Letter

Conductance Measurements of Polar Molecules in a Nonconducting Solvent

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A central goal of molecular electronics is to develop functional molecular-scale components that may one day serve as nanoscale electronic circuit elements.¹ To drive future advances, a deeper understanding of the complex interplay between electrode, linker group, and molecular backbone properties and the impact of this relationship on charge transport through metal-molecule-metal junctions is required. We focus here on carboxylic acids, well-recognized linkers that facilitate the spontaneous formation of singlemolecule junctions from aqueous solutions^{2,3} or components adsorbed/deposited on surfaces (Figure 1a).⁴⁻⁶ Their weakly



Figure 1. (a) Schematic of molecular junctions formed from linear oligoalkane α, ω -bis(carboxylic acids) bound between gold electrodes. (b) Molecular structures of cyclic α, ω -bis(carboxylic acid) junction components and isochroman, a high-boiling-point ethereal solvent.

acidic character, hydrophilicity, and reactivity have enabled studies exploring the pH-dependence of junction conductance,^{3,6,7} the use of electrochemically stabilized silver, copper, and palladium electrodes,^{8–10} or the reversible formation of ester-containing molecular circuits.⁵ However, the capacity of bis(carboxylic acids) to generate extended hydrogen-bonding networks in the solid state¹¹ can limit their solubility in

commonly used electrically insulating scanning tunneling microscope (STM) solvents. To date, only two single-molecule conductance studies in such solvents have been reported: one using tetradecane (TD), restricted to compounds containing a *single* carboxylic acid,⁷ and one of bis(carboxylic acids) in toluene.⁸

Here we report that nonconducting organic solvents comprising oxygen functionalities can improve the solubility of polar compounds of interest, presumably by disrupting analyte—analyte intermolecular interactions. Critically, the use of isochroman (IC; Figure 1b) enables conductance measurements of α, ω -bis(carboxylic acids) with different backbones using uncoated¹² STM tips. These experimental studies, supported by *first-principles* calculations of model junctions, provide important new insights into the nature of charge transport across the -AuOC(O)- interfacial contact.

We perform conductance measurements using the STMbased break junction (STM-BJ) method (see the SI for more details).^{13,14} This technique involves repeatedly pushing an uncoated gold tip in and out of a gold substrate while applying a voltage bias (V_{bias}) between these electrodes and measuring the current (I) as a function of the tip-substrate displacement. Step features observed in the resulting conductance ($G = I/V_{\text{bias}}$)-displacement traces correspond to the formation of Au-Au point contacts at close to integer multiples of 1 G_0 (= $2e^2/h$) and molecular junctions at lower conductance (after

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the addition of a suitable analyte in solution). Thousands of these traces are compiled, without data selection, into 1D conductance and 2D conductance-displacement histograms. The resulting histogram features reveal the most probable properties of the junctions studied.

The utility of four different high-boiling-point ethereal solvents for STM-BJ studies was first evaluated using 4,4'bipyridine (bipy). Two solvents are cyclic ethers: IC (boiling point (BP) ~214 °C) and 2,3-dihydrobenzofuran (188 °C). The other two are acyclic ethers: dioctylether (286 °C) and cyclopentyl methyl ether (106 °C). Explicit molecular structures for these compounds are provided in Figure S1a. Each measurement provides histograms containing the characteristic two-peak feature of bipy junctions, corresponding to N lone pair–Au and pyridyl π –Au contact geometries,¹⁵ indicating that these solvents do not impede junction formation (Figure S1b-e). The conductance of junctions measured in these solvents typically lie between or above those obtained from measurements in TD or 1,2,4-trichlorobenzene, further indicating these new solvents only weakly interact with the gold surface (Table S1).^{16,17} The low instrument noise floor in these, and subsequent, measurements illustrate the electrically insulating nature of these solvents; no significant electrochemical currents are observed. We select IC for additional studies given its moderate BP and mixed aliphaticaromatic structure, which we reason will help solubilize a wider range of compounds.

We subsequently perform conductance measurements using IC solutions of alkane α, ω -bis(carboxylic acids) (Cn, where n is the number of carbon atoms between HOC(O)- linkers; Figure 1a). We plot, in Figure 2, overlaid 1D histograms for



Figure 2. (a) Overlaid 1D histograms for *Cn* measured in IC ($V_{\text{bias}} = 250 \text{ mV}$, 5000 traces). Inset: plot of the experimental single-molecule conductance against n ($\beta = 0.90/n$). (b) 2D histogram for C6.

these diacids. All histograms contain a sharp peak feature toward lower conductance, assigned to single-molecule junctions. In each case, we also observe an additional peak or shoulder at ~2× the conductance of the first peak, which we tentatively attribute to the formation of junctions with two molecules in parallel. These distinct features are also clearly observed in the corresponding 2D histograms and individual conductance–displacement traces (Figure 2b and S2). A semilog plot of the most probable conductance for each single-molecule junction against *n* shows that these values exhibit an exponential length dependence indicative of tunneling transport (Figure 2, inset). We obtain a tunneling decay constant (β) of 0.90/*n* and a contact conductance (G_c) of 7.2 × 10⁻³ G_0 from a linear fit to this data using $G = G_c \exp(-\beta n)$. This β is consistent with values obtained for other series of oligoalkanes

with different linker groups, and the low G_c (e.g., relative to 4.8 $\times 10^{-2} G_0$ for –SMe), attributed to the additional carbon atom in the linker group, also agrees well with previous reports.^{3,7,8,18} Conductance data for all diacid junctions are provided in Table S2. Additional discussion regarding studies of ethanedioic acid (C0) in IC, and C6 introduced using different methods, is provided in the SI.

We next evaluate the conductance of junctions formed from α, ω -bis(carboxylic acids) comprising cyclic 1,4-phenylene (**Ph**, n = 4), 1,4-cyclohexane (**Cy**, n = 4), and 1,4-xylylene (**Xy**, n = 6) backbones (for molecular structures, see Figure 1b). Here, values of n indicate the number of carbon atoms between linkers through a single branch of the backbone ring. In Figure 3a we plot overlaid 1D conductance histograms for these cyclic



Figure 3. (a) Overlaid 1D histograms for measurements of **Ph**, **Cy**, and **Xy** in **IC** ($V_{\text{bias}} = 250 \text{ mV}$, $\geq 5000 \text{ traces}$). (b) Au₁-cluster junction geometries that provide the highest tunnel couplings for each system.

diacids, in which we again typically observe two overlapping peaks characteristic of one and two molecular junctions. Surprisingly, while their hydrocarbon bridges differ in the number and structure (aliphatic, aromatic) of carbon atoms, these diacid junctions each exhibit a conductance within a factor of 1.1 from each other and between 60% and 68% of the conductance of C4 (Table S2). Repeated measurements of these analytes highlight the reproducibility of this result (Figure S3a–c). Our findings contrast with the ~10× higher conductance reported for Ph compared to Xy junctions formed using copper electrodes,¹⁹ although the conductance data obtained for Ph in that study included additional peak features at lower conductance.

To provide additional insights into the electronic properties of these junctions, we turn to *first-principles* calculations based on density functional theory (DFT; see the SI for further details and extended discussion). Each contact is initially modeled in Au₁ cluster junctions using a κ^1 (O-monodentate) coordination mode. This geometry is observed in molecular structures of Au(I)–carboxylate complexes determined from single-crystal X-ray diffraction (although Ag(I) and Cu(I) complexes with κ^2 (O,O-bidentate) or bridging coordination geometries are known).²⁰ Given that rotations about the unconstrained and sterically unimpeded single bonds in each system (e.g., AuOC(O)-aryl) are expected to be soft degrees of freedom,^{16,21} we performed geometry optimizations of each junction using input structures with different dihedral angles (Figure S5).

In Figure 3b we plot illustrative optimized geometries for Ph-Au₁, Cy-Au₁, and Xy-Au₁, which support a qualitative rationalization of the measured conductance for these

junctions. For Cy, transport is through a sp³-hybridized n = 4backbone, providing a similar but lower conductance compared to C4. This conductance ordering may be rationalized given that the structural constraints of the cyclohexane group ensure the alkane chain between each linker cannot adopt a more conductive all-trans configuration,²² and transmission through this cyclic hydrocarbon backbone may be further reduced through destructive σ interference effects.²³ Transport through conjugated phenylene backbones may be expected to be more efficient than that through nonconjugated alkanes due to their smaller HOMO-LUMO gaps that better align frontier orbitals with the electrode Fermi level $(E_{\rm F})$. However, we note that the geometry of the carboxylic acid linker in Ph orients the Au-O bond into the plane of the benzene group, perpendicular to the conjugated *p*-orbitals of the ring. This reduces electronic coupling between the delocalized backbone π -system and the electrode, forcing transport through the sp^2 -hybridized σ framework.²⁴ For Xy it is possible to access junction geometries that better align the terminal AuOC(O) – groups with the conjugated phenylene backbone, apparently providing a junction conductance comparable to those of Ph and Cy despite the longer, n = 6, conduction path.

We apply these and additional Au₁ cluster junctions to calculate tunnel couplings (4t²), a metric that correlates well with experimental conductance.^{25,26} For these neutral carboxylate-linked models, we obtain 2t from the energy difference between LUMO and LUMO+1 (see Computational Methods in the SI for justification), which provides the expected symmetric and antisymmetric orbital pair exhibiting Au *s*-O *p*-antibonding character (Figure 4a; analogous orbitals are provided for other junctions in Figure S7).²¹ We validate our



Figure 4. (a) Isosurface plots (isovalue of 0.06 Å⁻³) of the tunnelcoupled LUMO and LUMO+1 orbitals for C4-Au₁. (b) A plot of the calculated tunnel couplings for C*n*-Au₁ against *n* ($\beta = 0.86/n$), overlaid with the largest tunnel coupling for Ph-Au₁, Cy-Au₁, and Xy-Au₁ (all κ^1). (c) (Left) Overlaid transmission calculations for selected junctions, comprising the same (frozen) geometries used to calculate 4t² in panel (b). (Right) The different contact geometries evaluated with C4 junctions.

approach by plotting, in Figure 4b, $4t^2$ versus *n* for Cn-Au₁ models.²⁵ From a linear fit to these data, substituting $G = 4t^2$ into $G = G_c \exp(-\beta n)$, we obtain $\beta = 0.86/n$, in good agreement with our experimental result (Figure 2a). In Figure 4b, we also overlay the largest tunnel couplings obtained from all of the geometries evaluated for Ph-Au₁, Cy-Au₁, and Xy-Au₁ junctions (corresponding to the geometries in Figure 3b). While the tunnel coupling for each system is clearly dependent on its conformation (Table S4), the similarity of each of these maximum couplings to each other, and to the value found for C4-Au₁, supports our experimental finding that junctions formed from this series of molecules can exhibit a similar conductance. The potential energy landscapes of these junctions could be explored in future studies to evaluate the relative probabilities and thermal population of each conformation.

To further rationalize these findings, in Figure 4c we plot overlaid calculated transmission functions for C4, Ph, Cy, and Xy junctions with the same (frozen) geometries used to calculate 4t² in Figure 4b. These quantum transport calculations are performed within the framework of DFT and the nonequilibrium Green's function (NEGF) formalism using FHI-aims²⁷ combined with the AITRANSS transport module (see the SI for additional details).²⁸⁻³¹ Our calculations support the conclusions obtained from the tunnel coupling analysis, with each junction exhibiting similar zero-bias conductance due to strong $E_{\rm F}$ pinning to the HOMO. In this sense, transport is dominated by weakly coupled occupied orbitals, and no significant midgap states³² are observed for these chemisorbed Au-O contacts. Analogous calculations for fully relaxed junctions exhibit qualitatively similar features (see Figure S9 and the associated discussion). Interestingly, both tunnel coupling and transmission calculations reveal that the conductance of C4 junctions varies by only a factor of 3-4 when both carboxylates are contacted through κ^1 or κ^2 coordination modes (Figure 4c and Tables S4 and S5), in agreement with previous studies focused on junctions with only a single carboxylate linker. We suggest that both modes are experimentally accessible and contribute, along with changes in the backbone geometry, to the width of the conductance peaks observed.

Independent of the precise coordination geometry, these linker groups are widely considered to bind in junctions as carboxylates $(-COO^{-})$.^{3,7} The apparently spontaneous deprotonation of -COOH upon binding to gold is a process that is also important for thiols³³ and terminal alkynes,³⁴ although the larger pK_a of the alkyne C–H group may suggest a distinct deprotonation mechanism (pK_a^{DMSO} : PhCO₂H = 11.1, PhSH = 10.31, PhCCH = 28.7).³⁵ Inspired by their plausible κ^1 coordination mode, we propose that carboxylic acids could be considered simply as -OH linkers comprising an acidic proton. While -OH groups have to date scarcely been explored as junction linkers, they may yet be widely utilized after deprotonation with an appropriate base (as recently reported for phenol)³⁶ or at a metal surface after incorporating adjacent chemical functionality that lowers their pK_{a} . Such contact chemistries could prove useful for forming junctions with oxophilic metal electrodes,³⁷ or to evaluate in situ chemical reactions thought to result in alkoxide-terminated junctions.³

Together, the results of this study motivate additional investigations using **IC** or related solvents to probe the properties of carboxylic acid-linked single-molecule junctions.

We note our approach may be expanded to characterize other families of compounds with polar or charged³⁶ backbones/ contact groups. By substituting polar solvents with non-conducting analogues that can perform a similar solubilizing function, we greatly simplify STM-BJ experiments that typically otherwise require coated STM tips to minimize background electrochemical currents.¹² This may prove valuable for glovebox-based STM-BJ studies useful for investigations of air-sensitive electrode metals³⁸ and molecules,³⁹ in which the frequent use of coated STM tips would likely present a substantial experimental burden.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsphyschemau.5c00021.

Additional experimental details and conductance and computational data (PDF)

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Notes

The authors declare no competing financial interest.

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