Triplet-singlet relaxation in semiconductor single and double quantum dots

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We study the triplet-singlet relaxation in two-electron semiconductor quantum dots. Both single dots and vertically coupled double dots are discussed. In our work, the electron-electron Coulomb interaction, which plays an important role in the electronic structure, is included. The spin mixing is caused by spin-orbit coupling which is the key to the triplet-singlet relaxation. We show that the selection rule widely used in the literature is incorrect unless near the crossing/anticrossing point in single quantum dots. The triplet/singlet relaxation in double quantum dots can be markedly changed by varying barrier height, inter-dot distance, external magnetic field and dot size.

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I. INTRODUCTION

The application of semiconductor quantum dots (QDs) in generating spin-based qubits\textsuperscript{3} is one of the focuses in the field of spintronics.\textsuperscript{3} There are two types of qubits investigated extensively recently.\textsuperscript{2} One is based on the transition between single-electron Zeeman sublevels and the other is based on two-electron triplet-singlet (TS) states.\textsuperscript{1,2,4,5,6,7,8,9,10,11,12,13,14,15,16,17,18,19,20,21,22,23} Among these works, the decoherence time of the spin states, including both the spin dephasing time and spin relaxation time has attracted much attention as a thorough understanding of it is one of the prerequisites of the application. There are many works on spin relaxation reported, especially in single-electron QDs.\textsuperscript{2,4,5,6,7,8,9,10,11,12,13,14,15,16,17,18,19,20,21,22,23} Recently, the TS relaxation time of two-electron system has also been investigated.\textsuperscript{24,25,26,27} It was proposed that various mechanisms, such as the electron-phonon scattering together with the spin-orbit coupling,\textsuperscript{28,29} the hyperfine interaction,\textsuperscript{30,31} and the cotunneling effect, could induce TS relaxation.\textsuperscript{22} However, the mechanism involving electron-phonon scattering is usually treated as the key one because the nuclei-mediated relaxation\textsuperscript{32} and the cotunneling can be weakened via tuning external magnetic field and tunneling rates respectively.\textsuperscript{24} Specifically, Climente et al. used exact diagonalization technique to calculate the two-electron spectrum structure and the phonon induced TS relaxation in parabolic single QDs.\textsuperscript{22} They demonstrated the crucial role of the excited states on spectrum structure and showed a slow decrease of the relaxation time away from the TS crossing in contrast to a sharp increase in the vicinity of the crossing point, when the magnetic field is increased from zero Tesla. This feature agrees qualitatively with the recent measurement.\textsuperscript{22,23} Furthermore, their results indicated that the spin-down triplet state coupled with the singlet ground state through the spin-orbit coupling has a much shorter lifetime compared to the other two triplet states. This was understood by the so called “selection rule” based on the perturbation using the lowest two single electron levels. Similar perturbative discussion was also given in Ref.\textsuperscript{25}. Meunier et al. obtained perturbative wave functions from the selection rule and treated the spin-orbit coupling coefficient as a fitting parameter.\textsuperscript{33} Using these functions, they fit their experiment data with electron-phonon-scattering-induced TS relaxation and obtained a particularly small spin-orbit coupling coefficient. They attributed the reduction of the coupling coefficient to the neglect of high excited states. Sasaki et al. pointed out that the selection rule was correct only in the vicinity of the TS crossing point,\textsuperscript{10} which seems to be more correct intuitively. According to the previous work by one of the authors and confirmed by Destefani and Ulloa,\textsuperscript{34} the spin-orbit coupling in quantum dots is very strong and a large number of basis functions are needed in order to achieve convergence even for the lowest few states. Therefore, whether the selection rule based on the lowest few levels remains unchanged when many upper levels are involved remains questionable to us. Therefore, in this work we will first reinvestigate the selection rule based on exact diagonalization method, jointly with perturbation method with many basis functions.

The investigation on TS relaxation in double QD architectures is very limited. Recently, Wang and Wu studied the single-electron spin relaxation in vertically coupled double QDs and showed that the spin relaxation can be efficiently manipulated electronically by the inter-dot barrier.\textsuperscript{20} This suggests that the two-electron TS relaxation should also be manipulated by tuning inter-dot barrier height. This is another issue we are going to explore in this work.

We organize the paper as following. In Sec. II we set up the model and lay out the formalism. Then in Sec. III we show our numerical results. We discuss the single dot case in Sec. III A. We first show the exact diagonalization results with sufficient basis functions. We then reexamine the selection rule by using more basis functions instead of the lowest two, both perturbatively and exactly. We show the selection rule widely used in the literature is not correct except near the TS crossing/anticrossing points. In Sec. III B we show the results of double QDs. We summarize in Sec. IV.
II. MODEL AND FORMALISM

We start our investigation from a vertically-coupled double QD: Electrons are confined by a parabolic potential \( V_c(x, y) = \frac{1}{2} m^* \omega_0^2 (x^2 + y^2) \) (corresponding to the effective dot diameter \( d_0 = \sqrt{\hbar \pi / m^* \omega_0} \)) along the \( x \) - \( y \) plane \cite{33, 34, 35} with \( m^* \) representing the effective mass. Along the \( z \)-axis, a strong confinement is given by

\[
V_z(z) = \begin{cases} 
V_0, & \left| z \right| \leq \frac{a}{2} \\
0, & \frac{a}{2} < \left| z \right| < \frac{a}{2} + d \\
\infty, & \text{otherwise,}
\end{cases}
\]

with \( V_0 \), the inter-dot barrier \cite{35}. By taking \( a = 0 \), one comes to the single dot configuration. The single-electron Hamiltonian with magnetic field along the growth direction \( (\hat{z}) \) is given by

\[
H_e = \frac{\mathbf{P}^2}{2m^*} + V(\mathbf{r}) + H_{so}(\mathbf{P}) + H_Z ,
\]

in which \( V(\mathbf{r}) = V_z(z) + V_c(x, y) \) and \( \mathbf{P} = -i \hbar \nabla + e/\epsilon \mathbf{A} \) with \( \mathbf{A} = (B/2)(-y, x, 0) \). \( H_{so} \) represents the spin-orbit coupling which is the key to the spin flip. In this work, we only consider the Dresselhaus spin-orbit coupling \cite{36} as the Rashba coupling \cite{33} is comparably small in GaAs QDs. \( H_Z \) is the generalized Laguerre polynomial. In these equations, \( n, l, k \) are the radial quantum numbers of the first and the second electron, \( \ell = 0, \pm 1, \pm 2, \ldots \) is the azimuthal angular momentum quantum number. By solving the \( z \)-component of Eq. \ref{eq:1}, we obtain the lowest two electronic states along the \( z \)-axis as following:

\[
\phi_z = \begin{cases} 
C_0^0 \sin[k(z - \frac{d}{2} - d)], & \frac{d}{2} < z < \frac{\ar}{2} + d, \\
C_0^1 \cosh(\beta z), & \left| z \right| \leq \frac{a}{2}, \\
C_0^1 \sin[k(-z + \frac{d}{2} + d)], & -\frac{d}{2} - d < z < -\frac{\ar}{2},
\end{cases}
\]

\begin{equation}
\tag{7}
\end{equation}

and

\[
\phi_z = \begin{cases} 
C_1^1 \sin[k(z - \frac{d}{2} - d)], & \frac{d}{2} < z < \frac{\ar}{2} + d, \\
C_2^1 \sinh(\beta z), & \left| z \right| \leq \frac{a}{2}, \\
C_1^1 \sin[k(z + \frac{d}{2} + d)], & -\frac{d}{2} - d < z < -\frac{\ar}{2},
\end{cases}
\]

\begin{equation}
\tag{8}
\end{equation}

in which \( k^2 = 2m^*E_z/\hbar^2 \) and \( \beta^2 = 2m^*(V_0 - E_z)/\hbar^2 \) with \( E_z \) denoting the energy along this direction. We use the superscripts “0” and “1” to denote the even and odd parity respectively. The total spatial wave function is then denoted by \( |nl_nz⟩ \), with \( n_z = 0 \) and \( l_z = 1 \) in this work to distinguish the above even and odd states along the \( z \)-axis. Due to the strong confinement along the \( z \)-axis, levels higher than \( n_z = 1 \) are neglected. It is noted that when we refer to the single QDs, we only keep the lowest state (the even one) due to the small well width.

For two-electron system, the total Hamiltonian is written as

\[
H_{tot} = (H_1^1 + H_2^2 + H_C) + H_{ep}^2 + H_{sp}^2 + H_p .
\]

\begin{equation}
\tag{9}
\end{equation}

In this equation, the third term \( H_C = \frac{\alpha^2}{4\epsilon_0\epsilon \hbar |r_1 - r_2|} \) describes the Coulomb interaction between the two electrons with \( \kappa \) representing the static dielectric constant. \( H_{ep} = \Sigma_{\lambda\lambda'}b_{\lambda}a_{\lambda'}^\dagger \) represents the phonon Hamiltonian, and \( H_{sp} = \Sigma_{\lambda\lambda'}M_{\lambda\lambda'}(a_{\lambda}^\dagger + a_{\lambda}) \exp(iq \cdot r) \) is the Hamiltonian of the electron-phonon interaction. The superscripts “1” and “2” label the two electrons.

We construct two-electron basis functions from the single electron wave functions. To see the physics clearly, we construct our two-electron basis functions in either singlet or triplet forms. Taking two single-electron spatial wave functions \( |n_1l_1n_1z⟩ \) and \( |n_2l_2n_2z⟩ \) (denoted as \( |N_1⟩ \) and \( |N_2⟩ \) for short) as an example, the singlet functions can be constructed by

\[
|S⟩ = (|↑↓⟩ - |↓↑⟩)
\]

\[
\otimes \left( \frac{1}{\sqrt{2}}(|N_1N_2⟩ + |N_2N_1⟩), \frac{1}{2}(|N_1N_2⟩ + |N_2N_1⟩) \right) \tag{10}
\]

and the triplet functions for \( N_1 \neq N_2 \) by

\[
|T_+⟩ = \frac{1}{\sqrt{2}}(|N_1N_2⟩ - |N_2N_1⟩) \otimes |↑↓⟩ \tag{11}
\]

\[
|T₀⟩ = \frac{1}{2}(|N_1N_2⟩ - |N_2N_1⟩) \otimes (|↑↓⟩ + |↓↑⟩) \tag{12}
\]

\[
|T−⟩ = \frac{1}{\sqrt{2}}(|N_1N_2⟩ - |N_2N_1⟩) \otimes |↓↓⟩ \tag{13}
\]

Here, \( N \) and \( N' \), in the ket \( |NN'⟩ \), represent the spatial quantum numbers of the first and the second electrons respectively. We define the total angular momentum \( L = l_1 + l_2 \) and denote the total spin \( (S, \mathbf{S}) \) with \( S_z \) representing the \( z \)-component of the total spin \( \mathbf{S} \).

Then, we calculate the matrix elements of the Coulomb interaction and the spin-orbit coupling \cite{31} under these ba-
sis functions. The Coulomb matrix elements can be expressed in the form

$$\langle N_1 N_2 | H_C | N'_1 N'_2 \rangle = \frac{e^2}{4\pi^2 \epsilon_0 \kappa} \delta_{N_1+N_2} \delta_{N'_1+N'_2} \times Q(N_1, N_2, N'_1, N'_2),$$

in which $Q$ is given in detail in Appendix A. Thus we obtain the two-electron Hamiltonian. By diagonalizing the two-electron Hamiltonian, one obtains all the energy levels and eigenfunctions. We identify a state as singlet/triplet if its amplitude of singlet/triplet components is larger than 50%. We rewrite the spin-orbit coupling Hamiltonian [Eq. (3)] using the ladder operators as

$$H_{so} = \gamma_c (P^+ S^+ + P^- S^-),$$

with the coupling coefficient $\gamma_c = \frac{\pi}{\hbar^2}(P_z^2)$. Then it is noted that $P^\pm$ and $S^\pm$ change $L$ and $S_z$ by one unit, respectively. It suggests that a state with $(L, S_z)$ can only be coupled with the states with $(L+1, S_z+1)$ and $(L-1, S_z-1)$.

Treating $|i\rangle$ and $|f\rangle$ as the initial and final states, we can calculate the phonon-induced relaxation rate from the Fermi Golden Rule

$$\Gamma_{i\rightarrow f} = \frac{2\pi}{\hbar} \sum_{\mathbf{q}\lambda} |M_{Q\lambda}|^2 \langle f | (\mathbf{q}, \mathbf{r}_1, \mathbf{r}_2) = e^{i \mathbf{q} \cdot \mathbf{r}_1} + e^{i \mathbf{q} \cdot \mathbf{r}_2} \rangle = \langle \hat{n}_{\lambda} \delta (\epsilon_f - \epsilon_i - \hbar \omega_{\lambda}) \rangle,$$

in which $\chi(q, \mathbf{r}_1, \mathbf{r}_2) = e^{i \mathbf{q} \cdot \mathbf{r}_1} + e^{i \mathbf{q} \cdot \mathbf{r}_2}$ comes from the total electron-phonon interaction Hamiltonian $H_{ep} = H_{ep}^1 + H_{ep}^2$. Here, $\hat{n}_{\lambda}$ represents the Bose distribution of phonon with mode $\lambda$ and momentum $\mathbf{q}$. In our calculation, the temperature is fixed at 0 K. Therefore only the phonon emission process occurs.

### III. NUMERICAL RESULTS

In the numerical calculation, we include the electron-acoustic phonon scattering due to the deformation potential with $|M_{Q\alpha}|^2 = \hbar \kappa^2 \epsilon_2 q^2 / 2 D v_{st}^2$, and due to the piezoelectric field with $|M_{Q\alpha}|^2 = (32 \hbar \kappa^2 \epsilon_2 \epsilon_1 / \kappa^2 D v_{st}^2) [\langle 3q_2 q_0 q_2 \rangle / q^2]$ for the longitudinal modes and $\sum_{j=1,2} |M_{Q\alpha j}|^2 = (32 \hbar \kappa^2 \epsilon_2 \epsilon_1 / \kappa^2 D v_{st}^2) [\langle q_2^2 + q_0^2 + q_2^2 - 3 \delta q_2 q_0 q_2 \rangle / q^2]$ for the transverse modes. Here, $\kappa = 7$ eV stands for the acoustic deformation potential; $D = 5.3 \times 10^3$ kg/m$^3$ is GaAs volume density; $\epsilon_1 = 1.41 \times 10^9$ V/m denotes the piezoelectric constant and the static dielectric constant $\kappa$ is 12.9; $v_{st} = 5.29 \times 10^3$ m/s corresponds to the longitudinal sound velocity and $v_{st} = 2.48 \times 10^3$ m/s corresponds to the transverse one.

In our calculation, $g$ factor is $-0.44 \pm 0.17$ and the Dresselhaus coefficient $\gamma$ is $21.5 \ \text{A}^2 \cdot \text{eV}$. The typical electron effective mass $m^*$ in GaAs is $0.067 m_0$ with $m_0$ being the free electron mass.

![FIG. 1: (Color online) (a) The lowest four energy levels vs. magnetic field $B$ in single QD. The TS anticrossing point between $T_-$ and $S$ is shown and the range near this point is enlarged in the inset. (b) $\tau^{-1}$ of the three transition channels vs. the magnetic field. In the calculation, $d = 5 \text{ nm}$ and $d_0 = 30 \text{ nm}$.](image)

**A. Single dot**

We first set $a = 0$ to investigate the single dot case by exact diagonalization method with the lowest 800 singlet and 2220 triplet basis functions. Under the basis, the energy levels and the TS relaxation rates are well converged. The magnetic field dependence of the first four levels and that of the TS relaxation rates are plotted in Fig. 1. In the calculation, we take the well width $2d = 10 \text{ nm}$ and the effective diameter $d_0 = 30 \text{ nm}$. From Fig. 1(a), one notices that the ground state is a singlet denoted as "S", in a wide range of the magnetic field (from 0 T to 2.6 T approximately). In this region, the first three excited states are triplet states, labeled as $|T_+\rangle$ (spin-up), $|T_0\rangle$ (spin-zero) and $|T_-\rangle$ (spin-down), and the energy of $|T_-\rangle$ is the highest one among the three because of the Zeeman effect. When the magnetic field increases from 2.6 T, one further observes a TS crossing between the singlet and the two triplets $(|T_+\rangle$ and $|T_0\rangle$). Moreover, a TS anticrossing point (with a small energy gap shown in Fig. 1(a)) also exists between the singlet and $|T_-\rangle$ triplet state due to the Dresselhaus spin-orbit coupling. From the calculation, we notice that the major components of $|S\rangle$, $|T_\pm\rangle$ and $|T_0\rangle$ are $|S\rangle$, $|T_\pm\rangle$ and $|T_0\rangle$, which are the lowest singlet and triplet basis fun-
tions. Specifically, using the lowest two single-particle wave functions, $|nl_n\rangle$ with $n = n_z = 0$, $l = 0$ and $-1$, one can construct $|S^1\rangle$ with $(00)$ and $(00)$, and $|T^1_\perp\rangle$ and $|T^1_\parallel\rangle$ with $(00)$ and $(0 - 10)$ according to Eqs. (10)-(13). Therefore, the quantum numbers $(L, S_z)$ of $|S^1\rangle$, $|T^1_\perp\rangle$, $|T^1_0\rangle$ and $|T^1_\parallel\rangle$ are different, i.e., $(0, 0), (-1, 1), (-1, 0)$ and $(-1, -1)$ respectively.

From Fig. 1(b) one observes that the TS relaxation rates increase slowly with the magnetic field away from the crossing/anticrossing points, but decrease dramatically in the vicinity of the crossing/anticrossing points, in agreement with the measurement qualitatively. The relaxation rate reaches maximum where the wave length of the emissive phonon is comparable with the dot size. In our calculation, the TS splitting, i.e., the energy between the triplet and the singlet, $\Delta_{TS} \sim 0.2$ meV. The corresponding half-wavelength of the transverse phonon is therefore about 30 nm as the dot diameter $d_0$. This feature was interpreted as the competing effects of the magnetic field on the electron-phonon coupling and the spin-orbit coupling. Actually, the strength of the spin-orbit coupling is proportional to $\alpha$ [see Eq. (6)] which increases with the magnetic field whereas the electron-phonon scattering becomes rather weak when the emissive phonon momentum decreases.

Surprisingly, our results are very different from those shown in the previous work, where the transition rate of $|T_-\rangle$ is much larger than that of the other two triplet states $|T_+\rangle$ and $|T_0\rangle$. In that work, the authors interpreted their results by the selection rule based on the perturbation method including the lowest four basis functions, i.e., $|S^1\rangle$, $|T^1_\perp\rangle$, $|T^1_0\rangle$ and $|T^1_\parallel\rangle$. Under that basis, only $|T_-\rangle$ is coupled with $|S\rangle$ through the Dressehaus spin-orbit coupling according to Eq. (15). So only the transition from $|T_-\rangle$ to $|S\rangle$ can occur. Thus they concluded that the transition rate from $|T_-\rangle$ to $|S\rangle$ is much larger than those of the other channels even though much more (instead of four) basis functions are included. In fact, this selection rule is widely used in the literature. However, as one needs many basis functions to achieve convergence even in the single-electron QD system, whether the selection rule from the lowest four basis functions is robust against the inclusion of higher basis functions remains an open question. Here we reexamine the selection rule with more basis functions. Assuming the perturbation based on the lowest four states $|S^1\rangle$, $|T^1_\perp\rangle$ and $|T^1_0\rangle$ is adequate to describe the real physics, we expect the selection rule should always be valid when more basis functions are included. Specifically, we now use four single-electron functions $|00\rangle$, $|0 - 10\rangle$, $|010\rangle$ and $|0 - 20\rangle$ to construct the two-electron basis functions. Keeping only the index of $l$ from $|nl_n\rangle$ since the other two are fixed, the six lowest singlet states are constructed by $|00\rangle |00\rangle$, $|00\rangle |01\rangle$, $|00\rangle |-1\rangle$, $|01\rangle |00\rangle$, $|01\rangle |-1\rangle$, $|-1\rangle |00\rangle$ separately and the three lowest triplet states are constructed by $|00\rangle |-1\rangle$ in the way of Eqs. (10)-(13). We denote these nine basis functions as $|S^1\rangle$, $|S^2\rangle$, $|S^3\rangle$, $|S^4\rangle$, $|S^5\rangle$, $|T^1_+\rangle$, $|T^1_0\rangle$ and $|T^1_\parallel\rangle$ in sequence, and the quantum numbers $(L, S_z)$ are $(0, 0)$, $(-1, 0)$, $(1, 0)$, $(-2, 0)$, $(0, 0)$, $(-1, 1)$, $(-1, 0)$ and $(-1, -1)$ respectively. Therefore, only the singlet states $|S^1\rangle$ and $|S^5\rangle$ can mix with $|T^1_+\rangle$; $|S^4\rangle$ and $|S^5\rangle$ can mix with $|T^1_0\rangle$, according to Eq. (15) under these basis functions. No mixing occurs to the state $|T^1_\parallel\rangle$.

As the Coulomb interaction is too strong to treat perturbatively, we first diagonalize the Hamiltonian with the Coulomb interaction included to obtain a new set of basis functions, i.e., $|S^1\rangle = a_1|S^1\rangle + b_1|S^6\rangle$, $|S^2\rangle = a_2|S^1\rangle + b_2|S^6\rangle$, $|S^3\rangle = a_3|S^4\rangle + b_3|S^5\rangle$, $|S^4\rangle = a_4|S^1\rangle + b_4|S^6\rangle$, $|S^5\rangle = |S^4\rangle$, $|S^6\rangle = |S^1\rangle$, $|T^1_+\rangle = |T^1_0\rangle$, $|T^1_0\rangle = |T^1_\parallel\rangle$, and $|T^1_\parallel\rangle = |T^1_\parallel\rangle$. Here $a_i$ and $b_i$ are obtained from the numerical diagonalization. The corresponding eigenvalues are $E_1$ to $E_6$, $E_+$. $E_0$ and $E_-$. Respectively. Then we treat the spin-orbit coupling as perturbation under the new basis functions. The lowest four states then read

\[
|T^1_+\rangle = |T^1_+\rangle + \theta_+^1|S^4\rangle + \theta_+^2|S^5\rangle, \tag{17}
\]
\[
|T^1_0\rangle = |T^1_0\rangle, \tag{18}
\]
\[
|T^1_-\rangle = |T^1_0\rangle + \theta_-^1|S^1\rangle + \theta_-^2|S^6\rangle, \tag{19}
\]
\[
|S\rangle = \theta_+^1|S^4\rangle + \theta_+^2|S^6\rangle + \theta_-^2|T^1_\perp\rangle. \tag{20}
\]

with $\theta_+^1 = \sum_{i=1.4} \frac{(a^2_i - a^2_i)\Delta_i\Theta_i^2}{E_i - E_0}$, $\theta_+^2 = \Theta_1^2$, $\theta_-^2 = \sum_{i=1.2} \frac{(a^2_i + b^2_i)\Theta_i^2}{E_i - E_0}$ and $\theta_0^2 = \frac{a_1^2 + b_1^2}{E_+ - E_0}$. Here $\Theta_1^2 = a_1(b_1)$, $A = -i\gamma^*\alpha(1 - eB/2ha^2)$ and $B = -\frac{\gamma^*\alpha(1 + eB/2ha^2)}{2}$ with $\gamma^*$ being $\gamma/2d^2$.

Obviously, the transitions from both $|T^1_+\rangle$ and $|T^1_-\rangle$ to $|S\rangle$ can occur according to Eqs. (17)-(20). The matrix

\[
\begin{align*}
\text{FIG. 2:} \text{ (Color online)} \text{ Comparison of the TS relaxation rates of } T_+ \text{ and } T_- \text{ in single QD calculated from the perturbation method and the exact diagonalization method limited within nine basis functions } (|S^1\rangle \text{ to } |S^6\rangle, |T^1_\perp\rangle \text{ and } |T^1_0\rangle). \text{ Dashed curve and } \bullet: 1/\tau_{T+\rightarrow S} \text{ from the exact diagonalization and perturbation methods respectively; Solid curve and } \circ: 1/\tau_{T-\rightarrow S} \text{ from the exact diagonalization and perturbation methods.}
\end{align*}
\]
elements $\langle f|\chi|i\rangle^2$ in Eq. (16) now read

$$
|\langle S|\chi|T_+\rangle|^2 = |\theta_1^{s*}\theta_2^s(S^1|S^2) + \theta_1^{s*}\theta_2^s(S^6|S^6)|^2
= (xt)^2|\xi|^2I^2(q_z),
$$

with $x = \hbar^2/4\alpha^2$, $t = e^{-x}$. $I(q_z) = \pi^2\sin(dq_z)/\{dq_z[\pi^2 - (dq_z)^2]\}$, $C_1 = \theta_1^s + \theta_1^{s*}\theta_1^1 + \theta_2^s\theta_2^1$, $C_2 = \theta_1^{s*} + 2\theta_2^{s*}\theta_2^1$ and $\xi = \theta_1^{s*}\theta_2^1 + \sqrt{2}\theta_2^{s*}\theta_1^1$. We calculate the relaxation rates of these two channels and plot the results in Fig. 2. One notices that the two sets of dots (blue ▲ for $|T_+\rangle$ and red solid curve for $|T_-\rangle$) are close to each other and even show a crossing. In other words, the selection rule is violated. We also present the exact diagonalization results under the same basis functions $|S^1\rangle$-$|S^6\rangle$, $|T_0\rangle$ and $|T_+\rangle$ in Fig. 2 (blue dashed curve for $|T_+\rangle$ and red solid curve for $|T_-\rangle$). It is seen that the diagonalization results almost exactly match the perturbation results. This match further confirms that both our exact diagonalization and the perturbation calculations are correct. Compare Fig. 2 with Fig. 1(b), it is obvious that the high excited levels manifest themselves markedly in the relaxation rates.

From our calculation, we notice that the coefficients in Eqs. (17) and (19) are comparable. This is because that the denominators $E_+ - E_i$ in $\theta_{1,2}^{S}$ are close to $E_+ - E_3,4$ in $\theta_{1,2}^{T}$. This explains the reason why the curve of $|T_+\rangle$ is close to that of $|T_+\rangle$ in Fig. 2.

However, it is noted that the selection rule works well in the vicinity of the crossing/anticrossing points both in Figs. 1(b) and 2. This can be understood from Eqs. (17) and (19). Near the TS crossing point where $E_+ \sim E_1$, the energy splitting $E_+ - E_3,4$ is finite. Therefore $\theta_{1,2}^{S}$ only changes slightly compared with the region away from the TS crossing. Similar is true for the coefficients of $|T_0\rangle$. In contrast, $\theta_{1,2}^{T}$ is very large when $E_- \sim E_1$. Therefore the transition rate from $|T_-\rangle$ would be much larger than those from $|T_+\rangle$ and $|T_0\rangle$, i.e., the selection rule is valid in the vicinity of the TS crossing/anticrossing point. Moreover, the effect of the Zeeman splitting also makes the transition rate of $|T_-\rangle$ larger than those of $|T_+\rangle$ and $|T_0\rangle$ because of the larger phonon momentum $q$. Specifically, the energy splitting between $|T_-\rangle$ and $|S\rangle$ is about 0.18 meV at $B = 2.5$ T in Fig. 1(a), which is much larger than that between $|T_+\rangle$ ($|T_0\rangle$) and $|S\rangle$, i.e., $\sim 0.06$ meV (0.12 meV). As the transition rates are proportional to $q^n$ with $m > 0$ varying for different mechanisms, the rate of $|T_-\rangle$ is much larger than those of $|T_+\rangle$ and $|T_0\rangle$.

B. Double dot

Now we turn to study the TS relaxation rate in weakly coupled double QDs using a basis functions including 400 singlet and 1080 triplet states. In the calculation, $a = 8$ nm and $d = 7$ nm. In this part we still use $|T_\pm\rangle$ and $|T_0\rangle$ ($|S\rangle$) to denote eigenfunctions of the lowest three triplet states (lowest singlet state). To determine the contribution of the energy levels along the $z$-axis, we take the barrier height $V_0 = 0.25$ meV, the lowest one in our calculation, as an example. In this configuration the splitting between the first and the second levels along the $z$-axis is about 1 meV and that between the second and the third levels is much larger, about 0.2 eV. Compared with the lateral confinement (about 4 meV for $d_0 = 30$ nm), we only need to include the lowest two in our calculation.

We first investigate the TS relaxation rate as a function of the barrier height. In the calculation, $d_0 = 30$ nm and $B = 0.5$ T. As shown in Fig. 3, each transition rate first increases slowly until it reaches the maximum around $V_0 \sim 0.28$ eV where the TS splitting $\Delta_{TS} \sim 0.4$ meV corresponding to the wavelength of the emissive phonon being comparable with the dot size $d_0$. After that, the TS relaxation rate decreases rapidly with the barrier height. This would offer us a scheme to manipulate the TS relaxation in double QDs. Similar features (not shown here) are obtained when we increase the inter-dot distance. The dramatic decrease of the relaxation rate can be understood as following. When the barrier height becomes higher or the inter-dot distance becomes larger, the inter-dot coupling is weakened and the energy splitting between the lowest two levels along the $z$-axis becomes smaller. As a result, the splitting between $|T_\pm\rangle$ ($|T_0\rangle$) and $|S\rangle$ decreases too. This causes the decrease of the TS relaxation rate as discussed in the previous subsection.

![Fig. 3: (Color online) $\tau_{\text{TS}}^{-1}$ vs. barrier height in double QDs.](image)

In the calculation, $a = 8$ nm, $d = 7$ nm, $d_0 = 30$ nm and $B = 0.5$ T.

To have a look at the role of the magnetic field, we calculate the average relaxation rate $1/\tau = (1/\tau_{T_+\rightarrow S} + 1/\tau_{T_-\rightarrow S} + 1/\tau_{T_0\rightarrow S})/3$ as function of the barrier height at different magnetic field in Fig. 4(a), but with the dot size $d_0 = 30$ nm fixed. It is seen from the figure that higher magnetic field leads to relatively larger transition rate. It
is due to the enhanced spin-orbit coupling in strong magnetic field. The influence of the effective diameter of QDs with $V_0 = 0.35 \text{ V}$ and $B = 1 \text{ T}$ is also shown in Fig. 4(b). One finds the transition rates increase with the effective diameter $d_0$. The reason lies on the different symmetry properties of the singlet and triplet states. For the singlet state, the inter-electron distance decreases with the decrease of the dot size. The coulomb repulsion therefore lifts the corresponding energy levels. However, the energy lifts of the triplet states are smaller due to the antisymmetry property of the triplet states which prevents the electrons to be close to each other. Therefore, the TS splitting becomes smaller with the decrease of the dot size. This leads to the rapid decrease of the TS relaxation rates.

IV. SUMMARY

In summary, we have investigated the TS relaxation in single and double QDs. For the single dot case, we find that the average relaxation rate first slowly increases with magnetic field until it reaches the maximum where the wavelength of emissive phonon is comparable with the dot size. Then it drops sharply. This result qualitatively agrees with the recent measurement. Furthermore, our result shows the transition rates of the triplet $|T_+\rangle$ and $|T_0\rangle$ can be comparable with that of $|T_-\rangle$, which violates the selection rule in the literature. We show that the selection rule obtained from the lowest four basis functions does not hold in general cases where much more basis functions are needed to converge the triplet/singlet states. This is shown perturbatively by calculating the TS relaxation rates based on nine basis functions. Comparable transition rates of $|T_+\rangle$ and $|T_-\rangle$ are immediately obtained away from the TS crossing point. The perturbation results are in good agreement with the exact diagonalization results under the same basis functions. We also show that the selection rule works well in the vicinity of the TS crossing/anticrossing point due to the effects from the Zeeman splitting and the anticrossing. For the double QD case, we demonstrate that the TS relaxation rates vary more than two orders of magnitude by tuning the inter-dot barrier. This offers a feasible scheme to manipulate the TS relaxation in double QDs. The relaxation rates also sensitively depend on the dot size and magnetic field.

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APPENDIX A: Q IN COULOMB INTERACTION

Following Ref. 14, we obtain $Q$ in Eq. (14) as

$$Q(N_1, N_2, N'_1, N'_2) = \int_{-\infty}^{\infty} dk_z W_{N_1, N'_1}(k_z) W_{N_2, N'_2}(k_z) \frac{k^2}{2},$$

(A1)

where $P_{N,N'}$ and $W_{N,N'}$ come from the lateral and vertical parts of the matrix element $\langle n, l, n_z | \exp(i\mathbf{k} \cdot \mathbf{r}) | n', l', n'_z \rangle$, respectively. $P$ is given by

$$P_{N,N'}(k_z) = \sqrt{\frac{n!n'!}{(n+|l|)!(n'+|l'|)!}} \exp\left(-\frac{k_z^2}{4\alpha^2}\right) \times \sum_{i=0}^{n} \sum_{j=0}^{n'} C_{n,|l|}^{i} C_{n,|l'|}^{j} \tilde{n}! L_{n}^{|l'-l'|} \left(\frac{k_z^2}{4\alpha^2}\right) \times \left(\text{sgn}(l'-l) \frac{k_z^2}{2\alpha}\right)^{|l'-l'|},$$

(A2)

with $C_{n,|l|}^{i} = \frac{(-1)^i}{i!} \frac{(n+|l|)}{n!}$ and $\tilde{n} = i+j+(|l|+|l'|-|l'-l|)/2$. $\text{sgn}(x)$ represents the sign function. $W$ reads

$$W_{N,N'} = \langle n_z | \exp(i k_z z) | n'_z \rangle.$$  

(A3)
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33. Under the basis for Fig. 2, the TS crossing/anticrossing occurs near $B = 1.6$ T (not shown). Therefore, the selection rule already works at $B > 1.3$ T.