Cathodoluminescence monitoring of quantum emitter activation in hexagonal boron nitride

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The ability to locally activate or generate quantum emitters in two-dimensional materials is of major interest for the realization of integrated quantum photonic devices. In particular, hexagonal boron nitride (hBN) has recently been shown to allow a variety of techniques for obtaining quantum emitters at desired locations. Here, we use cathodoluminescence (CL) to monitor in situ the local activation of color centers by an electron beam in hBN. We observe that the CL signal saturates at a given surface dose, independently of the electron current density. Based on photoluminescence and photon correlations, we show that the number of photoactive color centers is proportional to the CL signal, and we estimate the maximum density of quantum emitters that can be generated by our technique. Our results provide insights about the activation mechanism and could help to optimize the controlled generation of single photon sources in hexagonal boron nitride.

The recent discovery of quantum emitters in two-dimensional materials1–6 has unveiled new roads towards integrated quantum photonics. Among the most promising 2D platforms, hBN has been shown to host a broad diversity of high-quality color centers, emitting in various wavelength ranges, from ultraviolet to near infrared.7–10 A large fraction of the current research effort is devoted towards control of the position and wavelength of these single-photon emitters (SPEs) for integration into photonic structures.11–13 In this context, we have previously demonstrated the possibility to locally activate a recently discovered family of quantum emitters using an electron beam.14–16 These blue-emitting color centers – abbreviated B-centers in the following – have a reproducible emission in the blue range, with very attractive optical properties, such as a high stability and a remarkably low inhomogeneous broadening of the SPE ensembles.14–16 However, their microscopic structure, as well as the mechanisms at the origin of their generation, remains to be understood. The current understanding is that they stem from the modification of a complex that preexists in the as-grown hBN crystal.15,16 We therefore refer to the generation process as an activation. Here, we demonstrate the possibility to measure the cathodoluminescence (CL) of the B-centers in situ during the irradiation process. We subsequently characterize the sample in photoluminescence (PL) to estimate the number of optically stable SPEs generated by our technique and therefore to infer the efficiency of the method. We expect this work to provide insights on the color center activation mechanism.

We exfoliated a hBN flake grown using the high-pressure high-temperature technique17 on a SiO2/Si substrate. Figure 1a shows a scanning electron microscope (SEM) image of the flake. Its thickness has been measured by atomic force microscopy to be 160 nm. CL is measured by collecting the sample emission by an aluminium parabolic mirror, with a hole to accommodate for the electron beam (figure 1b). The bare CL spectrum of the flake is shown on figure 1b, where the free exciton can be observed at 215 nm, as well as three peaks around 305 nm often attributed to emission from carbon defects.18 The presence of this sub-bandgap emission has recently been identified as a prerequisite for the generation of B-centers in hBN.19 No PL emission from B-centers is observed prior to the irradiation process.

We have then irradiated the hBN crystal using a defocused electron beam of 1 µm diameter in a SEM. The irradiation conditions are kept constant during the whole process, with a voltage of 15 kV and a current of 9 nA. We continuously recorded the CL spectra during the irradiation, using a long-pass filter to eliminate the second order diffraction of the exciton luminescence. Figure 2a shows the CL spectra taken at time intervals of 3 s. The emergence of the typical B-center emission peaks can be clearly observed. Their room-temperature spectra are composed of a zero-phonon-line at 440 nm and a red-shifted optical phonon replica ∼160 meV away. Figure 2b shows the integrated intensity as a function of time. After a stark linear increase, the emission rapidly reaches a plateau, which we attribute to saturation of the SPE activation.

We repeated the experiment at various positions on the crystal while varying the beam diameter between 170 nm and 7.5 µm. The voltage and current are kept identical – therefore the current density varies by more than three orders of magnitude. Figure 2c shows the CL intensity as a function of the dose per unit area for four irradiation spots. Despite some slight variations between the different irradiations, it can be seen that the CL signal follows the same general trend, with a linear increase for surface doses smaller than ∼1013 e·µm−2.
followed by a saturation around $10^{14} \, \text{e} \cdot \mu \text{m}^{-2}$. The CL signal per unit area at saturation is almost independent of the beam diameter. These observations further suggest that electron-beam activation of B-centers is bounded to a maximum density of emitters.

In order to estimate the number of photostable emitters generated by the irradiations, we measured the PL of the flake in a confocal microscope, by exciting with a circularly polarized laser of wavelength 405 nm and power 400 $\mu$W. The PL signal is collected by a high-NA (0.95) air objective, filtered using a bandpass filter centered around the zero-phonon-line (ZPL) of B-centers, and conveyed to avalanche photodiodes. Figure 3a shows a room-temperature PL confocal map measured around a zone irradiated by a 1 $\mu$m diameter beam, denoted by the red circle. The PL signal is the highest within the irradiation spot, and rapidly decreases with increasing distance. A horizontal and a vertical cut of the confocal map are shown on figure 3d and e. These PL profiles are well fitted by Gaussian functions centered at the position of the electron beam. Accordingly, the SPE density decreases exponentially with distance from the spot center. Therefore, at a distance of 1 to 2 $\mu$m away from the center area, the density is low enough that individual SPEs can be observed. No SPE are found at distances greater than 3 $\mu$m from the spot center. We note that back-action from the substrate during irradiation could also play a role in the low but finite density of SPEs around the spot center. Further studies based on suspended hBN flakes as well as numerical simulations may help determine this possibility. The color scale of figure 3a is magnified in regions where SPEs can be individually measured, as marked by the orange dashed circles. Figure 3b shows a low-temperature spectrum of the center area. The characteristic ZPL at 436 nm can be observed, flanked by the acoustic phonon shoulder. The narrow ensemble distribution ($<0.3$ nm) is typical of ensembles of electron-activated B-centers. Figure 3c shows the second-order correlation function of a representative individual emit-
FIG. 3. (a) Confocal PL map of one of the irradiated zones. The red circle denotes the position and size of the electron beam. Inside the orange circles, a second color scale with 10× smaller magnitude was used to reveal the weak luminescence of individual emitters. (b) Low temperature spectrum of the center area. (c) Second-order correlation function of one of the highlighted individual emitters, yielding $g^{(2)}(0) = 0.25 \pm 0.02$. (d), (e) Horizontal and vertical cuts of the center spot. The orange lines are Gaussian fits to the data. (f), (g) Horizontal and vertical cuts around an individual emitter. The orange lines are Gaussian fits to the data.

FIG. 4. Blue dots: Number of SPEs estimated from PL measurements as a function of the CL signal at the end of the irradiation process. Orange line: Linear fit to the data.

In order to estimate the number of individual emitters in the whole irradiated area, we have estimated the integrated intensity associated with individual emitters\(^{19}\). A horizontal and a vertical cut of the confocal map around a single color center are shown figure [3] and g. These profiles are then fitted with Gaussian functions to extract the integrated PL intensity associated with the SPE. We have repeated this estimation for the five highlighted SPEs to obtain the typical integrated PL intensity associated with individual SPEs as well as its standard deviation. We have then estimated the total integrated intensity of the SPE ensemble using a similar procedure, based on the PL intensity cuts shown figure [3]d and e. The ratio between the total and the individual intensities then provides an estimation of the number of emitters\(^{19}\), which we found to be $528 \pm 190$ for this particular spot. Knowing the spot area as well as the flake thickness, we can deduce the SPE density in the irradiated zone, which we find to be $\sim 3 \cdot 10^{15} \text{ cm}^{-3}$.

We have repeated the process for overall six irradiation spots of different diameters on the same flake. Figure [4] shows the estimated number of SPEs as extracted from the PL measurements shown figure [3] as a function of the CL signal as measured at the end of the irradiation process. The number of SPEs is found to be proportional to the CL signal, showing that the latter constitutes an in situ indicator of SPE activation. Through a slight improvement of our collection efficiency, it should be possible to monitor the CL signal at the scale of individual emitters. This could be used to further investigate the SPE activation process. The linear relation between the CL and PL also allows us to infer the electron dose per SPE below saturation, which we found to be in the order of $4 \cdot 10^{10}$ electrons per activated SPE at our voltage conditions (15 kV). This offers the possibility to fix a priori the value of the irradiation dose in order to obtain a certain number of SPEs at a given...
position.

In summary, we have demonstrated that CL allows us to monitor the electron-beam activation of single photon emitters in hBN. Our measurements have revealed a saturation of the activation process, which could help to understand the physical origin of the color centers as well as the irradiation mechanism. The fact that the SPE number saturates at a relatively low density of $3 \cdot 10^{15}$ cm$^{-3}$ is compatible with the current understanding that the irradiation modifies a pre-existing specie of finite density, rather than creating an intrinsic defect from pristine crystal. This limit density corresponds to a mean distance of 70 nm between neighboring centers. This length-scale gives the typical spatial accuracy with which a single color center can be activated. While it may not be sufficient for some near-field optics applications such as controlled coupling to a plasmonic resonator, it is on the other hand compatible with deterministic integration into dielectric photonic structures.

SUPPLEMENTARY MATERIAL

See supplementary material for a description of the characterization of the electron beam diameter and additional PL confocal maps.

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1. C. Chakraborty, L. Kinnischtzke, K. M. Goodfellow et al., Nat. Nanotechnol. 10, 507 (2015).
2. Y.-M. He, G. Clark, J. R. Schaibley et al, Nat. Nanotechnol. 10, 497 (2015).
3. M. Koperski, K. Nogajewski, A. Arora et al, Nat. Nanotechnol. 10, 503 (2015).
4. A. Srivastava, M. Sidler, A. V. Allain et al., Nature Nanotechnol. 10, 491 (2015).
5. P. Tonndorf, R. Schmidt, R. Schneider et al, Optica 2, 347 (2015).
6. T. T. Tran, K. Bray, M. J. Ford et al, Nat. Nanotechnol. 11, 37 (2016).
7. R. Bourelloir, S. Meuret, A. Tararan et al., Nano Lett. 16, 4317 (2016).
8. L. J. Martínez, T. Pelini, V. Waselowski et al., Phys. Rev. B 94, 121405 (2016).
9. M. Kianinia, Z.-Q. Xu, M. Toth, and I. Aharonovich, Appl. Phys. Rev. 9, 011306 (2022).
10. A. Gottscholl, M. Kianinia, V. Soltanov et al., Nat. Mater. 19, 540 (2020).
11. J. Ziegler, R. Klaiss, A. Blaikie et al., Nano Lett. 19, 2121-2127 (2019).
12. N. V. Proscia, Z. Shotan, H. Jayakumar et al., Optica 5, 1128-1134 (2018).
13. X. Xu, Z. O. Martin, D. Sychev, et al., Nano Lett. 21, 8182 (2021).
14. C. Fournier, A. Plaud, S. Roux et al., Nat. Commun. 12, 3779 (2021).
15. A. Gale, C. Li, Y. Chen et al., ACS Photonics 9, 2170 (2022).
16. B. Shevitski, S. M. Gilbert, C. T. Chen et al., Phys. Rev. B 100, 155419 (2019).
17. T. Taniguchi, and K. Watanabe, J. Cryst. Growth 303, 525 (2007).
18. M. Onodera, K. Watanabe, M. Isayama, et al., Nano Lett. 19, 7282 (2019).
19. M. Hollenbach, N. Klingner, N. S. Jagtap, et al., arXiv:2204.13173 (2022).