

# Low-temperature control of nanoscale molecular dynamics

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A novel *in situ* probe of the nanoscale molecular dynamics of organic-molecule and fullerene-tube nanostructures is proposed. General and consistent results for the nonlinear-current coupling to the nanostructure excitations are presented to document a frequency-selective electrostatic control of this current stimulation and optimal operation as a local source of current-induced molecular excitations  $\Omega_j$ . The control is possible for temperatures  $T \ll \Omega_j$ . Finally, it is explained in detail how Raman measurements of this molecular dynamics would probe the nanoscale excitations within organic and fullerene nanostructures under non-linear transport conditions.

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A successful future molecular-electronics technology requires an understanding of the fascinating nanoscale molecular devices themselves, of their nonlinear and interacting transport properties, and of the current-induced molecular dynamics. Fullerene tubes (nanotubes) [1] and organic molecules [2], which assemble between metal contacts to form organic nanostructures, offer interesting candidates for such a molecular-transport program. These organic nanostructures and the fullerene tubes implement, for example, the molecule resonant-tunneling diode [3], the single-Bucky-ball ( $C_{60}$ ) transistor [4], and the nanotube field-effect transistor [5], which achieves room-temperature operation and a nanoscale-feature size in all but the transport direction. Fullerene-tube heterostructures can be identified by experimentally observed kinks [6] and permit additional nanoscale molecular-electronic devices by combining sections of different local chirality [7] and thus a different nature of electron conduction [8]. The experimental selection of single-kink heterojunctions produce the nanotube equivalent of current-rectifying diodes [6]. A corresponding selection [6], and/or proposed engineering [9], of double-kink nanotube samples produces either the nanotube quantum dot [10] or the robust, i.e., temperature- and scattering-insensitive, nanotube resonant-tunneling transistor [11], which achieves a nanoscale-feature size in all dimensions.

Nanostructure device robustness is of central importance as, e.g., experiments on single-Bucky-ball transistors [4] and on current-induced atom/molecule manipulation [12], document significant molecular excitation induced by the nonlinear transport [13]. The coupling to this molecular excitation can even provide novel transport mechanisms as in the electron shuttle [14]. A molecular-electronics program must characterize devices both in terms of the nonlinear molecular current and in terms of the nonlinear-transport coupling to the molecule-structural dynamics — the molecular excitation.

Here I document frequency-selective electrostatic control of the current-induced molecular excitations and propose a probe of this local nanostructure dynamics. The control and suggested molecular probe work for characteristic excitation frequencies  $\Omega_j = \Omega_{0,1,\dots}$  that are larger than the temperatures  $T$  and would operate at the relevant nonequilibrium device condition, i.e., with the nonlinear transistor current enabled. For use in the molecular-dynamics probe, furthermore detail optimal operation as a strong frequency-selective source of current-induced molecular excitations at the large frequencies  $\Omega_j \gg T_{\text{room}}$  that characterize the local fullerene-tube nanostructure dynamics [15]. The suggested nanostructure probe extends an earlier proposal by Narayanamurti [16] that used a burst of incoherent (acoustic) phonons to identify and map defects deep

inside semiconductor heterostructures. Here, instead, propose (1) to exploit the nonlinear nanostructure transport conditions for a direct *in situ* and controlled excitation of the relevant high-energy vibrations  $\Omega_j - \Omega_{0,1,\dots}$  and (2) to use surface-enhanced Raman spectroscopy [15,17] to measure the resulting molecular excitation  $\delta N_{\text{vib}}(\Omega_j)$ , establish the associated decay,  $1/\tau_j$ , and thus probe the density of material defects [16], the mutual coupling between such vibrations, and the intrinsic nature of the excitation (phonon) propagation [18,19].

Figure 1 illustrates a pair of resonant-tunneling systems that could produce a strong source of molecular excitations and hence the proposed nanostructure-dynamics probe. The upper scheme involves an organic nanostructure [3] (ORTN) — an organic molecule assembled between and connected both mechanically and electrically to the source and drain metal contact through the synthesized inclusion of sulphur atoms S. The lower scheme involves a double-kink nanotube resonant-tunneling heterostructure (NRTN) [11], in which metallic nanotube leads (grey tubes) connect to the metal contacts (wedges), e.g., scanning-tunneling microscope tips. In both schemes the central barrier region traps a single resonant level of energy  $E_{\text{orb}}$  and connected by tunneling rates  $\Gamma_{L/R}$  to the surrounding metal leads or contacts. A close metal gate, e.g., another metallic nanotube (grey ring at potential  $\Phi_{\text{gate}}$ ) adjusts  $E_{\text{orb}}(\Phi_{\text{gate}})$  and enables a gate-controlled resonant-tunneling transport [4,11, 20,21].

The proposed nanostructure-dynamics probe exploits this gate-control of the nanostructure tunnel-

ing transport by a selective optimization of the current-induced molecule stimulation. To document the suggested operation, this paper (i) provides a nonequilibrium conserving Green-function calculation [13,22] of the current-induced spontaneous emission (net absorption) rates  $R_{\text{sp}}(E_{\text{orb}}; \Omega_j)$  [ $R_{\text{ab}}(E_{\text{orb}}; \Omega_j)$ ] as a function of  $E_{\text{orb}}(\Phi_{\text{gate}})$  and at a set of characteristic excitation frequencies; (ii) details how phase-space restrictions that rest on the Pauli exclusion principle permit a frequency-selective gate control of  $R_{\text{sp,ab}}(E_{\text{orb}}; \Omega_j)$ ; and (iii) identifies nonequilibrium tunneling conditions that can maximize and/or inhibit the resulting effective current excitation  $\delta N_{\text{vib}}(\Omega_j)$ . At a given gate voltage and thus resonant- energy position  $E_{\text{orb}}(\Phi_{\text{gate}})$ , determine an excess nanostructure vibrational population [23]:

$$\frac{1}{\tau_j} \delta N_{\text{vib}}(E_{\text{orb}}(\Phi_{\text{gate}}); \Omega_j) = R_{\text{sp}}(E_{\text{orb}}; \Omega_j) - R_{\text{ab}}(E_{\text{orb}}; \Omega_j) \delta N_{\text{vib}}(E_{\text{orb}}(\Phi_{\text{gate}}); \Omega_j). \quad (1)$$

The local electrostatic-field control  $E_{\text{orb}}(\Phi_{\text{gate}})$  [4,11,20] can, for example, produce a strong (frequency-selective) burst of nanostructure excitations  $\delta N_{\text{vib}}(\Omega_0)$ . Simultaneous Raman measurements [15] of the strength of the anti-Stoke Raman signal at nanostructure-vibration frequency  $\Omega_0$  can thus determine the decay  $1/\tau_0$  of this excess population [17]  $\delta N_{\text{vib}}(\Omega_0)$  and probe intrinsic mechanisms [16,18,19] affecting the nanoscale molecular dynamics.

### Electrostatic control of resonant-tunneling transport

A theory description of the electrostatic gate control  $E_{\text{orb}}(\Phi_{\text{gate}})$  exists for both the organic nanostructure [20] and the fullerene heterostructure [11] scheme (Fig. 1, upper and lower schemes). In both transport schemes this local electrostatic control permits current-switch and transistor effects [4,11,20] in which transport is focused onto a single molecular level. Such nanostructure transistors improve the semiconductor resonant-tunneling transistor design [21] through a dramatic miniaturization to nanoscale dimensions. For a calculation of the noninteracting resonant- tunneling transport in an organic-nanostructure transistor, refer to the analysis in Ref. 20. For a nonequilibrium Green function [13,22] calculation of the interacting transport in the nanotube heterostructure transistor (Fig. 1, lower scheme) refer to Ref. 11. Before reporting

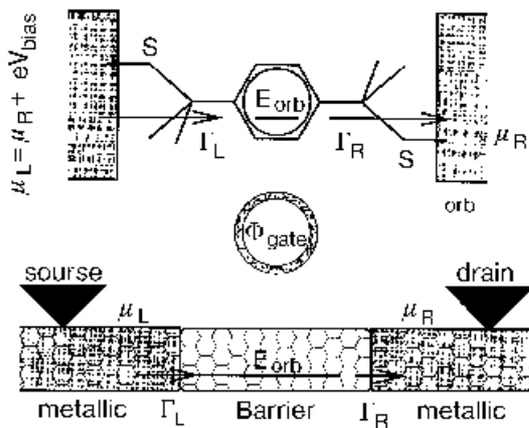


Fig. 1. Schematics of transport and local vibration-source realizations as organic resonant-tunneling nanoparticle (top panel) or as resonant-tunneling nanotube heterostructure (bottom panel). A metal gate (grey ring) at voltage  $\Phi_{\text{gate}}$  controls the current and molecular excitation in either nanostructures by adjusting the resonant-level energy position  $E_{\text{orb}}(\Phi_{\text{gate}})$ .

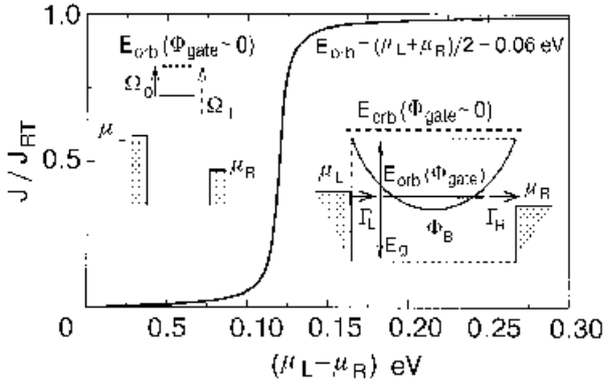


Fig. 2. Gate control of nonequilibrium resonant-tunneling current  $J < J_{RT}$ . The inset panels contrast transport conditions at (i)  $\Phi_{\text{gate}} \sim 0$ , when the molecular gap  $E_g$  forces a resonance-level energy position  $E_{\text{orb}}(\Phi_{\text{gate}} \sim 0)$  (dashed line in right insert) and vibration satellites  $E_{\text{orb}}(\Phi_{\text{gate}} \sim 0) - \Omega_{0,1}$  (left insert) far above the lead chemical potentials  $\mu_{L/R}$ ; And at (ii)  $\Phi_{\text{gate}} \sim 2$  V, when the adjusted electron potential  $\Phi_B$  (solid curve in right insert) positions  $E_{\text{orb}}(\Phi_{\text{gate}}) \approx (\mu_L + \mu_R)/2$  (solid line). The main panel assumes such a fixed  $E_{\text{orb}}(\Phi_{\text{gate}})$ : a moderate bias  $\mu_L - \mu_R \approx 0.3$  eV then saturates the current  $J \approx J_{RT}$ .

calculations of gate control in the current-induced molecular excitation, however, summarize the description of the important tunneling-transport mechanisms [11,20].

Figure 2 illustrates the electrostatic-gate effects on the nonlinear resonant-tunneling current. A finite applied bias  $eV_{\text{bias}}$  maintains the left and the right metallic-nanotube leads at the different chemical potentials  $\mu_L$  and  $\mu_R - \mu_R - eV_{\text{bias}}$ , respectively. The main panel in Fig. 2 assumes that a finite gate voltage  $\Phi_{\text{gate}}$  maintains a fixed resonant-level position  $E_{\text{orb}}(\Phi_{\text{gate}}) = (\mu_L + \mu_R)/2 + 0.06$  eV and it documents how the application of a moderate bias can then saturate a significant resonant-tunneling current

$$eJ < eJ_{RT} = e \left( \frac{4\Gamma_L \Gamma_R}{\Gamma_L + \Gamma_R} \right) \sim \begin{cases} 1 \text{ nA for ORTN,} \\ 5 \text{ } \mu\text{A for NRTTH.} \end{cases} \quad (2)$$

The current is characterized by the resonance width  $\Gamma = \Gamma_L + \Gamma_R$ . At low temperatures this tunneling current [13,22,24]

$$J[E_{\text{orb}}(\Phi_{\text{gate}}); \mu_{L/R}] = J_{RT} [P_{\text{occ}}^{\mu_L}(E_{\text{orb}}) - P_{\text{occ}}^{\mu_R}(E_{\text{orb}})] \quad (3)$$

results as a difference between contributions

$$P_{\text{occ}}^{\mu}(E_{\text{orb}}) = \left[ \frac{1}{2} + \frac{1}{\pi} \arctg \left( \frac{\mu - E_{\text{orb}}}{\Gamma} \right) \right] \quad (4)$$

evaluated at  $\mu = \mu_L$  and  $\mu_R$ , respectively. I focus on the NRTTH transport realizations where it is possible to achieve  $\Gamma \sim 10$  meV [11].

The pair of inset diagrams illustrate the gate operation and contrast the transport conditions in the absence and presence of a finite voltage  $\Phi_{\text{gate}}$  applying to the metal gate. At  $\Phi_{\text{gate}} \sim 0$  (left inset) the molecular gap  $E_g$  ( $\geq 1$  eV for the previously investigated NRTTH transistor [11]) forces a resonance-level energy position  $E_{\text{orb}}(\Phi_{\text{gate}} \sim 0) \approx \mu_L + E_g/2$  (dashed line in right inset). No current results, since both  $E_{\text{orb}}$  and the vibration satellites [13]  $E_{\text{orb}}(\Phi_{\text{gate}} \sim 0) - \Omega_{i=0,1,\dots}$  remain far above the chemical potentials of the leads. However, a voltage  $\Phi_{\text{gate}} \sim 2$  V suppresses the electron potential  $\Phi_B$  (solid curve, right inset), within the fullerene barrier and adjusts the resonance-level position  $E_{\text{orb}}(\Phi_{\text{gate}}) \approx (\mu_L + \mu_R)/2$  (solid line) to enable the tunneling processes (arrows). Below I assume a fixed bias  $eV_{\text{bias}} = 300$  meV and use the resonant-level gate control  $E_{\text{orb}}(\Phi_{\text{gate}})$  also to optimize the current stimulation of molecular excitations.

#### The molecular excitations

For the NRTTH it is relevant to consider current-induced excitation at energies  $\Omega_i = 100\text{--}200$  meV ( $800\text{--}1600$   $\text{cm}^{-1}$  [15]). I concentrate on a pair of high-energy modes, at assumed frequencies  $\Omega_{1(0)} = 200$  meV ( $120$  meV)  $\gg \Gamma$  and describe the current stimulation at zero temperature to illustrate on.. results and predictions.

The top and middle panels in Fig. 3 contrast the independent gate-voltage control of the electron-vibration interaction effects in  $R_{\text{sp}}$  and  $R_{\text{ab}}$  for a pair of vibration energies  $\Omega_0$  (black curve) and  $\Omega_1$  (dashed curve). The gate control is implicit as the gate voltage  $\Phi_{\text{gate}}$  adjusts the resonant level energy position  $E_{\text{orb}}(\Phi_{\text{gate}}) - \mu_L$ . The fixed applied bias satisfies  $2\Omega_1 > eV_{\text{bias}} > 2\Omega_0$  and  $eV_{\text{bias}} > \Omega_1$ . The documented current-excitation gate control arises within the region  $\mu_L > E_{\text{orb}} > \mu_r$  (identified by vertical lines) where the electrostatic gate enables a strong resonant-tunneling current Eq. (3). The excitation transition rates are illustrated for equal electron tunneling rates  $\Gamma_L = \Gamma_R$ .

I determine the magnitude and gate dependence of the current-induced molecular-excitation transition rates  $R_{\text{sp}}$  and  $R_{\text{ab}}$  through a separate nonequilibrium Green function calculation similar to that for the nonequilibrium defluctuations (shot noise) [25]. The calculation involves a determination of the nonequilibrium density-correlation components:

$$\Pi_r^0(\omega) = -i \int_0^{\infty} dt' \exp(i\omega t') \langle [n(t+t')n(t)] \rangle, \quad (5)$$

$$\Pi_{<}^0(\omega) = - \int_{-\infty}^{\infty} dt' \exp(i\omega t') \langle n(t+t')n(t) \rangle, \quad (6)$$

where  $n$  denotes the electron density at the resonance and where the notations and conventions introduced in Ref. 13 are followed. The retarded correlation component includes a commutator [...] and defines the frequency shift [13] and vibration decay [26] due to the electron-vibration (phonon) interaction. Here I evaluate both components (5), (6) out of equilibrium to establish the current-induced excitation level, Eq. (1).

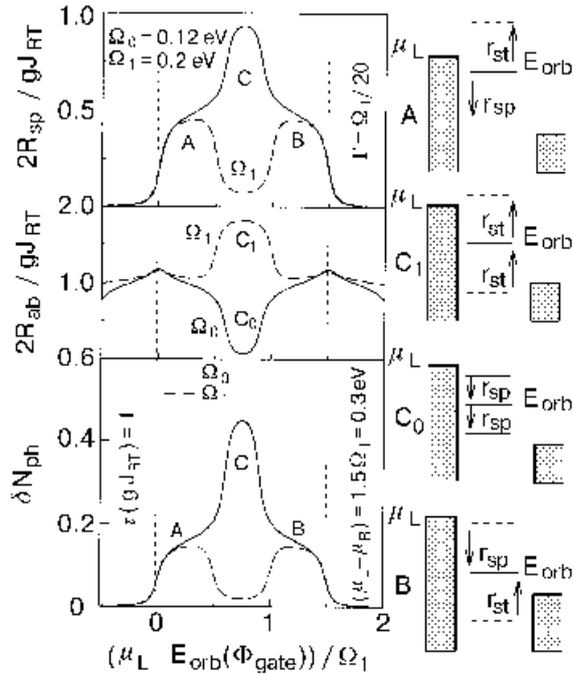


Fig. 3. Frequency selective current-stimulation of molecular vibrations. The left-most top and middle panels contrast the gate variation of the spontaneous phonon emission rate  $R_{sp}$  and net absorption rate  $R_{ab}$  at two frequencies  $\Omega_{0,1}$ . Both rates are proportional to  $J_{RT}$  and the electron-vibration coupling constant  $g$ . The gate variation is implicit and defined through  $E_{orb}(\Phi_{gate})$ . The set of four right-most schematics illustrates the Pauli exclusion mechanism responsible for the frequency-selective control. The set of downward (upward) arrows labeled by  $r_{sp}$  ( $r_{ab}$ ) identify inelastic tunneling events that contribute to the spontaneous emission (to the net stimulated absorption). Finally, the left-most bottom panel contrast the current-induced increase in excitation level  $\delta N_{vib}(\Omega_0)$  (solid curve) and  $\delta N_{vib}(\Omega_0)$  (dashed curve) for a given intrinsic vibration decay time  $\tau$ .

The spontaneous vibration emission rate  $R_{sp}$  is to lowest order in the dimensionless electron-vibration coupling strength  $g$  [13], given by the < correlation component, Eq. (6). The rate is proportional to  $J_{RT}$  but limited to the availability of phase space [27]:

$$\frac{R_{sp}[E_{orb}(\Phi_{gate}); \Omega_i]}{gJ_{RT}} - \frac{[-\Omega^2 \Pi_{<}^0(\Omega_i)/2]}{J_{RT}} = \frac{\Theta(\mu_L - \mu_R - \Omega_i) \Omega_i^2}{4(\Omega_i^2 + 4\Gamma^2)} \times$$

$$\times [\Delta P_{occ}^{prb}(E_{orb}; \Omega_i) + \Delta P_{log}(E_{orb}; \Omega_i)]. \quad (7)$$

A significant variation and structure arises in the vibration emission (7) when  $\Omega_i \gg \Gamma$  condition relevant for the local molecular modes of the NRTH. The spontaneous vibration emission is then dominated by the simple phase-space measure

$$\Delta P_{occ}(E_{orb}; \Omega_i) = [P_{occ}^{\mu_L}(E_{orb}) - P_{occ}^{\mu_R}(E_{orb} - \Omega_i)] + [P_{occ}^{\mu_L}(E_{orb} + \Omega_i) - P_{occ}^{\mu_R}(E_{orb})], \quad (8)$$

while the logarithmic correction term  $\Delta P_{log}$  can be ignored [28]. The phase-space measure (8) is specified by the nonequilibrium contributions (4) at the resonance level and at the vibration satellites  $E_{orb} \pm \Omega$ .

The net vibration absorption is at  $\Omega_i \gg \Gamma$  is given by [29]

$$\frac{R_{ab}[E_{orb}(\Phi_{gate}); \Omega_i]}{gJ_{RT}} - \frac{\Omega^2 [-\text{Im} \Pi_r^0(\Omega_i)]}{J_{RT}} \approx \frac{\Gamma \Omega_i^2}{4(\Omega_i^2 + 4\Gamma^2)} \left( \Gamma_R^{-1} [P_{occ}^{\mu_L}(E_{orb} - \Omega_i) - P_{occ}^{\mu_L}(E_{orb} + \Omega_i)] + \Gamma_L^{-1} [P_{occ}^{\mu_R}(E_{orb} - \Omega_i) - P_{occ}^{\mu_R}(E_{orb} + \Omega_i)] \right). \quad (9)$$

This rate is defined by a phase-space measure, which in contrast to Eq. (8), involves differences of contributions, Eq. (4) evaluated at the same chemical potential (at  $\mu_L$  or  $\mu_R$ ). Thus it is possible to achieve an important independent control of  $R_{sp}[E_{orb}(\Phi_{gate}); \Omega_i]$  and  $R_{ab}[E_{orb}(\Phi_{gate}); \Omega_i]$ .

The Pauli-exclusion principle explains the phase-space limitations on the current-induced spontaneous and net absorption, Eqs. (8) and (9). The four right-most panels A,  $C_0$ ,  $C_1$ , and B in Fig. 3 illustrate the set of different transport conditions

(all with  $J \sim J_{RT}$ ) which characterize the gate-control regimes identified in the left set of panels.

The presence of downward arrows  $r_{sp}$  identify conditions when the current flow can stimulate a spontaneous vibration emission as specified by the phase-space measure, Eq. (8). In region A (B) this emission arises when an electron tunnels into  $E_{orb}$  (into  $E_{orb} + \Omega_{i=0,1}$ ) but leaves at energy  $E_{orb} - \Omega_{i=0,1}$  (at  $E_{orb}$ ). For an applied bias which satisfies  $2\Omega_1 > eV_{bias} > \Omega_1$  (panel  $C_1$ ) neither type-A nor type-B vibration-emission processes are possible for local mode  $\Omega_1$ . However, both types of spontaneous emission processes remain possible for a vibrational mode at  $\Omega_0 < eV_{bias}/2$  (panel  $C_0$ ).

The presence of upward arrows  $r_{ab}$  instead identifies conditions for a net current-induced absorption  $R_{ab} \neq 0$ . In section A (B) a net absorption arises, when the electron enters at  $E_{orb}$  (at  $E_{orb} - \Omega_{i=0,1}$ ) but leaves at  $E_{orb} + \Omega_{0,1}$  (at  $E_{orb}$ ). Tuning  $E_{orb}$  to the central region C causes an enhanced absorption for mode  $\Omega_1$ , as both type-A and type-B absorption processes become possible (panel  $C_1$ ). However, for the lower mode at  $\Omega_0 < eV_{bias}/2$  I find an effective cancellation (panel  $C_0$ ) as the energies  $E_{orb}$  and  $E_{orb} \pm \Omega_0$  all carry a partial electron occupation and thus produces a vanishing net absorption rate,  $R_{ab} \rightarrow 0$ .

#### Frequency-selective molecular-vibration stimulation

The lower-left panel in Fig. 3 documents how an optimization of current-induced molecular excitation is possible. The panel contrasts the calculated gate-variation of the increase in the molecular excitation level, Eq. (1),  $\delta N_{vib}(\Omega_0)$ , and  $\delta N_{vib}(\Omega_1)$ , and details methods to enhance the current stimulation of mode  $\Omega_0$  at the expense of mode  $\Omega_1 > \Omega_0$ . Such selective excitation is possible even when  $eV_{bias} > \Omega_1 \gg \Gamma$ , and arises when  $2\Omega_1 > eV_{bias} > 2\Omega_0$  and  $E_{orb}$  is tuned to region C ( $E_{orb} \approx (\mu_L + \mu_R)/2$ ). These nonequilibrium transport conditions simultaneously minimize  $R_{st}(\Omega_1)$  towards zero and maximize the ratio  $R_{st}(\Omega_0)/R_{ab}(\Omega_0)$  to extinguish  $\delta N_{vib}(\Omega_1)$  and dramatically enhance the lower-frequency current stimulation  $\delta N_{vib}(\Omega_0)$ .

#### The molecular-dynamics source and probe.

The lower-left panel in Fig. 3 also details the suggested operation as a molecular-excitation source. The panel documents a crisp electrostatic gate control for the current-stimulation  $\delta N_{vib}(\Omega_1)$  which arises through an adjustment of the resonant-level energy position  $E_{orb}(\Phi_{gate})$  [11]. This implicit gate control permits a switch between enabling and

disabling the current-stimulation (1). Such operation can produce a frequency-selective molecular-vibration source and even a strong nonequilibrium burst  $\delta N_{vib}(\Omega_0) \gg 0$  of high-energy nanostructure vibrations.

Nanoscale molecular-dynamics probing is then possible with simultaneous Raman measurements of the anti-Stokes signal at  $\Omega_0$ , because the anti-Stokes strength is directly sensitive [17] to the excess molecular-excitation burst  $\delta N_{vib}(\Omega_0) \gg 0$ . Such Raman measurement can through Eq. (1) determine the decay  $1/\tau_i$  that characterize these nanostructure molecular excitations and thus probe mechanisms [16,18,19] which help determine the intrinsic nanoscale molecular dynamics. The suggested *in situ* molecular-dynamics probe could realize an important strong testing of our theoretical descriptions for both the nanostructure atomic configurations [10,11,30] and for the current-induced structural dynamics [4,12–14].

In summary, I have suggested a novel *in situ* probe of the nanoscale molecular dynamics of organic-molecule and fullerene-tube heterostructures. General nonequilibrium Green function results for the current coupling to local nanostructure excitations were presented to document a frequency-selective electrostatic control and optimal operation as a necessary current-excitation source. Raman measurements of the anti-Stokes signal can then permit an in-situ probe of the local nanostructure molecular dynamics at nonequilibrium conditions.

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23. Our results apply to linear order in the current and describe the stimulation of local high-energy nanostructure vibrations. Cases where the tunneling rate coincides with standing-wave resonances requires a treatment of also nonlinear excitation effects; See, e.g., Ref. 12.
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27. The rate  $R_{sp}$  is just the excitation increase  $\delta N_{ph} - \delta \langle b^\dagger b \rangle$  as one enables the current-vibration coupling  $g > 0$ . I introduce  $A_0(\omega) \equiv (2\tau_i)[1 + \tau^2(\omega - \Omega_i)^2]$  as the unperturbed vibration-spectral function and find to lowest order in  $g$  the current stimulation  $2\delta N_{ph} \approx g\Omega_i^2 \int \Pi_{<}^0(\omega)A_0(\omega)d\omega$ . The result Eq. (7) applies for  $\tau_i^{-1} < \Gamma$ .
28. The logarithmic correction to the phase-space measure is given by  $\Delta P_{\log}(E_{orb}; \Omega_i) - (\Gamma/\pi\Omega_i) \sum_{\alpha=1,2} \ln(\Delta_\alpha^2 + \Gamma^2) - (\Gamma/\pi\Omega_i) \times \sum_{x=L,R} \ln[(\mu_x - E_{orb})^2 + \Gamma^2]$ , where  $\Delta_{1(2)} = (\mu_{L(R)} - E_{orb} - (+)\Omega_i)$ ; The prefactor  $(\Gamma/\pi\Omega_i)$  ensures a smooth behavior at general  $\Gamma/\Omega_i$  and a vanishing one  $\Delta P_{\log} \rightarrow 0$  at  $\Gamma \ll \Omega_i$ .
29. A small logarithmic correction to  $R_{ab}$  is given through the full nonequilibrium determination of  $\Pi_r^0(\omega)$  [13]. I stress that  $-\text{Im} \Pi_r^0 > 0$  and that singlt-resonance tunneling never produces coherent emission.
30. See, e.g., J. M. Seminario, A. G. Zacarias, and J. M. Tour, *J. Am. Chem. Soc.* **120**, 3970 (1998); A. Rubio, D. Sanchez-Portal, E. Artacho, P. Ordejion, and J. M. Soler, *Phys. Rev. Lett.* **82**, 3520 (1999).