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Solvatomorph and polymorph screening of clopamide drug and its copper(II) complex crystals†

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Clopamide ((4-chloro-*N*-2,6-dimethylpiperidin-1-yl)-3-sulfamoylbenzamide, LH) drug is used worldwide in the treatment of hypertension and oedema. Despite its medical application its structure has not yet been published. We report the crystal structure of clopamide anhydrate and hemihydrate and a structural landscape of different polymorphs and solvatomorphs of its copper(II) complexes. Our main challenge was the growth of single crystals for X-ray diffraction experiments and then the analysis of the molecular arrangements of the pure drug as well as the series of complexes organised by the coordination bond and by different secondary interactions. We present how the inclusion of water contributes to the crystal perfection of the drug crystals (**1** and **2**). The newly defined chalcogen bond is recognised in the clopamide anhydrate crystals being in competition with an intramolecular halogen bond. All Cu(II) complexes have a square-planar coordination geometry, in which copper(II) centres are surrounded by piperidine-N and carbonyl-O donor atoms in a five-membered chelate ring. The two ligands are in *trans* positions. The solvatomorph crystals, containing small homologous alcohols (**3**–**6**), have been found to be isostructural (*P*1̄) with increasing size of void and unit cell volumes which terminates using *n*-butanol. In the solvent-free polymorph [CuL₂] crystals (**7**–**9**), the complex molecules form parallel columns strengthened by N–H...O interactions between the ligand sulfonamide groups. The arrangement of these columns in **7** (*P*1̄) is similar to that of the solvate **3** (*P*1̄), while **8** (*P*1̄) transitions to **9** (*R*3̄), where perfect threefold symmetry is realised, although leaving large empty spaces in the lattice. The columnal arrangement is preserved in the crystal containing dichloromethane (**10**) but the columns are no longer parallel. The solvate content and thermal behaviour of the crystals were studied by IR and thermoanalytical methods. The solution structure of the [CuL₂] complex was investigated in DMSO by EPR spectroscopy at 77 K. These results of the systematic comparison of several polymorphic and solvatomorphic crystal arrangements contribute to the better understanding of the structure-directing effects, which provide strategies to control the molecular self-assembly.

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1 Introduction

Understanding the molecular conformation and crystal structures (including polymorphs and solvatomorphs) is imperative in the use and development of active pharmaceutical ingredients (APIs). Polymorphism plays a very important role in the bioavailability of a drug¹ as the physicochemical properties (*e.g.*, solubility, stability) of polymorphs can differ significantly.² In many cases, a serious

limitation is that only drugs that have GRAS (generally recognised as safe) compounds in their crystal lattice can obtain a marketing authorisation. It has to be considered at the time of the selection of the solvent from which the API is crystallized.^{3,4} Special attention should be paid to the crystallization conditions and the crystal structures of the APIs.

Clopamide ((4-chloro-*N*-2,6-dimethylpiperidin-1-yl)-3-sulfamoylbenzamide) is an API used currently in the treatment of hypertension and oedema. It is an oral thiazide-like diuretic agent with antihypertensive activity.⁵ It has two interconnected functional groups: a piperidine and a sulfamoylbenzamide group. Despite the medical application of clopamide, its crystal structure has not yet been published. It has been found that the most widely used sulfonamide-type antihypertensive agents cause significant urinary potassium losses (hypokalemia).^{6,7} This side effect also exists

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when applying clopamide as well.⁸ It has also been shown that thiazide diuretics may also cause urinary loss of certain trace elements or modify their levels in blood and may induce changes in the levels of copper in normal hypertensives.^{9,10} According to the molecular structure it is noticeable that the carbonyl oxygen and the piperidine nitrogen of the ligand are able to coordinate to metal centres; thus the complexation of clopamide with metal ions in the human body may be responsible for this side effect. On the other hand, various metal complexes can advantageously modify the biological activity of an API compared to the free ligand. There are references which report an increased bioactivity by forming metal complexes.^{11,12}

To gain a better insight into the structure and interaction of this drug molecule, the structure of clopamide and its coordination compounds with copper(II) has been studied (Scheme 1) using different crystallization solvents under different conditions. Our aim was to obtain insights into the solvatomorph and polymorph crystal forms of clopamide and its copper(II) complexes using single-crystal X-ray diffraction (SXRD). The outcome of successful crystallization experiments from a homologous series of alcohols allowed us to harvest single crystals of a series of clopamide derivatives (Table 1). The crystal structures of the anhydrous LH (1) and hemihydrate LH·0.5H₂O (2) forms of clopamide have been determined. The bis-ligand copper(II) complex crystals were synthesized from a homologous series of alcohols, with the inclusion of solvent molecules, resulting in [CuL₂]·2(MeOH)·2(H₂O) (3), [CuL₂]·2(EtOH)·2(H₂O) (4), [CuL₂]·2(PrOH)·2(H₂O) (5) and [CuL₂]·2(iPrOH)·2(H₂O) (6). From organic solvents, three solvent-free polymorphs of [CuL₂] 7–9 and a crystal containing dichloromethane [CuL₂]·CH₂Cl₂ (10) were prepared. It has opened up a way to determine how far the molecular conformation and the packing arrangement can preserve isostructurality.^{13,14} The thermodynamic and kinetic stability of these complexes was investigated by thermoanalytical studies. The solution structure of the clopamide bis-ligand copper(II) complex was tested using electron paramagnetic resonance (EPR) spectroscopy.

A better understanding of the structure-directing effects in crystallization contributes to the development of strategies to control the process of molecular self-assembly.^{15,16} The obtained results contain valuable information with regard to

Table 1 The successful crystallisation experiments of clopamide ((4-chloro-*N*-2,6-dimethylpiperidin-1-yl)-3-sulfamoylbenzamide, LH) drug, which were followed by single-crystal X-ray structure determination. Details of crystallization conditions can be found in section 2.2

1	LH
2	LH·0.5H ₂ O
3	[CuL ₂]·2(MeOH)·2(H ₂ O)
4	[CuL ₂]·2(EtOH)·2(H ₂ O)
5	[CuL ₂]·2(PrOH)·2(H ₂ O)
6	[CuL ₂]·2(iPrOH)·2(H ₂ O)
7	CuL ₂ form I
8	CuL ₂ form II
9	CuL ₂ form III
10	[CuL ₂]·CH ₂ Cl ₂

synthon engineering,¹⁷ as we performed a systematic comparison of several solvatomorphic and polymorphic crystal arrangements. These investigations enrich the knowledge on the aspects which contribute to the development of materials with specific properties.¹⁸

2 Materials and methods

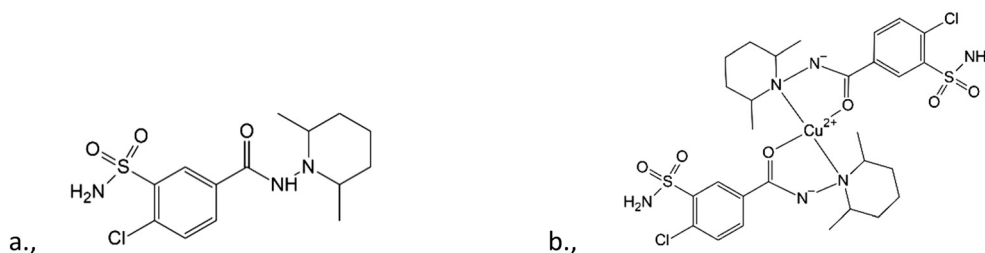
2.1 Reagents and materials

All chemicals were of reagent grade and used as purchased without any further purification. Clopamide was received from EGIS Pharmaceuticals Private Limited Company and used without further purification. All solvents were of analytical grade and used without further purification. CuCl₂·2H₂O, CuSO₄·5H₂O and NaOH were purchased from Sigma-Aldrich in analytical quality.

2.2 Synthesis of complexes and single crystal growth

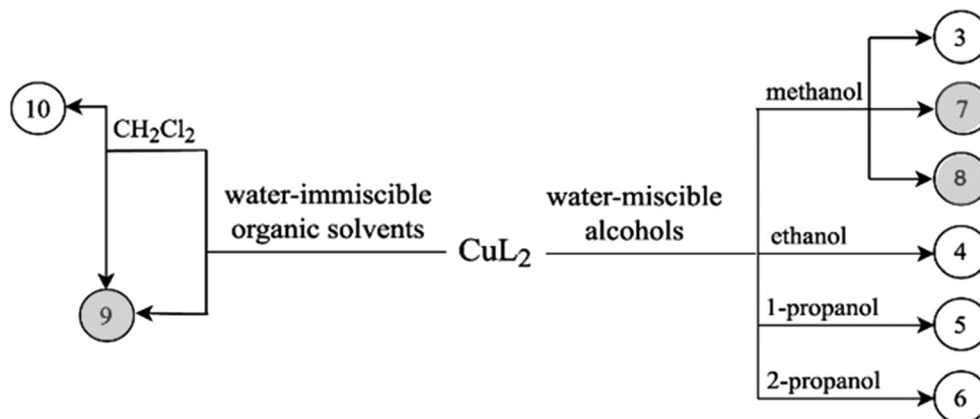
Crystallization experiments were performed in order to obtain single crystals suitable for SXRD measurements. Crystals were kept in their mother liquor to preserve their quality. A scheme of the determined copper(II) complex crystals obtained by solvatomorph and polymorph screening is depicted in Scheme 2.

2.2.1 Crystallization of HL (1). Clopamide (10 mg, 0.03 mmol) and CuSO₄·5H₂O (15 mg, 0.06 mmol) were added to 0.6 mL of methanol in a small ampule. A dark brown solution was formed, which was stirred at 65 °C for 1 hour. The solution was stored in a refrigerator at 5 °C. By decreasing the temperature, the solution became pale blue.



Scheme 1 General formula diagram of the clopamide (LH) molecule (a) and bis-ligand copper(II) complex [CuL₂] of clopamide (b). The carbonyl oxygens and the piperidine ring nitrogen atoms are coordinating to the copper centre.





Scheme 2 Crystallization experiments used to obtain crystals 3–10. Compound numbers with white background: crystals which enclathrate solvents; compound numbers with grey background: different polymorphic forms of CuL_2 .

Slow evaporation of the solvent (two weeks) in the fridge resulted in colourless clopamide anhydrate (**1**) single crystals.

2.2.2 Crystallization of $\text{HL} \cdot 0.5\text{H}_2\text{O}$ (2**).** Clopamide (5.1 mg, 0.015 mmol) was added to 0.6 mL of methanol in an ampule. The solution was stirred at room temperature for 5 minutes. Colourless single crystals were obtained by slow evaporation of the solvent after two days.

2.2.3 Syntheses and single-crystal growth of $[\text{CuL}_2] \cdot 2(\text{MeOH}) \cdot 2(\text{H}_2\text{O})$ (3**), $[\text{CuL}_2] \cdot 2(\text{EtOH}) \cdot 2(\text{H}_2\text{O})$ (**4**), $[\text{CuL}_2] \cdot 2(\text{PrOH}) \cdot 2(\text{H}_2\text{O})$ (**5**) and $[\text{CuL}_2] \cdot 2(\text{iPrOH}) \cdot 2(\text{H}_2\text{O})$ (**6**).** Clopamide (10–10 mg, 0.03 mmol) and CuCl_2 (0.1 mL, 0.1 M) were dissolved in 0.4–0.4 mL of methanol in the case of **3**, ethanol in the case of **4**, 1-propanol in the case of **5** and 2-propanol in the case of **6**. Dark green coloured solutions were formed. Each mixture was stirred at 65 °C for 5 minutes. The pH was set to approximately 9 by adding 0.02 mL 2 M NaOH solution in order to enhance the deprotonation of the hydrazine nitrogen atom and maximise the formation of the bis-ligand complex. Then, the reaction mixture was heated and became dark brown after a few minutes. Brown single crystals (**3**–**6**) were obtained at room temperature by slow evaporation of the solvents after four days.

2.2.4 Syntheses and crystallization of solvent-free polymorphic forms of $[\text{CuL}_2]$ (7**, **8** and **9**).** Clopamide (31 mg, 0.10 mmol) and $\text{CuSO}_4 \cdot 5\text{H}_2\text{O}$ (25 mg, 0.09 mmol) were dissolved in 0.5 mL of methanol. A dark brown coloured solution was formed. The mixture was stirred at 65 °C for 5 minutes. The pH of the reaction mixture was set to ~9 with 0.02 mL 2 M NaOH solution. The mixture was kept in the fridge at 5 °C. Brown single crystals of **7** were obtained after one year.

Clopamide in excess as ligand (23 mg, 0.07 mmol) and $\text{CuSO}_4 \cdot 5\text{H}_2\text{O}$ (5 mg, 0.02 mmol) were dissolved in 0.5 mL methanol. The pH of the reaction mixture was set to ~9 with 0.02 mL 2 M NaOH solution. The mixture was kept in the fridge at 5 °C. Brown single crystals of **8** were obtained after one year.

Clopamide (10 mg, 0.03 mmol) and an aqueous solution of CuCl_2 (0.1 mL, 0.1 M, 0.01 mmol) were added to water (0.3

mL). Clopamide is insoluble in water; thus a white suspension formed. After adding ethyl acetate (0.5 mL) two phases formed. Aqueous NaOH (0.02 mL, 2 M) was added dropwise to adjust the pH to approximately 9. The ampule was heated to 40 °C in order to increase the solubility of the complex in the organic solvent. The organic phase was separated and brown single crystals of **9** were obtained after one week by slow evaporation of the solvent in the fridge at 5 °C.

2.2.5 Synthesis and single-crystal growth of $[\text{CuL}_2] \cdot 2(\text{CH}_2\text{Cl}_2)$ (10**).** Crystals of **10** were prepared in the same way as crystal **9** but instead of ethyl acetate dichloromethane (DCM) was used. After the separation of the DCM solution brown single crystals of **10** were obtained after one day by slow evaporation of the solvent at room temperature.

2.3 Single-crystal structure determination

The single crystals were mounted on loops and transferred to a goniometer. X-ray diffraction data were collected at 103 K (**1**–**7**), 123 K (**8**) 158 K (**9**) and 128 K (**10**) on a Rigaku RAXIS-RAPID II diffractometer. The diffraction measurements were performed using $\text{Mo-K}\alpha$ radiation in all cases, except (**1**), (**9**) and (**10**), where a $\text{Cu-K}\alpha$ source was used. Numerical absorption corrections were carried out using the software CrystalClear.¹⁹ Sir2018 (ref. 20) and SHELXL²¹ under WinGX²² software were used for structure solution and refinement, respectively. The structures were solved by direct methods. The models were refined using the full-matrix least squares method on F^2 . Refinement of the non-hydrogen atoms was carried out with anisotropic thermal parameters. Hydrogen atoms were placed into geometric positions (except alcohol and water hydrogens). The atomic positions of alcohol and water hydrogens were located in difference electron density maps, whereas all the other hydrogens were placed in geometric positions. Hydrogens were included in structure factor calculations, but they were not refined. The isotropic displacement parameters of the hydrogen atoms were approximated from the $U(\text{eq})$ value of the atom they were



bonded to. In the case of crystal **7**, several data collections have been performed but the high R_{int} value could not be lowered (Table S9†), which is most probably due to crystal defects. After the removal of the crystals of **10** from the mother liquor the enclathrated dichloromethane molecules evaporate rapidly causing partial degradation of the single crystal, resulting in a higher R_{int} value. The details of data collection and refinement parameters are collected in Table S1.† Selected bond lengths and angles were calculated using the software PLATON.²³ The graphical presentations were prepared using Mercury²⁴ and the editing of CIF files was done using the software EnCifer.²⁵

2.4 NMR measurements and evaluation of the spectra

¹H and ¹³C NMR spectra of clopamide in deuterated DMSO were recorded using a Bruker Avance 400 (400 MHz spectrometer) at 25 °C. The spectra were evaluated using the Mestrenova 9 NMR program.²⁶

2.5 EPR measurements and evaluation of the spectra

X-band CW-EPR spectra were recorded on a BRUKER EleXsys E500 spectrometer (microwave frequency 9.4 GHz, microwave power 13 mW, modulation amplitude 5 G, modulation frequency 100 kHz). For the frozen solution measurement, a 0.2 mL sample of crystal **9** was prepared in 3.0 mM concentration, in pure DMSO, and placed into a quartz EPR tube. The EPR tubes were placed into a Dewar flask containing liquid nitrogen to perform the measurement at 77 K.

The spectrum was simulated using the EPR program.²⁷ Rhombic or axial *g* tensors and copper hyperfine *A* tensors were used for the simulation of the components. The splitting caused by the two coordinated nitrogen donor atoms is clearly visible in the spectrum and this superhyperfine coupling were taking into account by rhombic nitrogen tensor *a*. The linewidths were fitted with the help of orientation-dependent relaxation parameters (α , β , and γ) through the equation $\sigma_{M_I} = \alpha + \beta M_I + \gamma M_I^2$, where M_I denotes the magnetic quantum number of the copper nucleus. Since natural CuCl₂ was used for the measurements, the spectra were calculated as the sum of the spectra of ⁶³Cu and ⁶⁵Cu weighted by their natural abundances. The copper and nitrogen coupling constants and the relaxation parameters were obtained in field units (Gauss = 10⁻⁴ T).

2.6 Thermoanalytical methods

Thermal measurements were performed on a Setaram LabsysEvo (Lyon, France) TG–DSC system, in a flowing (90 mL min⁻¹) helium gas (99.9999% purity) atmosphere. The sample was weighed directly into a 100 μ L aluminium crucible (the reference cell was empty) and was heated from 20 °C to 500 °C at a heating rate of 20 °C min⁻¹. After loading the sample into the thermal analyzer and prior to running the measurement, a 10 min purging step was run with a high helium flow rate (200 mL min⁻¹). The obtained data were

baseline corrected and further processed using the thermoanalyzer's processing software (Calisto Processing, ver. 2.092). The thermal analyzer (both the temperature scale and calorimetric sensitivity) was calibrated using a multipoint calibration method, in which seven different certified reference materials (CRMs) were used to cover the thermal analyzer's entire operating temperature range. In parallel with the thermal measurements, the analysis of evolved gases/volatiles was performed on a Pfeiffer Vacuum Omni Star™ mass spectrometric evolved gas analysis (MS-EGA) system, which was connected to the above-mentioned thermal analyzer. The gas splitter at the thermal analyzer's furnace exit was thermostated to 220 °C, while the transfer line to the mass spectrometer was thermostated to 200 °C in order to avoid condensation of some analytes. The temperature of the mass spectrometer gas inlet was programmed to 120 °C. In order to compensate for the dead time (the time elapsed from the release of any volatile until it reached the ion source of the MS) between the thermal analyzer and the mass spectrometer at a given gas flow rate, both instruments were electronically synchronized. The volume of gas entering the mass spectrometer is 1–2 standard cubic centimeters per minute. EGA measurements were done in SEM bar graph cycle acquisition mode, where the *m/z* interval of 10–170 was continuously scanned with a speed of 20 ms amu⁻¹. The spectrometer was operated in electron impact mode.

2.7 Infrared measurements

IR spectra were collected by means of a Varian 2000 (Scimitar Series) FT-IR spectrometer (Varian Inc., US) equipped with a broad-band mercury–cadmium–telluride (MCT) detector and with a ‘Golden Gate’ single reflection diamond attenuated total reflection (ATR) unit (Specac Ltd, UK). The ATR technique allows the measurement of the samples without any sample preparation; for comparability, all samples were pressed with a constant pressure of 70 cNm at the diamond surface with a sapphire anvil. For all spectra, 64 scans at a nominal spectral resolution of 4 cm⁻¹ were applied. For evaluation, the GRAMS/AI software was used.

3 Results and discussion

3.1 Characterisation of clopamide anhydrate LH (1) and clopamide hemihydrate LH·0.5H₂O (2) crystals

Screening various crystallization conditions (polar and apolar solvents, temperature *etc.*), the clopamide (LH) molecule was found to form two different types of crystal structures: a solvent-free (**1**) and a hemihydrate form (**2**). Selected crystal data and parameters of structure refinement of crystals **1** and **2** are collected in Table 2, while more details can be found in Table S1.† Single crystals of **1** of acceptable quality suitable for SXRD measurements were obtained during a complex formation study. Although the colour of the brown copper complex could be observed in the methanol solution at room temperature, cooling to 5 °C and evaporation of the solvent



Table 2 Selected crystal data and structure refinement of crystals **1** and **2**

	(1)	(2)
Formula	LH	LH·0.5H ₂ O
Shape/colour	Needle/colourless	Prism/colourless
Crystal system	Monoclinic	Monoclinic
Space group	<i>P</i> 2 ₁ / <i>n</i>	<i>C</i> 2/ <i>c</i>
<i>a</i> (Å)	18.7914(4)	14.5166(8)
<i>b</i> (Å)	9.2921(4)	9.3747(7)
<i>c</i> (Å)	19.0621(4)	24.7064(17)
β (°)	99.322(7)	100.965(2)
<i>V</i> (Å ³)	3284.51(18)	3300.9(4)
<i>Z</i> / <i>Z'</i>	8/2	4/1
<i>D</i> _x (Mg m ⁻³)	1.399	1.428
Temperature (K)	128(2)	103(2)
Data/restraints/parameters	5936/0/401	3762/0/210
<i>R</i> _{int}	0.0740	0.0564
Final <i>R</i> indices [<i>I</i> > 2σ(<i>I</i>)]	0.0998, 0.2229	0.0440, 0.1076
<i>R</i> indices (all data)	0.1199, 0.2315	0.0510, 0.1122
Max. and mean shift/esd	0.000; 0.000	0.000; 0.000
Δρ _{max} , Δρ _{min} (e Å ⁻³)	0.617, -0.494	0.376, -0.412

resulted first in colourless needle-shaped crystals having the lowest solubility and identified as the crystals of clopamide anhydrate (**1**), then the blue CuSO₄·5H₂O crystallized. The colour changes of the solution during the crystallization processes can be followed in Fig. S1†. All other attempts, although they were numerous, to produce single crystals of **1** failed. **1** crystallizes in the monoclinic *P*2₁/*n* space group with two clopamide molecules in the asymmetric unit (Fig. 1a). The hemihydrate crystals of **2** were obtained from methanol, although clopamide dissolves poorly in methanol, even if it is heated. However, in the presence of some water the solubility increases. Clopamide hemihydrate crystallizes in the monoclinic *C*2/*c* space group. The asymmetric unit contains one clopamide molecule and one disordered water

molecule (with site occupancy of 0.5, Fig. 1b). The void size is large enough (61.75 Å³) to allow two positions of the water molecule to be hydrogen bonded to two different clopamide molecules. The methyl groups of the 2,6-dimethylpiperidine moiety have a larger space available for motions due to the presence of solvent cavities in **2**; this results in more elongated ellipsoids, shown on the ORTEP diagram in Fig. 1b. This hemihydrate form occurred during the different crystallization experiments using different non-absolute organic solvents such as methanol, ethanol, 1-propanol and acetone. Clopamide is poorly soluble in water, but it can be dissolved by the addition of some base (e.g., NaOH, diethylamine) and then it crystallizes as a hemihydrate. The hemihydrate form is present in medicaments, for example in Brinaldix tablets. The X-ray powder diffractogram of the clopamide sample used in the crystallization experiments agrees well with the calculated pattern based on the clopamide hemihydrate structure **2** (Fig. S2 and S3†).

In order to compare the two different conformations of molecules A and B in crystal **1** with that of molecule C determined in crystal **2**, conformational analysis was performed. The overlaid molecules are shown in Fig. 2. The root-mean-square-deviation (rmsd) and maximum atomic distances (maxD) were calculated using Mercury and found to be 0.0728 Å and 0.1542 Å for A and B, 0.1137 Å and 0.2450 Å for A and C and 0.0936 Å and 0.2699 Å for B and C. Based on the calculated values the best fit was found between molecules A and B, the two molecules in the asymmetric unit of **1**. The clopamide molecule C shows a slightly larger difference from both A and B molecules but still the conformation is very similar taking into account that the molecule contains three single bonds, N9–N8, N8–C7 and

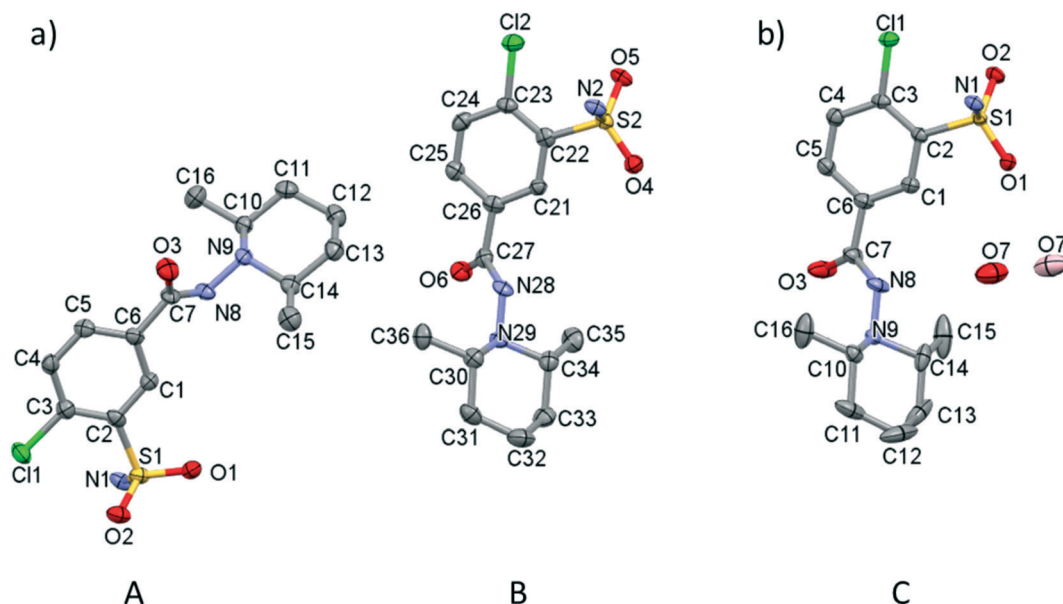


Fig. 1 (a) ORTEP presentation of the asymmetric unit of crystal **1** (A and B molecules) and (b) **2** (C molecule) with atom labelling. The asymmetric unit of crystal **2** contains one water molecule (O7) in two disordered positions with 0.5 site occupancy (coloured light and dark red). Displacement ellipsoids are drawn at 50% probability level. Hydrogen atoms are omitted for clarity.



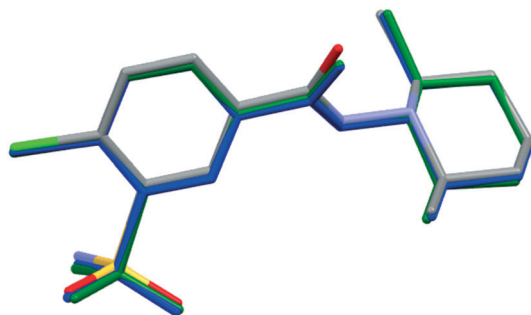


Fig. 2 Overlaid structures of clopamide molecules A (blue) and B (green) from crystal 1 and molecule C (coloured by elements) from crystal 2.

C6–C7, which allow high rotational freedom. A detailed comparison of the bond distances and angles can be found in Tables S2 and S3.†

The very similar systems of secondary interaction formed among the molecules in the crystal lattices result in the very similar molecular conformations in 1 and 2. The sulfonamide group on the clopamide molecule contains two =O and an –NH₂ group; consequently these molecules are able to form dimers *via* N–H···O type interactions. In both 1 and 2 crystal structures, isostructural planes of molecules are formed from these dimers by additional N–H···N, N–H···O and C–H···π interactions. Fig. 3a shows that the planes composed of molecules A (blue) and molecules B (green) are alternating in 1. There are no hydrogen bonds between layer A and layer B. The distance between the layers formed by molecule A is 14.503(4) Å in crystal 1, while the analogous distance is somewhat larger, 14.517(8) Å in crystal 2 (Fig. 3b). The water molecules in structure 2 act as spacers keeping the layers more distant than the layers are in structure 1. Thus a more favourable molecular conformation and higher symmetry can be realised in 2. The pseudo twofold axis in the $P2_1/n$, $Z' = 2$ (1) structure becomes a real symmetry in 2, and the space group of $C2/c$ is realised with $Z' = 1$ (2), as $C2/c$ is the minimal non-isomorphic supergroup of $P2_1/n$ with a difference of a twofold axis.

An intramolecular halogen bond can be observed between the chlorine and the oxygen of the chloro-sulfonamoylbenzamide moiety (3.148(6) Å 1A, 3.096(4) Å 1B), while an intermolecular chalcogen bond is present between a sulfoxo oxygen as donor and the lone pair of the carbonyl group as acceptor ((3.036(7) Å 1A, 3.233(7) Å 1B)), showing a competition between the two types of supramolecular interactions in the two layers of the two crystallographically independent molecules in 1. Although the chlorine and the sulfonamide oxygen come closer (3.0581(16) Å C) in 2, there is no chance to form a chalcogen bond because of the presence of the water molecule, which forms a bridge between the neighbouring molecules. In the absence of the chalcogen bond in 2, the intramolecular halogen bond becomes the shortest here.

The distances and angles of selected interactions in crystals 1 and 2 are collected in Table S4.† Comparisons of the unit cells from different crystallographic views and the packing arrangements are shown in Fig. S4–S6.†

3.2 Characterization of solvatomorph $[\text{CuL}_2] \cdot 2(\text{MeOH}) \cdot 2(\text{H}_2\text{O})$ (3), $[\text{CuL}_2] \cdot 2(\text{EtOH}) \cdot 2(\text{H}_2\text{O})$ (4), $[\text{CuL}_2] \cdot 2(n\text{PrOH}) \cdot 2(\text{H}_2\text{O})$ (5) and $[\text{CuL}_2] \cdot 2(i\text{PrOH}) \cdot 2(\text{H}_2\text{O})$ (6) complex crystals

The outcome of crystallization experiments performed to obtain copper(II) complexes of clopamide is summarized in Scheme 2. At pH ~9 the hydrazine N8 nitrogen deprotonates along with the increase in electron density on the N9 atom, and as a result the ligand is able to coordinate to the metal ion bidentately *via* N9 and the neighbouring carbonyl O3 atom. The bis-ligand copper(II) complexes $[\text{CuL}_2]$ could be obtained from different water-miscible and -immiscible solvents, producing solvatomorphic (3–6 and 10) and solvent-free polymorphic (7–9) crystals (Scheme 2).

Solvatomorphic crystals of 3–6 were obtained by using a homologous series of alcohols: methanol, ethanol, 1-propanol and 2-propanol. All four solvatomorphs crystallize in the $P\bar{1}$ space group with similar unit cell parameters and gradually increasing unit cell volume. Selected crystal data and parameters of structure refinement of crystals 3–6 are

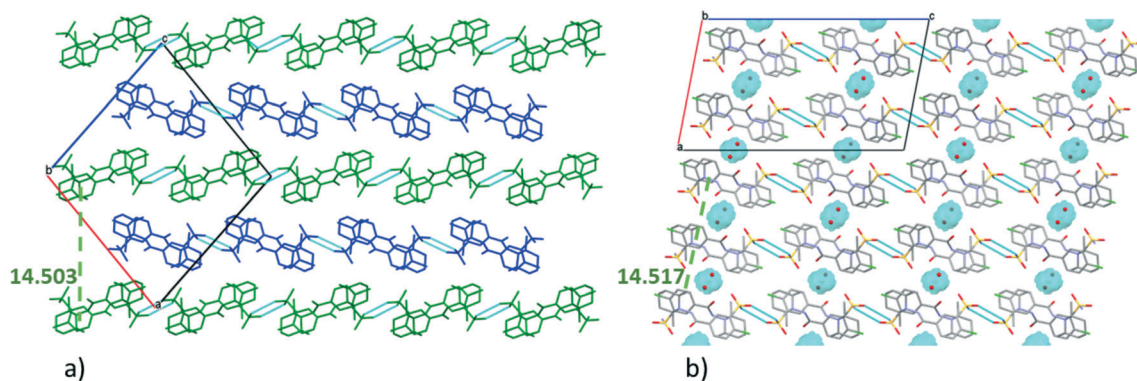


Fig. 3 Packing arrangement of molecules in the crystal lattice of 1 (a) and 2 (b) viewed from the crystallographic *b* axis. Molecules are coloured A (blue) and B (green) in crystal 1. The cavities containing the water molecules are highlighted in blue in crystal 2.



Table 3 Crystal data and refinement parameters of crystals 3–6

	3	4	5	6
Moiety formula	[CuL ₂] $\cdot$ 2MeOH $\cdot$ 2H ₂ O	[CuL ₂] $\cdot$ EtOH $\cdot$ 2H ₂ O	[CuL ₂] $\cdot$ 2PrOH $\cdot$ 2H ₂ O	[CuL ₂] $\cdot$ 2iPrOH $\cdot$ 2H ₂ O
Shape/colour	Prism/brown	Prism/brown	Prism/brown	Prism/brown
<i>M_r</i>	853.32	881.37	909.42	909.42
Crystal system	Triclinic	Triclinic	Triclinic	Triclinic
Space group	<i>P</i> 1	<i>P</i> 1	<i>P</i> 1	<i>P</i> 1
<i>Z</i> , <i>Z'</i>	1, 0.5	1, 0.5	1, 0.5	1, 0.5
<i>a</i> (Å)	7.7518(3)	8.2195(7)	8.404(4)	8.3107(14)
<i>b</i> (Å)	8.0901(4)	8.2439(7)	8.497(5)	8.7469(14)
<i>c</i> (Å)	16.3683(7)	16.4822(14)	16.310(8)	16.387(3)
α (°)	83.1130(10)	85.598(2)	79.445(10)	79.713(5)
β (°)	78.1130(10)	78.212(2)	83.113(12)	87.099(4)
γ (°)	71.6590(10)	68.273(2)	67.366(10)	65.629(4)
<i>V</i> (Å ³)	953.96(7)	1015.62(15)	1055.2(10)	1067.3(3)
<i>D_x</i> (Mg m ^{−3})	1.485	1.441	1.431	1.415
Temperature (K)	103(2)	103(2)	103(2)	103(2)
Data/restraints/parameters	6813/0/236	4634/0/244	3846/0/253	5523/0/368
<i>R</i> _{int}	0.0600	0.0918	0.1536	0.0884
Final <i>R</i> indices [<i>I</i> > 2σ(<i>I</i>)] <i>R</i> ₁ , <i>wR</i> ₂	0.0472, 0.1021	0.0972, 0.2386	0.0849, 0.2092	0.0690, 0.1530
<i>R</i> indices (all data), <i>R</i> ₁ , <i>wR</i> ₂	0.0684, 0.1284	0.1179, 0.2561	0.1143, 0.2328	0.0816, 0.1599
Max. and mean shift/esd	0.000; 0.000	0.000; 0.000	0.000; 0.000	0.000; 0.000
$\Delta\rho_{\text{max}}$, $\Delta\rho_{\text{min}}$ (e Å ^{−3})	0.832, −0.888	1.573, −1.098	0.852, −0.963	0.644, −0.708

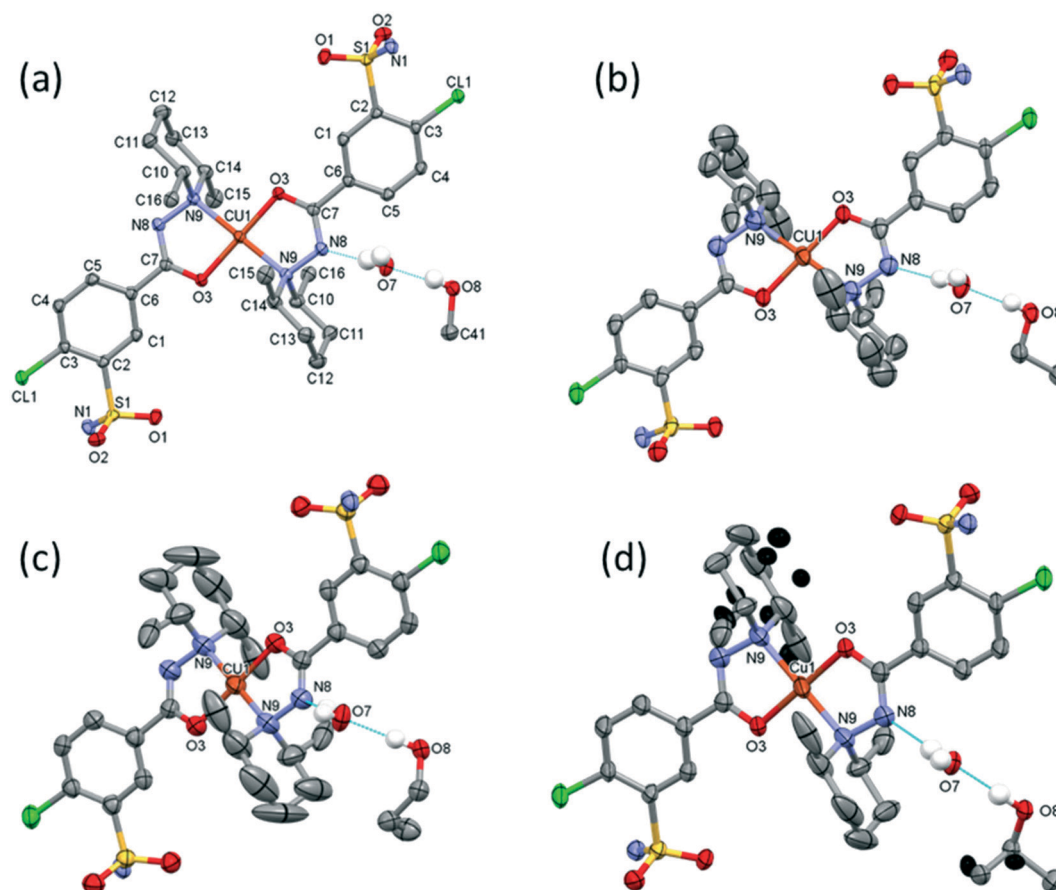


Fig. 4 Molecular structure of the homologous series of solvatomorphic complexes of **3** (a), **4** (b), **5** (c) and **6** (d). Hydrogen atoms involved in H bonds are shown; all other ones are omitted for clarity. Atomic displacement parameters are presented at 50% probability level. One ligand is in the asymmetrical unit, the copper(II) ion is on the symmetry centre, and the other parts of the complexes are generated by inversion symmetry. Solvent molecules are shown on one side only for clarity. Labels of atoms are fully listed for crystal **3**; the same was used for **4–6**. The increasing displacements of the piperidines' carbon atoms can be observed with the increasing size of the solvent molecule. The disordered atoms of **4** are marked with black ellipsoids in crystal **6**.



collected in Table 3, while more details can be found in Table S5.† All four crystals contain half of the $[\text{CuL}_2]$ complex, one alcohol and one water molecule in their asymmetric unit (Fig. 4). The copper(II) ion is coordinated by the N9 and O3 donor atoms of the ligand, forming a five-membered chelate ring in square planar geometry (Table S6†). The two ligands coordinate symmetrically in *trans* positions as the copper ion is located on an inversion centre. The complex formation reduces the strength of the intramolecular halogen bond, and the $\text{Cl}\cdots\text{O}$ distance increases to 3.208(2) Å in 3, 3.233(4) Å in 4, 3.189(5) Å in 5, and 3.204(3) Å in 6. The ORTEP diagrams show that the atomic displacement parameters presented by ellipsoids are growing from 3 to 6 with the increasing size of the solvent void, reflecting that the degree of freedom becomes higher especially for the piperidine rings of the hosts. In crystal 6 the piperidine ring could be split into two distinguishable disordered positions (occupancy: 2/3 : 1/3) (Fig. 4).

The $[\text{CuL}_2]$ complexes connected by alcohol and water molecules simultaneously in similar positions resulted in isostructural crystal structures, all in the $P\bar{1}$ space group. The cell similarity indexes ($I = |(a + b + c)/(a' + b' + c') - 1|$) were found to be between 0.01 and 0.04 for the pairs of crystals (see Table S7†), which proves their very high similarity. The conformations of the molecules together with the similar solvent positions are compared in Fig. S7†. The comparison of packing arrangements is shown in Fig. 5 and S8†. The copper–copper distances between neighbouring complexes increase in the case of methanol 7.752(1) Å, ethanol 8.220(1) Å, 1-propanol 8.497(5) Å and 2-propanol clathrates to 8.747(1) Å. The complex molecules form an isostructural three-dimensional network connected by the solvent molecules in all structures from 3 to 6.

The solvent molecules stabilize the crystal lattice similarly in all methanol (3), ethanol (4), 1-propanol (5), and 2-propanol (6) containing crystals, forming secondary interactions on both sides of the complex molecules: the alcohol oxygen atom O8 is an acceptor of adjacent sulfonamide NH_2 protons with $\text{N1-H1NA}\cdots\text{O8}$ and $\text{N1-H1NB}\cdots\text{O8}$, and the alcohol proton connects to the neighbouring water oxygen atoms with $\text{O8-H1O8}\cdots\text{O7}$. A water proton forms a H bond with the deprotonated ligand nitrogen $\text{O7-H2O7}\cdots\text{N8}$, while the other one connects to sulfonamide oxygen $\text{O7-H1O7}\cdots\text{O2}$ (Fig. 6). The details of H-bond interactions are collected in Table S8.† The distances between $\text{N8}\cdots\text{O7}$ and $\text{O7}\cdots\text{O8}$ atoms show no significant increase with the increasing size of the solvent molecules or with the increasing size of the available void volume (Table 4). As the volume of the unit cell and the volume of the void containing the solvent molecules increase in the order methanol, ethanol, 1-propanol and 2-propanol solvates from 3 to 6, the density of the crystals decreases, having an inverse relationship between the cell volume and the crystal density (Table 4 and Fig. 7).

Crystallization experiments have been performed with longer chain homologous alcohols but crystals could not be obtained from *n*-butanol and longer chain alkylalcohols (*n*-pentane *etc.*), which shows that the above presented crystal lattice can no longer accommodate larger size molecules. As the length of the *n*-alcohol increases in its homologous order, the molecular size increases, and the strength of the intermolecular interaction to the bridging water decreases, and thus the isostructural crystal series of the Cu-clopamide complex could not be completed with more members.

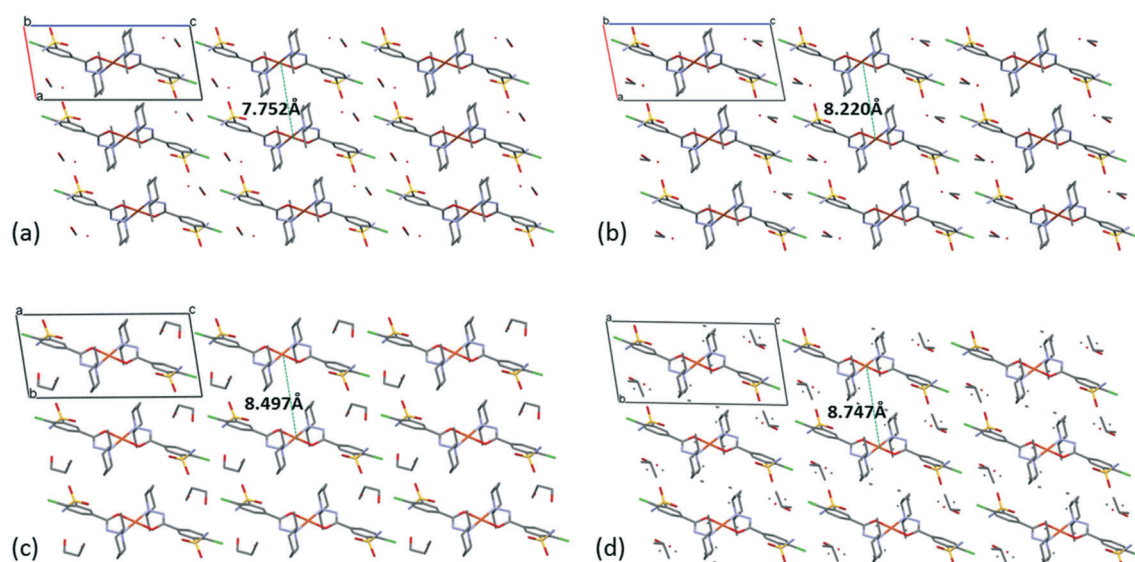


Fig. 5 Packing arrangements in crystal 3 (a), 4 (b), 5 (c) and 6 (d) viewed from the crystallographic *b* axis, presenting also the increasing distances of the metal Cu(II) centres with the increasing size of the solvent guest molecules.



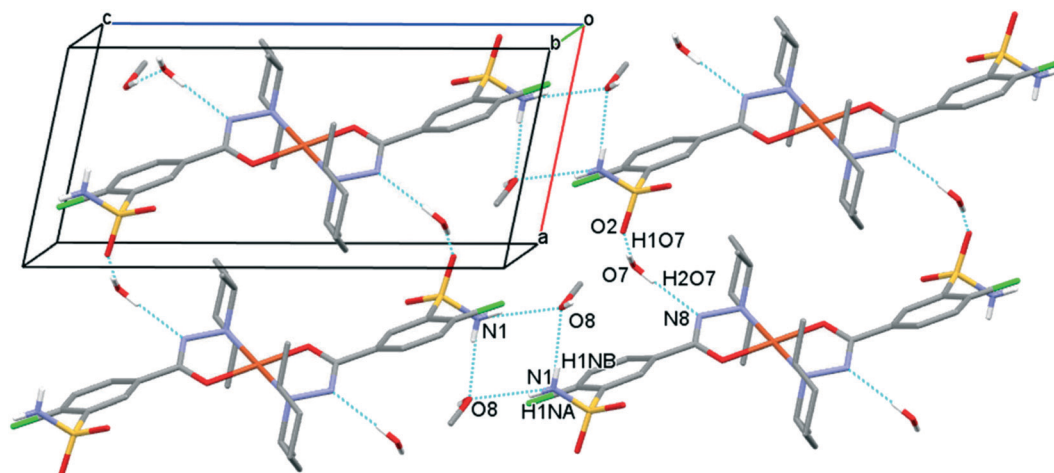


Fig. 6 Packing arrangements in crystal **3** showing the most important H-bond interactions. Similar synthons can be found in crystals **4**, **5** and **6**. Hydrogen atoms not involved in the H-bond system are omitted for clarity.

Table 4 Calculated structural parameters of crystals **3–6**

Crystal number	V_{cell} ($\text{\AA}^3$)	V_{void} ($\text{\AA}^3$)	$V_{\text{void}}/V_{\text{cell}}$ (%)	Density (g cm^{-3})	d ($\text{N8}\cdots\text{O7}$) ($\text{\AA}$)	d ($\text{O7}\cdots\text{O8}$) ($\text{\AA}$)
3	953.96(7)	117.56	12.3	1.485	2.845(3)	2.643(2)
4	1015.62(15)	158.00	15.6	1.441	2.857(7)	2.673(6)
5	1055.2(10)	202.08	19.2	1.431	2.846(6)	2.626(6)
6	1067.3(3)	193.73	18.2	1.415	2.882(4)	2.707(4)

3.3 Characterization of the polymorphic solvent-free $[\text{CuL}_2]$ (**7–9**) and $[\text{CuL}_2]\cdot\text{CH}_2\text{Cl}_2$ (**10**) complex crystals

Solvent-free $[\text{CuL}_2]$ crystals (Fig. 8a–c) were obtained from methanol solution using CuSO_4 instead of CuCl_2 salt. Use of CuSO_4 in the complexation reaction with clopamide resulted in the brown colour of the methanol solution, unlike when using CuCl_2 , where the colour of the solution was blue. Two polymorphic solvent-free $[\text{CuL}_2]$ crystals were obtained from methanol solution: crystal **7** was harvested from a solution containing a slight ligand excess (10:9) and crystal **8** was harvested from a solution with a strong ligand excess (7:2), though the composition of the obtained complexes is the same. From water-immiscible organic solvents (e.g. ethyl acetate) another solvent-free polymorph, **9**, was crystallized. Crystals **7** and **8** were found to crystallize in the triclinic

crystal system in the $P\bar{1}$ space group, while crystal **9** crystallizes in the trigonal crystal system $R\bar{3}$ space group.

Nevertheless, from dichloromethane solution crystal **10** (Fig. 8d) was obtained (besides **9**) with the inclusion of a CH_2Cl_2 guest molecule in a 1:1 complex-to-solvent ratio. **10** crystallizes in the monoclinic crystal system in the $C2/c$ space group. Selected crystallographic parameters are collected in Table 5, while detailed crystallographic data are given in Table S9.† Packing arrangements of crystals **7–10** are shown in Fig. S9.† In all cases a square planar coordination geometry with *trans* orientation of the two clopamide molecules was found, similarly to crystals **3–6**; however, significant differences can be seen in the conformation of the piperidine ring and the packing arrangement of the complexes in the crystals.

The main conformational differences of the complexes in their different crystals **6–10** are shown in Fig. 9. The root-mean-square deviation (rmsd) and maximum atomic distance (maxD) values were calculated using Mercury and are compared in Tables S10 and S11.† The coordination spheres of the central Cu(II) cations are highly similar in all crystals; however, the conformation of the ligands may differ mainly because of two reasons: (1) the six-membered piperidine ring in chair conformation can be placed in two different ways, it can be parallel or perpendicular to the Cu-N9 bond and (2) the phenyl ring may be at different angles to the coordination plane defined by atoms Cu1 , O3 , C7 , N8 , and N9 , according to different secondary interactions of the sulfonamide group in the different crystal lattices. These two characteristic values are compared in Table 6.

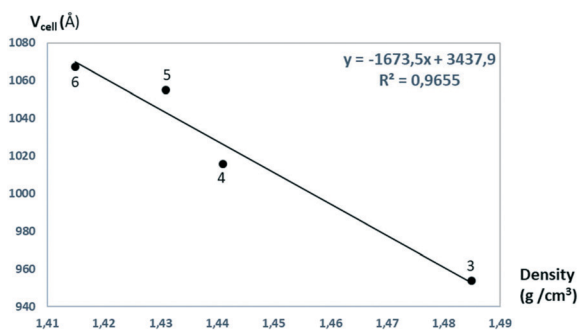


Fig. 7 The unit cell volumes are plotted as a function of the calculated densities obtained for crystals **3–6**.



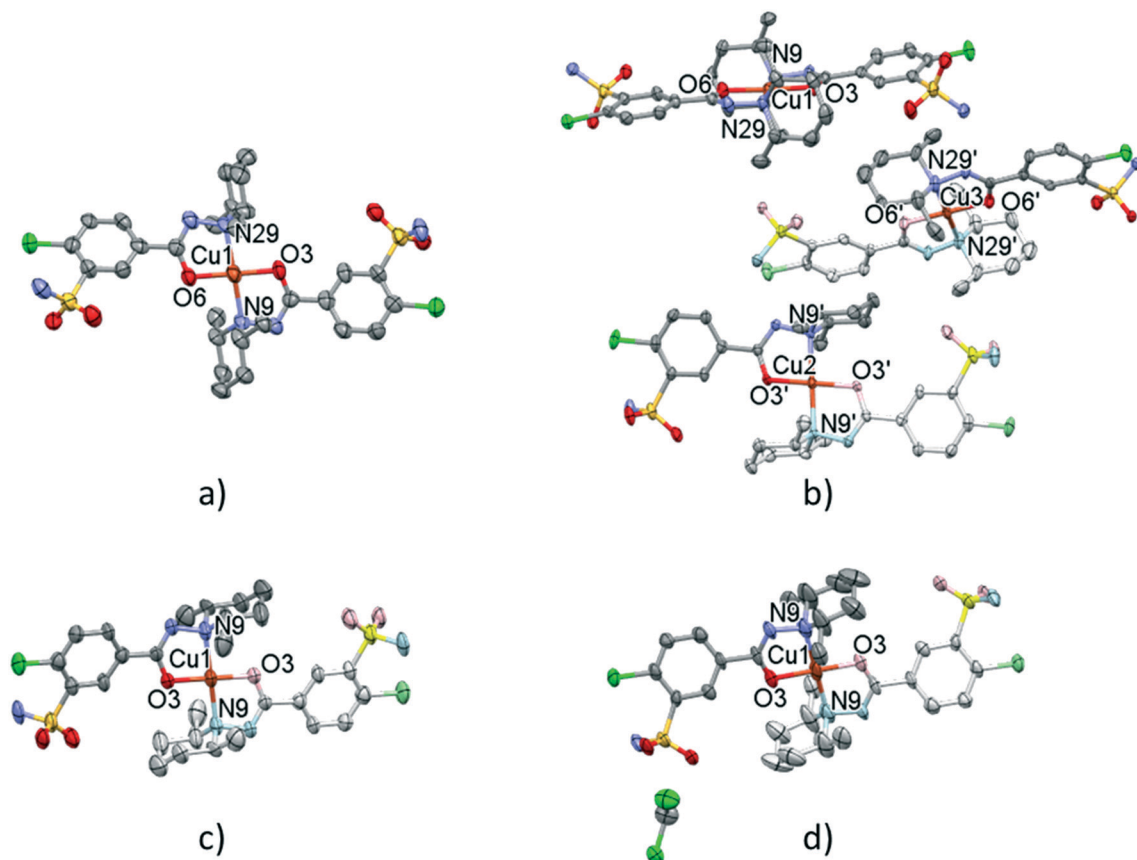


Fig. 8 Molecular structure of the solvent-free polymorphic complexes of $[\text{CuL}_2]$ **7** (a), **8** (b), and **9** (c). Molecular structure of the solvate crystal $[\text{CuL}_2] \cdot \text{CH}_2\text{Cl}_2$ **10** (d). Hydrogen atoms are omitted for clarity, atomic displacement parameters are presented at 50% probability level. The light-coloured atoms of the molecules are generated by inversion symmetry (full labelling can be found in Fig. S10 and S11†).

Table 5 Selected crystal data and refinement parameters of crystals **7–10**

	7	8	9	10
Moiety formula	$[\text{CuL}_2]$	$[\text{CuL}_2]$	$[\text{CuL}_2]$	$[\text{CuL}_2] \cdot \text{CH}_2\text{Cl}_2$
Shape/colour	Chunk/brown	Chunk/brown	Chunk/brown	Chunk/prism
M_r	753.20	753.20	753.20	923.05
Crystal system	Triclinic	Triclinic	Trigonal	Monoclinic
Space group	$P\bar{1}$	$P\bar{1}$	$R\bar{3}$	$C2/c$
a (Å)	7.8247(14)	13.4600(10)	23.078(3)	26.699(2)
b (Å)	13.829(2)	15.2424(11)	23.078(3)	9.8941(8)
c (Å)	15.156(3)	18.1855(16)	18.3291(12)	15.6876(14)
α (°)	78.981(4)	114.627(6)	90	90
β (°)	81.309(4)	93.362(8)	90	104.750(7)
γ (°)	85.041(4)	90.021(6)	120	90
V (Å ³)	1588.5(5)	3384.5(5)	8454(2)	4007.5(6)
Z , Z'	2, 1	4, 1 + 0.5 + 0.5	9, 0.5	4, 0.5
D_x (Mg m ⁻³)	1.575	1.478	1.331	1.530
Temperature (K)	103(2)	123(2)	153(2)	128(2)
Data/restraints/parameters	5798/0/410	12 256/0/822	3429/0/207	3661/0/234
R_{int}	0.1852	0.1301	0.1176	0.0901
Final R indices $[I > 2\sigma(I)]$, R_1 , wR_2	0.1495, 0.3073	0.0777, 0.1198	0.1033, 0.1672	0.1231, 0.2351
R indices (all data), R_1 , wR_2	0.2415, 0.3583	0.1375, 0.1366	0.2142, 0.2052	0.1623, 0.2539
Max. and mean shift/esd	0.000; 0.000	0.001; 0.000	0.000; 0.000	0.000; 0.000
$\Delta\rho_{\text{max}}$, $\Delta\rho_{\text{min}}$ (e Å ⁻³)	1.205, -0.647	0.564, -0.576	0.553, -0.264	0.887, -0.619

The conformations of the complexes in **3–6** are very similar; the two ligands coordinate symmetrically to the copper ion owing to an inversion centre. The piperidine ring

is parallel to the Cu–N9 binding axes and the angle between the coordination plane and the phenyl ring was found to be between 31.27(10)° and 35.60(18)°. Interestingly, in crystal **6**



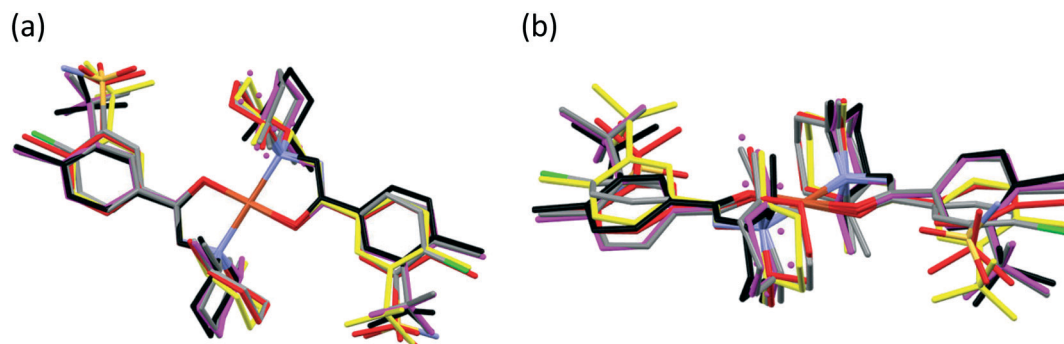


Fig. 9 Top view (a) and side view (b) of selected $[\text{CuL}_2]$ complexes overlaid by the coordinating atoms of clopamide and the central metal ion. Colour code: **6** (magenta), **7** (black), molecule A in **8** (red), **9** (coloured by element) and **10** (yellow). Solvents and hydrogen atoms are omitted for clarity.

Table 6 Angles ($^\circ$) of the plane formed by the coordinating atoms to the metal centre and the phenyl ring as well as the geometry of the piperidine ring in chair conformation to the Cu–N9 bond in crystals **3–10**

Crystal number	Angle between plane1 and plane2 ^a ($^\circ$)	Piperidine ring placement to Cu–N9 ^b
3	31.27(10)	Parallel
4	34.9(3)	Parallel
5	34.6(3)	Parallel
6	35.60 (18)	Parallel in major disordered positions Perpendicular in minor disordered positions
7	21.9(6) and 37.2 (6)	Parallel
8A	13.3(3) and 9.6 (3)	Perpendicular
8B	15.5(2)	Perpendicular
8C	3.8(3)	Perpendicular
9	13.7(4)	Perpendicular
10	23.9(4)	Perpendicular

^a Plane1 is the coordination plane defined by atoms Cu1, O3, C7, N8, and N9 atoms. Plane2 is the plane of the phenyl ring. ^b The piperidine ring in chair conformation can be placed parallel or perpendicular to the Cu–N9 bond.

the disordered positions of the piperidine ring appeared in both parallel (occupancy 2/3) and perpendicular placement (occupancy 1/3) to the Cu–N9 axis. In the case of **7** (Fig. 8a) the inversion symmetry is missing and the conformation of the two ligands is different. The angles between the phenyl rings and the coordination plane are 21.9(6) $^\circ$ and 37.2(6) $^\circ$, respectively, for the two ligands, and the piperidine rings are parallel to the Cu–N bond (Table 6). In crystal **8** with one asymmetrical $[\text{CuL}_2]$ and two different symmetrical $[\text{CuL}_2]$ complexes (1 + 0.5 + 0.5) in the asymmetric unit (Fig. 8b), in all three complexes the piperidine ring has been found perpendicular to the Cu–N bond, and the difference between them is primarily originating from the conformation of the phenyl ring, which predicts that their sulfonamide groups are likely to form different types of secondary interactions in this crystal (see also the overlaid structures in Fig. S12[†]; H-bond interactions are presented in Fig. S13[†]). There is half of the complex $[\text{CuL}_2]$ in the asymmetric unit in **9** and **10**, and an inversion centre is found where the metal ion is present; thus the two ligands are coordinated symmetrically. Similar to **8** the piperidine ring conformation is perpendicular to the Cu–N bond and differences in the phenyl ring angles have been observed. Comparing the distances and angles of the coordination spheres (Table S6[†]), it is noticeable that Cu–O coordination bonds are

significantly shorter, while Cu–N bonds are longer in the complexes in crystals **8**, **9** and **10** compared to the solvatomorphic crystals **3–6** or **7** which coincides with the perpendicular (shorter Cu–O) or parallel (longer Cu–O) conformation of the piperidine ring. The perpendicular position of the piperidine ring is stabilized by C–H $\cdots$ O intramolecular interactions between C13–H13A $\cdots$ O3 and C11–H11B $\cdots$ O3 (Table S12[†]). Although there are S–N–H $\cdots$ O=S intermolecular interactions formed around the inversion centres in **7**, **8** and **9**, like in the pure clopamide crystals of **1** and **2**, the N–H $\cdots$ Cl intramolecular hydrogen bonds become more important in the complexes and no halogen intramolecular interactions are observed.

Fig. 10 presents the similar arrangement of the molecules in the crystal lattices of **3** and **7**. While the sulfonamide groups connect two columns through methanol solvent molecules (with N–H $\cdots$ O and O–H $\cdots$ S bonds) in **3**, in the solvent-free form **7** the complexes are directly connected to each other through N–H $\cdots$ O bonds. An asymmetrical intermolecular interaction between a sulfonamide NH₂ and the deprotonated N8 hydrazine nitrogen atom was found among neighbouring complexes, which causes the asymmetric coordination of the two ligands; therefore there is the whole complex molecule in the asymmetric unit in **7**. Details of the H-bond interactions are collected in Table



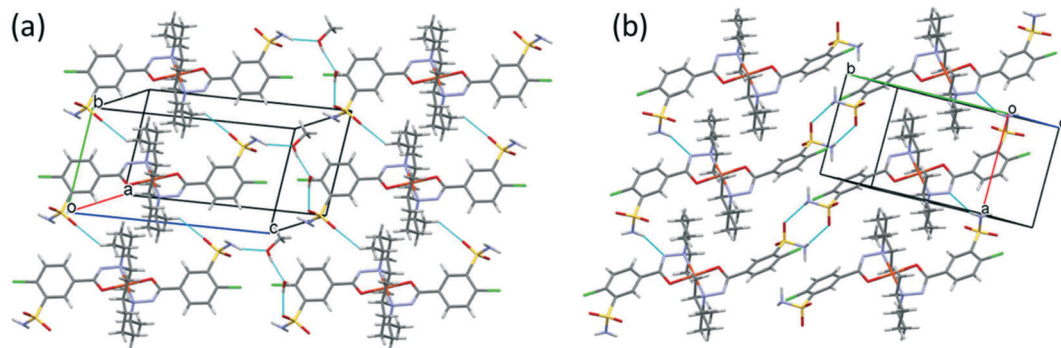


Fig. 10 Analogous packing arrangements in [CuL₂]·CH₃OH·H₂O crystal **3** (a) and in the solvent-free [CuL₂] crystal **7** (b), showing the columns supported by the systems of secondary interactions.

S12.† In crystals **8** and **9** the intermolecular sulfonamide N–H⋯O bonds generate linear columns of the complexes and adjacent columns then turn in a way that the angles between the columns of complexes are 62.7° in the case of **8** and 60° in the case of **9** (Fig. 11).

In crystal **8** the three different [CuL₂] complexes from the asymmetric unit (coloured red, blue and green) are arranged in an approximate triangle (Fig. 11a). This becomes a perfect threefold symmetry in **9**; there is one half complex molecule in the asymmetric unit in space group R3̄ (Fig. 11b). This

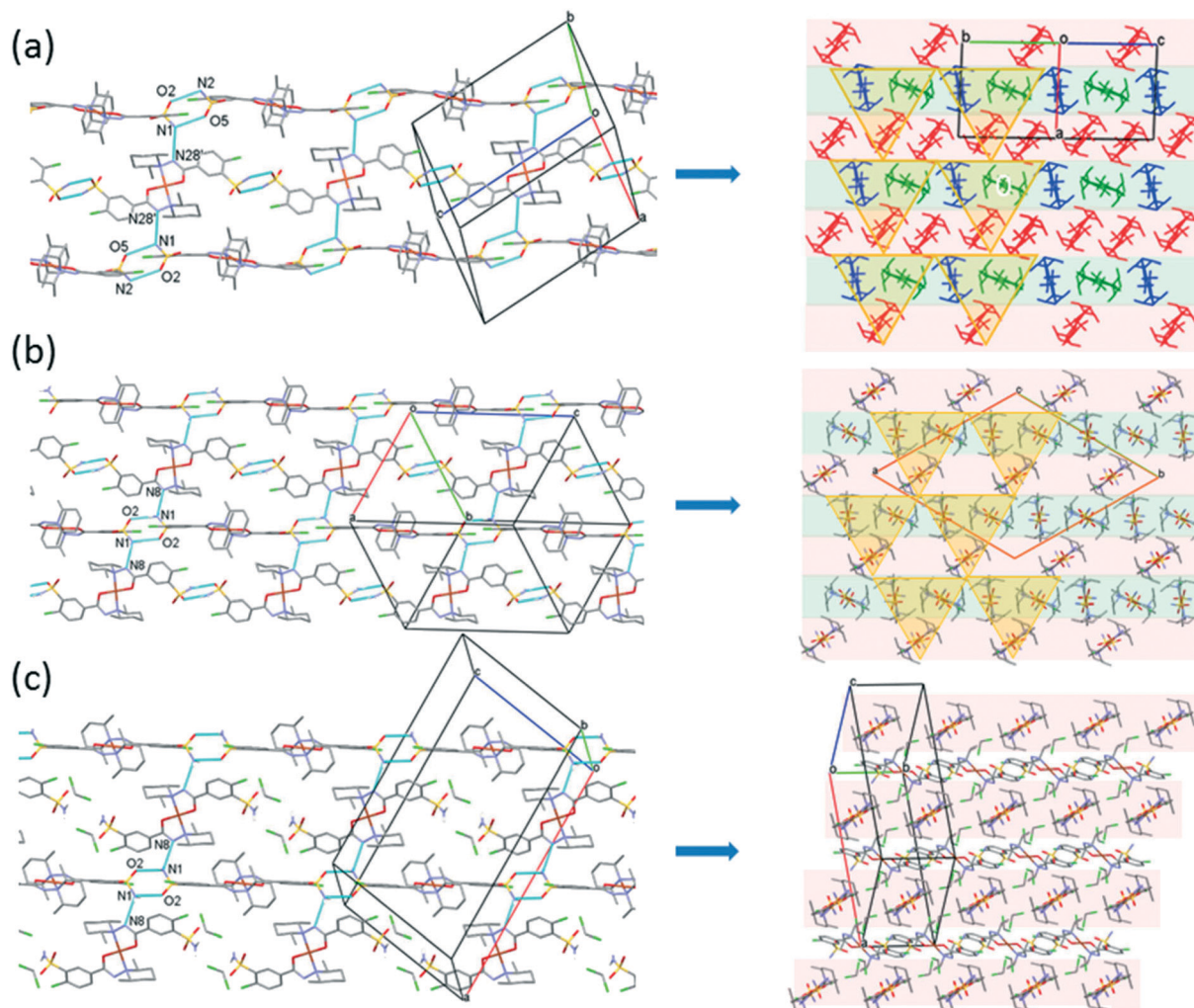
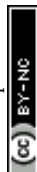


Fig. 11 Packing arrangements in crystal **8** (a) (molecule A is red, B is green and C is blue), crystal **9** (b) and crystal **10** (c) showing the columns formed by N–H⋯O type hydrogen bonds viewed from the side and also from the top of the columns. The light pink and green colours show the different sheets; the yellow triangles highlight the triangular arrangements of the molecules in the different crystals.



molecular arrangement creates channels; the void size calculated by Mercury (probe radius 1.2 Å, approximate grid spacing 0.7 Å) is 1452.26 Å³ (16.8% of the unit cell volume). Solvent inclusion (residual electron density) could not be located during refinement. This type of complex crystal **9** crystallized from different water-immiscible organic solvents (*e.g.* ethyl acetate, 1-butanol, 1-pentanol, 1-hexanol, toluene, chloroform, *etc.*) and no solvent was found in either case in the channels. Small molecules like water would however fit into the channels by size, but the piperidine moieties create a hydrophobic surface of the channel. A similar Cu(II) complex, the structure (tetrakis(μ^2 -vanillato-*O,O'*)-bis(nicotinamide-*N*)-di-copper(II)), was found in the literature (Ref. code DOHWON) having even larger empty channels.²⁸ TG-DSC-MSEGA coupled measurements have been performed on the crystals of **9** (Fig. 12), and around 3% solvent mixture (ethyl acetate and 1-butanol) loss was observed up to 224 °C without the collapse of the lattice. No structural change was observed by SXRD in the experiment when a single crystal of **9** was heated at 200 °C for an hour. The sample melts at 240.4 °C. Comparing the calculated crystal density of the three polymorphs it decreases in the

order **7** (1.575 g cm⁻³), **8** (1.478 g cm⁻³) and **9** (1.331 g cm⁻³), with **9** being the lowest one.

From the DCM solution, besides crystal **9** prismatic brown crystals of **10** could also be harvested in which the inclusion of one DCM molecule per asymmetric unit has been found (Fig. 8). We have seen that the conformation of the [CuL₂] complex in this structure is very similar to that of **9** (Fig. 9 and S14†). The secondary interactions of the DCM molecules in the lattice are much weaker in **10** compared to the alcohol-containing complex crystals of **3–6**. Crystal **10** decomposes rapidly when it is removed from its mother liquor. The packing of the molecules resembles that of crystal **9** as columns are formed by the sulfonamide connections; however, the angle between the Cu coordination planes of neighbouring complexes is much higher (80.7°) (Fig. 11c). The channels occurring in crystals **9** and **10** are compared in Fig. 13. The void volume occupied by the DCM molecules has grown from the empty 16.8% in **9** to 21.4% of the unit cell volume in **10**.

The difference in the conformation of clopamide in its pure crystal (**2**) and as a ligand in the complex crystal (**3**) is shown in Fig. 14. The sulfonamide group was found in the opposite direction in the complex as compared to the free

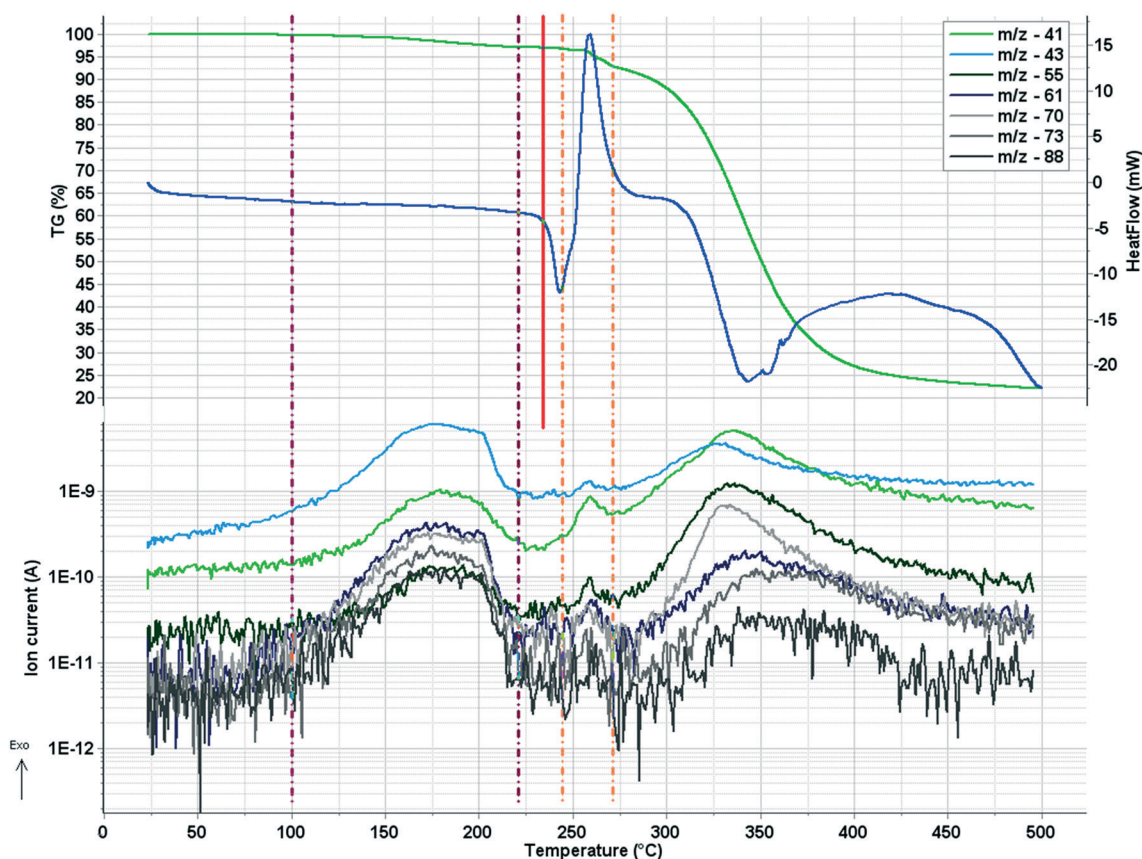


Fig. 12 TG-DSC-MSEGA diagram of sample **9**. On the upper part of the figure, the mass loss (TG – green curve) and heat flow (DSC – dark blue curve) signals are plotted, while on the lower part the ion current curves of some specific solvent fragments (ethyl acetate – *m/z* 88, 73, 70, 61, 43; 1-butanol – *m/z* 55 and 41) are plotted against temperature. The two vertical purple dash-dot lines mark the temperature interval, where the retained solvents (3%) are lost from the channels, while the red line shows the start of the melting of the sample. The orange dash-dot lines are marking a second small temperature interval, where traces of solvents are also lost and the decomposition starts.



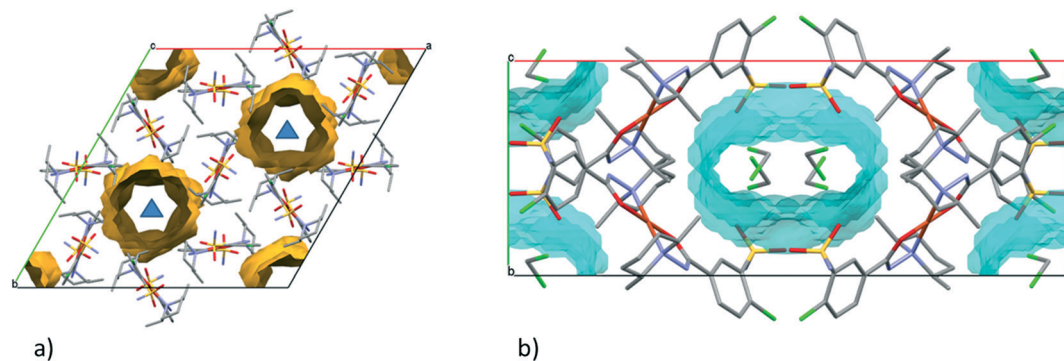


Fig. 13 (a) Empty channels in the crystal structure of **9** viewed from the crystallographic *c* axis. The three-fold axes are shown as blue triangles. (b) Unit cell of crystal **10** viewed from the crystallographic *c* axis. The channels in the crystal lattice (highlighted by cyan) are filled by DCM molecules which are surrounded by four complex molecules.

ligand crystals, as the chloro-sulfamoylbenzamide group turns around the flexible C6–C7 single bond. Although in solution the molecular freedom is higher, comparing the molecular conformations may provide insights into the reaction mechanism of complex formation.

3.4 Results of EPR spectroscopy measurements

In order to investigate the structure of the clopamide complex $[\text{CuL}_2]$ in solution, crystal **9** (a solvent-free polymorphic crystal form) was dissolved in pure DMSO in 3.0 mM concentration. The solution was poured in a quartz EPR tube and placed into a Dewar flask containing liquid nitrogen (77 K). The measured and simulated frozen solution spectra can be seen in Fig. 15. Depending on the stability of the complex when a crystal is dissolved, it may dissociate, thus the measured EPR spectrum is the superposition of the components appearing in the solution. When crystal **9** was dissolved in DMSO and EPR spectrum was measured for the frozen solution, the spectrum simulation showed the superposition of 80% bis-ligand and 20% mono-ligand complex. This means that in 20% of the complex, a ligand was released from the coordination sphere, resulting in the appearance of the mono-complex $[\text{CuL}]^+$. The well-resolved

superhyperfine line in the perpendicular range of the $[\text{CuL}_2]$ spectra originated from two nitrogen atoms coordinated to the copper ion. The *trans* position of the nitrogen atoms can be suggested based on the fact that magnetically equivalent couplings with the highest value in the *y* direction (referred to as the *g* tensor orientations) were obtained. The very low g_z (2.185) value and strong copper hyperfine coupling A_z (206 G) support the highly symmetrical square planar geometry of the complex with a high ligand field detected in the equatorial plane. In the case of the mono-complex, DMSO coordination beside one clopamide molecule in the equatorial plane can be presumed, and the higher g_z and lower A_z values reflect a lower ligand field compared to the bis-ligand complex. Nitrogen hyperfine couplings were not resolved for this complex.

3.5 Results of the infrared spectroscopy measurements

Based on the IR spectra, the stretching vibration band of the –OH group is missing from crystal **1**, while it appears at 3629

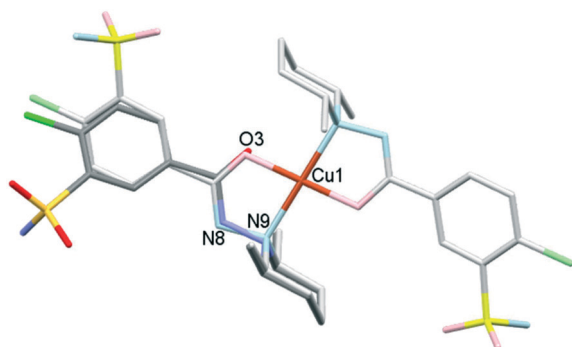


Fig. 14 Comparison of the conformations of the clopamide molecule in its hemihydrate crystal **2** and in its complex molecule in MeOH containing crystal **3**. The clopamide from **2** is shown in dark colours, while the complex from **3** is shown in light colours.

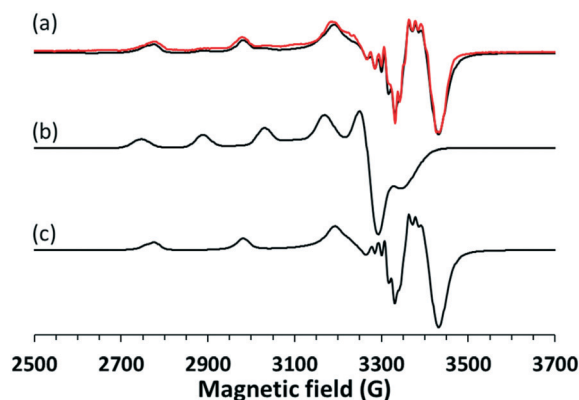


Fig. 15 (a) Experimental (black) and simulated (red) EPR spectra recorded at 77 K for crystal **9** dissolved in pure DMSO. The spectra were fitted with the superposition of (b) $[\text{CuL}]$ in 20% and (c) $[\text{CuL}_2]$ in 80%. The EPR parameters of $[\text{CuL}_2]$ are $g_x = 2.032$, $g_y = 2.047$, $g_z = 2.185$, $A_x = 33$ G, $A_y = 48$ G, $A_z = 206$ G and two nitrogens with $a_x = 5$ G, $a_y = 15$ G, $a_z = 5$ G, and of $[\text{CuL}]$ are $g_x = 2.060$, $g_y = 2.075$, $g_z = 2.279$, $A_x = 16$ G, $A_y = 54$ G, $A_z = 137$ G.



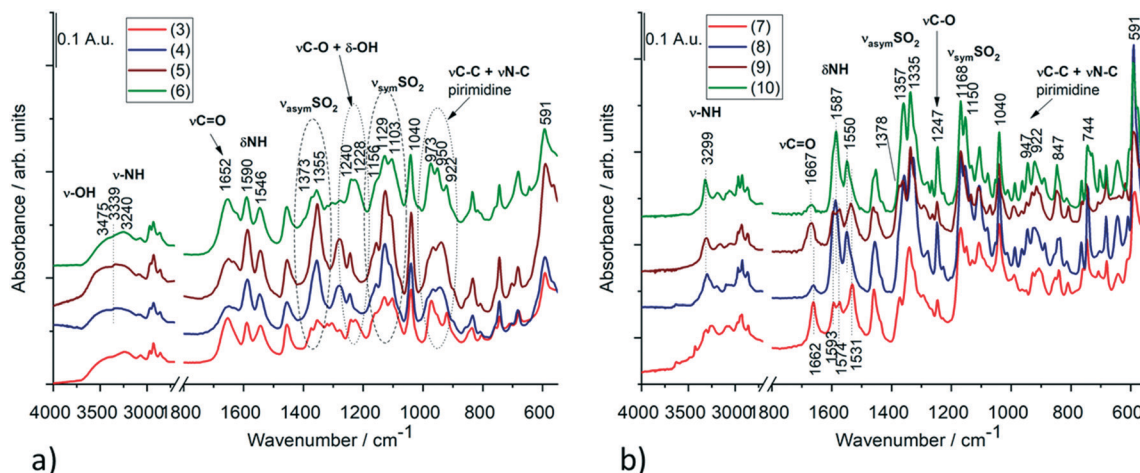


Fig. 16 IR spectra of (a) the solvatomorph crystals of **3**, **4**, **5** and **6** and (b) the solvent-free polymorphs (**7–9**) as well as the DCM containing crystal **10** of the complexes. Selected IR bands of these compounds with tentative assignments are presented in Tables S14 and S15.†

cm^{-1} in crystal **2** (Fig. S15 and Table S13†), which also shows a good agreement with the SXRD structures of the anhydrate and hemihydrate crystals of clopamide (Fig. S14†). In the case of the solvatomorph series, the shape (intensity and position) of the C=O and NH vibrational bands change in the complexes (Fig. 16a). SO_2 vibrations also alter; it is probably because of the violation of symmetry. In crystals **3**, **4**, **5** and **6**, the bands assigned to C–O and –OH vibrations are amplified due to the presence of bound alcohol molecules. Interestingly, samples **3–6** and **4–5** can be paired based on their spectra. The spectra of solvent-free polymorphs exhibit sharper bands (Fig. 16b), suggesting a more ordered crystal structure. This is further supported by the splitting of the more intense SO_2 stretching bands (Davydov splitting)²⁹ compared with the spectra of the solvatomorph series. It cannot be determined from the IR spectra how many molecules are present in a given unit cell; however, the appearance of Davydov splits in the spectra indicates a good agreement with the SXRD structures. In the case of crystal **10**, the bands of CH_2Cl_2 can be clearly seen (Fig. S16†). These extra bands do not appear in the other samples. A tentative assignment of the IR bands is presented in Tables S14 and S15.†^{30–33}

3.6 Results of the thermoanalytical measurements

The melting points of all crystalline samples were determined (Tables S1, S5 and S9†) by DSC. TG, DSC and DTG curves of **1** in inert atmosphere and in synthetic air atmosphere are presented in Fig. S17–S22.† The evolved sulfur dioxide is a very characteristic marker of the thermal degradation of clopamide, as the degradation of both the pure clopamide compound and its complexes starts at the $-\text{SO}_2\text{NH}_2$ group (Fig. S21†). The thermoanalytical results confirmed that 3% of water leaves crystal **2** up to 187.8 °C, followed by water leaving between 187.8 and 226.8 °C, with a weight loss of 26.7% accompanied by an endotherm peak. It means that a

large amount of water is strongly bound in the crystal lattice (Fig. S23 and S24†). For samples **7** and **8**, the mass spectrometric gas analysis results confirmed that no significant amounts of solvents were present in the crystals; these are solvent-free polymorphs (Fig. S25–S32†).

4 Conclusions

Clopamide ((4-chloro-*N*-2,6-dimethylpiperidin-1-yl)-3-sulfamoylbenzamide, LH) drug is used worldwide in the treatment of hypertension and oedema. It has also been shown that thiazide diuretics may induce changes in the level of copper in blood. Despite its medical application its structure has not yet been published, most probably because of the difficulties in single-crystal growth. We have successfully crystallized and determined the crystal structures of clopamide anhydrate and hemihydrate and a structural landscape of different polymorphs and solvatomorphs of its copper(II) complexes, revealing the molecular arrangements, supramolecular interactions and coordination properties of the complexes.

It is proved by PXRD that the drug Brinaldix is the hemihydrate form. As we were able to crystallize the anhydrate form of clopamide it is presented how the inclusion of water contributes to the crystal perfection of the drug (**1** and **2**). The water molecule acts as a spacer, keeping a distance between the molecular layers; thus a more favourable molecular conformation and higher symmetry of the space group can be realized.

The newly defined chalcogen bond is recognised in the clopamide anhydrate crystals being in competition with the intramolecular halogen bond present between the chlorine and the oxygen of the sulfonamide group. The intermolecular chalcogen bond is found between a sulfoxy oxygen as a donor and the lone pair of the carbonyl group as an acceptor. The longer the halogen bond, the shorter the chalcogen bond and *vice versa*. There is no chalcogen bond in **2** because the water



molecule acts as a bridge between the neighbouring molecules. In the absence of the chalcogen bond in **2**, the intramolecular halogen bond becomes the shortest.

All Cu(II) complexes have a square-planar coordination geometry, in which copper(II) centres are surrounded by piperidine-N and carbonyl-O donor atoms in a five-membered chelate ring. The two ligands are in *trans* positions. The 2,6-dimethylpiperidine units of the ligand molecules are found to be in parallel or in perpendicular positions to the Cu–N9 bond, which is in correspondence with the angle of the plane formed by the coordinating atoms to Cu(II) and the phenyl ring whose substituents form varying intermolecular interactions. The Cu–O coordination bonds are significantly shorter, while Cu–N bonds are longer in the complex crystals of **8**, **9** and **10** with perpendicular piperidine conformation compared to the solvatomorph crystals **3–6** or solvent-free **7** with parallel conformation of the piperidine ring.

The solvatomorph crystals, containing small homologous alcohols (**3–6**), have been found to be isostructural (*P*1) with increasing size of void and unit cell volumes which terminates using *n*-butanol. In the solvent-free polymorphic [CuL₂] crystals (**7–9**) the complex molecules are forming parallel columns strengthened by N–H⋯O interactions between ligand sulfonamide groups. The arrangement of these columns in **7** (*P*1) is similar to that of solvate **3** (*P*1), while **8** (*P*1) with its triangular arrangement transitions to **9** (*R*3), where perfect threefold symmetry is realised, although leaving large empty spaces in the lattice. Thermal investigations prove that there is no more than about 3% remaining solvent in the voids. The columnal structure is preserved, although the columns become farther away in the complex crystal (**10**), notwithstanding that the columns are no longer parallel and enclathrate dichloromethane solvent molecules in the crystal lattice.

The solvent content and thermal behaviour of the crystals were studied by IR and thermoanalytical methods, whose results are in good agreement with the structural results. The solution structure of the [CuL₂] complex was investigated in DMSO by EPR spectroscopy at 77 K. Upon dissolving the crystals they partly dissociate; 80% bis-ligand complex remained in the solution with a highly symmetrical square planar geometry but in 20% one ligand was released, resulting in the appearance of the mono complex [CuL]⁺ presuming the coordination of DMSO in the equatorial plane.

In addition to the structural description of a drug on the market, the systematic comparison of several polymorphic and solvatomorphic crystal arrangements provides valuable information to synthon and crystal engineering in order to develop new materials with specific properties.

Conflicts of interest

There are no conflicts to declare.

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