Energetics, Structures, and Phase Transitions of Cubic and Orthorhombic Cesium Lead Iodide (CsPbI₃) Polymorphs

Bin Wang, Novendra Novendra, and Alexandra Navrotsky

Supporting Information

ABSTRACT: Cesium lead iodide (CsPbI₃) perovskite has shown great potential as a light absorbing material for solar cell applications. Despite intense research leading to increasing power conversion efficiency, a major problem concerning CsPbI₃ lies in the long term stability and interconversion between different CsPbI₃ polymorphs, a subject barely studied from the thermodynamic perspective. We report the formation enthalpies of two CsPbI₃ polymorphs, α and δ CsPbI₃, using a combination of room temperature solution calorimetry in dimethyl sulfoxide (DMSO) and differential scanning calorimetry. We show that both polymorphs are stable with respect to their binary halides and confirm that the α-phase is a high temperature polymorph, metastable under ambient conditions. This work sheds light on patterns in polymorphism, possible decomposition reactions, materials stability, and compatibility within halide perovskites and related systems. Thermodynamic instability near ambient temperature of functional perovskites may be a general phenomenon related to their vibrational density of states.

Inorganic halide perovskite (IHP) materials have captured broad research interests and shown increasing power conversion efficiency (PCE) in perovskite solar cells (PSC). The generic formula for a 3D-structured inorganic halide perovskite (HOIP) where A is usually methylammonium (CH₃NH₃⁺), MA⁺ or formamidinium (HC(NH₂)₂⁺, FA⁺), IHPs exhibit superior thermal and chemical stabilities under exposure to external humidity, oxygen and heat, which is rendered by the replacement of organic with intrinsically more stable inorganic cations. Among various inorganic halide perovskites, cesium lead iodide (CsPbI₃) draws special attention for its desirable bandgap, ease of synthesis and tunable optical properties. Tremendous efforts have been devoted to the synthesis, modification and application of CsPbI₃ materials. Despite this, studies on the thermodynamic properties of CsPbI₃ materials remain scarce. Yet the underlying thermodynamic properties determine thermal and atmospheric stability, possible decomposition reactions, and materials compatibility.

There appear to be several polymorphs within the cesium lead iodide system, whose interconversion and thermodynamic and thermal stabilities have not been well characterized.

The one used as a light harvesting material in perovskite solar cells is α-CsPbI₃ ($E_g = 1.73$ eV, black), which adopts a cubic perovskite crystal structure (space group: $Pm3m$). However, a nonperovskite, δ-CsPbI₃, with wide band gap ($E_g = 2.82$ eV, yellow) and orthorhombic crystal structure (space group: $Pna21$), was reported to be thermodynamically more stable below $\sim 320$ °C. When exposed to ambient conditions, α-CsPbI₃ undergoes an immediate phase transition to its orthorhombic polymorph (δ-CsPbI₃). A spontaneous phase transition from δ- to α-CsPbI₃ was also reported when the temperature was ramped to $\sim 320$ °C. Thus, the α-phase appears to be the high temperature stable polymorph, but this hypothesis needs thermodynamic verification.

Here we report the formation enthalpies of α- and δ-CsPbI₃ polymorphs from lead iodide (PbI₂) and cesium iodide (CsI) using a combination of room temperature solution calorimetry and differential scanning calorimetry (DSC), confirming the energetic stability of both polymorphs with respect to their binary halide components and the energetic metastability of α-CsPbI₃ relative to the δ-phase. We propose some energetics–structure correlations in different polymorphs in perovskite-forming halide systems.

The schematic crystal structure and powder X-ray diffraction pattern of the synthesized orthorhombic δ-CsPbI₃ are shown in Figure 1, the latter confirming phase pure product. Structural details of δ-CsPbI₃, as well as those of α-CsPbI₃, are shown in Table 1.

The measured enthalpies of solution in dimethyl sulfoxide (DMSO) of δ-CsPbI₃ and constituent iodides are given in Table 2. The thermochemical cycle used to calculate enthalpy of formation is also shown. The formation enthalpy of δ-CsPbI₃ from binary iodides (PbI₂ and CsI) is significantly exothermic, $-16.93 \pm 0.87$ kJ/mol, in contrast to the positive value for the hybrid perovskite MAPbI₃. This more favorable formation enthalpy of CsPbI₃ indicates the greater stability rendered by the symmetrical, spherical Cs⁺ ion compared to the volatile MA⁺ ion due to its polarity and asymmetry. Indeed in general organic–inorganic hybrid perovskites are less thermodynamically stable than purely inorganic ones, giving the latter advantages in long-term use. Furthermore, the “yellow phase” with edge sharing polyhedra seems to be confined to the inorganic systems and does not appear to occur in the hybrid systems. This energetic stability is unlikely to be counteracted by entropy terms for formation from binary iodide constituents. Therefore, δ-CsPbI₃ is expected to have a
suggests a continuous curve could relate the heat capacity of enthalpy and entropy of transition at ambient conditions. The values can also be taken as a good approximation of the transition (as it does not exist), but the DSC curve (Figure 2) phenomena between 25 and 320°C expected as there are no phase transitions or disordering making it just barely stable energetically. However, the large negative free energy of formation from its iodide components, which explains the long-term stability of δ-CsPbI₃ with no obvious decomposition or change in color or morphology when held in ambient conditions for months.

The black cubic perovskite phase (α-CsPbI₃), with a desirable band gap, transforms rapidly to δ-CsPbI₃ below ~320°C. Thus, α-CsPbI₃ can not be obtained for room temperature solution calorimetry. However, the thermodynamics of the phase transition from δ→α CsPbI₃ near 320°C can be measured directly by differential scanning calorimetry (DSC), providing a means of relating the energetics of the two phases.

The DSC thermogram of δ-CsPbI₃ (Figure 2) shows an endothermic peak starting at 320°C and centered at 325°C, corresponding to the δ→α-CsPbI₃ phase transition with an enthalpy change of 14.10 ± 0.24 kJ/mol. The entropy of transition is 14100/593 = 23.78 J/mol·K. By neglecting the difference in heat capacities between the two phases, these values can also be taken as a good approximation of the enthalpy and entropy of transition at ambient conditions. The heat capacity of the α-phase cannot be measured below the transition (as it does not exist), but the DSC curve (Figure 2) suggests a continuous curve could relate the heat capacity of the δ-phase below the transition and the α-phase above, supporting similar heat capacities. Similar heat capacities are expected as there are no phase transitions or disordering phenomena between 25 and 320°C, and the vibrational heat capacities of the two phases are expected to be similar in that range. With these approximations, the enthalpy of formation of the α-phase at ambient conditions is ~2.83 ± 0.90 kJ/mol, making it just barely stable energetically. However, the large positive entropy of the δ→α transition clearly stabilizes α-CsPbI₃ above 320°C. Thus, the δ-phase is confirmed to be

<table>
<thead>
<tr>
<th>Reaction Scheme</th>
<th>Enthalpy Measurements</th>
<th>ΔH (kJ/mol)</th>
<th>Note</th>
</tr>
</thead>
<tbody>
<tr>
<td>δ-CsPbI₃ (sl, 25°C) → PbI₂ (sln, 25°C) + CsI (sln, 25°C)</td>
<td>ΔH₁ = ΔHᵣ₋₃ (δ-CsPbI₃, xl, 25°C)</td>
<td>-46.12 ± 0.81 (8)</td>
<td>c</td>
</tr>
<tr>
<td>CsI (sl, 25°C) → CsI (sln, 25°C)</td>
<td>ΔH₂ = ΔHᵣ₋₃ (CsI, xl, 25°C)</td>
<td>-12.64 ± 0.21 (8)</td>
<td>c</td>
</tr>
<tr>
<td>PbI₂ (sl, 25°C) → PbI₂ (sln, 25°C)</td>
<td>ΔH₃ = ΔHᵣ₋₃ (PbI₂, xl, 25°C)</td>
<td>-50.41 ± 0.24 (8)</td>
<td>c</td>
</tr>
<tr>
<td>CsI (sl, 25°C) + PbI₂ (sl, 25°C) → δ-CsPbI₃, (sl, 25°C)</td>
<td>ΔH₄ = ΔHᵣ₋₃ (δ-CsPbI₃, xl, 25°C)</td>
<td>-16.93 ± 0.87 (8)</td>
<td>ΔHᵣ = (-ΔH₁) + ΔH₂ + ΔH₃</td>
</tr>
<tr>
<td>δ-CsPbI₃ (sl, 25°C) + α-CsPbI₃ (sl, 25°C)</td>
<td>ΔH₅ = ΔHᵣ₋₃ (α-CsPbI₃, xl, 25°C)</td>
<td>14.10 ± 0.24 (8)</td>
<td>d</td>
</tr>
<tr>
<td>CsI (sl, 25°C) + PbI₂ (sl, 25°C) → α-CsPbI₃, (sl, 25°C)</td>
<td>ΔH₆ = ΔHᵣ₋₃ (α-CsPbI₃, xl, 25°C)</td>
<td>-2.83 ± 0.90 (8)</td>
<td>ΔHᵣ = ΔH₅ + ΔH₆</td>
</tr>
</tbody>
</table>

*State symbol: sln (DMSO solution), xl (crystalline solid). The mean of values calculated from solution calorimetry or differential scanning calorimetry, unless otherwise specified. Uncertainty is two standard errors of the mean. Numbers of measurements indicated in parentheses.
1. Calculated from solution calorimetry. Temperature: 25°C. Solvent: 25 g of 99.9% anhydrous DMSO.
2. Approximated to ΔHᵣ₋₃ (CsPbI₃, xl, 320°C) from differential scanning calorimetry.
thermodynamically stable with respect to its binary iodides below 320 °C and the α-phase above that transition temperature. Because of the rapid reversibility of the transition, the α-phase with desirable solar cell applications is metastable and generally not attainable under ambient conditions.

The ideal cubic perovskite structure of α-CsPbI₃ consists of a three-dimensional network of corner-sharing BX₆ octahedra ([PbI₆]⁴⁻) with the Cs⁺ cation in the 12-fold coordination site formed in the middle of the cube of eight such octahedra.¹⁵–¹⁷ δ-CsPbI₃ is not a perovskite. Its structure includes double chains of edge-sharing [PbI₆]⁴⁻ octahedra linked by 9-fold coordinated Cs⁺ cations (Figure 1A).¹⁵–¹⁷ The edge-sharing [PbI₆]⁴⁻ octahedra are tilted relative to the corner-sharing octahedra in α-CsPbI₃.¹⁶ The difference in structures is manifested in the rather large enthalpy and entropy change associated with the α–δ transition, much larger than those associated with typical distortional transitions in perovskites.¹⁸,¹⁹ It is interesting that, despite such structural and energetic differences, the high temperature α-phase reverts rapidly to the δ-phase upon ambient cooling. The rapid reversibility suggests the possibility of a concerted, rather than reconstructive, mechanism for the transition, but this has not been investigated.

An analogous stability trend was observed in CsSnI₃, CsPbI₃’s tin counterpart, which undergoes a phase transition between orthorhombic and cubic polymorphs as temperature changes.²⁰ Likewise, cubic CsSnI₃ is a high temperature phase while the orthorhombic polymorph is a low temperature one, and a reversible displacive phase transition between the two polymorphs was reported. Yet there is no comprehensive investigation of CsSnI₃ from the energetics perspective, suggesting a potential subject for future research.

This octahedral tilting in the δ-phase CsPbI₃ may cause the band gap (E_g) to rise from 1.71 to 2.82 eV. Such octahedral tilting is not uncommon: in hybrid organic— inorganic perovskites, similar octahedral tilting and corresponding band gap increase were also widely observed.²¹,²² Octahedral tilting was reported to reduce metal-halide orbital overlap, moving the valence band to lower energy and increasing the band gap.²¹ However, since the δ-phase is not, strictly speaking, a perovskite, additional factors, including the edge-sharing of octahedra, may be important in determining the band gap.

Clearly, as manifested in the 320 °C phase transition, there is a thermodynamic competition between the high temperature α-phase (more symmetrical and higher enthalpy and entropy perovskite) and the low temperature δ-phase (structurally more complex and less symmetrical nonperovskite of lower enthalpy and entropy). Such low temperature instability of perovskites relative to other phases may be a general phenomenon. Even within perovskites, transitions to more distorted polymorphs at lower temperature are common. Such distorted structures may have desirable properties resulting from their lower symmetry (including loss of a center of symmetry, leading, for example, to ferroelectric behavior, or they may compromise desirable band structure, as is the case for CsPbI₃. The distorted structures are of lower energy and lower entropy than the cubic aristotype, making the latter a high temperature polymorph. Similar arguments hold for stability relative to nonperovskite polymorphs. Such thermodynamic relations have been studied extensively in the context of high pressure geoscience, where MgSiO₃ based perovskite dominates the mineralogy of the Earth’s lower mantle. This silicate perovskite is of higher

enthalpy and entropy than polymorphs like MgSiO₃ ilmenite (which contains edge sharing polyhedra).²³ The reason for this behavior arises in large part from the vibrations of relatively loosely bound cations in the perovskite A-site which provide a large contribution to the low frequency vibrational density of states and lead to higher heat capacity and entropy.²³–²⁶ The present study suggests similar behavior in halide perovskites. Future computational studies could address this phenomenon in CsPbI₃ perovskite despite the inability to maintain the perovskite at low temperature. It is also possible that cryogenic quenching in the absence of air and water vapor could preserve enough perovskite for experimental studies by low temperature heat capacity measurements, spectroscopy, and inelastic neutron scattering.

In summary, by using a combination of room temperature solution calorimetry and differential scanning calorimetry, we have obtained the enthalpies of formation of α- and δ-CsPbI₃, shown that both are stable with respect to their binary halides, and obtained the thermodynamics of the α–δ transition. The desirable α-phase is metastable with respect to the δ-phase under ambient conditions. The effects of structural change on the thermodynamics and band gap have been discussed. The thermodynamic instability of perovskite phases at low temperature, reflecting their high vibrational entropy, may be a general phenomenon for halides as well as oxides.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.9b05924.

Experimental details of material synthesis, structural characterization, phase transition measurement, and solution calorimetry, thermal analysis (full pattern DSC and TG) plot (PDF)

**AUTHOR INFORMATION**

**Corresponding Author**

*ananvrotsky@ucdavis.edu*

**ORCID**

Novendra Novendra: 0000-0003-0927-5759
Alexandra Navrotsky: 0000-0002-3260-0364

**Notes**

The authors declare no competing financial interest.

**ACKNOWLEDGMENTS**

The authors thank Lili Wu for helpful discussions in designing experiments, Shuhao Yang for assistance with crystal structure visualization, and Tamilarasan Subramani for discussion during paper revision. The work was supported by the U.S. Department of Energy Office of Basic Energy Science, Grant DE-FG02-03ER46053.

**REFERENCES**

(3) Hoffman, J. B.; Schleper, A. L.; Kamat, P. V. Transformation of Sintered CsPbBr₃ Nanocrystals to Cubic CsPbI₃ and Gradient


