Length-dependent conductance and thermopower in single-molecule junctions of dithiolated oligophenylene derivatives

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(Dated: May 5, 2008)

We study theoretically the length dependence of both conductance and thermopower in metal-molecule-metal junctions made up of dithiolated oligophenlene derivatives contacted to gold electrodes. We find that while the conductance decays exponentially with increasing molecular length, the thermopower increases linearly as suggested by recent experiments. We also analyze how these transport properties can be tuned with methyl side groups. Our results can be explained by considering the level shifts due to their electron-donating character as well as the tilt-angle dependence of conductance and thermopower. Qualitative features of the substituent effects in our density-functional calculations are explained using a tight-binding model. In addition, we observe symmetry-related even-odd transmission channel degeneracies as a function of molecular length.

PACS numbers: 85.65.+h, 65.80.+n, 73.23.Ad, 73.63.Rt

I. INTRODUCTION

In the field of molecular electronics, research has so far mostly concentrated on the dc electrical conduction properties of single-molecule contacts. By now it is known that the charge transport through organic molecules is typically due to electron tunneling. This is evidenced in particular by the exponential decay of the conductance $G$ with increasing molecular length in contacts formed from oligomers with varying numbers of units. Considering the statistical nature of experiments at the molecular scale, a conclusive comparison to theory is presently difficult. Nevertheless, associated decay coefficients appear to be reproduced by theoretical calculations. For a deeper understanding of molecular-size contacts, it is useful to analyze also other observables in parallel with the dc conductance. Emerging new lines of research involve the photoconductance and heat transport. In this paper we concentrate on the thermopower. For metallic atomic contacts, this quantity was studied experimentally already some years ago, but for molecular contacts only very recently.

The thermopower $Q$, also known as the Seebeck coefficient, measures the voltage $\Delta V$ induced over a conducting material at vanishing steady state electric current $I$, when a small temperature difference $\Delta T$ is applied: $Q = -\Delta V/\Delta T |_{I=0}$. It is known that in bulk materials the sign of the thermopower is a hint of the sign of the main charge carriers. If $Q < 0$ ($Q > 0$), charge is carried by electron-like (hole-like) quasiparticle excitations, as in an $n$-doped ($p$-doped) semiconductor. Analogously, the thermopower of the molecular junction gives information about the alignment of the energies of the highest occupied and lowest unoccupied molecular orbitals (HOMO and LUMO) with respect to the metal’s Fermi energy $E_F$. In the experiment, $Q$ was measured for gold electrodes bridged by dithiolated oligophenylene molecules. It was found to be positive, which indicates that $E_F$ lies closer to the HOMO than to the LUMO. Also, it was observed that $Q$ grows roughly linearly with the number $N$ of the phenyl rings in the molecule. More recently, the effects of substituents and varied end groups have been analyzed for the benzene molecule.

Using transport calculations based on density-functional theory (DFT) we investigate in this paper the length dependence of the conductance and the Seebeck coefficient for dithiolated oligophenylene derivatives contacted to gold contacts. For the molecules studied in Ref. 14, we find that the conductance decays exponentially with increasing molecular length. Decay coefficients compare reasonably with previous DFT calculations and also with experiments considering differences in contact configurations, molecular end groups, and uncertainties with respect to environmental effects. In addition, the thermopower increases linearly, with a magnitude coinciding with the measurements. We also study how the results change as various numbers of methyl substituents are introduced to the molecules. The effect of these substituents is twofold: (i) they push the energies of the $\pi$ electrons up as a result of their electron-donating behavior and (ii) they increase the tilt angles between the phenyl rings through steric repulsion. The latter effect tends to decrease both $G$ and $Q$ due to a reduction of the degree of $\pi$-electron delocalization, while the former opposes this tendency by bringing the HOMO closer to $E_F$. A simplified $\pi$-orbital model is used to explain essential features of the DFT results.

The paper is organized as follows. Sec. III explains details of our DFT-based approach. Then, in Sec. III we present the molecular contacts, whose charge transport properties we determine in Sec. IV. In Sec. V we show,
II. METHODS

The general formulas for the treatment of thermoelectric effects, based on the Landauer-Büttiker formalism, are discussed in detail in several references.20,21,22,23 By expanding the expression for the current \( I \) to linear order in \( \Delta V \) and \( \Delta T \) and considering the cases \( \Delta T = 0 \) and \( I = 0 \), respectively, one arrives at

\[
G = G_0 K_0(T), \quad Q = -\frac{K_1(T)}{eT K_0(T)},
\]

(1)

with \( K_0(T) = \int \tau(E) \left[ -\frac{\partial f(E, T)}{\partial E} \right] dE \), \( K_1(T) = \int (E - \mu) \tau(E) \left[ -\frac{\partial f(E, T)}{\partial E} \right] dE \), and \( G_0 = 2e^2/h \). Here \( \tau(E) \) is the transmission function, \( f(E, T) = \left( \exp\left[ \frac{E - \mu}{k_B T} \right] + 1 \right)^{-1} \) the Fermi function, and \( \mu \) the chemical potential, \( \mu \approx E_F \). At low temperature, the leading-order terms in the Sommerfeld expansions yield

\[
G = G_0 \tau(E_F), \quad Q = -\frac{k_B \pi^2}{6} \frac{\tau'(E_F)}{\tau(E_F)} k_B T,
\]

(2)

where prime denotes a derivative. In our DFT-based results presented below, we calculate \( G \) and \( Q \) according to Eq. (1). However, Eq. (2) approximates the results to within a few percent at room temperature, since \( \tau(E) \) is smooth around \( E_F \) due to the off-resonant situation. It should be noted that the equations neglect electron-vibration interactions. These could in principle be included by adding the inelastic corrections to the expression for the current, but we expect also these contributions to be relatively small even at room temperature.

The transmission functions are computed with the help of Green’s function techniques. The electronic structure is described in terms of DFT as implemented in the quantum chemistry program TURBOMOLE V5.7, where we employ the BP86 exchange-correlation functional and the standard Gaussian basis set of split valence quality with polarization functions on all non-hydrogen atoms.24 For further details on our method, see Refs. 20,27,28,29.

Figure 1: (Color online) The molecules studied in this work. When they are contacted to the gold electrodes, sulfur atoms replace the terminal hydrogen atoms.

Figure 2: (Color online) HOMO and LUMO energies of the molecules of Fig. 1.

III. CONTACTS

The molecules studied are shown in Fig. 1. Those labeled with R1 to R4 are the pure oligophenylenes, S2 to S4 denote oligophenylenes where the hydrogen atom in one of the two ortho positions with respect to each ring-connecting carbon atom is substituted with a methyl group,20,21 and D2 to D4 have substituents in both ortho positions. Here, the numbers \( N = 1, \ldots, 4 \) refer to the number of phenyl rings in the molecule. The tilt angles for the R, S, and D molecules vary between \( 33.4^\circ \leq \varphi_R \leq 36.4^\circ, \quad 84.8^\circ \leq \varphi_S \leq 90^\circ, \) and \( 88.8^\circ \leq \varphi_D \leq 90^\circ, \), and the distances between the terminal carbon atoms of the molecules are described to a good accuracy by \( d = a + bN \), with \( a = -0.154 \text{ nm} \) and \( b = 0.435 \text{ nm} \).

The HOMO and LUMO energies of the molecules are shown in Fig. 2. It can be noticed that the HOMO-LUMO gaps of the S and D series are larger than those of the R series.

To form the junctions, each molecule is coupled to the hollow position of the tips of two gold [111] pyramids via a sulfur atom. The atomic positions of the molecule and the first gold layers of the tips are then relaxed. This is depicted in Fig. 3 for S3. The relaxed part is also the “central region” in the transport calculations.20 The tilt angles \( \varphi \) and the distances \( d \) of the contacted molecules are not essentially different from those of the isolated molecules.

Figure 3: (Color online) (a) The contacts to the gold electrodes are formed through sulfur atoms bonded to the hollow position of the tip of a [111] pyramid. (b) For the transport calculations, the tips of the pyramids are part of the extended molecule (C). The more remote parts (blue) are absorbed into ideal semi-infinite left (L) and (R) surfaces (gray).
IV. DENSITY-FUNCTIONAL-BASED TRANSPORT

The results for the transmission, its logarithmic derivative, the conductance, and the thermopower for all of the R, S, and D type molecular junctions are collected in Fig. 4. In order to compare with the room-temperature experiments, we set \( T = 298 \) K in Eq. 4. The Seebeck coefficients are displayed in Fig. 4(d) together with the experimental results for R (solid line), S (dashed line), and D (dash-dot-dotted line).

![Figure 4](image-url)

Figure 4: (Color online) (a,b) Transmission function and the negative of its logarithmic derivative. (c,d) The corresponding conductance, and the thermopower for all of the R, S, and D type molecules with \( N = 1, \ldots, 4 \) rings.

Although the conductances for S and D are very similar, their thermopowers are rather different. Furthermore, the magnitudes for \( N > 2 \) follow the surprising order \( Q_S < Q_R < Q_D \). As we will discuss below, these observations can be understood through the two competing substituent effects: (i) a change in the alignment of the π-electron levels with respect to \( E_F \) as a result of the electron-donating nature of the methyl group and (ii) an increase in the ring-tilt angles. For the isolated molecules the second effect results in the opening of the HOMO-LUMO gap when going from molecules R to S or D, while the first effect causes the difference between the HOMO and LUMO energies of the S and D series (Fig. 2).
are indicated. From left to right the molecular structure is displayed together with isosurface plots of the HOMO-1, HOMO, LUMO, and LUMO+1 wave functions. Below these plots the common name of the molecules and the energies of the molecular levels are indicated.

Figure 6: (Color online) Influence of methyl substituents on the level alignment of frontier molecular orbitals in benzene. We extract the parameters of our model as follows. In order to gain some understanding of the general features of the dependence of $G$ and $Q$ on the number of substituents, we study a simple tight-binding (TB) model, which describes the $\pi$-electron system of the oligophenylenes (Fig. 5). The onsite energies $\epsilon_{j}$ ($j = 1, \ldots, N$) are equal on all carbon atoms of phenyl ring $j$, the intra-ring hopping $t$ is assumed to be the same everywhere, and the inter-ring hopping $u$ is parametrized through $u = t \cos \varphi$. We assume the effect of the side groups to come into play only through $\varphi$ and $\epsilon_{j}$. The leads are modeled by "wide-band" self-energies $\Sigma_{L,R} = -i\Gamma$, acting on the terminal carbon atoms.

We extract the parameters of our model as follows. For the R molecules, we set $\epsilon_{j} = 0$. In the S and D molecules, the $\epsilon_{j}$’s for rings with one, two, or four methyl groups are obtained by performing DFT calculations of methylbenzene, dimethylbenzene, and tetramethylbenzene (Fig. 6). We find that the HOMO and LUMO energies in these molecules shift upwards monotonously with the number of methyl substituents. This can be attributed to the electron-donating character of the methyl groups, and the resulting increase of Coulomb repulsion on the phenyl ring. We use the averages of the HOMO and LUMO shifts relative to benzene, and obtain, respectively, $\epsilon^{(one)} = 0.17$, $\epsilon^{(two)} = 0.33$, and $\epsilon^{(four)} = 0.58$ eV. The onsite energies of the different phenyl rings of the R, S, and D molecules with $N = 1, \ldots, 4$ are schematically represented in the lower part of Fig. 5. The hopping $t$ is set to half of (the negative of) the HOMO-LUMO gap of benzene, with the result $t = -2.57$ eV (Fig. 5). The scattering rate $\Gamma$ and the tilt angles for the R, S, and D molecules are chosen to reproduce approximately the minimal transmissions for R1, R2, S2, and D2 in the DFT results of Fig. 5. This yields $\Gamma = 0.64$ eV, $\varphi_{R} = 40.0^\circ$, $\varphi_{S} = 85.5^\circ$, and $\varphi_{D} = 86.5^\circ$. The last free parameter is $E_{F}$, which should be determined by the overall charge-transfer effects between the molecule and the electrodes. We choose its value of $E_{F} = -1.08$ eV close to the crossing points of the transmission curves of the S and D molecules [cf. Figs. 4(a) and 7(a)].

Similar features can be recognized between the DFT results and our model. In particular, as shown in Fig. 7(b), the correct order of the thermopowers is reproduced. In going from R to S, the increase in the tilt angle and the associated breaking of the $\pi$-conjugation dominates over other side group effects. As a result both $G$ [Fig. 7(c)] and $Q$ [Fig. 7(d)] decrease. In going from S to D, an interplay of the sidegroup-induced level shifts and a small residual increase in $\varphi$ raises $Q$ above the value for R, while $G$ remains almost unchanged. We note that for an $N$-ring junction the lowest-order terms in an expansion of the transmission in powers of $u$ yield $\varphi$-dependences of the form $\tau(E) \approx c_{1}(E) \cos^{2(N-1)} \varphi + c_{2}(E) \cos^{2N} \varphi$, for $E \approx E_{F}$. Independently of $N$, this results in $Q \approx q_{1} + q_{2} \cos^{2} \varphi$, where we find $q_{1}, q_{2} > 0$ in our case. For $N = 2$, the neglect of the $c_{2}$-term leads to the well-known "cos$^{2} \varphi$ law" of $\tau$, but we see that for the tilt-angle depen-
dence of $Q$ the presence of this term is significant, since $q_2 = 0$ if $c_2 = 0$. When $E_F$ is close to a resonance, further higher-order terms become increasingly important and deviations from the $q_1 + q_2 \cos^2 \varphi$ law result. This is seen most clearly as the non-monotonous $\varphi$-dependence of $Q$ for molecule D3 in Fig. 7(d).

VI. EFFECTS BEYOND THE $\pi$-ORBITAL MODEL

Close to perpendicular ring tilts results from the $\pi$-orbital model should be taken with care. At $\varphi = 90^\circ$ the $\pi-\pi$ coupling $u$ vanishes, and any other couplings between the rings will become important. Let us analyze this for the biphenyl molecules R2, S2, and D2. For them, we have varied the tilt angle between the rings, and have determined the charge transport properties for every $\varphi$ using our DFT-based approach. Further details of the procedure are described in Ref. 29. For R2 (and similarly for S2 and D2) we observe that for most tilt angles ($\varphi \lesssim 80^\circ$) the transmission is dominated by a single channel [Fig. 8(a)] of $\pi-\pi$ character. However, at large angles two transmission channels of the same magnitude are observed, which become degenerate at $\varphi = 90^\circ$. They arise from $\sigma-\pi$ couplings between the two rings, whose strengths are proportional to $\sin \varphi$. Thus the two degenerate channels are of the $\sigma-\pi$ and $\pi-\sigma$ type. These features are obviously not accounted for by the TB model, where only a single $\pi-\pi$ channel is present independently of $\varphi$. The $\sigma-\pi$ couplings should also modify the tilt-angle dependence of the thermopower, which is plotted in Fig. 8(b). Due to the electron-donating nature of the methyl groups the thermopower at fixed $\varphi$ increases from R2 to S2 and D2. While the curve for R2 can be described by the law $q_1 + q_2 \cos^2 \varphi$ for $\varphi \lesssim 80^\circ$, a dip is observed for larger $\varphi$. Similar deviations in $Q$ are also present for S2 and D2, where we have investigated a smaller tilt-angle interval because of the steric repulsion of the methyl groups. 28

VII. DISCUSSION AND CONCLUSIONS

As is well known, there are many theoretical uncertainties involved in the determination of transport properties based on DFT calculations, and improvements for the methods of molecular-scale transport theory are currently being sought. Indeed, it is typical to find order-of-magnitude differences between measured conductances and those computed within DFT. It is also known that atomic configurations can have a strong influence on the conductance and hence a conclusive comparison with experimental data would require the statistical analysis of a large number of contact geometries. However, models predict $Q$ to be insensitive to changes in the lead couplings and hence it is expected to be a more robust quantity that $G$. Although the measurements of Ref. 14 were carried out at room temperature, we do not expect molecular vibrations to play an essential role in the results due to the weakness of the electron-vibration coupling. Thus, the comparison we have made with our elastic transport theory seems justified. Nevertheless, the investigation of the effect of molecular vibrations is an interesting direction for future research, also in view of optimizing the properties of molecular thermoelectric devices for potential applications. As far as gaining a basic understanding is concerned, low-temperature measurements would be desirable to remove uncertainties related to the thermal excitation of vibrations. The purpose of the present work was to study the general trends for a series of molecules that
are similarly coupled to the electrodes, and considering all the potential uncertainties, the agreement we obtain for the thermopower appears to be quite reasonable.

In conclusion, we have analyzed the length dependence of conductance and thermopower for oligophenylene single-molecule contacts. While we found the conductance to decay exponentially with the length of the molecule, we observe that the thermopower increases linearly. For possible future applications it is interesting to know, how the magnitudes of these quantities can be tuned. We analyzed how this can be achieved by chemical substituents, which control the Fermi-level alignment and the molecular conformation. We demonstrated that a simple π-electron tight-binding model can help to understand basic substituent effects. In addition, we observed an even-odd effect for transmission channel degeneracies upon variation of the number of phenyl rings, which we explained by molecular symmetries.

VIII. ACKNOWLEDGMENTS

We acknowledge stimulating discussions with M. Mayor. The Quantum Chemistry group of R. Ahlrichs is thanked for providing us with TURBOMOLE. This work was financially supported by the Helmholtz Gemeinschaft (Contract No. VH-NG-029), the EU network BIMORE (Grant No. MRTN-CT-2006-035859), and DFG SPP 1243. F. P. acknowledges the funding of a Young Investigator Group at KIT.
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