

Effect of Ring Strain on the Charge Transport of a Robust Norbornadiene-Quadricyclane-Based Molecular Photoswitch

Downloaded from: https://research.chalmers.se, 2025-12-04 20:03 UTC

Citation for the original published paper (version of record):

Tebikachew, B., Li, H., Pirrotta, A. et al (2017). Effect of Ring Strain on the Charge Transport of a Robust Norbornadiene-Quadricyclane-Based Molecular Photoswitch. Journal of Physical Chemistry C, 121(13): 7094-7100. http://dx.doi.org/10.1021/acs.jpcc.7b00319

N.B. When citing this work, cite the original published paper.

research.chalmers.se offers the possibility of retrieving research publications produced at Chalmers University of Technology. It covers all kind of research output: articles, dissertations, conference papers, reports etc. since 2004. research.chalmers.se is administrated and maintained by Chalmers Library



THE JOURNAL OF PHYSICAL CHEMISTRY

Article

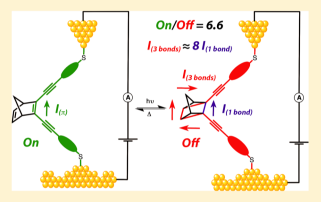
pubs.acs.org/JPCC

Effect of Ring Strain on the Charge Transport of a Robust Norbornadiene-Quadricyclane-Based Molecular Photoswitch

Behabitu E. Tebikachew,^{†,⊥} Haipeng B. Li,^{‡,⊥} Alessandro Pirrotta,^{§,} Karl Börjesson, Gemma C. Solomon,^{*,§,} Joshua Hihath,^{*,‡} and Kasper Moth-Poulsen^{*,†,}

Supporting Information

ABSTRACT: Integrating functional molecules into single-molecule devices is a key step toward the realization of future computing machines based on the smallest possible components. In this context, photoswitching molecules that can make a transition between high and low conductivity in response to light are attractive candidates. Here we present the synthesis and conductance properties of a new type of robust molecular photothermal switch based on the norbornadiene (NB)quadricyclane (QC) system. The transport through the molecule in the ON state is dominated by a pathway through the π conjugated system, which is no longer available when the system is switched to the OFF state. Interestingly, in the OFF state we find that the same pathway contributes only 12% to the transport properties. We attribute this observation to the strained tetrahedral



geometry of the QC. These results challenge the prevailing assumption that current will simply flow through the shortest through-bond path in a molecule.

INTRODUCTION

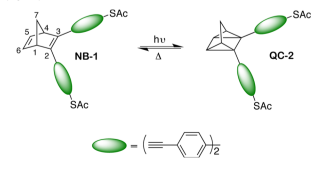
Integrating functional molecules into single-molecule devices is a key step toward the realization of future computing machines based on the smallest possible component, single molecules. 1-4 In this context, molecular switches that can make a transition between high and low conductivity in response to external stimuli have attracted significant attention since they can add a unique electronic function to molecular-scale devices and at the same time provide insight into the charge transport mechanism at the nanoscale.5-10

Photoswitching molecules, molecules that undergoes a welldefined structural isomerization upon photoexcitation, are appealing candidates for single-molecule electronics.^{3,11} There are several examples of photoswitching molecules. From a charge transport perspective diarylethenes 12-15 and azobenzenes^{5,16,17} have been the two major molecular platforms rigorously studied through various experimental approaches in the past decade. Recently, Jia et al. demonstrated a robust diarylethene-based device with remarkable stability and bidirectional in situ switching with graphene electrodes. 15 Other photoswitchable molecules with more complicated structures have also been investigated. 7,18,19 Here, we designed and synthesized a norbornadiene-based photoswitch (NB-1) with

embedded oligo(phenylene ethynylene) arms end-capped with thioacetate groups to anchor it to gold electrodes (Scheme 1).

Norbornadiene is a bicyclic hydrocarbon that undergoes photoisomerization to the metastable quadricyclane form upon light stimulation. The quadricyclane form can relax back to

Scheme 1. Photochemical Isomerization of the Norbornadiene Form (NB-1) to the Quadricyclane Form (QC-2) and the Thermal Relaxation



January 11, 2017 Received: Revised: March 1, 2017 Published: March 6, 2017

Department of Chemistry and Chemical Engineering, Chalmers University of Technology, 41296 Gothenburg, Sweden

[‡]Department of Electrical and Computer Engineering, University of California Davis, Davis, California 95616, United States

[§]Nano-Science Center and Department of Chemistry, University of Copenhagen, 2100, Copenhagen Ø, Denmark

Department of Chemistry and Molecular Biology, University of Gothenburg, 41296 Gothenburg, Sweden

norbornadiene by thermal activation. Hence, it is one of the few examples of a T-type negative photochrome along with dimethyldihydropyrene. 18 The photoswitching properties of norbornadiene have been demonstrated and utilized to study intramolecular energy transfer in solution. In 1994, Bonfantini et al. proposed norbornadiene as a potential candidate system for molecular electronics.²⁰ The prospect of using norbornadiene in electronic applications has also been discussed by Löfås et al.²¹ Yet, to the best of our knowledge, there are no examples of experimental conductance studies regarding the evaluation and use of the norbornadiene system as a photoswitch in molecular electronics. Hence, NB-1 is characterized in solution, revealing robust photoswtiching properties, and its transport properties are investigated in STM-break junction measurements. Furthermore, the transport properties of the system are analyzed in depth using DFT modeling and local current analysis.

Electron transport through NB-1 is dominated by the π system. However, upon photoexcitation the central unit undergoes an intramolecular [2 + 2] cycloaddition, rehybridizing the orbitals from sp² to sp³, to give the photoisomer, QC-2 (Scheme 1).²² In this state, the conductance must be dominated by a pathway involving the σ system, as no fully conjugated path remains. As a result, our measurements yield a larger on/off ratio than expected from breaking a single π -bond in the transport pathway. The DFT calculations show that the conductance in QC-2 is characterized by a path through the longer cyclobutane three σ -bonds over the shorter single σ bond path, and the resulting switching ratio is in good agreement with the experimental results. This work brings more insight into the effect of quantum interference on charge transport in strained ring structures and suggests new candidates and perspectives on engineering molecular-level photoswitches.

■ RESULTS AND DISCUSSION

The photoswitching molecule (NB-1) was synthesized through Sonogashira²³ cross-coupling reaction starting from 2,3dibromonorbornadiene. Full synthetic details and characterization are presented in the SI. The photoisomerization process was examined in solution using NMR and UV-vis spectroscopy. The proton NMR for the photoisomerization of NB-1 to QC-2 showed an alkenyl proton for NB-1, whereas in QC-2 the alkenyl protons disappeared and new alkyl protons emerge (SI: Figure S3). UV-vis absorption spectroscopy of NB-1 was carried out in toluene, and the onset of absorption wavelength was found to be around 460 nm (SI: Figure S4(a)), which is significantly red-shifted from unsubstituted norbornadiene. This is attributed to the extended π -conjugation in NB-1, which enabled the photoisomerization with visible light. Irradiating a toluene solution of NB-1 with visible light resulted in its photoisomerization to QC-2 with isosbestic points in the absorption spectra indicating a clean photoconversion (Figure 1(a)).

In order to examine the robustness and fatigue resistance of the photoswitch, a toluene solution of NB-1 (\sim 44 μ M) was irradiated with a 405 nm laser diode at 50 °C for 2 min in order to fully convert the NB-1 to QC-2. The ensuing QC-2 was allowed to relax for 28 min at this temperature prior to the next irradiation cycle. The conversion was continuously probed at 430 nm. Under nitrogen atmosphere, ²⁴ the compound showed no sign of degradation even after well over 100 switching cycles (Figure 1(c)). In air, more than 100 switching cycles were

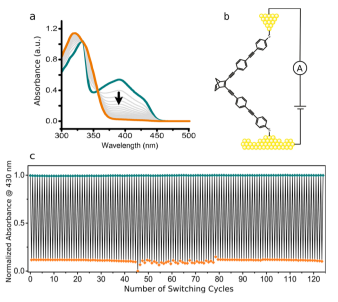


Figure 1. (a) Photoisomerization of NB-1 (dark cyan) to QC-2 (orange) indicated by the black arrow followed by UV/vis spectroscopy. The photoisomerization was performed using a 405 nm laser diode, and the sample was exposed to the irradiation every 10 s until the complete conversion of NB-1 through the gray plots to QC-2 was reached. (b) Sketch of device geometry under test in the STM break junction measurements. (c) No sign of degradation was observed in the accelerated stability test of the NB-1 (dark cyan maxima)/QC-2 (orange minima) system at elevated temperature (50 °C) sealed under nitrogen atmosphere; 124 photothermal switching cycles are shown.

performed with only 16% loss (SI: Figure S4(f)), corresponding to 99.8% yield per conversion cycle. This experiment reveals the robustness of the photoswitch, even at elevated temperatures.

The thermal relaxation of QC-2 to NB-1 over time was further investigated at different temperatures and found to follow first-order kinetics. The rate constants for the thermal switching reaction at a range of temperatures were extracted from the exponential fitting of the relaxation of QC-2 to NB-1. Rate constants obtained at four different temperatures (SI: Figure S4(b)) were fitted to the Arrhenius equation yielding a straight line (SI: Figure S4(c)). From the plot, the activation energy (E_a) as well as the pre-exponential factor (A) were determined for the thermal back conversion. The extracted values are summarized in Table 1.

Having demonstrated that the molecule effectively switches from the π -conjugated NB-1 state to the QC-2 state and vice versa in solution, we turn our attention to the charge transport properties of the system. STM-break junction (STM-BJ) experiments have been proven to be a robust method to obtain conductance values of single molecular junctions.^{26–33} This technique was used to examine the change in conductance at the single-molecule level between molecules NB-1 and QC-2 in order to gain insight into the details of the charge transport mechanism³⁴ (Figure 2). In STM-BJ a bias is applied between the tip and substrate, and the current is measured continuously as the tip is being retracted. Steps on the decaying conductance traces appear whenever molecules are bound between the two electrodes. With thousands of conductance vs distance curves collected, the most probable conductance value of the singlemolecule junction can be determined by statistical analysis.

Table 1. Photochemical and Thermodynamic Parameters for NB-1 and QC-2^{a,b}

$(\lambda_{\mathrm{NB-1}_{\mathrm{max}}}, \varepsilon) \; (\mathrm{nm}, \mathrm{M}^{-1} \; \mathrm{cm}^{-1})$	$(\lambda_{\text{QC-2max}'\varepsilon}) \text{ (nm, M}^{-1}\text{cm}^{-1})$	$E_{\rm a}~({\rm kJ~mol^{-1}})$	$A (s^{-1})$	$t_{1/2,25^{\circ}\mathrm{C}}\ (\mathrm{min})$	ϕ^*
$(332, 8.3 \times 10^4), (391, 4.3 \times 10^4)$	$(320, 9.1 \times 10^4)$	100.7	1.6×10^{14}	78.6	0.15

"Summary of calculated Arrhenius parameters for the thermal relaxation and photoisomerization quantum yield of QC-2 to NB-1. $^b\varepsilon$, molar extinction coefficient of NB-1 (two maxima) and QC-2 (one maximum); E_a , activation energy for the thermal relaxation of QC-2 to NB-1 derived from the Arrhenius equation; A, pre-exponential factor; $t_{1/2}$ 25 °C, half-life time of the thermal relaxation of QC-2 at 25 °C; ϕ^* , photoisomerization quantum yield of NB-1 to QC-2 obtained from the average of three separate measurements.

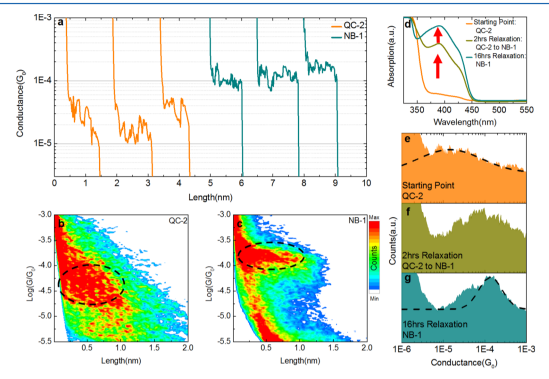


Figure 2. (a) Conductance vs length traces of the NB-1 state and the QC-2 state. Traces are displaced horizontally for clarity. (b) and (c) 2D histogram for QC-2 and NB-1 demonstrating a large change in the distribution of conductance values for the two states. (d) UV-vis spectrum for the relaxation of QC-2 to NB-1 in 16 h. (e)–(g) Conductance histograms of QC-2 before relaxation (e), after 2 h of relaxation (f), and after 16 h relaxation (g). All the experiments were carried out at room temperature \sim 23 °C.

To measure the conductance value of the norbornadiene-based photoswitching molecule at both NB-1 and QC-2 states, NB-1 was dissolved in mesitylene. The solution was then added to the STM cell (\sim 1 μ M in mesitylene), and a 50 mV bias was applied between the gold tip and the substrate, and the tapping process as described above was initiated with a tip speed of \sim 80 nm/s. For the QC-2 state, a UV-LED (centered at 405 nm) was employed to switch the molecule from NB-1 to QC-2 in solution phase, and the absorption spectrum was measured to verify the full conversion from NB-1 to QC-2. The STM-BJ conductance measurement for QC-2 was performed under continuous irradiation of UV-light to hinder the relaxation from QC-2 back to NB-1, and data collection was finished within 30 min after adding QC-2 to the cell.

Figure 2a shows individual decay curves for both QC-2 (orange traces) and NB-1 (dark cyan traces). It is observed that the steps of QC-2 are at lower conductance values compared to NB-1. By automatically selecting decay curves with clear steps, 2D histograms are constructed for QC-2 and NB-1 in Figure 2(b),(c). NB-1 yields a conductance value on the order of 10^{-4} G_0 , while QC-2 has a lower value on the order of 10^{-5} G_0 . Five sets of STM-BJ conductance measurements have been performed for both NB-1 and QC-2 states. Each measurement is based on the newly prepared substrate, solution, and gold tip.

The data are plotted in the 1D conductance histogram and fitted with a Gaussian distribution. The average conductance values each from the five experimental data sets are summarized in Table 2. The conductance values indicate that the change in molecular configuration directly leads to a conductance switch of molecular junction, with an ON/OFF ratio of 6.6.

Table 2. Conductance Measurement Results for NB-1 and QC-2 Forms $\,$

NB-1 conductance	QC-2 conductance	conductance ratio
$1.2 \pm 0.1 \times 10^{-4} G_0$	$1.9 \pm 0.8 \times 10^{-5} G_0$	6.6

To further confirm the correlation between states of the molecule and the junction conductance, in situ conductance measurements were performed in parallel with UV–vis absorption spectroscopy as the molecules relaxed from QC-2 to NB-1 (Figure 2(d)–(g)). A solution of QC-2 was initially added to the STM cell (~1 μ M in mesitylene), and three consecutive in situ conductance measurements were performed: immediately after molecular addition, 2 h later, and 16 h later. The system was maintained in a dark environment at room temperature throughout the experiment. Figure 2(e) shows a peak in the conductance histogram centered at ~1.9 × 10^{-5} G_0

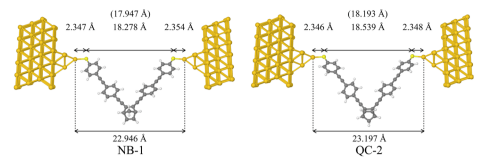


Figure 3. Optimized structure of the device region (scattering region) for NB-1 and QC-2 molecular junctions. The lengths in brackets correspond to the S-S distance of the gas-phase relaxed structures.

immediately following the addition of the QC-2 solution. Figure 2(f) shows the conductance histogram obtained after 2 h of relaxation and shows an evolution of the conductance distribution. The diminished low conductance peak and increased counts at high conductance range indicate the relaxation process from QC-2 to NB-1 is occurring. Finally, Figure 2(g) was obtained after 16 h of relaxation. A more distinct peak with a value around $1.2 \times 10^{-4} G_0$ is observed, indicating a complete relaxation. A control experiment with absorption spectrum measured in the solution phase was performed at the same time, and as demonstrated in Figure 2d, QC-2 underwent essentially complete relaxation process to NB-1 in 16 h, which is 99.99% back conversion calculated based on the values presented in Table 1.

The *in situ* relaxation STM break junction experiment suggests that the direct cause of the conductance change is the isomerization of the photoswitching molecule. We also tested switching NB-1 molecules to the QC-2 state by irradiating UV light onto the gold surface after molecules bound to gold, but no obvious conductance change was observed. We attribute this phenomenon to the rapid quenching of the excited states due to electronic coupling³⁵ once the molecule is coupled to the gold surface. Perhaps, this effect can be alleviated by inserting a saturated spacer between the photoactive unit and the electrode¹⁵ or by using a different anchoring group.³⁶ Detailed data for the STM-BJ measurements can be found in the SI Section III.

One of the advantages of the NB-1/QC-2 system is that the bonds that are broken and formed upon switching along the main conduction path undergo no skeletal rearrangement.³⁷ Hence, we expect the end-to-end length to remain largely unaffected. This will also yield junctions that are mechanically stable during the switching event. However, the limited structural change presents a potential problem for maximizing the conductance ON/OFF ratio.

It is possible to make an estimate of the conductance change upon switching when only a single bond changes from conjugated to saturated. By assuming that current only flows through the shortest through-bond path, knowledge of the characteristic conductance decay with length for conjugated and saturated systems allows us to estimate the difference between the two systems (full details in SI, section IV). This calculation yields a first estimate for the ON/OFF ratio of about 2, a very modest switching ratio. This calculation takes a simple view of conduction through the molecule and neglects through-space interactions or any contributions from longer pathways through the cyclic unit. While these assumptions might seem reasonable as a starting point, this raises a common challenge when trying to rationalize molecular electronics

measurements: whether molecules can be considered as simply a number of bonds/resistors in series.

To understand the ON/OFF ratio observed, we simulated the transport properties for NB-1 and QC-2 using quantum chemical modeling as shown in Figure 3. Full details of the simulation method are given in the SI (Section IV).

The calculated transmission for the two molecular junctions using DFTB+ is shown in Figure 4(a). The transmission through molecules terminated with thiols is usually dominated by a transmission resonance associated with the highest occupied molecular orbital (HOMO), whose energy lies close to the Fermi energy of the gold electrodes.^{38,39}

Our findings are in accordance with this: for both systems, the transmission peak close to the HOMO energy lies near the Fermi energy, and its tail is responsible for the transmission at the Fermi energy. Due to the smaller energy gap between occupied and virtual orbitals of π -conjugated molecules, the decay of the tail of the HOMO peak in NB-1 is slower than in QC-2 where the gap is larger. This results in a higher transmission for NB-1 at the Fermi energy. The calculated transmission ratio between NB-1 and QC-2 at the Fermi energy is 12. The trends observed in the transmission were also reproduced in DFT transmission calculations as shown in the SI (Figure S7).

While the transmission calculations confirm that we expect an ON/OFF ratio of significantly more than 2, the total transmission alone does not give us any insight into why this is the case. To examine this point, we probe the transport pathways through NB-1 and QC-2 using local current analysis, 40 the results of which are shown in Figure 4(b).

Local currents of NB-1 show that the current goes through the C–C double bond as expected, with no evidence of the current being dominated by anything other than the shortest through-bond path. However, the situation with QC-2 is somewhat more complicated. The simple picture of the current pathway had one double bond being replaced by one single bond, but this is not the dominant pathway we observe. In fact, the double bond is "replaced" by three σ bonds, with 88% of the current flowing out through the longer cyclobutane unit of QC-2 and only 12% taking the shortest through-bond path. Interestingly, if we perform the same back-of-the-envelope calculation assuming that we replace one conjugated bond with three saturated bonds we obtain an ON/OFF ratio of 28.

Clearly, the assumption that the current will always favor the shortest through-bond path breaks down in QC-2. Previous theoretical work on fully conjugated molecules has shown that the current may take a longer path when the shortest path is dominated by quantum interference.⁴¹ The geometry of the shortest path in QC-2 is indeed very close to a *cis* defect, which

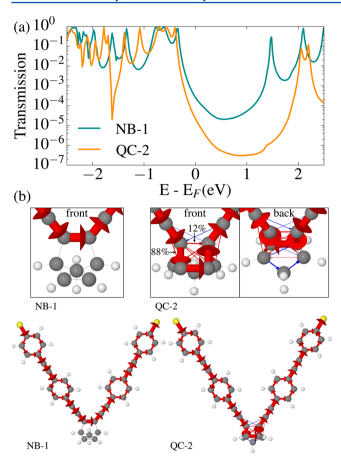


Figure 4. (a) Transmission calculated with DFTB+ and the auorg parameters for the molecules NB-1 and QC-2 sandwiched between the gold electrodes. (b) Local currents calculated with DFTB+ at the Fermi level energy with a symmetric bias of 0.1 eV. The red arrows indicate a positive contribution to the current density, while the (small) blue arrows indicate a negative contribution (*vide* zoomed out front and back insets Figure 4(b)). The photochemically active region affects the electron transport properties: a fully conjugated path in NB-1 is replaced by a saturated ring, which reduces the electron transmission. Moreover, the saturated region in QC-2 perturbs the current pathway: only 12% of the current goes through the shortest σ path, while the 88% goes through three σ bonds.

has been shown to reduce the current⁴² and exhibit interference effects.^{38,40}

When looking for ring-current reversals as the signature of interference, 40 we see some evidence of interference in QC-2 (Figures S8 and S9); however, the picture is not so definitive. What is clear is that some local current elements reverse direction, and the current through the shortest pathway goes through a minimum (significantly below the 12% seen at the Fermi energy) around the same energy as the transmission minimum. What this picture does not tell us is why the longer path is favored, as this is also a σ pathway with a dihedral close to zero.

We can probe this question by computing the local currents through a series of molecules with and without cyclopropane-strained carbons to determine which geometric features control the balance of current through the long and short paths (Figure 5). These molecular structures have been generated by simply removing atoms from the QC-2 relaxed structure and replacing them with hydrogen atoms that are then relaxed while keeping all parts of the original QC-2 structure fixed. We find that the

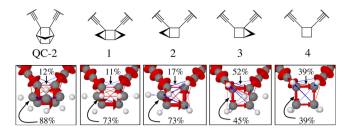


Figure 5. DFTB+ local currents for a series of modified **QC-2** structures. This analysis shows that the strained geometry of the cyclopropane rings significantly affects the local features of the current. Specifically, the strained geometry changes the balance between the short and long path.

symmetry of the carbons connecting the quadricyclane unit to the π -conjugated arms changes the balance between the current densities going through the long and the short path. When the carbons assume a strained sp³ symmetry (Figure 5 structure 1) the longer path is preferred. When the cyclopropane rings are removed, we find a balance close to 50/50 between the long and short path (Figure 5 structure 4).

While theory and experiment both demonstrate that the ON/OFF ratio should be higher than 2, the agreement between the two is certainly not quantitative. As always with these types of transport calculations, band line-up between the electrode Fermi energy and the molecular energy levels is far from perfect. Both DFTB+ and DFT calculations suffer from this problem, and it is clear from the transmission that the ON/OFF ratio predicted is sensitive to the position of the Fermi level. Beyond this issue, it is also known that treating the effective single-particle Hamiltonian from a DFT calculation as a true single-particle Hamiltonian for the purposes of a transport calculation may not always be a good approximation. Sec. 2019.

Chemically, we are also left with the question of whether the partitioning of the current through a polycyclic system into interatomic "bond current" contributions effectively describes the transport. Saturated systems are known to exhibit both surface and volume delocalization, ⁴⁴ which suggests the current might also flow through the space within the QC-2 core. It remains a question for future work as to when a "bond current" type representation is accurate, as opposed to a real-space local current representation. ^{45,46}

CONCLUSION

In conclusion, we have designed, synthesized, and characterized a thiol end-capped norbornadiene-based photoswitch, NB-1. The NB-1 photoswitch is found to have high fatigue resistance and excellent cyclability. Under nitrogen no sign of degradation was observed during well over 100 switching cycles at 50 °C. An STM-BJ conductance study revealed two conductance states, high (1.2 \pm 0.1 \times 10 $^{-4}$ G_0) and low (1.9 \pm 0.8 \times 10 $^{-5}$ G_0) conductances. While the switching ratio of the NB-1/QC-2 photoswitch pair is somewhat lower than some other well-studied systems, 15,17 it discloses a significantly different approach for charge transport mechanisms.

Through DFT calculations, we have shown that the NB-1/QC-2 photoswitch system effectively modulates the conductance through a junction, not only by breaking the conjugation but also by forcing the current to take a longer three-bond path in QC-2 through the saturated cyclobutane moiety. This result challenges the assumption that current will

simply flow through the shortest through-bond path and reminds us that the intricacies of quantum systems should not be ignored; i.e., it can be problematic to consider molecules and especially cage-like structures such as the QC-2 core as a sum of bonds. This is not to say that molecules generally cannot be considered as a sum of their parts; in fact, this has been shown to be an effective picture in many cases. It simply raises the question of how far one can break down the system, and clearly there are functional groups that need to be considered as single units, rather than a sum of bonds.

In general, despite the need to improve the quantum yield for real-world applications as is the case with other photoswitch systems, ¹² this study opens up a new perspective on tuning molecular conductance: using interference effects to effectively shuttle current between different parts of a molecule.

ASSOCIATED CONTENT

S Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcc.7b00319.

Detailed synthetic procedure, NMR spectra, absorption spectra, electrochemcial measurement, STM-BJ measurements, and the supporting quantum chemical calculations (PDF)

AUTHOR INFORMATION

Corresponding Authors

*E-mail: gsolomon@chem.ku.dk. *E-mail: jhihath@ucdavis.edu. *E-mail: mkasper@chalmers.se.

ORCID

Alessandro Pirrotta: 0000-0001-5487-1299 Gemma C. Solomon: 0000-0002-2018-1529 Kasper Moth-Poulsen: 0000-0003-4018-4927

Author Contributions

¹B.E.T. and H.B.L. contributed equally. The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

BET and KMP acknowledge funding from European Research Council (ERC-StG #337221, SIMONE). AP thanks Gabriele Penazzi for help with the transport calculations using DFTB+. AP and GCS received funding from the Danish Council for Independent Research | Natural Sciences and the Carlsberg Foundation. JH acknowledges support from the U.S. Office of Naval Research and the National Science Foundation. KB acknowledges funding from the Swedish Foundation for Strategic Research.

REFERENCES

- (1) Aviram, A.; Ratner, M. A. Molecular Rectifiers. Chem. Phys. Lett. 1974, 29, 277-283.
- (2) Haddon, R. C.; Lamola, A. A. The Molecular Electronic Device and the Biochip Computer: Present Status. *Proc. Natl. Acad. Sci. U. S. A.* **1985**, 82, 1874–1878.
- (3) Xiang, D.; Wang, X.; Jia, C.; Lee, T.; Guo, X. Molecular-Scale Electronics: From Concept to Function. *Chem. Rev.* **2016**, *116*, 4318–4440.

- (4) Sun, L.; Diaz-Fernandez, Y. a; Gschneidtner, T. a; Westerlund, F.; Lara-Avila, S.; Moth-Poulsen, K. Single-Molecule Electronics: From Chemical Design to Functional Devices. *Chem. Soc. Rev.* **2014**, *43*, 7378–7411
- (5) Zhang, C.; Du, M. H.; Cheng, H. P.; Zhang, X. G.; Roitberg, A. E.; Krause, J. L. Coherent Electron Transport through an Azobenzene Molecule: A Light-Driven Molecular Switch. *Phys. Rev. Lett.* **2004**, *92*, 158301.
- (6) Taherinia, D.; Frisbie, C. D. Photoswitchable Hopping Transport in Molecular Wires 4 Nm in Length. *J. Phys. Chem. C* **2016**, *120*, 6442–6449.
- (7) Broman, S. L.; Lara-Avila, S.; Thisted, C. L.; Bond, A. D.; Kubatkin, S.; Danilov, A.; Nielsen, M. B. Dihydroazulene Photoswitch Operating in Sequential Tunneling Regime: Synthesis and Single-Molecule Junction Studies. *Adv. Funct. Mater.* **2012**, 22, 4249–4258.
- (8) Tao, N. J. Electron Transport in Molecular Junctions. *Nat. Nanotechnol.* **2006**, *1*, 173–181.
- (9) Guédon, C. M.; Valkenier, H.; Markussen, T.; Thygesen, K. S.; Hummelen, J. C.; van der Molen, S. J. Observation of Quantum Interference in Molecular Charge Transport. *Nat. Nanotechnol.* **2012**, 7, 305–309.
- (10) Lindsay, S. M.; Ratner, M. A. Molecular Transport Junctions: Clearing Mists. *Adv. Mater.* **2007**, *19*, 23–31.
- (11) Zhang, J. L.; Zhong, J. Q.; Lin, J. D.; Hu, W. P.; Wu, K.; Xu, G. Q.; Wee, A. T. S.; Chen, W. Towards Single Molecule Switches. *Chem. Soc. Rev.* **2015**, *44*, 2998–3022.
- (12) Irie, M. Diarylethenes for Memories and Switches. *Chem. Rev.* **2000**, *100*, 1685–1716.
- (13) Matsuda, K.; Irie, M. Diarylethene as a Photoswitching Unit. *J. Photochem. Photobiol.*, C **2004**, *5*, 169–182.
- (14) Kim, Y.; Hellmuth, T. J.; Sysoiev, D.; Pauly, F.; Pietsch, T.; Wolf, J.; Erbe, A.; Huhn, T.; Groth, U.; Steiner, U. E.; et al. Charge Transport Characteristics of Diarylethene Photoswitching Single-Molecule Junctions. *Nano Lett.* **2012**, *12*, 3736–3742.
- (15) Jia, C.; Migliore, A.; Xin, N.; Huang, S.; Wang, J.; Yang, Q.; Wang, S.; Chen, H.; Wang, D.; Feng, B.; et al. Covalently Bonded Single-Molecule Junctions with Stable and Reversible Photoswitched Conductivity. *Science* **2016**, 352, 1443–1445.
- (16) Kumar, A. S.; Ye, T.; Takami, T.; Yu, B. C.; Flatt, A. K.; Tour, J. M.; Weiss, P. S. Reversible Photo-Switching of Single Azobenzene Molecules in Controlled Nanoscale Environments. *Nano Lett.* **2008**, *8*, 1644–1648.
- (17) Mativetsky, J. M.; Pace, G.; Elbing, M.; Rampi, M. A.; Mayor, M.; Samori, P. Azobenzenes as Light-Controlled Molecular Electronic Switches in Nanoscale Metal—molecule—metal Junctions. *J. Am. Chem. Soc.* **2008**, *130*, 9192–9193.
- (18) Roldan, D.; Kaliginedi, V.; Cobo, S.; Kolivoska, V.; Bucher, C.; Hong, W.; Royal, G.; Wandlowski, T. Charge Transport in Photoswitchable Dimethyldihydropyrene-Type Single-Molecule Junctions. J. Am. Chem. Soc. 2013, 135, 5974—5977.
- (19) Battacharyya, S.; Kibel, A.; Kodis, G.; Liddell, P. A.; Gervaldo, M.; Gust, D.; Lindsay, S. Optical Modulation of Molecular Conductance. *Nano Lett.* **2011**, *11*, 2709–2714.
- (20) Bonfantini, E. E.; Officer, D. L. The Synthesis of Norbornadienes Conjugatively Linked to Tetraphenylporphyrin and Anthracene: Towards a Norbornadiene-Derived Molecular Electronic Device. *J. Chem. Soc., Chem. Commun.* 1994, 1445–1446.
- (21) Löfås, H.; Jahn, B. O.; Wärnå, J.; Emanuelsson, R.; Ahuja, R.; Grigoriev, A.; Ottosson, H. A Computational Study of Potential Molecular Switches That Exploit Baird's Rule on Excited-State Aromaticity and Antiaromaticity. *Faraday Discuss.* **2014**, *174*, 105–124
- (22) Cristol, S.; Snell, R. Bridged Polycyclic Compounds. VI. The Photoisomerization of Bicyclo Hepta-2, 5-Diene-2, 3-Dicarboxylic Acid to quadricyclo[2,2,1,0] Heptane-2, 3-Dicarboxylic Acid. *J. Am. Chem. Soc.* **1958**, *80*, 1950–1952.
- (23) Chinchilla, R.; Nájera, C. Recent Advances in Sonogashira Reactions. Chem. Soc. Rev. 2011, 40, 5084–5121.

- (24) Dreos, A.; Borjesson, K.; Wang, Z.; Roffey, A.; Norwood, Z.; Kushnir, D.; Moth-Poulsen, K. Exploring the Potential of a Hybrid Device Combining Solar Water Heating and Molecular Solar Thermal Energy Storage. *Energy Environ. Sci.* **2017**, *10*, 728–734.
- (25) Stranius, K.; Börjesson, K. Determining the Photoisomerization Quantum Yield of Photoswitchable Molecules in Solution and in the Solid State. *Sci. Rep.* **2017**, *7*, 41145.
- (26) Xiao, X.; Xu, B.; Tao, N. J. Measurement of Single Molecule Conductance: Benzenedithiol and Benzenedimethanethiol. *Nano Lett.* **2004**, *4*, 267–271.
- (27) Venkataraman, L.; Klare, J. E.; Nuckolls, C.; Hybertsen, M. S.; Steigerwald, M. L. Dependence of Single-Molecule Junction Conductance on Molecular Conformation. *Nature* **2006**, *442*, 904–907.
- (28) Aradhya, S. V.; Meisner, J. S.; Krikorian, M.; Ahn, S.; Parameswaran, R.; Steigerwald, M. L.; Nuckolls, C.; Venkataraman, L. Dissecting Contact Mechanics from Quantum Interference in Single-Molecule Junctions of Stilbene Derivatives. *Nano Lett.* **2012**, *12*, 1643–1647.
- (29) Agraït, N.; Yeyati, A. L.; van Ruitenbeek, J. M. Quantum Properties of Atomic-Sized Conductors. *Phys. Rep.* **2003**, 377, 81–279.
- (30) Gittins, D. I.; Bethell, D.; Schiffrin, D. J.; Nichols, R. J. A Nanometre-Scale Electronic Switch Consisting of a Metal Cluster and Redox-Addressable Groups. *Nature* **2000**, *408*, 67–69.
- (31) Cui, X. D.; Primak, A.; Zarate, X.; Tomfohr, J.; Sankey, O. F.; Moore, A. L.; Moore, T. A.; Gust, D.; Harris, G.; Lindsay, S. M. Reproducible Measurement of Single-Molecule Conductivity. *Science* **2001**, 294, 571–574.
- (32) Rascón-Ramos, H.; Artés, J. M.; Li, Y.; Hihath, J. Binding Configurations and Intramolecular Strain in Single-Molecule Devices. *Nat. Mater.* **2015**, *14*, 517–522.
- (33) Aragonès, A. C.; Aravena, D.; Cerdá, J. I.; Acís-Castillo, Z.; Li, H.; Real, J. A.; Sanz, F.; Hihath, J.; Ruiz, E.; Díez-Pérez, I. Large Conductance Switching in a Single-Molecule Device through Room Temperature Spin-Dependent Transport. *Nano Lett.* **2016**, *16*, 218–226.
- (34) Xu, B.; Tao, N. J. Measurement of Single-Molecule Resistance by Repeated Formation of Molecular Junctions. *Science* **2003**, *301*, 1221–1223.
- (35) Dulić, D.; van der Molen, S. J.; Kudernac, T.; Jonkman, H. T.; de Jong, J. J. D.; Bowden, T. N.; van Esch, J.; Feringa, B. L.; van Wees, B. J. One-Way Optoelectronic Switching of Photochromic Molecules on Gold. *Phys. Rev. Lett.* **2003**, *91*, 207402.
- (36) Su, T. A.; Neupane, M.; Steigerwald, M. L.; Venkataraman, L.; Nuckolls, C. Chemical Principles of Single-Molecule Electronics. *Nat. Rev. Mater.* **2016**, *1*, 16002.
- (37) Dauben, W. G.; Cargill, R. L. Photochemical transformations—VIII The Isomerization of bicyclo[2.2.1]heptadiene to quadricyclo[2.2.1.0.0]heptane (Quadricyclane). *Tetrahedron* **1961**, 15, 197–201.
- (38) Tan, A.; Balachandran, J.; Sadat, S.; Gavini, V.; Dunietz, B. D.; Jang, S.-Y.; Reddy, P. Effect of Length and Contact Chemistry on the Electronic Structure and Thermoelectric Properties of Molecular Junctions. J. Am. Chem. Soc. 2011, 133, 8838–8841.
- (39) Malen, J. A.; Doak, P.; Baheti, K.; Tilley, T. D.; Segalman, R. A.; Majumdar, A. Identifying the Length Dependence of Orbital Alignment and Contact Coupling in Molecular Heterojunctions. *Nano Lett.* **2009**, *9*, 1164–1169.
- (40) Solomon, G. C.; Herrmann, C.; Hansen, T.; Mujica, V.; Ratner, M. a. Exploring Local Currents in Molecular Junctions. *Nat. Chem.* **2010**, 2, 223–228.
- (41) Tsuji, Y.; Movassagh, R.; Datta, S.; Hoffmann, R. Exponential Attenuation of through-Bond Transmission in a Polyene: Theory and Potential Realizations. *ACS Nano* **2015**, *9*, 11109–11120.
- (42) Li, C.; Pobelov, I.; Wandlowski, T.; Bagrets, A.; Arnold, A.; Evers, F. Charge Transport in Single AulalkanedithiollAu Junctions: Coordination Geometries and Conformational Degrees of Freedom. *J. Am. Chem. Soc.* **2008**, *130*, 318–326.

- (43) Strange, M.; Rostgaard, C.; Häkkinen, H.; Thygesen, K. S. Self-Consistent GW Calculations of Electronic Transport in Thiol- and Amine-Linked Molecular Junctions. *Phys. Rev. B: Condens. Matter Mater. Phys.* **2011**, 83, 1–12.
- (44) Cremer, D. Pros and Cons of σ -Aromaticity. *Tetrahedron* 1988, 44, 7427–7454.
- (45) Xue, Y.; Ratner, M. A. Local Field Effects in Current Transport through Molecular Electronic Devices: Current Density Profiles and Local Nonequilibrium Electron Distributions. *Phys. Rev. B: Condens. Matter Mater. Phys.* **2004**, *70*, 81404.
- (46) Walz, M.; Bagrets, A.; Evers, F. Local Current Density Calculations for Molecular Films from Ab Initio. *J. Chem. Theory Comput.* **2015**, *11*, 5161–5176.
- (47) Manrique, D. Z.; Al-Galiby, Q.; Hong, W.; Lambert, C. J. A New Approach to Materials Discovery for Electronic and Thermoelectric Properties of Single-Molecule Junctions. *Nano Lett.* **2016**, *16*, 1308–1316.