Dynamic vibronic coupling in InGaAs quantum dots

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The electron-phonon coupling in self-assembled InGaAs quantum dots is relatively weak at low light intensities, which means that the zero-phonon line in emission is strong compared to the phonon sideband. However, the coupling to acoustic phonons can be dynamically enhanced in the presence of an intense optical pulse tuned within the phonon sideband. Recent experiments have shown that this dynamic vibronic coupling can enable population inversion to be achieved when pumping with a blue-shifted laser and for rapid de-excitation of an inverted state with red detuning. In this paper we confirm the incoherent nature of the phonon-assisted pumping process and explore the temperature dependence of the mechanism. We also show that a combination of blue- and red-shifted pulses can create and destroy an exciton within a timescale ∼ 20 ps determined by the pulse duration and ultimately limited by the phonon thermalisation time.

INTRODUCTION

Vibronic sidebands are observed in the optical spectra of many solid-state materials [1], with Ti:sapphire [2] and nitrogen-vacancy (NV) centres in diamond [3,5] being good examples. The key point about a vibronic transition is that it involves the simultaneous absorption or emission of a photon and a phonon (or phonons) as the electron jumps between two electronic states. The sideband spectra are continuous bands, although sub-structure can frequently be identified due to the involvement of specific phonon modes, especially at low temperatures. For example, in the case of NV centres in diamond, clear structure can be identified that corresponds to the coupling of the electronic state to the A1 mode (ℏω = 65 meV), giving rise to resolved side-peaks in both absorption and emission at integer multiples of the phonon energy [3,4]. The coupling to the vibrational modes is so strong that only a few percent of the emission occurs in the zero-phonon line, with most of the photons emitted from the sidebands [5]. The relatively weak intensity of the zero-phonon line has serious consequences for practical applications of NV centres in optical quantum information processing.

The reason why the phonon sidebands in materials like diamond NV centres and Ti:sapphire are so strong is that both the electronic and vibrational modes are strongly localized on length scales similar to the unit cell size. This means that the overlap between the electronic wave functions and the phonon modes is large, and hence the vibronic coupling strong. By contrast, InGaAs quantum dots (QDs) have envelope wave functions localized on much larger length scales that are determined by the size of the dot, i.e. ∼ 10 nm. This means that the coupling to phonons is relatively weak, with the dominant interaction being to longitudinal-acoustic (LA) phonons via deformational potential scattering. The weak vibronic coupling in these dots gives rise to very strong emission in the zero-phonon line, with only ∼8% in the sideband at cryogenic temperature [6,7]. This makes InGaAs QDs excellent single-photon sources [8]. It also ensures that the light-matter coupling is strong, with optical dipole moments typically in the range of ∼ 30 Debye (∼ 1×10^{-28} Cm) [9]. Moreover, the relatively weak electron-phonon coupling leads to long coherence times that are ultimately limited only by the radiative lifetime [10] which facilitates their application in coherent control experiments [11,14]. This contrasts with bulk and quantum well samples, where the coherence time is only a few picoseconds at best [15,17].

The vibronic sideband in InGaAs quantum dots manifests itself in a number of important experimental situations. One example is the emission of photons from a nano-cavity when the dot and cavity are out of resonance, with the emission involving either the absorption or emission of a phonon depending on the sign of the detuning [8,18]. Another case is the observation of QD emission when pumping in the phonon sideband in a resonance fluorescence geometry [19]. The difference in the frequency of the pump laser and the QD exciton line facilitates spectral selection of the QD photons [20] and can be exploited for stabilizing the frequency of the zero-phonon line [21]. The sideband also appears in four-wave mixing experiments [22].

A key point about the experiments described above is that the optical pumping is relatively weak, so that the QD-laser system is in the weak-coupling limit. In this paper we explore the other limit where the QD-laser coupling is strong, such that a dressed-state picture is appropriate. This limit was addressed in a theoretical paper in 2013 by Glässl et al, where it was predicted that strong driving in the phonon sideband could lead to exciton populations approaching full inversion [23]. The predictions of the theory were confirmed independently by three experimental groups in 2014–15, both for pump-
MECHANISM OF DYNAMIC VIBRONIC COUPLING

The starting point for DVC is the coupling of excitons to the acoustic phonon bath by the deformation potential [22, 23]. In the absence of a laser field this leads to non-exponential pure dephasing of the excitonic dipole [22]. The behaviour of the coupled system becomes somewhat more interesting however when a strong laser field is applied. We consider a laser pulse with energy detuning $\hbar \Delta$ and area $\Theta$: 

$$\Theta = \int_{-\infty}^{+\infty} \Omega_R(t) \, dt, \quad (1)$$

where $\Omega_R(t)$ is the time-dependent Rabi frequency determined by the optical dipole moment and the time-varying electric field amplitude of the pulse. Using the rotating wave approximation, the Hamiltonian in the rotating frame becomes time independent and one can thus define the dressed states of the laser-QD states as the corresponding eigenstates. These are given by:

$$|\alpha\rangle = \sin (\theta) |0\rangle + \cos (\theta) |X\rangle \quad (2)$$

$$|\beta\rangle = \cos (\theta) |0\rangle - \sin (\theta) |X\rangle, \quad (3)$$

where $\theta$ is an admixing angle defined by:

$$\tan (2\theta) = -\Omega_R/\Delta, \quad 0 \leq 2\theta \leq 180^\circ. \quad (4)$$

The energies of the dressed states are given by:

$$E_{\pm} = \frac{\hbar}{2} (-\Delta \pm \Lambda(t)), \quad (5)$$

where $\Lambda(t)$ is an effective Rabi frequency defined as:

$$\Lambda(t) = \sqrt{\Omega_R(t)^2 + \Delta^2}. \quad (6)$$

The significance of these dressed states is that, in the presence of a driving laser such that $\theta > 0$, both eigenstates of the system contain an excitonic component and may couple to the acoustic phonon bath. This enables phonon-mediated relaxation from $|\alpha\rangle$ to $|\beta\rangle$. However, this relaxation is only possible when the states are admixed by the laser, giving rise to the dynamic nature of the vibronic coupling.

Figure 1 illustrates the dressed state energies (green lines), admixture angle $\theta$ (black line) and occupancy of $|\beta\rangle$ (blue line) against time. At $t \to -\infty$, the states are split by the laser detuning $\hbar \Delta$. During the passage of the pulse the states are admixed and the splitting becomes $\hbar \Delta$ as shown by the shift centred at $t = 0$. This enables relaxation from $|\alpha\rangle$ to $|\beta\rangle$ (blue arrows) by emission of an LA phonon with energy $\hbar \Delta$.
\[ |\beta\rangle \text{ only occurs whilst the states are admixed, turning the system from weakly to strongly vibronic and back again in approximately 20 ps.} \]

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure2}
\caption{(a) Illustration of the energies of the dressed states (\(|\alpha\rangle\) and \(|\beta\rangle\) plotted against detuning \(\Delta\) at a fixed time. The bare QD states in the frame of the laser are plotted for reference as dotted black lines. At \(\Delta = 0\) the states anti-cross with a splitting of \(\hbar \Omega_{0R}\). It can be seen that \(|\alpha\rangle\) is dominated by the ground state at \(\Delta > 0\), leading to phonon-assisted excitation of the exciton. Conversely, \(|\alpha\rangle\) is primarily excitonic in character at \(\Delta < 0\), and thus phonon-assisted de-excitation of the exciton occurs. The blue arrows illustrate the relaxation process corresponding to the emission of an LA phonon whilst the red arrows show the competing phonon absorption process. (b-e) Phenomenological energy level diagrams of the phonon emission and absorption processes for both positive and negative detunings. For simplicity, the phonon relaxation is incorporated as a virtual state (dashed lines).}
\end{figure}

The transfer between the dressed states shown in Fig. 1 can be exploited to achieve ultrafast incoherent excitation and de-excitation of the exciton by detuned laser pulses. Beginning with excitation, in the case of \(\Delta > 0\) the higher energy dressed state \(|\alpha\rangle\) is dominated by the crystal ground state whilst the lower energy state \(|\beta\rangle\) is primarily excitonic in character as illustrated to the right of Fig. 2(a). During the passage of the laser pulse, admixing of the states allows relaxation to occur from \(|\alpha\rangle\) to \(|\beta\rangle\) by emission of an LA phonon (\(\Gamma_{ph}^+\) - blue arrows).

For a sufficiently strong pulse, most of the population relaxes into \(|\beta\rangle\) and the exciton-dominated nature of \(|\beta\rangle\) means that the population of the exciton state after the passage of the pulse is high. Fig. 2(b) shows a phenomenological level diagram of the phonon-assisted excitation process whilst Fig. 2(c) illustrates the competing process whereby a phonon is absorbed (\(\Gamma_{ph}^-\) - red arrows) and a photon emitted (purple arrow). The latter process is weak at low temperatures.

In the case of \(\Delta < 0\) the main difference is that the characteristics of the dressed states are now exchanged. During the passage of the laser pulse, relaxation by phonon emission again occurs into the \(|\beta\rangle\) state. However, as this state is now dominated by the crystal ground state, the effect of the relaxation is instead to de-excite the exciton with most of the population left in the ground state after the passage of the laser pulse. Fig. 2(d) shows an illustrative schematic of this process with the emission of a photon (purple arrow) accounting for the rest of the energy difference. This emission is stimulated by the laser and hence the process is termed LA-phonon stimulated emission (LAPSE). Fig. 2(e) again illustrates the competing phonon absorption process.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure3}
\caption{Plot of the exciton-phonon interaction strength \(J(\omega)\) as determined by fitting data measured in Ref. [21]. The interaction strength peaks at a cut-off of around 2 meV and rolls off rapidly beyond this.}
\end{figure}

The physical factors that determine the efficiency of phonon-assisted excitation and de-excitation may be considered by analysing the parameters that enter eq. (4). The degree of admixing increases by increasing the driving strength (\(\Omega_R\)) or decreasing the detuning (\(\Delta\)) (noting that \(\Delta\) should exceed the laser linewidth to exclude resonant coherent driving), leading to more efficient relaxation. However, both of these dependencies are modified by the properties of the phonon bath. The exciton-phonon interaction strength is characterised by the function \(J(\omega)\) which increases with \(\omega\) at first due to the rising phonon density of states, and then rolls-off rapidly beyond a cut-off frequency which is typically around 1-2 meV [12]. The physical origin of this cut-off is the point at which the phonon wavelength is comparable to the spatial FWHM of the carrier wavefunction. As a result, the cut-off frequency depends strongly on the height of the QD and phonon sideband measurements may be used to probe the confinement potential. A plot of \(J(\omega)\) derived from fitting data measured in Ref. [21] is shown in Fig. 3. The form of \(J(\omega)\) modifies the detuning dependence of the DVC and also weakens the exciton-phonon interaction for very strong driving where the effective Rabi frequency \(\Lambda\) exceeds the cut-off frequency (although this regime has not been reached in present experiments).

The processes of interest for DVC both rely on phonon emission rather than phonon absorption, and are there-
fore highly sensitive to temperature. Increasing the bath temperature increases the phonon occupation leading to a higher probability of phonon emission and thus faster relaxation. However, the probability of phonon absorption generally increases more than that of emission. As illustrated by the red arrows in Fig. 2(a), phonon absorption \( (\Gamma_{\text{ph}}^-) \) causes the opposite population transfer between the dressed states to phonon emission \( (\Gamma_{\text{ph}}^+) \) and results in a lower final occupation of \( |\beta\rangle \). As such, DVC processes are generally optimised by low bath temperatures in order to maximise the difference between phonon emission and absorption rates. The influence of temperature on phonon-assisted excitation is studied both experimentally and theoretically in Section.

It is only by combining all of these influences that the full spectral and power dependence of the DVC emerges. Analytical approximations do not accurately predict these dependencies; hence numerical methods such as path integral calculations \[23, 24\] or the master equation formalism \[14\] are employed. The result of such calculations is a broad (few meV) spectral sideband feature that appears at high driving strengths and persists until the Rabi splitting exceeds the cut-off energy for phonon coupling.

**EXPERIMENTAL METHODS**

Figure 4 gives a schematic diagram of the apparatus used in the experiments. A mode-locked Ti:sapphire laser with 76 MHz pulse repetition rate is passed through two pulse shapers to obtain independently tunable pulses from within the \( \sim 10 \text{ meV} \) bandwidth of the \( \sim 100 \text{ fs} \) pulses \[35\]. The spectral FWHM is selected to be either 0.2 or 0.42 meV corresponding to a Fourier-transform-limited pulse duration of 16.8 or 8 ps. Both pulses pass through wave-plates to permit independent control of their polarisations, and one of them follows a variable path length controlled by a delay stage to enable pump-probe experiments with precise relative time control.

The pulses are incident on the sample in a variable temperature liquid He cryostat with piezo actuators for precise positioning of the sample relative to the focussed beams. The sample consists of a layer of InGaAs QDs embedded in a Schottky diode. The exciton created by the laser pulses can be measured by detecting the photocurrent (PC) from the QD when a reverse bias is applied to the diode \[11\]. Further details of the experimental methods may be found in ref. \[36\].

**FIG. 4.** Experimental arrangement. A femtosecond pulsed laser beam is split into two paths. The wavelength, FWHM, arrival time and polarization of the two pulses are controlled independently by the pulse shapers, delay stage and wave plates (WP). Finally the two beams are combined together and sent to the sample which is kept in a variable temperature liquid-He cryostat. The exciton created by the laser pulse is measured by detecting the photocurrent generated from the QD.

**COHERENCE AND SPIN PERSISTENCE OF DYNAMIC VIBRONIC COUPLING**

In ref. \[24\], Quilter et al. demonstrated the DVC by creating a QD exciton using a slightly blue-shifted laser pulse via emitting an LA phonon. Since it involves emission of a phonon, in theory the DVC should be incoherent, namely the phase of the exciton is random relative to that of the blue-shifted laser pulse. The exciton coherence time in this case is limited by the phonon relaxation time (few ps \[37\]). To demonstrate the incoherent nature of the DVC, we performed Ramsey-like interference measurements \[38–40\] using an unstabilized interferometer.

The QD is pumped with a blue-detuned pulse with \( \hbar \Delta = 1 \text{ meV} \) at \( t = 0 \). The pulse area (8.4\( \pi \)) is chosen to give an exciton population of \( C_X = 0.5 \). (Note that this is slightly higher than in ref. \[24\] due to different laser detunings.) Once the QD has been excited, the state is then probed by a \( \pi/2 \) pulse resonant with the exciton at \( t = \tau_{\text{delay}} \). The relative phase between the pump and probe is proportional to \( \tau_{\text{delay}} \). If the system is coherent, the probe pulse drives the QD to either \( C_X = 0 \) or \( 1 \) depending on the phase of the exciton with respect to the probe. The degree of coherence can then be determined by measuring the visibility of the Ramsey fringes as a function of \( \tau_{\text{delay}} \). In our experiment, the relative phase between the pump and probe is unstabilized, and so the final state fluctuates randomly within the visibility envelope.

Figure 5 shows the time-integrated PC signal versus \( \tau_{\text{delay}} \). The red squares show the results for phonon-sideband pumping with \( \hbar \Delta = 1 \text{ meV} \). A flat line is observed, with no Ramsey-like fringes, indicating that the phase of the exciton is random relative to the probe pulse.
COHERENCE AND SPIN PERSISTENCE OF DYNAMIC VIBRONIC COUPLING

The slow increase of the PC signal in time is most probably related to the decay of the exciton population created by the pump via electron/hole tunnelling before the arrival of the probe. By contrast, the black dots show the results measured when the pump pulse is tuned to resonance ($\Delta = 0$) and the pulse area is set to $\pi/2$. In this fully resonant situation, Ramsey-like fringes are observed provided $\tau_{\text{delay}}$ is shorter than or comparable to the exciton coherence time. The exciton coherence time of $\sim 40$ ps can be estimated from the dashed line envelope and is limited by the electron tunnelling rate [40]. The incoherent nature of the DVC is clearly demonstrated by the absence of the fluctuations in the PC signal in the red data compared with that in the black data. Our result is consistent with that reported by Weiler et al. [19] showing that the coherence properties of the emitted photon created via DVC is much worse than resonant CW excitation. However, Boumouar et al. [25] demonstrate that the photons generated by quasi-resonantly pumping the biexciton state using pulsed excitation show similar coherence properties as measured in the resonant two-photon scheme. The relation of the coherence properties of the emitted photon and exciton prepared via DVC is still unclear and require further investigations.

It should be noticed that the fact that the DVC process is incoherent does not imply that the exciton spin is random. The results published in ref. [23] clearly shows that the spin of the exciton is the same as that of the photon in the blue-shifted laser pulse. In theory the spin of the emitted photon in the LAPSE process should also be the same as that of the laser photon, since the phonon-assisted excitation and deexcitation are two opposite processes in the dressed state picture [see Fig. 2(a)].

Directly demonstrating the spin-preserving nature of the LAPSE process is beyond the scope of this paper, but it has been observed that the LAPSE process only occurs when the spin of the exciton and the LAPSE pulse are the same. This spin selectivity of the LAPSE process can be demonstrated by preparing a spin-up exciton using a circularly-polarized $\pi$ pulse and then trying to deexcite this exciton using a (a) co- or (b) cross-polarized red-shifted laser pulse. To isolate the PC signal of the QD under study from other QDs in the same sample, a reference spectrum measured with only a red-shifted laser pulse is subtracted. The pulse area $\Theta$ of the red-shifted laser pulse is (black) $\pi$ or (red) $7.35\pi$. The delay time between the $\pi$ pulse and the red-shifted pulse is 17 ps. XX: biexciton.

![Figure 5](image1.png)

**FIG. 5.** Comparison of Ramsey-like fringe data on the exciton for pumping in the phonon sideband (red) and at the exciton (black) at $T = 4.2$ K. In both cases the second $\pi/2$-pulse is resonant with the exciton. The pulse area $\Theta$ of the $+1$ meV detuned pump pulse is set to be $8.4\pi$ to generate $C_X = 0.5$. The dashed line is a guide to the eye.

![Figure 6](image2.png)

**FIG. 6.** Demonstration of the spin selectivity of the LAPSE process. Differential photocurrent spectra $\Delta PC$ were obtained by preparing an exciton in the QD using a circularly-polarized $\pi$ pulse and then measuring the PC as a function of a (a) co- or (b) cross-polarized red-shifted laser pulse. To isolate the PC signal of the QD under study from other QDs in the same sample, a reference spectrum measured with only a red-shifted laser pulse is subtracted. The pulse area $\Theta$ of the red-shifted laser pulse is (black) $\pi$ or (red) $7.35\pi$. The delay time between the $\pi$ pulse and the red-shifted pulse is 17 ps. XX: biexciton.
the co-polarized spectrum [see red lines in Fig. 6] ambiguously proves the spin selectivity of the LAPSE process. Based on this observation and the fact that the spin is preserved while the emission of the LA phonon as demonstrated in ref. [24], we conclude that DVC is incoherent, but spin-preserving.

**TEMPERATURE DEPENDENCE OF DYNAMIC VIBRONIC COUPLING**

![Graph](image)

**FIG. 7.** (a) Exciton population vs. laser detuning ($\Delta$) at 15 K (red triangles) compared to 4.2 K (black circles) for excitation with pulse area $\Theta = 4.4\pi$. The dashed line indicates a detuning of $\hbar \Delta = +0.83$ meV. (b) Calculated variation of the exciton population versus pulse area ($\Theta$) for $T = 4.2$ K (black), $T = 15$ K (red) and $T = 25$ K (blue) for positive detuning of $\hbar \Delta = +0.83$ meV. The dashed line shows $\Theta = 4.4\pi$.

In previous studies DVC processes have mainly been studied at $T = 4.2$ K [14, 24, 25], the temperature of liquid-He. It is interesting however to consider the influence of temperature on the mechanism. Increasing the temperature increases the strength of the exciton-phonon interaction, observed for instance by stronger damping and frequency renormalization of excitonic Rabi rotations [12]. As discussed in section [11] this increases the rate of phonon relaxation between the dressed states, and, for relatively weak excitation, increases the resulting exciton population as more population relaxes during the pump pulse. This is shown by the higher population observed at positive detuning for $T = 15$ K (red triangles) compared to $T = 4.2$ K (black circles) in Fig. 7(a) and also by the steeper initial gradient for higher temperatures observed in the calculations of Fig. 7(b).

However, this does not illustrate the complete picture. At higher bath temperatures the increased number of phonons increases the probability of the competing process at positive detuning, namely the annihilation of an exciton by absorption of a phonon (see Fig. 2(c)). This is shown in our measurements by the reduced detuning asymmetry of the phonon sidebands at $T = 15$ K in Fig. 7(a). The origin of the negative detuning signal is the creation of exciton population by phonon absorption, illustrating the increased probability of the competing absorption processes discussed in section [11]. As such, the reduced asymmetry in the phonon sidebands can be considered equivalent to the reduced difference between the probability of phonon emission and absorption.

The consequence of this competition is that for driving that is sufficiently strong to allow complete relaxation (i.e. in the plateau region of Fig. 7(b)), the final occupancy of the exciton state is in fact lower at higher temperatures owing to the thermalisation of the dressed states. This is illustrated by the reduced maximum exciton population attained at higher bath temperatures in Fig. 7(b). In summary, a higher bath temperature will lead to a faster rise with pulse area, but a lower maximum population transfer by DVC. As such, increasing the bath temperature may provide a means to enhance DVC processes when the driving is relatively weak as in Fig. 7(a).

**PICOSECOND TIMESCALE EXCITATION AND DE-EXCITATION**

The three first experimental observations of DVC in quantum dots [14, 24, 25] focused on the LA-phonon-assisted excitation process [see Fig. 2(a), (b)]. Recently Liu. et al. have demonstrated the reverse process - LA-phonon-assisted de-excitation [see Fig. 2] - by observing the reduction of the exciton population of an excited QD by a red-shifted laser pulse [26]. A combination of the LA-phonon-assisted excitation and de-excitation processes allows us to create and destroy an exciton within picosecond time scales, enabling ultrafast incoherent optical switching with a single QD much faster than the exciton radiative lifetime ($\sim$ ns [24, 11]).

To demonstrate this, a two-color pump-probe experiment is performed with a blue-shifted (excitation) and
red-shifted (de-excitation) pulse with the same pulse area. The pulses are equally detuned ($\hbar \Delta = \pm 0.7$ meV) from the exciton. At negative delay times, the excitation pulse creates a certain exciton population corresponding to the background PC level. The de-excitation pulse arrives before excitons are created, and therefore does nothing. When the two pulses overlap, the PC signal decreases to a minimum within $20$ ps, as shown in Fig. 8. This occurs due to erasure of the exciton created by the excitation pulse by the de-excitation pulse. The PC signal then slowly recovers due to the electron tunnelling out from the QD before the de-excitation pulse arrives. The increase of the PC signal at $\sim -10$ ps is most probably related to the phonon-assisted excitation of biexciton by a very small portion of imperfectly circularly polarized light in the excitation/deexcitation pulses. Fig. 8(b) shows the calculated exciton population with similar parameters as Fig. 8(a). The phase relation between the two pulses has been assumed to be random and we show the average of multiple repetitions of the simulation. The numerically complete path-integral method takes into account the electron tunnelling in the PC experiment by incorporating it as an Lindblad-type relaxation term [26]. Before the arrival of the pulses the QD is assumed to be in the ground-state and the phonon modes follow a thermal distribution. This model well reproduces the de-excitation of the QD and the subsequent recovery of the photo current signal. The oscillation of the exciton population at $\sim 5$ ps is not clearly visible in the measured data in Fig. 8(a) due to the limited time resolution of the experiment.

The $\sim 20$ ps excitation/de-excitation time in our experiment is determined by the laser pulse duration. The ultimate limit, is set by the phonon thermalisation time (few ps [37]). We note that in contrast to the optical switching scheme employing resonant coherent excitation [2], our scheme is robust against the fluctuation of laser power and detuning [28]. Furthermore, the incoherent nature of the phonon-assisted relaxation process determines that no phase locking is needed, in contrast to fully coherent excitation schemes.

CONCLUSIONS AND OUTLOOK

In summary, we have investigated the dynamic vibronic coupling in a single InGaAs QD. The DVC enables the population inversion and ultrfast depopulation of a QD by incoherent quasi-resonant excitation with the assistance of phonons [24, 26]. We prove that the DVC is an incoherent process by performing Ramsey-like interference measurements. The temperature dependence study shows that at high temperature the phonon-assisted absorption induced by both the blue- and red-shifted laser pulses are enhanced and the strong asymmetry of the phonon sidebands observed at low temperatures disappears. Furthermore we demonstrate that it is possible to create and destroy an exciton within a pulse-width limited ps time scale by combining a blue-detuned and a red-detuned pulses, opening the door to ultrafast incoherent optical switching with a single QD [12, 44]. The DVC may also be used to reduce the timing jitter of single/entangled photon sources [8, 45] or make tunable single QD lasers [46].

Compared with resonant coherent excitation, one factor which limits the potential applications of DVC is the high laser power required to achieve an efficient phonon-assisted excitation/deexcitation. This problem may be solved by incorporating the QD into a nanocavity with a small mode volume where the light matter interaction is strongly enhanced [8, 17]. When cavities are involved, the knowledge of phonon interactions in the strong QD-phonon coupling regime becomes an essential requirement for a full understanding of the behaviour of optically driven QDs. Various recent studies have shown new physical phenomena that arise from this coupling [15, 52].
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