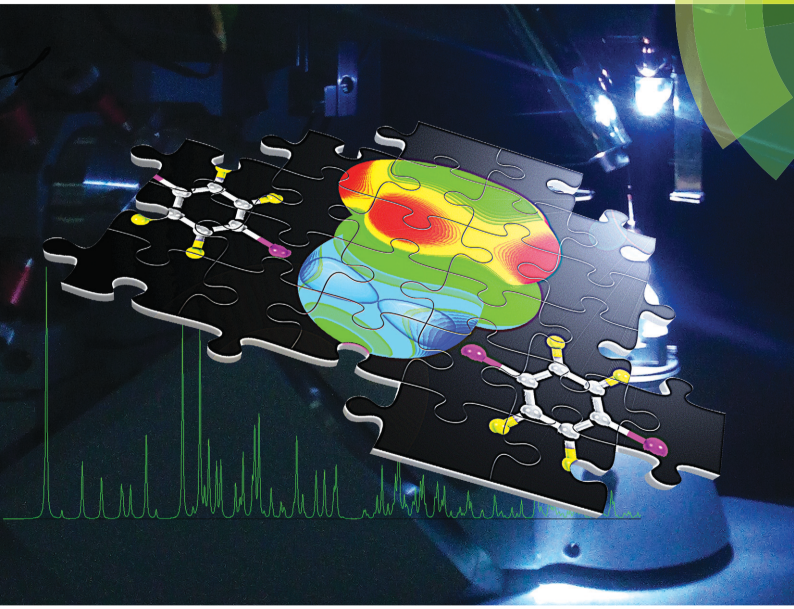
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# Organometallic halogen bond acceptors: directionality, hybrid cocrystal precipitation, and blueshifted CO ligand vibrational band<sup>†</sup>

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Iron cyclopentadienyl carbonyl-halide and -chalcogenolate complexes CpFe(CO)<sub>2</sub>X (X = Cl, Br, I, TePh, SPh) readily afford cocrystals with the bidentate halogen bond donor 1,4-diiodotetrafluorobenzene (*p*-DITFB) under slow evaporation or vapor diffusion conditions. The same microcrystalline [CpFe(CO)<sub>2</sub>TePh](*p*-DITFB) product instantly precipitates upon mixing *p*-DITFB and CpFe(CO)<sub>2</sub>TePh in hexane solution. Supramolecular [CpFe(CO)<sub>2</sub>X(*p*-DITFB)]<sub>*n*</sub> chains in the cocrystals are assembled by halogen bonds (XB) between the electrophilic area of iodine atoms of *p*-DITFB and the nucleophilic area of X in CpFe(CO)<sub>2</sub>X(*p*-DITFB)] cocrystals is explained by the pronounced electron-withdrawing effect of halogen bonding (XB), as supported by DFT calculations. The observed influence of the nature of the XB acceptor (X) on the XB geometry is described in terms of hybridization and electrostatic surface potential (ESP) mapping.

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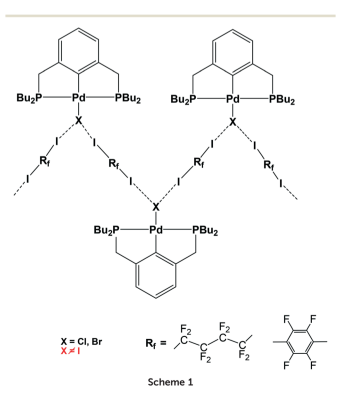
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# Introduction

Halogen bond  $(XB)^1$ -assisted supramolecular self-assembly of transition metal complexes is an attractive research area, developing along with the intensive exploration of XB and other  $\sigma$ - and  $\pi$ -hole<sup>2</sup> bonding interactions. Recent advances in this field have been recently reviewed.<sup>3</sup> In the XB-assisted supramolecular design, the ditopic diiodides of perfluorated alkyl and aromatic hydrocarbons are common XB-donor building blocks used as linkers between XB-acceptor fragments.<sup>4</sup> The same applies to hybrid XB-supported organometallic systems of Pt and Pd affording 1D straight<sup>5</sup> and zigzag chains<sup>6</sup> *via* I–N and I–X halogen bonds, respectively (see Scheme 1).

The formation of the L  $\rightarrow$  M bond results in the strengthening of the X-N halogen bond in M  $\leftarrow$  Py-X-NH<sub>3</sub> (M = Zn, Ag, Co) model systems.<sup>7</sup> However, the M  $\rightarrow$  L back-donation may produce the opposite effect and therefore the L  $\rightarrow$  M and M  $\rightarrow$  L interplay deserves special attention in the case of halogen-bonded X-L-M systems. A comparison of electrostatic surface potential (ESP) maps of halogen atoms linked to electron-withdrawing organic groups with those of metal-bonded halide ligands<sup>8</sup> shows the latter to have XB-acceptor character in their p-belt area, and



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at best a weak XB-donating apical area. The shape of a ligand L p-belt depends upon its anionic character and the extent of mixing of its p- and s-orbitals. Therefore, additional control over the directionality of XB may be achieved through the degree of XB-acceptor hybridization.

Spectroscopic studies including IR, <sup>19</sup>F, <sup>15</sup>N and <sup>13</sup>C NMR, in solution and in the solid phase, are effective in identifying the XB and quantifying its energy.<sup>9–12</sup> IR spectroscopy data for Lewis bases in dilute solution, in the presence of iodoethynyl R–C==C–I XB donors, gives a good correlation between the association constant ( $K_a$ ) and Hammett parameter ( $\sigma_{para}$ ) for diverse substituents R.<sup>13</sup> Using IR spectroscopy, it has also been shown that hydrogen bonding (HB) between a halide and the OH group of CpMn(CO)<sub>2</sub>(hydroxypyridine) induces a redshift of the v(CO) bands (max. 12 cm<sup>-1</sup>) owing to the electron donation from the halide anion *via* a HB–ligand–metal–CO chain.<sup>14</sup> In contrast, the electronwithdrawing effect of an I<sub>2</sub> molecule on the iodido ligand in [(Me<sub>5</sub>C<sub>5</sub>)Fe(CO)<sub>2</sub>I–I<sub>2</sub>–IFe(CO)<sub>2</sub>(Me<sub>5</sub>C<sub>5</sub>)] leads to a 7 cm<sup>-1</sup> blueshift of the v(CO) bands with respect to (Me<sub>5</sub>C<sub>5</sub>)Fe(CO)<sub>2</sub>I.<sup>15</sup>

As the shift of the v(CO) band(s) of metal carbonyls is known to be a sensitive probe for the changes in the electron density at the metal centre, we were interested in investigating the halogen-bonded complexes formed from halide- or organochalcogenolate-substituted metal carbonyls (XB acceptor) and a typical XB donor (*p*-DITFB). For this purpose, we have assembled organometallic halides and organochalcogenides with *p*-DITFB in a series of novel cocrystals and studied their crystal and electronic structures by means of XRD methods, IR spectroscopy and DFT calculations.

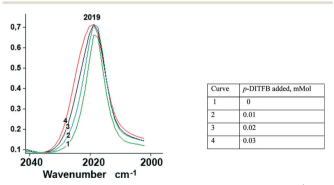
## Results and discussion

In order to assess the v(CO) shift in halogen-bonded complexes, we have simulated the  $CpFe(CO)_2X$ -DITFB (X = Cl, I, TePh; DITFB = 1,4-diiodotetrafluorobenzene) system in the gaseous state. Our computational study correctly reproduced the experimentally observed geometry of the complexes and led us to anticipate a 3-4 cm<sup>-1</sup> hypsochromic shift of the v(CO) bands with respect to free CpFe(CO)<sub>2</sub>TePh (see Table S1\_es of the ESI<sup>†</sup>). The nonpolar nature of hexane and its ability to solubilize CpFe(CO)2TePh seemed to be ideal for the experimental measurement of the XB-induced v(CO) shift in solution. However, during our first attempt to measure the IR spectra of the (CpFe(CO)<sub>2</sub>TePh)(p-DITFB) adduct, we observed that the addition of *p*-DITFB to the green hexane solution of CpFe(CO)<sub>2</sub>TePh unexpectedly resulted in fast decolorization and precipitation of a dark-green material. Its FTIR-ATR spectrum corresponds to the sum of the IR spectra of  $CpFe(CO)_2$ TePh and *p*-DITFB, with a 4 cm<sup>-1</sup> hypsochromic shift of the v(CO) bands when compared to crystalline CpFe(CO)<sub>2</sub>TePh. Precipitation of XB-assembled cocrystal microparticles by addition of water to a solution in a waterimmiscible organic solvent has been reported,<sup>16</sup> but to the best of our knowledge, the direct precipitation of twocomponent crystals just upon mixing of the XB donor and metal-halide or chalcogenolate XB acceptor solutions is unprecedented. Cyclohexane appeared to be an even more suitable nonpolar solvent because it can solubilize CpFe(CO)<sub>2</sub>TePh in the presence of DITFB. The IR-monitored titration of a 0.025 M solution of CpFe(CO)<sub>2</sub>TePh in cyclohexane with a 0.01 mmol portion of solid *p*-DITFB produced a gradual hypsochromic broadening (*ca.* 4 cm<sup>-1</sup> half-width) of the 2019 cm<sup>-1</sup> v(CO) band (see Fig. 1 and the ESI†).

To investigate the possible influence of direct attractive interactions between *p*-DITFB and carbonyl groups, we simulated the IR spectra of the I–CO-bonded CpFe(CO)<sub>2</sub>TePh–*p*-DITFB adduct (optimized in the gas phase) and found a redshift of the v(CO) band (see the ESI†). Moreover, the IR-monitored titration of its halogen-free analog CpMn(CO)<sub>3</sub> against *p*-DITFB under the same conditions also showed a slight bathochromic broadening of the v(CO) bands (see Table S1\_es of the ESI†). From these studies, it is clear that a direct I–CO interaction can only redshift the v(CO) band or reduce its blueshift. The complex interplay between these two opposite effects may be partly responsible for the absence of a clear correlation between the nature of X and the CO blueshift values.

The powder XRD pattern of the CpFe(CO)<sub>2</sub>TePh/*p*-DITFB green precipitate from the hexane solution matched well with the simulated powder XRD pattern<sup>17</sup> (see Fig. S3\_es in the ESI<sup>†</sup>) based on single crystal XRD data for the dark-green  $P_{2_1/c}$  cocrystals 1 prepared by slow vapor diffusion in CpFe(CO)<sub>2</sub>TePh-*p*-DITFB 1:1 mixtures in DCM/hexane (see Fig. 1).

Although the C–O distances in the carbonyl ligands of 1 (1.140(3)–1.146(3) Å) are not significantly different from those in the parent CpFe(CO)<sub>2</sub>TePh (1.140(6)–1.145(4) Å), the  $\nu$ (CO) frequencies in the FTIR-ATR spectrum of 1 are blueshifted (4 cm<sup>-1</sup>) with respect to the parent CpFe(CO)<sub>2</sub>TePh and match well with those of the green precipitate resulting from the interaction of CpFe(CO)<sub>2</sub>TePh with *p*-DITFB in hexane (see Table 1). The crystal structure of 1 is revealed to be of type II, having a genuine I–Te halogen bond with a distance of 3.4863(2) Å which is 0.5 Å shorter than the sum of the corresponding van der Waals (vdW) radii. The trimolecular (CpFe(CO)<sub>2</sub>TePh)<sub>2</sub>( $\mu_2$ -DITFB) assemblies are linked by Te–I chalcogen bonds at a distance of 4.2059(2) Å, which is close to the sum of the Te–I vdW radii (see Fig. 2).



**Fig. 1** Unsymmetrical hypsochromic broadening (*ca.* ~4 cm<sup>-1</sup> half-width) of the  $\nu$ (CO) band at 2019 cm<sup>-1</sup> as a result of gradual addition of *p*-DITFB to a 0.025 mmol cyclohexane solution of CpFe(CO)<sub>2</sub>TePh.

Table 1FTIR-ATR  $\nu$ (CO) frequencies in the neat crystalline samples LM(CO)<sub>2</sub>X and their cocrystals with p-DITFB

		Parent complex, $v$ , cm <sup>-1</sup>	Cocrystal with DITFB, $v$ , cm <sup>-1</sup>	Shift, cm <sup>-1</sup>
1	CpFe(CO) <sub>2</sub> TePh	2007, 1993, 1936	2013, 1993, 1941 $^a$ 2011, 1993, 1940 $^b$	4-6, 0, 4-5
3	CpFe(CO) <sub>2</sub> Cl	2048, 1990	2048, 2003	0, 13
4	CpFe(CO) <sub>2</sub> Br	2035, 1984	2044, 1990	9, 5
5	CpFe(CO) <sub>2</sub> I	2026, 1966, 1946	2032, 1982, 1962	6, 16, 16
6	CbCo(CO) <sub>2</sub> I	2040, 1992	2049, 2003 <sup><i>a</i></sup> 2047, 2001 <sup><i>b</i></sup> 2047, 2002 <sup><i>c</i></sup>	9, 11

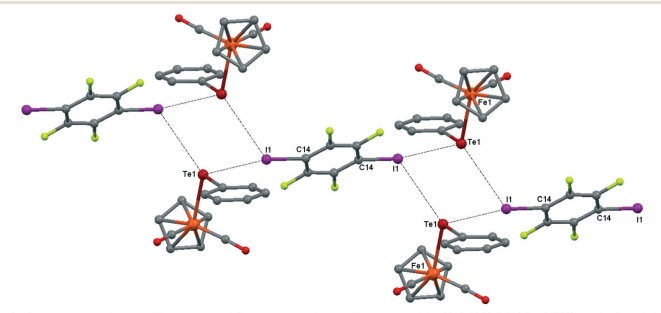
<sup>*a*</sup> Cocrystals obtained by vapor diffusion (see the Experimental section for details). <sup>*b*</sup> Precipitate from *cyclo*-hexane solution. <sup>*c*</sup> Precipitate from *n*-hexane solution.

It should be noted that the parent CpFe(CO)<sub>2</sub>TePh crystal exhibits significant static positional disorder with Te–Fe distances of 2.538–2.617 Å and a disordered Cp ring,<sup>19</sup> while in 1, the CpFe(CO)<sub>2</sub>TePh molecules are perfectly ordered leading to a well-defined Te–Fe distance of 2.5934(3) Å (see Fig. 3). Apparently, cocrystallization with an "innocent" XB (or other secondary bonded) counterpart can, in some cases, be an instrument to "fix" disordered structures.

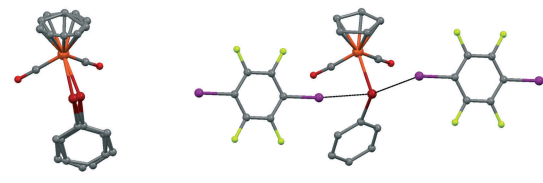
Using the same vapor diffusion technique, which offers the advantage of growing crystals under an inert atmosphere, we prepared the following cocrystals of *p*-DITFB with  $\mu_2$ -SPh, halides (Cl, Br, I) and (tetramethylcyclobutadiene)cobalt-(dicarbonyl)iodide [CbCo(CO)<sub>2</sub>I], which are congeners of CpFe(CO)<sub>2</sub>TePh: [CpFe(CO)(SPh)]<sub>2</sub>–DITFB (2), CpFe(CO)<sub>2</sub>Cl– DITFB (3), CpFe(CO)<sub>2</sub>Br–DITFB (4), CpFe(CO)<sub>2</sub>I–DITFB (5) and CbCo(CO)<sub>2</sub>I–DITFB (6) (see Fig. 4–8, respectively). The interatomic distances in the cyclodienyl–metal–carbonyl moieties in complexes 1–6 are equal or close to those in their parent structures. The solid-state FTIR-ATR data for their  $\nu$ (CO) frequencies (see Table 1) demonstrate the general blueshift, ranging from 5 to 16 cm<sup>-1</sup> when compared to the parent LM(CO)<sub>2</sub>X crystals. The hypsochromic shift in 5 is also in agreement with the 7 cm<sup>-1</sup> blueshift registered in the IR spectrum of the charge transfer complex (Me<sub>5</sub>C<sub>5</sub>)Fe(CO)<sub>2</sub>I-I<sub>2</sub> (1965, 2014 cm<sup>-1</sup>) compared to the parent complex ( $\eta^{5}$ -Me<sub>5</sub>C<sub>5</sub>)Fe(CO)<sub>2</sub>I (1957, 2007 cm<sup>-1</sup>).<sup>15</sup> It is noteworthy that this 7 cm<sup>-1</sup> blueshift can be correctly reproduced by the same computational method as used for the evaluation of the  $\nu$ (CO) blueshift in our complexes 1, 3, and 5 (see the ESI†).

Electron donation from ligand X of  $LM(CO)_2X$  to the iodine of *p*-DITFB appears sufficient to withdraw electron density in the X  $\rightarrow$  M bond and reduce M  $\rightarrow$  CO back-bonding.<sup>20</sup> This results in a decrease of the occupancy of the  $\pi^*C$ -O LUMO and, therefore, a slightly stronger CO bond.

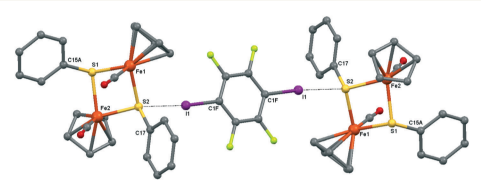
The solid-state structures of cocrystals 1–6 (see Fig. 4–8) are stabilized by short contacts between the halide ligand X of  $CpFe(CO)_2X$  (X = Cl, Br, I) and the iodine atoms of the



**Fig. 2** Crystal structure diagram of **1**, showing the I–Te halogen-bonded trimolecular assemblies  $(CpFe(CO)_2TePh)_2(\mu_2-DITFB)$  and the Te–I chalcogen interactions between them. Hydrogen atoms are omitted for clarity. Selected distances (Å): I(1)–Te(1), 3.4863(2); Te(1)–Fe(1), 2.5934(3); Te(1)–I(1), 4.2059(2). Selected angles (°): I(1)–Te(1)–I(1), 90.55(1); Fe(1)–Te(1)–I(1), 108.64(1); Fe(1)–Te(1)–I(1), 95.17(1); Fe(1)–Te(1)–C(8), 102.16(6); Te(1)–I(1)–C(14), 169.71(6). Dotted lines link Te and I atoms at distances shorter than the sum of Te–I vdW radii.<sup>18</sup>







**Fig. 4** Crystal structure diagram of **2**, showing its I–S halogen-bonded trimolecular assemblies  $([CpFe(CO)(\mu_2-SPh)]_2)_2(\mu_2-DITFB)$ . Hydrogen atoms are omitted for clarity. Selected distances (Å): I(1)-S(2), 3.130(1). Selected angles (°): C(1F)-I(1)-S(2), 173.9(1); Fe(1)–S(2)–I(1), 111.91(4); I(1)–S(2)–C(17), 102.5(1); I(1)–S(2)–Fe(2), 118.86(4). Dotted lines link S and I atoms at a distance shorter than the sum of the S–I vdW radii.<sup>18</sup>

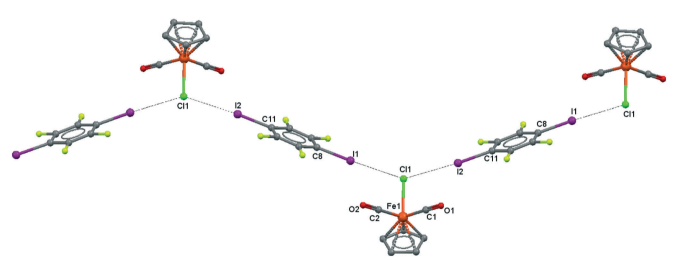
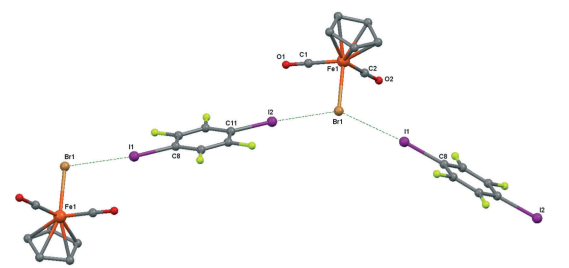


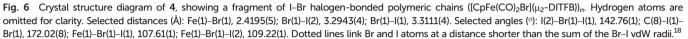
Fig. 5 Crystal structure diagram of 3, showing a fragment of I–Cl halogen-bonded polymeric chains  $([CpFe(CO)_2Cl](\mu_2-DITFB))_n$ . Hydrogen atoms are omitted for clarity. Selected distances (Å): Fe(1)–Cl(1), 2.318(3); Cl(1)–I(1), 3.219(5); Cl(1)–I(2), 3.229(5). Selected angles (°): I(2)–Cl(1)–I(1), 141.48(3); Fe(1)–Cl(1)–I(1), 110.02(4); Fe(1)–Cl(1)–I(2), 108.33(4); Cl(1)–I(2)–C(11), 173.4(1); Cl(1)–I(1)–C(8), 175.8(1). Dotted lines link Cl and I atoms at distances shorter than the sum of Cl–I vdW radii.<sup>18</sup>

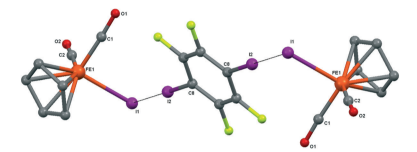
ditopic XB acceptor (*p*-DITFB). The dependence of the I-X-Fe angle on the nature of X is a remarkable illustration of the directionality of XBs.

NBO/PBE0 calculations for  $[CpFe(CO)_2X](p-DITFB)$  (X = Cl, I, TePh) adducts demonstrate the dominant contribution from the unhybridized apical p-electron pair of iodine in DITFB (see Tables 1 and S1†\_es) and p-electrons of the ligand X p-belt. The relative contribution of XB acceptor orbitals to X–I XB and their hybridization do not show much dependence on the nature of ligand X.

The geometry of a pure p-orbital implies certain demands for the XB geometry, thus providing the XB directionality,







**Fig. 7** Crystal structure diagram of **5**, showing its I–I halogen-bonded trimolecular assemblies (CpFe(CO)<sub>2</sub>I)<sub>2</sub>( $\mu_2$ -DITFB). Hydrogen atoms are omitted for clarity. Selected distances (Å): Fe(1)–I(1), 2.6025(4); I(1)–I(2), 3.5488(3). Selected angles (°): Fe(1)–I(1)–I(2), 89.06(1); C(8)–I(2)–I(1), 175.50(7). Dotted lines link I atoms at a distance shorter than the sum of I–I vdW radii.<sup>18</sup>

which was reproduced in our DFT simulation and is visualized in the ESP maps (see Fig. 9).

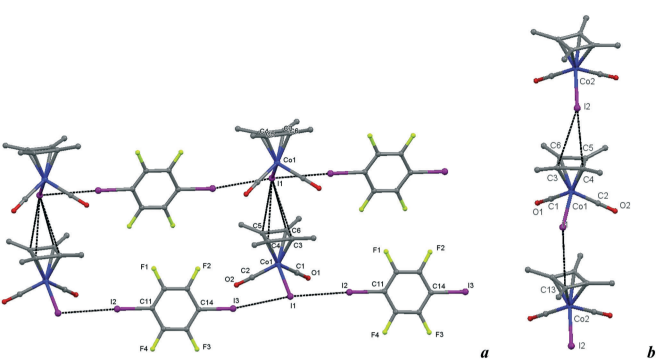
ESP mapping for CpFe(CO)<sub>2</sub>I (Fig. 9a) shows the nucleophilic area at the specific equatorial "belt" of the iodine ligand, which is orthogonal to the Fe-I bond and has two distinct maxima matching with the p-orbital as well as with the "docking" sites for XB with electrophilic iodine functions of p-DITFB. One of the equatorial p-orbitals of the Te atom in the TePh ligand of complex CpFe(CO)<sub>2</sub>TePh is involved in covalent bonding with a carbon atom of the Ph group and therefore Te is left with only one XB-suitable p-orbital, resulting in two clearly visible spots on the ESP map (Fig. 9b) and directing the XB bonding with electrophilic iodine functions of *p*-DITFB in line with the *x*-axis, as suggested by the symmetry of the p orbital of Te. The more anionic, and therefore having more isotropic ESP, chloride ligand in CpFe(CO)<sub>2</sub>Cl does not reveal specific nucleophilic areas, which makes XB bonding more electrostatic and therefore less angular demanding, so that the electrophilic iodines of *p*-DITFB may exist at angles exceeding 90° (see Fig. 9c).

This increasing directionality of XB bonding in the sequence Cl > Br > I can be an additional reason for the low

affinity of the iodine ligand in a sterically crowded (2,6bis[(di-*t*-butylphosphino)methyl]phenyl)–Pd–I complex toward its interaction with DITFB and  $I(CF_2)_8I$  XB donors, noted in ref. 6. As shown in Fig. 10, the *tert*-butyl groups of a 2,6bis[(di-*t*-butylphosphino)methyl]phenyl pincer provide efficient shielding of the 5p equatorial "belt" of iodine (which is perpendicular to the Pd–I bond) but allow space for the interaction with the more isotropic and less directionally demanding "sphere" of the Cl ligand.

We can describe the packing patterns of 3 and 4 as isotypically packed zigzag chains, where the X ligand (Cl, Br) accommodates two *p*-DITFB molecules, with an average I–X–I angle of 140°, while their TePh,  $\mu_2$ -SPh and iodo congeners **1**, **2** and **5** are associated into trimolecular units (CpFe(CO)<sub>*n*</sub>X)<sub>2</sub>( $\mu_2$ -DITFB). In contrast, iodide **6** forms polymeric chains [(Cb\*Co(CO)<sub>2</sub>I)( $\mu_2$ -DITFB)]<sub>*n*</sub>, with a nearly linear I<sub>DITEB</sub>-I<sub>M</sub>-I<sub>DITFB</sub> angle (see Fig. 2–8).

Finally, it is worth noting that the same phases 1–6 are formed irrespective of the ratio of the starting reagents. With respect to the competition between XB and hydrogen bonding (HB),<sup>22</sup> we should also mention that under similar conditions, no supramolecular assembly was observed when 1,4-



**Fig. 8** a) Crystal structure diagram of **6**, showing a fragment of I–I halogen-bonded polymeric chains ([Cb\*Co(CO)<sub>2</sub>I]( $\mu_2$ -DITFB))<sub>n</sub> linked by I– $\pi(\eta_4$ -Me<sub>4</sub>C<sub>4</sub>) interactions. Hydrogen atoms are omitted for clarity. Selected distances (Å): intramolecular Co(1)–I(1), 2.619(1); intermolecular I(3)–I(1), 3.589(1); I(1)–I(2), 3.585(1); I(1)–C(3–6) ring of the  $\eta^4$  = Me<sub>4</sub>C<sub>4</sub> ligand, 3.781(7)–3.866(7). Selected angles (°): I(3)–I(1)–I(2), 171.51(2); Co(1)–I(1)–I(3), 93.00(2); Co(1)–I(1)–I(2), 95.16(2); C(14)–I(3)–I(1), 169.6(2); C(11)–I(2)–I(1), 177.2(2). b) Comparison of the Cb\*Co(CO)<sub>2</sub>I stacks in **6** with those in the parent crystal. Selected distances (Å): intermolecular I(2)–C(4), 3.725(8); I(2)–C(3), 3.87(1); I(1)–C(13), 3.842(9), and similarities of the intramolecular distances Co(2)–I(2), 2.598(1); Co(1)–C(2), 1.781(7); C(2)–O(2), 1.157(9); Co(1)–C(1), 1.79(1); C(1)–O(1), 1.16(1). Dotted lines link specific atoms at a distance shorter than the sum of their vdW radii.<sup>18</sup>

hydroquinone (a HB donor) was used instead of 1,4-DITFB as a counterpart to the  $CpFe(CO)_2X$  acceptor.

#### architectures. The investigation of this complex interplay may be one of the future perspectives opened by this work.

# Summary

We have established that the terminal halogen (Cl, Br, I), TePh, and bridging  $\mu_2$ -SPh ligands in the complexes CpFe(CO)<sub>n</sub>X and Cb\*Co(CO)<sub>2</sub>X can act as XB acceptors. The XB they form with the iodine atoms of the bidentate XB donor *p*-DITFB assembles them into trimolecular entities  $(CpFe(CO)_2X)_2(\mu_2$ -DITFB) or into polymeric  $([CpFe(CO)_2X](\mu_2$ - $DITFB)_n$  chains, packed in the cocrystals, which may precipitate instantly from hexane solutions solely upon addition of p-DITFB. These cocrystals exhibit a hypsochromic shift of the carbonyl group vibration bands in their IR spectra of ca. 4-12 cm<sup>-1</sup>, indicating that the electron-withdrawing effect of the iodine of p-DITFB appears sufficient enough to affect the occupancy of the  $\pi^*C$ -O LUMO through I-X-M- $\pi^*C$ -O interactions. These experimental findings are in good agreement with the computational simulation performed on CpFe(CO)<sub>2</sub>TePh-DITFB, CpFe(CO)<sub>2</sub>Cl-DITFB and CpFe(CO)<sub>2</sub>I-DITFB complexes. The symmetry, electrostatic surface potential distribution and availability of the s- and p-orbitals of the XB acceptor ligand in the metal complexes dictate the definite directionality of XBs and the resulting supramolecular

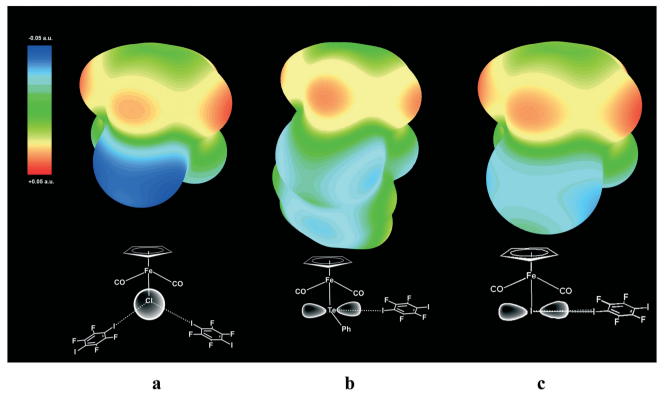
#### Experimental

All reactions and manipulations were performed using standard Schlenk techniques under an inert atmosphere of pure nitrogen or argon. Solvents were purified, dried and distilled under a nitrogen atmosphere prior to use. Commercial reagents (*p*-DITFB) were used without additional purification. CpFe(CO)<sub>2</sub>X (X = Cl, Br, I),<sup>23</sup> CpFe(CO)<sub>2</sub>TePh, CpFe(CO)( $\mu$ -SPh)]<sub>2</sub> (ref. 24) and Cb\*Co(CO)<sub>2</sub>I (ref. 25) were prepared following the reported procedures. Slow vapour diffusion techniques were used for the preparation of the cocrystals **1–6**.

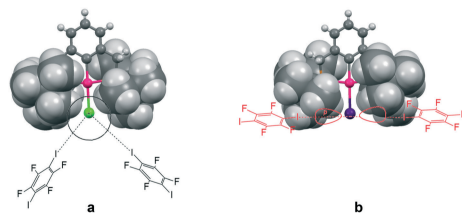
IR spectra were recorded on a Bruker Alpha FTIR spectrometer equipped with an ATR facility (for solid samples) and a  $0.1 \text{ mm CaF}_2$  cell for hexane and cyclohexane solutions.

#### Preparation of 1-6 (general procedure)

0.1 mmol of CpFe(CO)<sub>2</sub>X and 0.1 mmol of *p*-DITFB were dissolved in 0.05 ml of  $CH_2Cl_2$  in a 5 mm glass tube, and the mixture was placed into a 20 mm/10 ml test tube containing 1 ml of *n*-heptane in an argon atmosphere, which was closed and left for 48 h at 4 °C. Crystals formed which were used for XRD and IR investigations.



**Fig. 9** Electrostatic surface potential (ESP) maps for free parent molecules  $CpFe(CO)_2I$  (a),  $CpFe(CO)_2TePh$  (b) and  $CpFe(CO)_2CI$  (c) showing the nucleophilic areas at the iodine, tellurium and chlorine atoms, respectively. Note that the spherical potential distribution at the Cl ligand in  $CpFe(CO)_2CI$  does not dictate specific directionality of I–Cl XB (a), while unhybridized 5p orbitals of Te (b) and I (c)<sup>21</sup> direct the X–I XB at the right angle to Fe–TePh and Fe–I bonds.



**Fig. 10** The *tert*-butyl fragment of a (2,6-bis[(di-t-butylphosphino)methyl]phenyl)PdX (ref. 6) (X = Cl (a) and I (b)) complex is shown as a space filling model to demonstrate the shielding of the 5p equatorial "belt" of the iodide ligand (b), as compared to the spherical chloride "anion" (a), escaping the shield.

#### Precipitation of CpFe(CO)<sub>2</sub>TePh-DITFB

To an emerald-green solution of 2 mg ( $\sim 0.005$  mmol) of CpFe(CO)<sub>2</sub>TePh in *n*-hexane (1 ml) in a 2 ml Eppendorf tube, 0.008 g (0.02 mmol) of *p*-DITFB was added and the mixture was shaken several times to ensure complete dissolution. After several seconds, a green precipitate formed, leaving a pale greenish-yellow solution. The green solid residue was washed with hexane, dried in a flow of argon and subjected to IR measurement using the ATR attachment.

# Crystal structure determination of compounds 1–7

#### Powder XRD

The powder diffraction pattern of the sample was recorded at room temperature on a Bruker D8 Advance Vario powder diffractometer, equipped with a copper anode (Cu  $K_{\alpha 1}$ ), a Ge(111) monochromator, and a position-sensitive LynxEye detector (6–90°  $2\theta$  interval, 0.01° steps). The diffraction data

was processed and analyzed using Bruker Topas5 (ref. 26) software. The investigated sample is single-phase. The simulation of the diffraction pattern by the Rietveld method is confirmed by the phase composition of the sample phase, which is determined from the single crystal diffraction data. The discrepancy parameters are:  $R_{\rm wp} = 3.29\%$ ,  $R_{\rm p} = 2.51\%$ ,  $R'_{\rm wp} = 12.52\%$ ,  $R'_{\rm p} = 14.48\%$ , and  $R_{\rm Bragg} = 2.266\%$ . The unit cell parameters are as follows: a = 13.73614(15) Å, b = 11.69728(14) Å, c = 11.85569(14) Å, and  $\beta = 15.2909(8)^{\circ}$ .

#### Single crystal XRD

Crystals of 1–7 suitable for XRD investigation were grown by slow evaporation of a dichloromethane/*n*-hexane solvent mixture by varying the temperature from 25 °C to 0 °C. Relevant crystallographic data and the details of measurements are given in the ESI.†

A Bruker APEX II CCD area detector diffractometer equipped with a graphite-monochromated Mo K $\alpha$  radiation source (0.71070 Å) was used for cell determination and intensity data collection for compounds 1–6. The structure was solved by direct methods and refined by full-matrix least squares against  $F^2$  using SHELXL-97 and Olex2 software.<sup>27,28</sup> Non-hydrogen atoms were refined with anisotropic thermal parameters. All hydrogen atoms were geometrically fixed and refined using a riding model. Atomic coordinates and other structural parameters of 1–7 have been deposited with the Cambridge Crystallographic Data Centre (CCDC no. 1587662 (1), 1587659 (2), 1587660 (3), 1587664 (4), 1587663 (5), 1587661 (6), and 1587665 (7)).

#### **Calculation details**

Theoretical calculations were carried out with the ORCA 4.01 program package.<sup>29</sup> A non-hybrid PBE functional,<sup>30</sup> dispersion correction with Becke-Johnson damping (D3BJ)<sup>31</sup> and a def2-TZVP basis set<sup>32</sup> with a small-core pseudopotential for Te and I atoms<sup>33</sup> were used for geometry optimization and Hessian calculation. The def2/J auxiliary basis set34 was used for Coulomb fitting. Single point energies were calculated on these geometries using the hybrid functional PBE0 (ref. 35) with the same basis set and dispersion correction. NBO analysis was performed at this level of theory using the NBO 6.0 program.<sup>36</sup> Single point energies of DITFB adduct formation were also evaluated by means of new domain-based local pair natural orbital coupled cluster theory with single, double, and perturbative triple excitations (DLPNO-CCSD(T)<sup>37</sup> using def2 triple and quadruple-zeta basis sets<sup>29</sup> and appropriate correlation fitting basis sets).<sup>38</sup> Extrapolation to the complete basis set limit<sup>39</sup> was performed using parameters  $\alpha = 10.39$ ,  $\beta$  = 2.4 and  $\alpha$  = 7.88,  $\beta$  = 3 for "cardinal numbers" 2, 3 and 3, 4, respectively.<sup>40</sup>

# Conflicts of interest

There are no conflicts to declare.

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## References

- (a) G. R. Desiraju, P. S. Ho, L. Kloo, A. C. Legon, R. Marquardt, P. Metrangolo, P. Politzer, G. Resnati and K. Rissanen, *Pure Appl. Chem.*, 2013, 85, 1711–1713; (b) G. Cavallo, P. Metrangolo, R. Milani, T. Pilati, A. Priimagi, G. Resnati and G. Terraneo, *Chem. Rev.*, 2016, 116, 2478–2601.
- 2 (a) P. Politzer, J. S. Murray and T. Clark, *Phys. Chem. Chem. Phys.*, 2013, 15, 11178–11189; (b) H. Wang, W. Wang and W. J. Jin, *Chem. Rev.*, 2016, 116, 5072–5104.
- 3 (a) B. Li, S.-Q. Zang, L.-Y. Wang and T. C. W. Mak, *Coord. Chem. Rev.*, 2015, 308, 1–21, DOI: 10.1016/j.ccr.2015.09.005;
  (b) R. W. Troff, T. Mäkelä, F. Topić, A. Valkonen, K. Raatikainen and K. Rissanen, *Eur. J. Org. Chem.*, 2013, 1617–1637, DOI: 10.1002/ejoc.201201512; (c) R. Bertani, P. Sgarbossa, A. Venzo, F. Lelj, M. Amati, G. Resnati, T. Pilati, P. Metrangolo and G. Terraneo, *Coord. Chem. Rev.*, 2010, 254, 677–695.
- 4 C. Perkins, S. Libri, H. Adams and L. Brammer, *CrystEngComm*, 2012, 14, 3033–3038.
- P. Sgarbossa, R. Bertani, V. Di Noto, M. Piga, G. A. Giffin, G. Terraneo, T. Pilati, P. Metrangolo and G. Resnati, *Cryst. Growth Des.*, 2012, 12, 297, DOI: 10.1021/cg201073m.
- 6 M. T. Johnson, Z. Džolić, M. Cetina, O. F. Wendt, L. Öhrström and K. Rissanen, *Cryst. Growth Des.*, 2012, 12, 362–368.
- 7 Y. Wang, W. Wu, Y. Liu and Y. Lu, *Chem. Phys. Lett.*, 2013, 578, 38–42, DOI: 10.1016/j.cplett.2013.06.005.
- 8 L. Brammer, G. M. I. Espallargas and S. Libri, *CrystEngComm*, 2008, **10**, 1712–1727, DOI: 10.1039/ b812927d.
- 9 C. B. Aakeröy, M. Baldrighi, J. Desper, P. Metrangolo and G. Resnati, *Chem. Eur. J.*, 2013, **19**, 16240–16247.
- M. G. Sarwar, B. Dragisic, L. J. Salsberg, C. Gouliaras and M. S. Taylor, *J. Am. Chem. Soc.*, 2010, 132, 1646–1653.
- 11 A. Bauzà, D. Quiñonero, A. Frontera and P. M. Deyà, *Phys. Chem. Chem. Phys.*, 2011, 13, 20371–20379.
- 12 R. A. Thorson, G. R. Woller, Z. L. Driscoll, B. E. Geiger, C. A. Moss, A. L. Schlapper and N. P. Bowling, *Eur. J. Org. Chem.*, 2015, 1–12, DOI: 10.1002/ejoc.201403671.
- (a) C. Laurence, M. Queignec-Cabanetos, T. Dziembowska, R. Queignec and B. Wojtkowiak, J. Am. Chem. Soc., 1981, 103, 2567–2573; (b) C. Laurence, M. Queignec-Cabanetos

and B. Wojtkowiak, *J. Chem. Soc., Perkin Trans.* 2, 1982, 1605–1610; (*c*) C. Laurence, M. Queignec-Cabanetos and B. Wojtkowiak, *Can. J. Chem.*, 1983, **61**, 135–138.

- 14 J. W. Kee and W. Y. Fan, J. Organomet. Chem., 2013, 729, 14–19.
- 15 N. Masuhara, S. Nakashima and K. Yamada, *Chem. Lett.*, 2005, 34, 1352.
- 16 (a) X. Pang, H. Wang, X. R. Zhao and W. J. Jin, *CrystEngComm*, 2013, 15, 2722–2730; (b) H. Y. Gao, Q. J. Shen, X. R. Zhao, X. Q. Yan, X. Pang and W. J. Jin, *J. Mater. Chem.*, 2012, 22, 5336.
- 17 Mercury 3.6 (Build RC6) CCDC 2015.
- 18 S. Alvarez, Dalton Trans., 2013, 42, 8617.
- 19 Yu. V. Torubaev, A. A. Pasynsky and I. V. Skabitsky, *Russ. J. Coord. Chem.*, 2009, 35, 347.
- 20 G. Bistoni, S. Rampino, N. Scafuri, G. Ciancaleoni, D. Zuccaccia, L. Belpassi and F. Tarantelli, *Chem. Sci.*, 2016, 7, 1174–1184.
- 21 I. V. Alabugin, S. Bresch and M. Manoharan, J. Phys. Chem. A, 2014, 118, 3663–3677.
- 22 S. Panikkattu, P. D. Chopade and J. Desper, *CrystEngComm*, 2013, 3125–3136.
- 23 Synthetic Methods of Organometallic and Inorganic Chemistry, ed. W. A. Herrmann and G. Thieme, Stuttgart, Germany, vol. 82, p. 107; E. D. Scheremer and W. H. Baddley, J. Organomet. Chem., 1971, 27, 83–88; E. V. Mutseneck, D. A. Loginov, D. S. Perekalin, Z. A. Starikova, D. G. Golovanov, P. V. Petrovskii, P. Zanello, M. Corsini, F. Laschi and A. R. Kudinov, Organometallics, 2004, 23, 5944–5957.
- 24 E. D. Scheremer and W. H. Baddley, J. Organomet. Chem., 1971, 27, 83–88.
- 25 E. V. Mutseneck, D. A. Loginov, D. S. Perekalin, Z. A. Starikova, D. G. Golovanov, P. V. Petrovskii, P. Zanello, M. Corsini, F. Laschi and A. R. Kudinov, *Organometallics*, 2004, 23, 5944–5957.

- 26 Bruker TOPAS 5 User Manual.—Bruker AXS GmbH, Karlsruhe, Germany, 2015.
- 27 G. M. Sheldrick, Acta Crystallogr., Sect. A: Found. Crystallogr., 2008, 64, 112.
- 28 O. V. Dolomanov, L. J. Bourhis, R. J. Gildea, J. A. K. Howard and H. Puschmann, J. Appl. Crystallogr., 2009, 42, 339–341.
- 29 F. Neese, "Software update: the ORCA program system, version 4.0" WIREs Comput Mol Sci 2017, p. e1327, DOI: 10.1002/wcms.1327.
- 30 (a) J. P. Perdew, K. Burke and M. Ernzerhof, *Phys. Rev. Lett.*, 1996, 77, 3865; (b) J. P. Perdew, K. Burke and M. Ernzerhof, *Phys. Rev. Lett.*, 1997, 78, 1396.
- (a) S. Grimme, S. Ehrlich and L. Goerigk, J. Comput. Chem., 2011, 32, 1456; (b) S. Grimme, J. Antony, S. Ehrlich and H. Krieg, J. Chem. Phys., 2010, 132, 154104.
- 32 F. Weigend and R. Ahlrichs, *Phys. Chem. Chem. Phys.*, 2005, 7, 3297.
- 33 K. A. Peterson, D. Figgen, E. Goll, H. Stoll and M. Dolg, J. Chem. Phys., 2003, 119, 11113–11123.
- 34 F. Weigend, Phys. Chem. Chem. Phys., 2006, 8, 1057.
- 35 C. Adamo and V. J. Barone, *J. Chem. Phys.*, 1999, 110, 6158–6170.
- 36 E. D. Glendening, J. K. Badenhoop, A. E. Reed, J. E. Carpenter, J. A. Bohmann, C. M. Morales, C. R. Landis and F. Weinhold, Theoretical Chemistry Institute, University of Wisconsin, Madison, WI, 2013, http://nbo6.chem.wisc.edu/.
- 37 C. Riplinger, P. Pinski, U. Becker, E. F. Valeev and F. Neese, *J. Chem. Phys.*, 2016, 144, 024109.
- 38 A. Hellweg, C. Hattig, S. Hofener and W. Klopper, *Theor. Chem. Acc.*, 2007, 117, 587.
- 39 (a) W. Klopper and W. Kutzelnigg, *THEOCHEM*, 1986, 135, 339–356; (b) W. Kutzelnigg, *Int. J. Quantum Chem.*, 1994, 51, 447–463; (c) S. J. Zhong, E. C. Barnes and G. A. Petersson, *J. Chem. Phys.*, 2008, 129, 184116.
- 40 D. G. Truhlar, Chem. Phys. Lett., 1998, 294, 45-48.