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Optoelectronic characteristics and stability evaluation of  $Ba_2TiMxO_6$  (Mx = Ge, Sn, Se, Te) p-type semiconductors as candidates for functional layers in optoelectronic devices

Driven by the escalating demand for direct band gap semiconductors, this study elucidates for the first time the potential of Ba<sub>2</sub>TiMxO<sub>6</sub> (Mx = Ge, Sn, Se, Te) perovskites in the realm of advanced photovoltaics and optoelectronics. The scope of this study spans the cutting-edge density functional theory computations of the structural, optoelectronic, and thermodynamic characteristics of specified materials. Additionally, it includes a comprehensive stability analysis that delves into their geometrical parameters  $(\tau$  and  $\mu$ ), cohesive, formation, and decomposition energies, and their temperature-dependent phonon spectra, which provide critical insights into the stability of probed materials. The analyzed materials are from a notable class of direct gap semiconductors, as determined within the PBE-GGA framework, showcasing band gaps of 1.915, 2.315, 1.963, and 1.433 eV at 300 K for the investigated Ge-, Sn-, Se- and Tebearing perovskites, respectively. Furthermore, band gap modulation with temperature is studied, denoting a stable optoelectronic behavior of these materials. Probing the optical characteristics reveals UV range optical spectra of absorption and extinction coefficients, dielectric constants, refractive indices, and optical band gaps. This study observed the enticing optoelectronic characteristics and robust stabilities of the materials studied, accentuating their potential as candidates for UV-based optoelectronic devices.

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### 1 Introduction

Perovskite materials with multifunctional facets have emerged with great potential for a myriad of applications, spanning the development of third-generation solar cells, sophisticated optoelectronic and thermoelectric devices, efficient high-temperature solid oxide cells, and photo-catalytic water splitting systems. 1-3 Besides, these versatile materials are driving innovation in designing and enhancing the efficiency of electromagnetic devices, such as next-generation light-emitting diodes, lasers, spintronic devices, optical sensors, and high-performance transistors.<sup>4-7</sup> The materials science community has extensively investigated single perovskite materials to explore their scope and transformative capabilities in diverse applications. Siddique et al. analyzed AeVH<sub>3</sub> (Ae = Be, Mg, Ca, Sr) halide perovskites using firstprinciples methods to reveal their suitability for advanced hydrogen storage systems,8 while Fang et al. leveraged Ti-based SrTiO<sub>3</sub> perovskites to develop a low cost and mass-scale producible optoelectronic synaptic device (In<sub>2</sub>O<sub>3</sub>·SnO<sub>2</sub>/Nb: SrTiO<sub>3</sub>(ITO/

NSTO)) capable of processing multimodal signals.9 Recently, double perovskites have received significant attention due to their superior stability and enhanced performance. Mustafa et al. revealed the potential of vacancy-ordered double perovskites Tl<sub>2</sub>WX<sub>6</sub> (X = Cl and Br) for spintronic and thermoelectric applications<sup>10</sup> and further analyzed  $K_2ScAuX_6$  (X = Cl, Br) double perovskites for next-generation optoelectronic devices. 11 These studies underscore that double perovskite materials exhibit outstanding structural, electronic, optical, and thermoelectric properties, such as higher conductivities and absorption coefficients, longer diffusion lengths, enhanced carrier mobilities, higher figure of merits, and tunable band gaps. Moreover, they can be produced easily using solution processing techniques.

In the realm of perovskite-based solar cells and optoelectronics, multiple studies have uncovered a class of barium-based double perovskite oxides (DPOs), Ba<sub>2</sub>BB'O<sub>6</sub>, with improved structural stability and desirable optoelectronic properties. In these DPOs, B and B' ions occupy the corner positions of the unit cell. These ions are octahedrally coordinated with six oxygen anions, forming BO6 and B'O6 octahedra, and A ions are located at the voids between octahedra and are coordinated with 12 oxygen anions. 12-15

Lately, titanium, with its appealing natural abundance, nontoxic nature, and stable tetravalent oxidation state, has been used in Ba<sub>2</sub>BB'O<sub>6</sub> DPOs alongside various elements having an

<sup>&</sup>lt;sup>a</sup> Department of Physics, University of Science and Technology of China, Hefei 230026, China. E-mail: ziadesk ustc@mail.ustc.edu.cn, zjlin@ustc.edu.cn

b Hefei National Laboratory, University of Science and Technology of China, Hefei

oxidation number of +4 at B or B' sites. 16,17 Having Zr at the Bsite, barium-titanium based Ba<sub>2</sub>ZrTiO<sub>6</sub> exhibits ultraviolet (UV) light absorption characteristics in experimental and theoretical studies of their electronic and optical properties. Its high absorption coefficient for UV radiation and figure of merit (ZT) indicate its suitability for optical sensors and thermoelectric devices. 18-20 In experimental studies by Tahir Ghrib et al., the optical and thermoelectric properties of cerium-doped Ba<sub>2</sub>Ti-MoO<sub>6</sub> double perovskite evidenced its potential in photocatalytic applications and a new generation of thermal barrier coatings. 21,22 Likewise, various studies underscore the potential of Ba<sub>2</sub>TiMnO<sub>6</sub> for use in solar cells and optoelectronics due to its high optical conductivity, absorption coefficient, and dielectric constant.23-25 Recently, employing TB-mBJ functionals, Shah et al. explored the optoelectronic prospects of Ba<sub>2</sub>HfTiO<sub>6</sub>, Ba<sub>2</sub>Ce-TiO<sub>6</sub>, and Ba<sub>2</sub>TiTeO<sub>6</sub> DPOs that exhibit band gaps of 3.44, 2.96, and 1.26 eV, respectively. As it has a band gap energy close to the red edge of the visible spectrum, Ba<sub>2</sub>TiTeO<sub>6</sub> exhibits absorption and optical conductivity peaks in the visible region. The figure of merit values of these three oxides are 0.79, 0.75, and 0.85, respectively. The author deduced that the higher visible absorption and ZT value make Ba<sub>2</sub>TiTeO<sub>6</sub> more suitable for solar cells.<sup>26</sup> These investigations demonstrate that barium-titanium DPOs have great potential in various photovoltaics, photo-catalytic, thermoelectric, and vis-UV optoelectronic applications.

Inspired by these studies and pursuing lead-free, inorganic double perovskites characterized by dynamic stability and compelling optoelectronic properties, this study explores the potential of novel Ba<sub>2</sub>TiMxO<sub>6</sub> (Mx = Ge, Sn, Se, Te) DPOs in solar cells and optoelectronic applications. It presents the innovative concept of integrating titanium at the B-site alongside p-block tin (Sn) metal, germanium (Ge), and tellurium (Te) metalloids and selenium (Se) non-metal at the B' site of barium-based DPOs. This deliberate incorporation of metal, metalloids, and non-metal at the B' site allows us to examine the changes in optoelectronic properties contingent upon B' elements belonging to the same block, different groups, and different periods. Likewise, incorporating titanium at the B sites is strategically adopted to harness its 3d electronic states to enhance the conduction band states. Given its natural abundance and potential to support a transformative approach to cost-effective technology, this integration presents an innovative solution, paving the way for sustainable advancements in the field.

This study meticulously analyzes the structural, electronic, and optical properties at temperatures of 0 and 300 K, and thermodynamic properties in the pressure range of 0 to 20 GPa and temperature range of 0 to 700 K. Additionally, the dynamic stability of the considered DPOs is examined through geometric factors, structural energetics, and temperature-dependent phonon spectra. Moreover, the temperature-induced narrowing and widening of band gaps are elucidated to analyze these perovskites' optoelectronic behavior under temperature effects. Prior research about the selected DPOs has not involved a similar investigation, except for a recent theoretical study of Ba<sub>2</sub>Te-TiO<sub>6</sub>, <sup>26</sup> which made a comparison of its structural, electronic, and optical properties. In this study, a comparison is provided of the previously reported experimental or theoretical findings for the resembled double perovskites. This study underscores the scope of the selected DPOs and serves as an outline for experimentalists and theorists to fabricate these novel materials and investigate their efficiencies further.

# 2 Methodology

This investigation employed the Vienna Ab Initio Simulation Package (VASP) to perform first-principles calculations. 27-29 To address ion-electron interactions, projector-augmented wave potentials are used, and to incorporate the exchange-correlation effects, Perdew-Burke-Ernzerhof (PBE) formalized generalized gradient approximation (GGA)<sup>30</sup> is applied. However, the Heyd-Scuseria-Ernzerhof (HSE06) hybrid functional, 31 which incorporates the Hartree-Fock exchange, is also utilized to acquire another theoretical estimation of the electronic band structure for comparative purposes. Shape, volume, and ionic positions are relaxed until the static energies converge up to  $1 \times 10^{-7}$  eV, and the maximum Hellmann forces on each atom are less than 0.001 eV  ${\rm \mathring{A}}^{-1}$ . A 6  $\times$  6  $\times$  6 Gamma scheme sampled k-point grid and a plane-wave basis with a cutoff energy value of 450 eV as determined by a convergence test are set for the energy calculations. The k-point sampling is performed using the Gaussian smearing method with a smearing width of 0.02 eV. However, for the density of states, a fine k-point grid of 12  $\times$  12  $\times$ 12 is used, and the partial occupancies are treated with the tetrahedron method along with the Blöchl corrections.

The employed method is borrowed from previous research demonstrating that density functional theory (DFT) is a powerful approach for accurately predicting material properties. Recently, multiple studies have shown the usefulness of the PBE-GGA approach for obtaining insights into the diverse facets of materials. Hasan et al. have published impactful research elucidating the efficacy of DFT calculations within the framework of the PBE-GGA method. The authors conducted a comprehensive investigation of the pressure-varied optoelectronic properties of  $ASnX_3$  (A = Li, Na, X = Cl, Br, I) halide perovskites, portraying the competitive structural properties, band structures, density of states, and optical spectra under 0, 1, 2, 3, 4, and 5 GPa pressures for every studied material.<sup>32</sup> Sun et al. demonstrated the suitability of the PBE-GGA method by scrutinizing band gaps of GeS, GeSe, SnS, and SnSe semiconductors. The authors underscore that the employed method resulted in band gaps without overestimation that closely match the experimental values.33 Recent investigations of double perovskite oxides have also utilized the PBE-GGA exchange-correlation functional. The optoelectronic properties of La<sub>2</sub>FeWO<sub>6</sub> and A<sub>2</sub>NbTbO<sub>6</sub> (A = Ca, Sr, Ba) DPOs have been analyzed using the PBE-GGA method to determine their potential for use in solar cells and optoelectronics.34,35 In the same regard, Mera et al. examined Sr<sub>2</sub>MgWO<sub>6</sub> and Ba<sub>2</sub>BiVO<sub>6</sub> DPOs for their photocatalytic applications, employing the PBE-GGA method.<sup>36</sup> Karlicky et al. compared the experimental band gap of fluorographene (3 eV) with the results of various functionals. They found that the PBE-GGA result (3.1 eV) closely aligns with experimental values compared to the HSE06 (5.1),  $G_0W_0$  (7.3), and BSE- $G_0W_0$  (5.4) results, highlighting its reliability and effectiveness for accurate predictions in material science.<sup>37</sup> These studies and others from the literature 38-40 underline the reliability of DFT calculations using the GGA-PBE functional, establishing a solid basis for our methodology.

The optical properties of the studied DPOs are computed from the optical response theory as implemented in VASP, which illustrates that the frequency-dependent complex dielectric function  $\varepsilon(\omega)$  has a pivotal role in characterizing the interaction of a material with incident photons. <sup>41,42</sup> The  $\varepsilon(\omega)$ consists of a real part  $\varepsilon_r(\omega)$  and an imaginary part  $\varepsilon_i(\omega)$ , as eqn (1) describes.43

$$\varepsilon(\omega) = \varepsilon_{\rm r}(\omega) + i\varepsilon_{\rm i}(\omega) \tag{1}$$

Here  $\omega$  is the photon frequency. The more significant part is the imaginary part of the dielectric function that represents the permissible intra- and inter-band optical transitions from the valence to conduction band.  $^{43,44}$  Mathematically, a 3  $\times$  3 Cartesian optical tensor of  $\varepsilon_i(\omega)$  can be written as, <sup>45</sup>

$$\varepsilon_{i}(\omega) = \frac{4\pi^{2}e^{2}}{\Omega} \lim_{q \to 0} \frac{1}{q^{2}} \sum_{c,v,k} 2w_{k} \delta\left(\varepsilon_{c_{k}} - \varepsilon_{v_{k}} - \omega\right) \times \left\langle u_{ck+e_{x}q} \middle| u_{vk} \right\rangle \left\langle u_{vk} \middle| u_{ck+e_{\beta}q} \right\rangle,$$
(2)

where  $\Omega$  is the primitive cell volume,  $w_k$  are the k-point weights whose sum is equal to 1,  $e_{\alpha}$  and  $e_{\beta}$  are the unit vectors for  $\alpha$  and  $\beta$  bands, respectively,  $c_k$  and  $v_k$  are the conduction and valence bands states from the k-space of the Brillouin zone, and  $u_{ck}$  and  $u_{\nu k}$  are the cell periodic functions of the conduction and valence band states. At any k-point, c runs over the conduction band empty states, and  $\nu$  runs over the valence band occupied states. This tensor becomes a scaler for the cubic symmetry of the structure as the selected structures exhibit. The real part of the dielectric function can be obtained from the imaginary part by utilizing the Kramers-Kronig transformation, 43,45 described as follows.

$$\varepsilon_{\rm r}(\omega) = 1 + \frac{2P}{\pi} \int_0^\infty \frac{\omega' \varepsilon_i(\omega')}{\omega'^2 - \omega^2} d\omega'$$
 (3)

Here, P stands for the principal value. The absorption coefficient  $\alpha(\omega)$ , extinction coefficient  $k(\omega)$ , refractive index  $n(\omega)$ , reflectivity  $R(\omega)$ , and the energy-loss function  $L(\omega)$  are derived from the dielectric functions, as illustrated by the expressions below.45-47

$$\alpha(\omega) = \frac{\sqrt{2\omega}}{c} \left[ \sqrt{\varepsilon_{\rm r}^2(\omega) + \varepsilon_{\rm i}^2(\omega)} - \varepsilon_{\rm r}(\omega) \right]^{1/2} \tag{4}$$

$$k(\omega) = \frac{1}{\sqrt{2}} \left[ \sqrt{\varepsilon_{\rm r}^2(\omega) + \varepsilon_{\rm i}^2(\omega)} - \varepsilon_{\rm r}(\omega) \right]^{1/2}$$
 (5)

$$n(\omega) = \frac{1}{\sqrt{2}} \left[ \sqrt{\varepsilon_{\rm r}^2(\omega) + \varepsilon_{\rm i}^2(\omega)} + \varepsilon_{\rm r}(\omega) \right]^{1/2}$$
 (6)

$$R(\omega) = \frac{(n(\omega) - 1)^2 + k^2(\omega)}{(n(\omega) + 1)^2 + k^2(\omega)}$$
 (7)

$$L(\omega) = \frac{\varepsilon_i(\omega)}{\varepsilon_r^2(\omega) + \varepsilon_i^2(\omega)}$$
 (8)

Quasi-harmonic Debye model (QHDM)<sup>48</sup> is employed to compute the pressure- and temperature-dependent thermodynamic properties. The predictability of this model is precise and dependable when determining thermodynamic properties, particularly at low temperatures. 49 In this model, the vibrational Helmholtz free energy  $(F_{vib})$  can be computed from the following ean (9).48

$$F_{\text{vib}}(T,\theta_{\text{D}}) = 3nk_{\text{B}}T\left(\frac{3\theta_{\text{D}}}{8T} + \ln\left(1 - e^{-\theta_{\text{D}}/T}\right) - \frac{D(\theta_{\text{D}}/T)}{3}\right) \tag{9}$$

where n denotes the number of atoms in the formula unit, and  $k_{\rm B}$  represents Boltzmann's constant.  $\theta_{\rm D}$  is the Debye temperature, described as a function of volume in eqn (11). <sup>50</sup> The Debye integral, D, is calculated from eqn (10).50,51

$$D(T_{\rm o}) = \frac{3}{T_{\rm o}^3} \int_0^{T_{\rm o}} \frac{s^3}{e^s - 1} \mathrm{d}s \tag{10}$$

where  $T_0 = \frac{\theta_D}{T}$ . Debye's theory views vibrations in a solid as elastic waves, and the Debye temperature of these waves depends on their average sound velocity. The Debye model describes the Debye temperatures of the isotropic solids as given in eqn (11),50

$$\theta_{\rm D}(V) = \frac{\hbar}{k_{\rm B}} (6n\pi^2 V^{0.5})^{1/3} f(\sigma) \left( \sqrt{\frac{B_{\rm st}(V)}{M}} \right)$$
 (11)

where M represents the molecular mass per formula unit and  $\sigma$ is the Poisson ratio. The static bulk modulus  $B_{st}(V)$  and the function  $f(\sigma)$  are determined using eqn (12) and (13).<sup>48</sup>

$$B_{\rm st}(V) = V\left(\frac{\mathrm{d}^2 E_{\rm st}(V)}{\mathrm{d}V^2}\right) \tag{12}$$

$$f(\sigma) = \left[ 3 \left\{ 2 \left( \frac{2(1+\sigma)}{3(1-2\sigma)} \right)^{3/2} + \left( \frac{1+\sigma}{3(1-\sigma)} \right)^{3/2} \right\}^{-1} \right]^{1/3}$$
 (13)

A commonly used value of the Poisson ratio for Cauchy solids, 48 equal to 0.25, is adopted in this calculation.

The Debye temperatures at the equilibrium volumes for given temperatures and pressures are used to compute other thermodynamic properties by employing eqn (14)-(18) respectively.48,50

$$C_{\rm v} = 3nk_{\rm B} \left( 4D(\theta_{\rm D}/T) - \frac{3\theta_{\rm D}/T}{{\rm e}^{\theta_{\rm D}/T} - 1} \right) \tag{14}$$

$$\gamma = -\frac{\mathrm{d}\ln\theta_{\mathrm{D}}(V)}{\mathrm{d}\ln V} \tag{15}$$

$$\alpha = \frac{\gamma C_{\rm v}}{B_T V} \tag{16}$$

$$C_{\rm p} = C_{\rm v} (1 + \alpha \gamma T) \tag{17}$$

$$B_T = -V \left(\frac{\mathrm{d}P}{\mathrm{d}V}\right)_T \tag{18}$$

### 3 Results and discussion

### 3.1 Structural properties

In the cubic ( $Fm\bar{3}m$ ) crystal structures of  $Ba_2TiMxO_6$  (Mx = Ge, Sn, Se, Te) DPOs, consisting of 40 atoms per unit cell, as depicted in Fig. 1, the  $Ba^{+2}$ ,  $Ti^{+4}$ ,  $Mx^{+4}$ , and  $O^{-2}$  ions are located at 8c (0.25, 0.25, 0.25), 4a (0, 0, 0), 4b (0.5, 0.5, 0.5), and 24e (x, 0, 0) Wyckoff positions, respectively. Depending on the ionic radii of  $Mx^{+4}$  ions, the value of x varies, and is 0.748, 0.756, 0.755, and 0.761 for the optimized structures of the Ge, Sn, Se, and Te bearing DPOs, respectively. Table 1 lists the interatomic bond lengths between  $Mx^{+4}$ ,  $Ba^{+2}$ , and  $Ti^{+4}$  cations and  $O^{-2}$  anions at both 0 and 300 K. It shows that the Ba–O bond length is greater than that of the Mx-O and Ti-O bonds in these structures, which is due to the lower electronegativity and lower affinity of  $Ba^{+2}$  for the  $O^{-2}$  anion compared to  $Ti^{+4}$  and  $Mx^{+4}$  cations. This implies the presence of large BO6 octahedra in these DPOs, which are essential for perovskite structures.

All the calculated bond lengths of the studied structures are higher at 300 K than at 0 K. These bond lengths closely match those of Ba<sub>2</sub>ZrTiO<sub>6</sub> and Ba<sub>2</sub>TiMnO<sub>6</sub> structures calculated by Naseri *et al.*, and Deluque-Toro *et al.*<sup>20,23</sup> Bramnik *et al.* reported the experimental bond lengths of Ba<sub>2</sub>MgReO<sub>6</sub>, demonstrating that the Ba–O bond length is greater than the others.<sup>52</sup> The fourth-order Birch–Murnaghan's equation of state,<sup>53</sup> as

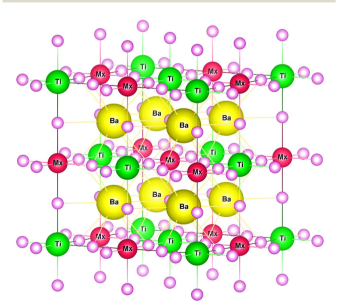
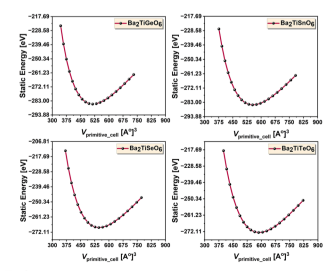


Fig. 1  $\,$  3D unit cell structure of crystalline  $\rm Ba_2TiMxO_6$  (Mx = Ge, Sn, Se, Te) compounds.

**Table 1** Bond lengths between  $Mx^{+4}$  cations and  $O^{-2}$  anions in  $Ba_2TiMxO_6$  (Mx = Ge, Sn, Se, Te)

		Bond lengt	Bond lengths [Å]		
DPOs	Bonds	0 K	300 K		
Ba <sub>2</sub> TiGeO <sub>6</sub>	Ge-O	1.982	1.994		
	Ва-О	2.832	2.848		
	Ti-O	2.023	2.034		
Ba <sub>2</sub> TiSnO <sub>6</sub>	Sn-O	2.104	2.117		
	Ва-О	2.906	2.923		
	Ti-O	2.004	2.016		
Ba <sub>2</sub> TiSeO <sub>6</sub>	Se-O	2.081	2.094		
2 0	Ва-О	2.891	2.909		
	Ti-O	2.007	2.019		
Ba <sub>2</sub> TiTeO <sub>6</sub>	Те-О	2.181	2.196		
	Ва-О	2.959	2.979		
	Ti-O	2.002	2.015		
Ba <sub>2</sub> ZrTiO <sub>6</sub> <sup>20</sup>	Zr-O	2.14			
2 0	Ti-O	1.99			
$Ba_2TiMnO_6^{23}$	Mn-O	1.903			
2 0	Ва-О	2.735			
	Ti-O	1.965			
Ba <sub>2</sub> MgReO <sub>6</sub> expt., 52	Mg-O	2.08			
2 0 0	Ва-О	2.85			
	Re-O	1.96			

described in eqn (19), is utilized to fit the energy-volume (E-V) data of Ba<sub>2</sub>TiMxO<sub>6</sub> structures. Fig. 2 displays the fitted E-V curves. The fitting yields the equilibrium volumes ( $V_o$ ), corresponding equilibrium energies ( $E_o$ ), bulk moduli (B), and pressure derivatives of bulk moduli (B'). Using the values of  $V_o$  and B along with other constants in eqn (11), the Debye temperatures of these structures are determined. Table 2 presents the static and equilibrium structural properties per unit cell of the selected DPOs. The higher bulk moduli of 160.41, 151.55, 145.55, and 141.02 GPa and higher Debye temperatures of 549.59, 517.50, 525.67, and 499.79 K for the Ge-, Sn-, Se-, and Te-bearing structures, respectively, indicate the hardness of these materials. Table 2 also portrays that our results are comparable to the theoretical and experimental findings for



**Fig. 2** Energy–volume curves – visualizing the structural optimization of the analyzed compounds.

similar DPOs.

$$E(V) = E_o + \frac{3}{8} V_o B_o f^2 \left[ (9H - 63B'_o + 143)f^2 + 12(B'_o - 4)f + 12 \right]$$

$$H = B_o B''_o + (B'_o)^2, \quad f = \frac{1}{2} \left[ \left( \frac{V}{V_o} \right)^{-2/3} - 1 \right]$$
(19)

### 3.2 Electronic properties

Utilizing the optimized crystallographic configurations of the examined materials, the electronic properties are determined at 0 K and ambient 300 K temperatures. To understand these characteristics, the Fermi energy level is set to 0 eV in the results displayed. The band structures are computed along the high symmetry points of the k-path in the first Brillouin zone of the reciprocal lattice as shown in Fig. 3. These band gap penned results unveil the direct band gap semiconductor nature of the studied DPOs, displaying that the band gaps in the Ge- and Sn-containing structures are located at the  $\Gamma$  high symmetry point, and in the Se- and Te-bearing structures, these are positioned at the L high symmetry point. As expected, the HSE06 method yields higher band gap values than PBE-GGA, consistent with its incorporation of exact exchange. Although the HSE06-calculated band gaps are higher than PBE-GGA ones - the two results are qualitatively the same, portraying the same pattern in band structures with differences in gap energies. Both calculations depict the smallest band gap of Ba<sub>2</sub>TiTeO<sub>6</sub> and that of Ba<sub>2</sub>TiSnO<sub>6</sub> is the largest. In the TB-mBJ-based DFT calculations, Shah et al. determined the band gap energy of Ba<sub>2</sub>TeTiO<sub>6</sub> to be 1.26 eV. <sup>26</sup> Meanwhile, Mustonen et al. reported that the GGA-calculated band gap of the Te-containing Ba<sub>2</sub>Mn-TeO<sub>6</sub> perovskite is 1.75 eV. 56 Compared to these findings, which involve different B- or B'-cations, our GGA-computed band gap of Ba<sub>2</sub>TiTeO<sub>6</sub> is 1.394 eV, providing a good point of comparison. Shah et al. also reported the band gaps for barium-titanium based Ba<sub>2</sub>HfTiO<sub>6</sub> and Ba<sub>2</sub>CeTiO<sub>6</sub> DPOs, with values of 3.44 eV and 2.96 eV, respectively. These values are comparable to our findings for Ba<sub>2</sub>TiGeO<sub>6</sub>, Ba<sub>2</sub>TiSnO<sub>6</sub>, and Ba<sub>2</sub>TiSeO<sub>6</sub> double perovskites. The materials exploration utilizing different functionals theoretically yields varying results, each offering a unique perspective for comparison against experimental findings. Nonetheless, no exact theoretical or experimental data regarding the electronic properties of Ba<sub>2</sub>(Ge,Sn,Se)TiO<sub>6</sub> DPOs is found for

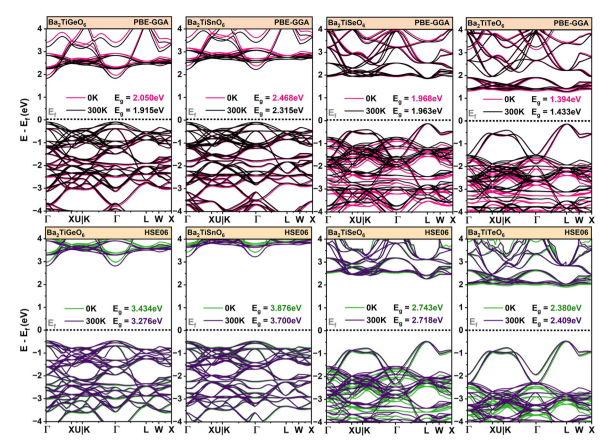
comparative reference, making this research pioneering. Therefore, our PBE-GGA and HSE06 calculated band gaps both provide worthwhile reference values that can guide future investigations.

The partial density of states (PDOS) presented in Fig. 4 shows that the valence band maximum (VBM) in the studied perovskites containing Ge, Sn, and Se is primarily occupied by O-p states, with some minor contributions from Ba-p and Ba-d states in the cases of Ge- and Sn-DPOs. While, in the case of Ba<sub>2</sub>TiSeO<sub>6</sub>, the minor contributions to the VBM are from the Ses and Ti-p states. However, for Ba<sub>2</sub>TiTeO<sub>6</sub>, the VBM is predominantly occupied by Te-s states, with minor contributions from Ti-p states. Therefore, the VBM of Ba<sub>2</sub>(Ge,Sn,Se)TiO<sub>6</sub> exhibits mainly a p-orbital characteristic, while the VBM of Ba<sub>2</sub>TeTiO<sub>6</sub> shows an s-orbital characteristic. On the other hand, the conduction band minimum (CBM) in the Ge- and Sn-based studied perovskites is primarily occupied by the Ti-d states and contributions from the Ge-s and Sn-s states, while, for Ba<sub>2</sub>Ti-SeO<sub>6</sub> and Ba<sub>2</sub>TiTeO<sub>6</sub>, the CBM is predominantly determined by Ti-d states, with significant contributions from the Se-p and Tep states, respectively. This indicates that the CBM of Ba<sub>2</sub>TiGeO<sub>6</sub> and Ba2TiSnO6 have s-orbital characteristics, while that of Ba<sub>2</sub>TiSeO<sub>6</sub> and Ba<sub>2</sub>TiTeO<sub>6</sub> have p-orbital characteristics. Moreover, these findings align with the PDOS results of Shah et al., which underlined the main contribution of oxygen atoms to the VBM and that of Ti atoms to the CBM and also indicated that the Ba atom does not make a significant contribution to the electronic structure.26

The band structures of Ba<sub>2</sub>(Ge,Sn,Se)TiO<sub>6</sub> materials show smaller band gaps at 300 K than 0 K, with a prominent decline in the Ge and Sn-bearing DPOs and a minute decrease in the Ba<sub>2</sub>TiSeO<sub>6</sub>, as Fig. 3 depicts. In contrast, the Te-bearing perovskite exhibits a slightly enhanced band gap at 300 K. The temperature-dependent band gaps in the 0 to 1000 K range, as shown in Fig. 5, illustrate the narrowing of band gaps in Ba<sub>2</sub>(Ge,Sn,Se)TiO<sub>6</sub> DPOs while widening in Te-containing perovskite. The narrowing of band gaps agrees with the Ünlü's thermodynamic model that characterizes the effect of pressure and temperature on the band gap of semiconductors. It explains that at elevated temperatures, the interaction of free electrons, holes, and recombined electron-hole pairs with lattice phonons produces thermal vibration energy, and the linear expansion in the lattice induces thermal strain energy. The aggregate of these energies is equal to the narrowing of

**Table 2** Structural properties of  $Ba_2TiMxO_6$  (Mx = Ge, Sn, Se, Te)

Mx			- Ba <sub>2</sub> TiTeO <sub>6</sub> Ba <sub>2</sub> ZrTiO <sub>6</sub> <sup>19,20</sup>	Ba <sub>2</sub> CaTeO <sub>6</sub> <sup>54</sup>	Ba MoYO	Ba <sub>2</sub> MnTeO <sub>6</sub>	Ba <sub>2</sub> MgWO <sub>6</sub>			
Quantities (PBE)	Ge	Sn	Se	Те	(PBEsol) <sup>26</sup>	(mBJ)	(mBJ)			
a [Å]	8.01	8.22	8.18	8.37	8.25	8.154, 8.24	8.39	8.07, 8.29	8.2106	8.1345
$V_{\text{st}} \begin{bmatrix} \mathring{\mathbf{A}} \end{bmatrix}^3 V_{\text{o}} \begin{bmatrix} \mathring{\mathbf{A}} \end{bmatrix}^3$	513.76	554.77	546.44	585.49						
$V_{\rm o} \left[ {\rm \mathring{A}} \right]^3$	514.39	556.34	547.91	588.58	563.448	542.22, 559.48	590.342	525.83, 568.89		538.26
$E_{\rm st}$ [eV]	-285.458	-284.892	-268.735	-272.670						
$E_{\rm o}$ [eV]	-285.461	-284.898	-268.740	-272.689						
B [GPa]	160.41	151.55	145.55	141.02	148.98	168.98, 152	155.72			137
$B^{\prime}$	4.85	4.78	4.74	4.70		3.79, NA	10.76			3.9
$\theta_{\mathrm{D}}\left[\mathrm{K}\right]$	549.59	517.50	525.67	499.79						



Graphical representation of the band structures of the examined DPOs at 0 and 300 K.

band gap energies.<sup>58</sup> Linearity in temperature-dependent band gap curves is also aligned with a linear relationship proposed by O'Donnell and Chen between the band gap of semiconductors and temperature, 59 as given by eqn (20).

$$E_{\rm g}(T) = E_{\rm g}(0) - 2SkT, \quad \frac{{\rm d}E_{\rm g}}{{\rm d}T} = -2Sk$$
 (20)

In this relationship,  $E_g(0)$  is the band gap at 0 K, k is the Boltzmann constant, and S is the dimensionless coupling constant. In their study, the band gaps of GaAs, GaP, and Si exhibited a negligible reduction up to a certain temperature and then decreased prominently. Likewise, the band gaps of Ba<sub>2</sub>TiGeO<sub>6</sub> and Ba<sub>2</sub>TiSnO<sub>6</sub> DPOs display a negligible decrease up to around 100 K and then decrease noticeably with the temperature rise.

In optical experiments conducted by Tauber et al., the direct band gap energy of the Te containing PbTe semiconductor at the L high symmetry point increases linearly with the temperature rise from 80 to 350 K with a temperature coefficient ( $\delta E_o$ )  $\delta T$ ) of 4.1  $\times$  10<sup>-4</sup> eV K<sup>-1</sup>,<sup>60</sup> as in the case of our studied Tecontaining Ba<sub>2</sub>TiTeO<sub>6</sub>. Tsang et al., in a theoretical band structure calculation based on the empirical pseudopotentials, found almost the same positive temperature coefficient as 3.9  $\times$ 10<sup>-4</sup> eV K<sup>-1</sup> of the PbTe semiconductor for the band gaps determined at the L high symmetry point for 40 to 200 K. They asserted the volumetric expansion and thermal vibrations of

the ionic cores are two main effects that alter the band gaps with temperature. They referred to these effects as the lattice effect and the Debye-Waller effect. The lattice expansion, which reduces the kinetic-energy term  $\hbar^2 |\vec{K} + \vec{G}|^2 / 2m$  in the pseudopotential Hamiltonian matrix element, accounts for the negative temperature coefficient. However, the Debye-Waller factor, which relates to the average squared displacement of the ion cores from their lattice positions, can cause a positive temperature coefficient depending upon the s-like or p-like symmetries in the conduction band minimum (CBM). The p-like elliptic symmetry is more sensitive to temperature than the s-like spherical symmetry, for example, the p-like CBM of PbTe causes Debye-Waller factors that lead to the positive temperature coefficient of PbTe.

Similarly, at elevated temperatures, the pronounced symmetry of the s-like characteristic in the conduction band of Ba<sub>2</sub>(Ge,Sn)TiO<sub>6</sub> and the valence band of Ba<sub>2</sub>(Se,Te)TiO<sub>6</sub> structures results in an isotropic Debye-Waller effect of thermal vibrations that induces a prominent shift of the density of states towards lower energy in Ge- and Sn- and towards higher energy in Se- and Te-bearing structures exhibited by both the band structures and total density of states depicted in Fig. 3 and 6. In contrast, the low symmetry p-like characteristic in the valence band of Ba<sub>2</sub>(Ge,Sn)TiO<sub>6</sub> and the conduction band of Ba<sub>2</sub>(Se,Te)TiO<sub>6</sub> structures results in unsymmetrical thermal vibrations and causes no significant shift. Moreover, the lattice

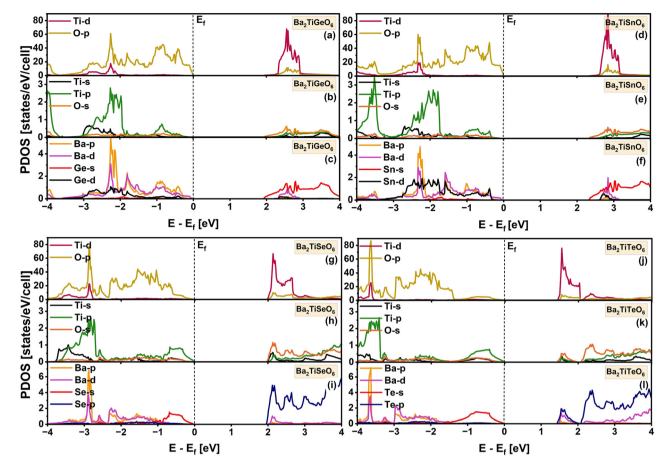


Fig. 4 The partial density of states of the examined DPOs at 300 K.

expansion diminishes the electron screening effects, thus causing a redistribution of the electronic density of the valence band to higher energy states and that of the conduction band electron density to lower energy states. These shifts render the negative temperature coefficients as  $-3.8\times10^{-4}$  and  $-4.0\times10^{-4}$  eV K $^{-1}$  of Ge- and Sn-keeping structures, respectively, and a little negative temperature coefficient as  $-2.78\times10^{-5}$  of Se- and a little positive temperature coefficient as  $7.94\times10^{-5}$  eV K $^{-1}$  of

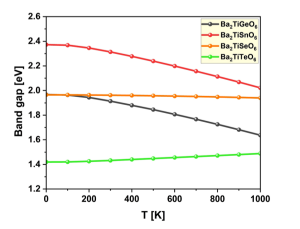


Fig. 5 Temperature-induced variations in the band gaps of  $Ba_2TiMxO_6$  (Mx = Ge, Sn, Se, Te) DPOs.

Te-bearing DPOs. These smaller coefficients denote the stable optoelectronic behavior of these materials under temperature change, which extends their application to high-temperature optoelectronic synaptic devices.

In semiconductors, the Fermi levels are typically positioned between the valence and the conduction band. The Fermi level's proximity to the VBM signifies p-type semiconductors, while its closeness to the CBM indicates n-type semiconductors. <sup>61–64</sup> Petit *et al.* and Park *et al.* noted a Fermi level in p-type ZnO semiconductors near the VBM. <sup>65,66</sup> Nolan *et al.* noted that a Fermi level closer to the VBM is a characteristic of p-type semiconductors. <sup>67</sup> Likewise, as Fig. 3 depicts, the Fermi levels of all the selected DPOs are situated near the VBM and indicate their p-type semiconducting properties.

The band gap energies of both PBE–GGA and HSE06 calculations indicate the UV range optical absorption of the studied compounds, denoting the applicability of studied materials in UV-driven optoelectronic devices. However, the PBE–GGA calculation for the  $\rm Ba_2TiTeO_6$  perovskite shows a 1.433 eV band gap at ambient temperature that not only indicates its potential for UV optoelectronics but also falls within the ideal range of 1.1–1.5 eV suited for the solar cells. Cao *et al.* derived the band gap temperature coefficient of the Te-based PbTe semiconductor as  $4.4 \times 10^{-4}$  eV K<sup>-1</sup> and witnessed that this positive coefficient is advantageous in attaining a high thermoelectric performance.<sup>68</sup>

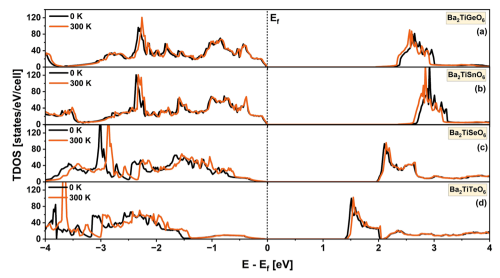


Fig. 6 The total density of states of the examined DPOs at 300 K.

Therefore, the positive coefficient of Ba<sub>2</sub>TiTeO<sub>6</sub> as 7.94  $\times$ 10<sup>-5</sup> eV K<sup>-1</sup> also indicates the benefit of its thermoelectric characteristics.

3.3 Optical properties

This study computed the optical properties at 300 K. The computation reveals that the selected DPOs exhibit isotropic optical properties and show symmetrical spectra in the x, y, and z directions due to their cubic structures. The frequencydependent real part of the dielectric function  $\varepsilon_{\rm r}(\omega)$  describes the polarization and dispersion of the light. <sup>69,70</sup> Under static conditions, at zero frequency, the  $\varepsilon_{\rm r}(0)$  represents the dielectric constant or the static dielectric function. A higher dielectric constant suggests a lower recombination rate of the charge carriers,<sup>71</sup> a lower band gap,<sup>72,73</sup> and a better optoelectronic performance of a material.<sup>74</sup> As seen in Fig. 7, the chosen perovskites bearing Ge, Sn, Se, and Te have dielectric constants of 5.13, 4.90, 6.75, and 6.15, respectively. So, Ba<sub>2</sub>TiTeO<sub>6</sub> with the lower band gap of 1.433 eV and a higher dielectric constant of 6.15 is indicated to be a better optoelectronic material.

Several studies reported an inverse relationship between the band gap and the dielectric constant as  $\varepsilon_{\rm r}(0) \approx 1 + (\hbar\omega_{\rm p}/E_{\rm gap})^2$ , where  $\hbar\omega_{\rm p}$  denotes the plasma frequency and  $E_{\rm gap}$  is the Penn band gap which is the energy equivalent frequency of the absorption peak. 72,73,75 The studied perovskites appear to follow this inverse relationship, where the Ba<sub>2</sub>TiSnO<sub>6</sub> with the highest band gap of 2.315 eV among the selected DPOs exhibits the lowest dielectric constant of 4.90, Ba<sub>2</sub>TiGeO<sub>6</sub> and Ba<sub>2</sub>TiTeO<sub>6</sub> with 1.915 and 1.433 eV band gaps exhibit comparatively higher dielectric constants as 5.13 and 6.15, and differently Ba<sub>2</sub>TiSeO<sub>6</sub> exhibits a higher band gap of 1.963 eV and a higher dielectric constant of 6.75 due to its lower Penn band gap and higher plasma frequency, examined as follows. It is mentioned that the Penn band gap corresponds to the frequency of the absorption peak,<sup>72</sup> and a comparatively low-frequency absorption peak of Ba<sub>2</sub>TiSeO<sub>6</sub> close to the violet-edge (3.26 eV) of the visible spectrum indicates its lower Penn band gap and that denotes its higher dielectric constant. The plasma frequency is

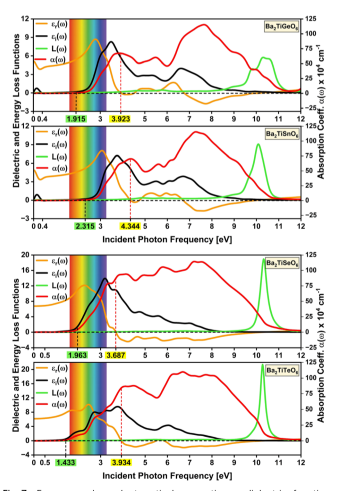


Fig. 7 Frequency-dependent optical properties - dielectric functions  $(\varepsilon_r(\omega), \varepsilon_i(\omega))$ , energy loss functions  $L(\omega)$ , and absorption coefficients  $\alpha(\omega)$ of the analyzed DPOs at 300 K.

a characteristic frequency at which free electrons collectively oscillate, and below this frequency, semiconductors exhibit metallic behavior and reflect most of the incident light.<sup>76-78</sup>

The energy loss function, which is the imaginary part of the inverse of the complex dielectric function  $(-Im[1/\varepsilon(\omega)])$  and describes the energy loss of an electron on interaction with the material, peaks at the plasma frequency.<sup>79</sup> Given these definitions, the plasma frequencies of the Ge-, Sn-, Se-, and Tecontaining studied DPOs are 10.34, 10.08, 10.32, and 10.28 eV, respectively, as evident in Fig. 7 for the peaks of energy loss functions. These values are comparable to the reported ones for double perovskites. The plasma frequencies of Sr<sub>2</sub>FeReO<sub>6</sub> and Ba<sub>2</sub>FeReO<sub>6</sub> DPOs are cited as 12.0 and 11.4 eV, respectively, 80 and that of Ba2ScNbO6 and Ba2ScTaO6 are noted as 10.16 and 10.24 eV, respectively. 81 Like the lower Penn band gap, the higher plasma frequency of the Ba<sub>2</sub>TiSeO<sub>6</sub> perovskite also suggests its higher dielectric constant in connotation to Penn's model.

As seen in Fig. 7, the  $\varepsilon_r(\omega)$  functions reach maximum values of 8.69, 8.04, 12.14, and 10.16 at frequencies of 2.79, 3.07, 2.28, and 2.45 eV for the Ge-, Sn-, Se-, and Te-bearing studied DPOs respectively. These values exhibit similar patterns to the dielectric constants, showing a higher peak value of the dielectric function for Ba<sub>2</sub>TiSeO<sub>6</sub>. All these peaks are within the visible light range, indicating maximum polarization and dielectric response in this range. Above critical frequencies of 3.923, 4.344, 3.687, and 3.934 eV for the studied Ge, Sn, Se, and Te-containing DPOs, as highlighted in yellow in both Fig. 7 and 8, the  $\varepsilon_r$  functions become negative and exhibit a metallic reflective response. The corresponding peaks in the reflectivity spectra are apparent. However, in the case of Ge- and Sn-containing chosen perovskites, the  $\varepsilon_r$  functions fluctuate around zero value above the critical frequencies, and the negative region begins at around 6.76 and 6.72 eV, indicating their subtle metallic responses at higher frequencies. Fig. 7 depicts that, after minima, the reflective response region ends near the plasma frequencies. Correspondingly, peaks in the reflectivity spectra can be observed before plasma frequencies.

Moreover, the Ge and Sn-containing studied perovskites show a valley region in their  $\varepsilon_r$  functions starting from zero frequency with a dip at 0.31 and 0.27 eV, respectively. These dips characterize the Drude peaks in the spectra of  $\varepsilon_i$ , absorption coefficients, and extinction coefficients. These peaks express that Ba<sub>2</sub>TiGeO<sub>6</sub> and Ba<sub>2</sub>TiSnO<sub>6</sub> are degenerate semiconductors, showing the intraband optical transitions for the intrinsic far infra-red (Far-IR) absorption within Ge-3d and Sn-3d conduction bands states of the Ba<sub>2</sub>TiGeO<sub>6</sub> and Ba<sub>2</sub>TiSnO<sub>6</sub>, respectively. 82-86 Islam et al. observed this far-IR peak for Cs2AgBiBr6 double halide perovskites below 1.5 eV due to the intra-band optical transitions within the Bi-6p and Ag-3d conduction band states.87 Nguyen et al. observed this peak at 0.14 eV for Ba<sub>2</sub>TiMnO<sub>6</sub> that occurs due to the intra-band excitation of Mn-3d orbitals of the conduction band. <sup>25</sup> Bulutay has discussed the intra-band transition peaks at around 0.5 eV for Si and Ge nano-crystals and underlined the use of these transitions in mid- and near-infrared photodetection.82

The imaginary part of the dielectric function  $\varepsilon_i(\omega)$  is an essential function, describing the optical transitions that

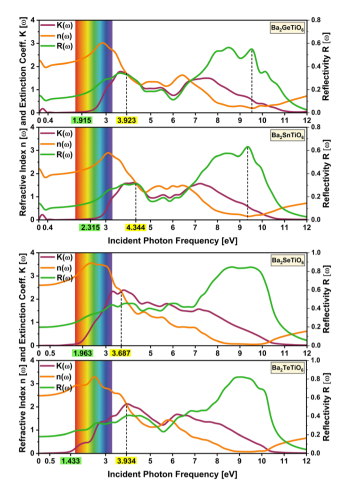


Fig. 8 Frequency-dependent optical properties – extinction coefficients  $K(\omega)$ , refractive indexes  $n(\omega)$ , and reflectivities  $R(\omega)$  of the analyzed DPOs at 300 K.

correspond to the band structures and indicating the light absorption characteristics. <sup>69,70,88</sup> Initial peaks in the  $\varepsilon_i$  functions of Ba<sub>2</sub>TiGeO<sub>6</sub> and Ba<sub>2</sub>TiSnO<sub>6</sub> depict the far-IR absorption peaks, as discussed earlier, showing the electronic excitation of Ge-3d and Sn-3d conduction band states, respectively. The spectra of  $\varepsilon_i$ , absorption coefficients, and the extinction coefficients exhibit the onset of the inter-band transitions in the visible energy range almost at the electronic band gap energies, highlighted green in Fig. 7 and 8. This indicates the visiblelight absorption of the chosen DPOs. However, in the case of Ge- and Sn-bearing perovskites, these spectra peak a little ahead of the violet edge of the visible spectrum, denoting their pronounced UV absorption. Instead, the  $\varepsilon_i$  spectra of Se- and Te-bearing perovskites peak in the visible range and span over the UV region, indicating their vis-UV absorption.

Interestingly, the absorption spectra of all studied DPOs exhibit peaks at the critical energies in the UV range, showing the absorption coefficients of  $67.196 \times 10^4$ ,  $70.269 \times 10^4$ ,  $87.260 \times 10^4$ , and  $84.557 \times 10^4 \ \mathrm{cm^{-1}}$  for the studied Ge-, Sn-, Se-, and Te-containing DPOs. Besides, the studied DPOs exhibit around 15.02%, 14.26%, 19.72%, and 18.10% reflection at low frequencies that increase to 34.37%, 31.26%, 45.36%, and

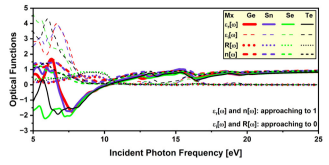


Fig. 9 Optical response of dielectric functions, reflectivities, and refrac tive indexes at higher frequencies of the studied Ba<sub>2</sub>TiMxO<sub>6</sub> DPOs.

40.89% near the critical energies of Ge-, Sn-, Se-, and Te-bearing perovskites, respectively.

The refractive index  $n(\omega)$  characterizes the transparent nature of a material. As depicted in Fig. 8, the static refractive indexes,  $n(0) = \sqrt{\varepsilon_r(0)}$  as implied by eqn (6), are 2.27, 2.21, 2.60, and 2.48 for the studied Ge, Sn, Se, and Te-containing DPOs. The refraction spectra manifest peak indexes of 3.02, 2.89, 3.56, and 3.30, at 2.84, 3.11, 2.36, and 2.51 eV within the visible frequencies. High refractive indexes within the visible spectrum demonstrate that these materials are less transparent and more prone to light absorption in this range. Also, the refraction spectra of Ge- and Sn-containing perovskites display a trough at low frequency, indicative of the far-IR light absorption.

Notably, beyond plasma frequencies, the  $\varepsilon_r$  functions approach an asymptotic value of one, while the  $\varepsilon_i$  functions approach a null value,  $R \approx 0$ , and  $n \approx 1$ , as depicted in Fig. 9. These elucidate that the studied materials tend toward transparent behavior beyond plasma frequencies.

Another essential optical parameter is the optical band gap  $(E_{\text{opt}})$  that represents the energy threshold for optical absorption that merely creates a bound electron-hole pair (exciton). In contrast, the electronic band gap  $(E_g)$  represents the energy required to excite an electron from the VBM to the CBM and create an unbound electron-hole pair that necessarily includes the binding energy of the exciton.<sup>89,90</sup> In solar cell or optoelectronic studies, the optical gaps are more important for indicating the

**Table 3** Electronic and optical band gaps of  $Ba_2TiMxO_6$  (Mx = Ge, Sn, Se, Te) at 300 K

Band gap [eV]	Ba <sub>2</sub> TiGeO <sub>6</sub>	$Ba_2TiSnO_6$	Ba <sub>2</sub> TiSeO <sub>6</sub>	Ba <sub>2</sub> TiTeO <sub>6</sub>
$\frac{E_{ m g}}{E_{ m opt}}$	1.915	2.315	1.963	1.433
	2.311	2.603	1.681	1.325

minimum energy absorption requirement.<sup>91</sup> Herein, the optical band gaps are determined using Tauc plot technique. 92,93 The values of  $(\alpha h \nu)^{1/2}$  as a function of incident photon frequency are plotted and then extrapolate the linear region in the rise of  $(\alpha h\nu)^{1/2}$ to the abscissa, where the intersections provide us with the optical band gaps, as shown in Fig. 10. Table 3 lists the electronic and optical band gaps, illustrating that the optical band gaps of Ba<sub>2</sub>TiGeO<sub>6</sub> and Ba<sub>2</sub>TiSnO<sub>6</sub> are higher, while those of Ba<sub>2</sub>TiSeO<sub>6</sub> and Ba2TiTeO6 are lower than their electronic band gaps.

Typically, optical band gaps are lower than electronic band gaps. To understand the higher optical band gaps of the studied Ge- and Sn-containing DPOs, the transition probabilities  $(P^2)$  that represent the sum of the squared transition dipole moments (TDM) are calculated along the k-path, as shown in Fig. 11. Moreover, the contribution weights of the atomic orbitals of each element for the VBM and CBM are calculated and presented as histograms in Fig. 12. The contribution weights of the atomic orbitals for the VBM and CBM dictate that the O-py and O-px states constitute the VBM, while Ti- $dz^2$ , Ti- $dx^2-y^2$ , and Ge-s and Sn-s constitute the CBM in the studied Ge- and Sn-containing materials.

Selection rules govern dipole transitions, allowing transitions between states of opposite parity. Although the transitions from O-p states to Ge-s or Sn-s states are allowed transitions, these are not witnessed in Fig. 11. This is because the lack of sufficient overlap, as indicated by the contribution weights of orbitals, prevents these transitions from occurring. Transitions from O-p to Ti-d states are also permitted. However, the inversion symmetry in the perovskite structures of Ba<sub>2</sub>Ti- $GeO_6$  and  $Ba_2TiSnO_6$  prevents these transitions at the  $\Gamma$  point, which is a location of their band gaps. This is evident from the absence of peaks in  $P^2$  at the  $\Gamma$  point, as depicted in Fig. 11. Thus, for Ba<sub>2</sub>TiGeO<sub>6</sub> and Ba<sub>2</sub>TiSnO<sub>6</sub>, the optical transitions are

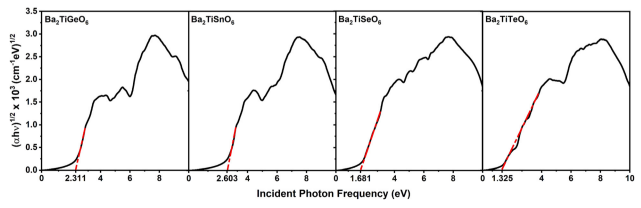


Fig. 10 Optical band gap calculation – graphical representation of Tauc plot for  $Ba_2TiMxO_6$  (Mx = Ge, Sn, Se, Te) DPOs.

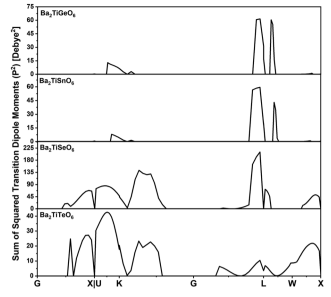


Fig. 11 Probabilities of transitions from the valence band to the conduction band across the high symmetry k-path for the chosen DPOs.

symmetry-forbidden, and the presence of transition peaks at other k-points accounts for the higher optical gaps observed in the Ge- and Sn-containing materials. In contrast, for Ba<sub>2</sub>TiSeO<sub>6</sub> and Ba2TiTeO6, the VBM is primarily occupied by Op, Se-s, and Te-s states, and the CBM is predominantly Ti-d states. This electronic structure suggests the potential for optical transitions from s to d and p to d states. However, at the L high symmetry point—which corresponds to the band gap of these materials—the  $P^2$  plot expresses the weak transitions. This is due to the imperfect spatial overlap between the O-p and Ti-d states, particularly in the case of Ba<sub>2</sub>TiSeO<sub>6</sub> perovskite.

These optical properties demonstrate that Ba<sub>2</sub>TiSeO<sub>6</sub> and Ba<sub>2</sub>TiTeO<sub>6</sub> exhibit low optical band gaps of 1.681 and 1.325 eV within the visible energy range, higher dielectric constants of 6.75 and 6.15, higher absorption coefficients of  $87.260 \times 10^4$ 

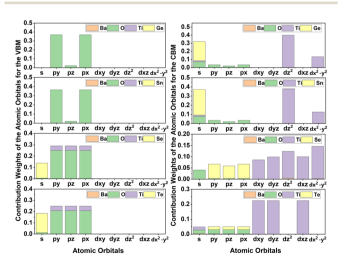


Fig. 12 Contribution weights of each atom's electronic orbitals to the valence and conduction band edges

and  $84.557 \times 10^4 \text{ cm}^{-1}$  at critical energies, higher static refractive indexes of 2.60 and 2.48 and higher peak values of 3.56 and 3.30 at 2.36 and 2.51 eV within visible frequencies. Consequently, these perovskites exhibit more suitable optical characteristics for vis-UV absorption.

### Thermodynamic properties

Employing QHDM, the thermodynamic properties of the investigated DPOs are computed for temperature variations ranging from 0 to 700 K at constant pressures of 0 and 15 GPa and pressure variations from 0 to 20 GPa at temperatures of 300 and 500 K. Fig. 13 demonstrates that the cell volume of all structures decreases under pressure and increases with temperature, regardless of the specific temperature or pressure selected. The curves of the isothermal bulk moduli show an increasing trend under pressure for both 300 and 500 K temperatures and a decrease for both 0 and 15 GPa pressures. Under pressure, atoms are forced closer together, reducing the overall volume of the crystal and increasing its hardness, as shown by an increase in bulk modulus. Instead, an increase in temperature causes the lattice parameters to expand, resulting in an overall increase in the crystal's volume. This expansion is due to the thermal vibration of atoms within the lattice, leading to a decrease in material hardness, evident with decreasing bulk modulus. However, at very low temperatures below 90 K, the cell volume and the bulk moduli exhibit a static behavior. This is because the interatomic forces that tightly pack atoms in a crystal lattice are strong enough to resist significant expansion at low temperatures. The characteristics of both cell volumes and bulk moduli of all studied DPOs are smooth with slight gradients, and no abnormalities are apparent, demonstrating their thermal stability.

The molar heat capacities per formula unit, displayed in Fig. 14, decrease as pressure increases for each studied DPO, exhibiting more steepness at 300 K compared to 500 K. At high

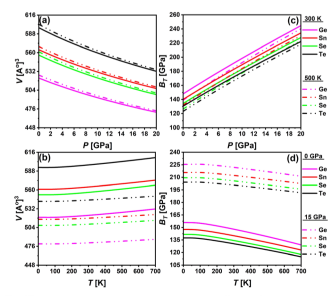


Fig. 13 Pressure and temperature-restrained effects of crystal volumes and isothermal bulk moduli of  $Ba_2TiMxO_6$  (Mx = Ge, Sn, Se, Te).

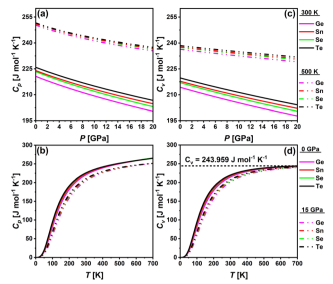


Fig. 14 Pressure and temperature-restrained effects of molar heat capacities of  $Ba_2TiMxO_6$  (Mx = Ge, Sn, Se, Te)

temperatures, the atoms already possess a significant amount of kinetic energy, so the additional compression from increased pressure has a minor impact on the heat capacities, and vice versa. Essentially, the O-p states along with Ge-s, Sn-s, Se-p, and Te-p states of the Ge-, Sn-, Se-, and Te-bearing studied perovskites, respectively, near the Fermi level provide a bunch of electronic states that make a reasonable electronic contribution to heat capacities. Besides, these curves manifest that the values of  $C_p$  are consistently higher than those of  $C_v$  for all selected materials. This observation aligns with the established understanding that  $C_v$  does not account for the expansion work associated with volume, while  $C_p$  values include work done in the expansion. 94 As noticed, the pressure-dependent isotherms of  $C_p$  of the studied DPOs are closer at high temperatures, highlighting that their values are approaching the same limit, and the trend is the same for  $C_v$ . This is because, as temperature surpasses the Debye temperatures, which are 549.59, 517.50, 525.67, and 499.79 K for Ge-, Sn-, Se-, and Te-bearing DPOs, respectively, C<sub>v</sub> curves approach a Dulong-Petit limit equal to 3NR, which is equal to 249.3 J mol<sup>-1</sup> K<sup>-1</sup> for structures of ten atoms in their formula unit. Seen here, the temperaturedependent isobars of  $C_v$  exhibit the values of 242.96, 243.64, 243.51, and 244.03 J mol<sup>-1</sup> K<sup>-1</sup> at 0 GPa and 700 K, however, the values approach 246.48, 246.78, 246.73, and 246.96 at 0 GPa and 1000 K for the Ge-, Sn, Se-, Te-bearing studied DPOs, respectively.

The pressure and temperature-varied thermal expansion coefficients ( $\alpha$ ), presented in Fig. 15(a) and (b), exhibit a substantial parabolic reduction in  $\alpha$  with increasing pressure and a rapid rise in  $\alpha$  with temperature up to approximately 250 K. Following 250 K, a low gradient rising linear function  $\alpha(T)$  is evident. At 300 K, the pressure-dependent  $\alpha$  values are lower than those at 500 K, while the temperature-dependent  $\alpha$ exhibits greater values at lower pressure. Eqn (16) elucidates the dependency of  $\alpha$  on bulk moduli and  $C_v$  values. Higher bulk

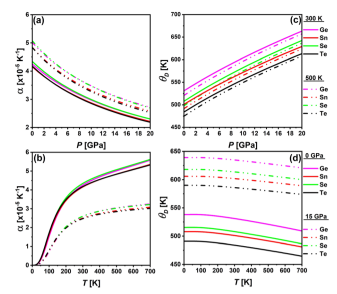


Fig. 15 Pressure and temperature-restrained effects of thermal expansion coefficients ( $\alpha$ ), and Debye temperatures ( $\theta_D$ ) of Ba<sub>2</sub>TiMxO<sub>6</sub> (Mx = Ge, Sn, Se, Te).

moduli and lower  $C_v$  at 300 K, as evidenced in Fig. 13(c) and 14(c) respectively, yield lower  $\alpha$  at 300 K, with the converse is observed at 500 K. Likewise, the lower bulk moduli and elevated values of  $C_v$  at low pressure, as demonstrated in Fig. 13(d) and 14(d), respectively, lead to heightened  $\alpha$  at lower pressure, and conversely.

The Debye temperature  $(\theta_D)$  represents the upper limit of atomic vibrational frequencies in crystalline materials. It provides a correlation between thermal and elastic properties and assists in characterizing phonons and the thermal behavior of materials, like specific heat, thermal conductivity, and enthalpies. It reflects the stiffness of solids and thus correlates with melting temperatures, such as a higher Debye temperature indicating a higher melting point. 95-98 The investigated materials exhibit rising  $\theta_{\rm D}$ under pressure and decreasing  $\theta_D$  with temperature rise, as shown in Fig. 15(c) and (d). Primarily, eqn (11) governs the dependence of  $\theta_{\rm D}$  on bulk moduli. A pressure rise increases the bulk modulus and implies higher  $\theta_D$ , which raises the stiffness and the melting points. While temperature reduces the bulk modulus and accordingly the  $\theta_D$ . Typical solids show the values of  $\theta_D$  in the 200 to 400 K range. The studied DPOs with the Ge-, Sn-, Se-, and Tecontaining structures exhibit  $\theta_D$  values of 531.09, 507.96, 501.03, and 484.05 K, respectively, at 0 GPa and 300 K. These values are higher than the typical range, indicating the stiffness and hardness of the studied perovskites.

Table 4 compares our results with theoretical and experimental findings at an ambient pressure of 0 GPa and a temperature of 300 K. The comparison demonstrates that our results agree with the reported data overall. The bulk modulus of Ba2TiGeO6 is very close to those of Ba2CeCoO6 and Ba2MgReO<sub>6</sub> DPOs, and the values of  $C_p$ ,  $C_v$ , and  $\alpha$  closely match the theoretical values, while the experimental  $C_p$  and  $\alpha$  values are slightly lower. The slight differences adhere to the diversity of material compositions.

DPOs  $V [\mathring{A}]^3$  $C_{\rm p}$  [J mol<sup>-1</sup> K<sup>-1</sup>]  $C_{\rm v}$  [J mol<sup>-1</sup> K<sup>-1</sup>]  $\alpha \; [\times \; 10^{-5} \; K^{-1}]$  $B_T$  [GPa]  $\theta_{\rm D} [K]$ Ba<sub>2</sub>TiGeO<sub>6</sub> 522.16 147.93 220.65 214.33 531.09 4.25 Ba<sub>2</sub>TiSnO<sub>6</sub> 564,40 140.12 224.01 217.84 507.96 4.16 Ba<sub>2</sub>TiSeO<sub>6</sub> 556.39 134.35 223.41 217.04 501.03 4.35 Ba<sub>2</sub>TiTeO<sub>6</sub> 597.17 130.51 225,91 219.76 484.05 4.19  $Ba_2^T IMnO_6^{0.23}$ 219.06 213.90 540.98 4.12 Ba<sub>2</sub>NaIO<sub>6</sub><sup>12</sup> 97.76 612.89 234.42 225.69 428,93 5.68 Ba<sub>2</sub>CeCoO<sub>6</sub><sup>99</sup> 147.32 496.56 Ba<sub>2</sub>CeCoO<sub>6</sub>
Ba<sub>2</sub>MgReO<sub>6</sub><sup>100</sup>
Ba<sub>2</sub>HoRuO<sub>6</sub><sup>expt., 101</sup>
Ba<sub>2</sub>HoIrO<sub>6</sub><sup>expt., 101</sup> 146 99 220.01 489,77 4.14 204.92 204.30 Ba<sub>2</sub>MgWO<sub>6</sub> expt., 102 3.22

Thermodynamic properties of  $Ba_2TiMxO_6$  (Mx = Ge, Sn, Se, Te) at the ambient temperature and pressure (300 K, 0 GPa)

#### Stability analysis 3.5

### 3.5.1 Stability analysis based on the geometric parameters. The formability and stability of perovskite structures can be assessed using the geometric parameters, known as tolerance ( $\tau$ ) and octahedral factors ( $\mu$ ). <sup>103,104</sup> Goldschmidt proposed an empirical relationship for the tolerance factor, as given in eqn (21), to predict the formability of perovskites based on the sizes of ionic radii that constitute the perovskite structure. 105

$$\tau = \frac{1}{\sqrt{2}} \frac{r_{\rm A} + r_{\rm X}}{r_{\rm B} + r_{\rm X}} \tag{21}$$

Here,  $r_A$ ,  $r_B$ , and  $r_X$  are the ionic radii of A and B cations, and X anions, respectively. In the case of double perovskites, the average ionic radii of B and B' ions are equivalent to  $r_{\rm B}$ . Goldschmidt documented that a structure is a stable cubic perovskite structure in the limit  $0.8 < \tau < 1$ . The dataset of perovskites, including double perovskites, has accumulated over the decades, leading to frequent fluctuations in this limit. A recent research study stated that DPOs (A2BB'O6) are stable in the limit  $0.82 < \tau < 1.1.^{106}$  The tolerance factors of the selected DPOs are computed using the Shannon revised data for ionic radii corresponding to the oxidation state and the coordination number of the individual ion. 107 Additionally, the tolerance factors according to Goldschmidt's non-rattling principle, which minimizes the ionic rattling and ionic space such that the cation's surface touched the anion's surface, 103 are calculated. For this purpose, the cation-anion bond lengths provided in Table 1 are utilized for  $r_A + r_X$  and  $r_B + r_X$  sums. Tolerance factors of both calculations are presented in Table 5 which indicates that the selected DPOs are stable structures both ways, as the values are lying within the limit 0.82  $< \tau <$ 1.1, where the calculation based on the non-rattling principle reveals the perfect cubic structure of these DPOs with  $\tau$  equal to 1 at both 0 and 300 K. The stability as examined with the nonrattling principle which does not account for the octahedral tilting and distortions are further evaluated by an octahedral factor ( $\mu = r_B/r_X$ ) as Li et al. proposed. This factor limits the octahedral tilting or distortions and together with the tolerance factor is necessary to predict the formability of perovskites. By Li et al.,  $\mu > 0.425$  endorses the formability of perovskite oxides, for which  $r_X = r_O$ . As demonstrated in Table 5, the octahedral factors of the BaSnTiO6 and BaTeTiO6 structures

surpass the lower limit, signifying their superior structural robustness and formability. Conversely, the octahedral factors of the other two structures marginally fall short of the limit, underscoring their diminished structural stability. In contrast to these two descriptors  $(\tau, \mu)$  stability analysis that has 80% accuracy as Marina et al. reported, 103 the new tolerance factor  $(\tau_n)$  recently introduced by Bartel *et al.*, which has integrated  $\mu$  in its formulation described in eqn (22), exhibited 92% accuracy solely sufficient in predicting stable perovskite structures. 109

$$\tau_{\rm n} = \frac{r_{\rm B}}{r_{\rm X}} - n_{\rm A} \left[ n_{\rm A} - \frac{r_{\rm A}/r_{\rm B}}{\ln(r_{\rm A}/r_{\rm B})} \right]$$
(22)

The author reported the stability of perovskites for  $\tau_{\rm n}$  < 4.18. Listed in Table 5, our calculated values of  $\tau_n$  being less than 4.18 indicate the stable structures of the selected DPOs.

In multiple studies, the authors reported the structural stability of Ba<sub>2</sub>CaTeO<sub>6</sub>, Ba<sub>2</sub>MnTeO<sub>6</sub>, and Ba<sub>2</sub>ZrTiO<sub>6</sub> DPOs, exhibiting tolerance factors within the range of  $0.82 < \tau < 1.1$  as mentioned in Table 5. Landinez et al. asserted the possibility of structural distortions in Ba<sub>2</sub>ZrTiO<sub>6</sub> for showing a tolerance factor greater than 1.18 Deluque-Toro et al. discussed that the computed  $\tau = 1.0778$  for Ba<sub>2</sub>TiMnO<sub>6</sub> disallows the perfect packing of Mn<sup>+4</sup> and Ti<sup>+4</sup> ions and indicates a distorted cubic structure.<sup>23</sup> In a recent study, Dai et al. described the stable cubic structures of Ba<sub>2</sub>AsXO<sub>6</sub> DPOs for demonstrating the tolerance and octahedral factors within the stability limits. 110 These studies ascertain our deductions about the stability of the chosen DPOs based on the geometric parameters.

Table 5 Geometric parameters of Ba<sub>2</sub>TiMxO<sub>6</sub> (Mx = Ge, Sn, Se, Te)

		τ (no rattl	ing)		
DPOs	τ	0 K	300 K	$ au_{\mathbf{n}}$	$\mu$
Ba <sub>2</sub> TiGeO <sub>6</sub>	1.082	1.00001	0.99992	3.908	0.405
Ba <sub>2</sub> TiSnO <sub>6</sub>	1.039	1.00041	1.00018	3.621	0.463
Ba <sub>2</sub> TiSeO <sub>6</sub>	1.090	1.00012	1.00023	3.983	0.395
Ba <sub>2</sub> TiTeO <sub>6</sub>	0.973	1.00039	1.00046	3.495	0.563
Ba <sub>2</sub> CaTeO <sub>6</sub> <sup>54</sup>	0.93				
Ba <sub>2</sub> MnTeO <sub>6</sub> <sup>56</sup>	1.00				
Ba <sub>2</sub> ZrTiO <sub>6</sub> <sup>18</sup>	1.031				
Ba <sub>2</sub> ZrTiO <sub>6</sub> <sup>19</sup>	1.1				
Ba <sub>2</sub> ZrTiO <sub>6</sub> <sup>20</sup>	0.85	0.94			0.47
$Ba_2TiMnO_6^{23}$	1.0778				
Ba <sub>2</sub> AsVO <sub>6</sub> <sup>110</sup>	1.057			3.590	0.467
Ba <sub>2</sub> AsNbO <sub>6</sub> <sup>110</sup>	1.067			3.652	0.452

### 3.5.2 Stability analysis based on the structural energetics

Cohesive and formation energies. The stability of the studied materials is further assessed from the computations of cohesive and formation energies, which are key factors in determining atomic forces in solids, as well as phase stability and formability. The formulation of these energies is described in eqn (23) and (24). It defines the cohesive energies as determined by subtracting the equilibrium energies of primitive lattices from the sum of elemental energies of isolated constituent atoms. Otherwise, the formation energies are calculated by subtracting the sum of the bulk energies of the individual constituent atoms from the equilibrium energies of the primitive lattices. These are divided by the total number of atoms in the primitive lattices to obtain the energy per atom. <sup>12,111,112</sup>

$$\Delta H_{c}^{\text{Ba}_{2}\text{TiMxO}_{6}} = \frac{\left(N_{\text{Ba}}E_{i}^{\text{Ba}} + N_{\text{Ti}}E_{i}^{\text{Ti}} + N_{\text{Mx}}E_{i}^{\text{Mx}} + N_{\text{O}}E_{i}^{\text{O}}\right) - E_{b}^{\text{Ba}_{2}\text{TiMxO}_{6}}}{N}$$
(23)

$$\begin{split} \Delta H_{\rm f}^{\rm Ba_2 TiMxO_6} &= \\ \underline{E_{\rm b}^{\rm Ba_2 TiMxO_6} - \left(N_{\rm Ba} E_{\rm b}^{\rm Ba} + N_{\rm Ti} E_{\rm b}^{\rm Ti} + N_{\rm Mx} E_{\rm b}^{\rm Mx} + N_{\rm O} E_{\rm b}^{\rm O}\right)} \\ N \end{split} \tag{24}$$

$$N = N_{\rm Ba} + N_{\rm Ti} + N_{\rm Mx} + N_{\rm O} \tag{25}$$

Here,  $E_{\rm b}$  represents the bulk energies per atom, calculated from the stable bulk phases of each element or structure, 112 while  $E_i$  refers to the energies of isolated constituent atoms. Table 2 lists the calculated cohesive and formation energies that witness the structural phase stability of Ba<sub>2</sub>TiMxO<sub>6</sub> (Mx = Ge, Sn, Se, Te) DPOs for exhibiting positive cohesive energies and negative formation energies in each case. More positive cohesive energies and more negative formation energies of Ba<sub>2</sub>Ti(Ge,Sn)O<sub>6</sub> DPOs demonstrate more tight atomic bonding in these structures and their higher phase stability compared with Ba<sub>2</sub>Ti(Se,Te)O<sub>6</sub> DPOs. Nevertheless, the Ba<sub>2</sub>TiSnO<sub>6</sub> exhibits superior stability with the lowest formation energy value of -2.789 eV per atom. Kazim et al. and Naseri et al. indicated the stability of the Ba2ZrTiO6 cubic structure based on their energies. This structure resembles our selected structures, showing a very low negative formation enthalpy of -14.991 eV per atom and positive cohesive energy of 4.43 eV per atom. 19,20 Ishfaq et al. recently studied Te-based Ba<sub>2</sub>CaTeO<sub>6</sub> double perovskite with the same  $Fm\bar{3}m$  cubic structure as in our calculations. The author described its stability for exhibiting the negative enthalpy of formation as -3.604 eV per atom. <sup>54</sup> Our results align with these findings. However, the condition  $E_{\rm f} \leq 0$  does not express the overall thermodynamic stability of a compound. Instead, it only evaluates the stability of a compound in terms of its ability to resist dissociation into elemental constituents. Besides cohesive and formation energies, decomposition energies of compounds, which indicate the possible decomposition of a compound into its parent phases, 113 are crucial in determining the stability of compounds.

Decomposition energies. Like cohesive and formation energies, decomposition energies can be utilized to evaluate the thermodynamic phase stability of a compound. Higher decomposition energies indicate more stable compounds, while lower decomposition energies suggest less stable compounds. Moreover, the compounds that exhibit negative decomposition energies in any of their possible decomposition reaction paths are supposed to be thermodynamically unstable, and they will spontaneously break down into their binary or ternary phases. Conversely, the positive decomposition energies signify the thermodynamic stability of the structure. 114-116 This investigation focused on the decomposition of the selected DPOs into their possible solid binary (MxO<sub>2</sub>) and ternary oxide phases (BaMxO<sub>3</sub>, BaTiO<sub>3</sub>, Ba<sub>2</sub>TiO<sub>4</sub>). Considering these binary and ternary oxide phases, the decomposition energies are computed for the following potential reaction pathways.

$$Ba_2TiMxO_6 \rightarrow BaTiO_3 + BaMxO_3$$
 (I)

$$Ba_2TiMxO_6 \rightarrow Ba_2TiO_4 + MxO_2$$
 (II)

The crystal structures of the experimentally observed solid phases of the considered binary and ternary oxides, as listed in Table 6, are utilized for computing the decomposition energies. Employing eqn (26) and (27) and after computing the static energies per formula unit of the selected DPOs and their decomposition phases, decomposition energies are determined.

$$\Delta H_{\rm d}^{\rm I} = E_{\rm BaTiO_2} + E_{\rm BaMxO_2} - E_{\rm Ba,TiMxO_6} \tag{26}$$

$$\Delta H_{\rm d}^{\rm II} = E_{{\rm Ba_2TiO_4}} + E_{{\rm MxO_2}} - E_{{\rm Ba_2TiMxO_6}}$$
 (27)

As listed in Table 6, the decomposition energies indicate that Ba<sub>2</sub>(Ge,Sn)TiO<sub>6</sub> are thermodynamically stable among the selected DPOs exhibiting all positive decomposition energies in both reaction pathways. In contrast,  $\Delta H_{\rm d}^{\rm I}$  energies of Ba<sub>2</sub>(Se,Te)TiO<sub>6</sub> structures are positive while their  $\Delta H_{\rm d}^{\rm II}$  energies are negative except one computed with tetragonal  $P4_2/mm$  TeO<sub>2</sub>. These negative energies indicate that Ba<sub>2</sub>(Se,Te)TiO<sub>6</sub> are thermodynamically unstable.

Sun et al. analyzed the decomposition energies of barium, strontium, and calcium-based perovskites. They noted that all perovskite compounds with positive decomposition energies demonstrated stability under experimental conditions, and the Ca<sub>2</sub>BiVO<sub>6</sub> double perovskite characterized by negative decomposition energy was deemed unstable in experiments. They also found that barium-based perovskites exhibit higher stability than the strontium- or calcium-based ones, with the calciumbased perovskites being the least stable. Additionally, they discovered that Ba<sub>2</sub>BiTaO<sub>6</sub>, Ba<sub>2</sub>SbTaO<sub>6</sub>, and Ba<sub>2</sub>BiVO<sub>6</sub> exhibit significant stability with decomposition energies of approximately 0.25, 0.2, and 0.195 eV per atom, respectively. 117 Given these results, our calculated decomposition energies indicate thermodynamically stable Ba<sub>2</sub>(Ge,Sn)TiO<sub>6</sub> structures which exhibit all positive decomposition energies and unstable structures of Ba<sub>2</sub>(Se,Te)TiO<sub>6</sub> DPOs with negative  $\Delta H_d^{\rm II}$  decomposition energies.

3.5.3 Stability analysis based on phonon spectra. The temperature-dependent phonon dispersion spectra of the

Table 6 Cohesive energies, formation energies, structural energies of the parent binary and ternary oxides, and the corresponding decomposition energies of  $Ba_2TiMxO_6$  (Mx = Ge, Sn, Se, Te)

Quantities	Ba <sub>2</sub> TiGeO <sub>6</sub>	$Ba_2TiSnO_6$	$Ba_2TiSeO_6$	$Ba_2TiTeO_6$	$\mathrm{Ba_2ZrTiO_6}^{19,20}$	Ba <sub>2</sub> CaTeO <sub>6</sub> <sup>54</sup>	
$\Delta H_{\rm c}$ [eV per atom]	6.946	6.933	6.527	6.626	NA, 4.43		
$\Delta H_{\rm f}$ [eV per atom]	-2.765	-2.789	-2.464	-2.552	−14.991, NA	-3.604	
Binary/ternary phase	Crystal structures	Static energy [eV f.u. <sup>-1</sup> ]	Ba₂TiGeO <sub>6</sub>	Ba <sub>2</sub> TiSnO <sub>6</sub>	Ba <sub>2</sub> TiSeO <sub>6</sub>	Ba <sub>2</sub> TiTeO <sub>6</sub>	
BaGeO <sub>3</sub>	Orthorhombic P2 <sub>1</sub> 2 <sub>1</sub> 2 <sub>1</sub>	-32.4444					
BaSnO <sub>3</sub>	Cubic <i>Pm</i> 3 <i>m</i>	-31.5522					
BaSeO <sub>3</sub>	Monoclinic P12 <sub>1</sub> /m1	-29.7556					
$BaTeO_3$	Monoclinic P12 <sub>1</sub> /m1	-29.6012					
			$\Delta H_{ m d}^{ m I}$ [eV f.u. $^{-1}$ ]				
BaTiO <sub>3</sub>	Tetragonal P4/mmm	-30.8188	8.1012	8.8520	6.6092	7.7473	
	Orthorhombic Amm2 (I)	-36.0201	2.8999	3.6508	1.4079	2.5460	
	Orthorhombic Amm2 (II)	-35.2095	3.7105	4.4614	2.2186	3.3567	
$Ba_2TiO_4$	Monoclinic P12 <sub>1</sub> /c1	-52.1318					
			$\Delta H_{ m d}^{ m II}$ [eV f.u. $^{-1}$ ]				
GeO <sub>2</sub>	Tetragonal P4 <sub>2</sub> /mnm	-19.1739	0.0587				
	Orthorhombic <i>Pnnm</i>	-19.1753	0.0574				
	Orthorhombic Pbcn	-19.0673	0.1654				
	Trigonal P3 <sub>2</sub> 21̄	-19.0097	0.2230				
	Cubic <i>Pa</i> 3̄	-18.5424	0.6903				
	Trigonal P3 <sub>1</sub> 21̄	-19.0103	0.2224				
	Tetragonal $P4_1\bar{2}_1\bar{2}$	-19.0025	0.2301				
$SnO_2$	Tetragonal P4 <sub>2</sub> /mnm	-18.7786		0.3127			
_	Orthorhombic Pnnm	-18.7796		0.3118			
	Cubic <i>Pa</i> 3̄	-18.4793		0.6121			
SeO <sub>2</sub>	Tetragonal P42/mbc	-15.4542			-0.4023		
	Orthorhombic Pmc2 <sub>1</sub> (I)	-15.4231			-0.3712		
	Orthorhombic Pmc2 <sub>1</sub> (II)	-15.4113			-0.3594		
	Orthorhombic Pham	-15.4663			-0.4144		
${ m TeO_2}$	Tetragonal P42/mnm	-15.4753				0.5604	
	Orthorhombic <i>Pbca</i>	-16.1341				-0.0985	
	Tetragonal $P4_3\bar{2}_1\bar{2}$	-16.1009				-0.0653	
	Tetragonal $P4_1\bar{2}_1\bar{2}$	-16.1011				-0.0655	
	Orthorhombic $P2_1\bar{2}_12\bar{1}$ (I)	-16.0962				-0.0606	
	Orthorhombic $P2_1\bar{2}_12\bar{1}$ (II)	-16.1004				-0.0648	

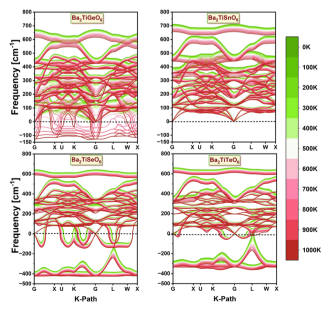


Fig. 16 Phonon dispersion spectra of the investigated Ba<sub>2</sub>TiMxO<sub>6</sub> (Mx = Ge, Sn, Se, Te) DPOs for the temperature range from 0 to 1000 K.

chosen DPOs, as displayed in Fig. 16, provide details about the stability concerning the structural distortions across a temperature range of 0 to 1000 K. Strict criteria are employed to obtain these spectra, including a fine k-mesh of  $12 \times 12 \times 12$  grid points and the structures are relaxed until the maximum Hellmann forces acting on each atom are less than  $10^{-6}$  eV Å<sup>-1</sup>. The phonon spectrum of Ba<sub>2</sub>TiGeO<sub>6</sub> indicates that this structure remains stable below 600 K, after which it becomes unstable, evident by the presence of soft modes with imaginary frequencies above this temperature. In contrast, the phonon spectrum of Ba<sub>2</sub>TiSnO<sub>6</sub> demonstrates a stable structure for all temperatures, apparent by the real frequency modes throughout the whole range of temperatures. However, the structures with Se+4 and Te<sup>+4</sup> cations are observed to be dynamically unstable at all temperatures, exhibiting soft modes across the entire temperature range. These findings are consistent with our conclusions drawn from the stability analyses based on the geometric factor and the structural energetics. The Ba<sub>2</sub>TiSnO<sub>6</sub> structure, which has a tolerance factor within limits, a higher octahedral factor than other selected DPOs, more positive cohesive energy, and more negative formation energy, and all positive decomposition energies, exhibits none of the negative frequency phonon modes

over an entire selected temperature range. Meanwhile, the Ba<sub>2</sub>TiGeO<sub>6</sub> structure, which has slightly lower cohesive energy and less negative formation energy than Ba2TiSnO6, exhibits dynamic stability up to 600 K, above which it shows soft phonon modes. Likewise, the Ba<sub>2</sub>(Se,Te)TiO<sub>6</sub> DPOs, which exhibit unstable structures in the energetic analysis, also demonstrate dynamic instability through their phonon spectra.

These stability analyses elucidate that Ba<sub>2</sub>TiGeO<sub>6</sub> and Ba<sub>2</sub>TiSnO<sub>6</sub> DPOs can be synthesized, while Ba<sub>2</sub>(Se,Te)TiO<sub>6</sub> perovskites will decompose spontaneously even after synthesis. In other words, the experimental reactions may result in a mixture of BaTiO<sub>3</sub>, Ba<sub>2</sub>TiO<sub>4</sub>, SeO<sub>2</sub> and BaSeO<sub>3</sub>, or TeO<sub>2</sub> and BaTeO<sub>3</sub> parent phases instead of the desired perovskites. However, the doping techniques with various concentrations of doped and dopant may enhance the structural stability of these DPOs.

### 4 Conclusions

This investigation uncovers the direct band gap p-type semiconducting characteristics of studied materials. Probing the temperature-modulated band gaps finds a narrowing of the band gap in Ba<sub>2</sub>Ti(Ge,Sn,Se)O<sub>6</sub> and a widening in Ba<sub>2</sub>TiTeO<sub>6</sub>. The temperature coefficients ( $\delta E_{\rm g}/\delta T$ ) are  $-3.8 \times 10^{-4}$ ,  $-4.0 \times 10^{-4}$  $10^{-4}$ ,  $-2.78 \times 10^{-5}$ , and  $+7.94 \times 10^{-5}$  eV K<sup>-1</sup>, respectively. These low-value coefficients infer the closely steady optoelectronic aspects under changing temperature conditions, indicating these materials' applicability for high-temperature optoelectronic synaptic devices like those used in aerospace and image processing.

The optical spectra of studied DPOs exhibit (1) the onset of optical transitions within the visible light spectrum near the band gap energies at 300 K and the absorption peaks at the critical frequencies of 3.923, 4.344, 3.687, and 3.934 eV in the UV region with absorption coefficients of 67.196, 70.269, 87.260, and  $84.557 \times 10^4 \, \mathrm{cm}^{-1}$  and reflection of 34.37%, 31.26%, 45.36%, and 40.89% for Ba<sub>2</sub>Ti(Ge,Sn,Se,Te)O<sub>6</sub> DPOs, respectively, highlighting the potential of these materials in UV-optoelectronics, and (2) the dielectric constants of 5.13, 4.90, 6.75, and 6.15 and optical band gaps of 2.311, 2.603, 1.681, and 1.325 eV, respectively, indicating their favorable optical performance. In addition, Ba<sub>2</sub>TiGeO<sub>6</sub> and Ba<sub>2</sub>SnTiO<sub>6</sub> exhibit Drude peaks at 0.31 and 0.27 eV, showing intraband transition within Ge-3d and Sn-3d conduction band states, respectively, demonstrating their application in mid- and nearinfrared photodetection systems.

In the span of stability, thermodynamic analysis shows that, under ambient conditions of 0 GPa and 300 K, the Ba<sub>2</sub>Ti(Ge,Sn,Se,Te)O<sub>6</sub> DPOs exhibit bulk moduli of 147.93, 140.12, 134.35, and 130.51 GPa, Debye temperatures of 531.09, 507.96, 501.03, and 484.05 K, and thermal expansion coefficients of 4.25, 4.16, 4.35, and  $4.19 \times 10^{-5} \text{ K}^{-1}$ , respectively, indicating the brittleness and hardness of these materials. The stability analyses established on geometric factors, cohesive energies, and formation energies demonstrate the dynamic stability of the chosen DPOs. Nevertheless, the decomposition energies and phonon spectra reveal the unstable

structures of Ba<sub>2</sub>(Se,Te)O<sub>6</sub> perovskites and the stable structure of Ba<sub>2</sub>TiSnO<sub>6</sub> across the entire temperature range of 0 to 1000 K, and the stable structure of Ba<sub>2</sub>TiGeO<sub>6</sub> up to 600 K. It establishes that among the materials studied, Ba<sub>2</sub>TiGeO<sub>6</sub> and Ba<sub>2</sub>TiSnO<sub>6</sub> stand out due to their stability. Therefore, with closely temperature-tolerant enticing optoelectronic characteristics and robust stabilities, these DPOs are promising candidates as a p-type functional layer in high-temperature UV-based photovoltaics and optoelectronics.

## Data availability

The authors confirm that the data supporting the findings of this study are available upon reasonable request.

### Conflicts of interest

There are no conflicts of interest.

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