Identification of the Atomic Scale Structures of the Gold-Thiol Interfaces of Molecular Nanowires by Inelastic Tunneling Spectroscopy

Firuz Demir$^{1,*}$ and George Kirczenow$^{1,†}$

$^1$Department of Physics, Simon Fraser University, Burnaby, British Columbia, Canada V5A 1S6

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We examine theoretically the effects of the bonding geometries at the gold-thiol interfaces on the inelastic tunneling spectra of propanedithiolate (PDT) molecules bridging gold electrodes and show that inelastic tunneling spectroscopy combined with theory can be used to determine these bonding geometries experimentally. With the help of density functional theory, we calculate the relaxed geometries and vibrational modes of extended molecules each consisting of one or two PDT molecules connecting two gold nanoclusters. We formulate a perturbative theory of inelastic tunneling through molecules bridging metal contacts in terms of elastic transmission amplitudes, and use this theory to calculate the inelastic tunneling spectra of the gold-PDT-gold extended molecules. We consider PDT molecules with both trans and gauche conformations bound to the gold clusters at top, bridge and hollow bonding sites. Comparing our results with the experimental data of Hihath et al. [Nano Lett. 8, 1673 (2008)], we identify the most frequently realized conformation in the experiment as that of trans molecules top-site bonded to both electrodes. We find the switching from the 42 meV vibrational mode to the 46 meV mode observed in the experiment to be due to the transition of trans molecules from mixed top-bridge to pure top-site bonding geometries. Our results also indicate that gauche molecular conformations and hollow site bonding did not contribute significantly to the experimental inelastic tunneling spectra. For pairs of PDT molecules connecting the gold electrodes in parallel we find total elastic conductances close to twice those of single molecules bridging the contacts with similar bonding conformations and small splittings of the vibrational mode energies for the modes that are the most sensitive to the molecule-electrode bonding geometries.

I. INTRODUCTION

Electrical conduction through single organic molecules bridging a pair of metal electrodes has been studied extensively, both experimentally and theoretically for more than a decade [1]. Knowledge of the microscopic details of the bonding between the molecule and the electrodes is required in order to understand the transport properties of these systems. For molecules amine-linked to gold electrodes, it has been argued [2] that only one bonding geometry should be important. However, for the more commonly studied molecular wires with gold-thiolate bonding, it has been suggested that a variety of molecule-metal interface geometries may occur and influence electrical conduction through these nanowires [1, 3–7]. Some possible geometries that have been proposed are those in which a sulfur atom of the molecule is located at a top site over a particular gold surface atom or over a bridge site between two atoms of the gold surface or over a hollow site of between three gold atoms [8–11]. However, which (if any) of these possibilities are actually realized in metal-molecule-metal nanowires has remained unclear [1]. Since a single molecule located between two electrodes andchemically bonded to both of them is not accessible to scanning microprobes that are sensitive to atomic-scale structure, direct experimental determination of the atomic scale geometries of the molecule-metal interfaces of metal-molecule-metal nanowires has been elusive and continues to be an important goal in this field [1]. In this paper, we show how inelastic tunneling spectroscopy (IETS) combined with ab initio computer simulations can distinguish between the different possible atomic-scale structures of the gold-thiolate interfaces of single-molecule nanowires and can therefore be used to definitively identify the molecule-electrode bonding geometries that are realized experimentally. A brief account of some of our preliminary results has already been published [12].

When the bias voltage applied across a molecular wire increases beyond the threshold value for the excitation of a vibrational quantum of the wire (to be referred to as a phonon) at low temperatures, a conductance step is observed in the current-voltage characteristic of the system. IETS experiments have measured the molecular wire phonon energies in this way for various molecules. [13–43] Theoretical studies of the IETS of molecular nanowires [12, 44–79] have also been carried out and have accounted for many aspects of the IETS data. These IETS experiments and comparison with the theoretical work have demonstrated conclusively that particular molecular species are involved in electrical conduction through metal-molecule-metal junctions by exploiting the fact that the measured phonon ener-
gies provide a fingerprint of the molecule involved in the conduction process. The effects of changes in the molecular conformation\cite{32, 41, 77} and in the orientation of molecules relative to the electrodes\cite{59, 79} on the inelastic tunneling spectra have also been studied. However, the possibility that IETS might identify the bonding geometries at the molecule-metal interfaces and thus resolve the long standing “contact problem” of determining the atomic scale structure of the molecule-electrode interfaces has only very recently begun to be explored\cite{12, 78}.

We consider one of the simplest organic molecules, 1,3-propanedithiolate (PDT), bridging gold electrodes. Electrical conduction through this molecular nanowire and its IETS have recently been studied experimentally by Hihath et al.\cite{29} using the statistical STM break junction technique: By repeatedly forming a gold-PDT-gold molecular nanowire, measuring its conductance characteristic and then breaking the wire and reforming it, Hihath et al.\cite{29} collected a large body of conductance and IETS data representing many separate realizations of this system. However, in common with previous break junction experiments\cite{1}, the specific geometry corresponding to any particular measurement was not identified.

In the present study, we calculated the relaxed geometries, vibrational normal modes, zero bias conductances, and IETS of trans and gauche PDT molecules bonded to pairs of gold clusters in top, bridge, and hollow conformations. An important aspect of the methodology that we used is that we carried out relaxations of the entire extended molecules that we studied, relaxing the positions of all of the atoms of the gold clusters as well as those of the atoms of the PDT molecule without any imposed constraints. This is necessary in order to obtain reliable results for the vibrational modes that have the largest amplitudes on the sulfur atoms that bond to the gold clusters and are therefore the most sensitive to the sulfur-gold bonding geometry. We also studied the dependence of our results on the sizes of the gold clusters and found good convergence of our results with increasing cluster size. Thus our results are expected to be relevant to PDT molecules bridging macroscopic gold electrodes as in experiments, and we do indeed find excellent agreement between our results and the experimental data of Hihath et al.\cite{29}. We identify the most frequently realized conformation in the experiment\cite{29} as that of trans molecules top-site bonded to both electrodes. We find the switching from the 42 meV vibrational mode to the 46 meV mode observed in the experiment\cite{29} to be due to the transition of trans molecules from mixed top-bridge to pure top-site bonding geometries. We also conclude that gauche molecular conformations and hollow site bonding did not contribute significantly to the experimental inelastic tunneling spectra.\cite{29} For pairs of PDT molecules connecting the gold clusters in parallel we find total elastic conductances close to twice those of single molecules bridging the contacts with similar bonding conformations and small splittings of the vibrational mode energy of interest.

The remainder of this paper is organized as follows: In Section II A, we outline how we calculate the equilibrium geometries and vibrational modes for the gold-PDT-gold molecular wires. In Section II B, we describe how the low bias elastic tunneling conductances of these molecular wires are calculated. In Section II C, we derive the perturbative expression that we use to calculate the conductance step heights associated with inelastic electron scattering in the molecular wires. In Section III A, we discuss the relaxed molecular wire geometries that we find. In Section III B, we present the results of our low bias elastic conductance calculations. In Section III C, we present the results of our calculations of the inelastic tunneling spectra of PDT molecules connecting gold electrodes and compare them with the experimental measurements of Hihath et al.\cite{29} We present our conclusions in Section IV.

II. THEORY

A. Calculation of the low energy conformations and vibrational normal modes

Our goal is to study inelastic tunneling processes that are sensitive to the structure of the gold-thiol interfaces. These processes typically involve excitation of vibrational modes with strong amplitudes on the sulfur atoms. Therefore, it is necessary to calculate accurate equilibrium geometries of the entire system being studied, including both the molecule and the gold electrodes, and the phonon energies and atomic displacements from equilibrium for the vibrational normal modes of the whole system. We do this by performing our calculations for extended molecules that consist of the molecule itself and two clusters of gold atoms (with up to 14 gold atoms per cluster) that represent the pair of electrodes to which the molecule binds.

In order for this to be practical, we perform ab initio relaxations of these extended molecules based on density functional theory (DFT)\cite{80, 81} with the GAUSSIAN 09\cite{82} package. We obtain relaxed structures for the various different bonding geometries and molecular conformations that we study by carrying out relaxations starting from different initial trial geometries. The relaxations are carried out with no constraints imposed on the geometries so that the relaxed geometries correspond to local minima of the total energy and therefore all of the computed vibrational modes about the relaxed geometries are well defined.

We note that in most previous calculations of the inelastic tunneling spectra of molecules bridging gold electrodes some or all of the gold atoms were frozen in the geometry of an ideal bulk gold crystal lattice. In that case the calculated geometries of the extended molecules were only partly relaxed and did not correspond to minima of the total energy. Therefore calculations of vibrational modes based on these partly relaxed geometries...
suffered from a lack of rigor. Furthermore in the experiment of Hihath et al.\cite{29} the molecular junctions were formed by crashing a gold STM tip into a gold substrate and then breaking the junction and repeating this process many times at cryogenic temperatures. This experimental “statistical STM break junction” methodology makes it very unlikely that the gold has the structure of an ideal bulk gold crystal anywhere near the PDT molecule that bridges the gold electrodes in the experiment. Thus models in which some of the gold atoms are frozen in the geometry of an ideal bulk gold crystal lattice are not expected to be appropriate for experiments of this type. This may explain why the authors of Ref. 78 had difficulty modeling the results of IETS experiments on molecular junctions formed by the mechanically controlled break junction technique.

In the present work, the atomic displacements from equilibrium in the normal modes and the corresponding frequencies and phonon energies are also calculated using DFT\cite{82}. By carrying out systematic calculations for gold clusters of different sizes, we have checked that our conclusions are independent of the cluster size for the larger clusters that we study and thus are applicable to molecules bridging the nanoscale tips of experimentally realized macroscopic gold electrodes.

We use the B3PW91 density functional (for the exchange-correlation energy) and the LanL2DZ pseudopotentials and basis sets.\cite{82, 83} However, we have repeated the calculations for some cases using the functional PBE0(PBE1PBE)\cite{82, 84} for comparison and found little change in the result. We calculated the vibrational frequencies without using any scaling factors and will, in what follows, compare the corresponding calculated phonon energies with the experimental data of Hihath et al.\cite{29} without making any adjustments.

### B. Calculations of the elastic conductances in the limit of low bias

Conduction through molecular nanowires in the low bias limit does not involve inelastic processes. We calculate the low bias conductances in this limit for the \textit{ab initio} extended molecule geometries obtained as described in Section II A using the extended Hückel model with the parameters of Ammeter et al.\cite{85} to estimate the electronic structures for these geometries. As is discussed in detail in Refs. 1 and 86, this methodology involves no fitting to any experimental data relating to transport in molecular wires or PDT molecules with gold contacts. Previous transport calculations based on extended Hückel theory have yielded elastic tunneling conductances in agreement with experiment for other molecules thiol bonded to gold electrodes\cite{86–90} and have also explained a number of transport phenomena observed in experiments on molecular arrays on silicon\cite{91–93} and graphene nanoribbons\cite{94} as well as electroluminescence data,\cite{95} current-voltage characteristics\cite{95} and STM images\cite{96} of molecules on complex substrates. Our calculated elastic tunneling conductances (at zero bias) for the gold-PDT-gold molecular nanowires are in good agreement with the values measured by Hihath et al.\cite{29}

We calculate the zero bias tunneling conductances $g$ for gold-PDT-gold molecular wires from the Landauer formula

$$
  g = g_0 \sum_{ij} \left| t_{ji}^v(i) \right|^2 \frac{2v_j}{v_i},
$$

where $g_0 = 2e^2/h$ and $t_{ji}^v(i)$ is the elastic transmission amplitude through the extended molecule at the Fermi energy $E_F$ of bulk gold calculated within extended Hückel theory using the standard parameter set of Ammeter et al.\cite{85} In the transmission amplitude, $\{0\}$ means that the extended molecule is in a relaxed geometry calculated as is discussed in Sec. II A. $i$ denotes an electronic state of a carrier that is coming from the left lead, and $j$ is the electronic state of a carrier that has been transmitted to the right lead. $v_j$ and $v_i$ are the corresponding electron velocities. The coupling of the extended molecule to the electron reservoirs was treated as in previous work\cite{86, 91–93, 97–101} by attaching a large number of semi-infinite quasi-one-dimensional ideal leads to the valence orbitals of the outer gold atoms of the extended molecule. This method of coupling the extended molecule to the reservoirs is successful because most of the resistance of this system is due to the weakly transmitting molecule itself (as distinct from the coupling between the extended molecule and the ideal leads) and because the many ideal leads behave similarly to a large number of phase-randomizing Büttiker probes\cite{102} in minimizing the influence of electronic dimensional resonances due to the finite sizes of the gold clusters of the extended molecule.

The transmission amplitudes $t_{ji}$ were found by solving the Lippmann-Schwinger equation

$$
  |\Psi\rangle = |\Phi_0\rangle + G_0(E)W|\Psi\rangle,
$$

where $|\Phi_0\rangle$ is an electron eigenstate of an ideal semi-infinite one dimensional left lead that is decoupled from the extended molecule, $G_0(E)$ is the Green’s function of the decoupled system of the ideal leads and the extended molecule, $W$ is the coupling between the extended molecule and leads, and $|\Psi\rangle$ is the scattering eigenstate of the complete coupled system. Since the basis set used in extended Hückel theory is non-orthogonal, the orthogonalization procedure introduced in Ref. 103 was used to facilitate the above calculations.

### C. Perturbative theory of the IETS of molecular wires

When the bias voltage $V$ applied across a molecular wire exceeds $\hbar \omega/e$ (where $\omega$ is the frequency of a vibrational mode of the system), it becomes energetically possible for electrons passing through the molecular wire to
emit phonons of that mode and a conductance step develops in the low temperature current voltage characteristic of the wire. Since the bias voltages involved are small, the mode frequencies that we calculate at zero bias using the ab initio approach described in Section II A are appropriate for estimating (via $|V| = \hbar \omega / e$) the values of the bias voltage at which conductance steps due to inelastic processes occur.

In order to estimate the heights of these conductance steps, we adopt a perturbative approach in the spirit of that proposed by Troisi et al. [49] who transformed the problem of calculating IETS intensities into an elastic scattering problem. However, unlike Troisi et al. [49], we formulate the present theory and results explicitly in terms of elastic electron transmission amplitudes $t^\alpha_{ji}$ through the molecular wire. Such perturbative theories are appropriate at low bias voltages provided that the elastic electron tunneling channel is well off resonance. [49] We shall assume this to be the case in the following derivation of the theory of IETS on which our numerical calculations of IETS intensities presented in Section III C are based.

In the harmonic approximation the vibrational Hamiltonian of an extended molecule is

$$H_{vib} = \sum_{in} \frac{\mathbf{p}_n^2}{2m_n} + \frac{1}{2} \sum_{injm} V_{injm} q_{in} q_{jm} ,$$

where $\mathbf{p}_n$ and $q_{in}$ are the $i^{th}$ atomic momentum and coordinate of atom $n$. They obey the commutation rules $[q_{in}, p_{jm}] = i\hbar \delta_{nj} \delta_{nm}$. The position coordinate $q_{in}$ is measured from the equilibrium position of the atom $n$.

In terms of the creation and destruction operators $a^\dagger_\alpha$ and $a_\alpha$ for phonons of the normal modes $\alpha$ of the extended molecule with frequencies $\omega_\alpha$, the harmonic Hamiltonian takes the form

$$H_{vib} = \sum_{\alpha} \hbar \omega_\alpha \left( a^\dagger_\alpha a_\alpha + \frac{1}{2} \right)$$

and the atomic position coordinate operators are

$$q_{in} = \sum_{\alpha} \sqrt{\frac{\hbar}{2\omega_\alpha}} (a^\dagger_{i\alpha} a_\alpha + d_{i\alpha} a^\dagger_{i\alpha}) .$$

(3)

Here the coefficients $d_{i\alpha}$ are the displacements from their equilibrium positions of the atoms $n$ of the extended molecule in normal mode $\alpha$ normalized so that $\sum_n m_n d_{i\alpha}^* \cdot d_{i\alpha} = \delta_{i\alpha}$ and $m_n$ is the mass of atom $n$.

Let us now approximate the interaction Hamiltonian $U(\mathbf{r})$ describing the interaction between an electron at position $\mathbf{r}$ and the extended molecule by a sum of terms

$$U(\mathbf{r}) = \sum_{n} u_n (\mathbf{r} - \mathbf{R}_n) ,$$

(4)

corresponding to the interactions of the electron with the individual atoms $n$ of the extended molecule whose nuclei are located at $\mathbf{R}_n$.

Then the change in the Hamiltonian $U(\mathbf{r})$ if the atoms are displaced from their equilibrium positions $\mathbf{R}_n^0$ is

$$\Delta U(\mathbf{r}) = \sum_{n} u_n (\mathbf{r} - \mathbf{R}_n) - u_n (\mathbf{r} - \mathbf{R}_n^0) .$$

(5)

To leading order in the displacements $q_{in}$ of the atoms from their equilibrium positions, this becomes

$$\Delta U(\mathbf{r}) \cong \sum_{in} -q_{in} \frac{\partial u_n (\mathbf{r} - \mathbf{R}_n^0)}{\partial r_i} .$$

(6)

Using Eq. (3), this yields the electron-phonon interaction Hamiltonian $H_{ep}$

$$H_{ep} = \Delta U(\mathbf{r}) \cong -\sum_{\alpha} \sqrt{\frac{\hbar}{2\omega_\alpha}} u_{in}^\alpha \sum_{in} d_{i\alpha} \frac{\partial u_n (\mathbf{r} - \mathbf{R}_n^0)}{\partial r_i} + \text{h.c.}$$

If $A$ is a suitable scale factor and $d_{i\alpha}$ is the vector with components $d_{i\alpha}$ this becomes

$$H_{ep} \cong \sum_{\alpha} \tilde{A}_{i\alpha} H_{\alpha}(\mathbf{r}) + \text{h.c.}$$

where

$$H_{\alpha}(\mathbf{r}) = \sqrt{\frac{\hbar}{2\omega_\alpha}} \lim_{A \to 0} \sum_{n} u_n (\mathbf{r} - (\mathbf{R}_n^0 + A \tilde{d}_{\alpha})) - u_n (\mathbf{r} - \mathbf{R}_n^0) \frac{A}{A} .$$

For the system of interest in this paper, the electron-phonon interaction $H_{ep}$ is a small perturbation. The operators $a^\dagger_{i\alpha}$ and $a_{\alpha}$ induce transitions between states with differing numbers of phonon quanta while $H_{\alpha}(\mathbf{r})$ simultaneously gives rise to transmission of electrons through the molecular wire in the inelastic channel with scattering amplitude $t^\alpha_{ji}$. Assuming that the system is cryogenic so that the presence of any ambient phonons may be ignored, and that $H_{\alpha}$ is so weak that $t^\alpha_{ji}$ is effectively linear in the strength of $H_{\alpha}$, and that $A$ is small enough that

$$H_{\alpha}(\mathbf{r}) = \sqrt{\frac{\hbar}{2\omega_\alpha}} \sum_{n} u_n (\mathbf{r} - (\mathbf{R}_n^0 + A \tilde{d}_{\alpha})) - u_n (\mathbf{r} - \mathbf{R}_n^0) \frac{A}{A}$$

to a good approximation, it follows that

$$\tilde{t}_{ji}^\alpha \cong \sqrt{\frac{\hbar}{2\omega_\alpha}} \frac{1}{A} \tilde{t}_{ji}^\alpha ,$$

(7)

where $\tilde{t}_{ji}^\alpha$ is the inelastic electron transmission amplitude that would be found if $H_{\alpha}$ were

$$\tilde{H}_{\alpha}(\mathbf{r}) = \sum_{n} u_n (\mathbf{r} - (\mathbf{R}_n^0 + A \tilde{d}_{\alpha})) - u_n (\mathbf{r} - \mathbf{R}_n^0) .$$
That is, \( \tilde{t}_{ji}^\alpha \) is the change in the elastic electron transmission amplitude when the perturbation \( \tilde{H}_\alpha \) is added to the electronic Hamiltonian \( H_{\text{equil. static}} \) of the static molecular wire in its equilibrium geometry. But \( H_{\text{equil. static}} + H_\alpha \) is simply the electronic Hamiltonian of a static molecular wire whose nuclear positions \( R_n \) are \( R_n = R_n^0 + \mathbf{A} \mathbf{d}_{\alpha n} \), i.e., they are shifted from their equilibrium positions by vectors \( \mathbf{A} \mathbf{d}_{\alpha n} \) that are proportional to the displacements of the atoms from their equilibrium positions in normal mode \( \alpha \).

This means that
\[
\tilde{t}_{ji}^\alpha = t_{ji}^{\text{el}}(\{ \mathbf{A} \mathbf{d}_{\alpha n} \}) - t_{ji}^{\text{el}}(\{ 0 \}),
\]
where \( t_{ji}^{\text{el}}(\{ \mathbf{A} \mathbf{d}_{\alpha n} \}) \) is the elastic electron transmission amplitude through the molecular wire with each atom \( n \) displaced from its equilibrium position by \( \mathbf{A} \mathbf{d}_{\alpha n} \) where \( A \) is a small parameter and \( t_{ji}^{\text{el}}(\{ 0 \}) \) is the elastic electron transmission amplitude through the molecular wire in its equilibrium geometry from state \( i \) with velocity \( v_i \) in the electron source to state \( j \) with velocity \( v_j \) in the electron drain. It then follows from Eq. (7) that
\[
t^\alpha_{ji} \approx \sqrt{\frac{\hbar}{2\omega_\alpha}} \lim_{A \to 0} \frac{t_{ji}^{\text{el}}(\{ \mathbf{A} \mathbf{d}_{\alpha n} \}) - t_{ji}^{\text{el}}(\{ 0 \})}{A}.
\]
The analog of Eq. (1) for the inelastic electron transmission probability associated with the emission of a phonon of mode \( \alpha \) is then
\[
I^\alpha_{ji} = \sum_{ij} \frac{|v_i| |v_j|^2}{|v_i|} = \frac{e^2}{2\omega_\alpha} \lim_{A \to 0} \frac{\sum_{ij} \frac{v_i}{v_j} \left| t_{ji}^{\text{el}}(\{ \mathbf{A} \mathbf{d}_{\alpha n} \}) - t_{ji}^{\text{el}}(\{ 0 \}) \right|^2}{A},
\]
From this it follows immediately that the IETS intensity (i.e., the conductance step height) \( \delta g^\alpha \) associated with the emission of phonons of mode \( \alpha \) is
\[
\delta g^\alpha = g_T T^\alpha, \quad i.e.,
\]
\[
\delta g^\alpha = \frac{2e^2}{\hbar} \frac{\hbar}{2\omega_\alpha} \lim_{A \to 0} \frac{\sum_{ij} \frac{v_i}{v_j} \left| t_{ji}^{\text{el}}(\{ \mathbf{A} \mathbf{d}_{\alpha n} \}) - t_{ji}^{\text{el}}(\{ 0 \}) \right|^2}{A}, \quad \text{or}
\]
\[
\delta g^\alpha = \frac{e^2}{2\pi\omega_\alpha} \lim_{A \to 0} \frac{\sum_{ij} \frac{v_i}{v_j} \left| t_{ji}^{\text{el}}(\{ \mathbf{A} \mathbf{d}_{\alpha n} \}) - t_{ji}^{\text{el}}(\{ 0 \}) \right|^2}{A}.
\]
Eq. (8) expresses the IETS intensities \( g^\alpha \) explicitly in terms of elastic electron transmission amplitudes \( t_{ji}^{\text{el}} \) through the molecular wire. It states that, in the leading order of perturbation theory, the scattering amplitude for inelastic transmission of an electron through the molecular wire is proportional to the change in the elastic amplitude for transmission through the wire if its atoms are displaced from their equilibrium positions as they are when vibrational mode \( \alpha \) is excited. We evaluate \( t_{ji}^{\text{el}} \) in Eq. (8) numerically to find the heights \( \delta g^\alpha \) of the conductance steps that arise from inelastic tunneling processes due to emission of phonons of vibrational mode \( \alpha \) by applying the methodology described in Section II B. These calculations are carried out at the zero bias Fermi energy since the values of the bias at which the inelastic transmission occurs in the experiments of Hihath et al. [29] are low.

### III. RESULTS

#### A. Low energy conformations of the extended molecules

As has been discussed in Section II A, we considered extended molecules, i.e., molecules together with finite clusters of gold atoms to which the molecules bond, and relaxed these entire structures (the molecules and gold clusters) using density functional theory-based calculations.[80–84] No constraints were applied to the extended moleucle geometries during the relaxation. This yielded calculated geometries that correspond to local energy minima of the extended molecules.

Representative examples of the relaxed geometries that we obtained for extended molecules in which the PDT molecule has the trans conformation are shown in Fig. 1. Top, bridge and hollow-site geometries where each sulfur atom of the molecule bonds to one, two, or three gold atoms, respectively, are shown in Fig. 1 (a), (c) and (d). A mixed bridge-top structure where one sulfur atom bonds to two gold atoms and the other bonds to one gold atom is shown in Fig. 1 (b).

For relaxed extended molecules having structures of the types depicted in Fig. 1, we found the extended molecules with the sulfur atoms binding to the gold clusters in bridge geometries to have the lowest energies. The hollow site bonding geometries had the highest energies. The top site bonding geometries had energies between those of the bridge and hollow cases. For example, the energy of the relaxed extended molecule in the bridge
geometry was 0.46 eV lower than that with the top geometry and 1.33 eV lower than that with the hollow geometry for our molecular wires with 13 Au atoms in each gold cluster. These numbers varied depending on the gold cluster size. But in all studied cases, the bridge geometry was more stable than the top geometry which was more stable than the hollow geometry for relaxed structures.

We found that extended molecules for which DFT geometry relaxations were started with the sulfur atoms over hollow sites on the surfaces of close packed gold clusters having atomic geometries resembling that of bulk fcc gold invariably relaxed to bridge bonding site geometries. This is reasonable physically since the bridge site is not very far from the hollow site and a drastic rearrangement of the gold clusters is not required for a transition from hollow site bonding to a lower energy bridge site bonding geometry to occur. Thus while we were able to generate examples of relaxed extended molecule geometries (such as that in Fig. 1 (d)) with each sulfur atom bonding to three gold atoms (i.e., hollow site bonding) the structures of the gold clusters near these bonding sites were much more open than that of a defect-free surface of a fcc gold crystal near a hollow site. Because of the much greater fragility and the higher energies of hollow site bonded structures relative to bridge and top site bonding, it is reasonable to expect hollow site bonding to be realized much less frequently than bridge and top site bonding (if it is realized at all) in statistical STM break junction experiments such as those of Hihath et al. As will be seen below, comparison of our theoretical inelastic tunneling spectra with the experimental data of Hihath et al supports this expectation.

**B. Elastic conductances in the limit of low bias**

Our calculated zero bias conductance values for top, bridge, and hollow site bonded trans PDT molecules are in the ranges $g_{trans} = 0.0012 - 0.0013g_0$, $0.0015 - 0.0019g_0$, and $0.0032 - 0.0035g_0$, respectively. Gauche1: One gauche C-C bond. Gauche2: Two gauche C-C bonds. Numbers in the configuration column indicate the numbers of atoms in the gold clusters. Notice that the low bias conductance decreases with increasing numbers of gauche bonds for similar bonding geometries, consistent with previous theories for longer molecular wires.

**TABLE I.** Calculated IETS intensities and phonon energies for Mode-I for one or two PDT molecules connecting gold clusters together with the calculated low bias conductances. T: top site bonding, B: bridge site bonding, and H: hollow site bonding of a molecule to a gold clusters. TT and BB indicate two molecules connecting the gold clusters in parallel and bonding in top and bridge geometries, respectively. Gauche1: One gauche C-C bond. Gauche2: Two gauche C-C bonds. Numbers in the configuration column indicate the numbers of atoms in the gold clusters. Notice that the low bias conductance decreases with increasing numbers of gauche bonds for similar bonding geometries, consistent with previous theories for longer molecular wires.

| Configuration | Phonon Energy of Mode-I (meV) | IETS Intensity of Mode-I ($10^{-6}g_0$) | Low Bias Conductance ($g_0$) |
|---------------|-------------------------------|----------------------------------------|-----------------------------|
| 14T–14 Trans  | 44.88                         | 10.03                                  | 0.0013                      |
| 14TT–TT14 Trans | 44.88                         | 11.06                                  | 0.0029                      |
| 14B–B14 Trans | 40.05                         | 15.97                                  | 0.0016                      |
| 14BB–BB14 Trans | 40.67                         | 9.76                                   | 0.0031                      |
| 14B–14 Trans  | 42.03                         | 9.54                                   | 0.0016                      |
| 13H–H13 Trans | 42.40                         | 34.02                                  | 0.0095                      |
| 13H–T13 Trans | 43.52                         | 11.74                                  | 0.0029                      |
| 14B–B14 Gauche1 | 39.06                         | 13.44                                  | 0.00133                     |
| 14B–B14 Gauche2 | 36.58                         | 104.97                                 | 0.00071                     |
| 14T–14 Gauche1 | 38.44                         | 1.80                                   | 0.00034                     |
| 14T–14 Gauche2 | 36.58                         | 1.54                                   | 0.00007                     |

* Experimental conductance measurement of Hihath et al. is about 0.006 $g_0 \pm 0.002$.

**C. Inelastic Tunneling Spectroscopy of the Gold-Sulfur Interface**

In this Section we present the results of our calculation of the inelastic tunneling spectra for trans PDT
molecules bonded to gold clusters in top, bridge, and hollow site conformations as well as for molecules with gauche bonds. We also consider some examples of pairs of molecules bridging the gold clusters. We compare our results with the experimental data of Hihath et al. [29] and deduce which molecular and bonding conformations were most commonly realized in the experiment. In the calculations reported below, density functional theory was used to find the relaxed extended molecule geometries, vibrational modes and their frequencies as described in Sections II A and III A. The inelastic tunneling intensities \( \delta g_{\alpha} \) corresponding to the various modes were calculated from Eq. (8), evaluating the elastic scattering amplitudes \( t_{ji}^{el} \) that enter Eq. (8) as is described in Section II B. We will focus primarily on those vibrational modes that have the strongest amplitudes of vibration on sulfur atoms and the strongest IETS intensities since these modes are the most sensitive to the molecule electrode bonding geometries and are also prominent in the experimental IETS spectra.

1. **Top and bridge site bonding of trans PDT molecules**

   Our calculations show that for trans PDT molecules the modes with strong amplitudes of vibration on the sulfur atoms that have the largest IETS intensities fall within the phonon energy range of a prominent feature of the experimental IETS phonon histogram [29] (shown in grey in Fig. 2(b)) that extends from 39 to 52 meV.

   The vibrational normal modes in this energy range and their phonon energies are shown in Fig. 2(a) for examples of extended PDT molecules with sulfur atoms bonded to 14-atom gold clusters in top and bridge geometries. The vibrational modes are labelled I and II according to the nature of the atomic motion. The corresponding calculated IETS spectra (IETS intensities vs. phonon energy) are shown in color in Fig. 2(b) for extended molecules with gold clusters of various sizes together with the experimental IETS phonon mode histogram [29].

   It is important to note that that vibrational modes I and II are strongly localized to the molecule itself and the gold atoms in its immediate vicinity. Their amplitudes of vibration decay rapidly in the gold clusters as the distance from the molecule increases. This is due to the fact that the frequencies of these modes are well above the phonon frequencies of bulk gold so that in the gold these modes are evanescent in character. For the largest gold clusters that we studied, the vibrational amplitudes of modes I and II are very small already before the farthest gold atoms from the molecule are reached. Therefore, the calculated properties of these modes are not impacted significantly by the finite sizes of the gold clusters in the present study.

   The mode with the strongest calculated IETS intensities in Fig. 2(b) is Mode-I that is depicted in the top row of Fig. 2(a). In this mode (also known as the Au-S symmetric stretch mode) the sulfur atoms have the strongest vibrational amplitudes and move in antiphase, approximately along the axis of the molecule. Mode-II (also known as the Au-S antisymmetric stretch mode) is shown in the lower row of Fig. 2(b). It is similar to Mode-I except that in Mode-II the sulfur atoms move in phase with each other. As seen in Fig. 2(b) the calculated IETS intensities for Mode-II are much weaker than those for Mode-I for both top and bridge site bonding.

   This difference between the IETS intensities of modes
I and II can be understood physically as follows. Since in Mode-I the two sulfur atoms move in antiphase the gold-sulfur distances for both sulfur atoms either increase or decrease together as the extended molecule vibrates. These distances can be regarded as the widths of tunnel barriers between the molecule and the two gold electrodes. The motions of the two sulfur atoms act in concert to widen or narrow both tunnel barriers together and thus to weaken or strengthen the electron transmission amplitude through the molecular wire. Therefore, the magnitude of the difference between the elastic electron transmission amplitudes of equilibrium and vibrating geometries through the molecular wire in Eq. (8) is enhanced. By contrast, in Mode-II, while the gold-sulfur distance for one sulfur atom increases, the gold-sulfur distance for the other sulfur atom decreases. These effects of the motions of the two sulfur atoms on the elastic transmission amplitude through the molecular wire tend to cancel. As a result, the magnitude of the difference $\delta_{ji}^{el}(\{A_{na}\}) - t_{ji}^{el}(\{0\})$ that appears in Eq. (8) is smaller for Mode-II than for Mode-I and consequently the IETS intensity $\delta g_{\alpha}$ is seen in Fig. 2(b) to be much weaker for Mode-II. For this reason, although the amplitudes of the motion of the sulfur atoms in modes I and II are similar and in both cases the motion is approximately along the molecular axis, Mode-I completely dominates the calculated IETS spectrum in the phonon energy range that corresponds to the main feature of the experimental IETS phonon histogram [29] in Fig. 2(b).

That the IETS intensity for mode I is much larger than that for mode II can also be understood in terms of the Troisi-Ratner propensity rules [57, 58] for inelastic tunneling intensities. For a very simple model of a perfectly symmetric molecule with symmetric electrodes, Troisi and Ratner [58] found the IETS intensity for mode II to be exactly zero as a consequence of symmetry. The IETS intensity for mode II is not (in general) exactly zero in the present model because the electrodes are not symmetric. We note that the electrodes are also not expected to be symmetric in current experimental molecular wire devices such as that of Hihath et al. [29].

Notice that our theoretical results for phonon Mode-I of PDT molecules bonded to both electrodes in the top site geometry, bonded to both electrodes in the bridge geometry, and bonded to one electrode in the bridge and to the other in the top geometry (the features around 45.5, 40.5, and 42 meV in the theoretical spectra in Fig. 2(b), respectively) are reasonably well converged with respect to increasing gold cluster size. Also the Mode-I phonon energy ranges of the bridge site-bonded and bridge/top site-bonded wires do not overlap and are well separated from the phonon energy range of Mode-I for pure top site bonding.

Based on these theoretical results, we propose the following identification of the bonding geometries between the PDT molecules and gold electrodes that gave rise to the main features of the experimental IETS histogram [29] that is reproduced in Fig. 2(b): We identify the main peak in the experimental histogram [29] in Fig. 2(b) that is centred at ~46 meV as being due to trans-PDT molecules that bonded to both gold electrodes in the top site geometry. We thus attribute the majority of the counts in the histogram to molecules in the pure top site bonding geometry. The weaker peak centred near 42 meV in the experimental histogram matches our predictions for molecules that bond to gold electrodes in the mixed bridge/top site geometry. The shoulder of the experimental histogram [29] at lower phonon energies centered near 40.5 meV corresponds to our results for molecules bonding to both electrodes in the bridge site geometry. Thus, our results show that IETS experiments can identify experimental realizations of PDT molecular wires in bridge, bridge/top, and top site bonding geometries by measuring the energies of the phonons emitted during electron transport through these systems.

Although the feature near 42 meV in the experimental histogram is not very prominent, Hihath et al. [29] observed a transition in which the IETS spectrum switched from exhibiting a peak near 42 meV to one near 46 meV as the molecular junction was stretched. They therefore argued that the 42 meV and 46 meV vibrational modes correspond to distinct conformations of the molecular junction. However, they were not able to identify the conformations that give rise to these modes. Our theoretical results indicate that the switching from the 42 meV to the 46 meV mode corresponds to the junction switching from trans PDT in a mixed top/bridge bonding geometry to trans PDT in a pure top bonding geometry as the junction is stretched. This is consistent with our finding that the pure top bonding geometry corresponds to a larger separation between the gold electrodes than does the top/bridge bonding geometry.

In addition to modes I and II, the theoretical IETS spectrum in Fig. 2(b) shows other vibrational modes (labelled “Mixed Modes”) at lower energies, between 28 and 36 meV. The calculated IETS intensities of these modes are weaker than those of Mode-I but in most cases not as weak as those of Mode-II. In the same energy range, the experimental histogram in Fig. 2(b) [29] shows very few counts. This suggests that the measurements of Hihath et al. [29] were not able to detect features in the inelastic tunneling spectrum having weak intensities and that the intensities of the “Mixed Modes” between 28 and 36 meV in Fig. 2(b) (as well as those of Mode-II) were at or below the detection threshold.

2. Hollow site bonding

For alkanedithiolates with longer chains of (6, 8, or 10) carbon atoms bridging gold electrodes, the low bias conductances have been observed experimentally to drop by a factor of ~4 from the second lowest conductance plateau to the lowest as the molecular junction is stretched. [5] It was conjectured [5] that this change in conductance may be due to the molecules switching
details in the hollow site bonding and the resulting greater variability structures (see Table I), this greater sensitivity is due to the calculated elastic conductances of hollow site bonded pure bridge or mixed bridge/top site bonding. As for the sizes of the gold clusters than they are for pure top, IETS intensities for these structures are more sensitive to are shown in Fig. 3. The calculated phonon energies and molecules hollow site-bonded to one or both gold clusters
mentally) than bridge or top site bonding.
more fragile (and thus less likely to be realized experi-
tries. This indicates that the hollow site bonding is much
started in hollow site bonded trial geometries in most
cases relaxed immediately to bridge-site bonded geome-
tion, in our DFT calculations PDT extended molecules
of the gold-PDT-gold system? As we have already men-
ted, in our DFT calculations PDT extended molecules
the change between gauche and gauche bonds would affect the inelastic tunneling spectra in the vicinity of either 42 or 46 meV mode (as their gold-PDT-gold junction was stretched) might have been due to a change between gauche and trans configurations of the molecular junction.

We have investigated theoretically how the presence of gauche bonds would affect the inelastic tunneling spectra of PDT molecules bridging gold contacts and the possible implications for the experiment.[29] Representative results of our calculations are presented in Fig 4 (a) and (b) for PDT molecules with one and two gauche bonds connecting pairs of gold clusters in bridge site and top site bonding configurations. For the bridge site bonding configurations, the calculated energies of the vibrational modes of the gauche PDT molecules are not in the vicinity of either 42 or 46 meV. However, the calculated spectra exhibit vibrational modes with very strong IETS intensities at phonon energies between 32 and 38 meV, a range where the experimental histogram[29] in Fig. 4 (b) shows very few counts. Thus it appears unlikely that these structures played a significant role in the experiment of Hihath et al.[29].

In Fig. 3 the energy range in which we find Mode-I phonons for pure hollow site bonding overlaps with the
~42 meV feature of the experimental IETS histogram of Hihath et al.[29]. However, we find the structures with pure hollow site bonding to also support phonon modes with strong calculated IETS intensities (similar in strength to the Mode-I intensities for the top and bridge site bonded structures in Fig. 2(b)) in the mixed mode region below ~38 meV in Fig. 3 where the experimental IETS histogram shows few or no counts. This suggests that structures with pure hollow site bonding were realized rarely (if at all) in the experiment of Hihath et al.[29].

For mixed hollow/top bonding the calculated Mode-I phonon energy in Fig. 3 does not match either the ~42 or the ~46 meV feature in the experimental histogram [29] but falls roughly half way between the two. Thus, in addition to the formation of bridge/top structures being much more likely (based on our ab initio simulations) than the formation of hollow/top structures, the bridge/top bonding geometries account for the phonon energy of the 42 meV feature of the experimental[29] IETS spectra while the hollow/top bonding geometries do not. Thus it appears unlikely that hollow/top structures played a significant role in the experiment of Hihath et al.[29].

3. Gauche molecular conformations

A gauche conformation of an alkanedithiolate molecule can be obtained starting from the trans conformation by rotating part of the molecule through ~60° about a C–C or C–S bond. Detection of gauche conformations of PDT molecules adsorbed on gold has been reported[108]. Also it has been conjectured [29] that the switching that Hihath et al. observed[29] in their inelastic tunneling spectra from the 42 meV vibrational mode to the 46 meV mode (as their gold-PDT-gold junction was stretched) might have been due to a change between gauche and trans configurations of the molecular junction.

In Fig. 4 (b) for PDT molecules with one gauche bond and top site bonding does exhibit a vibrational mode

![Fig. 3. (Color online.) Calculated IETS intensities (colored) vs. calculated phonon energies for trans-PDT molecules linking pairs of gold clusters with 12 (yellow) and 13 (green) Au atoms in each cluster for pure hollow site bonding and with 13 Au atoms in each cluster for mixed hollow/top site bonding (magenta). The experimental IETS phonon mode histogram of Hihath et al.[29] is shown in (darker, lighter) grey for (positive, negative) bias voltages.](image-url)
FIG. 4. (Color online.) (a) Representative examples of calculated sulfur-related vibrational Mode-I and II of gauche-PDT molecules connecting gold nanoclusters each with 14 gold atoms, and the vibrational mode phonon energies. G1 and G2 label molecules with one and two gauche bonds respectively. The molecules connect to both gold clusters in pure bridge and pure top site bonding configurations. Carbon, hydrogen, sulfur, and gold atoms are black, grey, yellow, and amber, respectively. Red arrows show un-normalized atomic displacements. (b) IETS intensities of gauche-PDT molecules linking pairs of gold clusters (14 Au atoms in each cluster) vs. the vibrational mode phonon energy. Sulfur atoms bond to the gold in top-top, bridge-bridge site geometries. The experimental IETS phonon mode histogram of Hihath et al. [29] is shown in (darker, lighter) grey for (positive, negative) bias voltages. Modes of the types I and II are indicated by arrows.

* Scaled down by a factor of 10.

with a phonon energy close to 46 meV. However, this is

a type II mode with a low calculated IETS intensity. It is therefore unlikely to have been detected in the IETS experiment.[29] Other calculated vibrational modes of the same top-site bonded gauche structure in the phonon energy range from 27 to 36 meV in Fig. 4 (b) have somewhat stronger but still weak IETS intensities and occur at energies at which the experimental histogram shows few if any counts. The PDT molecule with one gauche bond and top site bonding in Fig. 4 (b) also exhibits a vibrational mode with a phonon energy close to 46 meV but the calculated IETS intensity for this mode is much weaker still. These results indicate that gauche PDT molecules in top site bonding configurations also appear unlikely to have contributed significantly to the experimental IETS histogram of Hihath et al.[29]. We have also carried out calculations for gauche PDT molecules bonded to the gold electrodes in mixed top-bridge geometries. The results of these calculations (not shown in Fig. 4) are qualitatively similar to those for the pure bridge bonding geometries of gauche molecules discussed above. Namely, these structures were also found not to have vibrational modes with calculated energies in the vicinity of either 42 or 46 meV but they were found to exhibit vibrational modes with very strong IETS intensities at phonon energies between 32 and 38 meV, where the experimental histogram[29] shows very few counts. Thus it appears unlikely that these mixed bridge-top bonded gauche molecules played a significant role in the experiment of Hihath et al.[29].

The low bias elastic conductance measurements reported by Hihath et al.[29] also provide additional evidence indicating that switching between gauche and trans conformers is unlikely to have been responsible for the experimentally observed transition[29] from the structure with the 42 meV vibrational mode to that with the to 46 meV mode, as will be discussed in the following two paragraphs.

We find the calculated sulfur-to-sulfur distances in PDT molecules with gauche bonds to be shorter than those for purely trans PDT molecules. Thus, if the transition from the 42 meV vibrational mode to the 46 meV mode (that occurred as the junction was stretched) were due to switching between gauche and trans conformations, then the 42 meV structure (being the shorter one) would correspond to the gauche geometry and the 46 meV structure would have the trans geometry. However, the measured conductances of the PDT molecules exhibiting the 42 meV vibrational mode were larger than those of the PDT molecules exhibiting the 46 meV mode, as can be seen in Fig.4 of Ref. 29. By contrast, recent theoretical work[77, 107] on other alkane-dithiolate molecules bridging gold electrodes found conductances of pure trans-molecules to be larger than those of molecules with gauche bonds, a finding consistent with the results of our calculations of the low bias conductances for gauche and trans configurations of gold-PDT-gold molecular wires bonded to the electrodes in the same way; see Table I. These theoretical results suggest that if the...
switch from the structures with the \( \sim 42 \) meV phonon to those with the \( \sim 46 \) meV phonon were due to a change from a gauche to a trans molecular configuration, then the switch would be accompanied by an increase of the low bias conductance of the junction, contrary to what was observed experimentally.[29]

By contrast, our calculations show the low bias conductances of gold-trans-PDT-gold molecular wires in the bridge/top bonding configurations (with the \( \sim 42 \) meV phonon mode) to be higher than those for trans molecules in top bonding configurations (with the \( \sim 46 \) meV phonon mode); see Table I. Thus our interpretation of the observed transition from the \( \sim 42 \) meV phonon mode to the \( \sim 46 \) meV phonon mode being due to trans molecules switching from bridge/top bonding to pure top bonding is consistent with the experimental low bias conductance data[29].

4. Parallel Transport Through Pairs of Molecules

![Relaxed Structures](image)

**FIG. 5.** (Color online.) The relaxed geometries and calculated conductances for pairs of trans PDT molecules connecting gold clusters in parallel. 14 gold atoms per cluster. (a)-I Top site bonding. (a)-II Bridge site bonding. Calculated mode frequencies and IETS intensities are shown in (b) for the systems depicted in (a). The calculated mode frequencies and IETS intensities for single molecules similar to those in (a) connecting two gold clusters with the same kinds of molecule-gold bonding geometries are also shown for comparison. The experimental phonon mode histogram of Hihath et al. [29] for single molecules bridging the electrodes is also shown. The darker (lighter) grey represents the positive (negative) bias voltage.

In the preceding discussion we have focussed on cases where a single molecule connects the two electrodes. However in statistical STM break junction experiments, such as that of Hihath et al.[29], the electrodes can also be bridged simultaneously by two (or more) molecules in parallel. In this Section, we report theoretical results for the inelastic tunneling spectra of some examples of systems of this kind.

In Figs. 5(a) I and II, we show the relaxed geometries for pairs of top site and bridge site bonded trans PDT molecules, respectively, connecting two gold clusters. As can be seen in Table I, in each case the calculated low bias elastic conductances of these pairs of molecules are roughly twice that of a single molecule connecting the contacts and bonded to them in a similar configuration. This means that the molecules in these systems are behaving as nearly independent parallel quantum conductors in a way reminiscent of parallel semiconductor quantum wires[109] connecting a pair of electron reservoirs. Similar, roughly additive, behavior of the low bias conductances of other molecules bridging pairs of metal electrodes in parallel has also been found in previous theoretical studies, for not too small intermolecular separations [110–118].

The calculated inelastic tunneling spectra for these pairs of molecular wires are shown in Fig. 5(b). The inelastic tunneling spectra for single molecules connecting the two electrodes with similar bonding geometries are also shown for comparison. The Mode I inelastic tunneling spectra for the pairs of molecules are very similar to the corresponding spectra of the individual molecules apart from small phonon energy shifts and splittings, indicative of weak effective vibrational coupling between the molecules. However, for the pair of bridge site bonded molecules, the two Mode I features have noticeably differing calculated IETS intensities.

We also studied pairs of molecules with even smaller spatial separations than those of the molecules in Fig. 5(a). Namely, we considered pairs of top site bonded trans PDT molecules binding to nearest neighbor atoms of both gold clusters, as distinct from the second neighbor gold atoms to which the molecules bind in Fig. 5(a)-I. As expected based on previous studies [110–118], the calculated low bias conductances of these systems differed more from twice the conductance of a single molecule connecting the gold clusters than was the case for the more widely spaced molecules in Fig. 5(a). The splittings of the Mode-I phonon energies for these more closely spaced molecules were also considerably larger. However, we found that relaxed structures with such closely spaced molecules could only be generated for the smaller gold clusters that we studied; for the larger clusters DFT relaxations resulted instead in structures similar to those in Fig. 5(a). This suggests that structures with such closely spaced pairs of molecules are unlikely to be realized in experiments such as those of Hihath et al.[29].

Experimental inelastic tunneling spectra for molecules bridging the contacts in parallel were not reported in
Ref. 29. Thus comparison between these theoretical results and experiment, while of considerable interest, is not possible at present.

IV. CONCLUSIONS

We have calculated the relaxed geometries, zero bias conductances, and inelastic tunneling spectra of trans and gauche PDT molecules bonded to pairs of gold clusters in various ways and compared our results with the experimental STM break junction transport data of Hihath et al. [29] Although the gold clusters that represent the macroscopic electrodes in our calculations were not large (up to 14 gold atoms per cluster) they were sufficiently large for accurate modeling of the properties of the vibrational modes that we studied (as is discussed in Section III C 1) and our calculated inelastic tunneling spectra converged well with increasing size of the gold clusters. Thus, we expect our findings to be applicable to molecular wires bridging macroscopic gold electrodes. Consistent with this, our theoretical results agree well with the experimental data of Hihath et al. [29] We conclude that IETS when combined with such calculations is able to reveal detailed, previously inaccessible, information about the atomic scale structures of the molecule-metal interfaces and thus resolve the long standing “contact problem” of single-molecule nanoelectronics.

The success of our approach rests on the fact that ab initio density functional theory calculations of vibrational modes and their frequencies are believed to be reasonably accurate. The reason for this is that in the Born-Oppenheimer approximation they are electronic ground state total energy calculations and density functional theory has been designed and carefully optimized for calculations of the total ground state energies of interacting electron systems. [1] While the results of transport calculations for molecular nanowires are well known to be less accurate, due in part to the well known difficulty of estimating the alignment of the molecular energy levels relative to the Fermi energy of the electrodes [1], we rely on transport calculations only for qualitative information such as the identification of the phonon mode in a particular frequency range that has the largest IETS intensity. Our identification of this mode is also supported by physical reasoning.

We have definitively identified particular realizations of gold-propanedithiolate-gold molecular wires in a recent experiment [29] in which a trans PDT molecule bonded to a single gold atom of each electrode. We showed that this bonding geometry and molecular conformation was the one realized most frequently in the experiment. [29] Our theoretical results also showed that the switching from the 42 meV to the 46 meV mode observed in the experiment as the molecular wire was stretched [29] corresponds to the junction switching from trans PDT in a mixed top/bridge bonding geometry to trans PDT in a pure top bonding geometry.

Our results also showed that PDT molecular wires with gauche conformations connecting gold electrodes should have inelastic tunneling spectra that differ significantly from those with trans conformations. Comparing our results with the inelastic transport data of Hihath et al. [29], we concluded that gauche conformations were realized at most rarely in their experiment [29].

We also found that if a pair of PDT molecules connects two gold electrodes in a parallel geometry the coupling between the molecules should affect the low bias elastic and inelastic transport characteristics of the system only weakly, unless the molecules are extremely close together, for example, if they bond in a top site geometry to adjacent gold atoms of both electrodes. Structures with such closely spaced molecules were found to be unstable for the larger gold clusters in our DFT calculations.

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