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# Hierarchy of Supramolecular Arrangements and Building Blocks: Inverted Paradigm of Crystal Engineering in the Unprecedented Metal Coordination of Methylene Blue

Stefano Canossa,<sup>†</sup> Alessia Bacchi,<sup>†</sup> Claudia Graiff,<sup>†,‡</sup> Paolo Pelagatti,<sup>†,‡</sup> Giovanni Predieri,<sup>\*,†,‡</sup> Andrea Ienco,<sup>§</sup> Gabriele Manca,<sup>§</sup> and Carlo Mealli<sup>\*,§</sup>

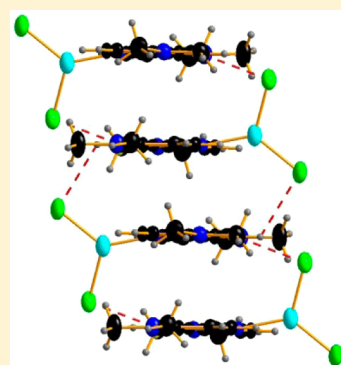
<sup>†</sup>Dipartimento di Chimica, Università degli Studi di Parma, Area delle scienze 17/A, 43124 Parma, Italy

<sup>§</sup>ICCOM-CNR, Via Madonna del Piano 10, 50019 Sesto Fiorentino, Florence, Italy

<sup>‡</sup>Consorzio Interuniversitario Reattività Chimica e Catalisi, Via Celsio Ulpiani 27, 70126 Bari, Italy

**S** Supporting Information

**ABSTRACT:** The aromatic methylene blue cation (MB<sup>+</sup>) shows unprecedented ligand behavior in the X-ray structures of the trigonal-planar (TP) complexes MBMCl<sub>2</sub> (M = Cu<sup>I</sup>, Ag<sup>I</sup>). The two isostructural compounds were exclusively synthesized by grinding together methylene blue chloride and MCl solids. Only in the case of AuCl did the technique lead to a different, yet isoformular, Au<sup>I</sup> derivative with separated MB<sup>+</sup> and AuCl<sub>2</sub><sup>-</sup> counterions and no direct N–Au linkage. While the density functional theory (DFT) molecular modeling failed in reproducing the isolated Cu and Ag complexes, the solid-state program CRYSTAL satisfactorily provided for Cu the correct TP building block associated with a highly compact  $\pi$  stacking of the MB<sup>+</sup> ligands. In this respect, the dispersion interactions, evaluated with the DFT functional, provide to the system an extra energy, which likely supports the unprecedented metal coordination of the MB<sup>+</sup> cation. The feature seems governed by subtle chemical factors, such as, for instance, the selected metal ion of the coinage triad. Thus, the electronically consistent Au<sup>I</sup> ion does not form the analogous TP building block because of a looser supramolecular arrangement. In conclusion, while a given crystalline design is generally fixed by the nature of the building block, a peculiarly efficient supramolecular packing may stabilize an otherwise unattainable metal complex.

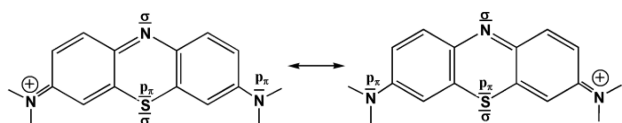


## INTRODUCTION

Methylene blue (MB) is a salt of the formula [(C<sub>16</sub>H<sub>18</sub>SN<sub>3</sub>)<sup>+</sup>Cl<sup>-</sup>] [IUPAC name: 3,7-bis(dimethylamino)-phenothiazin-5-ium chloride]. The MB<sup>+</sup> cation consists of three condensed six-membered rings and two coplanar NMe<sub>2</sub> substituents. The aromaticity is suggested by the total 18 p $\pi$  electron count, as indicated from the resonant structures in Scheme 1.

A qualitative molecular orbital (MO) picture<sup>1</sup> corroborates the donor weakness of the phenothiazine N atom, given that its in-plane  $\sigma$  lone pair lies well below the frontier MO region. The lowest unoccupied MO is instead an accessible  $\pi^*$  level, favoring the N-atom reduction to an amide, with the system's puckering due to the lost aromaticity. The acquired N

Scheme 1. Resonant Structures of the 18e<sup>-</sup> Aromatic MB<sup>+</sup>



<sup>a</sup>Electron pairs at the heteroatoms are distinguished for their p $\pi$  or  $\sigma$  character.

basicity<sup>2,3</sup> is also consistent with the known N–H/N–R<sub>40</sub> derivatives,<sup>4</sup> also stable in the monoxidized form.<sup>5,6</sup> For the redox reversibility, promoted by the alternative glucose and O<sub>2</sub> 42 reactants and the associated color change, MB is often used as an indicator.<sup>7</sup> Other applications are in photocatalysis<sup>8,9</sup> and 44 enzyme-catalyzed redox reactions. As a drug, MB is employed for treating diseases such as methemoglobinemia,<sup>10</sup> cyanide,<sup>11,12</sup> carbon monoxide poisoning,<sup>13</sup> malaria,<sup>14–16</sup> and 47 Alzheimer's.<sup>17</sup> In these fields, the combination of MB with a 48 metal center could be, in principle, advantageous, but its 49 coordination capabilities remain elusive in the literature. 50 Perhaps, the closest example of bonding to a metal is in the 51 crystal structure of MB<sup>+</sup>[HgCl<sub>3</sub>]<sup>-</sup>,<sup>18</sup> although the N–Hg 52 distance of 2.779 Å exceeds the sum of the covalent radii as well as the average value, evaluated from the Cambridge 54 database.<sup>19</sup> Other invoked cases of MB<sup>+</sup> coordination to a metal 55 are even more speculative in the lack of structural characterization.<sup>20,21</sup> 57

This paper provides the first evidence of the MB<sup>+</sup> ligand 58 behavior based on the structures of the three-coordinated 59 complexes [(MBMCl<sub>2</sub>)] with M = Cu<sup>I</sup> or Ag<sup>I</sup>. Support from 60

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61 suitable density functional theory (DFT) calculations is also  
 62 provided when taking into account crystalline supramolecular  
 63 interactions. Among the latter, the London dispersion forces  
 64 play a decisive role in some sterically encumbered organic  
 65 compounds, as emphasized by a recent review article.<sup>22</sup> The  
 66 role is instead much less considered for transition-metal  
 67 complexes, concerning, in particular, possible effects on metal  
 68 coordination. A conclusion reached by this study is that the  
 69 supramolecular interactions, among which is the  $\pi$  stacking of  
 70 aromatic units, can end up conferring unknown coordination  
 71 capabilities to an unsuited species such as  $\text{MB}^+$ .

## 72 ■ RESULTS AND DISCUSSION

73 Only by using a mechanochemical strategy was the synthesis of  
 74 complexes  $[(\text{MBCuCl}_2)]$  and  $[(\text{MBAgCl}_2)]$  (simplified as  
 75 MBCu and MBAg) successfully achieved. This implied the  
 76 grinding together of the solid methylene blue chloride  
 77 pentahydrate and CuCl or AgCl. Conversely, the equivalent  
 78 combination of reactants in solution failed. The progress of the  
 79 reactions was monitored by powder X-ray diffraction (PXRD)  
 80 analysis. In the case of Cu, the traces of the starting reagents  
 81 disappeared completely after 30 min of grinding without the  
 82 addition of a solvent's drop. In the case of Ag, a few drops of  
 83 acetone had to be added to complete the reaction within 20  
 84 min. In both cases, the color changed from dark green to  
 85 reddish brown, while some PXRD traces of the final powders  
 86 indicated the attainment of new crystalline phases (Figure 1).

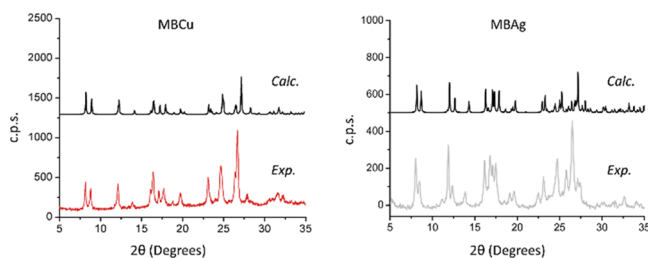
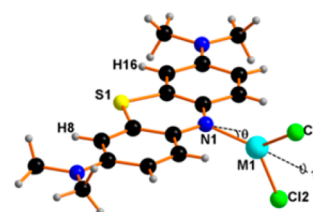


Figure 1. Experimental and calculated PXRD patterns of MBCu and MBAg products. Distinct positions of the experimental and calculated signals likely depend on the temperature of the data collection (298 and 100 K for PXRD and single-crystal diffraction, respectively).

87 Suitable crystals for X-ray analyses were isolated upon slow  
 88 evaporation of the dimethylformamide (MBCu) and acetonitrile  
 89 (MBAg) solutions of the ground solids. As shown in  
 90 Figure 1, the recorded PXRD patterns coincide with those  
 91 calculated from the determined single-crystal X-ray structures.<sup>23</sup>  
 92 Select geometric parameters of the isostructural MBCu and  
 93 MBAg compounds are shown in Table 1, where the values in  
 94 brackets are those of the MBCu optimization with CRYSTAL.<sup>24</sup>  
 95 Approximate trigonal-planar (TP) metal complexes are  
 96 formed by the two  $\text{Cl}^-$  and the  $\text{MB}^+$  ligands. This result is  
 97 surprising because, in general, a cationic ligand hardly forms an  
 98 uncharged metal complex.<sup>25–28</sup> The behavior of  $\text{MB}^+$  as a  $2e^-$   
 99 donor is corroborated by the relatively short N–M linkages of  
 100 1.987(4) and 2.329(5) Å in the Cu and Ag compounds,  
 101 respectively. The Ag coordination is comparatively less effective  
 102 because the N–Ag bond is larger than the sum of the covalent  
 103 radii and the average value from all of the corresponding  
 104 CCDC structures.<sup>19</sup> The point is also supported by the  
 105 somewhat more open Cl–Ag–Cl angle (119.48° vs 113.73°).  
 106 In no case is the  $\text{Cl}_2\text{M}$  fragment either coplanar or orthogonal  
 107 to the  $\text{MB}^+$  plane, thus excluding any higher symmetry of the

Table 1. Experimental Geometric Parameters (Å and deg) of the Isostructural Complexes  $(\text{MB})\text{MCl}_2$  ( $\text{M} = \text{Cu}, \text{Ag}$ )<sup>24</sup>



	MBCu	MBAg
M1–Cl1	2.2220(5) [2.18]	2.4425(19)
M1–Cl2	2.2494(5) [2.19]	2.4659(18)
M1–N1	1.9874(13) [1.91]	2.329(5)
Cl1–M1–Cl2	113.739(18) [111.0]	119.48(6)
Cl1–M1–N1	129.45(4) [126.8]	125.09(13)
Cl2–M1–N1	116.70(4) [122.2]	115.32(13)
$\theta$	11.8(4) [2.5]	9.5(2)
$\tau$	75.2(2) [57.6]	66.8(2)

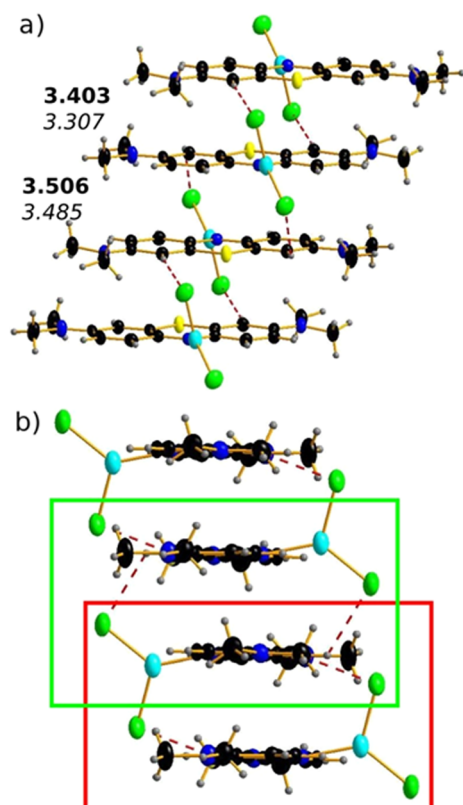
<sup>a</sup>The values in square brackets are for the Cu model, optimized with CRYSTAL.<sup>24</sup>

complexes. The corresponding  $\tau$  rotation of  $\text{Cl}_2\text{M}$  about the  
 N–M linkage has intermediate values of 75.2 and 66.8° for  
 MBCu and MBAg, respectively. Another important parameter,  
 defined in Table 1, concerns the out-of-plane shift of the metal  
 from the  $\text{MB}^+$  plane, which is almost equivalent in the two  
 complexes (angles 11.8° and 9.5°). However,  $\tau$  rotation has  
 consequences on the extended heap, formed by the parallel  
 $\text{MB}^+$  units in an alternating head-to-tail arrangement (Figure

2).

First of all, the view down the N–M vector (Figure 2a)  
 highlights the “leaning tower” shape of the heap, consistent  
 with the continuous lateral sliding of each complex unit by a  
 half condensed hexagon. In this manner, the atoms of two  
 adjacent  $\text{MB}^+$  cations do not eclipse each other. Conversely,  
 Figure 2b clarifies the role of the  $\tau$  and  $\theta$  angles for the whole  
 system. The  $\text{Cl}_2\text{M}$  rotation shortens the contact between any  
 Cl atom and one H–C group of one adjacent  $\text{MB}^+$  cation. This  
 allows single but continuous hydrogen-bonding interactions  
 with an apparent stabilization of the heap. On the other hand,  
 the latter cannot be pairwise equivalent because of the  $\theta$   
 deviation, which favors one  $\text{Cl}\cdots\text{H}-\text{C}$  interaction with respect  
 to the subsequent one. In fact, the  $\text{Cl}\cdots\text{H}$  distances are as  
 different as 2.614 and 2.89 Å and 2.622 and 2.816 Å in MBCu  
 and MBAg, respectively. As a major consequence, the heap’s  
 building block cannot be a single complex but one of two  
 alternative dimers. The first one features two N–M vectors,  
 which simultaneously point toward the associated  $\text{MB}^+$  unit  
 (green box in Figure 2b), while the vectors diverge in the dimer  
 formed by the two central complexes (red box). The two  
 possible building blocks have different interplanar distances of  
 3.506 vs 3.403 Å and 3.485 vs 3.307 Å in MBCu and MBAg,  
 respectively. Remarkably, the dimer with the closer contacts  
 between the cations is the one with larger  $\text{Cl}\cdots\text{H}-\text{C}$  contacts.  
 This suggests an inverse correlation between the main  
 supramolecular forces ( $\pi$ – $\pi$  and hydrogen bonding), as further  
 discussed in the computational part.

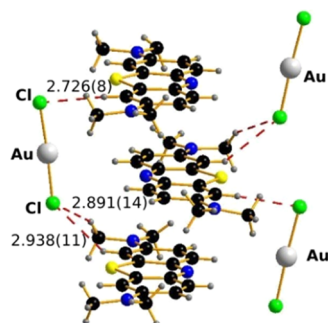
The compound MBAu, formed by the third coinage metal  
 (gold), was similarly synthesized by grinding together with the  
 solid reactants MB and Au(tht)Cl (tht = tetrahydrothiophene).  
 Five cycles of solvent-assisted grinding were necessary to 147



**Figure 2.** Layered MBCu and MAg complexes. (a) View down the N–M linkages, highlighting the “leaning tower” shape of the heap. (b) Orthogonal view down the largest MB<sup>+</sup> dimension, showing the two possible dimeric building blocks in the boxes.

148 completely convert the starting materials into a new crystalline  
 149 phase, as evidenced by the PXRD monitoring of the reaction  
 150 (see the [Supporting Information](#), PXRD3–5). In this manner,  
 151 the concomitant formation of a crystalline gold metal phase  
 152 also emerged, as shown by new signals in place of those  
 153 attributed to the target phase. Crystals suitable for X-ray  
 154 analysis were obtained upon the slow evaporation of a filtered  
 155 dichloromethane solution of the powder.

156 The X-ray structure of MBAu in [Figure 3](#) is indicative of a  
 157 salt of formula [(MB<sup>+</sup>)(AuCl<sub>2</sub><sup>−</sup>)]. In fact, the N–Au separation  
 158 of 4.201 Å excludes MB<sup>+</sup> coordination because it is  
 159 corroborated by the quasi-linearity of the AuCl<sub>2</sub><sup>−</sup> anion  
 160 (angle of 177.57°). Discrete ion pairs may also be excluded  
 161 because the metal fragments form hydrogen bonds with distinct



**Figure 3.** Packing features of the MBAu crystalline phase,<sup>23</sup> showing Cl⋯H contacts between linear AuCl<sub>2</sub><sup>−</sup> fragments and MB<sup>+</sup> molecules.

counterions and  $\pi$  stacking holds the parallel MB<sup>+</sup> units  
 162 together in a head-to-tail arrangement. 163

The now larger interplanar separation of 3.599 Å is indicative  
 164 of weaker interactions between the planes, also because the  
 165 adjacent cations are now shifted by one entire six-membered  
 166 ring. The reduced lattice energy is therefore likely insufficient to  
 167 compensate for the difficult formation of the N–Au<sup>I</sup>  
 168 coordination bond.<sup>29</sup> Examples of the latter are rare and not  
 169 well-defined, with one example being a T-shaped species with a  
 170 P–Au–S linear moiety associated with an orthogonal pyridine  
 171 ligand, which remains 2.56 Å far from the metal.<sup>30</sup> On the other  
 172 hand, the propensity of the d<sup>10</sup> coinage metals to add a third  
 173 coplanar ligand diminishes down the triad, as found for a  
 174 P<sub>2</sub>MCl series (P<sub>2</sub> = chelate diphosphine) with the progressively  
 175 larger Cl separation.<sup>31</sup> 176

Some authors attributed the different behavior of Au to  
 177 relativistic effects,<sup>32</sup> which remained elusive when included in  
 178 some of our DFT calculations. The latter were initially based on  
 179 the molecular modeling of the single MBMCl<sub>2</sub> complexes (M =  
 180 Cu, Ag) carried out with the B3LYP-DFT<sup>33</sup> method both in the  
 181 gas phase and in a CHCl<sub>3</sub> solvent.<sup>34,35</sup> Unfortunately, these  
 182 optimizations were not consistent with the X-ray structural  
 183 data. Only Cu apparently formed a three-coordinated complex  
 184 with the MB<sup>+</sup> ligand, but its geometry was clearly incorrect. Not  
 185 only were the N–Cu distance and the Cl–Cu–Cl angle  
 186 overestimated by about +0.15 Å and +32°, respectively, but the  
 187 metal was far from lying on the MB<sup>+</sup> plane ( $\theta = 49^\circ$ ). MAg  
 188 optimization was even less performing because a N⋯Ag  
 189 separation of >6.0 Å excluded the existence of the coordination  
 190 bond, consistently with a quasi-linear Cl<sub>2</sub>Ag<sup>−</sup> fragment (170°).  
 191 In this respect, MAg seemed more similar to MBAu than the  
 192 MBCu compound, which was similarly optimized. At this point,  
 193 it became evident that the neglected intermolecular forces  
 194 could have a role in the MB<sup>+</sup> coordination, as was already  
 195 implicit in the spectroscopic analysis of the MBCu and MAg  
 196 compounds in an acetonitrile solution. In fact, no NMR or MS  
 197 signal was attributable to the possible N–Cu or N–Ag  
 198 coordination. 199

Evidence for supramolecular forces at work in the extended  
 200 heap of MBCuCl<sub>2</sub> units emerged from the solid-state  
 201 calculations with the program CRYSTAL.<sup>24</sup> Before going into  
 202 the details, it is worth mentioning that the stacking had been  
 203 explored at the molecular level for the single dimeric building  
 204 block of [Figure 2b](#). Remarkably, the standard B3LYP<sup>33</sup>  
 205 calculations indicated immediate scission of the single  
 206 complexes, which attain the mentioned incorrect geometry.  
 207 Conversely, the dispersion correction in the B97D functional<sup>36</sup>  
 208 afforded an acceptable optimization of the (MBCuCl<sub>2</sub>)<sub>2</sub>  
 209 assembly with two in-pointing N–Cu vectors, although with  
 210 a  $\theta$  angle of 23.8° ([Figure S4](#)), which shortens the Cl⋯H–C  
 211 interactions (ave. 2.48 Å). Interestingly, a third stacked complex  
 212 ([Figure 4](#)) restores the experimental  $\theta$  value of 11° at the  
 213 central unit but not at the terminal ones (16° and 29°). This  
 214 problem of boundary conditions disappears in the solid-state  
 215 approach. 216

The CRYSTAL optimization with the B97D functional shows  
 217 an acceptable geometry of the stacked complex units (see the  
 218 values in brackets in [Table 1](#)). Also, the “leaning tower” aspect  
 219 of the heap is well reproduced ([Figure S5a](#)) with distinct and  
 220 alternating dimeric building blocks, although with smaller and  
 221 similar interplanar separations (3.10 and 3.06 Å). To evaluate  
 222 the effect, likely attributable to the dispersion forces, a new  
 223 CRYSTAL optimization was attempted at the BLYP level. At 224



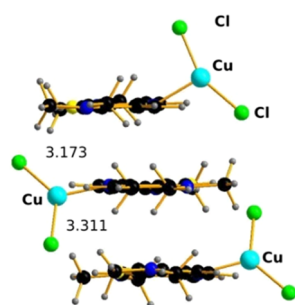


Figure 4. DFT-optimized assembly of three  $\text{MBCuCl}_2$  complexes.

225 first sight, the structure (Figure S5b) was better behaving,  
 226 especially for the larger and more asymmetric interplanar  
 227 separations of 3.41 and 3.30 Å. However, this result is  
 228 associated with a questionable stereochemistry, given that the  
 229 rotation of each  $\text{Cl}_2\text{Cu}$  fragment about the N–Cu linkage is in  
 230 the opposite sense (negative  $\tau$  angle). Consequently, the Cl–S  
 231 contacts are preferentially shortened with respect to the Cl···  
 232 H–C ones. This suggests a residually positive S atom in  $\text{MB}^+$ ,  
 233 possibly consistent with resonance structures rather than those  
 234 in Scheme 1. In any case, the stereochemical conflict between  
 235 the optimized and experimental structures kept us from further  
 236 exploring the problem.

237 The combined strategy of molecular and solid-state modeling  
 238 offered the possibility of deriving useful energy data on the  
 239 interaction between adjacent complexes.<sup>37</sup> On average, the  
 240 enthalpy of the attraction is about  $-40 \text{ kcal mol}^{-1}$ , mostly  
 241 because of the dispersion forces. In fact, the dismissal of the  
 242 latter in the BLYP approach transforms the interaction into a  
 243 minor repulsion ( $\sim +2 \text{ kcal mol}^{-1}$ ). Therefore,  $\pi$  stacking has a  
 244 governing role and overwhelms other contributions such as  
 245 hydrogen-bonding or residual electrostatic attraction between  
 246 differently charged counterions at different complexes.  
 247 Consider in this respect that the components of the latter  
 248 type are equally included in the two adopted functionals,  
 249 without any major evidence of their specific contribution in  
 250 favoring the assembly of the planar and formally closed-shell  
 251 complexes.

252 To support the still scarcely documented role of the packing  
 253 forces in assisting metal coordination, we mention a previous  
 254 study reported by some of us (C.M. and A.I.)<sup>38,39</sup> on  $\text{Mn}^{\text{II}}$   
 255 dimers with four carboxylate bridges and a terminal bipyridine  
 256 (bipy) chelate per metal. In these species, the local metal  
 257 coordination was trigonal-prismatic, while in the crystal, the  
 258 unsubstituted bipy ligands were involved in extended and  
 259 compact  $\pi$  stacking. However, the introduction of some methyl  
 260 substituent at bipy induces a major structural effect, such as the  
 261 forcing of an octahedral  $\text{Mn}^{\text{II}}$ , also because the  $\text{CH}_3$   
 262 substituents do not allow the same effective  $\pi$  stacking of the  
 263 planar chelates. In fact, the loss of lattice energy (although  
 264 unvalued by calculations) cannot support the trigonal-prismatic  
 265 coordination anymore. With respect to the latter case, the  
 266 present  $\text{MB}^+$  behavior seems even more remarkable because  $\pi$   
 267 stacking controls not only the coordination geometry but also  
 268 Werner's coordination number<sup>40</sup> (from 2 to 3).

## 269 CONCLUSION

270 In conclusion, the coordination capabilities of the aromatic  
 271  $\text{MB}^+$  cation have been assessed for the first time. Analyses have  
 272 shown that the supramolecular forces govern the overall  
 273 stereochemistry to the point of affording in some cases

otherwise disfavored and still unknown complexes of MB. 274  
 Because the latter compound already has a relevant number of 275  
 applications in different areas of chemistry, it cannot be 276  
 excluded that some new fruitful usage may be derived in 277  
 combination with a transition-metal center. For this reason, the 278  
 synthetic procedures, stereochemistry, and electronic under- 279  
 pinnings of our solid-state systems may represent a useful 280  
 guideline in the search for new supramolecular systems, where 281  
 MB (or a similar synthon) is associated with various metal 282  
 centers. In this respect, an added value of this paper is the 283  
 mechanochemical technique used, in place of other standard 284  
 synthetic strategies in solution. Hopefully, the formation of 285  
 other highly energetic supramolecular patterns may offer 286  
 important perspectives for chemistry. 287

## 288 ASSOCIATED CONTENT

### 289 Supporting Information

The Supporting Information is available free of charge on the 290  
 ACS Publications website at DOI: 10.1021/acs.inorg- 291  
 chem.6b02980. 292

X-ray crystallographic data in CIF format (CIF) 293

Experimental details on the syntheses of the compounds 294  
 at issue, IR and PXRD characterization data, the CCDC 295  
 search outcome on the N–Cu and N–Ag coordination 296  
 length bonds, and drawings and coordinates of computed 297  
 structures (PDF) 298

## 299 AUTHOR INFORMATION

### 300 Corresponding Authors

\*E-mail: giovanni.predieri@unipr.it. 301

\*E-mail: carlo.mealli@iccom.cnr.it. 302

### 303 ORCID

Alessia Bacchi: 0000-0001-5675-9372 304

Paolo Pelagatti: 0000-0002-6926-2928 305

Giovanni Predieri: 0000-0002-7912-641X 306

### 307 Author Contributions

This paper received contributions of all the authors. These 308  
 authors contributed with the DFT calculations and the further 309  
 interpretation of the computed results. 310

### 311 Notes

The authors declare no competing financial interest. 312

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## 319 DEDICATION

Dedicated to Antonio Tiripicchio on the occasion of his 80th 320  
 birthday. 321

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