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Influence of nitrogen donor ligands on the coordination modes of copper(II) 2-nitrobenzoate complexes: structures, DFT calculations and magnetic properties†

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Three novel copper(ii) 2-nitrobenzoate complexes, $[Cu(\gamma-pic)_2(2-nitrobenzoate)_2]$ **1**, $[Cu(\beta-pic)_2(2-nitrobenzoate)_2]$ benzoate)₂(H₂O)₂] **2** and $[Cu_2(H_2tea)_2(2-nitrobenzoate)_2]\cdot 2(H_2O)$ **3** (where γ -pic = 4-methylpyridine, β -pic = 3-methylpyridine and H₂tea = mono deprotonated triethanolamine), were isolated by addition of γ-pic, β-pic and H₃tea to the hydrated copper(II) 2-nitrobenzoate, [Cu₂(2-nitrobenzoate)₄(H₂O)₂]·2H₂O, suspended in a methanol:water (4:1) mixture. The newly synthesized complexes have been characterized by elemental analyses, thermogravimetric analysis (TGA), spectroscopic techniques (EPR, IR and UV/Visible), magnetic susceptibility measurements, single crystal X-ray structure determination and DFT study. All compounds crystallize in the monoclinic crystal system with the $P2_1/c$ space group. X-ray structure determination revealed the presence of monomers in both 1 and 2 and dimer in 3 with the deprotonated oxygen atom of the H₂tea ligand bridging two Cu(II) atoms. Two co-crystallized water molecules are also present in 3. The crystal lattice is stabilized by C-H...O hydrogen bonding interactions in 1 and O-H···O, C-H···O hydrogen bonding interactions in 2 and 3. The dimeric complex exhibits relatively strong ferromagnetic exchange with $J = -100 \text{ cm}^{-1}$ (corresponding to $H = JS_1S_2$). The zero-field splitting parameters (zfs) of the dimer triplet states D and E were derived from HF EPR spectra recorded at moderately low temperatures. The sign of D was determined from low-temperature HF EPR spectra.

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Introduction

In recent years, crystal engineering based on organic and/or inorganic molecular building blocks has been rapidly expanding owing to the novel and diverse topologies and potential applications in host-guest chemistry, catalysis, electrical conductivity and magnetism. 1-6 One of the important strategies is to establish the likely connections between organic and/or inorganic molecular building blocks by exploiting non-covalent interactions.^{7,8}

In the case of transition metal complexes, these interactions are established mainly among the 'organic' parts of the molecules, and can be modulated through a suitable choice of the ligands. Copper(II) complexes are among the most widely studied coordination compounds because of their wide range of applications, 9-15 in particular when the ligands are alkyl/aryl carboxylate anions, which can exhibit a wide variety of coordination modes (monodentate, symmetric\asymmetric chelating, and bidentate\ monodentate bridging). Moreover, properties of copper(II) carboxylates and the coordination abilities of carboxylate ions can be varied to a large extent using nitrogen donor ligands with appropriate functionalization.¹⁶⁻¹⁹ A combined stereoelectronic effect that emerges as a result of both the carboxylate anion and the ancillary N-donor ligand would dictate the final nature and structure of the complex formed. That is, in a nutshell, the chosen carboxylate ligand may show a propensity to exhibit different coordination modes depending upon the type of N-donor ligand chosen.

In this direction, with the aim of comparing the structural and packing features of complexes in which different N-donor ligands participate, we have chosen two very similar nitrogendonor ligands γ -pic (4-methyl pyridine) and β -pic (3-methyl

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[†] Electronic supplementary information (ESI) available: TGA, IR and UV characterization details, and figures depicting TGA curves, IR spectra and crystal packing for the three complexes. CCDC 915710 (1), 915711 (2) and 915712 (3). For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/ c3nj00736g

pyridine) which are rigid in their molecular topology along with a third N-donor ligand triethanolamine (H₃tea) which is highly flexible in terms of molecular conformation. This ligand after deprotonation (i.e. H₂tea) will be endowed with two -OH groups which may take part either singly or in a combined way in the hydrogen bonding as effective proton donors. In addition, a theoretical approach that can model these structures and their bonding features would be rewarding. Thus, this paper reports the synthesis, characterization and DFT study of three novel copper(II) complexes with different nitrogen donor ligands, $[Cu(\gamma-pic)_2(2-nitrobenzoate)_2]$ (1), $[Cu(\beta-pic)_2(2-nitrobenzoate)_2]$ $2(H_2O)_2$ (2) and $[Cu_2(H_2tea)_2(2-nitrobenzoate)_2]\cdot 2(H_2O)$ (3).

Particular attention has been devoted to complex 3 where the presence of the H2tea ligand favours the formation of a dinuclear molecule in which two Cu atoms are bridged by two oxygen atoms. In general, the magnetic behavior of transition metal complexes containing more than one metal atom with unpaired electrons is affected by the metal-metal interaction. The case discussed in this paper is that of weakly interacting metal ions (with the Cu...Cu distance in the range 3-5 Å), where the coupling between their electrons leads to low-lying excited states of different spin which can be thermally populated. The resulting magnetic behavior can be antiferromagnetic or ferromagnetic, and the ferromagnetic/antiferromagnetic coupling is strictly related to structural parameters such as the Cu-O-Cu angle and the Cu···Cu distance. Accordingly, the magnetic characterization of 3 has been performed both from a theoretical and experimental point of view.

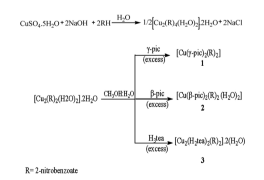
Experimental

Three copper(II) carboxylate complexes were synthesized by reacting copper(II) 2-nitrobenzoate with an appropriate N-donor ligand in a methanol-water mixture and allowing the solution to evaporate at room temperature as shown in Scheme 1.

All three complexes have been characterized and investigated by elemental analyses, TGA, spectroscopic (FT-IR, UV/Visible, EPR) and magnetic susceptibility studies, conductance measurements, as well as by single-crystal X-ray crystallography and DFT.

Syntheses

Synthesis of $[Cu_2(2-nitrobenzoate)_4(H_2O)_2] \cdot 2H_2O$. Sodium 2-nitrobenzoate was prepared by dissolving 2 g (0.012 mol) of



Scheme 1 Schematic representation of the synthetic procedure.

2-nitrobenzoic acid and 0.479 g (0.012 mol) of sodium hydroxide in a minimum amount of water. 1.490 g (0.0059 mol) of copper sulphate was dissolved in a minimum amount of water. Upon mixing both solutions at room temperature a green coloured precipitate appeared immediately. The precipitated product was filtered through fine paper, washed with water and dried at room temperature.

Synthesis of $[Cu(\gamma-pic)_2(2-nitrobenzoate)_2]$. 1: 0.05 g of the green coloured copper(II) 2-nitrobenzoate complex was suspended in a 4:1 methanol: water mixture and γ -pic was added till a clear dark blue coloured solution was obtained. When the solution was allowed to evaporate at room temperature, blue coloured crystals appeared after 3 days. Complex 1 is soluble in methanol and insoluble in water and acetone. Complex 1 decomposed at 190 °C. Anal. calcd for C₂₆H₂₂CuN₄O₈ (%): C, 53.65; H, 3.78; N, 9.63; Cu, 10.92. Found: C, 52.45; H, 3.66; N, 9.52; Cu, 10.73. IR/cm⁻¹ (KBr): 3565(m), 3317(m), 3128(b), 3208(m), 3085(w) 1607(s), 1564(ms), 1524(s), 1443(w), 1389(s), 1350(m), 1151(m), 1071(ms), 1034(ms), 977(w), 959(w), 862(m), 835(m), 805(m), 795(m), 785(m) 752(m), 700(m), 648(m), 589(w), 554(m), 493(ms), 459(m). UV/Visible (CH₃OH) λ_{max} = 732 nm ($\varepsilon_{\text{max}} = 94.07 \text{ mol}^{-1} \text{ L cm}^{-1}$).

 $[Cu(\beta-pic)_2(2-nitrobenzoate)_2(H_2O)_2].$ **Synthesis** of Complex 2 was synthesized in a similar manner to complex 1 by adding β -picoline. When the deep blue colored solution was allowed to evaporate at room temperature, blue coloured crystals appeared after 3 days. Complex 2 is soluble in methanol and insoluble in water and acetone. Complex 2 melted at 210 °C. Anal. calcd for C₂₆H₂₆CuN₄O₁₀ (%): C, 50.52; H, 4.21; N, 9.07; Cu, 10.28. Found: C 50.45; H, 4.16; N, 8.72; Cu, 10.08. IR/cm^{-1} (KBr): 3524(m), 3371(b), 3235(w), 1593(s), 1567(s), 1530(s), 1481(m), 1393(s), 1367(sh), 1194(w), 1130(w), 1064(w), 862(w), 836(m), 797(m), 744(m), 698(ms), 649(b), 598(w), 513(w), 468(ms). UV/Visible (CH₃OH) $\lambda_{\text{max}} = 738 \text{ nm} (\epsilon_{\text{max}} = 102.5 \text{ mol}^{-1} \text{ L cm}^{-1}).$

Synthesis of $[Cu_2(H_2tea)_2(2-nitrobenzoate)_2]\cdot 2(H_2O)$. 3: Complex 3 was synthesized in a similar manner to complexes 1 and 2 by adding H₃tea. When the deep blue colored solution was allowed to evaporate at room temperature, turquoise blue coloured crystals appeared after four days. The crystals were washed with hexane and vacuum dried. Complex 3 is soluble in methanol and insoluble in water and acetone. Complex 3 melted at 135 °C. Anal. calcd for C₂₆H₃₆Cu₂N₄O₁₄·2(H₂O) (%): C, 39.44; H, 5.05; N,7.07; Cu, 16.05. Found: C, 39.25; H, 4.46; N, 6.92; Cu, 15.96. IR/cm⁻¹ (KBr): 3418(b), 3225(b), 2944(w), 1601(m), 1566(m), 1548(s), 1537(s), 1450(m), 1385(s), 1171(w), 1137(w), 1084(s), 1053(m), 1024(m), 918(m), 904(m), 813(m), 780(m), 749(w), 705(w), 695(m), 681(w), 648(m), 568(m), 502(w), 466(m). UV/Visible (CH₃OH): $\lambda_{\text{max}} = 741 \text{ nm} (\varepsilon_{\text{max}} = 63.5 \text{ mol}^{-1} \text{ L cm}^{-1})$.

Details of the TGA and spectroscopic characterization of complexes 1-3 are available in the ESI.†

Materials and physical measurements

Analytical grade reagents were used without any further purification. Carbon, hydrogen and nitrogen were measured micro-analytically using an automatic Perkin Elmer 2400 CHN elemental analyzer and copper was determined gravimetrically.20

FT-IR spectra were recorded as KBr pellets using a Perkin Elmer Spectrum RXFT-IR system. The electronic spectrum was recorded in H₂O using a Hitachi 330 spectrophotometer. The thermogravimetric analysis (TGA) was conducted using an SDT Q600 instrument. The samples contained in the alumina pan were heated from 25 °C to 1000 °C at a constant rate of 10 °C min⁻¹ in a nitrogen environment with a flux (flow rate) of 10 mL min⁻¹. Conductance measurements were performed using a Pico Conductivity Meter (Model CNO4091201, Lab India) in aqueous medium at 25 °C by using double distilled water. Magnetic measurements for 1 and 2 at room temperature were carried out using the Gouy's method with following specifications: pole face diameter: 10.2 cm; pole gap: 4.0 cm; current: 7.0 & magnetic induction: 0.6 Tesla. Magnetic susceptibility measurements for 3 over the temperature range 1.8-300 K were performed at a magnetic field of 0.5 T using a Quantum Design SQUID MPMSXL-5 magnetometer. Correction for the sample holder as well as the diamagnetic correction γ_D which was estimated from the Pascal constants were applied.²¹

X-band EPR High-field and high-frequency EPR spectra at temperatures ranging from ca. 6 K to 290 K were recorded for 3 on a home-built spectrometer at the EMR facility of the NHMFL.²²

The instrument is a transmission-type device in which microwaves are propagated in cylindrical lightpipes. The microwaves were generated by a phase-locked Virginia Diodes source generating a frequency of 13 \pm 1 GHz and producing its harmonics of which the 2nd, 4th, 6th, 8th, 16th, 24th and 32nd were available. A superconducting magnet (Oxford Instruments) capable of reaching a field of 17 T was employed. The EPR spectra at X- and Q-bands were measured using a Bruker Elexsys E500 spectrometer equipped with a NMR teslameter and frequency counters. The simulations of the experimental spectra were performed using computer programs employing full diagonalization of the spin Hamiltonian matrix, written by one of us.

Crystallography

The crystallographic data for 1, 2, and 3 were collected using a Nonius Kappa CCD diffractometer at room temperature using

graphite-monochromated MoK α radiation ($\lambda = 0.71073$ Å). Data sets were integrated using the Denzo-SMN package²³ and corrected for Lorentz-polarization and absorption effects.²⁴ The structures were solved by direct methods (SIR97)²⁵ and refined by full-matrix least-squares methods with all nonhydrogen atoms being anisotropic. As for hydrogen atoms, those belonging to methyl groups in 1 and all the C-H in 2 and 3 were included on calculated positions, riding on their carrier atoms; the others were located in the difference-Fourier map and refined isotropically. All other calculations were performed using SHELXL-97²⁶ implemented in the WINGX system of programs.²⁷ The experimental details are reported in Table 1.

Quantum mechanical calculations

The 'in vacuum' fully optimized geometries of $[Cu(\gamma-pic)_2 (2-nitrobenzoate)_2$ (1) and $[Cu(\beta-pic)_2(2-nitrobenzoate)\cdot$ 2(H₂O)₂] (2) complexes have been obtained by density functional theory (DFT) calculations carried out at the B3LYP/6-31G(d,p) level of theory using the Firefly system of programs. 28,29

ORCA software³⁰ was used to perform the "broken symmetry" DFT calculations of exchange parameters J for 3. All atoms were retained at the positions determined by the X-ray structures. The "broken symmetry" procedure applied to a dimeric copper(II) complex first performs a Self-Consistent Field (SCF) calculation for a high-spin molecule (HS) with the spin equal to 1. In the next stage, another SCF calculation is performed taking the spin on one Cu atom 'up' and 'down' on another, which is referred to as the broken symmetry state (BS). Finally, the magnitude of J (for the spin Hamiltonian 1) is evaluated as $J = 2(E_{HS} - E_{BS})/(\langle S^2 \rangle_{HS} - \langle S^2 \rangle_{BS})$, where E are the energies and $\langle S \rangle^2$ are the expectation values of the spin-squared operator in the HS and BS states.

Ahlrichs-type basis set TZVPP³¹ for copper(II) and all coordinated atoms, and VDZ311a for other atoms were used, combined with the B3LYP³² functional. Ahlrichs polarisation functions from basis H-Kr R and auxiliary bases from the TurboMole library were also used. 31b

Table 1 Experimental de	etails
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	1	2	3
Chemical formula	C ₂₆ H ₂₂ CuN ₄ O ₈	C ₂₆ H ₂₆ CuN ₄ O ₁₀	$C_{26}H_{36}Cu_2N_4O_{14}\cdot 2(H_2O)$
$M_{ m r}$	582.02	618.05	791.70
Crystal system, space group	Monoclinic, $P2_1/c$	Monoclinic, $P2_1/c$	Monoclinic, $P2_1/c$
$a, b, c (\mathring{A})$	7.8506 (1), 21.8740 (4),	8.0020 (2), 12.2397 (3),	7.6633 (2), 27.2585 (7),
	8.1508 (2)	14.5180 (3)	8.5840 (2)
β (°)	103.0120 (6)	105.1790 (11)	117.5840 (11)
Z	2	2	2
$V(\mathring{A}^3)$	1363.75 (5)	1372.32 (6)	1589.29 (7)
$\mu (mm^{-1})$	0.86	0.86	1.42
Crystal size (mm)	$0.52 \times 0.47 \times 0.17$	$0.26 \times 0.20 \times 0.18$	$0.55 \times 0.26 \times 0.12$
No. of measured, independent and observed	16 268, 3883, 3325	16 455, 3294, 2519	6658, 3325, 2884
$[I > 2\sigma(I)]$ reflections			
$R_{ m int}$	0.034	0.039	0.056
$R[F^2 > 2\sigma(F^2)], wR(F^2), S$	0.038, 0.115, 1.08	0.043, 0.136, 1.02	0.045, 0.142, 0.99
No. of reflections	3883	3294	3325
No. of parameters	211	196	249
$\Delta \rho_{\rm max}$, $\Delta \rho_{\rm min}$ (e Å ⁻³)	0.31, -0.44	0.32, -0.61	0.54, -1.03

Results and discussion

The composition of 1, 2 and 3 was established by elemental analysis. The mode of coordination of the carboxylate ligand in these complexes was indicated by IR spectral criterion $\Delta v = v_{\text{carboxylato(asym)}} - v_{\text{carboxylato(sym)}}$ as reported in literature. ^{33,34}

Molecular geometry and coordination

To determine the structures unambiguously and to analyse the packing modes, single X-ray crystal structure determination of complexes 1, 2 and 3 was undertaken. Fig. 1-3 report the ORTEPIII diagrams for 1-3, respectively.

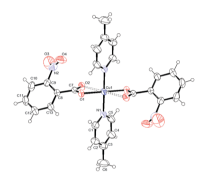


Fig. 1 ORTEPIII view and the atom numbering scheme for 1. Thermal ellipsoids are drawn at the 50% probability level. Contact distances are shown as dashed lines

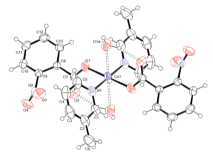


Fig. 2 ORTEPIII view and the atom numbering scheme for 2. Thermal ellipsoids are drawn at the 40% probability level. Contact distances and hydrogen bonds are shown as dashed lines

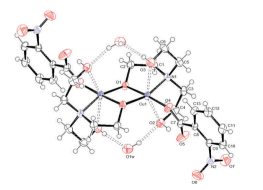


Fig. 3 ORTEPIII view and the atom numbering scheme for 3. Thermal ellipsoids are drawn at the 40% probability level. Contact distances and hydrogen bonds are shown as dashed lines

Table 2 Selected geometric parameters (Å, °)

1 Cu1-N1	2.005(1)	Cu1O2	2.518(1)
Cu1-O1	1.987(1)	04102	2.310(1)
N1-Cu1-O1	89.24(5)	N1-Cu1-O2	88.19(5)
Cu· · · Cu contacts	03.21(0)	111 041 02	00.13(0)
Cu···Cu	7.8506(1)	8.1508(2)	
2			
Cu1-N1	1.997(2)	Cu1···O1W	2.739(2)
Cu1-O1	1.971(2)		, , ,
N1-Cu1-O1	91.02(7)	N1-Cu1-O1w	91.10(8)
O1-Cu1-O1w	96.00(7)		()
CuCu contacts			
CuCu	8.0020(2)	9.4945(1)	
3			
Cu1-O1	1.946(2)	Cu1···O3	2.478(2)
Cu1-N1	2.042(2)	Cu1-O4	1.966(3)
Cu1···O2	2.576(2)	$Cu \cdot \cdot \cdot Cu^a$	2.9187(4)
Cu1-O1 ^a	1.934(2)		. ,
O1-Cu1-N1	85.00(9)	N1-Cu1-O3	76.3(1)
O1-Cu1-O2	90.59(9)	N1-Cu1-O4	98.70(9)
O1-Cu1-O3	97.2(1)	O2-Cu1-O4	88.83(9)
N1-Cu1-O2	78.40(9)	O3-Cu1-O4	85.0(1)
O1-Cu1-O1 ^a	82.46(9)	Cu1-O1-Cu1 ^a	97.5(1)
^a Symmetry code: (1	-x,-y,-z).		

Relevant geometrical parameters are reported in Table 2. In 1 and 2, the Cu atom lies on an inversion center, while in 3 a symmetry center is positioned just in the middle of the binuclear complex. In both 1 and 2 complexes, the Cu atom is linked to two picoline and two monodentate benzoate ligands, bonded via O1. In addition, it is involved in two further second-order interactions, giving a final (4 + 2) coordination: in 1 the carboxylic oxygens (O2) are at a Cu-O distance of 2.518(1) Å, while in 2 two centrosymmetrically related water molecules are at a distance of 2.739(2) Å (Table 2). The angles formed by the O2-Cu1 and O1w-Cu1 lines with the normal to mean basal planes measure 32.60(4) and 6.08(7)° in 1 and 2, respectively. Complex 3 is dimeric with the de-protonated oxygen atom of the H2tea ligand forming a bridge between two centrosymmetrically related Cu(II) atoms. The coordination geometry of 3 can be described, in analogy with those reported in the literature for analogous Cu(II) benzoate,35 3-chlorobenzoate36 and 4-methylbenzoate³⁷ complexes, as distorted tetragonal bipyramidal (or highly distorted octahedral), with each H2tea molecule acting as a tetradentate ligand or as square-planar with two additional interactions like in 1 and 2.

Actually, the equatorial plane in 3 is defined by one carboxylate oxygen atom of 2-nitrobenzoate, two bridging alkoxo oxygen atoms and one nitrogen while O2 and O3 atoms of H2tea are located at much longer distances (Table 2); the Cu atom is displaced by 0.023 Å from the least-square plane passing through (O1, N1, O4) and the angles formed by Cu1-O2 and Cu1-O3 lines with the normal to this plane are 16.44(9) and 11.19(8)°, respectively. For each dimeric moiety, two water molecules of crystallization are present in the crystal.

The Cu-N distances are similar in all the three complexes and do not show any significant discrepancy from structural parameters of related Cu compounds. The Cu-O distances

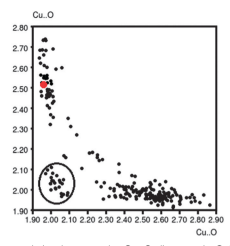


Fig. 4 Intercorrelation between the Cu-O distances in Cu(II) benzoate complexes. Data for compound 1 are reported as red points.

display a wide range of values, from 1.934(2) Å (oxo-bridge, complex 3) to 2.739(2) Å (water molecule, complex 2), that can be more appropriately considered as a contact distance. The histogram of the Cu-O(benzoate) distances, obtained from a search on the CSD database (278 hits) considering both the oxygens linked to the metal, is clearly bimodal showing two distinct bond length distributions around 2.0 and 2.6 Å, respectively, in perfect agreement with the structural data of 1 and 3. Moreover, Fig. 4, obtained by the same CSD search, suggests that these two distances are intercorrelated; if one increases, the other decreases. Geometrical data of complex 1 (red point) are perfectly in line with the others. The points enclosed in a circle, which do not follow the curve trend, refer to particular structures: of polymeric type, with huge ligands, or having Cl⁻ anions as ligands.

As far as the binuclear complex 3 is concerned, a comparison with the similar Cu-H₂tea complexes above mentioned^{35–37} shows that the molecular arrangement around the Cu atoms and the geometrical parameters are strictly comparable in all of them; benzoates are acting as monodentate ligands, with the oxygen of anionic H2tea bridging two metals, and one OH group of the H2tea being involved in an intramolecular hydrogen bond with the free benzoate oxygen. Moreover, two cocrystallized water molecules for each binuclear complex are always present connecting different molecules in exactly the same way as in complex 3 (vide infra), suggesting that in these structures, the presence of water is essential for the robustness of the crystalline framework.

It is interesting to note that complexes 1 and 2 present a different Cu environment in spite of having almost identical ligands (p-methyl and o-methyl pyridine, respectively). In order to explore whether the choice of different picoline molecules can affect the possibility of water inclusion in the first or the second coordination sphere, the equilibrium geometry of both complexes has been obtained by DFT calculations starting from the same initial point, i.e. by adding two Cu-coordinated water molecules to the picoline/benzoate skeleton at a distance of 2.40 Å or using an initial geometry with no coordinated water molecules.

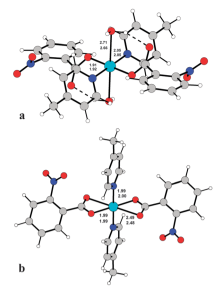


Fig. 5 DFT equilibrium geometries of complex 2 (a) in diaquo-form and (b) without coordinated water molecules. Relevant distances are shown together with the corresponding calculated bond lengths for complex 1 (reported underneath).

The results are reported in Fig. 5. For the diaquo complexes, the Cu-O, Cu-N distances are strictly comparable and the energies of the calculated molecules are practically identical: the γ -picoline complex is more stable than the β -picoline one by 0.36 kcal mol⁻¹. The same results have been obtained for complexes without water, but in this case the γ-picoline compound is more stable by 2.13 kcal mol⁻¹; this extra-stability is however not sufficient to explain the different coordination of the two compounds, which probably depends mostly on the solvent used (or in general on the crystallization conditions).

As mentioned at the outset, one of our objectives was to delineate the variation/change in the conformation of the carboxylate ligand upon the addition of three different N-donor ligands, two rigid (picolines) and one flexible (H3tea). Indeed, as anticipated, the -NO2 moiety is out of plane from the aromatic ring in all three, but the extent of variation is quite large, ranging from 42.4(3), 53.9(3) and 81.6(4)° for 1, 2 and 3 respectively (note that this variation is highest with the most flexible N-donor ligand H_3tea).

In an analogous manner, the -COO- moiety also gets rotated with respect to the aromatic plane, the highest rotation being observed in 1 and 2 (42.6(2) and 42.3(3)°) and almost in-plane of the aromatic ring in 3 $(5.2(2)^{\circ})$. Here also that distinction prevails; the rigid N-donor ligands are similar in behavior and the flexible one shows considerable deviation. It is clear that each of these structures would try to maximize the resonance energy by keeping both -NO2 and -COO groups in plane with the aromatic ring, but the opposing steric repulsion needs to be compensated by allowing them to move away from the plane to the extent that the lattice energy is minimum (most stable arrangement). Thus, in the most flexible deprotonated ligand H2tea (3) -COO is only few degrees away from the plane, but at the same time the -NO2 is almost orthogonal.

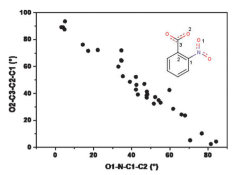


Fig. 6 Scatter plot of the out of plane angles for NO₂ and COO⁻ groups in Cu(2-nitrobenzoate) complexes.

In the rigid N-donors 1 and 2 the situation is in between, both the groups deviate to the extent of some 45°. These findings are supported by the survey of the conformation of the nitrobenzoate moiety in Cu complexes with N-donor ligands already present in literature. 21 structures of this type were found in the CSD; for each of them the NO2- and COO- out-of-plane was evaluated through the measure of the torsion angles O1-N-C1-C2 and O2-C3-C2-C1 (see Fig. 6). As shown in the plot, the two angles are strongly intercorrelated in the same way as found for the present complexes; the COO⁻ group is rarely coplanar with the phenyl moiety, and this happens when both the oxygens are 'busy' in covalent or strong H-bonding interactions within a same molecular complex, in which the Cu atom is linked to small or highly conformationally adaptable N-donor ligands.

This observation becomes interesting when we consider the mode of coordination in 1, 2 and 3. In 1, the carboxylate group is monodentate but the second oxygen is still interacting with the Cu atom, (Table 2) whereas in 2 one of the Cu-O distance becomes so large (3.185(2) Å) that the coordination is obviously monodentate and it is much higher yet in 3 (3.378(3) Å). That is, the nature of the N-donor ligand definitely exerts influence on the coordination mode and this in turn affects the structure and the packing architecture of the crystallized complex.

Packing

The hydrogen bonding structural parameters for the three complexes are reported in Table 3. In 1, the presence of a number of good hydrogen bonding acceptors but a lack of good hydrogen bonding donors allows only the formation of C-H- \cdots O bonds. The C-H- \cdots O1 contact on the two sides of the molecule generates ribbons linking the complexes along the c direction. These ribbons are in turn connected along the b direction through C-H \cdots O2 interactions. Conversely, in 2 and 3 the presence of water molecules or hydroxyl groups leads to strong O-H···O interactions, both intermolecular and intramolecular. In 2, the coordinated water molecules, besides being involved in intramolecular interactions, act as H-bond donors towards the O2 atoms of an adjacent molecule, giving rise to the $R^2_2(12)$ ring as shown in Fig. 7.

The resulting overall packing is made of parallel ribbons running along the a direction. In 3, the interaction network is

Table 3 Hydrogen bonding parameters; D = donor, A = acceptor (Å, °)

D-H···A	D-H	$\mathbf{D}{\cdot}{\cdot}{\cdot}\mathbf{A}$	$H\!\cdot\!\cdot\!\cdot\! A$	Angle
Complex 1				
$C2-H2\cdots O2^a$	0.95(2)	3.268(2)	2.36(3)	160(2)
$C4-H6\cdots O1^b$	0.90(3)	3.451(2)	2.56(3)	168(2)
Complex 2				
$O1W-H2W \cdot \cdot \cdot O2^{c}$	0.93(4)	2.728(3)	1.83(4)	163(4)
$O1W-H1W \cdot \cdot \cdot O2^d$	0.89(4)	2.892(3)	2.06(4)	154(4)
$C3-H3\cdots O1W^e$	0.93	3.426(4)	2.53	161
C6−H62···O3 ^f	0.96	3.440(5)	2.56	151
C12–H12···O2 g	0.93	3.397(3)	2.49	163
Complex 3				
O2-H2···O5	0.85(5)	2.631(4)	1.80(5)	165(4)
O1W-H2W···O2	0.84(5)	2.870(4)	2.05(6)	164(5)
$O1W-H1W\cdots O1^h$	0.82(7)	2.997(4)	2.28(7)	146(6)
$O3-H3\cdots O1W^{i}$	0.71(6)	2.711(4)	2.01(6)	171(6)
C1-H1B···O1W j	0.97	3.397(4)	2.61	138

 $\int_{0}^{a} \frac{x-1}{x}, y, z, \int_{0}^{b} \frac{x}{x}, y, z+1, \int_{0}^{c} \frac{1-x}{x}, -y, -z, \int_{0}^{d} \frac{x+1}{x}, y, z, \int_{0}^{e} \frac{x}{x}, \frac{1/2-y}{x}, z+\frac{1/2}{x}, \int_{0}^{e} \frac{x+1}{x}, y, z, \int_{0}^{e} \frac{x+1}{x}, -y, -z, \int_{0}^{e} \frac{x+1}{x}, -y, \int_{0}^{e} \frac{x+1}{x}, \int_$ -z. j x + 1, y, z + 1.

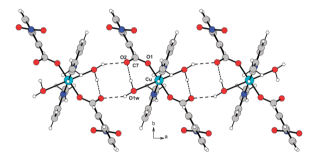


Fig. 7 O-H···O hydrogen bond arrangement in 2. Intermolecular and intramolecular hydrogen bonds are shown as dashed and dotted lines, respectively

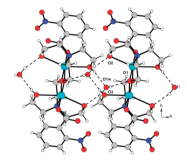


Fig. 8 O-H···O hydrogen bond arrangement in 3. Intermolecular hydrogen bonds are shown as dashed lines. For the sake of clarity, intramolecular O-H···O interactions are not shown.

somewhat similar but more complex than 2 due to the presence of a large number of hydrogen bond donor and acceptor groups. Water molecules of crystallization act as linkers by bridging the adjacent molecules; they behave as hydrogen bond donors towards O1 and O2 atoms and as acceptors to the O3 atom. This results in formation of molecular ribbons running along the a direction (Fig. 8). In both structures the crystal lattice is further stabilized by weak C-H···O interactions.

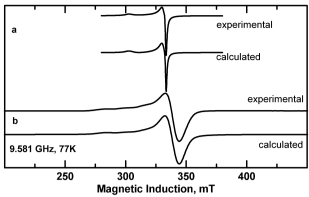


Fig. 9 X-band EPR spectra of powdered complexes (a) 1 and (b) 2 at 77 K (experimental and calculated with the parameters given in the text).

Magnetic and EPR measurements

Magnetic moments higher than the spin-only value of 1.73 are observed in copper(II) complexes (3d⁹), due to spin-orbit coupling. Monomeric copper(II) complexes generally show magnetic moments in the range 1.75 to 2.2 $\mu_{\rm B}$ at room temperature. The EPR spectra of the powdered complexes with $g_x = 2.056$, $g_y = 2.066$ and $g_z = 2.258$ parameters for 1 (Fig. 9a) and $g_x = 2.057$, $g_y = 2.057$ and $g_z = 2.262$ for 2 (Fig. 9b) are typical for axial symmetry of Cu(II) coordination polyhedron (with a very weak rhombic distortion) for 1 and axial symmetry for 2. The gparameter values for 1 and 2 correspond to the $d_{x^2-\nu^2}$ orbital of the unpaired electron ground state and are characteristic of the N2O2 donor set provided by the ligands in the xy Cu(II) plane. The low field "parallel" part of the spectrum for 2 exhibits signals due to hyperfine interaction between unpaired electrons and copper nucleus (nuclear spin I = 3/2) with $A_{\parallel} = 169 \times 10^{-4}$ cm⁻¹ (Fig. 9b). It is a rare case when the hyperfine splitting is resolved in a powder EPR spectrum of a Cu(II) complex suggesting a greater isolation of Cu(II) centers than in 1 due to water molecules at axial positions which are involved in hydrogen bonds with carboxyl groups. The Cu-Cu distances are in fact greater in 2 than in 1 (Table 2).

Magnetic susceptibility and EPR spectra of 3

The temperature dependence of the effective magnetic moment of 3 is shown in Fig. 10.

The moment is seen to increase with decreasing temperature, to reach a plateau between 25 and 4.5 K, followed by a decrease at the lowest temperatures. This is a characteristic pattern of a ferromagnetically coupled dimer. The exchange interactions, responsible for that behavior, can be described by the classical Heisenberg-Dirac-VanVleck Hamiltonian

$$\hat{\mathbf{H}} = J\hat{\mathbf{S}}_1\hat{\mathbf{S}}_2 \tag{1}$$

where $\hat{\mathbf{S}}_1$ and $\hat{\mathbf{S}}_2$ are the spin operators of the interacting ions, and I is the exchange integral, negative for the ferromagnetic interactions in our notation. The exchange interactions lead to the spin triplet state (S = 1) and spin singlet state (S = 0). The energy difference between these states equals J.

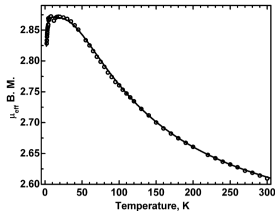


Fig. 10 The effective magnetic moment of 3 as a function of temperature. Dots: experimental, solid line – calculated with $J = -100 \text{ cm}^{-1}$ (see the text)

The magnetic susceptibility per mole of a dimeric copper(II) complex can be calculated from

$$\chi_{\rm d} = \frac{N\mu_{\rm B}^2 g^2}{3kT} \frac{6 \exp(-J/kT)}{1 + 3 \exp(-J/kT)}$$
 (2)

However, this equation cannot reproduce the magnetic moment drop at the lowest temperatures. That drop is caused by the combined Zeeman and zero-field splitting of the triplet state being comparable to the thermal energy kT. To account for this, terms expressing those interactions must be added to the HDVV Hamiltonian:

$$\hat{\mathbf{H}} = J\hat{\mathbf{S}}_{1}\hat{\mathbf{S}}_{2} + D_{12}\{\hat{\mathbf{S}}_{z1}\hat{\mathbf{S}}_{z2} - \hat{\mathbf{S}}_{1}\cdot\hat{\mathbf{S}}_{2}/3\} + E_{12}(\hat{\mathbf{S}}_{x1}\hat{\mathbf{S}}_{x2} - \hat{\mathbf{S}}_{y1}\hat{\mathbf{S}}_{y2}) + \mu_{\mathrm{B}}B\{g_{1}\}\hat{\mathbf{S}}_{1} + \mu_{\mathrm{B}}B\{g_{2}\}\hat{\mathbf{S}}_{2}$$
(3)

The spin Hamiltonian matrix was diagonalised to find the energy levels and the magnetic susceptibility per mole of the dimer was calculated from

$$\chi_{\rm d} = -\frac{N}{B} \frac{\sum_{i} \frac{\partial E_{i}}{\partial B} \exp(-E_{i}/kT)}{\sum_{i} \exp(-E_{i}/kT)}$$
(4)

The derivatives $\partial E_i/\partial B$ can be evaluated numerically by calculating the energy levels slightly below and slightly above $(\pm 5 \text{ Gauss})$ the operational magnetic field of a SQUID magnetometer (5000 G in our case). The D and E parameters have been determined from the EPR spectra (see below) and were not allowed to vary in the fitting of the magnetic data. The complex is strongly hygroscopic resulting in some uncertainty of the molar mass at the time of the magnetic measurement. To account for the observed slightly too low magnetic susceptibility, the g values were allowed to vary, but the ratio of g_z to g_x and g_{ν} was fixed as observed in EPR. The fitting procedure resulted in J = -100 cm⁻¹, representing relatively strong ferromagnetic interactions. This I value is slightly greater than that observed for an analogous dimer derived from benzoic acid.35 It is known since 197638 that in bis(hydroxo) and bis(alkoxo) bridged copper(II) complexes a relationship exists

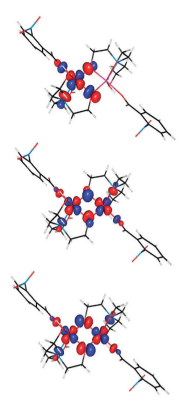


Fig. 11 Top: the "magnetic orbital" of one of the copper ions in 3. An orbital of the same shape is located also on the other copper ion. Center and bottom: the symmetric and antisymmetric combinations of the magnetic orbitals of two copper ions. The energy of the symmetric combination is lower by 2320 cm⁻¹ than that of the antisymmetric combination. The overlap integral between the magnetic orbitals is 0.076. $E(\text{High-Spin}) - E(\text{BrokenSym}) = -88.68 \text{ cm}^{-1}$.

between structural parameters, in particular the Cu-O-Cu bond angle, and the J value. In 3, the obtained $J = -100 \text{ cm}^{-1}$ is coupled with a Cu-O-Cu angle of 97.5(1)°, in line with what was found for similar dimeric complexes.39

In order to estimate the magnitude of the isotropic exchange integral I in eqn (1)–(3), a DFT calculation has been performed using the procedure described in the previous section. Correct sign of J was found while its magnitude was overestimated $(-176 \text{ cm}^{-1}).$

The interactions which contribute to ferromagnetism of dinuclear complexes involve pairs of overlapping 'magnetic orbitals' localized on each metal ion. 40 Representative orbitals are plotted in Fig. 11.

As for EPR spectra, a spin-triplet (S = 1) spectrum was seen at all temperatures over the range 3-300 K. Spectra were of the best quality at moderately low temperatures of about 200 K. Because of the large energy difference between the triplet and singlet states, the spectra could be interpreted in terms of the coupled-spin Hamiltonian with S = 1 (instead of using eqn (3)):

$$\hat{H} = \mu_{\rm B} B\{g\} \hat{\mathbf{S}} + D\{\hat{\mathbf{S}}_z^2 - S(S+1)/3\} + E(\hat{\mathbf{S}}_x^2 - \hat{\mathbf{S}}_y^2)$$
 (5)

Interestingly, the zero-field splitting tensor and the g tensor were found to be non-coaxial. Spectra at all frequencies (Fig. 12)

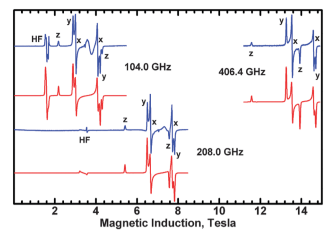


Fig. 12 High-frequency EPR spectra recorded at 200 K with the microwave frequencies as indicated. The blue and red traces are the experimental and simulated spectra, respectively. Labels x, y and z designate the magnetic field orientation at which the respective transitions occur. HF is the so-called half-field transition ($\Delta M_S = 2$). A feature between the X resonances in the 104 GHz spectrum is due to monomeric Cu²⁺ contamination. The spin Hamiltonian parameters are g_x = 2.060, g_y = 2.060, g_z = 2.303, D = -1.135 cm⁻¹, E = -0.0430 cm⁻¹.

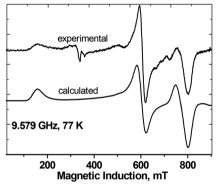


Fig. 1.3 The X-band EPR spectra of powdered complex 3 at 77 K experimental and calculated with the parameters given in the text.

were successfully simulated assuming that the zfs tensor is rotated versus the g tensor by 11 degrees about the X axis. At 200 K, the following parameters were found: $g_x = 2.060$, $g_y =$ 2.060, $g_z = 2.303$, D = -1.135 cm⁻¹, E = -0.0430 cm⁻¹. The parameters derived from the X-band spectra (Fig. 13) at 77 K are very similar: $g_x = 2.06$, $g_y = 2.06$, $g_z = 2.30$, |D| = 1.125 cm⁻¹, |E| =0.040 cm⁻¹, with signals due to the monomeric impurities present at 300 mT. The spin Hamiltonian parameters are temperaturedependent and at 10 K (Fig. 14) they are $g_x = 2.056$, $g_y = 2.045$, $g_z =$ 2.320, D = -1.042 cm⁻¹, E = -0.0075 cm⁻¹, with the same angle of rotation of the zfs and g tensors as found at 200 K (11 deg). The reason for the deviation between the axes of the g and zfs tensors observed in the EPR spectra is the misalignment between the equatorial ligand plane and the Cu-Cu direction.

Since the complex is ferromagnetic, EPR spectra could be detected at very low temperatures allowing determination of the sign of D and E^{41} . This is possible thanks to the Zeeman splitting being comparable to the thermal energy, kT (Fig. 15).

Fig. 14 Determination of the sign of the zero-field splitting parameters from the high-frequency EPR spectra recorded at 10 K with ν = 406.4 GHz. The sign of D and E does not affect the positions of the resonance lines, but it does affect the intensity pattern at low temperatures and high frequencies. The spin Hamiltonian parameters at 10 K are: g_x = 2.056, g_y = 2.045, g_z = 2.320, D = -1.042 cm⁻¹, E = -0.0075 cm⁻¹.

Magnetic Induction, Tesla

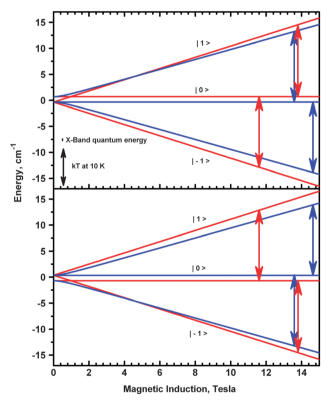


Fig. 15 Energy levels of the triplet state calculated with D < 0 (top) and with D > 0 (bottom). The red and blue lines represent the Z and X molecular orientation, respectively. The red and blue arrows have the length of the 406.4 GHz microwave quantum energy (13.55 cm $^{-1}$). With D negative, the high-field transition at the Z orientation (red arrow at 13.94 T) starts from an excited level and its intensity will be suppressed at low temperatures, while the low-field transition intensity (11.57 T) will be enhanced. An opposite relationship holds for the transitions at the X orientation (blue arrows). If D is positive, the energy diagram is reversed and all intensity relationships change to the opposite, while all resonance fields remain unchanged. Comparison of the X-band microwave quantum energy (0.32 cm $^{-1}$) to the thermal energy kT at 10 K (6.95 cm $^{-1}$) indicates that no such effects can be observed in X-Band EPR.

In copper(II) dimers, there are two contributions to the zerofield splitting parameters D and E. One is the magnetic dipoledipole interaction, while the other one is associated with the anisotropy of the exchange interactions. These contributions cannot be determined separately in an experiment. The dipolar part depends on the Cu-Cu separation. Its largest component is oriented along the Cu-Cu direction, while the largest component of the exchange-related zfs is expected to be parallel to the g_{z} direction. The latter should be at least approximately perpendicular to the plane of the equatorial ligands. The angle between the Cu-Cu vector and the normal to the equatorial ligand plane is 86 deg. We assume for simplicity that this angle equals 90 deg and define a system of coordinates in which the axis X is along Cu-Cu, and the axis Z is perpendicular to Cu-Cu and is tilted 4 deg from the normal to the plane defined by the equatorial ligands. In these coordinates one can write formulae for the dipole–dipole related D_{dip} and E_{dip} parameters using the point-dipole approximation:

$$D_{\text{dip}} = (2g_z^2 + 2g_x^2 - g_y^2)\mu_B^2/4R_{\text{Cu-Cu}}^3$$

$$E_{\text{dip}} = -(2g_x^2 + g_y^2)\mu_B^2/4R_{\text{Cu-Cu}}^3$$
 (6)

 $D_{
m dip}$ and $E_{
m dip}$ above are pertinent to spin Hamiltonian (5). (Note that they need to be multiplied by 2 to be used in eqn (3).) With $R_{
m Cu-Cu}=2.92$ Å and our g values we obtain $D_{
m dip}=0.0646$ cm $^{-1}$ and $E_{
m dip}=-0.0553$ cm $^{-1}$. These values should be considered as the upper limits, because the electron delocalization may considerably reduce them. 42

The experimentally determined *D* and *E* parameters are the sum of the dipole–dipole and exchange-related parts, therefore

$$D_{\text{ex}} = D - D_{\text{dip}} = -1.20 \text{ cm}^{-1}$$

 $E_{\text{ex}} = E - E_{\text{dip}} = 0.007 \text{ cm}^{-1}$

The exchange-related zero-field splitting thus appears to be remarkably axial ($|E| \ll |D|$). Maurice *et al.* derived formulae for $D_{\rm ex}$ and $E_{\rm ex}$ for two coupled ${\rm Cu}^{2^+}$ ions with the ${\rm d}_{x^2-y^2}$ ground state orbital. 42

$$D = 2\frac{\xi^{2}J_{x^{2}-y^{2},xy}}{\Delta E_{x^{2}-y^{2},xy}^{2}} - \frac{1}{4}\frac{\xi^{2}J_{x^{2}-y^{2},xz}}{\Delta E_{x^{2}-y^{2},xz}^{2}} - \frac{1}{4}\frac{\xi^{2}J_{x^{2}-y^{2},yz}}{\Delta E_{x^{2}-y^{2},yz}^{2}}$$

$$E = \frac{1}{4}\frac{\xi^{2}J_{x^{2}-y^{2},xz}}{\Delta E_{x^{2}-y^{2},xz}^{2}} - \frac{1}{4}\frac{\xi^{2}J_{x^{2}-y^{2},yz}}{\Delta E_{x^{2}-y^{2},yz}^{2}}$$
(7)

 $\xi = -828 \,\mathrm{cm}^{-1}$ is the spin-orbit coupling constant for Cu^{2^+} and symbols like $J_{x^2-y^2,xy}$ are the exchange integrals between the $\mathrm{d}_{x^2-y^2}$ ground orbital of one copper ion and the excited d_{xy} orbital of another. The d_{xz} and d_{yz} orbitals of one copper are oriented similarly *versus* the $\mathrm{d}_{x^2-y^2}$ orbital of another copper atom. Accordingly, the terms in equations above involving $J_{x^2-y^2,xz}$ and $J_{x^2-y^2,yz}$ are expected to be of similar magnitude resulting in a small or equal to zero E_{ex} parameter. With a different arrangement of the bridging group and a d_{z^2} ground state of copper, very large E values were observed.⁴³

The quantities $\xi/\Delta E_{x^2-y^2,xy}$ *etc.* in eqn (7) appear also in the theory of the g tensor in Cu(II). For example, $\Delta g_z = g_z - 2.0023 = 8\xi/\Delta E_{x^2-y^2,xy}$ and it is possible to eliminate the unknown ΔE values from eqn (7):

$$D_{\text{ex}} = \left[J_{x^2 - y^2, xy} \Delta g_z^2 - 2J_{x^2 - y^2, xz} \Delta g_y^2 - 2J_{x^2 - y^2, yz} \Delta g_x^2 \right] / 32$$
(7a)

Also, $J_{x^2-y^2,xy}$ is expected to be the largest of the three $J_{x^2-y^2,n}$ exchange integrals. When the two remaining ones are neglected, one can estimate $J_{x^2-y^2,xy} = 32D_{\rm ex}/\Delta g_z^2 = -425~{\rm cm}^{-1}$. That exchange interaction is thus ferromagnetic, as expected, since the overlap integral of the copper orbitals is zero (a lobe of the d_{xy} orbital of one copper atom points between the positive and negative lobes of the $d_{x^2-y^2}$ orbital of the other copper atom). $J_{x^2-y^2,xy}$ of the same order of magnitude were observed in copper dimers with similar bridging arrangement.⁴⁴

Conclusions

Paper

Three novel complexes were isolated by addition of γ -picoline, β-picoline and triethanolamine to the hydrated copper(II) 2-nitrobenzoate, [Cu₂(2-nitrobenzoate)₄(H₂O)₂]·2H₂O suspended in a 4:1 methanol-water mixture. It has been found that the addition of different types of N-donor ligands (rigid and flexible) has a profound influence on the orientation of -NO2 and -COOmoieties in these complexes and, above all, on the coordination geometry around the Cu atom. Detailed structural analysis of 1, 2 and 3 confirms that the addition of different N-donor ligands to copper(II) carboxylates may affect their physico-chemical behaviour such as magnetism. The dimeric complex 3 exhibits relatively strong ferromagnetic exchange with $J = -100 \text{ cm}^{-1}$ (corresponding to $H = IS_1S_2$). The zero-field splitting parameters (zfs) of the dimer triplet state D = -1.135 cm⁻¹, E = -0.043 cm⁻¹ were derived from the HF EPR spectrum recorded at moderately low temperatures. The sign of D was determined from lowtemperature HF EPR spectra.

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