

## REVIEW

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# Selenoborates: a latent structure resource for infrared functional crystal materials

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Optoelectronic functional crystal materials are crucial components in laser systems and could be applied as polarized light converters, modulators, photorefractive devices, window materials etc. Borates and chalcogenides are well-established sources of ultraviolet (UV) and infrared (IR) optoelectronic functional crystal materials. The selenoborate system combines the structural diversity of borates with the extensive optical transmission range of metal selenides, showing potential as IR functional materials. However, research on selenoborates remains limited. This review aims to stimulate further research into IR optoelectronic functional crystal materials within the selenoborate system. First, this work offers a comprehensive review of all selenoborates (21) reported to date, categorizing them according to their structural motifs from zero to three dimensions. Second, the unique synthesis methods of selenoborates are summarized. Additionally, first-principles calculations are utilized to analyze the bandgaps and nonlinear optical (NLO) properties of these materials. As a result, it is concluded that selenoborates possess considerable potential for further investigation as IR functional materials, especially in terms of the structures and properties regulated by excellent microstructural groups.

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## 1. Introduction

The development of laser technology is highly regarded by scientific researchers, and has propelled rapid advancements in various fields, such as industrial manufacturing, medicine, communication and the military.<sup>1–3</sup> Optoelectronic functional crystal materials are essential components of laser technology, with applications extending to laser crystals, modulators, frequency multipliers, and optical window materials.<sup>4–7</sup> Currently, prevalent systems of optoelectronic functional crystal materials include borates, phosphates, carbonates, nitrates, chalcogenides, phosphides and so on.<sup>8–24</sup>

Exploration of suitable systems may promote the development of target materials in this field. Since the determination of the first borate single crystal structure,  $\text{Be}_2\text{BO}_3(\text{OH})$ , in 1931, a growing number of well-characterized borates have been discovered and utilized as ultraviolet (UV) and deep-ultraviolet (DUV) optoelectronic functional crystals. Notable examples include  $\text{KBe}_2\text{BO}_3\text{F}_2$  (KBBF),  $\beta\text{-BaB}_2\text{O}_4$  ( $\beta$ -BBO) and  $\text{LiB}_3\text{O}_5$  (LBO).<sup>25–36</sup> Recently, Pan's group advanced research in

the fluoroborate system, reaching significant milestones in DUV nonlinear optical (NLO) crystals. Phosphate crystals such as  $\text{KH}_2\text{PO}_4$  (KDP) and  $\text{KTiOPO}_4$  (KTP) have been employed as optoelectronic functional crystal materials in the visible to near-infrared (IR) region. Meanwhile, chalcopyrite-type chalcogenides  $\text{AgGaS}_2$  (AGS) and  $\text{AgGaSe}_2$  (AGSe) have been widely used as commercial optoelectronic functional crystal materials in the mid- to far-IR region.<sup>37–41</sup> Combining the advantages of borates and chalcogenides, thioborates possess structural diversity and suitable transmission ranges. The exploration of the thioborate system by the Pan and Mao groups has led to the discovery of high-performance IR optoelectronic functional materials, such as  $\text{BaB}_2\text{S}_4$  and  $\text{LaBS}_3$ .<sup>42–55</sup> Thus, exploring and developing specific systems are crucial for discovering new high-performance materials.

Metal chalcogenides represent one of the most extensively studied families for obtaining IR optoelectronic functional crystal materials, demonstrating significant importance in both civil and military fields.<sup>56–65</sup> Compared to isostructural sulfides, metal selenides exhibit wider IR transmittance and larger SHG responses.<sup>66–68</sup> Besides, they have lower melting points in most cases, which is beneficial for crystal growth. With the expectation of obtaining high-performance IR functional crystals, selenoborates derived from thioborates naturally attract our attention. Up to now, there are few cases of selenoborates, most of which are reported by Krebs from the 1990s to the 2000s. In the selenoborate system,  $[\text{BSe}_3]$  and

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[BSe<sub>4</sub>] are the basic structural units, which could be interconnected to form zeolitic [B<sub>7</sub>Se<sub>13</sub>] and hypertetrahedral [B<sub>10</sub>Se<sub>20</sub>] units.<sup>69</sup> This phenomenon indicates greater space for discovering structural diversity in the selenoborate system (Fig. 1).

In this paper, the calculated microstructural performances of related units indicate that the vibrational frequencies of [BSe<sub>3</sub>] and [BSe<sub>4</sub>] show obvious shifts to lower absorption energies (maximum 725 cm<sup>-1</sup>) in comparison with those of [BS<sub>n</sub>] (*n* = 3, 4) (maximum 911 cm<sup>-1</sup>), which indicates that selenoborates have a long IR absorption edge (725 cm<sup>-1</sup>, >12 μm) that covers two atmospheric transparent windows (3–5 and 8–12 μm). In addition, [BSe<sub>3</sub>] demonstrates the highest polarizability anisotropy and hyperpolarizability (Fig. 2) among related units (BQ<sub>x</sub> (Q = S, Se, *x* = 3, 4), GaQ<sub>4</sub> (Q = S, Se), ASe<sub>4</sub> (A = Zn, As, Sn, Ge, Si, Al)). The highest polarizability anisotropy and hyperpolarizability values suggest the ability of selenoborates demonstrating large birefringence and SHG responses. The above results theoretically indicate that selenoborates may be potential IR functional materials.

Besides, we present a comprehensive structural review of the 21 reported selenoborates cataloged in the Inorganic Crystal Structure Database (ICSD) with version 5.2.0 being the latest release as ICSD-2024/01. These selenoborates are classified into four categories based on the polymerization degree of their anionic units (Table 1). We analysed their structural characteristics and found that the majority of these selenoborates were formed from [BSe<sub>4</sub>], with only three consisting of [BSe<sub>3</sub>]. This is different from thioborates, among which most of them are composed of [BS<sub>3</sub>]. Furthermore, we summarized the synthesis methods employed for these selenoborates. The challenge of B–Si exchange occurring at temperatures above 300–400 °C is addressed by using carbon-coated silica tubes and glassy carbon crucibles during the synthesis process. In addition, we calculated the bandgaps and SHG coefficients of selected selenoborates using density functional theory (DFT) calculations to investigate the properties of selenoborate system. The paper also discussed future directions for enhancing the structural diversity and optimizing the second-order

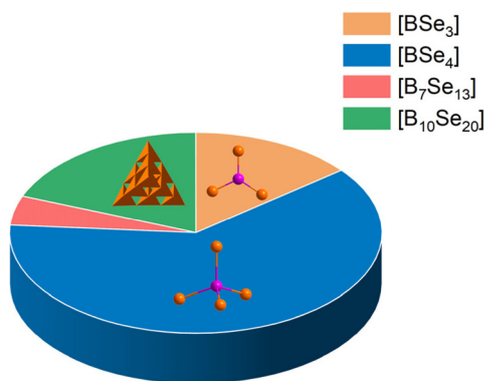


Fig. 1 The building units of reported selenoborates (based on the ICSD).

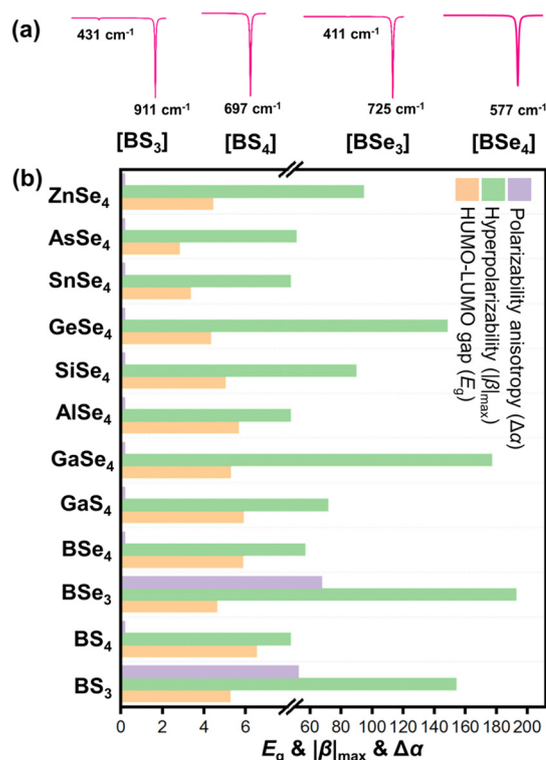


Fig. 2 The comparison of the microstructural performances of related units. (a) Vibrational data; (b) Polarizability anisotropy, hyperpolarizability and HUMO-LUMO gap.

nonlinearity of selenoborates, aiming at advancing the development of these material systems.

## 2. Syntheses

At present, the discovery of selenoborates remains limited, primarily due to the difficulty of their synthesis. This section provides an overview of the commonly used synthesis methods, the prevailing difficulties in this field and distinctive synthesis methods for reference. Similar to that of thioborates, most selenoborates are synthesized *via* solid-state reactions involving metal selenides, amorphous boron, and selenium as raw materials. These reactions are typically conducted in evacuated carbon-coated silica tubes under precisely controlled conditions, often at temperatures ranging from 600 to 950 °C. However, obtaining highly pure selenoborates with well-defined structures remains a formidable challenge. This difficulty is largely attributed to the high reactivity of boron chalcogenides with various container materials at elevated temperatures. In particular, the widely employed fused silica tubes prove vulnerable to the corrosive effects of boron selenide when temperatures surpass the threshold of 300–400 °C. This interaction prompts the exchange of boron with silicon, instigating the formation of selenosilicate.

To overcome these challenges and procure pure selenoborate samples, two distinctive methods could be employed:

Table 1 Structural information regarding reported selenoborates based on ICSD

Formula	Crystal system	Point group	Space group	Symmetry	Lattice parameters (Å, °)	Building unit	Ref.
<b>Zero dimensional</b>							
Tl <sub>3</sub> BSe <sub>3</sub>	Monoclinic	2/m	P2 <sub>1</sub> /m (No. 11)	CS	$a = 5.547(2), b = 10.099(3), c = 6.852(2), \beta = 97.59(3)$	BSe <sub>3</sub>	70
Ba <sub>7</sub> (BSe <sub>3</sub> ) <sub>4</sub> Se	Monoclinic	2/m	C2/c (No. 15)	CS	$a = 10.513(2), b = 25.021(5), c = 10.513(2), \beta = 90.10(3)$	BSe <sub>3</sub>	71
Ba <sub>3</sub> (BSe <sub>3</sub> ) <sub>2</sub> (SbSe <sub>3</sub> )	Hexagonal	6/m	P62m (No. 189)	NCS	$a = 17.720(4), c = 11.251(3)$	BSe <sub>3</sub>	73
Li <sub>6-2x</sub> Sr <sub>2-2x</sub> B <sub>10</sub> Se <sub>20</sub> ( $x \approx 0.7$ )	Tetragonal	4/m	I4 <sub>1</sub> /a (No. 88)	CS	$a = 14.735(6), c = 14.145(12)$	B <sub>10</sub> Se <sub>20</sub>	72
<b>One dimensional</b>							
Na <sub>2</sub> B <sub>2</sub> Se <sub>7</sub>	Monoclinic	2/m	C2/c (No. 15)	CS	$a = 11.863(4), b = 6.703(2), c = 13.811(2), \beta = 109.41(2)$	BSe <sub>4</sub>	75
K <sub>2</sub> B <sub>2</sub> Se <sub>7</sub>	Monoclinic	2/m	C2/c (No. 15)	CS	$a = 12.092(4), b = 7.054(2), c = 13.991(5), \beta = 107.79(3)$	BSe <sub>4</sub>	75
CsBSe <sub>3</sub>	Monoclinic	2/m	P2 <sub>1</sub> /c (No. 14)	CS	$a = 7.570(2), b = 12.791(4), c = 6.171(2), \beta = 107.09(2)$	BSe <sub>4</sub>	74
RbBSe <sub>3</sub>	Monoclinic	2/m	P2 <sub>1</sub> /c (No. 14)	CS	$a = 7.2789(15), b = 12.385(3), c = 6.1690(12), \beta = 105.67(3)$	BSe <sub>4</sub>	74
TlBSe <sub>3</sub>	Monoclinic	m	Cc (No. 9)	NCS	$a = 6.1662(12), b = 12.109(2), c = 7.0311(14), \beta = 113.88(3)$	BSe <sub>4</sub>	74
Cs <sub>3</sub> B <sub>3</sub> Se <sub>10</sub>	Triclinic	1	P1 (No. 2)	CS	$a = 7.5831(15), b = 8.4643(17), c = 15.276(3)$	BSe <sub>4</sub>	76
Tl <sub>3</sub> B <sub>3</sub> Se <sub>10</sub>	Triclinic	1	P1 (No. 2)	CS	$a = 107.03(3), \beta = 89.29(3), \gamma = 101.19(3)$	BSe <sub>4</sub>	76
Tl <sub>2</sub> B <sub>2</sub> Se <sub>7</sub>	Monoclinic	2/m	C2/c (No. 15)	CS	$a = 7.0989(14), b = 8.0724(16), c = 14.545(3)$	BSe <sub>4</sub>	76
Rb <sub>2</sub> B <sub>2</sub> Se <sub>7</sub>	Monoclinic	2/m	C2/c (No. 15)	CS	$a = 105.24(3), \beta = 95.82(3), \gamma = 92.79(3)$	BSe <sub>4</sub>	76
<b>Two dimensional</b>							
Ba <sub>2</sub> B <sub>4</sub> Se <sub>13</sub>	Monoclinic	2/m	P2 <sub>1</sub> /c (No. 14)	CS	$a = 12.790(3), b = 11.560(2), c = 12.862(3), \beta = 103.22(3)$	BSe <sub>4</sub>	77
Ba <sub>2</sub> Se <sub>6</sub>	Orthorhombic	mmm	Cmca (No. 64)	CS	$a = 11.326(2), b = 7.6587(15), c = 10.315(2)$	BSe <sub>4</sub>	77
Li <sub>6-x</sub> Cs <sub>x</sub> B <sub>10</sub> Se <sub>18</sub> ( $x \approx 1$ )	Tetragonal	4/mmm	I4 <sub>1</sub> /amd (No. 141)	CS	$a = 10.0285(14), b = 10.0285(14), c = 27.743(6)$	B <sub>10</sub> Se <sub>20</sub>	78
<b>Three dimensional</b>							
CuBSe <sub>2</sub>	Tetragonal	I42m	I42d (No. 122)	NCS	$a = 5.539, b = 5.539, c = 10.734$	BSe <sub>4</sub>	79
Li <sub>2</sub> B <sub>2</sub> Se <sub>5</sub>	Monoclinic	2/m	C2/c (No. 15)	CS	$a = 10.616(2), b = 5.3181(11), c = 12.382(3), \beta = 109.96(3)$	BSe <sub>4</sub>	81
Na <sub>6</sub> B <sub>10</sub> Se <sub>18</sub>	Tetragonal	4/mmm	I4 <sub>1</sub> /acd (No. 142)	CS	$a = 15.128(2), b = 15.128(2), c = 27.955(6)$	B <sub>10</sub> Se <sub>20</sub>	83
Li <sub>6+2x</sub> [B <sub>10</sub> Se <sub>18</sub> ] <sub>x</sub> Se <sub>x</sub> ( $x \approx 2$ )	Monoclinic	2/m	C2/c (No. 15)	CS	$a = 17.411(1), b = 21.900(1), c = 17.820(1), \beta = 101.6(1)$	B <sub>10</sub> Se <sub>20</sub>	84
Li <sub>7</sub> B <sub>7</sub> Se <sub>15</sub>	Tetragonal	4/mmm	P4 <sub>2</sub> /mbc (No. 133)	CS	$a = 11.4107(4), c = 16.4251(5)$	B <sub>7</sub> Se <sub>13</sub>	82

(i) Coating the inner surfaces of the containers with glassy carbon:

The silica tubes are coated on their inner surface with glassy carbon, which can be manufactured by slowly rotating the ampoule filled with acetone vapour helically through the flame of an oxygen–hydrogen welding torch at about 1300 K.

(ii) Utilizing glassy carbon or boron nitride crucibles:

The crucibles, fashioned from glassy carbon or boron nitride, measuring 100 mm in length with cap/outer/inner diameters of 18/12/5 mm, are meticulously sealed with a screw cap. These crucibles function as robust sample containers that are resistant to the corrosive tendencies of selenoborate synthesis.

After these preparatory measures, the crucibles are strategically positioned within outer silica tubes, whose inner walls are lined with glassy carbon. Then the tubes are fused. Given the air and moisture sensitivity inherent in selenoborates, all manipulations are conducted within a controlled environment – a glove box saturated with dry argon. To further shield against oxidation, the synthesized materials are placed within steel or tantalum ampoules, which are enveloped by an argon atmosphere. These ampoules are then carefully placed inside evacuated silica tubes, creating a multi-layered protective system. In addition to solid-state reactions,  $\text{CuBSe}_2$  was synthesized *via* solvothermal methods. The reagents were placed in a 50 mL Teflon-lined autoclave filled to 80% with anhydrous diethylenetriamine, sealed and heated at 180 °C for 36 hours before cooling naturally to room temperature. Despite notable progress in synthesis techniques, obtaining highly pure selenoborates remains a formidable task. One possible method is to perform various purifications on the initially synthesized sample, such as cleaning, recrystallization, and so on. Successful cases are rare, underscoring the intricate nature of this process. With continuous exploration, progress is expected in the synthesis of selenoborates.

### 3. Structural classification

A comprehensive review of the structural motifs of all known selenoborates based on the ICSD database was summarized. Similar to thioborates, selenoborates are characterized by two fundamental building units: the trigonal-planar  $[\text{BSe}_3]$  unit and the tetrahedral  $[\text{BSe}_4]$  unit (Fig. 3a and b). These units can interconnect to form a variety of structures. Additionally, multiple  $[\text{BSe}_4]$  units could be connected with each other to form the  $[\text{B}_7\text{Se}_{13}]$  unit and the supertetrahedral  $[\text{B}_{10}\text{Se}_{20}]$ . Structures containing  $[\text{BSe}_3]$  units are predominantly observed in low-dimensional selenoborates (constituting more than half of the 0D selenoborates). All 1D structures have  $[\text{BSe}_4]$  as the building unit.  $[\text{B}_7\text{Se}_{13}]$  units and supertetrahedral  $[\text{B}_{10}\text{Se}_{20}]$  make up most high-dimensional selenoborates. It is worth noting that, unlike thioborates, selenoborates tend to form high-dimensional structures such as chains and layers, which may be related to the fact

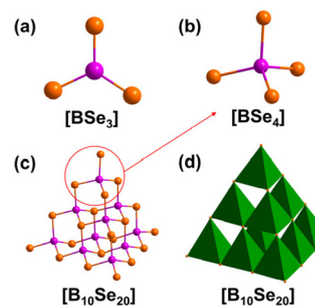


Fig. 3 The structures of (a)  $[\text{BSe}_3]$ , (b)  $[\text{BSe}_4]$ , and (c) and (d)  $[\text{B}_{10}\text{Se}_{20}]$ .

that  $[\text{BSe}_4]$  appears more commonly than  $[\text{BSe}_3]$  in selenoborates. Specifically, only 3 out of 21 known selenoborates are composed of  $[\text{BSe}_3]$ .

#### 3.1. Selenoborates with zero-dimensional structures

To the best of our knowledge, the four 0D selenoborates predominantly feature two types of structural units:  $[\text{BSe}_3]$  and  $[\text{B}_{10}\text{Se}_{20}]$ . Notably, the supertetrahedron  $[\text{B}_{10}\text{Se}_{20}]$  is constructed from corner-sharing  $[\text{BSe}_4]$  tetrahedra (Fig. 3c and d).

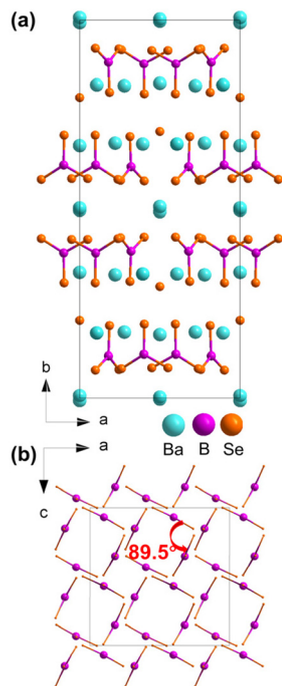
Among these compounds, three exhibit isolated  $[\text{BSe}_3]$  units with varying dihedral angles: 0° for  $\text{Tl}_3\text{BSe}_3$ , 89.5° for  $\text{Ba}_7(\text{BSe}_3)_4\text{Se}$  and 90° for  $\text{Ba}_3(\text{BSe}_3)(\text{SbSe}_3)$ . The other one is composed of the  $[\text{B}_{10}\text{Se}_{20}]$  supertetrahedron. In these selenoborate structures, the B–Se bond lengths range from 1.91(2) to 2.06(2) Å.

**3.1.1. Selenoborates with mononuclear  $[\text{BSe}_3]$  clusters.**  $\text{Tl}_3\text{BSe}_3$ <sup>70</sup> crystallizes in the monoclinic space group  $P2_1/m$  and represents the first reported selenoborate containing only isolated  $[\text{BSe}_3]$  as the anion units. The isolated  $[\text{BSe}_3]$  units are arranged parallel to each other, and are interconnected *via*  $\text{Tl}^+$  cations to form the overall framework. In the structure, the B–Se bond lengths are 1.94(2) Å and 1.97(4) Å. The slight elongation of the bond lengths in  $[\text{BSe}_3]$  may be caused by the relatively high charge on this anion. The Se–B–Se bond angles range from 119(2)° to 121(2)°.

$\text{Ba}_7(\text{BSe}_3)_4\text{Se}$ <sup>71</sup> crystallizes in the monoclinic space group  $C2/c$ , which is the first reported barium selenoborate. The dihedral angle between planar  $[\text{BSe}_3]$  units is 89.5°, and isolated  $[\text{BSe}_3]$  units are interconnected *via*  $\text{Ba}^{2+}$  cations to form the entire structure (Fig. 4). In the structure, the B–Se bond lengths are 1.938(10) Å and 1.992(10) Å, and the Se–B–Se bond angles range from 115.2(5)° to 123.4(5)°.

**3.1.2. The selenoborate with multinuclear  $[\text{B}_{10}\text{Se}_{20}]$  clusters.**  $\text{Li}_{6-2x}\text{Sr}_{2+x}\text{B}_{10}\text{Se}_{20}$  ( $x \approx 0.7$ )<sup>72</sup> crystallizes in the tetragonal space group  $I4_1/a$ , which is the first reported selenoborate containing isolated supertetrahedral  $[\text{B}_{10}\text{Se}_{20}]$ . In its structure, there are isolated  $[\text{B}_{10}\text{Se}_{20}]$  structures that are formed from tetrahedral  $[\text{BSe}_4]$  units, and these  $[\text{B}_{10}\text{Se}_{20}]$  units are connected by  $\text{Li}^+$  and  $\text{Sr}^{2+}$  cations to form the entire structure. The average B–Se bond length is 2.055 Å.

**3.1.3. The selenoborate with heteronuclear 0D clusters.**  $\text{Ba}_3(\text{BSe}_3)(\text{SbSe}_3)$ <sup>73</sup> crystallizes in the hexagonal space group

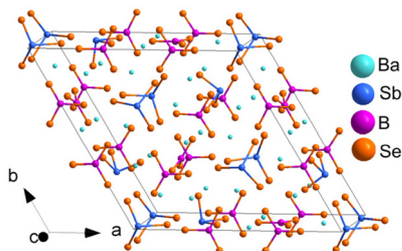


**Fig. 4** (a) The whole structure of  $\text{Ba}_7(\text{BSe}_3)_4\text{Se}$  and (b) arrangement and dihedral angle between planar  $[\text{BSe}_3]$  units in  $\text{Ba}_7(\text{BSe}_3)_4\text{Se}$ .

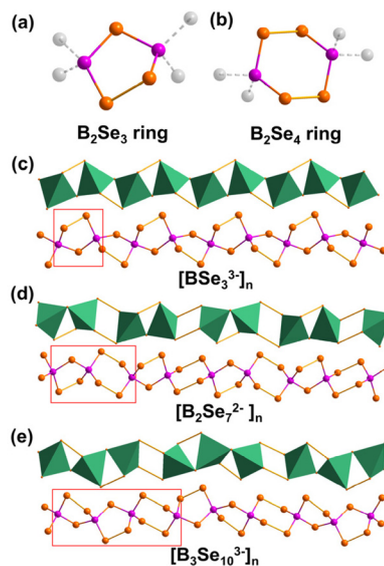
$P\bar{6}2m$ , which is the only example containing heteronuclear 0D clusters. In the structure, there are  $[\text{SbSe}_3]$  pyramids and isolated  $[\text{BSe}_3]$  units, and they are connected by  $\text{Ba}^{2+}$  cations (Fig. 5). The B–Se bond lengths range from 1.91(2) to 2.06(2) Å and the Sb–Se bond lengths range from 2.185(2) to 2.581(2). The Se–B–Se bond angles range from 115.2(9)° to 125.8(8)° and the Se–Sb–Se bond angles range from 91.0(4)° to 176.1(5)°.

### 3.2. One-dimensional selenoborates

All 1D selenoborates are constructed from the mononuclear cluster  $[\text{BSe}_4]$  and Se–Se bonds. Specifically,  $[\text{BSe}_4]$  and Se–Se bonds could form five-membered rings  $\text{B}_2\text{Se}_3$  and six-membered rings  $\text{B}_2\text{Se}_4$ . These units can be further constructed into polymeric chains  $[\text{BSe}_3^{3-}]_n$ ,  $[\text{B}_2\text{Se}_7^{2-}]_n$  and  $[\text{B}_3\text{Se}_{10}^{3-}]_n$  (Fig. 6). The B–Se bond lengths in 1D selenoborates range from 2.017(17) to 2.358(3) Å. The Se–B–Se bond angles range from 90.5(4)° to 118.2(4)°.



**Fig. 5** The structure of  $\text{Ba}_3(\text{BSe}_3)(\text{SbSe}_3)$ .



**Fig. 6** The structures of (a) the  $\text{B}_2\text{Se}_3$  five-membered ring, (b) the  $\text{B}_2\text{Se}_4$  six-membered ring, (c) polymeric chain  $[\text{BSe}_3^{3-}]_n$ , (d) polymeric chain  $[\text{B}_2\text{Se}_7^{2-}]_n$ , and (e) polymeric chain  $[\text{B}_3\text{Se}_{10}^{3-}]_n$ .

**3.2.1. Selenoborates with polymeric  $[\text{BSe}_3^{3-}]_n$  chains.**  $\text{CsBSe}_3$  and  $\text{RbBSe}_3$  crystallize in the monoclinic space group  $P2_1/c$ , and  $\text{TlBSe}_3$  crystallizes in the monoclinic space group  $Cc$ .<sup>74</sup> Besides, they are the first reported selenoborates containing polymeric anionic  $[\text{BSe}_3^{3-}]_n$  chains. In detail, the  $[\text{BSe}_4]$  units are connected both through Se–Se bonds and by sharing one vertex Se atom. First, two adjacent  $[\text{BSe}_4]$  units form a  $\text{B}_2\text{Se}_3$  five-membered ring, and then  $\text{B}_2\text{Se}_3$  five-membered rings helically connect and extend to become the polymeric  $[\text{BSe}_3^{3-}]_n$  chains. In their structures, the  $\text{Cs}^+$ ,  $\text{Rb}^+$  and  $\text{Tl}^+$  cations are inserted in the spaces between the  $[\text{BSe}_3^{3-}]_n$  chains in  $\text{CsBSe}_3$ ,  $\text{RbBSe}_3$  and  $\text{TlBSe}_3$ . The B–Se bond lengths in them range from 2.034(17) to 2.358(3) Å and the Se–B–Se bond angles range from 90.5(4)° to 113.4(4)°.

**3.2.2. Selenoborates with polymeric  $[\text{B}_2\text{Se}_7^{2-}]_n$  chains.**  $\text{Na}_2\text{B}_2\text{Se}_7$ ,  $\text{K}_2\text{B}_2\text{Se}_7$ ,<sup>75</sup>  $\text{Rb}_2\text{B}_2\text{Se}_7$  and  $\text{Tl}_2\text{B}_2\text{Se}_7$ <sup>76</sup> all crystallize in the monoclinic space group  $C2/c$ . In their structures, the  $\text{B}_2\text{Se}_3$  five-membered rings and six-membered rings connect alternately with a ratio of 1 : 1 to form  $[\text{B}_2\text{Se}_7^{2-}]_n$  chains, and the  $\text{Na}^+$ ,  $\text{K}^+$ ,  $\text{Rb}^+$  and  $\text{Tl}^+$  cations are situated between the polymeric  $[\text{B}_2\text{Se}_7^{2-}]_n$  chains. The B–Se bond lengths in this series of structures range from 2.021(8) to 2.062(14) Å. The Se–B–Se bond angles range from 94.3(4)° to 118.2(4)°. The Se–Se bond lengths in them range from 2.3630(14) to 2.3690(19) Å.

**3.2.3. Selenoborates with polymeric  $[\text{B}_3\text{Se}_{10}^{3-}]_n$  chains.**  $\text{Cs}_3\text{B}_3\text{Se}_{10}$  and  $\text{Tl}_3\text{B}_3\text{Se}_{10}$ <sup>76</sup> crystallize in the triclinic space group  $P\bar{1}$ , and they are the first reported selenoborates containing polymeric  $[\text{B}_3\text{Se}_{10}^{3-}]_n$  anionic chains. Therein, the  $\text{B}_2\text{Se}_3$  five-membered rings and six-membered rings occur in a ratio of 2 : 1, and two five-membered rings and one six-membered ring act as the repeatable section in the infinite  $[\text{B}_3\text{Se}_{10}^{3-}]_n$  chain. The  $\text{Cs}^+$  and  $\text{Tl}^+$  cations are located between the poly-

meric chains to balance the charge and hold the structure together. The B–Se bond lengths in them range from 2.017(17) to 2.097(17) Å. The Se–B–Se bond angles range from 94.4(5)° to 116.2(8)°. The Se–Se bond lengths in them range from 2.3396(14) to 2.380(2) Å.

### 3.3. Two-dimensional selenoborates

The 2D selenoborates are composed of only two types of building units: tetrahedral [BSe<sub>4</sub>] and supertetrahedral [B<sub>10</sub>Se<sub>20</sub>]. In these 2D selenoborates, the B–Se bond lengths range from 2.032(7) to 2.080(7) Å. The Se–B–Se bond angles range from 94.4(5)° to 116.2(8)°.

**3.3.1. Selenoborates with 2D polymers constructed from tetrahedral [BSe<sub>4</sub>].** Ba<sub>2</sub>B<sub>4</sub>Se<sub>13</sub><sup>77</sup> crystallizes in the monoclinic space group *P2<sub>1</sub>/c*, which is the first reported alkaline earth perselenoborate. In the structure of Ba<sub>2</sub>B<sub>4</sub>Se<sub>13</sub>, there are B<sub>2</sub>Se<sub>3</sub> five-membered and B<sub>2</sub>Se<sub>4</sub> six-membered rings composed of [BSe<sub>4</sub>] units and Se–Se bonds. Specifically, a segment of the B<sub>2</sub>Se<sub>3</sub> five-membered ring formally opens up, establishing diselenide bridges to neighboring chains. Consequently, 2D polymeric [(B<sub>4</sub>Se<sub>13</sub>)<sup>4-</sup>]<sub>n</sub> layers are formed (Fig. 7). The B–Se bond lengths range from 2.032(7) to 2.080(7) Å. The Se–B–Se bond angles range from 94.4(5)° to 116.2(8)°. The Se–Se bond lengths range from 2.370(1) to 2.380(1) Å.

BaB<sub>2</sub>Se<sub>6</sub><sup>77</sup> crystallizes in the orthorhombic space group *Cmca* and is the first reported selenoborate containing the B<sub>2</sub>Se<sub>2</sub> four-membered rings. In the structure of BaB<sub>2</sub>Se<sub>6</sub>, four-membered B<sub>2</sub>Se<sub>2</sub> rings and B<sub>6</sub>Se<sub>10</sub> macrocycles form voids, and Ba<sup>2+</sup> cations are located in the voids (Fig. 8). The B–Se bond lengths range from 2.038(3) to 2.073(4) Å. The Se–B–Se bond angles range from 101.4(3)° to 115.4(3)°. The Se–Se bond length is 2.361(1) Å. Specifically, the bond angle (76.7(3)°) within the B<sub>2</sub>Se<sub>2</sub> four-membered rings is much smaller than that in the B<sub>2</sub>Se<sub>3</sub> five-membered rings (94° in [B<sub>2</sub>Se<sub>7</sub><sup>2-</sup>]<sub>n</sub>), and the B–Se bond length within the B<sub>2</sub>Se<sub>2</sub> four-membered rings (2.073(4) Å) is longer than that of the diselenide bridges (2.038(3) Å).

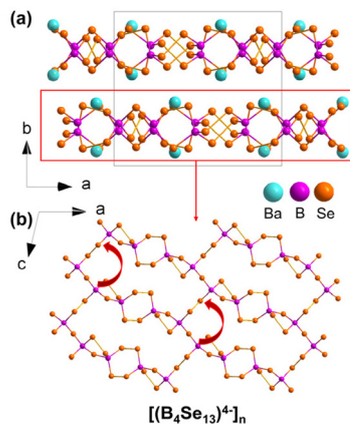


Fig. 7 The structures of (a) the Ba<sub>2</sub>B<sub>4</sub>Se<sub>13</sub> layer and (b) the [(B<sub>4</sub>Se<sub>13</sub>)<sup>4-</sup>]<sub>n</sub> layer.

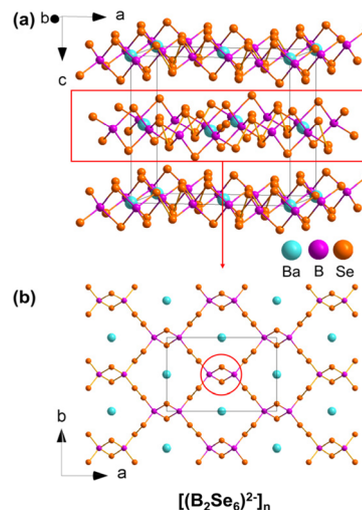


Fig. 8 The structures of (a) the BaB<sub>2</sub>Se<sub>6</sub> layer and (b) the [(B<sub>2</sub>Se<sub>6</sub>)<sup>2-</sup>]<sub>n</sub> layer.

**3.3.2. Selenoborates with 2D polymers constructed from supertetrahedral [B<sub>10</sub>Se<sub>20</sub>].** Li<sub>6-x</sub>Cs<sub>x</sub>B<sub>10</sub>Se<sub>18</sub> (*x* ≈ 1)<sup>78</sup> crystallizes in the tetragonal space group *I4<sub>1</sub>/amd* and is the only 2D selenoborate containing T3 supertetrahedral [B<sub>10</sub>Se<sub>20</sub>]. In its structure, the supertetrahedra [B<sub>10</sub>Se<sub>20</sub>] are corner-sharing to form layers (Fig. 9), between which are located the Li<sup>+</sup> and Cs<sup>+</sup> cations. The B–Se bond lengths range from 2.038(3) to 2.073(4) Å. The Se–B–Se bond angles range from 101.4(3)° to 115.4(3)°.

### 3.4. Three-dimensional selenoborates

In addition to the most common tetrahedral [BSe<sub>4</sub>] and supertetrahedral [B<sub>10</sub>Se<sub>20</sub>] units, the [B<sub>7</sub>Se<sub>13</sub>] unit represents a distinctive feature among 3D selenoborates, with no analogous configuration found in thioborates. The [B<sub>7</sub>Se<sub>13</sub>] unit is found in the zeolite-like compound Li<sub>7</sub>B<sub>7</sub>Se<sub>15</sub>. Fig. 10 illustrates the intricate constructional framework of B<sub>4</sub>Se<sub>13</sub> basic units. Originating from a central μ<sub>4</sub>-selenium atom, these units integrate with four [BSe<sub>4</sub>] tetrahedra in the initial shell. The interconnection of the tetrahedra in the second shell is facilitated by two [BSe<sub>4</sub>] entities from the first inner shell. Additionally, a bridging [BSe<sub>4</sub>] tetrahedron from this second outer shell establishes connectivity among the basic units within the anion network.

**3.4.1. Selenoborates with 3D polymeric networks constructed from [BSe<sub>4</sub>].** CuBSe<sub>2</sub><sup>79,80</sup> crystallizes in the tetragonal space group *I4̄2d* and is the only chalcopyrite-type semi-

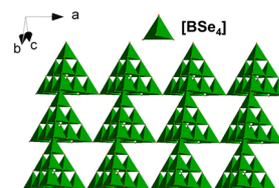
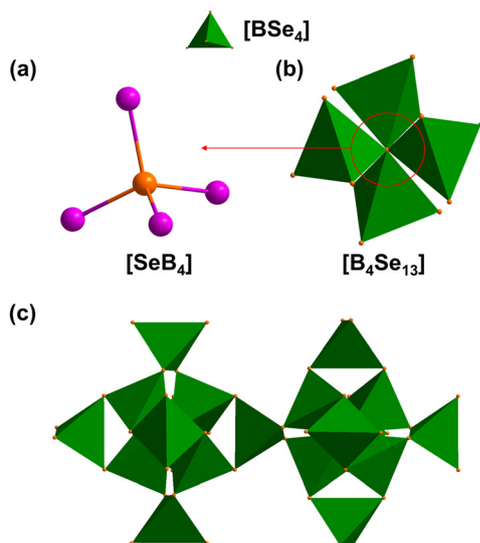
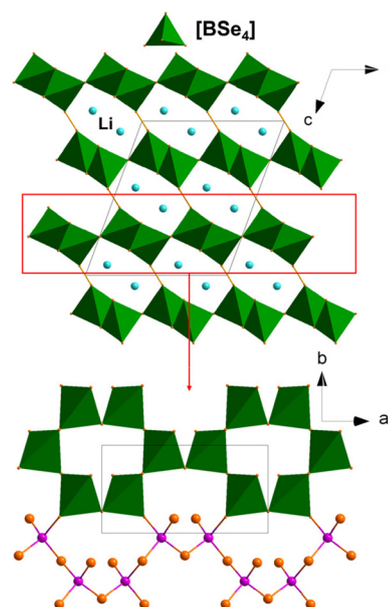


Fig. 9 The layer in the structure of Li<sub>6-x</sub>Cs<sub>x</sub>B<sub>10</sub>Se<sub>18</sub> (*x* ≈ 1).



**Fig. 10** Basic building block of the  $B_7Se_{13}^{5-}$  anion in  $Li_7B_7Se_{15}$ . (a)  $SeB_4$  tetrahedron forming (b)  $B_4Se_{13}$  basic units interconnected via additional BSe units to (c)  $B_7Se_{13}^{5-}$  entities forming the network.



**Fig. 11** The structure of  $Li_2B_2Se_5$ .

conductor among selenoborates. In its structure, there are  $[BSe_4]$  units connected by sharing corners to form a 3D network with  $Cu^{2+}$  cations distributed throughout. The band gap of nanocrystalline  $CuBSe_2$  has been studied, and the experimental band gap is 3.13 eV.

$Li_2B_2Se_5$ <sup>81</sup> crystallizes in the monoclinic space group  $C2/c$  and is the first perselenoborate with a 3D anionic network. In its structure, one  $[BSe_4]$  unit shares three vertex Se atoms with neighboring  $[BSe_4]$  units to form polymeric B–Se layers, which exhibit twelve-membered  $B_6Se_6$  rings. These layers are connected by Se–Se bonds to form a 3D network (Fig. 11). The average B–Se bond length is 2.061 Å.

**3.4.2. Selenoborates with 3D polymeric networks constructed from  $[B_7Se_{13}]$ .**  $Li_7B_7Se_{15}$ <sup>82</sup> crystallizes in the tetragonal space group  $P4_2/nbc$  and is a novel zeolite-like selenoborate. The structure of  $Li_7B_7Se_{15}$  comprises  $([B_7Se_{13}]^{5-})_n$ , diselenide  $Se_2^{2-}$  and  $Li^+$  cations. Alternatively, it could be described as an intercalation compound “ $Li_5B_7Se_{13}$ ” with additional diselenide “ $Li_2Se_2$ ”. The B–Se bond lengths range from 2.027(2) to 2.150(2) Å. The Se–B–Se bond angles range from 103.7° to 116.0°, which is consistent with the ideal tetrahedral angle (109.6°). The Se–Se bond length in “ $Li_2Se_2$ ” is 2.3643(5) Å.

**3.4.3. Selenoborates with 3D polymeric networks constructed from  $[B_{10}Se_{20}]$ .**  $Na_6B_{10}Se_{18}$ <sup>83</sup> crystallizes in the tetragonal space group  $I4_1/acd$ . In its structure, supertetrahedral  $[B_{10}Se_{20}]$  units are connected by sharing corners to form an anionic network and  $Na^+$  cations are located in the large voids between the anionic network to form a 3D network. The anionic network in  $Na_6B_{10}Se_{18}$  is composed of infinite polymeric anions of formula  $([B_{10}Se_{16}Se_{4/2}]^{6-})_{\infty}$ . The B–Se bond lengths range from 2.020(5) to 2.083(7) Å. The Se–B–Se bond angles range from 100.4(3)° to 114.0(4)°, which is nearly exactly the ideal tetrahedral angle (109.6°). Supertetrahedral

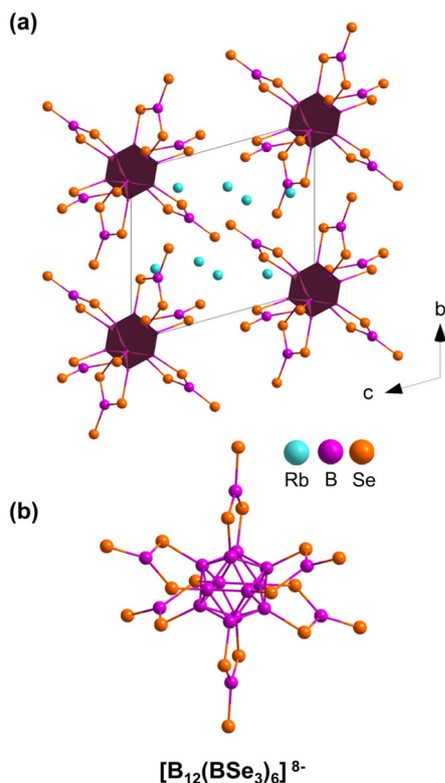
$[B_{10}Se_{20}]$  units can be found not only in 0D and 2D selenoborates, but also in selenoborates formed by a 3D polymeric network, which is special compared with the structure features of thioborates and borates.

$Li_{6+2x}[B_{10}Se_{18}]Se_x$  ( $x \approx 2$ )<sup>84</sup> crystallizes in the monoclinic space group  $C2/c$ . In its structure, corner-sharing supertetrahedral  $[B_{10}Se_{20}]$  units form a 3D anionic network, of which partially occupied  $Li^+$  and additional disordered  $Se^{2-}$  are located in the channels. The B–Se bond lengths range from 2.006(9) to 2.077(9) Å. The Se–B–Se bond angles range from 99.4(4)° to 116.3(4)°.

### 3.5. Selenoborate-closododecaborates

Advancements in boron chemistry have led to the discovery of selenoborate-closododecaborates as a distinct family within this field. Due to the lack of electrons in its outer shell, boron tends to form a framework that allows it to share electrons efficiently and achieve a more stable structure. The highly symmetrical  $B_{12}$  icosahedron is commonly found in many boron compounds because of its exceptional stability.  $Cs_8[B_{12}(BSe_3)_6]$ <sup>85</sup> and  $Rb_8[B_{12}(BSe_3)_6]$ <sup>86</sup> crystallize in the triclinic space group  $P\bar{1}$ , representing the first reported selenoborate-closododecaborates (Fig. 12). In their structures, the icosahedral  $B_{12}$  cluster is completely saturated with  $[BSe_3]$  units, resulting in a novel configuration within the selenoborate system.

$Rb_4Hg_2[B_{12}(BSe_3)_6]$  and  $Cs_4Hg_2[B_{12}(BSe_3)_6]$ <sup>86</sup> crystallize in the triclinic space group  $P\bar{1}$  and are the first transition metal selenoborates. In their structures, there is no direct linkage between isolated anion moieties. The B–Se bond lengths in them range from 1.915(11) to 2.006(8) Å. The B–B bond lengths in them range from 1.750(13) to 1.809(19) Å.



**Fig. 12** The structures of (a) the  $\text{Rb}_8[\text{B}_{12}(\text{BSe}_3)_6]$  and (b) the  $[\text{B}_{12}(\text{BSe}_3)_6]^{8-}$  unit.

$\text{Na}_6[\text{B}_{18}\text{Se}_{17}]^{87}$  crystallizes in the monoclinic space group  $C2/c$  and is the first boron selenium compound with a polymeric anionic cluster chain  $([\text{B}_{18}\text{Se}_{16}\text{Se}_{2/2}]^{6-})_{\infty}$ .

$\text{K}_8[\text{B}_{12}(\text{BSe}_3)_6]^{88}$  crystallizes in the monoclinic space group  $Cm$  and is the first selenoborate-*closo*-dodecaborate with three different anion substitution patterns.  $\text{Na}_8[\text{B}_{12}(\text{BSe}_3)_6]^{89}$  crystallizes in the monoclinic space group  $P2_1/c$ . There are isolated  $[\text{B}_{12}(\text{BSe}_3)_6]$  and  $\text{Na}^+$  cations in its structure.

$\text{Na}_2[\text{B}_{18}\text{Se}_{16}]^{90}$  crystallizes in the trigonal space group  $P\bar{3}$ . Formed by icosahedral  $\text{B}_{12}$  clusters and saturated  $[\text{BSe}_3]$  units, the cluster entities  $[\text{B}_{12}(\text{BSe}_3)_6]$  are connected to each other through Se–Se bonds, thus forming a 3D network with  $[(\text{B}_{18}\text{Se}_{12}\text{Se}_{6/2})_n]^{2-}$  and  $[(\text{B}_{18}\text{Se}_{15}\text{Se}_{6/2})_n]^{2-}$  anions in a ratio of 2 : 1.

## 4. Properties

For IR functional materials, the bandgap is one of the key performance indicators. Research on the properties of selenoborates remains relatively scarce, and only the bandgap of  $\text{CuBSe}_2$  has been estimated at 3.13 eV from its diffuse reflectance spectrum. To further explore the optical properties of the reported selenoborates, we calculated the bandgaps of selected alkali/alkaline earth metal selenoborates with ordered structures using the generalized gradient approximation (GGA) and the Heyd–Scuseria–Ernzerhof (HSE06) hybrid functional (Table 2).

**Table 2** The calculated bandgaps of selected selenoborates

Formula	Building unit	$E_g$ (GGA)	$E_g$ (HSE06)
$\text{Na}_2\text{B}_2\text{Se}_7^a$	$[\text{BSe}_4]$	1.67 eV	2.49 eV
$\text{K}_2\text{B}_2\text{Se}_7^a$	$[\text{BSe}_4]$	1.84 eV	2.73 eV
$\text{Ba}_7(\text{BSe}_3)_4\text{Se}$	$[\text{BSe}_3]$	2.20 eV	2.91 eV
$\text{CsBSe}_3^a$	$[\text{BSe}_4]$	1.84 eV	2.66 eV
$\text{RbBSe}_3^a$	$[\text{BSe}_4]$	1.68 eV	2.50 eV
$\text{Li}_2\text{B}_2\text{Se}_5^a$	$[\text{BSe}_4]$	1.72 eV	2.56 eV
$\text{Cs}_3\text{B}_3\text{Se}_{10}^a$	$[\text{BSe}_4]$	1.89 eV	2.75 eV
$\text{Rb}_2\text{B}_2\text{Se}_7^a$	$[\text{BSe}_4]$	1.93 eV	2.83 eV
$\text{BaB}_2\text{Se}_6^a$	$[\text{BSe}_4]$	1.78 eV	2.63 eV
$\text{Ba}_2\text{B}_4\text{Se}_{13}^a$	$[\text{BSe}_4]$	1.42 eV	2.18 eV

<sup>a</sup> The compounds with Se–Se bonds in the structures.

The majority of selected selenoborates exhibit HSE06 bandgaps exceeding 2.5 eV, thereby meeting the requirements for IR functional materials. Notably, the bandgap of the compound  $\text{Ba}_7(\text{BSe}_3)_4\text{Se}$ , which contains  $[\text{BSe}_3]$  as functional units, is larger than that of other compounds containing  $[\text{BSe}_4]$  units. Furthermore, among isomorphous selenoborates containing Se–Se bonds, those with heavier cations display larger bandgaps. This anomalous behavior may be attributed to the interaction between cations and Se–Se bonds.

In order to evaluate their potential as IR NLO materials, we calculated the SHG coefficients of  $\text{CuBSe}_2$  ( $I\bar{4}2m$ , NCS) and  $\text{TlBSe}_3$  ( $Cc$ , NCS). Specifically, the results show that  $\text{CuBSe}_2$  ( $d_{33} = -11.17 \text{ pm V}^{-1}$ ) and  $\text{TlBSe}_3$  ( $d_{13} = 35.39 \text{ pm V}^{-1}$ ) exhibit SHG properties of about  $0.8 \times \text{AGS}$  and  $2.6 \times \text{AGS}$ , respectively. This indicates the potential of selenoborates as IR NLO materials. Furthermore, their SHG properties may originate from the good arrangement of  $[\text{BSe}_3]$  units in their structures.

## 5. Summary and perspectives

In summary, this paper presents a comprehensive overview of the structures of reported selenoborates, along with their synthesis methods. The selenoborates are primarily based on two fundamental structural units:  $[\text{BSe}_3]$  and  $[\text{BSe}_4]$ , which, along with Se–Se bonds, form a variety of structural frameworks, including  $[\text{B}_{10}\text{Se}_{20}]$  supertetrahedra,  $[\text{BSe}_3^{3-}]_n$ ,  $[\text{B}_2\text{Se}_7^{2-}]_n$  and  $[\text{B}_3\text{Se}_{10}^{3-}]_n$  polymeric chains,  $[(\text{B}_4\text{Se}_{13})_n]^{4-}$  and  $[(\text{B}_2\text{Se}_6)_n]^{2-}$  layers and so on. Compared to borates and thioborates, selenoborates exhibit considerable potential for structural diversity, which remains largely unexplored and is worthy of further investigation.

Most selenoborates are synthesized *via* solid-state reactions involving metal selenides, amorphous boron, and selenium in evacuated carbon-coated silica tubes, typically at temperatures of 600–950 °C. Obtaining highly pure selenoborates with well-defined structures is challenging due to the reactivity of boron chalcogenides with container materials at elevated temperatures.

What is more, the analysis of microstructural performances of related units has shown that the  $[\text{BSe}_3]$  unit exhibits the

highest polarizability anisotropy and hyperpolarizability, highlighting its potential as a good seed for IR functional materials. The DFT calculations illustrated that selected alkali/alkaline earth metal selenoborates have large bandgaps and could meet the requirements for IR functional materials. Notably, reported NCS selenoborates exhibit significant SHG coefficients, highlighting the potential of selenoborates as IR NLO materials. Furthermore, selenoborates containing the [BSe<sub>3</sub>] unit may exhibit larger bandgaps. With targeted efforts in the following areas, further enhancements are anticipated.

(i) Current research on the properties of selenoborates is relatively sparse, highlighting the need for further investigation into their structure–property relationships. Notably, the common presence of Se–Se bonds within the selenoborate system has been discovered. While the effects of the Se–Se bonds on structures and properties remain not fully elucidated, these are worth exploring in the future.

(ii) The  $\pi$ -conjugated [BSe<sub>3</sub>] unit shows good microstructural performance with large polarizability anisotropy and hyperpolarizability. Considering thioborates such as NaBaBS<sub>3</sub>, LaBS<sub>3</sub>, NaSrBS<sub>3</sub> and so on that comprise planar triangular [BS<sub>3</sub>] as building units exhibit good properties, the exploration of more selenoborates containing [BSe<sub>3</sub>] units may help in the discovery of high-performance functional materials.

(iii) Structural analysis of selenoborates shows that none of the reported selenoborates are composed of mixed anionic groups [BSe<sub>3</sub>] and [BSe<sub>4</sub>]. It is known that thioborates composed of mixed anionic groups [BS<sub>3</sub>] and [BS<sub>4</sub>] all crystallize in NCS space groups. Therefore, synthesizing selenoborates with both [BSe<sub>3</sub>] and [BSe<sub>4</sub>] building units opens up vast possibilities for exploring novel NCS structures and enhancing the NLO properties of selenoborates.

(iv) The currently reported synthesis methods for selenoborates mainly use metal selenides as raw materials, which means high costs when growing crystals. The improvement of synthesis methods will bring new opportunities for the application of selenoborates.

(v) Among the 21 reported selenoborates, most are ternary compounds, predominantly featuring cations that are alkali or alkaline earth metals. In these structures, the arrangement of anions is primarily regulated by the presence of a single type of cation. Derived from this, the exploration of quaternary and multi-selenoborates warrants investigation. The incorporation of more diverse cations may provide more possibilities for regulating the structure, thereby enhancing the structural diversity of the selenoborate system.

## Data availability

All the data were extracted from cited articles.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

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