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# **Inorganic Chemistry**

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Article

## <sup>1</sup> Supramolecular Assemblies in Silver Complexes: Phase Transitions <sup>2</sup> and the Role of the Halogen Bond

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10 arrangement of silver supramolecular complexes. The complexes 11 are composed of X-phenyl(bispyrazolyl)methane (X = Br, I) and I-12 alkynophenyl(bispyrazolyl)methane ligands functionalized in meta 13  $(L^{3Br}, L^{3I})$  and para  $(L^{4Br}, L^{4I}, L^{4CCI})$  positions on a phenyl ring with 14 the purpose of providing different directionalities of the X function 15 with respect to the N,N coordination system. The obtained  $[Ag(L)_2]^+$  moieties show remarkable geometric similarities, and the 16



 $17 L^{4F}$ ,  $L^{4I}$ , and  $L^{4CCI}$  ligands exhibit the most conserved types of supramolecular arrangement that are sustained by XB. The increased 18  $\sigma$ -hole in L<sup>4CCI</sup> with respect to L<sup>4I</sup> leads to an occurrence of short (and strong) XB interactions with the anions. [Ag(L<sup>4I</sup>)<sub>2</sub>]PF<sub>6</sub> and  $19 \left[ Ag(L^{41})_2 \right] CF_3 SO_3$  are characterized by the presence of three different phases, and the single-crystal evolution from phase-1 (a 20 honeycomb structure with large 1D cavities) to phase-3 (solventless) occurs by a stepwise decrease in the crystallization solvent 21 content, which promotes an increase in XB interactions in the lattice. The present paper aims to provide useful tools for the selection 22 of appropriate components for the use of coordination compounds to build supramolecular systems based on the halogen bond.

#### INTRODUCTION 23

24 There is extreme variability in the 1D, 2D, and 3D structural 25 architectures that can be built by combining metal centers and 26 ligands with various functional groups.<sup>1-4</sup> In many cases, these 27 architectures are characterized by properties that depend on 28 structural cavities that can encapsulate additional components 29 such as solvent molecules, counterions, and small mole-30 cules.<sup>3,5-7</sup> Furthermore, these architectures have applications 31 in luminescent materials<sup>8,9</sup> and magnetic materials.<sup>10,11</sup> The 32 presence of metal centers is also a source of potential reactivity 33 toward the interaction and activation of small molecular 34 guests,<sup>12,13</sup> and the formation of channels favors transport to 35 and from the reactive metal centers. Moreover, structural 36 organization in a solid state may result in a close proximity of 37 specific molecular components, which can be made to react by 38 external stimuli, such as UV light, to form covalent bonds that <sup>39</sup> are otherwise difficult to obtain.  $^{14-16}$  In many cases, the ligand 40 is characterized by bridging functionalities; hence it is capable 41 of linking two, or more, metal centers in an extended network. 42 However, multidimensional architectures can be built by 43 combining coordination compound assemblies, which may be <sup>44</sup> connected due to weak interactions,<sup>17,18</sup> such as hydrogen <sup>45</sup> bonds,<sup>9,19–21</sup> halogen bonds,<sup>22,23</sup> and  $\pi - \pi$  stackings.<sup>24,25</sup> For 46 example, dimerization of carboxylic functional groups form

hydrogen bond (HB) dimers,<sup>26–28</sup> which represent a route for 47 expanding the dimensionality of these architectures.

In the present work, we wish to investigate the formation of 49 supramolecular architectures formed by silver bis(pyrazolyl)- 50 methane complexes functionalized with halogen atoms such as 51 bromine and iodine (L<sup>3Br</sup>, L<sup>3I</sup>, L<sup>4Br</sup>, L<sup>4I</sup>, L<sup>4CCI</sup>, Schemes 1 and 52 s1s2 2). The rationale behind the present work is to generate 53 s2 supramolecular architectures by exploring different types of 54 covalent and noncovalent interactions, particularly halogen 55 bonds.<sup>29</sup> The halogen bond is a weak interaction that has been 56 extensively investigated in recent decades,<sup>30-35</sup> and it occurs 57 due to the presence of a partial positive charge ( $\sigma$  hole, Cl < Br <sub>58</sub> < I) on the opposite side of the  $\sigma$  bond.<sup>36,37</sup> Halogen bond 59 interactions have been used to sustain the formation of 60 multidimensional networks based on coordination compounds 61 as repetitive units.<sup>38-43</sup> To compare the influence of halogen 62 atoms on the structural outcome, we investigate a parent 63

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Scheme 1. (a) Previously Reported Supramolecular Architecture Based on a Ag-S Coordination and (b) an Example of a Molecular Crystal Obtained by Exploiting Halogen Bonds and Other Weak Interactions, As Reported in This Work



Scheme 2. (a) Bis-pyrazolyl Ligands Used in the Present Study and (b) a Summary of the Complexes Synthesized in This Work



64 phenylbis(pyrazolyl)methane ligand that is devoid of halogen 65 atoms. However, we increase the XB donor potential of these 66 ligands with an alkyno group interposed by an iodine and a 67 phenyl ring. The resulting ligand ( $L^{4CCI}$ ) is characterized by a 68 large positive charge in the  $\sigma$ -hole, which is potentially capable 69 of forming strong XB interactions.

The complex building units are prepared by reacting the reacting the reacting the reacting with silver salts (AgPF<sub>6</sub> and AgCF<sub>3</sub>SO<sub>3</sub>). Silver usually 98

adopts coordination geometries that vary from linear to 72 tetrahedral.<sup>44–65</sup> Due to the lack of electronic stabilization ( $d^{10}$  73 metal ion), these geometries are usually distorted, and the 74 metal adapts to the steric requirements of the ligands. Thus, 75 two different counteranions are used to provide different 76 systems that can potentially act as halogen bond acceptors 77 (fluorine in  $PF_6^-$  or fluorine and oxygen in  $CF_3SO_3^-$ ) as well 78 as having different coordination capabilities toward a metal 79 center. Halogen bonds are present in all the structures 80 obtained with the halogenated ligands, and in two cases, 81 when crystallizing  $[Ag(L^{4I})_2]PF_6$  and  $[Ag(L^{4I})_2]CF_3SO_3$  in 82 tetrahydrofuran/hexane (THF/Hx), we identify three phases, 83 which differ by their solvent content. Phase-1 is characterized 84 by a honeycomb structure with large 1D channels (diameter of 85 30 Å), and they evolve into phase-2 after a spontaneous loss of 86 most of the solvent. Phase-2 presents smaller 1D channels that 87 confine THF molecules, and the evolution from phase-2 to a 88 solventless phase-3 can be promoted by a thermal treatment at 89 120 °C. The phase-1 to phase-3 transitions are investigated by 90 thermal methods and powder X-ray diffraction. The single- 91 crystal X-ray structures allow the role of the weak interactions 92 in the formation of these supramolecular arrangements to be 93 established. Overall, the analysis of the structural arrangements 94 wishes to provide useful hints for the selection of the 95 appropriate components in the formation of supramolecular 96 systems based on coordination compounds as repetitive units. 97

#### RESULTS AND DISCUSSION

The Ag<sup>+</sup> complexes are prepared by mixing AgPF<sub>6</sub> and 99 AgCF<sub>3</sub>SO<sub>3</sub> with different ligand (L) systems in a 1/2 ratio. The 100 choice of the two different counteranions is made by 101 considering their different symmetries and potential inter- 102 actions with metal centers. In particular, the highly symmetric 103 PF<sub>6</sub><sup>-</sup> anion exhibits little coordination capability toward Ag<sup>+</sup>. 104 In contrast, the CF<sub>3</sub>SO<sub>3</sub><sup>-</sup> anion has a more pronounced 105 tendency to interact with a metal center, usually as an O- 106 monodentate.<sup>49-52</sup> Both anions, having a net negative charge, 107 can also act as nucleophilic sites for interactions involving 108 positively charged molecular fragments, such as the  $\sigma$ -holes 109 present in halogen atoms or electrophilic CH groups. 110

The coordination geometries of all of the complexes 111 reported in the present work are depicted in Figures S26- 112 S58. In the structures, the metal appears to be in a distorted 113 tetrahedral environment, but it is difficult to assign a definitive 114 metal geometry to these four-coordinated silver complexes. 115 Hence, it is perhaps useful to employ a  $\tau_4$  index that takes into 116 account all the possible distortions from the square-planar to 117 the tetrahedral geometries.<sup>66,67</sup> In all complexes, the  $\tau_4$  and  $\tau_4'$  <sup>118</sup> geometry indices are in the ranges 0.70-0.61 and 0.65-0.46, 119 respectively, which are in line with a seesaw geometry (Table 120 S10). More specifically, two of the four Ag–N coordinative 121 interactions are longer (range 2.34–2.49 Å), and two are 122 shorter (2.21-2.31 Å, Tables S7-S9). As a general 123 observation, in all the reported systems, the metal environment 124 is quite conserved. Hence, we may assume that  $[Ag(L)_2]^+$  is a 125 preorganized synthon, with a relatively fixed orientation of 126 phenyl rings attached to a bis-pyrazole moiety. Consequently, 127 the different functionalizations of the aromatic ring, with 128 respect to the 3 (meta) and 4 (para) positions, is what dictates 129 the orientations of the halogen atoms (Figure 1). When the 130 fl halogen atoms are in the 4 position, the conformation of the 131 phenyl ring has little influence on the directionality of the C-X 132 vector. In contrast, functionalization of the 3 position leads to 133

f2



**Figure 1.** Schematic representation of the multiple experimental orientations, which can be experienced by the halogen atoms after the formation of the Ag complex. The indicated values refer to the observed angles between the C-halogen vector (color codes: C, gray; N, blue; Br, light brown; P, orange; I, purple).

134 different arrangements of the two C–X vectors located on the 135 two ligands. In Figure 2, we report an overview of the 136 intermolecular interaction exchange between the halogen 137 atoms and the surrounding molecules. More specifically, it 138 can be pointed out how the presence of the halogen atom at



**Figure 2.** Summary of the XB interactions exchanged by the halogen atoms in the reported structures. The inset describes a simplified representation of XB between two halides and one anion (color codes, C, gray; N, blue; O, red; H, white; F, green; S, yellow; Br, light brown; P, orange; I, purple).

position 4 gives rise to two crystal packing motifs involving the 139 anions. In the first one, one halogen atom acts as an XB donor 140 toward the anion, and it simultaneously acts as an XB acceptor 141 toward a second halogen atom (Figure 2h). In the second 142 motif, the anion bridges between two XB donor moieties 143 (Figure 2i,l).<sup>68</sup> The following discussion will focus on the 144 solid-state properties of the silver complexes with the L<sup>41</sup> and 145 L<sup>4CCI</sup> ligands (9–12; see Scheme 2), which are the systems 146 characterized by preorganized orientations of the two halogen 147 atoms within the  $[Ag(L)_2]^+$  moiety. The presence of the alkyne 148 group in L<sup>4CCI</sup> is devised to increase the XB donor properties 149 of the ligand with respect to L<sup>41</sup> while preserving the same 150 spatial orientation of the halogen.<sup>30</sup>

#### SOLVENT ROLE AND PHASE TRANSITIONS 152

The **9–12** complexes are crystallized in different solvent 153 mixtures since it was previously shown that the solvent can 154 have an important role in selecting the type of weak interaction 155 that occurs between two multifunctional molecular counter- 156 parts.<sup>69</sup> In our case, the choice of solvent is limited by 157 solubility issues related to the  $[Ag(L)_2]^+$ /anion complexes, 158 which can dissolve in chlorinated solvents (DCM, DCE) or 159 more polar solvents (acetone, THF, methanol). The 160 crystallizing conditions make use of hexane as an antisolvent. 161 The crystalline material for  $[Ag(L^{41})_2]PF_6$  (**9**) and  $[Ag(L^{41})_2]$ - 162 CF<sub>3</sub>SO<sub>3</sub> (**10**) in the form of different solvates is recovered 163 from the following crystallization conditions: dichloroethane/ 164 hexane (DCE/Hx), dichloromethane/hexane (DCM/Hx), and 165 THF/Hx. 166

More interesting results are obtained from the THF/Hx 167 crystallizing condition, since three different phases can be 168 observed, and the evolution from phase-1 to phase-3 occurs 169 upon decreasing the solvent content. 9-phase-1 and 10-phase-1 170 are characterized by long prismatic crystals (Figure 3 and 171 f3 Figure 4), which are stable in the presence of the mother liquor 172 f4 or when protected by a low temperature environment (less 173 than 200 K). The single-crystal X-ray characterization of these 174 systems shows that they are isostructural, forming large 1D 175 hexagonal cavities filled with solvent molecules (10-phase-1 176 will be described here). When proceeding from the molecular 177 unit to the crystal packing, it can be instructive to analyze the 178 hierarchical construction of the lattice based on the weak XB, 179 CH…F, CH…O, and CH… $\pi$  interactions. In particular, three 180 molecular units in the inner core are arranged around a  $PF_6^-$  or 181 CF<sub>3</sub>SO<sub>3</sub><sup>-</sup> anion (P18/S18) by means of CH···F or CH···O 182 interactions. Interestingly, toward the periphery of the trimer, 183 the inner iodine atoms of the molecular unit (I6) are directing 184 the  $\sigma$ -hole toward the negative corona of the outermost iodine 185 atom (I3). Even though the halogen-halogen intermolecular 186 distance is particularly long (~4.5 Å), the geometry of the 187 interaction is preserved, since the C-I6…I3 angle is 188 approximately  $160^{\circ}$  and close to the theoretical value of 189 180°. The supramolecular trimers are then piled one over the 190 other by means of CH…F and CH…O interactions mediated 191 by anions bridging different stacking levels along the c-axis 192 (Figure 3b). The expansion of the columnar stacking of trimers 193 along the *ab* crystallographic plane is promoted by the anion 194 (P17/S17) that links together two adjacent complex molecules 195 by means of CH···F interactions (Figure 3c). The final 196 supramolecular arrangement results in the formation of a 197 honeycomb structure characterized by hexagonal 1D cavities 198 that are parallel to the crystallographic *c*-axis. Even though the 199 data collection is performed at low temperature, it is possible 200



**Figure 3.** Crystal structure of **10**-phase-1 crystallized in THF/Hx. (a) Supramolecular trimer. (b) Trimeric units piled along the *c*-axis. (c) Expansion of the trimeric unit in the ab crystallographic plane. (d) Crystal packing highlighting the 1D channels.



Figure 4. (A)–(D) Polarized microscopy images collected at different points in time showing the phase transition from 10-phase-1 (needles) to 10-phase-2 (plates) at RT. (E,F) Hot-stage microscopy images showing the phase transition from 10-phase-2 to 10-phase-3.

to identify only some THF molecules in these cavities during 201 crystallization, which are close to the surface of the hexagonal 202 channel. One of these THF molecules exchanges an XB with 203 the I3 atom, which points toward the interior of the cavity 204 (I3…O 1s 3.24 Å, see Figure 2f). It is not possible to identify a 205 reasonable structural model of the solvent into the remaining 206 part of the large 1D cavity, and the Squeeze program is used to 207 determine the residual and diffuse electron density. By taking 208 into account only the complex molecular entity  $[Ag(L^{4I})_2]$ - 209 CF<sub>3</sub>SO<sub>3</sub>, which is the building unit of the hexagonal 210 framework, the cavity volume corresponds to 7300 Å<sup>3</sup>/cell 211 (56% of the unit cell volume), and the 1D cavity diameter is 212 approximately 30 Å (Figure 3d). When the prismatic needle- 213 like crystals of 10-phase-1 are taken out of the mother liquor, 214 they rapidly change phase and convert into 10-phase-2 (Figure 215 4A-D, a video of the single-crystal phase transition is available 216 as a web enhanced object). The time span of the conversion 217 process usually takes significantly less than a minute, and it can 218 be partially hampered by immersing the crystals into viscous 219 matrixes, but the conversion eventually goes to completeness. 220 The 10-phase-2 crystals exhibit a plate-like morphology, and 221 according to the single-crystal X-ray analysis, they contain one 222 THF molecule per  $[Ag(L^{4I})_2]CF_3SO_3$  complex (Figure 5). 223 fs The complex molecules and the CF<sub>3</sub>SO<sub>3</sub><sup>-</sup> anions delimit a 224 small channel-like cavity (~5 Å diameter) hosting the THF 225 molecules. All of the iodine atoms are engaged in XBs with the 226 oxygen or fluorine atoms of the anion, and the I…F interaction 227 distances (3.27 and 3.38 Å) are significantly longer than the I··· 228 O interaction distances (2.97 and 2.99 Å). 9-phase-2 is 229 isomorphous with 10-phase-2, as reported in the Supporting 230 Information (Table S1 and Figure S44). 10-phase-2 is stable 231 for several weeks at room temperature; however, when the 232 plate-like crystals are heated above 130 °C, they convert into 233 10-phase-3, as reported in Figure 4E,F (a video of the SC to 234 SC phase transition is available as a web enhanced object). 235 After the thermal treatment, the crystals were partially 236 fractured, but a small sample could be used for the single- 237 crystal X-ray data collection (see Figure 5f). The structural 238 analysis revealed that the system has experienced a significant 239 spatial reorganization resulting in a more compact structure 240 with negligible residual voids; hence, 10-phase-3 can be 241 considered the completely desolvated phase. Only I3 242 exchanges an XB with an oxygen atom of the anion (3.14 Å; 243 see Figure 2g), whereas the second iodine atom I6 interacts 244 with a methyl group by virtue of its negatively charged corona 245 surrounding the  $\sigma$ -hole. Interestingly, 10-phase-3 exhibits a 246 different structural organization than 9-phase-3. The latter 247 phase is also closely packed and forms without a solvent. 248 Nevertheless, in this case, both iodine atoms act as XB donors, 249 in line with one of the structural motifs involving the halogen 250 atoms reported in Figure 2h. In fact, I3 simultaneously acts as 251 an XB donor (toward the anion, I6…F17, 3.221 Å) and an 252 acceptor (toward I6, I3…I6, 3.646 Å). 9-phase-3 can be 253 obtained after a thermal treatment at 130 °C of 9-phase-2, but 254 interestingly, it can also be obtained by direct crystallization 255 from a DCM/Hx mixture (Figure 6). 256 f6

#### POWDER X-RAY DIFFRACTION

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Powder X-ray diffraction is performed for compounds **9** and **10** 258 to confirm the phase composition in the bulk and to monitor 259 the conversion between the three phases. For both 260 compounds, phase-1 is the most difficult to characterize 261 experimentally since the long prismatic crystals are very 262



Figure 5. Views of the intermolecular interactions (a) and crystal packing (c) for 10-phase-2 (THF molecules in a space-filling model). Single-crystal to single-crystal phase transition from 10-phase-2 to 10phase-3 due to the loss of solvent and thermal treatment (b, d, f). Representation of the interactions (dashed bonds) exchanged by the anions and the complex molecules for 9-phase-3 (e). The insets in (b) and (d) describe the decrease in the residual electron density associated with the THF molecules.

263 unstable once removed from the mother liquor (THF/Hx). 264 Wet crystals are placed onto the sample holder and covered <sup>265</sup> with a protective film as described in the experimental section. 266 The crystals are not ground to prevent phase transformations; 267 hence, the data collection is affected by significant preferential 268 orientation. Figure 7 shows the comparison between the 269 experimental PXRD of 10-phase-1 with the calculated patterns 270 derived by SC-XRD.<sup>70</sup> There is good agreement between the 271 predicted and experimental diffractograms, which is partic-272 ularly true for the peak positions, while the relative intensities 273 are affected by the cited strong preferential orientation and 274 sample roughness. The conversion of phase-1 into phase-2 is 275 very fast, and it went to completion in approximately 30 s after 276 the removal of the film and the evaporation of the solvent. The 277 rapid evolution from phase-1 to phase-2 implies that the 278 investigation based on the thermal analysis (DSC and TGA; 279 see below) exclusively involves phase-2 even when starting 280 from the long prismatic crystals of phase-1. The plate-like 281 crystals of phase-2 are stable for several days; however, it is 282 interesting to note that the solvent content of phase-2 is not



Figure 6. Schematic representation of the phase transitions and the solvates of 9 and 10 described in this work.

constant over time. In fact, the SC-XRD data collection of a 283 freshly mounted crystal shows one THF molecule per complex 284 molecular entity, whereas after the crystal is left for 24 h at 285 room temperature and in the absence of the crystallizing 286 solvent, the solvent content decreases to 0.25 per complex 287 molecule (compare Figure 5b,d). This observation is in line 288 with the presence of the narrow channels that characterize 289 phase-2 and that contain the THF molecules, which weakly 290 interact with complex cations and anions. The relatively small 291 size of the channels implies that the removal of the solvent can 292 be tolerated without the system undergoing structural 293 reorganization. Phase-2 of the two compounds is isostructural, 294 but after the thermal treatment, they give rise to two different 295 phase-3's (Figures 7g and S21). 296

## THERMAL ANALYSIS

As pointed out in the PXRD section, the transformation from 298 hexagonal phase-1 to phase-2 for both systems is very rapid 299 once they are removed from the mother liquor (THF/Hx), 300 and it occurs in a time span of a few seconds. Hence, the 301 thermal experiments monitor the events occurring between 302 phase-2 and phase-3. The hexagonal prims of 10-phase-1 are 303 dried prior to performing the experiments to remove the 304 solvent wetting the crystals. The DSC profile shows the 305 presence of an exothermic peak at 113 °C with a shoulder at 306 the beginning of the event (Figure 7B). The exothermic peak is 307 associated with the phase-2 to phase-3 conversion, whereas a 308 minor endothermic event can be associated with the residual 309 THF desorption within the channels of 10-phase-2. There are 310 no other events occurring when decreasing the temperature to 311 room temperature (RT) and then increasing the temperature 312 above 200 °C. The TGA profile of 10-phase-2 shows no 313 appreciable weight loss at approximately 120 °C, but the 314 compound undergoes decomposition at 221 °C. 315

For 9, we grew plate-like crystals corresponding to 9-phase- 316 2, which were then used for the thermal experiments (Figure 317 S17). In the DSC scan, there is an endothermic peak at 111 °C 318 associated with the loss of the solvent into the small channels, 319 and concomitantly, the systems experience a structural 320 reorganization to the more stable 9-phase-3. There are no 321 other events taking place when decreasing the temperature to 322

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Figure 7. (A) Overlap between the simulated and experimental XRD of 10: 10-phase-1 (a,b), 10-phase-2 (c,d), 10-phase-3 (f,g). (B) DSC and (C) TGA traces for 10-phase-2.

<sup>323</sup> RT and then increasing the temperature above 200 °C. At 264 <sup>324</sup> °C, the system undergoes decomposition. The loss of solvent <sup>325</sup> at 111 °C is confirmed by a 5% weight loss, in agreement with <sup>326</sup> 1 molecule of THF per  $[Ag(L^{4I})_2]PF_6$  (6% theoretical), Figure <sup>327</sup> S18.

## 328 A LONGER SYNTHON: [AG(L<sup>4CCI</sup>)<sub>2</sub>]<sup>+</sup>

329 As pointed out earlier in the discussion, the presence of an 330 iodoalkyne functional group confers to the ligand a greater 331 propensity to act as an XB donor with respect to the L<sup>41</sup> ligand, 332 with L<sup>4CCI</sup> having a more pronounced *σ*-hole and negative 333 corona on the halogen atom. The molecular structures of 334 [Ag(L<sup>4CCI</sup>)<sub>2</sub>](PF<sub>6</sub>)·THF (11·THF) and [Ag(L<sup>4CCI</sup>)<sub>2</sub>]-335 (CF<sub>3</sub>SO<sub>3</sub>) (12) are reported in Figure 8. In both compounds, 336 the anions bridge two complex cations by means of an XB with



Figure 8. Halogen bond interactions for  $11 \cdot \text{THF}$  (a) and 12 (c). Schematic representations of the supramolecular chains in  $11 \cdot \text{THF}$  (b) and 12 (d).

the peripheral iodine atoms, in agreement with the second 337 motif described above for the XB reported in this work (see 338 Figure 2l). In particular, in 12, CF<sub>3</sub>SO<sub>3</sub><sup>-</sup> acts as an XB acceptor 339 toward two opposite iodine atoms (I3…O1T, 2.844 Å; I6… 340 F2TB, 3.059 Å); in 11. THF,  $PF_6^-$  engages two cis-fluorine 341 atoms into an XB formation with symmetry-related iodine 342 atoms (I3…F24B, 2.987 Å; I3…F24A, 2.960 Å). The result is 343 the formation of supramolecular chains in both cases, even 344 though they each express a different structural packing. In fact, 345 in 12, the chains are more interwoven, leaving no residual void 346 for the presence of solvent during crystallization. In contrast, in 347 11. THF, the chains delimit channel-like cavities filled with 348 disordered THF molecules during crystallization. Obviously, in 349 11. THF and 12, the presence of the Ph—C $\equiv$ C—I group with 350 respect to the Ph-I moiety in 9 increases the dimension of 351 the supramolecular synthon, which is represented by the 352  $[Ag(L)_2]^+$  complex cation. However, by comparing the 353 structures of 11. THF, 12, and 9/10-phase-1, it is evident 354 that increasing the length of the XB donor moiety does not 355 produce systems characterized by large channels as in the 9/ 356 **10**-phase-1. 357

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The ligands described here are characterized by two functions, 359 one that can bind a metal center (N,N system) and the other 360 that can form directional supramolecular interactions (halogen 361 atoms). The assemblies generated with Ag<sup>+</sup> result in an almost 362 invariant molecular geometry, with the cation in a well-defined, 363 even though distorted, tetrahedral environment. The reason for 364 such behavior can be ascribed to the presence of the moderate 365 steric hindrance provided by the methyl groups attached to the 366 pyrazole rings. Consequently, when the two ligands are bound 367 to the metal center, they are interlocked to minimize steric 368 repulsion. The result is a preorganized XB donor directionality, 369 which exclusively depends on the position of the halogen 370 functionalization on the phenyl ring. When comparing all of 371

358

The [Ag(L<sup>4I</sup>)<sub>2</sub>]PF<sub>6</sub>/CF<sub>3</sub>SO<sub>3</sub> systems crystallized from THF/ 378 379 Hx are characterized by the presence of three phases. The 380 conversion from phase-1 to phase-3 occurs by a stepwise 381 decrease in the solvent content and by a reinforcement of the 382 XB interactions among the molecular components. 9/10-383 phase-1 presents a supramolecular structure with a large 384 columnar cavity filled with solvent molecules, and we have not 385 investigated the various methodologies that are suitable to 386 activate porous materials that are characterized by large cavities occupied with solvent-guest molecules. These methods are 387 usually applied to metal organic frameworks,<sup>71</sup> in which the 388 weakest type of interaction is the metal-ligand coordination 389 390 bond, which is significantly stronger than the XB or other weak 391 interactions found in the reported structures.

Even though the Ph-Br and Ph-I moieties may be 392 393 considered moderately good XB donors, the XB donor 394 potential of these ligands can be increased by introducing 395 additional electron withdrawing groups on the aromatic 396 fragment, hence increasing the positive charge of the  $\sigma$ -<sup>397</sup> hole.<sup>30</sup> L<sup>4CCI</sup> was therefore devised to increase the strength of 398 the XB interactions. A summary of the normalized 399 distances<sup>72-74</sup> pertaining to the XBs described here is provided 400 in Figure 9. Generally, it is clear that shorter interactions are



Figure 9. Plot of the normalized distance R for the XB interactions. A value of 1 corresponds to an XB donor-acceptor distance equal to the sum of the v.d.W. radii.

401 experienced with the oxygen atom of the CF<sub>3</sub>SO<sub>3</sub><sup>-</sup> anion and 402 iodine compared to those of the other interactions. This 403 observation is also in agreement with the propensity of 404 CF<sub>3</sub>SO<sub>3</sub><sup>-</sup> to act as an O-monodentate ligand toward a metal 405 center when compared to  $PF_6^-$ . It is also evident that there is a 406 significant decrease in the XB distance along the Br < I < C $\equiv$ 407 C—I series. This observation is consistent with a more 408 pronounced  $\sigma$ -hole on L<sup>4CCI</sup> with respect to L<sup>4I</sup>. As a result, 409 more robust interactions can potentially be formed with this 410 ligand, and the high directionality of the XB allows for 411 significant control over the supramolecular arrangements. The 412 studied systems provide insight into the choice of molecular 413 components for the construction of supramolecular assemblies 414 comprising coordination entities as building units.

#### ASSOCIATED CONTENT 415

## 416 Supporting Information

417 The Supporting Information is available free of charge at 418 https://pubs.acs.org/doi/10.1021/acs.inorgchem.0c00256.

Synthesis of the ligands and complexes, NMR spectra, 419 single-crystal structures, optical microscope images, 420 crystallographic tables, geometric parameters, au indices, 421 thermal analyses (DSC and TGA), and powder X-ray 422

## Web-Enhanced Features

diffraction spectra (PDF)

Single-crystal to single-crystal phase transitions 10-phase-1/10- 425 phase-2/10-phase-3 in an MPEG format are available in the 426 HTML version of the paper. 427

#### **Accession Codes**

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CCDC 1902455-1902465 and 1968843-1968848 contain 429 the supplementary crystallographic data for this paper. These 430 data can be obtained free of charge via www.ccdc.cam.ac.uk/ 431 data request/cif, or by emailing data request@ccdc.cam.ac. 432 uk, or by contacting The Cambridge Crystallographic Data 433 Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 434 1223 336033. 435

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