the sample that experience no Knight shift. A magnification of this region for the spectra collected at all six temperatures is shown in Figure 2(a). The considerably broader ⁷¹Ga signals dominating the spectra arise from regions of the sample having different Knight shifts [18]. Unlike doped Si above the MIT [28], the distribution of Knight-shifted intensity is essentially temperature-independent, as discussed below.

Figure 2(b) shows that the chemical shift of the zero Knight shift peak has a weak dependence upon tem-



FIG. 1. (Color online) MAS-NMR spectra of bulk GaN:Ge powder at 11.7 T: (a) Overlaid ⁷¹Ga spectra (5 kHz MAS, 15 s delay) at different temperatures from 123 K to 473 K; see Fig. 2(a) and Fig. 2(c) for expanded views of the peak at K=0 and of the left edge region indicated by the small box; (b) ¹⁴N rotor-synchronized Hahn-echo NMR spectrum (12 kHz MAS, 20 s delay) acquired at 295 K. Spinning sidebands (SB) are indicated by asterisks. (c) Comparison of the Knight shift distributions of ⁷¹Ga and ¹⁴N (obtained by summing the CB and 3 pairs of SBs), with normalized peak heights and with an x-axis corresponding to the ⁷¹Ga Knight shift and the ¹⁴N Knight shift scaled by a factor of 1.059 to give the best overlap.

perature, varying linearly with T^2 . This likely reflects the quadratic temperature dependence of the hexagonal GaN lattice parameters [29]. When this temperature dependence is taken into account in referencing the ⁷¹Ga Knight shift distributions to the position of this particular peak maximum at each temperature, the Knight shifts are clearly independent of temperature. This is shown by the comparison of the same region on the left edge of the



FIG. 2. (Color online) (a) ⁷¹Ga MAS NMR spectra of GaN:Ge at different indicated temperatures, showing an enlargement of the zero-Knight shift peak region for the spectra shown in Fig. 1(a). The six spectra are offset for clarity. (b) Plot of the peak position of the zero-Knight shift peak as a function of T^2 . The dotted line has a slope of -1.81 x 10^{-5} ppm/K² and represents a linear best-fit to the data points. (c) ⁷¹Ga MAS NMR spectra of GaN:Ge at different indicated temperatures, showing the temperature dependence of a high-frequency region of the spectra; (d) The same highfrequency region as (c), with the x-axis rescaled such that the maximum of the zero-Knight shift peak shown in (a) has a shift of 0 ppm at each temperature. After correction for the small temperature dependence of the zero-Knight shift peak, the ⁷¹Ga Knight shift is thus shown to be temperatureindependent across the inhomogeneously-broadened spectrum shown in Fig. 1(a).

broad 71 Ga peaks before (Figure 2(c)) and after (Figure 2(d)) application of the temperature-dependent correction.

We use this temperature-independence of the Knight shift K over the temperature range 123-473 K to estimate a lower bound for the Fermi level in GaN:Ge. We use Eq. 5 of reference 13, rewritten below, to obtain the factor (in square brackets) involving $(T/T_F)^2$ that scales the Knight shift at different temperatures T depending upon the value of the Fermi temperature T_F , for the situation where $T < T_F$:

$$K = 2\pi \left(\frac{\gamma_s^2 \hbar^2}{2}\right) \left(\frac{P_f}{k_B T_F}\right) \left[1 - \frac{\pi^2}{12} \left(\frac{T}{T_F}\right)^2\right]$$
(1)

Here γ_s is the electron magnetogyric ratio, \hbar is Planck's constant, and P_f is a measure of the average electronic probability density at the nucleus; we assume that the derivation of this equation applied to ³¹P in Si:P is valid for ⁷¹Ga in GaN. Assuming a T_F of 4500 K yields a scaling factor of 0.991 at 473 K and 0.999 at 123 K, the two extremes of temperature. The 0.008 difference in the scaling factors would result in a 1.2 ppm change in the Knight shifts at the 155 ppm edge shown in Figure 2(d). Since any actual change in the Knight shift with temperature must be smaller than the measurement uncertainty, ca. 1.2 ppm, we conclude that T_F is greater than 4500 K, which corresponds to a Fermi energy of ca. 400 meV above the bottom of the conduction band. Thus, the GaN:Ge doping is extremely degenerate $(T_F/T > 10)$ over the range of temperatures investigated, in agreement with further evidence from relaxation analyses below. A Fermi level at least 400 meV above the bottom of the conduction band is plausible, since DFT calculations for GaN doped with a carrier concentration of 1×10^{19} $\rm cm^{-3}$ give a Fermi level of 125 meV above the conduction band minimum [30], whereas the GaN:Ge sample is estimated to have a carrier concentration in the range $0.7-1.8 \times 10^{20}$ cm⁻³ [18]. These estimated values are well above the MIT for n-type GaN:Si of 1.6×10^{18} cm⁻³ [31] and theoretical predictions for h-GaN of 1×10^{18} cm⁻³ [32].

The ¹⁴N spectrum of GaN:Ge in Figure 1(b) corroborates and extends these conclusions. It has a centerband region showing a weak narrow signal at -302 ppm, the same shift as observed for undoped h-GaN [27]. A dominant inhomogeneously-broadened peak is also present at -202 ppm, reflecting a distribution of Knight shifts. In addition to the centerband, a set of spinning sidebands spaced at integer multiples of the spinning rate (12 kHz) reflect the ¹⁴N nuclear quadrupole coupling constant (NQCC) arising from (first order) interactions between the nuclear electric quadrupole moment and the electric field gradients at the nucleus. The NQCC is axially symmetric (asymmetry parameter $\eta=0$), with a quadrupole coupling constant C_q (= e^2Qq/h)of 24.14 kHz at 296 K [33].

The inset in Figure 1(b) shows a region centered on the +1 sideband superimposed upon an equivalent region of the centerband and normalized to the same peak height. The greater intensity on the left side of the sideband spectrum indicates that those regions of GaN:Ge having larger Knight shifts also have significantly altered anisotropic interactions that cause their sideband intensities to increase relative to the centerband. In principle these altered interactions could result from changes in the electric field gradients at the ¹⁴N sites that change the NQCC or from an increase in the ¹⁴N chemical shift anisotropy from its value of zero in an undoped GaN film [33]. Although the latter cause seems less likely, the corresponding increase in intensity of Knight-shifted regions seen for the -1 sideband is somewhat less than that for the +1 sideband, an asymmetric behavior that is not

expected for alterations of the NQCC alone. This correlation between the charge (NQCC, or possibly chemical shift as well) and the spin (Knight shift) effects of dopants has been seen in ⁷¹Ga spectra of GaN films [34] and suggests that the most Knight-shifted ¹⁴N and ⁷¹Ga nuclei are closest to the dopants.

Complementary analyses of the ⁷¹Ga and ¹⁴N Knightshift distributions directly relate the conduction electron density at Ga and N lattice sites. The Knight shift Kdue to the contact interaction in a Fermi gas of noninteracting electron spins, after replacing the spin contribution to the susceptibility χ_e^s by $\gamma_e^2 \hbar^2 \rho_0(E_F)/2$, is described by [17]:

$$K = \frac{4\pi}{3} \gamma_e^2 \hbar^2 \langle |u_k(0)|^2 \rangle_{E_f} \rho_0(E_f)$$
⁽²⁾

Here, $\langle |u_k(0)|^2 \rangle_{E_f}$ is the probability of an electron at the Fermi level in an *s*-orbital being at the nucleus, $\rho_0(E_f)$ is the density of states (DOS) of this *s*-orbital at the Fermi level, γ_e is the electron gyromagnetic ratio, and \hbar is Planck's constant. Interestingly, after scaling the ¹⁴N frequency axis by a factor of 1.059, the shapes of the ⁷¹Ga and ¹⁴N intensity distributions are nearly identical, Figure 1(c). This suggests that each ⁷¹Ga nucleus associated with a particular ⁷¹Ga Knight shift isochromat has neighboring ¹⁴N nuclei with correlated Knight shifts. The scaling factor provides a direct experimental means of comparing, at both atomic sites, the relative *s*-orbital characters of the conduction band electrons at the Fermi edge.

2. Comparison of electronic projected densities of states at ^{71}Ga and ^{14}N lattice sites

The probability $\langle |u_k(0)|^2 \rangle_{Ef}$ for electron wavefunctions at Ga and N atoms can be estimated from theoretical calculations of atomic hyperfine coupling constants of atoms [35], parameters of great interest in EPR spectroscopy. For unit spin density, the probabilities of an s-orbital electron being at the nucleus for ^{14}N (2s2p orbitals) and 69 Ga (4s4p orbitals) are 5.60 a.u.⁻³ and 10.18 a.u.^{-3} respectively [35]. Since the electron wavefunctions for ⁶⁹Ga and ⁷¹Ga atoms are the same, the ratio of the amplitudes of the s-orbital electron wavefunctions at ⁷¹Ga to those at ¹⁴N nuclear sites, $\langle |u_k(0)|^2 \rangle_{Ef}$, for unit spin density may be expected to approximate 1.82 (= 10.18/5.60). If the density of states $\rho_0(E_F)$ for the *atomic* s-orbitals of N and Ga were equal for all of the atoms whose nuclei experience varying Knight shifts across the distribution, the ratio of the widths of the ⁷¹Ga and ¹⁴N Knight shift distributions would be 1.82. Since the actual scaling factor to match the ⁷¹Ga spectrum with the ¹⁴N spectrum is 1.059, we conclude that the conduction band electrons at N atoms in Ge-doped GaN have a factor of 1.7 (= 1.82/1.059) more atomic sorbital character than those at the Ga atoms at each site in the distribution.

This detailed information about electronic structure at the two types of atoms in GaN doped n-type should be able to provide an experimental check of the accuracy of theoretical calculations of GaN electronic structure. Although there have been a number of such calculations, what is required is a projected partial Density of States (PDOS) of the wavefunctions at the conduction band edge onto the s-orbitals of the Ga and N atoms. One such local density-functional calculation on undoped h-GaN using the LMTO method in conjunction with the LDA approach has been carried out, and the results have been compared with experimental results from synchrotron X-ray emission and absorption spectra that yield information on the PDOS [36]. From the graphical presentation of the s-states of Ga and N at the conduction band edge (or 400 meV higher) given in Figure 1 of this reference, it appears that the N atoms have a ca. 1.0-1.1x greater PDOS than the Ga atoms. This factor is not exceedingly far from the 1.7x factor deduced from the Knight shift results. Exact quantitative agreement is not to be expected, since the symmetry-determined projections onto s-states forming the basis of the DFT results, while approximating 4s (for Ga) and 2s (for N) orbitals, are likely not identical to the atomic s-orbitals used to interpret the Knight shift results from hyperfine calculations. The absolute magnitude of the two PDOS is also eliminated as a concern, since it is the relative magnitudes at Ga and N sites that are being compared. We note that double-resonance EPR measurements of Overhauser shifts due to residual donors in h-GaN [37] provide information about the greater relative strength of the hyperfine interactions of an electron to ¹⁴N relative to ⁶⁹Ga or ⁷¹Ga nuclei (although the ratio given based on the Overhauser shifts can be questioned because of the neglect of the different magnetogyric ratios for the two nuclei). The hyperfine interactions reflect a spatial average over a donor hydrogenic 1s orbital, and thus cannot be directly compared to our results involving delocalized (metallic) conduction band electrons.

3. Korringa T_1 relaxation of ⁷¹Ga and ¹⁴N

Associated with Knight shifts is the Korringa spinlattice relaxation time $T_{1,K}$, which for the Fermi gas of non-interacting electrons assumed in Eq.(2) is given by [17]:

$$\frac{1}{T_{1,K}} = \left(\frac{\gamma_n}{\gamma_e}\right)^2 \frac{4\pi k_B T}{\hbar} K^2 \tag{3}$$

where k_B is Boltzmann's constant and γ_n is the nuclear gyromagnetic ratio. Figure 3 summarizes ⁷¹Ga $T_{1,K}$ measurements on GaN:Ge over the temperature range 123 K to 473 K. The inset to Fig. 3(a) depicts the two rate constants W_1 and W_2 characterizing the singleand double-quantum quadrupolar relaxation processes, respectively, and the single rate constant W characterizing the magnetic (Korringa) relaxation process. Since



FIG. 3. (Color online) Plots of ⁷¹Ga MAS-NMR spin-lattice relaxation times due to Korringa relaxation, $T_{1,K}$, for different temperatures and Knight shifts K in GaN:Ge. (a) $T_{1,K}K^2$ versus inverse temperature for three representative Knightshifted signals indicated by dashed lines in Fig. 1(a). The dashed line is the theoretical prediction from Eq.(3). The inset shows transition rates between the ⁷¹Ga I=3/2 Zeeman sublevels due to quadrupolar (W_1 and W_2) and magnetic (W) relaxation processes. (b) $T_{1,K}^{-1/2}$ plotted as a function of the ⁷¹Ga Knight shift K at three temperatures; dashed lines are theoretical predictions from Eq.(3).

 $W_1 \cong W_2$ in GaN [38], the single-exponential time constant T_1 obtained from fitting the saturation-recovery intensities (with complete saturation of all three transitions, a condition that was shown to be necessary to obtain physically-meaningful ⁷¹Ga relaxation results [26]) at each frequency in the spectrum obeys the equation [38] $1/T_1 = 2W_1 + 2W$. Thus, subtracting the quadrupolar relaxation term, measured at the zero-Knight shift peak, from the measured $1/T_1$ at any given frequency position in the spectrum yields $1/T_{1,K}(=2W)$, the Korringa relaxation rate [18] (making the reasonable assumption that the quadrupolar relaxation term is constant across the Knight-shifted spectrum) [39]. By Eq.(3), this rate is proportional to K^2 , and $T_{1,K}$ is expected to depend linearly upon 1/T with a zero intercept for all K.

Using a reduced variable for the ordinate, $T_{1,K}K^2$, we



FIG. 4. Plot of ¹⁴N $T_{1,K}^{-1/2}$ values as a function of the ¹⁴N Knight shift K for bulk GaN:Ge. The dashed line is the theoretical prediction from Eq.(3). Spin-lattice relaxation times were measured at 295 K, 9 kHz MAS, and 11.7 T.

compare in Fig. 3(a) the relaxation data at six different temperatures from 123 to 473 K for three representative Knight shifts to a single theoretical dashed line obtained from Eq.(3). The linearity, slope, and zero intercept of the data for the various K values agree very well with the theoretical prediction, which involves no adjustable parameters. Temperature-dependent Korringa relaxation in n-type GaN has been observed, under conditions where a Knight shift measurement was not possible [40]. Figure 3(b) shows the square root of these relaxation rates plotted against K for three representative temperatures, along with theoretical predictions based upon Eq.(3) (shown as dashed lines, and again involving no adjustable parameters). The deviations shown in the inset for small Knight shifts (< 60 ppm) at low temperature (173 K) may represent the effects of freezing out and localization of donor electrons onto Ge sites, leading to more rapid relaxation at lower temperatures than predicted by the Korringa relation.

The ¹⁴N Knight shifted signals in bulk GaN:Ge also exhibit Korringa relaxation. This is demonstrated by the Korringa relaxation plot for ¹⁴N, shown in Figure 4. The ¹⁴N $T_{1,K}$ values were obtained by least-squares singleexponential fitting of saturation-recovery intensities at equally-spaced intervals across the centerband region of the spectrum. The values for $T_{1,K}$ are obtained using Eq.(3) by setting $T_{1,0}$ to a value based on measurements of the peak at K = 0 (at -302 ppm). The agreement with the theoretical prediction for Korringa relaxation is reasonable. The systematic deviation from the theoretical prediction in the K range of 50 to 150 ppm, where fits are most reliable, is attributed to difficulties in properly removing the baseline contribution due to the overlap between the spinning sidebands and the centerband in the series of saturation-recovery ¹⁴N NMR spectra used to obtain the reported $T_{1,K}$ values.



FIG. 5. (a) X-ray diffraction pattern of nanocrystalline GaN. The black lines indicate reflections indexable to wurtzite GaN (JCPDS Card No. 76-073); the asterisks indicate reflections indexable to β -Ga₂O₃ (< 3%, JCPDS Card No. 43-1012). (b) Rotor-synchronized ⁷¹Ga MAS-NMR Hahn-echo spectrum (18 kHz MAS, 295 K, 18.8 T, 3 s delay) of nanocrystalline GaN; the inset shows a representative TEM image. (c) Plot of ⁷¹Ga $T_{1,K}^{-1/2}$ values as a function of the ⁷¹Ga Knight shift K; the dashed line is the theoretical prediction from Eq.(3).

B. Nanocrystalline GaN

1. ⁷¹Ga Knight shifts and Korringa T_1 relaxation

The ⁷¹Ga spectrum of nanocrystalline GaN in Fig. 5(b) exhibits similar evidence for distributions of local metallic environments. The sample (50-100 nm crystallite diameter by Scherrer analysis), produced by ammonolysis of GaO(OH) at 1100 °C with NH₃, is crystalline h-GaN, as evidenced by the XRD pattern in Fig. 5(a). Similar ⁷¹Ga spectral features as for bulk GaN:Ge are observed: a broad signal around 417 ppm and a narrow peak (11 ppm FWHM) at the shift position of undoped bulk h-GaN at 331 ppm [27]. The spin-lattice relaxation times obtained (using single-exponential fits) and analyzed as discussed above yield a plot of $(T_{1,K})^{-1/2}$ versus K, Fig. 5(c), in excellent agreement with the the-



FIG. 6. (Color online) (a) Two-dimensional ${}^{15}N{}^{71}Ga{}$ dipolar-mediated TEDOR correlation spectrum (20 kHz MAS, 298 K, 18.8 T) of ${}^{15}N$ -enriched GaN nanoparticles. A CPMG pulse sequence was used for detection to maximize the ${}^{15}N$ signal intensity [39], which yields a manifold of narrow peaks (spikelets) whose envelope reflects the ${}^{15}N$ shift distribution. A 1D ${}^{71}Ga$ echo spectrum is shown along the vertical axis for comparison. (b) Slices extracted from the 2D spectrum at positions indicated by the Roman numerals.

oretical line from Eq.(3). Similar distributions of ⁷¹Ga Knight shifts, but affecting a smaller proportion of Ga atoms, are observed for smaller-size GaN nanocrystals (8-11 nm) [22, 23]. The ⁷¹Ga NMR relaxation analyses establish conclusively that the nanocrystalline GaN is heavily doped above the MIT, and demonstrate that synthesis of nanocrystalline GaN by ammonolysis is an effective way to introduce electrically-active dopants into a semiconductor nanomaterial, which can otherwise be difficult [41]. Such NMR techniques provide a promising tool for understanding heavily doped semiconductor nanomaterials, which have interesting plasmonic properties for optoelectronic applications [42] but for which bulk electrical characterization methods are often infeasible due to the difficulties associated with making electrical contacts.

2. Two-dimensional NMR correlation of ^{71}Ga and ^{15}N Knight shifts

The ⁷¹Ga and ¹⁵N Knight shifts in nanocrystalline GaN are spatially correlated over atomic length scales. This is established by the 2D $^{15}N{^{71}Ga}$ transferredecho double resonance (TEDOR) spectrum in Fig. 6 of a ¹⁵N-labelled nanocrystalline GaN (Ga¹⁵N) sample [39]. The TEDOR pulse sequence is useful for transferring polarization from one type of nucleus to another type via relatively weak dipolar couplings [43, 44]. The two-dimensional version, which has been developed and applied to polarization transfer from quadrupolar nuclei to spin-1/2 nuclei [45–49], yields a 2D spectrum showing correlated signal intensity from spatially-proximate nuclear spin pairs that are dipole-dipole coupled. In our case the result is a 2D contour plot with 71 Ga and 15 N shift axes, and a 1D projection spectrum for ¹⁵N. The relative intensity at each ¹⁵N shift position reflects the number of ¹⁵N nuclei at a given ¹⁵N shift that are dipolecoupled (through space, <1 nm) to ⁷¹Ga nuclei having the range of ⁷¹Ga shifts depicted in the 2D contour plot. Perfectly correlated ⁷¹Ga and ¹⁵N shifts would yield a contour plot consisting of a narrow diagonal ridge, which resembles what is experimentally observed in Fig. 6(a).

The slope of the best-fit diagonal in Fig. 6(a) is 1.052 ± 0.007 (⁷¹Ga/¹⁵N), the same (within experimental uncertainty) as the scaling factor of 1.059 (⁷¹Ga/¹⁴N) determined above for bulk GaN:Ge, Fig. 1(c). This equivalence for the two different approaches indicates that the relative *s*-orbital characters of the conduction band electrons at the Ga and N sites are very similar in bulk Gedoped and nanocrystalline GaN. Representative vertical slices extracted from the TEDOR spectrum, Fig. 6(b), have widths that are considerably smaller than the overall 1D ⁷¹Ga spectrum, again indicating that the respective ⁷¹Ga and ¹⁵N Knight shifts from neighboring atoms in the nanocrystals are correlated.

Figure 7 compares the shift distributions of 71 Ga and 15 N for the nanocrystalline Ga 15 N sample, with the ppm axis for the latter scaled by the factor of 1.052 determined from fitting the TEDOR spectrum ridge, as described in the caption and text. As was the case with the analogous 71 Ga/ 14 N comparison for GaN:Ge shown in Fig. 1(c), the distributions are comparable in shape, although there is a slightly greater relative intensity for 15 N on the right side of the peak maxima.

It is noticeable in Fig. 6 that there is an increasing spread in the frequency range of the ⁷¹Ga nuclei in crosspeaks associated with a given ¹⁵N frequency as one moves to larger Knight shifts, a spread which reaches its maximum of about 60 ppm (⁷¹Ga) width in the asymmetric high-frequency tail of the distribution. A priori this spread could arise from three possible sources involving either chemical shifts, NQCCs, or Knight shifts. We rule out differential chemical shift changes of the two types of nuclei (e.g. from defects or surface effects) as the explanation, since Fig. 7 shows a good correlation be-



FIG. 7. (Color online) Comparison of the Knight shift distributions of 71 Ga and 15 N for nanocrystalline Ga 15 N, with normalized peak heights and with an x-axis corresponding to the 71 Ga Knight shift and the 15 N Knight shift scaled by a factor of 1.052, the same as the slope of the dashed line in the TEDOR spectrum (Fig. 6(a)). The 71 Ga spectrum (red) was acquired at 298 K, 18.8 T, 20 kHz MAS, and with a recycle delay of 10 s. The 15 N spectrum (blue) was acquired at 298 K, 7.4 T, and with a recycle delay of 240 s.

tween the ⁷¹Ga and ¹⁵N shifts (especially at larger shifts), as expected for a distribution arising solely from Knight shifts. We also rule out, on two grounds, NQCC effects due to Ga atoms in the direct vicinity of defects such as O_N substitutions (or N vacancies) having strongly asymmetric environments resulting in much larger ⁷¹Ga quadrupolar interactions and associated spectral broadening (so-called second-order quadrupolar broadening). Firstly, previous ⁷¹Ga MAS NMR results on similar GaN nanoparticle materials found little or no evidence for anisotropic second-order quadrupolar broadening effects across the Knight shift distribution [23]. Secondly, if such effects were responsible, then one would not expect Korringa-type relaxation behavior across the entire distribution of ⁷¹Ga Knight shifts, as observed in Fig. 5(c).

We are led to conclude that each ¹⁵N Knight shift isochromat is correlated with a distribution of ⁷¹Ga Knight shifts whose width increases for the larger ¹⁵N Knight shifts. To understand how this might happen, consider the inset to Fig. 6(a), which depicts a particular ¹⁵N nucleus and the associated distances and calculated dipolar couplings to both a directly- bonded ⁷¹Ga nucleus as well as to next-nearest-neighbor ⁷¹Ga nuclei. There are 4 closest sites for $^{71}\mathrm{Ga}$ nuclei, and 24 sites for ⁷¹Ga nuclei separated by a gallium and a nitrogen atom (having dipolar couplings to 15 N that are 19% or 8% of the directly-bonded value). Thus, both directly-bonded ⁷¹Ga nuclei as well as next-nearest-neighbor ⁷¹Ga nuclei will contribute to the observed signal intensity in the dipolar-coupling mediated ¹⁵N{⁷¹Ga} TEDOR spectrum shown in Figure 6(a).

The increasing spread in the frequency range of the 71 Ga nuclei in cross peaks associated with the most Knight-shifted 15 N nuclei can then be explained if those 15 N nuclei, such as depicted in the inset to Fig. 6(a),

are close to the shallow donors, and thus experience the largest Knight shifts due to peaking of the wavefunction near donor sites [50]. Their 4 closest sites for ⁷¹Ga nuclei, and their 24 sites for next-nearest-neighbor ⁷¹Ga nuclei would then have significantly different nanoscale proximities to the same shallow donor, and hence different ⁷¹Ga Knight shifts.

This interpretation in terms of varying local metallic properties is consistent with the larger ¹⁴N Knight shifts associated with N atoms closer to (Ge) shallow donors in GaN:Ge (Fig 1(b)). A rough estimate of the magnitude of this spatial variation of ⁷¹Ga Knight shifts can be made by assuming that the 60 ppm 71 Ga linewidth described above is due to a 60 ppm difference in Knight shifts between two ⁷¹Ga nuclei attached to the same N atom with a 3.20 Å separation (from the crystal structure of hexagonal GaN), and with the internuclear vector aligned with the dopant atom for maximal effect. This yields a Knight shift spatial variation in the regions nearest the dopant of ca. 20 ppm/A. The sensitivity of 2D NMR techniques to nanoscale variations of electronic structure thus provides a basis for spatially correlating the electronic properties of two atom types in compound semiconductors (bulk or nanocrystalline) doped above the MIT for comparison with theoretical calculations of such properties. The attribution of at least some part of the observed distribution of Knight shifts to nanoscale variations in electronic structure, based upon these experimental results, has significant implications. It indicates that interpreting observed Knight shift distributions in terms of macroscopic (rather than nanoscopic) probability distributions of carrier densities is not justifiable in the way originally proposed [14, 18].

3. Likely shallow donors in nanocrystalline GaN

Oxygen substitution on an N site (O_N) is likely to be present in the nanocrystalline GaN materials, which were prepared from an oxygen-containing precursor, GaO(OH). The XRD pattern (Fig. 5(a)) of the nanocrystalline natural abundance GaN shows weak reflections that are indexable to β -Ga₂O₃ (< 3%), consistent with oxygen being present in the sample. This impurity is not visible in the 71 Ga NMR spectrum in Fig. 5(b) due to its dilute quantity and inhomogeneously broadened signals. Oxygen doping in GaN is usually associated with Ga vacancies (V_{Ga}) , which act as triple acceptors and partially compensate the O_N shallow donors. Typically, \mathbf{V}_{Ga} and \mathbf{O}_N are assumed to form a complex which acts as a deep level, and only the uncompensated O_N will act as shallow donors [51]. Nitrogen vacancies could also be present, given that the synthesis temperature is high enough to dissociate GaN under vacuum [52]. However, nitrogen vacancies are reported to be energetically unfavorable under n-type conditions, with a formation energy that depends greatly on Fermi level [51].

The most energetically favorable shallow donors in n-

type material are O_N , Si_{Ga} , and Ge_{Ga} [51], of which O_N is the most physically reasonable given the presence of oxygen in the starting material. The GaN wurtzite lattice can accommodate large concentrations of O atoms while maintaining tetrahedral coordination of Ga atoms [53]. However, Si_{Ga} cannot be totally ruled out since the ammonolysis was done in alumina crucibles, and typical alumina ceramics use silica-based sintering aids in their manufacture. A recent modeling study based on hybrid functional calculations [54] predicted similar shallowdonor electronic behaviors for O_N and Ge_{Ga} substitution defects in GaN, which is fully consistent with our analyses of the similar NMR spectral features and corresponding electronic structures observed here for nanocrystalline GaN and bulk GaN:Ge. These results are also consistent with previous results [23] for nanocrystalline GaN synthesized differently, which exhibited a broad distribution of Knight-shifted intensities [39].

III. CONCLUSIONS

In summary, ⁷¹Ga, ¹⁴N, and ¹⁵N MAS-NMR experiments have provided detailed new insights on the similar local electronic structures of nanocrystalline and doped bulk GaN. They reveal the highly metallic character of both bulk GaN heavily doped n-type with Ge and nanocrystalline GaN made by high temperature ammonolysis, and conclusively demonstrate the origin of the shifts observed in the latter material to be conduction band electrons. Such high carrier concentrations in nanocrystalline GaN are most likely due to oxygen atoms from the GaO(OH) precursor, which are known to give rise to n-type GaN in bulk [51, 53].

Measurements of Korringa spin-lattice relaxation times for differing Knight shifts across the spectra prove that both forms of GaN are well above their MIT, with carriers exhibiting the near-ideal characteristics of a degenerate Fermi gas of non-interacting spins. This close approach to ideal behavior, which contrasts with deviations from ideal Korringa behavior often observed for true metals [55], can be attributed to the orders of magnitude lower carrier concentration in the degenerately-doped semiconductor.

Although the magnitude and character of the conduction electron wavefunction at the Group III and Group V nuclear sites have been measured in other bulk metallic semiconductors [56], this is the first determination of the Group V/III ratio of s-orbital carrier wavefunction probabilities at the Fermi edge across the entire distribution of Knight shifts in both bulk doped and nanocrystalline materials, with the N to Ga ratios equal to ca. 1.7 for both forms.

Two-dimensional ${}^{15}N{}^{71}Ga$ NMR results establish that the local metallic properties (Knight shifts) at Ga and N sites are spatially correlated on a sub-nanometer scale. They provide additional evidence that distributions of Knight shifts are due to electronic disorder arising from differing proximities to randomly located shallow donors. This conclusion has been reached more indirectly by a theoretical model accounting for ²⁹Si Knight shift distributions in doped silicon [9]. Such detailed insights on atomic-level electronic structures are expected to be relevant generally to Group III-V and other technologically important semiconducting solids. The present approach to observing spatial correlations in electronic structure between different atom types should thus prove valuable in future investigations of nanocrystalline semiconductors, where the effects of interfaces and randomlysubstituted dopants on the MIT are not well understood, and in comparisons of their properties to those in doped bulk analogs.

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