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Chemical durability and degradation mechanisms of CsPbI₃ as a potential host phase for cesium and iodine sequestration

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Effective nuclear waste management of radioactive cesium and off-gas iodine from complex waste streams of used fuels is essential for the sustainable development of advanced nuclear fuel cycles. Once cesium and iodine are separated from their respective waste streams, host phases are required to immobilize them into a durable waste form matrix for long-term disposition. The inorganic metal halide perovskite, CsPbI₃, has a unique crystal structure capable of incorporating both cesium and iodine simultaneously. Exposure to groundwater in geological repositories is a long-term concern for waste forms, as this may cause corrosion and decrease the waste form's ability to retain radionuclides. In this study, we explore the potential of CsPbl₃ perovskite as a promising host phase to incorporate Cs and I, and investigate its chemical durability and degradation mechanisms in an aqueous environment. CsPbI₃ was synthesized through a solution-based method and was consolidated into dense pellets by spark plasma sintering. The chemical durability of the CsPbI₃ pellets was evaluated by static leaching tests in deionized water at different temperatures of 25, 58, and 90 °C. The elemental release mechanisms and surface alteration of the monolithic CsPbI₃ pellets were investigated. Both I and Cs displayed a non-congruent leaching behavior and faster release rates as compared to Pb, particularly at longer leaching durations and higher temperatures. At the initial leaching stage, a PbI₂ alteration layer formed on the surface of the pellet due to the rapid release of Cs and I, followed by the formation of a PbI(OH) alteration layer. The activation energies for both dissolution and diffusion controlled mechanisms were determined to be $44.90 \text{ kJ mol}^{-1}$ and $45.40 \text{ kJ mol}^{-1}$ for Pb, $27.10 \text{ kJ mol}^{-1}$ and $40.82 \text{ kJ mol}^{-1}$ for I and $24.27 \text{ kJ mol}^{-1}$ and 23.86 kJ mol⁻¹ for Cs, respectively. These results show a clear decrease in activation energies from Pb to I and Cs, suggesting a preferential release of I and Cs. The solution-based synthesis of CsPbI₃ as a host phase for Cs and I and the fundamental understanding of the chemical durability and degradation behavior will be useful for further exploring its application for immobilizing iodine and cesium into final durable waste forms for long-term geological disposition.

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Introduction

Effective management and disposal of nuclear wastes are crucial to the development of advanced nuclear fuel cycles. Materials are required for the storage and disposal of radioactive fission products either through chemical reprocessing of used nuclear fuels or direct disposition of wastes in geological repositories. Among complex high level waste streams, I-129 is of significant concern for long term disposal due to its long half-life ($t_{1/2} = 1.6 \times 10^7$ years), radiotoxicity, and highly mobile nature, especially when concentrated in a localized geological repository. I-129 is water-soluble, highly environmentally

mobile, and can easily enter the biosphere, making it a serious environmental hazard. $^{4-6}$ In geological repositories, I-129 is one of the key dose contributors after several hundreds of thousands of years of nuclear waste storage. With an increase in temperature, I-129 becomes increasingly volatile, undergoing oxidation and reduction, and forming soluble salts. 6,7 In addition, radioactive cesium Cs-135 ($t_{1/2}=2.3\times10^6$ years, a β emitter) and Cs-137 ($t_{1/2}=30.08$ years, a β/γ emitter) are also problematic radionuclides in terms of waste storage and disposal. Particularly, long-lived Cs-135 is of great concern for the long-term performance of a geological disposal system, 8,9 and the short-lived Cs-137 is a major heat generator for interim storage of nuclear wastes. 10 Similar to iodine, cesium is also highly radiotoxic and water soluble, and needs to be confined in stable and durable matrices.

I-129 may be released after fuel assemblies are removed from the reactor core, usually in gaseous effluents as I₂, methyl iodide (CH₃I), bound to aerosols,¹¹ or as off-gas during chemical

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reprocessing of used fuels.12,13 Wet scrubbing methods, including alkaline scrubbing,14 Mercurex process,15 Iodox process,16 fluorocarbon solvents,17 and electrolytic scrubbing,18 can be used to capture gaseous iodine using a liquid solvent, and the resulting products must be further processed into a solid waste form. The solid adsorbents for iodine capture are usually silver-based compounds such as AgZ (silver exchanged mordenite),19,20 AgA (silver impregnated alumina)21 and silver impregnated silica gel,22,23 and carbon-based solid adsorbents such as coal24 and graphene.25 Solid adsorbents remove iodine through the adsorption of iodine onto a solid compound and generally have higher iodine removal efficiencies and are more easily converted into a final waste product. Cesium may be extracted from spent nuclear fuels dissolved in nitric acid using the Fission Product Extraction Processes (FPEX). 26,27 Synthetic zeolite can also be used to capture cesium from the waste streams of the chemical reprocessing.28,29 The adsorbed iodine after capture from the off-gas streams and extracted cesium from the waste streams of chemical reprocessing must be incorporated into an intermediate host phase or eventually be consolidated into durable solid waste forms for long-term disposition.

Different intermediate host phases or waste forms were proposed for iodine and cesium immobilization and disposal. Garino (2011) suggested the use of AgI and silver-exchanged MOR (Ag-MOR) for iodine immobilization, either through combining it with bismuth-silicon-zinc (Bi-Si-Zn)-oxide glass to form composites or a glass shell structure for storage.30 However, there are environmental concerns with silver in waste forms for long-term storage as it is commonly reduced in subterranean environments similar to geological repositories, potentially liberating iodine from the waste form.³¹ Therefore, the silver-based materials can only serve as intermediate phases, not an ideal waste form material. Ceramic waste forms for iodine immobilization have also been studied, and their chemical durability and the confinement of iodine in ceramic host phases were investigated by leaching experiments. For example, Audubert (1997) proposed a core shell composite with lead-vanadate iodoapatite (Pb10(VO4)6I2) as the core and lead orthovanadate (Pb₃(VO₄)₂) as the shell, showing a good chemical durability.32 More recently, Zhang (2019) investigated leadvanadate iodoapatite ($Pb_{9.85}(VO_4)_6I_{1.7}$), where iodine leaching was conducted in water and 0.1 mol L⁻¹ Na₂CO₃, NaCl, Na₂SO₄. and Na₃PO₄ solutions, and high iodine release rates were observed especially in the basic and acidic solutions.33 There have also been ceramic waste forms proposed for cesium immobilization such as hollandite (AxByC8-yO16) which is one of the phases in multiphase Synroc assembly. Wang (2020) investigated the structure and the aqueous durability of a complex hollandite ($[Cs_xBa_y][Fe^{3+}_{2y+x}Ti^{4+}_{8-2y-x}]O_{16}$) (0.3 $\leq x, y$ \leq 0.75) in water at 90 °C and observed Cs leaching rates in the order of 10⁻³ g m⁻² d⁻¹ after 28 days.³⁴ Various amounts of Ba, the Cs daughter product, could be accommodated on the Cs site by substituting Fe³⁺ unto the Ti⁴⁺ site. Hollandite displays great flexibility in its design and low leaching rates and thus is a promising candidate for Cs immobilization.34,35

Metal halide perovskites are another class of ceramics and are being investigated for simultaneous iodine and cesium immobilization with very high waste loadings. The metal halide perovskites developed recently have been widely used in the optoelectronic field due to the benefits of having excellent carrier mobility and suitable band gap, without the drawback of sensitive organic compounds found in organic and organicinorganic hybrid solar cells.36,37 These perovskites have not been widely researched towards their applications in nuclear waste management. Recently, Yang (2020) investigated the leaching behavior of Cs₃Bi₂I₉ in a silica composite (70 wt%) and silica core-shell structure (20 wt% Cs₃Bi₂I₉).³⁸ The glass-bonded perovskite structure displays remarkably high waste loadings and excellent chemical durability, outperforming the state-ofthe-art waste form materials for iodine and Cs immobilization. An innovative concept was also reported recently using metal halide perovskite Cs₂SnCl₆ as a potential host phase to incorporate alkali metals and halides (e.g., Cs and Cl) separated from complex waste streams associated with the pyroprocessing of used fuels or molten salt reactors.39 The glassbonded Cs₂SnCl₆ was also proposed as an advanced waste form material for chloride-based salt wastes and displayed unprecedented high waste loadings and chemical durability. These pioneering works highlight the immense potential of using inorganic metal halide perovskites either as host phases to immobilize waste elements (particularly alkali metals and halides) or embedded into a glass or ceramic composite for geological disposition.

Analog to Cs₃BiI₉ and Cs₂SnCl₆ for iodine and Cl incorporation, CsPbI₃ is another derivative within the family of metal halide perovskites with rich crystal chemistry and structural flexibility. Fig. 1 shows the crystal structure of the δ -CsPbI₃. An ideal perovskite structure would comprise of 8 corner-sharing [PbI₆]⁻⁴ octahedra forming a cube structure with the 12-coordinated Cs⁺ cation occupying the central position within the cube. 40 However, the δ-CsPbI₃ structure consists of double edge sharing $[PbI_6]^{-4}$ octahedra that are slightly tilted with respect to the ideal perovskite structure and connected by the 9-fold coordinated Cs+ cation. CsPbI3 can be used as a host phase for separating cesium and iodine through a simple solution based approach from the wet scrubbing solution or liquid waste streams from chemical reprocessing, and the waste-loading perovskite host phase can be embedded into the glass or ceramic matrix to form composite waste forms. However, the CsPbI3 faces a similar problem as silver-based host phases, as Pb is also a regulated contaminant according to the Resource Conservation and Recovery Act (RCRA) and could be potentially released from the host phase.

In this study, we investigated the chemical durability and phase degradation behavior of monolithic CsPbI₃ as a host phase to incorporate cesium and iodine *via* static leaching tests. Waste element-loaded CsPbI₃ powders were synthesized through a scalable solution-based approach by direct precipitation and consolidated into dense pellets using spark plasma sintering (SPS). SPS was used to sinter the CsPbI₃ pellets due to its fast and uniform sintering process as well as its improved energy efficiency and reproducibility compared to techniques

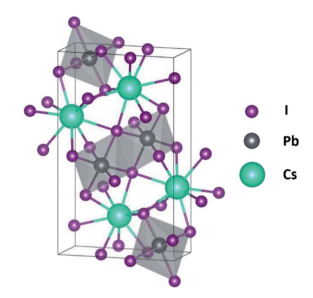


Fig. 1 Illustration of CsPbI₃ crystal structure, where Cs and I are indicated by the largest green sphere and smallest purple sphere respectively, and Pb is the grey sphere inside the octahedron.

such as hot press sintering and hot isostatic pressing.41 Static leaching tests were performed on the sample pellets in deionized water at three different temperatures (25, 50, and 90 °C) to measure the cumulative elemental release. The temperature dependent leaching rates were used to calculate the activation energies of the individual elemental release. Inductively coupled plasma-mass spectroscopy (ICP-MS) measurements were performed to quantify the elemental release in the sampled leachate, and post-leaching characterization was performed on the sample to identify the alteration layers that formed on the surface of the samples and investigate the mechanism of their formation. The results and analysis of these tests should provide a fundamental understanding of chemical durability and degradation mechanisms of the monolithic CsPbI₃ perovskite as the host phase of Cs and I, and will be useful for further exploring its application for advanced waste form design in composites.

2. Materials and methods

2.1 CsPbI₃ powder synthesis

The CsPbI $_3$ powders were synthesized by a chemical precipitation method using CsI (99.9%, Sigma Aldrich) and PbI $_2$ (99.9%, Sigma Aldrich) as precursor materials, and methanol was used as the solvent. 1 mmol of CsI was dissolved in 10 mL methanol and heated to \sim 60 °C in a water bath and 1 mmol of PbI $_2$ powders were dissolved in 4 mL n-butyl acetate in a separate beaker. The PbI $_2$ solution was then added to the warm CsI solution and stirred vigorously leading to a spontaneous precipitation of fine yellow CsPbI $_3$ powders. The chemical reaction equation during the process is as follows:

$$CsI + PbI_2 \rightarrow CsPbI_3$$
 (1)

The mixture was then stirred for 30 minutes with a stirring speed of 300 rpm to ensure the completion of the reaction in a 60 $^{\circ}$ C water bath. The solid CsPbI₃ precipitates were washed twice using *n*-butyl acetate *via* centrifugation and dried in an oven at 80 $^{\circ}$ C overnight.

2.2 CsPbI₃ pellet sintering

CsPbI₃ powders were then ground into fine powders and loaded into a 6 mm diameter cylindrical graphite die and consolidated into a dense pellet using SPS (Model 10-3 SPS system, Thermal Tech. LLC, Santa Rosa, CA). A steady flow of argon gas (1 L min $^{-1}$) was used during the sintering process to purge the chamber to avoid oxidation and degradation of CsPbI₃ and graphite die. CsPbI₃ samples were sintered at five different temperatures (300, 320, 350, 400, and 435 $^{\circ}$ C) all at 40 MPa for 1 minute to evaluate the effects of temperature on the microstructure of the CsPbI₃ pellets. The sintered pellets were then ground by SiC abrasive papers and then polished with 1 μ m diamond paste to achieve a mirror finish with anhydrous ethanol acting as the polishing agent. Both top and bottom surfaces were polished as the whole pellets were subjected to leaching conditions.

2.3 Crystalline CsPbI₃ sample structure and microstructure characterization

The phase composition of the CsPbI₃ pellet and synthesized powders were confirmed via X-ray diffraction (XRD) using a Panalytical X'Pert Pro system (Westborough, MA, USA) with a copper target ($K\alpha = 0.15406$ nm) and a step size of 0.013° . The physical density of the pellet was measured via the Archimedes technique using an Adam analytical scale (Danbury, NY, USA). The microstructures of the pellet samples were examined before and after the leaching experiments via digital photography, and scanning electron microscopy (SEM) using a FEI Versa (USA) with an energy-dispersive spectroscopy (EDS) system, conducted with an additional Oxford Instruments INCA detector (Abingdon, UK). Thermogravimetric analysis (TGA) was used to analyze the thermal stability of CsPbI3 using a TGA-Q50 system (TA instruments, New Castle, DE). ~20 mg of CsPbI₃ powders were weighed and placed in an alumina crucible, after which the sample was heated at a rate of 20 °C min⁻¹ in a steady flow of argon gas at 50 mL min⁻¹ until the temperature reached 1100 °C and then cooled at the same rate to 50 $^{\circ}\text{C}.$ A measurement of the Raman shift at the pellet surface was performed before and after the leaching experiment, using a Raman spectrometer with a 514 nm laser, an exposure time of 10 s, and an operating power of 20 mW.

2.4 Chemical durability test

Static leaching was conducted on the CsPbI $_3$ pellets sintered at 320 °C, where high-purity 18 M Ω deionized water was used as the leaching agent, following an ASTM C1220 leaching protocol. The pellet was placed in the leaching agent in a sealed PTFE container with the pellet surface area to volume ratio (S/V) kept at 5.0 m $^{-1}$ throughout the experiment (initial leachate volume is 46.1 mL based on the sample surface area). The static leaching experiments were conducted at room temperature (RT), 58 and 90 °C, and only 1 mL of leachate was

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removed from the PTFE container and replaced with the same amount of fresh leachate to minimize the impact on the concentration at the time intervals of 1, 2, 3, 4, 5, 6, 7, 8, 9, 10 and 68 h. After the experiment, all the leachates collected were filtered and acidified using 5 vol% HNO₃ for elemental analysis by ICP-MS. The ICP-MS Varian 820 was used to measure the concentrations of Cs, I, and Pb in the collected leachates to calculate the corresponding cumulative mass release m_i (g m⁻²) using eqn (2).

$$m_i = \frac{C_i V}{S} \tag{2}$$

where C_i is the concentration measured using ICP-MS (g L⁻¹), V is the leachate volume (L) and S is the surface area of the pellet (m²). The relation between the cumulative mass release and t_i leaching time (day) is given by an analytical Cote's model43 for the time evolution of the leaching behavior shown in eqn (3).

$$m(t) = k_1 t + k_2 t^{1/2} + k_3 (1 - e^{-k_4 t})$$
 (3)

Here, m(t) stands for the cumulative mass leaching out as mg m^{-2} . The first term on the right side of the equation, k_1t , represents a linear dissolution behavior. The second and third terms stand for the mass transport by diffusion and species exchange between the matrix surface and solution, respectively. The derivative of eqn (3) allows the determination of the leaching rate, and the parameter k_1 represents the long-term rate assuming an infinite leaching duration.

3. Results and discussion

3.1 Microstructure and thermal stability of as-sintered CsPbI₃ pellets under elevated temperatures

Fig. 2f presents the TGA analysis of CsPbI₃ powders, which shows a mass loss of greater than 90% at temperatures above 900 °C. The onset temperature of thermal instability for the assintered CsPbI₃ is approximately 510 °C at which materials start to experience phase decomposition and release of CsI, and almost all the weight was lost around 1000 °C. Therefore, all CsPbI₃ pellets were sintered at temperatures below the on-set point of 510 °C to avoid phase degradation during the sintering process. Fig. 2a-e shows SEM images of the fractured surfaces of the CsPbI₃ pellets sintered at 300 °C, 320 °C, 350 °C, 400 °C, and 435 °C, respectively, all at 40 MPa for 1 minute. The surfaces appear to be dense and relatively large grains \sim 2-10 μm can be seen, and grain sizes increase at increased sintering temperatures. While no apparent secondary phases are observed, intergranular cracking is apparent in the samples sintered at 350, 400, and 435 °C. The existence of intergranular cracking within the grain boundary might significantly reduce the grain stability as well as the sample integrity, which could further accelerate the water molecule diffusion into the pellets. Hence, the CsPbI₃ pellet sintered at 320 °C was used for the leaching experiment as it was sintered at the highest temperature without intergranular cracking.

3.2 Microstructure evolution and surface alteration of CsPbI₃ post static leaching

Fig. 3a shows the XRD patterns of the as sintered and post static leached CsPbI₃ pellets which can be indexed as the α-CsPbI₃ using PDF# 18-0376, and no impurity or secondary phases can be observed from the as-sintered pellet. The XRD patterns of the post leaching pellets at room temperature and 90 °C show strong PbI₂ (PDF# 07-0235) peaks and minor PbI(OH) (PDF# 22-0655) peaks, indicating an alteration of the CsPbI₃ structure with Cs depletion. Fig. 3b presents the Raman spectrum of the CsPbI₃ pellet after the 68 hour leaching test in deionized water. Strong Raman peak shifts are observed at 75 cm⁻¹ and 105 cm⁻¹, corresponding to the A1_g and E_g stretching modes of PbI₂ (ref. 44) respectively, as observed in the pellets leached at room temperature and 90 °C. Fig. 3c shows an optical image of the morphology of the leached surface and Fig. 3d is the EDS analysis of the alteration products. The quantitative EDS

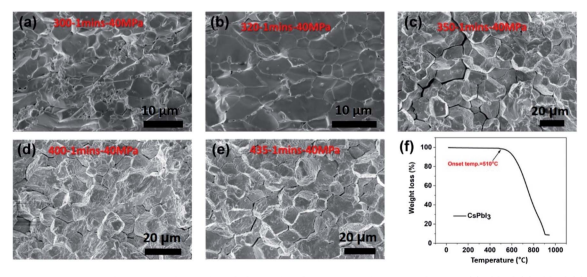


Fig. 2 SEM images showing the fractured surfaces of the SPS densified CsPbI₃ pellets for 1 min-40 MPa at (a) 300 °C (b) 320 °C (c) 350 °C (d) 400 °C and (e) 435 °C. (f) Thermogravimetric analysis of CsPbI₃ powders from room temperature to 950 °C

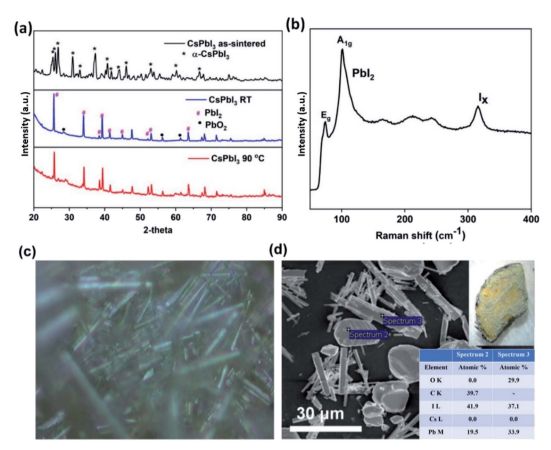


Fig. 3 (a) XRD diffraction patterns of CsPbI $_3$ pellets before and after leaching testing at RT and 90 °C; (b) Raman spectra of the leached pellet leached for 68 hours at 90 °C in deionized water; and (c) optical image of alteration product, (d) SEM image and EDS point analysis at the leached pellet surface post 90 °C leaching for 68 hours.

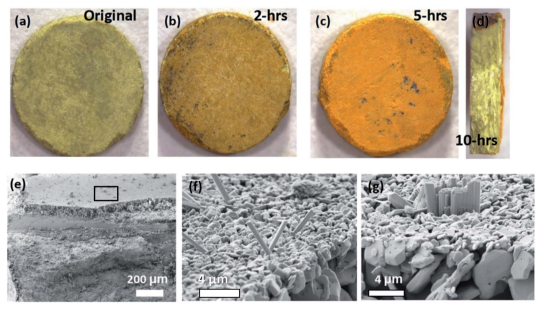


Fig. 4 Optical images showing the morphology, microstructure, and color evolution of the $CsPbl_3$ pellets (a) before leaching; (b) 2 hours; (c) 5 hours; and (d) 10 hours after leaching; (e) an SEM image of the fractured pellet cross-section; and (f and g) SEM images showing surface alteration for the post leaching pellet after 10 hours water exposure.

analysis of spectrum 2 indicates a Pb: I atomic ratio of 1:2; while spectrum 3 has a Pb: I: O ratio is approximately 1:1:1. No Cs is observed from both spectra confirming the depletion of Cs and the retention of Pb and I in the alteration layer, and the alteration products as inferred from spectra 2 and 3 correspond to PbI₂ and PbI(OH), respectively.

Fig. 4a–d are optical images showing the degradation morphologies of the CsPbI $_3$ pellets during various stages of the leaching experiments. The pellet changes in color from yellow to dark orange, suggesting severe phase degradation occurred as shown in Fig. 4a–c. The surface of the sample pellet post 10 hours leaching test is covered by an orange alteration layer as shown in Fig. 4d. The alteration layer uniformly covers the surface with a thickness of approximately 100 μ m, as

demonstrated in the SEM image in Fig. 4e. Fig. 4f–g show the surface microstructure of the altered CsPbI₃ pellets, indicating the formation of a loosely-connected alteration layer. From Fig. 4f–g, rod-like particles can be seen on a layer of small polyhedral grains which are layered on top of larger polyhedral grains. Based on the shape of these particles and EDS point analysis, it can be inferred the rod-shaped particles are PbI(OH) and the polyhedral particles are PbI₂ based on their crystal structures.

3.3 Chemical durability and leaching mechanisms of CsPbI₃

The water interaction induces the elemental release which can be quantified in Fig. 5. The elemental release rates (g m⁻² h⁻¹) for Cs, Pb, and I are shown in Fig. 5a, c and e; and the

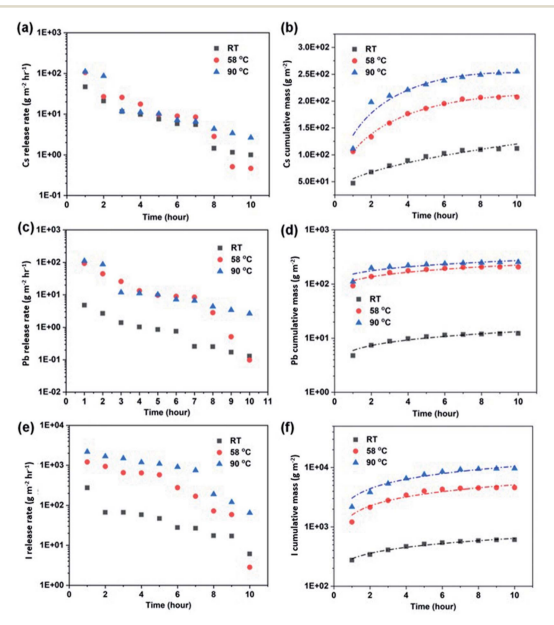


Fig. 5 Elemental release rates (a, c and e) and cumulative mass release rates (b, d, and f) at different temperatures for short-term static leaching for Cs, Pb, and I respectively.

cumulative mass release (g m $^{-2}$) of the corresponding elements are shown in Fig. 5b, d and f. The elemental release rate for CsPbI $_3$ increases when subjected to elevated temperature from 25 °C to 90 °C. For each element, Cs, Pb, and I, the elemental release rates gradually decrease with increased leaching duration, suggesting gradual saturation of the solution, commonly observed for static leaching tests. Moreover, I releases much faster than Cs and Pb, which can be attributed to the weak covalent bonding of Cs–I and Pb–I. Pb has the lowest elemental release rate, probably due to the formation of the surface alteration layer indexed as PbI(OH) (PDF# 22-0655), as confirmed by the XRD profile.

The corresponding atomic ratio can be seen in Fig. 6a–c, and an incongruent leaching mechanism dominates with the I/Cs ratio significantly higher than 3.0 for temperature ranges from 25 °C to 90 °C. The initial leaching of Cs and I can be attributed to the breakage of the weaker Cs–I bonds within the matrix. At higher temperatures, the atomic ratio of I : Cs as shown in Fig. 6b and c increases over time, with the I/Cs atomic ratio being the highest at 90 °C. The I/Cs ratio increases possibly due to the breaking of the stronger PbI $_2$ metal-halide bond. The formation of PbI(OH) alteration layer could lead to the decrease of the I : Cs ratio in the leachate. The initial atomic ratio of Cs/Pb is higher than the stoichiometric ratio of 1 : 1 for CsPbI $_3$ at 25 °C, indicating a lower release rate of Pb, consistent with the stronger covalent bonding of Pb–I compared to Cs–I. The

initial rapid dissolution of I and fast release of Cs compared to Pb leads to the formation of PbI₂ in the surface alteration layer as inferred from the XRD pattern of the leached surface. PbI₂ typically has a lower solubility compared to CsPbI₃, which may explain the lower Pb release rate compared to Cs. The bond breakage of Pb–I becomes more severe with time, leading to the change from non-congruent to congruent then back to a noncongruent release mechanism, as evidenced by the Pb/I ratio shown in Fig. 6.

Interestingly, the atomic ratio of Cs/Pb rapidly decreases past the stoichiometric ratio of 1.0 at elevated temperatures from 58 °C to 90 °C, indicating a preferential release of Pb at higher temperatures and longer duration, as the increased temperature and extended duration significantly accelerate the Pb-I bond breakage. The hydrolysis reaction between the PbI₂ surface product with water leads to the enhanced release of Pb and the formation of the hydrolysis product PbI(OH). Therefore, the Pb leaching rate increases and surpasses that of Cs as the surface layer is already depleted of Cs. For comparison, a transition from non-congruent to congruent, followed by a noncongruent leaching mechanism can be seen for I/Pb at all temperatures, with the initial I/Pb atomic ratio significantly higher than 3.0, since the covalent bonding of Cs-I is much weaker than Pb-I. The rapid initial release of I and Cs leads to a surface depleted in both elements, which leads to the initial formation of PbI₂ at the pellet surface, and the release of I from

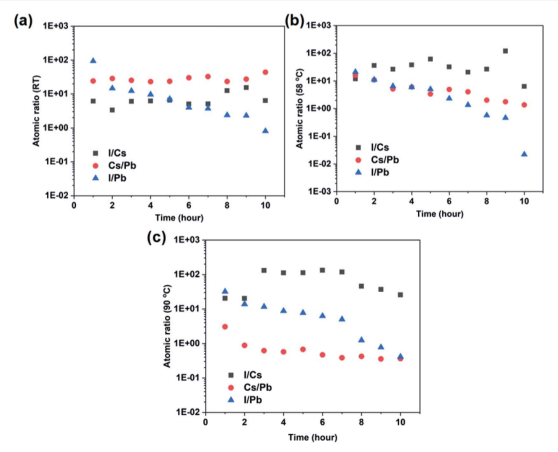


Fig. 6 The atomic ratios of I/Cs. Cs/Pb. and I/Pb in bulk solutions over time at RT (a), 58 °C (b), and 90 °C (c) as measured by ICP-MS.

 PbI_2 leads to the formation of PbI(OH). The insoluble PbI(OH) (PDF# 22-0655) alteration layer further decreases the release rate of I from the pellet surface.

The rate constants for both dissolution-controlled and diffusion-controlled mechanisms were determined by fitting the cumulative mass concentration using Cote's model with the corresponding fitting curves shown in Fig. 5b, d and f, where the fitting parameters are listed in Table 1. The cumulative mass curves are well fitted using Cote's model as supported by the R^2 values on average being higher than 0.9. The results and fitting parameters indicate that the CsPbI₃-water interaction behavior can be well explained by three distinct mechanisms, namely, dissolution, surface effect, and diffusion. The dissolution rate and diffusion rate are denoted by k_1 and k_2 in Table 1, respectively. The cumulative mass curves confirm the gradual reduction of elemental release rates due to the solution saturation during the static leaching test. Particularly for the iodine, this saturation might result in the reprecipitation as evidenced by the reduction in I concentration. With the re-precipitation of the hydrolysis product of PbI(OH) layer on the pellet surface, the elemental releases are continuously reduced (Fig. 5).

A key variable in understanding the diffusion-dominated process is its temperature dependence which may affect the diffusion coefficient in a similar manner to the reaction rate constant.⁴⁶ The variation of dissolution and diffusion coefficients with temperature can be well described using the Arrhenius equation below,⁴⁷

Table 1 Fitting parameters for Cs, Pb, and I calculated from the short-term cumulative mass release rates using Cote's model shown in eqn (3)

	Fitting parameters	RT	58 °C	90 °C
Cs	k_1	4.6	17.9	26.0
	k_2	29.9	123.2	162.3
	k_3	25.4	0.0	0.0
	k_4	1.0	1.0	1.0
	$rac{k_4}{R^2}$	0.9	1.0	0.9
Pb	k_1	0.6	1.0	17.0
	k_2	3.4	50.8	55.2
	k_3	2.6	64.4	98.0
	k_4	1.0	1.0	1.0
	R^2	0.9	0.9	0.9
I	k_1	16.4	52.0	115.7
	k_2	162.0	1626.9	2947.3
	k_3	127.5	0.0	0.0
	k_4	1.0	1.0	1.0
	R^2	1.9	0.9	1.0

$$R(T) = Ae^{-E_a/RT} (4)$$

where R(T) is the dissolution (k_1) or diffusion-controlled (k_2) rate constants at temperature T, which are obtained by fitting the short-term cumulative mass release rates (summarized in Table 2) for each element at all three temperatures (Fig. 5b, d and f) using Cote's model. A is an unknown constant and E_a (kJ) is the apparent activation energy, both of which are independent of temperature, and R is the gas constant $(8.3145 \times 10^{-3} \text{ kJ K}^{-1} \text{ mol}^{-1})$. By plotting the natural logarithm of the k_1 or k_2 rate constant versus the inverse of absolute temperature, a linear relationship is obtained (Fig. 7a–b). The slope of this linear relation is then used to calculate the activation energy associated with either the dissolution or diffusion mechanisms for each element, which are listed in Table 3.

Since the elements in the perovskite are not released congruently, it is expected that their activation energies vary accordingly, with Cs having the lowest values of 24.27 kJ mol⁻¹ and 23.86 kJ mol⁻¹ for dissolution control and diffusion control mechanisms, respectively. This is consistent with the experimental observation in which Cs is preferentially released quickly. Pb has the highest activation energy for both the dissolution and diffusion controlled mechanisms, supporting the observation of the high non-stoichiometric Cs/Pb atomic ratio at room temperature.

In contrast, the activation energy of dissolution for iodine release is significantly lower than that of a diffusion-controlled process, which indicates that elemental diffusion needs to overcome a higher energy barrier before it can occur. This suggests that the iodine release from the CsPbI₃ is mainly controlled by the dissolution process upon the bond breaking of Cs–I and Pb–I. The activation energy of dissolution for iodine release is also close to that of Cs, and thus the I/Cs atomic ratio remains relatively close to its stoichiometric value. For Cs and Pb, both dissolution and diffusion activation energies are close, indicating that both processes co-exist to dominate their release rates.

While CsPbI₃ has the benefit of over 50 wt% iodine waste loading, compared to other glass and apatite-based waste forms for iodine incorporation investigated by Coulon *et al.* (2017), Chong *et al.* (2018) and Lee *et al.* (2017), the iodine cumulative leaching from CsPbI₃ is significantly high.^{48–50} Cesium loading is also much higher in the CsPbI₃ perovskite than the more commonly studied hollandite structure, and the release rate of Cs is significantly higher than other hollandite waste forms previously studied by Wang *et al.* (2020) and Yang *et al.* (2019).^{34,51} Therefore, CsPbI₃ itself cannot be used as a durable

Table 2 Dissolution and diffusion-controlled release rates of Cs, Pb, and I at different temperatures

	Dissolution controlled			Diffusion controlled		
°C	Cs $(g m^{-2} h^{-1})$	Pb (g $m^{-2} h^{-1}$)	$I (g m^{-2} h^{-1})$	Cs (g $m^{-2} h^{-1}$)	Pb (g $m^{-2} h^{-1}$)	I (g m ⁻² h ⁻¹)
25	4.6	0.6	16.4	29.9	3.4	162.0
58	17.9	1.0	52.0	123.2	50.8	1626.9
90	26.0	17.0	115.7	162.3	55.2	2947.3

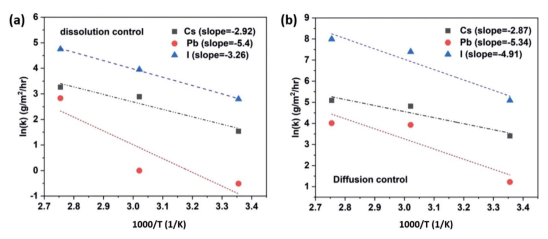


Fig. 7 (a and b) Plots of $\ln(k)$ vs. 1000/T (1/K), where k is the dissolution or diffusion-controlled rate constants determined by fitting of the Cote's model in the of units of g m⁻² h⁻¹.

Table 3 Calculated activation energies, $E_{\rm a}$, for each element in kJ mol $^{-1}$ based on the Arrhenius equation

	Dissolution	Diffusion	
Activation energy	kJ mol ⁻¹	kJ mol ⁻¹	
Pb	44.90	45.40	
I	27.10	40.82	
Cs	24.27	23.86	

waste form matrix for iodine and Cs immobilization because of its lower chemical durability. On the other hand, a high amount of Cs and I can be incorporated into the CsPbI₃ structure, after which CsPbI₃ can be embedded into either a low temperature glass or a ceramic matrix to form a composite waste form. This concept has already been demonstrated by Yang *et al.* (2021) and may further improve the chemical durability of CsPbI₃.³⁹ Additionally, it is expected that the formation of more water insoluble alteration layers such as PbI₂ or PbI(OH) will greatly benefit the retention of iodine and cesium and reduce their release rates. In conjunction with a composite matrix, this could unlock the potential of using inorganic metal halide perovskite as a promising host phase for the immobilization of cesium and iodine.

4. Summary

In this study, we explore the $CsPbI_3$ perovskite as a potential host phase for incorporating high amounts of Cs and iodine by investigating its chemical durability and degradation mechanisms. The waste-element loaded α -CsPbI $_3$ can be synthesized from a scalable solution-based approach at a low temperature, and consolidated into a dense pellet at 320 °C via SPS without phase decomposition and release of volatile Cs and I. Static leaching testing of the SPS-densified CsPbI $_3$ shows susceptibility of the material to water dissolution, leading to a significant release of waste elements. Mechanistic understandings of non-congruent dissolutions and time evolution of kinetics

based on solution chemistry analysis are achieved. This incongruent dissolution mechanism is also evident in the elemental release measurements as Cs is preferentially released compared to Pb at the initial stage of the leaching test at all temperatures. Surface alterations of a PbI₂ alteration layer upon initial rapid dissolution of Cs and I occur, followed by a hydrolysis reaction and the formation of PbI(OH) at longer durations. Activation energies for element releases are determined, being the highest for Pb followed by I and Cs. A dissolution dominated mechanism is identified for iodine release through bond-breaking of Pb–I and Cs–I. Advanced materials designs by embedding the waste-loaded host phase with high chemical loadings into a highly chemical durable matrix are required to utilize unique merits of perovskite structures for effective separation and immobilization of problematic Cs and I.

Conflicts of interest

There are no conflicts to declare.

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