

Letter pubs.acs.org/JPCL

# Simultaneous Suppression of $\pi$ - and $\sigma$ -Transmission in $\pi$ -Conjugated Molecules

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Cite This: J. Phys. Chem. Lett. 2020, 11, 7400-7406



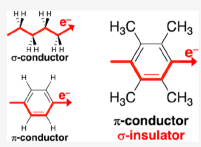
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Supporting Information

ABSTRACT: Molecular dielectric materials require ostensibly conflicting requirements of high polarizability and low conductivity. As previous efforts toward molecular insulators focused on saturated molecules, it remains an open question whether  $\pi$ - and  $\sigma$ -transport can be simultaneously suppressed in conjugated systems. Here, we demonstrate that there are conjugated molecules where the  $\sigma$ -transmission is suppressed by destructive  $\sigma$ -interference, while the  $\pi$ -transmission can be suppressed by a localized disruption of conjugation. Using density functional theory, we study the Landauer transmission and ballistic current density, which allow us to determine how the transmission is affected by various structural changes in the molecule. We find that in para-linked oligophenyl rings the  $\sigma$ -transmission can be suppressed by changing the remaining hydrogens to methyl groups due to the inherent



gauche-like structure of the carbon backbone within a benzene ring, similar to what was previously seen in saturated systems. At the same time, the methyl groups fulfill a dual purpose as they modulate the twist angle between neighboring phenyl rings. When neighboring rings are orthogonal to each other, the transmission through both  $\pi$ - and  $\sigma$ -systems is effectively suppressed. Alternatively, breaking conjugation in a single phenyl ring by saturating two carbons atoms with two methyl substituents on each carbon, results in suppressed  $\pi$ - and  $\sigma$ -transport independent of dihedral angle. These two strategies demonstrate that methylsubstituted oligophenyls are promising candidates for the development of molecular dielectric materials.

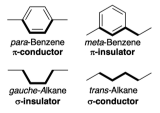
The capacitance of dielectric materials used in the semiconductor industry continues to be limited by leakage currents due to direct tunneling, and by the polarizability of the material.<sup>1-4</sup> This problem means that there is ongoing interest in molecules with extremely low conductance and high polarizability as researchers explore the possibility of molecular dielectric materials. 5-10 However, in a hydrocarbon backbone,  $\pi$ -systems provide the highest polarizability, while saturated molecules have hitherto been found to be the best insulators. 11-14

While the concept of a conjugated insulator sounds at first to be oxymoronic, there have been hints in this direction. In the low-bias regime, the single-molecule conductance is mediated by a coherent tunneling mechanism that can be suppressed by destructive quantum interference in either the  $\pi$ - or  $\sigma$ systems. 14–17 Consequently, the conductance of a meta-linked benzene is expected to be much lower than that of *para*- and *ortho*-linked benzenes. <sup>18–22</sup> However, destructive quantum interference has not been predicted or observed to occur in both the  $\pi$ - and  $\sigma$ -systems of a single molecule. For example, only the  $\pi$ -transmission is suppressed in meta-benzenes and similar cross-conjugated motifs; thus, Kiguchi et al. found the conductance of meta-linked benzenedithiol and benzenediamine to "only" be around half that of their para-linked counterparts as significant  $\sigma$ -transport remains. <sup>23</sup> In general, molecules with  $\pi$ -interference do not achieve lower conductance than alkanes of equivalent length because the  $\sigma$ transmission is not suppressed. 14,24-27 The conductance

difference between meta- and para-linked benzenes is therefore larger in longer molecules when  $\sigma$ -transport is naturally suppressed due to faster decay with length. 28-38 An optimal insulating/dielectric molecule will, however, deliver the lowest possible conductance with the shortest possible length. We consider the para-case the archetypical  $\pi$ -conductor and the meta-case the archetypical  $\pi$ -insulator, as shown in Scheme 1.

We and others recently uncovered chemical design rules for suppression of the  $\sigma$ -transmission in saturated molecules by destructive quantum interference.  $^{14,39-41}$  The  $\sigma$ -interference

Scheme 1. Transport Properties of Organic Molecular Wires



Received: June 4, 2020 Accepted: August 12, 2020 Published: August 12, 2020





effect can appear when there is a gauche defect in the molecular backbone, i.e., a dihedral angle of approximately  $60^\circ$  or less. Furthermore, the suppression is strongest in methyl-substituted species (as opposed to all substituents being hydrogen). Therefore, the interference effect is clearest in the single-molecule conductance of saturated silanes that are generally permethylated, though it is also significant in alkanes. By considering *trans*-alkanes the archetypical  $\sigma$ -conductor and *gauche*-alkanes the archetypical  $\sigma$ -insulator, we deduce from Scheme 1 that the structural requirements for  $\pi$ -and  $\sigma$ -interference appear to be mutually exclusive. The  $\sigma$ -path through *meta*-benzene is structurally akin to that of the *trans*-alkane, and the  $\sigma$ -path of *para*-benzene is structurally akin to that of the *gauche*-alkane. It is with this structural analogy in mind that we shall explore if both the  $\pi$ - and  $\sigma$ -transmission of conjugated molecules can be suppressed in a single system.

In this Letter, we explore how both the  $\sigma$ - and  $\pi$ transmission can be switched off in  $\pi$ -conjugated biphenyland triphenyl-based molecules using a combination of structural modifications that both disrupt  $\pi$ -conjugation and support  $\sigma$ -interference. These systems are well-studied both theoretically and experimentally in the context of molecular electronics and are ideal for exploring the  $\sigma$ -transmission in  $\pi$ conjugated systems because the  $\pi$ -transmission can be controlled by the torsion angle of the phenyl rings. $^{46-50}$  The torsion angle can be controlled by the choice of substituents on the phenyl rings, and several molecular systems covering the full range of angles from coplanar to perpendicular orientation have been synthesized. 47-55 This includes methylating the rings, the same substituent pattern that supports maximal destructive  $\sigma$ -interference in saturated molecules. <sup>47–55</sup> When two phenyl rings are orthogonal relative to each other, the  $\pi$ – $\pi$ coupling will be almost fully suppressed. This alters the electronic properties because  $\pi - \sigma$  and  $\sigma - \sigma$  electronic couplings become more important than the  $\pi$ - $\pi$  coupling, <sup>46,56</sup> e.g., the surprisingly high hyperpolarizability of twisted chromophores.57

We compute the Landauer transmission and ballistic current density using our in-house transport code, which utilizes the Atomic Simulation Environment (ASE) and GPAW. 60,61 Our code is available as freeware on https://github.com/ marchgarner/Current Density. The methods are described in detail in recent work and in Supporting Information part A. 62,63 Briefly, all molecules are optimized using density functional theory (DFT) with the PBE functional and 6-311G(d,p) basis set as implemented in Gaussian09.<sup>64,65</sup> The optimized structures are rotated without further optimization; thus all degrees of freedom are frozen except for the dihedral angle(s). The transmission and ballistic current density are calculated using DFT in combination with the nonequilibrium Green's functions approach using s-band approximated electrodes. The ballistic current density is calculated in the low-bias limit using a bias of 1 mV, which is opened symmetrically around the Fermi energy. Junction structures are included in xyz format as Supporting Information.

First, we take a simple biphenyl system with and without methyl substituents, as shown in Figure 1a. In agreement with previous work, the transmission at the Fermi energy follows a cosine-squared dependence with the dihedral angle between the phenyl rings. Shown in Figure 1b, the difference between BPh-H and BPh-Me is negligible on a log scale. In both systems the transmission drops rapidly as the dihedral angles approach 90°. While the methyl substituents are undoubtedly

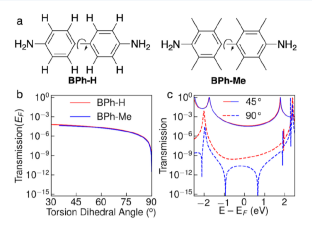


Figure 1. Transmission of biphenyl under axial torsion. (a) Chemical structure of nonmethylated (BPh-H) and permethylated (BPh-Me) biphenyl functionalized with amine groups. (b) Transmission of BPh-H (red) and BPh-Me (blue) at the Fermi energy as a function of dihedral angle. (c) Transmission as a function of energy for BPh-H (red) and BPh-Me (blue) calculated at 45° (full lines) and 90° (dashed lines) dihedral angles.

electron donating compared with hydrogen, this effect does not have a significant impact on the transmission at the Fermi energy. This is also the case if we look at the full transmission plots at 45° in Figure 1c; only small differences are evident and these are not near the Fermi energy. However, at 90° the transmission plots are remarkably different across a large energy range. Both **BPh-H** and **BPh-Me** have low transmission, but in the methylated case two sharp antiresonances appear close to the Fermi energy; a clear signature of suppression due to destructive quantum interference.

At 90° torsion the phenyl rings are orthogonal, and the  $\pi$ - $\pi$ coupling between the phenyl rings is negligible. Instead, the transmission is mediated by  $\pi$ - $\sigma$  coupling, i.e.,  $\pi$  through one ring and  $\sigma$  through the second (or vice versa). This can be deduced from the current density plots in Supporting Information part B and is in agreement with previous theoretical studies of biphenyl systems. 46,56 As the transmission is mediated in the  $\sigma$ -system in one ring that contains a gauchelike dihedral angle of 0°, the transmission is very low. However, the introduction of methyl substituents in BPh-Me induces destructive  $\sigma$ -interference further reducing the transmission compared with BPh-H. This behavior upon substitution is similar to what we have recently demonstrated in cyclic and bicyclic silanes and alkanes. 14,41 The appearance of two antiresonances near the Fermi energy is predicted by the Hückel-type Ladder C model (see Supporting Information part C) but is usually not seen in DFT calculations. 39,66,67 We note here that crystal structures obtained on similar biphenyl molecules found the dihedral angles for the hydrogen substituted system to be 36.4° while the methylated system was 89°. 48 The optimized dihedral angles of BPh-H and BPh-Me are in good agreement with these values, and the transmissions of the optimized structures are included in Figure S9. Thus, while the geometry we find to maximally suppress  $\sigma$ - and  $\pi$ -transmission represents a higher energy structure for BPh-H, it is very close to the minimum energy structure for BPh-Me.

The suppression of the  $\sigma$ -transmission is even clearer in the triphenyl systems, **TPh-H** and **TPh-Me**, shown in Figure 2a, where a significant reduction in the transmission at the Fermi energy is evident across a wider range of dihedral angles. We

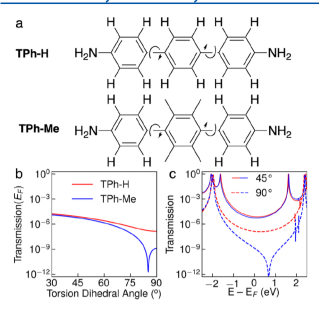
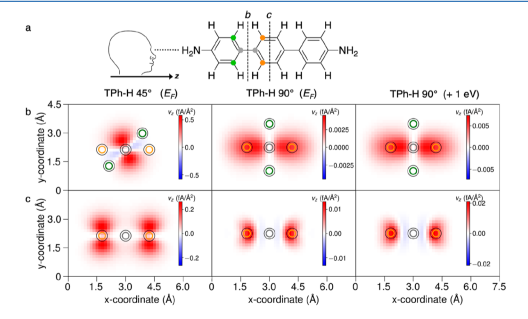


Figure 2. Transmission of triphenyl under axial torsion. (a) Chemical structure of amine-functionalized triphenyl (TPh-H) and centrally methylated (TPh-Me) triphenyl. (b) Transmission of TPh-H (red) and TPh-Me (blue) at the Fermi energy as a function of dihedral angle. (c) Transmission as a function of energy for TPh-H (red) and TPh-Me (blue) calculated at 45° (full lines) and 90° (dashed lines) dihedral angles.

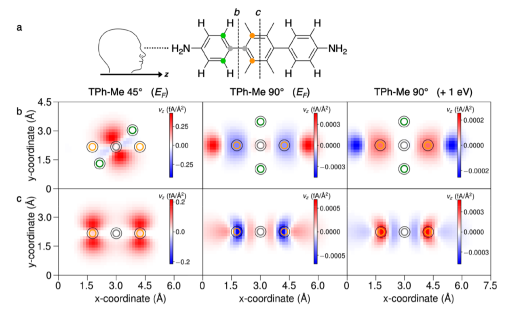
simultaneously rotate the two outer rings relative to the central ring. Only the central ring is methylated in **TPh-Me**. This may shift the minimum energy structure to a lower dihedral angle, and a related biphenyl system with a single methyl substituent on each ring was found from a crystal structure to have a dihedral angle of  $79.7^{\circ}.^{48}$  However, at the level of theory applied in this letter, we find the optimized dihedral angles of **TPh-Me** to be  $90.0^{\circ}$ , and the transmission plots for the

optimized conformations of **TPh-Me** and **TPh-H** are shown in Figure S9. Shown in Figure 2b, the transmission at the Fermi energy is similar at first but the transmission rapidly drops off in the methylated case as the dihedral angle approaches 90°. The transmission plots in Figure 2c reveal that at 90° the transmission of **TPh-Me** is suppressed in a broad energy range due to an antiresonance near the Fermi energy. The introduction of the otherwise benign methyl substituents on the central ring changes the transmission by several orders of magnitude because of the  $\sigma$ -interference effect. In good agreement with this result, Chen et al. recently reported the single-molecule conductance of a similar methyl-substituted triphenyl system to be surprisingly low. <sup>54</sup>

Let us take a closer look at the interactions that mediate the current in TPh-H and TPh-Me and in particular the role of the  $\sigma$  and  $\pi$  systems in mediating the current. In a molecule where the  $\pi$ -system dominates the current (oligo-phenyl systems with coplanar rings), we expect to see current density above and below the plane of the benzene rings. On the other hand, if the  $\sigma$ -system is carrying the current, we expect to see current density in the plane of the benzene ring directly through the  $\sigma$ bonds. These expectations follow the symmetry of the  $\sigma$  and  $\pi$ systems. In Figures 3 and 4, the current density is plotted through two select surfaces of TPh-H and TPh-Me in the transport direction. To provide navigation, the atoms near each cross-section are shown in the current density heat-maps as colored circles following the designations in Figures 3a and 4a. In Figures 3b and 4b (the first row of current density plots), the surface is chosen through the inter-ring carbon-carbon bond to provide information about the interactions that govern the coupling between the rings. In Figures 3c and 4c (the second row of current density plots), the surface is chosen through the central phenyl ring to provide information about the interactions that govern the transport through this ring.



**Figure 3.** Ballistic current density through select surfaces of **TPh-H** at 45° and 90° axial torsion. Surfaces are chosen perpendicular to the transport direction, and the current density is therefore plotted as the *z*-component of the current density vector field. (a) Designation of the two surfaces perpendicular to the transport direction and color code of atoms shown in the current density plots. The atom designation is read looking from left to right. (b) Current density through surface b for **TPh-H** at 45° at the Fermi energy, 90° at the Fermi energy, and 90° at +1 eV. (c) Current density through surface c for **TPh-H** at 45° at the Fermi energy, 90° at the Fermi energy, and 90° at +1 eV.



**Figure 4.** Ballistic current density through select surfaces of **TPh-Me** at  $45^{\circ}$  and  $90^{\circ}$  axial torsion. Surfaces are chosen perpendicular to the transport direction, and the current density is therefore plotted as the *z*-component of the current density vector field. (a) Designation of the two surfaces perpendicular to the transport direction and color code of atoms shown in the current density plots. The atom designation is read looking from left to right. (b) Current density through surface b for **TPh-Me** at  $45^{\circ}$  at the Fermi energy,  $90^{\circ}$  at the Fermi energy, and  $90^{\circ}$  at +1 eV. (c) Current density through surface c for **TPh-Me** at  $45^{\circ}$  at the Fermi energy,  $90^{\circ}$  at the Fermi energy, and  $90^{\circ}$  at +1 eV.

At 45° in Figures 3b and 4b, in both TPh-H and TPh-Me, there is clear forward current (red in the heat map) and two nodal planes, one for the  $\pi$ -system of each ring, thus indicating it is  $\pi$ - $\pi$  interactions mediate the electronic coupling between the two rings. At 45° in Figures 3c and 4c both TPh-H and TPh-Me show clear current through the  $\pi$ -system of the central phenyl ring (current density above and below the plane of the ring with a nodal plane in the plane of the ring). As expected, the current is mediated by the  $\pi$ -system throughout these molecules.

At 90° the pattern of the current density is different. Considering first the pattern of the current density at the Fermi energy, in TPh-H, it is clear that the  $\pi$ -system of the outer ring(s) matches the symmetry of the  $\sigma$ -system of the central ring (Figure 3b). That is, the current is mediated by the  $\pi$ -system of the first ring (current density above and below the plane of the ring), the  $\sigma$ -system of the central ring (current density in the plane of the ring, Figure 3c), and again the  $\pi$ system of the third ring. In TPh-Me at the Fermi energy, the current is quite different despite the structural similarity to TPh-H. There are strong backward currents (blue in the heatmap in both Figure 4b,c). We note here that destructive interference effects cannot be seen directly in the current density. The current density strictly only provides us with information about the (here small) current that continues to flow, rather than the current that has been suppressed due to interference. The evidence of destructive interference that can be found in the current density is ring currents that change direction.<sup>68</sup> Thus, while we see strong backward currents in the current density at the Fermi energy for TPh-Me, this is not in and of itself evidence of interference. Instead, we plot the current density at 1 eV above the Fermi energy, as it is clear from the transmission shown in Figure 2 that this energy is on the other side of the minimum transmission for both TPh-Me and TPh-H. The energy dependence of the current density is qualitatively different for the two systems. In TPh-H the

pattern of current density is essentially energy independent, whereas TPh-Me the forward and backward currents (red and blue regions) are reversed for 1 eV versus the Fermi energy. Beyond the qualitative differences in the transmission dips, sharp in the case of TPh-Me versus smooth for TPh-H, these differences in the energy dependence of the current density provides further evidence that the methyl substituents result in destructive  $\sigma$ -interference for these systems. Finally, we verify that it is the  $\pi$ -system that mediates the current in the first and last ring in all the triphenyl systems in Figure S4. Thus, it is the same  $\pi - \sigma - \pi$  interaction through the three rings that mediates the current in both TPh-Me and TPh-H at dihedral angles close to 90°. However, the methyl substituents effectively shut down the  $\sigma$ -transmission in the central ring due to destructive  $\sigma$ -interference. This is the cause of the large transmission difference of the two systems at 90° torsion that is seen in Figure 2.

As a final test of this comprehensive understanding of the conduction suppression in the triphenyl systems, we explore two similar systems where the central ring is partially saturated, thus providing an alternative approach to locally disrupting conjugation. Shown in Figure 5a, Sat-H has one double-bond less in the central ring as two hydrogens have been added to saturate two opposite carbon atoms; likewise, Sat-Me has two extra methyl groups on opposite carbon atoms in the central ring. In these systems there is no continuous  $\pi$ -system through the molecule, and the current will always be mediated to some extent by the  $\sigma$ -system of the central ring. Breaking the conjugation through the molecule in this way has the expected effect. In Figure 5b, the transmission of Sat-H is over an order of magnitude lower than TPh-H at dihedral angles below 45° due to the loss of a continuous  $\pi$ -system. But as the axial torsion of the two molecules approaches 90°, the transmission at the Fermi energy becomes very similar because transport through the  $\pi$ -system of the central ring is suppressed in both cases.

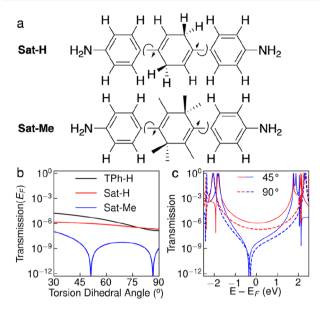


Figure 5. Transmission of triphenyl where two carbons have been saturated; i.e., the central ring is a cyclohexadiene. (a) Chemical structures of the partially saturated amine-functionalized triphenyl (Sat-H) and centrally methylated (Sat-Me) triphenyl. (b) Transmission of Sat-H (red) and Sat-Me (blue) at the Fermi energy as a function of dihedral angle. (c) Transmission as a function of energy for Sat-H (red) and Sat-Me (blue) calculated at 45° (full lines) and 90° (dashed lines) dihedral angles. The transmission of the optimized conformations are shown in Figure S9.

The transmission through **Sat-Me** is quite different. The transmission at the Fermi energy (Figure 5b) is suppressed over the full range of dihedral angles, even when the rings are near-planar. Looking at the full transmission plot in Figure 5c, it is clear that the antiresonance in the transmission is now almost independent of axial torsion. The transmission is always low for **Sat-Me** because the transmission is always mediated by the  $\sigma$ -system of the methylated central ring. The relatively small chemical difference of changing hydrogen to methyl substituents on the central ring, has a major effect on the electron transport properties because of the  $\sigma$ -interference effect.

In this and recent work our focus has been on the effect of methyl substituents, but of course other substituents may provide a similar effect. In Supporting Information part E, we have tested halide and phenyl substituted systems, which all show similar degrees of suppression as that in the methyl-substituted systems. Given that destructive quantum interference effects in both  $\sigma$ - and  $\pi$ -systems can be controlled with chemical design, we foresee that it will be possible to engineer highly insulating and polarizable molecules. Finally, we also note that our current density calculations of phenyl-substituted triphenyl, included in Supporting Information part F, show little indication of current through peripheral nonbonded paths due to through-space conjugation; recent experimental results may need to be reinterpreted.  $^{69,70}$ 

In summary, we have demonstrated that it is possible to simultaneously suppress  $\pi$ - and  $\sigma$ -transmission of biphenyl and triphenyl systems using destructive  $\sigma$ -interference and local disruption of conjugation. The interference effect is weak in hydrogen-substituted systems but can effectively suppress the  $\sigma$ -transmission in methyl-substituted phenyl rings, a substituent that also shifts minimum energy structures toward the

dihedral angles that support maximal suppression of the  $\pi$ -transport. Substituted oligophenyls are promising systems for further research in molecule-based dielectrics, as they indicate that it is indeed possible to realize conjugated insulators at the single-molecule level.

#### ASSOCIATED CONTENT

## Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpclett.0c01727.

Junction structures in xyz format (ZIP)

Computational methods, biphenyl current density, ladder C model, triphenyl current density convergence, effect of substituted triphenyl, hexaphenyl current density, and transmission of optimized conformations (PDF)

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#### Notes

The authors declare no competing financial interest.

### ACKNOWLEDGMENTS

M.H.G. and G.C.S. are grateful for funding from the Danish Council for Independent Research Natural Sciences and the Carlsberg Foundation.

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