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A weaker donor shows higher oxidation state upon aggregation⁺

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The charge-transfer between TTFs and I_2 shows that the stronger donor TTF1 is in a cation radical state and the weaker donor TTF2 is neutral in solution, whereas TTF1 exists as a cation radical and TTF2 is dicationic in complexes. The dicationic and neutral states of TTF2 are reversible upon aggregation and solvation.

Charge-transfer (CT) between an electron donor and acceptor plays the pivotal role in supramolecular assembly and creation of conducting materials. There remains a challenge in CT, that is, whether a weaker donor could show a positively charged state higher than a stronger donor through the CT with the same acceptor.

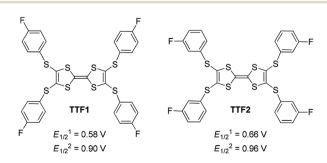
Iodine (I_2) can serve as an acceptor to prepare CT complexes. The CT complex pervlene-iodine is one of the earliest organic conductors.1 Upon gaining one electron from a donor molecule, iodine would form polyiodides,² which show diverse structures and have received growing interest in supramolecular architectures and materials science.3,4 Tetrathiafulvalene (TTF) is an electron donor with three reversible states, $(TTF)^0$, $(TTF)^+$, and (TTF)^{2+,5} TTF derivatives (TTFs) have been widely employed as building blocks for functional materials.6 The CT complexes of I2 and TTFs can be prepared by mixing these two species.7 Because I₂ is not a strong acceptor, TTFs are mainly in the cation radical or partially charged state in CT complexes.8 Ar-S-TTFs are derived from TTF by decorating four arylthio groups onto the peripheral positions (Scheme 1). Ar-S-TTFs can adjust their geometry and electronic state to adapt to a guest molecule,9 and they form CT complexes with various acceptors such as fullerene,10 heteropoly acid,11 and CuBr₂.12

The structures of polyiodides depend on the nature of the counter cations,^{3b} and Ar-S-TTFs can modulate the geometry and electronic state according to the guest. Therefore, the CT complex containing these two flexible components seems promising. Being continuous study on Ar-S-TTFs, herein we report the CT between Ar-S-TTFs (**TTF1** and **TTF2**) and I₂. It is found that a weaker donor **TTF2** carries the positive charge

higher than a stronger donor **TTF1** in their CT complexes with I₂. Meanwhile, the iodine atoms form polyiodides with different structures in CT complexes, *i.e.*, the infinite covalent chain of $[(I_n)^-]_{\infty}$ in **TTF2** complex and 2-D network comprised of $(I_3)^-$ and I₂ in **TTF1** complex.

Electrochemical analysis shows that both **TTF1** and **TTF2** have two reversible redox potentials. The first redox potential $(E_{1/2}^{-1})$ of **TTF2** (0.66 V vs. SCE in CH₂Cl₂) is higher than that of **TTF1** (0.58 V), and the second redox potentials $(E_{1/2}^{-2})$ show similar tendency (Scheme 1). Therefore, as donor molecule, **TTF2** is weaker than **TTF1**. Both donors display weak absorption band at 400–500 nm due to the intramolecular CT transition,⁹ whereas the cation radicals of them show broad absorption at 650–1100 nm.¹¹ For example, electrochemical oxidation of **TTF1** under constant potential of 0.75 V results in an absorption band in this region as proved by the spectroelectrochemical study (Fig. 1a).

By mixing **TTF1** and I_2 in CH₂Cl₂, an absorption band appears at 650–1100 nm (Fig. 1b), which is identical to that observed in the spectroelectrochemistry. The mixture of **TTF1** and I_2 in CH₂Cl₂ shows ESR signal with g = 2.006 (Fig. 1c). Therefore, the CT occurs between **TTF1** and I_2 in CH₂Cl₂ solution, and **TTF1** is at the cation radical state. While CT occurs between **TTF1** and I_2 in CH₂Cl₂, the thin layer chromatography reveals that the neutral **TTF1** remains in solution even though excess I_2 is added (>3 equiv.); this means I_2 cannot completely



Scheme 1 Chemical structures of the Ar-S-TTFs reported herein, along with their first $(E_{1/2}^{-1})$ and second $(E_{1/2}^{-2})$ redox potentials in CH₂Cl₂ recorded *versus* SCE.

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[†] Electronic supplementary information (ESI) available: CCDC 1818732 and 1818736 respectively for (**TTF1**)·(I₃)·(I₂) and (**TTF2**)·(I₅)·(I₂) contain the crystallographic data. The selected crystallographic data are supplied in Table S1 in ESI. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c8ra02956c

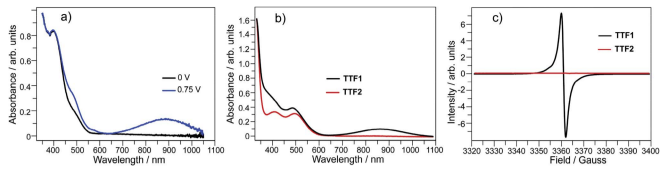


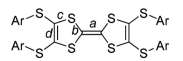
Fig. 1 (a) Spectroelectrochemistry of TTF1 in CH₂Cl₂ ($c = 5 \times 10^{-4}$ mol L⁻¹); (b) UV-Vis absorption spectra and (c) ESR spectra of TTF1 and TTF2 upon adding 3 equivalents of I₂ in CH₂Cl₂ ($c = 1 \times 10^{-5}$ mol L⁻¹).

transform **TTF1** into cation radical. On the other hand, there is no CT between **TTF2** and I_2 in CH_2Cl_2 solution, because the absorbance of (**TTF2**)⁺⁻ is not observed (Fig. 1b) and the mixture of **TTF2** and I_2 is ESR inactive (Fig. 1c).

Although **TTF1** and **TTF2** exhibit the different behaviors upon mixing with I₂ in CH₂Cl₂, they both afford CT complexes with I₂. The CT complexes are obtained as black block-like single crystals by evaporating the CH₂Cl₂/*n*-hexane (v/v, 1 : 1) solution of mixture of **TTF1** (or **TTF2**) and I₂ at room temperature. The compositions of complexes are determined on the basis of single crystal structure analyses to be (**TTF1**)·(I₃)·(I₂) and (**TTF2**)·(I₅)·(I₂).

 $(\mathbf{TTF1}) \cdot (\mathbf{I}_3) \cdot (\mathbf{I}_2)$ crystallizes in the $P\overline{1}$ space group. There are one TTF1 molecule and three pairs of iodine atoms (I1-I2, I3-I4, and I5-I6) in the asymmetric unit. The I3 and I5 locate on the inversion centres. The bond length of central C=C (bond *a* in Scheme 2) on TTF moiety can be used to estimate the charge on TTFs,⁷ *i.e.*, 1.34 Å, 1.39 Å, and 1.45 Å respectively for $(TTF)^0$, $(TTF)^+$, and $(TTF)^{2+}$. Referring Fig. 2a, the central C=C bond length in **TTF1** is 1.39 Å, same to that in $(TTF)^{+.7}$ The site charge (ρ) on TTF moiety also can be estimated *via* an empirical formula $\rho = 6.347 - 7.436\delta$,¹³ where $\delta = (b + c) - (a + d)$, and a, b, c, and d are bond lengths (Scheme 2). The calculated δ -value of TTF1 is 0.721 Å, which gives the site charge on TTF1 to be +1. The iodine atoms (I1-I6) form three tightly connected units, [I1-I2], [I4-I3-I4], and [I6-I5-I6] (Fig. 2c). The I1-I2 bond length (2.74 Å) is identical to that of neutral I₂ (2.74 Å), and the I–I bond lengths (2.91-2.93 Å) in both [I4-I3-I4] and [I6-I5-I6] are very close to that of triiodide (2.90 Å).14 Therefore, the [I4-I3-I4] and [I6–I5–I6] units are intrinsic $(I_3)^-$. These results indicate that TTF1 is at cation radical state in complex, which is reasonable according to the formation of (TTF1)⁺ in solution by mixing TTF1 and I₂.

The **TTF1** molecules are dimerized in complex (Fig. 2b). Within a dimer, there are S \cdots S contacts (3.45–3.53 Å) between TTF cores, and C \cdots S contacts (3.42–3.46 Å) between the



Scheme 2 The bonds (a-d) on Ar-S-TTFs for the estimation of charge ρ .

peripheral sulfur atoms and the phenyls. Meanwhile, the $(I_3)^$ anions and neutral I_2 together form the two-dimensional (2-D) sheet *via* multiple I···I contacts (3.32–3.96 Å). The 2-D sheet is not flat but shows a zig–zag shape along the *b*-axis direction (Fig. 2d). The dimers of **TTF1** are sandwiched by the neighbouring 2-D anion sheets. There are I···S contacts (3.69–3.78 Å) between the anion sheets and **TTF1** dimers. This type of 2-D polyiodide framework is rare in the CT complexes of TTFs and I_2 .¹⁵

 $(TTF2) \cdot (I_5) \cdot (I_2)$ crystallizes in the C2/c space group. The asymmetric unit contains half of TTF2, three tightly connected iodine atoms (I1, I2, I3) with I3 on the 2-fold screw axis, and one isolated iodine atom (I4) at the general position. Referring Fig. 3a, the central C=C bond length (1.45 Å) on TTF moiety is close to that observed in the dicationic salts of Ar-S-TTFs (1.42 Å).¹² The calculated δ value of **TTF2** is 0.573 Å, giving the site charge on TTF2 to be +2. These results firmly prove that TTF2 is dicationic in complex, against the neutral state of TTF2 by mixing it with I₂ in CH₂Cl₂. As shown in Fig. 3b, the I4-I4 bond length (2.79 Å) is close to that of I_2 (2.73 Å), thus the (I4)₂ is a neutral I2. The I1, I2, and I3 atoms form an infinite chain with a periodicity of -[I1-I2-I3-I2-I1]-. Regarding the charge on TTF2, a periodic unit [I1-I2-I3-I2-I1] has a charge of -2. The interatomic distances in [I1-I2-I3-I2-I1] unit vary from 3.04 Å to 3.19 Å, almost identical to those in the infinite polymeric $[(I_n)^-]_{\infty}$ (3.02–3.20 Å).^{3a} Therefore, the present polyiodide chain also would be a $[(I_n)^-]_{\infty}$ polymer, and all the iodine atoms in $[(I_n)^-]_{\infty}$ are partially charged.^{3a} The $[(I_n)^-]_{\infty}$ chains are connected by (I4)₂ through the I···I contacts (3.42 Å) to form a ladder-like structure. The TTF cores and peripheral aryls on TTF2 molecules together form a channel along the longitudinal axis of TTF2 (Fig. 3c), and the channel grows through the $C \cdots S$ contacts (3.34-3.48 Å) between the peripheral sulfur atoms and the phenyls. The $[(I_n)^-]_{\infty}$ chains penetrate into the channel. It is worth noting that $[(I_n)^-]_{\infty}$ chain has not been observed in the complexes comprised of TTFs and polyiodide.

The charged states of **TTF1/TTF2** in CT complexes are further proved by the spectroscopic studies. (**TTF1**)·(I_3)·(I_2) shows a ESR signal with g = 2.009 and (**TTF2**)·(I_5)·(I_2) is ESR inactive (Fig. 4a). This is consistent with crystallographic study, *i.e.*, **TTF1** and **TTF2** are respectively at cation radical and dicationic states. The UV-Vis absorption spectra of both complexes in solid state are distinct from those of neutral **TTF1** and **TTF2** (Fig. 4b). (**TTF1**)·(I_3)·(I_2) shows two absorption bands at the low energy

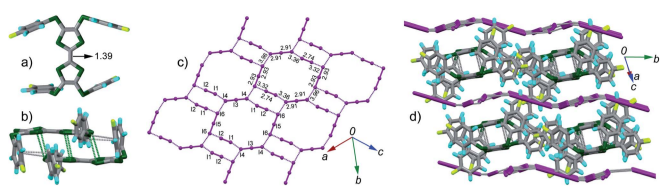


Fig. 2 Crystal structure of complex (TTF1) · (I_3)· (I_2): (a) top view of molecule TTF1 with the central C=C bond length shown in unit of Å; (b) TTF1 dimer with atomic short contacts shown in dashed lines (green for S…S and grey for C…S); (c) anion sheets composed of (I_3)⁻ and I_2 with the I–I bond length and I…I contacts (purple dashed lines) shown; (d) packing structure viewed along the longitudinal axis of TTF1 dimer with the I…I contacts shown in grey dashed lines.

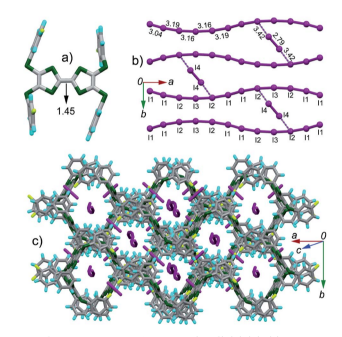


Fig. 3 Crystal structures of complex $(TTF2) \cdot (I_2) \cdot (I_2)$: (a) top view of molecule TTF2 with the central C=C bond length shown in unit of Å; (b) the $(I_3)^-$ anion chain with the I–I bond lengths and I···I contacts (purple dashed lines) shown; (c) packing structure projected along the longitudinal axis of the TTF moiety.

region. The band at 800–950 nm that belonging to absorbance of $(\mathbf{TTF1})^{+*}$. The band at 950–1400 nm ascribable to intermolecular CT transition between the **TTF1** cation radicals in a dimer, *i.e.*, $(\mathbf{TTF1})^{+*} + (\mathbf{TTF1})^{+*} \rightarrow (\mathbf{TTF1})^{2+} + (\mathbf{TTF1})^{0.8a}$ The $(\mathbf{TTF2}) \cdot (\mathbf{I}_{5}) \cdot (\mathbf{I}_{2})$ displays very broad absorption at 500–1400 nm, which is distinct from $(\mathbf{TTF1}) \cdot (\mathbf{I}_{3}) \cdot (\mathbf{I}_{2})$.

As aforementioned, **TTF2** is neutral upon mixing with I₂ in CH₂Cl₂, whereas it is dicationic in (**TTF2**)·(I₅)·(I₂). Moreover, **TTF2** is a donor weaker than **TTF1**, but it shows higher oxidation state in complex. This is against to the criteria for CT between TTF and acceptor, say, the charge on TTF in CT complex depends on the oxidation potential (E_D^{ox}) of TTF and the reduction potential (E_A^{red}) of acceptor.¹⁶ The TTF would be neutral, cation radical, and partially charged under the condition of $E_D^{ox} - E_A^{red} > 0.34$ V, $E_D^{ox} - E_A^{red} < -0.02$ V, and -0.02 V < $E_D^{ox} - E_A^{red} < 0.34$ V, respectively. In the present case, the E_A^{red} of **TTF2** is 0.69 V and the E_A^{red} of I₂ is 0.53 V (Fig. S4 in ESI†). Therefore, **TTF2** would be partially charged in CT complex. One may concern that the increment of charge transfer degree between I₂ and **TTF2** in (**TTF2**)·(I₃)·(I₂) would be attributed to the aggregation of donor and acceptor.

In this regard, the absorption spectra of complexes are studied by dissolving them in CH₂Cl₂. (**TTF1**)·(I₃)·(I₂) shows characteristic absorbance of (**TTF1**)^{+•} in CH₂Cl₂ (Fig. 4c), therefore the charged state of **TTF1** remain the same in solution and CT complex. On the other hand, the charge on **TTF2** is distinctly variated by dissolving (**TTF2**)·(I₅)·(I₂) in CH₂Cl₂. The **TTF2** is

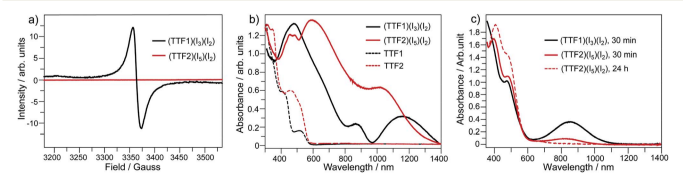
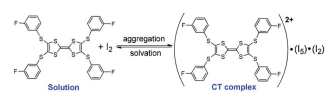


Fig. 4 (a) ESR spectra for the crystalline complexes of (TTF1)·(I_3)·(I_2) and (TTF2)·(I_2); UV-Vis absorption spectra of (TTF1)·(I_3)·(I_2) and (TTF2)·(I_3)·(I_2); UV-Vis absorption spectra of (TTF1)·(I_3)·(I_2) and (TTF2)·(I_3)·(I_2); UV-Vis absorption spectra of (TTF1)·(I_3)·(I_2) and (TTF2)·(I_3)·(I_2); UV-Vis absorption spectra of (TTF1)·(I_3)·(I_2) and (TTF2)·(I_3)·(I_2); UV-Vis absorption spectra of (TTF1)·(I_3)·(I_2) and (TTF2)·(I_3)·(I_2); UV-Vis absorption spectra of (TTF1)·(I_3)·(I_2) and (TTF2)·(I_3)·(I_2); UV-Vis absorption spectra of (TTF1)·(I_3)·(I_2) and (TTF2)·(I_3)·(I_2); UV-Vis absorption spectra of (TTF1)·(I_3)·(I_2) and (TTF2)·(I_3)·(I_2); UV-Vis absorption spectra of (TTF1)·(I_3)·(I_2) and (TTF2)·(I_3)·(I_2) in the (b) solid state, and (c) CH₂Cl₂ solution ($c = 10^{-5}$ mol L⁻¹) after standing under inert atmosphere for 30 min and/or 24 h.



Scheme 3 Reversible process upon aggregation and solvation of $(TTF2) \cdot (I_5) \cdot (I_2)$

reduced from $(\mathbf{TTF2})^{2+}$ to $(\mathbf{TTF2})^{+}$ in 30 min as proved by an absorption band at 700–1050 nm. And, the $(\mathbf{TTF2})^{+}$ disappears to give neutral **TTF2** when the solution is kept for 24 h under inert atmosphere. This means that the retro CT occurs from $[(\mathbf{I}_n)^-]_{\infty}$ to $(\mathbf{TTF2})^{2+}$ upon dissociation of $(\mathbf{TTF2}) \cdot (\mathbf{I}_5) \cdot (\mathbf{I}_2)$, and both anionic and cationic components return to the neutral state. Moreover, the absorbance of $(\mathbf{TTF2}) \cdot (\mathbf{I}_5) \cdot (\mathbf{I}_2)$ can be restored by evaporating the solution to gain solid complex. This process, exchanging the dicationic and neutral states of **TTF2**, is thus reversible upon aggregation and solvation of complex as shown in Scheme 3. These results prove that the dicationic state of **TTF2** in CT complex comes from the aggregation of donor and acceptor.

In summary, the CT between TTF1/TTF2 and I₂ is studied in both solution and solid state. The stronger donor TTF1 turns into cation radical and the weaker donor TTF2 remains neutral upon mixing with I₂ in solution. On the other hand, TTF2 shows an oxidation state (dicationic) higher than that of TTF1 (cation radical) in their CT complexes, which is unusual for CT between TTFs and acceptors. The high oxidation state of TTF2 in complex is due to the aggregation of donor and acceptor. The dicationic and neutral states of TTF2 are reversible upon aggregation and solvation of CT complex. Moreover, the structures of polyiodides in CT complexes can be finely tuned by varying the aryls on Ar-S-TTFs, to give infinite $[(I_n)^-]_{\infty}$ and 2-D network comprised of $(I_3)^-$ and I_2 . Along with previous report, this work further indicates that Ar-S-TTFs show unique feature, i.e., self-modulation of electronic states and molecular geometries according to guest molecules.

Conflicts of interest

There are no conflicts to declare.

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