Negative Differential Conductance in Nano-junctions: A Current Constrained Approach

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Abstract
A current constrained approach is proposed to calculate negative differential conductance in molecular nano-junctions. A four-site junction is considered where a steady-state current is forced by inserting only the two central sites within the circuit. The two lateral sites (representing e.g. dangling molecular groups) do not actively participate in transport, but exchange electrons with the two main sites. These auxiliary sites allow for a variable number of electrons within the junction, while, as required by the current constrained approach, the total number of electrons in the system is kept constant. We discuss the conditions for negative differential conductance in terms of cooperativity, variability of the number of electrons in the junction, and electron correlations.
I. INTRODUCTION

Understanding of charge transport through low dimensional electronic system has got tremendous impetus in recent years because of their huge potential in advanced electronic devices. Especially, the possibility to switch a molecule between off and on states and to support decreasing current with increasing voltage, the so called negative differential conductance (NDC) behavior, are hot topics in the field of single molecule electronics, mainly due to its credible applications. Since its first observation in the tunnel diode by Esaki in 1958, NDC has been subject of many experimental and theoretical studies. NDC has been observed in a variety of experimental systems and has been discussed theoretically, even if most of theoretical work was based on one-electron pictures. Recent experiments demonstrated NDC in double quantum dots junctions and have rekindled interest in the phenomenon, occurring in the low temperature, weak-coupling limit. Several theoretical studies on donor-acceptor double quantum dot systems address strong rectification and NDC features within many-electron pictures. However, all available theoretical studies of NDC are based on voltage constrained (VC) approaches, where the electric current is driven through the junction by imposing a finite potential drop between two applied electrodes. Two semi-infinite electronic reservoirs are unavoidably introduced in VC models, posing some fundamental problems and leading to a picture difficult to reconcile with correlated electrons.

As an alternative to VC approach, current constrained (CC) approaches have been developed to describe transport in both meso- and molecular junctions. CC approaches avoid any reference to reservoirs, and quite naturally apply to correlated electrons. On physical grounds, a current can be forced in a circuit by driving a time-dependent magnetic flux through the circuit. The model, originally proposed by W. Kohn to describe optical conductivity in extended systems, has been applied to describe electrical transport in mesoscopic systems, and more recently in molecular junctions. Alternatively, variational techniques can be adopted, and the system can be forced in a current-carrying non-equilibrium steady state introducing properly defined Lagrange-multipliers in the Hamiltonian. While offering an interesting alternative to the popular and successful VC methods, the CC approach suffers from several limitations that hinders its widespread use. Not explicitly accounting for contacts, the CC approach relies on phenomenological models for the relaxation dynam-
ics to describe dissipation. In polyatomic junctions, constraints must be explicitly enforced to avoid charge accumulation at atomic sites within the junctions, leading to a cumbersome numerical problem for large and asymmetric molecular systems. Finally, in VC approaches electrons can be exchanged between the electrodes and the molecular junction and charge injection/depletion phenomena can be observed upon changing the applied gate voltage. These phenomena are critical to many processes, including NDC, but can not be described in CC approaches where the current is forced through a closed circuit and the total number of electrons in the junctions is strictly constant.

Here we propose a simple and effective strategy to overcome the last general problem inherent to CC approach. Specifically, we demonstrate that if one or more auxiliary sites are attached to the main current-carrying nanojunction, they can exchange electrons with the junction, thus representing electron source or sink. Along these lines, we present for the first time a calculation of NDC within a CC-based model. Describing NDC in an alternative picture with respect to the widespread VC-based models deepens our current understanding of the basic physics underlying this strongly non-linear phenomenon, as required to control and optimize the performance of NDC-based devices.

II. THE MODEL AND METHOD

Previous work from two of the present authors based on the VC approach demonstrated interesting nonlinear behavior in the current-voltage characteristics of two-dot systems with correlated electron. Taking clue from this work, here we explore transport behavior in two-dots junctions within the CC approach. The study becomes interesting, especially in the context of the number of electrons within the junction, which plays an important role in the low-bias current-voltage characteristics.

Fig. 1(a) shows a rough sketch of the model. We consider a 4 sites Hubbard model as described by the Hamiltonian:

\[ H_0 = \sum_{i=1}^{4} \epsilon_i \hat{n}_i - \sum_{\sigma=\uparrow,\downarrow} t_{i,i+1}(a_{i,\sigma}^\dagger a_{i+1,\sigma} + h.c.) \]

\[ + \sum_{i=1}^{4} U_i \hat{n}_{i\uparrow} \hat{n}_{i\downarrow} \] (1)

where \( a_i \) (\( a_i^\dagger \)) annihilates (creates) an electron with spin \( \sigma \) on site \( i \) and \( \hat{n}_i = \sum_{\sigma} a_{i,\sigma}^\dagger a_{i,\sigma} \). For
the sake of simplicity in the following we set constant on-site repulsions, $U_i = U$, and hopping integrals $t_{ij} = t$, even if different choices have been explored. Moreover we consider an asymmetric junction with a weak bond: $-\epsilon_2 = \epsilon_3 = 2, t = 0.2$.

The eigenstates of $H_0$ are stationary states and do not sustain any current. To induce current through the central (2-3) sites we insert these two sites into a closed circuit through which we impose a current according to the CC prescription:

$$H(\lambda) = H_0 - \lambda \hat{j},$$

(2)

where $\hat{j} = -it_{23} \sum_\sigma (a_{2,\sigma}^\dagger a_{3,\sigma} - h.c.)$ measures the current flowing through the bond between sites 2 and 3 (the junction region). Here and in the following $e$ and $\hbar$ are adopted as units for charge and momentum, respectively. The field $\lambda$ coupled to the current enters the Hamiltonian as a Lagrange multiplier, whose value is fixed by the requirement that a finite current $I = \langle G(\lambda)|\hat{j}|G(\lambda) \rangle$ flows through the system. The eigenstate of $H(\lambda)$ describe electrons with a finite overall linear momentum which looks rather unphysical in an open circuit, as described by $H_0$. This paradox can be solved, according to Burke et al., connecting the two ends of the junction (sites 2 and 3 in our model system) through a long, thin and ideally conducting wire so that electrons escaping from the right side of the junction immediately appear on the left side as to maintain fixed number of electrons inside the junction. As schematically shown in Fig. 1, the junction is then inserted in a closed loop and this allows to assign a precise physical meaning to $\lambda$, that was introduced formally as a Lagrange multiplier. Following the seminal work of Kohn, in fact one can force a current through a closed ring threading an oscillating magnetic field of frequency $\omega$ through the center of the ring to generate a spatially uniform oscillating electric field $E$ across the junction. At the leading order in the field, the Hamiltonian in the presence of a magnetic flux can be written as in Eq. (2) with $\lambda$ proportional to the amplitude of the vector potential. Quite interestingly, a similar result has been recently reported via a general and exact projection operator technique.

The two central sites (sites 2 and 3) are inserted in a closed circuit where the current is forced (cf Fig. 1a) and hence take active part in transport. The lateral sites (1 and 4) represent auxiliary sites: the current does not flow through these sites, but they are connected to the circuit and, exchanging electrons with the main sites (2 and 3), provide
FIG. 1: (a) A schematic representation of the four-site system, where current flows through the two central sites in the closed circuit, while the lateral sites behave as sources/sinks of electrons. (b) The biphenyl molecule with two dangling groups, R and R’, represents an example of a molecule that can be roughly described by the four-site model in panel (a). The red arrows describe the direction of current flow.

a source/sink for the injection/removal of electrons within the circuit. Being interested in NDC we set $U >> t$: the model then describes four weakly connected quantum dots. Alternatively, the same model may represent a minimal model for a molecular junction with at least one weak (poorly conjugated) bond and with two dangling groups (R and R’) connected to the current carrying skeleton as shown, as an example in Fig. 1(b). From a chemical perspective, the groups R and R’, modeled by the auxiliary sites of Fig. 1a can behave as electron withdrawing/donating groups with respect to the main sites.

If the auxiliary sites (1 and 4) are disconnected from the main circuit ($t_{12} = t_{34} = 0$) the number of electrons participating in transport is strictly constant: having inserted the two central sites in an (ideal) closed circuit implies that electrons escaping from site 3 immediately enter on 2 (or vice versa). As discussed above, the obvious constraint of constant charge in a closed circuit hinders the description of phenomena related to charge injection/depletion processes, that are instead easily captured in VC approaches where electrons are exchanged between the junction and the leads. However, if the auxiliary sites are connected to the junction the number of electrons within the circuit, $n = \langle \hat{n}_2 + \hat{n}_3 \rangle$, becomes a variable: finite $t_{12}$ and/or $t_{34}$ allow for charge injection/depletion within the junction.

The Hubbard Hamiltonian in Eq. (1) and its current-carrying version in Eq. (2) can be written on a real-space basis and the resulting matrix can be diagonalized exactly. The
real space basis is defined by the complete and orthonormal set of functions ($\phi_i$) that specify the occupation of site spin-orbitals. Both the total number of electrons, $N$, and the $z$-component of the total spin, $S_z$, commute with the Hamiltonian, and the Hamiltonian matrix can be defined for a particular charge and spin sector. Here we set $N = 4$ and work in the subspace relevant to the ground state, i.e., $S_z = 0$, for a grand total of 256 basis states.

The definition of characteristic current-voltage curves is quite subtle in CC approaches. As discussed above, Hamiltonian in Eq. 2 describes (at the leading order in the vector potential) the effect of an oscillating magnetic flux driving a current through a circuit. In this view, $\lambda$ is simply proportional to the corresponding electrical field: $E \propto \omega \lambda$. However in the DC limit, $\omega \to 0$, the driving field, and hence the driving potential vanish, leading to the unphysical result of a finite current at zero bias. This zero-resistance picture emerges because relaxation dynamics is neglected: in fact a finite potential drop is required to sustain current just to overcome friction in the junction. The calculation of characteristic curves in the CC approach then requires a model for relaxation dynamics. In particular, the current-carrying state, $|G(\lambda)\rangle$, is a non-equilibrium state and the system relaxes back to the ground state, $|G(0)\rangle$. The amount of power spent to sustain the system in the non-equilibrium current-carrying state is set by the relaxation dynamics: the faster the relaxation the more power must be spent to sustain the current. The Joule law defines the relation between the electrical power spent on the molecule, $W$, and the potential drop needed to sustain the current: $W = IV$. Since $I$ is known, characteristic $I(V)$ curves can be obtained from $W$.

We adopt a simple phenomenological model for relaxation dynamics that is widely used in spectroscopy. The model is general and properly obeys basic physical requirements, including detailed balance. In particular relaxation is described by introducing the density matrix ($\sigma$) written on the basis of the eigenstates of $H(0)$. On this basis the equilibrium density matrix is diagonal, with diagonal elements describing the Boltzmann populations of the relevant states. Relaxation dynamics of digonal elements of $\sigma$ are only affected by inelastic scattering and are written as:

$$ (\dot{\sigma}_R)_{kk} = \sum_m \gamma_{km} \sigma_{mm} - \sum_m \gamma_{mk} \sigma_{kk} $$

(3)

where $\gamma_{km}$ measures the probability of the transition from $m$ to $k$ as due to inelastic scattering
events. Off diagonal elements of the relaxation matrix may be written as:

\[ (\dot{\sigma}_R)_{km} = -\Gamma_{km}\sigma_{km} \]  

where

\[ \Gamma_{km} = (\gamma_{kk} + \gamma_{mm})/2 + \gamma'_{km} \]

accounts for both inelastic scattering events, via the population inverse lifetimes \( \gamma_{kk} = \sum_m \gamma_{mk} \), and for elastic scattering events via \( \gamma'_{km} \) that describes the loss of coherence due to pure dephasing phenomena.\(^{39,40}\)

The total work exchanged by the junction, \( \text{Tr}(\dot{\sigma}_R\hat{H}) \), has 2 non-vanishing contributions: \( W = -\lambda \text{Tr}(\dot{\sigma}_R\hat{j}) \) that represents the work done on the junction to sustain the current and a second contribution, \( W_d = \text{Tr}(\dot{\sigma}_R\hat{H}_0) \), that measures the work dissipated by the junction. Since the current operator, \( \hat{j} \), has vanishing diagonal elements on the real eigenstates of \( H(0) \), \( W \) is only affected by the relaxation of off-diagonal elements of the density matrix. As a result, both elastic and inelastic scattering events affect \( W \). This is a physically relevant result; in fact, both phenomena concur to build up the resistance in the junction. On the opposite, the power dissipated by the junction, \( W_d = \text{Tr}(\dot{\sigma}_R\hat{H}_0) \), is only affected by inelastic scattering events. It represents just a fraction of the total power dissipated on the junction, thus suggesting that some dissipation must occur at the electrodes.\(^{35}\)

While the relaxation model is fairly crude and requires invoking electrodes to actually balance the energy, it has the advantage of setting on a firm basis the relation between the molecular resistance and dissipation phenomena. In particular we notice that the proposed model properly predicts the maximum conductivity of the simplest junction as \( e^2/\hbar \) as a result of the maximum lifetime of the electron within the junction which cannot be longer than the time required to the electron to cross the junction.\(^{35}\) This observation drives us to the non-trivial role of electrodes on relaxation dynamics. In the case of very weak contacts, the presence of metallic surfaces close to the molecular junction is expected to open new channel for energy dissipation then reducing the population lifetimes.\(^{41}\) On the opposite, in the strong contact regime the mixing of the discrete molecular eigenstates with the continuum of states of the metallic leads is responsible for increased dephasing rates.\(^{24}\) A very simple model for the relaxation matrix can then be obtained in the strong contact regime, where depopulation contributions to \( \Gamma_{km} \) can be neglected, and in the hypothesis that the effects of the electrodes on the dephasing rates is similar on all states, so that \( \Gamma_{km} = \Gamma \). In
III. RESULTS AND DISCUSSION

We consider two different choices for the energies of the auxiliary sites: in the first model (model a) we set $\epsilon_1 = -\epsilon_4$ to mimic groups with opposite electron donating/accepting characteristics. In the second model (model b) we set $\epsilon_1 = \epsilon_4$ mimicking two equivalent side-groups added to the main chain (R=R’ in Fig. 1(b)). Quite interestingly, the second model can also describe the effects of charge injection/depletion in the junction as a result of an applied gate voltage. The color maps in Fig. 2 show the differential conductance ($\frac{dI}{dV}$) calculated as a function of the potential drop ($V$) and of the energies of the auxiliary sites, $\epsilon_1 = -\epsilon_4$ (left panel) and $\epsilon_1 = \epsilon_4$ (right panel) for $U = 5$. NDC is not observed in either case. This result, confirmed at different $U$ values, is not surprising. In fact NDC suggests a largely non-linear behavior of the junction, with a bistable $V(I)$ characteristic. Such a behavior requires cooperativity and hence the introduction of some self-consistent interaction in the model.

With this requirement in mind, we modify the model to account for possible effects of the voltage drop on the auxiliary sites. In fact, sites 1 and 4 do not play any active role in transport, but can be affected by the voltage drop at the junction. In particular, being
directly connected to sites 2 and 3 (device region), they can experience at least a portion of
the electrical field required to sustain the current in the main junction. As a result, a term
is added to the Hamiltonian affecting the auxiliary-site energies as follows:

\[ x V (\hat{n}_4 - \hat{n}_1), \]  

where \( x \) measures the fraction of the potential drop that is felt by sites 1 and 4. We observe
that only the energy of auxiliary sites must be explicitly modified by the potential: the effect
of the potential on the junction (sites 2 and 3) is already accounted for by the Hamiltonian
in Eq. 2. The correction term in Eq. 6 introduces cooperativity in the model: the on-site
energies, \( \epsilon_1 \) and \( \epsilon_4 \), do depend in fact on the current flowing through the junction that, in
turn is affected by the on-site energies. Color maps in Fig. 3, obtained setting \( x = 0.5 \) show
NDC regions, demonstrating the cooperative nature of the phenomenon. Fig. 3 summarizes
the main results of this paper showing the differential conductance calculated for the two
choices, (a) \( \epsilon_1 = -\epsilon_4 \) (left columns), and (b) \( \epsilon_1 = \epsilon_4 \) (right columns) and \( U =5 \) and 2 (top
and bottom panels, respectively). NDC feature is definitely more prominent in model b
than in model a and is suppressed upon decreasing electron correlation strength, \( U \) (this
conclusion is also supported by extensive calculations run at different \( U \)). These findings
are in line with results obtained in the VC approach, showing that strong on-site \( e-e \)
correlations favor NDC in double-quantum dot systems\[26\] Note that, the \( (\frac{\partial I}{\partial V}) \) map (and
the \( I-V \) characteristics as well) are asymmetric because of the inherent asymmetry in the
device region, with sites 2 and 3 having different on-site energies.

Due to symmetry reasons, the number of electrons in the junction is constant in model
(a), \( n = 2 \). In model (b), \( n \) is instead variable. Since the hybridization energy, \( t \), is small
with respect to on-site energies, \( n(V, \epsilon_1) \) assumes almost integer and constant values (shown
as italic numbers in the figures) in different regions of the \( V, \epsilon_1 \) plane connected by very sharp
borders. Quite impressively, NDC features are observed for model (b) just at these borders,
suggesting a strong relation between NDC and sharp variations of \( n \). Upon decreasing \( U \), the
variation of \( n \) becomes smooth and the NDC regions become less prominent. Minor NDC
features are also observed in model (a) with constant \( n \), in regions where the occupancy of
the two auxiliary sites abruptly changes from \( \langle \hat{n}_1 - \hat{n}_4 \rangle = 0 \) to \( \pm 2 \).

To obtain a more detailed picture of the microscopic origin of NDC, we disentangle the
contributions to the current from the different states. Specifically we work a real-space basis,
FIG. 3: Color map showing differential conductance ($\frac{dI}{dV}$) as a function of potential drop ($V$) and the energies of the auxiliary sites for $\epsilon_1 = -\epsilon_4$ (left panels) and $\epsilon_1 = \epsilon_4$ (right panels), for two different $U$ values. The blue numbers in the right panels show the approximate value of the number of electrons in the junction, $n$, in the different regions. The same number is constant ($n = 2$) in the left panels.

where states are defined assigning electrons with a specified spin to each site in the junction. A generic state $|p\rangle = |0, \uparrow, \downarrow, \downarrow\uparrow\rangle$ then graphically represents the state where no electrons are found in the first site, one electron with spin $\alpha$ and one electron with spin $\beta$ are located on sites 2 and 3, respectively, and site 4 is doubly occupied. In more formal language: $|0, \uparrow, \downarrow, \downarrow\uparrow\rangle = a_{2,\alpha}^\dagger a_{3,\beta}^\dagger a_{4,\beta}^\dagger a_{4,\alpha}^\dagger |0\rangle$ where $a_{i,\sigma}^\dagger$ are the electron creation operators entering the Hamiltonian in eq. 1 and $|0\rangle$ is vacuum state- The current-carrying ground state has complex coefficients $c_p$ in the real space basis, $|G(\lambda)\rangle = \sum_p c_p |p\rangle$ and the expectation value of the current operator can be written as $I = \sum_{pr} I_{pr}$, where $I_{pr} = c_p c_r^* j_{pr}$, and $j_{pr}$ is an off-diagonal element of the current operator on the chosen basis $j_{pr} = \langle p | \hat{j} | r \rangle$ (diagonal elements of $\hat{j}$ vanish in the chosen basis). For the sake of clarity, we set $U = 5$ in the following discussion. Fig 4 shows the results relevant to model (a) with $\epsilon_1 = -\epsilon_4 = 3$. In this case, as noticed before, $n = 2$. Two main contributions to $I$ can be recognized as $I_1 = I_{|\uparrow, \downarrow, \uparrow\downarrow, 0\rangle, \downarrow\uparrow, \downarrow\uparrow, \downarrow\uparrow} + I_{|\downarrow, \uparrow\downarrow, \uparrow\downarrow, \downarrow\rangle, \uparrow\downarrow, \downarrow\uparrow, \downarrow\uparrow}$ and $I_2 = I_{|\uparrow, \downarrow\uparrow, 0, \uparrow\downarrow\rangle, \downarrow\uparrow, \downarrow\uparrow, \downarrow\uparrow} + I_{|\downarrow, \downarrow\uparrow, \uparrow\downarrow\rangle, \downarrow\uparrow, \downarrow\uparrow, \downarrow\uparrow}$. Quite interestingly, as shown in Fig. 4, the sum of these two contributions quantitatively matches with the total $I(V)$ characteristic. The states involved in $I_1$ and in $I_2$ only differ by the occupation at 1st and 4th sites: the states involved in $I_1$ show single occupancy of sites 1 and 4, while the states involved in $I_2$
FIG. 4: Results for $\epsilon_1 = -\epsilon_4 = 3$, and $U = 5$. The black line in the top panel shows the characteristic curve. The blue and green lines in the same panel show the two major contributions to the current as discussed in the text, the red dashed line shown the sum of the two contribution, and it is almost indistinguishable from the total current. In the bottom panel blue (green) lines show the weight of the two states responsible for the $I_1$ ($I_2$), as discussed in the text. The black dotted line is shown as a guide for the eyes have zero occupation at site 1 and double occupation at site 4. The position and height of the NDC peak of $I_2(V)$ exactly matches with the NDC peak of the $I(V)$ characteristics: the fall of $I_2$ and the rise of $I_1$ at $V \sim 0.75$ causes the NDC feature in $I(V)$ characteristics. For $V > 0.75$, $I_2$ starts decreasing to zero, while $I_1$ increases to meet $I_2$ at $V=1.04$ resulting a valley in $I(V)$. So, for $V > 1.04$, $I_1$ dominates over $I_2$ resulting in further increase of total $I$.

The bottom panel of Fig. 4 shows the weight of the states that mainly contribute to the current, as shown in the figure label. The probability of states contributing to $I_1$(blue lines) vanishes for $V < 0.75$: $I_1=0$ in that region. $I_1$ instead contributes significantly to total $I$ for $V > 0.75$, where the contribution from $I_2$ becomes negligible. The current flows only through the central bond, and, in a system with $n = 2$ and $\epsilon_2 = -\epsilon_3 = 2$,
FIG. 5: Results for $\epsilon_1 = \epsilon_4 = 2$, and $U = 5$. Panel (a): Characteristic curve (black line) and total number of electrons in the junction (blue line). Panel (b) the three main contributions to the current in the NDC regions, as discussed in the text. The red dashed lines shows their sum to be compared to the total current (black line). Panel (c) shows the weight of states contributing to $I_1$ (blue lines), $I_2$ (violet lines) and $I_3$ (green lines). More specifically the blue continuous and dashed line refer to states $|\uparrow, \downarrow, \uparrow, \downarrow\rangle$ and $|\uparrow, \downarrow, \uparrow, 0, \downarrow\rangle$, respectively; the green continuous and dashed lines refer to $|\downarrow, \downarrow, \uparrow, 0, \downarrow\rangle$, and $|\downarrow, \uparrow, \downarrow, \uparrow, 0\rangle$, respectively; the violet continuous and dashed lines refer to $|0, \downarrow, \uparrow, \downarrow\rangle$, and $|0, \uparrow, \downarrow, \downarrow\rangle$, respectively.

It is associated with off-diagonal elements of $\hat{J}$ mixing up states where both electrons reside on site 2 or are equally distributed in sites 2 and 3. NDC is observed when the occupancy of the auxiliary sites changes: the NDC feature occurs when the population of states contributing to $I_2$, characterized by $\langle \hat{n}_4 - \hat{n}_1 \rangle = 2$, start decreasing at the expense of the increasing population of states contributing to $I_1$, characterized by $\langle \hat{n}_4 - \hat{n}_1 \rangle = 0$.

Model b is more complex, as shown in Fig. 5, collecting the results obtained for $\epsilon_1 = \epsilon_4 = 2$ and $U = 5$. In this case a variable number of electrons is found in the junction, as shown by the blue lines in Fig. 5a. The $I - V$ characteristic (black line) shows NDC features both in the negative and positive $V$ regions, with a more
prominent feature at $V \sim 2$. In the region of interest for NDC the current is mainly due to three specific contribution, as shown in Fig. 5 b. The three contributions are:

$I_1 = I_{\uparrow\uparrow, \downarrow\downarrow, 0} + I_{\downarrow\uparrow, \uparrow\downarrow, 0} + I_{\downarrow\downarrow, \uparrow\uparrow, 0}$, $I_2 = I_{\uparrow\downarrow, \uparrow\downarrow, 0} + I_{\downarrow\uparrow, \uparrow\downarrow, 0}$, and $I_3 = I_{\downarrow\uparrow, \uparrow\downarrow, 0} + I_{\downarrow\downarrow, \uparrow\downarrow, 0}$. The states involved in $I_1$ have two electrons in the junction, $n = 2$, while the states contributing to $I_2$ and $I_3$ have $n = 3$. As expected, $I_2$ and $I_3$ contributions become sizable for large negative and positive $V$, respectively. However it is clear from the figure that the NDC features observed in the characteristic $I(V)$ are only due to the NDC features observed in the $I_1(V)$ contribution to the total current.

For a better understanding, Fig. 5c shows the probability of states playing a main role in transport. $I_1$ drops at the NDC features because the relevant states, having two electrons in the junction, lose their weight and the increasing contributions of either $I_2$ in the negative $V$ region or of $I_3$ in the positive $V$ region, related to states with $n = 3$, are not able to fully compensate for the loss.

IV. CONCLUSIONS

The current constrained approach offers an alternative picture of transport in nanojunctions with respect to the widespread voltage-constrained approaches. In the CC approach Lagrange multipliers are used to force the system into a current carrying state, then avoiding any explicit reference to semi-infinite electronic reservoirs. The method then applies quite naturally to describe transport in systems with strongly correlated electrons. However, the total number of electrons taking part in transport is fixed in CC approaches, and this has hindered so far the application of the method to problems where charge injection/depletion in the junction is important. This problem could be solved introducing an effective relaxation model that, accounting for the exchange of electrons between the junction and the electrodes, allows the system to relax towards states with different number of electrons. This approach however requires a fairly complex model for relaxation dynamics. Here we borrow from the field of optical spectroscopy a simple phenomenological model for relaxation that maintains constant the number of electrons, and overcome the limitation of fixed number of electrons connecting one or more auxiliary sites to the main junction: while the current does not flow through the auxiliary sites, they exchange electrons with the junction, allowing for a variable number of electrons taking part in transport.
Along these lines we attacked the problem of NDC in a very simple model with a two-site junction connected to two auxiliary sites. NDC is a highly non linear phenomenon: the bistable behavior of the $V(I)$ curve at NDC can only be observed in largely cooperative systems. We introduced cooperativity imposing a dependence of the on-site energies of the auxiliary sites on the voltage drop: since the voltage required to support the current depends in turn on the energy of the auxiliary sites, the problem becomes self-consistent and the model supports NDC. Quite interestingly, dangling bonds (side groups) connected to the main current-carrying unit have recently been discussed as responsible for non-linearity in the characteristic curves. Particularly side groups may be responsible for quantum interference phenomena that drastically suppress current.\textsuperscript{42} Our study suggests that a proper choice of the chemical nature of the side groups can not only reduce the current, but eventually lead to NDC.

Here we just investigated an asymmetric junction with weak bonds and a grand-total of four electrons, in two different cases. In the first case we choose equal and opposite on-site energies for the two auxiliary sites leading to a constant number of electrons in the junction ($n = 2$). Minor NDC features are observed in this case, mainly related to an abrupt variation of the occupation of the auxiliary sites. More prominent NDC features are observed in the second model, where, having fixed the energies of the auxiliary sites to the same value, the number of electrons involved in the transport is variable. NDC is related in this case to an abrupt variation with the applied potential of the number of electrons in the junction. In all cases, large $U$ and small $t$ are needed to observe NDC: weak bonds and large correlations in fact ensure the abrupt variation in the nature of the current carrying state $|G(\lambda)\rangle$, as required for to observe NDC.

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