Efficient calculation of inelastic vibration signals in electron transport: Beyond the wide-band approximation

Lu, Jing Tao; Christensen, Rasmus Bjerregaard; Foti, Giuseppe; Frederiksen, Thomas; Gunst, Tue; Brandbyge, Mads

Published in:
PHYSICAL REVIEW B

Link to article, DOI:
10.1103/PhysRevB.89.081405

Publication date:
2014

Document Version
Publisher's PDF, also known as Version of record

Link back to DTU Orbit

Citation (APA):
Efficient calculation of inelastic vibration signals in electron transport: Beyond the wide-band approximation

Jing-Tao Lü,1,2 Rasmus B. Christensen,2 Giuseppe Foti,1,4 Thomas Frederiksen,4,5 Tue Gunst,2 and Mads Brandbyge2,*

1School of Physics, Huazhong University of Science and Technology, Wuhan, China
2Department of Micro- and Nanotechnology, Technical University of Denmark, Ørsteds Plads, Bldg. 345E, DK-2800 Kongens Lyngby, Denmark
3Centro de Física de Materiales, Centro Mixto CSIC-UPV, Donostia-San Sebastián, Spain
4Donostia International Physics Center (DIPC) – UPV/EHU, Donostia-San Sebastián, Spain
5IKERBASQUE, Basque Foundation for Science, Bilbao, Spain

(Received 24 December 2013; revised manuscript received 24 January 2014; published 21 February 2014)

We extend the simple and efficient lowest order expansion (LOE) for inelastic electron tunneling spectroscopy (IETS) to include variations in the electronic structure on the scale of the vibration energies. This enables first-principles calculations of IETS line shapes for molecular junctions close to resonances and band edges. We demonstrate how this is relevant for the interpretation of experimental IETS using both a simple model and first-principles simulations.

DOI: 10.1103/PhysRevB.89.081405 PACS number(s): 73.63.-b, 68.37Ef, 61.48.—c

The inelastic scattering of electronic current on atomic vibrations is a powerful tool for investigations of conductive atomic-scale junctions. Inelastic electron tunneling spectroscopy (IETS) has been used to probe molecules on surfaces with scanning tunneling microscopy (STM) [1], and for junctions more symmetrically bonded between the electrodes [2–7]. Typical IETS signals show up as dips or peaks in the second derivative of the current-voltage (I-V) curve [8]. In many cases the bonding geometry is unknown in the experiments. Therefore, first-principles transport calculations at the level of density functional theory (DFT) in combination with nonequilibrium Green’s functions (NEGF) [9–14] can provide valuable insights into the atomistic structure and IETS. For systems where the electron-vibration (e-vib) coupling is sufficiently weak and the density of states (DOS) varies slowly with energy (comparable to typical vibration energies) one can greatly simplify calculations with the lowest order expansion (LOE) in terms of the e-vib coupling together with the wide-band approximation (LOE-WBA) [10,15]. The LOE-WBA yields simple expressions for the inelastic signal in terms of quantities readily available in DFT-NEGF calculations. Importantly, the LOE-WBA can be applied to systems of considerable size.

However, the use of the WBA cannot account for IETS signals close to electronic resonances or band edges, which often contains crucial information [16,17]. For example, a change in IETS signal from peak to peak-dip shape was recently reported by Song et al. [6] for single-molecule benzene-di-thiol (BDT) junctions, where an external gate enabled tuning of the transport from off-resonance to close-to-resonance. Also, high-frequency vibrations involving hydrogen appear problematic since the LOE-WBA is reported to underestimate the LOE intensity [18].

Here we show how the energy dependence can be included in the LOE description without changing significantly the transparency of the formulas or the computational cost. We describe how the generalized LOE differs from the original LOE-WBA, and demonstrate that it captures the IETS line shape close to a resonance. We apply it to DFT-NEGF calculations on the resonant BDT system and to off-resonant alkane-di-thiol junctions, and show how the improved LOE is necessary to explain the experimental data.

Method. We adopt the usual two-probe setup with quantities defined in a local basis set in the central region (C) coupled to left/right electrodes (α = L, R). We consider only interactions with vibrations (indexed by λ, with energies ℏωλ, and e-vib coupling matrices Mλ) inside C. To lowest order in the e-vib self-energies Σλ (second order in Mλ) the current can be expressed as a sum of two terms I(V) = Iε + I1, using unperturbed Green’s functions G0 = G–1 defined in region C [10,15],

\[ I_c = \frac{G_0}{e} \int_{-\infty}^{\infty} d\epsilon \{ f_L(\epsilon) - f_R(\epsilon) \} \text{Tr}[G^0 T^L G^0 T_R^R](\epsilon) + 2Re \text{Tr}[G^0 \Sigma^L G^0 \Sigma^R](\epsilon). \]

where G0 = 2e2/ℏ is the conductance quantum and summation over the vibration index λ is assumed. The e-vib self-energies Σλ are expressed as

\[ \Sigma^{\pm}(\epsilon) = M^{\pm}(\alpha)(N^{\pm}(\epsilon) + 1)G^{\pm}(\epsilon) + N^{\pm}(\epsilon)G^{\pm}(\epsilon)M^{\pm}(\alpha), \]

\[ \Sigma^{\pm}(\epsilon) = \pm \frac{1}{2} (\lambda(\epsilon) - \Sigma^{\pm}(\epsilon)) - \frac{i}{2} \lambda(\Sigma^{\pm}(\epsilon) - \Sigma^{\pm}(\epsilon)), \]

with ε± = ε ± ℏωλ, bosonic occupations Nλ, and \( \lambda \) denoting the Hilbert transform. Finally, the lesser/greater Green’s functions G± describing the occupied/unoccupied states

\[ G^{\pm}(\epsilon) = \mp \{ f_L(\pm \epsilon)A_L(\epsilon) + f_R(\mp \epsilon)A_R(\epsilon) \} \]

are given by the spectral density matrices Aα = G0 Tα G0 for left/right moving states with fillings according to the reservoir Fermi functions \( f_\alpha(\epsilon) = n_f(\epsilon - \mu_\alpha). \)
The above equations are numerically demanding because of the energy integration over voltage-dependent traces. In the following we describe how further simplifications are possible without resorting to the WBA. Here we are interested in the “vibration signal” that is the change in the current close to the excitation threshold \(|eV| \approx \hbar \omega_o\), with \(eV = \mu_L - \mu_R\). As IETS signals are obtained at low temperatures, we assume that this is the smallest energy scale \(k_BT \ll \hbar \omega_o, \Gamma\), where \(\Gamma\) is the typical electronic resonance broadening. The inelastic term \(I_i\) [Eq. (2)] then reduces to

\[
I_i \approx \frac{G_0}{2e} \sum_{\sigma=\pm} \left( \frac{\coth \frac{\hbar \omega_o}{2k_BT}}{2k_BT} - \frac{\coth \frac{\hbar \omega_o + \sigma eV}{2k_BT}}{2k_BT} \right) \int_{-\infty}^{\infty} d\epsilon \text{Tr} \times [M_{\sigma} \tilde{A}_{\lambda}(\epsilon) M_{\sigma} A_R(\epsilon)](f_L(\epsilon) - f_R(\epsilon)),
\]

where \(\tilde{A}_{\lambda} = G^T \Gamma_{\lambda} G^T\) is the time-reversed version of \(A_{\lambda}\). In the second derivative of \(I_i\) with respect to voltage \(V\), the coth parts give rise to a sharply peaked signal around \(|eV| = \hbar \omega_o\), with width of the order of \(k_BT\). If the electronic structure \(A_{\lambda}\) varies slowly on the \(k_BT\) scale, it can be replaced by a constant using \(\epsilon \approx \mu_L\) and \(\epsilon_R \approx \mu_R = \mu_L + \sigma \hbar \omega_o\). Thus, around the vibration threshold we get

\[
\delta_i^3 I_i \approx \gamma_{i,\lambda} \delta_i^3 T_{\text{sym}},
\]

where, as in the LOE-WBA, define the “universal” function

\[
T_{\text{sym}} = \frac{G_0}{2e} \sum_{\sigma=\pm} \sigma(\hbar \omega_o + \sigma eV) \times \left( \frac{\coth \frac{\hbar \omega_o}{2k_BT}}{2k_BT} - \frac{\coth \frac{\hbar \omega_o + \sigma eV}{2k_BT}}{2k_BT} \right).
\]

The elastic term \(I_e\) [Eq. (1)] can be divided into two parts, \(I_e = I_{e}^h + I_{e}^n\), where the first (latter) represents all terms without (with) the Hilbert transformation originating in Eq. (4). The “non-Hilbert” part \(I_{e}^n\) yields a coth factor and integral of similar in form to the one for \(I_i\). Both \(I_i\) and \(I_{e}^n\) thus yield an inelastic signal with a line shape given by the function \(\delta_i^3 T_{\text{sym}}\) and the sign/intensity governed by \(\gamma_e = \gamma_{i,\lambda} + \gamma_{n,\lambda}\), with \(\gamma_{n,\lambda} \approx \nu \Im B_{\lambda}\), and

\[
B_{\lambda} \equiv \text{Tr}[M_{\lambda} A_R(\mu_L) G(\mu_L) A_R(\mu_R) - M_{\lambda} G(\mu_L) \Gamma(\mu_L) A_R(\mu_R) M_{\lambda} A_R(\mu_L)].
\]

The “Hilbert” part \(I_{e}^h\) requires a bit more consideration. Besides terms which do not result in threshold signals [19], we have terms involving \(\mathcal{H}[A_{\lambda} f_{\sigma}]\). Again, if \(A_{\lambda}\) varies slowly around the step in \(f_{\sigma}\) we may approximate

\[
\mathcal{H}[A_{\lambda}(\epsilon) f_{\sigma}(\epsilon)](\epsilon) \approx A_{\lambda}(\epsilon) \mathcal{H}[f_{\sigma}(\epsilon)](\epsilon).
\]

The Hilbert transformation of the Fermi function is strongly peaked at the chemical potential, and again we evaluate the energy integral by evaluating all electronic structure functions \((A_{\lambda}, G^T, \Gamma_{\lambda})\) at the peak values, keeping only the energy dependence of the functions related to \(f_{\sigma}\) inside the integral. The result is

\[
\delta_i^3 I_{e}^h \approx \kappa_i \delta_i^3 T_{\text{sym}}.
\]

with \(\kappa_i = 2ReB_{i}\), and, again as in the LOE-WBA, the universal function

\[
T_{\text{sym}} = \frac{G_0}{2e} \int_{-\infty}^{+\infty} d\epsilon \mathcal{H}[f(\epsilon_-) - f(\epsilon_+)](\epsilon) [f(\epsilon - eV) - f(\epsilon)]
= -\frac{G_0}{2e\pi} \sum_{\sigma=\pm} \sigma(eV + \sigma \hbar \omega_o) \ln \left| \frac{eV + \sigma \hbar \omega_o}{\hbar \omega_o} \right|.
\]

Here the latter is for \(k_BT = 0\), while it can be expressed using the digamma function for finite \(k_BT\) [20]. In total we have written the IETS as a sum of individual vibration signals [10],

\[
\delta_i^3 I_i = \gamma_i \delta_i^3 T_{\text{sym}}(V, \hbar \omega_o, \Gamma, N_d)
+ \kappa_i \delta_i^3 T_{\text{sym}}(V, \hbar \omega_o, T).
\]

Equation (14) is our main formal result. As for the LOE-WBA we have expressed the vibration signals from the universal functions, and structure factors containing quantities readily obtained from DFT-NEGF. However, importantly, here we have generalized these to include the effect of \(\hbar \omega_o\), and thus the change in electronic structure over the excitation energy. Our LOE expressions for \(\gamma_{i,\lambda}\) and \(\kappa_i\) above simply reduce to the LOE-WBA when \(\mu_L = \mu_R = \mu_0\). We will now demonstrate some situations where the LOE expression Eq. (14) is crucial for detailed interpretation of experimental IETS line shapes.

**Simple model.** First we use a single-level model to illustrate how the “asymmetric” term contains important information about the energy dependence of the electrode couplings. In the LOE-WBA one always has \(\kappa_i = 0\) for symmetric junctions. This is not the case for the LOE expression Eq. (10). We therefore consider a symmetric junction containing a single electronic level at \(\varepsilon_0\) (with \(\mu_0 = 0\)), coupled to a local vibration \((\omega_0)\), and with energy-dependent electrode coupling rates. Assuming symmetrical potential drop, and using the notations \(\Gamma_i = \Gamma_L(\mu_L) + \Gamma_R(\mu_L)\) and \(\Gamma_r = \Gamma_L(\mu_R) + \Gamma_R(\mu_R)\) we can write the “symmetric,”

\[
\gamma = -C \left[ \Gamma_1^2 \Gamma_2^2 - (4\varepsilon_0^2 - \hbar^2 \omega_0^2)^2 \right],
\]

and “asymmetric” coefficients,

\[
\kappa = 4C (\delta \Gamma_1 \varepsilon_0 + \varepsilon_0 \Gamma_0) \left[ \Gamma_1 \Gamma_r - (4\varepsilon_0^2 - \hbar^2 \omega_0^2) \right],
\]

where \(\delta \Gamma = \Gamma_1 - \Gamma_r\), \(\Gamma = (\Gamma_1 + \Gamma_r)/2\), and \(C\) is a constant common to \(\gamma\) and \(\kappa\). In the typical case of transition metal electrodes the coupling can contain contributions both from a wide \(s\) band as well as from a narrow \(d\) band leading to a significant \(\delta \Gamma\) and finite \(\kappa\). To model the \(s\) band we use a constant \(\Gamma_0\), and to mimic the coupling (hopping \(t\)) to a \(d\) band we add the self-energy of a semi-infinite 1D chain, with bandwidth \(2W\) centered at \(\varepsilon_0 = 0\). Figures 1(a)–1(c) compare the signals corresponding to the case for the LOE expression Eq. (14). We will now demonstrate some situations where the LOE expression Eq. (14) is crucial for detailed interpretation of experimental IETS line shapes.
In the break-junction experiments the atomic structure of the junction is unknown. We anticipate that the gap between the electrodes is quite open and involves sharp asperities with low-coordinated gold atoms in order to allow for the external gating to be effective. In order to emulate this we consider BDT bonded between atoms on Au(111) surfaces [Fig. 2(a)], and employ only the \( \Gamma \) point in the transport calculations yielding sharper features in the electronic structure. We correct the HOMO-LUMO gap [22] and model the electrostatic gating simply by a rigid shift of the molecular orbital energies relative to the gold energies. In Figs. 2(b) and 2(c) we compare IETS calculated with LOE and LOE-WBA as a function of gating. As in the experiment, we observe three clear signals around \( \hbar \omega_0 = 95,130,200 \text{ meV} \) due to benzene vibrational modes. Off-resonance the LOE and LOE-WBA are in agreement as expected. But when the gate voltage is tuned to around \( V_g \approx -1 \text{ V} \) the methods deviate because of the appearance of sharp resonances in the transmission around the Fermi energy [Fig. 2(a)]. These resonances involve the \( d \) orbitals on the contacting gold atoms, as seen in the eigenchannel [23] plot in Fig. 2(a), and result in a peak-dip structure as seen in the experiment and anticipated by the simple model. Thus it is important to go beyond LOE-WBA in order to reproduce the peak to peak-dip transition taken as evidence for close-to-resonance transport.

**IETS of alkane-dithiol.** As another demonstration of the improvement of LOE over LOE-WBA, we consider molecular junctions formed by straight or tilted butane-dithiol (C4DT) molecules linked via low-coordinated Au adatoms to Au(111) electrodes, see inset to Fig. 3. Based on DFT-NEGF [21] we calculate elastic transmission and IETS for the periodic structure averaged over electron momentum \( k_x \) [24]. As shown in Figs. 3(a) and 3(b), transport around the Fermi level is off-resonance but dominated by the tail of a sulfur-derived peak centered at approximately 0.25 eV below the Fermi level. This feature introduces a relatively strong energy dependence into the electronic structure which makes the WBA questionable. Indeed, as shown in Fig. 3(c), LOE-WBA gives a smaller IETS intensity compared to the LOE for the energetic CH\(_2\) stretch modes (\( \hbar \omega_0 \sim 375 \text{ meV} \)). The WBA may thus be the reason why LOE-WBA calculations were reported to underestimate the IETS intensity for these energetic modes in comparison with experiments [18]. We note that the intensity enhancement is found to be more pronounced for the straight configuration, which we speculate may be related to the change in the sulfur density of states as a function of tilt (for the tilted geometry
The slope of the PDOS close to the Fermi level decreases. The intensity change reported in Fig. 3 thus suggests the relevance of going beyond LOE-WBA for simulations involving high-energy vibrational modes.

**Conclusions.** A generalized LOE scheme for IETS simulations with the DFT-NEGF method has been described. Without introducing the WBA, our formulation retains both the transparency and computational efficiency of the LOE-WBA. This improvement is important to capture correctly the IETS line shape in situations where the electronic structure varies appreciably on the scale of the vibration energies, such as near sharp resonances or band edges. Together with DFT-NEGF calculations we have discovered that the intricate experimental line shape of a gated BDT can be explained without the need to assume asymmetric bonding of the molecule to the electrodes. Also, simulations for C4DT junctions suggest that going beyond WBA is important to capture the IETS intensity related to energetic CH2 stretch modes.

**Acknowledgements.** We acknowledge computer resources from the DCSC, and support from Center for Nanostructured Graphene (Project DNRF58). J.T.L. acknowledges support from the National Natural Science Foundation of China (Grants No. 11304107 and No. 61371015), and the Fundamental Research Funds for the Central Universities (HUST:2013TS032). G.F. and T.F. acknowledge support from the Basque Departamento de Educación and the UPV/EHU (Grant No. IT-756-13), the Spanish Ministerio de Economía y Competitividad (Grant No. FIS2010-19609-CO2-00), and the European Union Integrated Project PAMS.

---


EFFICIENT CALCULATION OF INELASTIC VIBRATION . . .

PHYSICAL REVIEW B 89, 081405(R) (2014)