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The crystal structure and electronic properties of three novel charge transfer co-crystals TCNQF_n- triphenylene (n = 0, 2, 4)[†]

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The pure phase syntheses of three novel organic charge transfer complexes, obtained by combining the polyaromatic hydrocarbon triphenylene with 7,7,8,8-tetracyanoquinodimethane (TCNQ) and its fluorinated derivatives, are reported together with their crystal structures determined by 3D electron diffraction. The degree of charge transfer is estimated *via* examining intramolecular vibrational mode displacements between neutral TCNQF_n and its complexes with triphenylene using infrared spectroscopy. The direct optical band gaps of 1.87, 1.76 and 1.70 eV for triphenylene–TCNQF_n (n = 0, 2, 4) respectively, are determined *via* diffuse reflectance spectroscopy. The application of advanced structure determination techniques in combination with spectroscopic methods has allowed us to shed light on compelling charge transfer systems and to gain indications for the design of improved electrical organic conductors.

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1 Introduction

Co-crystals are multicomponent molecular systems that consist of two or more chemical species forming a single crystal structure.^{1,2} In particular, organic charge transfer complexes (OCTC) are co-crystals consisting of two chemically distinct organic species, a π -electron donor and a π -electron acceptor, which interact through strong Coulomb and dipolar forces, resulting in a degree of charge transfer (CT). The versatility of organic CT complexes is apparent via the rich array of physical phenomena they exhibit, including room temperature ferroelectricity,³ semiconductivity and superconductivity.⁴ These donor-acceptor (DA) systems have a distinct advantage conductive materials, as over monomolecular materials, in that they offer greater charge mobility potential⁵ and band gap tunability.⁶

The use of DA systems as organic field effect transistors (OFETs)⁷ is common and by creating molecular-level ordered hetero-junctions of segregated or mixed-stack architectures, crystals can be formed that show either n/p type or ambipolar transport respectively.⁸ In mixed-stack systems, the donor and acceptor molecules alternate along the stacking direction

-D-A-D-A-D-, meanwhile, in segregated-stack systems, they pack into columns separately -A-A-A- and -D-D-D-.8,9 Materials of the first category are limited by simple band theory to semiconducting behavior that usually results in lower conductivity compared to segregated-stack systems.¹⁰ On the other hand, mixed-stack OCTC crystals are of interest because of the potential for a new type of mechanism for ferroelectricity, estimated to be about 20 times larger than conventional mechanisms.¹¹ A prominent example in this regard is the tetrathiafulvalene:chloranil, 1:1 complex exhibiting ferroelectricity below 80 K.12 Generally, the electronic properties of OCTCs can be tuned by composition, stoichiometry, temperature or pressure.^{13,14} For example, by increasing acceptor electron affinity via fluorine substitution of hydrogen atoms about the aromatic core, crystal packing can be altered along with respective HOMO and LUMO energies tailoring electronic and optical properties to desired states.15 In recent years the combination of polycyclic aromatic hydrocarbons (PAHs) and halogenated acceptors have attracted considerable attention as functional materials in electronics and photonics.^{5,15-18} PAHs are typical examples of π -electron systems able to donate electrons,¹⁹ while 7,7,8,8-tetracyanoquinodimethane TCNQ is arguably the most investigated acceptor for the construction of CT complexes.²⁰ It has been shown that co-crystals containing PAHs and TCNQ molecules show conducting properties with a relatively small band gap and moderate CT between the chemical components.15,19,21

In this paper we describe novel DA complexes based on the polyaromatic hydrocarbon triphenylene and TCNQF_n,

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where n = 0, 2, 4. The effects of fluorine substitution on the TCNQ scaffold have been previously investigated in terms of molecular packing and electronic properties, in view of the possible tuning of conductivity,¹⁵ however this is the first time triphenylene as a co-former, has been investigated. Due to sub-micron crystalline domains, three-dimensional electron diffraction (3D ED), was employed to solve the crystal structures.²² Variations in the electronic and optical properties were evaluated by the degree of CT, estimated *via* infrared spectroscopy and the direct optical band gaps determined from diffuse reflectance spectroscopy.

2 Experimental

Materials were purchased from Fisher Scientific (triphenylene >99% purity) and Tokyo Chemical Industries (TCNQF_n >98% purity) and used as received with no further purification. Cocrystals were prepared by dissolving equimolar constituents in 1 mL of toluene left at 323.15 K for 72 h before cooling to room temperature under ambient conditions. Solutions were filtered through a 0.22 µm membrane into round bottom test tubes which were cleaned with dry toluene in an attempt to reduce the number of nucleation sites, a needle hole allowed the volatile solvent to escape.

Scanning electron microscopy (SEM) was performed on a JEOL SEM IT300 running an accelerating voltage of 20 kV. Electron diffraction data was recorded with a Zeiss Libra transmission electron microscope (TEM) operating at 120 kV and equipped with a LaB₆ source. 3D ED was performed in STEM mode after defocusing the beam in order to have a pseudo-parallel illumination on the sample, as described in Lanza *et al.*²³ ED patterns were collected in Kohler parallel illumination with a beam size of about 150 nm in diameter, obtained using a 5 μ m C2 condenser aperture. Data were recorded by a single-electron ASI MEDIPIX detector.²⁴ An extremely low dose illumination, corresponding to about 0.01 el Å⁻² s⁻¹, was adopted in order to avoid beam damage.

A needle of each co-crystal was gently crushed and directly loaded on a carbon-coated Cu TEM grid without any solvent or sonication. 3D ED acquisitions were performed by rotating the sample around the TEM goniometer axis in steps of 1°, with total tilt ranges up to 110°. A camera length of 180 mm was used, allowing resolution in real space up to 0.7 Å. After each tilt, a diffraction pattern was acquired and the crystal position was tracked by STEM imaging. During the experiment, the beam was precessed around the optical axis by an angle of 1°. Precession was obtained using a Nanomegas Digistar P1000 device. All data acquisitions were performed at room temperature.

3D ED data were analyzed using the software *PETS*2.0.²⁵ Structure determinations were obtained by standard direct methods (SDM) for triphenylene–TCNQF_n (n = 0, 2) and by simulated annealing (SA) for triphenylene–TCNQF_n (n = 4), as implemented in the software SIR2014.²⁶ Data were treated with a fully kinematical approximation, assuming that I_{hkl} was proportional to $|F_{hkl}|^2$. Least-squares structure refinement was performed with the software *SHELXL* using electron atomic scattering factors.²⁷

Geometry optimizations *via* the *Crystal*17 software package were obtained using the dispersion corrected PBE0 hybrid functional,²⁸ Grimme's empirical energy corrections,²⁹ bielectronic integral truncation parameters (TOLINTEG) 7 7 7 7 14, the 6-31g(d) elemental basis set and utilizing the Pack-Monkhorst method a shrinking factor 4 was used in reciprocal space and to define the sampling of *k* points about the "Gilat net".^{30,31}

The degree of CT was quantified with a powdered sample *via* Fourier-transformed infrared spectroscopy using a Perkin Elmer spectrum 2 FT-IR spectrometer, while direct optical band gap energies were acquired during diffuse reflectance spectroscopy using a Perkin Elmer lambda 650 series spectrophotometer and processed by applying a Kubelka-Munk transformation. Crystal structure visualisations and Hirshfeld surfaces were rendered *via* the CCDC *Vesta* (Ver. 3.3.1) and *CrystalExplorer* (Ver. 17.5) software packages respectively.

3 Results

All the triphenylene–TCNQF_n (n = 0, 2, 4) co-crystals grown, resulted in a needle like morphology (Fig. 1). Up to 8 mm long needles of the scarlet non fluorinated complex contrast the black needles of the fluorinated systems of analogous size. SEM imaging (Fig. 2) revealed that all needle-shaped cocrystals are, in fact, polycrystalline bunches of very thin platelets (Fig. 2c insert). In the case of triphenylene-TCNQ, all 3D ED data delivered a triclinic primitive unit cell with approximate parameters a = 6.8 Å, b = 11.2 Å, c = 17.7 Å and α = 78.4°, β = 83.9°, γ = 72.9°. Such a cell would conveniently host two pairs of molecules. Examination of the main planes of 3D ED reconstructions revealed no extinction features. In all data sets, diffuse scattering was present parallel to c^* . Structure solution of the triphenylene-TCNQ co-crystal was obtained ab initio in space group P1 (2). Ultimately, 30 out of 34 non hydrogen atoms of the structure were spotted ab initio by automatic routines (Fig. 3a and b).

Analysis of this structure show that the co-former molecules are weakly coupled in dimers and form two mixed-stack columns along [100] that are reciprocally rotated by 180 and alternate along [001] (Fig. 3a and b). The molecular



Fig. 1 Optical images of crystalline CT complexes against 5 mm squared graph paper for scale. (a) Triphenylene–TCNQ (b) triphenylene–TCNQF₂ and (c) triphenylene–TCNQF₄.



Fig. 2 SEM micrographs (a–c) and polarized light microscopy (d–f) of triphenylene–TCNQ (a and d), triphenylene–TCNQF₂ (b and e) and triphenylene–TCNQF₄ (c and f).

planes of triphenylene and TCNQ are always parallel to the (100) plane and in particular the TCNQ molecules lie along one of the three benzene fragments of the triphenylene molecule. Intra-stack dimerization of the unique unit group D–A–D is shown to have alternating face-to-face distances of 3.228 Å and 3.559 Å. In more detail, DA columns are repeating along [010] and stabilized by TCNQ interacting with triphenylene by C–H…N contacts along [010] (distances 2.86 Å and 3.04 Å) and along [001] (distance 3.06 Å).



Fig. 3 Intermolecular displacements next to an expanded unit cell view down DA stack axis: triphenylene–TCNQ (a and b), triphenylene–TCNQF₂ (c and d) and triphenylene–TCNQF₄ (e and f). Triphenylene donor molecules are in blue, while TCNQF_n acceptor molecules are in red.



Fig. 4 Percentage contributions to the Hirshfeld surface area for close intermolecular contacts.

Data collected on triphenylene–TCNQF₂ micrometric crystals consistently indicated a primitive monoclinic cell with approximate parameters a = 9.4 Å, b = 7.18 Å, c = 32.2 Å and $\beta = 98.4^{\circ}$.

Such a cell would likely contain four pairs of molecules. Upon inspection of the reciprocal space reconstruction, the extinction rules 0k0: k = 2n and h0l: l = 2n were observed, hence indicating crystallization in the space group $P2_1/c$ (14). The structure of the triphenylene–TCNQF₂ co-crystal was solved with the *ab initio* localization of 30 out of 36 non hydrogen atoms in the asymmetric unit (Fig. 3c and d).

As in the case of the non fluorinated analogue, the triphenylene–TCNQF₂ structure comprises alternating DA molecules (Fig. 3c and d). Dimers are however no longer present, as D–A and A–D distances are equivalent (3.35 Å), in addition to which, TCNQF₂ molecules are slightly rotated and shifted with respect to triphenylene molecules. Interestingly, DA molecules form in columns along [010]. Every second column, layers of DA molecules are kinked of about 42° along [001]. The overall structure is stabilized by lateral C–H…N and C–H…F contacts of 2.43 Å and 2.62 Å.

In the case of triphenylene–TCNQF₄, all 3D ED data sets were consistent with a *C*-centered monoclinic cell with approximate parameters a = 17.8 Å, b = 17.4 Å, c = 9.2 Å and $\beta = 130.3^{\circ}$. This cell would conveniently host four pairs of molecules. A close look on planar cuts from 3D ED reconstructions revealed that reflections *hol*: $l \neq 2n$ were missing or relatively weak.

Table 1 Summary of charge sensitive IR absorption peak displacements. Modes 1–3 track C=C markers and mode 4 tracks C=N markers. Each column tracks the displacement of one peak between each TCNQF_n system with a unique peak (1 cm⁻¹) tracked for (n = 0, 2, 4). Modes for (K)–TCNQF_n were taken from literature¹⁵

	Mode 1	Mode 2	Mode 3	Mode 4
TCNQ	1543			2226
Triphenylene-TCNQ	1543			2219
(K)-TCNQ	1509			
TCNQF ₂	1395	1550	1575	2230
Triphenylene-TCNQF ₂	1385	1547	1573	2220
(K)-TCNQF ₂	1348	1486	1524	
TCNQF ₄	1396	1550	1599	2227
Triphenylene-TCNQF ₄	1389	1548	1594	2226
(K)-TCNQF ₄	1353	1501	1540	



Fig. 5 Triphenylene–TCNQF_n unit cells, (1) $F_n = 0$, (2) $F_n = 2$ and (3) $F_n = 4$ with cell packing (centre) surrounded by asymmetric units of TCNQF_n (red) and triphenylene (black) viewed down the *a*, *b*, *c*, *a**, *b** and *c** cell axis.

Then, structure solution of $TCNQF_4$ -triphenylene was performed by SA using 3D ED data in space group *Cc* (9).

Molecular models for TCNQF₄ and triphenylene were taken from literature and used as separate fragments. TCNQF₄– triphenylene appears an ideal case for SA as both co-former molecules are entirely rigid. Solution revealed that the structure is actually centrosymmetric. Both TCNQF₄ and triphenylene molecules fall on conceivable symmetry elements after an appropriate cell origin shift is applied, the inversion center at 4*d* and the two-fold axis at 4*e*, respectively. The subsequent refinement confirms that the correct space group is C2/c, as only half of each molecule is crystallographically independent.

The structure of triphenylene–TCNQF₄ is made by mixedstack DA dimers along [010] with intra-stack separation of 3.27 Å, separating each other by a distance of 3.40 Å (Fig. 3e and f). The TCNQF₄ molecule lies along one of the three edges of the triphenylene molecule. All molecules lie in parallel planes, but alternating DA dimers are reciprocally rotated of about 60° along [010]. The structure is stabilized by lateral C–H…N contacts with distances 2.66–2.86 Å and by C–F…N distances of 3.20 Å.

A higher degree of CT is structurally dependent upon smaller interplanar distances (denser stacks) and upon the minimization of shift between in-plane molecules within each stack.¹⁹ In this regard assessing the degree of CT exhibited by these materials becomes more complex than just attributing it to fluorination of the acceptor. Any unexpected structural arrangements, noted within the F_n complexes, *i.e.* how the π -stack interaction and geometry of the donor triphenylene is expressed in alternating DA pairs, makes comparisons difficult.

Intermolecular bond analysis *via* the generation of Hirshfeld surfaces using *CrystalExplorer* (Ver. 17.5) for each



Fig. 6 IR absorption spectra identifying negligible charge sensitive mode displacements between TCNQ and triphenylene–TCNQ. Black, red and blue lines represent TCNQ, triphenylene and the co-crystal triphenylene–TCNQ respectively. Inset displays C \equiv N markers at higher wavenumbers.

constituent per complex shows that with the introduction of halogen bonding in the system, a stronger resultant π - π stack network is created as indicated by the increasing C···C interaction as quantified by the Hirshfeld surface analysis (Fig. 4).

This coincides with a suppression of H···H, C···H and H···N interactions as competing C···F and H···F networks are established. Empirically, the degree of CT, ρ , for PAH–TCNQ based complexes has been estimated by extracting a ratio of selected intramolecular bond lengths *via* the Kistenmacher method³² such that $\rho = \frac{c}{b+d} = 0.482e^{-1}$ for the non fluorinated complex (Table 1).

To quantify the degree of CT between these three systems, one can compare the room temperature infrared absorption spectra of triphenylene, TCNQF_n, triphenylene–TCNQF_n and a complex in which the electron donating species has ionized. A blueshift in absorption is associated with the shortening of covalent bonds, C=N and C=C can be compared to a greater blueshift of these markers in ionized systems, such as potassium (K)–TCNQF_n or the radical anions of TCNQF_n, from which a reference can be set for a degree of CT at $\rho = 1e^{-}$.

The ratio of blueshift, between triphenylene–TCNQF_n and (K)–TCNQF_n, can be used to estimate the degree of CT, on the assumption that bond length displacement observed *via* C=C and C=N markers, is linearly proportional to the degree of CT. Deviations in molecular configuration between triphenylene–TCNQF_n cells can not be exclusively attributed to, and therefore assessed by, the degree of CT. Discrepancies are expected due to the unique set of intermolecular forces intrinsic to the structure of each complex presented. Therefore estimates from C=N markers are subject to uncertainty, as the peripheral position of the C=N bonds,



Fig. 7 IR absorption spectra identifying significant charge sensitive mode displacements between $TCNQF_2$ and triphenylene– $TCNQF_2$. Black, red and blue lines represent $TCNQF_2$, triphenylene and the triphenylene– $TCNQF_2$ co-crystal respectively. Inset displays $C \equiv N$ markers at higher wavenumbers.



Fig. 8 IR absorption spectra identifying significant charge sensitive mode displacements between TCNQF₄ and triphneylene-TCNQF₄. Black, red and blue lines represent TCNQF₄, triphenylene and the triphenylene-TCNQF₄ co-crystal respectively. Inset displays C \equiv N markers at higher wavenumbers.

makes them susceptible to inter-stack coupling as depicted in Fig. 5.

As can be seen in Fig. 6, there is only negligible shift of the charge sensitive mode at 1543 cm⁻¹ in TCNQ; which is displaced to 1509 cm⁻¹ in (K)-TCNQ corresponding to $\rho =$ $1e^{-1}$. The C=N mode displacement from 2226 cm⁻¹ in TCNQ to 2219 cm⁻¹ in triphenylene–TCNQ gives $\rho = 0.38e^{-1}$ given the corresponding vibrational mode the TCNQ radical holds at 2208 cm⁻¹. However values of ρ determined *via* markers in peripheral C=N modes are likely inaccurate due to the interacting stack coupling C-H…N contacts present in system. The mode at 1543 cm⁻¹ shows negligible CT for the



Fig. 9 Ultraviolet-visible spectroscopic Tauc plot indicating direct optical band gaps for triphenylene–TCNQ (blue), triphenylene– $TCNQF_2$ (black) and triphenylene– $TCNQF_4$ (red).

complex with both markers, standing in disagreement to the empirical method. Charge sensitive modes of TCNQF₂ at 1395, 1550 and 1575 cm⁻¹ have respective wavelength absorbance shifts compared to (K)–TCNQF₂ of –44, –63 and –50 cm⁻¹. Correspondingly triphenylene–TCNQF₂ observed shifts to 1385, 1547 and 1573 cm⁻¹ yielding ρ = 0.23, ρ = 0.05 and ρ = 0.04*e*⁻ respectively (Fig. 7).

As can be seen in Fig. 8, C=N markers representing $\rho = 1e^{-1}$ correspond to a displacement from TCNQF₄ at 2227 cm⁻¹ to 2194 cm⁻¹ in the TCNQF₄ radical anion,³³ with triphenylene-TCNQF₄'s mode displaced to 2226 cm⁻¹ suggesting $\rho = 0.03e^{-1}$. C=C charge sensitive modes in TCNQF₄ at 1396, 1550 and 1599 cm⁻¹ are displaced in the complex to 1389, 1548 and 1594 cm⁻¹, yielding $\rho = 0.16$, 0.04 and 0.08 e^{-1} respectively, when compared to peak shifts in (K)-TCNQF₄.

Direct optical band gaps for triphenylene–TCNQ, triphenylene–TCNQF₂ and triphenylene–TCNQF₄ were measured as 1.877, 1.761 and 1.704 eV indicating optoelectronic applications at respective wavelengths of 660.54 nm, 704.06 nm and 727.06 nm (Fig. 9). Whether the band gap remains the same at 1.761 eV for n = 2 in the recently reported 2,6-difluoro-7,7,8,8tetracyanoquinodimethane molecule (where flourine is switched from a *-para* to an *-ortho* orientation)³⁴ if cocrystallised with triphenylene, remains to be seen.

Comparing the apparent degree of CT, determined from vibrational mode shifts, with structural analysis of all three complexes, it can be seen that generally, a higher relative displacement reduces the degree of CT. It has been argued that an increase in the degree of CT is structurally dependent with a reduction of the face-to-face molecular displacement and a minimization of the shift between D–A molecules.³⁵ However, in contrast to this, the triphenylene–TCNQF₂ complex, has a relatively high degree of CT which coincides with the greatest average displacement. The triphenylene– TCNQF₂ complex is unique in the three materials presented here in that molecules in the stack along the *c*-axis see a greater difference in molecular overlap on each side of a given molecule (Table 2).

4 Conclusions

We have reported three novel organic CT complexes based on the polyaromatic hydrocarbon triphenylene and $TCNQF_n$ (n =0, 2, 4) with 1:1 stoichiometry. Despite crystallization on sub-micron scale, all three structures were solved using 3D electron diffraction, that already has been successfully used for the determination of nanocrystalline organic cocrystals.36,37 recently In particular, reported acetamidophenol-TCNQ co-cocrystals^{38,39} revealed the possibility of a new family of organic CT complexes, based on acetamidophenol molecules. Estimations of the degree of CT show that it is relatively low, but increases with the degree of fluorination of the acceptor molecule. Furthermore, the optical band gaps of these complexes show that they exhibit semiconducting behaviour. The progressive fluorination of the TCNQ scaffold prior to solution based co-crystallization therefore offers an efficient means of tuning the band gap in this co-crystal system.

Table 2 Converged crystallographic data of triphenylene–TCNQF_{n=0,2}, F4 complex not converged, only optimized to a relative energy minima. CCDC deposition numbers for the experimentally derived structures are included</sub>

	Triphenylene-TCNQ	$Triphenylene-TCNQF_2$	$Triphenylene-TCNQF_4$
Crystallographic information			
CCDC number	2201747	2201748	2201749
Formula	$C_{30}H_{16}N_4$	$C_{30}H_{14}F_2N_4$	$C_{30}H_{12}F_4N_4$
Formula weight	432.47	468.46	504.44
Crystal system	Triclinic	Primitive monoclinic	C-Centered monoclinic
Space group	$P\bar{1}$	$P2_1c$	C2c
a (Å)	6.8(1)	9.4(2)	17.8(4)
b (Å)	11.2(2)	7.18(5)	17.4(3)
c (Å)	17.7(4)	32.2(2)	9.20(18)
α (°)	78.4(5)	90	90
β (°)	83.9(5)	98.4(5)	130.3(5)
γ(°)	72.9(5)	90	90
Cell volume (Å ³)	1261(44)	2156(50)	2173(76)
Temperature (K)	293	293	293
Structure determination information			
Tilt range (°)	110	105	100
Data resolution (Å)	1.0	1.0	1.0
Independent reflections (No.)	1747	1641	1648
R _{int}	21.2	23.8	22.1
Structure refinement information			
Reflections $> 4\sigma$ (No.)	712	955	477
$R1_{4\sigma}$ (%)	41.04	30.66	35.33
Goodness of fit	2.925	1.156	2.461
Optical information			
Band gap (eV)	1.877	1.761	1.704

Conflicts of interest

"There are no conflicts to declare".

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