Phonon-assisted tunneling regimes in diatomic molecules

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Electronic transport in diatomic molecules (two-level systems) connected to metallic contacts is analyzed in the presence of competing electron-electron and electron-phonon interactions. We show that phonon emission and absorption processes are strongly modified when a Coulomb energy \( U \) is included, as the phonons open channels that can result in destructive or constructive interference effects. Resonance conditions for these processes produce dramatic effects both in the density of states at the molecular sites, as well as in the conductance through the system. We find in particular an enhanced Rabi-assisted tunneling due to phonons, as the resonance conditions are met, which is made more evident for increasing temperatures. These effects are controllable by voltage gating of the molecular sites, and should be accessible in current experiments.

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I. INTRODUCTION

Electronic transport properties of nanometer scale structures such as natural and artificial molecules (e.g. quantum dot structures) attached to metallic reservoirs have received a great deal of attention recently. Three main reasons drive the active research in the area: i) these systems have a fundamental role in the development of the field of “molecular electronics” where molecules serve as electronic components, ii) they are relatively “simple” structures in which to address different conceptual issues regarding the nature of charge and spin transport, and iii) they represent the prototypical gateable structures at the nanoscale.

One of the most defining characteristics of these molecular systems is the strong spatial confinement that results in enhanced electron-electron interaction (EEI), and profoundly modifies their transport properties. Well-studied examples of the effect of EEI in confined systems are Coulomb blockade and Kondo effect in quantum dots. Molecular systems may also exhibit strong electron-phonon interactions (EPI), which play a significant role in their physical properties. Several works have addressed the effects of EPI in quantum-dot molecules for different geometrical arrangements, in a perturbative regime. It is known for example that phonons cause a broadening of Coulomb blockade peaks and the appearance of satellite peaks in the nonlinear transport regime. Phonons also provide relaxation mechanisms (through inelastic scattering processes) that affect the electronic conductance, specially in molecular systems as well as polaronic shifts of the electronic levels.

While the effect of these different interactions on electronic transport properties has been studied extensively in regimes where one of them is dominant, not much is known when both interactions become comparable. Recently, two works have addressed this issue. Al-Hassanieh et al. have studied the effect of electron-phonon interac-
metallic leads. We now explore the regime where the local electronic level spacing tends to zero, which makes the phase effects much more important in the coherent transport processes. We also study in detail all resonant regimes and conducting channels provided by the EPI. To solve the equations of motion, we use a Hubbard-like decoupling procedure for all the Green’s functions involved in the calculation. This approach has proven reliable in the description of properties of interacting systems at temperatures above their characteristic Kondo temperature. These calculations allow us to confirm the existence of resonance conditions affect the conductance through the many-body nature of the problem. We show that there are structures in the DOS dependences on site per site (or equivalently a quantum dot pair) with local energies $\epsilon^0_\alpha$ and $\epsilon^0_\beta$ (we assume the non-degenerate case with $\epsilon^0_\alpha > \epsilon^0_\beta$), as schematically shown in Fig. 1.

Each single-level site (or quantum dot) is connected independently to two external metallic leads. We include local (on-site) electron-electron interactions and an electron-phonon inter-site interaction. As described the model can represent various experimental geometries; in particular, this model was studied in the limit of non-interacting electrons, to explore interference effects.

The total Hamiltonian is then written as

$$H_T = H_{mol} + H_{leads} + H_{mol-leads},$$

where

$$H_{mol} = H_{el} + H_{ph} + H_{el-ph},$$

with

$$H_{el} = \sum_{\sigma,\gamma=\alpha,\beta} \left[ \epsilon^0_{\sigma\gamma} c_{\sigma\gamma}^\dagger c_{\sigma\gamma} + \frac{U}{2} n_{\sigma\gamma} n_{\bar{\sigma}\bar{\gamma}} \right],$$

$$H_{ph} = \left( b^\dagger b + \frac{1}{2} \right) \hbar \omega_0,$$

and the electron-phonon interaction is given by

$$H_{el-ph} = \lambda \sum_{\sigma} (b^\dagger c_{\sigma\alpha} c_{\bar{\sigma}\beta} + h.c.),$$

where $b^\dagger$ (b) creates (annihilates) a phonon with energy $\hbar \omega_0$, $\epsilon_i = \epsilon^0_i - eV^0_i$, and $i = \alpha, \beta$. We have assumed that $U_\alpha = U_\beta = U$, and included independent gate voltages $V^0_i$ that control the electron occupation of the molecule sites by shifting the localized energies with respect to the Fermi energy of the left and right electrodes. Also,

$$H_{mol-leads} = \sum_{i=\alpha,\beta} t_i c_{i\gamma}^\dagger c_{\gamma\alpha} + h.c.$$

and

$$H_{leads} = \sum_{k,\sigma} \epsilon_{\sigma\gamma} c_{\gamma\sigma}^\dagger c_{\gamma\sigma}.$$

$H_{mol-leads}$ connects the diatomic molecule to the leads; for simplicity, we consider $t_i = t'$, for all $\gamma = R, L$ and $i = \alpha, \beta$. Away from the Kondo regime, the main effect of the leads is to broaden the energy levels of the dots through the tunnel couplings $t'$. To describe the transport of electrons through the molecule, we calculate the local retarded Green’s functions defined in the usual way

$$\langle\langle A; B\rangle\rangle = \int i\theta(t) \langle\langle A(t), B(0)\rangle\rangle e^{i\omega t} dt,$$

where $A$ and $B$ are operators and $\langle\langle \cdot \cdot \rangle\rangle$ denotes the thermodynamic average (for $T \neq 0$) or the ground state expectation value (for $T = 0$). In the frequency domain, the equation of motion for Green’s functions is written as

$$\omega \langle\langle A; B\rangle\rangle = \langle\langle A, B\rangle\rangle + \langle\langle A, H; -; B\rangle\rangle,$$

where the subindex in the brackets denotes commutation ($-$) and anti-commutation ($+$) relations. The expression for the local electronic Green’s function is:

$$C_{ii}(\omega) = \langle\langle c_{i\sigma}; c_{i\sigma}^\dagger \rangle\rangle, \quad i = \alpha, \beta.$$

The assumption of identical leads allows the introduction of symmetric and antisymmetric fermionic operators $c_{Sk\sigma}, c_{Ak\sigma}$

$$c_{Sk\sigma} = \frac{1}{\sqrt{2}} (c_{Rk\sigma} + c_{Lk\sigma})$$

$$c_{Ak\sigma} = \frac{1}{\sqrt{2}} (c_{Rk\sigma} - c_{Lk\sigma}).$$

FIG. 1: Schematic representation of the model system. Only the electron-phonon interaction connects the local sites ($\epsilon^0_\alpha < \epsilon^0_\beta$) via phonons of frequency $\omega_0$ and with coupling constant $\lambda$. II. MODEL AND THEORETICAL METHOD

Our model consists of a diatomic molecule with a single level per site (or equivalently a quantum dot pair) with local energies $\epsilon^0_\alpha$ and $\epsilon^0_\beta$ (we assume the non-degenerate case with $\epsilon^0_\alpha > \epsilon^0_\beta$), as schematically shown in Fig. 1. Each single-level site (or quantum dot) is connected independently to two external metallic leads. We include local (on-site) electron-electron interactions and an electron-phonon inter-site interaction. As described the model can represent various experimental geometries; in particular, this model was studied in the limit of non-interacting electrons, to explore interference effects. The total Hamiltonian is then written as

$$H_T = H_{mol} + H_{leads} + H_{mol-leads},$$

where

$$H_{mol} = H_{el} + H_{ph} + H_{el-ph},$$

with

$$H_{el} = \sum_{\sigma,\gamma=\alpha,\beta} \left[ \epsilon^0_{\sigma\gamma} c_{\sigma\gamma}^\dagger c_{\sigma\gamma} + \frac{U}{2} n_{\sigma\gamma} n_{\bar{\sigma}\bar{\gamma}} \right],$$

$$H_{ph} = \left( b^\dagger b + \frac{1}{2} \right) \hbar \omega_0,$$

and the electron-phonon interaction is given by

$$H_{el-ph} = \lambda \sum_{\sigma} (b^\dagger c_{\sigma\alpha} c_{\bar{\sigma}\beta} + h.c.),$$

where $b^\dagger$ (b) creates (annihilates) a phonon with energy $\hbar \omega_0$, $\epsilon_i = \epsilon^0_i - eV^0_i$, and $i = \alpha, \beta$. We have assumed that $U_\alpha = U_\beta = U$, and included independent gate voltages $V^0_i$ that control the electron occupation of the molecular sites by shifting the localized energies with respect to the Fermi energy of the left and right electrodes. Also,

$$H_{mol-leads} = \sum_{i=\alpha,\beta} t_i c_{i\gamma}^\dagger c_{\gamma\alpha} + h.c.$$

and

$$H_{leads} = \sum_{k,\sigma} \epsilon_{\sigma\gamma} c_{\gamma\sigma}^\dagger c_{\gamma\sigma}.$$

$H_{mol-leads}$ connects the diatomic molecule to the leads; for simplicity, we consider $t_i = t'$, for all $\gamma = R, L$ and $i = \alpha, \beta$. Away from the Kondo regime, the main effect of the leads is to broaden the energy levels of the dots through the tunnel couplings $t'$. To describe the transport of electrons through the molecule, we calculate the local retarded Green’s functions defined in the usual way

$$\langle\langle A; B\rangle\rangle = \int i\theta(t) \langle\langle A(t), B(0)\rangle\rangle e^{i\omega t} dt,$$

where $A$ and $B$ are operators and $\langle\langle \cdot \cdot \rangle\rangle$ denotes the thermodynamic average (for $T \neq 0$) or the ground state expectation value (for $T = 0$). In the frequency domain, the equation of motion for Green’s functions is written as

$$\omega \langle\langle A; B\rangle\rangle = \langle\langle A, B\rangle\rangle + \langle\langle A, H; -; B\rangle\rangle,$$

where the subindex in the brackets denotes commutation ($-$) and anti-commutation ($+$) relations. The expression for the local electronic Green’s function is:

$$C_{ii}(\omega) = \langle\langle c_{i\sigma}; c_{i\sigma}^\dagger \rangle\rangle, \quad i = \alpha, \beta.$$

The assumption of identical leads allows the introduction of symmetric and antisymmetric fermionic operators $c_{Sk\sigma}, c_{Ak\sigma}$

$$c_{Sk\sigma} = \frac{1}{\sqrt{2}} (c_{Rk\sigma} + c_{Lk\sigma})$$

$$c_{Ak\sigma} = \frac{1}{\sqrt{2}} (c_{Rk\sigma} - c_{Lk\sigma}).$$
After this transformation, \( H_{\text{mol}} \) and \( H_{\text{mol-leads}} \) read as

\[
H_{\text{leads}} = \sum_{k, \sigma} \epsilon_k c_{k \sigma}^\dagger c_{k \sigma}.
\]

and

\[
H_{\text{mol-leads}} = \sum_{k, \sigma} \tilde{t} c_{\sigma}^\dagger \hat{S} c_{\sigma} + h.c.,
\]

where we have assumed \( \epsilon_{Rk} = \epsilon_{Lk} = \epsilon_k \), and \( \tilde{t} = \sqrt{2} t' \). In the symmetric-antisymmetric basis, the coupling between the molecule and the leads occurs only through the symmetric channel. The other terms in the Hamiltonian are not affected by this transformation. We obtain the relevant Green’s functions by using the equation of motion technique. As an illustrative example, let us describe the steps taken in the calculation of the Green’s function for level \( \alpha \):

\[
(\omega - \epsilon_\alpha)\langle c_{\alpha \sigma}; c_{\alpha \sigma}^\dagger \rangle = 1 + U\langle n_{\alpha \sigma} c_{\alpha \sigma}; c_{\alpha \sigma}^\dagger \rangle + \lambda\langle \langle b_{\beta \sigma}^\dagger c_{\alpha \sigma}; c_{\alpha \sigma}^\dagger \rangle \rangle + \sum_k \tilde{t}\langle \langle c_{\sigma \sigma}^\dagger; c_{\alpha \sigma}^\dagger \rangle \rangle.
\]

To obtain a closed set of equations, we apply a Hubbard-like decoupling procedure. In particular, for Green’s functions that involve operators for both levels \( \alpha \) and \( \beta \) (“mixed Green’s functions”), we close the equations by considering the following approximations:

\[
\langle b_{\alpha \sigma}^\dagger c_{\beta \sigma}^\dagger c_{\alpha \alpha}; c_{\alpha \alpha} \rangle \approx \langle b_{\alpha \sigma}^\dagger c_{\beta \sigma}^\dagger \rangle\langle c_{\alpha \sigma}; c_{\alpha \sigma}^\dagger \rangle, \tag{15a}
\]

\[
\langle b_{\beta \sigma} c_{\alpha \sigma}^\dagger c_{\alpha \alpha}; c_{\alpha \alpha} \rangle \approx \langle b_{\beta \sigma} c_{\alpha \sigma} \rangle\langle c_{\alpha \sigma}; c_{\alpha \sigma}^\dagger \rangle, \tag{15b}
\]

\[
\langle b_{\beta \sigma} c_{\alpha \alpha}; c_{\alpha \alpha}^\dagger \rangle \approx \langle b_{\beta \sigma} \rangle\langle c_{\alpha \sigma}; c_{\alpha \sigma}^\dagger \rangle. \tag{15c}
\]

The first two equations are the standard Hubbard decoupling procedure. At this level of the approximation, correlations between electron operators with opposite spins are neglected. The third equation decouples the phonon mode from the electron operators, as suggested by Zubarev. Finally, by keeping up to second order terms in \( \lambda \) in the expression of the self-energies, we obtain:

\[
G_{ii}(\epsilon) = \left\{ \frac{(\epsilon - \epsilon_i) - U(1 - \langle n_{i\sigma} \rangle) - \Sigma_{i\sigma}(\epsilon)}{\epsilon - \epsilon_i - U(1 - \langle n_{i\sigma} \rangle) - \Sigma_{i\sigma}(\epsilon)} \right\}^{-1}, \tag{16}
\]

with \( \Sigma_{i\sigma}(\epsilon) = \Sigma^{(el)}_{i\sigma}(\epsilon) + \Sigma^{(ph)}_{i\sigma}(\epsilon) \). Here \( \Sigma^{(el)}_{i\sigma}(\epsilon) \) is the electron self-energy in level \( i \) due to the leads, and \( \Sigma^{(ph)}_{i\sigma}(\epsilon) \) is the phonon-related self-energy. They have the following expressions:

\[
\Sigma^{(el)}_{i\sigma}(\epsilon) = \lambda^2 \tilde{t}^2 \tilde{g}(\epsilon) \left\{ 1 + \frac{[\epsilon - \epsilon_j - U(1 - \langle n_{j\sigma} \rangle)]\tilde{t}^2 \tilde{g}(\epsilon)}{(\omega - \epsilon_j)(\epsilon - \epsilon_j - U) - [\epsilon - \epsilon_j - U(1 - \langle n_{j\sigma} \rangle)]\tilde{t}^2 \tilde{g}(\epsilon)} \right\}, \tag{17}
\]

\[
\Sigma^{(ph)}_{i\alpha}(\epsilon) = \lambda^2 \frac{[\langle b_{\beta \sigma}^\dagger \rangle + \langle n_{\beta \sigma} \rangle][\epsilon - \epsilon_\beta + \hbar \omega_0 - U(1 - \langle n_{\beta \sigma} \rangle)]\tilde{t}^2 \tilde{g}(\epsilon + \hbar \omega_0)}{(\epsilon - \epsilon_\beta + \hbar \omega_0)(\epsilon - \epsilon_\beta - U + \hbar \omega_0) - [\epsilon - \epsilon_\beta + \hbar \omega_0 - U(1 - \langle n_{\beta \sigma} \rangle)]\tilde{t}^2 \tilde{g}(\epsilon + \hbar \omega_0)}, \tag{18}
\]

\[
\Sigma^{(ph)}_{i\beta}(\epsilon) = \lambda^2 \frac{[\langle b_{\beta \sigma}^\dagger + 1 \rangle - \langle n_{\alpha \sigma} \rangle][\epsilon - \epsilon_\alpha - \hbar \omega_0 - U(1 - \langle n_{\alpha \sigma} \rangle)]\tilde{t}^2 \tilde{g}(\epsilon - \hbar \omega_0)}{(\epsilon - \epsilon_\alpha - \hbar \omega_0)(\epsilon - \epsilon_\alpha - U - \hbar \omega_0) - [\epsilon - \epsilon_\alpha - \hbar \omega_0 - U(1 - \langle n_{\alpha \sigma} \rangle)]\tilde{t}^2 \tilde{g}(\epsilon - \hbar \omega_0)}. \tag{19}
\]

In all these expressions \( \tilde{g}(\epsilon) \) refers to the Green’s function of the leads at energy \( \epsilon \) (both leads have the same Green’s function in equilibrium). In Eq. \( \text{17} \), the expression for the self-energy \( \Sigma^{(el)}_{i\sigma}(\epsilon) \) of an electron in level \( i \), involves energies and occupation numbers of electrons in the \textit{other} level \( j \). These occupation numbers \( \langle n_{j\sigma} \rangle \) have to be calculated self-consistently. Since we are interested in the non-magnetic regime, the occupation numbers are taken to be spin-independent (\( \langle n_{j\sigma} \rangle = \langle n_{j\bar{\sigma}} \rangle \)) and they will be denoted just by \( \langle n_j \rangle \). Notice that the electron self-energy includes the \textit{indirect coupling} between the two levels due to the presence of the leads through the second term inside the brackets in the denominator of Eq. \( \text{17} \). The relevance of this term reduces as the inter-level separation in the dots increases. As a consequence, the net effect of the electron self-energy is to produce a finite width in the DOS peaks, to shift their positions by an amount \( \text{Re} \Sigma^{(el)}_{i\sigma}(\epsilon) \), and to give rise to transport interference effects due to the topology of the system.

Equations \( \text{17} \) and \( \text{18} \) are the phonon self-energies calculated up to second order in the electron-phonon interaction strength \( \lambda \). They involve phonon occupation numbers \( \langle b_{\beta \sigma}^\dagger b_{\beta \sigma} \rangle \), which for simplicity can be calculated from the free-boson Hamiltonian. Although this approximation is not essential, it is convenient to simplify the numerical calculation, and it does not introduce any artifact in the results, as it is a frequency independent \( \epsilon \)-number.

From Eq. \( \text{16} \), we obtain the total DOS by using \( \rho_{\sigma}(\epsilon) = (-1/\pi)[\text{Im}G^{(el)}_{i\sigma}(\epsilon) + \text{Im}G^{(ph)}_{i\beta}(\epsilon)] \). In the limit where the electron-phonon coupling is negligible (\( \lambda \to 0 \)), the DOS presents four Coulomb blockade peaks coming...
from the original atomic levels located at \( \epsilon_\alpha, \epsilon_\alpha + U, \epsilon_\beta \) and \( \epsilon_\beta + U \). As the electron-phonon interaction is turned on, these peaks show Rabi splittings, at characteristic energies, corresponding to phonon-absorption and/or phonon-emission processes at resonance conditions. These resonance conditions occur when the phonon energy \( \hbar \omega_0 \) matches the energy difference between the two energy levels. These resonance conditions occur when

\[
\begin{align*}
\hbar \omega_0 + U &= \Delta \epsilon \\
\hbar \omega_0 &= \Delta \epsilon \\
\hbar \omega_0 - U &= \Delta \epsilon ,
\end{align*}
\]

where \( \Delta \epsilon = \epsilon_\beta - \epsilon_\alpha \). Figure 2 shows a schematic representation of the effective levels active in each resonance. Notice that the resonance conditions do not explicitly depend on the electron occupation number (unlike the simpler approximation results in [18]). However, when the system is found in one of these resonances the splitting \( \delta_\alpha \propto \text{Re} \Sigma_{\alpha\beta}^{(ph)} \) for level \( \alpha \) (and \( \delta_\beta \propto \text{Re} \Sigma_{\beta\alpha}^{(ph)} \) for level \( \beta \), depends on the gate voltage through the occupation numbers.

Using the expressions above for the Green’s functions, we proceed to calculate the effect of electron-phonon interactions on the DOS and the conductance of the system. Transport properties are calculated in linear response in the fully interacting regime (\( U \neq 0 \)). The conductance has the following expression in the Keldysh formalism:

\[
G = 4\pi^2 t^4 \rho_L(\omega) \rho_R(\omega)|G_{11}(\omega)|^2 \big|_{\omega=\epsilon_F} .
\]

where

\[
G_{11}(\omega) = [\tilde{g}(\omega) t']^2 \left[ G_{\alpha\alpha} + G_{\beta\beta} + G_{\alpha\beta} + G_{\beta\alpha} \right] \quad \text{Eq. (22)}
\]

is the Green’s function that describes the dynamics of an electron with energy \( \omega \) from the site -1 to 1 (representing the leads, see Fig. 1) and \( \tilde{g}(\omega) \) is the Green’s function of the leads, calculated as disconnected from the molecule. As the four terms of Eq. (22) are complex quantities with different phases, their contributions to the current can interfere constructively or destructively as we discuss below. The off-diagonal Green’s functions (\( G_{\alpha\beta} \) and \( G_{\beta\alpha} \)) in the conductance are also calculated by equation of motion techniques at the same level of approximation described above. Finally, notice that we have neglected inelastic processes in the present calculation because we assume that the system is essentially at thermodynamical equilibrium, at the small bias voltages of interest.

### III. NUMERICAL RESULTS

Equations (13)-(19) are solved numerically and used to calculate the DOS and conductance in (21), as described above. For convenience, the Fermi energy is set to zero \( (\epsilon_F = 0) \). In what follows, all the physical quantities are given in units of the inter-level separation \( \Delta \epsilon \). The levels \( \epsilon_\alpha \) and \( \epsilon_\beta \) are set at 1.0 and 2.0, respectively (above \( \epsilon_F \)). The effect of an increase in the gate voltage \( V_g \) is lowering these levels (initially towards the Fermi energy). In the next few figures, we assume equal gating of both sites, so that \( V_g = V_g = V_g \).

Let us consider first the results for the interacting system in the absence of electron-phonon interaction (\( U \neq 0; \lambda = 0 \)). Figure 3 shows the results for the DOS (top) and the conductance (bottom) for the particular values \( U = 0.4 \) and \( T = 0.0025 \) (energies in units of \( \Delta \epsilon = \epsilon_\beta - \epsilon_\alpha \)). As expected, the conductance exhibits peaks located at gate voltages near \( \epsilon_\alpha, \epsilon_\alpha + U, \epsilon_\beta \) and \( \epsilon_\beta + U \); these are the well-known Coulomb blockade peaks that appear whenever each level is in resonance with the Fermi energy. Notice slight shifts from the anticipated energy values, caused by the non-vanishing real part of the electron self-energy discussed above.

Figure 4 shows the DOS as a function of gate voltage \( V_g \) and energy \( \omega \) in the presence of electron-phonon interaction. The different plots refer to different values of the phonon energy \( \hbar \omega_0 \). In all cases, the electron-electron interaction is set at \( U = 0.4 \), the electron-phonon coupling strength \( \lambda = 0.2 \), and \( T = 0.0025 \). The values for the phonon energies, \( \hbar \omega_0 = 0.6, 1.0, 1.4 \), have been chosen to fit the resonance conditions given by Eqs. (20a), (20b) and (20d), respectively. From the figure, it can be observed that in the first resonance condition for the smaller \( \hbar \omega_0 = 0.6 \) (top panel) the peaks corresponding to the levels \( \epsilon_\alpha + U \) and \( \epsilon_\beta \) are split, mixed by the phonon channels. The splitting for the peaks corresponding to the level \( \epsilon_\alpha + U \) is \( \approx 2 \delta_\alpha \) \((\approx 0 \) at a gate voltage \( V_g \approx 1.4 \)), while the splitting for the level \( \epsilon_\beta \) is \( \approx 2 \delta_\beta \) \((\approx 0.05 \) at \( V_g \approx 2 \), for example). Notice that an analysis of Eqs. (17)-(19) reveals that when level \( \epsilon_\alpha \) coincides with \( \epsilon_F \) the splitting \( \approx 2 \delta_\alpha \) becomes small. This can be understood in terms of the occupation number for the level \( \beta \) which in these conditions remains close to zero (this is seen by analyzing the value of the \( \text{Re} \Sigma_{\alpha} \), which...
approaches zero). A similar analysis shows that when the level \( \epsilon_\beta \) crosses \( \epsilon_F \), the splitting \( 2\delta_\beta \) is also small, since \( \text{Re}\Sigma_\beta \approx 0 \), a consequence of the levels \( \epsilon_\alpha \) and \( \epsilon_\alpha + U \) being far below \( \epsilon_F \) (almost full levels). The splitting of the levels in the DOS is clearly, and most importantly, manifested in the conductance as a function of the gate voltage \( V_g \), as shown in Fig. 5.

In the case of the second resonance condition, \( \hbar \omega_0 = 1.4 \) (middle panel in Fig. 4), there are two pairs of levels involved: \( \epsilon_\alpha, \epsilon_\beta \) and \( \epsilon_\alpha + U, \epsilon_\beta + U \). The effect of the resonance conditions on the conductance results to be much less important than in the previous case, as the splitting in the DOS at the Fermi level is much smaller here. Notice, however, that in this case all four conductance peaks exhibit a certain degree of splitting (see middle panel in Fig. 5), indicating that the interference effects involve two pairs of levels (as in Fig. 2b and 2c).

For the last resonance condition, \( \hbar \omega_0 = 1.4 \), the levels involved are \( \epsilon_\alpha \) and \( \epsilon_\beta + U \). These levels are quite far apart, separated by \( \Delta \epsilon + U \), which results in the splittings in the DOS and the conductance to be barely observable, except for the sharp dip/splitting in the first and last Coulomb blockade peaks (lower panel, Fig. 5). We notice that the Rabi splittings are generally less important for the last two resonance conditions. The reason for this behavior is that larger \( \hbar \omega_0 \) means that the original levels participating in the phonon processes are far apart, decreasing the splitting energies \( \delta_\alpha \) and \( \delta_\beta \). This is clearly reflected in the respective conductance curves, where the peak splittings are very small.

We have also analyzed the dependence of these results on temperature. The top panel in Fig. 5 shows the conductance as a function of gate voltage for three different temperatures, \( T = 0.0025, 0.025 \), and 0.1. A relatively weak dependence on temperature is observed, consistent with the fact that the resonance conditions are not explicitly dependent of the electron occupations. However, the self-consistency of the calculation induces an indirect occupation dependence, resulting in splittings that become more pronounced for increasing temperature (i.e., larger phonon numbers).

In order to fully probe all the resonance conditions, we fix \( \hbar \omega_0 \) and \( U \) and control the energy spacing \( \Delta \) by gating only one of the localized levels. Figure 5 shows a color map of the DOS as function of frequency and energy of the variable level \( \epsilon_\beta \). In this plot \( \epsilon_\beta \) is shifted down from above to below \( \epsilon_F \) as \( V_g \) increases; \( \epsilon_\alpha \) is kept fixed at a point just below the Fermi energy, \( \epsilon_\alpha = -0.001 \), so that \( \Delta \) also changes with \( V_g \). Since the energy spacing is not constant, we choose instead \( \Delta = D/8 \) as the energy unit, where \( D \) is the bandwidth in the leads. Let us analyze in detail the various peaks in this figure. Following a vertical line, such as \( \epsilon_\beta = 2.25\Delta \), one finds DOS peaks at \( \epsilon_\alpha, \epsilon_\alpha + \hbar \omega_0 \) and \( \epsilon_\alpha + U \) (\( \approx 0, 1, \) and 1.4), respectively. There is one more peak at \( \epsilon_\alpha + U + \hbar \omega_0 \) (\( \approx 2.4 \)), which is out of range of the vertical axis in this panel. Note that peaks at both \( \epsilon_\alpha \approx 0 \) and \( \epsilon_\alpha + U \) appear since the site \( \alpha \) is in the mixed valence regime. Now following a horizontal line, at \( \omega = -\Delta \), we find peaks corresponding to the levels \( \epsilon_\beta - \hbar \omega_0, \epsilon_\beta, \epsilon_\beta + U - \hbar \omega_0 \) and \( \epsilon_\beta + U \), respectively. One notices anticrossings of levels whenever one of the levels is a phonon replica, and crossings when both levels are “pure” electronic. The anticrossings reflect a direct interaction between the electronic levels mediated by phonons. The crossings appear because the levels interact only indirectly through the leads, an interaction that is energy dependent.

The bottom of Fig. 5 shows the conductance of the system. Note the appearance of antiresonances in the conductance whenever a crossing or anticrossing coincides with the Fermi level. The antiresonances localized near 1 and -0.4 are due to the anticrossings in the density of states discussed above. They result from the absence of...
FIG. 4: (Color online) Map of the DOS as function of gate voltage and energy in the EEI and EPI competing regime. From top (in units of $\Delta \epsilon$), with $\hbar \omega_0 = 0.6$, $\hbar \omega_0 = 1$ and $\hbar \omega_0 = 1.4$. In all panels, $U = 0.4$, $\tilde{p} = \sqrt{2}/5$, $\lambda = 0.2$, $\epsilon_0^\alpha = 1$, and $T = 0.0025$. The energy axes are shifted by $\epsilon_\alpha$, so that level $\epsilon_\alpha$ appears always at zero.

states at the Fermi level. On the other hand the dips near $-1.4$ and $0$ are due to level crossings. Notice that there is a non-zero contribution to the conductance coming from level $\epsilon_\alpha$, as it is always kept at the Fermi level. The full uninhibited $\alpha$-level conductance is evident for $\epsilon_\beta > 1.5$, where the $\beta$-level is empty. Notice yet, that the total conductance exhibits dips and even complete vanishing due to the complex pattern of interference arising from competing channels of electrons traversing the molecule through both arms of the structure whenever $\epsilon_\beta$ is near the Fermi level. In particular, the vanishing conductance at $\epsilon_\beta \simeq -0.5$ appears from the strong level anticrossing lying right at the Fermi level (see upper panel in Fig. 6).

In Fig. 7 we plot the conductance of the system as function of $\epsilon_\beta$, where the level $\alpha$ is shifted to a position below the Fermi energy ($\epsilon_\alpha = -0.1\Delta$), in such a way that it essentially does not contribute to the conductance. As we apply a gate voltage to change $\epsilon_\beta$, we observe a conductance curve consisting of three peaks. Two of them, located near $\epsilon_\beta = 0$ and $\epsilon_\beta = -1.4\Delta$ are the usual Coulomb blockade peaks, which are essentially not affected by the EPI. The effect of phonons, however, can be seen most clearly in the appearance of an additional peak near $-0.5\Delta$, due to phonon-assisted processes. This peak arises from the anticrossing of the levels $\epsilon_\alpha$ and $\epsilon_\beta + U - \hbar \omega_0$, opening a new channel for electron transport.

IV. CONCLUSION

In summary, we have studied in detail a diatomic model when both EEI and EPI are taken into account. We have presented an accurate description of the Rabi-assisted tunneling phenomena studying the Green’s functions of the system in the Coulomb blockade regime above the Kondo temperature. We have obtained well defined resonance conditions for the system. A detailed analysis of the various resonance conditions and interacting regimes shows that the most dramatic changes occur for weak EEI ($U, \hbar \omega_0 < \Delta \epsilon$), since the EPI is enhanced when the states involved in the phonon emission and absorption are close in energy. These results emphasize the rich physics involved in phonon-assisted phenomena, which we believe should be accessible in experiments. The fact that the phonon-induced resonance conditions are subtly temperature dependent makes the effect strong and sensitive to be probed in experiments.

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FIG. 5: (Color online) Conductance as function of gate voltage for system as in Fig. 4. From top to bottom, $\hbar \omega_0 = 0.6$, $\hbar \omega_0 = 1$ and $\hbar \omega_0 = 1.4$; parameters as above. Top panel also shows $T$ dependence of conductance for $\hbar \omega_0 = 0.6$; notice splittings in middle two peaks increase with increasing temperature. In the top panel the curves are offset for clarity.
FIG. 6: (Color online) Top: Map of the density of states as function of energy (vertical axis) and level position $\epsilon_\beta$ (horizontal axis). Bottom: Conductance as function of energy position $\epsilon_\beta$. Energies in units of $\Delta = D/8$. Other parameters are $\hbar \omega_0 = 1 \cdot \Delta$, and $U = 1.4\Delta$, $\tilde{V} = \sqrt{2}/5$, $\lambda = 0.2$, and $T = 0.0025\Delta$. Level $\alpha$ is kept fixed at $\epsilon_\alpha = -0.001\Delta$, essentially at the Fermi level and always contributing to the conductance and interference. Notice $\epsilon_\beta$ scale runs opposite each other in both panels.
FIG. 7: Conductance as function of position of level \(\epsilon_\beta\) for the same case as in Fig. 6 but with the \(\epsilon_\alpha\) level well below the Fermi energy, \(\epsilon_\alpha = -0.1/\Delta\), so that it does not contribute to the conductance.