Characterization of optical and spin properties of single tin-vacancy centers in diamond nanopillars

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DOI: 10.1103/PhysRevB.99.205417

I. INTRODUCTION

Color centers in diamond have attracted much interest as candidates for optically active, solid-state quantum bits. Of particular interest are inversion-symmetric color centers based on group-IV impurities in diamond because they emit strongly into their zero-phonon lines and are insensitive to electric field noise to first order. Early studies of the negatively charged tin-vacancy (SnV\(^{-}\)) center in diamond have found the SnV\(^{-}\) to be a promising candidate: it has high quantum efficiency, emits strongly into its zero-phonon lines, and is expected to have a long \(T_2\) spin coherence time at 4 K. To develop the SnV\(^{-}\) into a spin qubit requires further characterization, especially of the spin and optical properties of individual SnV\(^{-}\) in nanofabricated structures. In this work, we isolate single SnV\(^{-}\) centers in diamond nanopillars and characterize their emission properties and their spin response to a magnetic field. We observe narrow emission linewidths <250 MHz, as well as a strong polarization dependence of each transition. We also find the Zeeman splitting under a magnetic field to be in good agreement with theoretical prediction. Our results pave the way toward future employment of single SnV\(^{-}\) centers as optically accessible quantum memories.

II. ISOLATION OF SINGLE TIN-VACANCY CENTERS

Starting with electronic-grade, single-crystal diamond from Element Six, we perform a boiling tri-acid (1:1:1 nitric:sulfuric:perchloric acids) clean. We then etch 300 nm with oxygen (O\(_2\)) plasma etch. Using the patterned Si\(_x\)N\(_y\) as a spin qubit in a quantum network.

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PHYSICAL REVIEW B 99, 205417 (2019)
an etch mask, we etch 500 nm into the diamond with a directional O2 plasma etch. A scanning electron microscope (SEM) image of a resulting nanopillar can be found in Fig. 2(a).

We verify that we have isolated single emitters in the diamond nanopillars by performing second-order autocorrelation ($g^{(2)}[\tau]$) measurements. A measurement of $g^{(2)}[\tau]$ for an emitter in a nanopillar under 1 mW, 532 nm excitation is presented in Fig. 2(b). The data are fit to the function

$$g^{(2)}[\tau] = 1 - c[(1 + b)e^{-|\tau|/\tau_1} - be^{-|\tau|/\tau_2}],$$

(1)

where $b$, $c$, $\tau_1$, and $\tau_2$ are fitting parameters [12]. $\tau_1$ provides an estimate of the excited-state lifetime of the emitter. $\tau_2$ is the decay time on the photon bunching and relates to a third shelving state in the system. This third shelving state has been discussed for SnV− by Iwasaki et al. [7] and explored theoretically and experimentally for SiV− [8,13]. For the emitter presented in Fig. 2(b), $g^{(2)}[0] = 0.23 \pm 0.02$, indicating that this emitter is indeed a single emitter. $\tau_1$ and $\tau_2$ were found to be 4.8 ± 0.1 and 103 ± 10 ns, respectively. The estimated emitter lifetime $\tau_1$ of 4.8 ns is on par with previous measurements of ~5 ns [7].

Based on a survey of 200 pillars designed for a diameter of 200 nm, we estimate that 2% of such pillars contain low-strain single emitters. This estimate is based on the number of pillars with ground-state splitting $\sim$850 GHz and $g^{(2)}[0] < 0.5$. About 4% of 200-nm pillars contained single SnV−, strained or unstrained. Additionally, by counting the number of SnV− in 200-nm-diameter pillars and comparing the areal density to the implantation dose, we estimate the conversion efficiency to be 0.7%. This conversion efficiency may be improved by varying the implantation dose and annealing time.

For all photoluminescence (PL) studies, we use 532 nm laser light to excite the emitter above resonance. We measure our data with two home-built confocal microscopes with free-space paths into cryostats: a Montana Instruments Cryostation and an attoDRY2100. Further details on our setups can be found in the Supplemental Material [14].

III. OPTICAL PROPERTIES

PL spectra of a single SnV− in a nanopillar at 5 K are presented in Figs. 3(a) and 3(b). We observe strong emission into the C and D transitions, which present as two closely spaced, sharp lines around 620 nm in Fig. 3(a). As expected at this temperature, only the C and D transitions are visible [7]. The phonon sideband is also visible in the background-subtracted plot of Fig. 3(a). The background PL spectrum, which can be found in the Supplemental Material [14], was acquired with the same excitation power and integration time on another pillar of the same size that did not contain an emitter. Figure 3(b) is a higher-resolution PL spectrum of the C and D transitions only. From this spectrum, the ground-state splitting of the emitter is found to be 842 GHz, close to the ~850 GHz expected, indicating that this emitter is under minimal strain.

To obtain the linewidth of the emitters, we perform photoluminescence excitation (PLE) experiments. Here, we tune a laser across the ZPL and collect the photons emitted into a confocal microscope. We measure the C transition via PLE, shown in Fig. 3(c), and found the linewidth to be 231.9 ± 9.5 MHz at 1.7 K. The emitter measured for this experiment had a $\tau_1 = 3.8$ ns based on $g^{(2)}[\tau]$ data, which corresponds to a lifetime-limited linewidth of approximately 42 MHz. The discrepancy between the lifetime-limited and measured linewidths may be caused by a significant spectral diffusion that we observed during the measurements. The spectral diffusion may be mitigated by implanting with a lower dose.

We also study the polarization of the emission into the C and D transitions for a single SnV−. To do so, we use the setup illustrated in Fig. S1 of the Supplemental Material [14] and measure the spectrum with different half-wave-plate angles. The C and D transitions are then fit to Lorentzians, the areas under which are taken to be the emission intensity. The emission intensity of the C and D transitions is plotted as a function of the half-wave-plate angle in Fig. 4(a). The SnV− studied displays a clear polarization dependence in its emission. The strong polarization dependence is, as expected, similar to that of SiV− in diamond [15] and is consistent with previously reported room-temperature absorption data for SnV− [16]. From Fig. 4(a), we observe that the dipole
moments associated with the C and D transitions projected onto the (001) diamond surface are perpendicular to one another.

We estimate the spatial inhomogeneous broadening present in the emitters by plotting the average PL spectra of 44 pillars, as shown in Fig. 4(b). The C and D transitions appear in the plot as two distinct peaks and have been fit to Gaussian distributions. From the fits, we find the full widths at half maxima of the C and D transitions to be 271 ± 8 and 583 ± 27 GHz, respectively. Raman tuning has been used to tune the ZPL of a SiV− up to 100 GHz [17,18]. Thus, it may be possible in future works to overcome a significant portion of the inhomogeneous broadening found here.

IV. ZEEMAN SPLITTING

We explore the spin properties of a single SnV− and compare our experimental results to theoretical predictions. When a magnetic field is applied to a SnV− center, the twofold spin degeneracy is lifted and the two branches of the excited- and ground-state manifolds each split into two more states. Consequently, each of the original optical transitions splits into four: two spin-conserving transitions that are strongly allowed and two spin-nonconserving transitions that are only weakly allowed by magnetic field components orthogonal to the symmetry axis of the emitter. A schematic diagram of the energy levels and optical transitions is shown in Fig. 5(a).

For our theoretical model, we solve for the eigenstates of the Hamiltonian derived in Ref. [9]:

\[
\hat{H}_{\text{eff}}^{\sigma u} = -\hbar \lambda^{\sigma u} \hat{L}_z \hat{S}_z + \mu_B \gamma^{\sigma u} \hat{L}_z B_z + \mu_B g_\lambda \hat{S} \cdot \hat{B} + 2\mu_B \delta_\lambda^{\sigma u} \hat{S}_z B_z + \hat{\chi}_{\text{strain}},
\]

where the z direction is along the symmetry axis of the emitter. The superscripts g and u respectively denote the parity of the states: even (gerade) and odd (ungerade). For SnV− and similar negatively charged group-IV-based color centers, the ground state corresponds to g and the excited state corresponds to u. The sign on the effective spin-orbit-coupling (first) term opposes that of the Hamiltonians previously used to model other negatively charged group-IV-based color centers [9,15]. This Hamiltonian can be used to find the energy levels of a SnV− as a function of applied dc magnetic field B for the arbitrary magnetic field direction. The third term, which contains \( \hat{S} \cdot \hat{B} \), describes the Zeeman effect. In the form \( x = \{x^e, x^u\} \), where x is a parameter, the values given in Ref. [9] for the parameters in Eq. (2) are \( \lambda = \{850, 3000\} \) GHz, \( f = \{0.154, 0.098\} \), and \( \delta_f = \{0.014, 0.238\} \). We study an emitter under minimal strain, so we neglect the \( \hat{\chi}_{\text{strain}} \) term.

For the measurements in a magnetic field, our sample is cooled to 1.7 K in an attoDRY2100 that is equipped with a superconducting magnet. The magnetic field is applied out of the plane of the chip, along the (001) direction, which shall be denoted as \( z^\prime \). We apply magnetic fields \( B_z \) from 0 to 9 T, inclusive, and perform PL spectroscopy. To collect spectra of the inner transitions with finer resolution, we use a double monochromator. The resulting double-monochromator data as a function of field are presented in Figs. 5(b) and 5(d). A detailed explanation of how the data in Figs. 5(e) and 5(e) were extracted from the raw spectra can be found in the Supplemental Material [14].

We plot the experimental and theoretical frequencies of the four peaks comprising C and D in Figs. 5(c) and 5(e). To correct for a slow spectral drift, we center the inner and outer frequencies pairwise about their average frequency. We apply the same correction to the theoretical plots. As can be seen in Figs. 5(c) and 5(e), the experimental data are in good agreement with the predicted behavior based on Eq. (2).
We note, however, that two of the values used to generate the solid curves of Figs. 5(c) and 5(e) are dependent on $g_{L}^{u}u$ values for SiV$^-$; specifically, $f_{g}^{u}u = \delta_{g}^{u}u g_{L}^{u}u$ and $\delta_{f}^{u}u = \delta_{g}^{u}u \delta_{L}^{u}u$, where only the $\delta_{g}^{u}u$ and $\delta_{f}^{u}u$ values were calculated for SnV$^-$ by the authors of Ref. [9], while $g_{L}^{u}u$ values were found for SiV$^-$. Thus, we have tried scaling the $g_{L}^{u}u$ values by a constant $\alpha^{u}u$ to find the least-squares fit of the theory to the data. We have found the best-fit values of $\alpha^{u}u$ to be $\alpha = [0.98, 1.32]$. The theoretical curves that include this scaling are shown as dashed curves in Figs. 5(c) and 5(e) and yield an improved agreement between the theory and the data—the sum of the squared residuals is reduced by a factor of 2 compared to the original case. Thus, while the original model is in good agreement with our data, it can be further improved by changing the $g_{L}^{u}u$ values used.

V. SUMMARY AND OUTLOOK

We have studied the optical properties of single SnV$^-$ centers that have been isolated in diamond nanopillars. We were able to generate emitters with narrow linewidths of $\sim 232$ MHz. These linewidths are an improvement over previously measured linewidths in bulk diamond and demonstrate that emitters can be incorporated into nanostructures without degrading the optical properties. SnV$^-$ linewidths may be further improved with lower implantation dose and a longer high-temperature anneal. We also observed a strong polarization dependence of the emission, with the $C$ and $D$ emission being orthogonally polarized. Lastly, we observed the behavior of a single SnV$^-$ as a function of magnetic field and found it to be in good agreement with theoretical predictions presented by Thiering and Gali [9]. Our findings indicate that the SnV$^-$ is a promising candidate for a spin qubit and warrants further investigation. Future works include characterizing the spin-coherence time $T_{2}$ [5, 19, 20] of the isolated emitters and fabricating more sophisticated nanostructures [10] such as photonic cavities resonant with SnV$^-$ ZPLs, two necessary steps in order to implement the SnV$^-$ as an optically interfaced spin qubit.

Note added. Recently, we became aware of a very recent, similar work [21].

ACKNOWLEDGMENTS

We acknowledge Gergő Thiering and Ádám Gali for fruitful discussions about theory. We also acknowledge Linda Zhang and Greg Pittner for helpful advice on fabrication. This work is financially supported by the Army Research Office (ARO) (Award No. W911NF-13-1-0309), the National Science Foundation (NSF) RAISE TAQS (Award No. 1838976), and the Air Force Office of Scientific Research (AFOSR) DURIP (Award No. FA9550-16-1-0223). A.E.R. acknowledges financial support from the National Defense Science and Engineering Graduate (NDSEG) Fellowship Program, sponsored by the Air Force Research Laboratory (AFRL), the Office of Naval Research (ONR), and the Army Research Office (ARO). C.D. acknowledges financial support from the National Science Foundation under Award No. ECCS-1542152.

FIG. 5. Behavior of SnV$^-$ in an external magnetic field $B_{z}$. (a) Schematic energy-level diagram of SnV$^-$ when a magnetic field is applied. Each of the original optical transitions splits into four transitions. Two of each set of four transitions are spin conserving and thus strongly allowed (bold arrows), and the other two are weakly allowed (thin arrows). Transitions $C$ (red) and $D$ (dark red) are represented by arrows, numbered 0 to 3 in order of decreasing energy. (b) Stacked spectra of inner transitions $C_{1}$ and $C_{2}$ as a function of $B_{z}$ collected with a double monochromator. Transition frequency is with respect to the central transition frequency at 0 T. Solid red lines are double or single Lorentzian fits to the data (black points). (c) Location of central frequencies of all four, i.e., $C_{0}$, $C_{1}$, $C_{2}$, and $C_{3}$, transitions. Error bars on the data points are the estimated error based on the variance of the central frequency fit parameters estimated by curve_fit in PYTHON. Solid curves are functions derived theoretically from Eq. (2) and correspond to data of the same color. Dashed curves are functions derived from Eq. (2) when $f_{g}^{u}u$ and $\delta_{f}^{u}u$ values were calculated by the optimal $\alpha^{u}u$. (d) Same as (b) for transition $D$. (e) Same as (c) for $D$ transitions.


