Journal of Materials Chemistry A



PAPER

View Article Online
View Journal | View Issue



Cite this: J. Mater. Chem. A, 2024, 12, 18544

Received 15th February 2024

Accepted 20th June 2024
DOI: 10.1039/d4ta01060d

rsc.li/materials-a

Copper- and manganese-based layered hybrid organic-inorganic compounds with polymorphic transitions as energy storage materials†

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Solid—solid phase change materials (ss-PCM) have emerged as a promising alternative to traditional methods of thermal regulation, such as solid—liquid transformations. Due to their wide operational temperature range and competitive performance, ss-PCM materials are increasingly being explored for their potential in cooling electronic devices. Here, we explore the potential of layered hybrid organic—inorganic perovskites (LHOIPs) as thermal energy storage materials for passive cooling applications. Two formulations, bis(dodecylammonium) tetrachlorocuprate(ii) (CuC₁₂) and bis(dodecylammonium) tetrachloromanganate(ii) (MnC₁₂) were synthesised and comprehensively characterised. The analyses revealed that these materials present a two-dimensional structure with a triclinic conformation at 100 K. Notably, both materials exhibited a polymorphic transformation with low thermal hysteresis (1.3–2.5 K). These findings indicate that these materials hold significant potential as thermoregulator materials in cooling electronics. Furthermore, both CuC₁₂ and MnC₁₂ demonstrated good thermal stability compared to other types of ss-PCM. Overall, the findings of this study suggest that LHOIPs, particularly CuC₁₂ and MnC₁₂, are promising candidates for further exploration as thermal energy storage materials in electronic cooling applications.

1 Introduction

Nowadays, efficient heat management is crucial for portable electronic devices, especially due to their design requirements and growing power, which results in high power densities.¹ This makes heat dissipation difficult and shortens the lifespan of electronic devices. In fact, 55% of electronic device failure or damage is due to internal overheating.²,³ The primary heat sources of portable electronic devices are Central Processing Units (CPUs) and batteries, where the heat generation density of CPUs is 50 kW m⁻² to 1 MW m⁻².⁴ The methods for electronic devices' thermal regulation can be divided into two main groups: active cooling, where the system needs an external source of energy to improve heat dissipation and passive cooling, generally based on the natural convection of the heat.⁵ One way to reduce the electronic devices' internal overheating is to enhance the performance of passive cooling by integrating

latent heat storage materials into heat sinks. This would

Latent heat storage, also called phase change materials (PCMs), is one of the most promising technologies for thermal regulation in medium temperatures. A wide variety of PCMs can be found nowadays. However, solid-solid PCMs (ss-PCMs) are recommended for this specific application since they do not present leakage and lower corrosion issues than solid-liquid PCMs (sl-PCMs). Also, the ss-PCMs present minimal volume changes and, in some cases, better thermal conductivity than normal paraffin.

Despite this, ss-PCMs present attractive qualities over sl-PCMs as thermoregulating materials in electronics, and interest in these materials has increased in different fields in the last year. However, ss-PCMs are much less studied than sl-PCMs.^{6,7} According to their composition, the ss-PCMs can be grouped into three main groups: organic ss-PCMs, inorganic ss-PCMs, and hybrid ss-PCMs.8 In the organic ss-PCMs, we can find plastic crystals and polymeric ss-PCMs. The polymorphic transition in the plastic crystals relies on a shift of the layered or chain-like structure to an orientally disordered cubic structure. 9,10 However, in the polymeric ss-PCM, the transition relies on the non-crystallizable moieties whose structure is reorganized over the phase transition and the crystallizable moieties preventing melting.11,12 In the inorganic ss-PCMs, the metallic-based and the ceramic-based ss-PCMs are reported, where the metallic-based relates to the martensitic phase transition, and the ceramic-based relates to the inorganic salts,

increase the heat absorption in passive cooling and reduce the workload of active cooling components.

Latent heat storage, also called phase change materials (PCMs).

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[†] Electronic supplementary information (ESI) available. CCDC 2305993 and 2305992. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/d4ta01060d

whose presents polymorphic transformation at high temperatures and the ceramics present a polymorphic transition through pressure and temperature.8 Finally, hybrid ss-PCMs are in the spotlight due to their versatility and tuneable properties, as they can store energy through various driven forces (change of pressure, magnetic field, or temperature). As with the other groups, the hybrid ss-PCMs can be divided into: complex compounds and perovskite-based materials. The main characteristics of each group of ss-PCM are listed in Table 1.

Two-dimensional layered hybrid organic-inorganic perovskites (LHOIPs) materials have gained significant attention recently due to their unique properties in optoelectronics, solar cells field, light-emitting diodes, and photodetectors applications, which present high environmental stability and efficient photoluminescence.¹³ LHOIPs present a wide range of compositional and structure varieties, allowing for tuning their properties to find specific applications. 14 One of the LHOIP chemical composition subfamilies follows this chemical formula (C_n- $H_{2n+1}NH_3$ ₂ MX_4 (n = 8 to 18), being M a divalent transition metal (Cu²⁺, Mn²⁺, Co²⁺, ...) and X a halogen (Cl⁻ or Br⁻). This subfamily of LHOIP presents polymorphic transitions that are interesting from a thermal energy storage (TES) point of view. This family of materials will be an essential step up of TES technology since tailored materials may be proposed for specific applications, such as cooling electronics, thermal management in highly specialised equipment, or aerospace purposes. Also, these materials have an advantage over the commonly studied organic or inorganic phase change materials since the phase transition of LHOIPs happens in a solid state. Hence, no leakage or encapsulation problem must be addressed. 15,16 Moreover, these materials deal with comparative thermal conductivity and lower thermal expansion than solid-liquid phase change materials, as well as enthalpy values and a wide range of temperatures as a function of the structure used.17

Polymorphic transitions of these materials often occur in a sequence of two (or more), where one is more energetic than the other.18-21 Regardless of the number of polymorphic transitions, in all cases, the organic chain remains ionically bonded to the inorganic part, which last remains unaffected by increasing the temperature.22,23 Different studies explain that the main polymorphic transition of this subfamily of LHOIPs relies only on the hydrocarbon chain rearrangement, forming an ordered state where the hydrocarbon chain states in a zig-zag conformation and when the temperature increases, the chain freedom conformation.22 moves

rearrangement, some studies notice that gauche conformation arises in the hydrocarbon chain and an increase in the distance between the facing ends, changing then the volume cell.14

The synthesis optimisation of bis(dodecylamonium) tetrachlorocuprate was previously studied. The study involved examining several synthetic routes and solvent media to determine the appropriate protocol and evaluate the effect of the solvent media. The study concluded that a synthetic protocol that avoids filtration and recrystallisation resulted in a superior yield without decreasing the main key thermal properties for TES applications.24 Moreover, prior research has examined how the length of the carbon chain affects the thermal properties. The research indicates that a longer carbon chain leads to larger volume cells and components with higher transition temperatures and greater overall transition enthalpy. Also, the study confirmed the cyclic durability of the materials for up to 200 cycles.21

In this study, we choose the bis(dodecvlammonium) tetrachlorocuprate(II) (C₁₂H₂₅NH₃)₂CuCl₄ and bis(dodecylammonium) tetrachloromanganate(II) (C₁₂H₂₅NH₃)₂MnCl₄, in order to disclose the crystal structure using X-ray diffraction (XRD), and the molecular organisation using infrared spectroscopy (FTIR) and nuclear magnetic resonance (NMR). Additionally, the thermal properties, such as energy storage capacity, latent heat, and transition temperature stability over thermal cycles, were determined using differential scanning calorimetry (DSC). Moreover, complementary techniques, such as scanning electron microscope (SEM), helium pycnometer, elemental analysis, and thermogravimetric analysis, are used to characterise the material thoroughly and demonstrate its potential in thermal energy storage applications.

Experimental procedure

Synthesis of samples

The reactants copper(II) chloride dihydrated (No. CAS 10125-13-0) and manganese(II) chloride tetrahydrate (No. CAS 13446-34-9) were purchased from VWR international, n-dodecylamine 98% (No. CAS 124-22-1) was purchased from Acros Organics, hydrochloric acid 37% (No. CAS 7647-01-0) from Labbox and anhydrous methanol (max. 0.003% H₂O) (No. CAS 67-56-1) from Merck Group. The synthesis of this layered hybrid organicinorganic perovskite follows the reaction in eqn (1), where ncorresponds to the number of carbons of the amine and M, the corresponding metal employed (Cu or Mn).

$$2C_nH_{2n+1}NH_2 + 2HCl + MCl_2 \rightarrow (C_nH_{2n+1}NH_3)_2MCl_4$$
 (1)

Table 1 Summary of the main properties of ss-PCMs subgroups

Properties	Organic ss-PCMs	Inorganic ss-PCMs ^a	Hybrid ss-PCMs	
Transition temp. (°C)	11-190	680-988	32-160	
Energy density (MJ m ⁻³)	1-275	300-464	16-233	
Thermal conduction (W m ⁻¹ K ⁻¹)	0.10-0.20	0.32-5.00	0.20-2.00	
Volume change	Medium	Medium	High	
Stability	Medium	Medium	High	
a Onder markelling				

Only metallics.

The reactants were accurately weighted to follow the molar ratio presented in eqn (1). Firstly, the alkylamine was dissolved in 45 mL of anhydrous methanol and mixed with the proper content of metal chloride in a round bottom flask. Then, a previously prepared mixture solution with the proper amount of hydrochloric acid and 5 mL of anhydrous methanol was added dropwise to the solution. The mixture was heated to reflux for 4 hours at 337 K under constant stirring. The final solution was poured into a Petri dish and left to dry in a desiccator for seven days. The synthesis procedure in the present study followed the method proposed by R. Salgado-Pizarro *et al.*²⁴

The theoretical elemental composition of C, H, N, Cl and Cu/Mn in $(C_{12}H_{25}NH_3)_2CuCl_4$ is 49.87%, 9.79%, 4.85%, 24.53% and 10.99%, and in $(C_{12}H_{25}NH_3)_2MnCl_4$ 50.62%, 9.91%, 4.92%, 24.90% and 9.65%, respectively. The experimental content of C, H, N, Cl, and Cu/Mn in $(C_{12}H_{25}NH_3)_2CuCl_4$ is 50.06%, 9.87%, 4.86%, 25.38%, and 10.44%, and in $(C_{12}H_{25}NH_3)_2MnCl_4$ is 46.79%, 9.92%, 4.58%, 25.19% and 9.8%, respectively. The C, H, and N contents were measured by elemental analysis, the Cl content was measured by Mohr titration, and the Cu/Mn content was measured by inductively coupled plasma mass spectrometry (ICP-MS).

2.2 Crystallographic study

A colourless plate-like specimen of $(C_{12}H_{25}NH_3)_2MnCl_4$ (MnC₁₂) with approximate dimensions $0.090 \times 0.040 \times 0.020$ mm, and a yellow plate-like specimen of $(C_{12}H_{25}NH_3)_2CuCl_4$ (CuC₁₂), with dimensions $0.080 \times 0.050 \times 0.020$ mm, were used for the X-ray

crystallographic analysis. The X-ray intensity data were measured on a D8 Venture system equipped with a multilayer monochromator and a Mo microfocus ($\lambda = 0.71073 \text{ Å}$). Using a narrow-frame algorithm, the frames were integrated with the Bruker SAINT software package. The data integration using a triclinic unit cell yielded for MnC₁₂ a total of 21 843 reflections to a maximum θ angle of 26.60° (0.80 Å resolution), of which 3188 were independent (average redundancy 6.852, completeness = 98.8%, R_{int} = 7.36%, R_{sig} = 5.02%) and 2674 (83.88%) were greater than $2\sigma(F^2)$. On the other hand, the integration of the data for CuC₁₂ also using a triclinic unit cell yielded a total of 38 908 reflections to a maximum θ angle of 26.43° (0.80 Å resolution), of which 6220 were independent (average redundancy 6.255, completeness = 99.7%, $R_{\text{int}} = 10.84\%$, $R_{\text{sig}} =$ 7.34%) and 4723 (75.93%) were greater than $2\sigma(F^2)$. Data were corrected for absorption effects using the multi-scan method (SADABS). The calculated minimum and maximum transmission coefficients (based on crystal size) are 0.5925 and 0.7454 for MnC₁₂ and 0.5945 and 0.7454 for CuC₁₂. Both structures were solved and refined using the Bruker SHELXTL software package, using the $P\bar{1}$ space group, with Z=1 for the formula unit $C_{24}H_{56}Cl_4MnN_2$ and with Z=2 for the formula unit C24H56Cl4CuN2. The final anisotropic full-matrix leastsquares refinement on F^2 for MnC₁₂ with 172 variables converged at R1 = 8.33%, for the observed data and $wR^2 =$ 22.93% for all data, while for CuC₁₂ with 287 variables converged at R1 = 4.82%, for the observed data and $wR^2 =$ 10.10% for all data. The crystal structures were deposited in the CSD (Cambridge Structural Database) under the deposition

Table 2 Crystallographic data structure refinements for MnC₁₂ and CuC₁₂

	MnC_{12}		CuC_{12}		
Chemical formula	$C_{24}H_{56}Cl_4MnN_2$		$\mathrm{C}_{24}\mathrm{H}_{56}\mathrm{Cl}_4\mathrm{CuN}_2$		
Formula weight	$569.44 \text{ g mol}^{-1}$		$578.04 \text{ g mol}^{-1}$		
Temperature	100(2) K		100(2) K		
Wavelength	0.71073 Å		0.71073 Å		
Crystal system	Triclinic		Triclinic		
Space group	$Par{1}$		$Par{1}$		
Unit cell dimensions	a = 5.0677(5) Å	$\alpha = 90.026(4)^{\circ}$	a = 7.0943(3) Å	$\alpha = 90.911(2)^{\circ}$	
	b = 5.0725(5) Å	$\beta = 92.083(4)^{\circ}$	b = 7.4399(3) Å	$\beta = 92.693(2)^{\circ}$	
	c = 29.955(3) Å	$\gamma=90.807(10)^{\circ}$	c = 28.7460(12) Å	$\gamma=90.210(2)^{\circ}$	
Volume	769.43(13) Å ³		$1515.36(11) \text{ Å}^3$		
Z	1		2		
Density (calculated)	$1.229 { m g cm}^{-3}$		$1.267 \mathrm{\ g\ cm^{-3}}$		
Absorption coefficient	0.790 mm^{-1}		$1.088 \; \mathrm{mm^{-1}}$		
F(000)	307		622		
θ range of data collection	2.04 to 26.60°		2.13 to 26.43°		
Index ranges	$-6 \le h \le 6, -6 \le k \le 6$	$-37 \le l \le 37$	$-8 \le h \le 8, -9 \le k \le 9,$	$-35 \le l \le 35$	
Reflections collected	21 843		38 908		
Independent reflections	3188 [R (int) = 0.0736]		6220 [R (int) = 0.1084]		
Refinement method	Full-matrix least-squares	on F^2			
Data/restraints/parameters	3188/1/172		6220/0/287		
Goodness-of-fit on F^2	1.093		1.045		
Final <i>R</i> indices $[I > 2\sigma(I)]$	$R_1 = 0.0833, \text{ w}R_2 = 0.219$	$R_1 = 0.0833, \text{ w}R_2 = 0.2194$		$R_1 = 0.0482, wR_2 = 0.0925$	
Final R indices	$R_1 = 0.0965, wR_2 = 0.2293$		$R_1 = 0.0742, wR_2 = 0.1010$		
Weighting scheme	$w = 1/[\sigma^2(F_0^2) + (0.1183P)]$	(2 + 3.6984P]	$w = 1/[\sigma^2(F_0^2) + (0.0336P)^2]$	+ 0.9350P]	
	Where $P = (F_0^2 + 2F_c^2)/3$		Where $P = (F_0^2 + 2F_c^2)/3$		
Largest diff. peak and hole	2.616 and -1.010 e Å^{-3}		$0.508 \text{ and } -0.569 \text{ e Å}^{-3}$		
RMS deviation from the mean	0.160 e Å^{-3}		0.092 e Å^{-3}		

numbers 2305993 and 2305992 for MnC₁₂ and CuC₁₂, respectively.

The crystal data and details of data collection and refinements for both compounds are summarised in Table 2.

Variable temperature experiments were performed at the ALBA Synchrotron beamline BL04-MSPD using Mythen detector,25 a wavelength of 0.9544 Å and a Cyberstar hot gas blower with a Eurotherm temperature controller (Eurotherm Worthing, UK). The powder diffraction patterns were indexed using DICVOL06 and the obtained cell parameters were refined via pattern matching with Topas software. 26,27

2.3 Molecular structure determination

Fourier-transformed infrared (FT-IR) spectroscopy coupled with Attenuated Total Reflectance (ATR) (PerkinElmer Spectrum TwoTM) was used to determine the molecular structures and evaluate the chemical changes that originated during the synthesis. Nuclear magnetic resonance (NMR) spectra of ¹H (400 MHz) and ¹³C (101 MHz) in methanol-d₄ (CD₃OD) were recorded using a Bruker BioSpin GmbH 400 MHz NMR spectrometer, in order to validate the amine protonation during the synthesis.

2.4 Morphology study

A scanning electron microscope (SEM) was used to address the morphology of the synthesised structures, using an XTE 325/ D8395, Quanta 200 FEI. Several representative flakes were

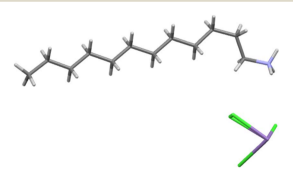


Fig. 1 Asymmetric unit of MnC₁₂.

mounted on a carbon adhesive and coated with carbon in order to acquire SEM images and perform energy-dispersive scattering (EDS) analysis of the surface.

Thermophysical study

Differential Scanning Calorimetry (DSC) was performed using a DSC822e from Mettler Toledo. Around 3.6 mg in Al crucible and a heating ramp from 30 to 80 °C was carried out to evaluate the polymorphic transition at 1 K min⁻¹. Ten cycles of heating and cooling ramps were performed at 10 K min⁻¹ for the stability evaluation. Specific heat (C_p) measurements were performed at 1 K min $^{-1}$ with an amplitude of 0.5 K and a period of 60 seconds in a Q2000 from TA Instruments. The tests performed with the DSC used a nitrogen atmosphere. Thermogravimetric analysis (TGA) was performed under an air atmosphere using a Q600 SDT from TA instruments to evaluate the thermal stability of the component.

3 Results and discussion

3.1 Crystal structure

The MnC₁₂ compound was crystallised from slow cooling to form white plate-like crystals of a triclinic system with space group $P\bar{1}$ and unit cell dimensions of a=5.0677(5) Å, b= $5.0725(5) \text{ Å, } c = 29.955(3) \text{ Å, } \alpha = 90.026(4)^{\circ}, \beta = 92.083(4)^{\circ}, \gamma =$ 90.807(10)° and Z = 1 (see Table 2). Its asymmetric unit is shown in Fig. 1. The ion Mn²⁺ adopts the six-coordinated configuration, forming an infinite net of [MnCl₄]²⁻ with the octahedral geometry. The monomeric [MnCl₄]²⁻ moiety binds with the monomeric [C₁₂H₂₅NH₃]⁺ through electrostatic interactions and several strong N-H···Cl and weaker C-H···Cl hydrogen bonds, which are shown in Table 3. These hydrogen bonds are the main interactions responsible for the sandwichlayered construction, which gives rise to a space supermolecule establishing a two-dimensional perovskite structure (see Fig. 2). The alkylammonium chain contains all trans C-C bonds except for a single gauche C-C bond, which is the one adjacent to the ammonium.

Compound CuC12 was also crystallised by slow cooling to form yellow plate-like crystals of a triclinic system with space group $P\bar{1}$ and unit cell dimensions of a = 7.0943(3) Å, b =

Table 3	Hydrogen	bonds	for	MnC_{12}
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Donor-H···acceptor [ARU]	d(D-H) (Å)	$d(H\cdots A)$ (Å)	$d(D\cdots A)$ (Å)	∠(D-H···A) (°)		
N1A-H1A···Cl1A $[x, -1 + y, z]$	0.91	2.43	3.289(5)	157		
N1A-H1A···Cl1B $\begin{bmatrix} -3 - x, -y, -1 - z \end{bmatrix}$	0.91	2.51	3.291(5)	145		
N1A-H1A···Cl2 $\begin{bmatrix} -2 - x, -y, -1 - z \end{bmatrix}$	0.91	2.73	3.361(5)	127		
N1A-H1B···Cl2 $[-2 - x, 1 - y, -1 - z]$	0.91	2.75	3.186(4)	111		
N1A-H1B···Cl3 $[-2 - x, 1 - y, -1 - z]$	0.91	2.29	3.186(4)	169		
N1A-H1C···Cl1A	0.91	2.41	3.314(5)	171		
N1A-H1C···Cl2 $[-2 - x, 1 - y, -1 - z]$	0.91	2.78	3.186(4)	109		
C1A-H1AA···Cl1A $\begin{bmatrix} 1 + x, -1 + y, z \end{bmatrix}$	0.99	2.76	3.679(10)	154		
C1A-H1AA···Cl1B $\begin{bmatrix} -2 - x, -y, -1 - z \end{bmatrix}$	0.99	2.22	3.096(10)	147		
C1A-H1AB···Cl1A $\begin{bmatrix} 1 + x, y, z \end{bmatrix}$	0.99	2.75	3.575(10)	142		
C1A-H1AB···Cl1B $[-2 - x, 1 - y, -1 - z]$	0.99	2.78	3.649(10)	147		

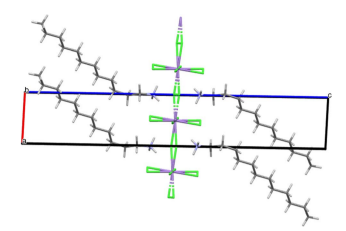


Fig. 2 Packing of MnC₁₂ viewed parallel to [010].

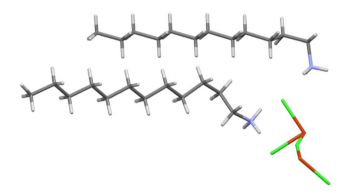


Fig. 3 Asymmetric unit of CuC₁₂.

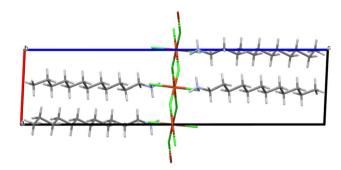


Fig. 4 Packing of CuC_{12} viewed parallel to [010].

7.4399(3) Å, c=28.7460(12) Å, $\alpha=90.911(2)^\circ$, $\beta=92.693(2)^\circ$, $\gamma=90.210(2)^\circ$ and Z=2 (Table 2). Its asymmetric unit is shown in Fig. 3, where two non-equivalent alkylammonium chains and two monomeric $[\text{CuCl}_4]^{2-}$ are observed. The steric configuration of $[\text{CuCl}_4]^{2-}$ is almost square, forming an infinite net of the inorganic moieties. Its packing is shown in Fig. 4, where the $[\text{CuCl}_4]^{2-}$ component binds with the organic $[\text{C}_{12}\text{H}_{25}\text{NH}_3]^+$ moiety through electrostatic interactions and several strong N-H···Cl and weaker C-H···Cl hydrogen bonds (see Table 4). Similar to the manganese compound, a sandwich construction is formed, which is extended to a space super-molecule, helping to establish the two-dimensional perovskite structure. One of the alkylammonium chains contains all trans C-C bonds, while the other chain contains all trans C-C bonds except for one single gauche C-C bond, the one adjacent to the ammonium.

The powder X-ray diffractogram of MnC₁₂ at 340 K was indexed to an orthorhombic unit cell with unit cell parameters $a = 66.25 \text{ Å}, b = 5.28 \text{ Å} \text{ and } c = 5.19 \text{ Å} \text{ and a volume of } 1816 \text{ Å}^3.$ On the other hand, the powder X-ray diffractogram of CuC₁₂ at 343 K was indexed to a tetragonal unit cell with unit cell parameters a = 10.49 Å and c = 31.92 Å and a volume of 3518 Å³. Taking into account the unit cell volume, the molecular weight of each compound, and an estimated density value of 1.2 mg m⁻³, the number of molecules in the unit cell was calculated to be Z = 2 for MnC₁₂ and Z = 4 for CuC₁₂. Different space groups could be assigned for both compounds based on the systematic absences; therefore, as an example, the space groups $P2_12_12_1$ and $P\bar{4}2_1m$ were assigned for MnC₁₂ and CuC₁₂, respectively, in order to perform the subsequent Pawley pattern matchings. These matchings fitted very well with the experimental X-ray powder diffractograms in both cases, being the agreement factors 4.33% (MnC₁₂) and 4.91% (CuC₁₂) (Fig. S1 and S2 in the ESI†).

Additionally, Fig. 5 presents the XRD profile of the two components before and after the phase transition. In both cases it can be observed an increase in the volume cell, as was reported previously, since the lower angle peaks are shifted to lower angles after the phase transition; this means that the distance between the perovskite layers increases due to the reorganisation of the carbon chain. Moreover, an amorphous halo around 11.5° is observed; this may be related to a loss of the order of the carbon chain, resembling the organisation of melting like a polymer after the phase transitions. 17,23

Table 4 Hydrogen bonds for CuC₁₂

Donor-H···acceptor [ARU]	d(D-H) (Å)	$d(H\cdots A)$ (Å)	$d(D\cdots A)$ (Å)	∠(D-H···A) (°)
N1-H1N1…Cl1	0.91	2.36	3.214(2)	157
N1-H2N1···Cl4 $[x, 1 + y, z]$	0.91	2.38	3.238(3)	157
N1-H3N3···Cl3 $[1 - x, 2 - y, 1 - z]$	0.91	2.48	3.365(3)	163
N2-H1N2···Cl1 $[1 + x, y, z]$	0.91	2.43	3.311(3)	162
N2-H2N2···Cl2	0.91	2.35	3.198(3)	156
N2-H3N2···Cl3 $[1 - x, 2 - y, 1 - z]$	0.91	2.32	3.205(2)	164
C10-H10B···Cl1	0.99	2.71	3.679(3)	166
C12-H12A···Cl1 $[x, 1 + y, z]$	0.99	2.66	3.576(3)	154
C24-H24B···Cl3 $[1 - x, 1 - y, 1 - z]$	0.99	2.77	3.687(3)	154

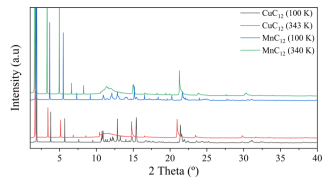


Fig. 5 XRD spectra of MnC₁₂ and CuC₁₂ before and after the thermal

Molecular structure

The FTIR-ATR spectra profiles for Cu and Mn components and the raw materials used are displayed in Fig. 6(a) and (b). The main bands of the raw materials are observed in two obtained

In both cases, the 2955, 2871, 2917, and 2849 cm⁻¹ bands correspond to asymmetric and symmetric stretching of R-CH₃ and R-CH₂-R, respectively, while the 1472 and 1463 cm⁻ bands are related to bending of R-CH₂-R. The 1377 cm⁻¹ band is assigned to the symmetrical bending of R-CH₃, and the 728 and 720 cm⁻¹ bands represent R-CH₂-R rocking, with the 891 cm⁻¹ band indicating terminal C-C stretching. 15,30-32

In the case of the primary amine vibration bands, include were detected at 3400-3250 cm⁻¹ for N-H stretching, 1581 cm⁻¹ for asymmetric deformation of NH₃⁺, 1491 and 1480 cm⁻¹ for symmetric deformation of NH₃⁺, 1215 cm⁻¹ for C-N stretching, and 769 cm⁻¹ for NH₃⁺ wagging.^{30,31}

Minor variations between the Cu and Mn compounds are observed in the FTIR spectra. In particular, the N-H stretching

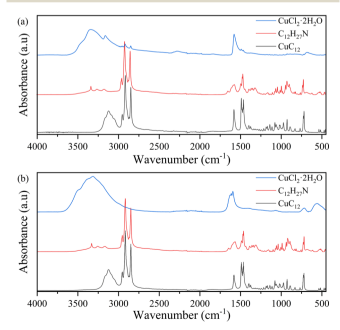
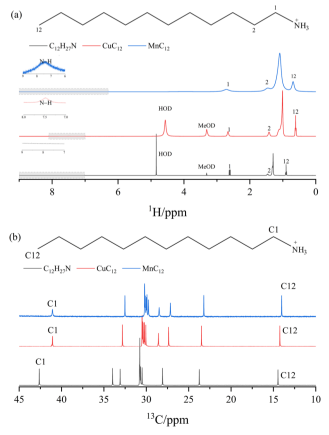


Fig. 6 Infrared spectra of CuC₁₂, MnC₁₂ and raw materials.

vibration band at 3400-3250 cm⁻¹ appears to shift to a lower wavenumber in the Mn compounds, while symmetric and asymmetric deformation of NH₃⁺ shift to higher wavenumber values in the Mn compounds due to polarity changes in the bonds.33

Fig. 7 presents the ¹H and ¹³C NMR spectra of CuC₁₂ and MnC₁₂ compounds, showing the expected peaks (¹³C or ¹H) corresponding to the alkylamine used. In the ¹H NMR spectrum (Fig. 7(a)), a slight shift (\sim 0.2 ppm) in the peak corresponding to the CH2 next to the N is observed in both CuC12 and MnC12 to lower fields concerning the parent unprotonated amine. Also, the alkylamine protonation is observable, detecting the shift of NH₃ at 7.5 ppm.34 Peaks assignable to remains of water and CH2DOD from the solvent are also detected in some spectra. In the ¹³C NMR spectrum (Fig. 7(b)), differences between the dodecylamine, $C_{12}H_{27}N$, and the obtained ammonium compounds are clearly observed. A shift to higher fields of the C1 in CuC₁₂ and MnC₁₂ pecks is detected regarding the parent C₁₂H₂₇N. The peak at 33.98 ppm detected in the $C_{12}H_{27}N$ seems to shift to lower ppm on the compounds' spectra. A 2D NMR experiment (gHSQC) correlating ¹H and ¹³C signals suggests that this peak is shifted to 27.20 ppm. Remarkably, in the CuC₁₂ spectrum, it is able to differentiate the shifts of all the 12 carbons (Fig. S6†) that cannot be determined in the C₁₂H₂₇N or the MnC₁₂ spectra. In order to validate the alkylamine protonation, it was prepared a protonated alkylamine with trifluoroacetic acid (TFA), and was analysed by the ¹³C and ¹H NMR spectra (Fig. S9 and S10†). It was detected that ¹³C and ¹H spectra of the protonated alkylamine with TFA



NMR spectra: (a) ¹³C-NMR and (b) ¹H-NMR.

present similar shift patterns to those obtained in our components, validating our hypothesis. Moreover, in both cases, slight differences in the signal positions between the CuC_{12} and MnC_{12} spectra are detected due to the difference in the polarity of the two metals used and due to the paramagnetic behaviour of the $(CuCl_4)^{2-}$ and $(MnCl_4)^{2-}$ moieties, the signals are not good enough to evaluate the NMR spectra correctly.

3.3 Morphology

The morphology of the resulting crystals was analysed by SEM, as shown in Fig. 8 and 9. Both samples exhibit similar morphologies, resembling flakes or stacking sheets, regarding a two-dimensional structure consistent with the XRD results. Furthermore, a composition analysis was conducted with EDS at different points of the solid to support the elemental analysis and to verify its homogeneity. The result indicates that the main components of the samples are C, Cl, and Cu/Mn, consistent with the elemental analysis. It should be noted that the carbon content is not comparable, as the samples are coated with graphite, and this technique cannot determine the hydrogen content. Furthermore, some potassium was detected in CuC_{12} , which can be attributed to the dust adhered during sample preparation, while oxygen was detected in the MnC_{12} sample, possibly due to the moisture of the sample.

3.4 Thermophysical evaluation

The main thermal properties of the two materials are shown in Table 5. They show similar transition temperature and enthalpy values, slightly higher values in the MnC_{12} sample. This effect could be attributed to the shorter distance between the ammonia and the $(MnCl_4)^{2-}$ that may be related to an increase

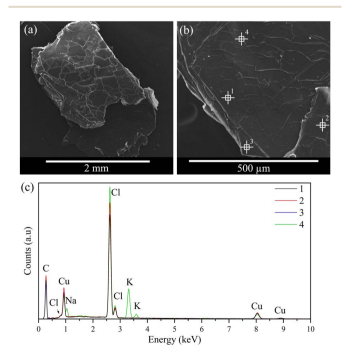


Fig. 8 (a) and (b) SEM images and (c) EDS spectra of CuC_{12} samples.

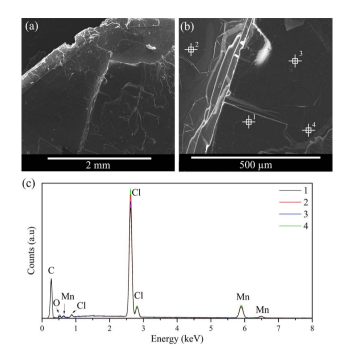


Fig. 9 (a) and (b) SEM images, and (c) EDS spectra of MnC_{12} samples.

Table 5 Transition temperatures and enthalpies of the CuC_{12} and MnC12

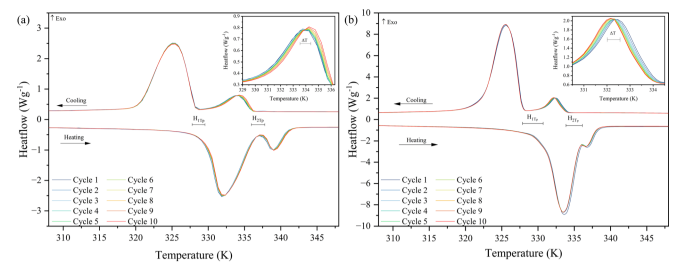
ss-PCM	T_1 (K)	ΔH_1 (J g ⁻¹)	T_2 (K)	ΔH_2 (J g ⁻¹)	$\Delta H_{\rm total}$ (J g ⁻¹)
12	330.9 331.7		338.1 336.0	11.3 7.4	74.5 79.9

in the polarity of the bond. Hence, higher energy will be needed to reorganise the structure.

Fig. 10(a) and (b) show the heat flow curves of ten heating and cooling cycles of the two materials studies performed in a DSC. In both cases, the curve profile is stable under cycles, and no decrease in the enthalpy values is observed. However, a slight curve shift is observed in both samples towards higher temperatures for the CuC_{12} sample and the opposite for the MnC_{12} , where the values of those temperature variations are in Table 6. This difference may be attributed to slight differences in the organisation of the alkylamine chain over the cycles that may be a consequence of the heating rate used. Moreover, in the heating cycles of MnC_{12} , the two transitions overlap, which may be attributed to the high heating rate.

Additionally, the phase change hysteresis was calculated between mean values of the onset temperature of the heating and cooling phase transitions (see Table 6). For the CuC_{12} , similar hysteresis values were obtained in the two transitions, 1.36 and 1.65 K, respectively. Slightly higher values were obtained in the MnC_{12} than CuC_{12} in both transitions, 2.55 and 2.15 K, respectively.

The specific heat measurements of the heating and cooling of both samples are presented in Fig. 11. A good overlap of the heating and cooling curves is observed for the two samples.



Heat flow curves of ten thermal cycles (a) CuC₁₂ and (b) MnC₁₂.

Table 6 Peak temperature variation and onset temperature hysteresis of the complex over ten cycles

	Heating		Cooling			
ss-PCM	$\Delta T_{\rm p1}$ (K)	$\Delta T_{\mathrm{p2}}\left(\mathrm{K}\right)$	$\Delta T_{\rm p1}$ (K)	$\Delta T_{\mathrm{p2}}\left(\mathrm{K}\right)$	$H_{1\text{To}}\left(\mathbf{K}\right)$	$H_{2\mathrm{To}}\left(\mathrm{K}\right)$
CuC ₁₂ MnC ₁₂	0.17 0.16	0.16 0.16	0.00 0.16	0.5 0.16	1.36 2.55	1.65 2.15

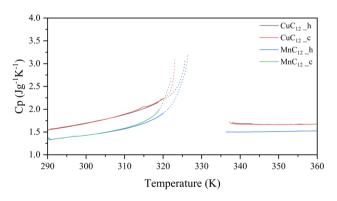


Fig. 11 Specific heat profile as a function of the temperature for CuC_{12} and MnC₁₂ (h: heating; c: cooling).

However, close to the transition temperature, the cooling curve increases its C_p value because of the hysteresis of the system. For the CuC₁₂, the curves remain overlapped up to 320 K, where the transition temperatures are 330.9 and 338.1 K for heating and cooling, respectively. In the case of MnC₁₂, the curves remain overlapped up to 310 K, where the transition temperatures are 331.7 and 336.0 K for heating and cooling.

Moreover, CuC_{12} has higher C_p values than the MnC_{12} sample, which goes from 1.5 to 2.2 J g^{-1} K⁻¹ and 1.3 to 1.5 J g^{-1} K⁻¹ before the transition. After the transition, the values remain constant at around 1.7 and 1.5 J g^{-1} K⁻¹. Comparing the results with other solid-solid PCM, which works in the same range of temperatures, they present values similar to the polymeric solid-solid PCM, which are around 0.8 to 1.5 J g⁻¹ K⁻¹.35,36

The thermogravimetric study performed is presented in Fig. 12, which reveals four different decomposition steps (I-IV). The first step (I), occurring up to 420 K, is attributed to the sample moisture and results in a weight loss of 1.5 wt%. This effect is observed mainly in the MnC₁₂ sample and is likely due to the sample preparation process. The second step (II) corresponds to the degradation of the carbon chain portion of the LHOIP, obtaining $C_xH_vN_z$ as an undecomposed organic residue and the formation of MCl2.37,38 This step shows a mass loss of 50 and 45 wt% up to 595 and 635 K for CuC₁₂ and MnC₁₂, respectively. The third step (III) involves decomposing the $C_xH_yN_z$ organic residue with a mass loss of 10 and 7 wt% up to 700 K.37 The final step (IV) involves the release of chlorides from the metal complex, which corresponds to a mass loss of 30 and 37 wt% up to 900 K.39 A

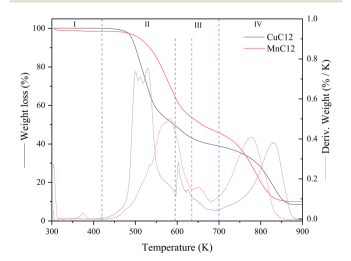


Fig. 12 Thermograms, weight loss as a function of temperature for CuC₁₂ and MnC₁₂ complexes.

schematic approach to the decomposition steps is presented in eqn (2).

$$(C_{12}H_{27}NH_3)_2MCl_4 \xrightarrow{II} C_xH_yN_z$$

+ $MCl_2 \xrightarrow{II} MCl_2 \xrightarrow{IV} M (M = Cu, Mn)$ (2)

Furthermore, the thermo-oxidative temperatures at 5% ($T_{5\%}$) and 10% ($T_{10\%}$) mass loss and the non-volatile inorganic residue at 900 K were determined. The thermos-oxidative temperatures of CuC_{12} at $T_{5\%}$ and $T_{10\%}$ are 492 and 500 K, and for the MnC_{12} , they are 520 and 540 K, respectively. The residue at 900 K was 8.5 and 9.4 wt% for the CuC_{12} and MnC_{12} , respectively. The MnC_{12} presents higher thermo-oxidative temperatures, possibly due to the higher polarity of the alkylamine and manganese chloride, resulting in a higher bond force. These results show that CuC_{12} and MnC_{12} present competent thermal stability compared to other families of solid–solid PCMs. The maximum working temperature for organic ss-PCMs is 423–523 K, and for polymeric ss-PCMs is 473-573 K. 36

4 Conclusions

In this work, we report the structural and thermal characterization of selected LHOIPs, CuC_{12} and MnC_{12} , to be considered as suitable solid–solid phase change materials. The main remarks of the study are the following:

- The crystal structure of CuC_{12} and MnC_{12} is resolved at 100 K for the first time, obtaining a triclinic structure at this temperature. Moreover, the structures were deposited in the CSD.
- The molecular structure of both structures is well described by FTIR and NMR, where the NMR results validate the amine protonation of the obtained components.
- The calorimetry results show that we are dealing with first-order structures, and polymorphic transitions and with low-hysteresis materials that present a specific heat around 1.5–2.2 J $\rm g^{-1}~K^{-1}$.

This study provides enough knowledge to consider these materials ideal candidates for cooling purposes in special applications like electronics or aerospace, among others. However, further investigation regarding the molecular motion of the hydrocarbon chain must be done in order to understand the polymorphic transitions and enhance them fully.

Data availability

Data will be made available on request. The crystallographic data for $[MnC_{12}]$ and $[CuC_{12}]$ has been deposited at the [CCDC] under [2305993] and [2305992].

Author contributions

R. Salgado: conceptualization, methodology, formal analysis, investigation, data curation, writing – original draft, writing – review & editing, visualisation. C. Puigjaner: formal analysis, investigation, data curation, writing – original draft, writing –

review & editing, visualisation. J. García: formal analysis, investigation, writing – review & editing, visualisation. A. I. Fernández: conceptualization resources, supervision, writing – review & editing, funding acquisition, project administration. C. Barreneche: resources, supervision, writing – review & editing, funding acquisition, project administration.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors would like to thank the Catalan Government for the quality accreditation given to their research group (DIOPMA 2021 SGR 00708). DIOPMA is a certified agent TECNIO in the category of technology developers from the Government of Catalonia. The authors are grateful to the CCiTUB for the equipment or measurements of XRD, ICP, elemental analysis and NMR, to the ALBA Synchrotron for the XRD at variable temperature and to the Condensed Matter Physics Department of the University of Barcelona for their contribution with some calorimetry measurements. This work is partially supported by the grant PID2021-123511OB-C32 funded by MCIN/AEI/10.13039/501100011033 and, as appropriate, by "ERDF A way of making Europe" and the PhD grant PRE2019-087336 funded by MCIN/AEI/10.13039/501100011033 and, as appropriate, by "ESF Investing in your future" of R. Salgado-Pizarro.

References

- 1 P. K. Schelling, L. Shi and K. E. Goodson, *Mater. Today*, 2005, 8, 30–35.
- 2 M. Ikhlaq, M. Yasir, M. Demiroglu and M. Arik, *IEEE Trans. Compon., Packag., Manuf. Technol.*, 2021, 11, 1156–1170.
- 3 L. T. Yeh, J. Electron. Packag., 1995, 117, 333-339.
- 4 A. Danowitz, K. Kelley, J. Mao, J. P. Stevenson and M. Horowitz, *Commun. ACM*, 2012, 55, 55–63.
- 5 A. R. Dhumal, A. P. Kulkarni and N. H. Ambhore, *J. Eng. Appl. Sci.*, 2023, **70**, 140.
- 6 T. Bo, T. Zhi-Cheng, L. Rui-Bin, M. Chang-Gong and Z. Jing-Nan, *Energy Convers. Manage.*, 2010, **51**, 1905–1910.
- 7 W. Li, D. Zhang, T. Zhang, T. Wang, D. Ruan, D. Xing and H. Li, *Thermochim. Acta*, 1999, **326**, 183–186.
- 8 A. Usman, F. Xiong, W. Aftab, M. Qin and R. Zou, *Adv. Mater.*, 2022, 34, 2202457.
- 9 J. Timmermans, J. Phys. Chem. Solids, 1961, 18, 1-8.
- 10 S. Santos-Moreno, S. Doppiu, G. A. Lopez, N. Marinova, Á. Serrano, E. Silveira and E. Palomo del Barrio, *Materials*, 2020, 13, 1162.
- 11 X. Lu, C. Fang, X. Sheng, L. Zhang and J. Qu, *Ind. Eng. Chem. Res.*, 2019, **58**, 3024–3032.
- 12 K. Chen, R. Liu, C. Zou, Q. Shao, Y. Lan, X. Cai and L. Zhai, Sol. Energy Mater. Sol. Cells, 2014, 130, 466–473.
- 13 S. Tan, N. Zhou, Y. Chen, L. Li, G. Liu, P. Liu, C. Zhu, J. Lu, W. Sun, Q. Chen and H. Zhou, *Adv. Energy Mater.*, 2018, 1803024.

- 14 J. Li, M. Barrio, D. J. Dunstan, R. Dixey, X. Lou, J.-L. Tamarit, A. E. Phillips, P. Lloveras, J. Li, D. J. Dunstan, R. Dixey, A. E. Phillips, X. Lou, M. Barrio, J.-L. Tamarit and P. Lloveras, Adv. Funct. Mater., 2021, 31, 2105154.
- 15 R. Salgado-Pizarro, J. A. Padilla, E. Xuriguera, C. Barreneche and A. I. Fernández, Energies, 2021, 14, 2621.
- 16 C. Barreneche, L. Navarro, A. de Gracia, A. I. Fernández and L. F. Cabeza, Renew Energy, 2016, 85, 281-286.
- 17 V. Busico, P. Corradini, M. Vacatello, F. Fittipaldi and L. Nicolais, in Thermal Storage of Solar Energy, ed. C. den Ouden, Springer, Dordrecht, 1981, pp. 309-324.
- 18 R. Kind, S. Pleko, H. Arend, R. Blinc, J. Seliger, B. Loar, J. Slak, A. Levstik, C. Filipi, G. Lahajnar, F. Milia and G. Chapuis, J. Chem. Phys., 1979, 71, 2118.
- 19 M. Vacatello, V. Busico and P. Corradini, J. Chem. Phys., 1982, 78, 590-591.
- 20 M. Bochalya, P. K. Kanaujia, G. Vijava Prakash and S. Kumar, AIP Conf. Proc., 2019, 2142, 2-6.
- 21 R. Salgado-Pizarro, A. Svobodova-Sedlackova, A. I. Fernández and C. Barreneche, I. Energy Storage, 2023, 73, 109156.
- 22 V. Busico, C. Carfagna, V. Salerno, M. Vacatello and F. Fittipaldi, Sol. Energy, 1980, 24, 575-579.
- 23 G. F. Needham, R. D. Willett and H. F. Franzen, J. Phys. Chem., 1984, 88, 674-680.
- 24 R. Salgado-Pizarro, C. Barreneche and A. I. Fernández, Mater. Today Chem., 2023, 30, 101491.
- 25 F. Fauth, I. Peral, C. Popescu and M. Knapp, Powder Diffr., 2013, 28, S360-S370.
- 26 A. Boultif and D. Louër, J. Appl. Crystallogr., 2004, 37, 724-731.

- 27 A. A. Coelho, J. Evans, I. Evans, A. Kern and S. Parsons, Powder Diffr., 2011, 26, S22-S25.
- 28 M. Bochalya, P. K. Kanaujia, G. Vijaya Prakash and S. Kumar, *J. Solid State Chem.*, 2019, 273, 219–225.
- 29 D. He, Y. Di, Y. Yao, Y. Liu and W. Dan, J. Chem. Eng. Data, 2010, 55, 5739-5744.
- 30 M. Bochalya, P. K. Kanaujia, G. Vijaya Prakash and S. Kumar, *J. Solid State Chem.*, 2019, 273, 219–225.
- 31 J.-K. Kang, J.-H. Choy and M. Rey-Lafon, J. Phys. Chem. Solids, 1993, 54, 1567-1577.
- 32 R. G. Snyder, J. Mol. Spectrosc., 1961, 7, 116-144.
- 33 E. Pretsch, T. Clerc, J. Seibl and W. Simon, Tabellen zur Strukturaufklärung organischer Verbindungen: spektroskopischen Methoden, Springer-Verlag, Heidelberg, 1976.
- 34 J. E. Sarneski, H. L. Surprenant, F. K. Molen and C. N. Reilley, Anal. Chem., 1975, 47, 2116-2124.
- 35 A. Fallahi, G. Guldentops, M. Tao, S. Granados-Focil and S. Van Dessel, Appl. Therm. Eng., 2017, 127, 1427-1441.
- 36 C. R. Raj, S. Suresh, R. R. Bhavsar and V. K. Singh, J. Therm. Anal. Calorim., 2020, 139, 3023-3049.
- 37 D. Wyrzykowski, A. Pattek-Janczyk, T. Maniecki, K. Zaremba and Z. Warnke, J. Therm. Anal. Calorim., 2008, 91, 279-284.
- 38 E. Styczeń, A. Pattek-Janczyk, M. Gazda, W. K. Jóźwiak, D. Wyrzykowski and Z. Warnke, Thermochim. Acta, 2008, **480**, 30-34.
- 39 E. Styczeń, W. K. Jóźwiak, M. Gazda, D. Wyrzykowski and Z. Warnke, J. Therm. Anal. Calorim., 2008, 91, 979-984.