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## Tuning quantum interference through molecular junctions formed from cross-linked OPE-3 dimers†

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This study highlights a novel strategy for tuning the electrical conductance of single molecules by cross linking the molecules to form a dimer. By studying the electrical conductance of dimers formed by cross linking OPE monomers, we demonstrate that the appearance of destructive or constructive quantum interference in cross-linked OPE-based dimers is independent of the nature of the molecular cross link. Instead, the type of the interference is controlled by the connectivity to external electrodes and is determined by the presence or otherwise of *meta*-connected phenyl rings in the transport path. This is expected to be an important design feature, when synthesising molecules with cross links of different stiffnesses for thermoelectric energy harvesting, since it shows that the stiffness (and hence phonon transport properties) can be tuned without affecting the nature of the electronic quantum interference.

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Comprehending charge transport through single molecules or self-assembled monolayers (SAMs), is an essential goal of molecular electronics research.<sup>1–3</sup> To realize this target, various strategies have been developed to measure the conductance, current–voltage characteristics, inelastic electron tunneling spectra, transition voltage spectra (TVS), and current-induced local heating of single molecules located between two electrodes.<sup>4–7</sup> Theories of electron transport in single-molecule junctions are built on the concept that electrons passing through a molecule from a source electrode to a drain electrode are phase coherent and that the energy  $E$  of the electron does not change during the passage.<sup>8–10</sup> Consequently, if the source–drain voltage is small, the electrical conductance  $G$  of the molecular junction is given by the Landauer formula  $G = G_0 T(E_F)$ , where  $G_0$  is the quantum of conductance and  $T(E_F)$  is the transmission coefficient, evaluated at the Fermi energy  $E_F$  of the electrodes.

Typically,  $E_F$  lies within the energy gap between the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) of the molecule and consequently certain simplifications arise, which would not occur if  $E_F$  coincided with either of the HOMO or LUMO energy levels. For example, the electrical conductance of OPE-like molecules is measured to be proportional to the cosine-squared of the dihedral angle between neighbouring phenyl rings<sup>8</sup> and theory shows that this occurs, because  $E_F$  lies within the HOMO–LUMO gap.<sup>11,12</sup> We have chosen OPE3 dimers as examples of cross-linked molecules, because their monomer counterparts have been studied extensively in the literature, both for their quantum-interference-derived properties<sup>13–18</sup> and their thermal properties.<sup>19,20</sup>

In the current work, we address the question of how electrical conductances of linear molecules change when a chemical cross link is introduced between pairs of molecules to form dimers. We study the conductance of molecular dimers composed of a series of cross-linked OPE3 molecules, and we derive useful rules for predicting their transport properties, based on identification of the dominant transport paths for electrons passing through the dimers. These transport properties are also rationalised by identifying quantum interference features associated with frontier orbitals, taking into account their energy level spacings and degeneracies.

To form a cross-linked dimer, we started from the well-studied oligophenyleneethynylene (OPE3) molecule, which is known to be a highly conjugated molecular conductor. Next, we

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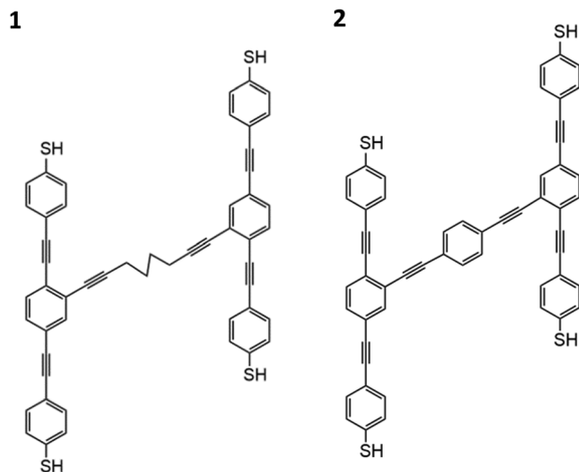


Fig. 1 Two OPE3 molecules cross-linked by two different bridges to form molecules **1** and **2**. Each molecule possesses four thiol terminal end-groups (SH).

linked two OPE3s together by bridges formed from either an alkane chain to create **1** or a highly conjugated phenyleneethynylene chain to create **2**, as shown in Fig. 1.

## Computational methods

All the theoretical simulations were carried out using the density functional (DFT) code SIESTA.<sup>21</sup> The optimised geometries of isolated cross-linked dimers were obtained by relaxing the cross-link dimers until all forces on the atoms were less than  $0.01 \text{ eV } \text{Å}^{-1}$  (for more detail see optimised DFT structures of isolated structures see Fig. S1, ESI†). We used a double-zeta plus polarization orbital basis set, norm-conserving pseudopotentials, the local density approximation (LDA) exchange correlation functional, and to define the real space grid, an energy cutoff of 250 Rydbergs. We also computed results using GGA and found that the resulting structures were comparable,<sup>22–24</sup> with those obtained using LDA. The electronic properties of the cross-linked (C-L), dimers were modelled using a combination of DFT and quantum transport theory. We are interested in computing the electrical conductance of each dimer when a source electrode makes contact with one of the thiols and a second (drain) electrode makes contact with a different thiol. Since these C-L dimers (labelled **1** and **2** in Fig. 2), possess four terminal thiol end-groups, we computed transport properties with 6 distinct pairs of contacts to electrodes. For each of the two C-L dimers, the highest occupied molecular orbital (HOMO), lowest unoccupied orbital (LUMO), and their neighbours (*i.e.*, HOMO–1, LUMO+1 *etc.*), along with their energies are shown in Fig. S3 and S4 (ESI†). In the absence of the bridges, since two isolated OPE3s each possess identical orbital energy levels, the energy levels of the decoupled dimer are doubly degenerate. In the presence of a bridge, each degenerate pair is weakly coupled and the degeneracy is slightly lifted. Consequently, as shown in Fig. S3 and S4 (ESI†), the LUMO and LUMO+1 of each dimer are almost degenerate and similarly the HOMO and HOMO–1 are almost degenerate.

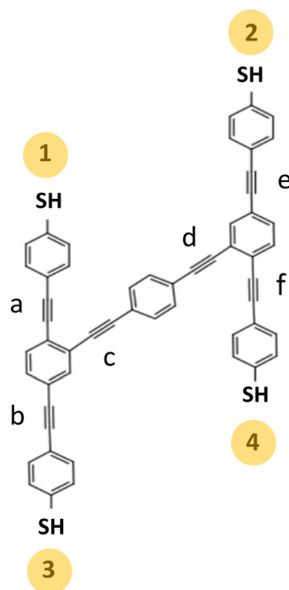


Fig. 2 Four different sulphur atoms can connect to external electrodes as 6 different pairs, namely **1–3**, **2–4**, **1–2**, **3–4**, **1–4**, **2–3**. **1–3** = **2–4**: *para-para* (P–P), **1–2** = **3–4**: *ortho-meta* (O–M), **1–4**: *ortho-ortho* (O–O), and **2–3**: *meta-meta* (M–M).

Our aim is to compute the electrical conductances of these OPE3 dimers (**1–2**), when only two of the four thiols are contacted to two metal electrodes and the current passes from one electrode to another *via* the dimer. As shown in Fig. 2, if the thiols are labelled **1** to **4**, then there are 6 choices of thiol pairs, namely **1–3**, **2–4**, **1–2**, **3–4**, **1–4** and **2–3**. To anticipate the role of quantum interference, we note that the triple bond labelled 'a' is *para* connected to triple bond 'b', and following the discussion in ref. 25 since *para* (and *ortho*) connectivity corresponds to constructive quantum interference (CQI), electrons travelling from thiol **1** to thiol **3** are expected to experience CQI and the corresponding electrical conductance is expected to be high. Similarly, since triple bond 'e' is *para* connected to triple bond 'f', electrons travelling from thiol **2** to thiol **4**, or from thiol **1** to thiol **3** are expected to experience CQI, leading to high electrical conductance. On the other hand, triple bond 'a' is *ortho* connected to triple bond 'c', and triple bond 'e' is *meta* connected to triple bond 'd'. Since *meta* connectivity corresponds to destructive quantum interference (DQI), electrons travelling from thiol **1** to thiol **2**, or from thiol **3** to thiol **4**, or from thiol **1** to thiol **2** are expected to experience DQI, leading to low electrical conductance. It should be noted that electrons travelling from thiol **1** to thiol **2** pass through the *ortho* path 'a'–'c', and the *meta* path 'd'–'e'. Similarly, from thiol **3** to thiol **4**, there is a *meta* path 'b'–'c' and an *ortho* path 'd'–'f'. Since the *meta* paths leads to DQI the conductance of molecules connected to electrodes *via* the thiol pairs **1–2** or **3–4** is expected to be low up to this point four thiol pairs are explored **1–3**, **2–4**, **1–2**, **3–4** and the predicted trend is either CQI (**1–3** and **2–4**) or DQI (**1–2** and **3–4**). Next we consider the electrical conductance when electrodes are connected to thiols **1** and **4** or **2** and **3**. Since the **1–4** path includes *ortho* 'a'–'c' and *ortho* 'd'–'f', we



expect it to exhibit CQI and high electrical conductance. In contrast, the 2–3 connectivity to electrodes involves the *meta*-connected paths ‘e’–‘d’ and ‘c’–‘d’, leading to DQI and therefore low electrical conductance is anticipated.

To examine the above quantum interference predictions, we construct a Hückel (*i.e.*, tight binding, TB) Hamiltonian,<sup>26–37</sup> which captures the dependence of the conductance for the two cross-linked dimers on their connectivity to electrodes. The Hamiltonian matrix comprises of diagonal elements  $H_{jj} = \varepsilon_j$ , which describe the energy  $\varepsilon_j$  of an electron on site  $j$  and nearest neighbour off-diagonal elements  $H_{ij}$ , which describe hopping integrals between neighbouring sites  $i$  and  $j$ . All other matrix elements are set to zero. If all the sites were identical, then the simplest model would be obtained by setting all  $\varepsilon_j = 0$  (which defines the zero of energy) and all nearest neighbour coupling equal to  $-1$ , which sets the energy scale. Such a Hamiltonian is a simple connectivity table, whose entries  $H_{ij}$  are equal to  $-1$  when two atoms  $i$  and  $j$  are connected and are zero otherwise.

When semi-infinite one-dimensional crystalline leads are coupled to sites **1** and **3**, the resulting transmission coefficient  $T(E)$  is shown as the red-solid curve in Fig. 3. The smooth nature of this curve near the middle of the HOMO–LUMO gap ( $E = 0$ ) indicates the presence of CQI. Similarly, coupling to sites **2** and **4**, leads to CQI, as indicated by the smooth nature of the red-dotted curve near  $E = 0$ . On the other hand, when electrodes are linked to sites **1** and **2** or **3** and **4**, the green-solid/-dotted curves are produced. These curves possess sharp dips near  $E = 0$ , signaling the presence of DQI.<sup>38–44</sup> For electrodes connected to sites **1** and **4** the purple-solid curve is produced and exhibits CQI, whereas for electrodes connected to sites **2** and **3**, the light green-solid curve is produced, which possesses a DQI dip. These results for a tight binding model with a single bond  $H_{ij}$  forming the bridge between the monomers, which is the

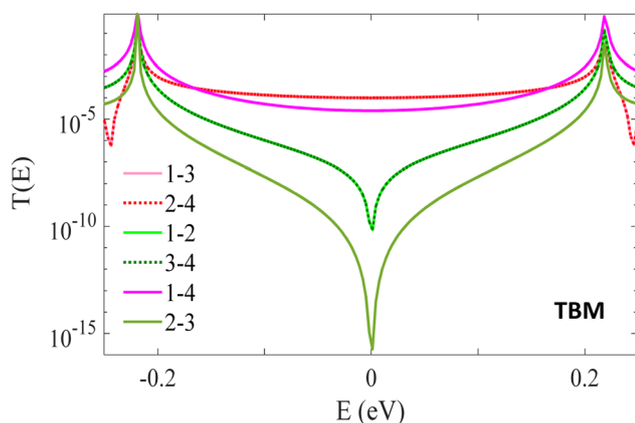


Fig. 3 Transmission coefficients, obtained from the tight binding model (TBM), for 6 different connectivities to electrodes (see Fig. 2) to cross-linked, dimer **1**. Curves for the *para* connected junctions **1–3** and **2–4** (red-solid and red-dotted lines) are identical, as are the curves for *ortho* connected and *meta* connected junctions **1–2** and **3–4** (green-solid and green-dotted lines). Transmission coefficients of the *ortho*–*ortho* connected and *meta*–*meta* connected junctions **1–4** and **2–3** are purple-solid and light green-solid lines. Note: TBM detail, the coupling parameter  $\gamma = -1$  and on energy sites of carbon and sulphur are  $\varepsilon_C = 0$  and  $\varepsilon_S = 0$ .

simplest model of **1**, in which an alkane chain bridges the dimer. Tight binding transmission coefficients of cross-linked dimers **1** and **2** demonstrate that the bridging linker has no influence on the presence of CQI or DQI in the transmission curves, because the two dimers exhibit roughly the same curves for the 6 different pairs of contacts, as demonstrated in Fig. S5 and S6 (ESI†).

To compare with the TBM results we performed DFT transport simulations for the two C–L dimers, each with 6 different pairs of connectivities to electrodes. To simulate the likely contact configuration during a break-junction experiment, we employed gold metal electrodes constructed from 6 layers of Au (111), each containing 30 gold atoms and further terminated with a pyramid of gold atoms. After relaxing each molecular junction, we calculated the transmission coefficient for each of the two molecules shown in Fig. 1, using the Gollum quantum transport code<sup>36</sup> (for more details, see DFT-based transport simulations section in the ESI†).

Fig. 4, shows zero bias transmission coefficients  $T(E)$ , obtained from density functional theory, for 6 different electrode connectivities to dimer **1**. This figure displays 6 transmission functions; three of them show CQI, while the other three show DQI. In agreement with the tight binding results and the above discussion, these show that DQI arises when the transport path contains one of more *meta* connected phenyl rings.

TBM results are in qualitative agreement with the DFT transmission coefficients of the studied molecules, as shown in Fig. S5 of the ESI.† Both TBM and DFT approaches demonstrate that in the presence of the 1–3, 2–4 and 1–4 connectivities, there is no signature of a DQI. However, the same dimers with 1–2, 3–4 and 2–3 connectivities switch from CQI to DQI regardless to the bridging linker chemical structure. The main qualitative difference between the DFT results and the TBM results arise from the fact that the tight-binding model is a nearest-neighbour bipartite lattice, in which atoms can be labelled, such

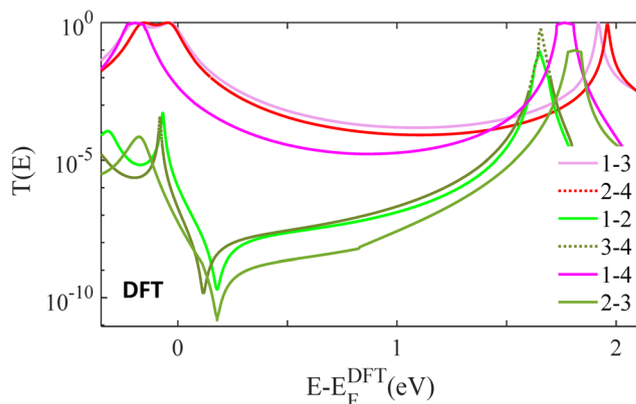


Fig. 4 Zero bias transmission coefficients  $T(E)$ , obtained from density functional theory (DFT), in 6 different possible pairs contact points (see Fig. 2). Cross-linked, dimer **1**: *para* connected junctions **1–3** and **2–4** (red-solid and red-dotted lines), *ortho* connected + *meta* connected junctions **1–2** and **3–4** (green-solid and green-dotted lines), *ortho* connected + *ortho* connected and *meta* connected + *meta* connected junctions **1–4** and **2–3** (purple-solid and light green-solid lines).



that odd numbered atoms interact with even numbered atoms only and *vice versa*. This chiral symmetry guarantees that the transmission coefficient is a symmetric function of the electron energy  $E$ . In the DFT simulation, this symmetry is not present, because next-nearest-interactions are present. The DFT results (S7–S20, ESI†) demonstrate that the presence or otherwise of DQI is insensitive to the nature of the bridging linker and confirm that 1–3, 2–4 and 1–4 are high conductance connectivities due to CQI, whereas 1–2, 3–4 and 2–3 are low-conductance connectivities, due to the presence of DQI.<sup>37–42</sup>

In summary, using density functional theory and a tight-binding model, we have demonstrated that the appearance of destructive or constructive quantum interference in cross-linked OPE-based dimers is independent of the nature of the molecular cross link. Instead, the nature of the interference is controlled by the connectivity to external electrodes and is determined by the presence or otherwise of *meta*-connected phenyl rings on the transport path. This is expected to be an important design feature, when synthesising molecules with cross links of different stiffnesses, because it means that the stiffness and hence phonon transport properties can be tuned, whilst maintaining CQI or DQI.

In the literature, and as discussed in a recent textbook,<sup>8</sup> there are many demonstrations of quantum interference effects in monomer molecules, starting with the first demonstration DQI<sup>42</sup> in 2011 and including more recent demonstrations of QI, such as ref. 43–45 switching in ref. 46–49 and QI in graphene junctions.<sup>50–52</sup> In contrast, the present paper breaks new ground, by investigating and demonstrating quantum interference effects in cross-linked dimers. These results have several experimental consequences. For example, when applied for experimental systems based on break junction experiments, where all connectivities would be sampled, our calculations lead to the prediction that conductance histograms with multiple peaks would be measured, corresponding to both CQI (high conductances) and DQI (low conductances). In addition, in a break junction experiment, they predict the appearance of qualitatively distinct pulling curves. In one type of a curve, electrodes will maintain *e.g.*, a 1–3 connectivity and the junction will break cleanly, leading to a single dominant conductance plateau. In another, the junction may jump from a 1–3 connectivity to a 3–2 connectivity before breaking, leading to two distinct plateaus in a pulling curve.

In the case of SAMs, if both thiols 3 and 4 are initially bound to a substrate and a top contact such as graphene is added, then the 1–3, 2–4 connectivities will act in parallel, leading to CQI and high conductance. In contrast, if the staggered conformations in Fig. 1 are realised, then only the 3–2 connectivity will be relevant, leading to CQI and low conductance. This leads to a new strategy for achieving novel memristive switching behaviour in SAMs, in which a functional cross-linker is employed, whose rigidity can be switched in response to an external stimulus, which in turn causes the molecule to switch between a staggered and non-staggered conformation. The associated switching from a high to low electrical conductance, with an on–off ratio of up to 6 orders of magnitude (see Fig. 4)

would be ideal for in-memory, vector-matrix multiplication in deep neural networks.

## Conflicts of interest

There are no conflicts to declare.

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