

1 **Copper(II) cubanes with a {Cu₄O} core and well defined S = 1 ground state†**
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18 **ABSTRACT:**
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20 The reaction of 2-pyridinemethanol with copper 4-fluorobenzoate has yielded a family of type II
21 cubanes with formula $[\text{Cu}_4(\text{pymO})_4(4\text{-F-PhCOO})_3(\text{NO}_3)]$ (1), $[\text{Cu}_4(\text{pymO})_4(4\text{-F-PhCOO})_4]$ (2) and
22 $[\text{Cu}_4(\text{pymO})_4(4\text{-F-PhCOO})_4(\text{H}_2\text{O})]$ (3). These systems exhibit an unexpected $S = 1$ ground state and
23 their magnetic properties have been unambiguously characterized and rationalized as a function of the
24 asymmetry of the $\{\text{Cu}_4\text{O}_4\}$ cage and Cu–O–Cu bond angles. Analysis of the coupling constants was
25 performed applying new interaction schemes. Magneto-structural correlations have been performed
26 from the analysis of previously reported type II copper cubanes..

27 **Introduciton**

28

29 Copper cubanes have been classified in structural types as a function of their topology and degree of
30 distortion. Mergehenn and Haase¹ proposed a classification based on the relative distribution of the
31 elongated Cu–O distances in the cube: if the four elongated distances are roughly parallel, the cube can
32 be envisaged as two weakly interacting dimeric subunits (named type I) whereas if the elongated
33 distances are distributed perpendicularly on two opposite faces of the cube (named type II), it can be
34 envisaged to be a folded Cu₄O₄ ring with four weak additional interactions (Scheme 1). More recently,
35 Alvarez et al.² made an alternative proposal based on the distribution of the six Cu···Cu distances, with
36 the (2 + 4) and (4 + 2) classes being equivalent to types I and II, respectively, and by adding a new class
37 in which the six Cu···Cu distances are similar, named the (6 + 0) class. The (2 + 4) and (4 + 2) cubes
38 usually corresponds to systems in which the Cu^{II} cations have a square pyramidal or elongated
39 octahedral environment whereas the (6 + 0) cubes should be assigned to the scarce cores with six
40 equivalent faces, in which the coordination polyhedron around the Cu^{II} cations is usually a trigonal
41 bipyramid. Obviously, the magnetic properties of the cubes are strongly dependent on the structure, the
42 (6 + 0) class being closer to a true cube while the type I or (2 + 4) class is closer to two more or less
43 weakly interacting dimers and the type II or (4 + 2) class is more related to a distorted Cu₄ ring.

44 In all reported cases, dominant anti- or ferromagnetic interactions mediated by the short Cu–O
45 superexchange pathways lead to the $S = 0$ or $S = 2$ typical ground states.

46 2-Pyridinemethanol (pymOH) and the closely related (*R/S*)- α -methyl-2-pyridinemethanol (MpymOH)
47 ligands are able to generate polynuclear systems linking up to three cations (Scheme 2). Their copper
48 chemistry has been poorly explored and only some dimers,³ isolated⁴ or linked cubanes,^{4c,5} single
49 chains⁶ and some heterometallic Cu^{II}–Gd^{III} clusters⁷ have been reported for pymOH and only one pair
50 of enantiomers⁸ have been described for (*R/S*)-MpymOH. Our initial target was to explore the reactivity
51 of these ligands in carboxylate-copper chemistry but unfortunately unambiguous characterization was
52 only possible for pymOH derivatives.

53 In this work we report the syntheses and characterization of three new cubanes with a Cu₄O₄ core
54 belonging to the (4 + 2) class, obtained from the reaction of copper(II) 4-fluorobenzoate and 2-
55 pyridylmethanol (pymOH) with formulas [Cu₄(pymO)₄(4-F-PhCOO)₃(NO₃)] (1), [Cu₄(pymO)₄(4-F-
56 PhCOO)₄] (2) and [Cu₄(pymO)₄(4-F-PhCOO)₄(H₂O)]·0.5MeOH·0.25H₂O (3·0.5MeOH·0.25H₂O).
57 Magnetic susceptibility and magnetization measurements prove that these systems possess a well
58 isolated $S = 1$ ground state. This unique property has been rationalized as a function of the cage bond
59 parameters. We also report a general study of the magnetic response of the previously reported (4 + 2)
60 class cubes and a critical analysis of the models usually applied to fit the magnetic data.

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62 Experimental

63

64 Materials and methods

65 The Cu(4-F-PhCOO)₂ starting reagent was synthesized in typical yields >70% mixing equimolecular
66 amounts of aqueous solutions of Cu(NO₃)₂·6H₂O and Na(4-F-PhCOO) salts. The copper carboxylate
67 was collected *via* filtration and washed with cold water. Samples for analysis were gently dried to
68 remove volatile solvents. The yield for 1–3 was around 25% of well formed crystals which were
69 employed for instrumental measurements.

70 IR spectra (4000–400 cm⁻¹) were recorded using a Bruker IFS-125 FT-IR spectrometer with samples
71 prepared as KBr pellets. Variable-temperature magnetic studies were performed using a MPMS-5
72 Quantum Design magnetometer operating at 0.03 T in the 300–2.0 K range. Diamagnetic corrections
73 were applied to the observed paramagnetic susceptibility using Pascal's constants.

74 Energy levels plotted in Fig. 4b–d and those in Fig. 6 have been calculated for an arbitrary J_2 value of
75 -50 cm^{-1} .

76

77 Single-crystal X-ray crystallography

78 Blue prism-like specimens of approximate dimensions 0.196 mm × 0.336 mm × 0.522 mm (1), 0.082
79 mm × 0.168 mm × 0.227 mm (2) and 0.390 mm × 0.397 mm × 0.508 mm (3) were used for X-ray
80 crystallographic analysis. The X-ray intensity data were measured on a D8 Venture system equipped
81 with a multilayer monochromator and a Mo microfocus ($\lambda = 0.71073 \text{ \AA}$). The frames were integrated
82 with the Bruker SAINT software package using a narrow-frame algorithm. The final cell constants were
83 based upon the refinement of the XYZ-centroids of reflections above 20 $\sigma(I)$. Data were corrected for
84 absorption effects using the multi-scan method (SADABS). The structures were solved using the Bruker
85 SHELXTL Software Package, and refined using SHELXL.9 Details of crystal data, collection and
86 refinement for 1–3 are summarized in Table 1. Analyses of the structures and plots for publication were
87 performed with the Ortep3¹⁰ and POVRAY programs.

88

89 Synthetic procedure

90 [Cu₄(pymO)₄(4-F-PhCOO)₃(NO₃)] (1). A few crystals of complex 1 were initially obtained from a
91 Cu(4-F-PhCOO)₂ starting reagent contaminated with nitrates. In light of the structural results, the
92 synthesis was repeated by dissolving Cu(4-F-PhCOO)₂ and Cu(NO₃)₂·6H₂O in a 3 : 1 ratio (0.375
93 mmol, 0.128 g : 0.125 mmol, 0.037 g) in methanol (5 mL) and the ligand pymOH in 5 mL of
94 acetonitrile. Both solutions were mixed and stirred for three hours.

95 Complex 1 crystallizes as blue crystals *via* vapour diffusion with diethyl ether. Anal. Calcd for
96 C₄₅H₃₆Cu₄F₃N₅O₁₃ (1): C, 46.35; H, 3.11; N, 6.01%. Found: C, 46.92; H, 3.4; N, 5.88%. Relevant IR
97 bands: 3440 (s, broad), 3077(w), 2835 (w), 1620(s), 1580(s), 1506(s), 1440(s), 1360(s), 1310 (s),
98 1210(w), 1150(w), 1050(s), 860(w), 785(w), 760(w), 630(w) cm⁻¹.

99 [Cu₄(pymO)₄(4-F-PhCOO)₄] (2). Cu(4-F-PhCOO)₂ (0.5 mmol, 0.170 g) was dissolved in methanol (5
100 mL) and the ligand pymOH was dissolved in 5 mL of acetonitrile. Both solutions were mixed and stirred
101 for three hours. Slow evaporation of the resulting solution yields complex 2 as blue crystals. Anal. Calcd

102 for $C_{52}H_{40}Cu_4F_4N_4O_{12}$ (2): C, 50.24; H, 3.24; N, 4.51%. Found: C, 49.32; H, 3.10; N, 4.31%. Relevant
103 IR bands: 3440 (s, broad), 3077(w), 2835 (w), 1620(s), 1440 (w), 1400(s), 1360(w), 1250(w), 1210(w),
104 1150(w), 985(w), 630(w), 480(w), 411(w) cm^{-1} .

105 $[Cu_4(pymO)_4(4-F-PhCOO)_4] \cdot 0.5MeOH \cdot 0.25H_2O$ ($3 \cdot 0.5MeOH \cdot 0.25-H_2O$). $Cu(4-F-PhCOO)_2$ (0.5
106 mmol, 0.170 g) was dissolved in methanol (5 mL). The ligands pymOH (0.75 mmol, 0.081 g) and (S-
107 pyeOH) (0.25 mmol, 0.035 g) were dissolved in acetonitrile (5 mL). The mixture of both solutions was
108 stirred for three hours, filtered and layered with diethyl ether. Well formed blue crystals suitable for X-
109 Ray analysis grew after two weeks. Anal. Calcd for $C_{52.5}H_{44.5}Cu_4F_4N_4O_{13.75}$
110 ($3 \cdot 3 \cdot 0.5MeOH \cdot 0.25H_2O$): C, 49.20; H, 3.50; N, 4.37%. Found: C, 49.73; H, 3.41; N, 4.16%. Relevant
111 IR bands: $\nu = 3440$ (s, broad), 3160(w) 2835 (w), 1610(s), 1550(s), 1400(s), 1210(s), 1080 (s), 860(s),
112 780(s), 618(s), 530(w) cm^{-1} .

113

114 Results and discussion

115

116 Structural description

117

118 [Cu₄(pymO)₄(4-F-PhCOO)₃(NO₃)] (1). The molecular structure consists of isolated cubanes with a
119 {Cu₄O₄} core. A view of the structure is shown in Fig. 1 and the main bond parameters are summarized
120 in Table 2. One pymO⁻ ligand is coordinated to each copper cation, providing the four μ₃-alcoxo
121 corners of the cube. The pymO⁻ ligands are placed roughly perpendicular to two opposite faces of the
122 cube whereas three of the remainder four faces are occupied by the three bidentate carboxylates. The
123 nitrate anion acts as monodentate ligand, coordinated to Cu₄. Cu₂ shows a square pyramidal CuNO₄
124 environment whereas Cu_(1,3,4) exhibit an axially elongated octahedral CuNO₅ coordination
125 polyhedron. The elongated axial bond distances involve one Cu–O cage bond for each copper cation and
126 one Cu–O bond with one O-carboxylate or O-nitrate for Cu_(1,2,4). The equatorial bond distances are in
127 the short 2.000–1.905 Å range whereas the axial Cu–O bond distances are relatively large, ranging
128 between 2.344(3)–2.697(3) Å.

129 [Cu₄(pymO)₄(4-F-PhCOO)₄] (2). A view of the structure is shown in Fig. 2 and the main bond
130 parameters are summarized in Table 2. The structure is very similar to 1 but in this case the nitrate
131 ligand has been substituted by a fourth carboxylate. In this case two carboxylates act as bidentate ligands
132 coordinated to the neighbor {Cu₁/O₄/Cu₄/O₁₀} and {Cu₁/O₁₀/Cu₃/O₁} faces whereas the other two
133 carboxylates act as monodentate ligands coordinating Cu₂ and Cu₃. Cu₁ and Cu₄ show an elongated
134 octahedron coordination polyhedron whereas Cu₂ and Cu₃ exhibit a square pyramidal environment. The
135 core of the cube is more distorted than complex 1 as is reflected in the large Cu₄–O₇ distance of
136 2.962(7) Å or the Cu₄–O₄–Cu₂ bond angle of 120.1(3)°.

137 [Cu₄(pymO)₄(4-F-PhCOO)₄(H₂O)]·3·0.5MeOH·0.25H₂O (3·3·0.5MeOH·0.25H₂O). A view of the
138 structure is shown in Fig. 3 and the main bond parameters are summarized in Table 2. The structure of 3
139 is closely related to compound 2 but now there is an additional water molecule coordinated to Cu₂,
140 which turns to be hexacoordinated. The coordinated water molecule establishes two strong H-bonds
141 with the noncoordinated O₆ and O₁₂ atoms belonging to the monodentate carboxylates and also
142 interacts with the crystallization water molecule. O₆···O_{3w} and O₁₂···O_{3w} distances are 2.685(3) and
143 2.780(2) Å respectively. The presence of this new ligand on Cu₂ increase the distance between the
144 monodentate carboxylates and as a consequence, displaces Cu₄ with the concomitant increase of the
145 Cu₄–O₇ distance (up to 3.253 Å) and the Cu₂–O₄–Cu₄ bond angle, which reaches 124.32(6)°. As can
146 be seen in Table 2, the three cubes are quite similar in their general trends, increasing the distortion of
147 the cage from the least (1) to most distorted (3) cube.

148 Spin levels and ground state for the Cu₄ (4 + 2) cubane topology

149 The magnetic properties for the (4 + 2) copper cubane topology have been widely studied via DFT
150 calculations^{2,11} and all studied cases lead to the S = 0 or S = 2 ground state. As can be expected, it was
151 also stated that the axial-equatorial interactions involving often very large Cu–O distances (on two
152 elongated opposite faces, Scheme 1) must always be weak.² Surprisingly, the susceptibility
153 measurements performed for complexes 1–3 clearly suggest an unprecedented “anomalous”
154 intermediate spin ground state S = 1 (see further magnetic properties discussion), apparently
155 incompatible with a Cu^{II} cubane topology.

156 To have a clear picture of the magnetic properties of all previously reported cubanes with a (4 + 2)
 157 shape, a search in the CCDC database was performed and 119 entries were obtained for Cu^{II} cubes with
 158 four elongated Cu–O bonds larger than 2.100 Å as the only restraint. Cubes for which the complete
 159 magnetic analysis was not reported or the coordination polyhedron around the Cu^{II} cations was a
 160 trigonal bipyramid were discarded from this study. The magnetic data for the 43 (4 + 2) Cu^{II} cubes with
 161 reported magnetic data and a square pyramidal or elongated octahedral environment around the Cu^{II}
 162 cations are summarized in Table 3.

163 The next step was to check which models were applied to fit the experimental data and up to five models
 164 were found to describe the magnetic response of these systems (Scheme 3).

165 Despite the evidence that the superexchange interaction mediated by the opposite faces with exclusively
 166 axial–equatorial (Jahn–Teller) pathways is usually poorly effective in comparison with the four faces
 167 with equatorial–equatorial pathways,^{2,11} some coupling constants analysis were performed assuming a
 168 regular model (Scheme 3, model (6) and Table 3), for which the corresponding Hamiltonian is:

$$169 \quad H = -J_1(S_1 \cdot S_2 + S_1 \cdot S_3 + S_1 \cdot S_4 + S_2 \cdot S_3 + S_2 \cdot S_4 + S_3 \cdot S_4) \quad (1)$$

170
 171 On the other hand, the magnetic properties for most of the
 172 reported systems were calculated with the (2:4) or (0:4) models
 173 (Scheme 3 and Table 3) for which the Hamiltonians are:

$$174 \quad H = -J_1(S_1 \cdot S_2 + S_3 \cdot S_4) - J_2(S_1 \cdot S_3 + S_1 \cdot S_4 + S_2 \cdot S_3 + S_2 \cdot S_4) \quad (2)$$

$$175 \quad H = -J_2(S_1 \cdot S_3 + S_1 \cdot S_4 + S_2 \cdot S_3 + S_2 \cdot S_4) \quad (3)$$

177 For a reduced number of asymmetric cubes, models taking into account different interactions for each
 178 pair of opposite

179
 180
 181
 182 faces of the cube were applied (models (0:2:2) and (2:2:2), Scheme 3 and Table 3), applying the
 183 Hamiltonians:

$$184 \quad H = -J_2(S_1 \cdot S_3 + S_2 \cdot S_4) - J_3(S_2 \cdot S_3 + S_1 \cdot S_4) \quad (4)$$

$$185 \quad H = -J_1(S_1 \cdot S_2 + S_3 \cdot S_4) - J_2(S_1 \cdot S_3 + S_2 \cdot S_4) \quad (5)$$

$$186 \quad -J_3(S_2 \cdot S_3 + S_1 \cdot S_4)$$

187 Hamiltonians (3) and (4) are the limit of (2) and (5) when J_1 was neglected assuming $J_2, J_3 \gg J_1$. The
 188 reported ground state for all cubes applying Hamiltonians (1)–(5) is systematically $S = 0$ for negative
 189 $J_{2,3}$ values or $S = 2$ for positive ones. This experimental feature can be easily rationalized plotting the
 190 energy of the six spin levels of the cube (one $S = 2$, three $S = 1$ and two $S = 0$), as a function of the
 191 coupling constant.

192 If we assume that the interaction between the copper centers through the elongated (Jahn–Teller) faces
 193 are negligible and the other four interactions are identical, model (0:4) and Hamiltonian (3), we obtain
 194 the spin level distribution shown in Fig. 4a, which evidences that $S = 0$ and $S = 2$ are the only possible

195 ground states as a function of the sign of J_2 . If we take into account the elongated faces, model (2:4) and
 196 Hamiltonian (2), we realize that for a dominant antiferromagnetic interaction J_2 the ground state is
 197 always $S = 0$ (the first $S = 1$ excited state has the same slope) and for a positive sign of J_2 , the ground
 198 state can switch from $S = 2$ to $S = 0$ for J_2/J_1 ratios lower than -0.5 , Fig. 4b and c respectively).

199 In a few cases, the fit of the experimental data was performed assuming a set of two or three J values for
 200 opposite faces of the cubes. Neglecting the interaction mediated by the opposite elongated faces,
 201 Scheme (0:2:2) and Hamiltonian (4), we realize that if one of the interactions is antiferromagnetic then S
 202 $= 0$ is the ground state for any positive or negative J_2/J_3 ratio (Fig. 4d). As in the (0:4) case, the addition
 203 of the weak interactions mediated by the elongated faces, model (2:2:2), only produces very small
 204 changes in the energy of the spin levels.

205 The above calculations exclude these models to analyse compounds 1–3 and then, the origin of their
 206 intermediate ground state must be found in other structural facts, neglected until now. The dependence
 207 of the magnitude and the sign of the coupling constants as a function of the Cu–O–Cu bond angle has
 208 been demonstrated via theoretical calculations and has been the preferred parameter to correlate the
 209 magnetic properties.^{2,11} The Cu–O–Cu bond angles involved in equatorial–equatorial bridges for the (4
 210 + 2) class of cubes (Fig. 5) take values comprised between 100° – 115° , being exceptional to find Cu–O–
 211 Cu bond angles below or above these limits. The border between the ferromagnetic–antiferromagnetic
 212 response is unclear because it can depend on the characteristics of the bridging ligand that provides the
 213 μ^3 -O linkage among other factors² but, always assuming that there are compounds out of the rule,
 214 around 108° is a roughly reasonable limit. As general rule, copper cubanes with these four Cu–O–Cu
 215 bond angles clearly lower than 108° tend to be ferromagnetic with a $S = 2$ ground state and those with
 216 these bond angles clearly larger than 108° tend to be antiferromagnetic with a $S = 0$ ground state (ESI,
 217 Table S1†). In light of these previous data, the detailed analysis of the structures of complexes 1–3
 218 unveils an uncommon feature: the three cages are very asymmetric as a consequence of the coordination
 219 of the bidentate carboxylates on the *contiguous* faces. As a consequence, the Cu–O–Cu bond angles
 220 involving short Cu–O distances are also more similar on the *contiguous* faces instead of the *opposite*
 221 faces as it is common (Table 2). Taking the parameters defined in Fig. 5 as a reference, complex 1 has
 222 one large δ bond angle of 112.2° , two short α and β of 100.3° and 102.6° and one intermediate γ of
 223 108.7° . The Cu–O–Cu bond angles for complexes 2 and 3 exhibit two large *contiguous* α and δ angles
 224 ($110.9^\circ/120.1^\circ$ and $110.8^\circ/124.4^\circ$) and two short β and γ Cu–O–Cu bond angles ($101.5^\circ/104.3^\circ$ and
 225 $100.8^\circ/104.2^\circ$).

226 In basis to these structural parameters we attempted the analysis of the energy of the spin levels for the
 227 new models plotted in Scheme 4, which describe the interactions as three similar and one different
 228 interaction, (0:1:3) model, and two similar interactions on *contiguous* faces, (0:2:2c) model. The
 229 corresponding Hamiltonians are:

$$230 \quad H = -J_2(\mathbf{S}_1 \cdot \mathbf{S}_4) - J_3(\mathbf{S}_1 \cdot \mathbf{S}_3 + \mathbf{S}_2 \cdot \mathbf{S}_3 + \mathbf{S}_2 \cdot \mathbf{S}_4) \quad (6)$$

$$231 \quad H = -J_2(\mathbf{S}_1 \cdot \mathbf{S}_3 + \mathbf{S}_1 \cdot \mathbf{S}_4) - J_3(\mathbf{S}_2 \cdot \mathbf{S}_3 + \mathbf{S}_2 \cdot \mathbf{S}_4) \quad (7)$$

232 The plots of the energies of the six spin levels of the cubane topology for these models are shown in Fig.
 233 6. Obviously, if the sign of both the J_2 and J_3 constants is the same, the ground states will be newly $S =$
 234 0 or 2 . However, by forming these plots we realize that for the (0:1:3) model $S = 2$ is the ground state if
 235 the $-J_3/J_2$ ratio is lower than $1/3$ but one of the $S = 1$ spin levels becomes clearly the ground state for
 236 larger ratios. Equally for the (0:2:2c) model, for negative J_3/J_2 ratios (i.e. different sign for the two
 237 coupling constants) one well isolated $S = 1$ spin level becomes the ground state.

238

239 Magnetic properties

240 χ_{MT} vs. T plots for 1–3 are shown in Fig. 7. χ_{MT} at room temperature for 1 is $1.86 \text{ cm}^3 \text{ Kmol}^{-1}$. Upon
241 cooling, the χ_{MT} value decreases continuously down to a plateau value of $1.18 \text{ cm}^3 \text{ Kmol}^{-1}$ around 12
242 K. Below this temperature χ_{MT} raises slightly to decrease finally to $1.12 \text{ cm}^3 \text{ Kmol}^{-1}$ at 2 K. Complexes
243 2 and 3 exhibit χ_{MT} values of 1.55 and $1.51 \text{ cm}^3 \text{ Kmol}^{-1}$ at room temperature. For decreasing
244 temperatures, the χ_{MT} value decreases continuously down to a well defined minimum of 1.08 cm^3
245 Kmol^{-1} at 60 K for 2 and 70 K for 3. At low temperature, the χ_{MT} value slightly increases prior to the
246 final decrease to and 1.07 and $1.16 \text{ cm}^3 \text{ Kmol}^{-1}$ at 2 K.

247 According the crystallographic data and the above proposed models, the susceptibility data was fitted
248 with the PHI program¹² applying the (0:1:3) model for 1 (Scheme 4, Hamiltonian (6)) and (0:2:2c)
249 model (Scheme 4, Hamiltonian (7)) for 2 and 3. R quality factors were calculated as $R = (\chi_{MT_{\text{exp}}} -$
250 $\chi_{MT_{\text{calc}}})^2 / (\chi_{MT_{\text{exp}}})^2$. Excellent fits nicely reproducing the experimental data, including the χ_{MT}
251 minima, were obtained for the parameters $J_2 = -71.4 \text{ cm}^{-1}$, $J_3 = 17.2 \text{ cm}^{-1}$, $g = 2.24$ ($R = 3.44 \times 10^{-5}$)
252 for 1, $J_2 = -153 \text{ cm}^{-1}$, $J_3 = +22 \text{ cm}^{-1}$, and $g = 2.22$ ($R = 1.17 \times 10^{-5}$) for 2 and $J_2 = -164 \text{ cm}^{-1}$, $J_3 =$
253 $+30 \text{ cm}^{-1}$, and $g = 2.21$ ($R = 9.80 \times 10^{-6}$) for 3, Fig. 7.

254 As was indicated above, one of the Cu–O–Cu bond angles of compound 1 is $\gamma = 108.7^\circ$ and it should be
255 close to the FM-AF limit and consequently with a low absolute value. A second simulation discarding
256 this interaction was performed to prove this assumption applying the simplified Hamiltonian:

$$257 \quad H = -J_2(S_1 \cdot S_3) - J_3(S_2 \cdot S_3 + S_2 \cdot S_4) \quad (8)$$

258 obtaining an equally good fit for the parameters $J_2 = -64.8 \text{ cm}^{-1}$, $J_3 = 12.8 \text{ cm}^{-1}$, $g = 2.26$, which
259 probably are more reliable.

260 The values for the antiferromagnetic interactions are in good agreement with the increase in the largest
261 Cu–O–Cu bond angle of 112.2° for 1, 120.1° for 2 and 124.4° for 3.

262 Ground state for the three complexes is then $S = 1$, with a 7.5 , 29.8 and 30.7 cm^{-1} gap to the first $S = 0$
263 excited spin level for 1–3 respectively (Fig. 8). As a consequence of this spin level distribution, the
264 magnetization of these complexes must be similar, following an $S = 1$ Brillouin shape. Magnetization
265 experiments performed in the 0 – 5 T range of an applied external field nicely confirm this assumption,
266 tending in all cases to a quasi saturated magnetization value equivalent to two electrons (Fig. 8).

267

268 Comments to the bibliographic data. Overlooked $S = 1$ cubes

269 The analysis of the bibliographic data of the magnetic properties of Cu^{II} cubes belonging to the $2 + 4$
270 class, published along more than twenty years, reveals to be extremely confused. Some relevant
271 magnetic features for 43 of those cubanes are summarized in Table 3. In this table the cubes with all of
272 the elongated Cu–O distances larger than an arbitrary value of 2.60 \AA (for which negligible magnetic
273 interactions through the elongated opposite faces must be assumed) are tabulated separately. In addition
274 to these 37 complexes, there are six other cubes for which disputable (but relevant) magnetic data were
275 reported, that will be discussed specifically. An overview of the data collected in Table 3 evidences that
276 the models applied in the magnetic analysis are not always justified. For cubes with large elongated Cu–
277 O distances, the most reasonable approach seems to be the one J model (0:4), and effectively, most of
278 these cubes were fitted according this model. However, in spite of the structural evidence, in some few
279 cases the authors assumed the regular cube model (6) (ELEYIE, NILDAP or WEMSUE).

280 As can be expected, J_1 usually shows low values for all cubes fitted with the (2:4) model and $S = 0$
281 ground state but in contrast, the large values of J_1 reported for QOMRAL or POLKEH seem to be
282 clearly overestimated. In this sense, the fits performed for LOCPIE and NODHEV become interesting,
283 for which the authors compared the fits with the (2:4) and (0:4) models obtaining minimal deviation in
284 J_2 , evidencing that for strongly AF coupled cubes the calculated value for J_1 is poorly reliable.

285 Much more interesting is the analysis of the models applied for ferromagnetic cubes. These kind of
286 systems can give a χ_{MT} plot that suggests the expected value for a $S = 2$ total spin but often, a decay at
287 low temperature or a continuous increase in χ_{MT} up to a value slightly lower than the expected for $S = 2$
288 has been reported. These plot shapes can be due to intercluster interactions or weak anisotropy in the
289 ground state, as has been demonstrated for FEVYAH by Ozarowski et al.²¹ When the isotropic (2:4)
290 model was applied to fit cubes with a ferromagnetic response, a systematic error is often produced, that
291 consists of the obtention of a pair of coupling constants with characteristic values very close to $-2J_1 =$
292 J_2 as occurs for ASUPEJ01, BUFTUR, CAQDAZ, IHELOX, NAXBET, SAPYUE, XOVVUA or
293 XOXGEY. The reason for this can be found in the plot of the spin levels for a ferromagnetic cube in
294 Fig. 4c: for a $-J_1/J_2 = 0.5$ ratio there is a crossing between the $S = 0$ and the $S = 2$ spin levels and the
295 population of both levels produces a decay of the χ_{MT} plot at low temperature (Fig. S1†).

296 The $S = 1$ ground state was erroneously claimed for ASUPEJ01, SAPYOY and SAPYUE despite their
297 χ_{MT} plots showing a continuous increase for decreasing temperatures. Fits were performed with the
298 (2:4) model that never can lead to the $S = 1$ ground state and these three cubes are obviously
299 ferromagnetic, with a $S = 2$ ground state.

300 In contrast, reviewing the χ_{MT} plots reported for this kind of cubes, we realized that the $S = 1$ ground
301 state is unusual but not unprecedented. DIBTAL shows a χ_{MT} response very similar to that of complex
302 1 but the authors reported the magnetic properties of this cube as being unexplainable and no fit was
303 tried. From its χ_{MT} shape and the low temperature value (with a plateau at around $1.1 \text{ cm}^3 \text{ Kmol}^{-1}$), the
304 $S = 1$ ground state becomes evident. The reason for this magnetic response lies in their α - δ Cu-O-Cu
305 bond angles which follows the sequence 112.3° - 111.2° - 99.7° - 105.5° , corresponding to the (0:2:2c)
306 model with two ferromagnetic and two antiferromagnetic contiguous interactions.

307 Other cubes with a probable $S = 1$ ground state are the enantiomers MOYJUH (*R*) and MOYKAO (*S*)
308 recently reported by S. Gao et al.;⁴³ only the (*R*) isomer MOYJUH was measured. Its low temperature
309 χ_{MT} plot tends clearly to $1.1 \text{ cm}^3 \text{ Kmol}^{-1}$ and the fit of the experimental data was performed with the
310 (2:4) model discarding the low temperature data. The reported values of $J_1 = -11.2 \text{ cm}^{-1}$ and $J_2 = +7.6$
311 cm^{-1} lead to a well defined $S = 0$ ground state with a gap of 11 cm^{-1} to the first $S = 1$ excited level,
312 which is not compatible with the experimental plot. The clearest proof for the $S = 1$ ground state for this
313 compound was provided by its magnetization, which follows an apparent Brillouin shape, tending to the
314 equivalent to two electrons.⁴³ The α - δ sequence of Cu-O-Cu bond angles for MOYJUH are comprised
315 between 100.5° - 107.4° and then does not follow the (0:1:3) nor the (0:2:2c) scheme. However, this
316 compound is extremely unusual because three Cu^{II} cations show a square pyramidal environment
317 whereas the fourth Cu^{II} cation has a trigonal bipyramidal coordination and then a new model and
318 probably DFT calculations would be necessary to explain its unusual magnetic response.

319 **Magneto-structural correlations.** Finally, an undesirable consequence of the employment of unreliable
320 J values should be pointed out, which were obtained by applying inappropriate models to fit the
321 experimental data: several trials to correlate the J values with the experimental Cu-O-Cu bond
322 angles^{2,11b,c} or more recently, the proposal of Boča et al.^{11a} as a basis of a chemometric analysis of the

323 Cu^{II} chromophores are far from being a linear relationship and partially it is due (as several authors have
324 pointed out) to the employment of unreliable experimental J values.

325 Along the paper we have assumed that the main parameter that determines the magnetic response of the
326 $(4 + 2)$ class of Cu^{II} cubes is the set of four $\alpha\text{-}\delta$ Cu–O–Cu bond angles. To perform a final check of the
327 validity of this very simplified model, we have selected a coherent group of cubes on the basis of the
328 following four conditions: (i) *comparable $\mu_3\text{-OR}$ bridging ligands*. Practically all complexes are linked
329 by alcoxo or phenoxo bridges but complexes as GIBHAC have been excluded because the bridging
330 ligands are $\mu_3\text{-OH}$, which gives a completely different magnetic response. (ii) S4 or quasi S4 symmetry.
331 It means that the four $\alpha\text{-}\delta$ Cu–O–Cu bond angles are identical or with a maximum tolerance of $\pm 1^\circ$. (iii)
332 *Square pyramidal or elongated octahedron environment around the four Cu^{II} cations*. It means to
333 discard complexes in which one or more copper atoms have a BPT environment. (iv) *To discard any*
334 *questionable value* (mainly for the cubes with an $S = 2$ ground state), indicated in the previous section.

335 The plot of the unfiltered J_2 values vs. the mean Cu–O–Cu $\alpha\text{-}\delta$ bond angles for all compounds reported
336 in Table 3 is very disperse and no conclusion can be extracted (Fig. 9). However, plotting the 20
337 selected cubes with the above criteria provides a clear indication of the dependence of the sign of the
338 magnetic interaction with this parameter and corroborates the assumption of the FM/AFM limit around
339 $108^\circ\text{--}110^\circ$. There are only two cubes BOGCOP and DARKUC that are clearly out of this correlation
340 without any apparent reason.

341

342 **Conclusions**

343 Three new Cu^{II} cubane-like complexes belonging to the (4 + 2) class have been characterised. From the
344 analysis of the susceptibility and magnetization data, the $S = 1$ ground state has been unambiguously
345 assigned for all of them together with the new coupling schemes that justify this unprecedented
346 response. A detailed analysis of the bibliographic data reveals that, to avoid overparametrization, often
347 oversimplified or inappropriate coupling schemes have been applied leading to a confuse landscape. The
348 reported 1–3 compounds are the first characterized cubes with an $S = 1$ ground state but they are not the
349 first compounds exhibiting this property, because in the literature we have found three unexplained
350 systems that belong to this unusual family.

351

352 **Acknowledgements**

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354

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429

430 **Legends to figures**

431
432 **Scheme 1** Schematic drawing of the cubane CuII complexes according the relative position of the
433 elongated Cu–O distances (red dashed bonds).

434
435 **Scheme 2** Ligands employed in this work and their coordination mode in the {Cu4O4} core of
436 compounds 1–3.

437
438 **Fig. 1** Top, a view of the molecular structure of compound 1. Bottom, the labeled core of the cubane.
439 Bonds depicted in orange correspond to the short Cu–O distances inside the {Cu4O4} cage

440
441 **Fig. 2** Top, a view of the molecular structure of compound 2. Bottom, the labeled core of the cubane.
442 Bonds depicted in orange correspond to the short Cu–O distances inside the {Cu4O4} cage

443
444 **Fig. 3** Top, a view of the molecular structure of compound 3. Bottom, the labeled core of the cubane.
445 Bonds depicted in orange correspond to the short Cu–O distances inside the {Cu4O4} cage and the
446 dashed red bonds show the H-bonds involving the coordinated water molecule

447
448 **Scheme 3** Interaction schemes for the CuII cubane topology according the literature. The models have
449 been named according to the number of identical faces and coupling constants.

450
451 **Fig. 4** Plot of the six spin levels of a CuII cubane for: (a) model (0:4) and Hamiltonian (3) for a +25 to
452 –50 cm⁻¹ range of J values, (b) model (2:4) and Hamiltonian (2) for an AF J2, (c) model (2:4) and
453 Hamiltonian (2) for a FM J2 and (d) model (0:2:2) and Hamiltonian (4). Color key of the spin levels: S =
454 2, red; S = 1, black; S = 0, blue; and degenerate levels, green. Arbitrary value for J2 = –50 cm⁻¹ in (b–
455 d).

456 **Fig. 5** Set of consecutive Cu–O–Cu bond angles involving the four short Cu–O distances in the cubane
457 core.

458
459 **Scheme 4** Low symmetry interaction schemes for the CuII cubane topology proposed for compounds 1–
460 3. The models have been named according to the number of identical faces and coupling constants.

461
462 **Fig. 6** Plot of the six spin levels of a CuII cubane for the low symmetry models (0:1:3) and Hamiltonian
463 (6) (left) and (2:2:2c) and Hamiltonian (7) (right), showing the S = 1 ground state for J3/J2 ratios lower
464 than –1/3 and 0 respectively. Bottom, the spin arrangement that allows to the S = 1 ground state. Color
465 key of the spin levels: S = 2, red; S = 1, black; S = 0, blue; degenerate levels, green. Arbitrary value for
466 J2 = –50 cm⁻¹.

467
468 **Fig. 7** Temperature dependence of χ_{MT} for compound 1 (circles), 2 (triangles) and 3 (squares). Solid
469 lines show the best obtained fits.

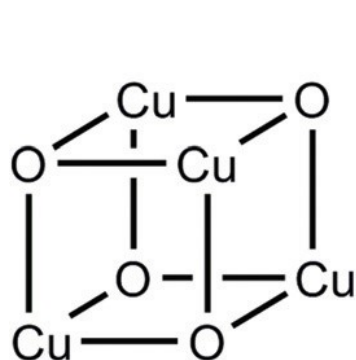
470
471 **Fig. 8** Left, energy levels calculated from the fit parameters for complexes 1–3. Right, magnetization
472 data for complexes 1 (squares), 2 (circles) and 3 (triangles) in agreement with the expected S = 1 ground
473 state.

474
475 **Fig. 9** Plot of the relationship between J and the mean $\alpha-\delta$ Cu–O–Cu bond angle for all samples
476 tabulated in Table 3 (left) or the 20 selected compounds according the criteria described in the text,
477 (right), (R factor = 0.76). The compounds represented as stars correspond to complexes BOGCOP and
478 DARKUC, which do not follow the correlation

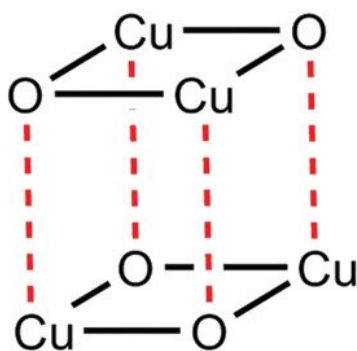
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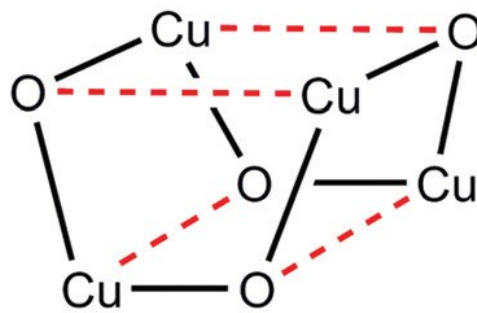
SCHEME 1



6+0 class



Type I
2+4 class

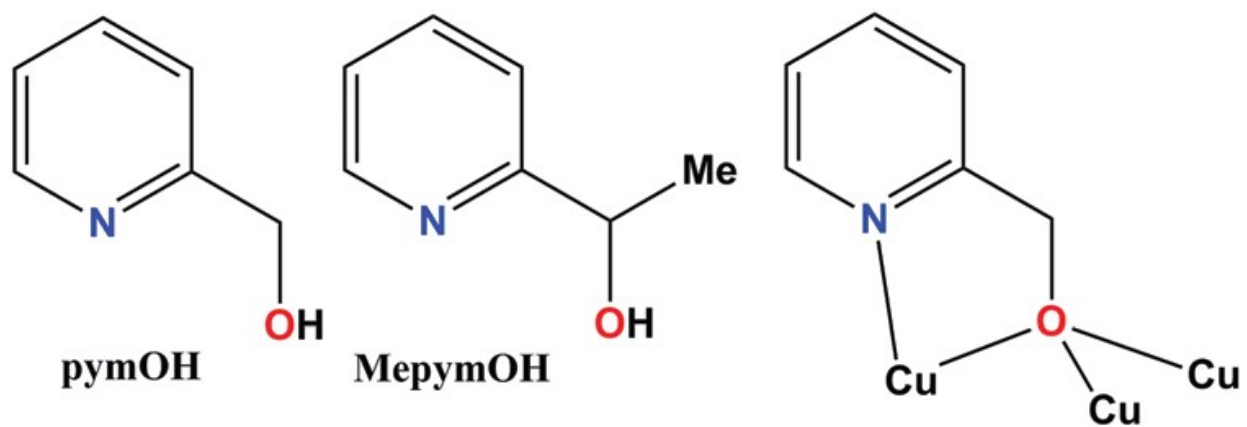


Type II
4+2 class

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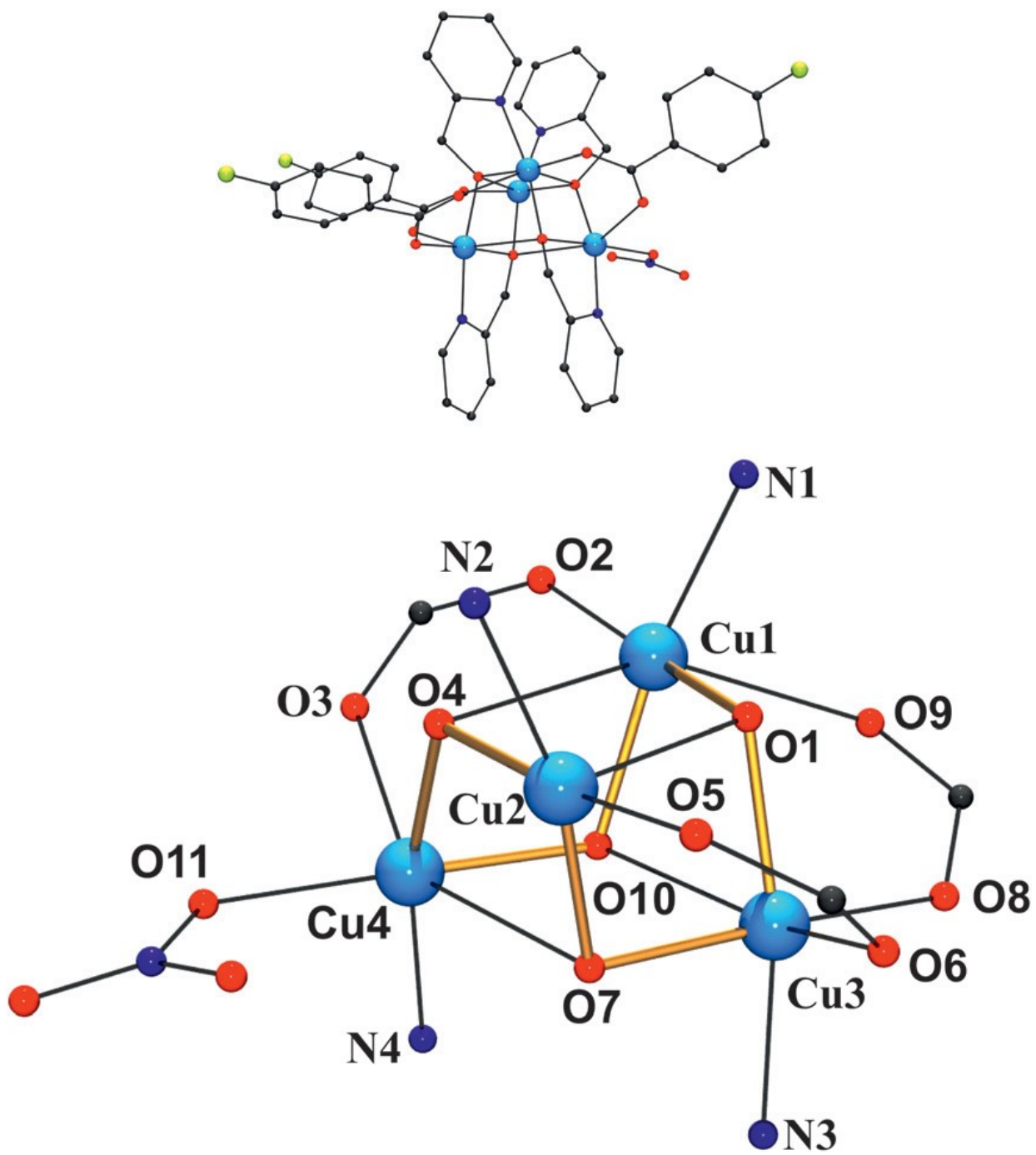
Scheme 2



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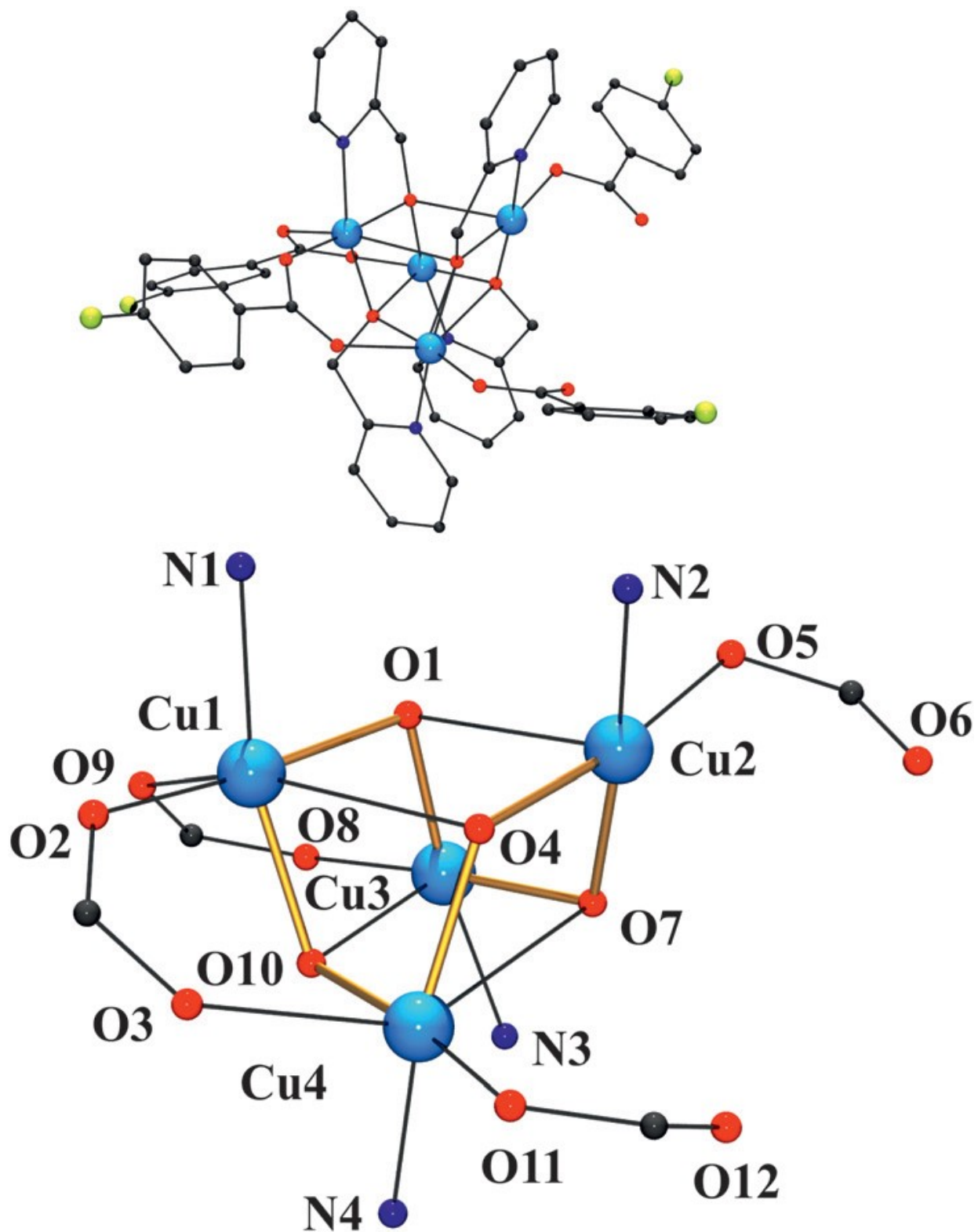
FIGURE 1



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FIGURE 2

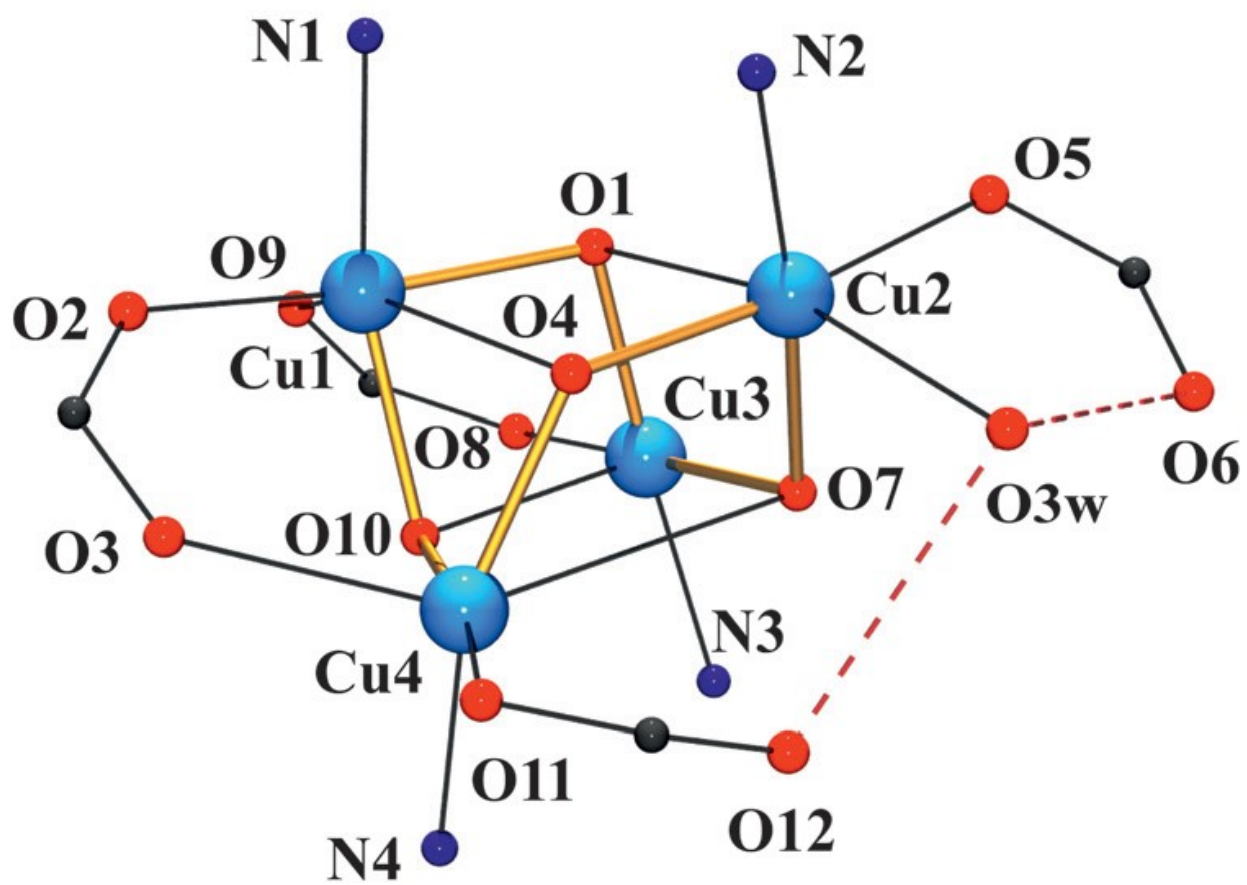
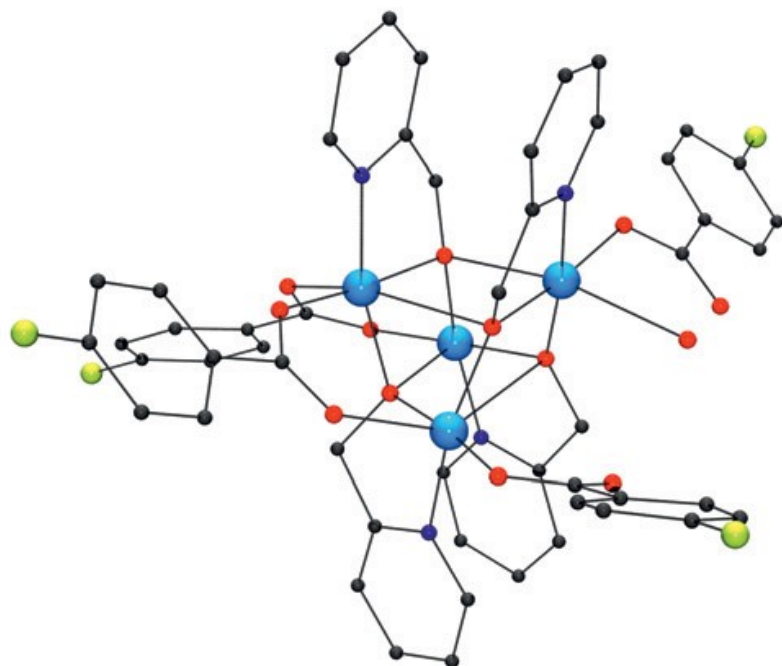


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FIGURE 3

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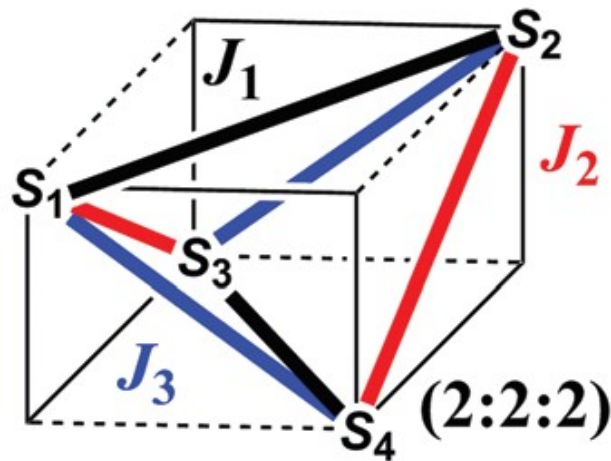
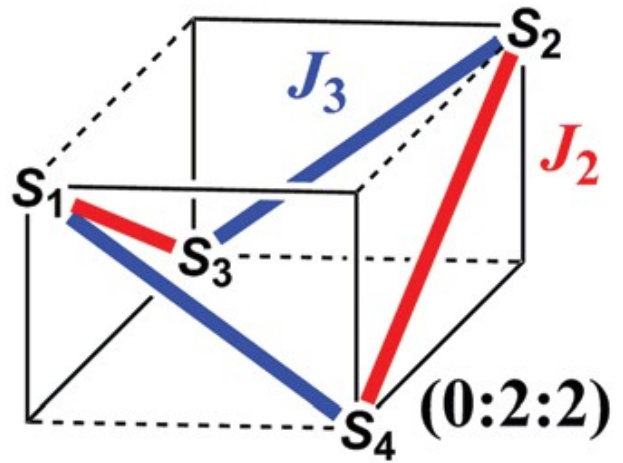
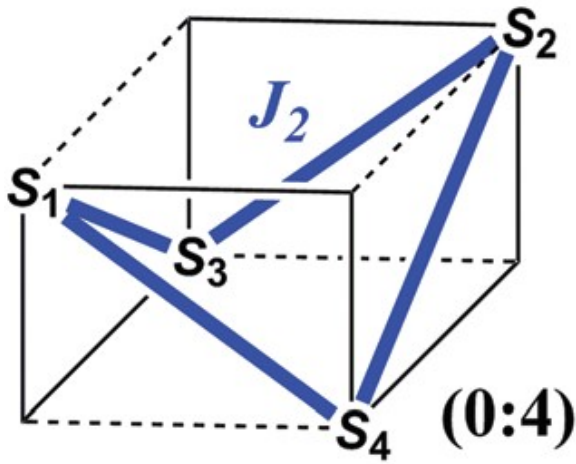
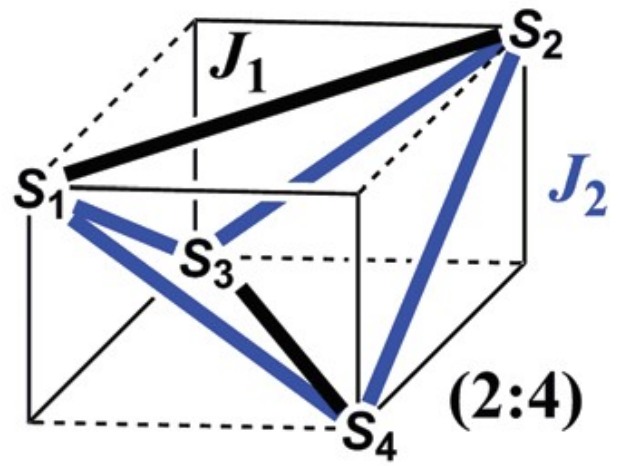
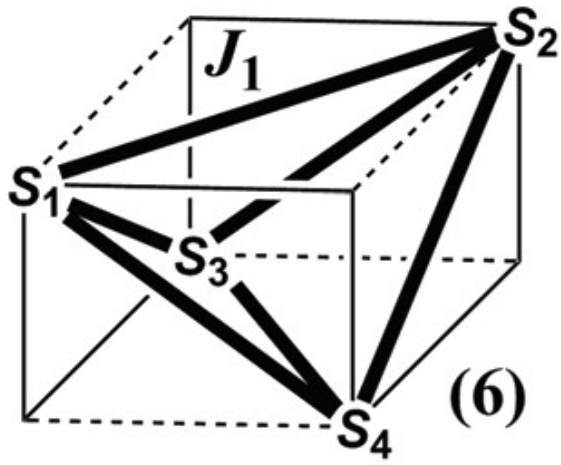
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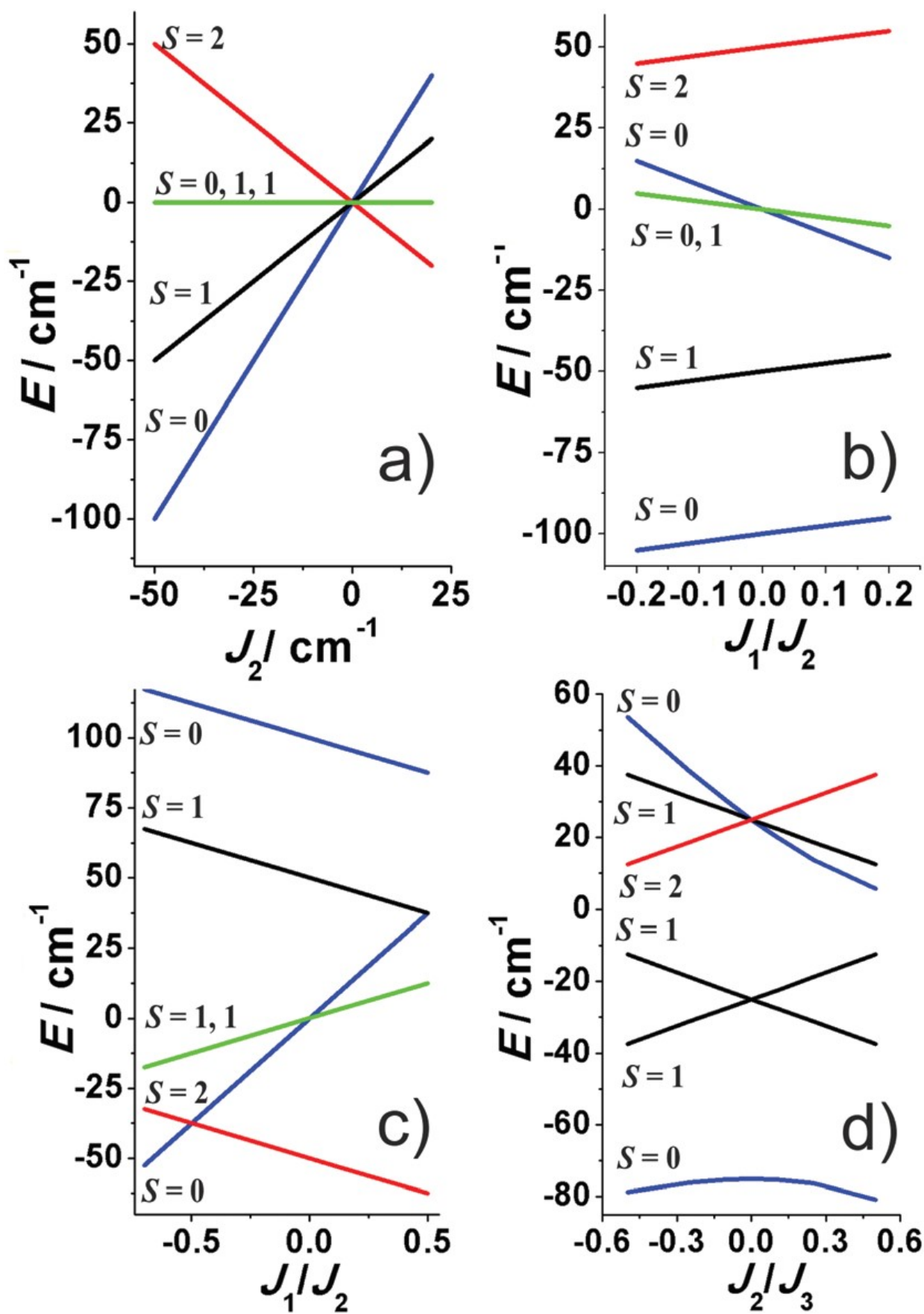
SCHEME 3

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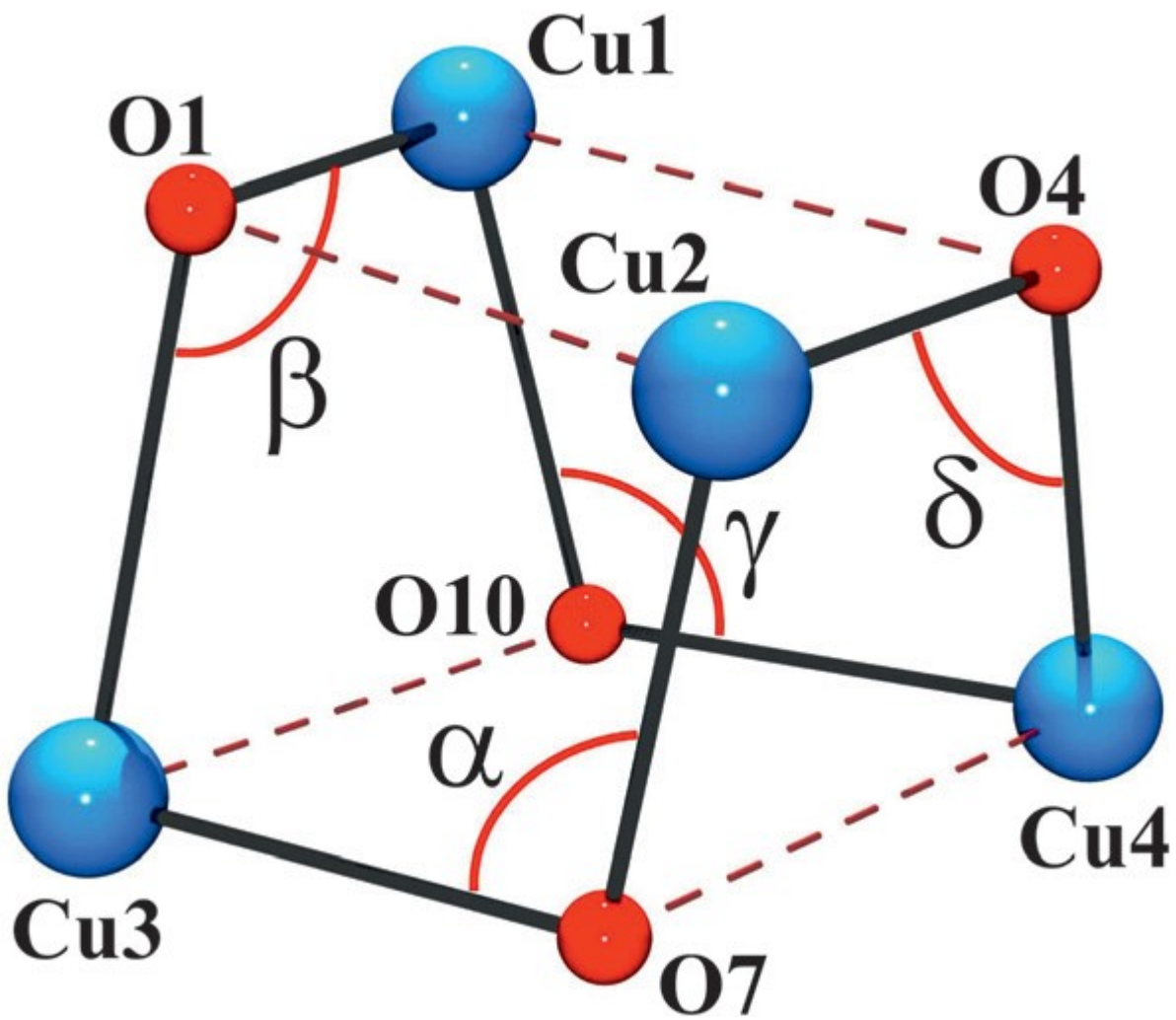
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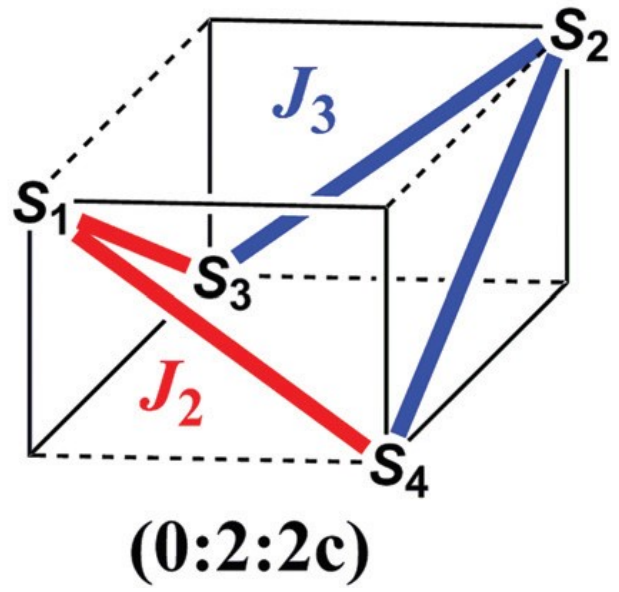
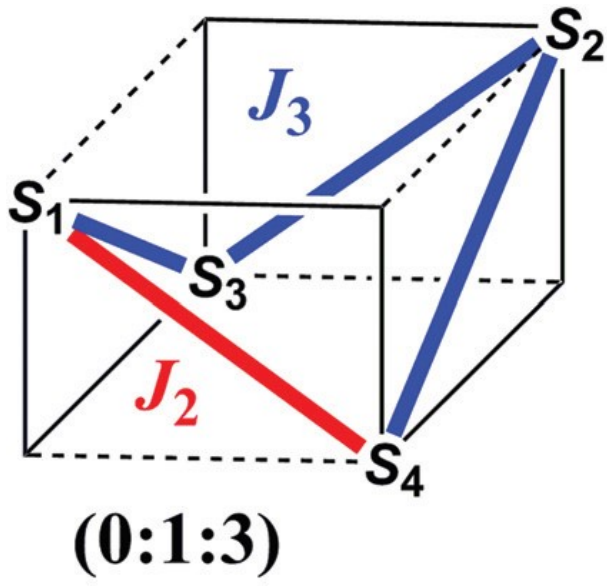
FIGURE 5



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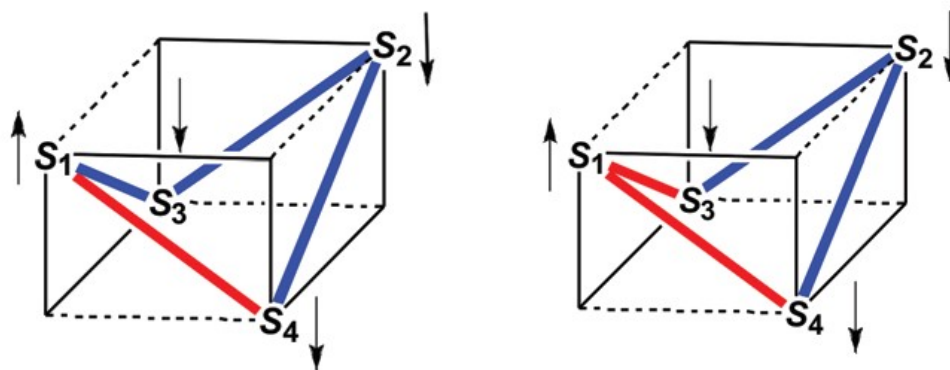
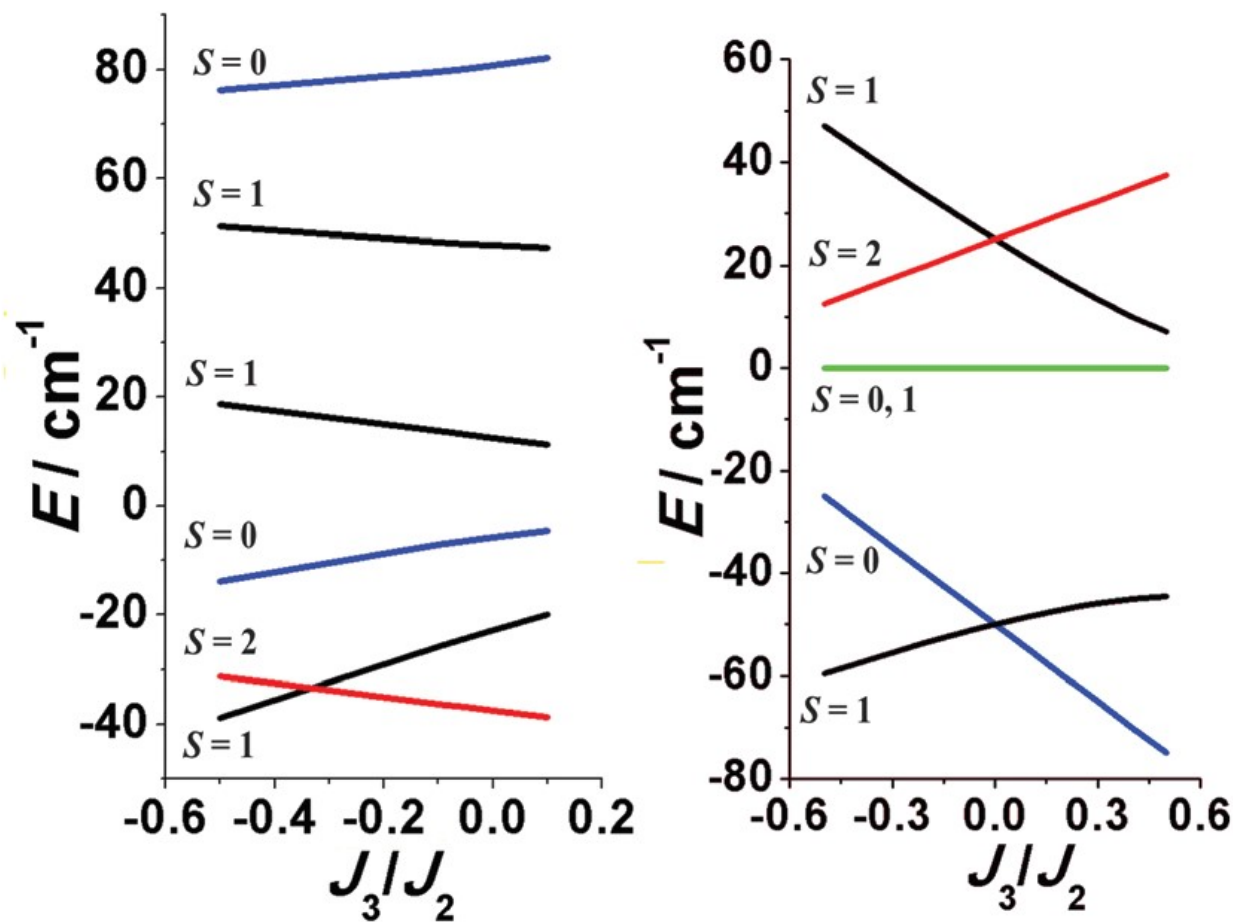
SCEHME 4



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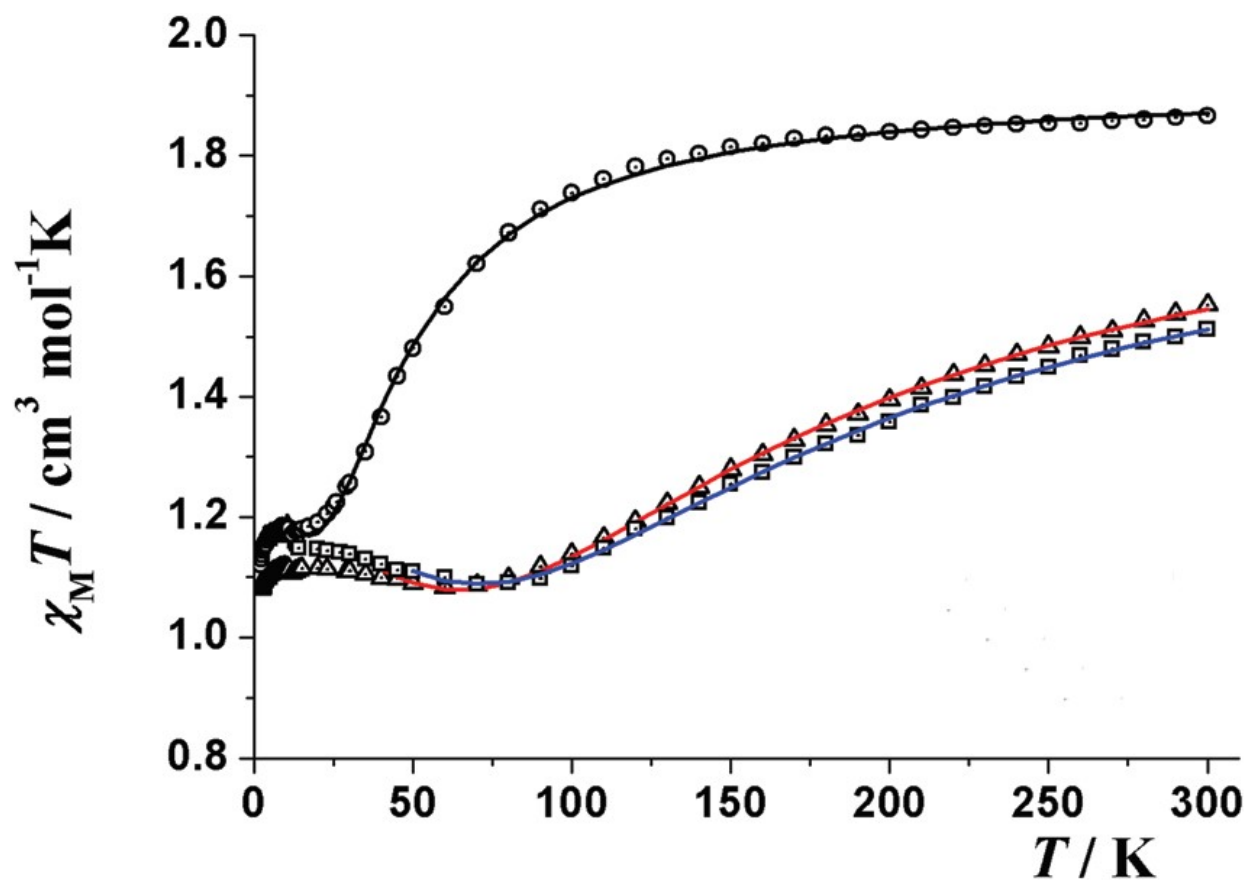
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FIGURE 6



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FIGURE 7

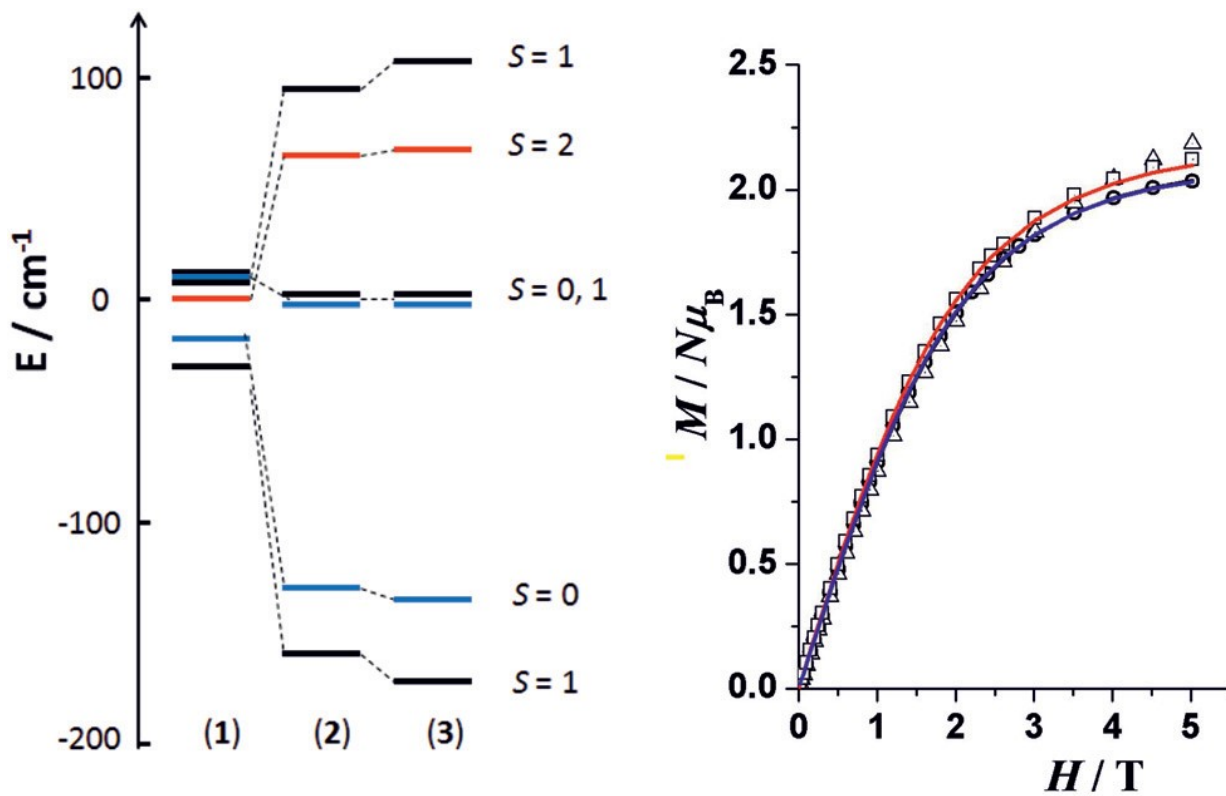


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FIGURE 8



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547 **Table 1** Crystal data, collection and structure refinement details for the X-ray structure determination of
 548 complexes 1–3

	(1)	(2)	(3)
Formula	C ₄₅ H ₁₆ Cu ₄ F ₃ N ₂ O ₁₃	C ₅₂ H ₄₀ Cu ₄ F ₄ N ₄ O ₁₂	C ₂₃₀ H ₁₇₈ Cu ₁₆ F ₁₆ N ₁₆ O ₂₅
<i>F</i> _w	1165.95	1243.04	5126.31
System	Monoclinic	Monoclinic	Monoclinic
Space group	<i>P</i> 2 ₁ / <i>c</i>	<i>C</i> 2/ <i>c</i>	<i>C</i> 2/ <i>c</i>
<i>a</i> /Å	13.344(13)	38.530(3)	41.241(2)
<i>b</i> /Å	18.937(2)	13.529(11)	13.4071(6)
<i>c</i> /Å	19.212(16)	19.446(14)	19.5890(9)
<i>α</i> /°	90	90	90
<i>β</i> /°	115.388(5)	92.806(4)	107.999(2)
<i>γ</i> /°	90	90	90
<i>V</i> /Å ³	4385.9(7)	10 125.0(14)	10 301.1(8)
<i>Z</i>	4	8	2
<i>T</i> , K	293(2)	302(2)	100(2)
<i>λ</i> (MoK α), Å	0.71073	0.71073	0.71073
ρ_{calc} , g cm ⁻³	1.766	1.631	1.653
μ (MoK α), mm ⁻¹	2.001	1.740	1.715
<i>R</i>	0.0676	0.0485	0.0326
ωR^2	0.1683	0.1265	0.0837

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552 **Table 2** Main bond distances (Å) and angles (°) for complexes 1–3
 553

Distance Å	(1)	(2)	(3)
Cu1–O1	1.942(2)	1.961(7)	1.969(1)
Cu3–O1	1.983(2)	1.926(7)	1.931(1)
Cu3–O7	1.947(2)	1.929(7)	1.911(1)
Cu2–O7	1.914(2)	1.914(8)	1.948(1)
Cu2–O4	1.916(3)	1.945(6)	1.959(1)
Cu4–O4	1.929(3)	1.931(7)	1.937(1)
Cu4–O10	1.931(2)	1.980(6)	1.980(1)
Cu1–O10	1.922(2)	1.960(7)	1.970(1)
Cu1–O4	2.697(3)	2.651(6)	2.697(1)
Cu2–O1	2.344(3)	2.462(7)	2.417(1)
Cu3–O10	2.410(3)	2.359(6)	2.378(1)
Cu4–O7	2.594(3)	2.962(7)	3.253(1)
Angles (°)	(1)	(2)	(3)
Cu1–O10–Cu4	108.7(1)	104.3(3)	104.23(5)
Cu4–O4–Cu2	112.2(1)	120.1(3)	124.32(6)
Cu2–O7–Cu3	100.3(1)	110.9(4)	110.77(6)
Cu3–O1–Cu1	102.6(1)	101.5(3)	100.80(5)

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557 **Table 3** Magnetic data for the (4 + 2) copper cubes reported in the literature. All J values have been
 558 normalized to the $-JS_x \cdot S_y$ Hamiltonian
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560	561	562	563	564	565
CCDC	Model/ Hamiltonian	$J_1/J_2/J_3$ ^a (cm ⁻¹)	Ground state	Ref.	
563 Cubes with at least one Jahn-Teller Cu-O distance <2.6 Å					
564	ELEYIE (6)/(1)	-20.8	S = 0	13	
565	WEMSUE (6)/(1)	-4.5	S = 0	4c	
566	GIBHAC (2:4)/(2)	+7.6/-21.7	S = 0	14	
567	NINPEG (2:4)/(2)	-5.2/-74.8	S = 0	15	
568	QOMRAL (2:4)/(2)	-2.6/-50	S = 0	16	
569	VEGROP (2:4)/(2)	-6.4/-10.9	S = 0	17	
570	BUFTUR (2:4)/(2)	-19.8/+41.0	S = 2	18	
571	CAQDAZ (2:4)/(2)	-32.6/+89.8	S = 2	19	
572	DARKOW (2:4)/(2)	-14.2/+57.0	S = 2	20	
573	FEVYAH (2:4)/(2)	-1/+65.0	S = 2	21	
574	IHELOX (2:4)/(2)	-31.8/+66.0	S = 2	22	
575	LITXOD (2:4)/(2)	+10.2/+39.8	S = 2	11c	
576	NAXBET (2:4)/(2)	-35.2/+7.2	S = 2	23	
577	QOMREP (2:4)/(2)	+15.2/-9.4	S = 2	16	
578	XEXZUX (2:4)/(2)	-21.4/+54.6	S = 2	24	
579	XINYUP (2:4)/(2)	-10.5/+61.0	S = 2	25	
580	XOVVUA (2:4)/(2)	-33.5/+67.0	S = 2	26	
581	XOXGEY (2:4)/(2)	-20.6/+41.2	S = 2	27	
582	FEJMIS (0:4)/(3)	-99.2	S = 0	28	
583	HAKXIB (0:4)/(3)	+10.4	S = 2	29	
584	MUGWUH (2:2:2)/(5)	+3.0/+24.4/+64.8	S = 2	11b	
585	WEMTAL (2:2:2)/(5)	-1.2/-1.2/+74.2	S = 2	4c	
586 Cubes with all Jahn-Teller Cu-O distances >2.6 Å					
587	NILDAP (6)/(1)	-118.8	S = 0	30	
588	BAQYAV (0:4)/(3)	-27.1	S = 0	31	
589	JELPUL (0:4)/(3)	-130.0	S = 0	32	
590	UFATOL (0:4)/(3)	-117.0	S = 0	33	
591	UFATUR (0:4)/(3)	-111.0	S = 0	33	
592	BOGCOP (0:4)/(3)	+34.2	S = 2	34	
593	DARKUC (0:4)/(3)	+34.8	S = 2	20	
594	BOQZUD (2:4)/(2)	+27.2/-69.8	S = 0	35	
595	LOCPIE (0:4)/(2)	-73.6	S = 0	36	
596	NODHEV (2:4)/(2)	+6.2/-80.5	S = 0	11a	
597	POLKEH (2:4)/(2)	-46.0/-136.0	S = 0	37	
598	VAVGUW (2:4)/(2)	-5/-75.0	S = 0	38	
599	LIJQEA-IE	Uncoupled	—	39	
600	GUFJEX01	-72.6	S = 0	40	
601 Disputable interesting cases					
602	ASUPEJ01 ^a (2:4)/(2)	-18/+38.4	S = 2	41	
603	SAPYON ^a (2:4)/(2)	-18.4/+14.7	S = 2	41	
604	SAPYUE ^a (2:4)/(2)	-15.6/+33.3	S = 2	41	
605	DIBTAL ^b No fit	—	S = 1 ?	42	
606	MOYJUH ^b (2:4)/(2)	-11.2/+7.6	S = 1 ?	43	
607	MOYKAO ^b		S = 1 ?	43	

608 ^a Complexes for which an S = 1 ground state was erroneously assigned.

609 ^b Complexes that probably have an S = 1 ground state.