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**S-(CF<sub>3</sub>)Thianthrenium and S-(CF<sub>3</sub>)dibenzothiophenium cations form potent chalcogen bonds (ChBs) with [Mo(CO)<sub>5</sub>CN]<sup>-</sup>, yielding S<sub>2</sub>N<sub>2</sub> supramolecular motifs. Crystal structures reveal shorter S···N contacts opposite the CF<sub>3</sub> group compared to the aryl substituents. The energetic features of the ChBs have been studied using DFT calculations demonstrating the structure guiding role of ChBs.**

The chalcogen bond (ChB) is an attractive interaction characterized by an electrophilic chalcogen atom (ChB donor) interacting with an electron-rich site (ChB acceptor).<sup>1</sup> In the solid state, ChBs have been compared to halogen bonding in terms of directionality and supramolecular synthons.<sup>2</sup> Understanding the physical nature of ChB interactions is complex, since it involved electrostatic effects, polarization, dispersion contributions, and n → σ\* orbital delocalization. In recent years, ChB has gained prominence in synthesis, catalysis, material design, and coordination chemistry, particularly for crystal engineering.<sup>2,3</sup> Supramolecular hierarchies and design strategies have been devised to create molecules with large σ-holes and high electrostatic potentials. Achieving this typically involves incorporating multiple electron-withdrawing substituents.<sup>4</sup>

Fluorine's significance in pharmaceuticals and agrochemicals has led to extensive research on methodologies for introducing fluorine or fluorinated groups into organic molecules.<sup>5</sup> Notably, the incorporation of CF<sub>3</sub> groups is of particular interest.<sup>6</sup> Umemoto's work in the 1990s introduced dibenzothiophenium cations as effective electrophilic trifluoromethylating agents, with reactivity fine-tuned by varying the chalcogen atom and incorporating substituents in the aryl rings.<sup>7</sup> Recently, Ritter<sup>8</sup> introduced

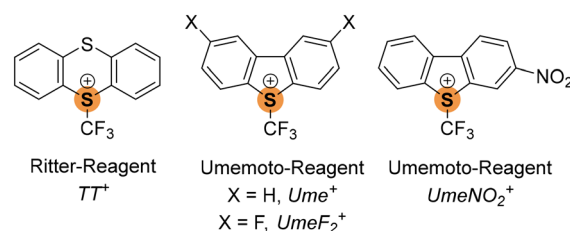
## CF<sub>3</sub>-substituted sulfonium cations as efficient chalcogen bond donors towards cyanometalates†

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S-(trifluoromethyl) thianthrenium triflate as a readily available trifluoromethylating agent. Given the pronounced σ-holes in sulfonium cations, we explored whether these cationic S-CF<sub>3</sub> reagents could serve as potent ChB donors. As ChB acceptors, we considered cyanometalates due to their high Lewis basicity<sup>9</sup> of the negatively polarized nitrogen atoms and their suitability as halogen bond acceptors.<sup>10</sup>

Surprisingly, this approach remains relatively unexplored in the context of ChB. Only two X-ray structures<sup>11,12</sup> have been reported in the CCDC involving sulfonium cations R<sub>3</sub>S<sup>+</sup> and cyanometalate anions, where the S···N contacts fall below the sum of van der Waals radii ( $\sum$ vdW(SN): 3.35 Å).<sup>13</sup>

We envisaged the use of the Ritter reagent [TT]<sup>+</sup>[CF<sub>3</sub>SO<sub>3</sub>]<sup>-</sup> as a soluble ChB donor (Scheme 1). To fine-tune the ChB donor ability, a series of Umemoto reagents with different substituents was also used (Scheme 1). To avoid the formation of polymeric network structures, [NEt<sub>4</sub>]<sup>+</sup>[Mo(CO)<sub>5</sub>CN]<sup>-</sup> was used as a building block. Salt metathesis of [NEt<sub>4</sub>]<sup>+</sup>[Mo(CO)<sub>5</sub>CN]<sup>-</sup> and [TT]<sup>+</sup>[CF<sub>3</sub>SO<sub>3</sub>]<sup>-</sup> in CH<sub>2</sub>Cl<sub>2</sub> led to [TT]<sup>+</sup>[Mo(CO)<sub>5</sub>CN]<sup>-</sup> which crystallizes in the triclinic space group *P*1. The sulfur atom carrying the CF<sub>3</sub> substituent acts as a biaxial ChB donor whereas the nitrogen of the cyanide ligand acts as a bifurcated ChB acceptor, resulting in the formation of an intermolecular S<sub>2</sub>N<sub>2</sub> ring motive with two different S···N distances (Fig. 1). The shorter contact of 2.775(2) Å is situated in elongation of the S-CF<sub>3</sub> group, whereas the longer contact of 3.212(1) Å is in the plane of the



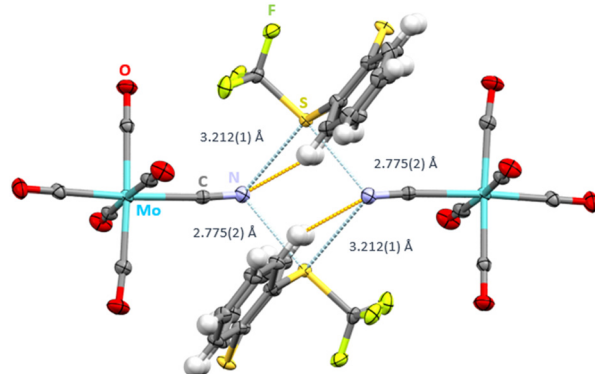
**Scheme 1** Available sulfonium-type chalcogen bond donors containing S-CF<sub>3</sub> moieties.

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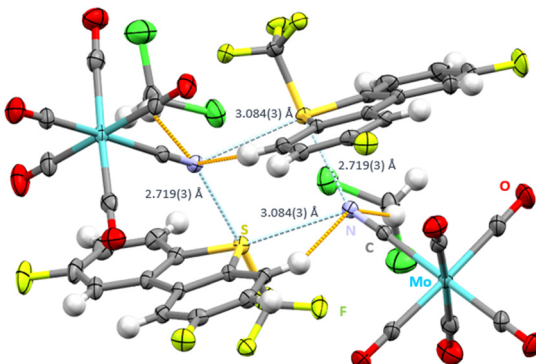




**Fig. 1** Intermolecular S···N contacts (light blue, distances given) and hydrogen bonds (orange) in the crystal structure of  $[\text{TT}]^+[\text{Mo}(\text{CO})_5\text{CN}]^-$ . Thermal ellipsoids at the 50% probability level.

aryl rings. Both values are below the sum of van der Waals radii of 3.35 Å of sulfur and nitrogen.<sup>13</sup> Whereas the C-S···N angles are close to linear for the short S···N contact ( $\text{C}_{100}-\text{S}_1-\text{N}_1$ :  $172.31(22)^\circ$ ) the deviation from linearity is greatly pronounced for the longer S···N interaction ( $\text{C}_{25}-\text{S}_1-\text{N}_1$ :  $95.95(19)^\circ$ ). The bifurcation of the cyanide ligand leads to  $\text{C}\equiv\text{N}\cdots\text{S}$  angles of  $117.91(37)^\circ$  and  $123.59(39)^\circ$ . Surprisingly, the second sulfur atom of the *S*-(trifluoromethyl) thianthrenium cation is also a ChB donor. Here, the S···O contact is 3.264(4) ( $\sum_{\text{vdW}}(\text{SO})$ : 3.32 Å (ref. 13)) to the negatively polarized oxygen atom of the carbonyl ligand (Fig. S1†). This interaction is furthermore supported by short contacts between the hydrogen atoms in  $\beta$ -position to sulfur to the oxygen atoms of the anions (H–O distances 2.524(1) and 2.588(1) Å).

Interestingly, when salt metathesis between the unsubstituted  $[\text{Ume}]^+[\text{CF}_3\text{SO}_3^-]$  and  $[\text{NET}_4]^+[\text{Mo}(\text{CO})_5\text{CN}]^-$  was attempted, only single crystals of both starting materials were observed, without any salt metathesis. Consequently, derivatives of the Umemoto cation containing electron-withdrawing substituents were prepared. Salt metathesis of the double fluorinated Umemoto cation  $[\text{UmeF}_2]^+[\text{B}(\text{C}_6\text{F}_5)_4^-]$

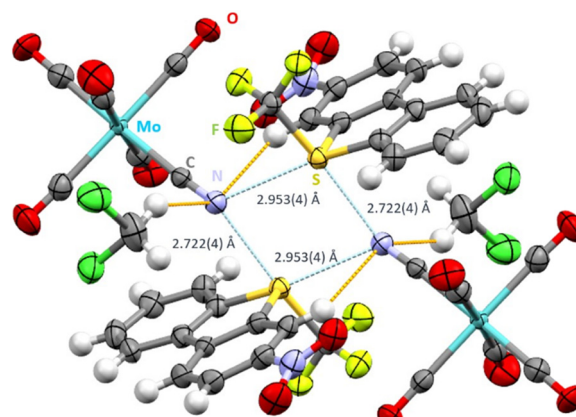


**Fig. 2** Intermolecular S···N contacts (light blue, distances given) and hydrogen bonds (orange) in the crystal structure of  $[\text{UmeF}_2]^+[\text{Mo}(\text{CO})_5\text{CN}]^- \cdot \text{CH}_2\text{Cl}_2$ . Thermal ellipsoids at the 50% probability level, solvent molecules not shown.

with  $[\text{NET}_4]^+[\text{Mo}(\text{CO})_5\text{CN}]^-$  in  $\text{CH}_2\text{Cl}_2$  gave single crystals of  $[\text{UmeF}_2]^+[\text{Mo}(\text{CO})_5\text{CN}]^- \cdot \text{CH}_2\text{Cl}_2$  (space group  $C2/c$ , Fig. 2). In agreement with the previous structure the contact *trans* to the  $\text{CF}_3$  group (2.719(3) Å) is shorter than the contact *trans* to the aryl rings (3.084(3)). The C-S···N ChB is strongly directional with an angle of  $176.68(23)^\circ$ . Bifurcation around the cyanide nitrogen atoms leads to  $\text{C}\equiv\text{N}\cdots\text{S}$  angles in the range of  $129.64(38)^\circ$ – $141.48(41)^\circ$ . As in all other structures the S– $\text{CF}_3$  bonds are longer (1.884(5)–1.886(7) Å) than the *S*-aryl bonds (1.776(6)–1.782(6) Å).

Salt metathesis of nitro-substituted  $[\text{UmeNO}_2]^+[\text{SbF}_6]^-$  with  $[\text{NET}_4]^+[\text{Mo}(\text{CO})_5\text{CN}]^-$  in  $\text{CH}_2\text{Cl}_2$  proved challenging due to extensive decomposition reactions. In one case, the desired cocrystal was obtained, but it was later discovered that the old batch of the molybdenum compound used was contaminated with around 18% of the analogous tungsten compound. Experiments with the pure molybdenum compound were unfortunately not successful despite numerous attempts.  $[\text{UmeNO}_2]^+[\text{Mo}_{0.822}\text{W}_{0.178}(\text{CO})_5\text{CN}]^- \cdot \text{CH}_2\text{Cl}_2$  crystallizes in the space group  $P\bar{1}$ , displaying the same intermolecular  $\text{S}_2\text{N}_2$  motive (Fig. 3). Interestingly, shortening of the S···N contacts to 2.722(4) Å (*trans* to the  $\text{CF}_3$  group) and 2.953(4) Å (*trans* to the aryl group) was observed. In contrast to the previous structure, both ChB are more linear ( $\text{C}_{13}-\text{S}_1-\text{N}_2$   $174.36(25)^\circ$  and  $\text{C}_4-\text{S}_1-\text{N}_2$   $172.01(21)^\circ$ ). The bifurcated nitrogen atom of the cyanide group shows  $\text{C}_{15}-\text{N}_2-\text{S}_1$  angles of  $130.92(46)^\circ$  and  $127.24(42)^\circ$ . Furthermore, the nitrogen atoms display close contacts to two hydrogen atoms of neighboring cations ( $\text{N}_2-\text{H}_9$  2.681(7) Å) and solvent molecules ( $\text{N}_2-\text{H}_{20b}$  2.728(7) Å). Additionally, three carbonyl oxygens and both oxygen atoms of the nitro group display O–H contacts in the range of 2.543(7)–2.683(9) Å and 2.678(7) Å, respectively. The oxygen atoms of the nitro group form also intramolecular O···H bonds to the hydrogen atoms in *ortho*-position (2.433(6) Å and 2.455(7) Å).

In order to rationalize the strong ChB donor ability of the  $\text{CF}_3$ -substituted sulfonium cations, electrostatic potential



**Fig. 3** Intermolecular S···N contacts (light blue, distances given) and hydrogen bonds (orange) in the crystal structure of  $[\text{UmeNO}_2]^+[\text{Mo}_{0.822}\text{W}_{0.178}(\text{CO})_5\text{CN}]^- \cdot \text{CH}_2\text{Cl}_2$ . Thermal ellipsoids at the 50% probability level.

maps (MEPs) were calculated, allowing to establish a hierarchy between them. In this context, the influence of electron-withdrawing groups (F, NO<sub>2</sub>) on the strength of the  $\sigma$ -holes was of particular interest. Sulfonium cations (R<sub>3</sub>S<sup>+</sup>) exhibit three  $\sigma$ -holes extending from the three C-S bonds, as previously demonstrated.<sup>14</sup> This configuration results in a substantial region of maximum electrostatic potential centered around the sulfur atom. In the [TT]<sup>+</sup> cation (MEP surface in Fig. 4a), all three  $\sigma$ -holes extending from the C-S bonds exhibit MEP maxima with almost identical values, ranging from 107 to 109 kcal mol<sup>-1</sup>. However, in the case of the two Umemoto cations (shown in the lower part of Fig. 4), the maximum MEP values at the  $\sigma$ -holes significantly increase, ranging from 117 to 130 kcal mol<sup>-1</sup>.

The  $\sigma$ -holes opposite the S-C<sub>Ar</sub> bonds are more positive than those opposite the S-CF<sub>3</sub> bond due to the influence of the adjacent hydrogen atoms. Notably, in the [UmeNO<sub>2</sub>]<sup>+</sup> cation (Fig. 4b), the most positively charged  $\sigma$ -hole is opposite to the unsubstituted aromatic ring, driven by the enhanced acidity of the H-atom adjacent to the nitro group. The  $\sigma$ -hole opposite the S-CF<sub>3</sub> bond is less positive due to  $\pi(\text{arene}) \rightarrow \sigma^*(\text{S-C})$  charge transfer through hyperconjugation, reducing the MEP.

ChBs were further analyzed using DFT calculations and quantum theory of atoms-in-molecules (QTAIM), coupled with noncovalent interaction plot (NCIPlot) for real-space visualization. For [UmeNO<sub>2</sub>]<sup>+</sup>[Mo(CO)<sub>5</sub>CN]<sup>-</sup> assembly, only the major component [Mo(CO)<sub>5</sub>CN]<sup>-</sup> was considered in the calculations. Fig. 5 displays the results for the three tetrameric assemblies. Within these analyses, we observe four bond critical points (BCPs, red spheres) and bond paths (solid orange lines) connecting S and N atoms, confirming the existence of S<sub>2</sub>N<sub>2</sub> supramolecular rings (pink in Fig. 5) in all three assemblies. These ChBs are

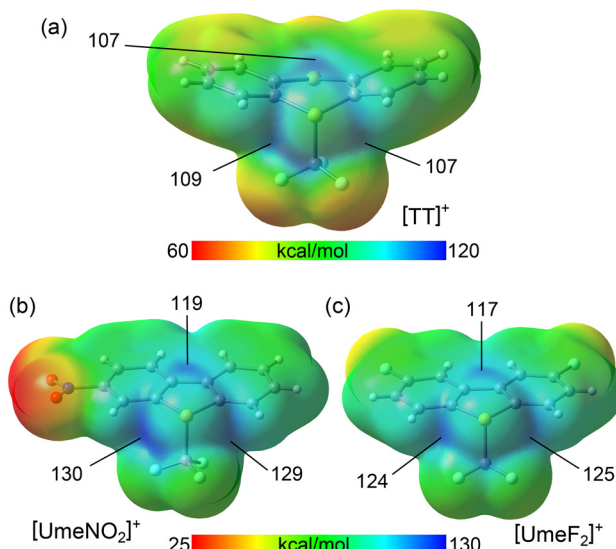


Fig. 4 MEP surfaces (isosurface 0.001 a.u.) of the Ritter (a) and Umemoto (b and c) reactants. The values at the  $\sigma$ -holes are indicated in kcal mol<sup>-1</sup>.

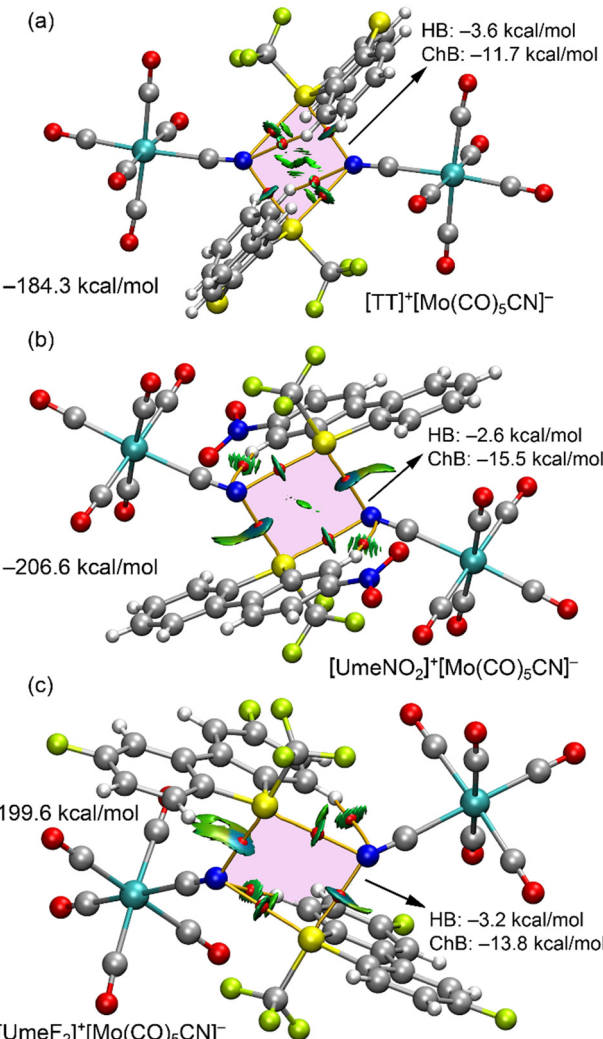
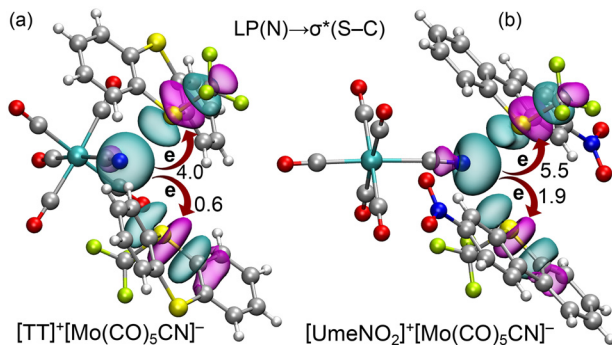


Fig. 5 QTAIM/NCIplot analyses for the tetrameric assemblies of [TT]<sup>+</sup>[Mo(CO)<sub>5</sub>CN]<sup>-</sup> (a), [UmeNO<sub>2</sub>]<sup>+</sup>[Mo(CO)<sub>5</sub>CN]<sup>-</sup> (b) and [UmeF<sub>2</sub>]<sup>+</sup>[Mo(CO)<sub>5</sub>CN]<sup>-</sup> (c). Only intermolecular HB and ChBs are represented for clarity. The energies of the assemblies are also indicated. The contributions of HB and ChB interactions derived from the QTAIM parameters are indicated. NCIplot settings: RDG = 0.5,  $\rho$  cut-off = 0.04 a.u.; colour range  $-0.035 \leq \text{sign } \lambda_{2\rho} \leq 0.035$  a.u.

characterized by green/blue reduced density gradient (RDG) isosurfaces coinciding with the BCP locations. The colour of these RDG isosurfaces indicates stronger N $\cdots$ S-CF<sub>3</sub> interactions than N $\cdots$ S-C<sub>Ar</sub> interactions, consistent with the experimental distances but in disagreement with the MEP surface analysis (the  $\sigma$ -hole opposite the S-CF<sub>3</sub> bond is the least positive). This can be attributed to a shortening of the ChB distance due to the concurrent formation of anion- $\pi$  interaction (see ESI,<sup>†</sup> Fig. S3). Additionally, charge transfer effects play a more prominent role in N $\cdots$ S-CF<sub>3</sub> interactions than in N $\cdots$ S-C<sub>Ar</sub> interactions (*vide infra*). The QTAIM analysis also reveals two ancillary hydrogen bonds (HBs) between N-atoms of both anions and two aromatic C-H bonds, each characterized by a BCP, bond path, and green RDG isosurface.





**Fig. 6** Plot of the NBOs involved in the  $\text{LP}(\text{N}) \rightarrow \sigma^*(\text{S-C})$  for the tetrameric assemblies of  $[\text{TT}]^+[\text{Mo}(\text{CO})_5\text{CN}]^-$  (a) and  $[\text{UmeNO}_2]^+[\text{Mo}(\text{CO})_5\text{CN}]^-$  (b). One of the anions has been omitted for clarity. The second order perturbation energies are given in  $\text{kcal mol}^{-1}$  close to the curved arrows.

The assemblies' formation energies in Fig. 5 are large (ranging from  $-184$  to  $-207 \text{ kcal mol}^{-1}$ ) due to strong coulombic attraction between two anions and two cations. The strongest interaction corresponds to the  $[\text{UmeNO}_2]^+[\text{Mo}(\text{CO})_5\text{CN}]^-$  assembly, consistent with MEP analysis. To assess ChB and HB strength independently of coulombic forces, interaction energies based on QTAIM parameters at the BCPs were calculated (see Theoretical methods, ESI†). The results in Fig. 5 emphasize the dominance of ChBs. For the  $[\text{TT}]^+[\text{Mo}(\text{CO})_5\text{CN}]^-$  assembly, the ChB contribution is  $-11.7 \text{ kcal mol}^{-1}$ , with a HB contribution of  $-3.6 \text{ kcal mol}^{-1}$ . In the case of the  $[\text{UmeNO}_2]^+[\text{Mo}(\text{CO})_5\text{CN}]^-$  assembly (Fig. 5c), HB contribution is almost equivalent to that of  $[\text{TT}]^+[\text{Mo}(\text{CO})_5\text{CN}]^-$  assembly, while ChB contribution increases to  $-13.9 \text{ kcal mol}^{-1}$ . The  $[\text{UmeNO}_2]^+[\text{Mo}(\text{CO})_5\text{CN}]^-$  assembly presents the most substantial ChB contribution of  $-15.5 \text{ kcal mol}^{-1}$ , in agreement with experimental distances and MEP surface analysis.

Finally, we analyzed orbital charge transfer effects in the  $[\text{TT}]^+[\text{Mo}(\text{CO})_5\text{CN}]^-$  and  $[\text{UmeNO}_2]^+[\text{Mo}(\text{CO})_5\text{CN}]^-$  assemblies using the natural bond orbital (NBO) method. Fig. 6 illustrates the relevant orbitals in both assemblies and their associated stabilization energies. It is important to note that, for clarity, we depict only one anion in Fig. 6, though our calculations considered the entire tetrameric assembly. Consequently, only one set of ChB interactions is shown. In both assemblies, we observe electron donation from the LP located at the sp-hybridized N-atom of the anion to the antibonding S-C bonds of the cations. The corresponding  $E^{(2)}$  stabilization energies from the  $\text{LP}(\text{N}) \rightarrow \sigma^*(\text{S-C})$  charge transfer are shown in Fig. 6. In line with interaction energies and experimental distances, the  $[\text{UmeNO}_2]^+[\text{Mo}(\text{CO})_5\text{CN}]^-$  assembly displays higher stabilization energies than  $[\text{TT}]^+[\text{Mo}(\text{CO})_5\text{CN}]^-$ . Furthermore, we note that second-order stabilization energies  $E^{(2)}$  for  $\text{LP}(\text{N}) \rightarrow \sigma^*(\text{S-CF}_3)$  electron donation surpass those for  $\text{LP}(\text{N}) \rightarrow \sigma^*(\text{S-C}_\text{Ar})$ . This heightened electron transfer aligns with shorter equilibrium distances and stronger  $\text{N} \cdots \text{S-CF}_3$  ChBs compared to  $\text{N} \cdots \text{S-C}_\text{Ar}$  interactions which would not be expected if just ESP are taken into account.

In conclusion, our study reveals strong ChB donors in electrophilic trifluoromethylating agents with  $\text{S-CF}_3$  bonds. These compounds offer tunable  $\sigma$ -hole/ESP maxima through aryl ring substitutions on sulfur. We have demonstrated the relevance of supramolecular  $\text{S}_2\text{N}_2$  rings in three crystal structures through X-ray analysis and DFT calculations. These assemblies rely on ChB interactions, with nucleophilic cyanide groups of  $[\text{Mo}(\text{CO})_5\text{CN}]^-$  acting as bifurcated acceptors. Additionally, the QTAIM analysis uncovers weaker H-bonds within these assemblies compared to the chalcogen bonds.

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## Conflicts of interest

There are no conflicts to declare.

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