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Spin Multiplicity and Solid-State Electrochemical Behavior in Charge-Transfer Co-crystals of DBTTF/F4TCNQ

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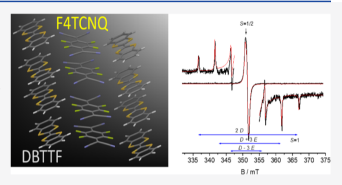
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ABSTRACT: Charge-transfer crystals exhibit unique electronic and magnetic properties with interesting applications. The charge-transfer single crystal formed by dibenzotetrathiafulvalene (DBTTF) together with 2,3,5,6-tetrafluoro-7,7,8,8-tetracyanoquinodimethane (F4TCNQ) presents a long-range ordered supramolecular structure of segregated stacks, with a unitary degree of charge transfer. Thus, the crystal structure is composed of dimerized radical molecules with unpaired electrons. The energy levels and the spin degrees of freedom of this material were investigated by solid-state electrochemistry and electron paramagnetic resonance (EPR) spectroscopy. The electrochemical data, supported by density functional theory calculations, show



how this organic Mott insulator has an electronic gap in the range of hundreds of meV. EPR experiments show the presence of a ground-state S = 1 triplet spin state along with localized S = 1/2 spins. The calculations also predict a ground-state triplet configuration, with the singlet configuration at 170 meV higher energy. DBTTF/F4TCNQ seems to be a candidate material for organic electronic and spintronic applications.

■ INTRODUCTION

Charge-transfer complexes arise from the combination of two neutral molecules with different electron affinities, which, by mutual polarization, induce the transfer of an electron from a donor (D) to an acceptor (A), commonly referred to as a D–A pair. The charge-transfer interaction is crucial in the solid state and leads to the formation of some typical crystal structures, where the donor and acceptor molecules stack with their π -conjugated orbitals facing each other in an alternating fashion. Another common solid-state arrangement is when the donor and the acceptor form parallel stacks with only a π -overlap between molecules having the same molecular structure, that is, only donors or only acceptors. These are referred to as segregated stack charge-transfer co-crystals. Recently, charge-transfer co-crystals have been used to demonstrate new interesting applications.

In the vast literature on these systems, one of the first organic metals, tetrathiafulvalene/2,3,5,6-tetrafluoro-7,7,8,8 tetracyanoquinodimethane (TTF/TCNQ), has emerged as a prototype system and played a central role in the development of the field of organic electronics.⁵ In this case, the TTF and TCNQ molecules stack into segregated stacks with a charge transfer that does not correspond to a complete unit of charge. This is a prerequisite for metallic behavior where a partially filled electronic band contributes to conduction.⁶ When

complete charge transfer occurs, that is, when both the donor and acceptor exchange a complete unit of charge, an ionic crystal is formed by the cation of the donor and the anion of the acceptor.^{7,8} Such systems have not received equally extensive attention in the literature, mainly because their performances as conductors were observed to be poor. In fact, the presence of ionized molecules in the lattice increases the on-site charge repulsion when electrons have to delocalize or move in response to an external field. This gives rise to what is commonly known as a Mott insulator or Mott gap behavior in electrical transport. 10 The unpaired electrons, on these molecular ions, within the crystal lattice open up interesting possibilities when looking at the spin degrees of freedom. This aspect has been recently recognized by Wudl and colleagues.¹¹ Thus, charge-transfer co-crystals combine the high conductivity or semiconductor behavior together with the unpaired or paired (singlet or triplet multiplicity) spins. The charge and

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spin degrees of freedom could be combined toward the design of new organic multiferroic materials. 12,13

In this paper, a charge-transfer co-crystal with peculiar electronic and magnetic properties based on DBTTF/ F4TCNQ is investigated. A detailed structural characterization by single-crystal X-ray diffraction reveals that the material unit cell is composed of homodimers of DBTTF and F4TCNQ which are arranged in segregated stacks. The spin degrees of freedom were investigated using electron paramagnetic resonance (EPR) spectroscopy, which show the coexistence of spin doublet (S = 1/2) and triplet (S = 1) states at room temperature. The material is an electroactive Mott insulator with well-defined frontier orbitals capable of accepting electrons and holes, as demonstrated by the solid-state electrochemistry data. These data were modeled using a density functional theory (DFT) cluster approach, which captures the EPR observations and is able to describe the novel electrochemistry data. Typically, electrochemical-based cyclic voltammetry (CV) measurements are performed for studies in solution; here, to probe the crystal directly and avoid the dissolution of the solid-state material in solution, a new custom-designed setup was adopted. The approach is based on a solid-state electrochemical configuration fully described in the Supporting Information. 14-16 The energies of the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) levels and of the related band gap, E_{BG} , were determined from the electrochemical results. EPR experiments reveal a rather complex picture for the electronic structure, where localized doublet radicals on DBTTF coexist along with the triplet species, found at a much lower concentration. Ab initio DFT calculations (CAMB3LYP/cc-pVTZ) were used to rationalize the experimental evidence in terms of a molecular perspective.

■ EXPERIMENTAL AND THEORETICAL METHODS

Chemicals. DBTTF (99% purity) was purchased from Sigma-Aldrich, and 2,3,5,6-tetrafluoro-7,7,8,8-tetracyanoquino-dimethane (98% purity) was purchased from TCI, UK. The materials were purified using a vacuum sublimation apparatus. Acetonitrile anhydrous, ACN (<99% purity), tetrabutylammonium hexafluorophosphate, TBAPF6 (99% purity), and 5% Nafion 117 solution were obtained from Sigma-Aldrich and were used for electrochemical studies. During all electrochemical analyses, alumina powder was used to renew the electrode surface. All glassware were cleaned by high-purity water (Millipore >20 $\mathrm{M}\Omega$ cm).

Electrochemical Setup. The electrochemical analysis was performed using a PalmSens3 portable potentiostat (PalmSens, Houten, Netherlands). It was interfaced to a tablet or laptop computer, and the software PSTrace 4.6 was used to set the process parameters. A conventional three-electrode cell was used for electrochemical studies of pristine solutions. The working electrode (WE) selected was a glassy carbon electrode (GCE) with a diameter of 3 mm. The counter electrode (CE) was a Pt electrode. The potentials were measured against a Ag/ AgCl/KCl_{sat} reference electrode (RE). A peculiar arrangement was adopted in the case of co-crystals to preserve their solid state: a cylindrical Teflon cell featuring a hole (0.8 cm diameter) in the bottom was used in a vertical configuration, where a glassy carbon plate (GCP) (size: 25 mm × 25 mm) was tightened from below; a Teflon ring was used to ensure no solution leakage from the cell. The cell was rested/leaning on a copper plate as the electrical contact. The GCP was used as

WE, while a Pt wire and Ag/AgCl/KCl_{sat} were used as CE and RE, respectively. CE and RE were placed at the top of the cylinder in contact with the electrolytic solution. The cocrystals were placed over the WE and covered with a Nafion membrane. Then, the membrane was left "to dry" for 5 min, and the electrolytic solution was added. The Nafion membrane prevents direct contact between the solid co-crystal and the electrolytic solution, avoiding the dissolution of the co-crystals but preserving the ion transfer. In fact, if dissolved in solution, the co-crystals would be divided, giving the individual species.

Charge-Transfer Co-crystal Preparation and X-ray Structure. DBTTF/F4TCNQ co-crystals were grown using a horizontal quartz tube under an argon stream. This growth method is widely used for organic materials¹⁷ and recently has been employed by our group and others to grow high-quality organic co-crystals.^{8,18,19} Argon gas was obtained from BOC, UK, with a purity of 6N. The tube was filled with quartz tubes of smaller diameter and 10 cm length, used to harvest the crystals at the end of the growth procedure. The tube, with 150 mg of each material in powder form, was placed in a Carbolite EHC 12/600B three-zone furnace, with the hot zone at 507 K, the center of the furnace at 466 K, and the cold zone at 420 K for the co-crystal DBTTF/F4TCNQ. The tubes were cleaned by ultrasonication in deionized water, 2-propanol, and acetone, sequentially. For structure determination, a single crystal was mounted on a Dual-Thickness MicroMount. Intensity data were collected on a Rigaku Supernova Dual, EosS2 system using monochromated Cu K α radiation (λ = 1.54184 Å). Unit cell determination, data collection, and data reduction were performed using the CrysAlisPro software. The structure was solved with SHELXT and refined by a full-matrix least-squares procedure based on F2(SHELXL-2018/3).²⁰ All non-hydrogen atoms were refined anisotropically. Hydrogen atoms were placed onto calculated positions and refined using a riding model.

EPR Spectroscopy. Room-temperature X-band CW EPR spectra were collected with a super-high Q resonator (ER4122 SHQE, operating at ~9.8 GHz). Low-temperature Q-band CW EPR spectra were collected using the EN 5107D2 Bruker resonator housed in an Oxford CF935 cryostat. Detailed experimental conditions are reported in the figure captions. All samples were finely ground in order to obtain a powder averaged spectrum. Spectra simulations were performed using EasySpin 5.2.28 toolbox 17 within the Matlab 2019b software suite (The Mathworks Inc., Natick, MA).

Calculation Details. Theoretical calculations have been performed within the framework of ab initio linear combination of atomic orbitals using the FireFly, GAMESS, and ORCA suite of programs. Concerning the DBTTF and F4TCNQ single-molecule constituents, calculations were performed at both the CAMB3LYP/6-31G(d) and CAMB3LYP/cc-pVTZ levels of theory for the sake of comparison to evaluate the reliability of 6-31G(d). DFT calculations concerning the crystal were performed by the extraction of a "cluster" of molecules, maintaining the experimental geometrical disposition exactly as obtained by experimental X-ray diffraction data analysis. In the case of the DBTTF/F4TCNQ co-crystal, the data reported in the following are obtained by considering explicitly a two (DBTTF)-by-two (F4TCNQ) cluster.

■ RESULTS AND DISCUSSION

Figure 1 shows the molecular structures and unit cell of DBTTF/F4TCNQ in different orientations. This material has

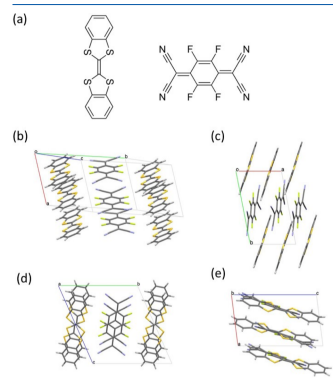


Figure 1. (a) Molecular structures of DBTTF and F4TCNQ. (b) Crystalline structure of DBTTF/F4TCNQ co-crystals, here showing a total of nine molecules with unit cell axes; the number of molecules per unit cell is four. The unit cell axes are labeled. (c-e) Projections of the unit cell along the c, a, and b unit cell axes. Unit cell axes at 170 K are a = 7.4984(2) Å, b = 13.0612(5) Å, and c = 13.6469(6) Å.

a triclinic unit cell with the space group $P\overline{1}$, and the stoichiometry of the co-crystal is 1:1. Our structure compares well with the first X-ray study on this system by Emge et al. Two peculiar features are the segregated stack of donors and acceptors and the dimerization of both DBTTF and F4TCNQ. The dimerization is clearly apparent in the F4TCNQ stack, as can be seen in Figure 1a,b. At 170 K, the interplanar distances between F4TCNQ molecules are 3.223 and 3.416 Å, whereas those between DBTTF molecules are 3.418 and 3.491 Å. At room temperature (300 K), the dimerization persists, and the distances become 3.281 and 3.456 Å for F4TCNQ and 3.477 and 3.526 Å for DBTTF. Therefore, F4TCNQ molecular planes are not only closer to each other on average but also show a larger degree of dimerization, both at 170 K and room temperature.

The crystal symmetry was investigated in a range of temperatures from 80 K to room temperature. The crystal remains triclinic with the space group P1. A detailed analysis of the single-crystal structures at different temperatures will be reported elsewhere.

The EPR powder spectra of the co-crystal were recorded at room temperature at X-band frequency and are shown in Figure 2. The spectrum of DBTTF/F4TCNQ is characterized by a relatively intense absorption line at $g \approx 2.00$ (magnetic field ≈ 352 mT), assigned to S = 1/2 radical species. Low-temperature Q-band experiments allow resolving the aniso-

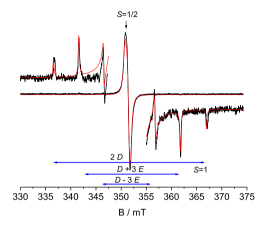


Figure 2. Room-temperature experimental (black curve) and simulated (red curve) CW X-band EPR spectra of DBTTF/F4TCNQ co-crystals in the powder form. The spectral features belonging to the triplet state (S = 1) in the spectrum are indicated by the blue double-sided arrows. The spectra were recorded at 13 dB attenuation and a modulation amplitude of 0.1 mT. The parameters used in the simulation with EasySpin are reported in the Supporting Information.

tropic **g** tensor (Supporting Information, Figure S1), which agrees well with the literature values for the DBTTF cation and with g-factors obtained for this radical from DFT calculations (Supporting Information Table S1). Based on these observations, the recorded doublet state can thus be assigned to isolated radical species, possibly defects, or highly localized spin 1/2 on DBTTF. Preliminary measurements on a single crystal reassure that the radical is not induced by grinding, in keeping with similar observation reported in the case of 5-alkylphenazinium—TCNQ charge-transfer crystals. Superimposed to the narrow doublet signal, a weaker six-line pattern is observed, which can be safely attributed to S=1 species. A convincing simulation of this signal (red curve in Figure 2) was obtained using the spin-Hamiltonian

$$\hat{\mathcal{H}} = \mu_{\rm B} S \mathbf{g} B + D \left(S_z^2 - \frac{2}{3} \right) + E (S_x^2 - S_y^2)$$

and assuming an isotropic g value (g = 2.0038) and the zero-field splitting parameters |D| = 426 MHz and |E| = 46.8 MHz. These values are very similar to those obtained for the triplet ground state observed for 5-alkylphenazinium—F4TCNQ ion pairs. ²³

Figure 3a shows the CV curve recorded for pristine F4TCNQ in AN solution, using tetrabutylammonium hexafluorophosphate (TBAPF6) as the supporting electrolyte and GCE as WE, at 50 mV s⁻¹ (other scan rates are reported in the Supporting Information). The CV curve of F4TCNQ is in good agreement with the data already reported in the literature.²⁴ Two reduction/oxidation peaks are present in the forward-negative/backward-positive potential scan direction; they are of reversible nature with a peak-to-peak separation of about 60 mV. As F4TCNQ is an electron acceptor, the reduction peaks, E_{PC} , are particularly interesting. These are found at 0.543 and 0.062 V and correspond to the formation of the anion F4TCNQ⁽⁻⁾ and dianion F4TCNQ⁽²⁻⁾, respectively, as indicated in Figure 3a. It was observed that F4TCNQ is characterized by chemical stability issues, likely related to the reactivity of the cyano groups with humidity and water traces in the ACN solutions.²⁴ A systematic study on

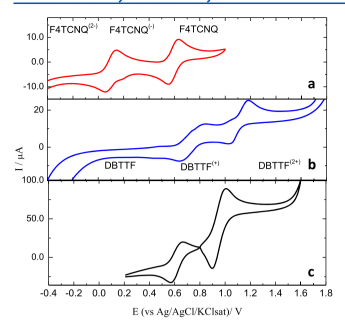


Figure 3. CV curves of the pristine organic compounds in 0.1 M TBAPF6 ACN solution containing (a) 1 mM F4TCNQ and (b) DBTTF powder with Nafion. (c) CV curve of DBTTF/F4TCNQ cocrystals. WE/GCE for (a) and GCP for (b,c), RE: Ag/AgCl/KCl_{sat}, CE: Pt wire. CV profiles were obtained at 50 mV s⁻¹ scan rate.

F4TCNQ stability as a function of time in the adopted experimental conditions was carried out (details are reported in the Supporting Information), and it allowed to ensure a total reliability of our electrochemical measurements.

Figure 3b shows the DBTTF solid-state CV performed on the powdered material. To the best of our knowledge, the DBTTF solid-state CVs have not yet been reported in the literature. Two clear oxidation peaks appear in the forward oxidation scan: the relevant underlying redox processes appear reversible, with a peak-to-peak separation of about 100 mV. The anodic peaks, $E_{\rm PA}$, are 0.85 and 1.183 V and correspond to the formation of the DBTTF mono- and dications. These anodic peaks are at a higher voltage compared to those recorded in solution (Supporting Information, Figure S3), pointing to the solid-state effect in the shift of HOMO levels. It is important to note that the redox potentials for F4TCNQ/F4TCNQ(-) and DBTTF/DBTTF(+) are very close. If we consider that in the solid state, the F4TCNQ LUMO is

predicted to shift by up to 1.3 eV with respect to the vacuum level, 25 the domain of electrical potentials corresponding to the stability of F4TCNQ $^{(-)}$ overlaps with that of the DBTTF monocation. Thus, the electrochemistry of the pure compounds suggests a favorable charge-transfer reaction and the formation of a co-crystal with integer charge transfer, in accordance to the analysis of the charge-transfer degree by IR spectroscopy. 25

Figure 3c shows the CV profiles of solid DBTTF/F4TCNO co-crystals, using GCP as WE, after covering it with the Nafion membrane (details on the solid-state CV method are provided in the Supporting Information). Panel (c) reports two CV curves, which are recorded starting from the open-circuit potential and are separated for the anodic and cathodic scans. The data show two quasi-reversible redox processes (two current peaks are evident in both the cathodic and anodic scans; moreover, they are, within each scan, almost of the same current intensity). It is possible to observe that the CV curve of the cocrystal (i) does not correspond to the sum of the CV curves of the individual compounds, (ii) is characterized by two redox processes occurring in the voltage ranges 0.5-0.7 and 0.9-1.1 V, and (iii) is characterized by larger current in the oxidation scan. As the co-crystal is characterized by a high degree of charge transfer, DB is considered as a cation and F4 as an anion, which means (DBTTF¹⁺/F4TCNQ¹⁻). This is supported by EPR and the arguments presented above on the CV curves of the single materials. The peak at about +1.0 V is assigned to the oxidation of DBTTF molecule to its dication $(DBTTF^{(+)}/F4TCNQ^{(-)})$ oxidation to $DBTTF^{2+}/F4^{TCNQ(-)}$. An almost symmetrical reduction peak is found in the backward curve at about +0.90 V, which is assigned to the relevant quasi-reversible reduction process. In the cathodic CV scan, the reduction peak centered at about +0.57 V corresponds to the reduction of the acceptor or the donor on the surface of the crystal, giving respectively DBTTF/ $F4TCNQ^{(-)}$ or $DBTTF^{(+)}/F4TCNQ^{(2-)}$.

Remarkably, the peak-to-peak current difference in the cathodic CV scan is about 60% smaller than the peak-to-peak current difference in the anodic one. This result suggests that the oxidation is more efficient than the two processes in reduction: the larger current can be due to a more effective ability of the DBTTF stack to exchange its spin 1/2 uncorrelated electrons. If, instead, the oxidation involves the F4TCNQ stacks, their close dimerization distance, cf. Figure 1,

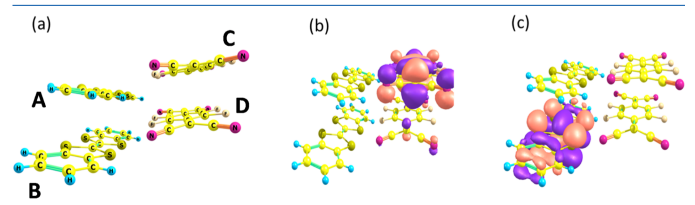


Figure 4. Graphical representation of the DBTTF/F4TCNQ 2×2 superstructures used in the theoretical calculations; atomic coordinates were obtained from the relevant experimental, X-ray, and co-crystal geometries of Figure 1. (a) Labeling of molecules in the cluster. (b) Calculated HOMO and (c) LUMO.

Table 1. DBTTF/F4TCNQ Cluster: LUMO, HOMO, Mulliken Net Charge, and Energy Stability Values for the Singlet, Triplet, and Quintet Electronic State Multiplicity^a

	singlet	triplet	quintet
	vacuum	vacuum	vacuum
LUMO	-4.622162376	-4.65182302	-4.664019096
НОМО	-5.90355662	-5.284764836	-5.000074301
HOMO/LUMO gap	1.281394244	0.632941816	0.336055205
charge DBTTF(A)	0.31376	0.911646	0.912747
charge DBTTF(B)	0.474524	0.873542	0.875194
charge F4TCNQ(C)	-0.44138	-0.90325	-0.85547
charge F4TCNQ(D)	-0.34691	-0.88194	-0.93247
energy tot	-6411.10742583	-6411.113703	-6411.106642
kcal mol ⁻¹	3.939300805	0.0	4.431201
S**2		2.8511	6.1011

 ${}^{a}S^{**2}$ is defined as S(S+1). Refer to Figure 4a for the relevant geometry.

could help charge delocalization. To help these assignments, quantum chemistry calculations were performed.

The theoretical results are presented here, focusing on the (i) band gap and frontier orbital energy variation of DBTTF and F4TCNQ as isolated molecules compared to the values of structures from the co-crystal, (ii) charge distribution within the donor—acceptor partners in the co-crystal, and (iii) the spin multiplicity of the energy levels for the superstructure representative for the co-crystal. For the co-crystal, theoretical calculations were carried out using a cluster strategy. The cluster shown in Figure 4a was considered, which is representative of one unit cell with a total of four molecules: two DBTTF and two F4TCNQ. In both cases, the geometry of the cluster was maintained fixed, according to the experimental X-ray data (see Figure 1).

Table 1 summarizes the theoretical values of HOMO, LUMO, the atomic Mulliken net charge, as well as the total molecular electronic energy. These physical quantities are reported for different values of the total spin in the cluster, that is, a singlet, triplet, or quintet spin state. The charge is factorized for each molecule forming the cluster; thus, the considered unit cell is neutral. Four molecules are labeled according to the scheme in Figure 4a. The degree of intermolecular charge transfer can be made quantitative, exploiting the Mulliken electron density decomposition scheme. Interestingly, a complete single-electron transfer occurs only in the triplet and quintet: -1.8 electrons is the net charge on two F4TCNQ molecules for the triplet and also quintet. The degree of charge transfer in the singlet state is smaller by about 50%, and one unit of charge is shared between two F4TCNQ acceptor molecules, which is inconsistent with the experimental evidence.²⁵ A comparison of the total electronic energies of the singlet, triplet, and quintet states shows that the triplet is the most stable. The singlet and quintet are just 16.32 and 18.41 kJ mol⁻¹ higher in energy, respectively. Thus, the theoretical results suggest a triplet ground state but with the singlet and quintet states that can be thermally populated.

Figure 4b,c shows the frontier molecular orbitals. It is possible to observe that the HOMO is localized on the F4TCNQ molecules, while the LUMO is on the DBTTF. This result is also consistent with the Mulliken net charge variation during simulated oxidation and reduction processes.

Table 2 shows that oxidation leads to a prominent decrease in the negative F4TCNQ charge, which is left almost neutral. In the reduction process, a larger variation is found for

Table 2. DBTTF/F4TCNQ 2 × 2 Cluster, CIF300K UCAM-B3LYP/6-31G*

	neutral	anion	$\begin{array}{c} \text{variation} \\ 0 \rightarrow \text{anion} \end{array}$	cation	variation $0 \rightarrow \text{cation}$
charge DB 1 st	0.911646	-0.0491	-0.9607	0.3477	-0.56395
charge DB 2 nd	0.873542	-0.0768	-0.9503	0.6498	-0.22374
charge DB both	1.785188	-0.1259	-1.9111	0.8781	-0.90709
charge F4 both	-1.785188	-0.8740	0.91119	0.1218	1.90699
charge F4 1 st	-0.90325	-0.4659	0.43735	0.0830	0.98625
charge F4 2 nd	-0.88194	-0.4080	0.47394	0.0388	0.92074

DBTTF, which results almost neutral in the reduced crystal. This analysis helps in assigning the experimentally determined redox processes of Figure 3c, where the reduction is likely to result in DBTTF/F4TCNQ⁽⁻⁾ and the oxidation peak in the formation of DBTTF⁽⁺⁾/F4TCNQ.

CONCLUSIONS

The comparison of data obtained by CV, EPR, and X-ray techniques, supported by theoretical DFT-based calculations, allowed us to obtain a clear picture of the electronic and spin phenomena occurring in the charge-transfer co-crystal formed by DBTTF and F4TCNQ at room temperature.

The degree of charge transfer in DBTTF/F4TCNQ is close to a single unit of charge, which means that the molecular units have unpaired spins in their frontier orbitals. The EPR results give a clear indication of the existence of a dominant DBTTF radical cation species in the solid state, together with a triplet signal due to spin—spin interactions between charges likely on the strongly dimerized F4TCNQ stacks. The solid-state electrochemistry data show reversible cycles, which, on the basis of the theoretical results, can be ascribed to the reduction of DBTTF⁽⁺⁾ in the co-crystal to neutral DBTTF and oxidation of F4TCNQ⁽⁻⁾ to neutral F4TCNQ on the anodic scan. The overall experimental picture is consistent with the theoretical results, which show that for a 2 × 2 cluster the triplet appears to be the most stable electronic state, and the comparison between experimental and calculated spin-Hamiltonian param-

eters allows us to attribute the spin 1/2 signal to a localized radical on the DBTTF stack.

 $E_{\rm BG}$ for the triplet state is in good agreement with the value obtained by elaborating the solid-state CV results. The work highlights DFTTF/F4TCNQ co-crystals as an interesting platform for spintronic devices because of the Mott electronic gap and the richness of spin degrees of freedom that could be controlled with temperature or external magnetic fields.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpcc.1c00020.

Estimation of HOMO and LUMO energy values from electrochemical results; EPR Q-band spectra and relevant DFT-calculated g-tensor; CV curves of F4TCNQ and DBTTF pristine and solid state; co-crystal CV curves as a function of the scan rate; analysis of the chemical stability (probed via UV—vis and CV measurements) of F4TCNQ and DBTTF in solution (PDF)

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Notes

The authors declare no competing financial interest.

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SUPPORTING INFORMATION

Spin Multiplicity and Solid State Electrochemical Behaviour in Charge Transfer Co-crystals of DBTTF:F4TCNQ

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4. Electrochemical behaviour of solid pristine F4TCNQ, DBTTF

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1. Estimation of HOMO and LUMO from E_{ONSET}

Energy of HOMO and LUMO were calculated from E_{ONSET} ¹ as

HOMO=- *Eonset,ox* (4,8 – *Eonset,ox fc*) [eV]

LUMO= - E onset, red (4.8 - Eonset, ox fc) [eV]

2. Q-band EPR spectrum

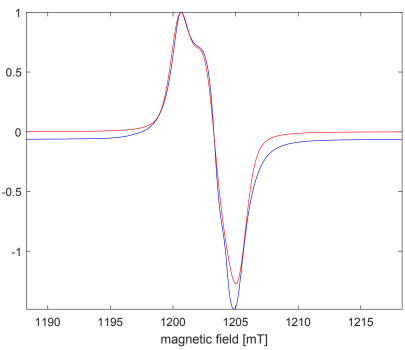


Figure S1. Experimental (blue) and simulated (red) Q-band CW EPR spectrum of DBTTF:F4TCNQ co-crystals recorded at 50 K. The spectrum was recorded at 25 dB attenuation and a modulation amplitude of 0.1 mT. The simulation of the spectrum was obtained using the following spin-Hamiltonian $\hat{\mathcal{H}} = \mu_B S \mathbf{g} B$ for a S=1/2 species with \mathbf{g} tensor $\mathbf{g}=[2.0027\ 2.0059\ 2.0105]$. DFT calculated value \mathbf{g} tensor (DBTTF)= [2.0021370\ 2.0064689\ 2.0106794]

Table S1. Calculated g-tensor values, CAMB3LYP/6-31(d,p) level of theory. DBTTF radical cation, F4TCNQ radical anion species.

	gl	g2	<i>g3</i>
DBTTF	2.0021370	2.0064689	2.0106794
F4TCNQ	2.0030361	2.0028716	2.0023441

3. Electrochemical behaviour of pristine F4TCNQ, DBTTF in solutions

This analysis was performed using PalmSens3 portable potentiostat (PalmSens, Houten, Netherlands) interfaced with the software PSTrace 4.6. Figure S2 and S3 show CV curves recorded for the pristine F4TCNQ and DBTTF organic compounds in 0.1 TBAPF6 AN solution, using a GCE as WE, with a diameter of 3 mm, a Pt electrode as CE and Ag/AgCl/KClsat as RE. CVs were recorded for three different scan rate, 20, 50 and 100 mV s⁻¹, respectively. This set up has been used for all anlysis in solution.

F4TCNQ. Figure S2 shows the CVs obtained for F4TCNQ. The profile is in good agreement with data already reported in the literature²⁻⁴. Two reduction/oxidation peaks are present in the forward-negative/backward-positive potential scan direction, they are of reversible nature with a peak-to-peak separation of about 60 mV, which remains almost constant at different scan rate values. All in all, the first two reduction processes appear fast and reversible. Table S1 describes oxidation-reductive processes of F4TCNQ solution and Table S2 summarizes the F4TCNQ peaks and standard potential values. Table S3 reports LUMO values comparable to those in literature²⁻⁴.

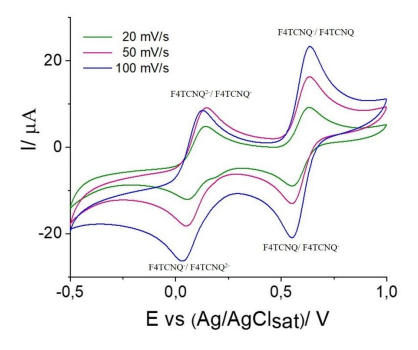


Figure S2. CV voltammogram of 1 mM F4TCNQ in 0.1 M TBAPF6 ACN solution, CE: Pt wire. Green, red and blue CV profiles were obtained at 20, 50, 100 mV s⁻¹ scan rate, respectively.

Table S2. Oxidation-reductive processes of F4TCNQ

$F4TCNQ_{sol} + e^{-}$	⇄	$F4TCNQ_{sol}^{1-}$	Ep ²
$F4TCNQ_{sol}^{1-} + e^{-}$	⇄	$F4TCNQ_{sol}^{2-}$	Ep^{I}

Table S3. E, ΔE and $E_{1/2}$ measured for F4TCNQ at 20, 50, 100 mV $s^{\text{-}1}$

Scan rate	E (V)	$\Delta E(V)$	E1/2 (V)	
20 mV s ⁻¹				
$E_{pa}{}^I$	0.153	0.091	0.108	
$E_{pc}{}^I$	0.062	0.091	0.108	
E_{pa}^2	0.640	0.097	0.597	
E_{pc}^{2}	0.543	0.097	0.397	
50 mV s ⁻¹				
$E_{pa}{}^I$	0.148	0.091	0.105	
$E_{pc}{}^I$	0.057	0.091	0.105	
E_{pa}^{2}	0.640	0.086	0.600	
E_{pc}^{2}	0.554	0.080	0.000	
100 mV s ⁻¹				
$E_{pa}{}^{I}$	0.118	0.096	0.000	
$E_{pc}{}^{I}$	0.032	0.086	0.088	
E_{pa}^{2}	0.645	0.001	0.600	
E_{pc}^{2}	0.554	0.091	0.600	

Table S4. $E_{onset,rid}$ and LUMO energy measured for of F4TCNQ at 20, 50, 100 mV s⁻¹

Scan rate	E _{onset, rid} F4TCNQ /F4TCNQ ⁻ (V)	LUMO (eV)
20 mV/s		
	0.667	-5.167
50 mV/s		
	0.656	-5.147
100 mV/s		
	0.664	-5.154

DBTTF. Figure S3 sets out DBTTF CV profiles. Two oxidation peaks are present in the forward oxidation scan: these redox processes appear fast and reversible, with a peak-to-peak separation of about 60 mV. Table S5 shows the relevant current peak and standard potentials. A third broad current peak is present at a potential around +1.4 V, that shows an irreversible nature (possibly due to the initial stage of a polymerization reaction, which is typical of thiophene derivatives^{5,6}. Few electrochemical studies about DBTTF are present in the literature; in any case the obtained results are in quantitative agreement with the data reported⁷. To the best of our knowledge, this paper reports for the first time CV voltammograms of DBTTF. The values of HOMO present in the Table S6 are in excellent agreement with those reported by Li et al., 2012 ⁸.

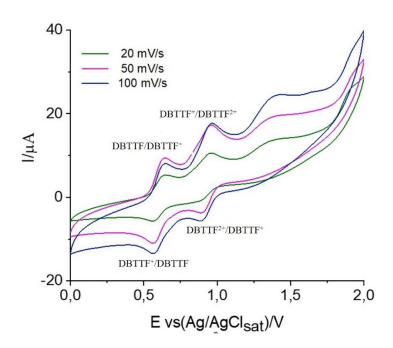


Figure S3. CV voltammograms of 1 mM DBTTF in 0.1 M TBAPF6 ACN solution, CE: Pt wire. Green, red and blue CV profiles were obtained at 20, 50, 100 mV s⁻¹ scan rate, respectively.

Table S5. Oxidation-reductive processes of DBTTF

$DBTTF_{sol}$	⇄	$DBTTF_{sol}^{1+} + e^-$	$E_p{}^l$
$DBTTF_{sol}^{1+}$	⇄	$DBTTF_{sol}^{2+} + e^-$	E_p^2

Table S6. E, $\Delta E,\,E_{1/2}$ to 20, 50, 100 mV $s^{\text{--}1}$ DBTTF

	E (V)	$\Delta E(V)$	$E_{1/2}(V)$	
20 mV s ⁻¹				
$E_{pa}{}^{l}$	0.639	0.074	0.602	
$\dot{E_{pc}}^{l}$	0.565	0.074	0.002	
E_{pa}^{2}	0.957	0.070	0.922	
E_{pc}^2	0.887	0.070	0.922	
$E_{pa}{}^3_{polymerization}$	1.398			
50 mV s^{-1}				
$E_{pa}{}^{l}$	0.648	0.076	0.610	
E_{pc}^{I}	0.572	0.076	0.610	
E_{pa}^{2}	0.967	0.060	0.027	
E_{pc}^{ra}	0.907	0.060	0.937	
$E_{pa}^{\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	1.400			
100 mV s ⁻¹				
$E_{pa}{}^{l}$	0,653	0.001	0.607	
E_{pc}^{I}	0.562	0.091	0.607	
E_{pa}^{2}	0.973	0.076	0.935	
$E_{pc}^{'}$	0.897	0.076	0.933	
$E_{pa}^{3}_{polymerization}$	1.403			

Table S7. $E_{onset,rid}$ and HOMO of DBTTF to 20, 50, 100 mV s⁻¹

Scan rate	$E_{onset, ox} \ DBTTF/DBTTF^+(V)$	HOMO (eV)
20 mV s ⁻¹		
	0.568	-5.068
50 mV s ⁻¹		
	0.553	-5.044
100 mV s ⁻¹		
	0.561	-5.051

Table S8. Summarizes the electrochemical results of the pristine F4TCNQ and DBTTF at 20 mV s $^{-1}$ potential scan rate.

20 (mV s ⁻¹) scan rate	$E_{pa}\left(V\right)$	$E_{pc}(V)$	$\Delta E(V)$	HOMO (eV)	LUMO (eV)
$F4TCNQ_{(sol)} + e^{-} \rightleftharpoons F4TCNQ_{(sol)}^{-}$	0.640	0,543	0.097		-5.167
$F4TCNQ_{(sol)}^{-} + e^{-} \rightleftharpoons F4TCNQ_{(sol)}^{2-}$	0.153	0.062	0.091		
$DBTTF_{(sol)} \rightleftharpoons DBTTF_{(sol)}^+ + e^-$	0.639	0.565	0.074	-5.068	
$DBTTF^{+}_{(sol)} \rightleftharpoons DBTTF^{2+}_{(sol)} + e^{-}$	0.957	0.887	0.070		

^{4.} Electrochemical behaviour of solid pristine F4TCNQ, DBTTF

This analysis was performed using PalmSens3 portable potentiostat (PalmSens, Houten, Netherlands) interfaced with the software PSTrace 4.6. Figure S4 shows CV curves recorded for the solid pristine DBTTF using a cylindrical teflon cell featuring a hole (0.8 cm diameter), the glassy carbon flat as WE, AN solution using TBAPF6 as supporting electrolyte and nafion membrane to prevent direct contact between the solid pristine and the electrolytic solution. CVs were recorded for three different scan rate, 20, 50 and 100 mV s⁻¹, respectively. The voltammogram in solution and solid state are comparable despite different working conditions: two pairs of peaks are always observed indicative of the two reversible reactions to an electron, this time shifted to more positive potentials. This highlights the reliability of solid state electrochemistry on flat cell. the DBTTF polymerisation peak observed in the 1.3 V solution voltammogram has disappeared. From this it is clear the advantage of an analysis in solid state that is the reduction of the phenomena of interference involved to solution.

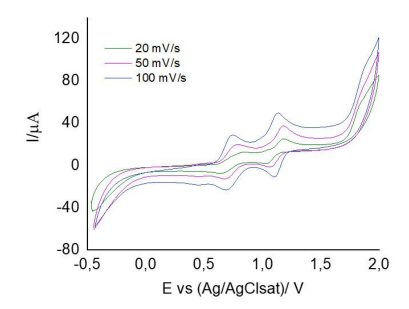


Figure S4. CV of 1 mg DBTTF powder covered with a nafion membrane, 0.1 M TBAPF6 AN solution, glassy carbon plate WE, Pt wire the CE, Ag/AgCl/KCl RE. 20 50 100 mV s⁻¹ scan rate.

Table S9. E, ΔE , $E_{1/2}$ to 20, 50, 100 mV s⁻¹ DBTTF powder

E(V)	$\Delta E(V)$	$E_{1/2}(V)$
------	---------------	--------------

20 mV s ⁻¹			
E_{pa}	0.821	0.152	0.745
E_{pc}	0.669		
$\mathrm{E}_{pa} \ \mathrm{E}_{pc}$	1.186 1.060	0.126	1.123
$\frac{\rm L_{pc}}{50 \rm mV s^{-1}}$	1.000		
30 III V S			
E_{pa}	0.792	0.102	0.741
E_{pc}	0.690	0.102	0.741
E_{pa}	1.197	0.121	1.137
E_{pc}	1.076	0.121	1.137
100 mV s ⁻¹			
E_{pa}	0.740	0.046	0.717
E_{pc}	0.694	0.040	0.717
E_{pa}	1.135	0.025	1.122
E_{pc}	1.110	0.023	1,122

In the case of F4TCNQ, registration of the CV in solid state was not possible. This was due to the p-type dopant effect of F4TCNQ against the nafion layer. This reaction is clearly visible by the sudden bright green colouring of the solution inside the cell. What happens is a doping of the F4TCNQ caused by the strong interaction with the side chains of the nafion.

Ion pairs are formed between the electron of F4TCNQ and the hole in the polymer which increases its conductivity.

5. Stability of pristine F4TCNQ, DBTTF through CVs and UV-Vis techniques

A systematic study on F4TCNQ, DBTTF time dependent stability in the adopted experimental conditions was carried out by recording both CVs and UV-Vis spectra as a function of time.

This analysis was performed using PalmSens3 portable potentiostat (PalmSens, Houten, Netherlands) interfaced with the software PSTrace 4.6, using a GCE as WE, with a diameter of 3 mm, a Pt electrode as CE and Ag/AgCl/KClsat as RE. CVs of F4TCNQ 1 mM in AN solution, 0.1 M TBAPF6 was monitored for 68 h.

The UV-Vis spectrum were obtained using a Jasco V-550 UV-Vis (JASCO Corporation, Tokyo, Japan) interfaced with a computer. UV-Vis spectrum of F4TCNQ 4x10⁻³ mM in AN solution, 0.4 mM TBAPF6 was monitored from 0 to 68 h.

The stability study allowed to ensure a total reliability of our electrochemical measurements, which were run on a much shorter time scale with respect to the start of any degradation reaction.

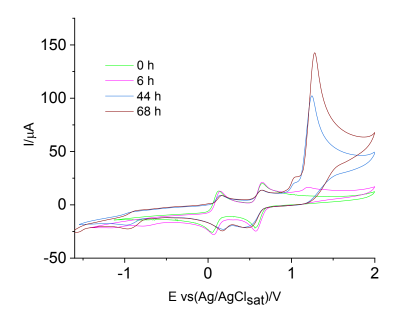


Figure S5. CVs of F4TCNQ 1 mM in AN solution, 0.1 M TBAPF6 monitored from 0 to 68 h

F4TCNQ is characterized by some chemical stability issue, likely related to the reactivity of the cyano groups with water. The Figure S5 sets out CVs of the monitored F4TCNQ solution in a time from 0 to 68 h. This analysis shows the degradation of F4TCNQ over time. This is observable with the formation of a third irreversible peak of oxidation at 1.4 V which corresponds to the rupture of the cyan group resulting in the formation of CN* radicals. The reduction peak moves to lower potentials sign of loss of cyan groups, strong electron-attractors⁵.

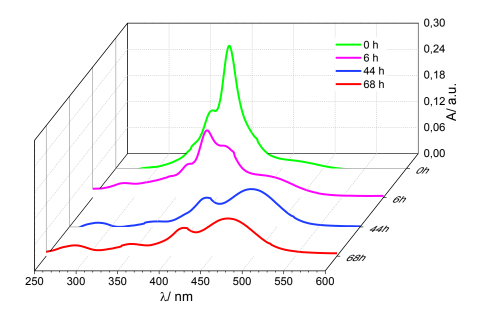


Figure S6. UV-Vis spectrum of F4TCNQ 4x10⁻³ mM in AN solution, 0.4 mM TBAPF6 monitored from 0 to 68 h

The Figure S6 shows the UV-Vis spectrum of the monitored F4TCNQ solution in a time from 0 to 48 h. The data in Figure S7 at 0 time are in good agreement with data already reported in the literature⁹. F4TCNQ degradation is confirmed by UV-Vis analysis where a substantial change to the characteristic absorption band can be detected.

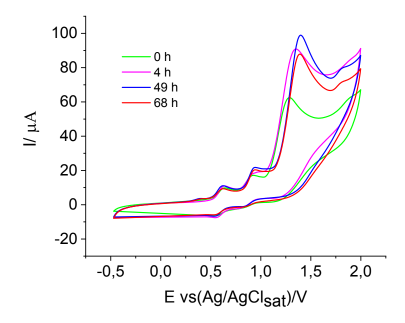


Figure S7. CVs of DBTTF 1 mM in AN solution, 0.1 M TBAPF6 monitored from 0 to 68 h

The Figure S7 shows CVs of DBTTF solution monitored for 68 h. The CVs indicate a considerable increase in the phenomenon of polymerization. This is observable from increase in peak current at a potential around +1.4 V, which basically of irreversible nature¹⁰.

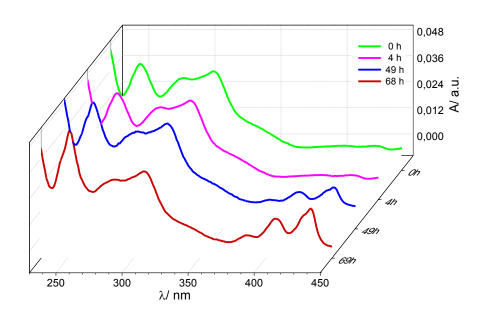


Figure S8. UV-Vis spectrum of DBTTF $4x10^{-3}$ mM in ACN solution, 0.4 mM TBAPF6 monitored from 0 to 68 h

In Figure S8 we show the UV-Vis spectrum of the DBTTF solution in a time interval from 0 to 48 h. The formation of two bands at 410 and 440 nm is probably linked to the polymerization phenomenon.

6. Affinity between donor and acceptor

Donor and acceptor were added simultaneously in the same AN solution, 0.1 M TBAPF6 to demonstrate their strong affinity. The solution was analyzed through a conventional three electrode cell using a GCE as WE, with a diameter of 3 mm, a Pt electrode as CE and Ag/AgCl/KClsat as RE.We observed, the instantaneous formation of a precipitate. This indicates the strong polarity of formed complex. The electrochemical (Fig. S9) and UV-Vis (Fig. S10) response of the materials left in solution seem to be the sum of the response from the two pristine compounds, indicating that some uncomplexed species remain in solution³.

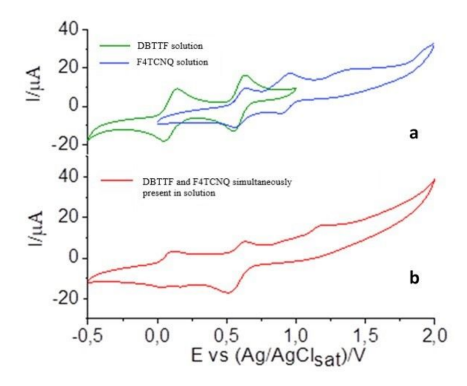


Figure S9. a) CVs of the 1mM F4TCNQ and 1mM DBTTF compounds in 0.1 M TBAPF₆ ACN solution. b) CV of the 0.1 mM DBTTF and 0.1mM F4TCNQ simultaneously in solution. WE: GCE, RE: Ag/AgCl/KCl_{sat}, CE: Pt wire. 50 mV s⁻¹ is the potential scan rate.

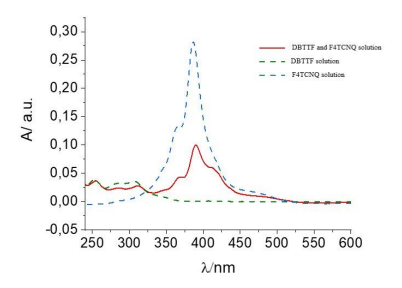


Figure S10. UV-Vis spectrum of green $4x10^{-3}$ mM F4TCNQ solution (green spectrum), $4x10^{-3}$ mM DBTTF solution (blue spectrum) and $2x10^{-3}$ mM DBTTF and F4TCNQ simultaneously in solution (red spectrum)

7. CVs of Co-crystals to 20 50 100 mV s⁻¹ scan rate

The following are the CVs of DBTTF:F4TCNQ co-crystal in the 0.0 to 1.6 V potential range, Figure S11 a), and CVs of DBTTF:F4TCNQ co-crystal recorded starting from the OCP to anodic and cathodic scans, Figure S11 b). CV curves recorded using a cylindrical teflon cell featuring a hole (0.8 cm diameter), the glassy carbon flat as WE, AN solution using TBAPF6 as supporting electrolyte and nafion membrane. CVs were recorded for three different scan rate, 20, 50 and 100 mV s⁻¹, respectively.

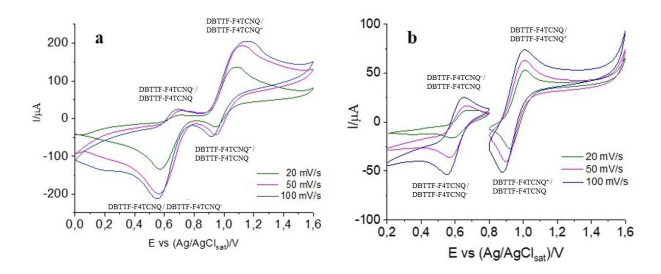


Figure S11. a) CVs of DBTTF:F4TCNQ co-crystal in the 0.0 to 1.6 V potential range $\frac{16}{5}$ CVs of DBTTF:F4TCNQ co-crystal recorded starting from the OCP to anodic and cathodic scans. 20 50 100 mV s⁻¹ scan rate.

Table S10. Oxidation-reductive processes of DBTTF:F4TCNQ co-crystal

$DBTTF:F4TCNQ_{(solid)}$	⇄	$DBTTF:F4TCNQ^{+}_{(solid)} + e^{-}$	E_p^2
$DBTTF:F4TCNQ_{(solid)} +$	⇄	$DBTTF:F4TCNQ_{(solid)}^{-}$	$E_p{}^I$

TableS11. E, ΔE , $E_{1/2}$ to 20, 50, 100 mV s⁻¹ of DBTTF:F4TCNQ co-crystal

Scan rate	E (V)	$\Delta E(V)$	$E_{1/2}(V)$
20 mV/s			
$rac{E_{pa}^{-1}}{E_{pc}^{-1}}$	0.667 0.587	0.080	0.627
$rac{E_{pa}^{-2}}{E_{pc}^{2}}$	1.017 0.925	0.100	0.971
50 mV/s			
$rac{E_{pa}^{-1}}{E_{pc}^{-1}}$	0.661 0.576	0.085	0.618
$rac{E_{pa}^{-2}}{E_{pc}^{2}}$	1.005 0.897	0.108	0.948
100 mV/s			
$rac{E_{pa}^{-1}}{E_{pc}^{-1}}$	0.650 0.551	0.099	0.600
$\frac{E_{pa}^{-2}}{E_{pc}^{2}}$	1.011 0.877	0.134	0.944

 $\textbf{Table S11}. \ Eonset, HOMO, LUMO \ and \ Band \ Gap \ of \ DBTTF: F4TCNQ \ co-crystal \ to \ 20, \ 50, \ 100 \ mV \ s-1$

Scan rate	E_{onset} (V)	HOMO (eV)	LUMO (eV)	Band gap (eV)	
20 mV s ⁻¹					
$DBTTF:F4TCNQ_{(solid)} \rightleftharpoons DBTTF:F4TCNQ_{(solid)}^{+} +$	0.919	-5.249		0.249	
$DBTTF:F4TCNQ_{(solid)} + e^{-} \rightleftharpoons DBTTF:F4TCNQ_{(solid)}$	oi 0.670		-5.000		
50 mV s ⁻¹					
$DBTTF:F4TCNQ_{(solid)} \rightleftharpoons DBTTF:F4TCNQ_{(solid)}^{+} +$	0.900	-5.222		0.228	
$DBTTF:F4_{(solid)} + e^{-} \rightleftharpoons DBTTF:F4_{(solid)}$	0.670		-4.994	5. 22 6	
100 mV s ⁻¹					
$DBTTF:F4TCNQ_{(solid)} \rightleftharpoons DBTTF:F4TCNQ_{(solid)}^{+} +$	0.899	-5.236		0.222	
$DBTTF:F4TCNQ_{(solid)} + e^{-} \rightleftharpoons DBTTF:F4TCNQ_{(solid)}^{-}$	oi 0.670		-5.003	0.233	

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