Non-coherent transport in serial/parallel quantum dots in self-consistent field regime

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Abstract—In this work we introduce a simple phenomenological model of the conduction between a couple of serial or parallel quantum dots. This model is extended to an arbitrary of number and to a square array of quantum dots. The local potential is computed taking into account the net charge at the quantum dot. Master equations are presented for total current and charge number \( N \) considering both the local potential and the local current. Interesting results are reported, namely the Negative Differential Resistance for serial quantum dots as well as resonant conductance. A paradoxical asymmetric current appears for the parallel case due to the capacitive coupling.

Index Terms—1. Modeling and simulation of systems and properties of matter at the nanoscale: I-V characteristic, non-coherent transport, quantum dot, rate equations, serial/parallel, self-consistent field,

I. INTRODUCTION

Single-electron devices are currently conceived to take advantage of the tunnel current between quantum states belonging to nano-scale particles [1, 2]. Up to now, the study of such tunnel current has been carried out using simple phenomenological (i.e. rate equation) approaches [3, 4] or more complex quantum theories including NEGFF [6, 7, 8], transfer Hamiltonian approach [9, 10]. Concerning the first case, simple rate equations give us useful and intuitive approaches. The effects of local potential due to selfcharge can be also included by means of a couple of master equations [4]. On the other hand, quantum study of non-equilibrium schemes results in a more complex approach. Until now research has been mostly concentrate on single dots, but rapid progress in microfabrication technology has made possible the extension to couple-dot system with aligned levels [11, 12, 13]. A very interesting result was obtained by Shanguan et al [14]. These authors describe the transport in one-dimensional quantum dot array by using many body approaches. Up to now the only computation of transport in an array of quantum dots was done by Carreras et al [15] but no local potential due to selfcharge was included.

In this work we present a semiclassical approach based on non-coherent rate equations [4, 5] that is extended from two to an arbitrary number of quantum dots either serial, parallel and square array. The whole array of quantum dots has been treated as a separated quantum system. Then the leads (reservoirs) were incorporated through the rate equations for single dots. In this approach, the coherent effects does not appear and the probabilities of finding an electron in two different dots are coupled by some rate \( \gamma \). This treatment is called classical (non-coherent) approach. The effects of the local potential are computed within the selfconsistent field (SCF) regime. The SCF method is widely used because the exact method based on a multielectron picture is usually impossible to implement. As example we compute the total current and charge transport distribution \( N \) for two parallel and serial quantum dots (Fig. 1). The model has been implemented in a home made program in MATLAB code that gives us the current voltage characteristics, the differential conductance and the population inside each dot for a range of applied voltage. Some known results as Negative Differential Resistance (NDR) and resonant conductance are well explained. Moreover we found that the coupling of parallel quantum dots leads to an asymmetric conductance.

II. THEORETICAL MODEL

When we have two leads coupled to a system with no voltage applied between them, the system is in equilibrium with a common electrochemical potential. In this equilibrium state, the average number of electrons in any energy level is given by the Fermi function:

\[
f(E) = \frac{1}{1 + e^{(E - \mu)/k_B T}}
\]
An applied bias voltage $V_G = \frac{\mu_L - \mu_R}{e}$ drives the system out of equilibrium. Each contact tries to bring the system into equilibrium with itself. The system is thus forced into a balancing act between two reservoirs with different population. Then, the average number of electrons $N$ inside the system will be something between $f_L$ and $f_R$. We can not use the Fermi distribution function anymore to calculate the carrier in the system. In order to obtain the new distribution function we need to apply a master equation.

Large classes of stochastic processes in physics are characterized by the Markov property. A Markov process is defined as a stochastic process where the transitions in successive time intervals are statistically independent or in other words the transition between two states does not take into account the prehistory of the process. Defining the conditional probability for a transition from state $\nu$ to a state $\nu'$ at times $t$ and $t'$, respectively: $p(\nu' | \nu, t)$, the Markov property can be derived which is called Master equation

\[
\frac{\partial}{\partial t} P_\nu(t) = \sum_{\nu' \neq \nu} [W_{\nu\nu'} P_{\nu'}(t) - W_{\nu'\nu} P_\nu(t)]
\]

For the occupation probabilities $P_\nu$ of state $\nu$. The $W_{\nu\nu'}$ are the rates for the transition from state $\nu'$ to state $\nu$ per unit time. This equation can be interpreted quite easily: the first term describes the gain of state $\nu$ due to transitions from other states $\nu'$ and the second term is due to the loss from state $\nu$ into other states $\nu'$ (therefore the minus sign).

The Qd system is described by a Fock state $(\nu_1, \ldots, \nu_N)$ with the occupation numbers $n_i \in \{0, 1\}$ and N single particle states. The single particle states $i$ correspond to the energy level in different Qds. The time evolution of these occupation probabilities $P_\nu$ is determined by sequential or single electron tunnelling of an electron into or out the source/drain contact. We use the Master equation and write it in the compact matrix form as

\[
\dot{P} = MP
\]

The diagonal elements $M_{\nu\nu}$ of the transition matrix $M$ contains the loss term and the non-diagonal elements correspond to the gain term. In order to fulfill the probability normalization $\sum_\nu P_\nu = 1$ the sum of the matrix elements in a column of $M$ has to be zero.

The central quantity of interest is the mean current through the respective Qd system. The current for the right and left lead, respectively, in the steady state are

\[
I = \sum_\nu P_{\nu\nu}W_R = \sum_\nu P_{\nu\nu}W_L
\]

where the sum is extended over states $\nu$ in which the dot adjacent to the lead is occupied.

The two systems that we have studied are in Fig. 1. For simplicity we assume only a single state with energy $\epsilon$ on each dot is important and the double occupancy of the dot can be ignored. $\gamma_i$ are the tunnelling rates.

![Figure 1: System composed of quantum dot 1 and 2, and two leads. $\gamma_i$ are the tunneling rates of dot 1 and 2, and two leads.](image)

**A. SERIAL QUANTUM DOT**

For the single particle levels in Qd1 and Qd2 it is assumed $\mu_L - \epsilon >> k_BT$ and $\epsilon - \mu_R >> k_BT$ such that $f_\nu \approx 1$ and $f_R \approx 0$. With $P = (P_{(0,0)}, P_{(1,0)}, P_{(0,1)}, P_{(1,1)})$ the first sub-index indicates the occupancy of the Qd1 and the second sub-index the occupancy of the Qd2. The matrix $M$ explicit reads

\[
\begin{pmatrix}
-\gamma_1 & \gamma_3 & 0 & 0 \\
0 & -(\gamma_1 + \gamma_2 + \gamma_3) & \gamma_2 & 0 \\
\gamma_1 & \gamma_2 & -\gamma_2 & \gamma_3 \\
0 & \gamma_1 & 0 & -\gamma_3
\end{pmatrix}
\]

We are interested in the steady state $MP = 0$, then we need to diagonalize the $M$ matrix and find the eigenvectors associated with the eigenvalues 0. But we can obtain the same result if we use a rate equation. Rate-equation calculation consider the dynamic behavior of
the number of electron $n_i$ at each quantum dot balanced with the incoming and outcoming flux either from a partially filled ($f_L, R$) lead or other quantum dot ($n_j$) [16, 17].

Rate-equation for serial Qd can be written as follow:

$$\frac{dn_1}{dt} = \gamma_1 f_L + \gamma_2 n_2 - (\gamma_1 + \gamma_2)n_1 \quad (6)$$

$$\frac{dn_2}{dt} = \gamma_3 f_R + \gamma_2 n_1 - (\gamma_2 + \gamma_3)n_2 \quad (7)$$

where coefficients $\gamma_i$ are the transition probabilities as described in Fig. 1a. $f_L$ and $f_R$ are the Fermi-Dirac distribution functions in the electrodes. Density of states at electrodes is considered constant in the energy range that transport takes place. $n_i$ is the average number of electrons at $i$-th Qd.

The steady state solution of (6) and (7) give as the expression

$$n_1 = \frac{\gamma_1(\gamma_2 + \gamma_3)f_L + \gamma_2 \gamma_3 f_R}{(\gamma_1 + \gamma_2)(\gamma_2 + \gamma_3) - \gamma_2^2} \quad (8)$$

$$n_2 = \frac{\gamma_3(\gamma_2 + \gamma_1)f_R + \gamma_2 \gamma_1 f_L}{(\gamma_1 + \gamma_2)(\gamma_2 + \gamma_3) - \gamma_2^2} \quad (9)$$

Besides the outcoming net flux from Qd2 to the right lead is

$$I = (-q)\frac{\gamma_3}{h}(f_R - n_2) \quad (10)$$

That help us to consider a n-independent way to express the current using (9) and (10)

$$I^* = \frac{\gamma_1 \gamma_2 \gamma_3}{\gamma_1 \gamma_2 + \gamma_1 \gamma_3 + \gamma_2 \gamma_3}(f_L - f_R) \quad (11)$$

The expression of the current can be physically interpreted quite easily: the three barriers in the Qd stack act as a serial connection of resistors corresponding to the inverse of tunnelling rates $R_{total} = \sum_i R_i$.

**B. PARALLEL QUANTUM DOT**

Using the same assumption as before the M-matrix can be written as

$$P = \begin{pmatrix} -2\gamma_1 & \gamma_3 & 0 \\ \gamma_1 & -(\gamma_1 + \gamma_3) & 0 \\ 0 & \gamma_1 & -2\gamma_3 \end{pmatrix}$$

with $P = (P_{(0,0)}, P_{(1,0)}, P_{(0,1)}, P_{(1,1)})^T$. We can propose the same procedure for parallel Qd. In this case rate-equations are

$$\frac{dn_1}{dt} = \gamma_1 f_L + \gamma_3 f_R + \gamma_2 n_2 - (\gamma_1 + \gamma_2 + \gamma_3)n_1 \quad (12)$$

$$\frac{dn_2}{dt} = \gamma_1 f_L + \gamma_3 f_R + \gamma_2 n_1 - (\gamma_1 + \gamma_2 + \gamma_3)n_2 \quad (13)$$

considering the Fig. 1b. The steady-state solution of (12) and (13) is

$$n_1 = \frac{\gamma_1(\gamma_1 + \gamma_3 + 2\gamma_2)f_L + \gamma_3(\gamma_1 + \gamma_3 + 2\gamma_2)f_R}{(\gamma_1 + \gamma_3)^2 + 2\gamma_2 \gamma_3 + 2\gamma_2 \gamma_1} \quad (14)$$

$$n_2 = \frac{\gamma_1(\gamma_1 + \gamma_3 + 2\gamma_2)f_R + \gamma_3(\gamma_1 + \gamma_3 + 2\gamma_2)f_L}{(\gamma_1 + \gamma_3)^2 + 2\gamma_2 \gamma_3 + 2\gamma_2 \gamma_1} \quad (15)$$

The net out coming flow is

$$I^* = (-q)\frac{\gamma_3}{h}(f_R - n_1) \quad (16)$$

for each quantum dot. And substituting (14) and (15) in (16) we obtain the n-independent expression of the current

$$I^* = \frac{\gamma_1 \gamma_3 (\gamma_1 + \gamma_3 + 2\gamma_2)}{(\gamma_1 + \gamma_3)^2 + 2\gamma_2 \gamma_3 + 2\gamma_2 \gamma_1}(f_L - f_R) \quad (17)$$

**C. EFFECT OF BROADENING**

Up to now we are considering only one energy level per dot. But we have missed the broadening of the level that inevitably accompanies any process of coupling to it. The standard way to introduce energy level broadening as a consequence of contacts is to assign a Lorentzian shape to density of states [18, 19]

$$\rho(E)_i = \frac{\gamma}{\pi} \frac{\gamma}{(E - \epsilon_i)^2 + (\frac{\gamma}{2})^2} \quad (18)$$

The broadening $\gamma$ is proportional to the strength of the coupling, $\gamma = \sum \gamma_i$, where $\gamma_i$ are the transition rates introduced before. This comes out of a full quantum mechanical treatment, but we could rationalize it as a consequence of the “uncertainty principle” that requires the product of the lifetime (= $\hbar/\gamma$) of a state and its spread in energy ($\gamma$) to equal $\hbar$. The current expression is modified into:

$$I = (-q)\frac{\gamma_1}{h}(f_R - n_1) \rightarrow I = (-q) \int \frac{\gamma_1}{\hbar} \rho_{RP_i}(E)(f_R - n_1) dE \quad (19)$$

where now $n_i$ can be interpreted as the non-equilibrium distribution functions. This allow us to compute the total number of electrons at each Qd as

$$N_i = \int_{-\infty}^{+\infty} dE \rho_{n_i}, \quad i = 1, 2 \quad (20)$$

This procedure allow to rewrite n-independent equations as

$$I^* = \frac{q}{\hbar} \int_{-\infty}^{+\infty} dE \frac{\gamma_1 \gamma_3 \rho_{PL} \rho_{PL} \rho_{PL} \rho_{PL}}{\gamma_1 \gamma_3 \rho_{PL} \rho_{PL} + \gamma_1 \gamma_3 \rho_{PL} \rho_{PL} + \gamma_2 \gamma_3 \rho_{PL} \rho_{PL}}(f_L - f_R) \quad (21)$$
The boundary conditions are parallel case, Fig. 2b, as before. We can obtain a recurrence for the case of $N$-$Q_d$ in a case. Conditions must be imposed in order to describe serial each dot. But due to the form of the system, boundary a recurrence in the average number of carriers inside.

$\rho_i$ presented. The outflow in each dot. Now the extended model is extended to the inflow and the outflow in each dot. Now the extended model is presented.

The most general system is consider an array of $N \times M$ Qd, as in Fig. 2c. Using the same approach we obtain

$$n_{i,j} = \frac{\gamma_{i-1,j} \rho_{i-1,j} + \gamma_{i+1,j} \rho_{i+1,j} + \gamma_{i,j} \rho_{i,j} + \gamma_{i+1,j} \rho_{i+1,j}}{\gamma_{i-1,j} + \gamma_{i+1,j} + \gamma_{i,j} + 1} \ldots$$

$$+ \frac{\gamma_{i,j} \rho_{i,j}}{\gamma_{i-1,j} + \gamma_{i+1,j} + \gamma_{i,j} + 1} + \gamma_{i,j} + 1 \ldots$$

Where the position of each dot is defined by $i,j$ sub-indexes and the boundary conditions are

$$\gamma_{N+1,j} = 0 ; \ \gamma_{0,j} = 0 ; \ \gamma_{i,0} = \gamma_{1} ; \ \gamma_{i,M+1} = \gamma_{M}$$

$$\rho_{N+1,j} = 0 ; \ \rho_{0,j} = 0 ; \ \rho_{i,0} = \rho_{L} ; \ \rho_{i,M+1} = \rho_{R}$$

$$n_{N+1,j} = 0 ; \ n_{0,j} = 0 ; \ n_{i,0} = f_L ; \ n_{i,M+1} = f_R$$

where $i = 1 \ldots N$ and $j = 1 \ldots M$.

This general expression of the non-equilibrium distribution function in each dot and the equation of the out coming flow (19) allow us to compute the total current in the steady state.

E. LOCAL POTENTIAL

Another factor that we have to take into account is how the voltage applied to the external electrodes change the electrostatic potential inside each dot. It is easy to see that this can play an important role to determining the shape of the current-voltage characteristics. The classical right solution of potential at each quantum dot involves Poisson equation:

$$\nabla^2 (\phi \nabla V) = -\frac{\Delta q}{\epsilon_0}$$

Where solution is

$$U = U_L + \frac{q^2}{C_{tot}} \Delta N$$

Being $U_L$ the Laplace solution of the system. Considering the capacitive coupling and boundary conditions

$$U_L = \sum_{i} \frac{C_i}{C_{tot}} (-qV_i)$$

Where $i$ covers all the elements of the system, including electrodes and arbitrary number of Qd, and $C_i$ is the capacitive coupling between $i$ and $i+1$ element. Finally $C_{tot} = \sum_i C_i$. The charge energy system constant $U_0 = \frac{q^2}{C_{tot}}$ gives us the potential increase as a consequence of the electron addition. We observe that potential depends on the increasing charge density, but at the same time charge density depends on energy level, a function of local potential. This considerations confers equation for $N$, eq. (20), and for $U$, eq. (27),

\[ f_i = \frac{\gamma_{i} \rho_{i} - 1 \rho_{i+1} + \gamma_{i+1} \rho_{i+1}}{\gamma_{i} \rho_{i+1} + \gamma_{i+1} \rho_{i} + 1} \quad i = 1 \ldots N \]

\[ \rho_0 = \rho L \quad \rho_{N+1} = \rho R \quad n_0 = f_L \quad n_{N+1} = f_R \]

We can obtain a recurrence for the case of $N$-$Q_d$ in a parallel case, Fig. 2b, as before.

\[ n_i = \frac{\gamma_{i} \rho_{i} \rho_{f} \rho_{f} + \gamma_{i} \rho_{i+1}}{\gamma_{i} \rho_{i} + \gamma_{i+1} \rho_{i} + 1} \quad i = 1 \ldots N \]

And the boundary conditions are

$$\gamma_0 = 0 \quad \rho_0 = 0 \quad \gamma_{N+1} = 0 \quad \rho_{N+1} = 0 \quad n_0 = 0 \quad n_{N+1} = 0$$
the position of master equations, as usual in similar cases (9) and (11).
For serial Qd we can consider that first Qd is capacitive coupled with left lead and second Qd, while the second Qd is only coupled with the first Qd and the right lead. Thus the Laplace potential results are
\[ U_{L1}^S = \frac{C_s}{C_{tot}} (-qV_G) + \frac{C_c}{C_{tot}} U_2 \]  
\[ U_{L2}^S = \frac{C_c}{C_{tot}} U_1 \]

where we consider \( V_G \) the potential of left electrode and 0 for the right one. \( C_s \) and \( C_c \) are coupling capacities between Qd and electrode and between Qds. \( U_i \) is the total potential in the \( i \) – th Qd.

Similar expressions can be derived for the parallel system:
\[ U_{L1}^P = \frac{C_s}{C_{tot}} (-qV_G) + \frac{C_c}{C_{tot}} U_2 \]  
\[ U_{L2}^P = \frac{C_c}{C_{tot}} (-qV_G) + \frac{C_s}{C_{tot}} U_1 \]

We shall solve \( U_1 \) and \( U_2 \) equations simultaneously in an iterative process forcing comparison of results [20] and thus ensuring simultaneous convergence.

III. RESULTS

For simplicity we consider fully symmetric system \( \gamma_i = \gamma_j \) and \( \epsilon_i = \epsilon_j \). The effects of local potential on DOS should be computed as \( \rho(E) \rightarrow \rho(E - U) \), which will modify computations for population \( N \) and currents \( I \). Transport requires energy levels of Qds to be between electrochemical potentials of leads. Moreover overlapping of the DOS is also required. Both conditions ensure electron transport and net current. In this model we consider \( \gamma_i << U_0 \) and \( K_B T << U_0 \). Both conditions are necessary for Coulomb Blockade [21]. The self-consistent charging model based on the Poisson equation represents a good zero-order approximation (called Hartree approximation) to the problem of electron-electron interactions, but it is generally recognized that it tends to overestimate the effect. Corrections for the so-called exchange and correlation effects are often added, but the description is still within the one-electron picture which assumes that a typical electron feels some average potential, \( U \), due to the other electrons. Despite our self-consistent potential model do not explain potential differences between spin up and down as consequence of broken spin degeneration [22, 23], we can obtain a first approach to Coulomb Blockade.

Fig. 3a and 3b show obtained intensities for parallel model in a range of capacities values. The \( I(V) \) curve is strongly dependent on \( C_s \) and \( C_c \) through the Laplace solution of the system. The value of the capacity indicates how the system is coupled. When \( C_s \) tends to cero, the two Qd are electrically decoupled with the leads and the potential has not Laplace term, this case implies a lightly coupled system. The other case is when the value of the capacity between Qd (\( C_c \)) goes to cero, which means no electrical influence between Qd. The Poisson term follows the charge \( N_i \) at each Qd and it is therefore always positive, inducing a shift on the potential. Conductance is depicted in Fig. 4a and 4b for the two ranges of capacities. In coherence with results shown in [22] conduction gap is found. The peak of conduction becomes finite as consequence of the temperature. Furthermore the charge effects tend to shift and widening the peak.
Figure 4: Conductance characteristics obtained in the parallel case, in the same cases as before. (a) Resonant conductance peak appears as a result of the rectifying effect. When $C_S$ increases the peak diminishes and the rectifying effect disappears. (b) Conductance gap is obtained and the symmetry breaks due to the electrostatic coupling between the Qds.

Concerning the symmetry let us take into account two considerations. First we compute the Laplace domain at each Qd, as follow

$$\epsilon_1 = \epsilon_0 + \frac{C_s}{C_{tot}} (qV_G) - \frac{C_c}{C_{tot}} U_2 \quad \text{where} \quad U_2 = \frac{C_s}{C_{tot}} (qV_G) \frac{C_c}{C_{tot}} U_1,$$

that define a recurrence (eq. (31) and (32)). In order to analyze the results we will consider second order term $\frac{C_s^2}{C_{tot}} \approx 0$. Thus energy level becomes

$$\epsilon_1 = \epsilon_0 + \frac{C_s}{C_{tot}} (qV_G) - \frac{C_c}{C_{tot}} \frac{C_s}{C_{tot}} (qV_G) \quad (33)$$

As previously stated, the necessary condition for transport is energy level must stay between the electrochemical potentials $\mu_1 > \epsilon_i > \mu_2$ of the leads. In parallel Qd system, there are multiple pathways possible for electron transport and thus the condition $\epsilon_1 = \epsilon_2$ is not required. Then, transport starts when lead electrochemical potential reach the minimum energy level $\epsilon_1$. The potential threshold becomes:

$$V_G^+ = \frac{\epsilon_0}{1 - \frac{C_s}{C_{tot}} + \frac{C_c}{C_{tot}}} \quad (35)$$

$$V_G^- = -\frac{\epsilon_0}{\frac{C_s}{C_{tot}} - \frac{C_c}{C_{tot}}} \quad (36)$$

Two potential thresholds appear, for positive and ne-
gative voltage, as a consequence of previous condition. This creates a region in which no transport occurs because the previous condition is not accomplished, hence a conductance gap appears. The symmetry of the $I(V)$ curves is reached when $V_G^+ = |V_G^-|$. This condition occurs when $\epsilon_0 = 0$ or $C_c = 0$. In the case of a non-degenerate semiconductor the only possibility of broken symmetry is the capacitive coupling between quantum dots.

Fig. 5a and 5b show obtained intensities for serie model in a range of capacity values, in the same cases than before. Now, the electron has single transport way and conditions for transport are more restrictive. Overlapping of the density of states between Qds is also required. Conductance is depicted in Fig. 6a and 6b for the two ranges of capacities.

As we did before, the evolution of the energy levels inside each Qd taking into account (21) and (22) are

$$\epsilon_1 = \epsilon_0 + \frac{C_s}{C_{tot}} (qV_G)$$  \hspace{1cm} (37)

$$\epsilon_2 = \epsilon_0 + \frac{C_s C_c}{C_{tot}^2} (qV_G)$$  \hspace{1cm} (38)

To break the recurrence we will consider $\frac{C^2_c}{C_{tot}} \approx 0$. Now the two conditions imposed before in order to transport occurs and can be summarized as $\mu_1 > \epsilon_1 = \epsilon_2 > \mu_2$. As we can see from the expressions (28) and (29), $\epsilon_1$ increases faster than $\epsilon_2$ and thus the condition $\epsilon_1 = \epsilon_2$ can not be accomplished. Introduced DOS broadening relaxes restrictive conditions for transport as a result of energy channel overlapping. When the voltage increases, the separation between $\epsilon_1$ and $\epsilon_2$ increases as well and then the overlap between DOS decrease closing some channels and decreasing the outcoming flow. As a consequence a NDR appears due to the different electrostatic coupling in each dot [24, 25, 26, 27]. The $I(V)$ peak is related with the maximum overlapping between the DOS of the first and the second Qd [12]. Moreover the width is related with the DOS broadening. The symmetry of the $I(V)$ curves is dominated by $\epsilon_0$.

IV. CONCLUSIONS

A simple phenomenological model of the conduction between serial or parallel quantum dots has been developed based in rate equations for non coherent Qd system. This approach provides the most simple and transparent way for a description of electron transport. In the beginning we consider a system of two Qd linked by a ballistic channel and connected with the emitter and collector reservoirs. Level broadening has been introduced in the DOS changing the expression for current and the population inside each dot. Moreover this model can be extended easily to an array of $N \times M$ Qd. An important point has been the calculation of the local potential, solving the Poisson equation with appropriate boundary conditions for each dot and configuration. Despite its simplicity the effect of self-charge has been taken into account. As we may expect the calculation of the local potential inside each dot is the most important point. This local potential dominates de current characteristics since overlapping of the DOS is imposed in order to have electron transport. $I(V)$ and conductance curves have been obtained in a range of capacity values. NDR and $I(V)$ resonant peak have been observed in the serial configuration due to the different electrostatic coupling. In parallel configuration a conduction gap has appeared as some other

Figure 6: Conductance characteristics obtained in the serial case, in the same cases as before. (a) NDR appears when $C_S$ increases, increasing the difference between the energy levels. (b) Symmetry breaking due to the electrostatic coupling between the Qds. Also NDR is obtained.
authors have reported, therefore two threshold voltages have been obtained. A paradoxical asymmetric current is obtained for the parallel case due to the capacitive coupling between Qds.

ACKNOWLEDGMENT

This work was supported by NASCENT FP7-NMP-245977 european project.

REFERENCES

[20] To solve the system we use the following procedure: we start with an initial guess for $U_{old}$, then calculate $N$ and the total potential $U$. Using the algorithm $U_{new} = U_{old} + \alpha(U_{old} - U_{old})$ we define a new potential $U_{new}$, where $\alpha < 1$. The iterative process has to be repeated until we find a $U_{new}$ which is sufficiently close to the original value. A simultaneous convergence of $U_1$ and $U_2$ is needed.
[23] The system can be described by Hamiltonian of the general form
\[
H = \sum_{k,\sigma} \epsilon_k^\lambda a_k^\dagger a_k + \sum_{p,\sigma} \epsilon_p^R b_p^\dagger b_p + \sum_{\sigma} \epsilon_1 c_1^\dagger c_1 + \sum_{\sigma} \epsilon_2 c_2^\dagger c_2 + U_{1n1n2} + U_{2n2n2} + \text{H.c}
\]
Where, $a_k^\dagger(a_k, \sigma), b_k^\dagger(b_k, \sigma)$ and $c_i^\dagger(c_i, \sigma)$ are the creation (annihilation) operators of an electron with the wave vector $k$ and spin $\sigma$ in the right lead, left lead and $i-th$ Qd respectively. $\epsilon_k^\lambda$ and $\epsilon_p^R$ are the energies levels inside the leads, $\epsilon_1$ and $\epsilon_2$ are the discrete energies levels of the $i-th$ dot. The fifth and sixth terms are the Coulomb integral for two electrons residing in the same dot. The last term describes electron tunnelling between the leads and dots.