Barocaloric Effects in Dialkylammonium Halide Salts

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ABSTRACT: Barocaloric effects—solid-state thermal changes induced by the application and removal of hydrostatic pressure—offer the potential for energy-efficient heating and cooling without relying on volatile refrigerants. Here, we report that dialkylammonium halides—organic salts featuring bilayers of alkyl chains templated through hydrogen bonds to halide anions—display large, reversible, and tunable barocaloric effects near ambient temperature. The conformational flexibility and soft nature of the weakly confined hydrocarbons give rise to order–disorder phase transitions in the solid state that are associated with substantial entropy changes (>200 J kg\(^{-1}\) K\(^{-1}\)) and high sensitivity to pressure (>24 K kbar\(^{-1}\)), the combination of which drives strong barocaloric effects at relatively low pressures. Through high-pressure calorimetry, X-ray diffraction, and Raman spectroscopy, we investigate the structural factors that influence pressure-induced phase transitions of select dialkylammonium halides and evaluate the magnitude and reversibility of their barocaloric effects. Furthermore, we characterize the cyclability of thin-film samples under aggressive conditions (heating rate of 3500 K s\(^{-1}\) and over 11,000 cycles) using nanocalorimetry. Taken together, these results establish dialkylammonium halides as a promising class of pressure-responsive thermal materials.

INTRODUCTION

Since the invention of vapor-compression refrigeration nearly two centuries ago, cooling technologies have played a critical role in driving economic growth and improving living conditions around the world.\(^1\)\(^-\)\(^3\) Indeed, vapor-compression systems are now ubiquitous, with over 4 billion air conditioning units in use throughout the world.\(^4\)\(^-\)\(^8\) As demand for cooling systems and heat pumps continues to increase,\(^7\)\(^,\)\(^8\) the direct emission of HFCs is projected to climb to upward of 20% of greenhouse gas emissions by 2050,\(^9\)\(^,\)\(^10\) and replacing high-GWP refrigerants with low-GWP refrigerants—such as hydrofluorocarbons, hydrocarbons, ammonia, or carbon dioxide—often involves complicated trade-offs in efficiency, performance, cost, and safety.\(^1\)\(^3\) Moreover, volatile molecules often react in unexpected ways once they inevitably reach the atmosphere,\(^1\)\(^4\)\(^-\)\(^16\) which can have unintended consequences for the environment that are difficult to predict.\(^1\)\(^7\)\(^-\)\(^20\) Alternative technologies that do not rely on volatile refrigerants are, thus, urgently needed to enable more sustainable cooling and heating.

Just as compressing and expanding a volatile refrigerant induces a phase transition between a gas and liquid that is accompanied by a large change in entropy and temperature, increasing and decreasing the pressure around a solid material can also induce phase transitions between high- and low-entropy states. Such solid-state transitions can, in principle, be leveraged to pump heat from a low-temperature heat source to a high-temperature heat sink.\(^2\)\(^1\)\(^-\)\(^2\)\(^4\) In most solid materials, pressure-induced thermal changes—known as barocaloric effects—are small unless very large pressures (well above 1 kbar) are applied.\(^2\)\(^5\)\(^-\)\(^2\)\(^7\) Certain classes of solid-state phase-change materials, however, can exhibit large barocaloric effects at low pressures that provide exciting opportunities for high-efficiency heating and cooling with nonvolatile refrigerants.\(^2\)\(^8\)

Phase transitions between crystalline and dynamically disordered bilayers of hydrocarbon chains in two-dimensional (2D) metal–halide perovskites of the form \([\text{Cs}_2\text{H}_{2\text{n}+1}\text{NH}_3]\text{MX}_3\) (M = Mn, Cu; X = Cl, Br) offer a...
particularly promising mechanism for inducing large, reversible barocaloric effects at low pressures. In 2D perovskites, inorganic layers of corner-sharing metal–halide octahedra template the packing of alkylammonium cations and keep the hydrocarbon chains confined to the solid state even when they take on liquid-like vibrational and rotational degrees of freedom in the partially disordered, high-temperature phase. The structural and chemical features of 2D perovskites...
are well-suited to promoting strong barocaloric effects: (i) the flexibility of hydrocarbon chains leads to strong coupling between structural degrees of freedom and volume, giving rise to high pressure sensitivity, and (ii) the organic−inorganic and organic−organic interfaces can be tailored to minimize hysteresis. Although the inorganic layers of 2D perovskites offer a unique handle for improving and optimizing materials properties important to high-efficiency barocaloric cooling—including hysteresis, thermal conductivity, and mechanical strength—the metal cations and charge-balancing halide anions within the layers also add substantial weight and volume without directly contributing to the phase-change entropy.

In pursuit of a metal−free platform for solid-state hydrocarbon transitions that produce strong barocaloric effects, we recognized that the organic bilayers within dialkylammonium halide salts of the form \((\text{C}_n\text{H}_{2n+1})_2\text{NH}_3\text{X}\) [denoted here as \((\text{C}_n)_2\text{X}\) where \(\text{X} = \text{Cl}, \text{Br}, \text{I}\)] adopt structures that are analogous to those of 2D perovskites. As illustrated in Figure 1, the structure of \((\text{C}_n)_2\text{X}\) can be derived from \((\text{C}_n\text{H}_{2n+1})_2\text{NH}_3\text{MX}\) without altering the overall hydrocarbon chain packing simply by removing the metal cations and merging axially adjacent ammonium groups and halides. Similar to 2D perovskites, dialkylammonium halides thus feature bilayers of hydrocarbon chains that are confined through hydrogen bonding interactions with 2D sheets of halide anions and can undergo reversible solid-state order−disorder transitions. Although the phase-change thermodynamics of many dialkylammonium halides has been investigated in detail at ambient pressure in the context of thermal energy storage, the structural information available for this class of compounds is limited, making it challenging to gain mechanistic insights into their phase transitions and to establish structure−property relationships. In addition, the effect of pressure on dialkylammonium halide phase transitions has not yet been investigated, though we note a somewhat similar compound (di-\(n\)-butylammonium tetrafluoroborate) was recently reported to exhibit large barocaloric effects. Herein, we report a detailed investigation of barocaloric effects in symmetric dialkylammonium halide salts and demonstrate that, when appropriately designed, these materials exhibit properties that make them promising solid refrigerants.

**RESULTS AND DISCUSSION**

**Solid-State Phase Transitions at Ambient Pressure.** To begin investigating barocaloric effects in dialkylammonium halide salts, we selected \((\text{C}_n)_2\text{Br}\) \((\text{C}_n)_2 = \text{di-}\(n\)-hexylammonium) as a representative compound known to undergo a high-entropy phase transition near ambient temperature. Specifically, an order−disorder transition occurs in \((\text{C}_n)_2\text{Br}\) at a transition temperature, \(T_{tr}\) of 293 K with an associated entropy change, \(\Delta S_{tr}\) of 300 J K\(^{-1}\) kg\(^{-1}\) \((315\: \text{J}\: \text{K}^{-1}\: \text{L}^{-1})\) and an enthalpy change, \(\Delta H_{tr}\) of 88 kJ kg\(^{-1}\) \((92\: \text{kJ}\: \text{L}^{-1})\) (Figure 2d). As anticipated, the magnitude of these entropy and enthalpy changes exceeds those of analogous 2D perovskites on both a gravimetric and volumetric basis (Table S1). Interestingly, the molar \(\Delta S_{tr}\) of \((\text{C}_n)_2\text{Br}\) \((80\: \text{J}\: \text{mol}^{-1}\: \text{K}^{-1})\) is more than twice as large as that of \((\text{C}_3\text{H}_{13}\text{NH}_3)_2\text{MnCl}_4\) \((37\: \text{J}\: \text{mol}^{-1}\: \text{K}^{-1})\) even though both compounds undergo transitions near 292 K. This can be partially attributed to a minor disordering transition that \((\text{C}_3\text{H}_{13}\text{NH}_3)_2\text{MnCl}_4\) undergoes at much lower temperatures, though the total hydrocarbon order−disorder transition entropy is still 21% lower for the 2D perovskite compared to \((\text{C}_n)_2\text{Br}\). This could result from a gradual change in hydrocarbon degrees of freedom for \((\text{C}_3\text{H}_{13}\text{NH}_3)_2\text{MnCl}_4\) outside of the first-order phase-transition regions or could reflect an intrinsic difference in the overall entropy change that may arise from differences in chain packing and confinement effects.

To gain additional insight into the structural nature of the \((\text{C}_n)_2\text{Br}\) phase transition, we performed a series of variable-temperature X-ray diffraction experiments. Despite the extensive investigations of their thermal properties, the only crystal structure previously reported for a dialkylammonium halide salt was, to the best of our knowledge, of \((\text{C}_18)_2\text{Br}\) at 288 K. Here, the structure of \((\text{C}_n)_2\text{Br}\) in the low-temperature phase was determined by single-crystal X-ray diffraction at 100 K (Figures 2a and S10). In this monoclinic \((\text{C}_2/c)\) phase, the dihexylammonium cations pack into noninterdigitated bilayers with an interlayer spacing of 13.1 Å. Each ammonium cation is confined by charge-assisted hydrogen bonds to four coplanar Br anions in a square planar arrangement. Note that the confinement area of the Br anions is 28.7 Å\(^2\) (Figure S8 and Table S2), which matches the area for the plane of axial Br anions in Cu−Br perovskites (29 Å\(^2\)). In addition, the electron-deficient carbon atoms (C1 and C1’') adjacent to the ammonium center also contribute to chain confinement by engaging in ion−dipole (Br−H−C) interactions. The two hexyl chains, which are oriented out of the Br plane with a dihedral angle of 72.8°, are tilted 43.5° with respect to the Br plane and have a single gauche C2−C3 bond (torsion angle of 65°) among trans C−C bonds (torsion angle of ∼180°) (Table S4). Again, these structural features are similar to those observed in Cu−Br perovskites that also display a tilt angle of 41−43° and a gauche C2−C3 bond at low temperatures.

Attempts to determine the crystal structure of \((\text{C}_n)_2\text{Br}\) above the phase transition temperature were unsuccessful owing to the high mechanical flexibility of the partially disordered crystal. Indeed, AFM force spectroscopy revealed that the elastic modulus of \((\text{C}_n)_2\text{Br}\) at 300 K is only 0.6 GPa (Figure S54). Nonetheless, variable-temperature powder X-ray diffraction (PXRD) experiments show that \((\text{C}_n)_2\text{Br}\) transitions to an expanded tetragonal phase \((\text{I}4/\text{mmm})\) at high temperatures with a large volume change \((\Delta V_v)\) of 10.4% (Figure 2c,2e). The volume change is highly anisotropic, largely occurring through a 13.4% increase in the interlayer distance (Figure S18). The coefficient of thermal expansion \((\alpha)\) for the high-temperature phase \((5.6\: (7) 10^{-4}\: \text{K}^{-1})\) is notably larger than for the low-temperature phase \((2.2\: (2) 10^{-4}\: \text{K}^{-1})\), indicating that strong anharmonicity is present in the disordered hydrocarbons—similar to the rotator phases of \(n\)-alkanes. In addition, there is a large contribution from diffuse scattering to the PXRD patterns of the high-temperature phase, which is consistent with the presence of substantial dynamic disorder (Figures 2c and S28). In fact, previous solid-state NMR studies have suggested that dialkylammonium cations in shorter-chain analogues of \((\text{C}_n)_2\text{Br}\) \((n = 2, 3, 4)\) rotate along the principal chain axis in the high-temperature phase, behaving as low-dimensional plastic crystals.

In the absence of an experimental crystal structure, we used density functional theory (DFT) calculations to model the high-temperature structure of \((\text{C}_n)_2\text{Br}\) with the 100-K single-
crystal structure serving as a starting point and adjusting the unit-cell dimensions to match those determined from PXRD at 330 K. In the resulting DFT model structure, which represents a snapshot of a potential local energy minimum in the partially disordered phase, the tilt angles of the hexyl groups increase to above 60° relative to the Br plane as the chains undergo conformational changes (Figure 2b). Specifically, the hydrocarbon chains now have an additional gauche C{3}−C{4} bond, along with small distortions (0−30°) in the trans C−C bonds across the rest of the chain (Table S4). The hydrocarbon chains are also engaged in weaker hydrogen bonds, as indicated by a Hirshfeld surface analysis showing elongated and distorted N−H···Br and C−H···Br contacts (Figure S9). Still, the square planar Br anions remain largely intact around the ammonium centers with minimal change in Br−Br distances (Figure S8).

The presence of in-plane hydrogen bonding in the high-temperature phase is further supported by absorption bands near 2800−3000 cm{−1} in the infrared spectrum (Figure S4), which can be assigned to red-shifted N−H stretching vibrations.40−42 Overall, these results suggest that hydrogen bonding between ammonium chains and Br anions persists in the high-temperature phase and imparts a degree of long-range order, highlighting the critical roles of halide anions in governing the internal degrees of freedom of the confined chains.

Further insight into the structural and chemical factors that influence phase-change thermodynamics can be gained by examining the effects of halide substitution, which has a direct impact on chain confinement as demonstrated in 2D perovskites.29,43 Similar to (C{6})_{2}Br, the Cl analogue (C{6})_{2}Cl, which was previously reported but not structurally characterized,2 and I analogue (C{6})_{2}I undergo reversible solid−solid transitions below room temperature at 279 and 286 K, respectively (Figure 3a).

Interestingly, the molar entropy changes of both compounds [45 J K{−1} mol{−1} for (C{6})_{2}Cl and 64 J K{−1} mol{−1} for (C{6})_{2}I] are lower than that for (C{6})_{2}Br. Note that (C{6})_{2}Cl has a minor phase transition at 115 K with a small ΔS{m} of 8 J K{−1} mol{−1}, which has been attributed to the onset of rotation of the dihexylammonium cations.32

We tentatively hypothesize that Br anions lead to the highest entropy change within the (C{6})_{2}X (X = Cl, Br, I) series because the ammonium confinement area is large enough to maximize hydrocarbon degrees of freedom in the high-temperature phase but small enough to prevent the hydrocarbons from partially disordering in the low-temperature phase. A single-crystal structure obtained at 100 K confirms that the area of the Cl plane is 13% smaller than that for the Br analogue (Figure S12 and Table S3). Moreover, the interlayer spacing decreases with increasing halide size in the high-temperature phase from 16.0 Å (Cl) to 14.3 Å (I; Figure 3b). This trend—also observed in 2D perovskites48,49—arises because the dihexyl chains adopt smaller tilt angles to maximize van der Waals interactions between chains as the area of the halide plane increases. Note that F analogues of (C{6})_{2}X do not undergo solid−solid transitions because the F anions are too small (r{F} \approx 1.33 Å) to accommodate disordered dialkylammonium cations that have rotational diameters on the order of 4−5 Å.33

Consistent with their smaller entropy changes, the (C{6})_{2}Cl and (C{6})_{2}I phase transitions are also associated with smaller—though similarly anisotropic—volume changes of 6.5 and 7.8%, respectively (Figures S16 and S17). The experimentally determined volume and entropy changes can be used to estimate the pressure dependence of the transition temperature, dT{tr}/dP, through the Clausius–Clapeyron relationship (dT{tr}/dP = ΔV{tr}/ΔS{m}). Also termed the barocaloric coefficient, the magnitude of dT{tr}/dP is a critical determinant of the performance and operating conditions of a barocaloric material as it dictates how large of a pressure shift (∆P) is needed to induce a given thermal change (∆T). As dictated by the Clausius–Clapeyron relation, dT{tr}/dP is highest when ΔS{m} is small and dV{tr} is large. The inverse relationship between ΔS{m} and dT{tr}/dP represents a challenge for barocaloric materials given the importance of maximizing both ΔS{m} and dV{tr}/dP. Owing to their large dV{tr}, the phase transitions for all three dialkylammonium halide compounds studied here are predicted to exhibit high sensitivity to pressure, with calculated barocaloric coefficients of 33, 30, and 30 K kbar{−1} for Cl, Br, and I analogues, respectively (Figure 3c).

These values are...
comparable to some of the highest coefficients reported for barocaloric materials of any type (Table S10 and Figure S52) and are larger than those associated with melting transitions in \( n \)-alkanes (20 K kbar\(^{-1} \)).\(^{44,45} \)

We note that the higher pressure sensitivity compared to \( n \)-alkanes results from the lower relative \( \Delta S_{\text{tr}} \) of dialkylammonium halides rather than a difference in \( \Delta V_{\text{tr}} \).

**Pressure–Temperature Phase Diagram.** To experimentally probe barocaloric effects in \((C_6)_2X\), we performed isobaric high-pressure differential scanning calorimetry (HP-DSC) measurements using He as the pressure-transmitting medium. In these experiments, phase transition temperatures \( \Delta T_{\text{hys}} \) as well as enthalpies and entropies \( \Delta S_{\text{tr}} \) were determined as a function of pressure up to 150 bar (Figure 4a). The resulting pressure–temperature \((P, T)\) diagrams reveal phase boundaries that allow pressure sensitivity and hysteresis to be evaluated (Figure 4b). As summarized in Figure 4b, \((C_6)_2X\) has a thermal hysteresis \( \Delta T_{\text{hys}} \) of 4 K, whereas \((C_6)_2Cl\) and \((C_6)_2I\) exhibit a lower \( \Delta T_{\text{hys}} \) of 2 K. In addition to thermal hysteresis, pressure hysteresis \( \Delta P_{\text{hys}} \) (defined as the difference in onset pressures between compression- and expansion-induced transitions) is an important metric for evaluating barocaloric materials. On a \((P, T)\) phase diagram, \( \Delta P_{\text{hys}} \) corresponds to the horizontal width of the hysteresis region defined by cooling and heating boundaries and thus can be calculated from \( \Delta T_{\text{hys}} \) and \( \Delta T_{\text{tr}}/\Delta P \) as

\[
\frac{dT_{\text{tr}}}{dP} = \frac{\Delta T_{\text{hys}}}{\Delta P_{\text{hys}}}
\]

Using eq 1, \( \Delta P_{\text{hys}} \) was calculated to be 76 bar (Cl), 149 bar (Br), and 98 bar (I). Since phase transitions of real materials are not perfectly sharp—occurring over a range of pressures rather than a single pressure—\( \Delta P_{\text{hys}} \) represents the minimum pressure required to begin to induce a thermodynamically reversible barocaloric effect,\(^{46,47} \) but higher pressures will be required to induce a large enough reversible temperature and entropy change to drive a practical cooling or heating cycle. Importantly, the \( \Delta P_{\text{hys}} \) values for the dialkylammonium halide salts are below 150 bar, which are among the lowest pressure

![Figure 4](https://doi.org/10.1021/jacs.3c12402)
hysteresis values that have been reported for barocaloric materials (Table S10). Moreover, it is worth highlighting that dicylaluminum halides, along with 2D perovskites, represent one of few barocaloric material classes that feature a combination of large entropy changes (>200 J K⁻¹ kg⁻¹) and low pressure hysteresis (<150 bar) (Figure S51).

Interestingly, as shown in Figure 4c, the calculated ΔP₁₅₀ values are correlated with ΔS₁₅₀ suggesting that there may be a trade-off between the reversibility and magnitude of barocaloric effects in (C₆)ₓX. Although ΔS₁₅₀ also appears to be correlated with ΔV₁₅₀ (Figure 3c), more detailed investigations into geometric phase compatibility are needed to provide insight into the entropy–hysteresis correlation and to better understand the influence of overall volume change and interfacial strain on phase-change reversibility. More broadly, investigations into nucleation mechanisms—at both molecular and microstructural levels—will be required to understand the origin of hysteresis and to help establish design principles for simultaneously maximizing entropy changes and minimizing hysteresis across hydrocarbon order–disorder transitions. Such design principles, which have received limited attention in the barocaloric field, will play a critical role in advancing the basic science needed to support practical applications of barocaloric materials.

**Evaluation of Barocaloric Effects.** To evaluate barocaloric effects in (C₆)ₓX (X = Cl, Br, I), we first constructed isothermal entropy curves through the so-called quasi-direct method. In this approach, isobaric entropy curves, Sｂ(P, T), are calculated from heat flow and heat capacity data obtained from isobaric DSC measurements (Figures S30–S34). Isothermal entropy curves, ΔSｂ, which describe entropy changes induced by a shift in pressure as a function of temperature, are then obtained by subtracting the isobaric entropy curves at two different pressures. For the disordering transition that is induced by decompression from applied pressure P to ambient pressure Pₐ, the ΔSｂ curve is calculated from Sｂ curves associated with heating transitions as

\[
\Delta S_b(P \rightarrow P_a) = S_{b,heating}(T, P_a) - S_{b,heating}(T, P)
\]  

Similarly, for the compression-induced ordering transition, Sｂ curves obtained from cooling transitions are used to calculate the ΔSｂ curve with

\[
\Delta S_b(P_0 \rightarrow P) = S_{b,cooling}(T, P) - S_{b,cooling}(T, P_0)
\]

Figure 5. (a–c) Isothermal entropy changes (ΔSₜ) calculated via the quasi-direct method for (a) (C₆)Cl, (b) (C₆)Br, and (c) (C₆)I. The shaded area indicates the cyclical isothermal entropy change (ΔSₜ,cyc) that is accessible for each operating pressure. The temperature window over which a nonzero ΔSₜ,cyc is possible is indicated by green dashed lines. ΔSₜ,direct values measured from quasi-isothermal DSC experiments are shown as green diamonds for comparison. (d) Maximum cyclical isothermal entropy change (ΔSₜ,cyc,max) calculated from isobaric HP-DSC measurements is plotted against operating pressure (ΔP). ΔP₁₅₀ values determined from isobaric HP-DSC are shown as dashed lines for (C₆)Cl (green), (C₆)Br (red), and (C₆)I (purple).

\[
\Delta S_t(P_0 \rightarrow P) = S_{b,cooling}(T, P) - S_{b,cooling}(T, P_0)
\]

The resulting ΔSₜ curves for (C₆)ₓX (X = Cl, Br, I) illustrate the impact of hysteresis on accessing reversible barocaloric effects (Figure 5a–c). Specifically, cyclical isothermal entropy changes (ΔSₜ,cyc) can be induced for (C₆)Cl and (C₆)I at a 150 bar operating pressure over a temperature range of 2 K. In contrast, compression and decompression ΔSₜ curves for (C₆)Br do not overlap below 150 bar because of a larger pressure hysteresis of 149 bar (Figure 5b). The lack of overlap in ΔSₜ curves indicates that there is no temperature range over which ΔSₜ,cyc can be induced from a cyclic change in pressure. At 150 bar, there is a small region of overlap in the ΔSₜ curves, yielding a maximum entropy change of 12 J K⁻¹ kg⁻¹ over an extremely narrow temperature range of around 293 K (Figure 5d).

Although a 150 bar driving pressure is above the pressure hysteresis for (C₆)ₓX, the maximum entropy changes accessible at 150 bar are still lower than the full ΔSₜ values (Figure 5d), indicating that ΔSₜ,cyc arises from partial phase transformations. This is because first-order phase transitions are associated with a finite peak width (ΔTₚ). To access the full phase-change entropy, a hysteresis loop created between heating and cooling entropy curves at an applied pressure needs to be fully separated from a hysteresis loop at ambient pressure to an extent that the temperature shift in the hysteresis loop surpasses the combination of ΔTₚ,ΔT_width,heating, and ΔT_width,cooling. Such a pressure, denoted as ΔP_rev,full, can be estimated from ΔTₚ/dP as

\[
\Delta P_{\text{rev,full}} = \frac{\Delta T_{\text{heating}} + \Delta T_{\text{width,heating}} + \Delta T_{\text{width,cooling}}}{\Delta T_{\text{rev,full}}}
\]

Above ΔP_rev,full reversible adiabatic temperature changes also become accessible. Assuming that dTₚ/dP and ΔTₚ are relatively constant at higher pressures, the quasi-direct analysis suggests that ΔP_rev,full of 253, 257, and 200 bar are required to fully overcome hysteresis for (C₆)Cl, (C₆)Br, and (C₆)I, respectively. Note that the maximum possible magnitudes of

\[
\Delta T_{\text{rev,full}} = \frac{\Delta T_{\text{heating}} + \Delta T_{\text{width,heating}} + \Delta T_{\text{width,cooling}}}{\Delta T_{\text{rev,full}}}
\]
adiabatic temperature changes \( (\Delta T_{\text{ad, max}}) \) can be estimated using the indirect calculation \( \Delta T_{\text{ad, max}}(T) = -T \Delta S_{\text{it, max}} / c_p \), where \( c_p \) is the ambient-pressure heat capacity and \( \Delta S_{\text{it, max}} \) is the maximum isothermal entropy change. \( \Delta T_{\text{ad, max}} \) values for \((C_2)X\) are 33 K (Cl), 59 K (Br), and 45 K (I), which—assuming constant \( dT_p/dP \)—would require operating pressures of at least 1150, 2187, and 1842 bar, respectively.

Although the quasi-direct analysis offers a facile method to estimate barocaloric properties, this approach inherently relies on phase-change properties measured for thermally induced transitions rather than pressure-induced transitions. As such, this can be misleading. For instance, the measurement of transition hysteresis and peak width—kinetic factors that directly influence reversible barocaloric effects—using isobaric HP-DSC can be influenced by extrinsic factors, such as the sample environment, scan rate, and thermal contact.\(^{35,46}\)

**Quasi-Isothermal Pressure-Swing DSC.** To more directly evaluate barocaloric effects in \((C_2)X\) (\( X = \text{Cl}, \text{Br}, \text{I} \)), we carried out quasi-isothermal, pressure-swing DSC measurements (Figure 6a–f). In these experiments, heat flow signals were measured as the hydrostatic pressure was continuously shifted between 1 and 150 bar. Although the use of He gas as the pressure-transmitting medium led to small temperature fluctuations \( (\Delta T_{\text{avg}} < 0.2 \text{ K}) \) during pressure cycling (Figures S38–S40), the sample temperature stabilized to within 0.4 K of the transition onset, indicating that quasi-isothermal conditions were maintained (Table S9). By capturing the compression-induced exotherms and decompression-induced endotherms, pressure-swing DSC enables the direct characterization of phase-change thermodynamics under more realistic conditions. Throughout the pressure swing, the sample temperature was set to about 1 K above the ambient-pressure \( T_{\text{set, heating}} \) (Figure S41). These set temperatures, \( T_{\text{set}} \), were selected based on the \( P-T \) diagrams and isothermal entropy curves to ensure that reversible transitions were accessible below 150 bar. From the isothermal cycling data, the amount of heat absorbed and released during the pressure swing, \( q_{\text{it}} \), were obtained by integrating the endotherms and exotherms. Using \( q_{\text{it}} \) isothermal entropy changes, \( \Delta S_{\text{it, direct}} \) were then calculated as \( \Delta S_{\text{it, direct}} = q_{\text{it}} / T_{\text{set}} \). Note that the experimental \( \Delta S_{\text{it, direct}} \) values tend to exceed the isothermal entropy changes calculated from the isobaric data because the temperature fluctuation during the pressure swing contributes to driving the phase transitions toward completion.\(^{53}\)

As shown in Figure 6d,e,f, pressure-induced phase transitions in \((C_2)\text{Cl}\) and \((C_2)\text{I}\) were reversible with \( \Delta S_{\text{it, direct}} \) nearly approaching the full transition entropy: 82% (compression) and 93% (decompression) of \( \Delta S_{\text{it}} \) for \((C_2)\text{Cl}\) (163–190 J K\(^{-1}\) kg\(^{-1}\)), and 92% (compression) and 98% (decompression) of \( \Delta S_{\text{it}} \) for \((C_2)\text{I}\) (189–200 J K\(^{-1}\) kg\(^{-1}\)). Note that the
compression $\Delta S_{\text{direct}}$ values were consistently smaller than the decompression $\Delta S_{\text{direct}}$, which can likely be attributed to a larger dissipation of heat signals during the slower compression step (6 bar min$^{-1}$) than during decompression (13 bar min$^{-1}$). Significantly, the $\Delta S_{\text{direct}}$ values remained unchanged over the three cycles, demonstrating that the 150 bar pressure swing provides a sufficient thermodynamic driving force for (C$_6$I)$_2$Cl and (C$_6$I)$_2$I to reversibly transition between low-entropy and high-entropy phases. Overall, these results are in agreement with the $\Delta S_v$ curves calculated from isobaric HP-DSC measurements (Figure 5a,5c), directly validating the reversible temperature window predicted from the quasi-direct analysis.

By contrast, (C$_6$I)$_2$Br displays limited reversibility under the 150 bar swing with isothermal entropy changes substantially lower than the full transition entropy: 74% of $\Delta S_v$ (223 J K$^{-1}$ kg$^{-1}$) and 58% of $\Delta S_v$ (174 J K$^{-1}$ kg$^{-1}$) for compression and decompression, respectively (Figure 6b,6e). Furthermore, the $\Delta S_{\text{direct}}$ values decreased upon cycling, ultimately approaching 33% $\Delta S_v$ (compression) and 50% $\Delta S_v$ (decompression). This result indicates that (C$_6$I)$_2$Br undergoes only partial phase transformations under these conditions, with the phase fraction available to transition decreasing after the first cycle (Figure S42). Consistent with the quasi-direct analysis predicting a limited temperature window for $\Delta S_v$ at 150 bar (Figure 5b), this experiment demonstrates that an operating pressure sufficiently larger than $\Delta P_{\text{hys}}$ is indeed required to fully drive reversible phase changes.

For (C$_6$I)$_2$Cl and (C$_6$I)$_2$I, heat flow traces were plotted as a function of pressure to directly evaluate pressure hysteresis (Figure 7), which was calculated for each cycle as the difference in onset transition pressure ($P_{tr}$) between compression and decompression, with $\Delta P_{\text{hys}} = P_{tr,\text{compression}} - P_{tr,\text{decompression}}$. This evaluation reveals that pressure-driven phase transitions in (C$_6$I)$_2$Cl and (C$_6$I)$_2$I are associated with $\Delta P_{\text{hys}}$ of 59 and 58 bar, respectively. Intriguingly, these experimental $\Delta P_{\text{hys}}$ values are smaller than the $\Delta P_{\text{hys}}$ values of 76 bar (Cl) and 98 bar (I) calculated from isobaric HP-DSC measurements. We speculate that the difference in $\Delta P_{\text{hys}}$ mostly arises from extrinsic factors, such as deviations from isothermality and changes in the quality of thermal contact during the pressure cycle. Although further studies are needed to fully elucidate the origin of such differences, these results highlight the importance of using both isothermal and isobaric measurements to evaluate the potential of new barocaloric materials.

**High-Pressure X-ray Diffraction and Raman Spectroscopy.** Synchrotron PXRD and variable-pressure Raman spectroscopy measurements were used to extend the phase diagram for (C$_6$I)$_2$Br, which has the highest entropy change of the dialkylammonium halide salts evaluated here, up to 1000 bar. Figure 8a shows a waterfall plot obtained for the PXRD patterns of (C$_6$I)$_2$Br collected from 360 to 220 K at 300 bar using He as the pressure-transmitting medium. As illustrated in the waterfall plot, the compound, initially in the high-entropy phase at 360 K, undergoes a phase transition to a contracted state upon cooling at 300 bar with an onset temperature of 296 K. Unit-cell parameters determined by Le Bail refinement show that the ordering transition is accompanied by a large anisotropic contraction of the lattice with $\Delta V_{\text{tr}}/V_{LT}$ of 10.4% (Figure S19), which is comparable to the volume change observed at ambient pressure.

Variable-pressure Raman spectroscopy experiments further show that this order−disorder transition persists at pressures as high as 900 bar. Specifically, Raman spectra were collected for a power sample of (C$_6$I)$_2$Br while pressurizing from 1 to 1000 bar and subsequently depressurizing to 1 bar at a constant temperature of 312 K. As depicted in Figure 8b, reversible transitions between ordered and disordered states were observed at 900 bar (compression) and 740 bar (decompression), the difference between which is in excellent agreement with the $\Delta P_{\text{hys}}$ of 149 bar predicted from HP-DSC experiments.

The transition onsets during the pressure-driven phase change were identified primarily by tracking C−H stretching modes. Peaks at 2920 and 2980 cm$^{-1}$—corresponding to symmetric and antisymmetric C−H stretches of the α-carbon—became more prominent and sharper upon the compression-induced ordering transition. Additionally, in the fingerprint region, a new peak emerged at 1340 cm$^{-1}$ in the high-pressure ordered phase (Figure S44). DFT calculations indicate that this peak is associated with a wagging mode of N−H bonds (Figures S49 and S50). In addition to identifying phase boundaries at higher pressures, these results provide direct experimental evidence that the compression-induced ordering transition is accompanied by an enhancement of C−H−Br and N−H−Br interactions. This observation aligns with the Hirshfeld surface analyses based on computational structural models (Figure S9).

It is worth noting that the changes to C−H stretching vibrations observed during the transition are comparable to those observed in phase transitions in aqueous phospholipid dispersions and n-alkanes. In addition, high-pressure Raman experiments used water as the pressure-transmitting medium. In contrast to 2D perovskites that typically degrade in water, this demonstrates that dialkylammonium halides are sufficiently hydrophobic for water to be used as a heat transfer fluid, which should provide valuable opportunities for thermal device engineering. Most importantly, the transition onset temperatures and pressures determined from high-pressure PXRD and Raman spectroscopy are in excellent agreement with the extrapolated phase boundaries obtained by extending...
temperatures and pressures identified from PXRD and Raman experiments are in excellent agreement with the phase boundaries determined from isobaric HP-DSC.

Moreover, owing to its high sensitivity of near 1 nJ K\(^{-1}\), the nanocalorimetry technique enables thermal characterization of thin-film samples at scan rates of up to 10 \(10^3\) K s\(^{-1}\). To perform nanocalorimetry measurements, we first deposited a 50-nm-thick film of \((\text{C}_{10})_2\text{Br}\) \([\text{(C}_{10}\text{)}_2 = \text{di-}\text{n-decylammonium}]\) onto a nanocalorimetry sensor with an area of \(1.9 \times 10^{-2}\) cm\(^2\). Note that the C\(_{10}\) analogue was used for these experiments instead of C\(_6\) because its transition temperature (328 K) is above room temperature, which simplified the nanocalorimetry measurements. Due to the longer, more flexible C\(_{10}\) chain, \((\text{C}_{10})_2\text{Br}\) displays a phase-change entropy of 123 J mol\(^{-1}\) K\(^{-1}\) that is 54% higher than the molar \(\Delta S_m\) of \((\text{C}_{6})_2\text{Br}\) (Table S5). This is consistent with the entire \((\text{C}_{n})_2\text{Br}\) \((n = 2-18)\) series of compounds for which the molar \(\Delta S_m\) values monotonically increase with increasing chain length (Figure S6). The order–disorder transition of \((\text{C}_{10})_2\text{Br}\) is also accompanied by a large volume change of 90.0 cm\(^3\) kg\(^{-1}\) (9.6%) (Figure S21), which gives rise to a high pressure sensitivity of 23 K kbar\(^{-1}\) (Figure S37 and Table S8).

To allow for rapid scan rates, the thermal mass of the thin-film sample was kept very low (on the order of 100 ng). Interestingly, the phase transition temperature of \((\text{C}_{10})_2\text{Br}\) was 10 K lower for the thin-film sample compared to a bulk powder sample (Figure 9a), suggesting that the phase behavior of dialkylammonium halides can be influenced by size and substrate effects. This result has implications for the design and optimization of composite materials based on dialkylammonium halides and highlights the importance of exploring how sample size and morphology influence phase-change properties. Although further studies are needed to elucidate the origin of the HP-DSC results at lower pressures (Figure 8c). As such, these experiments confirm that a pressure sensitivity of 27 K kbar\(^{-1}\) persists for \((\text{C}_{6})_2\text{Br}\) even at much higher pressures and that the compound still undergoes a single transition between a low- and high-entropy state.

**Nanocalorimetry.** In addition to water compatibility, dialkylammonium halides have several other features that are advantageous for device applications. In particular, because of their solution-processable nature, they can be readily deposited onto a wide range of substrates as films with controllable thicknesses (Figure S53). Beyond device engineering, this offers a unique opportunity to characterize phase-change phenomena under extreme, nonequilibrium conditions and can allow for rapid thermal cycling tests to be performed.\(^{56-58}\)

To perform nanocalorimetry measurements, we first deposited a 50-nm-thick film of \((\text{C}_{10})_2\text{Br}\) \([\text{(C}_{10}\text{)}_2 = \text{di-}\text{n-decylammonium}]\) onto a nanocalorimetry sensor with an area of \(1.9 \times 10^{-2}\) cm\(^2\). Note that the C\(_{10}\) analogue was used for these experiments instead of C\(_6\) because its transition temperature (328 K) is above room temperature, which simplified the nanocalorimetry measurements. Due to the longer, more flexible C\(_{10}\) chain, \((\text{C}_{10})_2\text{Br}\) displays a phase-change entropy of 123 J mol\(^{-1}\) K\(^{-1}\) that is 54% higher than the molar \(\Delta S_m\) of \((\text{C}_{6})_2\text{Br}\) (Table S5). This is consistent with the entire \((\text{C}_{n})_2\text{Br}\) \((n = 2-18)\) series of compounds for which the molar \(\Delta S_m\) values monotonically increase with increasing chain length (Figure S6). The order–disorder transition of \((\text{C}_{10})_2\text{Br}\) is also accompanied by a large volume change of 90.0 cm\(^3\) kg\(^{-1}\) (9.6%) (Figure S21), which gives rise to a high pressure sensitivity of 23 K kbar\(^{-1}\) (Figure S37 and Table S8).

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of this phenomenon, similar observations were previously made for martensitic transitions in alloy films\textsuperscript{59,60} and spin transitions in molecular thin films.\textsuperscript{61}

During over 11,000 thermal cycles with a heating scan rate of 3500 K s\textsuperscript{-1}, the phase transition temperature for (C\textsubscript{6}H\textsubscript{13})\textsubscript{2}Br remained constant with onset and peak temperatures of 311 and 318 K, respectively. Note that a decrease in transition enthalpy was observed during thermal cycling, which is attributed to sample sublimation under the ultrahigh vacuum conditions (1.3 × 10\textsuperscript{-5} Pa) used for the nanocalorimetry experiments (Figure 9b). This was confirmed by optical images of the sensor area before and after cycling (Figures S56 and S57).

\section*{CONCLUSIONS}

The foregoing results establish dialkylammonium halide salts as a new class of highly tunable barocaloric materials with excellent energy density, reversibility, processability, and stability. Through a systematic investigation of barocaloric effects in these organic materials, this work contributes to the rapidly growing knowledge base around pressure-induced phase transitions in organic salts,\textsuperscript{37,62,63} long-chain hydrocarbons,\textsuperscript{65,64--66} and plastic crystals.\textsuperscript{67--69} Efforts to further investigate thermal conductivity, dynamics of disordered phases, and barocaloric effects across a wider range of soft materials are currently underway in our laboratory.

\section*{ASSOCIATED CONTENT}

\section*{Supporting Information}

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.3c12402.

Additional experimental details, high-pressure calorimetry data, entropy curves used to evaluate barocaloric effects, powder X-ray diffraction data, single-crystal X-ray diffraction data, Raman spectroscopy data, and optical images and nanocalorimetry data, and a summary of all structural and thermodynamic data (PDF)

Accession Codes

CCDC 2171719–2171722 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, or by emailing data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.

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Notes

The authors declare the following competing financial interest(s): J.S., F.E.C., J.D.B., and J.A.M. are inventors of technology utilized in this research that is licensed to PasCal Technologies, Inc., of which J.S. and J.A.M. are Co-Founders. J.A.M. is a Fiduciary Board member of PasCal Technologies, Inc. By improving upon the licensed technology, the outcomes of this research could result in a financial benefit to J.S, F.E.C., J.D.B, and J.A.M. and PasCal Technologies, Inc.

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■ NOTE ADDED AFTER ASAP PUBLICATION

CCDC codes were inadvertently omitted in the version published January 16, 2024 and were correctly restored January 18, 2024.