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This is a pre print version of the following article:

Original Citation:

Availability:

This version is available <http://hdl.handle.net/2318/1770025> since 2022-01-19T12:43:37Z

Published version:

DOI:10.1039/D0CE00370K

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Investigation on the Interconversion from DMF Solvated to Unsolvated Copper(II) Pyrazolate Coordination Polymers

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Abstract

A series of copper(II) pyrazolates were synthesized employing different synthetic approaches. By refluxing either $[\text{Cu}_3(\mu_3\text{-OH})(\mu\text{-pz})_3(\text{CH}_3\text{COO})_2(\text{pzH})]$ or $[\text{Cu}(\mu\text{-pz})_2\cdot\text{H}_2\text{O}]_n$ in water, $[\text{Cu}(\mu\text{-pz})(\mu\text{-OH})]_n$ (**1**) was unexpectedly obtained. Whereas using microwave-assisted solvothermal synthesis, the compounds $[\text{Cu}(\mu\text{-4-Xpz})_2(\mu\text{-DMF})]_n$ (**2_{DMF}**-**5_{DMF}**) (4-Xpz = 4-Xpyrazolate, X = H, Cl, Br, I) were isolated. As unveiled by powder X-ray diffraction, all the compounds are 1-D coordination polymers showing square planar *trans*- CuN_2O_2 (**1**) or octahedral *trans*- CuN_4O_2 (**2_{DMF}**-**5_{DMF}**) metal centers, reasonably involved in through-bond magnetic interactions (magnetic susceptibility in the range 0.91-1.56 B.M.). Upon heating under inert atmosphere, **2_{DMF}**-**5_{DMF}** lose DMF to give the corresponding low-crystallinity $[\text{Cu}(\mu\text{-4-Xpz})_2]_n$ coordination polymers **2-5**, of which only **3** was characterized in the past. In order to shed light on the local-level structural features of **2**, **4** and **5**, UV-Vis-NIR absorption and fluorescence emission spectroscopy measurements were performed, strongly suggesting a tetrahedral stereochemistry for the metal centres.

Keywords Cu(II) pyrazolates, coordination polymers, microwave-assisted solvothermal synthesis, powder X-ray diffraction, electronic-state transition spectroscopy.

1. Introduction

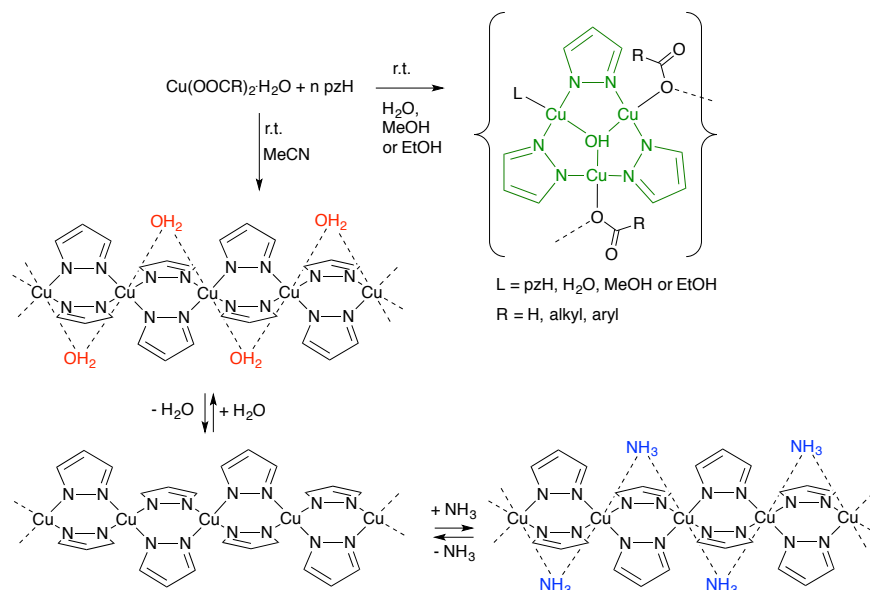
Even though about thirty years have passed from the intuition of Robson and Hoskins¹ that initiated the coordination polymers (CPs) age, the interest towards these compounds is far from being exhausted. Coordination polymers²⁻⁵ are inorganic-organic materials showing not only fascinating structural features, but also interesting functional properties and potential applications in relevant fields, such as luminescence,⁶⁻⁹ heterogeneous catalysis,¹⁰ magnetism,¹¹⁻¹⁴ conductivity,¹⁵ nanoelectronics,¹⁶ to quote a few.

In the crystal structure of these compounds, rigid or flexible polytopic ligands, behaving as spacers, bridge metal ions or oxo-metallic clusters, behaving as nodes. The organic spacers that have been commonly employed in the construction of first-generation CPs are polytopic carboxylates,^{17,18} pyrazines and bipyridines.¹⁹ To date, a great number of CPs containing other nitrogen-donor ligands, such as poly(azolates) (pyrazolates, imidazolates, triazolates and tetrazolates) have been reported.²⁰⁻²⁴

In the past years, we have deeply investigated the interactions of 3d transition metal ions with pyrazole (pzH) and monocarboxylates, synthesizing mono- and polynuclear complexes as well as CPs. Besides zinc(II), cadmium(II) and mercury(II) carboxylates,^{25,26} our interest has been mainly addressed to investigate how the reaction conditions between copper(II) carboxylates and pzH can influence the composition as well as the structural and functional properties of the final products. Noteworthy, we observed a different behaviour if the reactions among Cu(II) carboxylates and pzH were carried out in protic solvents (H₂O, alcohols) or in MeCN. In the first case, compounds based on the trinuclear triangular moiety [Cu₃(μ₃-OH)(μ-pz)₃]²⁺, the charge of which is balanced by two carboxylate ions coordinated to the metal centre, were obtained (Scheme 1). These trinuclear complexes self-assemble in hexanuclear clusters through carboxylate bridges and form extended 1-D, 2-D or 3-D CPs or supramolecular networks, displaying intriguing catalytic activity in the cycloalkane oxidation to alcohols.^{27-40,41} Completely different results were obtained when Cu(II) carboxylates were reacted with pzH in MeCN, yielding the pale-pink 1-D CP [Cu(μ-pz)₂·H₂O]_n independently from the carboxylate employed (Scheme 1).⁴²⁻⁴⁴ This compound reversibly loses water, yielding [Cu(μ-pz)₂]_n. Besides water, [Cu(μ-pz)₂]_n is able to reversibly bind other small molecules such as NH₃, changing its colour from beige to deep blue⁴² (Scheme 1).

Based on the above mentioned results, we decided to expand the investigation on novel coordination compounds having the {Cu₃(μ₃-OH)(μ-pz)₃}²⁺ cluster or {Cu(μ-pz)₂} fragment as secondary building unit (SBUs). Herein we report the results obtained upon refluxing either [Cu₃(μ₃-OH)(μ-pz)₃(CH₃COO)₂(pzH)] or [Cu(μ-pz)₂·H₂O]_n in water, and upon reacting copper(II) acetate hydrate and 4-Xpyrazole, (X = H, Cl, Br, I) along solvothermal, microwave-assisted

syntheses, using dimethylformamide (DMF) as solvent. The isolated compounds were characterized as *per* their crystal structure, thermal behaviour, under N₂ and in air, and electronic-state spectroscopic properties.



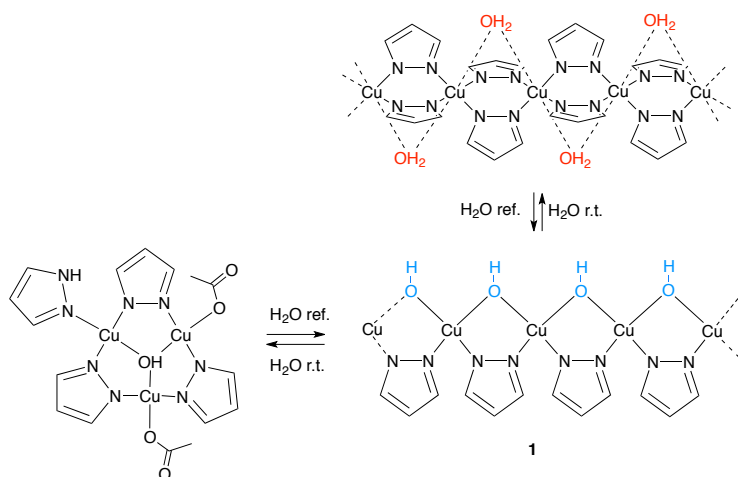
Scheme 1. Chemistry and reactivity of Cu(II) pyrazolate coordination compounds.

2. Results and Discussion

2.1. Synthesis and preliminary characterization

By refluxing either $[\text{Cu}_3(\mu_3\text{-OH})(\mu\text{-pz})_3(\text{CH}_3\text{COO})_2(\text{pzH})]$ or $[\text{Cu}(\mu\text{-pz})_2 \cdot \text{H}_2\text{O}]_n$ in water, compound $[\text{Cu}(\mu\text{-pz})(\mu\text{-OH})]_n$ (**1**) was obtained (Scheme 2) as a blue-purple microcrystalline powder insoluble in common solvents. This compound was previously isolated through a different procedure, but not completely characterized, by Mezei *et al.*^{45,46} Interestingly, the conversion of $[\text{Cu}_3(\mu_3\text{-OH})(\mu\text{-pz})_3(\text{CH}_3\text{COO})_2(\text{pzH})]$ or $[\text{Cu}(\mu\text{-pz})_2 \cdot \text{H}_2\text{O}]_n$ into **1** is reversible, the starting compounds being slowly reformed by cooling down the reaction mixtures and leaving them under stirring at room temperature for 12 hours (Scheme 2; IR evidences).

On the other hand, by reacting copper(II) acetate hydrate and pzH in DMF along a microwave-assisted solvothermal synthesis, a microcrystalline blue-violet powder was precipitated and characterized as $[\text{Cu}(\mu\text{-pz})_2(\mu\text{-DMF})]_n$, **2_{DMF}**, (Scheme 3). Similarly, copper(II) acetate hydrate and the substituted pyrazoles 4-XpzH (X = Cl, Br, I) afforded $[\text{Cu}(\mu\text{-4-Xpz})_2(\mu\text{-DMF})]_n$ (**3_{DMF}**, **4_{DMF}** and **5_{DMF}**, for X = Cl, Br, I, respectively; Scheme 3).



Scheme 2. Synthesis of compound 1 by using two different reagents.

It is important to underline that all the attempted reactions performed through conventional methods at room temperature or under reflux gave impure products difficult to separate and purify. By heating compounds 2_{DMF} - 5_{DMF} at 453 K for two hours, DMF could be completely removed (IR spectroscopy evidences; see below), yielding the corresponding non-solvated forms (**2-5**, respectively), showing a different coordination at the metal ion (see the sections dedicated to the crystal structure description and the spectroscopic studies), hence a different colour (Figure 1).

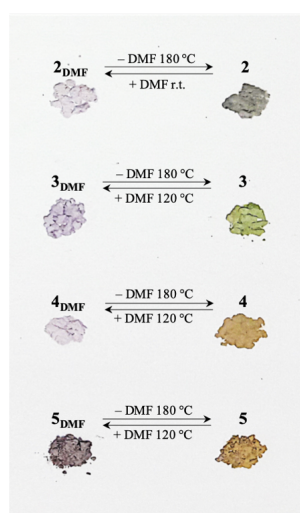
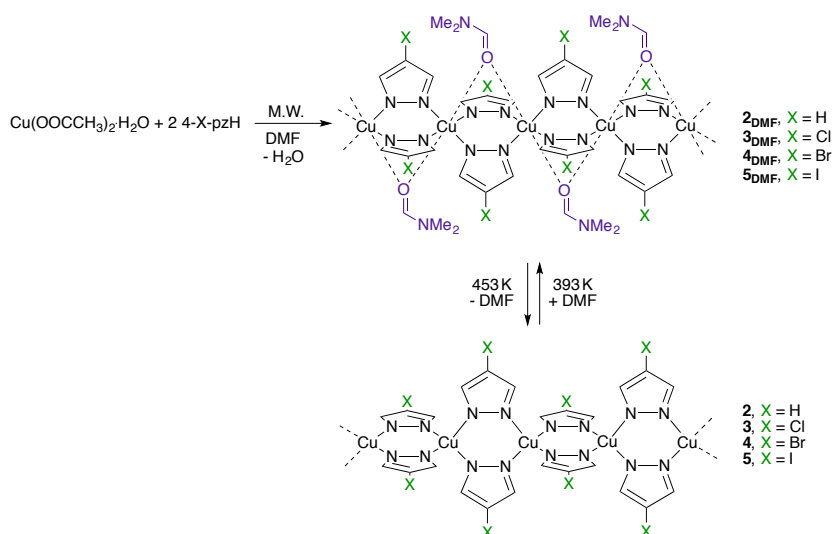


Figure 1. Colours of the solvated compounds 2_{DMF} - 5_{DMF} and the non-solvated compounds **2-5**. The colour change is induced by the (loss of) coordination of the DMF molecules to the copper(II) ions.

The non-solvated forms can reacquire the original colour (and composition; IR evidences) in a few minutes by suspending them in DMF at r.t. (**2**) or at 393 K (**3-5**) (Scheme 3).



Scheme 3. Synthesis of compounds **2**_{DMF}-**5**_{DMF} and **2-5**.

The IR spectrum of **1** (Figure S1) shows the O-H bond stretching as a medium-intensity band at 3602 cm^{-1} , the signal sharpness highlighting the absence of hydrogen-bond interactions, in agreement with what observed in the crystal structure (see section 2.2). The IR spectra of **2**_{DMF}-**5**_{DMF} and **2-5** (Figure 2 for **3**_{DMF} and **3**; Figures S2-S4 for **2**_{DMF} and **2**, **4**_{DMF} and **4**, **5**_{DMF} and **5**) exhibit similar patterns, in agreement with their comparable structural features (see section 2.2). The presence of coordinated DMF is confirmed, in all cases, by an intense band in the range $1651\text{-}1655\text{ cm}^{-1}$ due to the C=O stretching, shifted downfield with respect to free DMF (1661 cm^{-1}).^{REF} The $\nu(\text{C-Cl})$ stretching mode in the IR spectra of **3**_{DMF} and **3** is found at 969 and 972 cm^{-1} (Figure 2), respectively, the small shift being possibly related to the absence of DMF in **3**. As expected, in the IR spectra of **4**_{DMF}, **5**_{DMF}, **4** and **5** (Figures S3, S4) the absorption bands due to the $\nu(\text{C-halide})$ stretching modes (halide = Br or I) appear at lower frequencies (955 cm^{-1} for C-Br, 942 cm^{-1} for C-I) due to the increase of the halogen atomic mass.

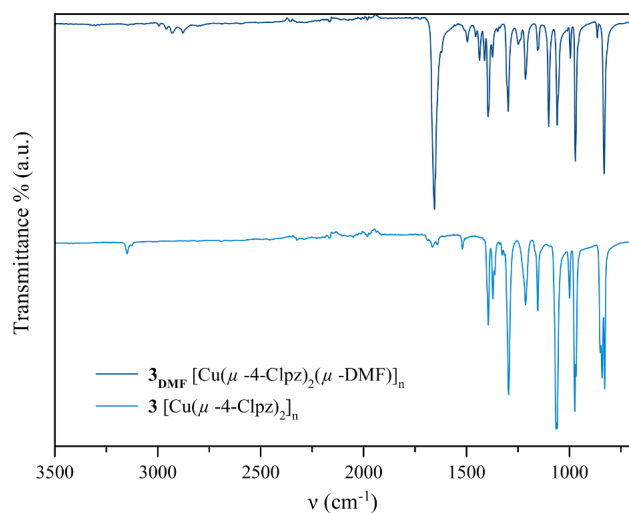


Figure 2. IR spectra of $[\text{Cu}(\mu\text{-4-Clpz})_2(\mu\text{-DMF})]_n$ ($\mathbf{3}_{\text{DMF}}$, blue) and $[\text{Cu}(\mu\text{-4-Clpz})_2]_n$ ($\mathbf{3}$, light blue).

2.2. Crystal Structure Analysis

Compound **1** crystallizes in the orthorhombic space group *Pbca*. The asymmetric unit comprises one copper(II) ion, one *pz'* ligand and one OH⁻ group, all in general positions. As assumed by Mezei and co-workers,⁴⁵ the metal centres show the *trans*- CuN_2O_2 square planar stereochemistry, defined by the nitrogen atoms of two *pz'* spacers and the oxygen atoms of two hydroxyl groups (Figure 3a; see the caption to the Figure for the main bond distances and angles at the metal ion). Both anions are exo-bidentate and define 1-D helices of collinear metal ions running along the [010] crystallographic direction (Figure 3b) with a $\text{Cu}\cdots\text{Cu}$ distance of 3.386(3) Å (half of the *b*-axis). The chains pack, staggered, along the [001] crystallographic direction defining a herringbone motif (Figure 3c). No hydrogen bond interactions are present, as witnessed by IR spectroscopy. The value of the magnetic susceptibility (0.91 B.M.), lower than expected for spin-only Cu(II) ions (1.73 B.M.), suggests the presence of through-bond interactions among the nearby metal centres along the helices.

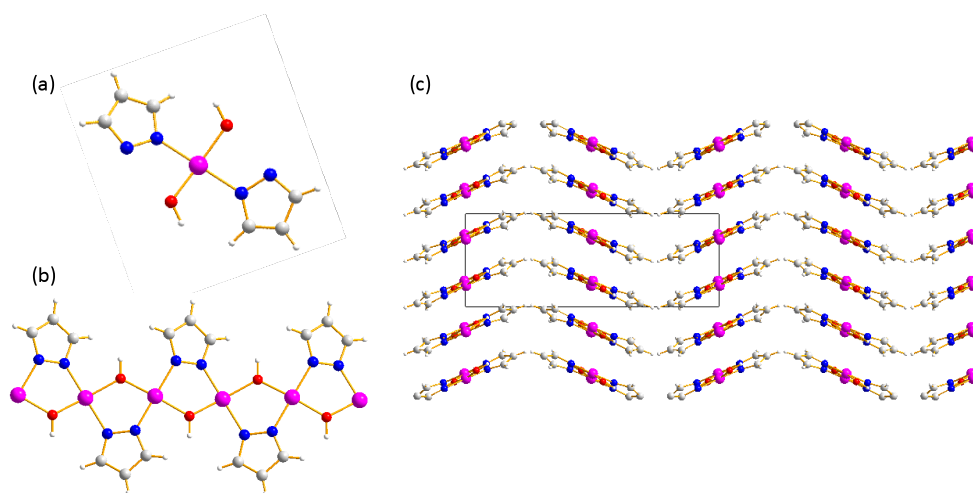


Figure 3. (a) Representation of the *trans*-CuN₂O₂ square planar stereochemistry of the copper(II) ion in **1**. (b) Representation of portion of the 1-D helix of collinear metal ions running along the [010] crystallographic direction in **1**. (c) Representation of portion of the crystal packing of **1** viewed along the [010] crystallographic direction. Horizontal axis, *a*; vertical axis *c*. Colour code: Carbon, grey; hydrogen, light grey; copper, fuchsia; nitrogen, blue; oxygen, red. Main bond distances (Å) and angles (°) at the metal ion: Cu-N 1.983(16), 2.0469(94); Cu-O 1.8831(86), 1.9563(88); intra-chain Cu...Cu 3.386(3); N-Cu-N 178.94(55); O-Cu-O 168.86(18); N-Cu-O, 87.88(39), 92.93(41).

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Compounds **2**_{DMF}-**4**_{DMF} crystallize in the orthorhombic space group *Pnam* and are isomorphous. In light of this, their structural features will be described jointly in the following. The asymmetric unit contains half of a copper(II) ion (on an inversion centre, *b* Wyckoff letter), two halves of two ligands and half of a DMF molecule (on *.m* positions, *c* Wyckoff letter). The metal ions show a *trans*-N₄O₂ octahedral stereochemistry, defined by the nitrogen atoms of four pyrazolate ligands and the oxygen atoms of two solvent molecules (Figures 4a, S5a and S6a; see the captions to the Figures for the main bond distances and angles at the metal ion). In all the compounds, a Jahn-Teller type distortion elongates the Cu-O bond distances [2.656(7)-2.780(13) Å] with respect to the Cu-N ones [1.782(9)-1.999(6) Å]. The two independent pz⁻ or 4-Xpz⁻ spacers (X = Cl, Br) and the DMF molecules are exo-bidentate, bridging the metal ions along 1-D helices running along the [001] crystallographic direction with a pitch amounting to the value of the *c*-axis (Figures 4b, S5b and S6b). The observed values of the magnetic susceptibility (1.31-1.56 B.M.), which are lower than expected for spin-only Cu(II) ions, are consistent with the formation of coordination polymers showing through-bond interactions among the neighbouring metal centres. The elongation direction of the helices is consistent with the observed preferred orientation pole ([110]), as inter-helix non-bonding interactions are at work in the *ab*-plane. Indeed, the samples of

2_{DMF} are characterized by thin needles (Figure S7). While in 3_{DMF} and 4_{DMF} the copper(II) ion is almost coplanar with the two independent ligands [distance among copper(II) and the r.m.s. planes defined by the penta-atomic rings: 0 and 0.25 Å in 3_{DMF} , 0.02 and 0.25 Å in 4_{DMF}], in 2_{DMF} the metal centre is almost coplanar with only one of the two spacers [distance among copper and the r.m.s. planes defined by the penta-atomic rings 0.13 and 0.86 Å]. The reciprocal disposition of the helices in the ab -plane defines a rhombic motif (Figures 4c, S5c and S6c) with an edge amounting to half of the diagonal of the C -face of the unit cell. No hydrogen bond interactions are present.

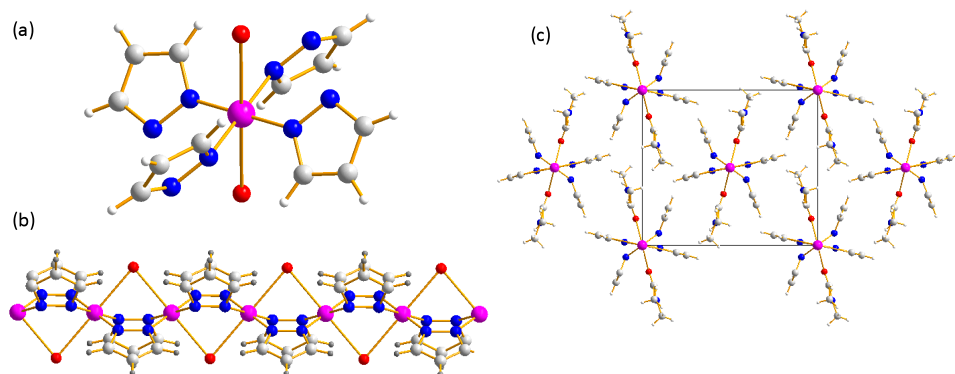


Figure 4. (a) Representation of the *trans*- CuN_4O_2 octahedral stereochemistry of the copper(II) ion in 2_{DMF} . (b) Representation of portion of the 1-D helix of collinear metal ions running along the [001] crystallographic direction in 2_{DMF} . (c) Representation of portion of the crystal packing of 2_{DMF} viewed along the [001] crystallographic direction. Horizontal axis, b ; vertical axis a . Colour code: Carbon, grey; hydrogen, light grey; copper, fuchsia; nitrogen, blue; oxygen, red. Main bond distances and angles at the metal ion: Cu-N 1.782(9), 1.927(14); Cu-O 2.780(13); intra-chain $\text{Cu}\cdots\text{Cu}$ 3.5500(2); N-Cu-N 83.45(62), 96.55(62), 180; O-Cu-O 180; N-Cu-O 81.63(45)-98.37(45). For the representation of the crystal structures of 3_{DMF} and 4_{DMF} the reader is referred to Figures S5 and S6.

As expected, upon increasing the van der Waals radius of the atom in position 4 of the pyrazolate ring, the a - and b -axis (hence the diagonal of the C -face) and the unit cell volume increase proportionally (Table S1 and Figure S8).

As synthesized 5_{DMF} (**m-5_{DMF}**) is characterized by a PXRD pattern which can be indexed by a monoclinic P unit cell [space group $P112_1/m$,^[1] $a = 11.1475(5)$ Å, $b = 9.5458(4)$ Å, $c = 7.3608(2)$ Å, $\gamma = 114.5194(19)^\circ$, $V = 712.64(5)$ Å³] describing all the observed Bragg reflections (Figure S9). The unit cell volume is compatible with the formula unit $[\text{Cu}(\mu\text{-4-Ipz})_2(\mu\text{-DMF})]_n$. As proved by PXRD monitoring (Figure S10), this monoclinic phase progressively transforms into an orthorhombic $Cmcm$ phase, **oC-5_{DMF}** ($Cmcm$ is a proper super-group of $P112_1/m$), which is stable indefinitely. The transformation is completed in about 3.5 hours. This occurrence prevented the structural characterization of the monoclinic phase. In the crystal structure of **oC-5_{DMF}**, the

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asymmetric unit comprises one quarter of the metal ion (on a $2/m$ position, b Wyckoff letter), half of a ligand (on a $..m$ position, g Wyckoff letter) and one quarter of a DMF molecule (on a $m2m$ position, c Wyckoff letter). As in compounds **2**_{DMF}-**4**_{DMF}, also in **5**_{DMF} the metal ion shows the *trans*-CuN₄O₂ octahedral coordination geometry, defined by the nitrogen atoms of four ligands and the oxygen atom of two solvent molecules (Figure 5a; the Figure caption provides the main bond distances and angles at the metal ion). A Jahn-Teller type distortion involving the Cu-O bonds [2.483(2) Å vs. 1.924(1) Å of the Cu-N bonds] is present. The metal ion is out of the r.m.s. plane defining the penta-atomic ring by 0.35 Å. μ -bridging by the ligands and the DMF molecules likewise brings about the formation of 1-D helices running along the 2_1 screw axes parallel to the [001] crystallographic direction (Figure 5b), with a Cu··Cu distance of 3.6793(1) Å (half of the value of the c -axis). Similarly, the elongation direction of the 1-D polymeric motifs is consistent with the observed preferred orientation pole ([100]). The helices pack, eclipsed, forming a rhombic motif, with diagonals defined by the a - and b -axis (Figure 5c).

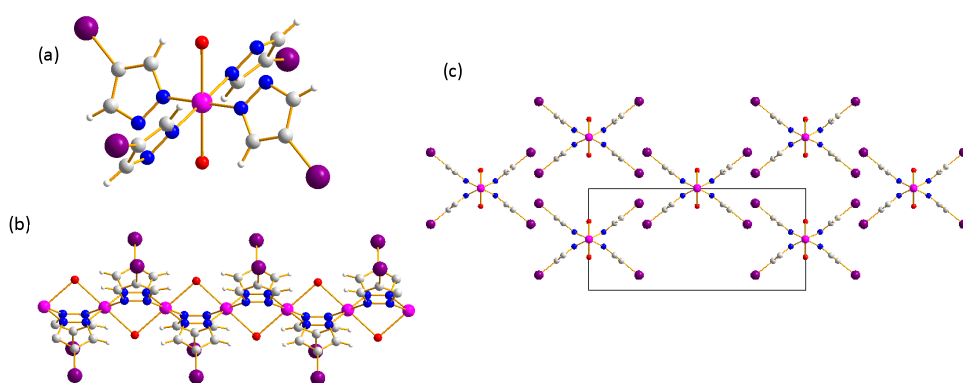


Figure 5. (a) Representation of the *trans*-CuN₄O₂ octahedral stereochemistry of the copper(II) ion in **5**_{DMF}. (b) Representation of portion of the 1-D helix of collinear metal ions running along the [001] crystallographic direction in **5**_{DMF}. (c) Representation of portion of the crystal packing of **5**_{DMF} viewed along the [001] crystallographic direction. Horizontal axis, a ; vertical axis b . For clarity, only the oxygen atom of DMF is represented. Colour code: Carbon, grey; hydrogen, light grey; copper, fuchsia; iodine, violet; nitrogen, blue; oxygen, red. Main bond distances (Å) and angles (°) at the metal ion: Cu-N 1.924(1); Cu-O 2.483(2); intra-chain Cu··Cu 3.6793(1); N-Cu-N 180; O-Cu-O 180; N-Cu-O 77.97(34)-102.03(34).

In **5**_{DMF}, neither the edges of the rhombi nor the unit cell volume (Table S1) follow the same trend of compounds **2**_{DMF}-**4**_{DMF} vs. the van der Waals radii of the atom in position 4 of the pyrazolate ring. Indeed, the atomic packing factor of **5**_{DMF} is comparable to that of **3**_{DMF} (Table S1).

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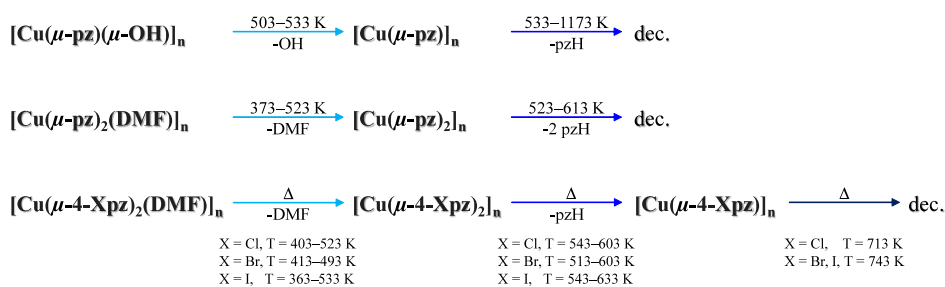
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2.3. Thermal behaviour

The thermal behaviour of compounds **1**, **2**_{DMF}-**5**_{DMF} was investigated by thermogravimetric analysis (TGA) and variable-temperature PXRD. According to the TGA trace (Figure 6a, Scheme 4), under nitrogen **1** is stable up to 503 K. The weight loss of 14.0% in the temperature range 503–533 K can be tentatively ascribed to the loss of one hydroxyl group *per* formula unit (calculated weight 11.0%), leading to the formation of [Cu(μ -pz)]_n. Decomposition starts immediately after this event. At 1173 K, the residual mass of 40.4% can be ascribed to metallic copper (calculated residual mass 43.0%). The VT-PXRD experiment carried out in air (Figure 6b) highlights that **1** decomposes, without prior transformations, in the range ~563–583 K.

According to its TGA trace (Figure S11a, Scheme 4), compound **2**_{DMF} first undergoes a weight loss of 26.2%, which can be interpreted as the loss of one mole of DMF *per* formula unit (calculated weight loss 26.9%), to yield the non-solvated form **2**. After this event, starting from 523 K, a weight loss of 51.0% is observed, corresponding to the loss of two ligands *per* formula unit (calculated weight loss 52.2%).

As evident from their TGA traces (Figures S12a–S14a), compounds **3**_{DMF}-**5**_{DMF} overall undergo the same sequence of transformations, namely (Scheme 4): *i*) complete loss of DMF to yield the non-solvated forms **3–5**, respectively; *ii*) loss of one mole of ligand *per* formula unit, affording the corresponding [Cu(μ -4-Xpz)] (X = Cl, Br, I) copper(I) compounds; *iii*) decomposition. Details on these transformations are provided in Table 1.



Scheme 4. Thermal behaviour of compounds **1** and **2**_{DMF}-**5**_{DMF}, as observed by TGA under a flow of N₂.

Table 1. Details of the thermal behaviour of compounds **2**_{DMF}-**5**_{DMF}, as observed by TGA.

		2 _{DMF}	3 _{DMF}	4 _{DMF}	5 _{DMF}
Loss of 1 mol of solvent per f.u.	T range (K)	373-523	403-523	413-493	363-533
	Obs.(Calc.) wt. (%)	26.2(26.9)	21.6(21.5)	17.9(17.1)	14.4(14.0)
Loss of 1 mol of ligand per f.u.	T range (K)	523-613	543-603	513-603	543-633
	Obs.(Calc.) wt. (%)	51.0(52.2) ^a	31.3(30.0)	32.6(34.0)	35.6(36.9)
Decomposition	Onset T [K]	673	713	743	743
	Residual mass (%)	9.2	19.8	13.0	20.0

^a In this case, the weight loss can be ascribed to the evolution of two moles of ligand *per* formula unit.

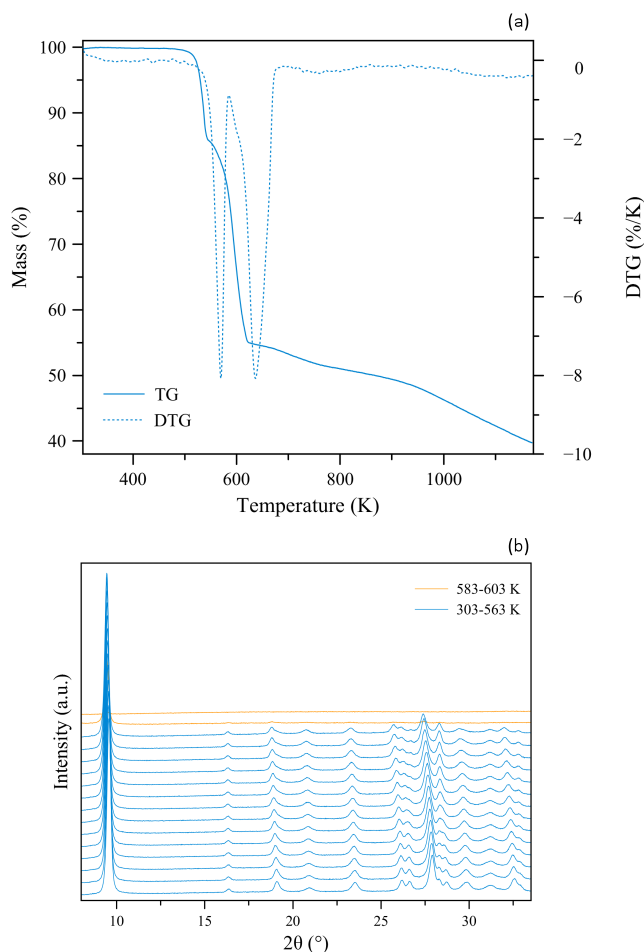
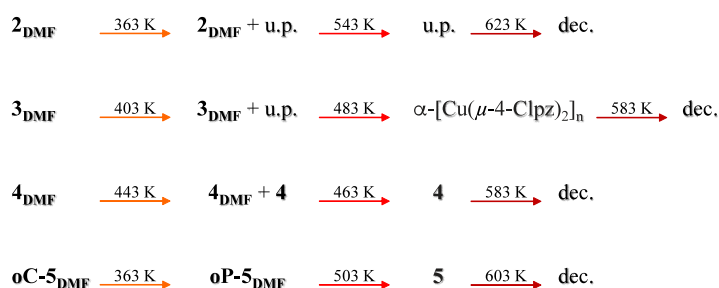


Figure 6. (a) TGA (blue continuous line) trace measured under a flow of N₂ on **1** and the corresponding DTG trace (blue dashed line). (b) Powder X-ray diffraction patterns acquired on **1** as a function of the temperature in the temperature range 303-603 K, with steps of 20 K.

The VT-PXRD experiment carried out on **2_{DMF}** (Figure S11b) shows the stability of this phase up to ~343 K. At ~363 K a new phase appears and coexists with **2_{DMF}** up to 523 K, imparting a beige colour to the sample. At ~543 K a new phase is observed, characterized by only one peak at ~8.4° (2θ), not ascribable to the already known phases α-Cu(pz)₂ or β-Cu(pz)₂⁴⁷ and disappearing at 663 K. In accordance with what disclosed by TGA, the VT-PXRD data acquired on **3_{DMF}** (Figure S12b) show that the compound is stable up to ~383 K. At ~403 K a new phase appears and coexists with **3_{DMF}** up to ~463 K, imparting a light green colour to the sample. At ~483 K the green α-[Cu(μ-4-Clpz)₂]_n polymorph⁴⁸ is formed, which starts losing crystallinity at ~583 K (vs. T_{dec} = 663 K).⁴⁸ The VT-PXRD experiment carried out on **4_{DMF}** (Figure S13b) unveils that the compound is stable up to ~423 K. At ~443 K a new low-crystallinity phase of beige colour appears which, based on the TGA, can be interpreted as the non-solvated form **4**. The latter coexists with **4_{DMF}** up to ~463 K. The powder pattern of **4** cannot be described by the unit cell parameters of the green [Cu(μ-4-Brpz)₂]_n polymorph.⁴⁸ At ~583 K, concomitant to the decomposition of **4**, CuBr₂ is formed (PXRD evidence, acquiring data in the 25.5-86.5° 2θ range). Finally, the VT-PXRD experiment carried out on the orthorhombic *C* phase **oC-5_{DMF}** (Figure S14b) highlights that it is stable up to 343 K. At ~363 K a light-purple phase appears (**oP-5_{DMF}**), which is stable up to ~483 K and is sufficiently crystalline to allow for the unit cell parameters retrieval. The obtained orthorhombic space group, *Pbnm*, is a proper subgroup of *Cmcm*. The unit cell parameters (Table S1) are comparable to those of the orthorhombic *C* phase. The colour change and symmetry decrease can be attributed to a partial loss of solvent, as witnessed by the structure determination (see below). At ~503 K **oP-5_{DMF}** greatly loses crystallinity and a new low-crystallinity, light brown phase is formed (**5**), which is stable up to ~583 K. At 603 K, a peak at ~25.1° appears, which belongs to CuI (PXRD evidence, acquiring data in the 23.5-86.5° 2θ range). A schematic representation of the behaviour of **2_{DMF}**-**5_{DMF}** as observed by VT-PXRD is shown in Scheme 5. By comparing the information provided by Schemes 4 and 5, it appears evident that all the coordination polymers show, under nitrogen, a behaviour different than in air.



Scheme 5. Thermal behaviour of compounds **2**_{DMF}-**5**_{DMF}, as observed by VT-PXRD. u.p. = uncharacterized phase.

In the crystal structure of **oP-5**_{DMF}, (Figure S15), the asymmetric unit contains half of a Cu(II) ion (on an inversion centre, Wyckoff letter *a*), the halves of two ligands and half of a DMF molecule (on *..m* positions, Wyckoff letter *c*). The mono-dimensional helices composed by the two μ - κ N: κ N'-pyrazolate ligands and the μ - κ O: κ O-DMF molecule run along the [001] crystallographic direction [Cu...Cu distance of about 3.6456(1) Å, half of the *c* axis]. These chains pack staggered, defining a rhombic motif having the *a* and *b* axes as diagonals. The apparent overlap, along the *c* axis, of different DMF molecules belonging to different chains, can be explained considering that the phase is only partially solvated, as can be seen from the value of 0.5 of the site occupation factor of DMF.

2.4. Electronic state transition spectroscopy

UV-Vis-NIR absorption and fluorescence emission spectroscopy measurements were performed in order to shed some light on the local-level structural features of the species **2**, **4** and **5**, obtained upon heating **2**_{DMF}, **4**_{DMF} and **5**_{DMF} and preliminary interpreted as non-solvated [Cu(μ -pz)₂]_n and [Cu(μ -4-Xpz)₂]_n (X = Br, I) compounds, the too low crystallinity of which prevented a structural determination from PXRD data. As anticipated above, the crystal structure of α -[Cu(μ -4-Clpz)₂]_n, compound **3** in the present work, was reported by Storr and co-workers as early as in 1991,⁴⁸ and consisted in 1-D helices with CuN₄ tetrahedral metal centres. Hence, we used **3** as a benchmark for the assessment of the tetrahedral stereochemistry spectral properties, as compared to **3**_{DMF}, which was used to typify the octahedral geometry. In Figure 7a the absorption spectra of **3**_{DMF} and **3** in the 200-1200 nm spectral band are shown. The UV portion of the **3**_{DMF} spectrum is dominated by two major bands, one peaked at 210 nm, which can be attributed to the pyrazolate chromophore π - π^* transition, the other one peaked at 330 nm, which we ascribe to a charge-transfer transition from the pyrazolate ligand to the Cu(II) metal ions. In the visible region, the Cu(II) *d-d* transitions (see⁴⁸ for the attribution) build up a broad band peaked at 550 nm, with a shoulder in the red, around 650 nm. The same transitions can be detected in the spectrum of **3**. However, they are all notably red-shifted (to 240 nm, 380 nm, and 630 nm, respectively), with the *d-d* transition band shoulder falling in the NIR, around 800 nm, in agreement with results previously reported by Storr.⁴⁸ This evidence suggests a completely different charge distribution in the octahedral versus tetrahedral geometry.

In Figure 7b, the fluorescence emission spectra obtained upon excitation of **3**_{DMF} and **3** at the respective π - π^* transition band peak are shown. While the former spectrum consists in a broad,

almost unstructured main band covering the 400–480 nm wavelength interval, and in a second, less intense band peaked at 545 nm, the latter spectrum is constituted by three well defined bands peaked at 400 nm, 480 nm and 535 nm, respectively. Moreover, the green fluorescence band is relatively more intense than the blue fluorescence band in the **3** than in the **3**_{DMF} spectrum. In Figure 7c, the fluorescence spectra obtained upon excitation of **3**_{DMF} and **3** at the respective charge-transfer transition band peak are displayed. While the former complex emits sizeable fluorescence peaked at 435 nm, the latter is almost non-fluorescent. In light of this, the observations made on the fluorescence spectral features upon excitation at the π - π^* transition band peak can be given an interpretation. Indeed, the continuous emission from 400 nm to 480 nm in **3**_{DMF} can be interpreted as the result of the emission bands peaked at 400 nm and 480 nm, detected also for **3**, superimposed with the emission band peaked at 435 nm due to out-of-peak excitation of the charge-transfer transition. In summary, desolvation of **3**_{DMF}, resulting in a change from an octahedral to a tetrahedral geometry of the metal ion, implies a general bathochromic shift of the UV-Vis absorption bands, combined to a substantial quenching of fluorescence emission, associated to the relaxation from the charge-transfer transition state.

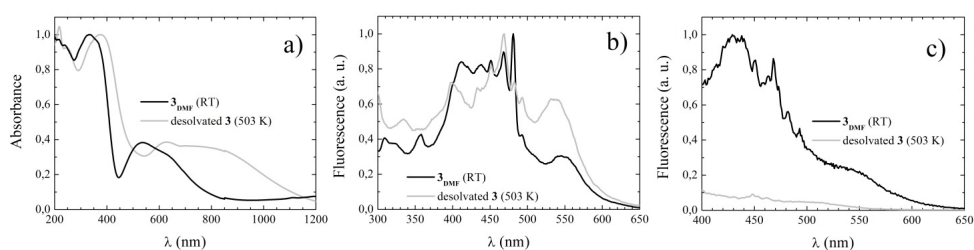


Figure 7. Comparison of the electronic-state transition spectroscopic properties of **3**_{DMF} and **3**: a) UV-Vis-NIR absorption spectra; b) fluorescence emission spectra upon excitation at the π - π^* transition band peak; c) fluorescence emission spectra upon excitation at the charge-transfer band peak. Colour codes are specified in the figure legends.

The above described spectral properties were used as benchmarks of the presence of the tetrahedral geometry in the powders of **2**, **4**, and **5**. Upon heating a sample of **2**_{DMF} at 463 K, a new phase appears and is present together with **2**_{DMF} (Figure S11): the absorption and fluorescence spectral features of **2**_{DMF} are conserved in this sample (see Figure 8), although we assist to both a widening of the absorption bands, and a partial quenching of the charge-transfer transition excited fluorescence, possibly due to local distortions. Heating to 583 K, temperature at which **2**_{DMF} has completely disappeared (Figure S11), results in complete loss of the absorption spectral line-shape structure (Figure 8a). Namely, the π - π^* charge transfer and d - d transition lines are integrated within a rather broad absorption band peaked at 355 nm. Thus, although a red shift in the main absorption

peak is actually observed, a convincing evidence in favour of the assumption of the tetrahedral geometry is lacking. Nonetheless, upon excitation at 355 nm, no fluorescence signal is detected (Figure 8c). Moreover, upon excitation at 240 nm, a fluorescence spectrum very similar to that observed for **3** is retrieved (Figure 8b).

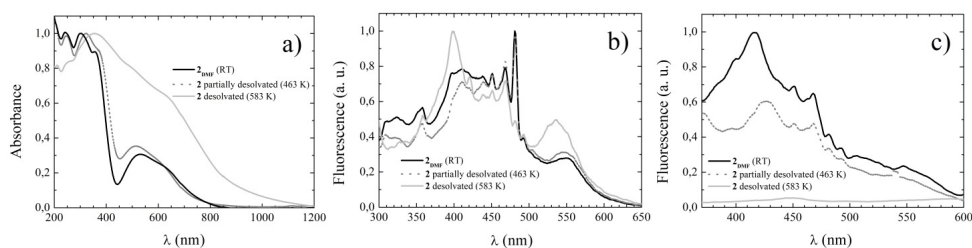


Figure 8. Comparison of the electronic-state transition spectroscopic properties of **2**_{DMF}, **2**_{DMF} heated at 463 K, and **2**: a) UV-Vis-NIR absorption spectra; b) fluorescence emission spectra upon excitation at the π - π^* transition band peak; c) fluorescence emission spectra upon excitation at the charge-transfer band peak. Colour codes are specified in the figure legends.

The situation is more suggestive in the case of **4**_{DMF} heated to 643 K (Figure 9) and **5**_{DMF} heated to 523 K (Figure 10), temperatures at which the low crystallinity phases **4** and **5** are detected by VT-PXRD (Figures S13 and S14, respectively).

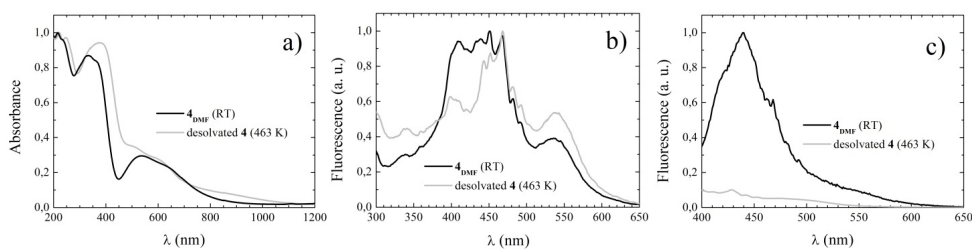


Figure 9. Comparison of the electronic-state transition spectroscopic properties of **4**_{DMF} and **4**: a) UV-Vis-NIR absorption spectra; b) fluorescence emission spectra upon excitation at the π - π^* transition band peak; c) fluorescence emission spectra upon excitation at the charge-transfer band peak. Colour codes are specified in the figure legends.

In these cases, both the π - π^* and the charge transfer bands are well resolvable within the absorption spectrum, and they are both significantly red shifted with respect to **4**_{DMF} (Figure 9a) and **5**_{DMF} (Figure 10a). Moreover, the shoulder above 800 nm distinctive of the *d-d* absorption of tetrahedral Cu(II) ions in **3** appears, although in the case of **4** also a blue-shifted *d-d* absorption is present. The fluorescence spectra of both **4** (Figure 9b) and **5** (Figure 10b) obtained upon excitation at the π - π^* band peak display the characteristic two-bands shape of **3**, and a minimum in correspondence of the fluorescence emission maximum detected upon excitation of the solvated

compounds at the charge-transfer band peak. Moreover, excitation in this band does not result in fluorescence emission for either of the two compounds (Figures 9c and 10c).

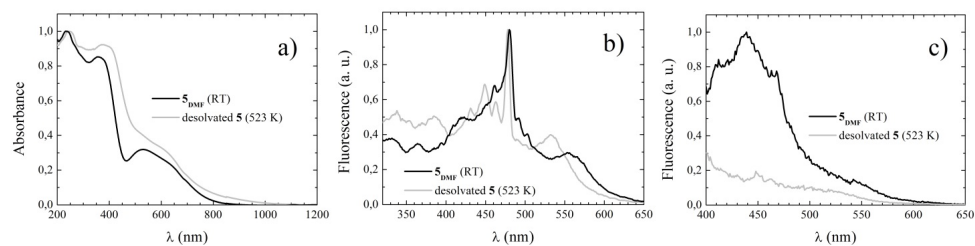


Figure 10. Comparison of the electronic-state transition spectroscopic properties of 5_{DMF} and 5 : a) UV-Vis-NIR absorption spectra; b) fluorescence emission spectra upon excitation at the π - π^* transition band peak; c) fluorescence emission spectra upon excitation at the charge-transfer band peak. Colour codes are specified in the figure legends.

These spectral features concur in indicating that the metal centre of the non-solvated complexes assumes a tetrahedral geometry, although the appearance of a blue-shifted d - d band and the lower intensity of the 800 nm d - d shoulder with respect to 3 might be the symptom of partial loss of crystallinity. The main spectroscopic data just discussed are compared in Table 2.

Table 2. Spectral properties of **2_{DMF}**-**5_{DMF}** and **2-5**. Peak wavelength of π - π^* transition absorption band, $\lambda_{\pi-\pi^*}$; peak wavelength of charge-transfer transition absorption band, λ_{charge} ; peak wavelength of d - d transition absorption band, λ_{d-d} ; peak wavelength(s) of fluorescence emission spectrum upon excitation at the π - π^* transition band peak, $\lambda_{\text{fluo},\pi-\pi^*}$; peak wavelength of fluorescence emission spectrum upon excitation at the charge-transfer transition band peak, $\lambda_{\text{fluo,charge}}$.

Compound	$\lambda_{\pi-\pi^*}$ (nm)	λ_{charge} (nm)	λ_{d-d} (nm)	$\lambda_{\text{fluo},\pi-\pi^*}$ (nm)	$\lambda_{\text{fluo,charge}}$ (nm)
2_{DMF}	242	300	531 (621)	411, 439, 451, 468, 481, 547	417
2	(259)	355	(620)	398, 469, 535	-
3_{DMF}	213 (238)	332	536 (625)	412, 437, 451, 468, 481, 545	434
3	217 (253)	374	629 (800)	400, 468, 536	-
4_{DMF}	215 (232)	332	535 (620)	411, 439, 451, 468, 536	439
4	223, 245	375	500 (590, 840)	403, 468, 540	-
5_{DMF}	236	357	528 (628)	420, 461, 480, 554	440
5	250	370	520 (605, 850)	431, 450, 463, 533	-

2.5. Theoretical characterization

The availability of accurate structural parameters has paved the way for the *ab initio* study of compounds **1** and **2_{DMF}**-**5_{DMF}**. Starting from the experimental crystal structure, each compound was fully relaxed in both its low- and high-spin configuration. For all compounds, the low-spin configuration is favoured with respect to the high-spin state by a quantity ranging from 1.13 (**2_{DMF}**) to 5.69 (**1**) kJ/mol, as evaluated at the B3LYP-D* level. In all the systems, the spin-only magnetic moment has a value of ~ 1.27 B.M., in rather good agreement with the observed ones.

The B3LYP-D* equilibrium structures of the compounds (see Tables S2 and S3 for the unit cell parameters) show a sensitive contraction of the unit cell volumes ($\sim 9\%$) with respect to the experimental values, which can be ascribed to an overestimation of the dispersive interactions in particular along the O \cdots Cu \cdots O direction, due to Grimme's correction. Nevertheless, in spite of the volume contractions, the mean bond distances and angles are overall preserved. In particular, in **2_{DMF}**-**5_{DMF}** the Cu \cdots Cu intra-chain distances have values of ~ 3.5 Å and the Cu-O bonds, which are shorter by ~ 0.2 Å with respect to the experimental values, are still 20% longer than the Cu-N ones, as predicted by the Jahn-Teller distortion. As regards **1**, the Cu-O (1.98-1.99 Å) and Cu-N (1.97-1.99 Å) distances are almost indistinguishable and extremely closed to the experimental values.

PBE geometries optimization (see Table S4 for the unit cell parameters) yielded a sensitive increase of the unit cell volumes, confirming the key role of dispersive interactions in determining the crystal structure of these compounds.

The B3LYP-D* optimized geometries were then used to calculate the harmonic IR frequencies, whose wavenumbers are all real, confirming that the periodic models are minima on the potential energy surface. Despite significant discrepancies in the absolute values of calculated *vs.* measured frequencies, due to the well-known difficulties of the harmonic approximation in describing light-atom modes, the main features of the experimental spectra are perfectly reproduced, as well as the general trends and peculiarities of each compound. The IR spectrum of **1** presents the O-H bond stretching mode at 3790 cm⁻¹ (*vs.* the experimental value of 3602 cm⁻¹) and confirms the assignment of the signal around 920 cm⁻¹ to the vibration of the Cu-O bond. The different coordination of the Cu(II) metal site in **1** and **2_{DMF}** clearly emerges from the spectral fingerprint of the two polymers, as shown in Figure S16. The IR spectra of **2_{DMF}**-**5_{DMF}** exhibit similar patterns, as already pointed out in section 2.1. The presence of coordinated DMF is confirmed by the intense band in the range 1713-1705 cm⁻¹, due to the C=O stretching and correctly red-shifted with respect to its calculated position in the free DMF molecule (1775 cm⁻¹). The effect of the halogen atomic mass is highlighted by the calculated $\nu(\text{C-X})$ stretching mode frequencies of **3_{DMF}**-**5_{DMF}** that are found at 1008, 975 and 959 cm⁻¹ for X = Cl, Br and I, respectively.

The thermal behaviour was simulated by totally removing DMF molecules from **3_{DMF}**-**5_{DMF}**. A subsequent complete relaxation of the structures has led to the theoretical coordination polymers **3-5**, whose structural parameters are summarized in Table S5. In all cases, the predictable structural effect is a dramatic reduction of the unit cell volumes that become almost 30% smaller than in the solvated compounds. From an energetic point of view, the non-solvated structures are 103.6, 115.5 and 164.4 kJ/mol less stable than the **3_{DMF}**, **4_{DMF}** and **5_{DMF}** systems, respectively, at the B3LYP-D* level. A similar estimate, performed with B3LYP only, shows that almost the 70% of the computed stability has to be ascribed to the Cu-O bond and that this essential contribution is mainly due to dispersive interactions, slightly overestimated by the *a posteriori* Grimme-D3 correction. The peaks of the $\nu(\text{C-X})$ stretching modes, due to the absence of the DMF molecules, are blue-shifted (~ 6 cm⁻¹ *vs.* the measured shift of 0-5 cm⁻¹) as can be seen in Figure S17. Interestingly enough, calculations indicate that the volume reduction is also driven by the incoming of new interplays involving the Cu(II) ions. In particular, in the Cl/Br compounds the Cu-X (X = Cl, Br) distances of 5.41/5.37 Å in **3_{DMF}**/**4_{DMF}** are shortened to 3.42/3.33 Å in **3/4**.

Unfortunately, simulations performed at 0 K prevent to overcome the local minimum in the potential energy surface represented by the square planar coordination of the metal site: a certain

amount of thermal energy is indeed needed to promote the concerted rearrangement of the pyrazole rings yielding to the observed tetrahedral coordination of Cu(II).

3. Experimental Section

3.1. Materials and Methods

All the chemicals were purchased from Sigma-Aldrich and used without further purification. $[\text{Cu}(\mu\text{-pz})_2\cdot\text{H}_2\text{O}]_n^{42}$ and $[\text{Cu}_3(\mu_3\text{-OH})(\mu\text{-pz})_3(\text{CH}_3\text{COO})_2(\text{pzH})]^{27}$ were prepared according to literature methods. All the reactions and manipulations were carried out in air, under either ambient or solvothermal conditions. Elemental analyses (C, H, N) were performed with a Fisons Instruments 1108 CHNS-O elemental analyser. Infrared spectra were recorded from 4000 to 650 cm^{-1} with a Perkin Elmer Spectrum One Model FTIR spectrometer with ATR mode. Thermogravimetric analyses on **2**_{DMF}-**4**_{DMF} and **m-5**_{DMF} were performed using a Thermogravimetric Analyzer Pyris 1 Perkin-Elmer heating from 303 to 1073 K with a speed of 10 K/min, under a flow of N₂ (30 mL/min). The accuracy of the sample temperature measurement was checked by the onset fusion temperature of indium (156.6, 0.2 K) and tin (232.0, 0.2 K) with heating/cooling dynamic segments. The experiments were repeated twice to confirm the reproducibility of the generated data. Simultaneous thermal analysis (simultaneous thermogravimetric analysis and differential scanning calorimetry) on **1** was carried out using a NETZSCH STA 409 PC instrument. 10 mg of as-synthesised **1** were placed in an alumina crucible; the temperature programme ranged from 303 to 1173 K with a heating rate of 10 K min⁻¹ under a flow of N₂ (40 mL min⁻¹). Raw data from TGA and DSC were corrected based on a background curve. Magnetic susceptibilities were measured at 293 K by the Gouy method with a Sherwood Scientific magnetic balance MSB-Auto, using HgCo(NCS)₄ as calibrant, and were corrected for diamagnetism with the appropriate Pascal constants. The magnetic moments (in Bohr Magnetons, B.M.) were calculated from the equation $\mu_{\text{eff}} = 2.84(\chi_{\text{m}}^{\text{corr}}T)^{1/2}$, where $\chi_{\text{m}}^{\text{corr}}$ is the corrected mass magnetic susceptibility, T = 293 K. The SEM images were acquired with a Field Emission Scanning Electron Microscope (FE-SEM) ZEISS SIGMA 300 FESEM equipped with Gemini column, able to reach a maximum resolution of 1.2 nm, and with three electron detectors, namely: high efficiency secondary "in-lens" electron detector; ETSE secondary electron detector (Everhart-Thornley Secondary Electrons); HDBSD backscattered electron detector with 4 independent sectors (High Definition Back Scattered Detector). Powder X-ray diffraction (PXRD) qualitative analyses were carried out with a Bruker D8 Advance diffractometer (see the section describing the structure characterization for the instrument specifics), acquiring data at room temperature in the 3–35° 2θ range, with steps of 0.02° and time *per* step of 1 s.

The electronic-state absorption spectroscopy measurements were undertaken by taking advantage of a Jasco V-770 UV-Vis-NIR spectrophotometer, equipped with a ISN-923 60 mm integrating sphere and an explicitly designed solid-state sample holder. The fluorescence emission spectra in solid state were recorded with a Jasco FP 8500 spectrofluorimeter. The powders were introduced in the same sample holder as above and submitted to the excitation beam with an inclination angle of 36° through a goniometric lodge, in order to minimize collection of excitation stray light.

3.2. Syntheses

*Synthesis of [Cu(μ -pz)(μ -OH)]_n, **1**.* [Cu(μ -pz)(μ -OH)]_n was isolated by following two synthetic approaches:

- a) A suspension of [Cu₃(μ_3 -OH)(μ -pz)₃(CH₃COO)₂(pzH)] (1.0 mmol, 0.595 g) in water (100 mL) was refluxed for 3 h, observing a colour change of the solid from blue to blue-violet. The blue-violet solid was filtered, washed with water and dried under vacuum at room temperature, recovering **1** in the form of powders. Yield 0.107 g, 72%. Melting point: > 573 K (decomposition).

Elemental analysis calculated for C₃H₄CuN₂O: C, 24.41, H, 2.73; N, 18.98%. found: C, 24.62; H, 2.68, N, 18.85%. IR (cm⁻¹): 3602m ν (O-H), 3117w ν (C-H_{arom}), 1489m, 1415m, 1378s ν (C=C + C=N), 1276m, 1182m, 1061s, 921s ν (Cu-O), 872m, 748vs. Magnetic susceptibility: μ_{eff} (293 K) = 0.91 B.M.

- b) A suspension of [Cu(μ -pz)₂·H₂O]_n (1.0 mmol, 0.217 g) in water (100 mL) was refluxed for 2 h. The obtained blue-violet solid was quickly filtered, washed with water, dried under vacuum and recognized as **1** by means of elemental analysis and infrared spectroscopy. Yield 0.115 g, 78%.

*Synthesis of [Cu(μ -pz)₂(μ -DMF)]_n, **2**_{DMF}.* Cu(CH₃COO)₂·H₂O (1.0 mmol, 0.199 g) was added to a pzH (2 mmol, 0.136 g) solution in DMF (20 mL). The reaction was carried out by the microwave-assisted solvothermal technique with the following reaction conditions: 0.5 h at 413 K, 2 h at 393 K and slow cooling at 8 K/minute to room temperature. The pale pinkish-violet precipitate thus obtained was filtered, washed with cold DMF and dried in air. Yield 0.223 g, 84%. Elemental analysis calculated for C₉H₁₃CuN₅O: C, 39.92; H, 4.84; N, 25.86%. found: C, 39.79; H, 4.79; N, 25.77%. IR (cm⁻¹): 3129w and 3106w ν (C-H_{arom}), 2927wbr ν (C-H_{aliph}), 1655vs ν (C=O), 1483m ν (C=C + C=N), 1383s 1095m, 1055s, 862m, 745vs. Magnetic susceptibility: μ_{eff} (293 K) = 1.56 B.M.

*Synthesis of [Cu(μ -4-Clpz)₂(μ -DMF)]_n, **3**_{DMF}.* The pale violet compound **3**_{DMF} was synthesized similarly to **2**_{DMF} by using 4-chloro-1H-pyrazole (4-ClpzH) instead of pzH. Yield 0.289

g, 88%. Elemental analysis calculated for: $C_9H_{11}Cl_2CuN_5O$: C, 31.82; H, 3.26; N, 20.62%. Found: C, 31.78; H, 3.13; N, 20.53%. IR (cm^{-1}): 3146vw $\nu(C-H_{arom})$, 2931wbr $\nu(C-H_{aliph})$, 1655vs $\nu(C=O)$, 1496m $\nu(C=C + C=N)$, 1394s, 1099m, 1058m, 969s $\nu(C-Cl)$, 830s, 662m. Magnetic susceptibility: μ_{eff} (293 K) = 1.32 B.M.

Synthesis of $[Cu(\mu-4-Brpz)_2(\mu-DMF)]_n$, 4_{DMF} . The pale violet compound 4_{DMF} was synthesized similarly to 2_{DMF} by using 4-bromo-1H-pyrazole (4-BrpzH) instead of pzH. Yield 0.383 g, 89%. Elemental analysis calculated for: $C_9H_{11}Br_2CuN_5O$: C, 25.22; H, 2.59; N, 16.34%. Found: C, 25.32; H, 2.57; N, 16.27%. IR (cm^{-1}): 3146vw $\nu(C-H_{arom})$, 2928wbr $\nu(C-H_{aliph})$, 1655vs $\nu(C=O)$, 1435m $\nu(C=C + C=N)$, 1395s, 1289m, 1100m, 1057s, 955s $\nu(C-Br)$, 829s, 663m. Magnetic susceptibility: μ_{eff} (293 K) = 1.33 B.M.

Synthesis of $[Cu(\mu-4-Ipz)_2(\mu-DMF)]_n$, 5_{DMF} . The violet compound 5_{DMF} was synthesized similarly to 2_{DMF} by using 4-iodo-1H-pyrazole (4-IpzH) instead of pzH. Yield 0.469 g, 90%. Elemental analysis calculated for: $C_9H_{11}CuI_2N_5O$: C, 20.69; H, 2.12; N, 13.40%. Found: C, 20.75; H, 2.09; N, 13.29%. IR (cm^{-1}): 3127vw $\nu(C-H_{arom})$, 2933wbr and 2855w $\nu(C-H_{aliph})$, 1651vs $\nu(C=O)$, 1431m, 1374s $\nu(C=C + C=N)$, 1281m, 1098m, 1053s, 942s $\nu(C-I)$, 841s, 662m. Magnetic susceptibility: μ_{eff} (293 K) = 1.38 B.M.

Synthesis of $[Cu(\mu-pz)_2]_n$, 2 . Compound 2_{DMF} (0.5 mmol, 0.135 g) was maintained at 453 K under atmospheric pressure for two hours, to completely remove DMF, obtaining compound 2 as a light grey powder. Yield 0.096 g, 98%. Elemental analysis calculated for: $C_6H_6CuN_4$: C, 36.45; H, 3.06; N, 28.35%. Found: C, 36.49; H, 3.09; N, 28.39%. IR (cm^{-1}): 3128w $\nu(C-H_{arom})$, 1487m $\nu(C=C + C=N)$, 1380s, 1276m, 1176m, 1055s, 888w, 784m, 745s, 757s. Magnetic susceptibility: μ_{eff} (293 K) = 1.59 B.M.

Synthesis of $[Cu(\mu-4-Clpz)_2]_n$, 3 . Compound 3_{DMF} (0.5 mmol, 0.169 g) was maintained at 453 K under atmospheric pressure for two hours, to completely remove DMF, obtaining compound 3 as a green powder. Yield 0.132 g, 99%. Elemental analysis calculated for: $C_6H_4Cl_2CuN_4$: C, 27.03; H, 1.51; N, 21.02%. Found: C, 26.92; H, 1.57; N, 20.97%. IR (cm^{-1}): 3149w $\nu(C-H_{arom})$, 1394m $\nu(C=C + C=N)$, 1293s, 1212m, 1153m, 1060vs, 973s $\nu(C-Cl)$, 840s, 828s, 669w. Magnetic susceptibility: μ_{eff} (293 K) = 1.44 B.M.

Synthesis of $[Cu(\mu-4-Brpz)_2]_n$, 4 . Compound 4_{DMF} (0.5 mmol, 0.214 g) was maintained at 453 K under atmospheric pressure for two hours, to completely remove DMF, obtaining compound 4 in the form of a light brown powder. Yield 0.176 g, 98%. Elemental analysis calculated for: $C_6H_4Br_2CuN_4$: C, 20.27; H, 1.13; N, 15.76%. Found: C, 20.31; H, 1.15; N, 15.77%. IR (cm^{-1}): 3149w $\nu(C-H_{arom})$, 1386m $\nu(C=C + C=N)$, 1287s, 1201m, 1153m, 1153m, 1058vs, 955s $\nu(C-Br)$, 843s, 826s, 667w. Magnetic susceptibility: μ_{eff} (293 K) = 1.43 B.M.

*Synthesis of [Cu(μ -4-*Ipz*)₂]_n.* **5**. Compound **5**_{DMF} (0.5 mmol, 0.261 g) was maintained at 453 K under atmospheric pressure for two hours, to completely remove DMF, obtaining compound **5** in the form of a light brown powder. Yield 0.223 g, 99%. Elemental analysis calculated for: C₆H₄CuI₂N₄: C, 16.03; H, 0.90; N, 12.47%. Found: C, 16.14; H, 0.88; N, 12.63%. IR (cm⁻¹): 3136w ν (C-H_{arom}), 1373m ν (C=C + C=N), 1279s, 1201m, 1157m, 1056vs, 942s ν (C-I), 839s, 826s, 664w. Magnetic susceptibility: μ_{eff} (293 K) = 1.47 B.M.

3.3. Powder X-ray Diffraction Structural Characterization

Microcrystalline powders of **1**, **2**_{DMF}-**4**_{DMF}, **oC**-**5**_{DMF} and **oP**-**5**_{DMF} were deposited in the hollow of a 0.2 mm deep silicon zero-background plate. Data acquisitions were performed on a vertical-scan Bruker AXS D8 Advance θ : θ diffractometer, equipped with a Lynxeye linear position-sensitive detector, an X-ray tube (Cu K α , λ = 1.5418 Å), a filter of nickel in the diffracted beam and the following optical components: primary beam Soller slits (2.5°), fixed divergence slit (0.5°), antiscatter slit (8 mm). The generator was set at 40 kV and 40 mA. After preliminary acquisitions for fingerprinting analysis, typically performed in the 3–35° 2 θ range, diffraction data sets for a full structure determination were collected from 5° up to 105° 2 θ , with steps of 0.02°, and an overall scan time of approximately 12 hours. In case of **1**, **2**_{DMF}, **oC**-**5**_{DMF} and **oP**-**5**_{DMF}, a peak search, followed by profile fitting, enabled us to estimate the low-to-medium-angle peak maximum positions which, through the Singular Value Decomposition algorithm⁴⁹ implemented in TOPAS-R V3,⁵⁰ provided approximate unit cell parameters for all the compounds. The space groups were assigned based on the observed systematic absences. Comparison of the PXRD patterns of **3**_{DMF} and **4**_{DMF} with that of **2**_{DMF} suggested that the three compounds share the same space group and similar unit cell parameters, as proved by whole powder pattern refinements carried out with the Le Bail method as implemented in TOPAS-R V3. Structure determination was carried out for **1**, **2**_{DMF} and **5**_{DMF} with TOPAS-R V3 by a combined Monte Carlo/Simulated Annealing approach. A rigid body was used to describe the crystallographically independent portion of the pz⁻ and 4-Xpz⁻ (X = Cl, Br, I) ligands, initially assigning average values to bond distances and angles.^[11] The position of their centre of mass and their orientation (when allowed by symmetry) were let to vary. Also DMF was modelled as a rigid body.^[11] Structure refinement was carried out with the Rietveld method as implemented in TOPAS-R V3. In the case of **3**_{DMF} and **4**_{DMF} the crystal and molecular structures were refined starting from the crystal structure of **2**_{DMF}, after introducing, in position 4 of the pyrazolate, a chlorine and bromine atom, respectively. During the final Rietveld refinement stages, the site occupation factor of the DMF molecule was refined without restraints, reaching a value

lower than 1 in the case of **2_{DMF}** (0.8) and **5_{DMF}** (0.9). In addition, ligand bond distances (except the C/N-H distances) were refined in a restrained range of values.^[III] In all cases, the background was modelled by a polynomial function of the Chebyshev type. A common, refined isotropic thermal factor (B_{iso}) was attributed to all atoms, except to the metal centres, to which the isotropic thermal factor $B_{\text{iso}}(\text{M}) = B_{\text{iso}} + 2.0 \text{ (\AA}^2\text{)}$ was assigned. The peak profile was described by the Fundamental Parameters Approach.⁵¹ Anisotropic peak broadening was successfully described using second- or fourth-order spherical harmonics. Preferred orientation was observed in the case of **2_{DMF}-4_{DMF}** (along the [110] pole) and **5_{DMF}** (along the [100] pole) and corrected by the March-Dollase approach.⁵² The final Rietveld refinement plots are shown in Figures S18 and S19 of the Electronic Supplementary Information. The pertinent CIF files are supplied as Electronic Supplementary Information.

Crystal data for **[Cu(μ -pz)(μ -OH)]_n, **1**: C₃H₄CuN₂O, FW = 147.62 g mol⁻¹, orthorhombic, *Pbca*, $a = 18.7113(9) \text{ \AA}$, $b = 6.7293(3) \text{ \AA}$, $c = 6.8436(1) \text{ \AA}$, $V = 861.71(6) \text{ \AA}^3$, $Z = 8$, $Z' = 8$, $\rho = 2.27 \text{ g cm}^{-3}$, $F(000) = 584$, $R_{\text{Bragg}} = 0.003$, $R_{\text{p}} = 0.013$ and $R_{\text{wp}} = 0.019$, for 4901 data and 42 parameters in the 7.0-105.0° (2θ) range. CCDC No. [?](#).**

Crystal data for **[Cu(μ -pz)₂(μ -DMF)_{0.8}]_n, **2_{DMF}**: C_{8.4}H_{11.6}CuN_{4.8}O_{0.8}, FW = 256.16 g mol⁻¹, orthorhombic, *Pnam*, $a = 12.1799(5) \text{ \AA}$, $b = 13.7527(6) \text{ \AA}$, $c = 7.0999(4) \text{ \AA}$, $V = 1189.28(9) \text{ \AA}^3$, $Z = 8$, $Z' = 4$, $\rho = 1.43 \text{ g cm}^{-3}$, $F(000) = 522.4$, $R_{\text{Bragg}} = 0.006$, $R_{\text{p}} = 0.026$ and $R_{\text{wp}} = 0.038$, for 4876 data and 53 parameters in the 7.5-105.0° (2θ) range. CCDC No. [?](#).**

Crystal data for **[Cu(μ -4-Clpz)₂(μ -DMF)]_n, **3_{DMF}**: C₉H₁₁Cl₂CuN₅O, FW = 339.67 g mol⁻¹, orthorhombic, *Pnam*, $a = 13.7038(3) \text{ \AA}$, $b = 14.4516(3) \text{ \AA}$, $c = 7.1384(2) \text{ \AA}$, $V = 1413.70(6) \text{ \AA}^3$, $Z = 8$, $Z' = 4$, $\rho = 1.60 \text{ g cm}^{-3}$, $F(000) = 684$, $R_{\text{Bragg}} = 0.015$, $R_{\text{p}} = 0.024$ and $R_{\text{wp}} = 0.034$, for 4876 data and 47 parameters in the 7.5-105.0° (2θ) range. CCDC No. [?](#).**

Crystal data for **[Cu(μ -4-Brpz)₂(μ -DMF)]_n, **4_{DMF}**: C₉H₁₁Br₂CuN₅O, FW = 428.57 g mol⁻¹, orthorhombic, *Pnam*, $a = 14.1228(4) \text{ \AA}$, $b = 14.4113(3) \text{ \AA}$, $c = 7.1279(2) \text{ \AA}$, $V = 1450.72(6) \text{ \AA}^3$, $Z = 8$, $Z' = 4$, $\rho = 1.96 \text{ g cm}^{-3}$, $F(000) = 828$, $R_{\text{Bragg}} = 0.019$, $R_{\text{p}} = 0.021$ and $R_{\text{wp}} = 0.031$, for 4851 data and 50 parameters in the 8-105.0° (2θ) range. CCDC No. [?](#).**

Crystal data for **[Cu(μ -4-Ipz)₂(μ -DMF)_{0.9}]_n, **oC-5_{DMF}**: C_{8.6}H₁₀I₂CuN_{4.9}O_{0.9}, FW = 513.76 g mol⁻¹, orthorhombic, *Cmcm*, $a = 20.2680(5) \text{ \AA}$, $b = 9.5313(2) \text{ \AA}$, $c = 7.3586(1) \text{ \AA}$, $V = 1421.53(6) \text{ \AA}^3$, $Z = 16$, $Z' = 4$, $\rho = 2.40 \text{ g cm}^{-3}$, $F(000) = 949.6$, $R_{\text{Bragg}} = 0.054$, $R_{\text{p}} = 0.046$ and $R_{\text{wp}} = 0.069$, for 4851 data and 45 parameters in the 8-105.0° (2θ) range. CCDC No. [?](#).**

Crystal data for **[Cu(μ -4-Ipz)₂(μ -DMF)_{0.5}]_n, **oP-5_{DMF}**: C_{7.5}H_{7.5}I₂CuN_{4.5}O_{0.5}, FW = 486.02 g mol⁻¹, orthorhombic, *Pbnm*, $a = 20.1716(4) \text{ \AA}$, $b = 9.2333(2) \text{ \AA}$, $c = 7.2913(1) \text{ \AA}$, $V = 1358.00(5)$**

\AA^3 , $Z = 8$, $Z' = 4$, $\rho = 1.19 \text{ g cm}^{-3}$, $F(000) = 892$, $R_{\text{Bragg}} = 0.024$, $R_p = 0.027$ e $R_{\text{wp}} = 0.036$, for 4851 data and 49 parameters in the 8-105.0° (2θ) range. CCDC No. [2](#).

3.4. m-5_{DMF} oC-5_{DMF} Phase Transformation Monitoring

A 20-mg sample of as-synthesized 5_{DMF} (already a mixture of the pristine monoclinic and orthorhombic phases) was deposited in the hollow of a 0.2 mm deep silicon zero-background plate. Consecutive PXRD data acquisitions were performed on the vertical-scan Bruker AXS D8 described above in the 2θ range 8.0-35.0° for a period of 5.4 hours.

3.5. Variable-temperature Powder X-ray Diffraction

To complement the thermal analysis, the thermal behaviour of **1** and 2_{DMF}-5_{DMF} was investigated *in situ* by variable-temperature PXRD. In the case of 5_{DMF}, the study was carried out starting from the oC-5_{DMF} phase. Using a custom-made sample heater (Officina Elettrotecnica di Tenno, Ponte Arche, Italy), 20-mg samples of the five compounds were heated in air with steps of 20 K and applying the experimental conditions collected in Table 3. A PXRD pattern was acquired at each step.

Table 3. Experimental conditions of the variable-temperature experiments carried out on compounds **1** and 2_{DMF}-5_{DMF}.

Compound	T range (K)	2 θ range (°)
1	303-623	8.0-33.5
2 _{DMF}	303-703	5.0-25.0
3 _{DMF}	303-683	5.0-27.5
4 _{DMF}	303-623	5.0-27.5
oC-5 _{DMF}	303-643	5.0-27.5

3.6. Computational Details

The theoretical characterization of the systems **1**, 2_{DMF}-4_{DMF} and oC-5_{DMF} was performed with the quantum-mechanical *ab initio* CRYSTAL⁵³ code which solves the Schrödinger equation for periodic systems adopting a basis set of localized orbitals, centred on each atom of the reference cell and obtained as linear combination of Gaussian-type functions (GTO). Both the Hartree-Fock (HF) and density functional methods (DFT) are available as well as several hybrid approaches.⁵⁴ In particular, the calculations were performed adopting the B3LYP-D* hybrid exchange-correlation functional,^{55,56} where D* stands for the D3 Grimme's *a posteriori* energy correction for dispersive interactions as

tuned for molecular crystals.⁵⁷ This approach enabled us to take into account weak binding effects that, however, can rule the interaction between the metal sites and coordinating molecular fragments, as in the case of Cu(II) and the oxygen atom of DMF. To avoid a possible overestimation of such effects and to determine their range of validity, some calculations were computed also at the B3LYP level. The basis set consisted of triple-z quality set (TZP) developed by Ahlrichs and co-workers⁵⁸ supplemented by a set of polarization functions on all atoms, and can be considered a good compromise between accuracy and computational time for solid state calculations. The level of accuracy in evaluating the Coulomb and exchange integrals was controlled by five parameters, T_1 – T_5 , that were set to 7, 7, 7, 12 and 24 in the present calculations. The Hamiltonian matrix was diagonalized in a set of reciprocal k points ranging from 27 to 36 as a function of the reciprocal lattice symmetry and corresponding to the common shrinking factor of 4.⁵⁹ Lattice constants and internal coordinates were simultaneously optimized using the analytical gradients and upgrading the numerical Hessian with the Shanno et al. algorithm.⁶⁰ Default convergence criteria for the geometry optimization on the root mean square and on the absolute value of the largest component of the gradient and the estimated nuclear displacement were used. Infrared intensity for each normal mode was obtained by computing the dipole moment variation along the normal mode, adopting the Berry phase method.⁶¹

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Funding Sources

Any funds used to support the research of the manuscript should be placed here (per journal style).

Foot Notes

- I. The non-standard space group $P112_1/m$ was preferred to $P2_1/m$ to have the 2_1 screw axis along which the 1-D helices run parallel to the [001] crystallographic direction, as in **2_{DMF}-4_{DMF}**.
- II. ^[II] Bond distances and angles for the rigid body describing the ligand: C-C and C-N of the pyrazole ring 1.36 Å; C-H of the pyrazole ring 0.95 Å; C-Cl 1.70 Å; C-Br 1.85 Å; C-I 2.05 Å; pyrazole ring internal and external bond angles 108° and 126°, respectively. Bond distances and angles for the rigid body describing the solvent: C-N, 1.45 Å; C-H, 0.95 Å; C-O, 1.25 Å; angles at the nitrogen and amidic carbon atoms, 120°; angles at the carbon atoms of the methyl groups, 109.5°.
- III. ^[III] C-C and C-N bonds in the pyrazolate ligand, 1.32-1.38 Å; C-N bonds in the DMF molecule, 1.40-1.45 Å; C-Cl, 1.65-1.77 Å; C-Br, 1.83-1.92 Å; C-I, 2.01-2.14 Å

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