Impact of capacitance and tunneling asymmetries on Coulomb blockade edges and Kondo peaks in non-equilibrium transport through molecular quantum dots

A. A. Aligia  
Centro Atómico Bariloche and Instituto Balseiro, 
Comisión Nacional de Energía Atómica, 8400 Bariloche, Argentina

P. Roura-Bas  
Dpto de Física, Centro Atómico Constituyentes, Comisión Nacional de Energía Atómica, Buenos Aires, Argentina

Serge Florens  
Institut Néel, CNRS et UJF, BP 166, 38042 Grenoble Cedex 9, France  
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We investigate theoretically the non-equilibrium transport through a molecular quantum dot as a function of gate and bias voltage, taking into account the typical situation in molecular electronics. In this respect, our study includes asymmetries both in the capacitances and tunneling rates to the source and drain electrodes, as well as an infinitely large charging energy on the molecule. Our calculations are based on the out-of-equilibrium Non-Crossing-Approximation (NCA), which is a reliable technique in the regime under consideration. We find that Coulomb blockade edges and Kondo peaks display strong renormalization in their width and intensity as a function of these asymmetries, and that basic expectations from Coulomb blockade theory must be taken with care in general, especially when Kondo physics is at play. In order to help comparison of theory to experiments, we also propose a simple phenomenological model which reproduces semi-quantitatively the Coulomb blockade edges that were numerically computed from the NCA in all regimes of parameters.

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I. INTRODUCTION

In the last decade, an enormous amount of research in the field of nanoelectronics was devoted to the study of electronic transport through quantum dots. A seminal and very versatile type of device is the gate-defined artificial atom created at the two-dimensional electron gas confined in semiconducting heterostructures. More recently, molecular quantum dot setups, several physical effects are generically observed when the system is cooled at cryogenic temperatures: Coulomb blockade is probably the major one, due to the large charging effects in these nanoscopic quantum dots. In addition, more subtle correlations show up due to the density of states of the Anderson model is two times wider when the dot is occupied. Coulomb blockade is therefore nearly five times more intense and nearly two times narrower than the peak at negative voltage. The authors assumed that for negative voltage, the width in the spectral density of the dot level is two times larger than for positive voltage. This is based on the fact that the charge-transfer peak (the one near the dot level $E_d$ in the density of states of the Anderson model is two times wider when the dot is occupied. However, as we shall show, the peak for negative voltage corresponds to the intermediate valence (IV) regime, in which the width of the spectral density is approximately 3/2 times larger than in the empty-orbital regime. In addition, the ratio of widths for negative and positive voltages depends also on the asymmetries of the source and drain capacitances. Our calculation will also be able to take into account the presence of a Kondo resonance at small bias, which in previous Coulomb blockade theory was clearly lacking.

1. Impact of Capacitance and Tunneling Asymmetries

Asymmetries between source and drain electrodes can also take place in transport features at finite bias voltage. For instance, Coulomb blockade edges were shown by Könenmann et al. to present a strong width renormalization in the situation of large tunnel asymmetries between source and drain electrodes, that is also quite typical in molecular quantum dots. Enhancement of cotunneling features of excitation levels (as precursor of the Kondo effect) were also described by Paaske et al. and later widely observed. All these features need to be incorporated in the current theoretical developments of non-equilibrium quantum transport, which has so far mainly focused in analyzing the main universal Kondo signatures in $G(V)$ under many simplifying assumptions (such as symmetric capacitive and tunnel coupling to the electrodes). In particular, the experiment by Könenmann et al. investigated a strongly confined semiconducting quantum dot coupled to two conducting leads with highly asymmetric tunnel couplings, and studied tunneling resonances at nearly opposite values of $V$. While in absence of interaction effects one expects nearly the same intensity and width of the two resonances (see Section III), the Coulomb blockade edge at positive voltage was however nearly five times more intense and nearly two times narrower than the peak at negative voltage. The authors assumed that for negative voltage, the width in the spectral density of the dot level is two times larger than for positive voltage. This is based on the fact that the charge-transfer peak (the one near the dot level $E_d$ in the density of states of the Anderson model is two times wider when the dot is occupied. However, as we shall show, the peak for negative voltage corresponds to the intermediate valence (IV) regime, in which the width of the spectral density is approximately 3/2 times larger than in the empty-orbital regime. In addition, the ratio of widths for negative and positive voltages depends also on the asymmetries of the source and drain capacitances. Our calculation will also be able to take into account the presence of a Kondo resonance at small bias, which in previous Coulomb blockade theory was clearly lacking.
In this paper, after a brief discussion of what is expected in the absence of Coulomb interactions for asymmetric coupling to the leads, we calculate the differential conductance $dI/dV$ in the Keldysh formalism within the non-crossing approximation (NCA).\textsuperscript{20,21} The NCA reproduces well the scaling relations with temperature $T$ and bias voltage in the Kondo regime\textsuperscript{22} and was for example also successfully used to interpret experimental results on a controlled crossover between SU(4) and SU(2) Kondo states driven by magnetic field in a nanoscale Si transistor\textsuperscript{23}, as well as quantum phase transitions involving singlet and triplet states in molecular quantum dots\textsuperscript{24}. It was also recently discussed as a possible model for non-equilibrium Kondo spectroscopy in a system of double quantum dots.\textsuperscript{24} For asymmetric coupling to the leads, we will also derive an analytical expression which describes semi-quantitatively the NCA results. This expression is used to interpret the experiments of Könenmann et al.\textsuperscript{15} providing a good agreement, and could be used for further studies of molecular systems.

The paper is organized as follows. In Section II we present the Hamiltonian and the equation describing the current in presence of a finite bias. In Section III, we compute for comparison the differential conductance $G(V)$ in absence of Coulomb interaction. Section IV contains the NCA results for the non-linear conductance, and presents an analysis of the positions and widths of the peaks displayed in $G(V)$ for different experimentally relevant conditions. In Section V, we discuss the experiment of Könenmann et al., and end up in Section VI with a summary.

II. MODEL

We consider the Anderson impurity model with infinite on-site repulsion $U$, an assumption that is realistic for most molecular quantum dots, but also well obeyed in very small semiconducting devices in the vicinity of a charge-degeneracy point. The impurity states thus involve a singlet configuration $|s\rangle$ together with a spin doublet $|\sigma\rangle (\sigma = \uparrow$ or $\downarrow$) corresponding to one additional electron (or hole) in the dot, taking this particle from the left ($L$) or right ($R$) conducting leads to which the dot is connected. The Hamiltonian thus reads:

$$H = \sum_{\sigma} E_d |\sigma\rangle \langle \sigma| + \sum_{\nu k \sigma} \varepsilon_{\nu k} c_{\nu k \sigma}^\dagger c_{\nu k \sigma}$$

$$+ \sum_{\nu k \sigma} (V_{\nu k}^e |\sigma\rangle \langle s| c_{\nu k \sigma} + \text{H.c.}),$$

where the constraint $|s\rangle \langle s| + \sum_{\sigma} |\sigma\rangle \langle \sigma| = 1$ is imposed. Here $c_{\nu k \sigma}^\dagger$ create conduction states in the lead $\nu$ with spin $\sigma$ and wave vector $k$. The tunnel couplings of the quantum dot to the leads are $\Delta_{\nu} = \pi \sum_{k} |V_{\nu k}^e|^2 \delta(\omega - \varepsilon_{\nu k})$, assumed independent of energy $\omega$, and we define the total hybridization $\Delta = \Delta_L + \Delta_R$. We take the sign of the bias voltage in such a way that $\mu_L - \mu_R = eV$, where $\mu_{\nu}$ is the chemical potential of lead $\nu$. Note that the interchange of right and left leads or electrons by holes is equivalent to a change of sign of $V$. We take the arbitrary origin of energies at $\mu_R = 0$. The capacitance effects modify the energy necessary to add an electron to the dot with the lever arm parameter $\alpha$ (which depends on the source, drain and gate capacitances)\textsuperscript{15,16}, defined as:

$$E_d = E_d^0 + \alpha eV,$$

where $0 \leq \alpha \leq 1$, and $e$ is the electron charge (taken positive). For the situation of symmetric voltage drop that is usually considered, $\alpha$ is often taken near 1/2, but this may not be the case in many experimental setups.

We obtain the conductance $G = dI/dV$ differentiating the current $I$ (numerically from NCA results), which is given by\textsuperscript{25}

$$I = C \int d\omega \rho(\omega, V, E_d) [f_L(\omega) - f_R(\omega)],$$

$$C = \frac{8\pi e \Delta_L \Delta_R}{h},$$

where $f_L(\omega) = f(\omega - \mu_L)$ is the Fermi distribution in each lead, with $f(\omega) = 1/(e^{\omega/kT} + 1)$, and $\rho(\omega, V, E_d)$ is the non-equilibrium (voltage-dependent) spectral density of the impurity level.

III. NON-INTERACTING RESONANT-LEVEL MODEL

Before discussing in more detail the complex effects of Coulomb interactions, we present in this section a simplified model that sheds light onto the expected behavior of the differential conductance $G(V)$. The first assumption that we make in this section is that the tunnel couplings to the leads are very asymmetric. This hypothesis is realistic for molecular quantum dots and also in the experiment of Könenmann et al. in Ref.\textsuperscript{13} We make here the choice $\Delta_R \gg \Delta_L$, so that the system is in equilibrium with the right lead and the explicit dependence on the voltage of the spectral density $\rho(\omega)$ disappears [the dependence of $\rho(\omega)$ with $V$ through $E_d$ given by Eq. 2 remains]. Then one has

$$\rho(\omega, V, E_d) \simeq \rho_R(\omega, E_d),$$

where $\rho_R(\omega, E_d)$ is the spectral density of the impurity states in equilibrium with the right lead, for a dot energy given by Eq. 2. Using equations 2, 4, one obtains at $T = 0$

$$G(V) = Ce\rho_R(eV, E_d^0 + \alpha eV)$$

$$+ \alpha Ce \int_0^{eV} d\omega \frac{\partial \rho_R(\omega, E_d^0 + \alpha eV)}{\partial E_d}.$$
density \( \rho_R(\omega, E^0_d) \), as is usual with experiments with a scanning tunneling microscope (STM). Therefore, in this limit, \( G(V) \) has a single peak located at \( V = E^0_d / e \) (the charge-transfer peak), and no peak for the opposite bias voltage. In addition, a Kondo peak at \( V = 0 \) can show up in presence of Coulomb interactions, if \( -E^0_d \gg \Delta \).

The main assumption that we make in this section is to take a non-interacting resonant level spectral density of width \( 2\Delta \) at half maximum, \( \rho_R(\omega, E_d) \approx \rho_0(\omega - E_d) \), with

\[
\rho_0(\omega - E_d) = \frac{\Delta/\pi}{(\omega - E_d)^2 + \Delta^2}. \tag{6}
\]

This is a crude approximation which is certainly not valid in presence of interactions (it misses the important Kondo anomaly), but it can describe qualitatively the position and width of the Coulomb peaks in \( G(V) \) derived from the charge-transfer peak in the spectral density of the dot states, which can be renormalized from bare values by the interaction, as we discuss in Section IV B 2 Assumptions, as in Eq. (6), that \( \partial \rho_R / \partial E_d = -\partial \rho_R / \partial \omega \) [but not \( \rho_R = \rho_0 \) yet], the second term of Eq. (6) can be easily evaluated and one obtains

\[
G(V) \simeq C e[(1 - \alpha)\rho_R(eV, E_d) + \alpha \rho_R(0, E_d)], \tag{7}
\]

noting that \( E_d \) displays an explicit dependence upon the voltage from Eq. (6). While the assumption made above for the derivatives of the current is not exactly true (even in equilibrium) because of the dependence of the weight and width of the peaks with \( E_d \), this simple expression reflects a general feature of the expected Coulomb peaks, in connection to the charge-transfer peak of the equilibrium spectral density \( \rho_R \). Indeed, now \( G(V) \) displays two finite bias peaks, one that is similar to the STM case and is located at voltage \( eV = E_d \) [namely \( eV = E^0_d / (1 - \alpha) \)], and a second “intermediate valence” (IV) peak at an opposite voltage value such that \( E_d = E^0_d + \alpha eV = 0 \). In this latter case, the impurity experiences indeed maximal charge fluctuations because the level is located exactly on-resonance, and thus one can expect a different width renormalization than for the STM peak, due to the different influence of the Coulomb interaction, as we will demonstrate later on. For positive \( E^0_d \) (corresponding to a vanishing charge in the dot, also called the empty orbital regime), the STM peak lies at positive bias and the IV peak is located at negative bias, and conversely for negative \( E^0_d \) (corresponding to a spin-degenerate odd charge in the dot, also called the local moment regime). Thus, if \( \rho_R = \rho_0 \) is assumed (which is valid only in absence of Coulomb interaction), the conductance \( G(V) \) displays two peaks as a function of voltage: i) an “STM-like” peak at \( V = E^0_d / (1 - \alpha)e \) of width \( 2\Delta/(1 - \alpha)e \) and maximum intensity \( (1 - \alpha)e C/\Delta \); ii) an IV peak at \( V = -E^0_d / (\alpha e) \) of width \( 2\Delta/(\alpha e) \) and maximum intensity \( \alpha e C/\Delta \). Therefore, in the absence of Coulomb interactions, the amplitude and width renormalizations are related, and cannot reproduce the experimental observations made in Ref. 15. We now turn to a full microscopic calculation of the problem, without recourse to the many assumptions made above, that incorporates the correct nature of the Coulomb blockade edges and Kondo resonances for all parameters in the device.

IV. NCA RESULTS FOR THE INTERACTING MODEL

A. The equilibrium spectral density

We consider now Hamiltonian (1) in its full complexity for the case of a finite bias voltage. The non-crossing approximation (NCA) has proved in the past an excellent technique to deal with large Coulomb interactions, and can be extended to the present non-equilibrium situation. In order to gain intuition, we first focus on the zero-bias \( (V = 0) \) spectral density of the localized level, which is shown on Fig. 1 for low temperatures and three values of the dot energy level \( E_d \), corresponding to the empty orbital \( (E_d = 4\Delta) \), local moment \( (E_d = -4\Delta) \), and mixed valence \( (E_d \sim 0) \) regimes. For positive \( E_d \) the result is very similar to the non-interacting resonant level prediction, Eq. (6), with a spectral width \( 2\Delta \), except for the fact that the position of the peak is slightly shifted to higher energy \( E_{d\text{eff}} \). This energy shift \( E_{d\text{eff}} - E_d \) is in very good agreement with the Haldane shift \( 2\pi D/\Delta \ln(D/\Delta) = 0.733\Delta \) for the half-band width \( D = 10\Delta \) that we have used in the calculations. This is expected because the NCA incorporates naturally the
lowest order perturbative terms of the development in the hybridization $V_i^\alpha$. Note that the small spike at the Fermi energy $\omega = 0$ is an artifact of the NCA in the empty orbital regime.

In the local moment regime at negative $E_d$, the charge-transfer peak is nearly two times wider due to the underlying extra spin degeneracy, as was discussed in the introduction and in previous works, and the NCA correctly reproduces this behavior. The total intensity is also reduced by correlation effects, as discussed phenomenologically in Section IV B.2 and the position of the peak is also shifted to higher energies. More importantly, the screening of the spin-degenerate states leads to a Kondo peak at the Fermi level, a non-trivial effect that the NCA is able to describe properly.

Finally, the density of states for the mixed valence situation ($E_d^{\text{eff}} = 0$) shows a peak centered at the Fermi level as expected, with a width of about $3\Delta$ that interpolates between the widths found in the empty orbital and local moment regimes. The temperature in this calculation has been increased to avoid a shortcoming of the NCA.

Regarding the conductance $G(V)$, these results anticipate similarities and differences with the naive expectations based on the non-interacting resonant level model discussed at the end of the previous section (apart from the obvious lack of Kondo resonance in the latter model). A minor point is that $E_d^{\text{eff}} = 0$ should be replaced by an effective level $E_d^{\text{eff}}$, which includes the Haldane shift. Based on Fig. 1 and Eq. (2), the width of the STM-like peak in the empty orbital regime should be $2\Delta/(1-\alpha)e$, thus $4\Delta$ at $\alpha = 1/2$ for a symmetric voltage drop (as for the non-interacting resonant level model). However, the width of the STM peak in the local moment regime should rather be $4\Delta/(1-\alpha)e$, thus $8\Delta$ at $\alpha = 1/2$. More subtly, the IV peak (located at opposite voltage to the STM peak) should have the same width for the empty orbital and local moment regimes, namely $3\Delta/(1-\alpha)e$, thus $6\Delta$ at $\alpha = 1/2$. These general trends, that we will now examine microscopically from full NCA calculations of the conductance, show that the experimental analysis of the width of Coulomb blockade peak must be done with care.

**B. Conductance for asymmetric tunnel coupling to the leads**

**1. Numerical NCA results**

In this subsection, we calculate the conductance using the non-equilibrium formalism within the non-crossing approximation (NCA) for a large choice of parameters. Details on the technique are given for example in Ref. We choose here asymmetric tunnel coupling to the leads $\Delta_R = 60\Delta_L$, and set the reference energy scale as $\Delta = 2\Delta_L + \Delta_R = 1$, with a temperature $T = 0.01$ and a half-bandwidth of the leads $D = 10$. Owing to the large ratio $\Delta_R/\Delta_L \gg 1$, the impurity is effectively in equilibrium with the right lead, although the calculations were in practice performed using the full out-of-equilibrium NCA. In Fig. 2 we show $G(V)$ in units of the conductance quantum $G_0 = 2e^2/h$ for positive $E_d$ (empty orbital regime) and several values of capacitance asymmetries $\alpha$. The small spike at $V = 0$ is again an artifact of the NCA and should be disregarded. For symmetric
voltage drop, $\alpha = 1/2$, the discussion of Section III based on the non-interacting resonant level predicts two peaks of equal width $4\Delta$ and equal height $eC/(2\Delta)$ at $eV = \pm 2E_d^{0,\text{eff}} \approx 9.4$. However, the more refined analysis of Section IV based on the general features of the interacting spectral density, predicts different width renormalization: a width $4\Delta$ for the STM peak, and a width $6\Delta$ for the IV peak. These trends are well described by the full NCA calculation shown in Fig. 2. However, the conductance $G = dI/dV$ is not directly given by the spectral density, but the dependences of its effective width and weight with $E_d$ affect its shape [second term of Eq. (5)]. This is particularly important for the IV peak, for which these dependences are larger. This will be discussed in more detail in the next subsection. The width of the IV peak within the NCA is about 20% larger than $6\Delta$. This can be due to the above mentioned effects or to the limitations of the NCA at low temperatures in the IV regime. In any case, our results demonstrate that the STM and IV peaks are generically asymmetrical, and that their widths depend sensitively on the charge state present on the quantum dot.

As $\alpha$ decreases, the tendencies predicted by the simple analysis of Section III remain qualitatively valid. The position of the STM peak tends to $E_d^0$ and its width to $2\Delta$ in the limit $\alpha \to 0$. In contrast, the IV peak broadens enormously and moves to more negative values of $V$ when the capacitances become very asymmetric.

In Fig. 3 we show the results for the local moment regime at negative $E_d^0$. In this case, both peaks are much broader than the predictions of the naive expectations based on a non-interacting resonant-level spectral density. Indeed, the STM peak now lying at negative voltage shows strong interaction effects (already witnessed in the spectral density in Fig. 1) that make it twice broader without interactions (the width is $8\Delta$ for symmetric voltage drop $\alpha = 1/2$). The IV peak, now located at positive bias voltage, displays again a width that is about $3/2$ wider than the simple expectations outlined at the end of Section III due to combined charge fluctuations and Coulomb effects (the width is $6\Delta$ for $\alpha = 1/2$). The IV peak is also shifted to the left, since for $\alpha = 1/2$ one would expect it to lie at $eV = 2E_d^{0,\text{eff}} \approx 6.5\Delta$, while it is located at $5.7\Delta$. While naively one would expect a ratio of maxima inversely proportional to the width, the intensity of the STM peak is smaller than expected, especially for $\alpha = 1/2$. This further correlation effect is discussed in the next subsection where we present a more refined analytical approach. Again, as $\alpha$ decreases, the width and intensity of both peaks evolve in agreement with the expectations of the non-interacting resonant-level model.

As expected, the most drastic difference with the non-interacting resonant-level model for negative $E_d^0$ is the appearance of the Kondo peak at $V = 0$. Note that its main features are independent of $\alpha$, which will be discussed in Section IV C 2. We now turn to a physical interpretation of the numerical NCA result.

2. Phenomenological approach

The main reason for the failure of the non-interacting resonant-level model to account for Coulomb blockade peaks at asymmetric tunnel coupling to the leads is that the correct width and weight of the peak in the impurity level spectral density $\rho(\omega)$ both depend in a non-trivial way on the occupancy, as discussed in Section IV A. One of the simplest ways to modelize this effect is the alloy analogy approach to periodic Anderson models, in which the electrons with spin down are frozen to calculate the dynamics of electrons with spin up and vice versa. This approach is also related to the so-called Hubbard-III approximation in which the weight of the charge-transfer peak near $E_d$ for spin $\sigma$ (or the lower Hubbard band in a Hubbard model) is proportional to the number of unoccupied states with the opposite spin, $1 - \langle n_{-\sigma} \rangle$.

To correct the width inconsistency in the non-interacting resonant level model, we have fitted the width of the NCA peak as a function of $E_d$ at a temperature $T = 2\Delta$ (such that the Kondo peak is absent and the width of the IV peak is well behaved), with other parameters taken as in Section IV B. For simplicity, in this section and in section V we replace the notation $E_d^{\text{eff}}$ by $E_d$, incorporating the Haldane shift in the latter. From this, we obtain that the effective half-width at half-maximum can be approximated as:

$$
\frac{\Delta_{\text{eff}}}{\Delta} = \frac{3}{2} - \frac{1}{2} \tanh\left( \frac{E_d}{2.77\Delta} \right).
$$

This simple formula nicely interpolates between the empty orbital case ($\Delta_{\text{eff}} = \Delta$ for $E_d \gg \Delta$) and the local moment regime ($\Delta_{\text{eff}} = 2\Delta$ for $-E_d \gg -\Delta$), passing through the mixed valence regime ($\Delta_{\text{eff}} = (3/2)\Delta$ for $|E_d| \ll \Delta$).

The above discussion suggests that a better description of the single peak in the equilibrium local density of state can be obtained using the improved formula:

$$
\rho_R(\omega, E_d) = \frac{(1 - \langle n_\sigma \rangle)\Delta_{\text{eff}}/\pi}{(\omega - E_d)^2 + \Delta_{\text{eff}}^2},
$$

where $\langle n_\sigma \rangle = \int d\omega \rho_R(\omega) f_R(\omega)$ is the occupation of the dot for a given spin, independent of the spin in the absence of an applied magnetic field. Integration gives $\langle n_\sigma \rangle$ in terms of the digamma function $\psi(x)$:

$$
\langle n_\sigma \rangle = \frac{1}{2} - \frac{\Psi_R}{3/2 - \Psi_R},
$$

$$
\Psi_R = \frac{1}{\pi} \text{Im} \psi\left( \frac{1}{2} + \frac{\Delta_{\text{eff}} + i(E_d - \mu_R)}{2\pi T} \right).
$$

Replacing then Eq. (3) in Eq. (1) gives the current:

$$
I = C(1 - \langle n_\sigma \rangle)(\Psi_R - \Psi_L),
$$

Finally differentiating this expression, one obtains an analytical although lengthy expression for the differential
FIG. 4: (Color online) Differential conductance in the empty orbital regime $E_d^0 = 4\Delta = 4$, for asymmetric tunnel couplings ($\Delta_R = 60\Delta_L$), low temperature $T = 0.01\Delta$, and two values of the capacitance ratio $\alpha = 1/2, 1/4$. Full lines are the NCA simulations, and dashed lines are the results from the phenomenological approach.

FIG. 5: (Color online) Differential conductance in the local moment regime $E_d^0 = -4\Delta = -4$, with the same parameters as in Fig. 4. Full lines are the NCA simulations, and dashed lines are the results from the phenomenological approach.

Conductance $G(V)$, which we do not reproduce here. At $T = 0$, $G(V)$ contains two terms which have the form of those included in Eq. (4), plus some additional terms due to the dependence of $\Delta_{\text{eff}}$ and \langle n_{d\alpha} \rangle with respect to $E_d$. These additional terms affect the IV peak, for which the variation of $\Delta_{\text{eff}}$ and \langle n_{d\alpha} \rangle is significant.

In Fig. 4 we compare the conductance $G(V)$ obtained through this phenomenological approach to the full NCA result in the empty orbital case $E_d^0 = 4\Delta$ (note that the Haldane shift is included by hand in the level position within the phenomenological approach). We remind the reader that for $\alpha = 1/2$, the non-interacting resonant-level model predicts a symmetric $G(V) = G(-V)$ conductance curve, which turns out to be incorrect in the presence of strong Coulomb interaction. The phenomenological approach is thus a considerable improvement for the IV peak at negative voltage, although some quantitative discrepancies remain. In particular, the analytical expression gives an intermediate valence peak a little bit narrower and shifted towards $V = 0$, but the general trends are well reproduced by the simple approach. In addition, this discrepancy might be due to limitations of the NCA at low temperatures in this regime.

In the local moment regime for negative $E_d^0$, as shown in Fig. 5, the comparison is less satisfactory. The Kondo peak is obviously missing in the phenomenological approach and the positions of the Coulomb peaks show also some deviations, particularly for the IV peak now located at positive $V$. Nevertheless, the intensities and the widths (which are larger than those predicted by the non-interacting resonant-level model) are semiquantitatively reproduced. The small discrepancy in the position of the STM peak at negative $V$ might be due to some dependence with $E_d$ of the Haldane shift, being smaller for negative $E_d$. Having clarified the physics at play in the asymmetric situation, which simplifies because the system is in equilibrium with a single lead, we now consider the problem in its full complexity for the regime of symmetric tunnel couplings.

C. Conductance for symmetric tunnel coupling to the leads

1. Numerical NCA results

For symmetric tunnel coupling to the leads, an approximation like Eq. (4), which would permit to reduce the problem to the calculation of the spectral density at equilibrium, is no more possible. In Fig. 4 we show the results of the non-equilibrium NCA calculation for symmetric tunnel barriers, $\Delta_L = \Delta_R = \Delta/2$, and parameters similar to those of the experiment by Könenmann et al.\[15\] but with negative $E_d^0$. In contrast to the results of Section IV B 1, the symmetry now imposes that $G(V) = G(-V)$ for $\Delta_L = \Delta_R$ and $\alpha = 1/2$.

While one would expect that the non-interacting resonant-level model is meaningless in this non-equilibrium interacting case, the symmetric coupling to the leads “blurs” in the conductance the asymmetry due to correlation effects. The simplest non-interacting resonant level model with renormalized parameters is indeed able to reproduce the Coulomb blockade peaks obtained with the NCA (of course the Kondo anomaly is not captured). Except for the small modification of the $\alpha$ value,
which is hard to justify, the changes in the other parameters of the model are expected: a width 4.2 meV nearly two times larger than expected (2.30 meV) because of correlation effects, an effective $E_d$ shifted upwards due to renormalization effects (the Haldane shift is 1.04 meV for the half band width $D = 20$ meV used in the figure) and smaller intensity also due to correlation effects (weight proportional to $1 - \langle n_--n_+ \rangle$). We now turn to a detailed description of the Kondo resonance for a large range of parameters.

2. On the width of the non-equilibrium Kondo peak

In this section, we study the relation between the width of the low bias Kondo resonance in the differential conductance $G(V)$ (which is present for negative enough $E_d^0$) and the Kondo peak in the equilibrium spectral density of the impurity level $\rho(\omega)$. We define $T^G_K$ (resp. $T^\rho_K$) as the energy scale corresponding to half the width at half maximum of the peak in $G(V)$ (resp. $\rho(\omega)$) near $V = 0$ (resp. $\omega = 0$). Both quantities are of the order of the Kondo temperature, but differ from a non-universal prefactor. Since the Kondo temperature is often extracted experimentally from the differential conductance curve, but theoretically the equilibrium density of states is more standardly computed, for instance with the Numerical Renormalization Group (which has difficulties in grasping the effect of finite bias voltage), we consider here systematically the ratio $T^G_K/T^\rho_K$, for several ratios of couplings to the leads $\Delta_R/\Delta_L$, keeping $\Delta = \Delta_R + \Delta_L = 1$ fixed, and also for several values of the level position $E_d$. In this section we keep $E_d$ fixed and assume a symmetric voltage drop $\mu_L = -\mu_R = eV/2$, as $V$ is increased, and take the half band width $D = 10$. Both the differential conductance $G(V)$ and equilibrium density of states $\rho(\omega)$ are computed with the NCA, the former with the non-equilibrium formalism, and the latter at equilibrium. The resulting ratio of Kondo temperatures is shown in Table I.

Table I: Ratio $T^G_K/T^\rho_K$ of the Kondo temperatures extracted from finite bias conductance $G(V)$ and equilibrium density of states $\rho(\omega)$, obtained for different values of the level energy $E_d$ and various asymmetry ratios $\Delta_R/\Delta_L$.

| $\Delta_R/\Delta_L$ | $E_d=-2$ | $E_d=-3$ | $E_d=-4$ | $E_d=-5$ | $E_d=-6$ |
|---------------------|-----------|-----------|-----------|-----------|-----------|
| 1                   | 1.59      | 1.45      | 1.33      | 1.31      | 1.30      |
| 2                   | 1.50      | 1.19      | 1.078     | 1.09      | 1.08      |
| 5                   | 1.29      | 1.03      | 1.008     | 1.003     | ~1        |
| 10                  | 1.29      | ~1        | ~1        | ~1        | ~1        |

FIG. 6: (Color online) Differential conductance at symmetric tunnel couplings, $\Delta_L = \Delta_R$, for parameters close to the experimental measurement of Königman et al., namely $\Delta = 1.15$ meV, $E_d^0 = -4.5$ meV, $T = 0.014$ meV and $\alpha = 0.425$. The full line is the non-equilibrium NCA calculation, while the dashed line corresponds to a fit to the non-interacting resonant-level model with renormalized parameters $\Delta_{\text{eff}} = 2.1$ meV, $E_d^0 = -3.44$ meV, $\alpha = 0.45$ and a scale reduction factor 0.355.

FIG. 7: (Color online) Differential conductance near $G(V)$ (broken lines in red, corresponding to the left vertical scale), and equilibrium spectral density of the quantum dot $\rho(\omega)$ (full line in black, corresponding to the right vertical scale), computed for $\Delta = 1$, $E_d = -4$ and several tunnel asymmetry ratios $R = \Delta_R/\Delta_L = 1, 5, 30$. The coefficient $A = 4\Delta_R\Delta_L/D^2$ is introduced to normalize the conductance to the density of states.
asymmetry ratio \( R = \Delta R / \Delta L \) is increased and the impurity level position \( E_d \) is decreased negatively, which can be understood as follows. For large asymmetry of the tunnel coupling to the leads, \( R \gg 1 \), the dot can be considered at equilibrium with the right lead, and Eq. (4) becomes valid. Furthermore, in the Kondo limit for fixed chemical potential \( \mu \), the Kondo peak is insensitive to a small change in the localized level \( E_d \), and remains close to the Fermi level. For \( |eV| < T_K^0 \) this is supported by Fermi-liquid approaches.\(^{32,33}\) Thus, one can drop the second term in Eq. (4) and \( G(V) \) for \( |eV| < T_K^0 \) gives directly the spectral density of the dot state \( \rho_R(eV, E_d^0) \). This effect is evident in Fig. 7, where the evolution of the Kondo peak in conductance \( G(V) \) with increasing asymmetry ratio \( R \) is shown and compared to the equilibrium density of states \( \rho_R(eV, E_d^0) \).

V. COMPARISON WITH EXPERIMENT

In this last section, we discuss the experiment of Könenmann et al.\(^{15}\). This experiment corresponds to a value of \( E_d^0 \) that is positive and very large (two orders of magnitude bigger than \( \Delta \)). This situation is very hard for the numerical solution of the self-consistent NCA equations, but since the Kondo effect does not take place, we can compare the experiment to the phenomenological approach developed in Section IV B 2, which takes into account the changes in both width and height of the charge-transfer peak in the spectral density with the dot level \( E_d \). The experiment was done for highly asymmetric tunnel coupling to the leads, so that the main assumption of Section IV B 2 is fulfilled.

In the experiment, there is a peak at \( V = -9.7 \) mV with a half width of 76 \( \mu \)V and another one at \( V = 8.9 \) mV with a half width of 35.5 \( \mu \)V which is about five times more intense. For \( \alpha \approx 1/2 \) (as suggested in Ref. 15), the non-interacting resonant level model incorrectly predicts two peaks of equal width and height at opposite voltages (see the end of Section III). For \( \alpha = 1/2 \), our more accurate phenomenological theory predicts a ratio of about 3/2 between the width of the IV peak at negative \( V \) and the STM peak at positive \( V \), and not a ratio of about two. However, the fact that the positions of the two peaks are not symmetric with respect to zero bias shows that \( \alpha \) is in fact different from 1/2. Using \( \alpha = 0.45 \) to adjust the correct peak position, this provides an additional contribution to the ratio in the peak widths. In Fig. 8 we display our results for parameters close to the experiment. While we are unable to reproduce exactly the intensities, positions and widths of both peaks, we obtain a semiquantitative agreement with experiment. Our conductance curves can be fitted with the sum of two Lorentzians: one located at \( V = -9.97 \) mV with half-width at half-maximum 64 \( \mu \)V and peak value 1.17\( A \), another one located at \( V = 8.18 \) mV, with half-width at half-maximum 36 \( \mu \)V and peak value 4.21\( A \). In the future, comparisons to further experimental data for molecular quantum dots deep in the Kondo regime would be interesting, in order to test our predictions.

VI. SUMMARY

Motivated by experiments with semiconducting and molecular quantum dot,\(^{15,16}\) we have analyzed the complete shape of the differential conductance \( G(V) \) in the infinite-\( U \) Anderson model connected to two leads, as a function of the ratio of the potential drops between the dot and both leads, and for various tunneling asymmetries. The simplest situation is the one found in scanning tunneling spectroscopies, where both the potential drop and tunneling rates are strongly asymmetric towards the metallic surface on which a molecule is deposited. As is well-known in this situation, the conductance provides exactly a measure of the equilibrium spectral density of the quantum dot and displays only one Coulomb peak when voltage is aligned to the charge-transfer peak in the spectral density (this peak position may contain renormalization effects). When the impurity level is deeply negative, a local moment forms, and the Kondo effect leads to an additional anomaly near zero bias, both in the density of states and differential conductance.

The situation of an arbitrary ratio for the capacitance to the source and drain is more complex (even still assuming very asymmetric tunneling rates). We find that the Coulomb peak position and width in this regime not only display trivial changes expected from the modifi-
cation of the capacitance ratio, but are also corrected by many-body effects. As it is known from previous calculations\textsuperscript{28,19}, and reproduced by our NCA results, the width of the charge-transfer peak is about two times larger for large and negative level position (local moment regime) than for large and positive position (empty orbital regime). In addition, an extra Coulomb peak appears at opposite voltage values to the peak found in the STM situation, as expected from Coulomb blockade theory. However, this peak has in general a different nature, and relates to the intermediate valence situation of the underlying quantum impurity model, and displays accordingly a width renormalization of about 3/2 compared to expectations based on a non-interacting effective resonant level model. All these effects can be taken into account by a simple phenomenological approach described in Section IV B 2, which is able to describe well our numerical simulations (except for the presence of the Kondo in Section IV B 2, which is able to describe well our numerical simulations (except for the presence of the Kondo resonance) and agrees semi-quantitatively with some published experimental results [see Section IV].

We also provided some general study of the Kondo temperature extracted from the differential conductance (as typically done in experiments), and made some precise connection to the Kondo scale extracted theoretically from the equilibrium spectral density. We find that the former scale can be up to 60\% larger than the latter one, and that they coincide only for asymmetric tunnel couplings deeply in the Kondo regime.

We hope that all these results will serve as guidelines for interpreting more quantitatively experimental data obtained from molecular quantum dots, since many-body effects can quantitatively alter the predictions of basic Coulomb blockade theory.

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