Room temperature negative differential resistance of a monolayer molecular rotor device

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An electrically driven molecular rotor device comprised of a monolayer of redox-active ligated copper compounds sandwiched between a gold electrode and a highly doped $P^+$Si substrate was fabricated. Current-voltage spectroscopy revealed a temperature-dependent negative differential resistance (NDR) associated with the device. Time-dependent density functional theory suggests the source of the observed NDR to be redox-induced ligand rotation around the copper metal, an explanation consistent with the proposed energy diagram of the device. An observed temperature dependence of the NDR behavior further supports this hypothesis. © 2009 American Institute of Physics. [DOI: 10.1063/1.3222861]

Negative differential resistance (NDR) is an essential property that allows fast switching in certain types of electronic devices such as Esaki diode and IMPATT diode.1,2 Beyond the traditional electronic devices, a number of nanoscale molecular devices have been studied and reported to exhibit NDR characteristics.3–7 Although various mechanisms have been proposed, including charging and discharging processes of electrons, chemical reaction, redox reaction, and the association-dissociation processes of molecules, their specific underlying physics remain unclear. These functional molecular units acting as state variables provide an attractive alternative to overcome the limits of conventional metal-oxide-semiconductor field-effect transistor technology due to their potential scalability, low cost, low variability, highly integratable characteristics, and the capability to exploit self-assembly processes.8–10 Thus the elucidation of switching mechanisms and the development of different operational approaches have drawn a lot of attention. In this letter, we demonstrate an electrically driven sandwich-type monolayer molecular rotor switch with NDR. The observed NDR behavior is attributed to rotational motion on solid support. Both calculations of time-dependent density functional theory and an observed temperature dependence of the NDR behavior support this hypothesis.

Synthesized using a self-assembly approach, the molecular switch device shown in Fig. 1 is comprised of a heteroleptic copper compound covalently bonded to a highly doped silicon substrate. Each complex contains three subunits: a bifunctional stator [a bidentate ligand bonded to both a solid support ($P^+$Si) and a Cu axle], the metal axle (Cu), and a diimine rigid rotator (2,9-dimethyl-1,10-phenanthroline). Preparation of the heteroleptic copper system was carried out through covalent grafting of a stator monolayer onto the hydroxylated surface of a $P^+$Si substrate using silanol bonds.11

The stators were then used to chelate a copper metal axle that subsequently bonds to the rotator subunit. The device was completed by deposition of a Ti/gold film through a shadow mask on top of the molecular layer to form the top electrode. The number of molecules per unit area and per unit devices tested is approximately $1.1 \times 10^{14}$ cm$^{-2}$ and $8 \times 10^{10}$, respectively. This copper compound exhibits two discrete, redox-dependent conformational states, Cu(I) and Cu(II). The Cu(I) form has tetrahedral geometry while the Cu(II) form is square planar.12 The compounds undergo a one electron redox-induced rotational conformational change depending on the oxidation state of the copper metal. Interconversion between these two states provides the basis for a controlled, bistable nanoswitch.

$I-V$ characteristics of the monolayer device are shown in Fig. 2. In this case, the silicon substrate represents the system ground and the dc bias applies from the top gold electrode. The bias sweep direction followed the arrows numerically as shown in Fig. 2. Unlike previously reported devices, the NDR observed here disappeared after reversal of the sweep.
ing direction in the negative voltage regime (arrow 7). At room temperature, the peak current density is approximately 0.1 A/cm² with a corresponding peak-to-valley ratio near 5. For a given device, the I-V characteristics are stable and reproducible with consecutive positive and negative bias sweeps. The magnitude of observed current fluctuations is smaller than 1% and errors are within 0.5%. Small fluctuations are observed among devices owing to the nonuniformity of the active molecular layer.

To further explore the correlation of the observed NDR effect with redox-induced conformational switching, devices whose molecular layer consisted of solely grafted stators or Cu ligated stators were prepared. Due to their lack of a ligated rigid rotator, these devices lack the capability for rotational motion upon oxidation or reduction of the Cu metal axle. The lack of observed NDR effects in these cases is due to the conformational change in the active molecular layer. As seen in Fig. 2, state “1” and state “2” represent the low conductivity state, while state “2” is determined by the redox energy of the copper system. The difference in value of the turning voltage between positive and negative ranges is due to the different contact energy barriers of the two electrodes. The arrows show the sequence of the voltage scan. Arrows “4” and “5” correspond to the band diagrams in Figs. 4(a) and 4(b), respectively (to be discussed later).

To gain a thorough understanding of electron transport during this rotation behavior, temperature dependent I-V measurements of the device were performed from 77 K to room temperature. The NDR effect disappeared at about <244 K. The peak minus background currents at multiple temperatures were extracted and plotted as shown in Fig. 3. Peak currents measured at $T$ > 244 K followed an exponential relationship with respect to the reciprocal of the temperature. Based on the well-known Arrhenius equation, we have

$$E_a = -R \left[ \frac{\partial \ln k}{\partial (1/T)} \right]_P,$$

where $E_a$ is the activation energy, $k$ is the rate constant of chemical reactions, $T$ is the temperature, and $R$ is the gas constant. Curve fitting of the data acquired for $T$ > 244 K yielded a rotational activation energy of approximately 0.3 eV, and this value is consistent with the theoretical quenching energy supports the proposal that the observed NDR effect in this device is due to rotational motion within the molecular layer. The peak and background currents at 0.5 and −0.5 V, respectively, were plotted versus 1000/$T$ and displayed different temperature dependences, as shown in the inset of Fig. 3. The weak temperature dependence of both background currents is indicative of electron tunneling between the two device electrodes.

The proposed mechanism for the observed device I-V characteristics shown in Fig. 2 is based on an electron tunneling induced molecular rotation behavior that serves to modify the band diagram of the active molecular layer. Density Functional Theory (DFT) calculations have been employed in an attempt to reconcile the sequence of carrier transport processes and the role of energy states, which helped formulate the proposed band diagram presented in Fig. 4. Application of a bias potential to the top gold electrode shifts the energy positions of molecular states relative to both gold electrode and $P$Si band edges. As the rotary motion around the molecular axle is controlled by electron transfer, a portion of the applied potential is used to initiate the redox process within the molecular layer. First, theoretical calculations of the highest occupied molecular orbital (HOMO)-lowest unoccupied molecular orbital (LUMO) band gap for the Cu(I) and Cu(II) complexes by time-dependent DFT (TD-DFT) yielded values of approximately 2.7 and 3.4 eV, respectively. Similar methods have been used in the study of organocopper complexes. Additional calculations of the molecular redox energy produced a value of
1.05 eV, which is close to 0.7 eV of the difference of band gaps between Cu(I) and Cu(II). The 0.35 eV difference between these two values may be attributed to the rotational energy contribution. Finally, the electron affinity of the Cu(I) complex is estimated to be 3.7 eV, which anchors the LUMO energy state with respect to the gold Fermi level.

The process begins with the low conductivity state of the Cu(II) complex under positive bias (arrow 1 in Fig. 2). The device remains in the low conductivity state until one electron from the P\( ^\circ \)Si substrate tunnels through the energy barrier and charges into the HOMO state of the active molecular layer. The resulting reduction of Cu(II) to Cu(I) shifts the relative position of the HOMO energy states upward and serves to shrink the band gap, resulting in a high current (arrow 2 in Fig. 2). Upon application of negative bias to the gold electrode, the conducting channel of the Cu(I) complex remains open and results in current flow through both HOMO and LUMO states shown as Fig. 4(a) (arrow 4 in Fig. 2). Subsequent oxidation of Cu(I) to Cu(II) occurs when one electron tunnels out of the HOMO state and discharges the active molecular layer. This process alters the band structure and increases the band gap due to a reduction in Coulomb energy by the removal of one electron shown as Fig. 4(b) (arrow 5 in Fig. 2). Current flow then stops, resulting in NDR. The return trace does not show an NDR behavior as the conduction path is closed. Continuing toward more negative bias produced an exponential increase in measured current with respect to the applied electric field, which is attributed to direct tunneling between the two electrodes. Since the model used in the calculations was based on the gas phase, discrepancies between the calculated and experimental values could be the result of deviations from the theoretical model due to adsorption of the molecules onto a solid support. There are other possible transport mechanisms such as Redox-based filament, hopping transport, which one might consider. For both of these two transport mechanisms, the current would show different temperature dependences. However for our device, the weak temperature dependence of the background current implies the tunneling transport. Furthermore, the thickness of the molecular layer in our device is only <5 nm, and thus the filament formation is not likely. In addition, the control samples with Zn as a metal axel, which cannot be rotated with the redox process, did not show any NDR effects. The above arguments further disqualify the filament transport as the conduction mechanism in our molecular rotor devices. Thus we may conclude that our proposed Redox-assisted tunneling transport is the most likely model to explain the observed phenomena. Clearly further experiments are needed in order to gain deeper understanding of the underlying mechanism for this molecular device.

In summary, an electrically driven molecular rotor device has been designed and fabricated. The NDR behavior was observed and attributed to the rotational motion about the copper metal axle. From temperature dependent measurements, the activation energy of the rotation was estimated to be 0.3 eV from the Arrhenius plot. The proposed band diagram was used to explain the electron transport behavior during the device operation.

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**References:**

15. All calculations were performed on model system [Cu(2,9-dimethyl-1,10-phenanthroline)] \( \text{II} \) in the gas phase with Gaussian 03 using the density functional B3LYP with the 6-31G(d) basis set. (Frisch, M. J. Gaussian 03, revision C.02; Gaussian, Inc.: Wallingford, CT, 2004.) The LANL2DZ basis set was used for Cu and the 6-31G(d,p) basis set for C, N, and H.