RSC Advances



PAPER

View Article Online



Cite this: RSC Adv., 2024, 14, 16886

Structural strategy for advancing nonlinear optical effects in 1D-[MX₂]_∞ chains: internal distortion and atomic types

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This work aimed to alleviate the limitations of existing mid-infrared nonlinear-optics (MIR-NLO) crystals by conducting theoretical research on $1D-[MX_2]_{\infty}$ (1D = one-dimensional; M = metallic element; X = anionicelement) structures in relation to NLO. An analysis was conducted on the electronic structure and optical properties of six selenides (BaZnGeSe₄, K_x Ba_{1-x/2}Ga₂Se₄, K_x Ba_{1-x/2}Ga_yIn_{2-y}Se₄, K_x Ba_{1-x/2}In₂Se₄, BaZnSiSe₄, $SrZnSiSe_4$) and three arsenides (Cs_2SnAs_2 , Rb_2SnAs_2 , K_2SnAs_2) using first principles. Afterwards, the inherent characteristics of these 1D-[MX₂]_∞ chains were further examined. The calculation results suggest that the presence of internal distortion (non-centrosymmetric) and high electronegativity in M elements can greatly enhance the NLO capability of $1D-[MX_2]_{\infty}$ structures. Furthermore, the first-ever prediction of K₂SnAs₂'s NLO ability has been made. Ultimately, the establishment of a theoretical structure (K₂BaSn₂As₄) provides guidance for the subsequent creation of high-performance MIR-NLO crystals.

Received 8th March 2024 Accepted 20th May 2024

DOI: 10.1039/d4ra01791a

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Introduction

Measurement and testing are essential prerequisites for the advancement of any nation, particularly in highly precise scientific domains. MIR lasers have exhibited significant luminosity in metrology domains, including chemical composition analysis, medical surgical procedures, and radar detection. 1-3 The quality of MIR-NLO crystals, which serve as the fundamental element for converting laser frequency, has a significant impact on the measurement accuracy of the entire laser system. The predominant commercially available MIR-NLO crystals on a large scale consist primarily of diamond-like structures or perovskite structures, such as AgGaS2 (AGS), AgGaSe2, GaAs, LiNbO₃, and so on.⁴⁻⁷ Advancements in synthesis technology and theory have led to the successful production of an increasing number of MIR-NLO crystals with distinctive characteristics, including InGa₂Cl₇, MoO₂Cl₂, NaMg₃Ga₃Se₈, KYGeS₄, and others.⁸⁻¹¹ Several new generation MIR-NLO crystals, including Li_{1-x}Ag_xGaSe₂, BaGe₄Se₇, La₃Nb_{0.5}Ga_{5.5}O₁₄, and others, have been progressively utilized in industrial production and practical applications. 12-14

Considering the increasing demand for high-performance MIR-NLO crystals, it is necessary to conduct first-principles research on materials. Alongside traditional research on electronic orbitals, recent years have witnessed the emergence of various sophisticated chemical computing software. 15 The rapid strides in computer science have propelled the development of high-throughput screening and material design, facilitating a more seamless integration of theoretical insights and experimental endeavors. Therefore, future material synthesis endeavors are poised to be more precise, leading to significant reductions in research time.

In recent years, a plethora of reports underscore the significance and triumphs of cutting-edge technology. Notably, advanced deep ultraviolet borate crystals have undergone meticulous screening using the Inorganic-Crystal-Structure-Database (ICSD).16 Moreover, a systematic theoretical analysis has delved into the intrinsic correlation between crystal structure and birefringence.17 Optical theoretical structure prediction stands out as the forefront of NLO advancements, primarily lauded for its efficacy in augmenting experimental success rates.18 A striking example is the recent success story of NaSrBS₃, the pioneer material predicted within the thioborate domain, thanks to refined theoretical guidance.19 In addition to the aforementioned borate/thioborate, this technology extends its reach into the realm of chalcogenides. Noteworthy theoretical inquiries have unfolded regarding the crystal structure and synthesis pathways of LiGaSe2, including theoretical predictions and successful preparation of MgGa₂Se₄.^{20,21} In addition, with the support of classical molecular dynamics, Density-Functional-Theory and other theories, significant breakthroughs have also been made in the field of organic luminescent molecules in recent times.22

In 2022, extensive theoretical research elucidated the crystal structure of 1D functional modules [BO₂]_∞ chains within the domain of deep ultraviolet NLO.23 These reports underscored

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the dynamic and thermodynamic stability of these theoretical crystals. Subsequently, in 2024, the successful synthesis of two high-quality NLO materials, NaLu(SeO₃)₂ and Cs₃In(In₄Se₇)(P₂-Se₆), based on theoretical designs, marked significant advancements.^{24,25} Notably, the prediction and realization of optical properties in nanoscale thin film materials further underscore the critical role of theoretical design in shaping the

future exploration of NLO materials.26

Beyond NLO, the integration of theoretical research into practical experimentation has reached a notable maturity, particularly evident in the medical domain.²⁷ In addition, the exploration of the superposition effect among two-dimensional layers, amalgamating theoretical analysis with experimental methodologies, has garnered widespread recognition, especially in the study of optics and topological structures.²⁸ Noteworthy is the successful prediction, through optical theory research, of the fluorescence capabilities of periodically stacked photonic crystal plates.²⁹ These reports underscore the paramount importance of theoretical predictions in various chemical fields.

Among these NLO systems, the MIR-NLO ability of the AII-M^{II}-M^{IV}-Ch (A^{II} = alkaline earth metal element; M^{II} = +2 valence metal element; $M^{IV} = +4$ valence metal element; Ch = S, Se, Te) system has recently been extensively investigated. Several optical materials with exceptional performance are currently being investigated. Sulfides like SrMgGeS₄, SrCdSiS₄, and $SrZnGeS_4$ exhibit extremely large band gap (E_{gap}) values and laser damage threshold (LDT).30-32 Selenides such as SrHgGeSe4, BaHgGeSe₄, and SrHgSnSe₄ demonstrate pronounced second harmonic generation (SHG) characteristics.33,34 The 1D-[MX₂]_∞ structure in this series of materials consistently achieves a favourable equilibrium between SHG and LDT. Both BaZnGeSe₄ and BaZnSiSe₄ exhibit wide E_{gap} values and possess strong SHG properties.35 In particular, the SrZnSiSe₄ possesses a 2 \times SHG and a 10 \times LDT for AGS.³⁶

Furthermore, alongside $A^{II}-M^{IV}-Ch$, a group of 1D-NLO materials containing $A^{I}-A^{II}-M^{III}-Se$ ($A^{I}=$ alkali metal element; $M^{III}=+3$ valence metal element) has been recently synthesized with great success. 37 The shared structural characteristic among $A^{II}-M^{IV}-Ch$ and $A^{I}-A^{II}-M^{III}-Se$ is the existence of $1D\cdot[MX_2]_{\infty}$ chains. 38 $A^{I}-A^{II}-M^{III}-Se$ contain a greater amount of alkaline ions compared to $A^{II}-M^{IV}-Ch$, making it a variant structure. The alteration in the structure has resulted in a rise in the A/M (A= alkali metal element and alkaline earth metal element) value. The precise methodology for calculating A/M has been elucidated in pertinent reports. 39 It is worth noting that among these 1D materials, $K_xBa_{1-x/2}Ga_2Se_4$ demonstrates excellent NLO ability, with a SHG of $1\times AGS$ and a LDT of $13.4\times AGS$. It is evident that $1D\cdot[MX_2]_{\infty}$ chains possess significant research value in the field of NLO. 40

The primary obstacle faced by current optical parametric oscillation (OPO) MIR lasers is the insufficient output energy of idle light, while attempts to enhance the energy of pump light often result in the fragmentation of MIR-NLO crystals.⁴¹ Furthermore, the majority of MIR lasers are constrained by phase matching in MIR-NLO crystals, thereby rendering them incapable of producing precise wavelengths of MIR light.⁴²

Hence, a superior MIR-NLO crystal typically necessitates the presence of several optical indicators, including robust nonlinear coefficients, high LDT, suitable phase matching, and sufficiently low absorption bands.⁴³

Non-centrosymmetric tetra-coordinated phosphorus (Pn; Pn = P, As) compounds are widely recognized as a highly soughtafter category of optical materials.44 This is primarily attributed to their exceptional upper limit in nonlinear coefficients. The ZnGeP2 crystal currently in use has a nonlinear coefficient of 70 pm V^{-1} , which is five times higher than that of AGS. Additionally, its LDT is about three times greater than AGS.45 The CdSiP₂ crystal is extensively utilized in medical devices for its capacity to produce intense MIR light at approximately 6200 nm.46 Furthermore, individuals have progressively uncovered a significant quantity of tetra-coordinated Pn-NLO materials that exhibit exceptional performance. Mg₂In₃Si₂P₇, MgSiP₂, and BaGe₂P₂ are examples of high-performance NLO materials that have garnered significant interest. 47-49 Ba₂Si₃P₆ exhibits high LDT and facilitates the growth of large single crystals in the context of 1D structure.50 The aforementioned diverse situations clearly showcase the exceptional characteristics of [MPn₄] in the realm of MIR-NLO.

Pn materials also contain an extensive variety of 1D-[MX₂]_∞ chains, including K2SiP2, Na2SiP2 and others. 51-53 Regrettably, the majority of these structures exhibit centrosymmetry and lack the capacity for second-order polarization. Consequently, it is critical to investigate the optical properties of [MPn₄] in MIR-NLO, specifically in 1D- $[MX_2]_{\infty}$ chains. This study commenced by conducting a systematic analysis of six existing 1D-[MSe₂]_∞ structures (BaZnGeSe₄, K_xBa_{1-x/2}Ga₂Se₄, K_xBa_{1-x/2}Ga_yIn_{2-y}Se₄, $K_xBa_{1-x/2}In_2Se_4$, $BaZnSiSe_4$, $SrZnSiSe_4$). The optical properties and electronic structure of the subject were examined by employing first principles. Furthermore, this study provided the first-ever prediction of the NLO capability of K2SnAs2 by analysing the electronic structures of three 1D-[SnAs₂]_∞ materials (Cs₂SnAs₂, Rb₂SnAs₂, K₂SnAs₂).^{54,55} This work systematically clarifies the diverse impacts of structural modifications on the optical properties of 1D-[MX₂]_∞ chains through the construction of a theoretical structure (K2BaSn2As4). This theoretical investigation has the potential to offer adequate structural theoretical backing for the fabrication of MIR-NLO crystals of superior quality.

2 Computational methods

The open-source computing software "Abinit" is well-known for its high computational accuracy and extensive functionality. The ground state Bloch wavefunctions utilized in this study were acquired via "Abinit", with the Perdew–Burke–Ernzerhof (PBE) functional serving as the pseudopotential. ^{56–59} PBE functional offers enhanced precision in delineating the wavefunction across the entirety of a crystal structure, particularly in capturing high-order derivatives of the wavefunction. The calculation of all ground state Bloch wavefunctions utilised the k-points of $4 \times 4 \times 4$ and the cut-off energy of 2200 eV. ^{60,61} The primary approach used to derive Partial Density of States (PDOS) values is via the Projector-Augmented-Wave (PAW)

method, which involves the individual calculation of atoms.⁶² The calculation of linear optics and NLO is performed utilizing the Independent-Particle-Approximation (IPA) theory.63 The number of conduction bands utilized in optical response calculations is typically two to three times greater than the number of valence bands. Other structures were evaluated using optical band gap (E_{opt}) values as scissors, whereas K₂SnAs₂ and K₂BaSn₂As₄ were evaluated using Heyd-Scuseria-Ernzerhof band gap ($E_{\rm HSE06}$) values as scissors.⁶⁴ All calculated structures have been optimized, and the stress tolerance has been set to 2.5×10^{-3} eV Å⁻¹.65,66</sup> The maximum number of iterations for all calculations is capped at 200. In the calculation of PDOS, the smear value is set to more than 10 times the default value. The structural optimization part mainly adopts the Broyden-Fletcher-Goldfarb-Shanno (BFGS) minimization method. In the process of calculating dipole hyperpolarizability and first dipole hyperpolarizability using "Gaussian", B3LYP functional and LanL2DZ basis set were adopted. In all calculation processes, the remaining calculation parameters are set to default values. 67-69 The calculated data is processed using recognized methods.

The calculation method for the effective nonlinear coefficient ($d_{\rm eff}$) is:⁷⁰

$$\left\langle \left(d_{\text{eff}}^{2\omega} \right)^{2} \right\rangle = \frac{19}{105} \sum_{a} \left(d_{aaa}^{2\omega} \right)^{2} + \frac{13}{105} \sum_{a \neq b} d_{aaa}^{2\omega} d_{abb}^{2\omega} + \frac{44}{105} \sum_{a \neq b} \left(d_{aab}^{2\omega} \right)^{2} + \frac{13}{105} \sum_{abc, \text{cyclic}} d_{aab}^{2\omega} d_{bcc}^{2\omega} + \frac{5}{7} \left(d_{abc}^{2\omega} \right)^{2}$$

$$(1)$$

The calculation method for the dipole hyperpolarizability difference ($\Delta \alpha$) is:⁷¹

significant NLO capabilities. Therefore, in the process of constructing theoretical crystals, it is advisable to have only one 1D- $[MX_2]_{\infty}$ chain wherever feasible. In other words, the simpler the structure of the non centrosymmetric theoretical crystal, the better. In addition, considering the excellent NLO performance of non-centrosymmetric Pn materials, it is necessary to use $[MPn_4]$. The above two points are the core ideas for constructing theoretical crystals, and additional attention should be paid to maintaining charge balance during this process.

Presently, centrosymmetric structures constitute the overwhelming majority of 1D-[MPn₂]_∞ structures uncovered (Fig. 1 and 2). To elucidate the relationship between the internal configuration and optical properties of 1D-[MPn₂]_∞ chains in their entirety, it is necessary to introduce a theoretical framework. In addition to conducting theoretical analyses of established 1D-[SnAs₂]_∞ structures, this work also develops a novel theoretical crystal structure. The artificial distortion within the 1D-[SnAs₂]_∞ chain primarily constitutes this theoretical structure. By introducing K \times 2, Ba \times 1, Sn \times 2, As \times 4 into a simple cube (P1) in order to achieve charge balance within the system. Next, establish the coordinates (0.500, 0.535, 0.535) and (0.000, 0.465, 0.465) for the two Sn atoms. Conduct structural optimization while accounting for variations in cell shape and the displacement of other ions, while disregarding symmetry (Fig. 2e and f). In conclusion, a theoretical auxiliary structure denoted as K₂BaSn₂As₄ (Cm) was acquired, representing the entire system with an absolute energy of around 3489.76 eV (Fig. 2d). Tables 1 and 2 present the lattice parameters and atomic coordinates pertaining to the theoretical structure.

4 Electronic structure

4.1 Band structure

A systematic analysis of the electronic structures of these 1D crystals was conducted utilizing first principles methods. Its low

$$\Delta \alpha = \frac{\sqrt{2}}{2} \times \sqrt{(\alpha_{xx} - \alpha_{yy})^2 + (\alpha_{xx} - \alpha_{zz})^2 + (\alpha_{yy} - \alpha_{zz})^2 + 6(\alpha_{xy}^2 + \alpha_{xz}^2 + \alpha_{yz}^2)}$$
 (2)

The calculation method for the total first dipole hyperpolarizability ($\beta_{\rm tot}$) is: 72

 $E_{\rm gap}$ value and direct band gap semiconductor nature are both inferred from the band structure of BaZnGeSe₄ (Fig. 3a). The band structure takes on an entirely new appearance as a result

$$\beta_{\text{tot}} = \sqrt{(\beta_{xxx} + \beta_{xxy} + \beta_{xyy})^2 + (\beta_{yyy} + \beta_{xxz} + \beta_{yyz})^2 + (\beta_{xzz} + \beta_{yzz} + \beta_{zzz})^2}$$
(3)

3 Theoretical structure

From the perspective of NLO, constructing a 1D- $[MX_2]_{\infty}$ chain with excellent NLO capability often requires a high degree of consistency in the $[MX_4]$ arrangement. In theory, such a structure would polarize the electron gas in one direction, producing

of the adjustment of A/M. The indirect band gap semi-conductors denoted as $K_xBa_{1-x/2}Ga_2Se_4$, $K_xBa_{1-x/2}Ga_yIn_{2-y}Se_4$, and $K_xBa_{1-x/2}In_2Se_4$ exhibit a decreasing order of $E_{\rm gap}$ values (Fig. 3b-d). In contrast, their $E_{\rm opt}$ and $E_{\rm gap}$ values exhibit a diametrically opposed trend, which is consistent with the findings of this article's calculations.

(a) BaZnGeSe₁ (b) K.Ba_{1-x-2}Ga₂Se₂ (c) K.Ba_{1-x-2}Ga₃In_{1-y}Se₄ (d) K.Ba_{1-x-2}In₂Se₄ A/M = 0.5Rotate A/M = 0.5 - 1.0

Fig. 1 (a)–(d) The crystal structures of BaZnGeSe₄, K_x Ba_{1-x/2}Ga₂Se₄, K_x Ba_{1-x/2}Ga_yIn_{2-y}Se₄, and K_x Ba_{1-x/2}In₂Se₄, as well as the arrangement of their 1D chains.

 Cs_2SnAs_2 and Rb_2SnAs_2 , which are both centrally symmetric 1D crystals, both have indirect band gap semiconductor characteristics (Fig. 4a and b). Although their structures are virtually identical, Rb_2SnAs_2 has a slightly lower $E_{\rm gap}$ value than Cs_2SnAs_2 . When the radius of alkali metals decreases, the intercrystal gaps become inadequate for the preservation of the space group of *Ibam*. At this juncture, the configuration of these 1D- $[SnAs_2]_{\infty}$ chains has undergone a transformation, as has the configuration of K_2SnAs_2 (C222). The atomic configuration in question exhibits characteristics reminiscent of a trigonal

system, thereby liberating it from the electronic confines imposed by indirect band gap. The calculation results indicate that the $E_{\rm gap}$ value of $K_2 SnAs_2$ is approximately 0.81 eV, which is higher than the $E_{\rm gap}$ values of $Rb_2 SnAs_2$ and $Cs_2 SnAs_2$ (Fig. 4c). The fundamental reason for such differences may come from the symmetry of the crystal, which changes the energy at point Z and causes Fermi level ($E_{\rm F}$) to regress to Γ Near the point.

 $\rm K_2BaSn_2As_4$ lacks an indirect band gap across the entirety of the energy space, with an estimated $E_{\rm gap}$ value of 0.80 eV (Fig. 4d). This is very close to the $E_{\rm gap}$ value of $\rm K_2SnAs_2$.

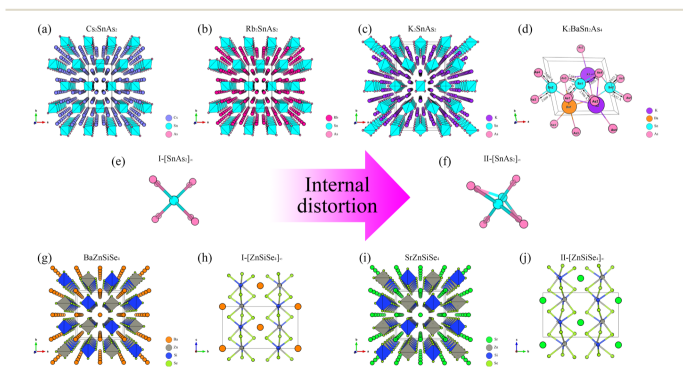


Fig. 2 (a)–(c) The crystal structures of Cs_2SnAs_2 , Rb_2SnAs_2 , and K_2SnAs_2 . (d) Theoretical structure of $K_2BaSn_2As_4$ and its bond length parameters. (e) and (f) The arrangement of two 1D–[$SnAs_2$] $_{\infty}$ chains results in internal distortion accompanied by changes in anions. (g)–(j) The crystal structures of $BaZnSiSe_4$ and $SrZnSiSe_4$, as well as their arrangement of 1D chains.

Table 1 The atomic coordinates of theoretical structure $K_2BaSn_2As_4$ (*Cm*). Its crystal parameters are: a=8.21387 Å, b=10.91161 Å, c=7.15652 Å, $\alpha=90.0000^\circ$, $\beta=106.2560^\circ$, $\gamma=90.0000^\circ$

Atom	x	у	z	
K	-0.06756	0.21776	0.67958	
Ba	0.09628	0.00000	0.21394	
Sn1	0.53500	0.00000	0.50000	
Sn2	0.46500	0.00000	0.00000	
As1	0.45759	0.18175	0.24359	
As2	0.24093	0.00000	0.65981	
As3	0.73448	0.00000	0.85842	

Table 2 The table lists the bond lengths of all Sn–As chemical bonds in the theoretical structure $K_2BaSn_2As_4$

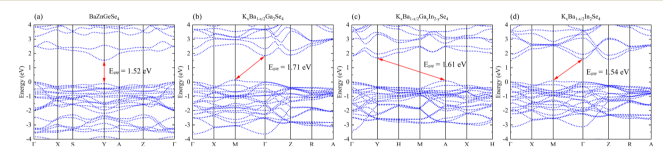
Chemical bond	Bond length		
Sn1-As1	2.65543 Å		
Sn1-As2	2.94772 Å		
Sn1-As3	2.62890 Å		
Sn2-As1	2.65237 Å		
Sn2-As2	2.60882 Å		
Sn3-As3	2.67987 Å		

Therefore, 1D chains may primarily regulate the $E_{\rm gap}$ value of this direct band gap structure. The $E_{\rm gap}$ value of BaZnSiSe₄ possessing a 1D-[MSe₂]_∞ structure is approximately 0.32 eV greater than that of SrZnSiSe₄ (Fig. 4e and f). Both substances have direct band gap semiconductor characteristics, with their $E_{\rm F}$ situated in close proximity to the Y point. The computed values of these band structures exhibit a high degree of congruence with the computed values documented in the scientific literature.

Research on 1D-[SnAs₂]_∞ and 1D-[ZnSiSe₄]_∞ indicates that in similar 1D-[MX₂]_∞ structures, the $E_{\rm gap}$ value typically decreases with a decrease in the $A^{\rm I}/A^{\rm II}$ ions radius. This may indicate that the stronger the electronegativity of $A^{\rm I}/A^{\rm II}$ ions, the higher the $E_{\rm gap}$ value of the material. In addition, combining the calculation results in Fig. 3 and 4, it can be found that the $E_{\rm gap}$ value of the 1D-[MX₂]_∞ structure is mostly dominated by the electronegativity difference between the M–X bonds, with the electronegativity potency of $A^{\rm I}/A^{\rm II}$ ions following closely behind.

4.2 Orbital structure

The purpose of this calculation was to perform a more comprehensive analysis of the electronic structures of these 1D structures. Zn-3d orbital electrons engage in valence band



 $\textbf{Fig. 3} \quad \textbf{(a)-(d) The band structures of BaZnGeSe}_4, \ \textbf{K}_x\textbf{Ba}_{1-x/2}\textbf{Ga}_z\textbf{Se}_4, \ \textbf{K}_x\textbf{Ba}_{1-x/2}\textbf{Ga}_y\textbf{In}_{2-y}\textbf{Se}_4, \ \textbf{and} \ \textbf{K}_x\textbf{Ba}_{1-x/2}\textbf{In}_2\textbf{Se}_4. \ \textbf{Se}_x\textbf{Se}_{1-x/2}\textbf{Ga}_y\textbf{In}_{2-y}\textbf{Se}_4, \ \textbf{Se}_x\textbf{Se}_{1-x/2}\textbf{In}_z\textbf{Se}_4. \ \textbf{Se}_x\textbf{Se}_{1-x/2}\textbf{Se}_{1-x/2}\textbf{Se}_{1-x/2}\textbf{Se}_{2-x/2}\textbf{Se$

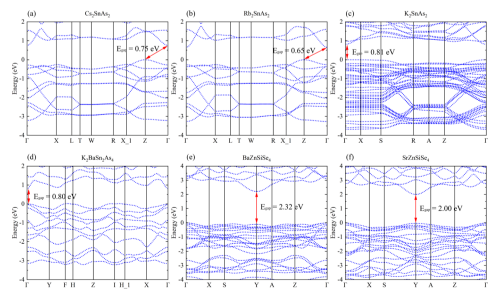
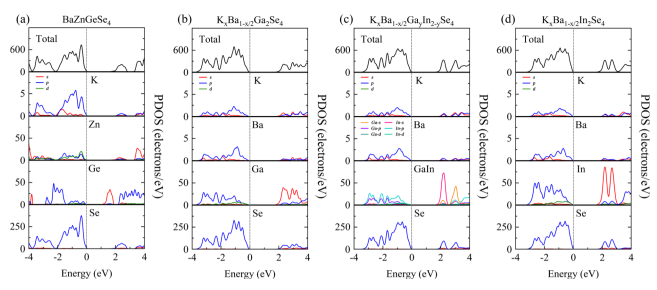


Fig. 4 (a) – (f) The band structures of Cs₂SnAs₂, Rb₂SnAs₂, K₂SnAs₂, K₂BaSn₂As₄, BaZnSiSe₄, and SrZnSiSe₄



(a)–(d) The PDOS of BaZnGeSe₄, $K_xBa_{1-x/2}Ga_2Se_4$, $K_xBa_{1-x/2}Ga_yIn_{2-y}Se_4$, and $K_xBa_{1-x/2}In_2Se_4$

bonding in close proximity to the E_F of BaZnGeSe₄, whereas Ge-4s orbital electrons are involved in conduction band bonding (Fig. 5a). From the PDOS of A^I-A^{II}-M^{III}-Se, some rules regarding the $E_{\rm gap}$ value can be found. As the participation of $M^{\rm III}$ -valence electrons increases near the valence band, the $E_{\rm gap}$ value gradually decreases. The degree of participation of MIII-valence electrons can also be explained by the binding ability of Se towards MIII-valence electrons (Fig. 5b-d). Considering that the electronegativity of In is weaker than that of Ga, the factor that affects the $E_{\rm gap}$ value is likely to come from the bond length.

The PDOS analysis reveals that the primary distinction between Cs₂SnAs₂ and Rb₂SnAs₂ lies in the Sn-As bond (Fig. 6a and b). The electron localization in the Sn-As bond of Cs₂SnAs₂ is marginally greater than that in the Sn-As bond of Rb₂SnAs₂. The disparity in E_{gap} values may primarily stem from the spatial separation between 1D-[SnAs2] chains. The degree of electronic localization of the Sn-As bond is significantly greater in K₂SnAs₂ compared to Cs₂SnAs₂ and Rb₂SnAs₂. The average Sn-As bond lengths of these three 1D-[SnAs₂]_∞ structures are 2.68 Å, 2.67 Å, and 2.62 Å, respectively. In these 1D structures, their bond lengths do not change in the order of E_{gap} values, so the correlation between E_{gap} values and bond lengths may not be significant.

Due to the fact that K₂BaSn₂As₄ and K₂SnAs₂ demonstrate an equivalent level of electronic localization, their E_{gap} values are comparable (Fig. 6c and d). To acquire more practical $E_{\rm gap}$ values, the HSE06 functional was employed to recalculate the $E_{\rm gap}$ values of both $K_2BaSn_2As_4$ and K_2SnAs_2 . The subsequent values of 1.77 eV and 1.49 eV for the calculated calibration band gaps may indicate that internal distortion positively influences the expansion of the $E_{\rm gap}$ value of the 1D-[SnAs₂]_{∞} chains. Particularly, the degree of Zn involvement near $E_{\rm F}$ distinguishes BaZnSiSe4 from SrZnSiSe4 with regard to the Zn-Se bond (Fig. 6e and f). In terms of bond length, the average Zn-Se bond lengths of these two structures are 2.48 Å and 2.53 Å, respectively, while their Si-Se average bond lengths are 2.28 Å and 2.32

Å, respectively. It can be seen that the compactness of 1D chains in BaZnSiSe₄ is significantly higher than that in SrZnSiSe₄.

Achieving a narrower E_{gap} value may be the consequence of decreasing the distance between 1D-[SnAs₂]_∞ without introducing internal distortion. Furthermore, a reduction in electronegativity between Ge and Se could potentially result in the development of narrow E_{gap} values. But in $A^{I}-A^{II}-M^{III}$ -Se, bond length also affects the bound state of the electron gas. This may indicate that the pattern of $E_{\rm gap}$ values varies when the type of anion changes. A slight increase in complexity may ensue when distortion is present within the 1D-[MX2]∞ chains. Internal distortion stretching substantially elongates the bond length of Sn-As in $1D-[SnAs_2]_{\infty}$ chains, an effect that substantially amplifies the E_{gap} values. However, for 1D-[ZnSiSe₄]_{∞} chains, $E_{\rm gap}$ values will be weakened as the bond length increases.

PDOS offers a lucid analysis of the contribution of each atomic orbital to each band, elucidating the optical characteristics of crystals primarily originating from various electronic transitions within the band spectrum. Therefore, by using PDOS, not only can we delve deeper into material E_{gap} , but we can also discern the contribution rates of individual functional groups to optical properties.

5 Optical properties

5.1 Linear optical properties

A systematic calculation of the linear optical properties of these eight structures was performed under the assumption of stable and convergent wavefunction conditions. The correlation between structure and linear optics is readily apparent, as the chemical environment of 1D-[ZnGeSe₄]∞ is essentially uniform along both the a- and b-axes (Fig. 7a). Phase matching occurs at a distance of approximately 520 nm to 1040 nm in BaZnGeSe4, and the disparity in refractive indices between the n_x and n_y directions can be minimal. Even under the same valence conditions, a change in atomic type can significantly alter the

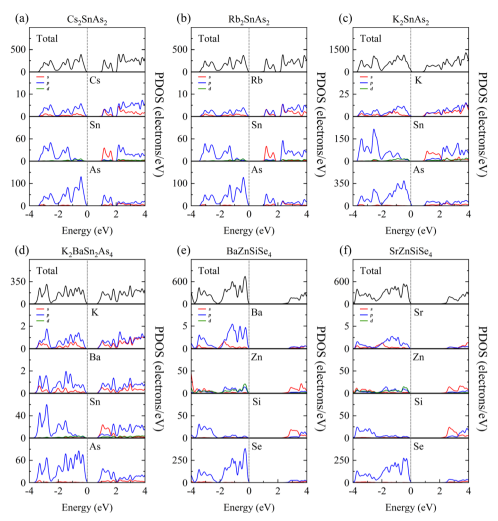


Fig. 6 (a)–(f) The PDOS of Cs₂SnAs₂, Rb₂SnAs₂, K₂SnAs₂, K₂BaSn₂As₄, BaZnSiSe₄, and SrZnSiSe₄.

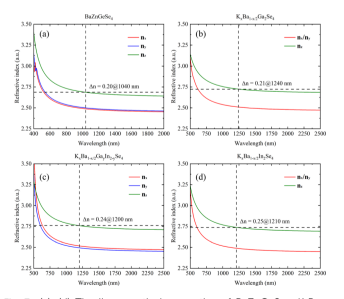


Fig. 7 (a)–(d) The linear optical properties of BaZnGeSe₄, $K_xBa_{1-x/2}Ga_2Se_4$, $K_xBa_{1-x/2}Ga_yIn_{2-y}Se_4$, and $K_xBa_{1-x/2}In_2Se_4$.

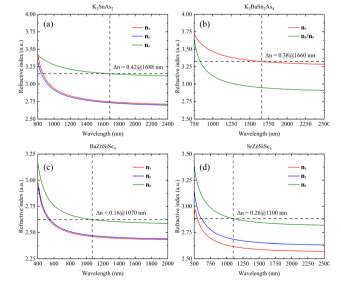


Fig. 8 (a)–(d) The linear optical properties of K_2SnAs_2 , $K_2BaSn_2As_4$, $BaZnSiSe_4$, and $SrZnSiSe_4$.

linear optical properties of the crystal. In the $A^{I}-A^{II}-M^{II}-Se$ system, the positions where these three materials generate phase matching are very close, but their Δn increases sequentially (Fig. 7b-d). This may be due to the larger atomic radius of the In atom, leading to an increased degree of charge shift.

The calculation results also indicate that K_2SnAs_2 has excellent birefringence and can achieve phase matching of $\Delta n = 0.42$ at 1688 nm. In contrast, the Δn of $K_2BaSn_2As_4$ at 1660 nm is about 0.38, which is slightly lower than the Δn of K_2SnAs_2 (Fig. 8a and b). The overall refractive index of $K_2BaSn_2As_4$ on the three axes is higher than K_2SnAs_2 , and their cut-off bands are positively correlated with the $E_{\rm gap}$ value. However, due to the generally low $E_{\rm gap}$ value of arsenides, their absorption cut-off band is larger than that of selenides.

The refractive index of BaZnSiSe $_4$ is significantly lower than that of 1D-[SnAs $_2$] $_\infty$ chains. BaZnSiSe $_4$ can achieve phase matching at 1070 nm, and its Δn is approximately 0.16. Although I-[ZnSiSe $_4$] $_\infty$ chains and I-[SnAs $_2$] $_\infty$ chains have similar configurations, there is a significant difference in their refractive indices (Fig. 8c). The influence of atomic type on refractive index is worth considering, especially for heavier atoms. SrZnSiSe $_4$ can achieve phase matching at 1100 nm, and the Δn between n_z and n_x reaches 0.26 (Fig. 8d). Considering the situation in 1D-[SnAs $_2$] $_\infty$ chains, the impact of internal distortion on the Δn of 1D-[MX $_2$] $_\infty$ crystals is different. To explain this phenomenon, a more detailed analysis of these 1D chains is needed.

5.2 Nonlinear optical properties

Systematically calculating the NLO properties of these structures is in addition to their linear optics properties. BaZnGeSe₄ has a $d_{\rm eff}$ of approximately 28.03 pm V⁻¹ (2050 nm), and its predominant second-order polarization is in the *xxz* direction (Fig. 9a). Particularly in the case of M disorder, the substitution

of elements with the same valence state has a significantly greater effect on second-order polarization than on first-order polarization. In $A^I-A^{II}-M^{III}-Se$, $K_xBa_{1-x/2}Ga_yIn_{2-y}Se_4$ has the maximum $d_{\rm eff}$, and its second-order polarization is mainly concentrated in the d_{33} direction (Fig. 9b–d). The relative difference between $K_xBa_{1-x/2}Ga_2Se_4$ and $K_xBa_{1-x/2}In_2Se_4$ is also worth noting, and the calculation results show that the former's $d_{\rm eff}$ is much greater than the latter's $d_{\rm eff}$. Whether this difference exists due to the $E_{\rm gap}$ value or the polarity of the 1D-[MSe₂] $_{\infty}$ chains themselves is unknown. Further elucidation of the aforementioned calculation outcomes requires a theoretical analysis on a more microscopic scale.

It has been possible to predict the NLO properties of K₂SnAs₂ for the first time, with a d_{eff} of approximately 1.76 pm V^{-1} (2050) nm). The primary cause of the low NLO in K₂SnAs₂ at the macroscopic level is the dearth of non-centrosymmetric components within the cell (Fig. 10a). Aside from the 1D-[SnAs₂]_∞ chains situated at the geometric center, which possesses non-centrosymmetric characteristics, the remaining components are centrosymmetric. In contrast to the intricate configuration of 1D-[SnAs₂]_∞ chains in K₂SnAs₂, the theoretical framework is considerably more straightforward. K2BaSn2As4 has a $d_{\rm eff}$ of up to 33.32 pm V⁻¹ (2050 nm) as a result of its noncentrosymmetric structure and strong consistent arrangement (Fig. 10b). Undoubtedly, significant distortion occurs not only in the relative positions of Sn⁴⁺ ions but also among As³⁻ ions. The anion theory posits that the predominant cause of nonlinear properties are these As³⁻ ions.⁷³

 $BaZnSiSe_4$ and $SrZnSiSe_4$ have the d_{eff} values 11.40 pm V^{-1} (2050 nm) and 26.01 pm V^{-1} (2050 nm), respectively (Fig. 10c and d). The close proximity between the calculated and experimental outcomes demonstrates that the parameter settings for the calculations are logical and convergent. Consequently, internal distortion improves the overall NLO characteristics of

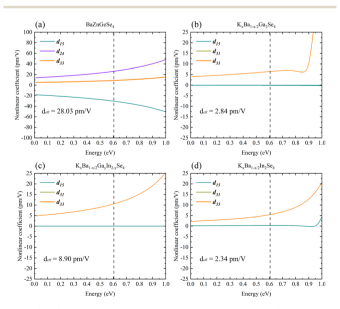


Fig. 9 (a)–(d) The nonlinear optical properties of BaZnGeSe₄, K_x Ba_{1-x/2}Ga₂Se₄, K_x Ba_{1-x/2}Ga_yIn_{2-y}Se₄, and K_x Ba_{1-x/2}In₂Se₄.

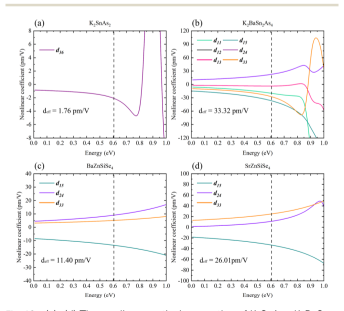


Fig. 10 (a)–(d) The nonlinear optical properties of K_2SnAs_2 , $K_2BaSn_2-As_4$, $BaZnSiSe_4$, and $SrZnSiSe_4$.

Table 3 The comparative summary table lists theoretical calculations in this study alongside experimental data. All experimental data on SHG are subject to variations due to size factors and powder effects, leading to relatively large fluctuations. The determined E_{gap} generally aligns with reported computational data. This discrepancy is primarily due to the intrinsic impact of discontinuities in the wavefunction, with E_{gap} being lower than E_{opt} , which is a normal occurrence

Crystal	This work			Experiment			
	$E_{\rm gap}$ (eV)	Δn	$d_{ m eff}~({ m pm}~{ m V}^{-1})$	E_{opt} (eV)	Δn	SHG	Ref.
BaZnGeSe ₄	1.52	0.20	28.03	2.46	_	<2.0 × AGS	35
$K_xBa_{1-x/2}Ga_2Se_4$	1.71	0.21	2.84	2.03	_	$< 0.9 \times AGS$	40
$K_x Ba_{1-x/2} Ga_y In_{2-y} Se_4$	1.61	0.24	8.90	2.34	_	$< 0.5 \times AGS$	37
$K_x Ba_{1-x/2} In_2 Se_4$	1.54	0.25	2.34	2.60	_	$< 0.5 \times AGS$	37
K ₂ SnAs ₂	0.81 (1.49)	0.42	1.76	_	_	_	55
K ₂ BaSn ₂ As ₄	0.80 (1.77)	0.38	33.32	_	_	_	_
BaZnSiSe ₄	2.32	0.16	11.40	2.71	_	$< 0.6 \times AGS$	35
SrZnSiSe ₄	2.00	0.26	26.01	1.95	_	$<$ 2.0 \times AGS	36

the crystal. Nonetheless, in light of the substantial disparity between the $E_{\rm gap}$ values of BaZnSiSe₄ and SrZnSiSe₄, additional investigation into the deep microscopic mechanism of 1D- $[{\rm MX}_2]_{\infty}$ chains is required (Table 3).

5.3 Inherent properties of $1D-[MX_2]_{\infty}$ chains

From a molecular standpoint, it is efficient to investigate the inherent characteristics of comparable 1D structures (characterized by identical elemental valence and configuration). The process commenced with the methodical calculation of the frontier molecular orbitals of the M_2X_6 molecules (Fig. 11). In accordance with the band structure, the $E_{\rm gap}$ values of molecules Ga_2Se_6 , $GaInSe_6$, and In_2Se_6 diminish in a sequential fashion. This result suggests that the variation in $E_{\rm gap}$ values is attributable to 1D- $[MX_2]_{\infty}$ chains as opposed to alkaline ions. Nevertheless, both calculation methods were unable to faithfully replicate the pattern of $E_{\rm opt}$ values; the same issue persists even in the reported calculation results. Moreover, the $E_{\rm gap}$ value of $ZnGeSe_6$ is considerably greater than that of Ga_2Se_6 , and this result of the calculation is more in line with the $E_{\rm opt}$ values.

In $\mathrm{Sn_2As_6}$, II- $\mathrm{Sn_2As_6}$ has a wider E_{gap} value than I- $\mathrm{Sn_2As_6}$. This corresponds to the calculated E_{HSE06} values, demonstrating the effect of internal distortion on expanding the E_{gap} values of 1D-[$\mathrm{SnAs_2}$] $_{\infty}$ chains. The calculation results also indicate that the E_{gap} value of II-ZnSiSe $_{6}$ is slightly lower than that of I-ZnSiSe $_{6}$. The energy difference between HOMO and LUMO is widened as a result of the contraction of bond length in ZnSiSe $_{6}$, this minute difference is greatly magnified in the crystal field. The band structure is consistent with the calculation results of ZnSiSe $_{6}$, suggesting that the majority of the variation in E_{gap} values originates from 1D-[ZnSiSe $_{4}$] $_{\infty}$ chains. The difference in E_{gap} values between I-ZnSiSe $_{6}$ and ZnGeSe $_{6}$ can be almost negligible, but their E_{gap} values are both higher than those of II-ZnSiSe $_{6}$.

In terms of optical properties, by using $\Delta \alpha$, it is possible to directly determine the polarity of similar molecules numerically (Fig. 12a). Corresponding to the refractive index, the $\Delta \alpha$ values of the three molecules Ga_2Se_6 , $GaInSe_6$, and In_2Se_6 increase sequentially. The $\Delta \alpha$ of $ZnGeSe_6$, I- $ZnSiSe_6$, and II- $ZnSiSe_6$ molecules also corresponds to their Δn (Fig. 12b). I- Sn_2As_6 has a visible central symmetry characteristic, resulting in a $\Delta \alpha$ of up to 61.17×10^{-24} esu. Benefit by the significant difference in

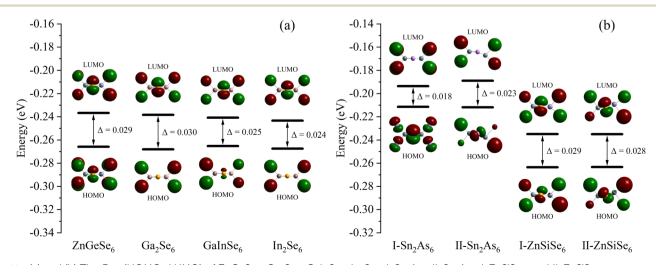


Fig. 11 (a) and (b) The $E_{\rm gap}$ (HOMO-LUMO) of ZnGeSe₆, Ga₂Se₆, GalnSe₆, In₂Se₆, I-Sn₂As₆, II-Sn₂As₆, I-ZnSiSe₆, and II-ZnSiSe₆.

Fig. 12 (a)–(d) The $\Delta \alpha$ and β_{tot} of ZnGeSe₆, Ga₂Se₆, GaInSe₆, In₂Se₆, I-Sn₂As₆, II-Sn₂As₆, I-ZnSiSe₆, and II-ZnSiSe₆.

position between Sn^{4+} , the central symmetry of II- $\mathrm{Sn}_2\mathrm{As}_6$ is much weaker. This structural change results in II- $\mathrm{Sn}_2\mathrm{As}_6$'s $\Delta\alpha$ (56.29 \times 10⁻²⁴ esu) being weaker than I- $\mathrm{Sn}_2\mathrm{As}_6$'s $\Delta\alpha$.

Although visually, II-ZnSiSe $_6$ has a more exaggerated degree of distortion than I-ZnSiSe $_6$. But in fact, in terms of the degree of central symmetry, II-ZnSiSe $_6$ is actually better than I-ZnSiSe $_6$. The $\Delta\alpha$ values for II-ZnSiSe $_6$ and I-ZnSiSe $_6$ are 43.94×10^{-24} esu and 42.88×10^{-24} esu, respectively. In the direction of these 1D chains, the length of II-ZnSiSe $_6$ should be greater than the length of I-ZnSiSe $_6$. In addition, the $\Delta\alpha$ of ZnGeSe $_6$ is also slightly smaller than that of I-ZnSiSe $_6$. These results all explain the underlying reason why SrZnSiSe $_4$ has a higher refractive index than BaZnSiSe $_4$, including ZnGeSe $_6$'s weak linear optical ability.

Additionally, the second-order optical properties of analogous molecules can be elucidated using $\beta_{\rm tot}$. It appears that $\rm M_2X_6$ molecules of identical elemental composition exhibit a delicate equilibrium between the dipole hyperpolarizability and the first dipole hyperpolarizability (Fig. 12c). It is noteworthy to mention that the $\beta_{\rm tot}$ of the dual element system is considerably lower than that of the multi-element system in $\rm M_2Se_6$. The $\beta_{\rm tot}$ of GaInSe $_6$ is several tens of times higher than the other two $\rm M_2Se_6$ (Ge $_2Se_6$ and In $_2Se_6$), although their molecular configurations are almost identical. But in terms of microstructure, the bond length of Ga–Se is smaller than that of In–Se. This causes the GaInSe $_6$ molecules to tilt to one side and produce good second-order polarization ability.

I-Sn₂As₆ has almost no second-order optical properties, while II-Sn₂As₆'s β_{tot} reaches an astonishing 11.55 \times 10⁻³⁰ esu (Fig. 12d). The same phenomenon also occurs in ZnSiSe₆, where I-ZnSiSe₆ has a higher β_{tot} than II-ZnSiSe₆. Although the difference in E_{gap} values between ZnGeSe₆ and I-ZnSiSe₆ is not significant, the former's β_{tot} is twice that of the latter. This also indicates the importance of highly electronegative M elements for second-order polarization. Pn materials' exceptional NLO

capability is, in fact, largely attributable to the electronegativity difference between M and Pn. As the binding ability of Pn ions to valence electrons (M) decreases, the material's $E_{\rm gap}$ value correspondingly decreases. This can also facilitate the influence of external electric fields on electrons within the crystal field. ${\rm Sn_2As_6}$ has a greater potential than ${\rm M_2Se_6}$ with regard to second-order optical properties.

However, a contradiction exists here. BaZnSiSe4 should possess a greater NLO capability than SrZnSiSe4, according to the results of the β_{tot} . In reality, SrZnSiSe₄ possesses an NLO ability that is far superior to BaZnSiSe4's. Without taking into account the scissor, the $d_{\rm eff}$ of BaZnSiSe₄ remains below 19.96 pm V⁻¹ (2050 nm). From the perspective of crystal field, the interaction between 1D chains is also a significant factor that cannot be ignored. In order to restore the environment of the crystal field, the β_{tot} values of [I-ZnSiSe₆] \times 4 and [II-ZnSiSe₆] \times 4 are additionally calculated. The pleasant surprise is the calculation results indicate that $\beta_{\rm tot}$ (31.96 \times 10⁻³⁰ esu) in [II- $ZnSiSe_6 \times 4$ far exceeds β_{tot} (1.19 × 10⁻³⁰ esu) in [I-ZnSiSe₆] × 4. The principal reflection of the component difference between the two is in the zzz direction, where the calculated data for the nonlinear coefficients are located. The aforementioned calculation outcomes methodically elucidate the intrinsic factors that contribute to the simultaneous existence of excellent Δn and outstanding $d_{\rm eff}$ in SrZnSiSe₄. This suggests that the configuration of 1D-[MX₂]_∞ chains significantly affects NLO, in addition to the second-order polarization capability of the functional groups themselves.

The contributions of orbitals in PDOS indicate that [ZnSe₄] group predominantly governs the optical properties near E_F , while [SiSe₄] and [GeSe₄] groups primarily control the optical properties of the intermediate bands. Additionally, the subtle optical properties at the lower levels are predominantly modulated by A^I/A^{II} ions. From the perspective of occupancy, [SiSe₄] and [GeSe₄] groups contribute significantly to the overall optical properties of the crystal and are the primary sites of charge transfer. Therefore, in regulating NLO capabilities, emphasis should be placed on the properties of high charge density groups and their arrangement.

6 Conclusions

To advance the NLO capability of 1D-[MX₂] $_{\infty}$ structures, this work suggests a number of efficient structural strategies. A decrease in the electronegativity difference (M–X) can, in general, cause the nonlinear coefficient to increase considerably. It is worth noting, nevertheless, that a low $E_{\rm gap}$ value can readily result in an excessively broad absorption cut-off band. A slight reduction in birefringence capability may occur as a result of the non-centrosymmetric distortion occurring within the 1D-[MX₂] $_{\infty}$ chains, which can substantially increase the material's nonlinear coefficient. Furthermore, it is noteworthy to consider the second-order polarization superposition effect that occurs between 1D-[MX₂] $_{\infty}$ chains. Finally, it is advantageous to incorporate [MPn₄] into 1D-[MX₂] $_{\infty}$ chains (while introducing internal distortions), particularly when the arrangement exhibits a strong tetrahedral consistency. In

summary, this theoretical investigation not only conducts a comprehensive examination of the optical properties and electronic configuration of 1D-[MX $_2$] $_\infty$ structures, but also offers structurally sound recommendations for the forthcoming fabrication of MIR-NLO optical crystals with exceptional performance.

Author contributions

Chaoyi Zhu: writing – original draft, conceptualization, data curation, methodology.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The author gratefully acknowledges financial support from the Shanghai Institute of Measurement and Testing Technology.

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