The conductance of molecular wires and DFT based transport calculations

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The experimental value for the zero bias conductance of organic molecules coupled by thiol-groups to gold electrodes tends to be much smaller than the theoretical result based on density functional theory (DFT) calculations, often by orders of magnitude. To address this puzzle we have analyzed the regime within which the approximations made in these calculations are valid. Our results suggest that a standard step in DFT based transport calculations, namely approximating the exchange-correlation potential in quasistatic nonequilibrium by its standard equilibrium expression, is not justified at weak coupling. We propose, that the breakdown of this approximation is the most important source for overestimating the width of the experimentally observed conductance peak and therefore also of the zero bias conductance. We present a numerical study on the conductance of the organic molecule that has recently been studied experimentally by Reichert et. al.\textsuperscript{1} that fully agrees with this conclusion.

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I. INTRODUCTION

Recently, several conductance measurements of single organic molecules have been reported\textsuperscript{1-5}. Fig. 1 illustrates a typical setup with one of the molecules used in experiments\textsuperscript{1}. In part the tremendous attention that this field “molecular electronics” has received in recent years is due to its possible technological impact. Indeed, a molecular storage device has already been built\textsuperscript{6} and also molecular switches have been realized\textsuperscript{7}. It is clear that for technological applications being able to systematically model and understand the charge transport properties of the combined system of molecule and contacts is of crucial importance.

Since the conductance is very sensitive to details of spectral and orbital properties of molecules and their wavefunctions, ab initio methods like density functional theory (DFT) are an indispensable tool not only for structure calculations but also for transport theory. Nowadays, DFT calculations including several thousand electrons are possible, which allows the treatment of large molecules and also to include parts of the contacts for more realistic calculations. Despite these enormous capacities, a quantitative description of transport for weakly coupled molecules with a conductance well below the conductance quantum, $g \ll 1$, has not been achieved. In fact, experimental and theoretical values for the zero bias conductance of organic molecules, e. g. benzene, often differ by 1-3 orders of magnitude\textsuperscript{8,9}.

In the present work we consider the organic molecule depicted in Fig. 1, which has been subject to experimental\textsuperscript{1} and DFT based theoretical investigations\textsuperscript{10}, before. In this case the experimental value for the zero bias conductance is smaller than the theoretical one by a factor $\sim 20$. Three different reasons that possibly could lead to this discrepancy should be mentioned and discussed\textsuperscript{11}.

First, Fig. 1 shows an idealized situation and actual microscopic conditions realized in the experiment are not known. However, we have chosen the particular case, Fig. 1, because the experimental results are well reproducible\textsuperscript{20}. If important variations in the atomic structure of the contact would exist, one should expect strong fluctuations in the conductivity from sample to sample. Indeed fluctuations are present, see Fig. 4 below, but their magnitude is relatively small so that strong structural variations are unlikely to occur. Small variations in structure, on the other hand, tend to have little impact on the theoretical conductance, only. Our calculations show, that the transmission is not very sensitive to changes in the bond angles or bond lengths as long as the change is within reasonable limits. Therefore, deviations from the assumed atomic arrangement depicted in Fig. 1 from its experimental realization do not seem to offer a plausible explanation of the large discrepancies observed.

Second, in theoretical calculations approximations have to be done when the molecule is coupled to the leads. Artefacts can be excluded only, when parts of the leads are included into the calculation (“extended molecule”), so that the following hierarchy of inequalities is met:

$$\delta_e M < \gamma_e M \ll \gamma_M$$

(1)
(δ_M: level spacing of extended molecule, γ_M (γ_eM) level broadening of bare (extended) molecule when coupled to the leads). Indeed, these conditions have not been met in previous calculations10, where δ_M ≈ γ_M. In order to improve upon this result, we have performed conductance calculations for the extended molecule, Fig. 1, where δ_M/γ_M ≈ 0.1 and δ_eM/δ_M ≈ 0.01 so that the hierarchy of inequalities is satisfied and artefacts from modeling of the leads can be excluded. We find that qualitative agreement of our data with the experiment is improved significantly as compared to the earlier calculation10. However, the quantitative disagreement is not decreased but rather increased by another factor of 10! This shows that insufficient modeling can be ruled out as a source for the discrepancy.

After we have discarded two obvious possibilities for explaining the observed discrepancy, we have to resort to more fundamental considerations. In this paper, we advocate a third possibility, namely that the standard implementation of DFT based transport calculations relies crucially on assumptions that are not justified in the limit of “weak coupling” where the spatial structure of molecular wavefunctions is strongly inhomogenous and transport is dominated by single resonances.12

We give a outline of our paper and a brief account of key results. In section II we discuss the derivation of the basic equation of DFT-transport, Eq. 11, and show that it has a wide range of applicability. This is true provided that the DFT-calculations are performed with an exchange-correlation potential appropriate for the nonequilibrium situation. However, since this potential is not known in general, in practice an “equilibrium exchange correlation approximation” (EXCA) is made in which the standard equilibrium potential is used. Apriori, this approximation is uncontrolled. In section II B we introduce an alternative – but equivalent – formulation of transport, the Kubo formalism. It enables us to give analytic arguments for the applicability of the EXCA. Our results suggest, that the EXCA can be trusted only in the case of non-resonant transport, when the molecular level broadening is strong and individual molecular levels strongly overlap. A typical example for this case is a chain of gold atoms. By contrast, if the transmission exhibits resonances, transport is determined by individual molecular orbitals and their broadening. In this case, the corrections to the EXCA become significant and in particular the level broadening and therefore the zero bias conductance may be severely overestimated. The molecule in Fig. 1 is a representative for these systems.

In section III, we present numerical transport calculations for a gold chain and the molecule, depicted in Fig. 1, that we have already alluded to above. In view of our theoretical analysis we propose that it is the breakdown of the EXCA in the weakly coupled limit that is the cause of the large discrepancy between theoretical and experimental molecular conductance. A discussion of our findings will be given in section IV.

II. TRANSPORT FORMALISMS

A. Landauer-Buttiker-Formalism for Interacting Electrons

Meir and Wingreen have derived a general expression for the current flowing through a region of space, where the charge carriers can interact – like a quantum dot or a molecule13. Specifically, the Hamiltonian has the structure

\[ H = H_0(\{d_m\};\{d_m\}) + \sum_{\alpha=l,r} \epsilon_{\alpha} c_{\alpha}^\dagger c_{\alpha} + \sum_{m a=l r} (t_{\alpha m} c_{\alpha}^\dagger d_m + \text{H.c.}) \]  \hspace{1cm} (2)

The first term describes the bare molecule which in general may include the electron-electron interaction. Its detailed structure will be of no importance in what follows. The molecule makes contact to two leads denoted left and right. The bare leads are assumed to be noninteracting and described by the second term in Eq. (2). The third term represents the contact. The Meir-Wingreen formula connects the retarded (advanced) Green’s functions \( G (G^\dagger) \) and the lesser function \( G^< \) of the full many-body problem (including the leads) with the \( dc \)-current

\[ I = -\int dE \text{ tr}(f_L \Gamma_L - f_R \Gamma_R)(G - G^\dagger) + \text{tr}(\Gamma_L - \Gamma_R) G^< \]  \hspace{1cm} (3)

where \( f_{L,R} = f(E - \mu_{R,L}) \) denotes the Fermi distribution functions for the leads at chemical potentials \( \mu_{R,L} \) and \( \Gamma_{L,R} \) is the imaginary part of the self energy that describes the coupling of the molecule to the external leads. (For the most part the dependence on energy \( E \) will be suppressed in our notation.) For noninteracting particles, in terms of these self energies we have \( G^{-1} = E_1 - H_0 - \Sigma_L - \Sigma_R \), so that \( \Gamma_{L,R} = i(\Sigma_L - R - \Sigma_L^I \Gamma_{L,R}) \) and

\[ G^< = iG ( f_L \Gamma_L + f_R \Gamma_R ) G^\dagger . \]  \hspace{1cm} (4)

In the appendix, we demonstrate that Eq. (4) is just the statement, that the density matrix can be constructed from left and right going scattering states \( \psi_{l,r} (x = (x, t)) \),

\[ G^<(x,x') = \sum_{l,r} f_l \psi_l(x) \psi_l^*(x') + f_r \psi_r(x) \psi_r^*(x'). \]  \hspace{1cm} (5)

From (3) together with (4) we find for the transmission in the limit of linear response and zero temperature

\[ T(E) = \text{tr} \Gamma_L G T_R G^\dagger . \]  \hspace{1cm} (6)

Eq. (6) is an incarnation of the familiar Landauer-Buttiker formula14 and valid for noninteracting electrons.
In this section, we argue that expression (6) continues to hold also for a much larger class of interacting problems. This is because according to the Runge-Gross theorem of time dependent DFT (TDDFT) the time evolution of any many-particle Hamiltonian can be calculated by solving a single particle problem in an appropriate effective potential\(^\dagger\). Since this point is of importance to us we elaborate on it in the following.

The effective time dependent single particle problem we should solve is of the form

\[ -i \frac{d}{dt} \theta_n(x) = \left( -\frac{1}{2m} \nabla^2 + V_n(x) \right) \theta_n(x) \quad (7) \]

where

\[ V_n(t) = V_i + V_H[n] + V_{xc}[n] + V_{ex}(t) \quad (8) \]

with \( V_i \) denoting the ion-core potential, \( V_H \) the Hartree-interaction, \( V_{xc} \) the exchange-correlation potential and \( V_{ex} \) the external probing field. (Notation suppresses the spatial index \( x \).

In addition to the explicit time dependence of the potential \( V_n \) imposed by \( V_{ex} \) an implicit dependence exists because \( V_n \) is a functional (in general nonlinear and nonlocal in time and space) of the density \( n(x,t) \). Let us assume that Eq. (7) describes the molecule together with the leads and an external perturbation \( V_{ex} \) that is switched on at \( t = t_0 \) and time independent thereafter:

\[ V_{ex}(x) = V_{ex}(x) \Theta(t - t_0) \]

At times prior to \( t_0 \) the system is assumed to be in thermal equilibrium (\( T = 0 \)), so that the reservoirs are characterized by occupation numbers \( f_n \). Then we have \( n(x) = G^\leq_{\uparrow}(x,x) \), with

\[ G^\leq_{\uparrow}(x,x') = \sum_n f_n \theta_n(x) \theta_n^*(x') \quad (9) \]

where \( \theta_n(x,t) \) evolves according to Eq. (7). Moreover, we specialize to the case, where \( V_{ex}(x) \) generates a monotonous electrical potential drop from \( V_{ex} = V \) in the asymptotic region of the left lead to \( V_{ex} = 0 \) in the right lead. In response to the potential drop a current is being generated. After an initial period exhibiting transient behavior, there will be a parametrically wide time interval in which the current and the density are quasistationary. This is precisely the situation for which also Eq. (3) has been derived\(^\dagger\). It is only at even much longer times, that the electrochemical potential becomes homogenous again and the current stops to flow. Formally speaking, we perform the order of limits in which the size of the reservoirs is send to infinity first and \( t_0 \to -\infty \), thereafter. We mention that, since we are interested in the long-time-behavior only, details of how the external potential is switched on, step-like or adiabatic, are unimportant. The corresponding memory is erased inside the reservoirs.

Since Eq. (9) describes the exact evolution of the time dependent density we can also calculate the (longitudinal) current density \( j = dn/dt \) and hence find an exact expression for the conductance. Indeed, the same reasoning that is used for noninteracting electrons to relate transport to a scattering problem can be employed for the present effective single particle problem as well. Consequently, a zero temperature description of the quasistationary region in terms of scattering states should exist with \( G^\leq_{\uparrow} \) taking a form similar to Eqs. (4) and (5)

\[ G^\leq_{\uparrow}(x,x') = \sum_{l,r} f_l \theta_l(x) \theta_r^*(x') + f_r \theta_r(x) \theta_r^*(x') \quad (10) \]

It involves the Kohn-Sham orbitals \( \theta_{r,l} \) representing the scattering states of the quasistationary nonequilibrium situation and their zero temperature occupation numbers \( f_{l,r} \) imposed by the left and right reservoirs. Since also the derivation of Eq. (4) can be repeated for an effective single particle problem, a relation analogous to (3) also holds for \( G^\leq_{\uparrow} \) and the corresponding retarded Green’s function \( G \). Together with the previous statement (10) this implies that the transmission is given by

\[ T = \text{tr} \, \Gamma_L \, G_{\uparrow} \, \Gamma_R \, G_{\downarrow} \quad (11) \]

with \( G^{-1}_{\uparrow}(E) = G_{0\uparrow}^{-1} - \Sigma_L - \Sigma_R \), and the resolvent matrix

\[ G_{0\uparrow}(x,x',E) = \sum_n \frac{\phi_n(x) \phi_n^*(x')}{E - \epsilon_n + i\delta} \quad (12) \]

where the sum is over KS-energies \( \epsilon_n \) and orbitals \( \phi_n(x) \) calculated for the uncoupled molecule.

Eq. (11) constitutes the main result of this section. Similar arguments can also be found in a recent communication by Stefanucci and Almbladh\(^\dagger\)

Several aspects of our finding should not pass by without further comment. a) Eq. (11) constitutes the generalization of the Landauer formula to interacting electron systems. In the special case, where \( \Gamma_L \) and \( \Gamma_R \) differ by a constant factor only, a derivation has already been given by Meir and Wingreen before\(^\dagger\). We emphasize, however, that this condition is extremely restrictive. It implies that every atom of the molecule is coupled in precisely the same way to the left lead as it is coupled to the right lead. Given the fact that physical couplings decay with increasing spatial distance, the “condition of proportional couplings” is violated for every realistic system with a finite extent.

b) As before the self energies \( \Sigma_{L,R} \) account for the coupling of the molecule to the leads. It is very well known that they can incorporate sophisticated many body effects, like i.e. the Abrikosov-Suhl resonance if Kondo-physics is present\(^\dagger\). However, one can restructure the problem and define an “extended molecule” that also comprises parts of the contacts, see Fig. 1. The new contact surface \( S_{cm} \) can be arranged sufficiently far away from the physical contact, so that the new self energy
depends on the type of lead only, but is totally inde-
pendent of which molecule is used. In fact, if a separ-
ation of energy scales exists such that the level spacing
of the extended molecule, $\delta_{IM}$, is much smaller than the
broadening, $\gamma_{IM}$, of the bare molecular levels upon cou-
ping to the leads the microscopic information carried by
$\Sigma_{L, R}$ becomes irrelevant. With any choice of $\gamma_{IM}$ in ac-
cord with Eq. (1) the simple replacement $\Sigma_{L, R}(x, x') =\gamma_{IM} \delta_{x x'}$ with $x, x'$ situated on $S_{IM}$ is justified. This
freedom merely reflects the experimentalists choice to at-
ach leads at convenience from any sort of shape or mate-
rial as long as the voltage drop is near the real molecule.
In this limit, all the detailed information about transport
properties of the molecule including correlation effects is
carried by the resolvent matrix (11) and inherited from
the exchange correlation potential.

c) Eq. 11 has been derived under the condition of van-
ishing temperature and frequency and in the regime of
linear response. Under these restrictions, scattering is
energy conserving and therefore an effective single parti-

cle scattering formulation of transport can exist. Upon
releasing the constraints the incoming particle can ex-
change energy with the molecule. Qualitatively new phe-
nomena can occur and in general the scattering problem
will become much more complicated and may not be un-
derstood in terms of a simple single particle picture. For
example memory effects appear, because here the incom-
ing electron sees the molecule in the state it has been left
in after interaction with the previous electrons.

d) Eq. (12) suggests that near equilibrium (i.e. for the
linear response) a variational principle may exist that al-

lows for the calculation of the voltage at a given current
or vice versa. The idea is to introduce a density operator
that maximizes the entropy under the constraint that the
current be finite. Work along this line has been done by
Kosov\textsuperscript{18}, recently.

B. Kubo Formalism of Linear Response

The practical usefulness of Eq. (11) is limited since
not much is known about the exchange correlation poten-
tial $V_{XC}(V)$ that defines the Kohn-Sham problem in
quasistatic nonequilibrium. Throughout all previous
works using the DFT approach to transport it has been
universally assumed that $V_{XC}(V)$ may be approximated by
its equilibrium form $V_{XC}(0)$ used in standard DFT-
calculations (EXCA). Let us discuss now, under which
conditions this approximation may be expected to hold.

Instead of solving a scattering problem one can also
find the conductance from the alternative, but equiva-

tent, Kubo formalism. The advantage of this starting
point for our purposes is that analytical statements about
the excitation frequencies are available which contain
information about the corrections to the EXCA.

We observe, that the current is related to the dynam-
ical polarization

$$I = dP/dt. \quad (13)$$

As a consequence the linear current response to a ho-
logenous external electric field is governed by the dy-
namical polarizability tensor of the entire system includ-
ing molecule and leads

$$I = -i\omega \alpha(\omega) E_{\text{ex}}, \quad (14)$$

which is closely related to the full density susceptibility

$$\alpha_{ij}(\omega) = \int dx \ dx' \ x_i \ x'_j \ \chi(x, x', \omega) \quad (15)$$

with the corresponding Lehmann representation

$$\alpha_{ij}(\omega) = \sum_{\mu=0} (\langle 0 | \hat{x}_i | \mu \rangle \langle \mu | \hat{x}_j | 0 \rangle / \omega - E_{\mu 0} + i0). \quad (16)$$

The sum is over all many body states with energies $E_{\mu}$
starting from the ground state $\mu = 0$ and the abbreviation
$E_{\mu 0} = E_{\mu} - E_0$ has been used. In the case of strongly reso-
nant transport the sum is dominated by the contribu-
tion of a few poles and the off-resonant current results from
the finite lifetime of the corresponding excitations which is
encoded in the dipole matrix elements. Clearly, this is the
regime in which molecules attached with thio-groups
find themselves.

The full density response contains a piece that de-
scribes dynamical screening. In terms of the bare KS-
response $\chi_{KS}(x, x', \omega)$ the relation

$$\chi^{-1}(\omega) = \chi_{KS}^{-1}(\omega) - v_{Q} - f_{XC}(\omega) \quad (17)$$

holds true, where the exchange-correlation kernel $f_{XC}(x, x') = dV_{XC}(x) / d\mu(x')$ has been introduced and
$v_{Q} = \| x - x' \|^\dagger$. In fact, Eq. (17) can be thought of as
a definition of $V_{XC}$. The crucial point is that the addi-
tional terms in (17) shift the excitation energies $\Omega_{\mu}$
away from their bare KS-values $\omega_{ij}^K = \epsilon_{i} - \epsilon_{j}$ (we introduce a multi-index $\mu = (i, j)$)

$$\Omega_{\mu} \approx \omega_{\mu}^{KS} + \langle \mu | v_Q | \mu \rangle + \langle \mu | f_{XC}(\omega_{\mu}^{KS}) | \mu \rangle \quad (18)$$

where $| \mu \rangle = | i, j \rangle$ denotes a single particle transition from
the occupied KS-orbit $j$ to the vacant orbit $i$. When $f_{XC}$
is decomposed into its exchange contribution and the cor-
relation part $f_{C}$ further analytical progress is possible\textsuperscript{19}
and one finds

$$\Omega_{\mu} \approx \omega_{\mu}^{KS} + \langle j | v_{x}^{HF} - v_{x}^{KS} | j \rangle - \langle ii \rangle v_{x}^{HF} - v_{x}^{KS} | i \rangle + \langle \mu | v_Q | \mu \rangle - \langle ii \rangle v_{x}^{Q} | j j \rangle + \langle \mu | f_{C} | \mu \rangle. \quad (19)$$

($v_{x}^{HF}$: exchange part of the bare Hartree-Fock (HF) oper-
ator, $v_{x}^{KS}$: exchange potential in the KS-equation). Eq. (19)
contains the plausible statement that the actual ex-
citation energies move away from their KS-values closer
to their Hartree-Fock estimates when dynamical screening
is taken into account\textsuperscript{19}. Note, that a corresponding
change must be expected to happen also in the dipole matrix elements of Eq. (16) which can be important since they control the lifetime of the excitations.

Equipped with this information we can proceed now and discuss the validity of the EXCA underlying previous DFT-transport calculations. Two limiting cases should be distinguished:

i) individual resonances strongly overlap, \( \gamma_M \gg \delta_M \). This is the limit of strong coupling, \( g \sim 1 \), where the wavefunctions \( |\psi_i\rangle \) are extended and the deviations from KS- and HF-excitation energies in Eq. (19) are small. In fact, for structureless, plane-wave-like states they vanish. In this case the standard approximation is justified. An example for a system in this class is a linear chain made from Au-atoms.

ii) individual resonances are fully developed, \( \gamma_M \ll \delta_M \). Here, the wavefunctions show pronounced localized features and in general the corrections to KS-excitation energies and matrix elements will not be small. While qualitatively correct results may still be found in many cases, a systematic quantitative analysis of the resonant structures in \( T(E) \) based on the EXCA is not possible, in general. A typical representative of weakly coupled systems is given in Fig. 1. The contact of the extended \( \pi \)-system to the leads is interrupted by an \( S \)-atom and therefore the molecular states are partly localized.

### III. NUMERICAL CALCULATIONS

The discussion presented in the previous section suggests that the experimental conductance of Au-chains is well described by the standard DFT-approach whereas the conductance of organic molecules coupled by thio-groups is only poorly captured. In the following section we present transport calculations making use of Eq. (12) together with the EXCA in order to corroborate this result.

#### A. Method

Our method is similar to approaches described in\textsuperscript{21–23}, however it has the advantage that using the program package TURBOMOLE we can include a considerably larger number of contact gold atoms. For the molecules of interest to us, this number (110) is sufficient so that the Fermi energy of the extended molecule is very close to the bulk value \( E_F \approx -5.1 \text{eV} \), even without attaching additional leads.

Since the jest of the method has been outlined previously\textsuperscript{21–23} we can limit ourselves to a brief description. The transmission is given by Eq. (11) but with a resolvent matrix \( G_0 \) that has been obtained employing the EXCA: KS-orbitals and KS-energies are taken from a standard DFT calculation\textsuperscript{24,25}. The self energies, \( \Sigma_{L,R} \), can be expressed in terms of the hopping matrix elements, \( t_{L,R}(X,N,xn) \) which describe a hopping process of an electron in an orbital state \( N \) of an atom at position \( X \) of the extended molecule to a state \( n \) of the atom at a location \( x \) in the leads: \( \Sigma_L = t_L g_L t_L^\dagger \) and similarly for \( \Sigma_R \). The hopping matrix elements we approximate by their bulk values that we obtain from an independent DFT calculation for a large gold cluster (146 atoms). Likewise, we replace the surface Green’s function \( \langle \phi_n|G_{L,R}|\phi_{n'}\rangle \) of the leads by a bulk one taken from the same calculation\textsuperscript{26}.

#### B. Results

Fig. 2 shows our results for the transmission of a linear chain of four equidistantly placed Au-atoms \( (d = 2.67 \text{\AA}) \). As a check we have performed calculations with leads that have been modeled by 54 (see Fig. 1) and 84 Au-atoms. Moreover the number of “surface” Au-atoms from the extended molecule that have been coupled to external leads has been varied: the self energy has been calculated with 29 and 41 atoms taken into account. It can be seen that the transmission is (essentially) independent of these parameters, as it should. Our results are in very good agreement with experiments and previous calculations\textsuperscript{27}.

Before we present our results for the transmission function the coupling of the organic molecule to the contacts should be discussed.

![Fig. 2. Transmission of a four atom gold chain. Upper panel: Contact was made of 54 gold atoms, see contacts in Fig. 1. Out of these 41 atoms (○, 5 × 5 and 4 × 4 bottoms layers) have been coupled to external leads. For comparison, results with 29 coupling atoms are shown as well (⊙). Lower panel: Same as upper panel with 84 gold atoms defining the contacts. \( (E_F \approx -5.1 \text{eV}) \) Traces are (nearly) independent of the modeling parameters chosen, and they agree well with experiments and previous calculations\textsuperscript{27}.](image-url)
In experiments a standard way to facilitate this coupling is to introduce a sulfur atom that forms a very strong bond with gold as well as with carbon atoms. Since the precise microscopic conditions of experiments are not known, several possibilities for the sulfur-gold bonds have to be considered. It is known, that sulfur tends to bind to three gold atoms on a plane Au(111) surface, i.e. the hollow site is the most stable one\(^\text{28}\). Binding to just one gold atom – on top position – corresponds to a local minimum of the free energy and binding to two gold atoms is unstable. We find that the situation on rough surfaces exhibiting edges, is different. Near an edge, the sulfur finds its most stable position by binding to two gold atoms (Fig. 1) in agreement with an earlier study on much smaller systems\(^\text{29,30}\). The on top position remains a minimum albeit at higher energy (+0.7eV) and the hollow position is unstable. Because in the break junction experiments of interest to us the sulfur atom is likely to be exposed to an irregular surface we consider all three cases.

In Figs. 3 and 4 we show the transmission and the I/V-characteristics of the molecule of Fig. 1. The traces correspond to the S-atom binding to one, two or three Au-atoms. These micro-deformations induce a slight shift of the transmission peak near −5.2eV by about 0.2eV.

Qualitatively, the observed features are in accord with experimental findings of [1], inset of Fig. 4: we find a conductance gap and the differential conductance exhibits a maximum at about 0.4V that stems from a resonant molecular level - the remnant of the HOMO of the bare molecule - about 0.2eV below the Fermi energy. The peak conductance and the zero bias conductance can differ by an order of magnitude and at larger voltages the conductance rapidly decreases before there is another increase, again. On the quantitative level serious discrepancies between our calculations and experiments persist: our value for the zero bias conductance 0.2e\(^2\)/h exceeds the experimental one, ∼ 10\(^{-5}\)e\(^2\)/h, by more than two orders of magnitude. Moreover, the width of the experimental peak is strongly temperature dependent and can decrease by an order of magnitude with temperature decreasing from 300K to 30K\(^\text{20}\). This strongly suggests that the experimental peak width at room temperature is determined by inhomogeneous broadening rather than a strong coupling to the leads.

The inhomogenous broadening can be understood on the basis of our results in Fig. 3. We have seen above that the position of the HOMO is roughly accounted for by our DFT procedure and it is mainly its damping that is overestimated. Therefore, the shift of the peak position upon changing the coupling of the molecule to the leads is indicative of a microscopic smearing mechanism: at sufficiently large temperatures the thermal average over the different types of S-Au-bonds leads to an effective broadening on the energy scale ≈ 0.3eV which indeed is in accord with experimental observations.

One might suspect that at least part of the reason for the large discrepancy between theoretical and experimental findings for the zero bias conductance is that in real experiments the molecule is exposed to various boundary conditions, e.g. stress, that prohibit the contact geometry to relax completely, which is what we have assumed in our calculation. Therefore, an important question is whether the conductance can be strongly affected by a slight change in microscopic degrees of freedom, like the S-Au-bond length or the bond angles defined in Fig. 5.

FIG. 3. Transmission of the molecule in Fig. 1 with sulfur atom coupling to one (solid), two (dashed) and three (dotted) gold atoms. (\(E_V \approx -5.1\text{eV}\)) Modifications in the atomic contact structure lead to a slight shift of the conductance peak that offer an explanation for the inhomogeneous peak broadening observed in experiments, see Fig. 4.

FIG. 4. Upper curves: dI/dV curve from data in Fig. 3. based on \(I(V) = \int dE T(E) (f(E + V/2) - f(E - V/2))\) with \(f(E)\) denoting the Fermi-function \((T = 300\text{K})\). Lower curves: experimental data from [1]. Different traces represent results from consecutive voltage sweeps. The theoretical and experimental traces exhibit the same qualitative features, but strong quantitative deviations in the conductance magnitude exist.

FIG. 5. Schematic representation of geometric degrees of freedom defining the sulfur-gold-coupling. (center atom: sulfur, left: gold surface, right: benzene-molecule)
FIG. 6. Transmission of benzene-1,4-dithiol for various parameters $\alpha$, $d$ and $\beta$ as defined in Fig. 4. Plot shows, that the transmission is robust against small structural modifications induced e.g. by strain. Upper panel: molecule, coupling to a Au(111) hollow site, fully geometry optimized (solid) and same molecule after rotation (dot-dashed, $\alpha = \pi/24$). Middle: change in bond length (optimal value: $d = 2.55$ Å), applied change 0.05 Å (dashed) and 0.1 Å (long dashed). Bottom: change in angle $\beta$ by $\pi/12$. ($E_F \approx -5.1$ eV)

Since our interest is in a qualitative question, we perform our study using the simpler molecule benzene-1,4-dithiol. Our findings for the transmission as a function of energy are in excellent agreement with recent results by Xue and Ratner\textsuperscript{31} and Stokbro \textit{et al.}\textsuperscript{32}. In Fig. 6, upper panel, we display the transmission of this molecule for the geometry optimized case and also after a subsequent rotation of the molecule about angle $\alpha = \pi/24$ defined in Fig. 4. The middle panel shows the impact of changing the sulfur-gold bond length $d$ by 0.05 Å and 0.1 Å, the lower channel exhibits the change upon changing the bond angle $\beta$ by $\pi/12$. These manipulations have only a small effect on the transmission confirming earlier findings\textsuperscript{31,33}. The example shows that the DFT-conductance does not change by orders of magnitude when changing details of the sulfur-gold bonds within reasonable limits.

Now, we are facing the following situation: On the one hand, the present calculations show that small variations in the nanostructure do not strongly influence the transmission. On the other hand, experiments are reproducible and do not show very strong fluctuations in the zero bias conductance, which implies that large variations in the atomic structure of molecule and contact do not exist. Furthermore, the chemical bond between S and Au is known to be very strong and stable – this is why sulfur has been chosen as the coupling element in the first place. Combining these three facts, we conclude that at present there is little evidence that modifications in the atomistic structure of the experimental contacts could exist, which are not properly included in the theoretical modeling, and therefore would hint to an explanation as to why the theoretical conductance exceeds the experimentally measured one so much.

FIG. 7. Transmission of benzene-1,4-dithiol calculated within DFT (solid Au54, as in Fig. 6; dot-dashed Au14), HF (dashed Au14) and extended Hückel (dotted Au14) with 14 and 54 atom gold clusters simulating the electrodes. DFT- and HF-calculations show large differences which indicates a breakdown of the EXCA explicated in section II B.

Instead, we advocate another resolution of the puzzle, namely that the mechanism described in section II becomes active because deviations from KS-states and HF-states are large. To illustrate that the proposed mechanism may indeed induce quantitative changes by orders of magnitude, we compare the transmission calculated within the approximations DFT, HF and extended Hückel. Fig. 7 exhibits the expected behavior: the DFT-transmission peaks are shifted as compared to the Hartree-Fock result and overlap much more strongly. In addition, the DFT-transmission has a broad background contribution only weakly depending on energy. It stems from electronic states that reside in the leads and “leak” into the molecule. The DFT-calculation overestimates tunneling of these states as well. The net result is a difference in the zero bias conductance by two orders of magnitude depending on whether the DFT-approximation or the HF-approximation is being used. Note, that because of the theoretical argument given in section II B it is not entirely clear, apriori, that it is necessarily DFT that gives the better approximation.

In order to reduce the numerical effort we have restricted ourselves to small electrodes consisting of 14 Au atoms each. By comparing the DFT-results for Au14 and Au54 depicted in Fig. 6 one can convince oneself, that this simplification leaves unaffected our main conclusion.

IV. CONCLUSIONS

The numerical results presented in the preceeding section are in full agreement with expectations based on the theoretical analysis performed in section II: while the
conductance of a Au-chain agrees well with experimental results, large quantitative discrepancies for organic molecule attached to gold electrodes via a thiol-bond exist. Arguments have been presented to the extent that these discrepancies cannot plausibly be explained by insufficient modeling.

The standard DFT-approach can be exact only in the case, where the ground state is represented by a single Slater determinant which in turn implies that HF is exact. For the molecule this is not the case, however. Since an approximation for the ground state based on the Slater-determinant of KS-orbitals is a mixture of the wavefunction of the exact ground state with its excitations, the character of the approximate state tends to be too delocalized. Consequently, the molecule-lead coupling comes out too strong and the level broadening is overestimated. We have given analytical arguments in favor of this picture.

In order to improve upon the standard DFT-approach to transport the exchange correlation potential should be replaced by the appropriate potential for the quasistatic nonequilibrium in order to include the corrections due to dynamical screening that are ignored otherwise. Unfortunately, at present such a potential is not known. However, our work suggests an alternative approach which is based on the Kubo formula. It enables us to to calculate the dynamical polarization of the molecule together with parts of the leads. Thereof we can obtain the dc-current in the limit of zero frequency. Work in this direction is under way.

Finally, let us emphasize that our results have implications for the calculation of nonequilibrium effects like the polarization of those electronic states that do not carry the current. Two effects that work in opposite directions occur. For a given current the voltage drop at the molecule is underestimated by DFT-based calculations (since $g$ is too large) and therefore so is the induced polarizing field. On the other hand, the charge response of localized states to this field is overestimated since they appear too metallic. Consequently, interpreting DFT based transport calculations for nonequilibrium effects is not straightforward and possible asymmetries in the I/V-characteristics could be blurred.

V. APPENDIX

We derive Eq. 4 from Eq. 5. The effective single particle Hamiltonian of the scattering problem is of the form $H = H_0 + H'$, where

$$ H_0 = \begin{pmatrix} h_L & 0 & 0 \\ 0 & h_C & 0 \\ 0 & 0 & h_R \end{pmatrix} \quad \quad H' = \begin{pmatrix} 0 & t_L^\dagger & 0 \\ t_L & 0 & t_R \\ 0 & t_R^\dagger & 0 \end{pmatrix} $$ (20)

The Hamiltonian of the uncoupled left (right) lead has been denoted by $h_L$ ($h_R$), the Hamiltonian of the central unit (molecule) is $h_C$ and the matrices $t_{L,R}$ are the couplings of the left and right leads to the central unit. If the molecule is uncoupled, the scattering states are of the form

$$ \Phi_l = (\phi_l, 0, 0) \quad \Phi_r = (0, 0, \phi_r). $$ (21)

When the coupling has been switched on, the new scattering states $\Psi_{l,r}$ can be obtained from a Lippmann-Schwinger equation

$$ \Psi_{l,r} = \Phi_{l,r} + \mathcal{G}(E+i0) H' \Phi_{l,r} $$ (22)

where $E$ labels the energy of the state and the resolvent matrix $\mathcal{G}(z) = [z - H]^{-1}$ has been introduced. The lesser function is defined by $(x=(x,t))$

$$ \mathcal{G}^{<}(x,x',E) = \sum_{\alpha=l,r} \int d(t-t') \Psi_{\alpha}(x) \Psi^{\ast}_{\alpha}(x') f_{\alpha} e^{iE(t-t')} $$ (23)

Using Eq. 22 a matrix representation can be derived

$$ \mathcal{G}^{<}(x,x',E) = \langle x | [1 + \mathcal{G} H'] g_0^{<} [1 + H' \mathcal{G}] | x' \rangle $$ (24)

with

$$ g_0^{<} = \begin{pmatrix} g_L^{<} & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & g_R^{<} \end{pmatrix} $$ (25)

and

$$ g_L^{<}(x,x',E) = \sum_{l} \int d(t-t') \phi_l(x) \phi_l^{\ast}(x') f_{l} e^{iE(t-t')} $$ (26)

and a similar equation for $g_R^{<}$. In Eq. 3 only those matrix elements of $\mathcal{G}^{<}$ are needed, that connect the left and right boundaries of the central unit. These elements are given by the term in Eq. 24 quadratic in the external coupling. Since

$$ H' g_0^{<} H' = \begin{pmatrix} 0 & 0 & 0 \\ 0 & \Sigma^{<} & 0 \\ 0 & 0 & 0 \end{pmatrix} $$ (27)

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with \( \Sigma^C = t_L g_L^C t_L^\dagger + t_R g_R^C t_R^\dagger \) and since \( G_0 \) is diagonal one has for the matrix element representing the central unit
\[
G^C = G^C \Sigma^C G^C \tag{28}
\]
and thus recovers Eq. (4). A similar derivation has also been given by Brandbyge et al. \(^{22}\)

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