Coherent transient transport of interacting electrons through a quantum waveguide switch

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Abstract

We investigate coherent electron-switching transport in a double quantum waveguide system in a perpendicular static or vanishing magnetic field. The finite symmetric double waveguide is connected to two semi-infinite leads from both ends. The double waveguide can be defined as two parallel finite quantum wires or waveguides coupled via a window to facilitate coherent electron inter-wave transport. By tuning the length of the coupling window, we observe oscillations in the net charge current and a maximum electron conductance for the energy levels of the two waveguides in resonance. The importance of the mutual Coulomb interaction between the electrons and the influence of two-electron states is clarified by comparing results with and without the interaction. Even though the Coulomb interaction can lift two-electron states out of the group of active transport states the length of the coupling window can be tuned to locate two very distinct transport modes in the system in the late transient regime before the onset of a steady state. A static external magnetic field and quantum-dots formed by side gates (side quantum dots) can be used to enhance the inter-waveguide transport which can serve to implement a quantum logic device. The fact that the device can be operated in the transient regime can be used to enhance its speed.

Keywords: quantum transport, electron–electron interaction, quantum wire and dot

(Some figures may appear in colour only in the online journal)

1. Introduction

Various schemes associated with quantum computing have been proposed for quantum information storage and transfers, such as superconducting coplanar waveguide resonators and dual waveguide devices \[1, 2\]. A waveguide can be defined as a quantum wire in which the electron wave propagates in quantized modes without losing phase coherence at low temperature. Various proposals have been suggested to implement a semiconductor qubit \[3–5\]. Among these, one idea is to use parallel quantum waveguides with a coupling-window placed between them \[6\]. Tuning the window-coupling allows an electron wave interference between the waveguides and can switch the electron motion from one waveguide to the other \[7\].

The interference in the coupling-region specifies the possible types of qubit-operations: The electron wave can be transferred through either waveguide only (pass gate-operation). It can travel through both waveguides, split equally, implementing the so called square-root-of-Not (\(\sqrt{\text{NOT}}\)) operation \[8\]. It can switch totally from one waveguide to the other (Not-operation) \[9\]. Or it might switch from the first waveguide to the second one and then re-enter the first waveguide (CNot-operation) \[10\]. There are several parameters and phenomena that can modify the efficiency of the kind of qubit-operation that are mentioned above such as an
external magnetic field, the Coulomb interaction, electrostatic gate voltage, and the geometry of the coupling window between the waveguides.

Ferry’s group proposed magnetically switching transport in an asymmetric double quantum waveguide, in which an external magnetic field is applied to switch electron motion from one quantum-waveguide to the other \[7, 10\]. Another suggestion for a switching-qubit is to vary the length of the coupling window (CW) \(L_{CW}\). By varying the length of the CW, a maximum inter-waveguide tunneling can be found that increases the efficiency of the device \[7\]. Another approach for a switching-qubit is to define a saddle potential in the coupling window instead of a hard-walled well potential. The smoother potential defines a soft barrier between the two waveguides that washes out sharp resonance peaks, but decreases the efficiency of the qubit \[11\].

In the present work, we consider a double quantum waveguide (DQW) system with a CW between the waveguides in a static perpendicular magnetic field. Both ends of the DQW are connected to semi-infinite leads with an applied source-drain bias. A coherent electron transport is investigated in the system by using a non-Markovian master equation \[12–15\]. Smooth Gaussian potentials are used to define the barrier between the waveguides and the saddle-like CW potential. In addition, two dots are embedded in the waveguides close to the CW to enhance inter-waveguide transport. We will investigate coherent electron transport in the system by varying the CW length.

Our results lead to the following conclusions: First, we predict a maximum conductance at resonant energy-levels between the two waveguides. At a resonance energy, inter-waveguide forward scattering is enhanced and the charge density splits equally between the waveguides when two-electron states are activated in the transport. In this case, the DQW essentially works as a \(\sqrt{\text{NOT}}\)-operation qubit. The Coulomb interaction shifts the two-electron states above the selected bias window and the charge current density does not split equally between the waveguides at the resonance condition. The conductance peak is slightly reduced, but it remains a peak as backward scattering is absent. In the conductance dip that forms when the energy levels of the waveguides are not in resonance backward scattering is sharply increased with or without the Coulomb interaction. The strength of the Coulomb blocking can be varied by the geometry or the material of the system. The transient transport through two capacitively coupled parallel quantum dots has been investigated by Moldoveanu et al in a lattice model for a weaker Coulomb interaction than is assumed here \[16\]. Second, an increased external static magnetic field favors inter-waveguide backward scattering leading to a decrease in the efficiency of the qubit. These results make clear the difference to previous schemes where variable magnetic field was used to control the switching between asymmetric waveguides \[6, 11\]. Third, the influence of the side quantum dots (QDs) can be to induce more energy-levels into resonance, and consequently increase the coupling between the waveguides and enhance the inter-waveguide transport mechanism. In the presence of side QDs, a Not-operation qubit can be realized. The efficiency is slightly decreased in the presence of side QDs because inter-waveguide backward scattering is also enhanced. At high magnetic field such that the effective magnetic length is comparable to the radius of the dots, variation of length of the CW does not influence the electron transport characteristics further as the electrons tend to be localized in the side dots. Thus, the net charge current is extremely suppressed.

The paper is organized as follows: in section 2, we present the model describing the window-coupled double waveguide system based on a quantum master equation (QME) approach. Section 3 presents our numerical results and discussion. Concluding remarks are addressed in section 4.

2. Model and theory

In this section, we present a method for calculating the transient ballistic transport in the coupled-window quantum waveguides without and with side QDs. We consider two laterally coupled waveguides connected to two semi-infinite wires from both ends. The lower waveguide (control) and the upper waveguide (target) are coupled through a CW with length \(L_{CW}\) as is schematically shown in figure 1. We consider a control-QD (QDC) and a target-QD (QDT) embedded, respectively, in the control- and the target-waveguide near the CW. The total system is exposed to an external magnetic field in the \(z\)-direction \(B\hat{z}\). The system is designed such that the electrons are injected from the left lead to only the control-waveguide (red arrow). Our aim here is to study the charging of, and the output from each of the waveguides by varying the CW length \(L_{CW}\) (blue dashed arrow). In addition, the effects of the Coulomb interaction, the external magnetic field, and the side dots on the conductance will be explored.

The DQW is a two dimensional system with hard-wall confinement in the contact area to the external leads at \(x = \pm L_x/2\), with \(L_x\) the length of the waveguides in the transport direction, but parabolic confinement in the \(y\)-direction. The single-electron Hamiltonian of the DQW including side QDs in an external magnetic field is

\[
h_{S} = \frac{(\mathbf{p} + e\mathbf{A}_{\text{ext}})^2}{2m^*} + V_{c}(x, y) + V_{DW}(\mathbf{r}) + V_{QD}(\mathbf{r}),
\]

where \(\mathbf{p}\) indicates the canonical momentum, \(\mathbf{A}_{\text{ext}} = (0, -By, 0)\) is vector potential of the external magnetic field.
field, and \( m^* \) is the effective mass of an electron. The confining potential of the DQW is \( V_c(x, y) = V_c(x) + V_c(y) \), where \( V_c(x) \) stands for a hard-wall confining potential and \( V_c(y) = \frac{1}{2} m^* \Omega_0^2 y^2 \) refers to a parabolic confining potential with characteristic energy \( \hbar \Omega_0 \). The Gaussian potential barrier between the two waveguides is presented by \( V_{GB}(r) \) and the side QDs are defined by \( V_{QDs}(r) \). We diagonalize the Hamiltonian of the central system presented in equation (1) using a basis \([15]\) built with the eigenfunctions of the first two terms of (1) to find the single-electron energy spectrum \( E_n \), where \( n \) is the composite quantum number of a single-electron state. The eigenfunctions of (1) are consequentially used to introduce a reduced density operator (RDO), by tracing out the electron-lead coupling up to second order in the dissipation range parameters in the non-Markovian QME is derived keeping terms for the system-lead coupling up to second order in the dissipation term, i.e. the integral kernel. The explicit expressions for \( \Delta_{l}^{(q)}(q) = [E_n - \varepsilon(l)(q)] \) the energy affinity between states in the leads and the central system \([13]\), and \( \sigma \) is a skewing parameter that shifts the weight of the coupling to the lower waveguide for an appropriate value stated below.

Before coupling the central system to the external leads a many-electron Fock space with sectors for 0-, 1-, 2-, and 3- electron states is constructed for the first term of the total Hamiltonian \( (2) \). This noninteracting many-electron (ME) basis is then used to diagonalize the Hamiltonian of the central system, the first two terms in equation (2) obtaining a Coulomb interacting ME basis. This procedure of exact numerical diagonalization of a ME Hamiltonian in a truncated basis is used in order to capture correlation effects that are of crucial importance in small systems with few electrons \([17]\). The diagonalization and the truncation procedure together with the necessary basis changes is described elsewhere \([18,19]\).

2.1. DQW connected to leads

We connect the DQW to two leads that act as electron reservoirs via a contact region. The central system and the leads have the same confinement energy and the same strength of the perpendicular magnetic field. The total Hamiltonian of system is then

\[
H(t) = \sum_{n} E_n d_n^{\dagger} d_n + \frac{1}{2} \sum_{n \neq m} (V_{Coul}) d_n^{\dagger} d_m^{\dagger} d_m d_n \\
+ \sum_{l=L,R} \int dq \, \bar{\epsilon}(q) c_{l}^{\dagger} c_{l} \\
+ \sum_{l=L,R} \chi(t) \int dq \left[ c_{l}^{\dagger} T_{q}^{l} d_n^{\dagger} d_n + d_{n}^{\dagger} T_{q}^{l} c_{l} \right].
\]

(2)

The first term describes the central system, where \( E_n \) is the single-electron energy of state \( |n\rangle \) and \( d_{n}^{\dagger} (d_{n}) \) denotes the electron creation (annihilation) operator respectively. The second term is the mutual Coulomb interaction between electrons in the central system. The third term of the Hamiltonian denotes the semi-infinite \( lth \) lead, where \( l \) refers to the left \( L \) or the right \( R \) lead with the dummy index \( q \) representing the ‘momentum’ of an electron in the lead and its subband index \([15]\), \( \bar{\epsilon}(q) \) is the single-electron energy spectrum in lead \( l \), and the electron creation and annihilation operator in lead \( l \) are \( c_{l}^{\dagger} \) and \( c_{l} \), respectively.

The last term is the time-dependent coupling Hamiltonian that connects the central system, the DQW to the external leads via a coupling tensor \( T_{q}^{l} \), and \( \chi(t) \) dictates the time-dependence of the coupling. The coupling Hamiltonian describes the transfer of an electron between the single-electron state of the central system \( |n\rangle \) and the single-electron state of the leads \( |q\rangle \)

\[
T_{q}^{l} = \int dr d\tau \bar{\psi}_{q}^{l}(r, \tau) \bar{g}_{q}^{l}(r, \tau) \bar{\psi}_{n}^{l}(r).
\]

(3)

\( \bar{\psi}_{q}^{l}(r) \) and \( \bar{\psi}_{n}^{l}(r) \) are the corresponding single-electron wave functions of the DQW system and the leads. \( g_{q}^{l}(r, \tau) \) is a nonlocal coupling kernel

\[
g_{q}^{l}(r, \tau) = g_{0}^{l} \exp \left\{ -\delta_{l}^{q} (x - x')^{2} - \delta_{l}^{q} (y - y' - \alpha)^{2} \right\} \exp \left\{ -\Delta_{l}^{q}(q)/\Delta \right\}.
\]

(4)

where \( g_{0}^{l} \) is the coupling strength, \( \delta_{l}^{q} \) and \( \delta_{l}^{q} \) are the coupling range parameters in the \( x \)- and \( y \)-direction, respectively, \( \Delta \) controls together with \( \Delta_{l}^{q}(q) = [E_n - \varepsilon(l)(q)] \) the energy affinity between states in the leads and the central system \([13]\), and \( \alpha \) is a skewing parameter that shifts the weight of the coupling to the lower waveguide for an appropriate value stated below.

2.2. QME and Charge current density

In this subsection, we present how the time evolution of the open system is calculated in order to study time-dependent transport properties within the QME approach \([20]\). The time evolution of our system obeys the quantum Liouville–von Neumann equation

\[
\frac{dW(t)}{dt} = -i \frac{\hbar}{\hbar} [H(t), W(t)].
\]

(5)

where \( W(t) \) is the density operator of the total system. In order to focus our attention on the open central system we project the dynamical evolution of the whole system on the central system introducing a reduced density operator (RDO), by tracing out all variables of the leads, \( \rho(t) = \operatorname{Tr}_{R} \rho(t) \), with the initial condition \( W(t < t_0) = \rho_{d} \rho_{d}^{\dagger} \), where \( \rho_{d} \) and \( \rho_{d}^{\dagger} \) are the density operators of the isolated left and the right leads, respectively.

The density operator of the isolated \( lth \) lead can be defined as \( \rho_{l} = e^{-\beta(H_l - \mu_{N_l})}/\operatorname{Tr}_l [e^{-\beta(H_l - \mu_{N_l})}] \), where \( \beta = 1/(k_{B} T) \) with \( k_{B} \) the Boltzmann constant, and \( N_{l} \) is the number of electrons in the leads. The projection formalism delivers an equation of motion for density operator of the central system, the quantum master equation \([21,22]\)

\[
\frac{d\rho(t)}{dt} = -i L_{S} \rho(t) + \int_{t_0}^{t} dt' \mathcal{K}(t, t') \rho(t'),
\]

(6)

where \( L_{S}\cdot [H_{S}, \cdot]/\hbar \) is the Liouvillian with respect to the time-independent Hamiltonian \( H_{S} \) of the central system and \( \mathcal{K}(t, t') \) is an integral kernel describing the dissipative time-dependent coupling to the leads \([15,22]\). In our approach the non-Markovian QME is derived keeping terms for the system-lead coupling up to second order in the dissipation term, i.e. the integral kernel. The explicit expressions for the kernel have been published elsewhere \([12,13,18]\).
QME is appropriate to describe sequential tunneling between the central system and the leads in the weak coupling limit. The structure of the QME as an integro-differential equation with memory effects implies higher order sequential tunneling, but not cotunneling processes.

In order to explore the switching processes between the waveguides, we define the net charge current with respect to the external leads as

\[ I_Q(t) = I_L(t) - I_R(t), \]

where \( I_L(t) \) refers to the partial current from the left lead into the control-waveguide and \( I_R(t) \) indicates the partial current into the right lead from both waveguides [23]. In equation (7) the negative sign in front of \( I_Q \) is since a positive charge current is defined as the current into the central system, but the currents \( I_L \) and \( I_R \) are defined positive from left to right.

To monitor the dynamic evolution of the electrons inside the central system, we calculate the expectation value of the local charge current density operator

\[ \hat{J}(\mathbf{r}, t) = \text{Tr}(\hat{\rho}(t)\hat{J}(\mathbf{r})), \]

where operator is defined by

\[ \hat{J}(\mathbf{r}) = \sum_{nn'} \left( \frac{e\hbar}{2m^*} \left[ \psi_n^S(\mathbf{r}) (\nabla \psi_n^S(\mathbf{r})) - (\nabla \psi_n^S(\mathbf{r})) \psi_n^S(\mathbf{r}) \right] \right. 
\]

\[ \left. + \frac{\beta^2}{m^*} A_{\text{ext}}(\mathbf{r}) \psi_n^S(\mathbf{r}) \psi_n^S(\mathbf{r}) \right) d_{\mathbf{r}} d_{\mathbf{r}}'. \]

In the following, we shall demonstrate the influence of the length of the CW, external magnetic field, and the side QDs on the coherent electron transport through the system and the charge current density in the central system in order to investigate inter-waveguide forward and backward scattering processes.

3. Results

In this section, we shall discuss our numerical results of the ballistic transport properties of the double waveguide system made of a GaAs semiconductor with the electron effective mass \( m^* = 0.067m_e \). The length of the central system is \( L_\text{c} = 300 \text{ nm} \) which is assumed to be much smaller than the phase coherent length \( L_\phi \). At low temperature \( T \sim 0.1\text{--}2.0 \text{ K} \), the phase coherent length of a semiconductor based (GaAs/Al\textsubscript{1-x}Ga\textsubscript{x}As) electron waveguide is \( \sim (30 - 40) \times 10^3 \text{ nm} \) [24]. The coherence in the electron transport through the double waveguide is an essential requirement for constructing a qubit. For the system placed in a static or vanishing external magnetic field, its length scales can be characterized by the effective magnetic length \( a_w = \frac{\hbar}{(m^* \sqrt{\omega_c^2 + \Omega_0^2})} \), where the cyclotron frequency \( \omega_c = \frac{eB}{m^*} \), and the transverse confinement energy \( \hbar \Omega_0 = 1.0 \text{ meV} \). Numerically the effective magnetic length can be expressed as

\[ a_w = \left( \frac{\hbar}{m^* \Omega_0} \right)^{1/2} \left( \frac{1}{1 + (eB/(m^* \Omega_0))^2} \right)^{1/4} = 33.74 \text{nm}. \]

For the case of a vanishing magnetic field we assign to it a finite very small value in order to avoid numerical problems caused by exact degeneration of states with opposite spin.

We assume the central system is connected to the external leads acting as electron reservoirs with chemical potential in the left (right) lead \( \mu_L = 4.0 \text{ meV} \) \( \mu_R = 3.0 \text{ meV} \), implying an applied potential difference or bias window \( \Delta \mu = eV_\text{bias} = 1.0 \text{ meV} \). Furthermore, the temperature of the leads is fixed at \( T = 0.5 \text{ K} \). The potential of the double quantum waveguides is shown in figure 2 with the side QDs located near the CW in order to influence the inter-waveguide transport. In the following, we shall explore the influence of the CW and the side QDs on the charging of and the transport through the system.

3.1. DQW without side QDs

We start by considering an ideal window-coupled DQW without side QDs. The injected electrons are assumed to come from the left lead only into the control-waveguide. This asymmetry in the injection is accomplished with the choice of the skewing parameter \( \alpha = 4.0a_w \) defined in equation (4) and only used for the left lead. In addition, we assume the two quantum waveguides to be of the same width. They are separated by a Gaussian potential barrier, in which the control- and target-waveguide interact with each other via a CW. The DQW system is described by

\[ V_{\text{DQW}}(\mathbf{r}) = V_\text{b}(\mathbf{y}) + V_{\text{CW}}(x, y), \]

where \( V_\text{b}(\mathbf{y}) \) is the barrier potential between the quantum waveguides defined as

\[ V_\text{b}(\mathbf{y}) = V_0 \exp(-\beta_0^2 y^2), \]

with \( V_0 = 18.0 \text{ meV} \) and \( \beta_0 = 0.03 \text{ nm}^{-1} \), as well as a CW potential

\[ V_{\text{CW}}(x, y) = -V_0 \exp(-\beta_x^2 x^2 - \beta_y^2 y^2), \]

with \( V_0 = 18.0 \text{ meV} \) and \( \beta_x = 0.03 \text{ nm}^{-1} \) implying a barrier width \( W_0 \approx 66.5 \text{ nm} \) for the first subband which can prevent electron tunneling between the waveguides through the barrier. \( \beta_x \) is a parameter that determines the CW length. Thus, the length of the CW can be defined as \( L_{\text{CW}} = 2/\beta_x \).
We begin to analyze the results by presenting figure 3 showing the net charge current $I_Q$ as a function of the CW length $L_{CW}$ at $t = 200 \text{ ps}$ and $B = 0.0 \text{ T}$ for noninteracting electrons (solid blue), and for Coulomb interacting electrons (dashed red). The Coulomb interaction reduces the overall current slightly. The oscillations in the net charge current are indicative of a possible charge transport between the control-waveguide and the target-waveguide. The oscillations give rise to a peak and a dip in the net charge current at $L_{CW} \approx 40$ and $110 \text{ nm}$, respectively. Similar oscillation features have been found by Zibold et al and Gong et al in the absence of the Coulomb interaction, and put in relation to energy-dependent inter-waveguide transmission in the quantum regime. They pointed out that the inter-waveguide transmission can be enhanced when the energy levels of the coupled-waveguide system achieve a resonance conditions for a specific CW length [25, 26].

To find resonant energy levels of the two waveguides, we present figure 4 showing the many-electron (ME) energy spectra for the DQW system as a function of the CW length at $B = 0.0 \text{ T}$ including one-electron states (1ES, red dots) and two-electron states (2ES, blue dots). The ME spectrum without the mutual Coulomb interaction is displayed in figure 4(a), but the interacting spectrum in figure 4(b). The only difference between the energy spectra is that the Coulomb interaction has raised most of the 2ES well above the bias window in figure 4(b). The 2ES do not contribute to the electron transport in the presence of the Coulomb interaction because they are far from the chosen bias window, leading to a decrease in the net charge current.

In the case of no coupling window the two waveguides are only weakly coupled through tunneling and the Coulomb interaction [16] leading to nearly degenerate states. When the length of the CW $L_{CW}$ is increased we thus find: First, the energy spectrum decreases monotonically [25]. Second, the near degeneration of the energy-levels is lifted. As a result, this leads to a crossover of energy levels (resonance) between the control- and target-waveguide at $L_{CW} \approx 40 \text{ nm}$ (left green rectangular) which corresponds to the current-peak in figure 3. The resonance between the waveguides enhances the inter-waveguide electron transport and increases the net charge current. The splitting of the energy levels increases at a higher coupling window length such as $L_{CW} \approx 110 \text{ nm}$ (right green rectangular) indicating weaker resonances and more back-scattering inter-waveguide transport.

To further explain the peak and the dip in the net charge current, we present the charge current density at $B = 0.0 \text{ T}$ and $t = 200 \text{ ps}$ in figure 5 for the noninteracting system (left panels) and the interacting one (right panels) for the peak and the dip net charge current at $L_{CW} \approx 40 \text{ nm}$ (a)–(b) and $110 \text{ nm}$ (c)–(d), respectively, shown in figure 3. The dynamical evolution occurring in the CW exhibits different types of quantum logic gates actions in our system. In the absence of the Coulomb interaction the charge current density shown in figure 5(a) reveals the following electron motion in the waveguides at $L_{CW} \approx 40 \text{ nm}$: The charge is injected into the control input, then it exits equally from both control and target output. In the DQW the charge exhibits partial inter-waveguide forward scattering which is similar to the condition of having equal electron transmission ratio in the double waveguides at a resonance [25]. The DQW here can be defined as a beam splitter or a $\sqrt{\text{NOT}}$-operation quantum logic gate ($\sqrt{\text{NOT}}$-operation qubit) [8]. Figure 5(c) shows the charge current density in the current-dip at $L_{CW} \approx 110 \text{ nm}$: The charge current density switches from the control- to the target-waveguide with a small ratio of inter-waveguide backward scattering. The charge switching property of the system here can be introduced as a Not-operation quantum logic gate [11].

The net charge current is suppressed here due to the presence of...
Figure 5. Charge current density in the DQW at $t = 200$ ps without (left panels) with (right panels) electron–electron Coulomb interaction in the peak and the dip net charge current at $L_{CW} \approx 40$ nm ($a$, $b$) and 110 nm ($c$, $d$), respectively, shown in figure 3 in the case of $B = 0.0$ T (blue solid). The effective magnetic length is $a_w = 33.72$ nm. The effects of a static external magnetic field are presented in figure 6(a) which shows the net charge current $I_Q$ as a function of the CW length $L_{CW}$ at $t = 200$ ps in the presence of the Coulomb interaction for different values of the magnetic field $B = 0.0$ T (blue solid), 0.1 T (green dashed), and 0.2 T (red dashed). For comparison we display in figure 6(b) the left and the right currents $I_L$ and $I_R$ at $B = 0.0$ T and the same point in time. The comparison shows that at $t = 200$ ps the system is still in the charging phase, but the overall shape of the currents is similar. The role of the external magnetic field in our system is different from the role it plays in many models where researchers have applied a variable magnetic field to switch between different processes in the waveguides. The external magnetic field has been used to switch an electron from one quantum-waveguide to the other one in a window coupled double waveguide [6]. In our model, we observe switching in the absence of a magnetic field, i.e. for $B = 0.0$ T. In the presence of stronger magnetic field, the electron motion is affected by the Lorentz force in which the electrons tend to a circular motion in both control-
target-waveguides increasing the electron dwell-time in the system. The inter-waveguide backward scattering is enhanced while a suppression in the inter-waveguide forward scattering is observed in the presence of strong magnetic field. Therefore, the net charge current decreases as shown in figure 6(a) in the case of $B = 0.1$ T (green dashed) and 0.2 T (red dotted).

To understand the reasons for the suppression of the net charge current in the presence of a higher magnetic field, we show in figure 7 the charge current density at magnetic field $B = 0.2$ T in the peak at $L_{\text{CW}} \simeq 40$ nm (a) and the dip at $L_{\text{CW}} \simeq 110$ nm (b), of the net charge current shown in figure 6(a) (red dotted). An obvious explanation is: The perpendicular magnetic field reduces the effective barrier height and increases the splitting of the energy levels at the crossover energy or resonance energy (not shown). Therefore the charge current density does not split equally between the waveguides or localize in only control-waveguide, but rather an inter-waveguide backward scattering is observed due to the Lorentz force, consequently the net charge current is suppressed and we observe formation of edge states in each of the waveguides. In this case the double-waveguide system does neither play the role of a NOT-operation, nor a Not-operation qubit in the presence of a higher magnetic field.

3.2. DQW with side QDs

In this section, we consider a side dot embedded in each waveguide to enhance the inter-waveguide transport as is shown in figure 2. The dots to the sides of the CW may be expected to increase the dwell time in the coupling region and change the resonance condition between the two waveguides. The window-coupling DQW potential with side QDs is described by

$$V_{\text{DQW}}(x, y) = V_{\text{B}}(y) + V_{\text{CW}}(x, y) + V_{\text{Dot}}(x, y),$$

where $V_{\text{Dot}}(x, y)$ is the side dot potential that is defined as

$$V_{\text{Dot}}(x, y) = -V_0 \exp(-\gamma_x x^2 - \gamma_y y^2),$$

with $V_0 = 8.0\text{ meV}$, $\gamma_0 = 5.6a_w$, and $\gamma_x = \gamma_y = 0.04\text{ nm}^{-1}$ implying the radius of each side-dot $R_{\text{Dot}} \approx 25$ nm.
localized electrons in the dots as the effective magnetic length is close to the dot radius. We present figure 8(b) to compare the left partial current (pink dotted), the right partial current (blue dashed) to the net charge current (blue solid) at $B = 0.0$ T. It can be seen that the partial currents have the same oscillation features and develop a peak and a current dip at the same CW lengths.

In order to understand the characteristics of net charge current, we refer to the energy spectrum and the charge current density. Figure 9 shows the ME-energy versus the window-coupling length at $B = 0.0$ T including one-electron states (1ES, red dots) and two electron states (2ES, blue dots). It should be noted that more energy levels get into resonance at $L_{CW} \simeq 40$ nm (left green rectangular) in the presence of the side-gate dots which indicates a stronger coupling between the control- and the target-waveguide. At larger CW length such as $L_{CW} \simeq 110$ nm (right green rectangular), the energy splitting increases resulting in different electron motion in the system.

The electron motion in the waveguides is completely changed in the presence of the side QDs. Figure 10 displays the charge current density at $B = 0.0$ T and $L_{CW} \simeq 40$ nm (a) and current-dip at $L_{CW} \simeq 110$ nm (b) shown in figure 8(a) (blue solid). In figure 10(a) the charge current density is displayed for $L_{CW} \simeq 40$ nm. The charge in the control-waveguide is switched to the target-waveguide, thus a NOT-operation is realized. Comparing to the system without side-gate dots in figure 5(b), the inter-waveguide transport is enhanced due to a stronger coupling between the waveguides at the resonant-energy levels at $L_{CW} \simeq 40$ nm. Therefore, the side QDs play an essential role to convert the qubit operation in the system. In addition, the inter-waveguide backward scattering here reduces the efficiency of the qubit. As we mention previously, the existence of inter-waveguide backward scattering might be due to symmetric waveguides while the backward-scattering in an asymmetric waveguides can be avoided or reduced [27]. Figure 10(b) shows the charge current density at the current-dip $L_{CW} \simeq 110$ nm. The incoming charge into the control-waveguide encounters the control-QD leading to electron back-scattering in the control-waveguide, thus the net charge current is suppressed forming a current-dip. We can say that the side-dots play a scattering role in the waveguides in such way that the electrons partially get forward- and backward-scattered in and into both waveguides.

By tuning the external magnetic field to a higher value, $B = 0.2$ T, the electrons manifest different dynamical motion in the waveguides. In figure 11 the charge current density is shown in this stronger magnetic field $B = 0.2$ T at the CW length $L_{CW} \simeq 40$ (a) and $110$ nm (b). The graphs show circular motion, edge states, in each waveguide. Figure 11(a) shows the current charge density at $L_{CW} \simeq 40$ nm, the electrons in the control-waveguide do not totally switch to the target-waveguide, because of increased energy level splitting at the resonance-energy in the presence of higher magnetic field. In addition, charge is partially localized in the control-QD as the effective magnetic length approaches the radius of the quantum dot $a_w \simeq R_{DQW}$. In figure 11(b), the electrons are mostly localized in the control-QD corresponding to the current-dip at $L_{CW} \simeq 110$ nm. This electron localization is expected because at a larger CW length the splitting of energy levels increases and leads to suppression of the inter-waveguide scattering. Therefore, the electrons move along the control-waveguide and get localized in the control-QD at this higher magnetic field.

4. Concluding remarks

We have presented a model for a double quantum waveguide in an external perpendicular magnetic field. The waveguides are coupled via a window to facilitate inter-waveguide transport. The DQW is weakly connected to two leads with different chemical potentials in which a non-Markovian master equation is utilized to investigate coherent switching in electron transport between the waveguides for the implementation of quantum logic gates.

By tuning the length of the CW, we have demonstrated two important physical characteristics of the waveguide system: resonant energy-levels and oscillations in the net charge current. The resonant energy levels indicate a strong coupling between the DQW and the oscillations reflect that the resonance conditions are governed by the length of the CW.

In the case of an ideal non-Coulomb-interacting window-coupled DQW system without a magnetic field, the charge current density splits equally between the waveguides at resonant energies, therefore, the net charge current reaches its maximum value and the waveguide system works as a √NOT-operation qubit. The splitting of charge is due to the contribution of 2ES to the electron transport. The inter-waveguide forward scattering leading to the charge splitting at resonant levels is not seen without including two-electrons states in the model. It is strongly reduced by the Coulomb
interaction that lifts the 2ES out of the group of active states in the transport. This aspect can though be controlled by the exact system size and design as the Coulomb blocking is reduced in a larger system, or by using a higher bias window or different material. Our model only includes sequential tunneling of electrons from the leads to the system, but the addition of higher order tunneling would only increase the importance of the many-electron structure and the Coulomb interaction on the operation of the system. The charge splitting process can not be obtained in the presence of an external magnetic field because inter-waveguide backward scattering is enhanced leading to decrease in the efficiency of the logic gate.

Two parallel waveguides or dots can be coupled in many distinct ways. Moldoveanu et al. [16] attain it capacitively, while Zibold et al. [25] fine tune the system by tunneling coupling. Both groups stress the strong dependence of the device on its exact geometry, and the structure and correlations of many-electron states that are active in the transport. Our investigation with a different coupling scheme for the waveguides confirms this sensitive dependence.

In addition, we have embedded a quantum dot in each waveguide to enhance inter-waveguide transport. In the presence of side QDs, more energy-levels of the two waveguides are brought into resonance which can lead to a stronger coupling between them. Effectively, the dots can increase the density of states around the resonant transport states. The electrons from the control-waveguide switch to the target-waveguide indicating a Not-operation qubit. But on the other hand, the side-dots can also increase the inter-waveguide backward scattering and reduce the qubit efficiency. In a stronger external magnetic field, the electrons get localized in the dots as the effective magnetic length is comparable to the radius of the side dot. In this case, the geometry of the CW does not affect the electron transport anymore. The external magnetic field here is considered static, but one more variable to influence the transport is the shape of the side QDs in the waveguides. We show that the varying of the CW length between the waveguides can specify the type of the qubit logic gate to be implemented by the DQW system.

A comparison between the charge current and the left and right currents in figures 6 and 8 at the time point in time \(t = 200\) ps shows that the system has not completely reached a steady state yet, but the sought after function of the DQW is already present in the late transition regime. This is important. The exact steady state takes a long time to reach due to a low rate of charging for 2 and 3 electron states. Valuable time can be gained by operation of a device in the transition regime.

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