Low Tunneling Decay of Iodine-Terminated Alkane Single-Molecule Junctions

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Abstract
One key issue for the development of molecular electronic devices is to understand the electron transport of single-molecule junctions. In this work, we explore the electron transport of iodine-terminated alkane single molecular junctions using the scanning tunneling microscope-based break junction approach. The result shows that the conductance decreases exponentially with the increase of molecular length with a decay constant $\beta_N = 0.5$ per $-\text{CH}_2$ (or $4$ nm$^{-1}$). Importantly, the tunneling decay of those molecular junctions is much lower than that of alkane molecules with thiol, amine, and carboxylic acid as the anchoring groups and even comparable to that of the conjugated oligophenyl molecules. The low tunneling decay is attributed to the small barrier height between iodine-terminated alkane molecule and Au, which is well supported by DFT calculations. The work suggests that the tunneling decay can be effectively tuned by the anchoring group, which may guide the manufacturing of molecular wires.

Keywords: Electron transport, Barrier height, Single molecular junction, Iodine, Alkyl-based molecules

Background
Understanding the electron transport of single-molecule junctions is crucial for the development of molecular electronic devices [1–16]. The non-resonant tunneling model has often been used to describe the electron transport process through small molecule, where contact conductance, molecular length, and the tunneling decay constant are the main parameters [17, 18]. In most molecular systems, decay constant is highly related to the electronic properties of organic backbone. For example, the conjugated molecular systems have low tunneling decay, unlike non-conjugated ones [17, 19]. Since the tunneling decay is decided by the barrier height between the Fermi level of electrode and lowest unoccupied molecular orbital (LUMO) or highest occupied molecular orbital (HOMO) of molecular junctions [17, 20], it is possible to tune the molecular energy level towards the Fermi level to achieve the low decay [21–24].

In single-molecule junctions, the anchoring group plays an important role in the control of electronic coupling between the organic backbones with the electrodes [21, 23–25]. A series of conductance measurements for the alkane-based molecules have showed a significant effect of different anchoring groups on the binding geometry, junction formation probabilities, contact conductance, and even conductance channel (through LUMO or HOMO) of molecular junctions [21–25]. Since the anchoring group can regulate the frontier orbitals in the molecular junction, the tunneling decay of the molecule may also be tuned by the anchoring group [24]. However, limited study has been focused on this area.

Herein, we report the electron transport of alkane molecules terminated with iodine group by using scanning tunneling microscopy break junction (STM-BJ) (Fig. 1) [26, 27]. The single molecular conductance measurements show that the conductance decreases exponentially with the increase of molecular lengths and the decay constant of alkane molecules with iodine group is much lower than that of the analogues with other anchoring groups. The different
tunneling decay constants for alkane molecules with varied anchoring groups are explained by barrier height between molecule and electrode.

**Methods**

1,4-Butanediiodo, 1,5-pentanediiodo, and 1,6-hexanediiodo were purchased from Alfa Aesar. All solutions were prepared with ethanol. Au(111) was used as the substrate, while mechanically cut Au tips were used as the tips. Before each experiment, the Au(111) was electrochemically polished and carefully annealed in a butane flame and then dried with nitrogen.

The Au(111) substrate was immersed into a freshly prepared ethanol solution containing 0.1 mM target molecules for 10 min. The conductance measurement was carried out on the modified Nanoscope IIIa STM (Veeco, USA.) by using the STM-BJ method at room temperature [28–30], which simply measured the conductance of single-molecule junctions formed by repeatedly moving the tip into and out of the substrate at a constant speed. During the process, the molecules could anchor between the two metal electrodes and form single molecular junctions. Thousands of such curves were collected for statistical analysis. All the experiments were performed with a fix bias voltage of 100 mV. Since molecules with iodine as the anchoring group are a photosensitive material, the experiment was carried out under shading.

**Results and Discussion**

**Conductance Measurement of Iodine-Terminated Alkane Single Molecular Junctions**

The conductance measurements were first carried out on Au(111) with monolayer of 1,4-butanediiodo by STM-BJ. Figure 2a gives out the typical conductance traces exhibiting the stepwise feature. Conductance traces show plateau at $1 G_0$, indicating the formation of stable Au atomic contact. Plateau at a conductance value of $10^{-3.6} G_0$ (19.47 ns) is also found besides the $1 G_0$, owing to the formation of molecular junction. A conductance histogram could also be obtained by treating with logarithm and binning of conductance value from more than 3000 conductance traces, and then, the intensity of conductance histogram was normalized by the number of traces used and shows a conductance peak at $10^{-3.6} G_0$ (19.44 ns) (Fig. 2b). Those show that the iodine group can serve as an effective anchoring group forming molecular junction. However, this value is smaller than the single molecular conductance value of 1,4-butanediamine with amine as the anchoring group, which may stem from weak interaction between iodine and Au electrode [31].

In comparison with 1,4-diiodobutane, pronounced peaks at $10^{-3.8} G_0$ (12.28 ns) and $10^{-4.0} G_0$ (7.75 ns) are found for 1,5-pentanediiodo and 1,6-hexanediiodo, respectively (Fig. 3). The conductance values decrease with the increasing of molecule length. Meanwhile, the...
conductance values of 1,5-pentanediiodo and 1,6-
hexanediiodo are smaller than those of 1,5-
pentanediiodo and 1,6-hexanediiodo, respectively
[31], which may be caused by the different interaction in
alkane-based molecular junctions between iodine and
amine anchoring groups binding to Au electrodes [32].

The two-dimensional conductance histograms were also
constructed for those molecular junctions (Additional file 1:
Figure S1) and give out similar conductance values of
one-dimensional histograms. Typically, the breaking off
distance of molecular junctions increases with the increas-
ing of molecular length. We also analyze the distance from
the conductance value of $10^{-5.0} G_0$ to $10^{-0.3} G_0$ as shown
in Fig. 4, and rupture distances of 0.1, 0.2, and 0.3 nm are
found for 1,4-butanediiodo, 1,5-pentanediiodo, and 1,6-
hexanediiodo, respectively. Here, the rupture distances are
obtained from the maximum peak of the rupture distance
histogram [33]. It was reported that there is a snap back
distance of 0.5 nm for Au after the breaking of Au–Au
contact [34, 35]; thus, the absolute distances for those
molecular junctions between electrodes could be 0.6, 0.7,
and 0.8 nm which are found for 1,4-butanediiodo, 1,5-
pentanediiodo, and 1,6-hexanediiodo, respectively. Those
distances are comparable to the length of molecules.

Eder et al. reported that the adsorption of 1,3,5-tri
(4-iodophenyl)-benzene monolayer onto Au(111) may
cause partial dehalogenation [36]; however, a very
larger conductance value for those Au–C covalent
contact molecular junctions can be found for
molecules with four (around $10^{-1} G_0$) and six (bigger
than $10^{-2} G_0$) –CH2– units [37]. Thus, we propose
that the current investigated molecules contact to the
Au through the Au–I contact.

**Tunneling Decay Constant of Iodine-Terminated Alkane
Single Molecular Junctions**

Under the current bias, those molecule conductance can
be expressed as $G = G_c \exp(-\beta_N N)$. Here, $G$ is the
conductance of the molecule and $G_c$ is the contact con-
ductance and is determined by the interaction between
the anchoring group and the electrode. $N$ is the methy-
lene number in the molecule, and $\beta_N$ is the tunneling
decay constant, which reflects the coupling efficiency of
electron transport between the molecule and the elec-
trode. As show in Fig. 5, we plot a natural logarithm
scale of conductance against the number of methylene;
tunneling decay constant $\beta_N$ of 0.5 per $–\text{CH}_2$ is determined from the slope of linear fitting. This tunneling decay is very low in alkane-based molecules. For the alkane-based molecules, $\beta_N$ is usually found around 1.0 per $–\text{CH}_2$ for thiol (SH) [23, 38], while around 0.9 and 0.8 per $–\text{CH}_2$ are determined for amine (NH$_2$) [23, 31] and carboxylic acid (COOH), respectively [39]. Thus, the tunneling decay with iodine shows the lowest value among those anchoring groups with a trend $\beta_N$ (thiol) $>$ $\beta_N$ (amine) $>$ $\beta_N$ (carboxylic acid) $>$ $\beta_N$ (iodine), which may be due to the difference in the alignment of molecular energy levels to the Fermi level of Au electrode [23, 31]. The tunneling decay of 0.5 per $–\text{CH}_2$ can also be converted to 4 nm$^{-1}$, which is comparable to oligophenyls with 3.5$–$5 nm$^{-1}$ [40, 41].

The $\beta_N$ for the metal-molecule-metal junctions can be simply described by the below equation [17, 20, 38],

$$\beta_N \propto \sqrt{\frac{2m\Phi}{h^2}}$$

where $m$ is the effective electron mass and $\hbar$ is the reduced Planck’s constant. $\Phi$ represents the barrier height, which is decided by the energy gap between the Fermi level and the molecular energy levels in the junction. Obviously, the $\beta_N$ value is proportional to the square root of barrier height. Thus, we may propose that iodine-terminated alkane molecules have small $\Phi$ with the Au electrode.

**Barrier Height of Single Molecular Junctions with Different Anchoring Groups**

Taking the $–(\text{CH}_2)_n–$ as the backbone, we performed the rough calculations (see computational detail in Additional file 1) to investigate the frontier molecular orbitals of complexes with four Au atoms at the both ends, including 1,6-hexanedithiol (C6DT), 1,6-hexanediamineb (C6DA), 1,6-hexanedicarboxylic acid (C6DC), and 1,6-hexanediiodo (C6DI). As shown in Table 1, the HOMO and LUMO are $–$6.18 and $–$1.99 eV, respectively, for C6DT, while HOMO (6.02 eV) and LUMO ($–$1.85 eV) are found for C6DA. Meanwhile, HOMO and LUMO energy levels are calculated for C6DC ($–$6.33 and $–$2.58 eV) and C6DI ($–$2.22 and $–$2.61 eV).

For the Fermi level of Au electrode, we need to consider the influence of the adsorption of molecules. In the vacuum condition, clean Au gives out work function of 5.1 eV [42]; meanwhile, this value can be obviously changed by the adsorption of molecules. Kim et al. [43] and Yuan et al. [44] have found that the work function of Au is around 4.2 eV (4.0$–$4.4 eV) upon the adsorbed self-assembled monolayers (SAMs) measured by the ultraviolet photoelectron spectrometer (UPS). Low et al. also investigated the electron transport of thiophene-based molecules of TOTOT (LUMO $–$3.3 eV, HOMO $–$5.2 eV) and TTO$_p$TT (LUMO $–$3.6 eV, HOMO $–$5.1 eV) with Au as the electrode (T, O, and $O_p$ denote thiophene, thiophene-1,1-dioxide, and oxidized thienopyrrolodione, respectively) [45]. The results show that the Fermi level of Au is in the middle of LUMO and HOMO. Thus, we can infer the Fermi level of Au can be around the average energy level of LUMO and HOMO, which are $–$4.25 and $–$4.35 eV established from TOTOT and TTO$_p$TT, respectively. The Fermi level of Au $–$4.25 and $–$4.35 eV are similar to that measured by UPS with $–$4.2 eV [43]. According to the above, we will use the $–$4.2 eV as the Fermi level of Au electrode with the adsorption of molecule.

Assuming the Fermi level of $–$4.2 eV for Au with SAM, C6DT and C6DA are the HOMO-dominated electron transport, while LUMO-dominated electron transport is proposed for the C6DC and C6DI. Thus, the barrier height $\Phi$ can be established as 1.98 eV (C6DT), 1.82 eV (C6DA), 1.62 eV (C6DC), and 1.59 eV (C6DI) (Table 1).

The trend for the barrier height between the molecule and Au is $\Phi_{\text{C6DT}}$ (thiol) $>$ $\Phi_{\text{C6DA}}$ (amine) $>$ $\Phi_{\text{C6DC}}$ (carboxylic acid) $>$ $\Phi_{\text{C6DI}}$ (iodine), which is consistent with the trend

| Table 1 Energy levels of the frontier orbitals of molecules contacting with four Au atoms computed by DFT method |
|---------------------------------------------------------------|
| $E_{\text{HOMO}}$ | $E_{\text{LUMO}}$ | $E_{\text{HOMO}}$ | $E_{\text{LUMO}}$ |
| Au$_4$-C6DT-Au$_4$ (eV) | 1.99 | 2.21 |
| Au$_4$-C6DA-Au$_4$ (eV) | 1.85 | 2.35 |
| Au$_4$-C6DC-Au$_4$ (eV) | 2.58 | 1.62 |
| Au$_4$-C6DI-Au$_4$ (eV) | 2.61 | 1.59 |
of the tunneling decay ($\beta$). Thus, the unusual low tunneling decay can be contributed to the small barrier height between iodine-terminated alkane molecules and Au.

**Conclusions**

In conclusion, we have measured the conductance of alkane-based molecules with iodine group contacting to Au electrodes by STM-BJ at room temperature. A tunneling decay $\beta_N$ of 0.5 per $-\text{CH}_2$ was found for those molecules with Au electrodes, which is much lower than that of alkane-based molecules with other anchoring groups. This can be caused by the small barrier height between the iodine-terminated alkane molecule and Au. The current work shows the important role of the anchoring group in electrical characteristics of single molecular junctions, which can tune the tunneling decay of molecular junction and guide the manufacturing molecular wire.

**Additional file**

Additional file 1: Two-dimensional conductance histograms of molecular junctions and computational details. (DOCX 173 kb)

**Abbreviations**

HOMO: Highest occupied molecular orbital; LUMO: Lowest unoccupied molecular orbital; SAMs: Self-assembled monolayers; STM-BJ: Scanning tunneling microscopy break junction; UPS: Ultraviolet photoelectron spectroscopy

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**Availability of Data and Materials**

The datasets supporting the conclusions of this article are included within the article and its additional file.

**Authors’ Contributions**

LLP, BH, ZWH, and JFZ carried out the experiments; QZ, YS, and ZJN contributed to the analyzed the results. HJX performed the calculations. LLP, XSZ, and WB conceived and designed the experiments and analyzed the results. HJX performed the calculations. LLP, BH, ZWH, and JFZ carried out the experiments; QZ, YS, and ZJN contributed to the analyzed the results. HJX performed the calculations. LLP, BH, ZWH, and JFZ carried out the experiments; QZ, YS, and ZJN contributed to the analyzed the results. HJX performed the calculations. LLP, BH, ZWH, and JFZ carried out the experiments; QZ, YS, and ZJN contributed to the analyzed the results. HJX performed the calculations.

**Competing Interests**

The authors declare that they have no competing interests.

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