

## PAPER

[View Article Online](#)  
[View Journal](#) | [View Issue](#)Cite this: *J. Mater. Chem. C*,  
2024, 12, 12521Expanding the horizons for viable precursors and liquid fluxes for the synthesis of BaZrS<sub>3</sub> and related compounds†Kiruba Catherine Vincent, <sup>‡a</sup> Shubhanshu Agarwal, <sup>‡a</sup> Zirui Fan, <sup>a</sup>  
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Chalcogenide perovskites represent a prominent class of emerging semiconductor materials for photovoltaic applications, boasting excellent optoelectronic properties, appropriate bandgaps, and remarkable stability. Among these, BaZrS<sub>3</sub> is one of the most extensively studied chalcogenide perovskites. However, its synthesis typically demands high temperatures exceeding 900 °C. While recent advancements in solution-processing techniques have mitigated this challenge, they often rely on costly and difficult-to-find organometallic precursors. Furthermore, there is a notable gap in research regarding the influence of the Ba/Zr ratio on phase purity. Thus, our study explores solid-state reactions to investigate the impact of metal ratios and sulfur pressure on the phase purity of BaZrS<sub>3</sub>. Expanding upon this investigation, we aim to leverage cost-effective metal halide and metal sulfide precursors for the solution-based synthesis of BaMS<sub>3</sub> (M = Ti, Zr, Hf) compounds. Additionally, we have devised a bilayer stacking approach to address the halide affinity of alkaline earth metals. Moreover, we introduce a novel solution-chemistry capable of dissolving alkaline earth metal sulfides, enabling the synthesis of BaMS<sub>3</sub> compounds from metal sulfide precursors. While the BaS<sub>x</sub> liquid flux has shown promise, we identify the selenium liquid flux as an alternative method for synthesizing BaMS<sub>3</sub> compounds.

Received 3rd June 2024,  
Accepted 9th July 2024

DOI: 10.1039/d4tc02287d

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## 1. Introduction

Over the past two decades, there has been a notable surge in the development of new solar absorbers for photovoltaic applications. Lead halide perovskites have been particularly exciting due to their optoelectronic properties and the efficiencies they have achieved in photovoltaic devices.<sup>1–3</sup> However, they often suffer from increased instability to air, moisture, and heat, rendering them unsuitable for large-scale deployment in their current form.<sup>4</sup> Chalcogenide semiconductor materials, such as Cu(In,Ga)Se<sub>2</sub> and Cu<sub>2</sub>ZnSnSe<sub>2</sub>, have also shown promise as photovoltaic materials with suitable bandgaps for single-junction applications, but they lag behind lead halide perovskites in terms of efficiencies.<sup>5,6</sup> Nevertheless, these materials offer increased stability to air, moisture, and thermal degradation compared to lead halide perovskites.<sup>7</sup> They could also be

readily mass-produced and deployed in photovoltaic farms, provided their device efficiencies are improved.

The possible research avenues in solar absorber research include improving the stability of halide perovskites, enhancing the device efficiencies of existing chalcogenide materials, or exploring stable, high-performing alternatives to lead halide perovskites. The latter has drawn the attention of the photovoltaic research community toward chalcogenide perovskites. These materials crystallize in a similar ABX<sub>3</sub> structure as lead halide perovskites, where A represents an alkaline earth metal (Ca, Sr, Ba), B denotes an early transition metal (Ti, Zr, Hf), and X is predominantly sulfur. While the Zr and Hf versions crystallize in a distorted perovskite structure, with the [BX<sub>6</sub>]<sup>2–</sup> octahedra slightly tilted, the Ti versions crystallize in a distinct hexagonal structure.<sup>8–10</sup> Chalcogenide perovskites exhibit extremely high light absorption coefficients (>10<sup>5</sup> cm<sup>–1</sup>), surpassing even lead halide perovskites, high dielectric constants, reasonable carrier mobilities, and strong photoluminescence.<sup>8,11–13</sup> Some of them feature suitable bandgaps for the top layer in tandem solar cell applications, while others show promise for solar water splitting and light-emitting diode applications.<sup>8,12–15</sup> ATiS<sub>3</sub> compounds possess a quasi-1-D structure with low thermal conductivity, making them promising candidates for thermoelectric applications.<sup>16,17</sup>

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Despite their considerable promise, progress in researching these materials has been slow, primarily due to the high synthesis temperatures, often exceeding 900 °C.<sup>12,18–21</sup> Among these materials, BaZrS<sub>3</sub> has been the focus of much attention owing to its particularly suitable bandgap for photovoltaic applications and its synthesis at relatively lower temperatures than other chalcogenide perovskites.<sup>22,23</sup> However, research on BaZrS<sub>3</sub> has predominantly centered on solid-state forms, with few reports on film synthesis, most of which rely on high-temperature vacuum deposition and annealing processes, which are less appealing.<sup>23–26</sup>

Recent studies from our group and others have made significant strides in lowering the synthesis temperature of BaMS<sub>3</sub> (M = Ti, Zr, Hf) compounds through moderate-temperature solution processing. These methods use a BaS<sub>x</sub> liquid flux in a sulfur-rich environment but still rely on complex and expensive organometallic precursors for film casting.<sup>27–31</sup> Historically, solution-processed synthesis of chalcogenide materials has utilized cost-effective metal salts and metal sulfide precursors.<sup>32–35</sup> However, challenges such as the high oxophilicity of transition metals, the strong affinity of alkaline earth metals for halides, and the low solubility of alkaline-earth and transition metal sulfides in traditional solution chemistries have complicated these efforts.<sup>30,36,37</sup> A recent study from our group addressed the issue of oxophilicity by utilizing an HfH<sub>2</sub>-based oxygen trap to remove oxide impurities from the film and the ampule environment during synthesis.<sup>38</sup> However, other challenges persist, requiring further investigation.

In this study, we have developed innovative approaches to address the halide affinity of alkaline-earth metals and synthesize BaMS<sub>3</sub> compounds from cost-effective metal halide precursors. We also demonstrate chemistry to dissolve alkaline-earth metal sulfides and propose a strategy to utilize metal sulfides for casting BaMS<sub>3</sub> films. Additionally, this study provides an example of using a selenium liquid flux, which could facilitate the synthesis of BaMS<sub>3</sub> and related compounds. Our study explores cost-effective solution-deposition routes for BaMS<sub>3</sub> synthesis by leveraging critical insights from previous reports.

## 2. Results and discussion

### 2.1. Does BaZrS<sub>3</sub> allow off-stoichiometric compositions?

Chalcogenide semiconductors like Cu(In,Ga)Se<sub>2</sub> and Cu<sub>2</sub>ZnSnSe<sub>4</sub> exhibit off-stoichiometry in their structure, where Cu-poor compositions up to a certain I/III or I/(II + IV) ratio do not induce phase segregation.<sup>39,40</sup> This is significant because Cu vacancies are shallow acceptor defects, imparting p-type conductivity to these materials.<sup>41</sup> Notably, the record efficiency Cu(In,Ga)Se<sub>2</sub> device possesses a Cu/(In + Ga) ratio of 0.93.<sup>42</sup> Conversely, Cu-rich compositions often result in secondary Cu<sub>2</sub>Se impurities, which may pose challenges if not easily eliminated.<sup>41</sup> This spurred our investigation into off-stoichiometric compositions in BaZrS<sub>3</sub>, which remains largely unexplored. Similar to Cu-poor compositions in Cu(In,Ga)Se<sub>2</sub>,

**Table 1** Phases observed after sulfurization of BaS and ZrS<sub>2</sub> powder in the presence of sulfur at 575 °C for 12 h

| S. no. | Ba : Zr (molar ratio) | X-ray diffraction phases (without water wash)                                          | Raman phases (without water wash)                                                      | Estimated sulfur pressure (atm) |
|--------|-----------------------|----------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|---------------------------------|
| 1      | 2 : 1                 | BaZrS <sub>3</sub> + BaS <sub>3</sub>                                                  | BaZrS <sub>3</sub> + BaS <sub>3</sub>                                                  | 0.79                            |
| 2      | 1.5 : 1               | BaZrS <sub>3</sub> + BaS <sub>3</sub>                                                  | BaZrS <sub>3</sub> + BaS <sub>3</sub>                                                  | 0.79                            |
| 3      | 1 : 1                 | BaZrS <sub>3</sub>                                                                     | BaZrS <sub>3</sub>                                                                     | 0.79                            |
| 4      | 0.9 : 1               | BaZrS <sub>3</sub>                                                                     | BaZrS <sub>3</sub>                                                                     | 0.79                            |
| 5      | 0.8 : 1               | BaZrS <sub>3</sub>                                                                     | BaZrS <sub>3</sub>                                                                     | 0.79                            |
| 6      | 0.7 : 1               | BaZrS <sub>3</sub>                                                                     | BaZrS <sub>3</sub>                                                                     | 0.79                            |
| 7      | 0.6 : 1               | BaZrS <sub>3</sub> + ZrS <sub>3</sub>                                                  | BaZrS <sub>3</sub> + ZrS <sub>3</sub>                                                  | 0.79                            |
| 8      | 0.5 : 1               | BaZrS <sub>3</sub> + ZrS <sub>3</sub>                                                  | BaZrS <sub>3</sub> + ZrS <sub>3</sub>                                                  | 0.79                            |
| 9      | 0.4 : 1               | ZrS <sub>3</sub> + BaZrS <sub>3</sub>                                                  | ZrS <sub>3</sub>                                                                       | 0.79                            |
| 10     | 1.5 : 1               | BaZrS <sub>3</sub> + BaS <sub>3</sub>                                                  | BaZrS <sub>3</sub> + BaS <sub>3</sub>                                                  | 0.79                            |
| 11     | 1.5 : 1               | BaZrS <sub>3</sub> + BaS <sub>3</sub>                                                  | BaZrS <sub>3</sub> + BaS <sub>3</sub>                                                  | 0.59                            |
| 12     | 1.5 : 1               | BaZrS <sub>3</sub> + BaS <sub>3</sub> + Ba <sub>3</sub> Zr <sub>2</sub> S <sub>7</sub> | BaZrS <sub>3</sub> + BaS <sub>3</sub> + Ba <sub>3</sub> Zr <sub>2</sub> S <sub>7</sub> | 0.46                            |

Ba-poor compositions could potentially alter the carrier concentrations in BaZrS<sub>3</sub>. The defects arising from such conditions, including V<sub>Ba</sub>, Zr<sub>Ba</sub>, S<sub>Ba</sub>, and other defect complexes, warrant thorough examination. Preliminary indications from defect calculations suggest that these defects are either shallow or possess high defect formation energies.<sup>43,44</sup>

Table 1 enumerates the powder reactions where we mixed BaS and ZrS<sub>2</sub> in different molar ratios and sulfurized at 575 °C for 12 h in the presence of sulfur vapor. It is evident from Table 1 and Fig. S1 (ESI†) that increasing the Ba:Zr ratio above 1 in the presence of excess sulfur led to the formation of the BaS<sub>3</sub> secondary phase alongside crystalline BaZrS<sub>3</sub>. This indicates that BaZrS<sub>3</sub> does not accommodate Zr-poor off-stoichiometries, perhaps because V<sub>Zr</sub> would disrupt the [ZrS<sub>6</sub>]<sup>2-</sup> octahedra and destabilize the BaZrS<sub>3</sub> crystal lattice. It is plausible that in Ba-rich compositions, barium polysulfide was initially formed, but it decomposed into BaS<sub>3</sub> and sulfur upon cooling.<sup>27</sup> Consequently, crystalline BaS<sub>3</sub> and BaZrS<sub>3</sub> were observed under barium-rich conditions. Previous studies have demonstrated the solubility of BaS<sub>3</sub> in water.<sup>28</sup> Thus, a gentle water wash could eliminate all BaS<sub>3</sub> impurities from the sample. In our investigation, we followed a water-washing procedure outlined in the experimental section, resulting in the phase-pure synthesis of BaZrS<sub>3</sub>, as validated by X-ray diffraction in Fig. 1(a) and Raman spectroscopy in Fig. S2 (ESI†). Regarding the Ba-poor compositions, no secondary phases were observed for Ba:Zr ~ 0.9, 0.8, and 0.7, yielding phase-pure BaZrS<sub>3</sub> as confirmed by X-ray diffraction and Raman spectroscopy. However, a further decrease to 0.6 metal ratio resulted in slight ZrS<sub>3</sub> impurities, evident in the X-ray diffraction spectra and Raman analysis (see Fig. 1(b) and Fig. S3, ESI†). As the Ba:Zr ratio decreased, the prevalence of secondary phases increased as anticipated. Interestingly, our study indicates that BaZrS<sub>3</sub> can accommodate Ba-poor compositions up to Ba:Zr ~ 0.7. It would be intriguing to investigate the impact of these compositions on the optoelectronic properties, although this falls beyond the scope of our current work. The off-stoichiometry of BaZrS<sub>3</sub> could potentially impact the carrier concentration, which may vary depending on the Ba : Zr ratio in the material. Consequently, a wide range of



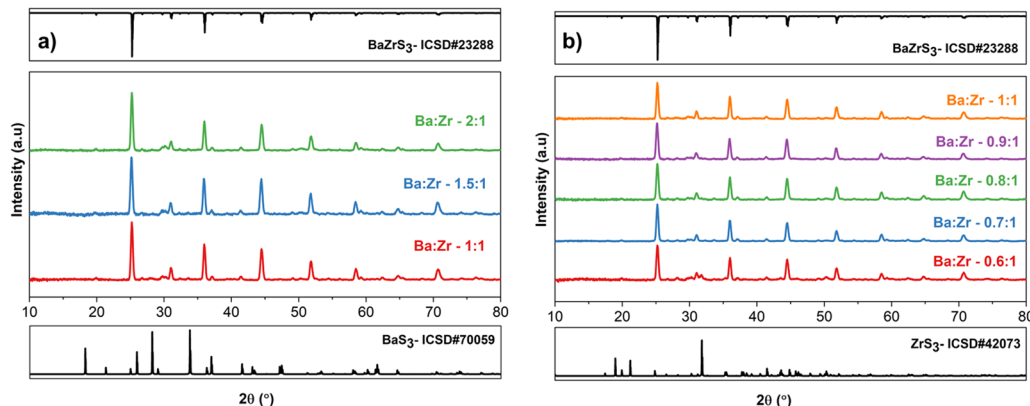


Fig. 1 (a) and (b) X-ray diffraction pattern of the Ba–Zr–S powder synthesized with different Ba:Zr ratios at 575 °C for 12 h with a sulfur pressure of 0.79 atm. BaS and ZrS<sub>2</sub> were used as barium and zirconium precursors. The synthesized powders in (a) were water-washed to remove any water-soluble impurities (BaS<sub>3</sub>). The sample with Ba:Zr 0.6 : 1 in (b) has ZrS<sub>3</sub> impurities.

carrier concentrations and both p-type and n-type conductivities have been reported for BaZrS<sub>3</sub>.<sup>12,45</sup> Nevertheless, these findings will be helpful in the subsequent section, where we will elaborate on our approaches to fabricating BaZrS<sub>3</sub> films.

It is essential to highlight that compositions with excess Ba under sulfur-poor conditions were also investigated. Despite the common expectation that Ruddlesden–Popper (RP) phases of BaZrS<sub>3</sub> would form in Ba-excess compositions due to their formation energies being similar to those of the perovskite phase,<sup>46</sup> our previous discussion elucidated that BaZrS<sub>3</sub>, along with BaS<sub>3</sub>, was formed in Ba-excess compositions under sulfur-rich conditions. However, as the sulfur pressure decreased during synthesis, RP phases, notably Ba<sub>3</sub>Zr<sub>2</sub>S<sub>7</sub>, reported to be the most stable among all Ba–Zr–S RP phases,<sup>45,47</sup> emerged alongside BaZrS<sub>3</sub>, as confirmed by X-ray diffraction and Raman spectroscopy (see Fig. S4 and S5, ESI†). At even lower sulfur pressures, starting binaries remained partially unreacted. Hence, it is imperative to operate under sulfur-excess conditions to achieve phase-pure BaZrS<sub>3</sub>.

These observations indicate that, in a saturated sulfur environment with barium above stoichiometric amount, BaZrS<sub>3</sub> does not accommodate Zr-deficient off-stoichiometric compounds. Instead, it leads to the formation of BaZrS<sub>3</sub> and BaS<sub>3</sub> secondary phases. Conversely, Ba-deficient off-stoichiometric compounds appear stable up to a Ba:Zr ratio of 0.7. Building on these findings, we have developed several solution processing routes for the synthesis of chalcogenide perovskites.

## 2.2. Stacked binary metal sulfides enabling the use of transition metal halides

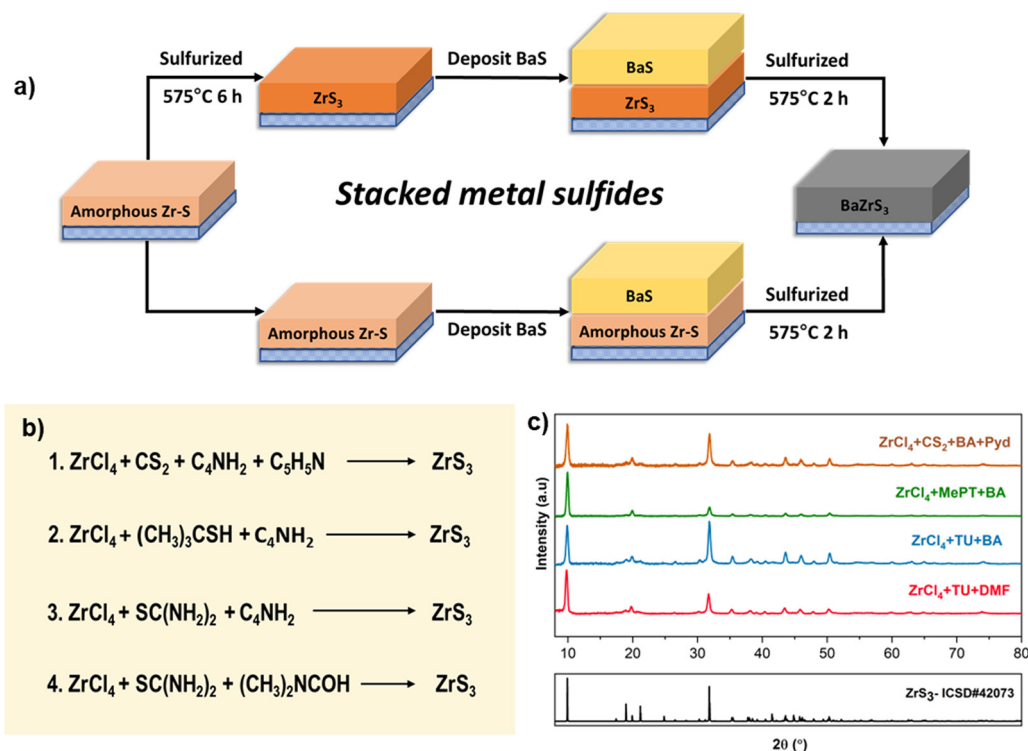
Metal halides are widely available and relatively inexpensive, making them preferred precursors for many material systems. For instance, metal chlorides of certain late and post-transition elements, dissolved in dimethyl formamide (DMF) with thiourea as a sulfur source, can be utilized to synthesize metal sulfide semiconductors.<sup>33,48</sup> However, the affinity of barium and other alkaline earth metals for chlorine hindered our

group's initial attempts to use these compounds in synthesizing BaZrS<sub>3</sub>. As further confirmation, we investigated using a dimethylformamide ink containing BaCl<sub>2</sub>, ZrCl<sub>4</sub>, and thiourea. As anticipated, the resulting film contained BaCl<sub>2</sub> and ZrS<sub>3</sub> after sulfurization, as depicted in Fig. S6 (ESI†), underscoring barium's affinity for chlorine. We encountered similar challenges with bromide and iodide precursors, discouraging using halides for mixed precursor inks. A potential alternative to bypass alkaline earth element's chlorine affinity involves employing a bilayer stacking approach with BaS<sub>x</sub>/ZrS<sub>x</sub> films, where ZrS<sub>x</sub> could be synthesized from zirconium halide precursors and then sulfurized to achieve BaZrS<sub>3</sub>.

While some researchers previously regarded ZrS<sub>3</sub> as a dissociation product of BaZrS<sub>3</sub>, our earlier suggestion posited ZrS<sub>3</sub> as a feasible zirconium precursor for synthesizing BaZrS<sub>3</sub>.<sup>27</sup> In this scenario, ZrS<sub>3</sub> could undergo a reaction with BaS in the presence of excess sulfur, leading to the formation of BaZrS<sub>3</sub>.<sup>27</sup> Based on this, our proposed method involves creating a binary sulfide stack with a bottom layer of ZrS<sub>3</sub> and a top layer of BaS, illustrated in Fig. 2(a). Subsequently, this stack is heated in a sulfur-rich environment to yield BaZrS<sub>3</sub>.

We dissolved Zr, Hf, and Ti halides using four distinct solution chemistries, resulting in metal–sulfur coordinated complexes, as depicted in Fig. 2(b). The confirmation of metal–sulfur interactions in these chemistries was studied through liquid Raman and <sup>1</sup>H-NMR (nuclear magnetic resonance) analyses. While amine–carbon disulfide chemistry has been extensively studied for dissolving various metal oxides, metal acetylacetonates, and metal hydroxides, limited research has focused on the dissolution of metal halides, particularly those of early transition metals. Liquid Raman analysis of a 0.3 M solution of ZrCl<sub>4</sub> in butylamine–CS<sub>2</sub>–pyridine (Fig. S7, ESI†) revealed broadening of the C=S peak at 656 cm<sup>−1</sup>, indicating potential metal–sulfur interaction, alongside a new peak at 2648 cm<sup>−1</sup>, possibly attributed to the S–H bond from CS<sub>2</sub> insertion in butylamine. Similarly, for ZrCl<sub>4</sub> in 2-methyl-2-propanethiol–butylamine solution, Fig. S8 (ESI†) shows a slight shift and broadening in the C–S peak at 590 cm<sup>−1</sup> and the S–H





**Fig. 2** (a) Reaction schematic depicting the synthesis of BaZrS<sub>3</sub> by layering binary sulfides. (b) Various molecular precursor chemistries employed for depositing ZrS<sub>3</sub> film. (c) Plot comparing the X-ray diffraction patterns of ZrS<sub>3</sub> film prepared using different solution chemistries, (i) ZrCl<sub>4</sub> + thiourea (TU) + dimethyl formamide (DMF), (ii) ZrCl<sub>4</sub> + TU + butylamine (BA), (iii) ZrCl<sub>4</sub> + 2-methyl-2-propanethiol (MePT) + BA and (iv) ZrCl<sub>4</sub> + BA + CS<sub>2</sub> + pyridine (Pyd).

peak at  $2577\text{ cm}^{-1}$ , indicating sulfur coordination to the zirconium through a dative bond.<sup>49</sup> However, the observed peak shift was minimal, and the ratio of the S–H peak to the C–S peak did not decrease significantly, suggesting incomplete substitution of chlorine with thiolate due to thiolate formation. This finding aligns with observations by Murria *et al.*, who reported similar incomplete substitution when dissolving copper chlorides in amine–thiol systems.<sup>32</sup>

<sup>1</sup>H-NMR studies also show a likely metal–sulfur interaction between ZrCl<sub>4</sub> and thiol (Fig. S9, ESI†). The Zr–Cl–thiolate, however, did not dissolve significantly in pyridine-d<sub>5</sub> initially. But, adding a stoichiometric amount of butylamine resulted in complete dissolution with a broadening in the <sup>1</sup>H-NMR peak of H bounded to sulfur, hinting towards some interaction between butylamine and 2-methyl-2-propanethiol likely forming  $[\text{CH}_3(\text{CH}_2)_3\text{NH}_4]^+[\text{SC}(\text{CH}_3)_3]^-$ . The <sup>1</sup>H-NMR results in Fig. S10 (ESI†) show the broadening of the thiol proton peak after adding ZrCl<sub>4</sub>, but the peak did not disappear. Notably, exchangeable protons, such as those in the thiol group, are not always accurately quantified through peak integration. The peak broadening indicates a dynamic thiol proton. We hypothesize that this suggests the thiolate is in equilibrium between its protonated and deprotonated forms. When it is deprotonated, we expect it to be bound to the Zr<sup>4+</sup>. This observation could also indicate that not all Zr–Cl bonds underwent replacement with thiol; alternatively, it might have resulted in the formation of an adduct.

Interactions were also observed in the ZrCl<sub>4</sub>–thiourea–dimethyl formamide and ZrCl<sub>4</sub>–thiourea–butylamine systems, leading to shifts and broadening in the C=S Raman peak, as illustrated in Fig. 3 and 4. Similar dissolutions can also be achieved with ZrBr<sub>4</sub> and ZrI<sub>4</sub>, yielding soluble complexes. All four Zr–S coordinated solutions were then doctor blade-coated onto alumina-coated Eagle XG glass substrates and annealed at  $350^\circ\text{C}$  on a hot plate to remove the excess solvent, resulting in an amorphous ZrS<sub>x</sub> compound. Subsequently, the resulting film underwent sulfurization in an ampule containing HfH<sub>2</sub> and sulfur for 6 h at  $575^\circ\text{C}$  to form a ZrS<sub>3</sub> film, as depicted in the X-ray diffraction patterns and Raman spectra in Fig. 2(c) and Fig. S11 (ESI†). The role of HfH<sub>2</sub> in removing any residual oxygen in the film is elaborated in our previous work.<sup>38</sup> It is important to note that although ZrS<sub>3</sub> could be synthesized at much lower temperatures of  $475^\circ\text{C}$ , we chose 6 h at  $575^\circ\text{C}$  to ensure complete removal of residual oxygen (see Fig. S12, ESI†). Further optimization strategies such as rigorous solvent drying, use of anhydrous precursors, processing in ultra-oxygen-free glove boxes, and avoiding exposure to air during the entire processing could significantly reduce the synthesis time and temperature. The X-ray diffraction patterns and Raman spectroscopy confirmed the production of phase-pure ZrS<sub>3</sub> in all cases. The secondary phase of ZrO<sub>2</sub> was eliminated with prolonged sulfurization time and careful handling of precursors and solutions. Similar to previous studies, the ZrS<sub>3</sub> films exhibited non-continuous morphology due to the ribbon structure of ZrS<sub>3</sub>.





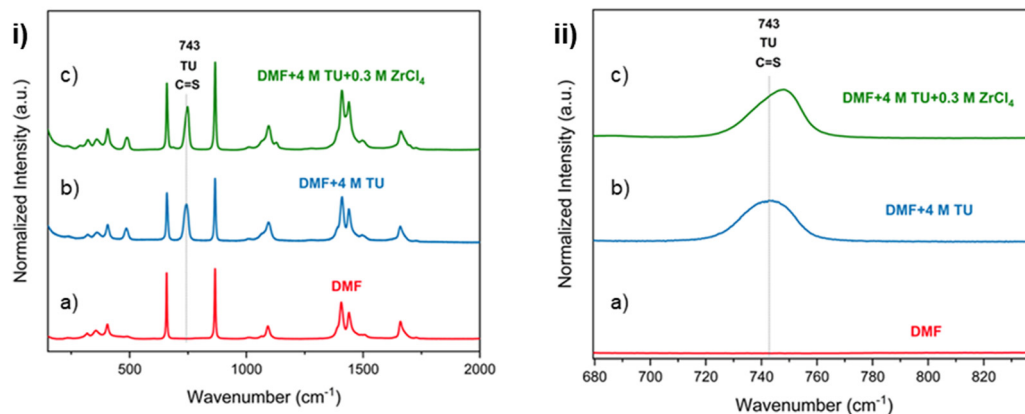


Fig. 3 Liquid Raman on only (a) dimethyl formamide (DMF), (b) thiourea (TU) + DMF, and (c) TU + DMF +  $\text{ZrCl}_4$  (0.3 M). The peak shift in the C=S bond stretch of TU suggests a metal–sulfur interaction. (ii) presents the zoomed-in representation of (i).

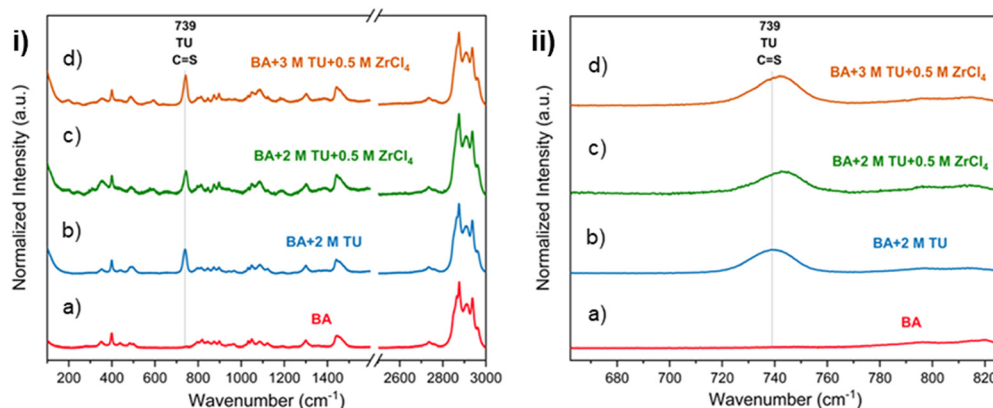


Fig. 4 Liquid Raman on only (a) butylamine (BA), (b) thiourea (TU) + BA, (c) TU (2 M) + BA +  $\text{ZrCl}_4$  (0.5 M), and (d) TU (3 M) + BA +  $\text{ZrCl}_4$  (0.5 M). The peak shift in the C=S bond stretch of TU suggests a metal–sulfur interaction. (ii) presents the zoomed-in representation of (i).

(refer to Fig. 5(b) and Fig. S13, S14, ESI†).<sup>38</sup> Energy dispersive X-ray spectroscopy (EDX) maps in Fig. S15 and S16 (ESI†) confirm the uniform distribution of Zr and S in the grains. Furthermore,  $\text{ZrS}_3$  synthesis from soluble inks of  $\text{ZrBr}_4$  and  $\text{ZrI}_4$  was also demonstrated (see Fig. S17, ESI†). It is worth noting that after annealing the film on the hot plate, trace amounts of residual chlorine, bromine, or iodine were detected in the film *via* X-ray Fluorescence measurements, but no halide impurities remained after sulfurization.

Subsequently, a thin layer of BaS was coated onto these  $\text{ZrS}_3$  films using the soluble barium thiolate ink obtained by dissolving Bis(pentamethylcyclopentadienyl)barium ( $\text{Cp}^*_2\text{Ba}$ ) in 2-methyl-2-propanethiol and butylamine (see experimental section for details), and the resulting stacked film of BaS/ $\text{ZrS}_3$  was sulfurized with sulfur and  $\text{HfH}_2$ . The as-annealed stacked films exhibit crystalline phases of both BaS and  $\text{ZrS}_3$ , as depicted in Fig. 5(a) and Fig. S18 (ESI†). While care was taken to ensure that approximately equal amounts of Ba and Zr molar ratios were present in the film by maintaining similar concentrations of Ba and Zr inks, previous discussions in the study indicate that a barium-excess film could also be utilized. This is because it

would form  $\text{BaZrS}_3$  and  $\text{BaS}_3$  upon sulfurization, and the  $\text{BaS}_3$  component could be easily washed away. During sulfurization at 575 °C, this process utilized the  $\text{BaS}_x$  liquid flux to accelerate mass transfer, and although shorter times should be feasible, sulfurization was performed for 2 h to minimize undesirable oxide phases. As illustrated in Fig. 5(a) and Fig. S18, S19 (ESI†), we obtained a pure  $\text{BaZrS}_3$  phase, as confirmed by X-ray diffraction and Raman spectroscopy. The diffuse reflectance spectroscopy also confirmed a bandgap of around 1.85 eV (see Fig. S20, ESI†). Nevertheless, the film exhibited cracking and isolated grains (see Fig. 5(c) and Fig. S21, ESI†), suggesting a need for further research to optimally control the  $\text{BaS}_x$  liquid flux. However, this could be an enticing way to make micron-scale single crystals without residual impurities. EDX analysis confirmed the uniform distribution of Ba, Zr, and S, as shown in Fig. S22 and S23 (ESI†). Additionally, the  $\text{BaZrS}_3$  grains show no residual chlorine (see Fig. S24, ESI†). This versatile method can be readily applied to Ba–Hf–S and Ba–Ti–S systems, yielding  $\text{Ba}_6\text{Hf}_5\text{S}_{16}$  and  $\text{BaTiS}_3$ , as demonstrated in Fig. S25–S27 (ESI†). However, achieving stoichiometric  $\text{BaHfS}_3$  using this route proved challenging due to the competing



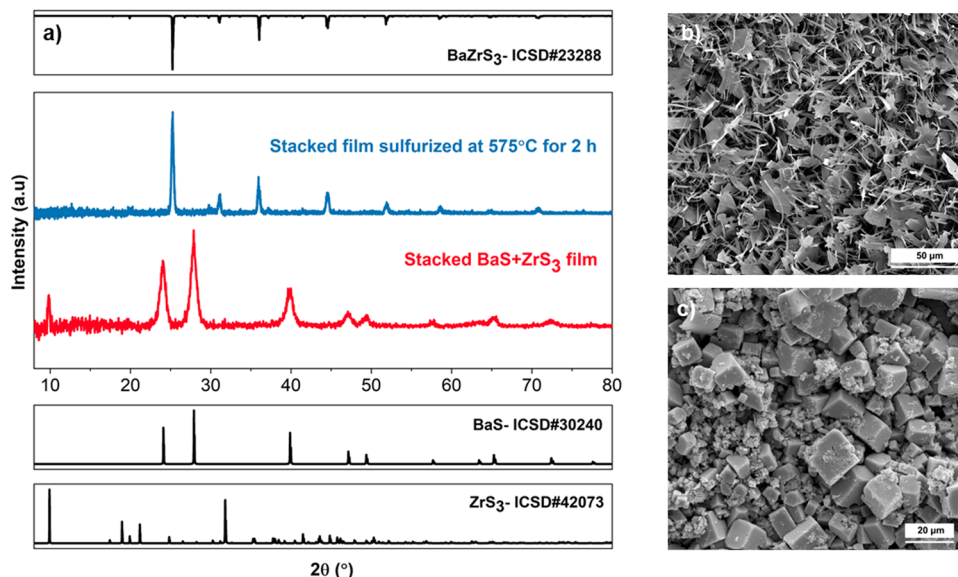


Fig. 5 (a) X-ray diffraction pattern showcasing the as-annealed stacked bilayer film of ZrS<sub>3</sub> and BaS before ampule sulfurization and the BaZrS<sub>3</sub> film after sulfurization. (b) SEM top-view image depicting the ZrS<sub>3</sub> film prepared from the ZrCl<sub>4</sub>–butylamine–CS<sub>2</sub>–pyridine solution chemistry. (c) SEM top-view image illustrating the BaZrS<sub>3</sub> film derived from the ZrS<sub>3</sub>–BaS bilayer stack.

Ruddlesden–Popper phases of the Ba–Hf–S system. In all the cases, an equal number of layers with equally concentrated barium and zirconium/hafnium/titanium solutions were blade-coated to ensure a nearly stoichiometric film.

The success of this approach is noteworthy as it enables the utilization of easily manageable and cost-effective halide precursors. While our primary focus was on one stacking method utilizing the halide precursors, alternative approaches could also be explored. In another instance, we applied a ZrCl<sub>4</sub>-containing ink at the bottom to form an amorphous ZrS<sub>x</sub> layer without sulfurizing it to ZrS<sub>3</sub>. The coated film underwent annealing at 450 °C for 30 minutes to eliminate residual chlorine. Subsequently, BaS was applied to the top using the barium thiolate solution, and the stacked film was sulfurized, resulting in BaZrS<sub>3</sub>, as shown in Fig. S28 (ESI†). In an alternate scenario, BaS was coated at the bottom, and ZrCl<sub>4</sub>-containing ink was applied at the top. After annealing and sulfurization of this stack, Ba<sub>3</sub>Zr<sub>2</sub>S<sub>7</sub> was obtained, albeit with significant secondary phases, including BaCl<sub>2</sub> and ZrS<sub>3</sub>, as illustrated in Fig. S29 (ESI†). This outcome was anticipated, as barium likely reacted with chlorine during the annealing process of the stacked film on the hot plate. These findings underscore the significance of selecting suitable precursors and optimizing the sequence of processing steps to establish a thermodynamic driving force for synthesizing chalcogenide perovskites.

### 2.3. Hybrid colloidal molecular precursor inks

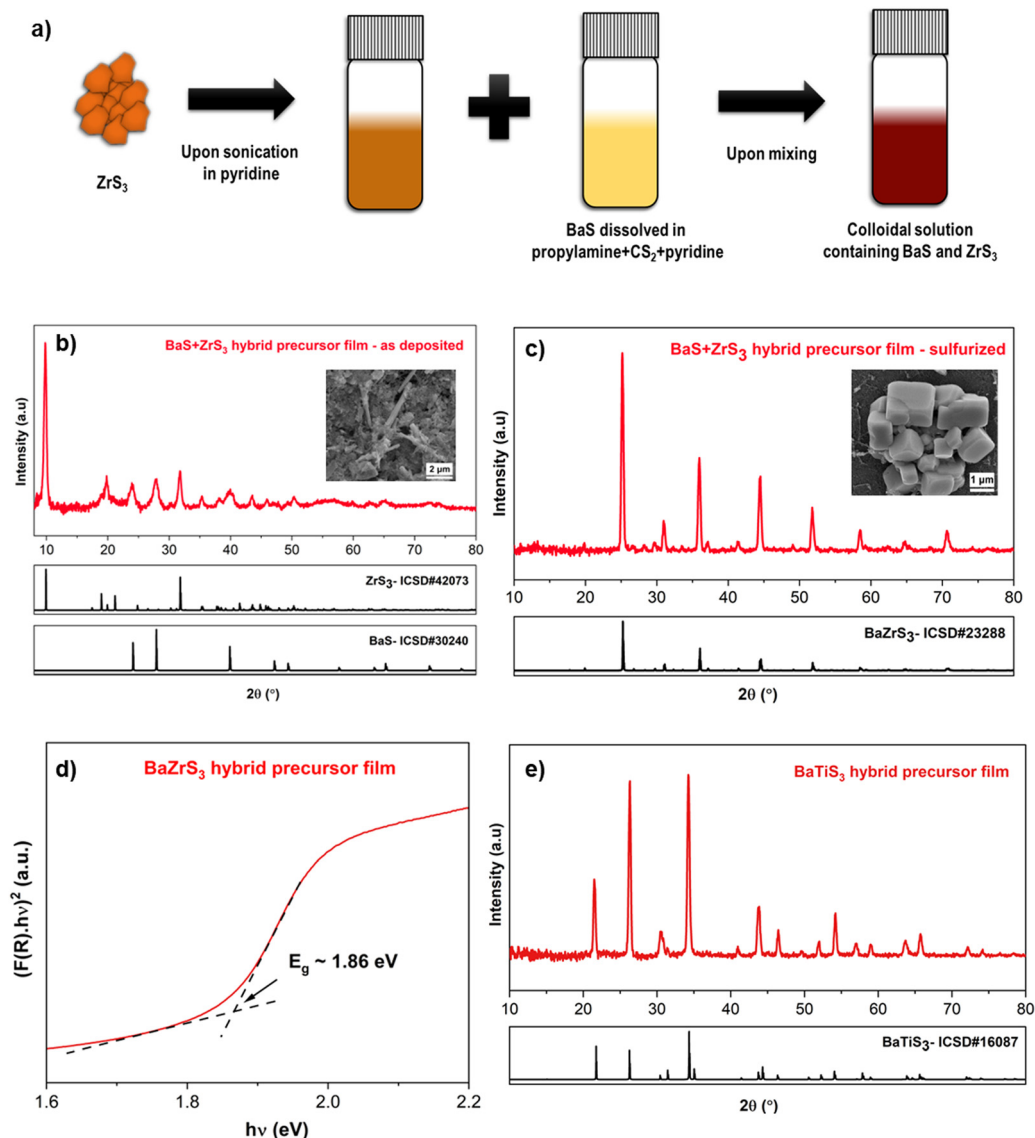
In addition to pure metals, metal chalcogenides are preferred as low-cost, anionic, impurity-free metal precursors for solution-processed chalcogenide thin films.<sup>50,51</sup> Employing metal sulfides as precursors for synthesis would restrict the incorporation of unwanted anionic impurities and offer a cost-effective, streamlined route to synthesize BaMS<sub>3</sub> compounds.

However, the dissolution of alkaline earth metal sulfides and transition metal sulfides has historically been challenging, with no reports of ZrS<sub>2</sub> or ZrS<sub>3</sub> dissolution in any solution chemistry and only one report on co-dissolution of BaS alongside Cu<sub>2</sub>S and SnO in ethylenediamine–ethanedithiol (EDA–EDT) at 60 °C for 11 days.<sup>52</sup> This dissolution was arduous, and attempts to dissolve only BaS in EDA–EDT were unsuccessful in our experiments, even after multiple attempts. Consequently, further investigation was necessary to identify solution chemistry capable of dissolving alkaline earth metal sulfides. As a result, various combinations of amine–thiol with different concentrations were explored, yet no success was achieved in room-temperature dissolutions. Notable solution chemistries such as hydrazine–sulfur also failed to dissolve BaS and ZrS<sub>2</sub>.

However, combining amine and CS<sub>2</sub> to produce alkylthiocarbamic acid opened new possibilities for reactive solvent systems as we investigated this less-studied chemistry. Previously, butyldithiocarbamic acid had been shown to dissolve metal salts such as metal oxides,<sup>34</sup> but we notably extended this chemistry by dissolving alkaline earth metal sulfides in propyldithiocarbamic acid in pyridine, resulting in a clear solution. This marks the first instance of standalone dissolution of BaS and SrS in solution chemistry and could play a pivotal role in the solution-processed synthesis of several alkaline earth metal-based chalcogenide compounds, including BaMS<sub>3</sub> chalcogenide perovskites, Cu<sub>2</sub>BaSnS<sub>4</sub>, and Cu<sub>2</sub>SrSnS<sub>4</sub>. As a side note, due to the current interest in forming thin films of Cu<sub>2</sub>BaSnS<sub>4</sub> for solar cells,<sup>53,54</sup> we demonstrate solution-deposited synthesis of Cu<sub>2</sub>BaSnS<sub>4</sub> using this chemistry in Fig. S30 (ESI†), where we co-dissolved Cu<sub>2</sub>S, BaS, and Sn in propylamine–CS<sub>2</sub>–pyridine to create a mixed precursor ink.

While our newly enhanced solution chemistry for BaS could be used with the previously described routes for homogeneous





**Fig. 6** (a) Reaction schematic illustrating the preparation process of a hybrid colloidal molecular precursor ink using binary metal sulfides. (b) X-ray diffraction pattern of the as-coated hybrid precursor film, with an inset displaying the SEM top view of the as-annealed film. (c) X-ray diffraction pattern of the  $\text{BaZrS}_3$  film obtained after the sulfurization of the as-coated film, with an inset showing the SEM top view of the sulfurized film. (d) Kubelka–Munk transformation applied to the diffuse reflectance spectra of the  $\text{BaZrS}_3$  film derived from hybrid precursor film (considering direct bandgap)<sup>8</sup> (e) X-ray diffraction pattern of the  $\text{BaTiS}_3$  film obtained after sulfurizing the as-coated  $\text{BaS-TiS}_2$  film from the hybrid precursor route.

precursor inks and stacked films as Ba source, here we describe a new hybrid molecular precursor route employing dissolved BaS and colloidal suspension of  $\text{ZrS}_3$ .  $\text{ZrS}_2$  and  $\text{ZrS}_3$  are van der Waals bonded 2D materials, suggesting the potential for colloidal suspensions in a suitable solvent. However,  $\text{ZrS}_2$  is prone to oxidation upon exposure to air, prompting us to utilize  $\text{ZrS}_3$  as the Zr precursor. Compared to a complete molecular precursor approach, colloidal particles in ink could facilitate nucleation in the coating process, leading to favorable film morphology. The hybrid molecular precursor route featuring dissolved BaS in an amine- $\text{CS}_2$  reactive solvent system with pyridine as the bulk solvent and colloidal  $\text{ZrS}_3$  flakes dispersed also builds upon the concept of  $\text{ZrS}_3$  as a plausible precursor outlined in our previous publication.<sup>27</sup> The resulting ink

underwent sonication for several days to achieve a homogeneous dispersion solution, with the Ba:Zr ratio adjusted to be stoichiometric. Subsequently, the ink was coated and sulfurized (refer to Fig. 6(a) for a schematic of the synthesis process). Despite the reasonable air stability of BaS and  $\text{ZrS}_3$ , the solution was coated in an oxygen-free glovebox in this study. The as-coated film contained both BaS and  $\text{ZrS}_3$ , as confirmed in Fig. 6(b), and upon sulfurization, this method yielded phase-pure  $\text{BaZrS}_3$  at 575 °C (refer to Fig. 6(c) and Fig. S31, ESI†). EDX measurements of this film (depicted in Fig. S32, ESI†) verified the presence of Ba, Zr, and S in the grains. However, we observed that the size of the  $\text{ZrS}_3$  colloidal particles influenced the reaction time. In one instance,  $\text{ZrS}_3$  powder was synthesized by sulfurizing Zr nanopowder, while in another case, bulk  $\text{ZrS}_2$

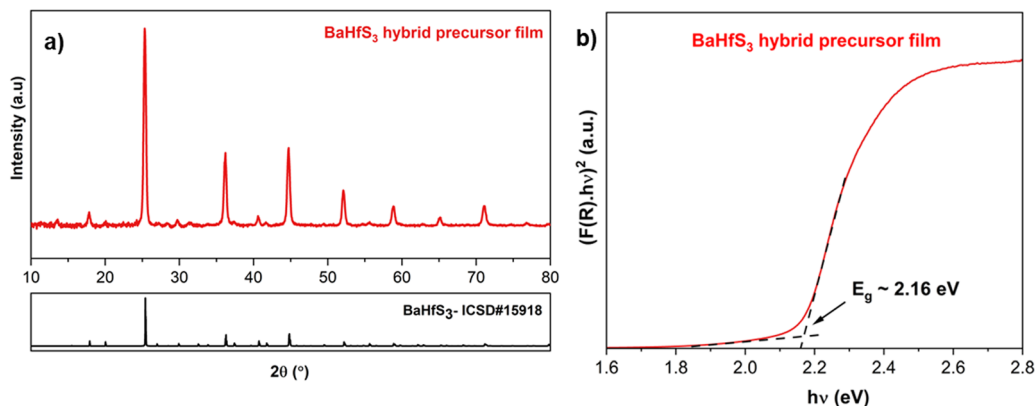


Fig. 7 (a) X-ray diffraction pattern of the BaHfS<sub>3</sub> film obtained after sulfurizing the as-coated BaS–HfS<sub>3</sub> film from the hybrid precursor route. (b) Kubelka–Munk transformation applied to the diffuse reflectance spectra of the BaHfS<sub>3</sub> film derived from hybrid precursor film (considering direct bandgap).<sup>8</sup>

powder was sulfurized to form ZrS<sub>3</sub> powder, both at 575 °C for 18 h. The average grain size of ZrS<sub>3</sub> produced from Zr nanopowder was significantly smaller than that from ZrS<sub>2</sub> powder, resulting in a faster reaction to form BaZrS<sub>3</sub>. With the larger ZrS<sub>3</sub> particles, we observed some unreacted BaS<sub>3</sub> and ZrS<sub>3</sub>, along with BaZrS<sub>3</sub>, in the film (refer to Fig. S33, ESI†). However, in both scenarios, the targeted continuous thin film was not achieved, leading to the formation of isolated large grains of BaZrS<sub>3</sub> (as indicated in the inset of Fig. 6(c) and Fig. S34, ESI†) rather than the intended continuous thin film. As previously noted, the overgrowth to form material islands is a common challenge for methods relying on a BaS<sub>x</sub> liquid flux. Therefore, controlling the quantity of liquid flux is crucial for this approach. Additionally, an alternative synthesis of ZrS<sub>3</sub> to produce smaller particles in the suspension could further decrease reaction time and improve film morphology. Nevertheless, the diffuse reflectance measurements confirmed a bandgap of 1.86 eV for the BaZrS<sub>3</sub> film, promising its potential utilization in tandem solar cell applications (depicted in Fig. 6(d)). Notably, this method was successfully extended for synthesizing related chalcogenide perovskite BaHfS<sub>3</sub> and hexagonal BaTiS<sub>3</sub>, as confirmed by the X-ray diffraction and Raman spectra (see Fig. 6(a), 7(a) and S35, S36, ESI†). Moreover, the diffuse reflectance indicated a bandgap of approximately 2.16 eV for BaHfS<sub>3</sub>, promising its potential utility in optoelectronic applications (shown in Fig. 7(b)).

This study represents the first account of a pathway solely employing metal sulfide precursors for the solution-processed synthesis of BaZrS<sub>3</sub> and other related compounds. Furthermore, the dissolution of BaS in propylamine–CS<sub>2</sub> exhibits novelty. Subsequent studies could explore solution chemistry aimed at co-dissolving BaS and ZrS<sub>2</sub>/ZrS<sub>3</sub> to establish a fully dissolved molecular precursor route, which would also aid in limiting excess BaS<sub>x</sub> flux and controlling grain growth.

#### 2.4. Utilizing selenium liquid flux: solid-state synthesis of BaMS<sub>3</sub>

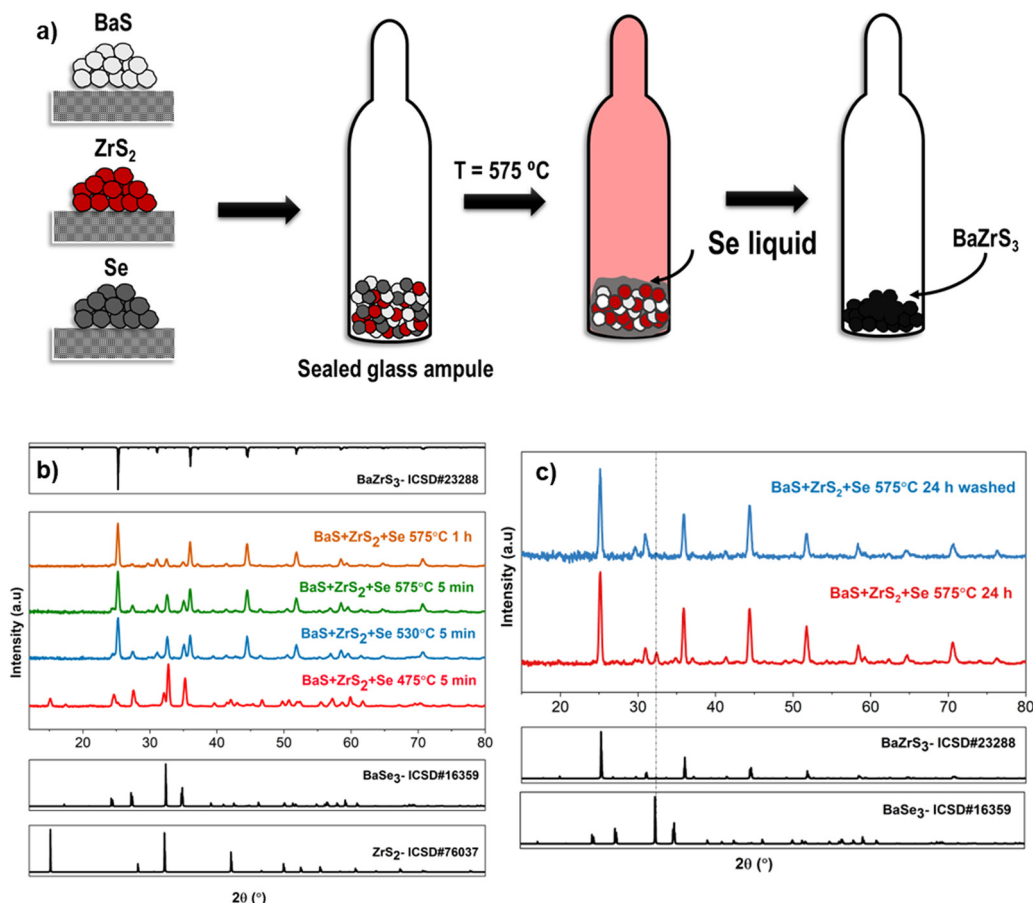
From the above discussions, it is evident that a liquid flux is paramount in synthesizing chalcogenide perovskites at lower

temperatures. BaS<sub>x</sub> liquid flux has been pivotal in the previously discussed solution-processing routes for synthesizing BaMS<sub>3</sub> compounds. This section explores other potential liquid fluxes for synthesizing BaMS<sub>3</sub> and related compounds. Prior studies on chalcogenide materials have employed various liquid fluxes to lower synthesis temperatures. Examples include Cu<sub>2–x</sub>Se flux in Cu-based chalcogenide semiconductors, Sb<sub>2</sub>S<sub>3</sub> liquid flux in antimony-based semiconductors, and BaS<sub>x</sub> in barium-based chalcogenide semiconductors.<sup>27,29,55–57</sup> As mentioned earlier, an ideal liquid flux should meet strict requirements: it should be in a liquid state at the reaction temperature, partially dissolve the metal precursors to reduce diffusion barriers, and avoid reacting with precursors to form undesired secondary phases. A selenium liquid flux can likely fulfill these requirements—it remains liquid above 220 °C, partially dissolves metal sulfides, and the instability of BaMSe<sub>3</sub> compounds could potentially prevent the formation of ternary selenide secondary phases. As a proof of concept, solid-state samples were synthesized by combining BaS, ZrS<sub>2</sub>, and elemental Se in a sealed ampule and heating to a target temperature (refer to Fig. 8(a)). X-ray diffraction measurements from samples quenched at different temperatures showed that BaS and ZrS<sub>2</sub> dissolved in the selenium liquid, as evidenced by the absence of BaS diffraction peaks at 475 °C and diminished ZrS<sub>2</sub> peaks. However, the emergence of BaSe<sub>3</sub> peaks indicated the partial breakdown of BaS to form BaSe<sub>3</sub>. As the temperature increased to 530 °C, intense peaks of BaZrS<sub>3</sub> were observed, accompanied by a diminished secondary phase of BaSe<sub>3</sub>. This increase in BaZrS<sub>3</sub> intensity continued to 575 °C, see Fig. 8(b). One hypothesis for this success is that BaS and ZrS<sub>2</sub> do not significantly break down into BaSe/BaSe<sub>3</sub> and ZrSe<sub>2</sub>/ZrSe<sub>3</sub> at the temperatures used. If they did, at least some byproduct sulfur would have escaped to the vapor phase, resulting in insufficient sulfur for BaZrS<sub>3</sub> formation. The absence of significant binary selenides supports this hypothesis—binary sulfides simply dissolve as binaries and reconstitute as BaZrS<sub>3</sub> without the need for additional sulfur in the mixture.

With increased reaction time, the relative peak intensity of BaZrS<sub>3</sub> to BaSe<sub>3</sub> continued to rise, albeit with a small amount of







**Fig. 8** (a) Reaction schematic depicting the synthesis of BaMS<sub>3</sub> utilizing a selenium liquid flux. (b) X-ray diffraction patterns illustrating the evolution of intermediate phases with temperature for binary sulfides when subjected to the selenium liquid flux. (c) Comparison plot of the X-ray diffraction pattern for the water-washed and unwashed BaZrS<sub>3</sub> powder.

residual BaSe<sub>3</sub> secondary phase remaining after 24 h of reaction at 575 °C. This residual BaSe<sub>3</sub> was found to be readily soluble in water. Consequently, the BaZrS<sub>3</sub> powder, containing the BaSe<sub>3</sub> secondary phase, underwent two rinses with deionized water to achieve a pure BaZrS<sub>3</sub> phase, as confirmed by X-ray diffraction analysis (see Fig. 8(c)). Interestingly, in the absence of excess sulfur and the presence of a liquid flux, BaZrS<sub>3</sub> formed instead of the Ruddlesden–Popper phase. This finding and previous results in this study suggest that Ruddlesden–Popper phases form in a sulfur-poor environment only when there are diffusional limitations. However, Raman analysis in Fig. S37 (ESI†) suggested the presence of residual melt from the selenium liquid flux, encapsulating the BaZrS<sub>3</sub> grains. EDX analysis of the water-washed BaZrS<sub>3</sub> powder further supported this observation. The BaZrS<sub>3</sub> powder exhibited lumping and possibly contained agglomerates of finer BaZrS<sub>3</sub> grains in the nanometer range, as depicted in Fig. S38 (ESI†), with small selenium-rich regions illustrated in Fig. S39 (ESI†). However, this excess selenium could be readily dissolved in various solution chemistries, with amine–thiol mixtures and trioctylphosphine being some solvents.<sup>58,59</sup> Fig. S40 (ESI†) highlights our attempt to dissolve this residual selenium. Notably,

compositional analysis of the BaZrS<sub>3</sub> powder revealed negligible selenium content (see Fig. S41, ESI†).

A similar procedure led to the synthesizing of ternary perovskite BaHfS<sub>3</sub> and hexagonal BaTiS<sub>3</sub>, as evidenced in Fig. S42 and S45 (ESI†). This approach's success lies in synthesizing sulfide perovskites and in the potential for selenium alloying at moderate temperatures by heating a mixture of BaS, ZrS<sub>2</sub>, Zr, and selenium powder together. However, this process would require optimization and, therefore, would be investigated as part of a separate study. Although BaMS<sub>3</sub> compounds have been successfully synthesized at temperatures below 600 °C, attempts to synthesize SrMS<sub>3</sub> compounds *via* sulfurization of respective binary sulfides with excess sulfur have not yielded successful results (see Fig. S46, ESI†). One reason could be that, unlike BaS<sub>x</sub> liquid flux at temperatures above 525 °C, Sr has not been reported to form any liquid strontium polysulfides at temperatures below 600 °C.

When the binary sulfides of strontium and zirconium were heated with selenium at 575 °C, we observed the emergence of α-SrZrS<sub>3</sub> peaks (needle-like crystal structure), as depicted in Fig. S47 (ESI†). However, achieving phase pure α-SrZrS<sub>3</sub> proved to be difficult in this case. It should be noted that single crystal



analysis on a similar sample subjected to 650 °C revealed a striking similarity in the diffraction pattern between  $\alpha$ -SrZrS<sub>3</sub> and the somewhat unlikely compound SrSeS<sub>3</sub>. Rigorous elemental characterizations are needed in the future to differentiate between these phases accurately. We noted a gradual increase in peak intensity from 48 h to 7 days and 21 days, as illustrated in Fig. S47 (ESI†). These results are promising and encourage further exploration of other liquid fluxes. Moreover, this outcome underscores the utility of selenium liquid flux and highlights that the synthesis of Sr ternaries might be thermodynamically favorable at these temperatures but suffers from diffusional limitations.

### 3. Conclusions

Chalcogenide perovskites, while promising, have posed challenges in the synthesis, hindering progress in this material system. BaZrS<sub>3</sub>, particularly suitable for photovoltaic applications, has been synthesized at temperatures unsuitable for solar cell integration. Our study aimed to ascertain the optimal metal ratios and sulfur pressure during the synthesis of BaZrS<sub>3</sub> to prevent phase disintegration. Subsequently, we proposed methods to address limitations arising from the halide affinity of alkaline earth metals and the unavailability of solution chemistries for alkaline earth metal sulfides to fabricate solution-deposited BaZrS<sub>3</sub> thin films.

We successfully dissolved alkaline earth sulfides such as BaS and SrS, demonstrating their utility in synthesis. Additionally, we showcased the successful synthesis of BaMS<sub>3</sub> by employing stacked films of appropriate precursors. Notably, the sequencing of stacked layers allowed the use of ZrCl<sub>4</sub> precursor without forming BaCl<sub>2</sub> impurity in the final BaZrS<sub>3</sub> film. Furthermore, we identified an alternative liquid in Se that facilitates the formation of BaMS<sub>3</sub> at temperatures below 600 °C. The versatility of our methods extends to the synthesis of other BaMS<sub>3</sub> (M = Hf, Ti) compounds, enabling low-cost and facile production of this class of chalcogenide perovskites.

## 4. Experimental methods

### 4.1. Materials and characterization

Barium sulfide (BaS, 99.9%), titanium(IV) sulfide (TiS<sub>2</sub>, 99.9%), selenium powder (99.99%), zirconium(IV) chloride (ZrCl<sub>4</sub>, anhydrous powder, > 99.99%), hafnium(IV) chloride (HfCl<sub>4</sub>, purified by sublimation, 99.9%), 2-methyl-2-propanethiol (MePT, 99%), butylamine (BA, 99.5%), carbon disulfide (CS<sub>2</sub>, anhydrous, > 99%), pyridine (Pyd, anhydrous, 99.8%), *N,N* dimethyl formamide (DMF, anhydrous, 99.8%), thiourea (TU, ACS Reagent, > 99%), sulfur flakes (> 99.99%), molecular sieves (Type 3A, bead size 8–12 mesh), propylamine (PA, > 99%) and alumina dispersed in isopropanol (20 wt%) were purchased from Sigma Aldrich. Bis(pentamethylcyclopentadienyl)barium (Cp\*<sub>2</sub>Ba), zirconium(IV) sulfide (ZrS<sub>2</sub>, 99%), zirconium(IV) bromide (ZrBr<sub>4</sub>, 98%) and zirconium(IV) iodide (ZrI<sub>4</sub>, 99.5%) were purchased from STREM chemicals. Titanium(IV) chloride (TiCl<sub>4</sub>, 99.99%)

and pyridine-d{5} (99.5% Isotopic) were purchased from Fisher Scientific. Titanium hydride (99.9%) and hafnium hydride (99.9%) were purchased from Nano Research Elements. Borosilicate glass ampules were procured from Chemglass. Hafnium disulfide (> 99.995%) was purchased from Ossila.

Solvents without sure seal were stored over molecular sieves for several days before use. Thiourea was recrystallized through a two-step process, using 18.2 MΩ deionized water and was dried under vacuum overnight. All other chemicals were used as received.

Alkali-free Eagle XG glass substrates were purchased from Stemmerich. It should be noted that the glass substrate itself contains traces of zirconium. Selenium powder was vacuum dried at 100 °C overnight before use. Sulfur flakes were finely ground inside a nitrogen-filled glovebox and vacuum-dried overnight at room temperature. All chemicals were stored in nitrogen-filled glove boxes. Sulfurization at 650 °C was carried out in a carbon coated quartz ampule sealed using a propane-oxygen torch.

Raman spectroscopy was performed using a Horiba/Jobin-Yvon HR800 Raman spectrometer with a 632.8 nm excitation laser wavelength. Spectra on liquid solutions were collected using a quartz cuvette enclosed in a nitrogen atmosphere.

X-ray diffraction analysis was conducted using a Rigaku SmartLab Diffractometer under ambient conditions, employing parallel beam geometry with an incident beam angle of 0.5 degrees. Data collection utilized a Cu Kα source ( $\lambda = 1.5406 \text{ \AA}$ ) operated at 40 kV and 44 mA.

Scanning electron microscopy (SEM) and energy dispersive X-ray measurements were performed using the FEI Nova three-dimensional system equipped with an Everhart Thornley detector. The measurements were carried out at an accelerating voltage of 10 kV with a working distance of 5 mm. The samples were coated with ~10 nm of platinum for better image resolution.

Reflectance data were obtained employing a PerkinElmer Lambda 950 spectrometer equipped with an integrating sphere.

NMR spectra were recorded on a Bruker AVANCE III 400 MHz spectrometer, and chemical shifts were referenced to residual solvent signals for both <sup>1</sup>H. For <sup>1</sup>H NMR, the zg30 pulse sequence was utilized with 9 scans, a 1-second relaxation delay, and an acquisition time of 2.1955 seconds.

### 4.2. Alumina deposition

A thin layer of alumina was coated onto the glass substrate to improve the surface wettability during the coating process. The alumina also acts as a barrier and prevents cationic diffusion from substrate. Eagle XG glass was thoroughly rinsed with DI water, 2-propanol, and methanol, followed by sonication in an alconox bath and DI water. The substrate was blow dried using a nitrogen gun. Subsequently, the glass was cleaned in a UV-ozone cleaner for 30 minutes. Commercial alumina nanoparticle solution was diluted by combining 0.2 ml of the commercial solution with 1.8 ml of Isopropanol. The resulting solution was spin-coated onto the cleaned EXG glass at 1000 RPM for



Table 2 Synthesis methods discussed in this work

| Method                 | Precursors used                                          | Powder/thin film | Product                          |
|------------------------|----------------------------------------------------------|------------------|----------------------------------|
| Solid-state synthesis  | BaS + ZrS <sub>2</sub> powders                           | Powders          | BaZrS <sub>3</sub> and RP phases |
| Stacked metal sulfides | MS <sub>3</sub> using halide precursors + BaS stack      | Thin film        | BaMS <sub>3</sub> M = Ti, Zr, Hf |
| Hybrid colloidal inks  | Colloidal MS <sub>3</sub> + BaS using sulfide precursors | Thin film        | BaMS <sub>3</sub> M = Ti, Zr, Hf |
| Selenium flux          | BaS + MS <sub>2</sub> powders with Se                    | Powders          | BaMS <sub>3</sub> M = Ti, Zr, Hf |

1 minute, followed by annealing at 100 °C for 1 minute and then at 500 °C for 30 minutes.

### 4.3. Sample preparation

All chemical storage, solution preparation, thin film coating, and sample handling were conducted inside nitrogen- and ultra-high purity (UHP) argon-filled gloveboxes. The ampules were sealed using a Schlenk line with UHP argon as the purging gas. Table 2 summarizes the four different synthesis methods discussed in this work.

**Solid-state synthesis.** BaS, ZrS<sub>2</sub> and S were loaded into borosilicate ampules in the desired ratios. Before sealing with a butane–air torch, the ampules underwent three cycles of vacuum–argon purging. The heat treatment was performed in a refractory tube furnace at the designated temperature and duration, followed by natural cooling.

The sulfur pressure was calculated at the synthesis temperature using the ideal gas law, with S<sub>6</sub> assumed to be the predominant sulfur species. Borosilicate ampules with a nominal volume of 10 ml were used; however, for the calculations, the volume was considered slightly larger to account for the additional volume contributed by the ampule stem.

The synthesized powder was subjected to two rinses with DI water to eliminate water-soluble secondary phases such as BaS<sub>3</sub>. The powder was then dried with flowing argon gas.

#### Stacked metal sulfides method

**Amine–thiol.** A 0.3 M solution of ZrCl<sub>4</sub> was prepared with the butylamine: 2-methyl-2-propanethiol molar ratio of 1:2.2. 2-methyl-2-propanethiol was added first to the metal precursor, followed by butylamine. The solution was stirred overnight at 35 °C and used for coating.

**Amine–CS<sub>2</sub>.** A 0.3 M solution of ZrCl<sub>4</sub> was prepared with a butylamine:CS<sub>2</sub>:pyridine molar ratio of 1:1:1. Butylamine was added first to the metal precursor, followed by CS<sub>2</sub> and pyridine. The solution was stirred overnight at 35 °C and used for coating. Similar dissolution procedures were followed for ZrBr<sub>4</sub>, ZrI<sub>4</sub>, TiCl<sub>4</sub>, and HfCl<sub>4</sub>.

**Amine–thiourea.** A 0.3 M solution of ZrCl<sub>4</sub> and 2 M thiourea in butylamine was stirred overnight at 35 °C and used. ZrCl<sub>4</sub> and thiourea were mixed together, and butylamine was added.

**DMF–thiourea.** A 0.3 M solution of ZrCl<sub>4</sub> and 3 M thiourea in dimethylformamide (DMF) was stirred overnight at 35 °C and used. ZrCl<sub>4</sub> and thiourea were mixed together, and DMF was added.

**Cp\*<sub>2</sub>Ba dissolution.** A 0.3 M solution of Cp\*<sub>2</sub>Ba was prepared with the butylamine: 2-methyl-2-propanethiol molar ratio of

1:1. 2-methyl-2-propanethiol was added first to the metal precursor, followed by butylamine.

**Bilayer fabrication.** In one route, the four ZrCl<sub>4</sub> solutions routes discussed above were coated, annealed, and sulfurized to fabricate the bottom layer of ZrS<sub>3</sub>. The samples were sealed in an ampule with HfH<sub>2</sub> and sulfur and heated to 575 °C for 6 h to synthesize the bottom ZrS<sub>3</sub> layer. Cp\*<sub>2</sub>Ba was blade coated on top of ZrS<sub>3</sub>, sealed in an ampule, and heated at 575 °C for 2 h to synthesize BaZrS<sub>3</sub>. Similar procedure was adapted for the iodide and bromide salts as well as BaHfS<sub>3</sub> and BaTiS<sub>3</sub>.

In another route, the ZrCl<sub>4</sub>–amine–CS<sub>2</sub> ink was coated and annealed on the hotplate at 450 °C for 30 minutes. Cp\*<sub>2</sub>Ba was blade coated on top, sealed in an ampule, and heated at 575 °C for 2 h to synthesize BaZrS<sub>3</sub>.

**Hybrid colloidal precursors method.** ZrS<sub>3</sub> powder was synthesized by heating Zr powder with excess sulfur sealed in an ampule at 575 °C for 16 h. A 0.2 M suspension of this ZrS<sub>3</sub> powder was sonicated in pyridine for 2 days to achieve a stable colloidal solution. Simultaneously, a 0.2 M solution of BaS was prepared with propylamine:CS<sub>2</sub>:pyridine molar ratio of 1:1:1.5. Propylamine was added first to the metal precursor, followed by CS<sub>2</sub> and pyridine. The solution was stirred for 3–4 days at 35 °C and combined with the ZrS<sub>3</sub> colloidal solution. The solution was then blade-coated, sealed, and sulfurized.

Similar procedures were repeated for Ba–Hf–S and Ba–Ti–S synthesis, with the only difference being the starting precursors. HfH<sub>2</sub> and TiH<sub>2</sub> were used as precursors to synthesize binary sulfides.

**Sulfurization.** The following procedure was followed for the thin film deposition in the stacked metal sulfides and hybrid colloidal precursors method.

Solutions were blade-coated using an automated blade coater at a speed of 15 mm per second with a single pass, followed by annealing at 350 °C for 2 minutes on a hot plate and cooling for 1 minute. The coating was repeated 6–8 times on alumina-coated Eagle XG substrates. The resulting films were transferred to a 5 ml ampule containing 5 mg of HfH<sub>2</sub> and 15 mg of sulfur. The ampules underwent three purging cycles with vacuum-UHP argon and were sealed using a butane-fueled blowtorch. Subsequently, the ampules were heated in a refractory furnace to the desired temperature and cooled naturally under a slow argon flow.

**Selenium liquid flux.** The powder synthesis experiments were conducted using borosilicate ampules, which were loaded with either 0.2 mmol BaS, 0.2 mmol ZrS<sub>2</sub>, and 30 mg selenium in a 20 ml ampule, or 0.1 mmol BaS, 0.1 mmol ZrS<sub>2</sub>, and 15 mg selenium in a 10 ml ampule. Before sealing with a butane–air



torch, the ampules underwent three cycles of vacuum–argon purging. The heat treatment was performed in a refractory tube furnace at the designated temperature and duration, followed by natural cooling.

The synthesized BaMS<sub>3</sub> (M = Ti, Zr, Hf) powder was subjected to two rinses with DI water to eliminate any water-soluble secondary phases. The water-washed powder was further washed in a 1 : 5 vol : vol mixture of ethanedithiol–propylamine to wash away any residual selenium. The powder was stirred in amine–thiol solvent overnight at room temperature. The powder was afterward dried with flowing argon gas.

## Data availability statement

The data supporting this article have been included in the ESI.†

## Conflicts of interest

The authors declare no conflict of interest.

## Acknowledgements

The authors acknowledge the National Science Foundation's financial support through grants 1735282-NRT (SFEWS) and 10001536 (INFEWS). Special thanks are extended to Prof. Suzanne C. Bart, Dr Apurva A. Pradhan, Dr Jonathan W. Turnley, and Dr Madeleine C. Uible for their valuable discussions regarding the project.

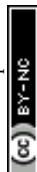
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