One of the core issues of modern optics is the subject of photon interaction with matter. In the Wigner-Weisskopf approximation, the emission rate is directly proportional to the local density of optical states (LDOS) [1]. Over the past decade, photonic resonators with increased LDOS have been exploits to enhanced the emission rate for improving numerous quantum optical devices (e.g., [2,3]). Single photon sources, in particular, promise to see large improvements [4]. While more attention has been given to increasing emission rate, the reverse is also possible in an environment with a decreased LDOS.

Here we demonstrate that by designing a photonic crystal structure with a modified single-defect cavity, we can significantly increase or decrease the spontaneous emission (SE) rate of embedded self-assembled InAs quantum dots (QDs) [5]. Photonic crystals (PCs), periodic arrays of alternating refractive index, are near-ideal test beds for such experiments. Their electromagnetic band structure modifies the LDOS relative to free space and hence the SE rate of embedded QD emitters. We demonstrate that SE of cavity-coupled QDs is enhanced up to 8 times compared to QDs in bulk GaAs. This coupling paves the way to single photon sources with higher out-coupling efficiency and visibility. On the other hand, decoupled QDs emit at up to fivefold decreased rate compared to bulk. In contrast with previous reports of SE rate modification in photonic crystals [6–8], these results represent the first direct measurements of a large lifetime modification of single emitters.

The total SE rate of a QD at position $\mathbf{r}_A$, spectrally detuned from the cavity resonance wavelength by $\lambda - \lambda_{\text{cav}}$, can be expressed as the sum of rates into cavity modes and all other modes, $\Gamma = \Gamma_{\text{cav}} + \Gamma_{\text{PC}}$. In the weak-coupling regime where the cavity decay rate $\kappa = \frac{\pi}{\lambda_0}$ exceeds the QD-cavity coupling strength $|g_{\text{cav}}(\mathbf{r}_A)|$, the SE rate can be calculated from Fermi’s golden rule. The cavity density of states follows a Lorentzian and gives a simple expression for $\Gamma_{\text{cav}}$ [1]. Similarly, $\Gamma_{\text{PC}}$ is related to the LDOS, which in the PC band gap is reduced relative to the bulk semiconductor.

Comparing $\Gamma$ to the bulk semiconductor emission rate $\Gamma_0$ gives the SE rate enhancement factor:

$$\frac{\Gamma}{\Gamma_0} = F_{\text{cav}} \left( \frac{\mathbf{E}(\mathbf{r}_A) \cdot \mathbf{\mu}}{|\mathbf{E}_{\text{max}}| |\mathbf{\mu}|} \right)^2 \frac{1}{1 + 4Q^2(\frac{\lambda}{\lambda_{\text{cav}}} - 1)^2} + F_{\text{PC}}. \quad (1)$$

In this equation, the factors $\left( \frac{\mathbf{E}(\mathbf{r}_A) \cdot \mathbf{\mu}}{|\mathbf{E}_{\text{max}}| |\mathbf{\mu}|} \right)^2$ and $1/[1 + 4Q^2(\frac{\lambda}{\lambda_{\text{cav}}} - 1)^2]$ describe the spatial and spectral mismatch of the emitter dipole $\mathbf{\mu}$ to the cavity field $\mathbf{E}$, and $F_{\text{PC}} \neq \Gamma_{\text{PC}}/\Gamma_0$ is the base SE rate in the photonic crystal without cavity. For perfect alignment, the rate enhancement is given by $F_{\text{cav}} + F_{\text{PC}}$, where the cavity Purcell factor

$$F_{\text{cav}} \equiv \frac{3}{4\pi^2} \frac{\lambda_{\text{cav}}^3}{n^3} \frac{Q}{V_{\text{mode}}}. \quad (2)$$

The refractive index $n = 3.6$ in GaAs and the cavity mode volume $V_{\text{mode}} \equiv \int V \epsilon(\mathbf{r}) |\mathbf{E}(\mathbf{r})|^2 d^3\mathbf{r} / \max[\epsilon(\mathbf{r}) |\mathbf{E}(\mathbf{r})|^2]$. We designed a single-defect cavity in a 2D-photonic crystal to maximize SE enhancement (Fig. 1). The structure was introduced earlier [9] and modified slightly for fabrication purposes. Briefly, we use finite difference time domain (FDTD) to design high $Q$, low $V_{\text{mode}}$ cavities with maximal QD-field interaction $(\mathbf{\mu} \cdot \mathbf{E})$. To better meet fabrication constraints, the design was adjusted based on FDTD analyses of fabricated structures [10]. The cavity supports an $x$-dipole mode with predicted $Q_{\text{pred}} = 45000$ and volume $\frac{1}{2}(\lambda/n)^3$. We patterned this structure on...
160 nm-thick GaAs membranes by a combination of electron beam lithography and dry and wet etching. Grown by molecular epitaxy, these membranes include a single central layer of self-assembled InAs QDs with density $200/\mu\text{m}^2$. The QD emission is inhomogeneously distributed about 920 nm with 50 nm linewidth [inset of Fig. 3(a)]. The cavity resonance frequency was chosen to fall near the middle of the QD distribution.

Cavity resonances of fabricated structures were measured by photoluminescence (PL) spectroscopy at 5 K. In the confocal microscope setup of Fig. 2, the above-bandgap pump beam at 750 nm excites QDs within the ~600 nm focal spot. This spot is much smaller than the 5 \, \mu\text{m}-diameter PC so that only QDs within the crystal are addressed. The fraction of photoluminescence that originates from the cavity is enhanced over the background in two ways: through enhanced emission rate into the cavity mode and increased coupling efficiency between cavity mode and collection optics. The latter efficiency was estimated by FDTD analysis of the cavity mode radiation pattern and equals $\sim 0.07$ for our objective lens with $NA = 0.6$. This enhancement allows us to map out the cavity by pumping the QDs at high intensity, resulting in a broad inhomogeneous emission spectrum that mimics a white light source. The cavity $Q$ then follows directly from a fit to the Lorentzian in Eq. (1). The resonance shown in Fig. 3(a) is a typical example for a fabricated cavity. It matches the polarization dependence predicted by FDTD. The quality factor $Q = 5000$ misses the predicted value $Q_{\text{pred}}$ by an order of magnitude. We believe this decrease to result largely from fabrication inaccuracies as FDTD simulations of fabricated structures, which take these errors into account, do match the measured resonances [10]. We note also that we observed no cavity linewidth narrowing at high pump intensity, so the cold-cavity $Q$ is not overestimated as a result of stimulated emission.

These typical high-$Q$ cavities have the disadvantage that spectral coupling is unlikely. A calculation of the odds for spatial and spectral alignment predicts $P > 20$ in only 4% of cavities with $Q = 4000$. This estimate was derived by Monte Carlo simulation of Eq. (1), relying on the actual QD distributions and FDTD-calculated mode pattern. To analyze a larger data set, we must therefore focus on the easier coupling to low-$Q$ cavities. At present, we will consider three different structures, labeled 2, 3, and 4, with the same cavity design of Fig. 1. Figures 3(b)–3(d) show their PL spectra. High pump-intensity spectra (insets) indicate $x$-dipole resonance modes with quality factors $Q_2 = 200$, $Q_3 = 250$, $Q_4 = 1600$.

![FIG. 1 (color online). FDTD-assisted design of the photonic crystal cavity. The periodicity is $a = 0.27\lambda_{\text{cav}}$, hole radius $r = 0.3a$, and thickness $d = 0.65a$. (a) Electric field intensity of $x$-dipole resonance. (b) Scanning electron microscopy image of a fabricated structure.](image1)

![FIG. 2. Apparatus for photoluminescence and autocorrelation measurements. The incident Ti:sapphire laser beam (continuous or 160 fs-pulsed) pumps QDs at 5 K. The emission is either directly analyzed by the spectrometer (75 cm, N$_2$-cooled CCD), or spectrally filtered and sent to the HBT-type setup for coincidence measurements.](image2)

![FIG. 3 (color online). PL measurements. (a) Structure 1 at high pump intensity (1.5 kW/cm$^2$) shows the polarized $x$-dipole cavity with $Q \approx 5000$. No single exciton lines match the cavity frequency in this case. Inset: Bulk QD spectrum. Peak 1: GaAs band gap transitions. Peak 2: GaAs free and impurity-bound exciton emission. Peak 3: Broadband QD emission. (b),(c) The high pump-power spectra (insets) for structures 2 and 3 reveal resonances with $Q = 250$ and $Q = 200$, respectively. The low-power ($\sim 0.5$ kW/cm$^2$) spectra show coupled single exciton lines that match the cavity polarization. (d) Structure 4 with single exciton line A coupled to dipole cavity mode with $Q \approx 1600$ (inset). Temperature tuning from 6 to 40 K allows improved spectral alignment (shaded line). In all measurements, the Ti:sapphire laser was pulsed at 160 fs with $\lambda = 750$ nm.](image3)
In structure 2, the low-intensity spectrum reveals a single QD exciton line \( A \), spectrally matched with the cavity [Fig. 3(b)]. Because of the low QD-cavity coupling probability, further evidence is required to confirm QD-cavity coupling. This comes from polarization matching. If the dot is coupled, only one of its two near-degenerate orthogonal emission lines [11] is enhanced. Figure 3(b) indicates this is the case. For the coupled system, we expect the excitonic lifetime in bulk semiconductor, which has a distribution of \( \tau_0 = 1.7 \pm 0.3 \) ns, this indicates a rate enhancement by \( F_A = 2.6 \pm 0.5 \). In striking contrast is the lifetime of QD excitons not coupled to a cavity. The cross-polarized emission \( A90^\circ \) has a lifetime of 2.9 ns, far longer than in bulk. As can be seen in Fig. 3(b), this line is much weaker than \( A \) due to low out-coupling and SE rate, as expected. Lines \( B \) and \( C \), which are spectrally far detuned, show lifetimes extended even further to 3.8 and 4.2 ns, respectively.

To verify that the observed emission results from single emitters, we measured photon statistics by the autocorrelation function \( g^{(2)}(t) = (I(t)I(t + t'))/(\langle I(t) \rangle^2) \). For short time scales, this function is measured from the coincidence rate between the two detectors of a Hanbury Brown–Twiss (HBT) interferometer. A start-stop scheme measures time delay \( t' = t_1 - t_2 \) between detection events. The laser repetition period is 13 ns (Fig. 2). Figure 4(a) presents the coincidence histogram for the cavity-coupled emission line \( A \). The antibunching of \( g_A^{(2)}(0) = 0.14 < \frac{1}{2} \) at zero delay time indicates that the emission is indeed from a single emitter. Lines \( B \) [Fig. 4(b)] and \( C \) are also single emitters with \( g_B^{(2)}(0) \sim 0.04 \) and \( g_C^{(2)}(0) \sim 0.03 \) (estimated by exponential fits to the autocorrelation data).

This pattern of short-lived coupled and long-lived decoupled excitonic lines was observed in many other structures. For instance, for structure 3, the PL spectrum [Fig. 3(c)] shows a coupled single exciton line \( A \), while a second line \( B \) (not shown) at 932 nm is clearly decoupled. Line \( A \) again has a short lifetime \( \tau_A = 1.70 \) ns, while line \( B \) has a very long lifetime \( \tau_B = 7.96 \) ns [Figs. 4(c) and 4(d)]. Again, single photon behavior is apparent, with \( g_A^{(2)}(0) = 0.23 \) and \( g_B^{(2)}(0) = 0.05 \). As an example of a strongly shortened lifetime, we show line \( A \) of structure 4. The low pump-intensity spectrum of Fig. 3(d) shows coupling to the \( x \)-dipole cavity. To increase spectral coupling, we shifted line \( A \) from 884.67 to 885.54 nm by temperature tuning from 6 to 40 K [12]. The lifetime is sharply reduced to 210 ps [Fig. 5(b)], roughly 8 times below average bulk GaAs, and photon coincidence is antibunched to \( g^{(2)}(0) = 0.16 \).

Off-resonant dots see up to a fivefold lifetime enhancement. To explain these increased lifetimes, we again turn to Eq. (1). The first term for emission into the cavity mode vanishes as the QD exciton line is far detuned from \( \lambda_{\text{cav}} \). The second term \( f_{\text{PC}} \) is reduced below unity due to the diminished LDOS in the PC for emission inside the PC band gap, leading to longer lifetime. This SE rate modification is illustrated in Fig. 6(b).

We verified the lifetime modification theoretically by FDTD analysis of a classical dipole in the PC with the \( x \)-dipole cavity. The simulation is based on the result that the quantum electrodynamical and classical treatments of radiated power yield proportional results, so that the SE rate is related to the classical dipole radiation power by \( \Gamma_{\text{SE}}/\Gamma_{\text{SE}}^{\text{bulk}} = \rho_{\text{PC}}^{\text{classical}}/\rho_{\text{bulk}}^{\text{classical}} \) for bulk GaAs and the PC [13]. In these simulations, we replicated the conditions under which the experimental data were obtained: 200 dipoles were placed at random positions and orientations in a photonic crystal structure, roughly covering the focal size over which we collected in the experiment. The same type
FIG. 6 (color online). SE rate modification. (a) Experimental (circles) and calculated (bars) data of $\Gamma/\Gamma_0$ of single QD exciton lines vs spectral detuning (normalized by the cavity linewidth $\Delta \lambda$). Coupling was verified by spectral alignment and polarization matching. (b) Illustration of the predicted SE rate modification in the PC as a function of normalized spatial and spectral misalignment from the cavity ($a$ is the lattice periodicity). This plot assumes $Q = 1000$ and polarization matching between the emitter dipole and cavity field. The actual SE rate modification varies significantly with exact QD location.

of single-defect cavity with low $Q \approx 320$ was at the center. With the dipoles radiating at frequency $\frac{2\pi a}{\lambda}$ and detuned from the cavity, we simulated the averaged emitted power and were thus able to calculate the spatial average over all dipoles (QDs) of $\Gamma/\Gamma_0 = \Gamma_{SE}/\Gamma_{SE}^{bulk}$. To find the variance, we repeated this simulation for single emitters at a range of locations, one at a time. These results, plotted in Fig. 6(a), agree well with our experimental observations and confirm earlier theoretical predictions of SE lifetime suppression inside the PC band gap of a similar structure [14]. The TE band gap of our structures extends from $\sim 784$ to $\sim 1045$ nm, calculated by FDTD simulations. Outside the band gap, we expect $\Gamma/\Gamma_0$ to return closer to the bulk value, though we cannot observe this effect as the QD distribution does not reach outside the band gap.

Because of spatial misalignment between the QD and cavity, most of the resonant dots presented here have moderate emission rate enhancement. This misalignment is estimated at about one lattice spacing based on the cavity mode pattern (Fig. 1). Considering the low probability of QD-cavity coupling, these modest rate enhancements are not surprising. With better spatial and spectral matching, Eq. (1) predicts that a rate enhancement of up to $\sim 400$ is possible. Beyond that, Eq. (1) is no longer valid as the system approaches the strong coupling regime near $k \sim [g_{\text{em}}(\mathbf{p}_{\text{em}})]$.

In conclusion, we show that by designing a suitable photonic crystal environment, we can significantly modify the SE rate of embedded quantum dots. For QDs coupled to a PC defect cavity, we observe an up to 8 times faster SE rate and demonstrate photon antibunching. This coupled system promises to increase out-coupling efficiency and photon indistinguishability of single photon sources. In contrast, for individual off-resonant QDs in the PC, we show an up to fivefold SE rate quenching as a result of the diminished LDOS in the photonic band gap. This reduction of SE rate may find applications in QD-based photonic devices (e.g., switching), including quantum information processing (e.g., quantum memory). It also shows that reported QD lifetime reduction due to surface proximity effects [15] should not limit the performance of PC-QD single photon sources. We also note that the simultaneous enhancement of coupled and suppression of uncoupled SE rates results in very high cavity coupling efficiency, $\beta = \frac{\Gamma_{\text{cav}}}{\Gamma_{\text{cav}} + \Gamma_{\text{PC}}}$, even for moderate Purcell enhancement. We find good agreement between experimental SE rate modification and FDTD simulations of SE in the photonic crystal.

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