Conversion of neutral nitrogen-vacancy centers to negatively charged nitrogen-vacancy centers through selective oxidation

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The conversion of neutral nitrogen-vacancy centers to negatively charged nitrogen-vacancy centers is demonstrated for centers created by ion implantation and annealing in high-purity diamond. Conversion occurs with surface exposure to an oxygen atmosphere at 465 °C. The spectral properties of the charge-converted centers are investigated. Charge state control of nitrogen-vacancy centers close to the diamond surface is an important step toward the integration of these centers into devices for quantum information and magnetic sensing applications. © 2010 American Institute of Physics. [doi:10.1063/1.3364135]

The combination of a long ground-state electron coherence time and the ability to perform optical spin readout in the negatively charged nitrogen-vacancy center (NV−) in diamond has motivated proposals to use this center for both quantum information processing (QIP) (Refs. 3 and 4) and magnetic sensing applications. The best optical and spin properties have been observed in NV− centers deep within the diamond lattice, however for both applications it will be highly desirable to use NV centers very close to the diamond surface. For QIP applications, surface NV centers can be coupled to on-chip waveguides and microcavities. For magnetometry applications, NV centers embedded in either diamond nanoparticles or two-dimensional sensing surfaces will combine high magnetic sensitivity with high spatial resolution.

One common technique to create NV centers near a surface is through ion implantation and high temperature annealing. Since the best spin and optical properties have been observed in high-purity diamond, it would seem advantageous to begin with this material and implant with nitrogen ions. Upon annealing, the vacancies created during implantation migrate to form NV centers with the implanted nitrogen. Recent reports, however, indicate that in high purity diamond the preferred charge state of NV centers created very close to the surface (within 200 nm) is the neutral (NV0) charge state. To date, the attractive optical and spin properties observed in the NV− center have not been observed in its neutral counterpart. Thus the first step toward engineering near-surface NVs for QIP and magnetometry is to demonstrate charge state conversion.

A study of the vertical distribution of NV centers created by ion implantation and annealing indicated that the neutral charge state observed near the diamond surface was due to electronic depletion effects consistent with an acceptor layer at the diamond surface. In this depletion region, which can extend several microns in high purity diamond (<<5 ppb nitrogen), nitrogen donors are ionized and thus cannot donate an electron to the NV center. One possible acceptor candidate is graphitic defects created during the ion implantation. Selective oxidation in air of sp2 bonded carbon (graphite or amorphous) over sp3 bonded carbon (diamond) has previously been demonstrated in detonation-synthesized nanodiamond. Here we show that the same technique can be used to convert NV0 to NV− for centers created 10–75 nm from a single-crystal diamond surface. Additionally we compare the low temperature optical properties of the near-surface NV centers to centers found deep within the diamond matrix.

Three high purity commercial diamond samples grown by chemical vapor deposition (E6, CVD electronic grade) were masked with a TEM grid and implanted with nitrogen (CORE Systems). Implantation conditions are given in Table I. Each sample was next cut into two pieces. All samples were annealed at 900 °C for 1 h in an Ar/H2 forming gas. One of the two pieces was reserved as a control sample. Half of the surface of the second piece was masked with a 125 nm thick SiO2 film deposited by e-beam evaporation. The three half-masked samples were then annealed at 465 °C in an oxygen atmosphere in steps of 15, 15, 60, and 60 min.

Before, between, and after annealing steps, confocal imaging was used to locate the same implantation region for spectral measurements. An example of a confocal image before and after annealing showing the same implantation square is shown in Figs. 1(a) and 1(b). To obtain these images the sample was excited with a 532 nm laser with 1 mW excitation power focused to a spot with diameter less than 1 μm while photoluminescence (PL) from the phonon sidebands (650–800 nm) was collected. After imaging, optical spectra were taken both in the clear and masked regions of each sample for several excitation powers. Representative spectra of the unmasked region both at room temperature and 10 K before and after annealing are shown in Figs. 1(c) and 1(d). The NV0 zero phonon line (ZPL) and phonon sidebands are clearly dominant before the O2 anneal. However after

TABLE I. Implantation conditions for the three N-implanted samples studied. Implantation depths are calculated using SRIM software (Ref. 13).

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Energy (keV)</th>
<th>Dose (cm−2)</th>
<th>Depth (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>10</td>
<td>1 × 1010</td>
<td>14 ± 5</td>
</tr>
<tr>
<td>S2</td>
<td>10</td>
<td>1 × 1011</td>
<td>14 ± 5</td>
</tr>
<tr>
<td>S3</td>
<td>50</td>
<td>1 × 1011</td>
<td>62 ± 14</td>
</tr>
</tbody>
</table>

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150 min of annealing the centers are predominantly NV−. We note that in most diamond samples both charge states are generally observed in the PL spectra with 532 nm excitation. This is due to optically induced charge state conversion and is even observed for single NV centers in high-purity diamond. The effect of the O2 anneal is to change the relative probability for the NV center to be in the NV− state.

The ratio of the room temperature NV− ZPL intensity to the total ZPL intensity as a function of annealing time for all three samples is plotted in Fig. 2(a). In all samples the charge state of the NV center switches from predominantly NV0 to predominantly NV− within the first 90 min of annealing. This charge state conversion is not observed in the masked regions of the sample indicating that the oxygen atmosphere is necessary for charge state conversion. The two 10 keV samples behave similarly, initially exhibiting a very small NV− component which rapidly increases. The 50 keV sample begins with a much larger NV− component and a slower conversion is observed.

The excitation power dependence of the NV− ZPL component at 10 K for the two implantation energies is plotted in Fig. 2(b). Very little dependence on excitation power is observed in the O2-annealed sample as the excitation power is decreased indicating that the measured ratio is stable and the centers would remain predominantly NV− in the dark. In contrast, a strong power dependence is observed in both control samples. In these samples the NV− component is small at low excitation power and increases with power. This "photochromatic" effect has been reported previously and was attributed to electron excitation and capture dynamics between nearby nitrogen donors and NV centers. We note that even at the lowest excitation power used, the NV− component still appears to be decreasing in the 50 keV (S3) sample.

The larger NV− component observed in the 50 keV case compared to 10 keV case could be due to a difference in the optically excited electron dynamics for the two implantation depths.

Oxygen annealing at elevated temperatures results in both surface termination with oxygen-containing functional groups and the selective removal of sp2 carbon from diamond samples. Hydrogen surface termination, which is often observed in CVD-grown samples, results in a p-type surface conductive layer15 which could be the source of an electron depletion layer before the O2 anneal. This H-terminated surface exhibits both significant surface conductivity [10−4–10−5 Ω−1 (Ref. 15)] and a water contact angle θ > 80°. However in both the control and annealed samples, the surface conductivity was less than 10−10 Ω−1. Additionally, the water contact angle before O2 annealing was θ=60°. Both measurements indicate the absence of a purely hydrogen-terminated surface. Upon O2 annealing the water contact angle decreased to less than 30°. This effect has been observed previously in ion-irradiated samples and is attributed to the conversion of irradiation-created sp2 carbon to sp3 carbon. As discussed in Ref. 12, once sp2 bonds are removed, the surface becomes more accessible to oxygen-containing functional groups. The contact angle may additionally be affected by the removal of hydrogen-containing functional groups.

Once charge state conversion has occurred it is possible to study both the spin and optical properties of the near-surface NV− centers. Here we briefly discuss the low temperature optical properties of the O2-annealed NV− centers. For many QIP applications it is desirable for NV centers to emit identical photons. Two factors which contribute to the spectral broadening of the emitted photons in the NV system are spectral diffusion and phonon broadening. High resolution PL excitation (PLE) spectra of NV centers in sample S1 under different conditions are shown in Figs. 3(a)–3(c). In each case a tunable red diode laser is scanned over the NV− ZPL line while PL from the phonon sidebands are detected (650–800 nm). Measurements were performed at 10 K where
phonon broadening of the ZPL is greatly reduced.

In Fig. 3(a), a single NV which formed during the CVD growth process ~60 μm beneath the diamond surface is studied. Before each scan a green repump laser is applied in order to reverse photoionization which eventually occurs with the red laser alone. During the measurement a single NV track blinks on and off. This blinking is likely caused by charge state conversion. Little spectral diffusion (~200 MHz) is observed between scans which is typical in the highest quality CVD samples. In Fig. 3(b), the NV centers formed by implantation and annealing are studied. Single PL tracks are not observed in this measurement and the NV centers randomly blink on and off throughout the entire scan range. If the repump laser is applied approximately every 20 scans, as in Fig. 3(c), it is possible to observe single NV tracks. However once a particular track goes dark and the repump is applied, the new track may be tens of gigahertz away from the previous one. The extreme sensitivity of the NV− transition frequency to the repump laser indicates the presence of charge traps close to the NV center. Future studies, including the effect of surface treatments, will be required to determine if it is possible to reduce the large spectral diffusion observed in these near-surface NV centers.

In summary, we have shown that it is possible to convert the charge state of NV centers created by ion implantation and annealing from NV0 to the desired NV− charge state through an oxygen anneal at 465 °C. The results suggest that the electronic depletion effect observed in high-purity implanted samples is related to graphitic damage which can be removed by selective oxidation. This is an important step toward engineering NV centers near the diamond surface suitable for integration with optical and scanning probe devices for both QIP and magnetometry applications.

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