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Purcell-Induced Bright Single Photon Emitters in Hexagonal Boron Nitride

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reveals the beneficial role of an alumina spacer between hBN and gold, mitigating the electronic broadening of emission from defects proximal to the metal. Our results offer arrays of bright, heterogeneously integrated single-photon sources, paving the way for robust and scalable quantum information systems.

KEYWORDS: boron nitride, single photon emitters, plasmonic resonators, Purcell enhancement, density functional theory, quantum optics

Single photon emitters (SPEs) are essential in quantum
photonic technologies, offering ways to revolutionize quantum computation, communication, and sensing systems.[1](#page-7-0)[−][5](#page-7-0) To enable the practical integration of SPEs into quantum photonic integrated circuits, these nonclassical light sources must possess a combination of properties that are challenging to achieve, including a high spontaneous emission rate, $6-8$ $6-8$ $6-8$ highly radiative quantum efficiency,^{[9](#page-7-0)} room-temperature operation, 10 and deterministic emitter placement.^{[11](#page-7-0)} Hexagonal boron nitride (hBN), a van der Waals material with a wide indirect bandgap of approximately 6 eV, is known to host room-temperature SPEs exhibiting zero-phonon line (ZPL) energies in the 1.5−2.2 eV range.^{[10,12](#page-7-0)−[18](#page-7-0)} The process of defect creation and activation usually requires techniques such as ion bombardment,^{[19](#page-7-0)} irradiation and plasma engineering,^{[20,21](#page-7-0)} thermal annealing, 14 and UV-ozone treatment 22 and results in the random spatial arrangement of defects. Achieving sitecontrolled activation and large-scale integration of SPEs into quantum photonic integrated circuits is, therefore, an active area of research. Previous works have attempted to deterministically place the emitters using focused ion beams, 19,23 19,23 19,23 electron beams, 11,24 11,24 11,24 femtosecond laser pulses, 25 25 25 and nanoindentation.^{[26](#page-7-0)} An alternative approach to inducing

defects by site damage is to activate and couple naturally occurring hBN defects with nanophotonic structures.[27](#page-7-0)−[31](#page-7-0)

Efforts to improve hBN SPEs often entail designing nanoscale cavities that utilize quantum electrodynamic effects such as the Purcell effect, which alters their spontaneous emission rate.^{[31](#page-7-0)−[45](#page-8-0)} Previous works have used plasmonic nanostructures to create Purcell-enhanced emitter-cavity coupled systems.^{46−[49](#page-8-0)} However, since plasmonic metals suffer from scattering and nonradiative losses, field-enhancing nanoantennas must meet stringent design and fabrication criteria for achieving an effective coupling to an emitter. Some of these requirements include spatial alignment of an emitter to the resonant cavity electric fields and an optimized gap between the emitter and nearby metallic surface to avoid quenching of fluorescence.^{34,50} An efficient plasmonic nanocavity-integrated antenna architecture using radiative decay

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Figure 1. PNR-induced SPEs in hBN at room temperature. (a) Schematic of multilayer hBN coupled to a PNR. The hBN is separated from the plasmonic cavity by a 5 nm alumina spacer layer. The wave function in green and red indicates the Purcell-enhanced SPE absorbing and then emitting a photon. (b) Scanning electron microscopy image of a single PNR draped around with multilayer hBN. Scale bar is 500 nm. (c) A confocal scanning photoluminescence (PL) intensity map of a 6 × 7 emitter array containing hBN draped PNRs. Scale bar is 2 *μ*m. Inset shows the optical image of the device; the scale bar is 5 *μ*m. (d) Schematic cross-section of the hBN draped PNR (e) Cross-sectional view of the simulated electric field enhancement |*E*|/|*E*0| distribution profile under normal illumination. The E-field distribution illustrates the confinement of the plasmonic hotspots at the resonator perimeter containing the draped hBN. (f) PL spectra for the device (hBN on PNR), background (PNR only), and hBN on Si pillar and (g) corresponding *g*(2)(*τ*) from the emitter recorded from the green circled regions of the PL intensity mapping from (d). (h) PL spectra and (i) corresponding $g^{(2)}(\tau)$ from the emitter recorded from the blue circled regions of the PL intensity mapping from (d).

engineering approaches could have immense potential and farreaching implications in both fundamental and applied research.

In this work, we achieve the on-site creation of optically activated, bright SPEs in a plasmonic nanoresonator (PNR) antenna architecture draped with unprocessed multilayer hBN. The gold-coated silicon nanopillars act as antenna nanoresonators supporting broadband surface plasmons that create hot spots with strong E-field confinement.^{[51](#page-8-0)} The emitters, placed into near contact with the gold nanoparticles, experience a reduced spontaneous emission lifetime of less than 0.5 ns, while the typical nonresonant hBN defect emission lifetime lies in the 2−5 ns range.^{[32](#page-7-0)−[34](#page-8-0),[52](#page-8-0)−[54](#page-8-0)} Additionally, we measured an average Purcell factor improvement of 2.46 with the incorporation of an alumina spacer. Radiative enhancement of photoluminescence (PL) resulted in bright room-temperature SPEs with sharp emission bandwidth down to sub-30 meV along with an average saturated photon count rate of more than 5 million counts/s. We also observed an SPE yield, determined by zero-delay second-order autocorrelation $g^{(2)}(0)$ < 0.5, of approximately 29%. We performed density functional theory (DFT) calculations on SPEs hosted within hBN layers placed on gold. These calculations, supported by the experimental PL and $g^{(2)}(\tau)$ measurements, reveal electronic quenching for intrinsic defects in close proximity to gold. This work demonstrates a viable path toward realizing scalable,

room-temperature quantum photonic devices utilizing Purcellenhanced bright SPEs in hBN.

PNR consists of a 30 nm-thick gold and a 5 nm-thick alumina spacer layer deposited on top of silicon pillar arrays with a height of 100 nm and a diameter of 160 nm. First, largescale pillar arrays are fabricated in an intrinsic silicon substrate using electron beam lithography and reactive plasma etching. The pillars were coated with gold by e-beam evaporation and alumina by atomic layer deposition. A multilayer 11 nm-thick hBN film was wet-transferred onto the PNR array; see [Supplementary](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02581/suppl_file/nl4c02581_si_001.pdf) Section S1. The hBN received no annealing, radiation, or other processes commonly used to create emitters. Upon the wet transfer, the hBN drapes around the PNR in a "tent-pole" profile (Figure 1a). Figure 1b shows a scanning electron microscope image of a representative PNR draped with hBN. Figure 1c represents a confocal scanning PL intensity map of the final assembled structure recorded at room temperature. Here, the spots with point-like emission marked with green and blue circles correspond to the regions where the transferred hBN is successfully draping the PNRs. The structural parameters with a cross-sectional geometry are illustrated schematically in Figure 1d. In Figure 1e, the fundamental plasmonic mode of hBN coupled to aluminacoated PNR is numerically calculated by the finite-difference time-domain (FDTD) method [\(Supplementary](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02581/suppl_file/nl4c02581_si_001.pdf) Section S2). The simulated electric field distribution shows that at 614 nm,

the gold nanoantenna stimulates E-fields in subwavelength volumes and gives rise to 10-fold enhanced plasmonic hot spots along the gold edges. This resonance wavelength is primarily determined by the size of the nanoantenna and its environment.^{7,[55](#page-8-0)} When hBN is draped on PNRs, the emitters could potentially be placed in the vicinity of the PNR hot spots. When the emitters have a spectral and spatial overlap with the plasmonic modes, near-field enhancement takes place.^{32,[52](#page-8-0)} According to the Purcell effect, in the weak-coupling regime, this helps in establishing an effective out-coupling between the emitter and the cavity and considerably increases the spontaneous emission rate of the systems. 32 The alumina spacer layer can help in suppressing the emitter quenching. $34,50$

[Figure](#page-2-0) 1f represents PL recorded from the resonator site marked with the green circle in [Figure](#page-2-0) 1c. The spectrum shows a sharp ZPL at 618 nm (2.01 eV) with a full width at halfmaximum (fwhm) of 21 meV ([Supplementary](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02581/suppl_file/nl4c02581_si_001.pdf) Section S3). This ZPL is accompanied by a 158 meV red-shifted phonon sideband (PSB) at 671 nm (1.85 eV). Second-order autocorrelation $g^{(2)}(\tau)$ measurements were performed at room temperature with a Hanbury-Brown and Twiss setup to record the nonclassical nature of SPEs from PNR-coupled hBN emitters. [Figure](#page-2-0) 1g shows that the antibunching dip is at $g^{(2)}(0) = 0.29(\pm 0.02)$, proving that this hBN defect site is an SPE. Similarly, the PL spectra from the nearby emission site (blue circle in [Figure](#page-2-0) 1d) are shown in [Figure](#page-2-0) 1h. Two ZPL lines appear at 576 nm (ZPL₁ at 2.15 eV) and 643 nm (ZPL₂ at 1.93 eV), respectively. $ZPL₂$ had a PSB of 162 meV situated at 702 nm (1.76 eV). In both cases, the energy difference between ZPL and PSB lies within 160 ± 5 meV as reported previousl[y14](#page-7-0) [\(Supplementary](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02581/suppl_file/nl4c02581_si_001.pdf) Sections S4−S6). Various ZPL positions may be attributed to the variations in the local mechanical draping profile of the hBN along the different resonators. The antibunching measurement ([Figure](#page-2-0) 1i) represents its SPE behavior with a $g^{(2)}(0) = 0.48(\pm 0.04)$. For the case of hBN on Si pillars, due to the absence of plasmonic resonance modes, the defect sites on Si pillars do not show quantum emission ([Supplementary](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02581/suppl_file/nl4c02581_si_001.pdf) Section S2). We attribute this to the high refractive index and absorptivity of silicon that could potentially lead to emitter quenching.

Figure 2a shows the results of picosecond laser-assisted timeresolved PL measurements. We measured the emission lifetime across three samples with hBN draped over (1) silicon pillars (green), (2) PNRs with no alumina coating (blue), and (3) PNRs with a 5 nm-thick alumina spacer coating (red), respectively. The highlighted recorded lifetimes for hBN on PNRs with alumina coating (red) and without any alumina coating (blue), are 998 and 473 ps, respectively. The recorded emission lifetime for hBN on silicon pillars reaches 2.3 ns. We measured the fluorescence lifetime across 23 different emitters both in the presence and in the absence of an alumina layer (Figure 2b). Without alumina, the average recorded average lifetime is 1180 ps. For alumina-coated PNRs, the average lifetime is 480 ps, showing a 2.46-fold reduction. Note that the typical defect lifetimes reported earlier for hBN land in the range of 2−5 ns,^{[10](#page-7-0),[11,14](#page-7-0),[21](#page-7-0),[26,27](#page-7-0),[31](#page-7-0)} which is ≈4−10 times longer than our observations for the PNR-coupled hBN, denoting significant Purcell enhancement of spontaneous emission (F_p) .^{[7](#page-7-0),[32](#page-7-0)} We express F_p as the ratio of the emitter transition rate (Γ) with the hBN on PNRs to each the native emitter transition rate and the nonalumina emitter transition rate (Γ_0) . Here, the transition rate is the sum of the radiative and nonradiative transition rates. F_p can be defined as the inverse

Figure 2. Purcell-enhanced photophysical characteristics of SPEs in hBN. (a) Time-resolved PL decay showing radiative transition lifetimes recorded from three different device configurations with hBN draped on alumina coated PNR (red dot), PNR without alumina (blue dot), and silicon pillar (green dot). Solid lines represent corresponding fits to a biexponential decay function model. (b) Histogram showing the statistical distribution of radiative transition lifetimes recorded from time-resolved photoluminescence measurements of 23 photon emitters recorded from hBN on alumina-coated PNR (red box) and PNR without alumina (blue box) (c) Fluorescence saturation measurements recorded from three different device configurations mentioned in (a) . (d) - (e) saturation measurements recorded from a total of 12 photon emitters across devices with hBN on PNRs (d) without alumina and (e) with alumina. Insets in (d) - (e) show corresponding histogram of saturated photon count rates. Corresponding fits in solid lines are plotted using a first-order saturation model.

ratio of the lifetime of the emitter (τ) with hBN on PNRs to each the native emitter lifetime and the nonalumina emitter lifetime (τ_0) :

$$
F_p = \frac{\Gamma}{\Gamma_0} = \frac{\Gamma_r + \Gamma_{nr}}{\Gamma_0} = \frac{\tau_0}{\tau}
$$
\n(1)

where Γ*^r* and Γ*nr* are radiative and nonradiative rates, respectively. This reduction in lifetime suggests a spontaneous emission Purcell-enhancement by a factor of 2.46 when comparing only the improvement by the alumina and up to ∼10 when comparing to the nonplasmonic pillar cases in the literature.^{[27](#page-7-0),3}

The presence of an alumina spacer layer allows for higher Purcell enhancement by suppressing emitter quenching. Figure 2 (c) - (e) characterize fluorescence saturation photon count rates across the three cases by measuring the PL intensity as a function of the excitation power. To extract the saturated single-photon count rates, we corrected the measured data sets for the background emission and fitted them using the firstorder saturation model:

Figure 3. Optical characterization of Purcell-induced SPEs. (a) Simplified schematic of the confocal PL setup integrated with a Hanbury-Brown and Twiss (HBT) interferometer that was used to record PL intensity and second-order autocorrelation $g^{(2)}(\tau)$ measurements at room temperature. The objective lens, reflecting mirror, dichroic mirror, flippable mirror, beam splitter, and single avalanche photon detectors are denoted by Obj., DM, FM, BS, and Detector, respectively. (b and c) Normalized PL spectra of six representative photon emitters and (insets) corresponding *g*⁽²⁾(τ) measurements recorded from hBN on PNR (b) with alumina and (c) without alumina. The spectra are normalized without background correction and offset vertically. The solid traces in the insets represent theoretical fits to the recorded $g^{(2)}(\tau)$ data. (d) The fwhm and the antibunching dip $g^{(2)}(0)$ values extracted from measurements recorded for 65 PNR-coupled emitters. (e) Scatter plot of the $g^{(2)}(0)$ values extracted from a total of 110 emitters from hBN on PNR with alumina (red dot) and without alumina (blue dot) device configurations. The error bars represent their corresponding fitting uncertainties. Background subtraction has not been applied to any recorded autocorrelation data.

$$
I(P) = \frac{I_{\infty}P}{P_{\text{sat}} + P} \tag{2}
$$

where I_{∞} and P_{sat} are fitting parameters corresponding to the saturated emission rate and saturation power, respec-tively.^{[11](#page-7-0),[14,](#page-7-0)[56](#page-8-0),[57](#page-8-0)} In the case of hBN on PNRs with alumina (red) and without alumina (blue), the saturated emission rates were 3.4 million counts/s (Mcps) $(P_{sat} = 1.4$ mW) and 1.3 Mcps (*Psat* = 1.8 mW), respectively. For hBN on Si pillars (green), I_{∞} was 0.18 Mcps (I_{∞} = 1.7 mW) [\(Supplementary](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02581/suppl_file/nl4c02581_si_001.pdf) [Section](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02581/suppl_file/nl4c02581_si_001.pdf) S8). The overall PL enhancement factor for hBN on PNR with alumina is 2.28 ([Supplementary](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02581/suppl_file/nl4c02581_si_001.pdf) Section S9), compared to the hBN on PNR without alumina suggesting the radiative enhancement for saturation intensities are comparable to the maximum Purcell enhancements achieved through radiative lifetime measurements [\(Figures](#page-3-0) 2a,b and [Supplementary](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02581/suppl_file/nl4c02581_si_001.pdf) Section S9). Overall, in the case of hBN coupled to alumina-coated PNRs, the efficiently localized Efield and suppressed nonradiative leakage allow the Purcellenhanced cavity dynamics to accelerate the spontaneous emission decay rate toward realizing fast and bright SPEs.

To gain insights into the Purcell-enhanced SPEs, we recorded PL and corresponding $g^{(2)}(\tau)$ measurements at room temperature from 65 emitters. The experimental setup is described in [Supplementary](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02581/suppl_file/nl4c02581_si_001.pdf) Section S11. We measured two

device configurations, one in which the hBN film is coupled to PNRs with alumina and one without, as shown in Figure 3a. Figures 3b,c show examples of PL spectra and corresponding $g^{(2)}(\tau)$ measurements for the two cases, respectively. In Figure 3b, PL spectra show that emission mostly occurs between 585 nm (2.11 eV) and 675 nm (1.83 eV). These spectra also show that emitters coupled to the alumina, on average, had narrower emission fwhm as compared to the latter case of PNRs without alumina. The alumina coating acts as a dielectric spacer preventing electromagnetic quenching and homogenizing the local density of photonic states. This effect has been theoretically predicted for deep-subwavelength nanoresonators.[32](#page-7-0),[33,52](#page-8-0) When the hBN was in direct contact with the gold without the spacer, the recorded PL spectra ($Figure 3c$) mostly consisted of multiple spectral peaks that had lower relative intensities as compared to the case with the spacer. In Figure 3d, the emission fwhm and corresponding $g^{(2)}(0)$ are plotted for 65 representative emitters recorded from hBN draped on PNRs with alumina (red dots) and without alumina (blue squares). In the presence of alumina, the emitters achieved an overall average sub-50 meV emission, with 20 out of the 69 emitters behaving as SPEs with $g^{(2)}(0) < 0.5$, providing a yield of approximately 29%. For the case without alumina, the emission becomes broader with an average fwhm of over 90 meV, and 3 out of 41 emitters are SPEs, which is less than

Figure 4. (a) Energy diagram depicting the effect of electronic broadening of the midgap states on single photon emission (top). Schematic illustration of defect structure V_N (bottom left) within hBN as analyzed through DFT calculations. V_N is considered at three positions (L₁, L₂, and $\rm L_3)$ above a gold (Au) slab. (bottom right). (b) fwhm of the $\rm V_N$ (red), $\rm N_BV_N$ (blue), and $\rm V_BC_N^-$ (brown) DFT midgap states as a function of layer number from the Au surface. Dashed lines correspond to defect structures in the absence of Au. The $\rm L_3$ position of $\rm V_{\rm B}C_{\rm N}^-$ is not considered due to significant out-of-plane deformation in defects on the top layer, which is not relevant to the study of proximity effects. (c) Illustration of the hBN on PNR without alumina and the corresponding structures used in DFT calculations. Partial density of states (DOS) integrated over the three-layer hBN/Au system in two configurations: with V_N on the top layer (L₃) and V_N on the bottom layer closest to Au (L₁). fwhm of the midgap states shaded in red shows significant broadening from the interaction with Au. (d) Similar DOS calculations as in (c) but evaluated for the case of hBN on PNRs with the alumina spacer layer.

10%. This observation suggests that the presence of the alumina spacer helps in boosting the success rate of SPE formation.

We note that emitters from samples without an alumina spacer exhibit broader average line widths [\(Figure](#page-4-0) 3d) and a degraded $g^{(2)}(0)$ ([Figure](#page-4-0) 3e). One reason could be the coupling to the image charge in gold, which may lead to damped single-photon transitions. However, the time-bandwidth product of the single photon peaks significantly exceeds unity, indicating that peak widths are not primarily determined by transition lifetimes. Thus, image-charge-induced damping does not fully explain the variation in the line widths. To better understand the line widths and SPE purity, we used DFT with a model of a hBN/gold interface. We perform DFT calculations for a model system of three-layer hBN supported on gold, with results separately obtained for defects on each layer to characterize proximity effects.

Our emitter creation technique does not involve any active preprocessing. The hBN is draped over PNRs without any specific defect activation process and thus emission is most likely to originate from native hBN defects. We recorded confocal PL measurements from 73 hBN-draped PNR sites that statistically suggest the emitter ZPLs are mostly in the 1.92−2.17 eV range [\(Supplementary](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02581/suppl_file/nl4c02581_si_001.pdf) Figure S14a) with an average separation energy between ZPL and PSB of 165 ± 10 meV [\(Supplementary](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02581/suppl_file/nl4c02581_si_001.pdf) Figure S14b). We find this experimentally acquired ZPL and PSB emission band statistics to be most consistent with hBN's intrinsic defects, such as a nitrogen vacancy (V_N) and an antisite complex in which the nitrogen occupies the boron site, and there is a missing atom at the nitrogen site (N_BV_N) .^{[10](#page-7-0)[,58](#page-8-0)−[60](#page-8-0)} It has been demonstrated, however, that the hBN SPEs at ∼2 eV likely come from carbon 61 61 61 and organic molecules-related 62 62 62 defects. These defect

candidates can result in significant out-of-plain deformation with sensitivity to the local environment. This environment sensitivity may help to explain the range of variations in the ZPL peak positions that are observed experimentally.

We perform DFT calculations on the less environmentsensitive charge-neutral defects V_N and N_BV_N to illustrate the proximity broadening effect in the absence of large structural deformations. Moreover, we also analyze the carbon-based defect $V_B C_N^-$ as it is believed to host a transition consistent with experimental observations of SPE.^{[61](#page-8-0)} Moreover, it has been reported that SiO_2 nanostructures^{[27](#page-7-0),[30](#page-7-0)} and mechanically strained polymer slabs 29 29 29 can induce strain that modifies the electronic band structure of hBN defects and enables single photon emission. Throughout our experiment, SPE behavior is only found in the case of hBN on PNRs ([Figures](#page-2-0) 1f-[1i](#page-2-0)). It should, however, be noted that for the case of hBN on Si pillars, no SPE emission is found. As a result, we attribute that in the case of hBN with PNRs, there could be a synergistic effect arising from a combination of a minimal strainperturbation and a plasmonic mode coupling mechanism likely playing an important role in achieving PNRs' Purcellinduced SPE phenomenon.

DFT calculations reveal that defects in proximity to gold may broaden the midgap states, facilitating numerous nearly degenerate single-photon transitions and thus broader line widths (see conceptual diagram in Figure 4a). The fwhm of midgap states for V_N , $N_B V_N$ and $V_B C_N^-$ are found to be significantly larger for defects positioned near Au (Figure 4b). For instance, the electronic density of states (DOS) for a V_N defected hBN/Au structure shows a narrow midgap peak when the defect is positioned on layer 3 above Au, and a much broader peak when the defect is on the closest layer, layer 1, above Au (Figure 4c). Additionally, charge transfer from the

Au to the hBN is evident, manifesting as demagnetization and the "filling in" of the band gap. While the influence of gold is found to rapidly diminish with layer depth, the electric field enhancement strongly localizes near the hBN/Au interface ([Supplementary](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02581/suppl_file/nl4c02581_si_001.pdf) Figure S18e), where electronic effects are likely still relevant. In contrast, an alumina spacer leads to a more evenly distributed field across hBN [\(Supplementary](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02581/suppl_file/nl4c02581_si_001.pdf) [Figure](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02581/suppl_file/nl4c02581_si_001.pdf) S18e), suggesting that the observed defect states are less likely to be positioned near an interface. We further observe negligible broadening when hBN is supported on alumina ([Figure](#page-5-0) 4d), in contrast to the significant broadening seen in the case of gold. These findings help explain the differences in the emission characteristics of SPEs with and without an alumina spacer and support the beneficial role of an insulating layer between hBN and Au.

In summary, we have demonstrated a PNR antenna architecture for the experimental realization of Purcellenhanced SPEs in hBN at room temperature. We have shown activation and enhancement of hBN SPEs by utilizing the coupling of natural hBN defects with the resonance modes of the PNR platform, accelerating the coupled emitters' spontaneous emission rate. An average of 2.46-fold reduction in SPE lifetime, down to an average of 480 ps, is observed for bright SPEs with an average saturated photon count rate of more than 5 million counts/s and a yield of 29%. We also find that suppression of line broadening by the use of an alumina spacer layer is favorable to the performance and yield of the emitters. DFT calculations further emphasize the beneficial role of the alumina spacer for intrinsic defects in hBN by SPE line width broadening mitigation. Our findings highlight the potential of natural defects in unprocessed hBN on resonant nanostructures as a versatile platform for Purcell-enhanced bright SPEs, offering stable operation at room temperature and emission across a broad spectral range. Our results represent an important step toward scalable room-temperature lightbased quantum information systems.

■ **ASSOCIATED CONTENT**

\bullet Supporting Information

The Supporting Information is available free of charge at [https://pubs.acs.org/doi/10.1021/acs.nanolett.4c02581.](https://pubs.acs.org/doi/10.1021/acs.nanolett.4c02581?goto=supporting-info)

> Details of sample fabrication, wet-transfer, Raman-, and AFM-characterization of hBN, details of FDTD simulations, experimental setup, and structural and optical characterization of various PNR devices draped with hBN; details of spectral analysis of SPE PL and *g* (2)(*τ*) measurements; laser power-dependent *g* (2)(*τ*) and fluorescence saturation; details of DFT calculation for native hBN defect models on gold; and references [\(PDF](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02581/suppl_file/nl4c02581_si_001.pdf))

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Notes

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