Phonon-assisted tunneling and two-channel Kondo physics in molecular junctions

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The interplay between vibrational modes and Kondo physics is a fundamental aspect of transport properties of correlated molecular conductors. We present theoretical results for a single molecule in the Kondo regime connected to left and right metallic leads, creating the usual coupling to a conduction channel with left-right parity (even). A center-of-mass vibrational mode introduces an additional phonon-assisted tunneling through the antisymmetric (odd) channel. A non-Fermi-liquid fixed point, reminiscent of the two-channel Kondo effect, appears at a critical value of the phonon-mediated coupling strength. Our numerical renormalization-group calculations for this system reveal non-Fermi-liquid behavior at low temperatures over lines of critical points. Signatures of this strongly correlated state are prominent in the thermodynamic properties and in the linear conductance.

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I. INTRODUCTION

Ground-breaking experimental results in single molecular transistors during the last decade1 have greatly expanded the field of molecular electronics, opening several possibilities for technological applications and investigations of fundamental aspects of the physics of these devices. By now it is established that strong correlation effects play a key role in the electronic transport through these systems, as evidenced by the observation of the Kondo effect2 in both break junctions3,4 and scanning tunneling microscope setups.5 A clear understanding of the mechanisms involved in the emergence of the Kondo effect in molecular systems is thus of primary importance.

One possibility to advance our knowledge of these devices is by establishing analogies with the well-known transport properties of semiconductor quantum dots in the Kondo regime. This, however, proves to be a challenging approach for several reasons: molecule-lead couplings are very sensitive to the particular configurations, charging energies are significantly larger, and, more importantly, deformations and vibrational modes in the molecule play an active role in transport.6

This variety of competing effects also brings theoretical challenges, such as the interesting prospect of investigating the interplay between vibronic states and Kondo physics. Different studies have investigated the effect of electron-phonon couplings in the charge degrees of freedom of the molecule, affecting the exchange correlations leading to the Kondo effect.7–9 This issue has also been highlighted in recent experiments reporting anomalous behavior in the Kondo transport4 which have been attributed to the “dressing” of the local energies by Holstein-type phonons.7

In addition to these local effects, considerable attention has been given to effects of vibrational modes in the tunneling from the molecule to the leads. Phonon-assisted couplings by “breathing”10,11 or “center-of-mass”10,12–14 molecular modes create additional correlations with the electrons in the leads. Such phonon-mediated tunneling processes will, in general, lead to novel features in the transport properties10–14 and can be experimentally probed by conductance measurements.

We address this subject in the present work by investigating a two-channel Kondo (2chK) effect15,16 in molecular systems with center-of-mass vibrational modes. Two-channel Kondo physics, originally investigated in the context of heavy-fermion materials,15,17 has been an active topic in the area of nanostructures. In semiconductor quantum dots, several theoretical predictions18 and a recent experimental observation19 of the 2chK have highlighted the renewed interest in such strongly correlated states. In addition, phonon-assisted 2chK behavior has been predicted in effectively non-interacting systems (e.g., metallic carbon nanotubes20 and metallic break junctions21) coupled to vibronic states with Kondo-type correlations appearing in orbital (pseudospin) degrees of freedom.20,21

In this paper, we consider the Kondo regime of a singly charged molecular level connected to metallic leads, fully including both electron-electron and electron-phonon interactions, as well as phonon-assisted tunneling processes arising from a center-of-mass vibronic mode. Our numerical renormalization-group (NRG) calculations show that the presence of the extra phonon-mediated conductance channel leads to NFL behavior at low temperatures, with prominent signatures in the thermodynamic properties and in the linear conductance. The 2chK fixed point occurs over lines of critical points covering a wide range of parameters, both away from particle-hole symmetry and in the presence of deformation-induced charge-phonon couplings.

The paper is organized as follows: the model is presented in detail in Sec. II, and a discussion on the 2chK regime and the dependence of the critical parameters is given in Sec. III. In Sec. IV, we discuss the NFL signatures in the transmission phase shift and in the conductance across the junction. We give our concluding remarks in Sec. V.

II. MODEL

We consider a molecular complex (e.g., an organometallic compound, C60, etc.) in the Coulomb blockade (CB) regime
connected to metallic leads (for instance, in a metallic break junction setup). Kondo correlations appear as the molecule is tuned into a CB valley with an odd number of electrons by a plunger gate voltage. We focus on the low-bias regime, in which electronic transport is dominated by a singly-occupied molecular level of energy $\epsilon_d$ (measured from the Fermi energy in the leads and tunable by the gate voltage) with a charging energy $U$ arising from the electron-electron interactions within the molecule.

In our model, the molecule is connected to left ($L$) and right ($R$) leads by tunneling couplings proportional to the overlap between the wave functions of the molecular level and the ($s$-like) metallic states in the leads. More importantly, we consider the effect of phonon-mediated coupling through a center-of-mass (c.m.) vibrational mode, as illustrated in Fig. 1. For small displacements, the molecule-lead tunneling can be effectively written as $\tilde{V}_L(1 \pm \alpha \hat{\xi})$, where $\hat{\xi}$ is a displacement operator in the direction of the motion and $\alpha$ is a system-specific parameter, being essentially proportional to the ratio between the overlap length and the oscillation amplitude of the vibrational mode. Additionally, deformations in the chemical bonding (also illustrated in Fig. 1) will, in general, introduce an effective coupling of the center-of-mass oscillating mode and the charge state of the molecule. We model this by a Holstein-type electron-phonon term with coupling $\lambda$.

The full Hamiltonian is given by

$$H = H_M + H_{\text{Leads}} + H_{\text{Dot-Leads}},$$

where

$$H_M = \epsilon_d n_d + U n_d n_d + \lambda (1 - n_d)(a + a^\dagger) + \omega_0 a^\dagger a,$$

$$H_{\text{Dot-Leads}} = \sum_k \tilde{V}_L(1 - \alpha \hat{\xi}) \hat{c}_k^\dagger \hat{c}_{L\kappa} + \text{H.c.}$$

$$+ \tilde{V}_R(1 + \alpha \hat{\xi}) \hat{c}_k^\dagger \hat{c}_{R\kappa} + \text{H.c.},$$

$$H_{\text{Leads}} = \sum_{k, \kappa=L,R} \epsilon_k \hat{c}_k^\dagger \hat{c}_{\kappa\kappa}.$$

In the above, $\hat{c}_k^\dagger(\hat{c}_k)$ are fermionic operators that create (destroy) electrons with spin $\sigma$ in the molecule and leads, respectively, $(n_{\kappa\kappa} = \hat{c}_k^\dagger \hat{c}_k$ is the electron number operator), and $\omega_0$ is the frequency of the local center-of-mass phonon mode, with $a^\dagger(a)$ being the phonon operators ($\hat{\xi}=a+a^\dagger$). We assume the wide band limit and $k$-independent dot-lead couplings ($\tilde{V}_L = \tilde{V}_R$).

Hamiltonian (1) can be written as an Anderson impurity model coupled to two independent fermionic channels. Defining symmetric (even) and antisymmetric (odd) combinations of the electronic operators in the left and right leads
coupled to two independent fermionic channels. Defining symmetric (even) and antisymmetric (odd) combinations of the electronic operators in the left and right leads $c_{\ell(k)\kappa} = (\tilde{V}_L c_{\ell(k)\kappa} \pm \tilde{V}_R c_{\ell(k)\kappa})/2\sqrt{\tilde{V}_L^2 + \tilde{V}_R^2}$, the Hamiltonian in Eq. (1) becomes

$$H_{\text{Dot-Leads}} = \sum_{\kappa, \sigma} \tilde{V}_\sigma \hat{c}_\kappa^\dagger \hat{c}_{\kappa\kappa} + \alpha(a + a^\dagger) \hat{c}_\sigma \hat{c}_{\kappa\kappa} + \text{H.c.},$$

where $\tilde{V} = 2\sqrt{\tilde{V}_L^2 + \tilde{V}_R^2}$. For $\alpha \neq 0$, a phonon-mediated coupling to the odd channel is present. As we shall see, this has important consequences in the physics of the ground state of the system. Notice that the odd-channel coupling is present even for molecules not symmetrically coupled to the leads (i.e., $\tilde{V}_L \neq \tilde{V}_R$), a more likely configuration in experiments. For $\alpha = 0$, this term vanishes and Hamiltonian (1) corresponds to the single-channel Anderson-Holstein model, previously investigated with NRG and analytical renormalization-group methods.

The eigenstates of Hamiltonian (1) can be labeled by total charge and SU(2) spin symmetry $(Q, S)$. At the molecule site, states are also labeled by the number of phonons $m(\forall(0) = m)$. Notice that both the electron-phonon $(\propto \lambda$) and the phonon-assisted tunneling $(\propto \alpha$) terms couples $m \pm 1$ states. This last term couples only states with a difference of one electron in the odd channel and, thus, commutes with a generalized parity operator, defined by $\hat{P} \equiv (-1)^{m+1} \hat{P}_Q$, where $Q$ is the total charge in the odd channel. Therefore, for $\lambda = 0$, Hamiltonian (1) has an additional O(1) parity symmetry (which is lost for $\lambda \neq 0$, as the electron-phonon term will couple states with different parity).

We solve the two-channel problem with Wilson’s NRG technique adapted to include the phonon degrees of freedom. As it is standard in the NRG method, we map Eq. (1) into a (two-band) tight-binding Hamiltonian by performing logarithmic discretizations of the continuum spectra in the even and odd fermionic channels. In the calculations, we use a discretization parameter $\Lambda = 3$ and keep up to 2500 states in the NRG iterations, which proves to be adequate for the calculation of thermodynamic properties. We have used a cutoff of $N_{\text{ph}} = 9$ in the maximum number of phonons and checked for convergence of the results with $N_{\text{ph}}$. We assume a constant (metallic) density of states $\rho_0$ in the leads with bandwidth $D$.

III. TWO-CHANNEL KONDO PHYSICS

From the NRG spectra, we calculated thermodynamic properties for this system. A particularly revealing quantity is the contribution to the total entropy coming from the “impurity” degrees of freedom (in the present case, the molecule), defined as the difference between the total entropy $S(T)$ and the entropy calculated in the absence of the molecule $S^{(0)}(T)$.

As it has been shown by Bethe ansatz and NRG calculations in two-channel Kondo models, $S_{\text{imp}}(T) = S(T)$
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FIG. 2. (Color online) [(a) and (c)] Impurity contribution to the entropy $S_{\text{imp}}$ at the particle-hole symmetric point $\epsilon_d=-U/2$ and [(b) and (d)] at $\epsilon_d=-0.9U$. In both cases, an unstable 2chK fixed point for which $S_{\text{imp}}-k_B \log 2$ is Boltzmann's constant by $T=0$. This is illustrated in Fig. 2(c) and (d). In both cases, $S_{\text{imp}}$ versus $\alpha$ at fixed temperatures.

This NFL fixed point is reached as $\alpha$ approaches a critical value $\alpha_c$, and at temperatures below a characteristic crossover energy scale $T^*$. Deviations from $\alpha=\alpha_c$ drive the system away from this state to a more conventional Kondo-screened state (characterized by $S_{\text{imp}}=0$), indicating the unstable nature of the fixed point. This is depicted in Fig. 2(c), showing $S_{\text{imp}}$ versus $\alpha$ at fixed temperatures. At lower temperatures, a narrow peak of height $n_{\text{imp}}=1/2 \ln 2$ pinpoints the critical value $\alpha=\alpha_c$. This marks the position of the fixed point in parameter space. At higher temperatures, the broadening of these peaks indicate that signatures of NFL behavior extend over a finite range of $\alpha$.

This behavior persists away from particle-hole symmetry, as shown in Figs. 2(b) and 2(d). Interestingly, the critical value $\alpha_c$ increases as the system approaches the mixed-valence regime ($\epsilon_d=0, -U$; $\langle n_\text{d}\rangle=0, 2$) as compared with the $\epsilon_d=-U/2$ case ($\langle n_\text{d}\rangle=1$). This contrasts with the (phonon-independent) two-channel Anderson model (2chAM), for which the critical couplings are $\epsilon_d$ independent. The crossover energy scale to the NFL fixed point, related to the Kondo temperature, decreases exponentially as the system enters the mixed-valence regime, as in the 2chAM.

In the mixed-valence regime, the system flows into a Fermi-liquid (FL) fixed point with $S_{\text{imp}}=k_B \ln(2)$ at higher temperatures before entering the NFL regime [as seen, e.g., in Fig. 2(b)]. This fixed point is nonmagnetic and characterized by a purity degeneracy in the ground state (rather than the usual spin degeneracy in similar models).

As previously discussed, the $S_{\text{imp}}$ vs $T/D$ curves display the NFL plateau at low temperatures for $-U<\epsilon_d<0$ at a critical $\alpha=\alpha_c$. The critical value $\alpha_c$ increases as the system is moved away from particle-hole symmetry, reaching its highest values at $\epsilon_d=0^-, -U^-$, as depicted in Fig. 3(a). In addition, at $\alpha=\alpha_c$ the crossover temperature sharply decreases as the systems are driven away from the particle-hole symmetric point. This is illustrated in Fig. 3(b), where we define the crossover temperature $T^*$ as $S_{\text{imp}}(T=T^*)=3/4 \ln 2$. This panel shows that $T^*$ decreases exponentially as $\epsilon_d=-U$, indicating that the 2chK is only reached within the range of gate voltages for which the molecular level is singly occupied. In fact, if we find the NFL fixed point in this “local-moment” range ($-U<\epsilon_d<0$) only, indicating that, along with susceptibility calculations, the Kondo screening occurs in the “real spin” as opposed to a “pseudospin” degree of freedom (e.g., those connected to charge states, or $|m\rangle$ and $|m\pm 1\rangle$ phonon states). In this range, the value $\alpha_c$ varies nearly quadratic with $\epsilon_d$, as shown in Fig. 3(a).

Furthermore, for fixed $\epsilon_d$, both $\alpha_c$ and $T^*$ decrease as the electron-phonon coupling $\lambda$ increases [Figs. 3(c) and 3(d)]. The decrease in $T^*$ is consistent with the fact that the electron-phonon coupling renormalizes the electron-electron interaction as $U_{\text{eff}}=U-2\lambda^2/\omega_0\xi^2$ effectively driving the system into the mixed-valence regime. The decrease in $\alpha_c$ with $\lambda$ indicates that not only the molecule-lead couplings but also the coupling of the vibrational mode with the charge state of the molecule plays a role in the mechanism leading to the two-channel Kondo effect.

IV. PHASE SHIFTS AND CONDUCTANCE

We now turn to the transport properties across the molecule. We first point out that the presence of the c.m. phonon...
term in Eq. (1) breaks the “proportionate coupling” condition between left and right leads (even in a rather unrealistic symmetric coupling configuration $V_L = V_R$). For this reason, a calculation of the linear conductance via a Landauer-type formula would involve not only the local density of states, as illustrated in Figs. 2(c) and 2(d). In this case, one expects a finite-temperature signature of the NFL regime.

The phase shifts $\delta_{c(i)}(\omega)$ can be obtained from the NRG spectra using the $(Q, S, P)$ quantum numbers to label the states. We note that the parity quantum number is strictly conserved only for $\lambda = 0$. Although it is possible, in some cases, to calculate the difference $\delta_{c} - \delta_{o}$ from the $(Q, S)$ NRG spectra, in the following we use $\lambda = 0$ as it retains most of the interesting physics. We should note that, away from FL fixed points, the correspondence between the excitations in the NFL spectra and the phase shifts entering the conductance formula is only approximate. Nevertheless, we expect the conductance obtained with this prescription to give a qualitatively accurate picture in the NFL state as well, as discussed below.

Results for the conductance are shown in Fig. 4(a). For $\alpha = 0$, the familiar shape is recovered: $G = G_0$ at the particle-hole symmetric point $\epsilon_d = -U/2$ and $G \to 0$ as $\epsilon_d \to 0$. As $\alpha$ increases, the peak narrows with $G = G_0$ at the $p-h$ symmetric point for $\alpha < \alpha_c$ and $\alpha > \alpha_c$. Interestingly, as $\alpha$ approaches the critical value $\alpha_c$ at $\epsilon_d = -U/2$, a dip appears in the conductance curve.

This is an indication of the NFL behavior and a signature of the two-channel fixed point. The behavior of individual phase shifts at the particle-hole symmetric point is illustrated in Figs. 4(b)–4(d). For $\alpha < \alpha_c$ [Fig. 4(b)], $\delta_{c} \to \pi/2$ and $\delta_{o} \to 0$ at low temperatures, indicating a decoupling of the odd channel while for $\alpha > \alpha_c$ the odd channel becomes strongly coupled at low temperatures [Fig. 4(d)]. In both cases, one expects a peak in the conductance.

At the critical point $\alpha = \alpha_c$ (NFL regime), the NRG spectra is identical for both even and odd parities, as predicted by conformal field theory. In this case, our prescription for obtaining the phase shifts gives $\delta_{c} = \delta_{o}$ at low temperatures, as depicted in Fig. 4(b), causing a destructive interference and suppressing the transmission. We thus expect this result to hold, even though the individual values of $\delta_{c(i)}(\omega)$ obtained from the NRG spectra in the NFL regime are only approximate.

These signatures in the low-temperature conductance versus gate voltage curves can, in principle, identify the 2chK regime in molecular junctions. Experimentally, a fine tuning of the microscopic parameter $\alpha$ to the critical value is nonetheless a challenging task. In general, the value of $\alpha$ will be determined by specific details of the junction, such as the ratio of amplitude of the center-of-mass vibration and the length of the molecule-lead overlap. It is interesting, however, to notice that, at higher temperatures, one might obtain $\delta_{c} = \delta_{o}$ over a wider range of $\alpha$ and $\epsilon_d$ near the critical values, as illustrated in Figs. 2(c) and 2(d). In this case, one expects a finite-temperature signature of the $T = 0$ non-Fermi-liquid point.

V. SUMMARY

In summary, we have studied center-of-mass vibrational effects and phonon-assisted processes in the transport properties of a molecular junction in the Kondo regime. The interplay between electron-electron and electron-phonon interactions in this system can be described by an effective two-channel Anderson model with phonon-assisted tunnel couplings. Our numerical renormalization-group calculations for the thermodynamic properties of the effective model show non-Fermi-liquid effects below a characteristic crossover temperature over critical lines in parameter space.

We find that the crossover temperature is at a maximum at the particle-hole symmetric point and rapidly approaches zero as the system enters the mixed-valence regime. Furthermore, we find distinct signatures of the non-Fermi-liquid phase in the linear conductance.

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