Controlled exchange interaction for quantum logic operations with spin qubits in coupled quantum dots

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Abstract

A two-electron system confined in two coupled semiconductor quantum dots is investigated as a candidate for performing quantum logic operations on spin qubits. We study different processes of swapping the electron spins by controlled switching on/off the exchange interaction. The resulting spin swap corresponds to an elementary operation in quantum information processing. We perform a direct time evolution simulations of the time-dependent Schrödinger equation. Our results show that – in order to obtain the full interchange of spins – the exchange interaction should change smoothly in time. The presence of jumps and spikes in the corresponding time characteristics leads to a considerable increase of the spin swap time. We propose several mechanisms to modify the exchange interaction by changing the confinement potential profile and discuss their advantages and disadvantages.

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I. INTRODUCTION

In quantum computation, one and two-qubit logic gates play a crucial role, since they allow us to perform arbitrary quantum logic operation [1]. Recently, a practical realization of these gates is a challenge for many physical laboratories. Among several propositions of constructing the systems performing the two-qubit gates, the most promising are those based on the mechanism of controlled switching on/off the exchange interaction between spin qubits [2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14]. Tuning the exchange interaction between electrons in coupled quantum subsystems can lead to the interchange of qubits and the performance of a designed quantum logic operation.

A physical implementation of the controlled exchange interaction is itself a difficult task. Moreover, the elements of a future quantum computer have to fulfill several conditions, known in the form of DiVincenzo criteria [15]. One of them is a scalability, which allows to join the elements into a large computational machine. It is expected that semiconductor nanodevices, in particular those consisting semiconductor quantum dots (QDs), should be scalable to large enough size. Recently, the electron spin states are the most promising candidates for qubits [6, 8, 9] due to the long coherence time [16]. The quantum information processing can be performed via the changes of electron spin states [8, 17]. The spin of the electron can be rotated as a result of precession in the external static magnetic field [14, 18]. The spin swapping can also be obtained when irradiating the electron system by the electromagnetic wave with the frequency adjusted to the Zeeman splitting in the external magnetic field [5, 19].

In the present paper, we study the process of swapping the electron spins, which results from the controlled switching of the exchange interaction between the electrons in coupled QDs. The tuned exchange interaction should lead to the rotation of electron spin [2, 3, 9, 14, 17, 20]. The theoretical model proposed takes into account all the three space dimensions and the electron spins. We apply the adiabatic approximation in order to decouple the transverse and longitudinal degrees of freedom and perform the integration over the transverse spatial coordinates. This leads to the effectively one-dimensional two-electron problem [21], which can be solved by the numerical method with an arbitrary precision [22]. The system under study is described by the two-particle wave function of the form of four-component state vector, which takes into account all possible spin configurations. We consider different
methods of turning on and off the exchange interaction between the electrons and investigate
the resulting rotation of electron spins. The present computer simulations are based the
accurate numerical solutions of the time-dependent Schrödinger equation, i.e., they allow us
to trace the time evolution of the electron system in a direct manner. Therefore, the present
results fill in a gap between the quantum information theory and experiment. We hope that
these results will serve as a guide for the experimental groups who are involved in designing
and constructing the nanodevices with spin qubits.

The paper is organized as follows: the theoretical model is presented in Sec. II, the
numerical method is described in Sec. III, the results are presented in Sec. IV (for the
vertically coupled QDs) and in Sec. V (for the laterally coupled QDs). Section VI contains
conclusions and summary.

II. THEORETICAL MODEL

The model applied in the present paper is an extension of that proposed by the present
authors in Ref. [23]. Contrary to Ref. [23], we explicitly take into account the spin states
of electrons. For sake of completeness, we repeat below the major steps of the approach
[23]. We study the two electrons localized in the double QD nanostructure. We assume
the cylindrical symmetry of the system with symmetry axis \((z\text{ axis})\) going through the
geometrical centers of the QDs. The potential confining the electrons in the \(x - y\) plane
is taken to be so sufficiently strong that the differences between the energy levels resulting
from the \(x - y\) space quantization are much larger than the energy of spin swapping. This
potential is usually called the lateral confinement potential. We approximate the lateral
potential by the two-dimensional harmonic oscillator potential and assume that both the
electrons occupy the ground state in the \(x - y\) motion. In other words, we assume that the
lateral electron degrees of freedom are frozen. The above assumptions allow us to reduce the
starting three-dimensional problem to the effectively one-dimensional problem [21] with the
electron-electron interaction being the following function of \(z_1\) and \(z_2\) coordinates of both
the electrons:

\[
U_{\text{eff}}(z_1 - z_2) = \frac{\varepsilon^2 \sqrt{\pi} \beta}{4 \pi \varepsilon_0 \varepsilon} e^{\beta |z_1 - z_2|^2} \text{erfc} \left( \sqrt{\beta} |z_1 - z_2| \right),
\]

where \(\varepsilon_0\) is the vacuum electric permittivity, \(\varepsilon\) is the static electric permittivity, \(\beta = m_e \hbar \omega_\perp / 2\), \(m_e\) is the electron effective conduction band mass, and \(\hbar \omega_\perp\) is the excitation
energy levels of the electron transverse motion. The Hamiltonian of the system takes the form \[21\]

\[
H = -\frac{\hbar^2}{2m_e} \left( \frac{\partial^2}{\partial z_1^2} + \frac{\partial^2}{\partial z_2^2} \right) + U(z_1) + U(z_2) + U_{\text{eff}}(z_1 - z_2) + 2\hbar\omega_{\perp},
\]

where \(U(z_i)\) is the potential energy of the vertical confinement of the \(i\)th electron. In the following, vertical confinement \(U(z)\) will be taken in a form of two potential wells separated by the potential barrier. The energy is measured with respect to the conduction band bottom of the quantum well material. In the calculations, we take on the parameters, which correspond to nanostructure based on GaAs: \(m_e = 0.067m_{e0}\), where \(m_{e0}\) is the free electron rest mass, and the static electric permittivity \(\varepsilon = 11.0\).

In the two-electron system, we are dealing with four independent spin states; therefore, the total wave function of the system can be represented by the following four-component vector:

\[
\Psi(z_1, \sigma_1, z_2, \sigma_2) = \begin{pmatrix}
p_{\uparrow\uparrow} \\
p_{\uparrow\downarrow} \\
p_{\downarrow\uparrow} \\
p_{\downarrow\downarrow}
\end{pmatrix},
\]

where \(\sigma_j (j = 1, 2)\) are the spin coordinates of the electrons, \(\psi_{nm} = \psi_{nm}(z_1, z_2)\) are the basis wave functions with the indices \(n\) and \(m = \uparrow\) and \(\downarrow\), which correspond to the \(z\)-spin component eigenvalues \(+\hbar/2\) and \(-\hbar/2\). Basis wave functions \(\psi_{nm}\) do not possess a well defined symmetry. However, total wave function (3) has to be antisymmetric with respect to simultaneous exchange of the space and spin coordinates. This property imposes the following conditions on the basis wave functions:

\[
\begin{align*}
\psi_{\uparrow\uparrow}(z_1, z_2) &= -\psi_{\uparrow\downarrow}(z_2, z_1), \\
\psi_{\downarrow\uparrow}(z_1, z_2) &= -\psi_{\downarrow\downarrow}(z_2, z_1), \\
\psi_{\downarrow\downarrow}(z_1, z_2) &= -\psi_{\uparrow\downarrow}(z_2, z_1).
\end{align*}
\]

In representation (3), the operators of the total spin are expressed by the following \(4 \times 4\) matrices:

\[
\sigma_x^{\text{tot}} = \begin{pmatrix}
\sigma_x & 1 \\
1 & \sigma_x
\end{pmatrix}, \quad \sigma_y^{\text{tot}} = \begin{pmatrix}
\sigma_y & -i1 \\
i1 & \sigma_y
\end{pmatrix}, \quad \sigma_z^{\text{tot}} = \begin{pmatrix}
\sigma_z + 1 & 0 \\
0 & \sigma_z - 1
\end{pmatrix},
\]
where $1$ denotes the unit $2 \times 2$ matrix and $\sigma_x, \sigma_y, \sigma_z$ are the spin Pauli matrices. The corresponding expectation values are calculated as $\langle \sigma_k^{\text{tot}} \rangle = \langle \Psi \sigma_k^{\text{tot}} \Psi \rangle$, where $k = x, y, z$. Multiplying $\langle \sigma_k^c \rangle$ by $\hbar/2$ we obtain the expectation values of the components of the total spin, i.e., $S_k^{\text{tot}} = (\hbar/2) \langle \sigma_k^{\text{tot}} \rangle$.

Due to their indistinguishability the electrons can not be numerated; therefore, we can not determine which electron is in a given spin state. However, we can distinguish the two quantum dots, determined by the corresponding potential wells. We shall call them the left (L) and right (R) quantum dot (potential well). If both the QDs are separated by the potential barrier, which prohibits a tunneling of electrons, then in the ground state of the system, each electron is localized in a single QD. Owing to this, we can determine the spins of electrons in the left and right QD. For this purpose, we introduce the auxiliary wave function $\varphi(z_1, z_2)$, which will be called the reference wave function. This wave function is a solution of the Schrödinger equation, for the two distinguishable particles, which – with an exception of indistinguishability – possess all the properties of the electrons. For the distinguishable particles we do no perform a symmetrization of the two-particle wave function. Instead the solutions found for $\varphi(z_1, z_2)$ correspond to the configurations, in which one electron (described by coordinate $z_1$) is localized in the left QD and the other ($z_2$) is in the right QD. This state always exists if the tunneling through the barrier is not possible. In each simulation performed, these configurations are realized at the initial ($t = 0$) and final ($t = T$) time instants. The reference wave function will serve to a construction of the initial state wave function and a determination of the spin states of electrons in both the QDs. The expectation values of spin components are calculated as follows

$$S_j^k = \frac{\hbar}{2} u^T \sigma_k^j u,$$

where $j = L, R$ corresponds to the electron localized in the left (L) and right (R) QD, and

$$u = \begin{pmatrix} \langle \varphi | \psi_{\uparrow \uparrow} \rangle \\ \langle \varphi | \psi_{\uparrow \downarrow} \rangle \\ \langle \varphi | \psi_{\downarrow \uparrow} \rangle \\ \langle \varphi | \psi_{\downarrow \downarrow} \rangle \end{pmatrix}.$$  

The spin matrix operators in Eq. (6) have the form

$$\sigma_x^L = \begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix}, \quad \sigma_y^L = i \begin{pmatrix} 0 & -1 \\ 1 & 0 \end{pmatrix}, \quad \sigma_z^L = \begin{pmatrix} 1 & 0 \\ 0 & -1 \end{pmatrix}.$$  

(8a)
\[ \sigma^R_x = \begin{pmatrix} \sigma_x & 0 \\ 0 & \sigma_x \end{pmatrix}, \quad \sigma^R_y = \begin{pmatrix} \sigma_y & 0 \\ 0 & \sigma_y \end{pmatrix}, \quad \sigma^R_z = \begin{pmatrix} \sigma_z & 0 \\ 0 & \sigma_z \end{pmatrix}. \quad (8b) \]

The operator of the \( k \)th component of the total spin is defined as \( S^\text{tot}_k = (\hbar/2)(\sigma^L_k + \sigma^R_k) \).

III. SETTING UP THE COMPUTER EXPERIMENT

We simulate the process of swapping the spins for two electrons localized in two QDs, which are separated by the potential barrier. We consider both the vertical and lateral geometry of the QD nanostructure. In the process studied, the spin of each electron localized in the left or right QD is flipped. The spins are swapped as a result of the controlled switching of the exchange interaction between the electrons. We discuss two methods to control the exchange interaction. According to the first method, the exchange interaction is switched on (off) by lowering (rising) the potential barrier. The second method is based on the changes of the depth of one of the potential wells, which leads to the flow of electrons and their localization in the same QD. In the QD-based nanodevices, both the methods can be implemented by changing the external voltages, which are the source of the electrostatic field forming the coupled QDs [13], or by locating the QD system in the electromagnetic field [7].

Based on the analogy with Rabi oscillations in a two-level quantum system, the process of switching on/off the exchange interaction that leads to the change of the initial spin orientation to the opposite one, will be called the \( \pi \) pulse. This process lasts for time \( T^\pi \). We shall also deal with processes, after which the expectation value of \( z \) electron spin component does not reach \( \pm \hbar/2 \), i.e., \( |\langle S^j_z \rangle| < \hbar/2 \). In such processes, the change of spin orientation is not complete. In electrostatic QDs, the \( \pi \) pulse can be realized by the proper change of gate voltages.

The additional purpose of the present study is the optimization of nanodevice parameters and time changes of the confinement potential in order to make the spin interchange time possibly short. The minimization of \( T^\pi \) is important since this time interval determines the duration of elementary quantum logic gate operation that should be much shorter than the coherence time of qubits. It is required [15] that the ratio of the gate operation time to the coherence time should be less than \( 10^{-4} \) in order to complete a computation and several
error correction runs \[24\] before the decoherence destroys the information stored in qubits. In GaAs, the spin coherence time has been estimated to be longer than 1 ms \[25\].

At the initial time instant in each simulated process, the system is always in the singlet ground state, in which the electron localized in the left (right) QD possesses the spin \( z \) component \( S^L_z(t = 0) = +\hbar/2 \) (\( S^R_z(t = 0) = -\hbar/2 \)). The two-electron wave function that satisfies these initial conditions has a form

\[
\Psi_i \equiv \Psi(z_1, \sigma_1, z_2, \sigma_2, t = 0) = \begin{pmatrix} 0 \\ \varphi(z_1, z_2) \\ -\varphi(z_2, z_1) \\ 0 \end{pmatrix},
\]

where \( \varphi(z_1, z_2) \) is the reference wave function defined in Sec. II. Wave function (9) fulfills symmetry constraints \(11\). After a normalization, the wave function (9) is subjected to a temporal evolution according to the time-dependent Schrödinger equation

\[
i\hbar \frac{\partial \Psi(z_1, z_2, t)}{\partial t} = H \Psi(z_1, z_2, t),
\]

where \( H \) is given by Eq. (2) and the time dependence of potentials \( U(z_1, z_2, t) \) and \( U_{\text{eff}}(z_1, z_2, t) \) will be determined in the following sections.

The spin expectation values are calculated using Eq. (6) as follows. First, the reference wave function \( \varphi(z_1, z_2) \) is calculated by the imaginary time step method \[22\]. The reference wave function is next applied to construct total wave function \( \Psi_i \) (9) for \( t = 0 \). Wave function \( \Psi_i \) is used as the initial condition to perform the time simulation of Schrödinger equation (11). In the last step, we apply \( \varphi(z_1, z_2) \) to calculate \( u \) (Eq. (7)) and \( S^L_k \) (Eq. (3)).

The results of simulations are presented as a function of duration time \( T \) of the process of switching the exchange interaction. We note that we are dealing with two time intervals \( T \) and \( T^\pi \), which have a different meaning. Time interval \( T \) determines the duration of the arbitrary process of changing the exchange interaction. Time interval \( T^\pi \) determines the duration of such process, during which the spins are fully swapped. At the end of each simulation, we record the expectation values of \( z \) spin component of the electron in the left (L) and right (R) QD. These final values of spins are denoted by \( S^L_{f,R} \), where \( S^L_{f,R} = S^{L,R}_{f,R}(t = T) \). Moreover, we record the energy difference \( \Delta E_{fi} = E_f - E_i \) between the energies of the final \( (E_f) \) and initial \( (E_i) \) states of the system. The non-zero value of \( \Delta E_{fi} = E_f - E_i \) gives the evidence that the process in non-adiabatic.
The numerical procedure applied allows us to observe the spins of single electrons in another manner. Namely, we can directly determine the temporal evolution of the second \( \psi_{\uparrow\downarrow} \) and third \( \psi_{\downarrow\uparrow} \) component of the total wave function \( \psi \), whereas – due to the symmetry – it is sufficient to detect one of these components. The full swapping of spins corresponds to the interchange of these two components, i.e.,

\[
\begin{pmatrix} 0 \\ \psi_{\uparrow\downarrow} \\ \psi_{\downarrow\uparrow} \\ 0 \end{pmatrix} \leftrightarrow \begin{pmatrix} 0 \\ \psi_{\downarrow\uparrow} \\ \psi_{\uparrow\downarrow} \\ 0 \end{pmatrix} \quad (11)
\]

IV. VERTICALLY COUPLED QUANTUM DOTS

The vertically coupled QDs [26] can be described by the simple model with two rectangular potential wells separated by the rectangular barrier (Fig. 1). The vertical QDs were fabricated on the basis of GaAs/InGaAs/AlGaAs heterostructure [26, 27]. The exchange interaction is switched on by lowering the height of potential barrier to the potential well bottom. The exchange interaction is switched off by raising the potential barrier to the initial height. The time dependence of the process of changing the barrier height can chosen in many ways. In the present paper, we have chosen the time dependencies described by the step, linear, and smooth functions.

We choose such values of the parameters of the nanostructure (Fig. 1) that the electron tunneling is negligibly small in the initial and final time instant. Then the electrons are well separated in space and localized in the left (right) QD. Owing to this the spin states of both electrons can be exactly known in both the initial and final instant. We choose the same widths of the potential wells and potential barrier \( d_L = d_R = d_B = 10 \) nm and the potential well depth \( V_L = V_R = -150 \) meV. At the initial time instant, we take on the top barrier energy \( V_B = 0 \) (the conduction band bottom of the barrier material is chosen as the reference energy and set equal to zero).
FIG. 1: Confinement potential profile $E \equiv V$ as a function of vertical coordinate $z$ and energy $V_B$ of the top of potential barrier. (a) $V_B = 0$, (b) $V_B = -50$ meV, (c) $V_B = -100$ meV, and (d) $V_B = -150$ meV.

A. Time changes of the potential barrier

First, we study the effect of the time step dependence of the barrier height [Fig. 2(a)]. In this process, the top of the barrier is rapidly lowered from $V_B = 0$ to $V_B = -150$ meV, i.e., the potential well bottom. We let the system to evolve for time $t = T$, after which the barrier is rapidly raised to the starting value. We have performed a series of simulations for different operation time $T$. At the beginning and the end of each process, we have determined $z$ spin component expectation values $S_{i,f}^{L,R}$.

FIG. 2: Time dependence of top barrier $E \equiv V_B$ changes: (a) step function, (b) linear function, (c) smooth (cosinus) function.

The results, displayed in Fig. 3(a), show that the spin expectation value detected at the final instant of each process rapidly oscillates as a function of time duration $T$ of the
process. However, the amplitude of these oscillations does not reach $\hbar/2$. Therefore, the sudden changes of the barrier do not lead to the full interchange of spins at any time interval $T$ studied. The behaviour of energy difference $\Delta E_{fi}$ between the final and initial state [Fig. 3(b)] allows us to explain this effect. Similar to the average electron spin, energy difference $\Delta E_{fi}$ exhibits rapid oscillations as a function of duration time $T$ of the process. We interpret the results of Fig. 3(b) as follows: during the sudden jumps of the barrier height, the electron system makes transitions to the excited states. As a consequence, the energy of electrons in the final state increases, i.e., the energy difference is always larger than zero and exhibits jumps, and the electron wave function spreads out over the two potential wells. Therefore, the switching of the interaction between the electrons in a rapid manner is not an adiabatic process.

![Graphs](image)

**FIG. 3:** (a) Expectation value $S_{L,P}^{z,f}$ of $z$ spin component detected at the end of the process of switching on and off the exchange interaction as a function of its duration time $T$ for step-like changes of the barrier. Solid (dashed) curve shows the results for the electron in the right (left) QD. (b) Energy difference $\Delta E_{fi}$ between the final and initial states energies as a function of duration time $T$ of the process.

Next, we study the two-step process, during which the potential barrier is a linear function of time [Fig. 2(b)]. In the first step, for $0 \leq t \leq T/2$, the top barrier energy is lowered according to

$$V_B(t) = V_B^{\text{max}} - 2(V_B^{\text{max}} - V_B^{\text{min}}) \frac{t}{T},$$

(12)

where $V_B^{\text{max}}$ ($V_B^{\text{min}}$) denotes the largest (smallest) energy of the barrier top in time interval
In the second step, for $T/2 \leq t \leq T$, the potential barrier is raised according to

$$V_B(t) = V_B^{\text{min}} + 2(V_B^{\text{max}} - V_B^{\text{min}}) \left( \frac{t}{T} - \frac{1}{2} \right).$$

(13)

The results (Fig. 4) show that spin expectation values $S_f^{L,R}$ are periodic functions of time $T$. Each maximal (minimal) value $S_f^R = +\hbar/2$ ($S_f^L = -\hbar/2$) of $z$ spin component corresponds to the state, in which both the electrons possess the spins of the opposite orientation with respect to that in the initial state. Therefore, there exists a series of duration times $T$, for which the $z$ spin component of each electron changes its sign, i.e., the electrons fully exchange their spins. The subsequent extrema in Fig. 4(a) correspond to the $\pi$ pulses with duration time $T_\pi^0 = nT_1^\pi + \text{const}$, where index $n$ numerates the extrema. If we increase process duration time $T$, we simultaneously increase the time interval during which both the electrons are localized in the same region of space. As a consequence, for the sufficiently long time $T$ the spin swapping can occur many times. If the process lasted for infinitely long time, the spin interchange would occur infinitely many times. This scenario would be realized for an isolated system (non-interacting with an environment). In this case, spin expectation values $S_f^{L,R}$ would oscillate with amplitude $\hbar/2$ infinitely many times, i.e., there exists an infinite series of time intervals $T_\pi^0$, for which the spins are fully interchanged. However, in real systems, we deal with the decay of quantum states, which leads to the energy dissipation, and the decoherence, which randomly changes the relative phase of spin qubits. These processes will lead to the decreasing amplitude of $S_f^{L,R}$ for increasing time interval $T$. Therefore, in order to perform successful quantum logic operation the spin swapping process should last for possibly short time. However, there exists the lower limit on time interval $T$, during which the barrier is changed. This limit results from the requirement of the adiabaticity of the process: the barrier has to be changed sufficiently slowly, i.e., in an adiabatic manner, in order to leave the system in the ground state. If time interval $T$ is too short, the system can be excited to higher-energy excited states, which leads to the increase of its energy and the non-adiabaticity of the process.

The non-adiabaticity of the spin interchange process can be also observed in the behaviour of $S_f^{L,R}(T)$ [Fig. 4(a)]. A closer look at the present computational results has led us to the observation that the first maximum is slightly lower than the other maxima. For the first maximum the exact spin expectation value reaches $\hbar/2$ with accuracy 98.7%. Similarly, due to the symmetry, for the first minimum, $S_f^L$ reaches the value $-\hbar/2$ with the same accuracy.
The extrema, which correspond to the second $\pi$ pulse, reach $\pm h/2$ with accuracy 99.4%. The next extrema tend to $\pm h/2$ with a high accuracy. Therefore, we deal with several processes of incomplete interchange of spins before $S_{f}^{L,R}$ start to switch between $\pm h/2$. The lowering of the amplitude of $S_{f}^{L,R}$ is more pronounced if the process of changing the potential barrier is more rapid. The present results show that several processes of incomplete spin swapping can occur before we achieve the full interchange of spins, which leads to an elongation of the spin exchange time. We note that this effect is not expected within the model, based on the effective Heisenberg Hamiltonian. In this model, the exchange interaction between the electrons is described by the Hamiltonian $H(t) = J(t)S_1 \cdot S_2$, where $J(t)$ is exchange interaction energy and the spin operators $S_j$ ($j = 1, 2$) are expressed in terms of the corresponding Pauli matrices. Our estimates show that – in real quantum systems – the spin interchange time will be longer than that resulting from the Heisenberg model, which do not take into account the physical implementation of switching on/off the exchange interaction.

![Graphs](image)

**FIG. 4:** (a) Expectation value $S_{f}^{L,P} = S_{f}^{L,R}$ of $z$ spin component detected at the end of the process of switching on and off the exchange interaction as a function of duration time $T$ for linear changes of the barrier. Solid (dashed) curve shows the results for the electron in the right (left) QD. (b) Energy difference $\Delta E_{fi}$ between the final and initial state energies as a function of duration time $T$ of the process. Inset zooms in the part of this figure.

Fig. 4(b) displays the dependence of energy separation $\Delta E_{fi}$ on time interval $T$. In the inset, we zoom in this dependence for the narrower scale of $T$, which allows us to observe the oscillations of $\Delta E_{fi}$. The amplitude of these oscillations decreases if the duration time of
spin interchange increases. Moreover, $\Delta E_{fi}$ asymptotically approaches zero for large $T$. For small $T$ the positions of the local minima of $\Delta E_{fi}$ do not coincide with $T^\pi_n$ [cf. Fig. 4(a)]. For the chosen nanostructure parameters, this coincidence appears for the third $\pi$ pulse at $T = 8.2$ ps. For linear changes of the barrier we deal with rapid turning on the process at $T/2$, which causes the non-adiabaticity of this method of spin interchange.

Based on the above results, we expect that we will obtain better results if the temporal changes of the potential barrier will be modelled by a smooth function of time. In the following, we study the smooth time dependence of the potential barrier, which is modelled by the following function [Fig. 2(c)]:

$$V_B(t) = \frac{1}{2}(V_B^{\text{max}} - V_B^{\text{min}})(1 - \cos \omega t) + V_B^{\text{max}},$$

(14)

where $\omega = 2\pi/T$. The results of simulations [Fig. 5(a)] show that the amplitude of $S_L^R(T)$ approaches $\hbar/2$ much faster than for step-like and linear changes of the barrier. Already the second maximum reaches $\hbar/2$ with accuracy 99.99%. Moreover, based on the results for energy difference $\Delta E_{fi}$ [Fig. 5(b)] we can regard this process to be adiabatic for $T \geq 2$ ps. The very fine jumps of $\Delta E_{fi}$ near zero, shown in the inset of Fig. 5(b), result from the small numerical errors. Therefore, the amplitude of the periodic changes of $S_L^R$ remains constant (and equal to $\hbar/2$) for $T > 2$ ps.

The spin of the electron localized in the left (right) QD can be determined from Eq. (6) only if the probability of finding both the electrons in the same QD is zero. This state is realized at the initial and final time instant in each process of the barrier changes. At the intermediate time instants, the electrons are not spatially separated; therefore, quantities $S_L^R \equiv S_L^R$, calculated from Eq. (6), are not equal to the eigenvalues of $z$ spin component of the electron in the left and right QD. Nevertheless, for arbitrary time, $S_L^R$ can serve as a control spin index.

In Fig. 6 we present the time dependence of control spin indices and energy of the system, which correspond to the pulses with duration times $T_{1}^\pi$ and $T_{2}^\pi$ for linear and smooth changes of the barrier. The energy is measured relative the ground state energy $E_0$ of two electrons in the lateral confinement potential ($E_0 = 2\hbar\omega_\perp$, where $\hbar\omega_\perp = 40$ meV). The behaviour of these time characteristics is different for the smooth and non-smooth changes of the barrier. In the case of the non-smooth process, the discontinuity of the first time derivative of the top barrier energy for $t = T/2$ leads to the corresponding discontinuity of the energy versus time
FIG. 5: (a) Expectation value $S_{L,P}^{L}$ of $z$ spin component detected at the end of the process of switching on and off the exchange interaction as a function of duration time $T$ for smooth changes of the barrier. Solid (dashed) curve shows the results for the electron in the right (left) QD. (b) Energy difference $\Delta E_{f,i}$ between the final and initial state energies as a function of duration time $T$ of the process. Inset zooms in the part of this figure.

plot [cf. dotted curves in Figs. 6(a) and 6(b)]. The consequences of this discontinuity can also be seen in the time dependence of the control spin indices [cf. solid and dashed curves in Figs. 6(a) and 6(b)]. For the pulse with time duration $T_{1}^{\pi} = 1.58$ ps the spin indices begin to oscillate after reversing the time evolution of the potential barrier [Fig. 6(a)]. The amplitude of these oscillations decreases with the increasing barrier height and falls down to zero when the barrier height reaches the starting level. For the three times longer pulse $T_{2}^{\pi} = 4.91$ ps reversing the time changes of the barrier leads to smaller oscillations [cf. Fig. 6(b)]. Nevertheless, the time dependence of the control spin indices is still slightly perturbed.

In the case of smooth time changes of the barrier [Eq. (14)], the energy of the system is also a smooth function of time [Figs. 6(c,d)]. In spite of this, for the first $\pi$ pulse with $T_{1}^{\pi} = 0.99$ ps the changes of the potential barrier are still seen by the system as rapid changes, which leads to bending the spin characteristics [Fig. 6(c)]. However, the time duration of the second $\pi$ pulse ($T_{2}^{\pi} = 3.04$ ps) is sufficiently long so that the spin characteristics [Fig. 6(d)] stay smooth during the full process of switching on/off the exchange interaction. In the cases of linear and smooth changes of the potential barrier, the time evolution of spin indices $S^{L,R}(t)$ for the second $\pi$ pulse with time duration $T_{2}^{\pi}$ suggests that the full interchange of spins occurs as a result of many-fold exchange of spins between the electrons. For the
FIG. 6: Spin control indices $S^{L,P}_f \equiv S^{L,R}$ and energy $E$ of the two-electron system as functions of time $t$. Plots (a) and (b) [(c) and (d)] correspond to the linear [cosinusoidal] changes of the potential barrier, plots (a) and (c) [(b) and (d)] correspond to the first [second] $\pi$ pulse with duration time $T^{\pi}_1$ [$T^{\pi}_2$]. Solid (dashed) curves display the control spin index of the electron in the right (left) QD and the dotted curves show the energy of the system.

second pulse the curves $S^{L,R}_f(t)$ go through zero three times [cf. Figs. 6(b,d)]. Thus, we conclude that in these processes we are dealing with the three-fold incomplete swapping of spins before we finally obtain the full interchange of spin orientations. In order to obtain an additional support for the occurrence of this process, we have observed the time changes of the second and third components of the total wave function [Eq. (3)]. The results of the simulations performed show the many-fold interchange of the second and third wave function components, which means that – in fact – the spins of the electrons change many
Based on these results, we have chosen to further studies the method of the smooth switching of the exchange interaction, which seems to be the most promising in a physical realization.

**B. Effect of quantum dot size and potential well asymmetry**

If the size of the coupled QD nanostructure is sufficiently large, the Coulomb interelectron repulsion leads to a localization of electrons at the opposite potential well boundaries \(32\). Then, the electrons form the Wigner molecule \(32\). In the case of Wigner localization, the lowering of the potential barrier can cause no interchange of spins during a time, which is shorter than the spin coherence time. We have checked the possibility of the occurrence of this effect by investigating the influence of the QD size of the time duration of spin interchange process. The calculations have been performed for the QD nanostructure modelled by rectangular potential wells and a barrier. The barrier thickness was fixed as \(d_B = 10\) nm, the top barrier energy was changed according to Eq. (14), the depths of both the potential wells were equal \((V_L = V_R = -200\) meV), and the thicknesses of both the potential wells were also the same \(d_L = d_R \equiv d\). We change potential well width \(d\) in the interval \(d \in [6, 50]\) nm. The size of the region, in which the spin exchange appeared, has the width \(2d + d_B\). Figure 7 depicts the calculated time duration of the first and second \(\pi\) pulse as a function of width \(d\) of the potential wells. We see that the time duration of spin interchange quickly increases with increasing QD size. The \(\pi\) pulse time changes with the QD size as follows: \(T^{\pi} \sim d^3\) for \(d < 20\) nm and \(T^{\pi} \sim d^6\) for \(d > 20\) nm. These results show that even for the QD with the size such small as \(d \simeq 50\) nm the full spin swapping can be not realized during the coherence time.

In the Heisenberg model of electron-electron interaction, the rate of spin interchange is determined by the exchange coupling constant, which is defined as the energy difference between the triplet and singlet states, i.e., \(J = E_T - E_S\). For the two identical QDs separated by the sufficiently thick and high potential barrier the triplet and singlet states are degenerate and \(J = 0\). If the potential barrier becomes thinner and lower, the wave functions of the electrons localized in both the QDs start to overlap and the energies of triplet and singlet states become different, i.e., \(J \neq 0\). The coupling constant \(J\) can also be
FIG. 7: Time duration $T_\pi$ of the first ($n = 1$) and second ($n = 2$) $\pi$ pulse as a function of thickness $d$ of the potential well. Inset zooms in the short time part of the figure.

changed if the QD potential wells are different, i.e., the QDs are asymmetric \[30\]. The larger the asymmetry of the potential wells the larger coupling constant $J$. Therefore, one could expect that the asymmetry of the QDs should lead to a shortening of the spin interchange time. However, it turns out \[31\] that this is not always the case.

We have investigated the influence of the asymmetry of potential well thicknesses on the duration time of spin interchange. For the fixed thickness of the left QD ($d_L = 10$ nm) we have performed the calculations for several values of thickness $d_R$ of the right QD. The results are plotted in Fig. 8 for $d_R = 5$ nm and $d_R = 15$ nm. For sake of clarity, we present in Fig. 8 only the results for the electron in the right QD. The spin expectation values ($S^L_f$) for the electron in the left QD can be obtained by the reflection of $S^R_f(T)$ curves with respect to the axis $S^R_f = 0$.

The results of calculations are displayed in Fig. 8. If the right potential well is thinner than the left one, energy difference $\Delta E_{fi}$ between the final and initial state tends to zero for long duration time $T$ of spin interchange [dashed curves in Fig. 8(a)]. However, if the right QD is thicker than the left one, $\Delta E_{fi}$ does not reach zero even for long time $T$ [Fig. 8(b)]. Moreover, it takes on negative values for some times $T$. The negative values of $\Delta E_{fi}$ can be explained by the shift of electron density toward the wider QD in which the final state energy is lower. Analyzing the plots of $S^R_f$ vs $T$ we have observed that the accuracy of the interchange of spins reaches 99.9 %, but is never equal to 100 %. We conclude that
the asymmetry of the QD confining potential results in the increasing multiplicity of spin exchange, which leads to the longer duration time of spin interchange.

FIG. 8: Expectation value $S^P_f \equiv S^R_f$ of $z$ spin component detected in the right QD at the end of the process of switching on/off the exchange interaction (solid curves) and energy difference $\Delta E_{fi}$ between the final and initial states (dashed curves) as functions of duration time $T$ of spin interchange for asymmetric QDs. The thickness of the left QD is fixed $d_L = 10$ nm, while the thickness of the right QD $d_R = 5$ nm in (a) and $d_R = 15$ nm in (b).

V. LATERALLY COUPLED QUANTUM DOTS

The electrostatic QDs with lateral interdot coupling are the subject of many recent studies [6, 28, 29]. In the laterally coupled QDs, the profile of the confinement potential can be modified by varying the voltages applied to the gates. This provides a fairly convenient way of tuning the potential barriers and wells. However, due to technological limitations, the size of the lateral QD is of the order of 100 nm [14], i.e., is larger than the size ($\sim 10$ nm) of the vertical QD [27] measured in the growth direction. The nanostructure consisting of two laterally coupled QDs separated by the potential barrier has a typical linear size $200 – 300$ nm. Due to this relatively large size, the duration time of the spin rotation is expected to be long (cf. see Subsection IV.B).

We still apply the theoretical model described in Section II with the frozen transverse motion of electrons. Now $(y, z)$ are the transverse coordinates and we assume that the
electrons move in the $x$ direction, e.g., the $x$ axis can be directed along the effectively one-dimensional flow of electrons between the two lateral QDs [14]. The larger size of the lateral QDs in comparison to that of vertical QDs leads to the weaker electron localization and the weaker electron-electron interaction. According to the theory [21], the strength of the potential confining the electrons in the transverse directions determines the strength of the effective electron-electron repulsion [Eq. (11)]. [We note that now $U_{\text{eff}}$ is a function of $x$.] The excitation energy $\hbar \omega_{\perp}$ is a measure of the strength of the transverse parabolic confinement. In the calculations for the lateral QDs, we take on $\hbar \omega_{\perp} = 5$ meV. Then, the transverse confinement potential rather well approximates the confinement in the double lateral QD fabricated by the TU Delft laboratory [33]. The present model is also applicable to quantum wires, which are composed from different semiconductors, e.g., InP and InAs [34], that form a structure of potential wells and barriers. In this case, we deal with quasi-one-dimensional coupled QDs. In the present section, we modify the exchange interaction between the electrons localized in laterally coupled QDs by tuning the potential barrier as well as the potential wells.

A. Symmetric quantum dots

The profile of the confinement potential for the laterally coupled symmetric QDs is assumed in the form (Fig. 9)

$$V(x) = kx^2 + V_2 \exp \left[-(x-x_0)^2/d^2\right]$$

where $k$ determines the strength of the parabolic confinement potential, parameter $V_2$ is the potential barrier height for $V_2 > 0$ (potential well depth for $V_2 < 0$), $x_0$ is the position of the center of the potential barrier (well), and $d$ is the range of the Gaussian potential.

The potential barrier separating the two QDs can be modified by changing $V_2$ [Eq. (15)]. In the calculations, parameter $V_2$ was changed in time in a smooth manner, i.e., according to Eq. (14), from 9 meV to 0 and next from 0 to 9 meV. Figure 9 shows the corresponding profiles of the confinement potential. We see that when changing the potential barrier we simultaneously change the potential wells. If the potential barrier is lowered, the electrons become localized closer to each other. The effective QD size ($d_{\text{eff}}^{L,R}$) can be defined as the width of the corresponding potential well determined at the half of barrier height. At the
initial time moment, the effective size of the QDs $d_{L}^{\text{eff}} = d_{R}^{\text{eff}} = 100 \text{ nm}$, the effective width of the barrier $d_{B}^{\text{eff}} = 60 \text{ nm}$, the electron in the left (right) QD has spin $+\hbar/2$ ($-\hbar/2$).

![Profiles of confinement potential](image)

**FIG. 9:** Profiles of confinement potential $E \equiv V$ vs $z \equiv x$ for two laterally coupled QDs for $V_{2} \geq 0$. Parameters of the confinement potential [Eq. (15)] are $k = 9.0 \times 10^{-7} \text{ meV(nm)}^{-2}$, $d = 185 \text{ nm}$, $x_{0} = 0$, $V_{2} = 9.0, 4.3$, and $0 \text{ meV}$ (solid, dashed, and dotted curves).

In Fig. 10, we display the time evolution of spin indices $S_{L,R}$ and parameter $V_{2}$. These results show that for the first $\pi$ pulse with time duration $T_{1}^{\pi} = 492 \text{ ps}$ the spin interchange occurs with 100% accuracy. We see that during the larger part of time evolution the spins do not change. The spins start to rotate when the barrier height $V_{2}$ decreases to $\sim 5 \text{ meV}$. This suggests that the lowering of the potential barrier should lead to a shorter spin rotation time. If, however, the initial barrier height decreases, the effective QD size decreases too. For $V_{2} = 5 \text{ meV}$ the effective size of the potential wells and the barrier decrease to $d_{L,R}^{\text{eff}} = 76 \text{ nm}$ and $d_{B}^{\text{eff}} = 42 \text{ nm}$. This means that the shortening of the spin rotation time results from the size effect, discussed in Subsection IV.B. The fastest rate of spin interchange is observed for $V_{2} < 1.5 \text{ meV}$. Then, the effective sizes are $d_{L,R}^{\text{eff}} = 40 \text{ nm}$ and $d_{B}^{\text{eff}} = 42 \text{ nm}$.

If the QD size is of the order of several ten nanometers, the electrons are spatially separated even in the absence of potential barrier [32]. For the QDs of this size the Coulomb repulsion is sufficiently strong so that the electrons are localized the QD boundaries and form Wigner molecules [32]. Only the localization of electrons close to each other can lead to the rotation of their spins. In the model of nanostructure (Fig. 9), the effective width of the potential barrier is much smaller than that of the double potential well. Therefore, we can force the electrons to be localized in the same region of the nanostructure, if we
allow the parameter $V_2$ in Eq. (15) to take on negative values during the time evolution. As a result, the potential barrier converts into the potential well (Fig. 11) and the electrons become localized in the same QD for some time during the $\pi$ pulse duration time.

FIG. 11: Profile of confinement potential $E \equiv V$ [Eq. (15)] as a function of coordinate $z \equiv x$ for $V_2 > 0$ and $V_2 < 0$. Shown are the plots for $V_2 = 9, 6, 3, 0, -3, -6, -9$ meV (from top to bottom). At the initial time instant (solid curve), $V_2 = 9$ meV, $d_B = 60$ nm, $d_L = d_R = 100$ nm.

In the first computer run, parameter $V_2$ was changed in a smooth manner from 9 meV to $-3$ meV. As expected, we obtained the considerably shorter duration time of the spin rotation: the first $\pi$ pulse lasted merely for 46.1 ps, i.e., it was more than ten times shorter than in the case of $V_2 \geq 0$. It appears a question: to what extent can we shorten the
duration time of spin rotation by lowering the potential well bottom? In order to answer this question, we have performed a series of simulations, in which the minimum value $V_{2}^{\text{min}}$ of parameter $V_2$ was lowered from 0 to $-9 \text{ meV}$. In each simulation, the initial value of parameter $V_2$ was the same: $V_2(t = 0) = V_{2}^{\text{max}} = 9 \text{ meV}$. The results are shown in Fig. 12. According to our expectation, the spin interchange duration time becomes shorter if $V_{2}^{\text{min}}$ decreases. However, we also observe an undesirable effect, namely, the process ceases to be adiabatic, which results from the increasing amplitude of changes of $V_2$ during time $T$.

The non-adiabaticity is visible in Figs. 12(a-d) as oscillations of energy difference $\Delta E_f(T)$ and deformations of plots $S_f^R(T)$ for small duration time $T$. The process can be treated as adiabatic for $T \geq 25 \text{ ps}$ ($T \geq 55 \text{ ps}$) if $V_{2}^{\text{min}} = 0$ ($V_{2}^{\text{min}} = -9 \text{ meV}$). We conclude that the lowering of $V_{2}^{\text{min}}$ below $-3 \text{ meV}$ is not advantageous. The shortest time of the full interchange of spins has been obtained for $V_{2}^{\text{min}} = -3 \text{ meV}$. Then, the orientation of spins was changed in the single swap after time $T_1^\pi = 46.1 \text{ ps}$ with accuracy 100%.

We note that for the laterally coupled QDs the change of the interdot potential barrier into the deep potential well means that a section of quantum wire is created below the gate electrodes, which are responsible for the formation of electrostatic QDs \footnote{Reference}. However, in this process, the spin qubits can be destroyed if the electrons will tunnel from the two-dimensional electron gas \footnote{Reference} to the QD. Therefore, it seems that the mechanism of conversion of the interdot potential barrier into the potential well will not lead to the required rotation of spins in the laterally coupled QDs. However, this mechanism can be applicable to the spin qubits in the vertically coupled QDs \footnote{Reference} and the coupled QDs formed in quantum wires \footnote{Reference}.

### B. Asymmetric quantum dots

The bias voltage applied to the coupled QDs leads to the asymmetry of the confinement potential profile, i.e., the potential well depths become different. This suggests another mechanism of swapping the spins, which can be modelled by raising and lowering the bottom of one potential well. Raising the potential well bottom of one QD will stimulate the flow of the electron to the second QD. Therefore, both the electrons will be localized in the same QD. The subsequent lowering of the potential well bottom back to the initial level will cause the flow of one of the electrons to the first QD. If the duration time of changing the
FIG. 12: Expectation value $S_f^P = S_f^R$ of $z$ spin component in the right QD (solid curve) and energy difference $\Delta E_{fi}$ (dashed curve) between the final and initial states as a function of time duration $T$ of the process. (a) $V_{2}^{\text{min}} = 0$, (b) $V_{2}^{\text{min}} = -3$ meV, (c) $V_{2}^{\text{min}} = -6$ meV, (d) $V_{2}^{\text{min}} = -9$ meV.

potential well will be suitably adjusted, the electrons will swap their spins. This mechanism is advantageous since the electrons meet in the same QD. According to the previous results, the stronger electron localization leads to the shorter time of spin swapping.

In order to simulate this process, we model the confinement potential by the linear combination of two Gaussians, i.e.,

$$V(x) = V_1 \exp \left[ -\frac{(x - x_1)^2}{d_1^2} \right] + V_2 \exp \left[ -\frac{(x - x_2)^2}{d_2^2} \right], \quad (16)$$

where $V_1$ and $V_2$ are the potential well depths, $x_1$ and $x_2$ are the positions of the centers of the QDs, and $d_1$ and $d_2$ determine the sizes of the QDs (Fig. 13).
FIG. 13: Profile of confinement potential $E \equiv V$ [Eq. (16)] as a function of coordinate $z \equiv x$ for several time instants. Parameter $V_1 = -10$ meV is fixed. At the initial moment (solid curve), $V_2(t = 0) = -10$ meV. Shown are also the potential profiles for $V_2 = -7, -5, -3,$ and $0$ meV.

We change parameter $V_2$ according to the smooth function of time [Eq. (14)]. At the initial time, the electron in the right (left) QD has spin $+\hbar/2$ ($-\hbar/2$). We have performed several simulations for different $\pi$ pulse duration time $T_\pi$ with the fixed parameters of confinement potential [Eq. (16)] $V_1 = -10$ meV, $d_1 = d_2 = 64$ nm, and $x_2 = -x_1 = 58$ nm. For $t = 0$ (solid curve in Fig. 13) the bottom of both the potential wells was at the level $-10.4$ meV, effective widths $d_1 = d_2 = 48$ nm, $d_B = 51$ nm, and the relative height of the barrier was $1.7$ meV. In each computer run, the initial (minimum) value of parameter $V_2$ was $V_2(t = 0) = V_2^{\text{min}} = -10$ meV. The potential profiles shown in Fig. 13 suggest that it is not necessary to remove the right potential well in order to localize both the electrons in the left QD. Again, we are looking for the answer to question: for which range of changes of the right potential well bottom the duration time of interchanging the spins is the shortest? The results of the simulations are displayed in Fig. 14. As in the previous figures, for sake of clarity, we present only the spin expectation values $S_f^R$ for the right QD, since $S_f^L$ can be obtained by the reflection with respect to zero ordinate axis. If parameter $V_2$ is changed from $V_2^{\text{min}} = -10$ meV to $V_2^{\text{max}} = 0$, the changes of the amplitude of the right potential well bottom are so large that even for $T \simeq 100$ ps the spin interchange process is non-adiabatic. In this case, we observe the spin exchange with accuracy $99.8 \%$ for $T_1^\pi = 90.4$ ps. The adiabatic process of full interchange of spins is observed for $T$ of the order of several hundred picoseconds. Therefore, the process, during which one of the
potential wells is entirely removed, is not advantageous. Therefore, we reduce the amplitude of changes of parameter $V_2$ by decreasing its maximum value $V_{2\text{max}}$. The useful results are obtained when raising the potential well bottom to $V_{2\text{max}} = -3 \text{ meV}$ [Fig. 14(a)]. However, we have to take into account that $V_{2\text{max}}$ can not be too small since then the potential well bottom is not raised high enough so that the probability of electron tunneling from the right to left QD is too small, which leads to the long spin swap duration time. For example, for $V_{2\text{max}} = -7 \text{ meV}$ [Fig. 14(c)] $T_{1\pi} = 713 \text{ ps}$. The shortest $\pi$ pulse time ($T_{1\pi} = 139 \text{ ps}$) has been obtained for $V_{2\text{max}} = -5.5 \text{ meV}$ [cf. Fig. 14(b)].

We conclude that the time changes of the confinement potential, which lead to asymmetry of QDs, result in the considerable shortening of the spin swap time if the amplitude of potential well changes is of the order of $\sim 5 \text{ meV}$.

VI. CONCLUSIONS AND SUMMARY

In the present paper, we have proposed a method for a theoretical quantitative description of spin swapping process in a double QD. Using this method we have estimated the spin swap duration time induced by the exchange interaction between electrons in vertically and laterally coupled QDs. The results obtained are also valid for quantum wires, which contain coupled QDs. The switching of the exchange interaction, considered in the present paper, was triggered by the corresponding time changes of the confinement potential profile. The time dependence of the confinement potential is of crucial importance in the process of spin interchange. We have shown that the confinement potential should change in time in a smooth manner and sufficiently slowly to ensure the adiabaticity of the spin swapping process.

We have also shown that the full spin interchange (with 100% accuracy) can hardly be achieved in the single spin swap process. In the coupled QD system, the full spin interchange occurs as a result of many-fold incomplete spin swaps. The most promising results, i.e., the shortest time of full spin swapping in the adiabatic process, have been obtained when smoothly changing in time the shape of the potential in the interdot region from the potential barrier to the potential well. However, it seems to be difficult to realize this process experimentally in the same nanodevice by changing the voltages applied to the gate.
FIG. 14: Expectation value $S^P_f \equiv S^R_f$ (solid curve) of $z$ spin component of the electron in the right QD and energy difference $\Delta E_{fi}$ (dotted curve) as functions of duration time $T$ of potential changes for asymmetric QDs. (a) $V_{2}^{\max} = -3$ meV, (b) $V_{2}^{\max} = -5$ meV, and (c) $V_{2}^{\max} = -7$ meV.

The results obtained for asymmetric QDs are rather disappointing: it turns out that the asymmetry of the confinement potential leads to the increase of the duration time of spin interchange, since in this case the full spin swapping is obtained after several incomplete spin swaps. Both in the vertically and laterally coupled QDs the spin swap duration time decreases if the range of the confinement potential becomes shorter. This leads to the conclusion that the size of the coupled QD system should be smaller than $\sim 100$ nm in order to obtain the sufficiently short spin swap time.
In summary, the exchange-interaction induced processes of spin rotation can serve as a mechanism to manipulate with electron spin qubits in coupled QDs. The spin swap duration time, which is several orders of magnitude shorter than the spin coherence time, can be obtained for sufficiently small coupled QDs, in which the gate voltages cause the smooth time changes of the confinement potential. A fabrication of the nanodevices, which satisfy these requirements, is a challenging task for the nanotechnology.

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