Long-distance radiative coupling between quantum dots in photonic crystal dimers

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(Received 14 August 2014; revised manuscript received 1 October 2014; published 21 October 2014)

We study the mutual interaction between two identical quantum dots coupled to the normal modes of two-site photonic crystal molecules in a planar waveguide geometry, i.e., photonic crystal dimers. We find that the radiative coupling between the two quantum emitters is maximized when they are in resonance with either the bonding or the antibonding modes of the coupled-cavity system. Moreover, we find that such effective interdot coupling is sizable, in the range of ~1 meV, and almost independent from the cavities’ distance, as long as a normal-mode splitting exceeding the radiative linewidth can be established (strong cavity-cavity coupling condition). In realistic and high-quality-factor photonic crystal cavity devices, such distance can largely exceed the emission wavelength, which is promising for long-distance entanglement generation between two qubits in an integrated nanophotonic platform. We show that these results are robust against position disorder of the two quantum emitters within their respective cavities.

DOI: 10.1103/PhysRevB.90.155436 PACS number(s): 42.50.Ct, 42.70.Qs, 78.67.Hc

I. INTRODUCTION

There is growing interest in controlling the radiative coupling between distant quantum emitters in integrated photonic technologies [1], with the main aim of realizing two-qubit gates in a reliable architecture for quantum information processing [2]. Semiconductor quantum dots (QDs) currently represent very promising candidates to implement single- and two-qubit operations, owing to their large oscillator strength and long decoherence time scales [3].

In general, the mutual interaction between two QDs decays rapidly when their distance is larger than the emission wavelength [4]. Possible ways to enhance the radiative coupling between two distant QDs have been proposed in the last few years, e.g., exploiting electromagnetic field confinement in planar microcavities [5] or photonic crystal (PC) integrated circuits [6]. In fact, PCs in planar waveguides, or PC slabs, have emerged as the preferential nanophotonic platform in view of realizing a fully integrated quantum photonic technology, owing to their engineering flexibility to tailor the propagation and confinement of light at optical or near-infrared wavelengths [7–9]. The strong light-matter coupling regime between a single QD and a cavity mode requires large oscillator strength of the dipole emitter, and a small cavity-mode volume [10]. Such characteristics can naturally be fulfilled in PC cavities [11], where QDs with large dipole moment can be deterministically positioned at the field antinodes [12]; the quantum nature of such a system has now been fully established [13–15]. In addition, PC cavities have already been assessed for broadband and fine postprocessing tuning [16], and tested against intrinsic disorder [17,18]. Recent advances in cavity design have led to remarkable figures of merit, such as flexible far-field engineering [19,20], as well as genetic optimization of ultrahigh $Q$ factors [21], with experimental values on the order of 2 million [22]. Proposals for increasing the mutual interaction distance between two QDs in a PC platform mainly considered using a waveguide as a bus for photon propagation [6,23,24]. The role of disorder on light localization was also addressed [25]. Alternatively, preliminary studies considered the mutual coupling between two QDs positioned at the field antinodes within the same PC cavity [24,26], for which early experimental evidence was shown [27]. The possibility of mediating the inter-QD coupling through the normal modes of a photonic molecule has been considered for coupled microdisks [28], where the distance is limited by evanescent intercavity coupling in free space.

Here we theoretically address the possibility of using strongly coupled PC molecules to efficiently increase the mutual QD coupling rate even at large distance, which has been overlooked in the literature, so far. A schematic representation of such a system is shown in Fig. 1. The photonic molecules we are interested in this work are composed of two coupled PC slab cavities, or PC dimers [29]. We treat the light-matter coupling in a semiclassical formalism based on Green’s tensors, following the formalism developed in Ref. [24] for generic PC structures. The classical electromagnetic fields are solved within a guided-mode expansion approach [30]. Interestingly, PC dimers present peculiar characteristics, such as coupling strength increasing with distance and switching of the fundamental mode bonding/antibonding character [31,32]. In particular, we will specifically consider coupled L3 cavities, i.e., three missing holes in a hexagonal lattice [7]. The coupling characteristics of the normal modes in such structures have already been addressed experimentally [33,34]. A high degree of control on mode tuning has now been demonstrated with a variety of fabrication techniques [35–38], and the possibility of reversing the mode symmetry at fixed intercavity distance has been shown [39,40], which places these systems among the best possible candidates to practically explore the rich physics of coupled quantum modes.
electromagnetic photonic modes in an arbitrary dielectric structure. Following this work, our starting point is the inhomogeneous wave equation for the electric field, with a polarization vector source that takes into account the linear optical response of the QDs (low-excitation regime) through a nonlocal susceptibility tensor. The electromagnetic problem is solved through the Green’s function approach. The field eigenmodes are used to expand the Green’s tensor in the resolvent representation, and the wave function \( \Psi_\alpha(r) \) of the QD \( \alpha \) is represented using a point dipole assumption, which is a good approximation when the electric field does not vary significantly in the region where \( \Psi_\alpha(r) \) is nonnegligible. Hence, the set of equations that define the complex frequency poles of the coupled QD-photonic system (polariton frequencies) can be given as [24]:

\[
(\omega^\beta - \omega)\tilde{\Omega}^\beta(\omega) = \sum_{a=1}^{N} \sum_{m=1}^{M} \mathbf{g}_m^\beta \otimes \mathbf{g}_{m}^{*\alpha} \tilde{\Omega}^\alpha(\omega),
\]

where \( \tilde{\Omega}^\alpha(\omega) = \mathbf{Q}^\alpha(\omega)/\omega(\omega^\alpha - \omega) \), and \( \mathbf{Q}^\alpha = \int dr \Psi_\alpha(r)\mathbf{Q}(r,\omega) \) are the overlap integrals between the wave function of the QD \( \alpha \) and the field \( \mathbf{Q}(r,\omega) \), with \( \mathbf{Q}(r,\omega) = \sqrt{\epsilon(r)}\mathbf{E}(r,\omega) \). In the present formalism, the transition frequencies of the QDs are denoted by superscripts, while the photonic eigenmodes frequencies are denoted by subscripts. The outer product \( \otimes \) is defined as

\[
A \otimes B = \begin{pmatrix} A_x B_x & A_x B_y \\ A_y B_x & A_y B_y \end{pmatrix}.
\]

The quantities \( g_{m}^{\alpha \beta} \) in Eq. (1) are the coupling strengths between the \( m \)th photonic mode and the \( \alpha \)th QD, and they are defined as follows:

\[
g_{m}^{\alpha \beta} = (g_{m,\alpha}^a g_{m,\beta}^a) / (2\pi \omega_0) \epsilon_{\infty} / \hbar \right)^{1/2} dQ_{m}(r),
\]

where \( \omega_0 \) is an average exciton transition frequency, \( d \) is the dipole moment of the QD, \( \epsilon_{\infty} \) is the dielectric constant of the semiconductor, and the \( m \)th mode is evaluated at \( r \), the position of the \( \alpha \)th QD. To obtain Eq. (3), we have used explicitly the point dipole assumption in the QD wave function, i.e., \( \Psi_\alpha(r) = C \delta(r - r_\alpha) \), with \( C \) a normalization constant. From Eq. (1) we define the following tensor:

\[
\tilde{G}^{\alpha \beta}(\omega) = \sum_{m=1}^{M} \mathbf{g}_m^\beta \otimes \mathbf{g}_{m}^{*\alpha} = \frac{d^2}{\epsilon_{\infty} \hbar \c} \tilde{G}(r_\alpha, r_\beta, \omega),
\]

where \( \tilde{G}(r_\alpha, r_\beta, \omega) \) is the Green’s tensor evaluated at the QD positions \( r_\alpha \) and \( r_\beta \). The components \( \tilde{G}_{xy}^{\alpha \beta}, \tilde{G}_{yx}^{\alpha \beta}, \tilde{G}_{xx}^{\alpha \beta}, \) and \( \tilde{G}_{yy}^{\alpha \beta} \) are interpreted as the effective radiative coupling strengths between the QD \( \alpha \) and QD \( \beta \) at the associated transition frequency \( \omega \). The complex frequency poles of the system are found by imposing the singular condition of the associated matrix of Eq. (1), which is mathematically equivalent to
diagonalizing the matrix [24]:

\[
\Lambda = \begin{pmatrix}
\omega_1^x & 0 & \ldots & 0 & g_{1,x}^1 & \ldots & g_{1,x}^M \\
0 & \omega_1^y & \ldots & 0 & g_{1,y}^1 & \ldots & g_{1,y}^M \\
\vdots & \vdots & \ddots & \vdots & \vdots & \ddots & \vdots \\
0 & 0 & \ldots & \omega_N^y & g_{N,y}^1 & \ldots & g_{N,y}^M \\
g_{M,x}^1 & g_{M,y}^1 & \ldots & g_{M,x}^N & \omega_1 & \ldots & 0 \\
\vdots & \vdots & \ddots & \vdots & \vdots & \ddots & \vdots \\
g_{M,x}^1 & g_{M,y}^1 & \ldots & g_{M,x}^N & 0 & \ldots & \omega_M \\
\end{pmatrix},
\] (5)

where possible deviations from the perfectly symmetrical QDs can be introduced in the model through different transition frequencies in the \(x\) and \(y\) directions, respectively. In this way, the real parts of the eigenvalues of \(\Lambda\) represent the polariton frequencies of the coupled system, and the imaginary parts determine their loss rates. The corresponding eigenvectors define the Hopfield coefficients [48], i.e., \(\lambda = (\lambda_1^1, \lambda_1^2, \ldots, \lambda_N^1, \lambda_M^1, \ldots, \lambda_M^N)\), whose square moduli can be directly interpreted as the bare-exciton (or bare-photon) fractions of the mixed polariton state.

For typical self-organized InGaAs QDs with exciton transition energy in the \(\sim 1.3\) eV range, the square dipole moment can be estimated to be \(d^2 \approx 0.51\) eV nm\(^3\), using the measured radiative decay rate of QDs with a radiative lifetime of \(1\) ns [24]. We adopt such value throughout this work. The last requirement of the model is the values of the normalized photonic eigenmodes \(Q_m\) at the QD positions, and their corresponding eigenfrequencies and losses, which are computed using standard methods to solve PC structures. In this work we employ the GME approach [30], which is the best compromise between computational effort and reliable results for strongly localized modes in high dielectric regions, as is the case of PC dimers.

III. PHOTONIC CRYSTAL DIMERS

The PC dimers that we address in this work are formed by two L3 cavities embedded in a PC slab with a hexagonal lattice.

![Graphs and diagrams](image_url)
of holes. The two cavities are nominally identical and have an outward displacement of the two lateral holes \( s = 0.15a \) along the cavity axis, where \( a \) is the PC lattice constant, and their radii are decreased to 80\% of the surrounding PC holes radius, which improves significantly the quality factor [49]. The L3 cavities are disposed in such a way that the line connecting their centers makes an angle of 0°, 30°, 60°, or 90° with the horizontal axis [34], respectively. We chose the same parameters as in Ref. [24] which are relevant in GaAs structures, namely, lattice constant \( a = 260 \text{ nm} \), hole radius 65 nm, and slab thickness 120 nm, with real part of the refractive index \( \sqrt{\varepsilon_{\infty}} = 3.41 \). Here, the photonic modes for the 0°, 30°, 60°, and 90° PC dimers are computed using the GME method [30]. We use a hexagonal supercell of superlattice parameter 24\( a \), and up to 11 025 total plane waves tested for convergence in the 30° and 60° cases; we use rectangular supercells of dimensions \( 27a \times 8\sqrt{3}a \) and \( 14a \times 19\sqrt{3}a \), and up to 11 881 and 14 641 total plane waves tested for convergence, in the 0° and 90° cases, respectively. Since we are interested in the frequency region below the second-order mode of the homogeneous slab, we consider only one guided mode in the expansion but we have checked that adding a second-order mode does not affect the results appreciably. Finally, the real part of the frequencies is averaged in the first Brillouin zone of the superlattice in order to smooth out finite supercell effects.

Figure 2 shows the results of the GME computations for the first two PC dimer modes, associated with the split states arising from the fundamental L3 cavity mode. The bonding states are labeled with the subscript \(+\), while the subscript \( -\) is used for the antibonding states. Panels (a)–(d) correspond to the 0°, 30°, 60°, and 90° PC dimers, respectively. For the 0° case we see a very small splitting between the normal modes, which does not change appreciably with the intercavity distance \( d_c \) (defined from center-to-center of the two PC slab cavities). The behavior of the normal-mode frequencies is quantitatively different for the other cases, in which a large splitting at small intercavity distances can be noticed. Such splitting decreases smoothly for the 30° dimer, and much more rapidly for the 60° and 90° dimers, on increasing \( d_c \). Nevertheless, between \( d_c = 4a \) and \( d_c = 5a \) for the 60° case, and between \( d_c = 7a \) and \( d_c = 8a \) for the 0° case, the splitting increases, which is a rather counterintuitive behavior and it typically occurs in PC molecules, as already evidenced [32,34]. In addition, the bonding \((+)\) and antibonding \((-)\) behavior of the modes changes as a function of distance, which is another interesting phenomenon already seen in experimental measurements on such systems [32]. As it is expected, the resonance frequencies of these PC dimers tend to the values of the isolated L3 cavity for large distances. In Figs. 2(e) and 2(f) we show the \( E_r \) patterns for the bonding and antibonding states, respectively, for the case of 30° at the intercavity distance \( d_c = 5\sqrt{3}a \). The symmetry point of these PC molecules is located at the center of the structure. The bonding (antibonding) mode has an even (odd) symmetry with respect to this point, as it can be seen in the figure. Since the 30° and 60° cases represent the most interesting physical behaviors, we will focus on these throughout the following.

The imaginary parts of photonic eigenfrequencies are calculated with the photonic Fermi’s golden rule using time-dependent perturbation theory [50], and averaging in the first Brillouin zone of the superlattice, in the same way as it was done for the real parts; the corresponding quality factor is computed with these averaged quantities through \( Q = \text{Re}(\omega)/2\text{Im}(\omega) \). Figures 3(a) and 3(b) show the quality factors and loss rates, \( \gamma_m = 2\text{Im}(\omega_m) \), of the split modes for the 30° and 60° PC dimers, respectively. The loss rates are relatively high for large splitting [see Figs. 2(b) and 2(c)] and decrease quickly when \( d_c \) increases, tending to the values of the isolated L3 cavity; this trend can also be verified in the quality factor, taking into account the inverse dependence with \( \text{Im}(\omega) \). Here, both quality factors and loss rates tend to the values of the isolated L3 cavity for large distances, as is expected. The calculations shown in Figs. 2 and 3 agree with previous work on similar systems [34].

IV. QUANTUM DOTS COUPLED TO PHOTONIC CRYSTAL DIMERS

After characterizing the photonic structures, we have all the parameters required to study the coupled QD-PC dimer system. In this section we study the coupling of two QDs coupled to the PC dimers studied in Sec. III, using the formalism presented in Sec. II.

A. Polariton states

We consider each QD positioned at the center of each L3 cavity, which simplifies considerably the problem since the QDs only couple with the \( y \) component of the electric field. This is due to the fact that the \( x \) field component is negligible for small intercavity distances \( d_c \), at the center of each L3 cavity, and eventually tends to zero for large \( d_c \) values (the \( E_x \) is exactly zero at the center of the isolated L3 cavity). We also consider that the loss rates through the PC dimer modes are...
FIG. 4. (Color online) (a) Real part of the eigenfrequencies of the PC-QD system for the 30° PC dimer at $d_c = 5\sqrt{3}a$, for zero dot-dot detuning. (b) The square modulus of the Hopfield coefficients associated with (a). (c) and (d) correspond to the same case as in (a) and (b), respectively, but with a finite dot-dot detuning of $\Delta = 300 \mu eV$. In panels (b) and (d), $\lambda^1_y$ and $\lambda^2_y$ are associated with the QDs 1 and 2, respectively, and $\lambda^1$ and $\lambda^2$ are associated with the PC modes with frequencies $\omega_1$ and $\omega_2$, respectively.

significantly larger than the QD loss rates $\gamma^\alpha$ through other channels; therefore, we set $\gamma^\alpha = 0$. With these conditions the $\Lambda$ matrix of Eq. (5) takes the following simplified form:

\[
\Lambda_y = \begin{pmatrix}
\omega^1 & 0 & g^1_{1,y} & g^1_{2,y} \\
0 & \omega^2 & g^2_{1,y} & g^2_{2,y} \\
g^{1*}_{1,y} & g^{2*}_{1,y} & \omega_1 - i\frac{\Delta}{2} & 0 \\
g^{1*}_{2,y} & g^{2*}_{2,y} & 0 & \omega_2 - i\frac{\Delta}{2}
\end{pmatrix},
\]

where $\omega^1 = \omega^1_{1,y}$, $\omega^2 = \omega^2_{2,y}$, $\omega_1 = \min(\omega_1, \omega_2)$, and $\omega_2 = \max(\omega_1, \omega_2)$. In Fig. 4(a) we show the real part of the eigenfrequencies from diagonalization of the matrix in Eq. (6), for the same dimer configuration shown in Figs. 2(e) and 2(f), as a function of the frequencies of the two QDs at zero dot-dot detuning. We see that vacuum Rabi splitting occurs in the frequency regions associated with the coupled mode resonances (the vertical dashed lines), which is the signature of the strong coupling between the excitonic QD states and the PC modes. Due to the opposite symmetry of the two photonic modes, i.e., symmetric (bonding) and antisymmetric (antibonding), the relevant QD states in the coupling are the excitonic states with the symmetry of the PC mode. In this way, the first anticrossing, which is associated with the bonding PC mode, corresponds to an excitonic symmetric state and the antisymmetric remains dark, while the second anticrossing, which is associated with the antibonding PC mode, corresponds to an excitonic antisymmetric state with the symmetric one remaining dark. When the first PC mode is antibonding, as in the case of some configurations of the 0°, 60°, and 90° PC dimers, the first anticrossing corresponds to an antisymmetric excitonic state, and the second one to a symmetric excitonic state, as verified in our calculations. The Hopfield coefficients are shown in Fig. 4(b), and we can see an interesting collective behavior suggesting that a strong radiative coupling between the QDs is present at each frequency of the PC dimer modes. In the region of the first strong coupling the polaritons 1 and 3 have comparably significant values of the coefficients $\lambda^1_y$ (QD 1), $\lambda^2_y$ (QD 2), and $\lambda^1$ (mode 1), and we see the same behavior for polaritons 2 and 4 in the region of the second strong coupling, but now with the mode coefficient $\lambda^2$ (mode 2), which corresponds to the antisymmetric mode. Usually, it is very likely that two QDs are detuned due to their inhomogeneous distribution of sizes. Therefore, we show in Figs. 4(c) and 4(d) the same analysis made in panels (a) and (b) but now introducing a finite and sizable detuning between the two QDs, $\Delta = \omega^1 - \omega^2 = 300 \mu eV$. We see that under such conditions, symmetric and antisymmetric excitonic states are possible and the dark mode is not present. All the Hopfield coefficients associated with the coupling of the QDs with each PC dimer mode are...
B. Long-distance radiative coupling between the quantum dots

We now try to give an answer to the question of how the radiative coupling depends on the intercavity distance in a PC dimer. The radiative coupling between the QDs is quantified by the tensor components of Eq. (4), which are proportional to the Green’s function evaluated at the QDs positions. Since the QDs are positioned in the center of each cavity, where the $E_y$ component is dominant, the dot-dot coupling is dominated only by the $G_{yy}^{12}$ component of the Green’s tensor, which can be written for the two PC dimer modes as [see Eq. (4)]

$$G_{yy}^{12}(\omega) = \frac{2\pi \omega d^2}{\hbar} \left( \frac{E_{1,y}(r_1)E_{1,y}^*(r_2)}{\omega_1 - i\frac{\gamma}{2} - \omega} + \frac{E_{2,y}(r_1)E_{2,y}^*(r_2)}{\omega_2 - i\frac{\gamma}{2} - \omega} \right).$$

(7)

We plot this effective coupling strength in Fig. 5(a) for the 30° PC dimer as a function of frequency, for all interdot distances studied in the present work. As is expected, when the QDs are in resonance with one of the PC dimer eigenfrequencies, the radiative coupling between the QDs is enhanced and it can remarkably reach values on the order of 1 meV or larger. However, the dot-dot coupling strength at these frequencies is surprisingly minimal at small intercavity distance, which are the cases where the coupling between the cavity modes is largest. As a counterintuitive consequence, the coupling strength between the QDs increases with the interdot distance after the smallest values of $d_c$, and remains relatively constant (small oscillations) at larger values of $d_c$. This can be easily verified in the Fig. 5(b), which shows the function $|G_{yy}^{12}(\omega)|$ at the values $\omega = \omega_1, \omega = \omega_2,$ and $\omega = \omega_{av} = (\omega_1 + \omega_2)/2$, respectively. The function $|G_{yy}^{12}(\omega)|$ evaluated at $\omega_{av}$ shows that even when the QDs are out of resonance from the dimer PC modes, the radiative dot-dot coupling is significant and increases with the interdot distance. For completeness, we show the respective calculations for the 60° dimer in Figs. 5(c) and 5(d). The behavior of $|G_{yy}^{12}(\omega)|$ is very similar to the 30° case, with the difference that the dot-dot coupling increases most rapidly with the interdot distance at small values of $d_c$, and due to the small splitting in the case of $d_c = 4a$ we see a pronounced peak in $|G_{yy}^{12}(\omega)|$ at the mean frequency $\omega_{av} = (\omega_1 + \omega_2)/2$. It is important to recall that these results are valid as long as the PC dimer mode splitting exceeds the photonic radiative linewidth, i.e., where the mode splitting can be spectrally resolved (strong cavity-cavity coupling condition). Qualitatively similar results have been obtained for PC dimers in the 0° and 90° configurations, respectively (not shown here).

These counterintuitive behaviors can be interpreted by analyzing the expression for the coupling constant $G_{yy}^{12}(\omega)$ in Eq. (7). Since the fields are strongly localized in both cavity regions for all intercavity distances in the strong cavity-cavity
coupling regime, the functions \( E_3(r_a) \) depend very weakly on the \( d_c \) parameter; furthermore, the functions \( E_{1,2}(r_1)E_{1,2}^*(r_2) \) and \( E_{2,3}(r_1)E_{2,3}^*(r_2) \) are real due to the point symmetry of the structure with respect to the origin of coordinates, and have approximately the same value with opposite signs due to the opposite symmetries of the two modes [see Figs. 2(e) and 2(f)]. In this way, we can approximate the \( |G_{12}^{12}(\omega)| \) function as

\[
|G_{12}^{12}(\omega)| \approx \frac{2\pi d^2 |g|^{12}}{h} \sqrt{\frac{\Delta_m - i \frac{2\pi}{\omega}}{(\omega_1 - i \frac{2\pi}{2} - \omega)(\omega_2 - i \frac{2\pi}{2} - \omega)}},
\]

where \( |g| = |E_{1,2}(r_1)E_{1,2}^*(r_2)| \approx |E_{2,3}(r_1)E_{2,3}^*(r_2)| \), and \( \Delta_m = \omega_2 - \omega_1 \). Neglecting the \( \gamma \) terms when they do not contribute significantly to the sums, as well as their second-order terms, we obtain the following trends for the effective coupling constant:

\[
|G_{12}^{12}(\omega_1)| \propto Q_1,
\]

\[
|G_{12}^{12}(\omega_2)| \propto Q_2,
\]

\[
|G_{12}^{12}(\omega_2)| \propto \frac{\omega_{2\omega}}{\Delta_m},
\]

which qualitatively explain the results shown in Figs. 3 and 5. Eventually, at very large intercavity distance, where the mode splitting tends to zero and is much smaller than the mode radiative linewidth, the coupling constant \( g^{12} \) tends to zero and consequently the dot-dot radiative coupling vanishes.

C. Nonideal quantum dot positioning

Coupling of the QDs with the PC dimer modes is maximized when the two QDs are exactly positioned at the centers of the two L3 cavities. The semiclassical formalism summarized in Sec. II allows us to study the effect of a nonideal positioning of the QDs through a statistical analysis. To accomplish this, the position of one QD is generated by a two-dimensional Gaussian probability distribution with variance \( \sigma \), and the corresponding position of the other QD is automatically determined since the dot-dot distance is maintained fixed. This model simulates in a realistic way possible misalignments in the writing stage of the photonic pattern with respect to the QDs, which can be deterministically positioned with a high degree of accuracy [12,13]. Figure 6 shows the results of this analysis, where the absolute value of the Green’s tensor component \( G_{12}^{12}(\omega) \) is studied as a function of the interdot distance for the 30° dimer. Panels (a) and (b) correspond to the cases \( |G_{12}^{12}(\omega)| \) and \( |G_{12}^{12}(\omega)| \), respectively. The curves were averaged over 1000 realizations for each value of \( \sigma \), and the standard error is explicitly shown in the figure with the corresponding error bars [51]. We see that the dot-dot coupling strength remains of the order of 1 meV for a statistical variance of \( \sigma = 26 \text{ nm} \), which is comparable to the precision that can be achieved with modern sample-fabrication techniques [15]. Furthermore, the coupling strength is sizable even for larger values of \( \sigma \), as can be seen from the figure. As an example, the coupling constant is of the order of 0.7 meV for \( \sigma = 52 \text{ nm} \). We point out that in the present analysis the QDs are not necessarily positioned at the cavity centers, since their positions are generated randomly with a Gaussian distribution, and consequently the QDs will also couple to the \( E_x \) component of the field. As a consequence, the other components of the Green’s tensor might be nonnegligible. Nevertheless, \( |G_{12}^{12}| \) is much larger and dominates for the values of \( \sigma \) considered here. As a final remark, we notice that the results for the 0°, 60°, and 90° PC dimers are equivalent to the 30° case, and coupling strengths of the same order of magnitude can be obtained.

V. CONCLUSION

We have studied the radiative coupling between two distant quantum dots embedded in the two cavities of photonic crystal molecules in planar waveguide geometry, or photonic crystal dimers, by using a semiclassical formalism based on the Green’s tensor. The photonic eigenmodes are found by guided-mode expansion, which allows us to estimate real and imaginary parts (losses) of the photonic eigenmodes, as well as the spatial mode profiles. Specifically, we have considered two L3 cavities made of three missing holes in a hexagonal lattice, in which field antinodes occur at each cavity center.
where the quantum dots can be placed. Parameters have been chosen to describe current systems typically fabricated with III-V semiconductors, such as InGaAs quantum dots in GaAs thin membranes.

Irrespective of the coupling angle between the two cavities (0°, 30°, 60°, 90°, respectively), we have shown that the effective interdot coupling is enhanced when the QDs are in resonance with either of the two normal modes of the PC dimer. Under such resonance conditions, and in the strong cavity-cavity coupling regime, the interdot coupling strength is actually proportional to the quality factors of the normal modes (bonding or antibonding), and it can be of the order of 1 meV for the cases considered here, which is at least an order of magnitude larger than typical values achieved in one-dimensional systems [24]. Since these quality factors can also increase as a function of the intercavity distance, tending to the limiting value set by the isolated cavity mode (in the $10^5$ range for the present case), then the radiative coupling can also increase with distance. Moreover, since the quality factors remain approximately constant at large distances, the radiative coupling can also remain constant at interdot separation that is significantly larger than their characteristic emission wavelength. In addition, we have also shown that when the QDs are out of resonance from the PC dimer modes, the interdot radiative coupling is still significant and inversely proportional to the normal-mode splitting between bonding/antibonding photonic states. Eventually, the mutual QD coupling goes to zero when such normal-mode splitting is blurred by their linewidth. Finally, with a statistical analysis of the nonideal positioning of the QDs within their respective cavities, we could quantitatively conclude that with Gaussian variances comparable to the precision of modern sample fabrication techniques, the coupling strength is not strongly affected and remains of the order of 1 meV.

We notice that part of the conclusions drawn in the present work can be extended to any type of photonic dimer, in principle. In fact, while the mentioned 1 meV coupling strength is quantitatively valid for the specific systems considered here, it is still the general conclusion that the radiative coupling between quantum dots in photonic dimers remains constant even at significantly large interdot distances when two identical cavities are in the strong cavity-cavity coupling regime. This is the main result of this work, which can be particularly relevant in view of possibly realizing PC dimers with normal-mode splitting resolved even at very large distances [41]. The latter could definitely solve the issue of entangling distant qubits for applications in quantum information processing in an integrated photonic platform [2], where independent optical manipulation of the two quantum emitters could be straightforward if their mutual distance significantly exceeds the resonant wavelength.

ACKNOWLEDGMENTS

We acknowledge financial support from FAPEMIG, CAPES, and CNPq, Brazil. This work was partially financed by PVE/CSF Project No. 407167/2013-7. The authors would like to thank L. C. Andreani, A. Badolato, M. Gurioli, M. F. Santos, and V. Savona for useful discussions. J.P.V. would like to acknowledge the Photonics and Nanostructures group for the wonderful reception during his stay in Pavia.

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