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Published in:
Journal of Materials Chemistry C

DOI:
10.1039/C8TC06412A

Published: 23/01/2019

Document Version
Peer reviewed version

Link to publication on the UWS Academic Portal

Citation for published version (APA):

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A 2-D π-π dimer model system to investigate structure-charge transfer relationships in rubrene

Sherlyn C. Jing, Callum J. McHugh and Jesus Calvo-Castro

Rubrene (5,6,11,12-tetraphenyltetracene) is undoubtedly one of the best performing organic charge transfer mediating materials, with experimentally determined mobilities up to 40 cm² V⁻¹ s⁻¹. Consequently, there has been increasing interest by means of crystal engineering in trying to generate rubrene-based materials with analogous or even superior conducting properties. Often, experimental measurements are carried out in thin film architectures of these materials, where measured properties can be detrimentally impacted by device manufacture rather than intrinsic charge transfer properties of the material. The latter results in discarding potential good performers. To address these concerns, we report a two-dimensional model system that will allow researchers to predict charge transfer properties of their materials solely requiring the coordinates of the π-π stacking motifs. We envisaged this study to be of significant interest to the increasingly large community of material scientists devoted to the realisation of improved organic charge mediating materials and particularly to those engaged in exploiting rubrene-based architectures.

Introduction

In the last decades, material scientists have devoted their efforts in achieving optimum charge carrier mobilities in small organic conjugated systems. Among the plethora of systems that have been explored, rubrene (5,6,11,12-tetraphenyltetracene) has attracted an increasing interest, primarily due to its exemplar field effect transistor properties, with experimentally determined values at room temperature of ca 20 cm² V⁻¹ s⁻¹ in single crystal organic field effect transistors (SC-OET) and a reported contact-free intrinsic mobility of ca 40 cm² V⁻¹ s⁻¹. Hitherto, three distinct polymorphs of rubrene have been reported, the monoclinic, triclinic and orthorhombic rubrene forms. Whilst high mobilities have been experimentally determined for the orthorhombic form, the triclinic form exhibits values which are one order of magnitude lower and no evidence of semiconducting behaviour has been reported for the monoclinic form. Although the mechanism responsible for these large mobilities is still a matter of significant debate, it is nowadays widely acknowledged that, over and above contributions such as interplanar distance, these remarkable experimental observations are associated to the supramolecular packing exhibited by orthorhombic rubrene, where the monomers self-assemble in a π-π slipped co-facial fashion that conform to one-dimensional stacking motifs along the crystallographic a-axis. This supramolecular arrangement confers anisotropic charge transfer behaviour to rubrene and is further characterised by large structural overlap between the monomers resulting in associated strong wavefunction overlap, critically important in achieving large charge transfer integrals. Minimal intermonomer displacements along the short molecular axis (also referred to as roll angle) are also considered to play a significant role in ensuring large mobilities in rubrene systems due to enhanced wavefunction overlap.

Figure 1. Capped sticks illustration of the crystal lattices for monoclinic (left), orthorhombic (centre) and triclinic (right) rubrene viewed along the a crystallographic axis.

More recently and inspired by these observations, crystal engineering has been exploited with a focus on improving the field-effect properties of rubrene derived materials by carrying out systematic substitutions on both the peripheral rings and the tetracene core that could ultimately result in different intermolecular slips along the long and short molecular axes and...
associated charge transfer integrals. Small conjugated organic materials have been reported to exhibit large sensitivity to small intermonomer displacements. In some cases, the rationale behind such systematic structural alterations has been to bypass the chemical oxidation that is detrimental for charge transfer processes in rubrene and rubrene based systems. Underpinned by the experimentally acquired mobilities in rubrene, it is commonly acknowledged that organic single crystals (OSCs) play a crucial role in maximising the performance in optoelectronic devices due to their higher purity and structural order when compared to crystalline and amorphous thin films. However, most reported experimental mobility data in the literature has been acquired employing thin film architectures where the presence of grain boundaries and defects are known to negatively influence the performance. As a consequence potentially good organic charge transfer mediating materials with charge transfer properties, which can equate or even improve upon existing ones, may be overlooked on the basis of poor preliminary data that might be ascribed to device manufacturing and not intrinsic material properties.

Motivated by these outcomes, in the following we report a comprehensive analysis of dimeric intermolecular interactions and associated charge transfer integrals for all reported rubrene-based single crystal structures that conform to one-dimensional slipped cofacial π-π stacking motifs. Of particular note is the effect that systematic substitution bears on the intermonomer displacements along the long and short molecular axes as well as their effects on disrupting the highly sought-after planarity of the tetracene core. To further the understanding of substitution-induced changes to the intermonomer displacements and the associated effect on the charge transfer properties, we report for the first time a two-dimensional dimeric model system. The model was generated by simultaneously modifying the displacements along the long and short molecular axes and computing the associated intermolecular interactions and charge transfer integrals. Remarkably, the generated model system is able to qualitatively predict observed device mobility data. As a result, we anticipate the work presented herein to be of critical interest to the large community of material scientists devoted to the development of rubrene based materials and more broadly to all material scientists engaged in the engineering of crystalline systems for optoelectronic applications.

**Experimental**

**Intermolecular interactions**

Dimer pair binding energies were all calculated employing Truhlar's density functional M06-2X at the 6-311G(d) level as implemented in Spartan 10 software. Truhlar’s density functional has shown to give good account of non-covalent intermolecular interactions in supramolecular organic conjugated systems. All computed intermolecular interactions were corrected for Basis Set Superposition Error (BSSE) by means of the counterpoise method of Boys and Bernardi.

**Charge transfer integrals**

Transfer integrals for hole (t_h) and electron (t_e) were computed by means of the energy splitting in dimer method for symmetric systems, with all dimer pairs investigated in this work being centrosymmetric. Within the framework of this method, t_h and t_e can be equated to half the splitting between the dimer HOMO/HOMO(-1) and LUMO/LUMO(+1) orbitals respectively. These calculations were carried out using Truhlar's density functional M06-2X at the 6-311G(d) level. We have previously reported17 that whilst the use of wB97XD density functional results in a 32% increase in the computed intermolecular interactions, the effect on the computer charge transfer integrals is negligible.

**Inner-sphere reorganisation energies**

The geometry of neutral (restricted) as well as radical anion/cation (unrestricted) tetracene and rubrene monomers were optimised by means of the density functional B3LYP at the 6-311G(d) level as implemented in Spartan 10 software. In all cases, optimised structures were confirmed by IR analyses that were characterised by the absence of any imaginary modes, hence denoting real equilibria minima. For all radical ion species, S^2 = 0.76, which indicates low spin contamination in all cases. Inner-sphere reorganisation energies on progression from neutral to radical species were calculated by subtracting the energy of the neutral species at its equilibrium geometry to that of the neutral species at the equilibrium geometry of the radical species. In turn, the inner-sphere reorganisation energies on going from the radical species to the neutral were calculated by subtracting the energy of the radical species at its equilibrium geometry to that of the radical species at the equilibrium geometry of the neutral species.

**Two-dimensional dimer model system**

The two-dimensional π-π dimer model system was generated employing two tetracene monomers. The geometry of the tetracene monomers was optimised using the density functional M06-2X at the 6-311G(d) level as implemented in Spartan 10 software.

![Figure 2](image-url)

Figure 2. Illustration of the long (blue) and short (red) intermonomer displacements in the generated two-dimensional model system. Grey filled circles illustrate the location of the single point calculations through the x/y map.

The two tetracene monomers in the dimer pair were mutually aligned in a fully eclipsed fashion (Δx = Δy = 0.0 Å) at the optimum computed interplanar distance Δz = 3.7 Å. The top tetracene monomer was then displaced, whilst fixing the coordinates of the...
bottom monomer, along the long (x) and short (y) axes simultaneously in 0.3 Å increments over a distance of 11.1 and 4.2 Å respectively as illustrated in Figure 2. The generated dimer model system bears more than 11 single point calculations per Å, which allows for the mapping of all substituted rubrene crystal structures reported to date (Table 1). Intermolecular interactions and transfer integrals for hole and electron at each of the 518 geometries of the model system were computed following the method described above.

Results and discussion

Intermolecular interactions and associated charge transfer integrals for reported rubrene based architectures

One-dimensional π-π stacking motifs are desirable charge propagation channels for effective charge transfer in organic semiconductors.\(^\text{14,15,16}\) On searching the Cambridge Structural Database (CSD) for rubrene-based single crystal structures, out of the 31 reported systems, 12 were observed to conform to one-dimensional π-π slipped cofacial stacking architectures (Table 1).\(^\text{1,13,37-41}\) Two substituted rubrene analogues, RAGCUA and RAGDEL, exhibit twisted tetracene cores as a result of the substitutions carried out on 2 of the 4 peripheral rings but were observed to also conform to the desired supramolecular stacking motif.\(^\text{46}\) In the following, we will first focus on the in-depth analysis of those architectures bearing a planar tetracene core and then focus on structures whereby the substitutions result in twisted tetracene cores, whilst maintaining desirable one-dimensional π-π stacking motifs.

Table 1. CSD identifier, measured intermonomer displacements, intermolecular interactions, and charge transfer integrals, \(t_a/\tau_a\) (k mol\(^{-1}\)) for π-π dimer pairs of reported rubrene based crystal structures.

<table>
<thead>
<tr>
<th>CSD identifier</th>
<th>(\Delta(xyz) / \text{Å})</th>
<th>(\Delta E_{CP})</th>
<th>(t_a/\tau_a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AXIDER(^\text{27})</td>
<td>8.43/3.47/3.76</td>
<td>-17.18</td>
<td>0.8/1.1</td>
</tr>
<tr>
<td>CIYXU(^\text{13})</td>
<td>6.21/0.16/3.59</td>
<td>-31.83</td>
<td>13.3/9.0</td>
</tr>
<tr>
<td>CIYXF(^\text{13})</td>
<td>6.37/0.32/3.51</td>
<td>-33.62</td>
<td>14.0/11.1</td>
</tr>
<tr>
<td>CIYYAM(^\text{11})</td>
<td>6.23/0.16/3.63</td>
<td>-29.12</td>
<td>13.0/7.4</td>
</tr>
<tr>
<td>INELUK02(^\text{42})</td>
<td>8.96/4.27/3.44</td>
<td>-10.71</td>
<td>0.4/0.4</td>
</tr>
<tr>
<td>KUSDEJ(^\text{18})</td>
<td>6.26/0.10/3.48</td>
<td>-36.66</td>
<td>14.9/10.6</td>
</tr>
<tr>
<td>MIVDOM(^\text{43})</td>
<td>6.62/0.02/3.59</td>
<td>-32.92</td>
<td>11.5/10.2</td>
</tr>
<tr>
<td>MIVDU5(^\text{41})</td>
<td>6.19/0.05/3.50</td>
<td>-38.93</td>
<td>14.4/9.2</td>
</tr>
<tr>
<td>PIFHIW(^\text{39})</td>
<td>6.32/0.16/3.50</td>
<td>-38.82</td>
<td>14.3/10.1</td>
</tr>
<tr>
<td>RAGCEK(^\text{40})</td>
<td>6.80/0.15/4.20</td>
<td>-38.60</td>
<td>3.8/7.4</td>
</tr>
<tr>
<td>RAGDIP(^\text{40})</td>
<td>7.14/0.33/3.57</td>
<td>-48.38</td>
<td>3.9/4.9</td>
</tr>
<tr>
<td>TOMVUM(^\text{41})</td>
<td>6.12/0.05/3.61</td>
<td>-36.37</td>
<td>12.8/7.0</td>
</tr>
</tbody>
</table>

It was observed that all systems in Table 1, with the exception of INELUK02 and RAGCEK bear planar tetracene cores and are furthermore characterised by similar intermonomer displacements along the long and short molecular axes to those of orthorhombic rubrene (\(\Delta x/\Delta y/\Delta z = 6.18/0.30/3.64 \text{ Å}\)). In the case of AXIDER and RAGDIP, although both exhibit planar tetracene cores, long molecular axis shifts of 8.43 and 7.14 Å were measured, respectively.

It was furthermore observed that in the eight systems exhibiting analogous displacements to those of orthorhombic rubrene (\(\Delta x/\Delta y/\Delta z = 6.18, 0.30 \text{ and } 3.64 \text{ Å}\) respectively), the largest shift along the short molecular axis is 0.32 Å, which ensures large structural overlap, critical in achieving effective wavefunction overlap and large mobilities. Strong intermolecular interactions among the dimer pairs within the one-dimensional stacks are desirable, conferring thermal integrity to the supramolecular architecture.

![Figure 3. Short interatomic contacts involving substituents in the π-π dimer pair of CIYXU (top left) and CIYYAM (bottom left) and crystal lattice of CIYXU (centre) and CIYYAM (right) along the crystallographic b-axis in both cases.](image-url)
Among the various substitutions explored in crystal engineering of organic systems, isosteric replacement of hydrogen for fluorine atoms has attracted increasing interest in rubrene based systems.\(^{38,40,41,43}\) Whilst both hydrogen and fluorine atoms are characterised by similar polarizabilities, their occupied volume is distinctly different.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure4}
\caption{Short interatomic contacts in the crystal lattices of MIVDUS (left) and TOMVUM (right) along the crystallographic a and b-axes respectively.}
\end{figure}

Therefore, the physicochemical properties of systems bearing this isosteric substitution can be critically affected, such as the reversal of the electronegativity of perfluorinated rings when compared to their non-fluorinated analogues. Although inherently weak, the number of intermolecular interactions in which organic fluorine can participate (i.e. C-F---H, C-F---F, C-F---\pi\_e and \pi---\pi) can induce significant structural changes,\(^{46-50}\) such as the cases of diketopyrrolopyrrole based materials.\(^{44,45}\) In rubrene based systems, it would be useful to rationalise the crystal lattice stabilisation in relation to the different fluoride based substitutions. The CIYAB/CIYAM trifluoromethyl substituted analogue, CIYXF does not bear any close interatomic contacts in the crystal lattice involving the -CF\(_3\) groups located at the para positions of the peripheral rings on positions 6 and 12 of the tetracene core. In turn, trifluoromethyl substitutions of the rings at positions 6 and 11 (MIVDUS) leads to close interatomic contacts (2.536 Å) between trifluoromethyl fluorine atoms and electropositive hydrogen atoms at the para positions of peripheral rings (Figure 4). Analogously, close C-F---F-C interatomic interactions at 2.771 Å apart were observed in TOMVUM, which bears fluorine substituents at the para positions of the four peripheral rings. It is anticipated that although systematic substitutions may not lead to close interatomic contacts in \pi---\pi dimer pairs, these substitutions often result in greater lattice stability which is beneficial in preserving the thermal integrity of the one-dimensional stacking motifs.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure5}
\caption{Short interatomic contacts involving substituents in the \pi-\pi dimer pair of AXIDER (top left) and RAGDIP (bottom left) and crystal lattice of AXIDER (centre) and RAGDIP (right) along the crystallographic a-axis in both cases. Both AXIDER and RAGDIP exhibit planar tetracene cores and larger displacements along their long molecular axes (\(\Delta x = 8.43\) and 7.14 Å respectively). In the case of AXIDER there is also a large short molecular axis shift (\(\Delta y = 3.47\) Å). As a result, the computed intermolecular interaction for AXIDER is the second lowest computed (\(\Delta E_{CP} = -17.18\) kJ mol\(^{-1}\)). Interestingly, we compute the largest intermolecular interaction for RAGDIP (\(\Delta E_{CP} = -48.38\) kJ mol\(^{-1}\)), despite its larger intermonomer displacement along the long molecular axis. This finding can be clearly ascribed to the short interatomic contacts between electronegative organic fluorine atoms at the meta and ortho positions of the perfluorinated peripheral rings and electropositive core hydrogen atoms at 2.644 and 2.546 Å respectively, which are enabled at the exhibited intermonomer orientation (Figure 5).}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure6}
\caption{Illustration of the computed Kohm-Sham supramolecular orbitals of orthorhombic rubrene (left) and RAGCEK (right). The remaining two systems in Table 1, namely INELUKO2 and RAGCEK display twisted tetracene cores. The former bears very slipped dimer pairs along both long (\(\Delta x = 8.96\) Å) and short (\(\Delta y = 4.27\) Å) molecular axes, resulting in negligible overlap between the tetracene cores and lowest computed interaction energy, \(\Delta E_{CP} = -10.71\) kJ mol\(^{-1}\). The contributions to the low stabilisation energy in the dimer pair can be attributed to interatomic C-F---F-C contacts at 2.909 Å. In turn, the intermolecular interaction energy computed for RAGCEK is \(\Delta E_{CP} = -38.60\) kJ mol\(^{-1}\).}
\end{figure}
Computed charge transfer integrals for holes and electrons for eight out of the 12 substituted rubrene systems in Table 1 are comparable or even superior to those computed for triclinic (t_s/t_o = 10.9/4.1 kJ mol\(^{-1}\)) and more importantly orthorhombic (t_s/t_o = 12.4/7.5 kJ mol\(^{-1}\)) rubrene forms. It is anticipated that fabrication of SC-OFTs employing these systems is warranted. The four outliers to this observed behaviour are characterised by larger intermonomer displacements and/or twisted tetracene cores, which results in lower wavefunction overlap. In this regard, critical changes to wavefunction overlap and associated computed transfer integrals on progression from orthorhombic rubrene to RAGCEK are observed, despite the negligible effects on the computed intermolecular interactions. These changes are manifest in a decrease in the computed hole transfer integrals whilst the integral for electron transfer remains unaltered (t_s/t_o = 12.4/7.5 and 3.8/7.4 kJ mol\(^{-1}\) for orthorhombic rubrene and RAGCEK respectively), which is remarkable given the small long molecular shift difference (\(\Delta x = 6.18\) and 6.80 Å for orthorhombic rubrene and RAGCEK respectively). These changes in electronic behaviour are associated to the significant decrease in the bonding/anti-bonding character of the ungerade HOMO(-1) and gerade HOMO on progression from orthorhombic rubrene to RAGCEK (Figure 6) and highlight the key role played by small intermonomer displacements on the computed charge transfer integrals (vide infra) and ultimately on potential device performance.

<table>
<thead>
<tr>
<th>CSD identifier</th>
<th>(\Delta(xyz))/Å</th>
<th>(\Delta E_{CP}) (kJ mol(^{-1}))</th>
<th>t_s/t_o</th>
</tr>
</thead>
<tbody>
<tr>
<td>RAGCUA(^{40})</td>
<td>7.69/0.50/4.05</td>
<td>-39.01</td>
<td>7.63/7.89</td>
</tr>
<tr>
<td></td>
<td>9.04/0.05/3.79</td>
<td>-42.98</td>
<td>0.31/1.36</td>
</tr>
<tr>
<td>RAGDEL(^{40})</td>
<td>7.83/0.40/3.77</td>
<td>-37.89</td>
<td>6.58/6.94</td>
</tr>
<tr>
<td></td>
<td>8.74/0.54/3.66</td>
<td>-44.16</td>
<td>0.73/0.01</td>
</tr>
</tbody>
</table>

RAGCUA and RAGDEL are both examples whereby the onedimensional π-π stacking motif is formed by two different sets of dimer pairs arranged in a fashion A-B-A-B (Figure 7). Their monomers are characterised by twisted tetracene cores as a result of the substitutions, which significantly increase intermonomer displacements along the long molecular axis and cause a lessening of the computed hole transfer integrals compared with orthorhombic rubrene. In turn, this larger shift imparts less significance on the electron transfer integrals in dimer pairs A of RAGCUA and RAGDEL, where ambipolar charge transfer properties are predicted. Nonetheless, the significantly lower computed integrals for dimer pairs B for both systems detracts any potential interest on these systems as charge transfer mediating materials based upon their computed electronic behaviour.

In summary, intermonomer displacements in Table 1 illustrate that irrespective of the substitution carried out, most (eight out of 12) systems exhibit shifts (\(\Delta x, \Delta y\) and \(\Delta z\) of 6.18, 0.30 and 3.64 Å respectively), as well as computed intermolecular interactions (\(\Delta E_{CP} = -35.60\) kJ mol\(^{-1}\)) and charge transfer integrals (t_s/t_o = 12.4/7.5 kJ mol\(^{-1}\)) which are comparable or even larger than those computed for orthorhombic rubrene. Accordingly, we anticipate that measurements in single crystals of these rubrene analogues are warranted and of interest towards development of superior alternative to the parent rubrene structure. Along these lines, it is of note that despite the negligible interest attracted by the other two known rubrene polymorphs, the triclinic form is also characterised by a supramolecular architecture where the monomers are stacked in a similar fashion to those of orthorhombic rubrene, with intermonomer displacements \(\Delta x, \Delta y\) and \(\Delta z\) of 5.94, 0.51 and 3.73 Å respectively, and computed intermolecular interactions (\(\Delta E_{CP} = -35.62\) kJ mol\(^{-1}\)) and charge transfer integrals (t_s/t_o = 10.9/4.1) comparable to the ones computed for its orthorhombic counterpart.

**Two-dimensional π-π dimer model system**

The remainder of the paper is devoted to in-depth analysis of intermolecular interactions energies and charge transfer integrals in rubrene-based π-π dimer pairs, as a function of intermonomer displacements, employing a two-dimensional model system formed by two planar tetracene monomers. Although one-dimensional model systems for tetracene have been previously reported,\(^{34,51}\) herein we report for the first time a two-dimensional model system for tetracene/rubrene which allows for a greater in-depth analysis and understanding of their charge transfer properties. This two-dimensional model is important in that it dispels popular misconceptions that short molecular axis displacements are more detrimental to charge transfer than analogous ones along the long molecular axis.

Our choice of tetracene instead of rubrene for the two-dimensional model system is justified on the basis of 1) computed intermolecular interaction energies in single crystal derived π-π dimer pairs of rubrene based systems do not often depend (vide supra) on the substitutions performed on the peripheral rings and 2) the negligible extension of the FMO wavefunction density onto the peripheral rings, irrespective of the substitutions. Furthermore, and to support the choice of tetracene and not rubrene for the calculation of charge transfer integrals in our model system, we went on to calculate inner-
sphere reorganisation energies ($\lambda$) for both systems. On comparing these reorganisation energies for both hole and electron transfer process, it was observed that $\lambda_h > \lambda_e$ for both systems\cite{43,52} and that there is a ca 2 kJ mol$^{-1}$ increase in both $\lambda_h/e$ on going from tetracene to rubrene ($\lambda_h/e = 11.2/16.0$ and 13.2/18.0 kJ mol$^{-1}$ for tetracene and rubrene respectively). Upon analysis of the 2-D model system for intermolecular interactions, it is observed that the potential energy surface (PES) for $\Delta y = 0.0$ Å is characterised by the presence of three clearly identified local minima at ca $\Delta x = 1.5, 3.6$ and 6.0 Å, coinciding with favourable local bond dipole/bond dipole and induced dipole interactions (Figure 8). We observed that whilst most investigated dimer pairs are located around the local minima at ca 6.0 Å, the $\pi$-$\pi$ dimer pair of triclinic rubrene ($\Delta y = 3.54$ Å) is closest to this energetically favourable location on the PES. It is noteworthy to observe that large binding energies are not solely restricted to close short molecular axis displacements in the model system. In fact, close values to that of the global minimum ($\Delta E_{CP} = -42.43$ kJ mol$^{-1}$ at $\Delta x = 1.2$ Å and $\Delta y = 0.6$ Å) were also observed at $\Delta x/\Delta y = 1.2/1.2$ Å ($\Delta E_{CP} = -41.06$ kJ mol$^{-1}$) and $\Delta x/\Delta y = 1.2/1.5$ Å ($\Delta E_{CP} = -39.26$ kJ mol$^{-1}$). The latter highlights that large thermal integrals of the $\pi$-$\pi$ dimer pairs can be obtained at short molecular axis shifts other than $\Delta y = 0.0$ Å.

![Figure 8](image-url) Three-dimensional plot of the computed intermolecular interactions as a function of the simultaneous displacements along the long (x) and short (y) molecular axes. Black filled circles denote the positions of reported rubrene-based systems.

Charge transfer integrals for both holes and electrons were computed for each single point calculation within the model system. Figure 9 illustrates larger computed transfer integrals for hole and electron at ca $\Delta x = 0.3/3.0/0.6/9.3$ and 0.3/2.1/4.5/6.6/8.7 Å respectively, which coincide in both cases with the nodal progressions for HOMO and LUMO wavefunctions. Interestingly, we observed $\Delta x/\Delta y$ regions with potential for ambipolar charge transfer behaviour and clearly distinct propagations of these regions along the short molecular axis for hole and electron transfer integrals. Consistent with the frontier molecular orbitals wavefunctions, propagation of the hole transfer integral along the short molecular axis exhibits a node at ca $\Delta y = 1.8$ Å. These observations reveal that large computed integrals for both hole and electron transfer can also be afforded in dimer pairs characterised by larger short molecular axis displacements (i.e. $\tau_h = 13.0$ Å and $\tau_e = 12.6$ kJ mol$^{-1}$ computed for $\Delta x/\Delta y = 0.0/3.0$ Å) and that dimeric systems exhibiting short molecular axis shifts should not be solely discarded. The large transfer integrals revealed by the generated model system, even for large displacements along the long/short molecular axis are anticipated to broaden the plethora of possibilities in the design, synthesis and uses of substituted rubrene systems as charge transfer mediating materials.

![Figure 9](image-url) Three-dimensional plot of the computed hole (top left) and electron (top right) transfer integrals as a function of the displacements along the long (x) and short (y) molecular axes.

Organic based optoelectronic materials are known for exhibiting dramatic changes on their computed charge transfer integrals upon small (ca 1 Å) intermonomer displacements. Aided by the generated two-dimensional model system, we observe that rubrene based materials also conform to these rules, with complete reversal of the $\tau_h/\tau_e$ on going from $\Delta x/\Delta y = 4.5/0.0$ ($\tau_h = 2.1$ and $\tau_e = 16.8$ kJ mol$^{-1}$) to 5.7/0.0 Å ($\tau_h = 12.8$ and $\tau_e = 1.8$ kJ mol$^{-1}$) as illustrated in Figure 9. In short, the gerade HOMO(-1) and ungerade HOMO supramolecular orbitals exhibit weak anti-bonding and weak bonding behaviour at $\Delta x/\Delta y = 4.5/0.0$ Å respectively.Remarkably, the displacement of 1.2 Å along the long molecular axis lowers/raises the energy of the ungerade HOMO/gerade HOMO(-1) that now becomes HOMO(-1)/HOMO with strong bonding/anti-bonding properties. Similarly, whilst the ungerade LUMO with strong bonding characteristics at $\Delta x/\Delta y = 4.5/0.0$ Å rises its energy at $\Delta x/\Delta y = 5.7/0.0$ Å exhibiting weak bonding behaviour, the gerade LUMO(+1) lowers its energy and exhibits a change in character from strong to weak anti-bonding upon the 1.2 Å displacement along the long molecular axis.

Lastly, we went on to test our generated model system by means of those computed charge transfer integrals for reported rubrene-based systems (vide supra). In all cases but one (RAGDEL), we report a successful agreement in the relative order of $\tau_h/\tau_e$ when computed charge transfer integrals for reported systems are compared to those predicted by the model system. In addition, it is striking that for all systems in Table 1 bar AXIDER, RAGCEK and RAGDIP, the predicted charge transfer integrals by our generated two-dimensional model system were within 10% of those computed for these $\pi$-$\pi$ dimer pairs. Finally, experimental mobilities have been reported for three of the investigated systems, namely KUSDJI ($\mu_{h/e} = 4.83/4.20$ cm$^2$ V$^{-1}$ s$^{-1}$), MVIDOM ($\mu_{h/e} = 0.07/0.01$ cm$^2$ V$^{-1}$ s$^{-1}$) and MIVDUS ($\mu_{h/e} = 0.25/0.12$ cm$^2$ V$^{-1}$ s$^{-1}$), which were observed in all cases to agree with the relative order of $\tau_h/\tau_e$ predicted by our reported model system.
Conclusions

In conclusion, π-π dimer pairs from rubrene-based crystal structures that conform to desirable one-dimensional π-stacking motifs have been investigated in detail and their structure-property relationships extracted in an effort to aid in the crystal engineering of superior alternatives to the parent material and its polymorphs. By means of computed charge transfer integrals, we observe that large computed values are not solely restricted to dimer pairs characterised by negligible displacements along the short molecular axis and that consequently, these systems should not be discarded until experimental mobilities are carefully examined in organic single crystal field effect transistors. We observe that contrary to other organic conjugated systems, a large number of rubrene based architectures exhibit, irrespective of their molecular substitutions, similar long and short molecular axes displacements in their π-π dimer pairs, analogous to those of orthorhombic rubrene (and to a lesser extent than triclinic rubrene). Consequently, their experimental charge transfer properties ought to be investigated. Importantly, we report a two-dimensional π-π dimer system that will allow material scientists devoted to the realisation of rubrene-based charge transfer mediating materials to screen any novel architectures by simply requiring the intermonomer displacements along the long and short molecular axes of π-π dimer pairs that conform to one dimensional stacking motifs. Underpinned by this model system, we anticipate that large hole transfer integrals can also be afforded by supramolecular motifs whereby short molecular axis displacements in the dimer pairs exceed 2 Å. In the case of electron transfer, our results reveal similar findings even for long molecular shifts beyond 6 Å. Along these lines, we envisage a large applicability of the reported model, irrespective of the substitution and intermonomer displacements. Consequently, it is anticipated that this study can play an important role in the design and engineering of next generation rubrene based charge transfer mediating materials and that the reported system will be widely employed in screening novel architectures that conform to desirable structural motifs tailored towards efficient charge transfer.

Notes and references

A 2-D \( \pi-\pi \) dimer model system to investigate structure-charge transfer relationships in rubrene

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A two dimensional \( \pi-\pi \) model dimer system that will enable researchers to predict charge transport properties for rubrene-based materials using their single crystal \( \pi \)-stacking data