



Cite this: DOI: 10.1039/d5el00157a

Evaluating the potential of CsBiSCl₂ as a solar absorber

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Efforts to develop lead-free and stable alternatives to halide perovskites have thus far mostly yielded materials with power conversion efficiencies (PCEs) well below 10% in solar cells. Recently, photovoltaics based on CsBiSCl₂ were reported to achieve 10.38% PCE. Still, the crystal structure is unknown, and it is unclear whether the reported thin film synthesis method could realize thin films with the desired phase and stoichiometry. Herein, we use *ab initio* Random Structure Searching (AIRSS) with a bespoke machine learned interatomic potential to explore the potential energy surface of CsBiSCl₂, finding the previously-proposed cubic perovskite structure to be implausible. The lowest-energy structure we find is a four formula unit orthorhombic structure (*Pnma* space group) that lies 2.4 meV per atom above the convex hull. There is strong competition in the Cs–Bi–S–Cl family, which can lead to phase impurities. By examining the reported solution synthesis method, we find that it is challenging to obtain the dimethylammonium bismuth sulfide intermediate product, and that Bi₂S₃ with dimethylammonium iodide on the surface likely forms instead. The significant I-containing residues in this intermediate results in Cs₃Bi₂I₉ being preferentially formed in thin films instead of CsBiSCl₂. Solid state synthesis without I present leads to phase impurities, consistent with the lowest-energy CsBiSCl₂ phase being metastable. Taking these experimental and computational results together, it is unlikely that >10%-efficient CsBiSCl₂ solar cells have been achieved.

Received 23rd September 2025
Accepted 26th October 2025

DOI: 10.1039/d5el00157a

rsc.li/EESolar

Broader context

Solar absorbers that are simultaneously efficient and able to be manufactured cost-effectively have long been sought after. Lead-halide perovskites (LHPs) were a surprising discovery that fulfils these requirements, and are now entering into commercial production. However, the toxicity and limited ambient stability of LHPs have prompted a search for alternatives that could mimic their exceptional optoelectronic properties, whilst overcoming their limitations. Most ‘perovskite-inspired’ materials thus far have yielded power conversion efficiencies (PCEs) <10% in solar cells. It is therefore remarkable that CsBiSCl₂ was recently reported to achieve 10.38% PCE under 1-sun illumination, and maintain 97% of its original PCE after 150 days of storage in air without encapsulation. CsBiSCl₂ is also free from any toxic elements, and is reported to have a bandgap of 2.0 eV, which is ideal for applications as the top-cell in tandem photovoltaics, for indoor photovoltaics, or for photoelectrochemical cells. However, the structure and synthesis of the material are poorly evidenced and understood. Herein, we use computational investigations to rigorously determine the lowest-energy structure for CsBiSCl₂, and we show that the original synthesis method reported does not yield CsBiSCl₂ thin films. We find the claimed 10.38% PCE to be spurious and recommend that the field avoids investing research efforts towards this material. In particular, we find that the lowest-energy structure of CsBiSCl₂ is above the convex hull, making it challenging to avoid phase impurities from the Cs–Bi–S–Cl family of materials.

Introduction

Developing solar absorbers that are simultaneously efficient, stable, comprised of nontoxic, earth-abundant elements, and can be produced cost-effectively has long been an elusive target of the broad thin film photovoltaics community. This effort was disrupted by the discovery that lead-halide perovskites (LHPs)

can achieve effective optoelectronic properties (long diffusion lengths, high luminescence quantum yields, and low Urbach energies) using low-cost, low-temperature fabrication methods. This strongly contrasts with traditional semiconductors, such as crystalline silicon (c-Si) or III–V compounds.^{1–3} Not only have LHPs now been realized in photovoltaics with similar power conversion efficiencies (PCEs) as industry-dominant c-Si solar cells (certified PCEs of 27.0% for LHP, and 27.8% for c-Si at present),⁴ LHPs have catalyzed the development of new classes of solar absorbers. These materials are termed ‘perovskite-inspired’, and include compounds that are structurally analogous (*e.g.*, halide elpasolites or double perovskites),⁵ chemically analogous (*e.g.*, tin perovskites),⁶ or electronically analogous

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(e.g., NaBiS₂).⁷ A key property sought for is defect tolerance, where low non-radiative recombination rates are achieved despite high defect concentrations through low capture coefficients of the most common point defects, self-healing, or the formation of benign defect pairs.^{8,9} There has especially been emphasis on heavy pnictogen-based compounds, namely Sb- and Bi-based materials,^{10–12} which have stable valence ns² electrons that are believed to be conducive towards achieving defect tolerance.¹³ Although these materials overcome the toxicity and stability limitations of LHPs, their performance thus far has been limited, with the majority of materials realized in photovoltaic devices having PCEs well below 10%.^{14–16}

Remarkably, Huang *et al.* reported a PCE of 10.4% from solar cells based on CsBiS₂Cl₂.¹⁷ Surprisingly, this high PCE was achieved in the first report of this material in solar cells. A PCE of 10% is considered a threshold for a new photovoltaic material to hold promise for commercialisation. Indeed, looking back at the development of LHPs: although the first publication of this material in solar cells was reported by Miyasaka and co-workers in 2009,¹⁸ it was not until Snaith and co-workers reported a PCE exceeding 10% in 2012¹⁹ that the field began to gravitate towards this material.^{20–22} It is therefore significant that CsBiS₂Cl₂ was reported to immediately achieve >10% PCE in the first publication on this material for solar cells. Furthermore, this material was claimed to be substantially more environmentally stable than LHPs, with the PCE maintaining 97% of the original value after storage for 150 days without encapsulation in ambient air. CsBiS₂Cl₂ is also free from any toxic elements regulated by the Restriction of Hazardous Substances (RoHS) directive.²³ The bandgap of 2.0 eV for this material is close to the ideal value for indoor photovoltaics,^{24,25} as well as X-ray detectors,²⁶ and could also be well suited for solar water splitting.²⁷

However, the crystal structure of CsBiS₂Cl₂ is not known, and the original report of CsBiS₂Cl₂ solar cells did not use any refinement methods to fit the measured X-ray diffraction pattern with a proposed structure.¹⁷ Understanding the structure of a material is critical. The crystal structure defines essential properties, including the stability, as well as the electronic structure, which in turn affects the bandgap and charge-carrier mobility (through the effective mass). Determining the structure is mandatory for performing atomistic simulations, including defect calculations, and determining the upper limit in efficiency of the material. Therefore, it is paramount to first understand the crystal structure of CsBiS₂Cl₂ and verify the stoichiometry of the compound formed.

Huang *et al.* suggested that CsBiS₂Cl₂ adopts an $n = 1$ Dion–Jacobson perovskite structure, comprised of layers of BiS₂Cl₄ corner-sharing units, with Cs⁺ cations in between these layers (Fig. 1a, left).¹⁷ However, there is no evidence for this structure. The general formula for the Dion–Jacobson homologous series is A'A _{$n-1$} Pb _{n} X _{$3n+1$} (A' = 1+ or 2+ cation, A = 1+ cation, X = 1– anion).²⁸ Adapting this formula to the Cs–Bi–S–Cl quaternary system depicted in Fig. 1a left, where $n = 1$, gives the stoichiometry CsBi(S,Cl)₄, not CsBiS₂Cl₂. This CsBi(S,Cl)₄ stoichiometry cannot be achieved with the standard oxidation states for the cations (I for Cs, III for Bi) and anions (II for S, I for Cl) without

violating the rules of charge neutrality (details in Table S1, SI). If we take the structure as drawn in Fig. 1a left, then S : Cl should be 1 : 3, but this requires Bi to be in the IV oxidation state, which is unlikely given the stability of the 6s² electron pair. Therefore, the suggestion that the polyhedra in CsBiS₂Cl₂ form layers with a corner-sharing Dion–Jacobson structure is unfounded.

Alternative structures for CsBiS₂Cl₂ may be more plausible. There have been studies into other I–III–VI–VII₂ materials, with a particular focus on lead-free analogues of CH₃NH₃PbI₃. In these materials, the Pb²⁺ cation is replaced with trivalent Sb³⁺ or Bi³⁺ and one I[–] anion is replaced with a chalcogenide anion (S^{2–} or Se^{2–}) to maintain charge neutrality, forming a split-anion perovskite.²⁹ Films of CH₃NH₃SbS₂ have been reported with the suggestion that the phase adopts a near-ideal perovskite structure based on a near-ideal calculated Goldschmidt tolerance factor of $t = 0.99$.³⁰ The Bi analogue, CH₃NH₃BiS₂, has also been reported, although it was proposed that this material preferentially adopts an orthorhombic or rhombohedral lattice as $t < 0.9$.³¹ However, further experimental and density functional theory (DFT) studies found no evidence for the formation of I–III–VI–VII₂ perovskites, instead showing that the formation of the ternary A₃M₂X₉ phase, along with binary chalcogenides and halides, is more energetically favourable.³²

Quarta *et al.* attempted to prepare nanocrystals of CsBiS₂Cl₂.³³ They proposed a cubic perovskite structure for this material (Fig. 1a, right) by using CrystaLLM, which is based on an autoregressive large language model, trained on millions of crystallographic information files.³⁴ However, their synthesis attempts were unsuccessful, and CsBiS₂Cl₂ with a perovskite phase was not obtained experimentally, instead forming a mixture of Cs₃BiCl₆, Cs₃Bi₂Cl₉ and Bi₂S₃. Another possibility is the UFES₃ structure type. This has been reported for both AgBiS₂Cl₂ nanocrystals and polycrystalline powders of CuBiS₂Cl₂.^{35,36} This is a layered structure with alternating slabs of [As₂Cl₄]^{7–} distorted octahedra, and [BiS₂Cl₆]^{7–} bi-augmented triangular prisms. However, this structure has not yet been investigated as a possibility for the CsBiS₂Cl₂ phase.

Already, several review papers have cited CsBiS₂Cl₂ as an important prospect in lead-free photovoltaics due to the reportedly high performance and excellent cell stability.^{9,37–40} However, such claims are, as-yet, premature without understanding what the structure adopted is, and whether the reported thin film synthesis route¹⁷ is a valid approach to obtain this structure with the intended stoichiometry. It is critical to address these questions before the photovoltaics field dedicates more effort into this material.^{19–23}

In this work, we first adopt a systematic global structure search to determine the ground state structure of CsBiS₂Cl₂. The approach is *ab initio* Random Structure Searching (AIRSS), coupled with a bespoke Ephemeral Data Derived Potential (EDDP) to determine whether (meta-)stable CsBiS₂Cl₂ structures exist and if previously-proposed perovskite structures are energetically accessible. Next, we repeat the synthesis method reported by Huang *et al.* for thin films. We examine the composition, and phases present in the reaction intermediate and final thin film product through energy-dispersive X-ray spectroscopy (EDX), as well as Pawley refinement of the



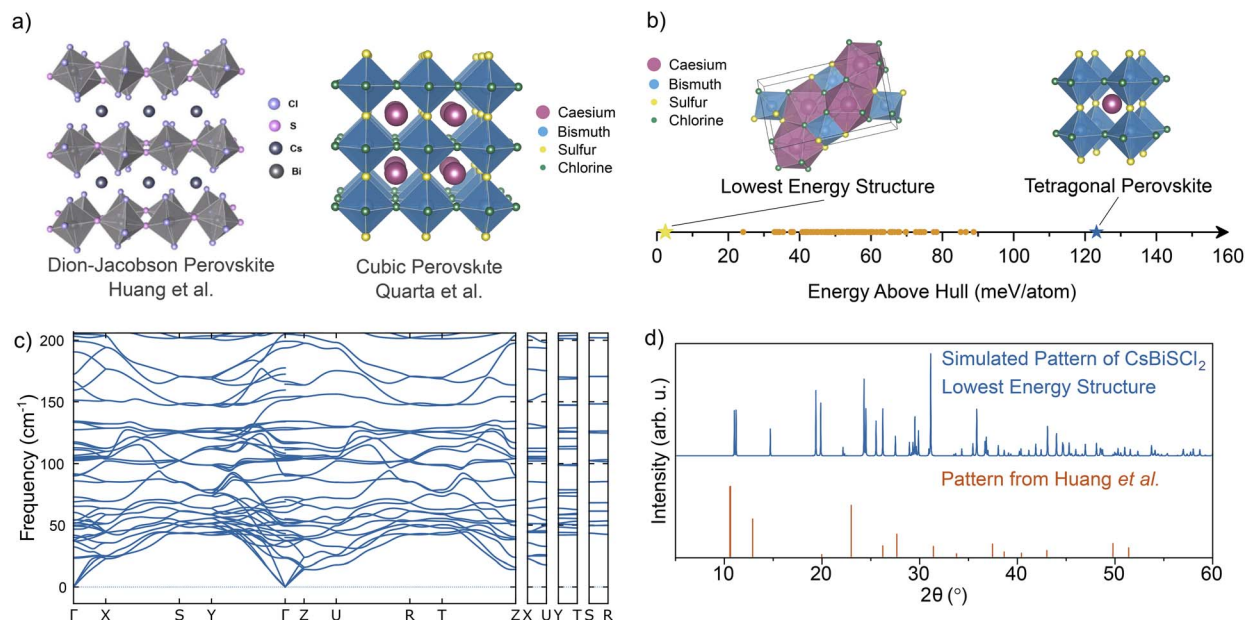


Fig. 1 Structure of CsBiSCl₂ (a) Proposed crystal structure of CsBiSCl₂ by Huang *et al.*¹⁷ and Quarta *et al.*³³ Illustration of structure proposed by Huang *et al.* reprinted with permission from J. Huang *et al.*, *J. Phys. Chem. Lett.*, 2024, **15**, 3383–3389. Cubic perovskite structure prepared from the information provided in ref. 33. (b) Energy above the Cs–Bi–S–Cl convex hull of CsBiSCl₂ structures selected for DFT refinalization. Shown are the lowest energy structure (yellow star), tetragonal perovskite structure (blue star), and remaining CsBiSCl₂ structures within 70 meV per atom of the hull (orange circles). (c) Phonon dispersion curve of the lowest-energy CsBiSCl₂ structure found from AiRSS. Lack of imaginary modes (*i.e.*, no negative frequencies) demonstrates the dynamic stability of this structure. (d) Comparison of the simulated powder X-ray diffraction (PXRD) pattern for the lowest energy *Pnma* structure and the CsBiSCl₂ PXRD pattern reported by Huang *et al.*¹⁷ Comparison with tetragonal perovskite reference pattern shown in Fig. S6, SI.

measured diffraction patterns with possible structures for CsBiSCl₂ and phase impurities. We use Fourier transform infrared spectroscopy (FTIR) to identify molecular species present, along with UV-visible (UV-vis) spectrophotometry to identify the absorption profiles of the thin film products formed. Given that the reported thin film synthesis route involves the use of HI, we compare with a solid-state melt synthesis approach that only uses the elements present in the target material, *i.e.*, using CsCl, BiCl₃ and Bi₂S₃ precursors. This work provides a thorough evaluation of the potential of CsBiSCl₂ as a solar absorber, and whether the wider photovoltaics field should dedicate further effort into this novel lead-free material.

Results and discussion

Identifying the lowest-energy structure of CsBiSCl₂

The AiRSS method involves iteratively generating ‘chemically sensible’ trial structures, relaxing them using an energy/force calculator, recording the result in a ranked database of energies, and repeating until some stopping criteria have been reached, *e.g.*, rediscovery of the same lowest energy structures multiple times (refer to Experimental section). Interesting structures, typically those within 100 meV per atom of the thermodynamic convex hull, can then be refined using precise density functional theory (DFT) settings.

For relatively simple materials (unaries, binaries) with small unit cells, it is tractable to carry out all geometry relaxations

using DFT. However, for compositionally complex materials where the search space is large, and/or those where a large unit cell is expected, traditional structure search soon becomes prohibitively expensive. Accelerating the structure search could be achieved using machine learning interatomic potentials (MLIPs). MLIPs can be classified as foundation or bespoke models, the former being trained on databases of diverse training data (*e.g.*, Materials Project’s MPtrj^{41–43}) with the goal of being applicable to arbitrary chemical systems, while the latter are trained on much smaller and narrower datasets, usually prepared by a single researcher, with a specific system in mind (*e.g.* Cs–Bi–S–Cl). Bespoke models are not transferable to arbitrary systems.

We initially attempted to use the foundation model Matter-Sim⁴⁴ to drive structural relaxations, but found it to have limited output structural diversity within the Cs–Bi–S–Cl system, as detailed in SI Note S1. We therefore trained a bespoke EDDP tailored to predicting the energy of CsBiSCl₂ structures. EDDPs are formulated specifically for random structure searching and, by design, avoid common pitfalls associated with modelling far-from-equilibrium structures using MLIPs.⁴⁵ To train the EDDP, the energies of 51 212 structures in the Cs–Bi–S–Cl space were calculated using DFT (PBEsol-D4). While the majority of structures were of the CsBiSCl₂ stoichiometry, the dataset also included all low-energy Materials Project structures within the Cs–Bi–S–Cl space, optimised perovskite CsBiSCl₂ structures, as well as a set of randomly generated unary, binary and ternary structures to improve the model fit. Crucially, this dataset

included structures spanning the entire search space, from high-energy to near-equilibrium ones. The as-trained EDDP had final training root mean squared error (RMSE) and mean absolute error (MAE) of 48.16 meV and 30.98 meV, while the testing RMSE and MAE were 235.94 meV and 42.40 meV.

The testing RMSE was raised by a single outlier with an error of >15 eV. While the global RMSE/MAE is quite high, this is mostly increased by non-stoichiometric and far-from-hull structures included in the dataset. The testing RMSE/MAE for CsBiSCL₂ structures within 1 eV of the hull are significantly improved at 22.7 meV per atom and 13.8 meV per atom respectively (Fig. S1, SI). Full details may be found in the Experimental section.

Using the as-trained EDDP, 92 770 CuBiSCL₂ structures with 2–10 formula units were generated and relaxed to their local minima. Of these, 3732 structures within 70 meV per atom of the convex hull were selected, and similar structures unified using a similarity threshold of 0.01, resulting in 118 unique low energy structures, after removal of one unphysical ‘low-energy’ structure (Fig. S2, SI). The lowest energy structure was rediscovered eleven times, and is a four formula unit *Pnma* structure (Fig. 1b). This structure was significantly lower in energy (by 28.2 meV per atom) than the next lowest-energy structure.

In an attempt to catch any low energy structures missed by standard AiRSS, 25 ps molecular dynamics (MD) simulations near the melting point of the *Pnma* structure were carried out for the top 10 000 structures (2778 after unification) using the bespoke MLIP and supercells of at least 40 atoms (so called ‘hot-AiRSS’ method).⁴⁶ While this reduced the number of unique structures after symmetrization and unification by 14% and resulted in the rediscovery of the *Pnma* structure another eight times; no lower energy structures were found. This supports the conclusion that the low-energy *Pnma* structure is dynamically stable. The structural density of states was flattened and broadened by this process, suggesting melting occurred in the far-from-hull regions (Fig. S3, SI).

The 118 unique low-energy structures, Materials Project (MP) near-the-hull structures, and the previously-proposed perovskite structures were selected for relaxation using precise DFT settings and more accurate Hamiltonian (*r*²SCAN-D3(BJ)+SOC). This high level of theory was chosen as the distance from the hull was within the margin of error set by our previous GGA/high-throughput DFT dataset. We found the lowest-energy *Pnma* structure predicted using EDDP-AiRSS remains the lowest energy CsBiSCL₂ structure in the refinalized DFT dataset and sits just 2.4 meV per atom above the Cs–Bi–S–Cl convex hull (Table S2, SI). This implies it is an energetically accessible structure, but no stable CsBiSCL₂ structures (*i.e.*, directly on the convex hull of the Cs–Bi–S–Cl system) were discovered.

To confirm the dynamic stability of the low-energy *Pnma* structure, a tight relaxation and subsequent harmonic phonon dispersion curve was calculated (Fig. 1c). We found that, despite the presence of low-lying optical modes near the *Z* point, the structure is dynamically stable with no imaginary modes present. Due to its proximity to the convex hull and its dynamic stability, the low-energy *Pnma* structure is a plausible meta-stable synthetic target. We estimate this structure has

a bandgap of >1.9 eV (Fig. S4, SI). Whilst this bandgap is suitable for top-cells in tandem photovoltaics, and for indoor photovoltaics, we note that using a meta-GGA functional would typically underestimate the bandgap, and there is a risk that the actual bandgap of the low-energy *Pnma* compound is substantially above 2 eV, making it not well suited for these applications. Furthermore, visual comparison of the simulated powder X-ray diffraction (PXRD) pattern of the low-energy *Pnma* structure with the PXRD pattern reported by Huang *et al.* for CsBiSCL₂ showed very little similarity (Fig. 1d).

Interestingly, we found the previously-proposed cubic perovskite structure to be energetically implausible. Upon relaxation, the cubic perovskite structure undergoes a second-order Jahn–Teller distortion, resulting in a tetragonal structure where bismuth atoms are off-centred in their S^{2−}/Cl[−] octahedra. This tetragonal perovskite has a large energy above hull of 123 meV per atom (Table S2, SI). In addition, visual comparison between the pattern reported by Huang *et al.* and the simulated pattern of the tetragonal structures shows little similarity (Fig. S6, SI). As such, we do not expect perovskite-structured CsBiSCL₂ to be experimentally accessible using standard synthetic approaches.

Investigation of the ‘DMABiS₂’ intermediate

The solution synthesis method reported by Huang *et al.* involved using CsCl and dimethylammonium bismuth sulfide (DMABiS₂) precursors, dissolved in a mixture of *N,N*-dimethylformamide (DMF) and dimethyl sulfoxide (DMSO), from which thin films of CsBiSCL₂ were spin coated.¹⁷ DMABiS₂ rather than Bi₂S₃ was used as the Bi and S precursor because it was suggested that this offers greater solubility in polar organic solvents.

The synthetic route to this ‘DMABiS₂’ intermediate involves introduction of the dimethylammonium cation (DMA⁺) to Bi₂S₃ through *in situ* acid hydrolysis of the DMF amide bond, catalyzed by HI (Fig. 2a). The generation of DMA⁺ cations through DMF hydrolysis has been previously applied to the synthesis of Cs_{1−x}DMA_xPbI₃ hybrid perovskites.⁴⁷ There was no suggested structure for the previously unreported DMABiS₂ intermediate. Bi₂S₃ has a 2D layered (lamellar) structure with layers connected by weak van der Waals interactions, with an interlayer spacing of 0.23 nm.^{48,49} Insertion of cations between the layers has been used to prepare intercalation compounds of Bi₂S₃ that have potential applications as battery materials.^{50,51} It may be the case that DMABiS₂ is an intercalation product, with DMA⁺ cations between the layers of Bi₂S₃. If this were the case, it is to be expected that PXRD peaks assigned to Bi₂S₃ should shift to lower 2θ values.

To synthesize the DMABiS₂ intermediate, we mixed 1.8 g Bi₂S₃ powder (3.5 mmol) with 3 mL DMF solvent in an N₂-filled glovebox, according to the method from Huang *et al.* We found that full dissolution of Bi₂S₃ did not occur, nor during the addition of 2.5 mL HI dropwise to this mixture. After mixing for 4 h at 500 rpm, we used Büchner filtration of the mixture inside a fume hood, obtaining a dark brown intermediate product, which is consistent with the report from Huang *et al.*¹⁷ But we also



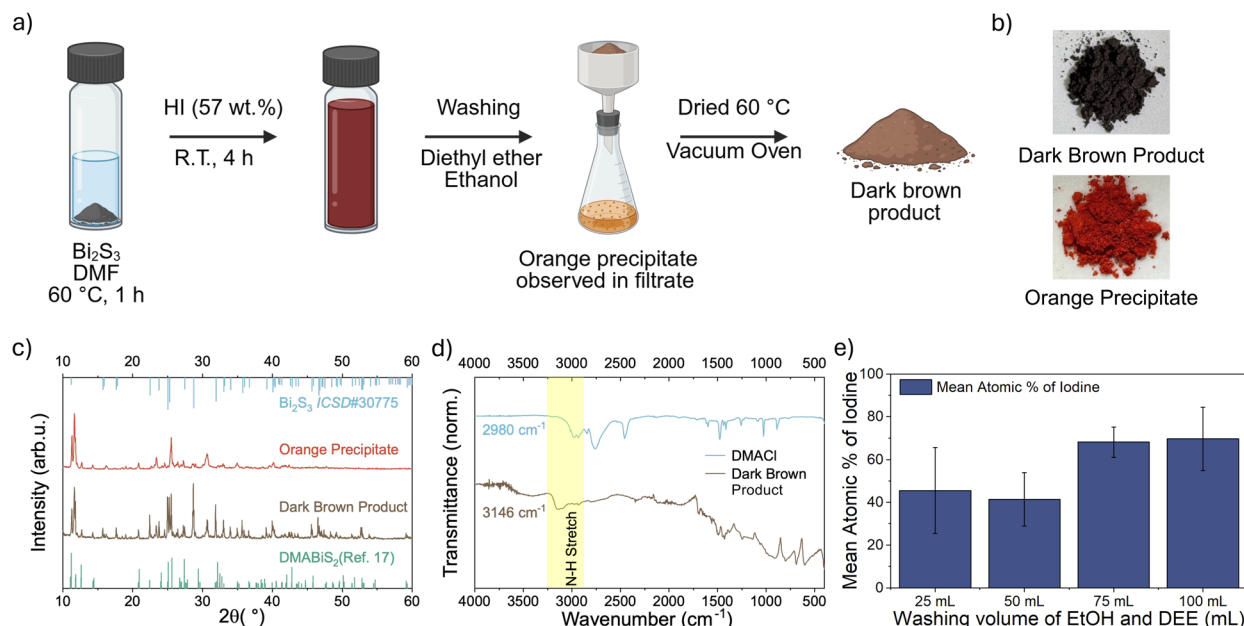


Fig. 2 Preparation of DMABiS₂ intermediate. (a) Illustration of the synthesis procedure for preparation of the DMABiS₂ intermediate. Created in BioRender. Quinn, E. L. (2025) <https://BioRender.com/xo6o9y9>. (b) Images of the dark brown product powder and the orange precipitate observed in the filtrate during the washing stage. (c) PXRD pattern of synthesized DMABiS₂ material. The light blue bars indicate the reference patterns of Bi₂S₃ (ICSD #30775), while the light green bars represent the reported pattern by Huang *et al.* for their claimed DMABiS₂ intermediate. (d) Infrared spectra of the dark brown powder and DMACI reference material. The peaks assigned to the ammonium N–H stretch are highlighted. The increase in peak wavenumber for the dark brown powder sample is attributed to the reduction in electronegativity of the environment leading to stronger N–H bonds. (e) The effect of solvent washing volume on the mean atomic percentage of iodine from EDS measurements of dark brown product powders. The error of atomic percentage is calculated using procedure described in Fig. S12, SI.

obtained an orange precipitate from the filtrate after washing with ethanol. This was not reported in the original synthesis. Fig. 2b shows photographs of both the dark brown product powder and the orange precipitate recovered from the filtrate.

We measured the PXRD pattern of the dark brown product and orange powder from the filtrate, and analyzed through Pawley fitting (Fig. 2c). This refinement method fits the peak positions according to the structure files input, but numerically fits the peak intensities. Pawley fitting is therefore suitable for obtaining lattice parameters and identifying whether the measured pattern is accounted for by the proposed structures, but does not solve the atomic coordinates. This is an appropriate step for determining the phase purity of the powder before performing Rietveld refinement to obtain atomic coordinates, ideally from a phase-pure sample that does not have preferred orientation.

Given that the structure of DMABiS₂ has not been solved, we first extracted the reported XRD peaks of this intermediate from ref. 17, and visually compared against the PXRD pattern for our dark brown product. As shown in Fig. 2c, there is a good match to all of the reference DMABiS₂ peaks, especially the strong peaks at 12° 2θ, which were claimed by Huang *et al.* to “confirm” the formation of DMABiS₂.¹⁷ There are, however, some peaks present in our dark brown product not present in the previously-reported pattern for DMABiS₂, suggesting that there are other phases present, or that we do not have a DMABiS₂ intercalation product.

We tested the hypothesis that the dark brown product is Bi₂S₃, performing Pawley fitting (Fig. S7a, SI) using the structure file from the Inorganic Crystal Structure Database (ICSD), collection code: 30775. As shown in Fig. 2a, there is a good match, with a goodness of fit (GoF) of 3.13. However, residuals are present, notably at 12°, 20°, and 30° 2θ. We therefore tested the hypothesis that a second phase was present: dimethylammonium iodide (DMAI). This improved the refinement (Fig. S7b, SI), with a goodness of fit of 2.99, although the key residuals at 12°, 20°, and 30° 2θ are still present. Therefore, we propose that the dark brown product we prepared is likely to contain two separate phases: Bi₂S₃ and DMAI, and possibly with a third unidentified phase. None of the peaks assigned to the Bi₂S₃ phase show a shift in 2θ position compared to the reference pattern. We therefore conclude that it is unlikely that there is an intercalation product.

SEM and EDX measurements were performed to aid the characterization of the prepared material. We observed two distinct phases: larger particles with a layered surface, and smaller particles dispersed across the surface of the larger particles. EDX analysis (Fig. S8 and Table S3, SI) suggested that the layered material was Bi₂S₃, with I impurities. The smaller particles contained a significant proportion of I and higher levels of N, suggesting that these smaller particles are likely to be primarily DMAI. This provides further evidence that no uniform DMABiS₂ phase was synthesized, but that the reported synthetic route prepares Bi₂S₃ with DMAI surface contamination, along with I residues throughout the sample. We



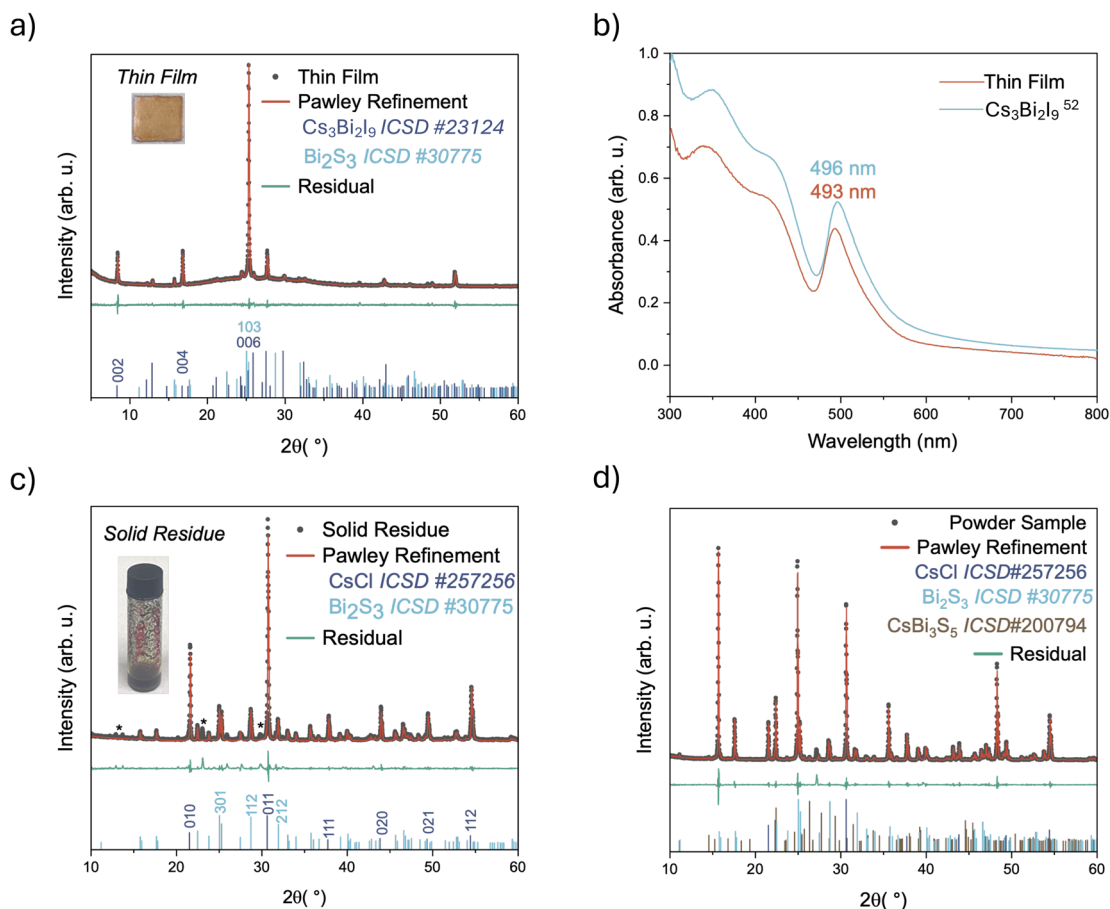


Fig. 3 Challenges with CsBiSbCl₂ synthesis (a) PXRD pattern of thin films synthesized according to Huang *et al.* (solid circles), with the corresponding Pawley refinement to Cs₃Bi₂I₉ and Bi₂S₃ (red line); $R_{wp} = 9.19$ GoF = 1.21. Inset: photograph of the as-prepared thin film on 1.4 cm² glass. (b) UV-vis absorption spectrum of prepared thin films and Cs₃Bi₂I₉ reference spectrum,⁵² with the characteristic Cs₃Bi₂I₉ excitonic peaks labelled. (c) PXRD pattern of solid residue from precursor solution from the attempted synthesis of CsBiSbCl₂ (solid circles), with the corresponding Pawley refinement to Bi₂S₃ and CsCl (red line); $R_{wp} = 13.00$ GoF = 2.14, unidentified peaks are labelled by *. Inset: photograph of the solid residue in a glass vial. (d) PXRD pattern of material from attempted solid-state synthesis of CsBiSbCl₂ (solid circles), with the corresponding Pawley refinement to CsCl, Bi₂S₃, and CsBi₃S₅ (red line); $R_{wp} = 10.97$, GoF = 1.74.

considered the possibility that BiSI may be present, however, Pawley refinement to BiSI structure file (ICSD collection code: 133376) revealed a poor match to this phase with a GoF of 4.79 (Fig. S9, SI). This allowed us to discount the presence of a BiSI phase.

Infrared (IR) spectroscopy data confirms the presence of an ammonium species in the dark brown product powder. Fig. 2d shows the IR spectrum of the material alongside the IR spectrum of reference DMAI material. The peak at 3146 cm⁻¹ is assigned to the N-H stretch of the DMA⁺ cation. This is at a higher wavenumber than the corresponding N-H stretch peak in the DMAI reference (2980 cm⁻¹). We attribute this observation to the reduction in electronegativity of the environment surrounding the DMA⁺ cation, since replacing Cl with I in DMAI would increase the electron density in the N-H bonds, making them stronger and oscillate faster.

We also analyzed the orange precipitate obtained from the filtrate. Its PXRD pattern is broadly similar to the pattern of the dark brown product, but significantly less crystalline, especially at higher 2θ values. We therefore concluded that this orange

precipitate was not the desired DMABiS₂ intermediate material. IR measurements of the orange precipitate (Fig. S10, SI) showed no distinct functional group features, providing further support that this precipitate was not the desired ammonium bismuth sulfide intermediate.

A significant limitation of this method for preparing the intermediate product is the high I contamination from DMAI. In an attempt to reduce this contamination, we increased the volumes of diethyl ether (DEE) and ethanol (EtOH) used to wash the powders from 25 mL to 50 mL, 75 mL, and 100 mL. Fig. 2e shows that despite increased washing volumes, iodine contamination was still significant, and no reduction in I wt% was observed from EDX of the dark brown powders. This may be because DMAI is strongly bound to the surface of Bi₂S₃, and cannot be removed through washing in polar solvents.

Challenges with CsBiSbCl₂ synthesis

We attempted to prepare CsBiSbCl₂ thin films from our intermediate dark brown product (details in Experimental). Fig. 3a shows the XRD patterns of the thin films obtained. Pawley



analysis showed that the final film was a mixture of phases: Bi_2S_3 and $\text{Cs}_3\text{Bi}_2\text{I}_9$. The stronger intensity of the peaks assigned to $\text{Cs}_3\text{Bi}_2\text{I}_9$ suggest that this is the main phase. Data from UV-visible spectrophotometry (UV-vis) support this. Fig. 3b shows the UV-vis absorption spectrum of the films prepared alongside the reported absorption spectrum of $\text{Cs}_3\text{Bi}_2\text{I}_9$.⁵² Both exhibit a distinct absorption onset peak centred at 493 nm wavelength. This peak is characteristic of $\text{Cs}_3\text{Bi}_2\text{I}_9$, and has been attributed to the first excitonic peak.⁵³ Furthermore, the fine structure of the absorption spectrum aligns with previously reported broader peaks at 426 nm, 359 nm, and 310 nm, associated with the higher-order excitonic states. This corroborates that the thin films were primarily $\text{Cs}_3\text{Bi}_2\text{I}_9$, as opposed to CsBiScl_2 . Iodine contamination of the intermediate materials has significant consequences for the preparation of the final thin films, resulting in preferential formation of iodide phases over the desired CsBiScl_2 product. Additionally, Pawley refinement of the PXRD patterns suggests no incorporation of Cl-based phases into the thin films, since the peak positions match those of the reference pattern for $\text{Cs}_3\text{Bi}_2\text{I}_9$. This is supported by EDX observations of prepared films (Fig. S11 and Table S4, SI).

Another problem with this solution synthesis approach is that the solid reagents did not completely dissolve in the DMF/DMSO mixture. The residual solid was collected, dried in a vacuum oven at 60 °C overnight, and analyzed by PXRD. Pawley refinement of this solid confirms that this powder is a mixture of Bi_2S_3 and CsCl (Fig. 3c).^{54–56} This is unsurprising given the poor solubility of CsCl in both DMF and DMSO due to its large lattice energy,⁵⁷ and provides further support for the dark brown intermediate product being primarily Bi_2S_3 with DMAI attached.

Alternative solid-state synthesis route

Given the challenges due to I contamination, we attempted solid state synthesis of CsBiScl_2 by sealing ground and pelletized CsCl , Bi_2S_3 , and BiCl_3 in quartz ampoules under vacuum. Due to the low boiling point of BiCl_3 , a synthesis temperature of 250 °C was chosen. While this is relatively low, there are reports of successful synthesis of ternary BiScl under similar conditions.⁵⁸

No single-phase powders were obtained from this solid-state synthesis. Rietveld refinement (Fig. 3d) of the powders revealed a mixture of binary (CsCl , Bi_2S_3) and ternary (Bi_3CsS_5) phases (goodness of fit of 1.74). As shown in Table S2, SI, all of these phases are on the convex hull of the Cs–Bi–S–Cl system. No peaks corresponding to a CsBiScl_2 perovskite phase were observed in the powders. A lower temperature synthesis was also run at 175 °C, but this contained binaries and ternaries, as well as some unidentified products. This shows that for low-temperature solid-state synthesis, binary and ternary phases form preferentially over the desired quaternary phase. Thus, although the lowest-energy CsBiScl_2 phase is 2.4 meV per atom above the DFT convex hull, it still proves challenging to access this phase.

Conclusions

In this work, we investigated the quaternary CsBiScl_2 system. Through a global structure search, we found that it is

energetically unfavorable for this compound to adopt a regular perovskite structure, in contrast to previous assumptions. Instead, we identified an orthorhombic structure within 2.4 meV per atom of the thermodynamic convex hull, which is dynamically stable, implying it is a plausibly accessible metastable target. We repeated the thin film synthesis reported by Huang *et al.*, demonstrating that the reported DMABiS_2 intermediate is not a uniform phase but rather DMAI bound to the surface of Bi_2S_3 . We attempted to prepare thin films, but were unable to prepare any phase pure films of CsBiScl_2 , instead primarily obtaining films of $\text{Cs}_3\text{Bi}_2\text{I}_9$ due to I contamination in the intermediate product that is difficult to remove through solvent washing. Finally, we showed through an I-free solid-state synthesis that it is challenging to obtain CsBiScl_2 , instead forming phases that are on the convex hull. We therefore find that the 10% PCE claim made in the original paper is spurious, and recommend directing efforts away from this material. We suggest that focus should be shifted toward known Bi-based quaternary mixed-metal chalcogenides, such as AgBiScl_2 and CuBiScl_2 ,^{35,36} wherein the structure of the material has been solved and there remains significant research scope surrounding the synthesis of these materials and optimization of device performance.

Experimental

Calculations

Structure search was carried out using the AiRSS method, with structural relaxations completed using an ephemeral data derived potential (EDDP), *i.e.*, a small bespoke MLIP iteratively trained on single point DFT energies of structures in the Cs–Bi–S–Cl space.

A typical AiRSS workflow involves four steps:

- (1) Randomly generating a ‘sensible’ Cs–Bi–S–Cl structure using simple rules derived from chemical intuition, *e.g.*, enforcing a minimum separation between atoms, fixed stoichiometry;
- (2) Use a (DFT) energy/forces calculator to optimize this structure until forces on atoms are below a desired threshold;
- (3) Add structure and to a ranked database of energies.

Steps 1–3 are repeated until some stopping criteria is reached, such as when the lowest energy structure has been rediscovered multiple times. At this point a subset of ‘interesting’ structures, usually those within 100 meV per atom of the convex hull, are refined using precise DFT settings.

Training of MLIP

The EDDP suite consists of four packages; ‘airss’, ‘ddp’, ‘repose’, and ‘nn’.^{46,59,60} These packages are used for random structure generation/search, training of (E)DDPs, structural optimisation, and implementing neural nets respectively.

An EDDP was iteratively trained using the ‘chain’ script and an initial dataset of 10 000 random CsBiScl_2 structures generated using AiRSS utility ‘buildcell’,⁶¹ as well as a ‘marker’ dataset of all <40 atom structures in the Cs–Bi–S–Cl phase space within 20 meV of the convex hull available on Materials Project.^{41,62,63}



Marker structures were randomly shaken 50 times each with an amplitude of 0.04 Å. DFT single-point energies were calculated for each structure in the training set using CASTEP 24.1⁶⁴ and the high-throughput QC5 ultrasoft pseudopotential set with a cut-off energy of 425 eV, a minimum Monkhorst-Pack grid density of 0.05 Å⁻¹, and a fine grid scale of 2. The PBEsol⁶⁵ functional with D4⁶⁶ dispersion correction was used.

Approximately 256 DDPs were trained on this dataset using the ddp utility 'forge' trained and of these a subset were selected and cast into a weighted ensemble of DDPs (EDDP) by 'flock'. DDPs were trained using up to 3-body terms and a cutoff of 8.0 Å, with a polynomial order of up to 16 for 2-body and up to 4 for 3-body terms, resulting in a feature vector with a length of 435. This was passed through a 5 node neural net and finally to the output layer.

Training, validation, and testing data was split in a ratio of 80 : 10 : 10. Using this EDDP and 'repose', 2000 random AiRSS structures were relaxed to their local minima. DFT total energies were calculated for these structures, added to the dataset, and used to train an improved EDDP. This iterative process was repeated a total of 5 times. To improve the performance of the MLIP, the training dataset was enriched with EDDP relaxed structures from the wider Cs–Bi–S–Cl space, as well as high and low density structures, resulting in a final dataset of 51 212 structures.

For the final MLIP, 213 DDPs were trained and 42 were selected and cast into an EDDP. To run the EDDP suite in a massively parallel way, the batch queuing utility 'DDP-batch' was utilised.⁶⁷ Random structure search was carried out using the as trained EDDP and 'repose', a structural optimiser native to the EDDP suite. This trained MLIP, training dataset, and the 'buildcell'/AiRSS input files used to generate it are available in the raw data repository. A total of 92 770 relaxations were carried out. Rediscovery rates were assessed by merging similar structures using the unite functionality of the 'cryan' tool in the 'airss' package, which compares unit cell volumes and inter-atomic distances within compared structures are within a specified threshold, *e.g.*, 1%.

Refinalization of low energy structures

Structures selected for refinalisation included the 118 unique low-energy structures found *via* random structure search, 36 MP near-the-hull structures, and proposed perovskite structure. Materials project structures included all Cs–Bi–S–Cl structures within 20 meV per atom of the r²SCAN⁶⁸ derived hull, which were also fewer than 50 atoms. Structures were relaxed using r²SCAN-D3(BJ)+SOC,^{68,69} as implemented in VASP 6.5.0^{70,71} and the accurate PAW PBE 'potpaw.64' pseudopotential set⁷² with semicore Cs s & p states treated as valence electrons. A basis set cutoff 450 eV was used, as well as a *k*-spacing of 0.25 Å⁻¹ for insulators and 0.15 Å⁻¹ for known metallic structures from Materials Project. Note, due to differences in definition, VASP grid spacings differ by a factor of 2π to the equivalent CASTEP grid spacing. The 'kgrid' ⁷³ utility was used for adjusting the *k*-point sampling density. Structures were optimised using the conjugate-gradient algorithm (IBRION = 2) until forces on each

atom were <0.03 eV Å⁻¹, and those near the hull (<30 meV per atom) were further optimised with a tighter force convergence criterion of <0.01 eV Å⁻¹.

Dynamic stability of low-energy CsBiSCL₂ structure

The low-energy *Pnma* structure was symmetrized using 'phonopy',^{74,75} further optimized until forces on each atom were <5 × 10⁻³ eV Å⁻¹. The orthorhombic primitive cell had dimensions of 9.45 × 4.13 × 16.08 Å³ and a 4 × 8 × 2 *I*-centered *k*-point grid was used during relaxation. 2 × 4 × 1 supercells and a commensurate 2 × 2 × 2 *k*-point grid were used for finite displacements with a displacement amplitude of 0.01 Å. Born effective charges were extracted from a finite differences (IBRION = 6) run on the primitive and used to estimate TO-LO splitting.

Materials

All chemicals were used as received without any further purification. Bismuth sulfide (Bi₂S₃, 99%), dimethyl sulfoxide (DMSO, ≥99.9%), hydroiodic acid (HI, 57 wt%, 99.95%), ethanol (≥99.8%), and diethyl ether (≥99.8%) were purchased from Sigma-Aldrich. Cesium chloride (CsCl, 99.999%), bismuth chloride (BiCl₃, 97%) and *N,N*-dimethylformamide (DMF, anhydrous, 99.9%) were purchased from ThermoFisher Scientific.

Preparation of DMABiS₂ intermediate

1.80 g Bi₂S₃ and 3 mL DMF were stirred at 60 °C for 1 h. 2.5 mL HI was added dropwise and the suspension was stirred for 4 h. The suspension was filtered *via* Büchner filtration and washed with diethyl ether and ethanol. The resulting powder was dried in a vacuum oven overnight at 60 °C.

Attempted synthesis of CsBiSCL₂ thin films

All steps, apart from substrate cleaning, were carried out inside N₂-filled glovebox.

0.16 g of the dark brown intermediate product and 0.34 g CsCl were mixed in DMF/DMSO (9:1) and stirred at room temperature for 2 h. The suspension was allowed to settle for 24 h before the liquid phase was filtered with a 0.2 μm PTFE filter. A 1.2 cm × 1.2 cm substrate was cleaned by ultrasonication in acetone for 15 min, followed by isopropanol for 15 min. After cleaning, the substrates were blown dry with N₂, followed by UV-ozone treatment (NanoBioAnalytics UV Ozone Cleaner UVC-1014) for 20 min. 40 μL of the filtered solution was spread onto the substrate followed by spinning at 1000 rpm for 10 s and 4000 rpm for 30 s. The deposited film was annealed at 220 °C for 1 h 20 min.

Attempted synthesis of CsBiSCL₂ powders

Polycrystalline samples of 'CsBiSCL₂' were prepared by a solid-state reaction. In an N₂-filled glovebox, BiCl₃, Bi₂S₃, and CsCl were weighed and wet ground in acetone for 30 min. 0.65 g aliquots were then pressed into 12 mm pellets by applying 9 ton. Pellets were sealed in 7 mm internal diameter quartz ampoules



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