Charge-state dynamics during excitation and depletion of the nitrogen-vacancy center in diamond

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The charge-state dynamics of the nitrogen-vacancy (NV) center in diamond play a key role in a wide range of applications, yet remain imperfectly understood. Using single picosecond pulses and pulse pairs, we quantitatively investigate the charge dynamics associated with excitation and fluorescence depletion of a single NV center. Our pulsed excitation approach permits significant modeling simplifications and allows us to extract relative rates of excitation, stimulated emission, ionization, and recombination under 531 and 766 nm illumination. By varying the duration between paired pulses, we can also investigate ionization and recombination out of metastable states. Our results are directly applicable to experiments employing stimulated emission-depletion imaging and can be used to predict optimal operating regimes where excitation and stimulated emission are maximized relative to charge-state-switching processes.

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I. INTRODUCTION

The nitrogen-vacancy (NV) center in diamond has become a workhorse for quantum information and metrology [1,2], with applications ranging from quantum networks [3] to geology [4] to fundamental spin-bath physics [5]. Such applications rely on the favorable spin properties of the negatively charged state (NV\(^{-}\)), which exhibits long coherence times and permits optical preparation and detection [1]. At the same time, controlled conversion to the neutral charge state (NV\(^{0}\)) offers opportunities for optical nanoscopy [6,7], charge-based memories [8,9], electrical spin detection [10,11], and improved spin detection via spin-to-charge conversion [12–15]. Understanding and controlling the charge state of the NV thus underlies a broad range of potential technologies, yet the dynamics of optically induced charge-state switching remains an area of active research [16–21].

In this paper, we focus on quantitative measurements of charge-state switching associated with pulsed optical excitation and depletion processes, where a pulse of green light excites the NV and a pulse of red or near-IR light causes stimulated emission. Such two-color pulsed illumination is employed in stimulated emission-depletion (STED) imaging, permitting imaging of NV centers with nanoscale resolution [22–24]. Understanding the associated charge-state dynamics provides further insight into the fluorescence depletion mechanism—stimulated emission or ionization—which also has implications for proposed laser threshold magnetometry [25]. Moreover, if it is possible to induce excitation and stimulated emission of NV\(^{-}\) without ionization, then several opportunities open up. Controlled transfer into and out of the NV\(^{-}\) excited state could enable fast electron-nuclear spin gates [26]. Alternately, if STED does not ionize and preserves the polarization [23] or coherence of surrounding spins, it may become possible to detect the joint state of several closely spaced, interacting spins [27].

Our experiments examine ionization (NV\(^{-}\) → NV\(^{0}\)) and recombination (NV\(^{0}\) → NV\(^{-}\)) induced by sequential pulses of green (531 nm) and red (766 nm) light, at wavelengths commonly used for STED experiments. Essentially, we ask the questions: (1) Can we directly compare the excitation rate of green light to rates of ionization or recombination out of the excited states? (2) How much does red light cause stimulated emission versus ionization or recombination? and (3) How does red-induced ionization out of the NV\(^{-}\) metastable singlet state compare to ionization out of the excited state?

We study single NV centers, where we can prepare and detect the charge state with high fidelity [12,17] and thereby determine ionization and recombination probabilities quantitatively. In contrast to previous work that examined longer-timescale dynamics of green and near-IR illumination [13,21], we examine single ~100 ps pulses or pulse pairs. Working in this single-pulse regime eliminates long-timescale dynamics associated with spin and metastable states, and reveals a simple picture of optically induced processes. We model the system with linear rate equations that we quantitatively fit for excitation, stimulated emission, ionization, and recombination rates. The model can then be used to predict optimal operating regimes and behaviors. We note that our approach of linear rate equations does not contradict the quadratic dependence of ionization on power observed at low intensities in, e.g., Ref. [17]. Because we explicitly include the excited states in our model (the \(3E\) excited state of NV\(^{-}\) and the \(2A_1\) state of NV\(^{0}\) [28]), each process (excitation followed by ionization) can be linear while the overall ionization process still requires two photons.

For example, our results corroborate ensemble measurements indicating that stimulated emission dominates over ionization [29], and additionally provide quantitative values that will make it possible to more accurately model potential NV lasing behavior. We also measure the rate of ionization by 766 nm light out of the NV\(^{-}\) excited and singlet states, which may inform spin-to-charge conversion approaches that utilize near-IR singlet ionization [13]. Perhaps disappointingly, our results indicate that excitation and depletion are inevitably accompanied by ionization, and at best an “ideal” excitation-stimulated emission cycle occurs with 82.5% ± 0.3%
probability. Our model and extracted rates thus provide a framework for understanding and designing a wide range of experiments.

II. EXPERIMENTAL METHODS

Our experiments employ four optical excitation pathways, allowing charge-state preparation, manipulation, and detection for a single NV in bulk diamond. Briefly, continuous-wave (CW) green (532 nm) excitation is used to initialize the distribution of NV charge and spin populations at the beginning of the experiment, while single-shot charge-state preparation and measurement is performed using yellow light (594 nm) [17]. We thereby extract the effects of picosecond pulses of green (531 nm) and red (766 nm) excitation on the NV charge state. Additional microwave control over the NV spin permits us to examine spin dependence of the charge dynamics.

More specifically, our laser sources comprise CW 532 nm (LaserQuantum Venus), CW 594 nm (Newport R-39582 HeNe), pulsed 531 nm and pulsed 766 nm lasers (both PicoQuant LDH-FA, with pulse widths <100 ps); each source is gated by an acousto-optic modulator (AOM). For simplicity, we will refer to the 532/531 nm lasers as “green,” the 594 nm laser as “yellow,” and the 766 nm laser as “red.” The two pulsed lasers are also separately controlled by a SEPIA II laser driver (PicoQuant PDL 828), allowing for tuning of pulse separation timing. The four optical excitation paths are combined in a homebuilt confocal microscope and focused through a 1.35 NA oil objective (Olympus UPlanSApo 60×) onto a single NV in a type Ia (111)-cut, chemical-vapor-deposition-grown diamond sample (Sumitomo). NV center fluorescence in the wavelength range 635–750 nm is detected with a single-photon counting module (PicoQuant -SPAD). The sample is mounted on an XYZ scanning piezo stage (nPoint NPXY100Z25-219), and a 20 μm copper wire soldered across the diamond allows application of microwave signals for driving spin transitions. Except for the pulsed laser driver, the timing of the experiments is determined by a field programmable gate array (FPGA) card (National Instruments 7841-R FPGA) that controls the scanning microscope and records photon counts; the FPGA card also outputs digital pulse patterns to rapidly turn on and off the laser AOMs and microwaves with 8.3 ns resolution [30,31].

Most of our experiments use a pulse sequence [see, e.g., Fig. 1(a)] that incorporates a spin and charge preparation step before pulsed illumination and a charge detection step afterwards. CW green illumination (2 μs, ~250 μW) is used to fix the initial charge-state distribution to approximately 70% NV− and also initializes the NV− spin primarily into mz = 0. Long, low-power CW yellow pulses (4 ms, 2 μW) are used to initialize and read out the NV center’s charge state before and after pulsed illumination. Yellow illumination efficiently excites NV−, while only weakly exciting NV0, leading to a high fluorescence contrast between the two charge states [17]. The number of photon counts collected during a yellow pulse can be used to determine the initial and final charge states of that pulse [12], with fidelities in the range 91%–97% in our experiments. These measurements allow us to extract the probability that the applied pulsed illumination caused ionization (NV− → NV0) or recombination (NV0 → NV−); see Appendix A for more information. In order to investigate single pulses we fix the pulsed lasers’ repetition rate to 1 MHz and turn on their acousto-optical modulators for a 1 μs illumination window so that on average only single pulses will reach the sample. Fluorescence counts are also collected during this 1 μs window, revealing charge-dependent fluorescence saturation behavior.

We acquire data with three types of measurements that give us information about different aspects of the system dynamics: (1) charge-state switching data (obtained using photon counts during yellow illumination) reveal ionization and recombination processes; (2) fluorescence counts (obtained during pulsed green illumination) provide information on the efficiency with which internal states are excited; (3) time-resolved fluorescence detection allows us to examine fluorescence depletion processes induced by red illumination. Due to slow data acquisition rates, this third measurement is performed with only green and red pulsed excitation [without CW green or yellow illumination; see Fig. 2(a)]. By simultaneously fitting all three types of measurement to a rate equation model, we can well constrain the rates of excitation, stimulated emission, ionization, and recombination under both red and green illumination.

FIG. 1. Excitation and charge-state switching under green illumination. (a) Pulse sequence used for data shown in (b) and (c). CW green (532 nm) is used to initialize the charge-state distribution; low-power CW yellow (594 nm) is used to measure the NV charge state before and after pulsed illumination. Between the yellow pulses, a green pulse is applied. (b) Charge-dependent fluorescence during pulsed green excitation as a function of pulsed green average power. Fluorescence for NV− initialization (red circles) is larger than for NV0 initialization (blue triangles) primarily due to spectral filtering. Black squares show total fluorescence without charge selection. (c) Single green pulse probability of ionization (blue triangles) and recombination (red circles). (d) Four-level rate equation model including transitions between the ground g− (g0) and excited states e− (e0) of NV− and NV0, respectively. Here we show the transition rates under green illumination. Solid lines in (b) and (c) are found by using this four-level model to simultaneously fit data shown in Figs. 1 and 2.
Efficient excitation with minimal ionization is often desired for NV\(^-\) emission or charge-state switching calculated by the four-level rate equation model simultaneously fit to data in Figs. 1 and 2. Since the data in Fig. 1 involve only green pulses, it can be modeled using the transition rates shown in Fig. 1(d). The model comprises the ground and excited states of NV\(^-\) and NV\(^0\), with the relevant rates shown for green-induced excitation (\(G\) for NV\(^-\) and \(cG\) for NV\(^0\)), ionization (\(aG\)) and recombination (\(bG\)). Note that all rates are linear in the green power \(\propto G\); because we explicitly include excited states in our model, it does not contradict nonlinear models that do not include excited states [13,17]. This model corresponds to proposed physical mechanisms for ionization (via excitation to the conduction band and Auger recombination) and recombination (via excitation from the valence band) [18,32]. Note that the NV\(^-\) singlet state is disregarded because it is populated on timescales (\(\sim 10–100\) ns [31]) much longer than the pulse widths (\(\sim 100\) ps) and pulse separations (\(\sim 0.6\) ns) relevant to the measurements in Figs. 1 and 2. Similarly, spontaneous emission can be neglected during the short optical pulses. We can also ignore dependence on the NV\(^-\) spin states in our model because the spin is initialized each time by the same CW green pulse and yellow pulse, spin-dependent singlet transitions are too slow to affect the dynamics, and ionization rates out of the excited state are measured to be spin-independent (data not shown). Similarly, we expect any slow processes within the NV\(^0\) state to be negligible with the consistent initialization procedure and on the sub-nanosecond timescales we study. It is worth emphasizing that this considerable simplification in modeling is obtained by working with single pulses, and ensuring that the charge and spin are identically initialized before each pulse.

We model the single-green-pulse, charge-dependent fluorescence data [Fig. 1(b)] by calculating the NV\(^-\) and NV\(^0\)
excited-state populations after the green pulse for each initialization condition. In particular, we start the model in a combination of NV$^-$ and NV$^0$ ground states with a charge-state distribution set by the initialization fidelities of the first yellow pulse (specifically, to model “NV$^-$” fluorescence, the initial NV$^-$ population is 91.25% ± 0.2% and the NV$^0$ population is 8.75% ± 0.2%, whereas for modeling “NV$^0$” fluorescence, the NV$^-$ population is 5.0% ± 0.2% and the NV$^0$ state, 766 nm illumination cannot cause further ionization or recombination. Working at the same fixed green and varying red powers. This occurs because the red light also induces stimulated emission; once the NV is in the ground state, 766 nm illumination can cause increased charge-state switching, above what was caused by the green excitation pulse; we quantify this by calculating the probability of ionization under green-red pulse pair illumination minus the probability of ionization with the green pulse alone. As shown in Fig. 2(c), the red-induced probability increases in ionization (blue triangles) and recombination (red circles) saturate with increasing red power. This occurs because the red light also induces stimulated emission; once the NV is in the ground state, 766 nm illumination cannot cause further ionization or recombination.

The solid lines in Figs. 2(b) and 2(c) are simultaneous fits to the data of Figs. 1 and 2 using the previously described four-level model, with the relevant rates for red-induced stimulated emission ($R$ for NV$^-$, $fR$ for NV$^0$), ionization ($dR$) and recombination ($eR$) shown in Fig. 2(d). As for green, all rates minimal ionization [29] but did not quantify the relative rates of stimulated emission and charge-state switching. Our pulsed-laser approach allows us to corroborate these ensemble measurements and quantify the effects of red illumination on both NV$^-$ and NV$^0$ states.

The energy of the red light is insufficient to promote either charge state of the NV center to an excited state, so another source of excitation is required before the red laser effects can be investigated. We use a single pulse of green excitation to probabilistically populate the excited states, and subsequently apply a single red pulse. Since we already have data constraining the effects of the green excitation, we can isolate the transitions induced by the red pulse. These green-red pulse pairs were initially chosen to have a 592 ± 2 ps separation time, so that minimal excited-state decay will occur between pulses.

We begin by investigating red-induced fluorescence-depletion as a function of red power. We measure the temporally resolved NV fluorescence using a time-correlated single-photon counting system (PicoHarp 300) synced to the pulsed laser output. In Fig. 2(a) we show example time-resolved traces for pulsed green-only illumination (green, top, 95 µW average green power), and green-red pulse pair illumination (red, bottom, 82 µW average red power) with a 1 MHz repetition rate. The green-only data reveal the expected exponential decay of fluorescence, with a lifetime of 12.2 ± 0.1 ns, while the additional red pulse induces a sharp drop in fluorescence. After subtracting off background counts, we fit the long-time exponential decays for both the green-only and green-red data sets. These fits [solid lines in Fig. 2(a)] provide the fluorescence values that would occur at the location of the red pulse [Fig. 2(d) dashed vertical line], labeled $A$ for green-only and $B$ for green-red. The proportional drop in fluorescence, given by $1 - \frac{A}{B}$, gives a measure of the probability that a single red pulse causes the system to leave the NV excited states, through either charge-state switching or stimulated emission, and will be referred to as the depletion probability. The depletion probability as a function of pulsed red power, shown in Fig. 2(b), was found to saturate near 100% depletion with increasing red power.

We now employ the pulse sequence shown in Fig. 2(c) to investigate whether red illumination causes ionization and recombination. Working at the same fixed green and varying red powers used in the fluorescence depletion measurements, we observe that 766 nm illumination can cause increased charge-state switching, above what was caused by the green excitation pulse; we quantify this by calculating the probability of ionization under green-red pulse pair illumination minus the probability of ionization with the green pulse alone. As shown in Fig. 2(c), the red-induced probability increases in ionization (blue triangles) and recombination (red circles) saturate with increasing red power. This occurs because the red light also induces stimulated emission; once the NV is in the ground state, 766 nm illumination cannot cause further ionization or recombination.

The IV. STIMULATED EMISSION AND CHARGE-STATE SWITCHING

We now move on to investigating the red-laser-induced effects of stimulated emission, ionization, and recombination. Red illumination (700–800 nm) is known to cause depletion of NV fluorescence, but this reduction could come from either stimulated emission or charge-state switching. Stimulated emission is the preferred mechanism as it is expected to conserve spin polarization in NV$^-$ [23], while ionization does not [19]; furthermore, stimulated emission underlies potential laser operation with NVs for which ionization would be a competing process. Earlier measurements on NV ensembles indicated
are linear in red power $\propto R$. Simultaneous fitting requires that we correlate the green powers used in the two data sets, and we find that the green power used in the measurements of Figs. 2(b) and 2(c) corresponds to an average green power of 95 $\mu$W as shown in Fig. 1 [34]. We model the depletion probability data shown in Fig. 2(b) by calculating the fluorescence intensity immediately before and after the red pulse. We begin by initializing the model populations in the ground states of NV$^-$ and NV$^0$, with an appropriate charge-state distribution found independently by applying a sequence of pulses with the same fixed green and varying red powers, and then reading out the charge state (data not shown). The model evolves under green illumination for 100 ps, probabilistically exciting both charge states. The excited states then relax for 592 ps, corresponding to the separation time between the green and red pulses. The green-only fluorescence level just before the red pulse [point A in Fig. 2(a)] is then given by $A = \alpha_+ e_-(t_A) + \alpha_0 e_0(t_A)$, where the excited-state populations $e_+(t)$ and $e_0(t)$ are evaluated at $t_A = 592$ ps after the green pulse and $(\alpha_+, \alpha_0)$ are the same fluorescence scaling factors used to model the data of Fig. 1(b). Next, the model evolves for 100 ps under red illumination [with rates as in Fig. 2(d)], causing stimulated emission and charge-state switching to occur. The fluorescence level immediately after the red pulse [point B in Fig. 2(a)] is $B = \alpha_- e_-(t_B) + \alpha_0 e_0(t_B)$, now with the populations $e_-(t)$ evaluated at $t_B$, immediately after the red pulse. The resulting value $1 - \frac{B}{A}$ is used to fit the measured depletion probability.

The red-induced increase in the charge-state switching probabilities shown in Fig. 2(c) is fit in a similar manner. The model is initialized entirely into one charge state (in the ground state), evolved under the calibrated green pulse, a 592 ps decay time, and a final red pulse, and the resulting population in the opposite charge state (ground plus excited states) is extracted. The simultaneous fit provides the red-dependent rates, which we again quote as a percentage of a given rate, here the NV$^-$ stimulated emission rate [$R$ in Fig. 2(d)]. We obtain relative rates of red-induced ionization $d = 7.1 \pm 0.3\%$, recombination $e = 22\% \pm 2\%$, and NV$^0$ stimulated emission $f = 74\% \pm 6\%$. These results indicate that our red pulses are mainly—but not entirely—inducing stimulated emission.

V. GREEN POWER DEPENDENCE OF RED BEHAVIOR

Recent literature results have shown that the effects of red (or near-IR) illumination on the NV center charge-state distribution strongly depend on green excitation power [13,21]. In particular, a red-induced increase of NV$^-$ population occurs when exciting with low green power, while an induced decrease occurs at high green powers.

To examine if any of these nonmonotonic changes occur in the single-pulse regime, we probe the charge-state switching probabilities of green-then-red pulse pairs [using the same pulse sequence as in Fig. 2(c)], with varying green and red power. Figure 3 shows the red-induced increase in charge-state switching probabilities [above the green-only probability, which follows Fig. 1(c)]. We find that the red-induced increase in ionization and recombination probabilities monotonically increases with green power. Physically, this makes sense: increased green power results in more excited-state population that the red illumination can affect. However, this monotonic behavior would predict a saturating NV charge-state distribution, which is not observed in the literature.

The resolution lies in the fact that we are working with single-pulse pairs. If we examine steady-state behavior instead, our results agree with previous measurements (see Appendix B). Thus, while the underlying ionization behavior is quite simple (as measured in our single-pulse experiments), longer timescale dynamics, such as those associated with the NV$^-$ singlet states and spin states, play a critical role in determining nontrivial steady-state charge distributions. These results thus highlight the utility of single-pulse experiments in isolating intrinsic optical excitation effects from slowly evolving internal dynamics.

The solid lines in Fig. 3 show predictions of our four-level model, using rates extracted from fits to Figs. 1 and 2. These predictions are in reasonable agreement with our data, and discrepancies are likely due to drifts in the alignment of the green laser during data acquisition over a period of a week. The model predicts that at even higher green powers, the maximum red-induced increase in the charge-state switching probabilities will decrease, as green-induced ionization and recombination reduces the population in excited states from which the red pulse can affect the system.

VI. MODEL PREDICTIONS

We now use the four-level model fit results to predict the effects of single green pulses on the NV$^-$ ground state, and single red pulses on the NV$^-$ excited state. Assuming that the system starts entirely in the NV$^-$ ground state, Fig. 4(a) shows the predicted probability to end up in the NV$^-$ excited state (green, top) versus ionizing to NV$^0$ (blue, bottom) after a single green pulse (shaded region shows one standard error). The excited-state probability initially grows due to an increased excitation rate, but eventually starts to decrease as ionization takes over. This leads to an optimum power for maximizing excited-state population after a single green pulse, which was found to be 158 $\mu$W average pulsed power (at 1 MHz repetition rate), leading to an excited-state probability of 88.4% $\pm$ 1.5%.
Starting entirely in the NV$^−$ excited state, Fig. 4(b) shows the predicted probability to end up in the NV$^−$ ground state via stimulated emission (red, top) versus ionizing to NV$^0$ (blue, bottom) after a single red pulse. Both stimulated emission and ionization saturate with increasing red power. At saturating powers (350 μW), there is a predicted 93.3% ± 0.2% probability of stimulated emission, and a 6.6% ± 0.2% probability of ionization.

Excitation followed by stimulated emission has potential applications as a proposed mechanism to engineer fast spin gates between electron and nuclear spins [26] and could also be useful in detecting joint states of closely spaced spins using stimulated emission depletion (STED) microscopy [22]. Such applications ideally seek to maintain spin coherence or polarization during the excitation and stimulated emission processes, which can be lost through ionization [19]. In Fig. 4(c) we examine the predicted probability to fully cycle within NV$^−$ (ground to excited state and back) for varying pulsed green and red powers. We find that a 153 μW pulsed green average power and a maximum pulsed red average power (350 μW) results in an optimum cycling probability of 82.5% ± 0.3%. As another example, our model predicts that with an initial 80% NV$^−$ charge-state population and 90% spin polarization, an optimized green-then-red pulse sequence will result in a final 78% ± 5% NV$^−$ population with 81% ± 9% spin polarization, in rough agreement with the preservation of spin polarization during STED found by Wildanger et al. [23].

We use the pulse sequence shown in Fig. 5(a) (top) to measure the red-induced increase in the ionization and recombination probabilities as a function of pulse separation time $\tau$. Due to the maximum 80 MHz repetition rate of the pulsed laser driver internal oscillator, the minimum step size for pulse separation is 12.5 ns. In order to fully explore the NV$^−$ singlet ionization dynamics, we use microwave $\pi$ pulses to flip the NV$^−$ spin, which is initially mostly polarized into $m_s = 0$, into the $m_s = ±1$ spin states. The $m_s = ±1$ spins have a much higher probability of decaying through the NV$^−$ singlet state, allowing us to observe the role of the singlet state in the ionization dynamics.

The results of this experiment are shown in Fig. 5(a) (bottom). The recombination probability out of NV$^0$ decreases exponentially with approximately the excited-state lifetime, indicating that there are no long-lived NV$^0$ metastable states from which our red laser can induce significant recombination. There is, however, a significant difference in the $\tau$ dependence of the red-induced ionization probabilities for the two NV$^−$ spin states. For $m_s = 0$, ionization decreases exponentially with the NV$^−$ excited-state lifetime, while the for $m_s = ±1$ a much longer decay time is observed, associated with the metastable singlet state.

We fit our results using a five-level rate equation model as shown in Fig. 5(b), now with the inclusion of a level associated with the NV$^−$ singlet state. Similar to the previously described fitting procedures, these data are fit simultaneously with the data shown in Figs. 1 and 2, and the green and red powers are correlated across data sets using ionization and recombination rates. The inclusion of the NV$^−$ singlet state and spin dependence requires that some parameters be fixed to literature values to reduce our parameter space. We fix the NV$^−$ $m_s = ±1$ excited-state lifetime to 6.0 ± 0.1 ns (using the ratio of $m_s = 0$ to $m_s = ±1$ lifetimes from [31]); we set the excited state to singlet state decay probability for $m_s = 0$ spins to 15% ± 5% and for $m_s = ±1$ spins to 55% ± 3% [31,36]; together with the measured $m_s = 0$ excited-state lifetime, these parameters determine the shelving rates $\sigma$ for the two spin states. Last, we fix the spin polarization to 90% ± 10% [1]. The errors on these fixed parameters are considered during error analysis procedures.

VII. VARYING PULSE SEPARATION

The short pulse separation time used during the above experiments ensured that we were primarily probing charge-state switching out of the NV$^−$ and NV$^0$ excited states. By increasing the separation time between the green and red pulses we can investigate ionization and recombination out of metastable states that exist in either charge state. Red-induced ionization out of the metastable singlet state in NV$^−$ has recently been of interest for use in spin-to-charge conversion [13]; our technique could also reveal recombination from metastable states in NV$^0$, such as the predicted quartet state [35].
We model our pulse separation data by initializing entirely into the ground state of either charge state, applying a 100 ps green pulse [using the four-level model in Fig. 1(d)], letting the excited states spontaneously decay for a duration equal to the pulse separation (and allowing decay into and out of the NV− singlet state), and then applying a 100 ps red pulse [relevant rates shown in Fig. 5(b)]. The fit results for the green and red parameters describing excitation, stimulated emission, and ionization or recombination out of the excited states all lie within error of the previously quoted results (see Table I).

The new fit results consist of the NV0 excited-state lifetime of \(1/\nu = 15.9 \pm 1.5\) ns, the NV− singlet lifetime of \(1/D = 141 \pm 20\) ns, and the red-induced ionization rate \(I_r\) out of the singlet state. Because we only measure singlet ionization at a single power, our data does not constrain the functional dependence of \(I_r\) on \(R\) (which likely varies from quadratic to linear as \(R\) increases, due to saturable absorption within the singlet manifold [37]). Nevertheless, we can compare \(I_r\) to the NV− stimulated emission rate at this specific power, where \(R = R_0 = 66.9 \pm 0.3\) GHz; we find that \(I_r = 2.15 \pm 0.5\%\) of \(R_0\). Comparing to the previously quoted red-induced ionization rate out of the excited state of 7.1\% ± 0.3\% of \(R\), one might expect that there would be more ionization occurring out of the excited state when compared to the singlet state. In fact this is not the case, as our model predicts that our red pulse induces an overall 6.6\% ± 0.2\% probability of ionization if the system starts in the NV− excited state and a 14\% ± 3\% probability of ionization out of the singlet state. The reason for this apparent discrepancy is that ionization out of the excited state has to compete with stimulated emission, while singlet ionization does not.

**VIII. CONCLUSION AND OUTLOOK**

Our experiments add to a growing body of work to understand excitation, depletion, ionization, and recombination of the NV center under green and red (or near-IR) illumination. Notably, by working with single green and red pulses, with control over the initial charge and spin states, we are able to greatly simplify the modeling and extract quantitative rates for optically induced charge-state switching out of the singlet and excited states. These models are simple to use and provide a basis for understanding more complex behaviors. For example, our model places constraints on proposed fast electron-nuclear spin quantum gates using enhanced excited-state hyperfine interactions [26], and indicates limits to the spin-polarization-preserving properties of STED [19,23]. We also make a quantitative measurement of the relative rates of stimulated emission and ionization induced by red illumination of NV−, which corroborates earlier ensemble measurements [29] and provides a parameter for calculating lasing thresholds [25]. Moreover, our measurement of relative ionization rates
out of NV\(^{-}\) excited and singlet states is potentially relevant to spin-to-charge conversion [13]. Ultimately, an improved understanding of the fundamental optically induced dynamics of the NV center can impact a broad range of current and future applications.

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APPENDIX A: EXTRACTING IONIZATION AND RECOMBINATION PROBABILITIES

High-fidelity charge-state initialization and readout is performed using yellow (594 nm) illumination. We model the resulting photon count distribution using the techniques described by Shields et al. [12]. This approach assumes that the dynamics can be fully described by the rates of ionization, recombination, and fluorescence for the two charge states, considering all possible charge-state switching sequences and weighting them by their probability to occur. For a given initial charge state that undergoes switching and spends a time \(\tau\) in the initial state and \(t_R - \tau\) in the other charge state, the photon count distribution is given by the convolution of two Poisson distributions, one corresponding to photons emitted during time \(\tau\) in the initial state, and the other corresponding to photons emitted during time \(t_R - \tau\) in the other charge state. The counts from each charge state follow a standard Poisson distribution, which we write as \(\text{Po}(n,x)\), representing the probability of obtaining \(n\) counts given the average number of counts \(x\). With a fixed measurement time \(t_R\), ionization rate \(\Gamma_+\), recombination rate \(\Gamma_0\), NV\(^{-}\) fluorescence rate \(\mu_-\), and NV\(^0\) fluorescence rate \(\mu_0\), we calculate the following photon count distribution given that we are initially in the NV\(^{-}\) charge state:

\[
P(n|\text{NV}^{-}) = P(n|\text{NV}^{-,even}) + P(n|\text{NV}^{-,odd}),
\]

where

\[
P(n|\text{NV}^{-,even}) = \sum_{m=0}^{\infty} \frac{\Gamma_0 \Gamma_+ e^{\tau} m^m (t_R - \tau)^m}{m!} e^{-\Gamma_+/\Gamma_0} \times \int_0^{t_R} dt e^{-\Gamma_+ \tau - \Gamma_0 (t_R - \tau)} P_{n-\mu_- \tau + \mu_0 (t_R - \tau)}
\]

\[
P(n|\text{NV}^{-,odd}) = \sum_{m=0}^{\infty} \frac{\Gamma_0 \Gamma_+ e^{\tau} m^m (t_R - \tau)^m}{m!} e^{-\Gamma_+/\Gamma_0} \times \int_0^{t_R} dt e^{-\Gamma_+ \tau - \Gamma_0 (t_R - \tau)} P_{n-\mu_- \tau + \mu_0 (t_R - \tau)}
\]

which has been written out for all even or all odd numbers of switching events (where \(m\) represents the number of even or odd switching events). The last term in the expression for

\[
P(n|\text{NV}^{-,even})
\]

is for zero switching events. These results can be rewritten using modified Bessel functions of the first kind to obtain the exact results quoted by Shields et al. [12]. The distribution for initialization in NV\(^0\) is found by simply switching \(-\leftrightarrow 0\) in the above equations. The final expected distribution is found by adding the distributions for an initial state of NV\(^0\) and NV\(^{-}\), scaled by the probability to start in each charge state:

\[
P(n) = P(n|\text{NV}^{-}) P(\text{NV}^{-}) + P(n|\text{NV}^0) P(\text{NV}^0),
\]

where \(P(\text{NV}^{-})\) and \(P(\text{NV}^0)\) are the probabilities that the NV starts in the NV\(^{-}\) or NV\(^0\) charge state at the beginning of the yellow pulse.

Using this model we can extract the charge-state switching rates \(\Gamma_-\) and \(\Gamma_0\) under continuous yellow illumination. This is done by collecting photon counts into 100 μs bins for one hour, with further binning performed during postprocessing. An example photon count distribution for 1 h of 2 μW yellow illumination with 50 ms binning is shown in figure 6 a. We fit the data to Eq. (A1) to extract the charge-state switching and photon emission rates, \(\Gamma_+ = 3.89 \pm 0.07\) Hz, \(\Gamma_- = 27 \pm 1\) Hz, \(\mu_0 = 48.87 \pm 0.09\) Hz, and \(\mu_- = 1870 \pm 30\) Hz. Here, since we are in a steady-state situation, the probability to be in NV\(^-\) is given by \(\frac{\Gamma_-}{\Gamma_- + \Gamma_0} = 12.6\%\). When doing these fits we

![FIG. 6. Extracting ionization and recombination probabilities. (a) Photon count distribution from 1 h of 2 μW continuous yellow illumination in 50 ms bins. Fitting (green solid line) allows for extraction of the yellow charge-state switching rates. (b) Typical photon count distribution of a 4 ms, 2 μW yellow pulse after initializing the charge-state distribution with a pulse of CW green. Fits allow us to extract the yellow readout fidelities. (c) Probability tree representing processes with no applied experiment between two yellow pulses. Expressions next to arrows indicate the probability of a given process; e.g., \(f^0\) is the probability to end up in state NV\(^0\) given an initial measurement of \(m^0\). (d) Probability tree representing measurements from two yellow pulses with an applied experiment (green row), allowing for extraction of the probability to ionize \((P_I)\) and recombine \((P_R)\). For both (c) and (d), a similar tree can be constructed for an initial measurement \(m^-\) by exchanging 0 ↔ −.](#)
ensure that the chosen bin time is longer than $1/\Gamma$, allowing for sufficient switching events to occur within the bin time to give a good estimate for $\Gamma$. We use this procedure primarily to characterize the yellow charge-state switching rates for later calculations.

The actual pulses used for charge initialization and readout are shorter and reflect a different initial charge-state distribution. In Fig. 6(b) we show a typical photon count distribution for a 4 ms, 2 $\mu$W yellow pulse (duration and power chosen to maximize fidelities while minimizing duration) after initializing the charge-state distribution with a pulse of CW green [i.e., from the first yellow pulse shown in Fig. 1(a)]. This distribution is fit using the same model discussed earlier but with fixed charge-state switching rates (extracted as described above) and varying charge-state distribution. For the example shown we obtain a 80.6% ± 0.2% NV$^-$ population. We now define a threshold, such that any counts less than or equal to the threshold give result $m^0$ and any counts above give result $m^-$. Using this threshold we want to determine the probability of correctly predicting the NV charge state at the beginning (readout) or end (initialization) of the yellow pulse.

The error in charge-state readout stems from the fact that there is some overlap between the two charge-state distributions, shown as shaded regions in Fig. 6(b). We define the blue shaded region (below the threshold and under the blue dashed curve) as $N^-e^-$ and the red shaded region (above the threshold and under the red dotted curve) as $(1 - N^-)e^0$, where $N^-$ is the NV$^-$ population, and we can extract both $N^-$ and $e^0$ from the fit. Our readout fidelity, depending on charge state, is then given by

$$R^{0(-)} = P(m^{0(-)}|NV^{0(-)}) = 1 - e^{0(-)} \quad (A3)$$

which is the probability to obtain the correct result $m^{0(-)}$ given an initial NV$^{0(-)}$ state. Since the initial state is a given, our extracted readout fidelities will not change with shifts in charge-state distribution, and will remain valid for the second yellow pulse in the sequence [Fig. 1(a)], regardless of experiment. We choose our threshold to maximize population-weighted average fidelity $1 - \frac{(1-N^-)e^0 + N^-e^-}{2}$.

In order to characterize our initialization fidelities, we measure the probability that two yellow pulses give the same result, with no applied experiment in between. We define $Q^{0(-)} = P(m^{0(-)}|m^{0(-)})$, which is the probability of obtaining the result $m^{0(-)}$ during the second yellow pulse given that we had the result $m^{0(-)}$ during the first yellow pulse.

We can expand $Q^{0(-)}$ in terms of initialization and readout fidelities, as shown in a probability tree in Fig. 6(c). We define the initialization fidelities as $I^{0(-)} = P(NV^{0(-)}|m^{0(-)})$, which is the probability of ending in the final charge state NV$^{0(-)}$ given that we measure $m^{0(-)}$. The final charge state of the first yellow pulse will be equal to the initial charge state of the second pulse due to slow charge-state relaxation in the dark (measured to be on the order of a second). Reading off the possible paths to get from result $m^0$ in the first pulse to result $m^0$ in the second pulse gives

$$Q^{0(-)}_Z = I^{0(-)}R^{0(-)} + (1 - I^{0(-)})(1 - R^{0(-)}), \quad (A4)$$

where $Q^{0(-)}_Z$ was found using a similar process, and the $Z$ indicates no applied experiment between pulses. Solving for the initialization fidelities, we find

$$I^{0(-)} = \frac{Q^{0(-)}_Z + R^{0(-)}(1 - Q^{0(-)}}{R^0 + R^{-1}}. \quad (A5)$$

We can now perform an experiment (e.g., apply pulsed illumination) between the two yellow pulses and use the characterized fidelities to calculate the probability that the experiment causes the NV center to switch charge states. Again, we compare two successive charge-state measurements; now there is some probability of switching charge states between the two yellow pulses due to the applied experiment, which we define as $P_I = P(NV^{0}|NV^-)$ for ionization and $P_R = P(NV^-|NV^0)$ for recombination. An example probability tree for finding $Q^0$ is shown in Fig. 6(d). Reading off the possible paths (with a similar method for finding $Q^-$) gives us

$$Q^0 = I^0(1 - P_R)R^0 + I^0P_R(1 - R^-)$$

$$+ (1 - I^0)P_I R^0 + (1 - I^0)(1 - P_I)(1 - R^-),$$

$$Q^- = I^-(1 - P_R)R^- + I^-P_R(1 - R^0)$$

$$+ (1 - I^-)P_I R^- + (1 - I^-)(1 - P_R)(1 - R^0),$$

which can be solved for $P_I$ and $P_R$:

$$P_I = \frac{I^0(Q^- - R^-) + (1 - I^-)(R^- + Q^0 - 1)}{1 - I^0 - I^-}(1 - R^-),$$

$$P_R = \frac{I^-(Q^0 - R^0) + (1 - I^-)(R^0 + Q^- - 1)}{1 - I^0 - I^-}(1 - R^0). \quad (A6)$$

Since we have determined our initialization and readout fidelities $I^{0(-)}$ and $R^{0(-)}$, and we directly measure $Q^{0(-)}$ from experiments using the pulse sequence shown in, e.g., Fig. 1(a), we can extract the ionization and recombination probabilities for an arbitrary experiment.

In summary our process of extracting charge-state switching probabilities consists of (a) characterizing the yellow charge-state switching rates by fitting the photon count distribution from an hour of continuous yellow illumination data, (b) determining the yellow readout fidelities by fitting the photon count distribution of the first yellow pulse [in the pulse sequences shown in Figs. 1(a), 2(c), and 5(a)], (c) finding the yellow initialization fidelities by measuring the probability that the two yellow pulses give the same result with no applied experiment, and finally (d) extracting the probability of ionization and recombination from charge-state measurements before and after the applied experiment. We use standard error and error propagation to obtain the final errors on these values.

**APPENDIX B: STEADY-STATE MEASUREMENTS**

In contrast to our single-pulse or pulse-pair experiments, we also measure steady-state values after a long duration of pulsed illumination. Here we measure the final NV$^-$ charge-state population after 150 $\mu$s of pulsed illumination (1 MHz
red illumination can cause either an increase or decrease in the NV⁻ population, depending on the green power. The amount of red-induced NV⁻ population increase is maximized at low green and red powers, while increasing the red power beyond this point decreases population at the same rate regardless of green power. These results agree with recent literature and point to long-timescale internal dynamics that lead to qualitatively different behavior than what we observe in the single-pulse regime.

FIG. 7. Steady-state measurements. (a) Steady-state NV⁻ charge-state population under pulsed green-only illumination, as a function of power. (b) Red-induced change in steady-state NV⁻ charge-state population during pulsed green-then-red illumination, as a function of green and red power. Repetition rates are 1 MHz.

the resulting fit parameters associated with green excitation, ionization and recombination all lie within error of previously quoted results. The parameters associated with red-induced stimulated emission, ionization and recombination were found to shift slightly to $d = 9.7\% \pm 0.4\%$, $e = 25\% \pm 2\%$, and $f = 84\% \pm 7\%$. The resulting weighted sum of squared errors when ionizing to the NV$^0$ ground state versus the NV$^0$ excited state were found to be 18.5 and 24.1, respectively, indicating that for our data a model consisting of ionization to the NV$^0$ ground state is more likely to be correct.


[34] Relating the green power used in Fig. 2 to the green powers in Fig. 1 is tricky because simple power-meter measurements may not accurately reflect the amount of power reaching the NV center due to drifts in alignment. Moreover, the fluorescence rate is relatively insensitive to green power at high power. Thus we use the green-induced ionization and recombination rates we record for the data of Fig. 2 to identify the corresponding power in Fig. 1.


