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# An anomalous ferroelastic phase transition arising from an unusual *cis-/anti*-conformational reversal of polar organic cations†

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Hybrid ferroelastics have attracted increasing attention for their potential application as mechanical switches. The sporadically documented anomalous ferroelastic phase transitions, *i.e.*, ferroelasticity that appears at a high-temperature phase rather than a low-temperature phase, are of particular interest but are not well understood at the molecular level. By judiciously choosing a polar and flexible organic cation ( $\text{Me}_2\text{NH}(\text{CH}_2)_2\text{Br}^+$ ) with *cis-/anti*- conformations as an A-site component, we obtained two new polar hybrid ferroelastics,  $\text{A}_2[\text{MBr}_6]$  ( $\text{M} = \text{Te}$  for **1** and  $\text{Sn}$  for **2**). These materials undergo distinct thermal-induced ferroelastic phase transitions. The larger  $[\text{TeBr}_6]^{2-}$  anions anchor the adjacent organic cations well and essentially endow **1** with a conventional ferroelastic transition ( $P2_1 \rightarrow Pm2_1n$ ) arising from a common order-disorder transition of organic cations without conformational changes. Moreover, the smaller  $[\text{SnBr}_6]^{2-}$  anions can interact with the adjacent organic cations in energetically similar sets of intermolecular interactions, enabling **2** to undergo an anomalous ferroelastic phase transition ( $P2_12_12_1 \rightarrow P2_1$ ) arising from an unusual *cis-/anti*-conformational reversal of organic cations. These two instances demonstrate the importance of the delicate balance of intermolecular interactions for inducing anomalous ferroelastic phase transitions. The findings here provide important insights for seeking new multifunctional ferroelastic materials.

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## Introduction

Ferroelasticity is a mechanical analog of ferromagnetism and ferroelectricity, which offers the unique mechanical properties of spontaneous strain, strain-stress hysteresis, and domain conversion. Extensive research on metal alloys and ceramics has revealed the important roles of ferroelasticity in mechanical switches and piezoelectric sensors.<sup>1–8</sup> Traditionally, displacive-type inorganic ferroelastics, such as  $\text{Gd}_2(\text{MoO}_4)_3$  and  $\text{Pb}_3(\text{PO}_4)_2$ , require solid-state synthesis at high temperatures.<sup>9–10</sup> Hybrid ferroelastics, as important supplements, could be assembled in solutions under moderate conditions from diverse organic and inorganic components, hence offering an eco-friendly and feasible approach for designing ferroelastic materials.<sup>11–14</sup> Recent years have witnessed the emergence of numerous hybrid ferroelastics,<sup>15–27</sup> such as  $(\text{Me}_3\text{NOH})_2[\text{ZnCl}_4]$  with a large spontaneous strain of 0.129,<sup>28</sup> (nortropinium)

$[\text{CdCl}_3]$  with the ultrahigh ferroelastic phase-transition temperatures of 463 and 495 K,<sup>29</sup> and the chiral (*R*-3-chloro-2-hydroxypropyltrimethylammonium) $_2\text{CuCl}_4$  with seven physical channel switches.<sup>30</sup>

According to tests on the 94 species of ferroelastic crystals suggested by Aizu, decreasing temperatures favor a symmetry reduction, and ferroelasticity generally appears at the low-temperature phase (LTP) during a conventional ferroelastic phase transition.<sup>31,32</sup> However, due to the complexity arising from the weak but abundant intermolecular interactions in molecule-based materials, some anomalous cases may occur, *i.e.*, ferroelasticity appears at the high-temperature phase (HTP) rather than LTP.<sup>33,34</sup> One well-known example is Rochelle salt ( $\text{KNaC}_4\text{H}_4\text{O}_6 \cdot 4\text{H}_2\text{O}$ ),<sup>35</sup> which undergoes a phase transition at 255 K from paraelastic LTP in an orthogonal space group,  $P2_12_12$  (point group 222), to ferroelastic HTP in the monoclinic space group,  $P2_1$  (point group 2). In 2021, a molecular rotor crystal,  $[\text{Zn}(\text{saloph})]_2(\mu\text{-dabco})$  ( $\text{H}_2\text{saloph} = N,N'$ -bis(3,5-di-*tert*-butylsalicylidene)-1,2-phenylenediamine,  $\text{dabco} = 1,4$ -diazabicyclo[2.2.2]octane),<sup>36</sup> was found undergoing an anomalous ferroelastic transition at 263 K from paraelectric LTP ( $Cmca$ ) to ferroelastic HTP ( $P2_1/c$ ); in this material the unequal motions of the rotators and anisotropic steric repulsion between the molecules make an important contribution to transition. Recently, we found that an anti-

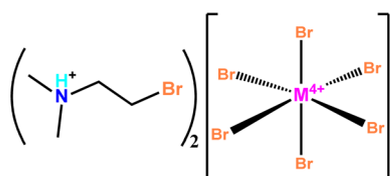
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XeF<sub>4</sub> molecular analog, (Me<sub>3</sub>NCH<sub>2</sub>CH<sub>2</sub>OH)<sub>4</sub>[Ni(NCS)<sub>6</sub>],<sup>37</sup> shows an unprecedented high- and low-temperature dual ferroelasticity with point-group changes of 2/m-4/mmm-mmm; in this material the anomalous ferroelastic HTP is strongly associated with the synergistic swaying of both organic and inorganic components through intermolecular hydrogen bonds. Understanding the origins of anomalous ferroelastic transitions at the molecular level in hybrid crystals is of particular importance for designing advanced high-temperature hybrid ferroelastics as it allows combining multiple functionalities, such as polar or chiral structures. However, among the sporadically documented anomalous ferroelastic transitions, only very rare cases involve polar structures.<sup>35</sup> Therefore, it remains a challenge to design anomalous ferroelastic transitions, owing to the scarcity of instances and the complexity of the molecular components.

In our ongoing studies on phase-transition hybrid crystals assembled by polar organic cations and inorganic components,<sup>11–13</sup> we found some organic cations with changeable *cis-anti*- conformations that can serve as unique components to induce polar structures, such as Me<sub>3</sub>N(CH<sub>2</sub>)<sub>2</sub>Br<sup>+</sup>. These polar structures bring about ferroelectric transition by changing conformations in the first hybrid postperovskite ferroelectric.<sup>38</sup> Considering that the discrete inorganic octahedra facilitate a more complicated and delicate interplay between the inorganic and organic components, we assembled Me<sub>2</sub>NH(CH<sub>2</sub>)<sub>2</sub>Br<sup>+</sup> with diverse discrete inorganic coordination octahedra to seek more uncommon phase transitions. Our efforts yielded two new polar hybrid crystals, (Me<sub>2</sub>NH(CH<sub>2</sub>)<sub>2</sub>Br)<sub>2</sub>[MBr<sub>6</sub>] (M = Te for **1**, and Sn for **2**, Scheme 1). We comprehensively investigated their structural phase transitions using thermal analyses, *in situ* variable-temperature single-crystal/powder X-ray diffraction, dielectric, and second-harmonic generation (SHG) measurements. We found that the synthesized materials undergo thermally induced ferroelastic crystalline phase transitions and solid–liquid transitions accompanied by similar thermally induced “low-high-low” SHG-switching behaviors. However, the slightly changed inorganic octahedra bring about distinct ferroelastic phase transitions, *i.e.*, a conventional ferroelastic transition ( $P2_1 \rightarrow Pm2_1n$ ) associated with a common order-disorder transition of organic cations for **1** and an anomalous ferroelastic transition ( $P2_12_12_1 \rightarrow P2_1$ ) associated with an unusual *cis-anti*-conformational reversal of organic cations for **2**. The underlying delicate balance of the intermolecular interactions for such distinct ferroelastic transitions was discussed *via* Hirshfeld surface analyses, which provides important clues for designing advanced hybrid ferroelastics.



Scheme 1 The structural formulae of **1** and **2** (M refers to either Te or Sn, respectively).

## Experimental section

### Materials and instruments

All chemicals were commercially purchased and were used without further purification. Powder X-ray diffraction (PXRD) patterns were recorded using a Bruker D8 ADVANCE X-ray powder diffractometer (Cu K<sub>α</sub>, λ = 1.54184 Å). Thermogravimetric analyses (TGA) were performed using a TA Q50 system at a heating rate of 10 K min<sup>−1</sup> under a nitrogen atmosphere. Differential scanning calorimetry (DSC) measurements were performed by heating and cooling the powdered sample at a rate of 10 K min<sup>−1</sup> on a TA DSC Q2000 instrument. The dielectric measurements were carried out on a TH2838 impedance analyzer at frequencies from 5 kHz to 2.0 MHz, with an applied voltage of 1.0 V and a temperature sweeping rate of 3 K min<sup>−1</sup> approximately in the range of 100–440 K in a Mercury iTC cryogenic environment controller of Oxford Instrument. The pressed-powder pellets were fixed by two magnetic sheets acting as electrodes. Variable-temperature polarization microscopy observations were carried out on a polarizing microscope OLYMPUS BX41-P equipped with a Linkam cooling/heating stage THMSE 600. The temperature was stabilized to an accuracy of ±0.1 K. The SHG effect was measured using an XPL1064-200 instrument at a heating/cooling rate of 2 K min<sup>−1</sup>.

### Synthesis of **1** and **2**

The red block crystals of **1** and yellow plate crystals of **2** were obtained by slow evaporation of mixed HBr aqueous solutions of (Me<sub>2</sub>NH(CH<sub>2</sub>)<sub>2</sub>Br)Br and corresponding metal oxide (TeO<sub>3</sub> and SnO for **1** and **2**, respectively) in a molar ratio of 1 : 2 for several days at room temperature. The yields were 90% and 95% for **1** and **2**, respectively. The phase purities of the poly-crystals were confirmed by PXRD analyses (Fig. S1, ESI†).

### X-ray crystallography of **1** and **2**

The variable-temperature single-crystal X-ray diffraction intensities were collected on an Agilent SuperNova single-crystal diffractometer with graphite monochromated Cu K<sub>α</sub> (λ = 1.54184 Å) radiation. Absorption corrections were applied by using the multi-scan program, CrysAlisPro. The structures were solved by the direct methods and refined by the full-matrix least-squares method with the SHELX program package on the Olex<sup>2</sup> program.<sup>39,40</sup> The hydrogen atoms were generated geometrically. All non-hydrogen atoms were refined using anisotropic thermal parameters. The positions of the hydrogen atoms were generated geometrically. Detailed crystallographic data and structural refinement parameters are listed in Table 1. The CCDC numbers were 2222445–2222446 for **1** and 2222447–2222448 for **2**, respectively.

## Results and discussion

### Thermal analyses

TGA measurements showed that **1** and **2** could be stable up to 453 K (Fig. S2†) under the N<sub>2</sub> atmosphere. DSC tests in the temperature range of 190–435 K revealed that **1** and **2** undergo



Table 1 Crystal data and structure refinement parameters for 1 and 2

Parameter	$(\text{C}_4\text{H}_{11}\text{NBr})_2[\text{TeBr}_6]$ (1)		$(\text{C}_4\text{H}_{11}\text{NBr})_2[\text{SnBr}_6]$ (2)	
	1_LTP	1_HTP	2_LTP	2_HTP
Formula weight	913.08		904.24	
Phase	1_LTP	1_HTP	2_LTP	2_HTP
Temperature (K)	150(2)	325(2)	293(2)	350(2)
Crystal system	Monoclinic	Orthorhombic	Orthorhombic	Monoclinic
Space group	$P2_1$	$Pm2_1n$	$P2_12_12_1$	$P2_1$
$a/\text{\AA}$	8.5323(5)	8.7336(8)	17.2270(2)	8.7521(7)
$b/\text{\AA}$	9.6457(6)	9.8400(8)	9.6868(1)	9.7104(6)
$c/\text{\AA}$	13.8826(8)	13.801(1)	14.0647(3)	14.0201(1)
$\beta/^\circ$	95.252(3)	90	90	94.430(8)
$V/\text{\AA}^3$	1137.7(2)	1186.1(2)	2347.04(5)	1188.0(2)
Z	2	2	4	2
$D_c/(\text{g cm}^{-3})$	2.665	2.557	2.559	2.528
$R_1 [I > 2\sigma(I)]^a$	0.0586	0.0540	0.0406	0.0796
$wR_2 [I > 2\sigma(I)]^b$	0.1714	0.0823	0.1121	0.2553
GOF	1.085	0.993	1.054	1.066
Flack parameter	$f^c$	0.02(3)	-0.001(6)	0.01(2)
CCDC number	2222445	2222446	2222447	2222448

<sup>a</sup>  $R_1 = \sum ||F_o| - |F_c|| / \sum |F_o|$ . <sup>b</sup>  $wR_2 = [\sum w(F_o^2 - F_c^2)^2 / \sum w(F_o^2)^2]^{1/2}$ . <sup>c</sup> For 1\_LTP: TWIN LAW (1 0 0 0 1 0 0.298 0 -1), BASF [0.016(2)].

reversible solid–liquid phase transitions at similar melting/solidification temperatures of 404/371 K and 399/373 K, respectively (Fig. S3†). In addition, as shown in Fig. 1 and S4,† the DSC tests at temperatures as low as 100 K detected a pair of endothermic/exothermic thermal anomalies at 314/306 K and 340/325 K in the heating/cooling runs, with thermal hystereses of 8 and 15 K for 1 and 2, respectively, indicating reversible one-step crystalline phase transitions; this was further confirmed by variable-temperature PXRD patterns (Fig. S5 and S6†). In the heating/cooling runs, the enthalpy changes ( $\Delta H$ ) and entropy changes ( $\Delta S$ ) during the crystalline phase transition in 1 were 2.6/3.4 kJ mol<sup>-1</sup> and 8.6/11.0 J mol<sup>-1</sup> K<sup>-1</sup>, respectively, both are higher than those observed in 2 (1.3/1.2 kJ mol<sup>-1</sup> and 4.0/3.6 J mol<sup>-1</sup> K<sup>-1</sup>, respectively), implying a more significant change in molecular dynamics for 1 (*vide infra*). For convenience, the phases of 1 below and above 314 K are labeled as 1\_LTP and 1\_HTP, and the phases of 2 below and above 340 K are labeled as 2\_LTP and 2\_HTP.

### Temperature-dependent dielectric responses

Such reversible crystalline phase transitions were further confirmed by a temperature-dependent dielectric constant ( $\epsilon'$ , Fig. 1b and d). For example, the  $\epsilon'$  measured at a frequency of 2 MHz (Fig. S7†) for 1 exhibits a steep increase from *ca.* 3.9 at 265 K (1\_LTP) to *ca.* 4.9 at 335 K (1\_HTP) in a heating run; a reverse drop then occurs at  $\sim$ 301 K in the cooling run that follows. In contrast, for 2,  $\epsilon'$  exhibits a slight jump from *ca.* 3.6 at 295 K (2\_LTP) to *ca.* 4.1 at 355 K (2\_HTP), maintains a gradual increase upon heating, and then exhibits a reverse gradual decrease in the cooling run that follows. Such different temperature dependence of  $\epsilon'$  values implies that the changes in the molecular dynamics of polar organic cations during the phase transition in 1 are much brisker than those in 2.

### Structural analyses

To investigate the structural phase transitions, *in situ* variable-temperature single-crystal X-ray diffraction measurements were performed for 1 and 2, respectively. As illustrated in Fig. 2, both 1 and 2 could be roughly described as a polar A<sub>2</sub>MX<sub>6</sub>-type structure, in which each M<sup>IV</sup> ion is coordinated with six Br<sup>-</sup> ions to form discrete [M<sup>IV</sup>Br<sub>6</sub>]<sup>2-</sup> octahedra separated by A-site (Me<sub>2</sub>NH(CH<sub>2</sub>)<sub>2</sub>Br<sup>+</sup>) cations with *cis*- and *anti*- conformations in 1 : 1 ratio. It should be noted that 1\_LTP and 2\_HTP crystallize in the same polar space group,  $P2_1$  (No. 4), with similar cell parameters. However, 1 and 2 exhibit distinct symmetry-breaking phase transitions upon heating, *i.e.*,  $P2_1 \rightarrow Pm2_1n$  for 1 and  $P2_12_12_1 \rightarrow P2_1$  for 2, respectively (Table 1).

In detail, 1\_LTP belongs to a monoclinic space group ( $P2_1$ , No. 4) with cell parameters of  $a = 8.5323(5)$  Å,  $b = 9.6457(6)$  Å,  $c = 13.8826(8)$  Å,  $\beta = 95.252(3)^\circ$ , and  $V = 1137.74(12)$  Å<sup>3</sup>. Its asymmetric unit contains a complete [TeBr<sub>6</sub>]<sup>2-</sup> octahedron and two crystallographically ordered organic cations in *cis*- and *anti*- conformations, respectively (Fig. S12a†). 1\_HTP belongs to an orthorhombic space group ( $Pm2_1n$ , No. 31), in

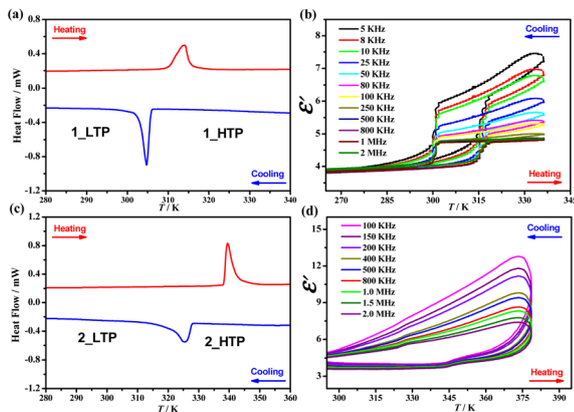


Fig. 1 DSC curves for 1 (a) and 2 (c). Dielectric constants were measured at different frequencies in a heating–cooling cycle for 1 (b) and 2 (d).



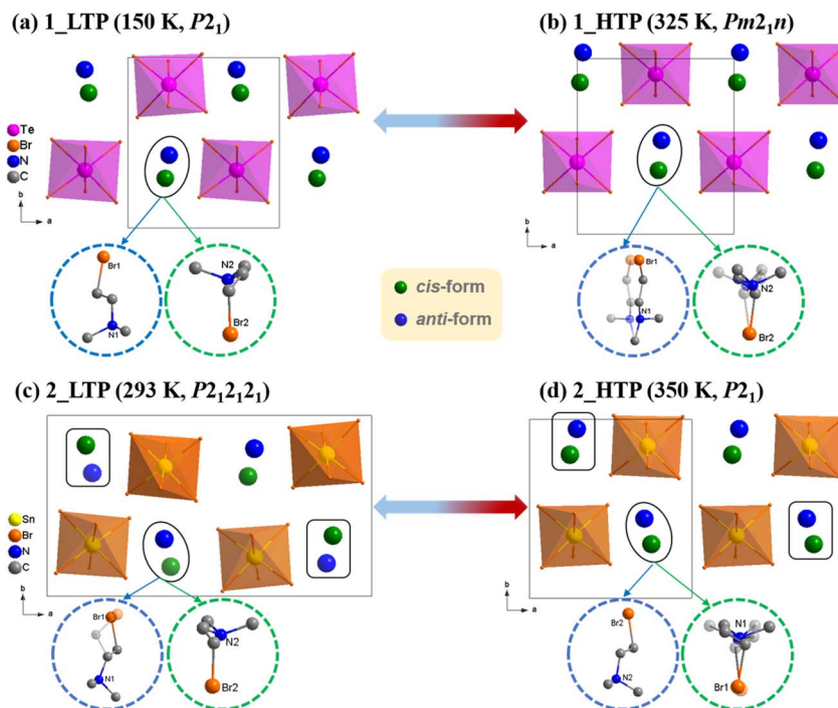


Fig. 2 Crystal packing structures in **1**\_LTP (a), **1**\_HTP (b), **2**\_LTP (c), and **2**\_HTP (d). For clarity, H atoms are omitted and the *cis*-form and *anti*-form conformations of the  $(\text{Me}_2\text{NH}(\text{CH}_2)_2\text{Br}^+)$  cations are represented as green and blue balls, respectively.

which the asymmetric unit becomes halved because of the additional crystallographic mirror symmetry (Fig. S12b<sup>†</sup>); in addition, each organic cation maintains its conformation but becomes 2-fold disordered about the mirror plane perpendicular to the *a*-axis (Fig. S13<sup>†</sup>). In short, the reversible **1**\_LTP  $\leftrightarrow$  **1**\_HTP phase transition can mainly be ascribed to the order-disorder transitions of organic cations, and it obeys a group-supergroup relationship with a normal symmetry-breaking change.

At 350 K (**2**\_HTP), **2** presented a similar structure as **1**\_LTP, except the *cis*-conformational organic cation was crystallographically 2-fold disordered on general position with site occupancies of 0.42 and 0.58 (Fig. S11d<sup>†</sup>). Upon cooling to **2**\_LTP, the space group changed from monoclinic  $P2_1$  into orthorhombic  $P2_12_12_1$  (No. 19) with the *Z* values varying from 2 to 4. In comparison with the **2**\_HTP, **2**\_LTP had a doubled *a*-axis (8.7521(7) Å vs. 17.2270(2) Å) as half of the organic cations reverse their conformations, while slightly changing positions to yield an additional  $2_1$  screw axis along the *a* axis. Moreover, for the two crystallographically dependent organic cations, the *anti*-form one was crystallographically ordered in **2**\_HTP but became 2-fold disordered on general position with the site occupancies of 0.48 and 0.52 in **2**\_LTP. Contrarily, the *cis*-form one being disordered in **2**\_HTP became ordered in **2**\_LTP. In short, differing from **1**, **2** underwent a much more complicated crystalline phase transition that obeyed an unusual group-supergroup relationship, *i.e.*, orthorhombic LTP to monoclinic HTP, which was accompanied by an uncommon *cis*-/*anti*-conformational reversal of flexible organic cations.

### Temperature-dependent lattice parameters

Such distinct phase transitions in **1** and **2** were also reflected in their differences in the temperature dependence of their cell parameters. Pawley refinements were performed on variable-temperature PXRD patterns to obtain the temperature-dependent lattice parameters (Tables S1–S4<sup>†</sup>) at 273–338 K for **1** and at 303–363 K for **2**. The PASCAL program<sup>41</sup> was then employed to calculate the thermal expansion coefficients for each phase (Tables S5 and S6<sup>†</sup>). The volume expansion coefficient slightly decreased from  $+488(8) \times 10^{-6} \text{ K}^{-1}$  (**1**\_LTP) to  $+394(12) \times 10^{-6} \text{ K}^{-1}$  (**1**\_HTP) for **1**, but significantly increased from  $+267(2) \times 10^{-6} \text{ K}^{-1}$  (**2**\_LTP) to  $+678(13) \times 10^{-6} \text{ K}^{-1}$  (**2**\_HTP) for **2**. During the transition from **1**\_LTP to **1**\_HTP at 314 K, the lengths of the *a*-axis and *b*-axis increased by 0.92% and 0.75%, respectively, and a decrease of 1.4% was recorded in the *c*-axis length, together with a slight volume expansion by 0.48% (Fig. 3a). In contrast, during the transition from **2**\_LTP to **2**\_HTP at 340 K, the *a*-axis length increased by 0.67%, the *b*-axis and *c*-axis lengths decreased by 0.21% and 0.58%, respectively, together with an obvious volume shrinkage by about 0.52% (Fig. 3b). Such an abnormal volume shrinkage phenomenon during the **2**\_LTP  $\rightarrow$  **2**\_HTP transition should be associated with the aforementioned *cis*-/*anti*-conformational reversal of the organic cations.

### Ferroelastic phase transitions and spontaneous strains

As indicated by the above structural analyses, during the phase transition,  $Pm2_1n$  (**1**\_HTP)  $\rightarrow$   $P2_1$  (**1**\_LTP) in **1** and  $P2_12_12_1$  (**2**\_LTP)  $\rightarrow$   $P2_1$  (**2**\_HTP) in **2**, the number of symmetry



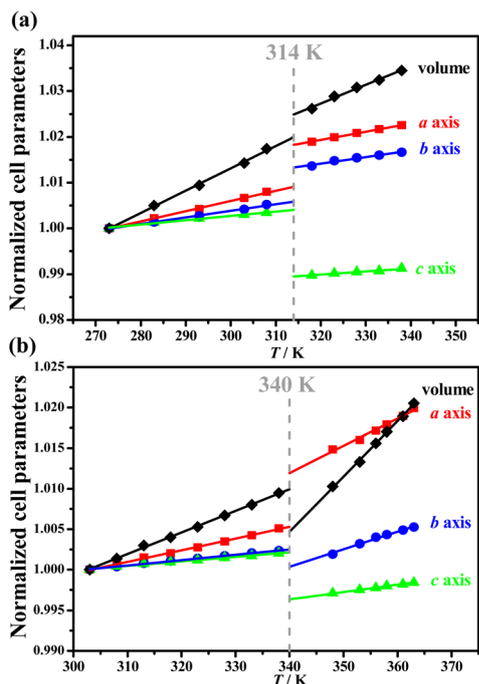


Fig. 3 Temperature dependence of normalized cell parameters for 1 (a) and 2 (b).

elements decreases from 4 ( $E$ ,  $C_2$ , and  $2\sigma_v$ ) to 2 ( $E$  and  $C_2$ ) for 1 and from 4 ( $E$ ,  $C_2$ , and  $2C_2'$ ) to 2 ( $E$  and  $C_2$ ) for 2. These transitions belong to ferroelastic transitions with Aizu notation of  $mm2F2$  and  $222F2$ , respectively.<sup>31</sup> To inspect these ferroelastic transitions, the variable-temperature polarization microscopy observations were performed on the single crystals of 1 and 2 by viewing them perpendicular to the (011) and (101) plane, respectively (Fig. S10 and S11†). As shown in Fig. 4, for 1, in the temperature range of 283–320 K, a stripe pattern could be distinguished on the crystal surface at 298 K, indicating the multiple ferroelastic domains with anti-parallel orientations for 1\_LTP. These striped domains fade out when the crystal is heated to 320 K, indicating a paraelastic state for 1\_HTP, and it

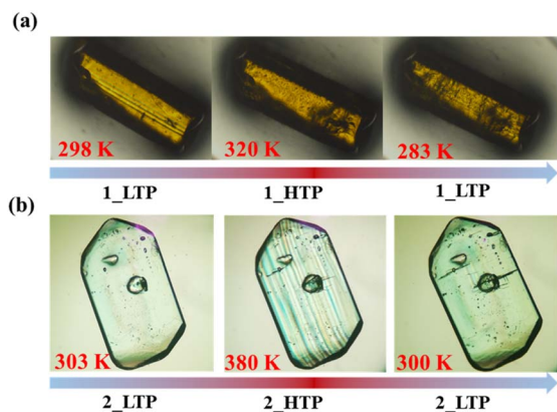


Fig. 4 Variation in the multi-domain structures in a heating-cooling cycle for 1 (a) and 2 (b).

could be recovered by cooling back to 283 K, indicating a recovered ferroelastic state for 1\_LTP. Such a domain revolution strongly verified the reversible and normal ferroelastic phase transition for 1, *i.e.*, between a ferroelastic LTP and a paraelastic HTP, which is similar to the behaviors of most known ferroelastic materials.<sup>17–29</sup> In contrast, in the temperature range of 300–380 K for 2, no multi-domain structure was observed at 303 K as it was in its paraelastic phase (2\_LTP). Upon further heating to 380 K (2\_HTP), the theoretically-predicted multiple domains with two anti-parallel orientation states could be clearly distinguished, which then completely disappeared after cooling back to 300 K. Therefore, as compared to 1 and most other ferroelastic materials, 2 is a rare ferroelastic material with a ferroelastic phase at HTP rather than at LTP.

Based on the cell parameters of 1 and 2 at 314 K and 340 K deduced by extrapolating the fitting lines of variable-temperature cell parameters, the total spontaneous strains were estimated as 0.059 and 0.062 for 1 and 2, respectively (for details, see ESI†).<sup>32,42</sup> These spontaneous strain values are of intermediate level for other hybrid ferroelastics,<sup>15–29</sup> but the value for 2 (0.062) was much larger than those observed in the documented anomalous ferroelastics with inverse symmetry breaking, such as  $[\text{Zn}(\text{saloph})]_2(\mu\text{-dabco})$  (0.02),<sup>36</sup>  $(\text{Me}_3\text{NCH}_2\text{-CH}_2\text{OH})_4[\text{Ni}(\text{NCS})_6]$  (0.0073),<sup>37</sup> and  $(\text{Et}_4\text{N})(\text{Me}_4\text{N})[\text{MnBr}_4]$  (0.0043).<sup>43,44</sup> These results implied that the anomalous ferroelastic transition and large spontaneous strain in 2 should be strongly associated with not only the order-disorder dynamic transition of cations but also the *cis-anti*-conformational variations in organic cations.

### Temperature-dependent SHG responses

The temperature-dependent SHG responses, which are sensitive to the inversion symmetry breaking,<sup>45–47</sup> were recorded by applying the Kurtz and Perry model to the polycrystalline samples ( $\lambda = 1064$  nm, Nd:YAG pulsed laser)<sup>48</sup> of 1 and 2. The SHG intensity at room temperature (Fig. 5) was about 0.06 and 0.13 times that of potassium dihydrogen phosphate (KDP) for 1\_LTP and 2\_LTP, respectively, and increased to 0.16 and 0.32 times that of KDP for 1\_HTP and 2\_HTP, respectively; it eventually dropped to 0.03 and 0.02 times that of KDP for molten 1 and 2, respectively (Fig. S8b and S9b†).<sup>49</sup> Such two-step “low-high-low” SHG-switching behaviors during one-step crystalline phase transition and one-step solid-liquid transition with a rather high switching contrast of 2.6 and 5.3 for 1, 2.5 and 16 for 2, respectively, have rarely been observed. They differ from most of the known SHG switches that present only a one-step “high-low” SHG switching upon heating.<sup>50–53</sup> As the inorganic anions are almost non-polar, the SHG intensification observed in the HTPs should be mainly ascribed to the polar arrangements and the molecular dynamics of the polar organic cations ( $\text{Me}_2\text{NH}(\text{CH}_2)_2\text{Br}^+$ ) with a terminal heavy bromine atom. In detail, these crystallographically disordered polar organic cations in the HTP phase make their dipole moment superimpose along the polarization direction (*i.e.*, the *b*-axis), hence enhancing the molecular polarization and



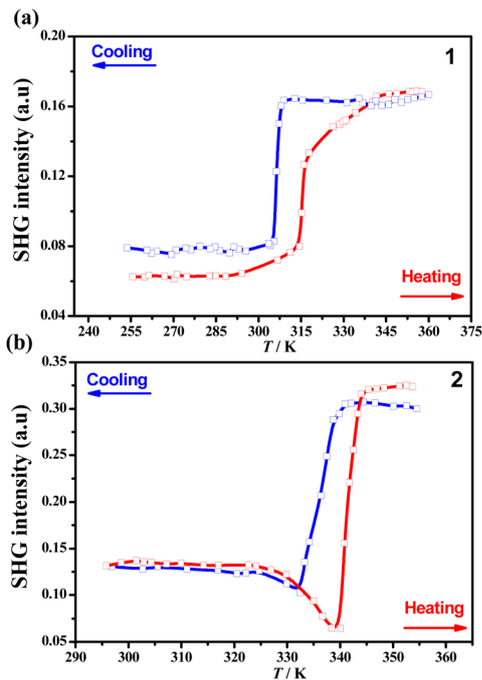


Fig. 5 Temperature dependence of normalized cell parameters for 1 (a) and 2 (b).

then the SHG signals. This observation is consistent with the theoretical calculation of the polarization intensity according to the point charge model<sup>54,55</sup> along the polarization direction for 1 and 2, which revealed that both 1\_HTP and 2\_HTP possess larger polarization than the 1\_LTP and 2\_LTP, respectively (Table S9†). It should be noted that, differing from the case in 1, the SHG signal did not monotonically increase during the 2\_LTP → 2\_HTP transition but temporarily decreased in the vicinity of the phase transition (Fig. 5b).<sup>56–58</sup> Such an abnormal temporary decrease in the SHG response should be strongly associated with the *cis-anti*-conformational reversal process of the organic cations, during which the polar arrangement of the organic cations may be temporarily destroyed to a certain degree.

### Hirshfeld surface analyses

Considering that 1 and 2 have the same organic cations ( $\text{Me}_2\text{-NH}(\text{CH}_2)_2\text{Br}^+$ ) but slightly different inorganic octahedra, it is reasonable to claim that the subtle change in metal ions makes an essential contribution to the distinct phase transitions of 1 and 2. This happens by changes in the conformations and orientations of organic cations as well as the intermolecular interactions involved. As shown by the DSC measurements, the melting points of 1 and 2 were almost the same, implying that they had similar intermolecular interactions between cations and anions. To find out the detail of the differences in their intermolecular interactions, Hirshfeld surfaces<sup>59</sup> were calculated for  $[\text{TeBr}_6]^{2-}$  in 1\_LTP,  $[\text{SnBr}_6]^{2-}$  in 2\_LTP and 2\_HTP, respectively. As shown in the decomposed fingerprint plots (Fig. 6d and e), approximately 93.4% and 94.2% of surface area

were associated with the  $\text{Br}\cdots\text{H-N/C}$  interactions for 1\_LTP and 2\_LTP, respectively. This made major contributions to the attractive intermolecular interactions between the inorganic octahedron and organic cations, while the rest of the surface areas associated with the  $\text{Br}\cdots\text{Br}$  weak interactions between the coordinated Br atom in the inorganic octahedra and the terminal Br atom in the organic cation made only minor contributions. Such similar percentages further confirmed the similarity of inorganic–organic intermolecular interactions in 1 and 2.

However, because of the larger ionic radius of  $\text{Te}^{\text{IV}}$  (97 pm) than  $\text{Sn}^{\text{IV}}$  (69 pm),<sup>60</sup> the  $[\text{TeBr}_6]^{2-}$  octahedron has a larger size than  $[\text{SnBr}_6]^{2-}$ , as reflected by the longer Te–Br bonds (2.670(2)–2.736(2) Å) than the Sn–Br ones (2.581(1)–2.619(1) Å) (Tables S7 and S8†).<sup>61</sup> For the same number of organic cations to interact, the smaller  $[\text{SnBr}_6]^{2-}$  anion was more crowded than  $[\text{TeBr}_6]^{2-}$  anion, and thus, brought about slightly different intermolecular interactions in 1 and 2. As indicated by the red spots on the Hirshfeld surface of  $[\text{TeBr}_6]^{2-}$  in 1\_LTP, besides the major strong inorganic–organic interactions, *i.e.*, the  $\text{N-H}\cdots\text{Br}$  ( $d_{\text{N}\cdots\text{Br}} = 3.41\text{--}3.52$  Å) weak hydrogen bonds, some weak  $\text{C-H}\cdots\text{Br}$  and  $\text{C-Br}\cdots\text{Br}$  ( $d_{\text{Br}\cdots\text{Br}} = 3.68\text{--}4.04$  Å) interactions were formed to complement the intermolecular interactions for both inorganic anions and organic cations. Namely, the larger  $[\text{TeBr}_6]^{2-}$  anions anchored with the adjacent organic cations to be in a crystallographically ordered state in 1\_LTP; it eventually yielded a one-step order-disorder 1\_LTP → 1\_HTP transition by swaying the organic cations along their long axes but maintaining their *cis-anti*-conformations.

By contrast, the smaller  $[\text{SnBr}_6]^{2-}$  anion was too crowded to satisfy all the interactions for the adjacent organic cations, bringing about more possible local structures with different but very similar intermolecular interactions. Accordingly, the organic cations in both 2\_LTP and 2\_HTP were neither crystallographically ordered (the case in 1\_LTP), nor crystallographically disordered (the case in 1\_HTP), but were only partially crystallographically disordered. The major attractive intermolecular interactions (Fig. 6d and f), *i.e.*,  $\text{N-H}\cdots\text{Br}$  and  $\text{C-H}\cdots\text{Br}$  interactions, in 2\_LTP and 2\_HTP, respectively, were similar in general but presented the relation of “as one falls, another rises”. In detail, as indicated by the red spots on the Hirshfeld surfaces of  $[\text{SnBr}_6]^{2-}$  anions (Fig. 6c and e), the  $\text{N-H}\cdots\text{Br}$  interactions in 2\_LTP ( $d_{\text{N}\cdots\text{Br}} = 3.44\text{--}3.58$  Å) are stronger than those in 2\_HTP ( $d_{\text{N}\cdots\text{Br}} = 3.53\text{--}3.55$  Å), while the  $\text{C-H}\cdots\text{Br}$  ones in 2\_LTP ( $d_{\text{C}\cdots\text{Br}} = 3.62\text{--}3.76$  Å) are weaker than those in 2\_HTP ( $d_{\text{C}\cdots\text{Br}} = 3.52\text{--}3.70$  Å). In addition, the relatively longer  $\text{Br}\cdots\text{Br}$  distance in 2\_LTP ( $d_{\text{Br}\cdots\text{Br}} = 3.81\text{--}4.07$  Å) and 2\_HTP ( $d_{\text{Br}\cdots\text{Br}} = 3.84\text{--}4.15$  Å) than that in 1\_LTP ( $d_{\text{Br}\cdots\text{Br}} = 3.68\text{--}4.04$  Å) indicates the weaker  $\text{Br}\cdots\text{Br}$  interactions in 2 (Fig. S14†), such that the smaller  $[\text{SnBr}_6]^{2-}$  anion could not anchor the terminal Br atom of the adjacent organic cations to maintain the conformations for organic cations. In short, the complicated and subtle intermolecular interactions between the organic cations and the  $[\text{SnBr}_6]^{2-}$  anion essentially brought about the complicated crystal structures for both 2\_LTP and 2\_HTP. More importantly, it could generate an



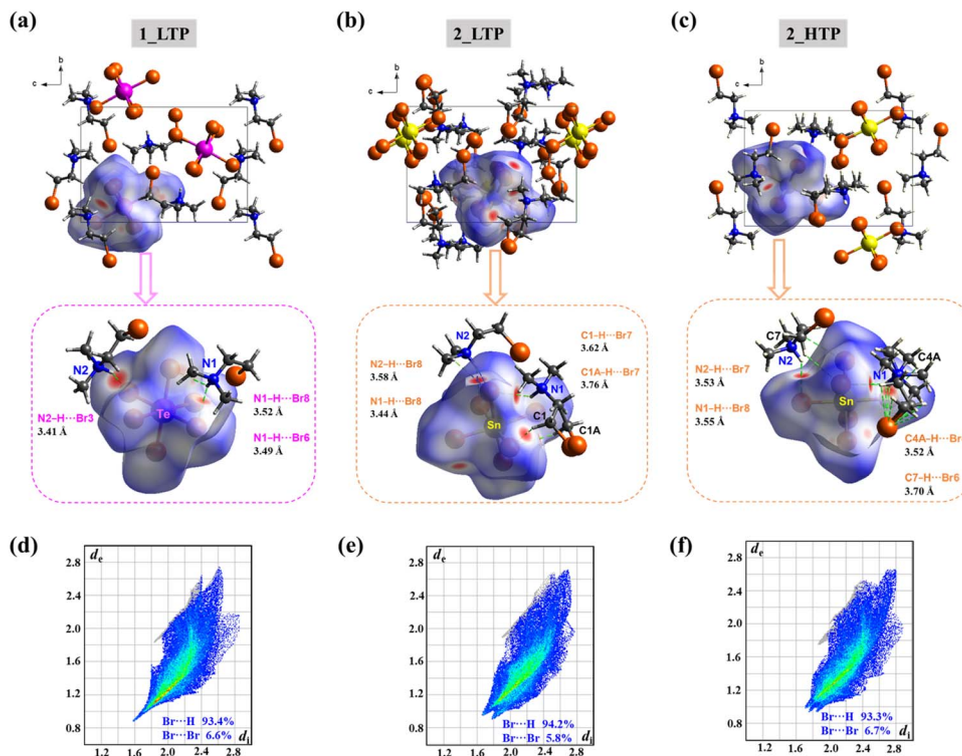


Fig. 6 The Hirshfeld surfaces and fingerprint plots for  $[\text{TeBr}_6]^{2-}$  and  $[\text{SnBr}_6]^{2-}$  in **1**\_LTP (a and d), **2**\_LTP (b and e), and **2**\_HTP (c and f), respectively. Inset: Hirshfeld surfaces show stronger short contacts (red spots) with neighboring atoms.

unusual *cis*-/*anti*-conformational reversal and anomalous ferroelastic phase transitions.

## Conclusions

To summarize, two new polar hybrid ferroelastics, *i.e.*, **1** and **2**, were synthesized successfully by the same organic cations ( $\text{Me}_2\text{NH}(\text{CH}_2)_2\text{Br}^+$ ) but slightly different inorganic octahedra, *i.e.*, relatively larger  $[\text{TeBr}_6]^{2-}$  anions in **1** and smaller  $[\text{SnBr}_6]^{2-}$  anions in **2**. With roughly similar intermolecular interactions, they undergo thermally induced reversible ferroelastic crystalline phase transitions and then solid-liquid phase transitions accompanied by similar and uncommon “low-high-low” SHG switches. However, the slightly different inorganic octahedra endow them with ferroelastic crystalline phase transitions with different symmetry-breaking behaviors, *i.e.*, a normal ferroelastic transition ( $P2_1 \rightarrow Pm2_1n$ ) resulting from a common order-disorder transition of the well-anchored adjacent organic cations without conformational changes in **1**, but an anomalous ferroelastic phase transition ( $P2_12_12_1 \rightarrow P2_1$ ) arising from an unusual *cis*-/*anti*-conformational reversal of organic cations involved in energetically-similar sets of intermolecular interactions in **2**. By presenting these two nice instances to understand the anomalous ferroelastic phase transition at the molecular level, this work demonstrates the crucial role of the delicate balance of intermolecular interactions in hybrid crystals in designing nontrivial phase transitions. The information here also provides important clues for the design of new high-temperature ferroelastic multifunctional materials.

## Data availability

All data included in this study are available upon request by contact with the corresponding author.

## Author contributions

W.-X. Zhang and B.-Q. Zhao conceived the idea, designed the experiments, and co-wrote the manuscript. B.-Q. Zhao performed the synthesis experiments, the DSC, dielectric, crystal structural analyses, ferroelastic domain observations, and the Hirshfeld surface analyses. X.-X. Chen, H. Ye, and Y.-P. Gong assisted the Pawley and Rietveld refinements. J. Wang and L. Ye provided help with the SHG measurements and Hirshfeld surface analyses.

## Conflicts of interest

There are no conflicts to declare.

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