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Vibration-Induced Kondo Tunneling through Metal-Organic Complexes with Even Electron Occupation Number

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We investigate transport through a mononuclear transition-metal complex with strong tunnel coupling to two electrodes. The ground state of this molecule is a singlet, while the first excited state is a triplet. We show that a modulation of the tunnel-barrier due to a molecular distortion which couples to the tunneling induces a Kondo-effect, provided the discrete vibrational energy compensates the singlet-triplet gap. We discuss the single-phonon and two-phonon-assisted cotunneling and possible experimental realization of the theory.

Tunneling through single-molecule devices is a complex phenomenon involving vibrational motion and many-particle processes in metallic leads. Vibrational effects have been observed in the sequential [1–3] and strong tunneling regime [4–6]. Phonon satellites coexist with resonance Kondo cotunneling [4] in transport through transition-metal (TM) organic complexes (TMOCs). The Kondo effect is a direct manifestation of strong correlation effects in tunneling [7]. Theoretical attention first focused on the weak coupling limit, where exchange of mechanical energy quanta with tunneling electrons (vibration assisted tunneling) [8–13] and modulation of the tunnel barriers (shuttling) [9,14] were discussed. In the strong tunneling limit the effect of vibrations on the Kondo anomaly in the linear conductance was discussed due to assisted tunneling [15–18] and also due to tunnel-barrier modulation [18]. In this Letter, however, we demonstrate that discrete vibrations through tunnel-barrier modulation can induce a Kondo effect in TMOCs with an even number of electrons by compensating for the singlet-triplet splitting at zero bias. Already for molecules of moderate size, many vibrational modes are available, underlining the importance of considering the above effect. Our effect is essentially different from the phonon-assisted Kondo effect in [17,18]: no strong electron-vibration coupling is required nor are special gate-voltage restrictions.

We study the transport through a TMOC with a TM ion secluded in a ligand cage. The cage is in tunnel contact with metallic reservoirs (surface, STM nanotip, or edges of metallic wire in electromigration or break junction geometry). Figure 1(a) illustrates this setup. We consider a TMOC with even electron number N fixed by charge and energy quantization. The ground state is supposed to be a spin singlet, and the energy of the lowest triplet excitation Δ exceeds Kondo temperature TK. The linear conductance is thus suppressed. To investigate how intramolecular vibrations may induce transport through a Kondo effect, first one should incorporate a vibronic mode in a generic tunneling Hamiltonian

\[ H = H_{\text{mol}} + H_{\text{res}} + H_{\text{tun}}. \]  

Here \( H_{\text{mol}} \) includes the 3d electron levels in a ligand field of the cage electrons, the molecular orbitals of these ligands, as well as interactions within the 3d shell and within the cage. One should take into account the three most relevant charge states including their dependence on the vibrational coordinate of the cage Q:

\[ H_{\text{mol}} = H_{\text{Q}}^{(N)} + H_{\text{Q}}^{(N+1)} + H_{\text{Q}}^{(N-1)} + T_n. \]  

FIG. 1. Schematic situation: (a) Electrode tunnel-coupled to a transition-metal-organic complex. Charging of the complex by a tunnel process deforms the outer part of the ligand cage without strongly affecting the direct coordination sphere of the metal ion and thereby the ligand-field splitting. We assume that the extra electron is localized mainly on the cage. Electrons tunnel onto the ion through the tails of the molecular state centered on the ion, which includes admixtures of the outer shell electronic states. Therefore the main effect of the charging is the modulation of the tunnel barrier between the ion-centered states and electrode. (b) Occupation of the two e-type orbitals (orbital splitting δ) discussed in the text in order of increasing many-particle energy going from left to right: 0, δ – I, δ, 2δ, where I is the exchange energy.
The last term $T_n$ is the kinetic energy of the cage distortion. The eigenstates of $H^{(N)}_Q$ are admixed to those of $H^{(N)}_R$ by the tunneling Hamiltonian $H_{\text{tun}}$ of electrons from the reservoir $H_{\text{res}}$, so that $H_{\text{tr}} = H_{\text{tun}} + H_{\text{res}}$ has the form

$$H_{\text{tun}} = \sum_{k, \sigma} E_k c_{k\sigma}^\dagger c_{k\sigma} + \hat{\nu}(Q) \sum_{k, \sigma} (d_{k\sigma}^\dagger c_{k\sigma} + H.c.). \quad (3)$$

Effectively a single electrode remains after the standard rotation of electron states [7]. We assume that there is no direct tunneling contact between the 3d electrons of the caged TM and the electrodes. However, the 3d orbitals are hybridized with ligand molecular orbitals, and tunneling becomes possible due to small overlap between the “tails” of distorted orbitals $d$ and the lead electrons [see caption of Fig. 1(a) for more details].

We write $H_{\text{mol}}$ for the low-lying many-particle states of the TMOC in the compact form $H^{(N)}_{\text{mol}} = \sum_{\Lambda, S, T_0, T} E_\Lambda(Q) X^{\Lambda, \Lambda'}$, where $X^{\Lambda, \Lambda'} = |\Lambda \rangle \langle \Lambda'|$ are so called Hubbard operators describing transitions between four eigenstates $\Lambda$, $\Lambda'$ of the TMOC corresponding to singlet ($S$) and triplet ($T_0, T = \pm$) states. All many-particle effects and their coordinate dependence are included in the energies $E_\Lambda(Q)$. Similarly, states of the charged TMOC with $N \rightarrow N \pm 1$ are denoted as $|\gamma\rangle$. The microscopic origin of the low-energy singlet ($S$) and triplet ($T$) is most easily conceived as follows. The evenly occupied ligand cage is in a singlet spin state, and it does not affect the structure of the spin multiplet. We assume that the ligand field of the cage has low symmetry, so that the fivefold orbital degeneracy is completely lifted. For instance, for a distorted tetrahedral symmetry of the ligand field we focus on the configuration $d^2(e^2)$ with two $e$-orbital levels split by $\delta$ due to this distortion. In case of distorted cubic symmetry, the same $S$-$T$ multiplet arises for the configuration $d^6(t^2e^2)$. Figure 1(b) illustrates the occupations of the orbital of the TMOC: it is seen that in the case of a weak intrashell exchange $I < \delta$ the ground state is singlet $S$ and the lowest excitation is the $S$-$T$ transition. The many-particle energy difference $\Delta = E_T - E_S = \delta - I$ results from competition between orbital energy and exchange energy gain. We assume $\Delta$ to be larger than the Kondo temperature $T_K$. The many-particle excitation energies of two singlet excited states $S^\prime$ and $S^\prime\prime$ are $\delta$ and $2\delta$, respectively. We ignore these singlet excitations since they are not involved in the Kondo tunneling.

We use the simplest approximation of a single harmonic vibration mode with frequency $\Omega$. We assume that the relative shifts of the harmonic potentials in different excited and charged states are negligibly small; i.e., for all $x = \Lambda, \gamma$ we have $E_x(Q) = E_x + \Omega Q^2 / 2$ and $T_n = \Omega P^2 / 2$. The coordinate $Q$ is normalized to the zero-point motion. In this weakly nonadiabatic limit, vibrations are slow compared to the electron motion on the molecule. No vibrational excitations can be induced by the tunneling in the linear transport regime, unless the distortion of the ligand cage $Q$ affects the tunnel amplitude $\hat{\nu}(Q)$. Then virtual local phonons can be emitted and absorbed. The tunneling Hamiltonian may be rewritten in the form

$$H_{\text{tun}} = \hat{\nu}(Q) \sum_{k, \Lambda, \sigma} [X^{\Lambda, \sigma} c_{k\sigma} + H.c.] \quad (4)$$

(here the second summation is restricted by spin selection rules). This situation is opposite to the antiadiabatic limit where the vibrations are fast and many phonon excitations are involved in the Kondo exchange [16].

Thus we consider the effect of the modulation of the tunnel amplitude on the linear transport in the Coulomb blockade regime. In this regime one has to eliminate the tunneling term from the Hamiltonian by summing over virtual processes where $N \pm 1$ electrons occupy the molecule and vibrational quanta are excited in these virtual states. This procedure known as a Schrieffer-Wolff (SW) transformation, leads us to a Kondo Hamiltonian [Eq. (6) below] for a $S$-$T$ multiplet [19] and an oscillator with electron-phonon interaction built into the exchange coupling. According to Ref. [19], transitions within the singlet-triplet spin manifold are described by two vectors $S$ and $R$, where $S$ is the usual spin 1 vector and $R$ is a vector describing $S$-$T$ transitions. These two vectors are constructed by means of Hubbard operators $X^{\Lambda, \Lambda'}$ in the following way [19]:

$$S^+ = \sqrt{2}(X^{10} + X^{0-1}), \quad S^z = X^{11} - X^{-1-1}, \quad R^+ = \sqrt{2}(X^{15} - X^{-5-1}), \quad R^z = -(X^{05} + X^{50}). \quad (5)$$

The effective Hamiltonian arising after the SW transformation has the form

$$H_{\text{eff}} = H_{\text{res}} + \frac{1}{2} \Delta S^2 + \tilde{J}_S S \cdot S + \tilde{J}_R R \cdot R + \Omega \frac{P^2}{2}. \quad (6)$$

The electron spin operator is given by the conventional expansion $S = \frac{1}{2} \sum_{ij} c_{i\sigma}^\dagger \sigma_{ij} \tau_{ij} c_{j\sigma'}$. The Pauli vector. The exchange coupling constants $\tilde{J}_S$ and $\tilde{J}_R$ are estimated as $\tilde{J}_S(Q) = \sum_{\gamma} |\hat{\nu}(Q)|^2 / |E_T - E_\gamma|$ and $\tilde{J}_R = \alpha \tilde{J}_S$. Here $\alpha < 1$ is a coefficient arising because of the admixture of singlet states $S^\prime, S^\prime\prime$ to the ground state [19]. In the weakly nonadiabatic regime the kinetic energy in the denominator has been neglected and the nearly identical Q dependence is practically cancelled out in the addition energies. Thus, no multiphonon replicas appear in the denominators of $\tilde{J}_{S,R}$, unlike in Ref. [16]. The main source of phonon emission or absorption in our case is the tunneling rate $|\hat{\nu}(Q)|^2$. Expanding it in the quantized displacement operator $Q = (b^\dagger + b)/\sqrt{2}$ we come to phonon-assisted exchange vertices presented in Fig. 3. In accordance with Fig. 2, we retain only single-phonon processes for $\tilde{J}_S(Q)$ and only two-phonon processes for $\tilde{J}_R(Q)$. $\tilde{J}_S(Q) = J_S + j_S Q^2$ and $\tilde{J}_R(Q) = J_R + j_R Q$. It is obvious that $j_S \ll j_R$.

To draw these vertices we used the fermionic representation for the operators $[(5)]$.
The Kondo temperature characterizing this channel depends on the phonon-assisted exchange constant $j_S$,
\[ T_K^{(2)} \sim D \exp\left(-\frac{1}{A\rho j_S}\right) \ll T_K^{(1)}. \]  
(11)

One concludes from these calculations that the single-phonon processes are sufficient to compensate the energy of the $S$-$T$ splitting and induce resonance tunneling through the TMOC provided a local vibration mode with appropriate frequency satisfying the condition that
\[ |\Omega - \Delta| \leq T_K^{(1)} \]  
(12)
exists in the cage. One can expect in this case a significant enhancement of the tunnel conductance already at $T > T_K^{(1)}$ according to the law $G/G_0 \sim \ln^{-2}(T/T_K^{(1)})$ [7], where $G_0$ is the conductance at unitarity limit $T \to 0$.

Thus we formulated in this Letter conditions under which phonons are not only involved in Kondo screening but even induce Kondo tunneling. In is worth mentioning that in spite of the fact that the Kondo effect exists in our case only under phonon assistance, the Kondo temperature (11) is the same as in the usual Kondo effect. Since $T_K$ is high enough ($\sim 10$ K) in electromigrated junction experiments with a TMOC deposited between contacts [4–6], the effect predicted in this work seems to be easily observable. The crucial point is the existence of the phonon satisfying condition (12) in a TMOC with the $S$-$T$ multiplet as a lowest spin excitation. One should note, however, that even if this condition is not exactly satisfied, one may tune the system by applying the magnetic field. Then the triplet is split, and only the level $E_T = E_T - E_Z$ is involved in the phonon induced Kondo tunneling ($E_Z$ is the Zeeman energy). Then $\Delta$ in (12) is substituted by $\Delta_Z = \Delta - E_Z$, and $E_Z$ may be tuned until the inequality is satisfied. Thus the vibration gives rise to a magnetic field induced Kondo effect at Zeeman energies which can be much smaller than $\Delta$. The Kondo screening takes place due to the processes presented in Figs. 2(a) and 3(d). The only difference is that in this case the effective spin of the TMOC is one-half instead of one [20]. The theory of this effect will be presented in a separate publication.

**FIG. 3** (color online). (a) Bare exchange vertices $J_{S,K}$; (b) - single-phonon correction $J_{S}$ to the vertex $J_{K}$; (c) two-phonon correction $J_{K}$ to the vertex $J_{S}$; (d),(e) renormalized vertices $\gamma_{1,2}$ corresponding to the processes illustrated by Figs. 2(a) and 2(b), respectively.
Another way to tune the condition (12) is to stretch the break junction and thereby distort the TMOC and change the frequency $\Omega$. Such mechanical control was demonstrated experimentally for the $H_2$ molecule [21]. There it was also shown that the isotope effect may be used for the same purpose.

The generic feature of the phonon-assisted Kondo screening discussed in this work is that only the virtual phonon excitations enter the cotunneling amplitude to compensate the $S-T$ energy gap $\Delta$, so that the Kondo effect manifests itself as a zero-bias anomaly. Another mechanism of such compensation was discussed in Ref. [22], where the energy deficit was covered by the conduction electron acceleration at finite bias. In that case, the Kondo regime arises in nonequilibrium condition, so it is fragile against dephasing effects [22,23]. In our case the system remains in thermodynamic equilibrium around a zero-bias anomaly, so that the only limiting factor of this sort is the lifetime of local vibration mode, which is usually long enough in comparison with $h/T_K$.

To conclude, we demonstrated in this work that phonon emission or absorption can induce Kondo tunneling in a transition-metal-organic complex with even electron occupation and a spin singlet ground state, when the conventional Kondo effect is suppressed. Unlike the situation studied in the current literature [15–18], where the influence of real phonon excitations on the conventional Kondo effect is discussed, and various kinds of sideband satellites due to the polaronic effect are considered, we appeal to virtual phonon excitation, so that the system remains in a quasielastic tunneling regime. Because the gate potential can tune the energetic costs of the various virtual processes that contribute to the ordinary Kondo effect, elastic cotunneling, vibrationally inelastic cotunneling, etc., further work beyond the scope of this Letter is required to assess the expected gate dependence of the Kondo resonance. One of essential ingredients of our theory is that we use the dynamical symmetry of the TMOC, which characterizes both the spin algebra of localized spin itself and transitions between various levels of different spin multiplets [19]. In our case the dynamical symmetry group is SO(4).

Since the tunnel contact between magnetic ion and metallic reservoir in TMOC is mediated by a ligand cage, one may be sure that the relevant vibration excitations are the local phonons characterizing this cage. Although we confined ourselves to a specific model with two electrons in an $e$ subshell, the mechanism is quite general. The theory may be easily modified for any system with the same structure of the lowest spin multiplets. One of such examples is the so-called “Fulde molecule” [24] schematically representing the spectra of lanthanocenes, where the rare-earth magnetic ion is sandwiched between two rings of CH radicals. In that case the number of electrons in the cage and in the $4f$ shell is odd. Another candidate is the endofullerene family with atoms [25] or magnetic ions within a carbon cage [26]. The theory of vibration-induced Kondo effect may be also generalized for the case of degenerate modes and for more complicated spin-multiplets including half-integer spins. These issues will be discussed in forthcoming publications.

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