The charge-memory effect, bistability and switching between charged and neutral states of a molecular junction, as observed in recent STM experiments, is considered within a minimal polaron model. We show that in the case of strong electron-vibron interaction the rate of spontaneous quantum switching between charged and neutral states is exponentially suppressed at zero bias voltage but can be tuned through a wide range of finite switching timescales upon changing the bias. We further find that, while junctions with symmetric voltage drop give rise to random switching at finite bias, asymmetric junctions exhibit hysteretic behavior enabling controlled switching. Lifetimes and charge-voltage curves are calculated by the master equation method for weak coupling to the leads and at stronger coupling by the equation-of-motion method for nonequilibrium Green functions.

Memory effects and switching at the molecular scale are in the focus of present experimental and theoretical studies within molecular electronics [1,2,3,4,5,6,7]. Beside stochastic switching in single-molecule junctions [4], recent STM experiments [2,3] show multistability and current-voltage curves. In this approximation it is not necessary to include Coulomb interaction explicitly, though one can additionally incorporate charging effects.

It was suggested [2,7] that bistability between charged and neutral states can be accounted for in a single-level model, when one electron level is coupled to one vibrational state, the free vibron of frequency \( \omega_0 \), the occupied (charged) state of the interacting system will have the energy \( \epsilon_1 = \epsilon_0 - \epsilon_p \), where \( \epsilon_p \) is so-called polaron shift (or recombination energy). Neutral and charged (polaron) states correspond to local minimums of the potential energy surface and are metastable, if the electron-vibron interaction is strong enough. Applying an external voltage, one can change the state of this bistable system, an effect that is accompanied by hysteretic charge-voltage and current-voltage curves. In this approximation it is not necessary to include Coulomb interaction explicitly, though one can additionally incorporate charging effects.

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Figure 1: (Color online) (a) The energy diagram of the single-level electron-vibron model, coupled to left and right lead (or tip and substrate in the case of STM). (b) Franck-Condon matrix elements \( M_{mn} \) for weak (\( g = 0.1 \), squares), intermediate (\( g = 1 \), triangles), and strong (\( g = 10 \), circles) interaction.
the left and right electrical potentials. \( \varphi_0 \) describes the shift of the molecular level by the bias voltage and can be written as \( \varphi_0 = \varphi_R + \eta(\varphi_L - \varphi_R) \), \( \eta \in [0, 1] \).

The coupling to the leads is characterized by the level-width function \( \Gamma_\ell(\epsilon) = 2\pi \sum_k |V_{\ell k}|^2 \delta(\epsilon - \epsilon_k) \), where the coupling \( V_{\ell k} \) is assumed to be energy-independent (wide-band limit). The full level broadening is given by the sum \( \Gamma = \Gamma_L + \Gamma_R \).

Consider first the case of very weak coupling to the leads, \( \Gamma \ll \omega_0, \epsilon_p \). Using the polaron (Lang-Firsov) \cite{13, 14, 15} canonical transformation, it is easy to show that the eigenstates of the isolated system \( (\Gamma = 0) \) are

\[
|\psi_{nm}\rangle = e^{-\frac{\lambda^2}{\omega_0}(a^\dagger - a)^d} \hat{d}^n \hat{d}^\dagger \frac{(a^\dagger)^m}{\sqrt{m!}} |0\rangle
\]

with the energies

\[
E_{nm} = \epsilon_1 n + \omega_0 m, \quad \epsilon_1 = \epsilon_0 - \frac{\lambda^2}{\omega_0}, \quad \epsilon_p = \frac{\lambda^2}{\omega_0}.
\]

When the system is weakly coupled to the leads, the polaron representation, Eqs. \((2, 3)\), is a convenient starting point. \( n \) denotes the number of electrons, while the quantum number \( m \) characterizes vibronic eigenstates, which are superpositions of states with different number of bare vibrons. The qualitative picture of the sequential tunneling through a polaronic state is given in Fig. \( \text{(a)} \). Here the potential energies of the neutral and charged states are sketched as a function of the vibronic coordinate \( x \). When the external voltage is applied, the energy levels are shifted depending on the asymmetry parameter \( \eta \). It should be noted that this type of the energy diagram is quite general for charge-controlled bistable systems.

In the sequential tunneling regime the master equation for the probability \( p_{nm}(t) \) to find the system in one of the polaron eigenstates \( (2) \) can be written as \( \text{(4)} \). Here the first term describes the tunneling transition \textit{into} the state \( |n, m\rangle \) and the second term the transition \textit{out} of the state \( |n, m\rangle \). \( I^V[p] \) is the vibron scattering integral describing the relaxation of vibrons to equilibrium. The transition rates \( \Gamma_{nm} \) are found from the tunneling Hamiltonian (the last term in Eq. \( (1) \)). Taking into account all possible single-electron tunneling processes, we obtain the incoming and outgoing tunneling rates at zero bias voltage as

\[
\Gamma_{nm}^{10} = \sum_{i=L, R} \Gamma_i (E_{1m} - E_{0m}) \frac{M_{nm}}{2} f^0(E_{1m} - E_{0m})^2,
\]

\[
\Gamma_{nm}^{01} = \sum_{i=L, R} \Gamma_i (E_{1m} - E_{0m}) \frac{M_{nm}}{2}
\]

\[
\times (1 - f^0(E_{1m} - E_{0m})^2) \quad \text{(5)}
\]

For energy-independent \( \Gamma_i \) (the wide-band limit) we obtain the simple analytical expression

\[
\tau_{00}^{-1} = \frac{1}{\Gamma} \sum_m e^{-\frac{g^m}{m!} f^0} \left( \epsilon_0 - \frac{\lambda^2}{\omega_0} + \omega_0 m \right) \quad \text{(9)}
\]

Figure 2: Inverse life-time \((\tau \Gamma)^{-1}\) of the neutral state (thin solid line) and the charged state (thick gray solid line) as a function of \( \lambda/\omega_0 \) at \( \epsilon_0 = \lambda^2/2\omega_0 \); and the same at \( \epsilon_0 = 0.9\lambda^2/\omega_0 \) (dashed lines), \( kT = 0.1\omega_0 \).
The corresponding expression for the life time of the charged state is (assuming that the equilibrium electrochemical potential in the leads is zero)
\[
\tau_{10}^{-1} = \Gamma \sum_m e^{-g\lambda m} \left( -\epsilon_0 + \frac{\lambda^2}{\omega_0} + \omega_0 m \right) .
\] (10)

The dependence of the tunneling rates on the scaled electron-vibron interaction constant \(\lambda/\omega_0\) is shown in Fig. 2. It is clearly seen that at large values of \(\lambda\) the tunneling from the neutral state to the charged state and vice versa is exponentially suppressed in comparison with the bare tunneling rate \(\Gamma\). Hence both states are (meta)stable at low temperatures and zero voltage.

Based on the experimental parameters of Ref. 2, the charged ground state is assumed to be below the equilibrium Fermi energy of the leads, while the neutral ground state is above it. In the experiments 2 the observed relaxation energy \(\epsilon_\sigma \approx 2.4\ eV\) leads to the parameter \(\lambda/\omega_0\) of the order 5 to 10. Thus the system is in the blockade regime at zero voltage, see Fig. 2.

Next we consider the other important question, whether fast switching between the two states is possible. At finite voltage the switching rates are
\[
\tau_{00}^{-1} = \sum_m e^{-g\lambda m} \left[ \Gamma_L f_0^0 (e_0 + \omega_0 m - (1-\eta)eV) 
+ \Gamma_R f_0^0 (e_0 + \omega_0 m + \eta eV) \right] ,
\]
\[
\tau_{10}^{-1} = \sum_m e^{-g\lambda m} \left[ \Gamma_L f_0^0 (-e_0 + \omega_0 m + (1-\eta)eV) 
+ \Gamma_R f_0^0 (-e_0 + \omega_0 m - \eta eV) \right] .
\] (11)

The voltage dependence of the inverse life time \((\tau\Gamma)^{-1}\) is shown in Fig. 3 for a junction with the same tunneling coupling, \(\Gamma_L = \Gamma_R\), but asymmetric electrical field \((\eta = 0)\), as well as for the completely symmetric junction \((\eta = 0.5)\). The results in Fig. 3 imply that in both cases one can tune \((\tau\Gamma)^{-1}\) upon sweeping the bias voltage and thereby control the timescales for switching between charged and neutral states. For the symmetric junction both switching rates, \(\tau_{00}^{-1}\) and \(\tau_{10}^{-1}\), (dashed lines) are simultaneously nonzero at finite voltage \((eV/\omega_0 \geq 40)\) for the parameters of Fig. 3 leading to random switching (noise) sketched as dashed line in the inset. On the contrary, for the asymmetric junction controlled switching into the neutral (black solid line) and charged (grey line) state can be achieved at large enough negative and positive voltage, respectively. This qualitatively different behaviour is a result of the distinct voltage asymmetry of the two inverse lifetimes which are never both finite. The further peculiar feature of the asymmetric case, namely that the switching rates of the neutral and charged states interchange their role as a function of bias, i.e., the neutral (charged) state is long-lived at negative (positive) bias, implies hysteretic behavior and a memory effect.

To this end we consider what happens, if one sweeps the voltage with different velocity (Fig. 4) for the asymmetric case \(\eta = 0\). If the voltage is changed fast enough, i.e. faster than the life time of charged and neutral states \((\tau \gg \tau_\sigma\) as discussed in the introduction), then both states can be obtained at zero voltage (hysteresis). In the opposite (adiabatic) limit the change is so slow that the system relaxes into the equilibrium state, and the population-voltage curve is single-valued. Note that this controlled switching is possible only for asymmetric junctions for the reason given above.

We finally compare the results with those of a further important limiting case, namely that the level width is finite (and possible finite dissipation of vibrons is taken into account). Then the master equation approach can no longer be used, and we apply alternatively the nonequilibrium Green function technique. Follow-
ing Refs. [21, 22], the average number of electrons is determined by the lesser Green function $G^< (t_1 - t_2) = i \langle d(t_2) d(t_1) \rangle$ as $\langle n \rangle = -i \int G^< (\epsilon) \frac{d\epsilon}{2\pi}$. The calculation of the Green function is a nontrivial task even in the single-level model. It is simplified in the important limit of the Green function is a nontrivial task even in the spectral and the distribution function

$$A(\epsilon) = \frac{2\Gamma}{(\epsilon - \epsilon_0 - \lambda \varphi_n)^2 + \Gamma^2},$$

$$f(\epsilon) = \frac{\Gamma_L f_R^0 (\epsilon - \epsilon \varphi_L) + \Gamma_R f_L^0 (\epsilon - \epsilon \varphi_R)}{\Gamma_L + \Gamma_R}.$$

The result is qualitatively the same as in the sequential tunneling case: For electrically asymmetric junctions two stable states exist at zero bias (memory effect), which can be switched by the voltage. The current shows similar hysteretic behaviour as a function of voltage. For the symmetric junction hysteresis is observed only at finite voltage (nonequilibrium bistability [9]). Hence, asymmetric junctions are again preferable for a memory effect.

Finally we note that in the case $\omega_0 \ll \Gamma$ we considered the stationary problem only, assuming that the switching rate between the two metastable states is small (compared e.g. to $\Gamma$) at large $\lambda / \omega_0$. The calculation of the life times of metastable states within the Green function approach and of dynamical effects arising from the competition between voltage sweeping and switching times, such as in Fig. 4, remains as a problem for the future.

To conclude, we considered a charge-memory effect and switching phenomena in single-molecule junctions taking into account dynamical effects such as the interplay between timescales of voltage sweeping and switching rates. We showed that bistability arises if quantum transitions between neutral and charged states involved are suppressed, e.g. due to Franck-Condon blockade. Different regimes, characterized by random mutual transitions and by single switching events into a stable configuration are identified. In the latter case controlled switching of the molecule is achieved by applying finite voltage pulses.

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Figure 5: Average number of electrons at $\Gamma_L = \Gamma_R = 5\omega_0$ as a function of normalized voltage $eV / \omega_0$ for the asymmetric junction, $\eta = 0$ (thin solid line), and for the symmetric junction, $\eta = 0.5$ (dashed line), for $\lambda / \omega_0 = 5$ and $\epsilon_0 = \lambda^2 / \omega_0$. 

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