Optical Gating of Photoluminescence from Color Centers in Hexagonal Boron Nitride

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ABSTRACT: We report on multicolor excitation experiments with color centers in hexagonal boron nitride at cryogenic temperatures. We demonstrate controllable optical switching between bright and dark states of color centers emitting around 2 eV. Resonant, or quasi-resonant, excitation of photoluminescence also pumps the color center, via a two-photon process, into a dark state, where it becomes trapped. Repumping back into the bright state has a step-like spectrum with a defect-dependent threshold between 2.25 and 2.6 eV. This behavior is consistent with photoionization and charging between optically bright and dark states of the defect. Furthermore, a second zero phonon line, detuned by +0.4 eV, is observed in absorption with orthogonal polarization to the emission, evidencing an additional energy level in the color center.

KEYWORDS: Color center, hexagonal boron-nitride, 2D materials, single photon source

Color centers in wide bandgap semiconductors, most notably diamond and SiC, with atom-like behavior and favorable spin properties, have long held potential for quantum photonic technologies, such as single photon sources and spin qubits.1−3 More recently, color centers in hexagonal boron nitride (hBN)4−6 have emerged as an alternative pathway for fundamental studies, and the rapid progress reflects the wide interest and potential of this material system. These defects span a wide spectral range, from the UV to near-infrared,5,7−10 and have attractive properties for quantum optics, including narrow line width, fast radiative recombination, stable emission, and a relatively high fraction of photons emitted into the zero phonon line (ZPL). A particular benefit is the layered structure of hBN, which allows the fabrication of two-dimensional samples and integration directly with other 2D materials11,12 and/or integrated photonic platforms.13 Furthermore, recent observations14,15 of an optically detected magnetic resonance suggests the potential to host spin qubits within a 2D material.

Typically, the excitation laser plays a crucial role in the optical control of quantum emitters. For example, it plays a role in stabilizing and enhancing the photoluminescence (PL) yield of color centers in diamond,16−20 SiC,21−23 and InGaAs quantum dots.24 In hBN there have been fewer studies in this area, but the choice of excitation energy has been shown to play a role in the absorption efficiency25,26 in stabilization and enhancement of the PL emission,27 and in photochemical modification of the defect.28

In this work, we use multicolor excitation to investigate the photophysical dynamics of yellow emitting color centers in hBN. Typically green excitation is used to study such defects. The addition of a blue repump laser dramatically enhances the PL yield of certain color centers but has a weak effect on others. We investigate two defects in detail using photoluminescence and photoluminescence excitation (PLE) spectroscopy. Through comparison with a rate equation model we find that the observed dynamics can be explained in terms of photoinduced switching between two states. The color centers are pumped into a dark state by resonant, or quasi-resonant, excitation and require a higher energy laser to repump back to the bright state. The repump spectrum is step-like with a defect specific threshold between 2.25 and 2.6 eV. This indicates a band to discrete state repump transition and points to charge state conversion as the origin of this behavior.

The sample consists of few-layer flakes of hBN drop-cast onto a silicon substrate coated with 5 nm of Al2O3 and annealed in nitrogen at 850 °C for 8 min. The optical experiments are performed at 5 K in a closed-cycle cryostat using a microphotoluminescence setup (see the Supporting Information).
Information (S.I.). Three excitation lasers are used; blue and green continuous wave (CW) excitation is provided by a 450 nm (2.76 eV) diode laser and a 532 nm (2.33 eV) diode pumped solid state (DPSS) laser, respectively, and for PLE experiments, a supercontinuum laser is filtered with an acousto-optic tunable filter (AOTF) to give a ~1 nm bandwidth and a pulselength of ~5 ps. The AOTF can be driven with a superposition of radio frequencies to create a filter with multiple pass bands that can be independently controlled (wavelength and transmittance). This enables PLE measurements with simultaneous resonant excitation, as shown in Figure 4(a).

Figure 1 illustrates the enhancement of photoluminescence from color centers in hBN using multicolor excitation. Figure 1(a) shows a typical PL map, recorded with green CW excitation, which shows a number of bright spots resulting from individual, and small clusters of, color centers in hBN flakes. In Figure 1(b), the same region is mapped using coaligned green and blue CW laser beams, resulting in an approximately 2-fold increase in the number of luminescent centers. Two representative emitters are highlighted in the PL maps and defined as defect-A and -B. Defect-A appears bright in both maps, whereas defect-B is dark under green excitation but bright when blue illumination is added. To verify that defects-A and -B are individual color centers, we measure the second-order autocorrelation function, $g^{(2)}(\tau)$ (see S.I.).

Further detail is shown in the PL spectra of the two defects (Figure 1(c) and (d)). Under green excitation the spectrum of Figure 1.
Figure 2. Time-resolved photoluminescence measurements of the zero-phonon line of defect-B. (a) Pulse-sequence used to excite photoluminescence. A blue pulse (450 nm, 2.76 eV) of width 0.5 ms is followed by a green pulse (532 nm, 2.33 eV) of 2.5 ms, with a repetition frequency of 208 Hz. (b) Example PL intensity time trace. Without the blue pulse, the green laser excites low intensity emission. The blue pulse weakly excites PL but also prepares the defect in the bright state. The subsequent arrival of a green pulse results in high-intensity PL, which rapidly decays, indicating the defect is pumped into the dark state. (c, d) PL decay curves resulting from the onset of a green pulse for different (c) blue pump powers (fixed green power of 2.2 mW) and (d) green pump powers (fixed blue power of 10.5 μW). Exponential decay rate as a function of (e) blue and (f) green pump power. (g, h) Peak ZPL intensity from the green laser pulse as a function of (g) blue and (h) green laser power. The red lines in (d–h) are calculated using the rate equation model (see Figure 3(c), Table 1). The gray line in (f) is a fit to the data assuming a one-photon bright-to-dark transition from G to C. The black line in (g) is a fit to the data assuming a two-photon dark-to-bright transition from S to C.
defect-A, in Figure 1(c), consists of a bright ZPL at 2.167 eV, with optical and acoustic phonon sidebands (PSB) that are consistent with previous reports.26,29,30 Addition of the blue laser results in a small enhancement of the overall PL intensity. With blue excitation alone, the ZPL intensity is reduced by a factor of ∼20, for the same excitation power. A comparison of the green laser power dependence of the ZPL of defect-A is made in Figure 1(e), with and without the blue laser. The presence of the blue laser increases the PL intensity at all powers of the green laser, indicating that the blue laser is not just providing extra power but increasing the PL yield of the defect.

In the case of defect-B, the choice of excitation conditions plays a much greater role. The PL spectrum, shown in Figure 1(d), is similar to that of defect-A, with a single ZPL at 2.107 eV and similar PSBs. However, this is only the case for simultaneous excitation with blue and green lasers. With only blue excitation, the ZPL intensity is reduced by a factor of ∼20, whereas for only green excitation the intensity is reduced by a factor of ∼400. The power dependence (Figure 1(f)) shows that even with a green laser power up to several milliwatts the PL yield is still hundreds of times less intense without the blue laser.

To further investigate this behavior, we used time-resolved photoluminescence to study the dynamics of defect-B. Figure 2(a) illustrates the experiment, where a train of alternating blue and green pulses excite the defect, while the PL from the ZPL is directed to an avalanche photodiode. Figure 2(b) plots a typical PL time trace recorded during this experiment, with and without the blue laser. With no blue laser pulses, the photoluminescence from the defect is weak and directly follows the intensity profile of the green laser. We term this the dark state of the defect.

When the blue laser pulses are applied, weak PL is observed for the duration of the pulse. However, the blue pulse also prepares the defect in a bright state, resulting in a strong PL signal at the start of the following green pulse. The peak intensity increases linearly with blue power and saturates at a few microwatts (Figure 2(g)). Under green illumination, the bright state lives for tens of microseconds. The PL decays exponentially with a rate that is quadratic with green laser power, indicating that the bright-to-dark transition is mediated by a two-photon process (Figure 2(f)).

For qualitative understanding, we construct a simple rate equation model motivated by the experimental observations and illustrated in Figure 3(c). The model consists of four energy levels. G and E are the ground and excited levels of the bright state, respectively; S represents the dark shelving state of the defect, and C the conduction band or some higher energy levels of the system. ΓR is the radiative decay of the bright state, and ΓCS and ΓCE are nonradiative transitions which link the dark and bright states. For each transition there is a corresponding optical pump. Fb is the bright state pump rate, Ff is the rate at which electrons are pumped out of the bright excited state, and Fr is the rate at which the color center is pumped out of the shelving state. The resulting system of four differential equations is solved numerically to calculate the photoluminescence, ∝ΔΓR (see S.I. for equations and further model details).

The model is compared to the data in Figure 2(e–h) and shows good agreement with the main experimental observations. First, the rate of the dark-to-bright process, plotted in Figure 2(f), has an exponent close to 2. The model, which has a two-step (two-photon) bright-to-dark transition, reproduces the data well (red-line), whereas a fit assuming a one-photon pump directly from G to C (gray-line) does not, showing a clear preference for two-step pumping from the excited bright state. Second, Figure 2(g) plots the peak PL at the beginning of the green pulse as a function of the average power of the blue laser, Pbd. At low powers, the exponent is ∼1, indicating a one-photon repump process that requires an energy >2.33 eV (green) but <2.76 eV (blue). Fits assuming a one-photon (Fr ∝ Pbd red-line) and two-photon (Fr ∝ Pbd2, black-line) pump from S to C are compared to data and verify that the one-photon model performs best (see S.I. for further discussion). Similar dynamics have been observed for germanium vacancies in diamond, where resonant excitation results in a two-step pumping of the system to a dark state, while a laser of higher energy recovers the system to a bright state with a one-photon process.20

To further verify this model, measurements are made of g(3)(τ) of defect-B for different blue and green excitation powers. In Figure 3(a), g(3)(τ) is plotted for fixed blue power of 0.36 mW and three different green powers. For low green power, g(3)(τ) is close to that expected for a two-level emitter. However, increasing the green power leads to pronounced bunching with an exponential decay on the order of microseconds. Similar bunching behavior has been observed previously for color centers in hBN5,31 and other wide bandgap semiconductors32,33 and attributed to the presence of metastable dark states. In Figure 3(b), g(3)(τ) is plotted for fixed green power of 4.3 mW and three different blue powers.

![Figure 3](https://example.com/figure3.png)

**Figure 3.** Second-order autocorrelation function g(2)(τ) of defect-B with (a) fixed blue (450 nm, 2.76 eV) power of 0.5 mW and varying green (532 nm, 2.33 eV) power and (b) fixed green power of 4.3 mW and varying blue power. The red lines show fits to the data using the model outlined in the text. (c) Rate equation model of two-photon photoionization and one-photon recharging.
When the blue power is low, the bunching is strong; but, increasing the blue power counteracts the tendency of the green laser to pump the defect into the dark state, and $g^{(2)}(\tau)$ close to the 2-level case is recovered. Again, similar observations have been made for the repumping of a color center in hBN but with different emission and pump energies. The $g^{(2)}(\tau)$ function is calculated from the rate equation model with the Quantum Toolkit in Python package and convolved with the instrument response function of the photon counting setup (see S.I. for details and example code). By keeping the decay rates constant and varying the pump rates in proportion to the experimental powers (see Table 1), good fits to the data could be found, as plotted in Figure 3(a) and (b).

Table 1. Rate Equation Model Parameters

| transition | rate (MHz) |
|------------|------------|
| $\Gamma_b$ | 380 ± 1.0  |
| $\Gamma_{cb}$ | 5.0 ± 0.4 |
| $\Gamma_{cs}$ | 2.75 ± 0.08 |
| $F_b$ | $(246.6 ± 0.3)P_c + (63.0 ± 0.4)P_b$ |
| $F_i$ | $(6.9 ± 0.3) \times 10^{-2}P_b$ |
| $F_k$ | $(0.30 ± 0.05)P_b$ |

“The radiative decay rate, $\Gamma_{br}$ is extracted from a fit to time-resolved PL data (see S.I.). All other rates are extracted from fits to $g^{(2)}(\tau)$, as shown in Figure 3. $P_c$ and $P_b$ are the power (in mW) of the green (532 nm) and blue (450 nm) lasers used in the experiments.

From the two-color experiments it is clear that the photon energy of the excitation laser determines if the color center is bright or dark. We therefore perform PLE spectroscopy to investigate how the optical pumping depends on photon energy. We first measure the energy threshold of the repump process for defect-B, plotted in Figure 4(b). The defect is excited by both the green CW laser and the tunable laser and reveals a clear energy threshold for the repump process of ~2.6 eV. To understand why defect-A is bright under green-only illumination, a similar measurement is made. The PLE spectrum is measured with and without the addition of the blue CW laser, as plotted in Figure 4(a), while recording the intensity of the PSB emission. Without the blue laser it is not possible to resonantly excite the defect. The threshold is measured as 2.25 eV by exciting the defect resonantly while simultaneously sweeping the repump laser, as shown in Figure 4(a). Although the ZPL energy of defect-A and -B differs by only ~50 meV, the repump threshold differs by ~350 meV. Nevertheless, the PLE measurements suggest that the two defects can be described by the same model but with different energy dependence of the repump rate, $F_k$.

The simple model describes the data well, and we propose the following interpretation in terms of charge state conversion. The green laser efficiently excites the defect from the ground (G) to excited state (E). A second green photon can further excite the electron into the conduction band, C. From C, the electron can be recaptured, recharging the color center into the bright state; or, the electron is lost or trapped by another defect state, shelving the color center in a dark charge state, S. In this picture, the bright state is negatively charged with respect to the dark state (a similar process involving the valence band is also compatible with the data). A similar two-photon ionization process is responsible for photoswitching from the negative to neutral charge state of NV-centers in diamond and out of the neutral state of the divacancy in SiC. In the case of NV-centers, a two-photon process is also responsible for repumping. However, in our measurements we find that the repump process is one-photon, assuming the photon energy is sufficiently large to overcome the photocharging threshold of 2.25 and 2.6 eV, for defects A and B, respectively. The sharp, step-like, repumping spectra (Figure 4(a) and (b)) indicate a transition between a discrete energy level and a continuum of states, i.e., the conduction or valence band, thus confirming this photocharging picture. There are two possible mechanisms underlying the charging process. The first is that the measured threshold corresponds directly to the charge transfer threshold of the color center itself, and we see reasonable agreement to calculations for the boron vacancy defects (1.5 to 2.4 eV) and CsY$_2$ (2.5 to 3.25 eV) defects (see S.I.). The second explanation is that the repumping threshold represents the charging threshold of a nearby trap state, or states, resulting in free carriers that can then be captured by the color center. We note that many defect species have charging thresholds in this region.

The PLE measurements also reveal further information. In Figure 4(c), the PLE spectrum of defect-A is measured for excitation polarization parallel and perpendicular to the ZPL emission (ZPL$_1$). When the polarization is parallel to ZPL$_1$, absorption resonances are observed in two bands detuned by ~170 meV and ~340 meV, which can be attributed to the one and two optical PSBs, respectively. With orthogonal polarization the PLE spectrum is very different, with only weak absorption into the PSB but with a strong and spectrally narrow absorption resonance detuned from ZPL$_1$ by 400 meV, evidencing a higher energy level in the electronic structure of the color center, labeled ZPL$_2$, in Figure 4(c). The PLE spectrum is similar to that of ZPL$_1$, including a PSB detuned by ~170 meV. Furthermore, the polar plot in Figure 4(c) shows that the two ZPLs are orthogonally polarized, with an 89° misalignment of the absorption dipoles of ZPL$_1$ and ZPL$_2$. Although the absorption resonance is clearly observed when detecting at the energy of ZPL$_1$, there is no PL emission at the energy of ZPL$_2$ when exciting with the blue laser (S.I. Figure S3). These observations explain previous measurements where, depending on the detuning of the laser energy from the ZPL, the absorption and emission dipoles can be aligned or misaligned by up to 90°. As shown in S.I. Figure S3, when exciting with energy above (below) ZPL$_2$, the absorption polarization is perpendicular (parallel) to the emission of ZPL$_1$. A similar PLE measurement of defect-B is presented in Figure 4(d), where low power blue excitation is used to maintain the color center in its bright state. With defect-A, when the polarization of the tunable laser is aligned parallel to the ZPL, absorption peaks corresponding to phonon-assisted processes are observed. In this case, the relative efficiency of the two PSB is greater than for defect-A, and the three PSB is also observed. For the orthogonal polarization the absorption is considerably less efficient at all energies, but there is a weak absorption peak at ~2.51 eV, labeled ZPL$_{PS}$ in Figure 4(d). The detuning of this absorption peak is 403 meV, suggesting a similar origin as in defect-A. Similar measurements of other defects are presented in the S.I. From a survey of 16 defects, 11
show a similar absorption peak with an average energy detuning of 0.5 ± 0.1 eV. Measurements of color centers in diamond again provide some clues as to the origin of ZPL2. In the photocharging cycle of NV centers, resonant excitation both at the ZPL energy of NV0 and at a higher energy level of the NV− results in an increase in the PL yield from NV−.17 In SiV− a higher energy level with perpendicular polarization was identified but was not observed in PL due to selection rules of the energy levels.42 Further work is required to confirm the origin of the transition reported here, but coupled with the measured energy threshold for repumping, it provides key extra information for the identification of the defect species. When compared to theoretical predictions36−38,43−47 and other experimental work,48,49 our results suggest that the defects studied here could be boron vacancies or related to carbon impurities (see S.I. for discussion).

In conclusion, illuminating a color-center in hBN on, or near, resonance drives the color center into a dark state. Applying a second laser, with a photon energy exceeding a defect-specific threshold, repumps the color center back to the bright state. Phenomenologically, the dynamics can be understood in terms of photoswitching between two different charge states of the defect, as is the case for color centers in diamond and SiC.21−23 Furthermore, PLE spectroscopy reveals a sharp absorption resonance, at approximately 400 meV higher energy than the ZPL. These results are of practical use for enhancing the PL yields in both future devices and experimental studies, particularly under resonant excitation,50 and can play an important role in the identification of color centers in hBN.

■ ASSOCIATED CONTENT

* Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.nanolett.0c00751.

Second-order autocorrelation measurements, additional photoluminescence measurements on defect-A, PLE measurements of ZPL1 in other defects, further model details, and example QuTiP notebook (PDF)

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