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Comprehensive first-principles analysis and device simulations of vacancy-ordered D₂CeX₆ double perovskites for high-efficiency lead-free solar cells

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The quest for efficient and durable absorber materials has steered attention toward vacancy-ordered double perovskites, which exhibit tunable band gaps and strong optical absorption, making them promising candidates for next-generation solar cell architectures. In particular, lead-free vacancyordered halide double perovskites have emerged as viable alternatives to toxic Pb-based counterparts. In this study, we systematically investigate the structural, electronic, charges density, mechanical, optical, phonon stability, molecular dynamics (MD), population analysis and photovoltaic properties of D₂CeX₆ (D = Ga, In, Tl; X = Cl, Br) compounds by employing first-principles calculations in conjunction with SCAPS-1D device simulations. Our results reveal that all compounds crystallize in a stable cubic phase with negative formation enthalpies, confirming their thermodynamic stability. Within GGA-PBE, the calculated direct band gaps are 1.733 eV (Ga₂CeCl₆), 1.276 eV (Ga₂CeBr₆), 1.555 eV (In₂CeCl₆), 0.859 eV (In₂CeBr₆), 1.755 eV (Tl₂CeCl₆), and 1.364 eV (Tl₂CeBr₆), placing all but In₂CeBr₆ within or near the optimal 1.1 to 1.8 eV range for single-junction solar cells. HSE06 hybrid functional results yield wider gaps of 1.776 eV, 2.843 eV, 2.261 eV, 2.170 eV, 3.632 eV, and 1.418 eV, respectively, suggesting suitability for both single-junction and tandem architectures. Optical analyses demonstrate high absorption coefficients (>10⁵ cm⁻¹), strong dielectric responses, and large refractive indices, particularly in In₂CeBr₆ and Tl₂CeBr₆. Mechanical evaluations confirm ductile behavior, with Tl₂CeBr₆ and In₂CeBr₆ exhibiting superior stiffness and near-isotropic mechanical stability. Molecular dynamics simulations (NPT ensemble, 50 ps) at room temperature confirm the excellent thermal robustness of the studied compounds. The phonon dispersion analysis further indicates full dynamical stability for Ga₂CeCl₆, In₂CeCl₆, and Tl₂CeCl₆, while Ga₂CeBr₆, In₂CeBr₆, and Tl₂CeBr₆ exhibit minor soft or near-zero modes that are likely stabilized by finite-temperature effects. Device-level simulations predict power conversion efficiencies (PCE) up to 25.29% for Ga₂CeBr₆, with In₂CeCl₆ and Tl₂CeBr₆ also exceeding 24%. These findings position the D₂CeX₆ family, especially bromide-based compounds, as promising candidates for efficient and stable lead-free perovskite solar cells.

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Introduction

The global energy landscape is undergoing a profound transformation as the urgency to reduce dependence on fossil fuels and mitigate the impacts of climate change continues to intensify. Among the portfolio of renewable technologies, solar photovoltaics (PVs) have emerged as one of the most promising candidates, offering clean, abundant, and sustainable electricity generation.1,2 While silicon-based solar cells dominate the market due to their maturity and long-term reliability, their high fabrication costs and rigid processing requirements highlight the necessity of exploring alternative materials.3 In recent years, halide perovskite solar cells (PSCs) have revolutionized the field of photovoltaics, achieving remarkable power conversion efficiencies (PCEs) through cost-effective solution

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processing and outstanding optoelectronic properties. 4,5 However, the toxicity and environmental instability of Pb-based perovskites remain significant obstacles to their commercialization, thereby motivating the search for sustainable, lead-free alternatives.6 Vacancy-ordered double perovskites (A2BX6) are derived from the ABX3 structure by removing half of the B-site cations, forming isolated BX6 octahedra surrounded by A-site cations (e.g., Ga⁺, In⁺, Tl⁺) and halides (Cl⁻, Br⁻). This structural modification enhances stability and makes them promising lead-free alternatives for optoelectronic and energy applications. Depending on the B-site cation (e.g., Ce⁴⁺), they exhibit strong optical absorption, indirect-to-direct band gaps, and low thermal conductivity, supporting applications in solar cells, LEDs, radiation detectors, and thermoelectric.8,9

Previous studies have provided important insights into related systems. For example, A. Jabar et al. investigated Cs2-CeCl₆ using LSDA + mBJ in Wien2k, reporting a semiconducting bandgap of \sim 1.828 eV along with its optical and thermoelectric behavior, though photovoltaic device-level performance was not assessed. 10 Earlier, Kaatz and Marcovich (in the year of 1966) established that Cs2CeCl6 crystallizes in a cubic K2PtCl6-type structure (Pm3m), where Ce^{4+} is octahedrally coordinated by Cl^{-} ions and Cs+ is twelve-fold coordinated, offering a useful structural reference for related Ce-based halide perovskites.11 Meanwhile, hybrid organic-inorganic Pb-halide perovskites have achieved record PCEs of up to 25.5%,12 but their deployment is hampered by inherent toxicity and poor environmental durability. Building on these motivations, we turned our attention to Ce-based vacancy-ordered double perovskites D_2CeX_6 (D = Ga, In, Tl; X = Cl, Br). Similar investigations, such as that of Redi Kristian Pingak et al., confirmed the structural stability and semiconducting behavior of Tl₂SnCl₆ and Tl₂-SnBr₆, with Tl₂SnBr₆ (1.48 eV) identified as a promising absorber due to its strong visible-light response.13 Beyond photovoltaics, Ce-based oxides and halides demonstrate intriguing functionalities. Mehwish K. Butt et al. reported ferromagnetic half-metallicity in CeAlO3 arising from Ce-4f states, suggesting potential in spintronic and optical applications.14 Likewise, Mariya Zhuravleva et al. showed that Cehalides such as Cs₃CeCl₆, Cs₃CeBr₆, CsCe₂Cl₇, and CsCe₂Br₇ exhibit high light yields and fast decay times, underlining their promise as scintillators for radiation detection. 15 Recent studies have shown that vacancy-ordered CsRbGeCl₆ and CsRbGeBr₆ possess direct band gaps of 2.60 eV and 1.78 eV, respectively, at the Γ -point, making them highly effective for light absorption across the visible to UV spectrum. Their excellent structural stability, non-toxic composition, and favorable electronic properties make these materials promising candidates for photovoltaic and other optoelectronic applications.¹⁶ Similarly, Rb₂GeCl₆ and Rb₂GeBr₆ have been identified as lead-free alternatives with direct band gaps, mechanical robustness, and thermodynamic stability, making them suitable for renewable energy applications.¹⁷ Further, N. V. Skorodumova et al. demonstrated that Ce-4f states in CeO2 and Ce2O3 play decisive roles in governing electronic structure and bonding.18 Tianyu Tang et al. reported that vacancy-ordered double halide perovskites A_2BX_6 (A = In, Tl; B = Pd, Pt; X = Cl, Br, I) are

structurally stable semiconductors with tunable band gaps, strong UV-IR absorption, and potential applications as solar absorbers (Tl₂PdBr₆) or transparent conductors (In₂PtCl₆, Tl₂-PtCl₆), yet their device-level performance and practical applicability remain unexplored.19

Additionally, vacancy-ordered halide perovskites A₂BX₆ (A = In, Tl; B = Pd, Pt; X = Cl, Br, I) offer extensive tunability of their band gaps (from 0.328 eV to 3.238 eV in HSE functional) and exhibit pronounced optical responses across the visible to infrared spectrum. Compounds such as Tl₂PdBr₆ and In₂PtCl₆ are emerging as leading candidates for solar absorber layers and transparent conductors, respectively, owing to their superior absorption coefficients and direct or quasi-direct band structures.20-22 Similarly, Showkat Hassan Mir et al. investigated Cerium-based double perovskites A_2CeX_6 (A = Cs, K; X = Cl, Br) and found them to be structurally stable, non-magnetic semiconductors with direct band gaps, anisotropic mechanical and electronic properties, and strong visible-light absorption.²³ Taken together, these studies suggest that incorporating Ce into vacancy-ordered double perovskites can yield a rich interplay of structural stability, optical absorption, and electronic diversity.

The present investigation focuses on the performance analysis of D₂CeX₆ (D = Ga, In, Tl; X = Cl, Br) double perovskite compounds, carried out through a combination of firstprinciples calculations within the CASTEP framework and SCAPS-1D device simulations. The study comprehensively examines their structural, electronic, optical, thermoelectric, and mechanical properties, phonon stability, molecular dynamics, which are critical for energy-harvesting and photovoltaic applications. Structural stability was confirmed under cubic symmetry and mechanical deformation, ensuring robustness for practical implementation. Among the investigated compounds, bromide-based perovskites, particularly Ga₂CeBr₆ and Tl₂CeBr₆, demonstrated superior optoelectronic performance, characterized by optimal band gaps, high absorption coefficients, and strong dielectric responses, while In₂CeBr₆ exhibited notable mechanical robustness and ductility. To the best of current knowledge, no prior theoretical or experimental study has systematically explored the D2CeX6 family in this context. The findings establish Ce-based halide double perovskites as promising lead-free alternatives to toxic Pb-based materials, offering a well-balanced combination of efficiency, stability, and sustainability for next-generation photovoltaic and advanced power-generation technologies.

2 Computational method

Density functional theory (DFT)

We employed Density Functional Theory (DFT) within the CASTEP package to systematically investigate the XRD, structural, electronic, mechanical, optical properties, population analysis, and phonon properties of D_2CeX_6 (D = Ga, In, Tl; X = Cl, Br) double perovskites.24-27 This ab initio approach is widely recognized for its reliability in describing the physicochemical characteristics of crystalline materials. Ultra-soft Vanderbilttype pseudopotentials28 were employed for Ga, In, Tl, Ce, Cl, and Br atoms to describe the electron-ion interactions while Paper RSC Advances

maintaining computational accuracy efficiently.29,30 A planewave cut-off energy of 700 eV was adopted to ensure precise representation of the electronic states and reliable energy convergence. The Brillouin zone was sampled using a Monkhorst-Pack k-point³¹ grid of $8 \times 8 \times 8$ for all compounds, guaranteeing consistent convergence of total energy, forces, and stress tensors across reciprocal space.32 Geometry optimizations were performed under stringent convergence criteria, with thresholds set to 1.0×10^{-6} eV per atom for total energy, 0.006 eV Å^{-1} for maximum force, $2.0 \times 10^{-4} \text{ Å for maximum}$ atomic displacement, and 0.02 GPa for maximum stress. The generalized gradient approximation with the Perdew-Burke-Ernzerhof (GGA-PBE) functional was mainly employed, as it ensures a reasonable compromise between accuracy and computational efficiency.33,34 In contrast, the Hevd-Scuseria-Ernzerhof (HSE06) hybrid functional, though more accurate,35 is highly time-consuming; thus, GGA-PBE was chosen as the primary functional, with HSE06 selectively applied to validate key geometry optimizations and electronic results. Calculations of the density of states (DOS), optical properties, X-ray diffraction (XRD) patterns, elastic constants, mechanical parameters, anisotropy index, and population analysis were performed within the GGA-PBE framework. The crystal structures of the compounds were modeled and visualized using VESTA software.36 Mechanical properties and elastic constants were analyzed through the Voigt-Reuss-Hill approximation, while Poisson's ratio, shear modulus, and Young's modulus for both 2D and 3D anisotropic structures were obtained using Pythonbased tools. Furthermore, to investigate the thermodynamic stability and dynamic behavior of the system, molecular dynamics simulations were carried out in the NPT ensemble at 298 K using a timestep of 1.0 fs, with the production run extended to 50 ps after equilibration.37

2.2. SCAPS-1D-based performance assessment for solar cell

The photovoltaic performance of D₂CeX₆-based solar cell devices was simulated using SCAPS-1D software (Version 3.3.09),³⁸ while the corresponding device architecture was designed with Adobe Illustrator (Version 2020). In this study, the SCAPS-1D-based performance evaluation of D₂CeX₆ double perovskites was carried out by numerically solving the fundamental semiconductor equations that describe charge carrier generation, distribution, and transport within the device.³⁹ These include the Poisson equation,

$$\frac{\partial^2 \varphi}{\partial x^2} = \frac{q}{\varepsilon} (n(x) - p(x) + N_{\mathsf{A}}(x) - N_{\mathsf{D}}(x) - p_{\mathsf{t}}(x) - N_{\mathsf{t}}(x)) \tag{1}$$

which describes the potential distribution, the continuity equations for holes and electrons, 40

$$\frac{\partial j_{\rm p}}{\partial_{\rm x}} = -q \left(R_{\rm p} - G_{\rm p} + \frac{\partial_{\rm p}}{\partial_{\rm t}} \right) \tag{2}$$

$$\frac{\partial j_{\rm n}}{\partial_{\rm x}} = q \left(R_{\rm n} - G_{\rm n} + \frac{\partial_{\rm n}}{\partial_{\rm t}} \right) \tag{3}$$

Drift-diffusion current equations,

$$J_{p} = -qD_{p}\frac{\partial p}{\partial x} + q\mu_{p}p\frac{\partial \varphi}{\partial x} \tag{4}$$

$$J_{\rm n} = q D_{\rm n} \frac{\partial n}{\partial x} + q \mu_{\rm n} n \frac{\partial \varphi}{\partial x} \tag{5}$$

Here, ε is the dielectric constant, n and p are carrier concentrations, N and $N_{\rm A}$ are donor and acceptor densities, D p/n are diffusion coefficients, and μ p/n are carrier mobilities. The program calculates key performance parameters such as open-circuit voltage ($V_{\rm OC}$),

$$V_{\rm OC} = \frac{n \times k \times T}{q} \ln \left(\frac{I_{\rm ph}}{I_{\rm O}} + 1 \right) \tag{6}$$

Short-circuit current density (J_{SC}) ,

$$J_{\rm SC} = q \times A \times G \times \eta \tag{7}$$

Fill factor (FF),

$$FF = \frac{P_{\text{Max}}}{P_{\text{in}}} = \frac{I_{\text{Max}} \times V_{\text{Max}}}{V_{\text{OC}} \times I_{\text{SC}}}$$
(8)

Power conversion efficiency (PCE)

$$PCE = \frac{I_{SC} \times FF \times V_{OC}}{P_{in}}$$
 (9)

where $P_{\rm in}$ is the incident solar power, n is the diode ideality factor, and k is Boltzmann's constant, ⁴¹ T is temperature, I_{ph} is the photo-generated current, I_0 is the reverse saturation current, A is the active area, G is the irradiance, and η is the quantum efficiency. By incorporating experimentally or theoretically derived material properties, including band gap, electron affinity, relative permittivity, and carrier mobility into these equations, SCAPS-1D enables the evaluation of recombination mechanisms, carrier transport, and defect tolerance. This facilitates the identification of optimal halide (Cl, Br) and D-site cation (Ga, In, Tl) combinations with balanced optoelectronic properties, reduced recombination losses, and enhanced efficiency, thus offering a cost-effective and accurate approach to designing high-efficiency, lead-free double perovskite solar cells before fabrication. These device-level simulations were complemented by first-principles calculations, which provided detailed insights into the structural, electronic, optical, mechanical, phonon, and population properties of vacancyordered D₂CeX₆ perovskites. The integration of DFT analyses with SCAPS-1D simulations delivers a comprehensive evaluation of the material's potential in optoelectronic and photovoltaic applications, offering valuable guidance for the development of lead-free vacancy-ordered perovskite solar cells.

3 Results and discussion

3.1. X-ray diffraction (XRD) analysis

X-ray diffraction (XRD) is a fundamental characterization technique used to determine the crystal structure, phase purity, and lattice parameters of materials by measuring the angles and

intensities of diffracted X-rays. 42 This analysis is essential as it enables precise identification of crystalline phases, detection of structural distortions, estimation of unit cell dimensions, and confirmation of material synthesis without destructive testing. For perovskites, XRD provides critical insights into how atomic arrangements vary with different cations or anions, reveals lattice strain and defects, and helps explain changes in electronic, optical, and mechanical properties. 43 Such structural information is indispensable for correlating crystal geometry with performance in photovoltaics, optoelectronics, and thermoelectric, thereby ensuring consistency between theoretical predictions and experimental results in optimized device design.44 The X-ray diffraction (XRD) patterns of vacancyordered halide perovskites D₂CeX₆ (D = Ga, In, Tl; X = Cl, Br) reveal sharp and intense peaks, confirming their high crystallinity and phase purity. The measured 2θ values show that Ga₂CeCl₆ has its first diffraction peak at 13.2° and the last prominent peak at 44.05°, while Ga₂CeBr₆ ranges from 12.7° to

42.3°, as shown in Fig. 1(a). Similarly, In₂CeCl₆ spans from 13.2° to 44.05°, and In₂CeBr₆ from 12.7° to 42.3°, as shown in Fig. 1(b). Tl₂CeCl₆ exhibits a slightly higher starting angle of 13.3° and extends to 44.4°, whereas Tl₂CeBr₆ lies between 12.7° and 42.6°, as shown in Fig. 1(c). The consistent observation that bromide-based compounds have their diffraction peaks shifted toward lower angles compared to chlorides indicates lattice expansion, attributable to the larger ionic radius of Br⁻, which increases the interplanar spacing (d-spacing) according to Bragg's law. 45 Differences among Ga, In, and Tl compositions, particularly at both lowest and highest 2θ positions, reflect the influence of A-site cation size, mass, and bonding characteristics on unit cell parameters. This variation in lattice dimensions directly impacts electronic band structure, phonon dynamics, and defect formation tendencies, which are crucial for optoelectronic and thermoelectric applications. The importance of XRD here lies not only in confirming the successful synthesis of the targeted phase but also in quantitatively linking structural

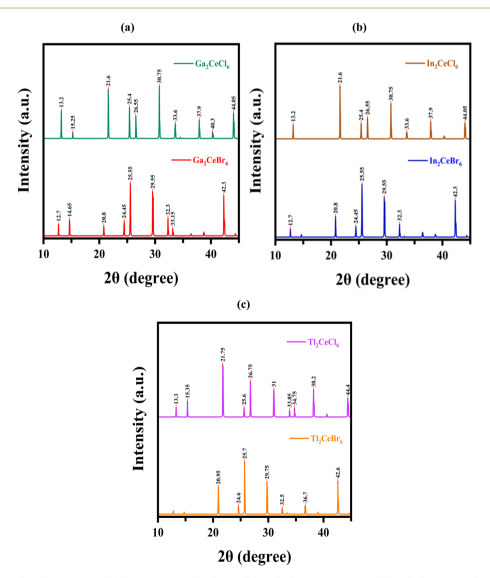


Fig. 1 X-ray diffraction (XRD) patterns of (a) Ga_2CeCl_6 and Ga_2CeBr_6 , (b) In_2CeCl_6 and In_2CeBr_6 , (c) Tl_2CeCl_6 and Tl_2CeBr_6 vacancy-ordered perovskites.

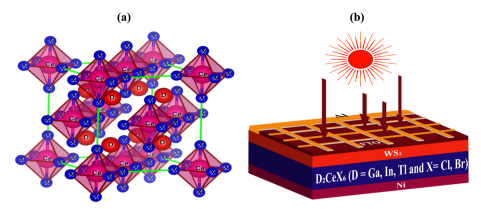


Fig. 2 The (a) crystal structure, (b) proposed solar cell device structure of D₂CeX₆ vacancy-ordered perovskites.

parameters to physical properties, thereby enabling predictive modeling in density functional theory (DFT) simulations and guiding experimental optimization for photovoltaic device integration.

3.2. Structural characteristics of vacancy-ordered perovskites

The D_2CeX_6 (D = Ga, In, Tl; X = Cl, Br) double perovskites adopt a highly symmetric cubic crystal structure, typically belonging to the space group $Fm\bar{3}m$ (#225). Within this framework, the Ce^{4+} cations occupy the octahedral coordination sites, where each Ce atom is surrounded by six halide ions (Cl⁻ or Br⁻), forming corner-sharing CeX_6 octahedra that serve as the fundamental building blocks of the lattice. The D-site cations (Ga, In, Tl) are accommodated in the interstitial cavities, stabilizing the perovskite structure through electrostatic interactions. In terms of crystallographic positions, the D-site atoms reside at the face-centered Wyckoff position 3c (0, 0.5, 0.5), the Ce atom is located at the corner site with Wyckoff position 1a (0, 0, 0), and the halide atoms occupy the body-centered Wyckoff position 1b (0.5, 0.5, 0.5).

This ordered arrangement reflects the vacancy-ordered nature of the perovskite, where one of the B-site cations in the conventional ABX₃ lattice is replaced by a structural vacancy,

giving rise to the distinctive A₂BX₆ stoichiometry. As illustrated in Fig. 2(a), the atomic configurations of analogous compounds such as Ga₂CeCl₆, Ga₂CeBr₆, In₂CeCl₆, In₂CeBr₆, Tl₂CeCl₆, and Tl₂CeBr₆ display a similar cubic symmetry and site occupancy pattern, highlighting the robustness of this structural motif across different compositions. The high degree of symmetry and well-defined cation coordination in these materials play a crucial role in determining their electronic structure, phonon dynamics, and defect tolerance properties that are directly linked to their optoelectronic and photovoltaic performance.

The optimized lattice constants of these vacancy-ordered perovskites are 7.071 Å to 7.784 Å, as shown in Table 1, primarily governed by the ionic radii of both the D-site cations and halide anions. Predictably, substitution of Cl⁻ with the larger Br⁻ ion results in a noticeable increase in lattice parameters. For instance, Ga₂CeCl₆ possesses a lattice constant of 7.410 Å, which expands to 7.784 Å for Ga₂CeBr₆. A similar lattice expansion trend is observed for the In- and Tl-based systems. The corresponding unit cell volumes reflect this trend, increasing with the incorporation of the larger halide anion. The highest unit cell volume of 333.524 Å³ is recorded for Ga₂CeBr₆, while In₂CeBr₆ and Tl₂CeBr₆ both display volumes of 249.999 Å³, indicating a consistent cubic framework for these bromide compounds. Furthermore, the calculated volumes obtained using GGA-PBE and HSE06 functionals show

 $\textbf{Table 1} \quad \text{The energy band gap, lattice constants, unit cell volume, density, and formation energy of D_2CeX$_6 vacancy-ordered perovskites}$

		GGA-PBE			HSE-06			
Ref.	Compounds	00		Unit cell volume, $V(\mathring{A}^3)$	Density, g cm ⁻³		Unit cell volume, $V(\mathring{A}^3)$	Formation energy, $\Delta E_{\rm f}$ (eV per atom)
This work	Ga ₂ CeCl ₆	1.733	7.410	287.748	2.841	1.776	287.775	-3.597
	Ga ₂ CeBr ₆	1.276	7.784	333.524	3.778	2.843	333.528	-3.227
	In ₂ CeCl ₆	1.555	7.473	295.156	3.277	2.261	249.999	-3.571
	In ₂ CeBr ₆	0.859	7.071	249.999	5.640	2.170	249.999	-3.064
	Tl_2CeCl_6	1.755	7.503	298.718	4.233	3.632	298.732	-3.699
	Tl ₂ CeBr ₆	1.364	7.071	249.999	6.830	1.418	249.999	-3.178
48	K ₂ CeCl ₆	1.742	7.522	300.945	2.378	_	_	-3.714
	Rb ₂ CeCl ₆	1.748	7.619	312.774	2.780	_	_	-3.720
	Cs_2CeCl_6	1.807	7.863	343.881	2.987	_	_	-3.746
	Fr_2CeCl_6	1.830	8.111	377.413	3.514	_	_	-3.705

negligible differences, indicating reliable structural optimization. Notably, the calculated densities are 2.841 g cm⁻³ for Ga_2CeCl_6 , 3.277 g cm⁻³ for In_2CeCl_6 , and 4.233 g cm⁻³ for Tl₂CeCl₆, while Ga₂CeBr₆, In₂CeBr₆, and Tl₂CeBr₆ exhibit 3.778, 5.640, and 6.830 g cm⁻³, respectively. These variations clearly demonstrate that both the nature of the D₂-site cation (Ga, In, Tl) and the X₆ anion (Cl, Br) strongly influence the structural packing and overall density, with heavier Tl and Br substitutions yielding denser frameworks. In addition to structural analysis, the electronic band gaps were calculated using both GGA-PBE and HSE06 functionals (Table 1). As anticipated, the HSE06 hybrid functional produced consistently higher band gaps owing to its more accurate treatment of exchange-correlation effects. For instance, Ga₂CeCl₆ exhibited an increase from 1.733 eV (GGA-PBE) to 1.776 eV (HSE06), with similar enhancements observed across all compounds. The bromide analogues displayed the lowest gaps, with In₂CeBr₆ showing the smallest values of 0.859 eV (GGA-PBE) and 2.170 eV (HSE06), consistent with the established trend of band gap narrowing as the halide ionic radius increases.

To further evaluate the thermodynamic stability of these perovskites, their formation energy ($\Delta E_{\rm f}$) was computed,⁴⁷ as in Table 1.

$$\Delta E_{\rm f} = E_{\rm tot} (A_2 \text{CeX}_6) - 2E(A) - E(\text{Ce}) - 6E(X)$$
 (10)

Negative formation energy indicates that a compound is thermodynamically stable and can form spontaneously, with more negative values reflecting stronger bonding and greater intrinsic stability. It's a key factor in assessing whether a material is synthesizable and suitable for applications like photovoltaics or catalysis. All compounds exhibit negative formation energy, confirming their thermodynamic favorability. Tl_2CeCl_6 displayed the most negative formation energy value of -3.699 eV per atom, indicating it as the most stable composition among the series. In contrast, Ga_2CeBr_6 showed the least negative value (-3.227 eV per atom), likely due to the combined effect of the lighter Ga cation and the larger, less tightly bound Br^- anion.

The calculated structural and electronic properties of Ga_2CeX_6 , In_2CeX_6 , and Tl_2CeX_6 (X = Cl, Br) exhibit close similarities with the reported results of K_2CeCl_6 , Rb_2CeCl_6 , Cs_2-CeCl_6 , and Fr_2CeCl_6 . In both cases, the GGA-PBE predicted band gaps are within the same semiconducting range. At the

Table 2 The variation of bond length within the atom of D_2CeX_6 (where D=Ga, In, Tl, and X=Cl, Br)

	Bond length, L (Å)						
Compounds	$L_{\mathrm{D-X}}$	$L_{\rm Ce-X}$	$L_{ m Ce-D}$				
Ga ₂ CeCl ₆	3.705	2.626	4.538				
Ga ₂ CeBr ₆	3.892	2.785	4.766				
In ₂ CeCl ₆	3.736	2.629	4.576				
In ₂ CeBr ₆	3.540	2.682	4.330				
Tl ₂ CeCl ₆	3.751	2.631	4.594				
Tl_2CeBr_6	3.540	2.682	4.330				

same time, the lattice constants and unit cell volumes follow a systematic increase with the incorporation of larger A-site cations. Furthermore, the negative formation enthalpies obtained in this work are comparable to those in ref. 48, confirming the thermodynamic stability of these Ce-based halide double perovskites.

3.2.1. Bond length variation analysis. The variation of bond lengths within D_2CeX_6 (D = Ga, In, Tl; X = Cl, Br) is important because it provides key insights into the structural stability, electronic interactions, and overall behavior of these vacancyordered double perovskites. Shorter Ce-X bonds indicate strong Ce-halide interactions, which help stabilize the crystal lattice, while variations in D-X and Ce-D distances reflect the influence of different cations and halides on the lattice geometry. These bond lengths directly affect orbital overlap, band structure, and optoelectronic properties, making them critical for accurately predicting electronic, optical, and mechanical behavior through computational methods. Overall, analyzing bond length variations connects atomic-scale geometry with material performance, providing a foundation for understanding and optimizing D₂CeX₆ compounds for applications such as solar cells and other optoelectronic devices.

Table 2 presents the bond length variations in D_2CeX_6 (D = Ga, In, Tl; X = Cl, Br), considering D-X, Ce-X, and Ce-D interactions. The D-X bonds range from 3.705 to 3.892 Å, with chlorides showing slightly shorter values (3.705 to 3.751 Å) than bromides, except for In₂CeBr₆ and Tl₂CeBr₆, where the D-X length decreases to 3.540 Å. The Ce-X bonds are consistently shorter (2.626 to 2.785 Å), indicating stronger Ce-X interactions compared to D-X; as expected, Ce-Br bonds (≈ 2.682 to 2.785 Å) are longer than Ce-Cl bonds (≈2.626 to 2.631 Å) owing to the larger ionic radius of Br-. The Ce-D bonds are the longest, ranging between 4.330 and 4.766 Å, with the maximum in Ga₂CeBr₆ and the minimum in In₂CeBr₆ and Tl₂CeBr₆. These variations reveal that the bond lengths are sensitive to both the halide ion and the D-site cation, where bromides generally favor longer separations than chlorides, and the shorter Ce-X bonds suggest stronger Ce-centered bonding, which may play a decisive role in stabilizing the crystal structure and influencing the electronic interactions within these vacancy-ordered double perovskites.

3.2.2. Geometric stability parameters. For vacancy-ordered perovskite structures, the stability and structural distortion are often evaluated using the Goldschmidt tolerance factor (t) and the octahedral factor (μ) . It is defined as:

$$t = \frac{(r_{\rm A} + r_{\rm X})}{\sqrt{2}(r_{\rm Ce} + r_{\rm X})} \tag{11}$$

$$\mu = \frac{r_{\rm B}}{r_{\rm X}} \tag{12}$$

The tolerance factor generally falls within the range of 0.8 to 1.0, where values close to 1.0 represent an ideal cubic configuration with minimal lattice distortion. Meanwhile, the octahedral factor typically lies between 0.41 and 0.90, indicating that the B-site cation is appropriately sized to fit within the BX₆

Table 3 Tolerance factor and octahedral factors variation in D_2CeX_6 vacancy-ordered perovskites

Materials	r_{A}	r_{Ce}	$r_{ m X}$	Tolerance factor (t)	Octahedral factor (μ)
Ga ₂ CeCl ₆	1.3	0.97	1.81	0.791	0.5359
Ga ₂ CeBr ₆	1.3	0.97	1.96	0.7867	0.4949
In ₂ CeCl ₆	1.44	0.97	1.81	0.8267	0.5359
In ₂ CeBr ₆	1.44	0.97	1.96	0.8205	0.4949
Tl_2CeCl_6	1.59	0.97	1.81	0.8648	0.5359
${ m Tl_2CeBr_6}$	1.59	0.97	1.96	0.8567	0.4949

octahedron. ^{49,50} When both factors fall within these ranges, the perovskite framework achieves a balanced geometric arrangement, leading to enhanced structural stability. Deviations outside these limits may result in lattice distortions, octahedral tilting, or complete structural instability, highlighting the critical role of ionic size matching in the design of stable vacancy-ordered perovskites.

The calculated tolerance factors, as summarized in Table 2, are 0.7867, 0.7867, 0.8267, 0.8205, 0.8648, and 0.8567 for Ga_2CeCl_6 ,

Ga₂CeBr₆, In₂CeCl₆, In₂CeBr₆, Tl₂CeCl₆, and Tl₂CeBr₆, respectively. All compounds exhibit values within the structural stability range, except Ga₂CeCl₆ and Ga₂CeBr₆, which fall slightly below the lower limit but remain close to the stability threshold. This indicates that most of these halide double perovskites possess geometrically favorable ionic packing. At the same time, the Ga-based variants, though marginally less ideal, are still likely to form stable frameworks with only minor octahedral tilting or distortion. The calculated octahedral factors, presented in Table 3, are 0.5359, 0.4949, 0.5359, 0.4949, 0.5359, and 0.4949 for Ga₂CeCl₆, Ga₂CeBr₆, In₂-CeCl₆, In₂CeBr₆, Tl₂CeCl₆, and Tl₂CeBr₆, respectively. All values fall within the established structural stability window (0.41 to 0.90), confirming that the B-site cations are appropriately accommodated within the BX6 octahedral framework. This suggests that these compounds are geometrically favorable for forming stable vacancyordered perovskite structures.

3.3. Electronic properties

The electronic properties of materials refer to how electrons behave within a solid, influencing key characteristics like

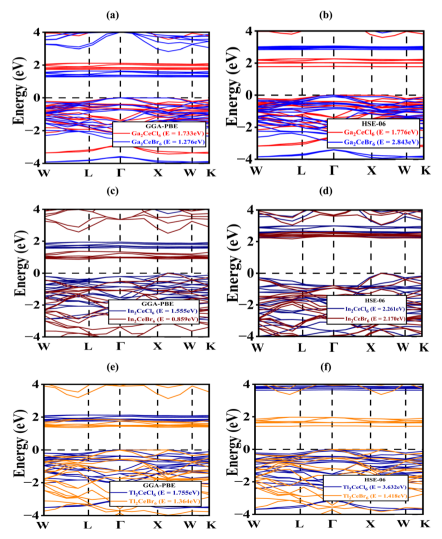


Fig. 3 Energy band gap of (a and b) Ga_2CeX_6 , (c and d) In_2CeX_6 and (e and f) Tl_2CeX_6 double perovskite materials using GGA-PBE and HSE06 function.

electrical conductivity, resistivity, carrier mobility, band structure, and band gap.51,52 These parameters determine whether a material behaves as a metal, semiconductor, or insulator and directly influence its suitability for electronic, optoelectronic, and energy-related applications. By analyzing electronic features, particularly the band structure, band gap nature (direct or indirect), and partial density of states, researchers can assess whether a specific D₂CeX₆ compound is more appropriate for light-emitting applications, such as LEDs, or for lightharvesting and charge-separating functions, such as photovoltaics. The band structure, a cornerstone of solid-state physics, illustrates the variation of electron energy with momentum in a crystalline material.53 For vacancy-ordered double perovskites D₂CeX₆, knowledge of the band dispersion and gap is essential, as these parameters dictate their electronic and optoelectronic performance, influencing applications in solar cells. Fig. 3

presents the calculated band structures using two computational approaches, GGA-PBE (Fig. 3(a, c and e)) and HSE06 (Fig. 3(b, d and f)), where the energy (E- E_X , in eV) is plotted along high-symmetry Brillouin zone paths (W-L- Γ -X-W-K), with the Fermi level fixed at 0 eV.

The band gap forms between the valence-band maximum (VBM) and conduction-band minimum (CBM). As shown in Fig. 3(a-f), DFT reveals a clear dependence on both the D-site cation (D = Ga, In, Tl) and halide (X = Cl, Br). In Fig. 3(a, c and e), within GGA-PBE, the calculated gaps are: Ga₂CeCl₆ of 1.733 eV, Ga₂CeBr₆ of 1.276 eV, In₂CeCl₆ of 1.555 eV, In₂CeBr₆ of 0.859 eV, Tl₂CeCl₆ of 1.755 eV, and Tl₂CeBr₆ of 1.364 eV, placing all but In₂CeBr₆ (too narrow) within or near the 1.1 to 1.8 eV single-junction "sweet spot." A clear anion dependence is observed: replacing Cl with Br consistently reduces the band gap due to the higher energy of Br-4p orbitals, which raises the

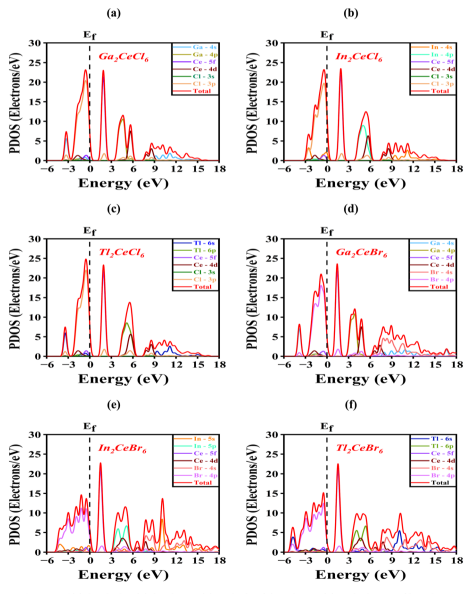


Fig. 4 The partial density of states of (a) Ga_2CeCl_6 , (b) Ga_2CeBr_6 , (c) In_2CeCl_6 , (d) In_2CeBr_6 , (e) Tl_2CeCl_6 , and (f) Tl_2CeBr_6 vacancy-ordered double perovskites.

VBM and narrows the gap. As expected, HSE06 increases the gaps because of its improved treatment of exchange: Ga_2CeCl_6 of 1.776 eV, Ga_2CeBr_6 of 2.843 eV, In_2CeCl_6 of 2.261 eV, In_2CeBr_6 of 2.170 eV, Tl_2CeCl_6 of 3.632 eV, and Tl_2CeBr_6 of 1.418 eV, as shown in Fig. 3(b, d and f). Under HSE06, Tl_2CeBr_6 (1.418 eV) and Ga_2CeCl_6 (1.776 eV) remain suitable for single-junction devices, whereas the wider-gap Ga_2CeBr_6 , In_2CeCl_6 , In_2CeBr_6 , and Tl_2CeCl_6 (>2.0 eV) are better suited as top cells in tandem architectures. In contrast, In_2CeBr_6 at the PBE level (0.859 eV) exemplifies the "narrow-gap" regime (<1.1 eV), which enhances IR absorption but suffers from higher thermalization losses.

The Partial Density of States (PDOS) is a powerful tool in materials science for identifying the orbital and atomic contributions to the electronic structure of compounds.54 Such analvsis provides key insights into the electronic behavior relevant for solar cells and light-emitting devices. For the D2CeX6 vacancy-ordered perovskites, the valence band (VB) extends from -6 to 0 eV, while the conduction band (CB) spans from 0 to +18 eV, as illustrated in Fig. 4. In the valence band region, the dominant contributions originate from halogen p-states hybridized with the D-site cation s-states: Cl-3p and Ga-4s in Ga₂CeCl₆ (Fig. 4(a)), Cl-3p and In-5s in In₂CeCl₆ (Fig. 4(b)), Cl-3p and Tl-6s in Tl₂CeCl₆ (Fig. 4(c)), Br-4p and Ga-4s in Ga₂CeBr₆ (Fig. 4(d)), Br-4p and In-5s in In₂CeBr₆ (Fig. 4(e)), and Br-4p and Tl-6s in Tl₂CeBr₆ (Fig. 4(f)). In all cases, Ce-4d states provide a minor but noticeable contribution to the VB. In the conduction band region, the primary states arise from Ce-5f orbitals, strongly mixed with the p-states of the D-site cations: Ce-5f and Ga-4p in Ga₂CeCl₆ (Fig. 4(a)), Ce-5f and In-5p in In₂CeCl₆ (Fig. 4(b)), Ce-5f and Tl-6p in Tl₂CeCl₆ (Fig. 4(c)), Ce-5f and Ga-4p in Ga_2CeBr_6 (Fig. 4(d)), Ce-5f and In-5p in In_2CeBr_6 (Fig. 4(e)), and Ce-5f and Tl-6p in Tl₂CeBr₆ (Fig. 4(f)).

Notably, localized peaks appear just above the Fermi level $(E_{\rm X})$, confirming that the conduction-band minimum (CBM) is dominated by Ce-derived states. Overall, the PDOS indicates that the VBM is mainly governed by halogen p-states hybridized with D-site s-states, while the CBM is strongly influenced by Ce-5f states, a characteristic feature of these vacancy-ordered double perovskites. It is worth noting that a slight mismatch between the band gap values obtained from the band structure (Fig. 3) and the PDOS (Fig. 4) is observed. This minor deviation originates from the intrinsic limitations of DFT within the PBE framework, differences in k-point sampling, and the broadening applied in DOS plotting. Nevertheless, all data have been carefully rechecked, and the overall electronic trends remain fully consistent, confirming the reliability of the presented results.

3.4. Homo-lumo analysis

HOMO-LUMO analysis investigates the energy difference between the Highest Occupied Molecular Orbital (HOMO) and the Lowest Unoccupied Molecular Orbital (LUMO), offering key insights into a material's chemical reactivity, stability, and electronic characteristics. ^{56,57} The HOMO corresponds to the molecule's ability to donate electrons, while the LUMO reflects its capacity to accept electrons. A smaller HOMO-LUMO energy

Table 4 Calculated HOMO-LUMO energies and electronic characteristics of D_2CeX_6 double perovskite compounds

Compounds	Type	N	s	Eigenvalue	Field
Ga ₂ CeCl ₆	НОМО	111	+	-0.263252	Yes
	LUMO	112	+	-0.219414	Yes
Ga ₂ CeBr ₆	HOMO	165	+	-0.244130	Yes
	LUMO	166	+	-0.212826	Yes
In_2CeCl_6	HOMO	129	+	-0.266703	Yes
	LUMO	130	+	-0.222910	Yes
In_2CeBr_6	HOMO	183	+	-0.227664	Yes
	LUMO	184	+	-0.185877	Yes
Tl_2CeCl_6	HOMO	161	+	-0.267885	Yes
	LUMO	162	+	-0.224097	Yes
Tl_2CeBr_6	HOMO	215	+	-0.229307	Yes
	LUMO	216	+	-0.186823	Yes

gap signifies higher chemical reactivity, lower stability, and improved electrical conductivity, whereas a larger gap indicates greater stability and reduced reactivity. Thus, this analysis is fundamental for understanding charge transfer, optical transitions, and bandgap behavior in both molecular and solid-state systems, serving as an essential tool for designing materials with tailored electronic and optoelectronic properties.⁵⁸

Table 4 presents the HOMO–LUMO energy values of six halide double perovskite compounds, namely Ga_2CeCl_6 , Ga_2CeBr_6 , In_2CeBr_6 , In_2CeBr_6 , Tl_2CeCl_6 , and Tl_2CeBr_6 , which help evaluate their electronic properties and chemical reactivity. The eigenvalues of both HOMO and LUMO orbitals are negative, indicating bound electronic states, while the difference between them represents the HOMO–LUMO energy gap (ΔE) , which reflects the ease of electronic excitation. Among these, bromine-based compounds (Br_6) exhibit slightly smaller energy gaps than their chlorine-based counterparts (Cl_6) , implying that Brbased systems are more reactive and less stable, with better potential for charge transfer and optical activity.

Conversely, Cl-based compounds show larger gaps, indicating greater chemical stability and lower electronic polarizability. Thus, the trend reveals that the HOMO–LUMO gap decreases in the order ${\rm Cl_6} > {\rm Br_6}$, suggesting that substitution of Cl by Br enhances the electronic softness and conductivity of the materials, which is beneficial for optoelectronic and semiconductor applications.

Fig. 5 presents the spatial distributions of the HOMO and LUMO for Ga_2CeCl_6 and Ga_2CeBr_6 double perovskites, illustrating their electronic charge density and orbital hybridization characteristics. In both compounds, the HOMO states are primarily localized around the halogen atoms (Cl or Br) and partially extend toward the Ce atoms, indicating that the valence band maximum (VBM) mainly originates from the porbitals of the halogens with minor Ce contributions.

Conversely, the LUMO orbitals are more delocalized over the Ce and Ga atoms, suggesting that the conduction band minimum (CBM) is predominantly governed by Ce 4f and Ga 4s/4p orbitals. This behavior highlights a charge transfer from the halogen sites to the metal centers during electronic excitation. Comparatively, Ga_2CeCl_6 exhibits a more compact and

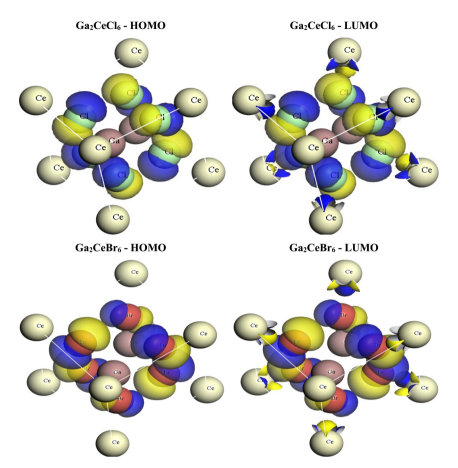


Fig. 5 HOMO and LUMO charge density distributions of Ga₂CeCl₆ and Ga₂CeBr₆ double perovskites.

symmetric orbital distribution owing to the smaller ionic radius and stronger bonding of Cl atoms, whereas Ga₂CeBr₆ shows broader and slightly distorted orbital features due to the weaker Ga–Br and Ce–Br interactions associated with the larger Br ions. Overall, both materials demonstrate a similar electronic transition pattern—from halogen-dominated HOMO to metalcentered LUMO—but Ga₂CeCl₆ displays stronger orbital overlap and higher electronic localization, resulting in a wider band gap and enhanced bonding strength. In contrast, Ga₂CeBr₆ exhibits narrower band gaps and improved charge transfer characteristics, making it potentially more favorable for optoelectronic applications.

Fig. 6 illustrates the HOMO and LUMO charge density distributions of In₂CeCl₆ and In₂CeBr₆ double perovskites, providing a detailed visualization of their electronic interactions, orbital hybridization, and charge transfer mechanisms. In both compounds, the HOMO orbitals are primarily localized around the halogen atoms (Cl or Br), with noticeable overlap extending toward the Ce atoms. This indicates that the VBM mainly originates from the halogen p-orbitals, slightly hybridized with Ce f-orbitals, suggesting a strong halogen–Ce bonding contribution in the occupied states. Conversely, the LUMO orbitals are predominantly distributed over the Ce and In atoms, revealing that the CBM is mainly derived from Ce 4f and In 5s/5p orbitals. This spatial separation between the HOMO and LUMO charge densities confirms a halogen-to-metal charge

transfer (HMCT) mechanism during electronic excitation, a typical feature of Ce-based halide perovskites. Comparatively, In₂CeCl₆ exhibits more compact and symmetric charge density lobes around the Ce-Cl-In framework, indicating stronger covalent bonding and tighter electronic confinement. This results in a larger band gap and enhanced structural rigidity. The smaller ionic radius and higher electronegativity of Cl promote stronger orbital overlap and increased charge localization. In contrast, In₂CeBr₆ displays more extended and diffuse charge distributions around the Br atoms, reflecting weaker In-Br and Ce-Br interactions due to the larger atomic radius and lower electronegativity of Br. This delocalization facilitates higher electron mobility and reduces the band gap, making the bromide compound more suitable for optoelectronic and photovoltaic applications that demand efficient charge transport. Overall, while both In₂CeCl₆ and In₂CeBr₆ exhibit a similar electronic transition pattern (halogen $p \rightarrow Ce/$ In f/s), In₂CeCl₆ is characterized by stronger bonding and greater electronic localization—ideal for structural and electronic stability—whereas In₂CeBr₆ shows enhanced delocalization and charge transport properties, making it a promising candidate for devices requiring improved conductivity and light absorption efficiency.

Fig. 7 presents the spatial charge density distributions of the frontier molecular orbitals HOMO and LUMO for Tl₂CeCl₆ and Tl₂CeBr₆, offering crucial insight into their electronic behavior

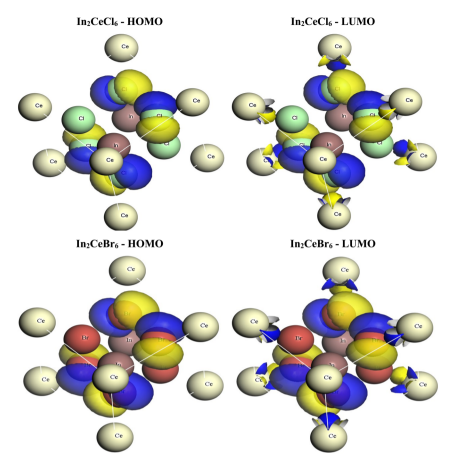


Fig. 6 HOMO and LUMO charge density distributions of In₂CeCl₆ and In₂CeBr₆ double perovskites.

and bonding nature. In Tl_2CeCl_6 , the HOMO (top left) shows electron density predominantly concentrated around the Cl and Ce atoms, suggesting strong Cl-p and Ce-f orbital hybridization within the valence band. This localization near the halide atoms indicates that the valence band maximum (VBM) is largely halogen-derived, with only a minor contribution from Ce orbitals.

Conversely, in the LUMO (top right), the charge density becomes more delocalized toward Ce and Tl centers, signifying that the conduction band minimum (CBM) arises mainly from Ce-d and Tl-p orbitals. This spatial transition of electron density from halogen to metal centers across HOMO-LUMO levels highlight a characteristic charge-transfer excitation process within the Tl-Ce-Cl framework. For Tl₂CeBr₆ (bottom row), a similar orbital distribution pattern is observed, yet with notable differences in the extent and nature of charge delocalization. The HOMO charge density around Br and Ce atoms appears more diffused and extensive than in Tl₂CeCl₆, reflecting the larger size and lower electronegativity of Br compared to Cl. This results in weaker bonding and reduced orbital confinement, which can lower the band gap energy. In the LUMO of Tl₂CeBr₆, electron density remains focused on Ce and Tl sites but extends more toward the Br orbitals, implying enhanced orbital overlap and electronic coupling between the halide and metal centers. Overall, both compounds share similar electronic topologies, where the valence band is dominated by halide-Ce interactions and the conduction band by Ce–Tl hybridization. However, the Br-based compound exhibits greater orbital delocalization, smoother charge transitions, and stronger metal-halide interaction continuity, all of which contribute to narrower band gaps and potentially improved optical absorption and charge transport. These electronic distinctions underline the role of halide substitution in tuning the band-edge characteristics and optoelectronic performance of Tl–Ce halide perovskites.

3.5. Charge density mapping

Charge density mapping is a powerful technique that visualizes the distribution of electronic charge within a crystal lattice, providing direct insight into the nature of chemical bonding and electronic interactions between constituent atoms. 59,60 For D_2CeX_6 vacancy-ordered double perovskites, charge density analysis is particularly important as it reveals bonding characteristics, charge localization around different atomic sites, and potential pathways for defect formation, all of which critically influence their optical, electronic, and structural properties. Fig. 8 elucidates the charge density maps of D_2CeX_6 (D = Ga, In, Tl; X = Cl, Br) double perovskites, where electron localization is visualized using a color scale. In all compounds, significant charge accumulation is observed at the Ce sites and around the

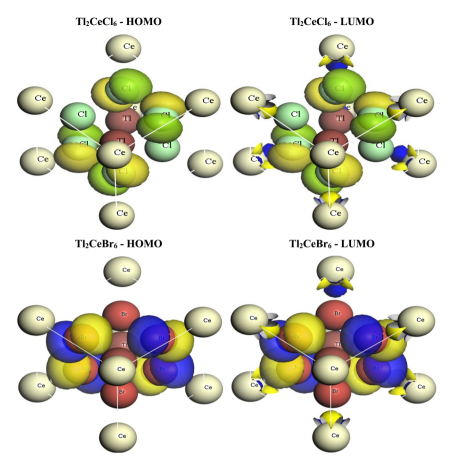


Fig. 7 HOMO and LUMO charge density distributions of Tl₂CeCl₆ and Tl₂CeBr₆ double perovskites

halogen atoms, with Cl exhibiting stronger localization than Br due to its higher electronegativity. For Tl₂CeCl₆, the maximum charge density reaches 1.61 \times 10 1 a.u. with a near-zero minimum of -3.97 \times 10 $^{-3}$ a.u., indicating strong electron localization at the Cl sites and predominantly ionic Tl–Cl and Ce–Cl interactions.

 Tl_2CeBr_6 shows a slightly lower maximum of 1.61 \times 10¹ a.u. with a minimum of 1.29 \times 10⁻² a.u., reflecting substantial accumulation around Br but limited interstitial density. In2- $CeCl_6$ exhibits a lower maximum of 1.09×10^1 a.u. and a very small minimum of 7.61 \times 10⁻⁴ a.u., while In₂CeBr₆ shows a comparable maximum with a slightly higher minimum (1.36 \times 10⁻² a.u.), suggesting more diffuse electron density around Br. The Ga-based systems display the strongest localization: Ga₂CeCl₆ reaches the highest maximum among all studied compounds at 3.95×10^{1} a.u. with a near-zero minimum of 2.92 \times 10⁻³ a.u., while Ga₂CeBr₆ retains similarly high values (3.94 \times 10^{1} a.u.; minimum 3.76×10^{-3} a.u.). Overall, the Ga-containing systems demonstrate the highest peak charge densities, whereas the In-based compounds show the lowest. A systematic trend emerges with A-site substitution, where electron density becomes more diffuse from Ga \rightarrow In \rightarrow Tl, consistent with increasing ionic size and reduced covalent interactions. Furthermore, Cl-based perovskites consistently exhibit stronger electron localization than their Br-based counterparts. These

results highlight the direct connection between charge density features, bonding character, and the electronic stability of the D_2CeX_6 family. Notable studies supporting these findings include work by Faizan *et al.*, Rifat *et al.*, and Saikot *et al.*, which collectively highlight the role of charge density analysis in advancing halide double perovskite research for next-generation optoelectronic applications.^{7,48,61}

3.6. Optical properties

The optical properties of a material define how it interacts with incident light, encompassing absorption, transmission, reflection, and emission processes across the ultraviolet (UV), visible, and infrared (IR) regions of the spectrum. ⁶² These characteristics are directly linked to the electronic structure and band gap, making them critical for determining a material's potential in optoelectronic applications. For double perovskites, analyzing the absorption coefficient, reflectivity, refractive index, dielectric function, and optical conductivity provides valuable insights into their suitability for device integration.

In this study, Fig. 9 presents the calculated optical properties of the D_2CeX_6 (D=Ga, In, Tl; X=Cl, Br) vacancy-ordered double perovskite compounds. The refractive indices of conventional double perovskites such as $(Cs_2, K_2, Rb_2)PbCl_6$ in the visible and ultraviolet regions offer significant advantages for solar cell applications, as their wide dielectric constants

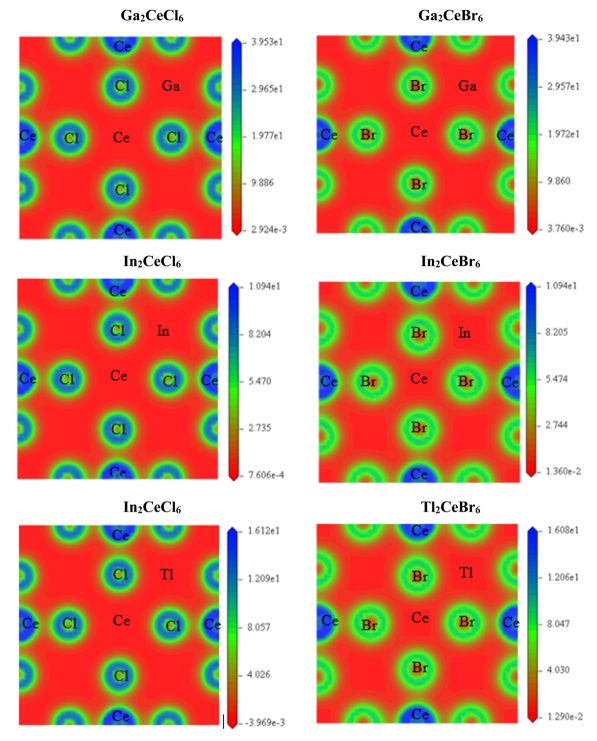


Fig. 8 Charge density mapping of D₂CeX₆ double perovskites.

enable strong photon absorption within the 1 to 5 eV energy range, making them suitable for efficient solar energy harvesting. Similarly, Cs_2PbX_6 (X = Cl, Br, I) double perovskites demonstrate excellent optoelectronic properties, with direct band gaps ranging from 0.45 to 2.54 eV and high absorption coefficients of approximately 5.95×10^5 cm⁻¹, which highlight their potential for photovoltaic and optical applications. However, the inherent toxicity and poor environmental stability

of lead severely limit its large-scale utilization. On the other hand, Although Si-based halide double perovskites A_2SiI_6 (A = K, Rb, Cs; X = Cl, Br, I) show suitable band gaps (0.84 to 1.15 eV), strong visible absorption, and good thermal stability, conventional silicon materials still suffer from an indirect 1.12 eV band gap, weak light absorption, high-temperature processing requirements, and limited flexibility, making them less efficient.⁶⁵ In contrast, our investigated D_2CeX_6 compounds

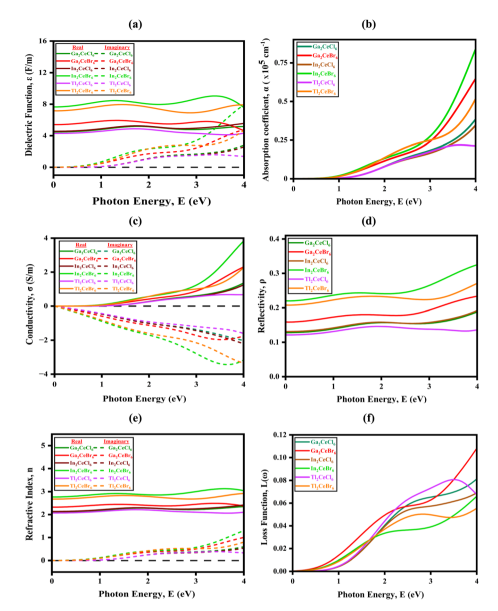


Fig. 9 Optical functions (a) dielectric function, (b) absorption coefficient, (c) conductivity, (d) refractive index, (e) reflectivity, (f) loss function of D_2CeX_6 double perovskites.

exhibit comparable or even superior optical characteristics, such as strong visible-light absorption, high dielectric constants, and large refractive indices, while eliminating the environmental and health hazards associated with lead. Moreover, their tunable band gaps within the optimal range for solar energy conversion, coupled with robust structural and mechanical stability, establish $D_2\text{CeX}_6$ materials as advanced, sustainable, and highly promising candidates for nextgeneration lead-free optoelectronic and photovoltaic devices.

3.6.1. Dielectric function. The dielectric function (ε) is a key parameter for understanding the optical and electronic behavior of materials. It consists of a real part (ε_1) , which describes the material's ability to store electric energy, and an imaginary part (ε_2) , which reflects energy loss due to absorption of electromagnetic radiation. For the D_2CeX_6 double

perovskites, the calculated dielectric function is shown in Fig. 9(a) and is expressed as: $\varepsilon(\omega) = \varepsilon_1(\omega) + i\varepsilon_2(\omega)$. ⁶⁶ Analyzing ε_1 and ε_2 provides insight into light–matter interactions, including polarization, charge screening, and optical absorption. These insights are essential for designing high-performance optoelectronic and photovoltaic devices, such as solar cells, LEDs, and photodetectors, where efficient light harvesting and minimal energy loss are critical. ⁶⁷ It is defined as, ⁶⁸

$$\varepsilon_{1}(\omega) = 1 + \frac{2}{\pi} P \int_{0}^{\infty} \frac{\omega^{'\varepsilon_{2}}(\omega^{'})}{\omega^{'2} - \omega^{2}} d\omega^{'}$$
 (13)

$$\varepsilon_2(\omega) = \frac{8}{2\pi\omega^2} \sum_{m'} |p_{nn'}(k)|^2 \frac{dS_k}{\nabla \omega_{m'}(ks)}$$
(14)

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Among the studied compounds, the static (0 eV) real part of the dielectric function (ε_1) is found to be 4.48 for Ga₂CeCl₆, 5.41 for Ga₂CeBr₆, 4.56 for In₂CeCl₆, 7.65 for In₂CeBr₆, 4.28 for Tl₂-CeCl₆, and 7.216 for Tl₂CeBr₆, as presented in Fig. 9(a). The ε_1 parameter reflects a material's capacity to store electric energy when subjected to an external electromagnetic field. With increasing photon energy, ε_1 initially increases due to enhanced polarization arising from interband electronic transitions, reaching a peak value before gradually declining. The relatively high ε_1 values suggest strong polarization capability and effective screening of internal electric fields, which are beneficial for lowering exciton binding energy and improving charge carrier separation in optoelectronic applications.

For all the investigated compounds, the imaginary part of the dielectric function (ε_2), which represents the material's ability to absorb electromagnetic energy, begins to increase from photon energies between 0.31 and 0.87 eV. As photon energy rises, ε_2 increases and reaches its maximum within the visible region (1.5–4 eV), with peak values of 2.83 for Ga₂CeCl₆, 4.74 for Ga₂CeBr₆, 2.67 for In₂CeCl₆, 8.15 for In₂CeBr₆, 1.59 (at 3.31 eV) for Tl₂CeCl₆, and 5.067 for Tl₂CeBr₆. These maximum values generally occur around 4.0 eV. The highest ε_2 peak observed in In₂CeBr₆ indicates significant energy loss due to strong absorption of electromagnetic radiation, whereas Tl₂-CeCl₆ exhibits the lowest ε_2 value, suggesting comparatively weaker absorption behavior.

The comparison between chloride and bromide analogs reveals that Br-based compounds generally exhibit enhanced imaginary dielectric components due to narrower band gaps, which facilitate easier electronic excitations. Conversely, Cl-based compounds show higher real parts, indicating stronger dielectric screening. The cation variation (Ga \rightarrow In \rightarrow Tl) affects both amplitude and energy dispersion, with heavier cations (Tl) contributing to stronger polarization and broader spectral response. Collectively, these dielectric characteristics reveal that Ga₂CeCl₆ and Tl₂CeBr₆ are particularly promising for optoelectronic applications requiring high dielectric screening and effective visible-light absorption.

3.6.2. Absorption coefficient. The absorption coefficient (α) is a critical parameter in materials science that quantifies how strongly a material absorbs light at specific photon energies. It is an important factor in assessing the appropriateness of materials for applications in optoelectronics, photovoltaics, scintillators, and photodetectors. ^{69,70} The absorption coefficient can be determined using eqn (15).

$$\alpha(\omega) = \frac{2\omega k(\omega)}{c} \tag{15}$$

The absorption coefficient, $\alpha(\omega)$, indicates how efficiently a material absorbs light at a particular angular frequency ω . It is commonly measured in cm⁻¹ and is directly related to the extinction coefficient $k(\omega)$, which represents the imaginary part of the complex refractive index.^{71,72} Fig. 9(b) shows that all six D₂CeX₆ compounds exhibit significant absorption coefficients, starting at photon energies of 0.8 to 1.7 eV. As the photon energy increases toward 3.0 eV, the absorption coefficients rise sharply,

reaching values of 18 524 cm⁻¹ (Ga₂CeCl₆), 24 394 cm⁻¹ (Ga₂-CeBr₆), 16 784 cm⁻¹ (In₂CeCl₆), 27 560 cm⁻¹ (In₂CeBr₆), 17 765 cm $^{-1}$ (Tl₂CeCl₆), and 25 221 cm $^{-1}$ (Tl₂CeBr₆). Such high absorption coefficients ($\alpha > 10^4 \text{ cm}^{-1}$)^{73,74} in the visible range are highly desirable for thin-film solar cells, as they allow efficient photon harvesting with minimal material thickness, indicating that all studied materials are promising candidates for photovoltaic applications. The absorption behavior is also influenced by the halide substitution (Cl to Br), which shifts the absorption edge toward lower energies due to the narrower band gap of Brbased compounds, thereby enhancing visible-light absorption. Additionally, the choice of D-site cation (Ga, In, Tl) affects the magnitude and position of absorption peaks, with In- and Tlbased compounds generally exhibiting higher absorption coefficients than Ga-based ones. These trends highlight the tunability of optical properties in D₂CeX₆ double perovskites through compositional engineering, which is crucial for optimizing performance in both solar energy conversion and optoelectronic applications.

3.6.3. Conductivity. Conductivity (σ) is a fundamental physical property that measures a material's ability to conduct electric current, especially under the influence of an electromagnetic field. In D₂CeX₆ double perovskites, the optical conductivity is frequency-dependent and complex, consisting of a real part, which represents the dissipative response or actual current flow, and an imaginary part, which corresponds to energy storage and polarization, as shown in Fig. 9(c). Calculating the optical conductivity is essential for understanding charge carrier dynamics, photon–electron interactions, and energy transport mechanisms, all of which are critical for the design of high-performance optoelectronic and photonic devices. The optical conductivity is obtained directly from the complex dielectric function through the following relation:

$$\sigma_{1}(\omega) = \frac{2\pi e^{2}\hbar}{m^{2}\omega\Omega} \sum_{k,v,c} |\langle \psi_{ck}|p|\psi_{vk}\rangle|^{2} \delta(E_{ck} - E_{vk} - \hbar\omega) \qquad (16)$$

In this expression, Ω denotes the volume of the unit cell, ψ_{vk} and $\psi_{\rm ck}$ represent the valence and conduction band wavefunctions at a given k-point, and δ is the Dirac delta function that maintains energy conservation, from the conductivity spectra. The real component of the optical conductivity exhibits a trend similar to that of the absorption coefficient, as both originate from electronic transitions between the valence and conduction bands. Higher values of real optical conductivity reflect stronger photon-electron interactions, leading to enhanced light absorption and more efficient generation of charge carriers. In D₂CeX₆ double perovskites, this relationship indicates that compounds with large absorption coefficients also display pronounced conductivity peaks, emphasizing their suitability for photovoltaic and optoelectronic applications, where effective light harvesting and rapid charge transport are crucial for device performance.

Fig. 9(c) shows the real and imaginary parts of optical conductivity (σ) as a function of photon energy (0 to 4 eV) for the double perovskites Ga₂CeCl₆, Ga₂CeBr₆, In₂CeCl₆, In₂CeBr₆, Tl₂CeCl₆, and Tl₂CeBr₆. Among all, Ga₂CeCl₆ exhibits the

highest real conductivity, rising sharply beyond 2.5 eV, indicating superior photon-induced carrier excitation and possible metallic-like behavior at high photon energy. Ga₂CeBr₆ also shows strong conductivity but slightly lower than Ga₂CeCl₆, suggesting that replacing Cl with Br slightly reduces the optical response, likely due to the larger ionic radius and weaker bonding of Br. The In-based compounds (In₂CeCl₆ and In₂-CeBr₆) show moderate conductivity values, implying less efficient charge carrier transition compared to Ga-based systems; however, In₂CeBr₆ displays a smoother rise, indicating better optical absorption at higher energies than In₂CeCl₆. In contrast, Tl-based compounds (Tl₂CeCl₆ and Tl₂CeBr₆) possess the lowest real conductivity across the entire photon energy range, reflecting their relatively poor electronic transition probability and higher resistive character. As shown in Fig. 6(c), the imaginary part of the optical conductivity initially starts with negative values, indicating a reactive polarization response. With increasing photon energy, the conductivity decreases further and reaches minimum values of -2.0 at 4.00 eV for Ga_2CeCl_6 , -1.95 at 3.40 eV for Ga_2CeBr_6 , -2.184 at 4.00 eV for In_2CeCl_6 , -3.43 at 3.42 eV for In₂CeBr₆, -1.57 at 4.00 eV for Tl₂CeCl₆, and −3.41 at 3.56 eV for Tl₂CeBr₆. This transient negative behavior reflects strong reactive polarization, which is advantageous for applications such as optical switching and memory devices. Notably, cation substitution (Ga \rightarrow In \rightarrow Tl) and halide replacement (Cl \rightarrow Br) systematically influence the peak positions and depth of the negative conductivity response.

Heavier cations and larger halides generally shift the spectral features toward lower energies and enhance polarization effects, thereby tailoring the material's suitability for frequency-selective optoelectronic and photonic applications.

3.6.4. Reflectivity. Reflectivity (ρ) is a material's ability to reflect incident electromagnetic radiation, expressed as the ratio of reflected power to incident power at a specific photon energy. It is a dimensionless quantity ranging from 0 to 1 (or 0% to 100%) and depends on the material's optical properties and surface characteristics. The reflectivity (ρ) is calculated from the complex dielectric function using the relation,

$$R(\omega) = \frac{(\eta - 1)^2 + k^2}{(\eta + 1)^2 + k^2} \tag{17}$$

Fig. 9(d) illustrates the variation of reflectivity (*ρ*) with photon energy (*E*) from 0 to 4 eV for six Ce-based double perovskite compounds: Ga₂CeCl₆, Ga₂CeBr₆, In₂CeCl₆, In₂-CeBr₆, Tl₂CeCl₆, and Tl₂CeBr₆. Reflectivity represents the fraction of incident light that is reflected from a material's surface and serves as an important indicator of its optical response and surface electronic structure. Overall, all compounds exhibit relatively low reflectivity (<0.4), suggesting strong light absorption and significant potential for optoelectronic and photovoltaic applications. At 0.1 eV, the initial reflectivity values are 0.128, 0.1589, 0.131, 0.220, 0.1213, and 0.1213 for Ga₂CeCl₆, Ga₂CeBr₆, In₂CeCl₆, In₂CeBr₆, Tl₂CeCl₆, and Tl₂CeBr₆, respectively. All materials show notable optical activity in the visible range (1.5–4 eV). As the photon energy increases, the reflectivity

rises to 0.189, 0.23, 0.185, 0.32, 0.135, and 0.27 for Ga₂CeCl₆, Ga₂CeBr₆, In₂CeCl₆, In₂CeBr₆, Tl₂CeCl₆, and Tl₂CeBr₆, respectively. Overall, these reflectivity characteristics highlight distinct, application-specific roles, underscoring the importance of reflectivity in optimizing these materials for optoelectronic technologies.

3.6.5. Refractive index. The refractive index (n) is a complex optical parameter that characterizes the propagation of light within a material.⁷⁷ It consists of a real part, indicating the phase velocity of light in the medium, and an imaginary part, which is associated with absorption losses (extinction coefficient). The refractive index of D_2CeX_6 double perovskites shown in Fig. 9(e) is essential for optimizing their performance in photonic, optoelectronic, and photovoltaic applications, since it governs light–matter interaction, reflection, refraction, and light confinement efficiency. The $n(\omega)$ is computed from the complex dielectric function, using the relation,⁷⁸

$$n(\omega) = \frac{1}{\sqrt{2}} \left[\left(\varepsilon_1(\omega)^2 + \varepsilon_2(\omega)^2 \right)^{\frac{1}{2}} + \varepsilon_1(\omega) \right]^{\frac{1}{2}}$$
 (18)

 $n(\omega)$ represents the frequency-dependent refractive index, indicating how the speed of light varies in the material at a particular photon energy.

Fig. 9(e) illustrates the variation of the refractive index (n), both real and imaginary components, with photon energy (0 to 4 eV) for six Ce-based double perovskite compounds: Ga₂CeCl₆, Ga₂CeBr₆, In₂CeCl₆, In₂CeBr₆, Tl₂CeCl₆, and Tl₂CeBr₆. The solid lines denote the real part of the refractive index, which signifies the phase velocity of light within the material, while the dashed lines indicate the imaginary part, which represents the extinction coefficient or absorption losses. Overall, the real part of n remains between approximately 2.0 and 4.5, showing moderate to high optical density typical of semiconducting materials. Ga₂CeCl₆ exhibits the highest real refractive index across most of the energy range, peaking near 4.5, implying stronger lightmatter interaction and higher optical polarization. Ga₂CeBr₆ also shows a comparatively high refractive index but slightly lower than its Cl counterpart, suggesting that Br substitution reduces the electronic polarizability due to its larger ionic radius and weaker bond strength. The In-based compounds (In₂CeCl₆ and In₂CeBr₆) have intermediate refractive indices around 2.5-3.0, indicating moderate optical response with more uniform energy dependence, suitable for balanced optical transmission and absorption. In contrast, Tl₂CeCl₆ and Tl₂-CeBr₆ display the lowest refractive indices (\sim 2.0 to 2.5), reflecting weaker electronic polarization and higher optical transparency. The imaginary components increase gradually beyond 2 eV for all compounds, highlighting the onset of interband electronic transitions that contribute to absorption. Among these, Ga-based compounds again exhibit the strongest imaginary peaks, confirming their intense optical activity, while Tl-based compounds show the weakest, consistent with their low absorption tendency. Overall, both halide and cation substitution significantly influence refractive behavior, with the comparative trend for real refractive index following Ga₂CeCl₆ >

Ga₂CeBr₆ > In₂CeBr₆ > In₂CeCl₆ > Tl₂CeCl₆ > Tl₂CeBr₆. This indicates that Ga-based systems, particularly Ga₂CeCl₆, possess superior optical confinement and are more suitable for light-modulating or refractive applications, whereas Tl-based perovskites are better for transparency-driven optoelectronic devices.

3.6.6. Loss function. The energy loss function, $L(\omega)$, defined as the imaginary part of the inverse dielectric function, describes the energy dissipated by fast electrons traversing a material and is essential for probing plasmonic behavior, electron scattering, and optical transitions. The optical loss function, $L(\omega)$, can be calculated using the following relation: $L(\omega)$

$$L(\omega) = -\operatorname{Im}\left[\frac{1}{\varepsilon(\omega)}\right] = \frac{\varepsilon_2(\omega)}{\varepsilon_1^2(\omega) + \varepsilon_2^2(\omega)} \tag{19}$$

As shown in Fig. 9(f), the relationship between the Loss Function, $L(\omega)$, on the y-axis and photon energy, E (in electron volts, eV), on the x-axis for six different materials: Ga₂CeCl₆, Ga₂CeBr₆, In₂CeCl₆, In₂CeBr₆, Tl₂CeCl₆, and Tl₂CeBr₆. Each material is represented by a distinct colored line as indicated in the legend: green for Ga₂CeCl₆, red for Ga₂CeBr₆, brown for In₂CeCl₆, cyan for In₂CeBr₆, magenta for Tl₂CeCl₆, and orange for Tl₂CeBr₆. The general trend for all materials shows that the loss function starts near zero at low photon energies (0 eV) and increases as the photon energy increases up to 4 eV. However, the rate and magnitude of increase differ for each compound. Ga₂CeBr₆ exhibits the highest loss function values, surpassing 0.10 at around 4 eV, indicating the strongest loss effect at high photon energies among the six. Tl₂CeCl₆ follows closely, peaking slightly below Ga₂CeBr₆ but still above 0.08. Ga₂CeCl₆ and In₂CeCl₆ show moderate increases, with their loss functions rising to approximately 0.08 and 0.07, respectively, at 4 eV. The In₂CeBr₆ and Tl₂CeBr₆ compounds display the lowest loss functions overall, with Tl₂CeBr₆ being the lowest, peaking near 0.06. Comparatively, the bromide variants (Ga₂CeBr₆, In₂CeBr₆, Tl₂CeBr₆) generally display higher loss functions than their chloride counterparts at higher photon energies, except for In₂CeBr₆, which is lower than In₂CeCl₆. Among the cations, Gabased compounds tend to show higher loss functions than Inbased compounds, while Tl-based compounds are intermediate but closer to Ga-based, especially in the chloride form.

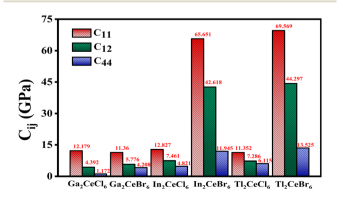


Fig. 10 Calculated elastic constants (C_{ij}) for D_2CeX_6 double perovskites.

This suggests that both the halide type and the central metal ion influence the loss function behavior, with bromides and Ga ions generally enhancing the loss function, except in specific cases like In_2CeBr_6 .

3.7. Elastic and mechanical properties

Mechanical properties define a material's response to stress, including its strength, elasticity, stiffness, and ductility. These are vital for assessing structural stability and durability. In solar cells, especially in thin-film or flexible applications, materials must endure mechanical strain during fabrication and operation without cracking or deforming. Elastic constants define the mechanical behavior of a solid under applied stress or strain, serving as essential indicators of its stability, rigidity, and bonding nature. In the framework of Hooke's law, the stress tensor (σ) is linearly related to the strain tensor (ε) through:

$$\sigma_{ij} = C_{ijkl} \varepsilon_{kl} \tag{20}$$

where C_{ij} are the components of the fourth-rank elastic stiffness tensor. The symmetry of the crystal determines the number of independent elastic constants. In cubic crystals, there are only three independent constants: C_{11} , C_{12} , and C_{44} . The mechanical stability of cubic crystals was evaluated using the Born–Huang criteria, expressed as:^{83–85}

$$C_{11} > 0$$
, $4C_{44} > 0$, $C_{11} - C_{12} > 0$ and $C_{11} + 2C_{12} > 0$ (21)

All the conditions of the Born–Huang stability criteria are satisfied and yield positive values, indicating that the $\mathrm{D}_2\mathrm{CeX}_6$ double perovskites are mechanically stable, as shown in Fig. 10. This confirms their ability to withstand external stress without structural failure, making them suitable for practical applications in devices where mechanical integrity is essential.

The mechanical properties of D₂CeX₆ double perovskites are critical for ensuring structural integrity, thermal stability, and resistance to external stress. The bulk modulus, shear modulus, Young's modulus, Poisson's ratio, and Pugh's ratio serve as indicators of ductility or brittleness, guiding decisions about the materials' suitability for device integration.^{48,61} Mechanical stability plays a direct role in the long-term reliability and lifespan of perovskite-based solar cells. Therefore, a thorough understanding and optimization of these properties are essential for developing mechanically robust, efficient, and commercially viable perovskite solar absorbers for future energy applications.

Fig. 11 illustrates the mechanical parameters of D_2CeX_6 double perovskites, including bulk modulus (B), shear modulus (G), Young's modulus (Y), Poisson's ratio (V), Pugh's ratio (B/G), and anisotropy factor (A^U), which are represented by equations as follows.

$$B_{\rm H} = \frac{B_{\rm H} + B_{\rm R}}{2} \tag{22}$$

$$G_{\rm H} = \frac{G_{\rm V} + B_{\rm R}}{2} \tag{23}$$

$$Y = \frac{9BG}{(3B+G)} \tag{24}$$

Bulk modulus (B) measures a material's resistance to uniform compression and indicates mechanical stiffness. Higher B values reflect strong interatomic bonding and limited compressibility, while lower values correspond to flexible or soft materials. Typically, materials with B > 40 GPa are classified as hard, whereas B < 40 GPa denotes soft or flexible behavior. As shown in Fig. 11(a), In₂CeBr₆ (50.30 GPa) and Tl₂CeBr₆ (52.72 GPa) are hard materials, while Ga₂CeCl₆, Ga₂CeBr₆, In₂CeCl₆, and Tl₂CeCl₆ are soft, capable of accommodating volume changes more easily. Shear modulus (G) quantifies resistance to shape (shear) deformation at constant volume. Higher G values indicate strong internal bonding and shear rigidity, while lower values correspond to easier shear deformation. As depicted in Fig. 11(b), Tl₂CeBr₆ (13.16 GPa) and In₂CeBr₆ (11.77 GPa)

exhibit the highest shear moduli, making them suitable for load-bearing and wear-resistant applications.

In contrast, Ga₂CeCl₆ (1.94 GPa) and Ga₂CeBr₆ (3.57 GPa) are more flexible, advantageous for damping, vibration isolation, or low-shear optical layers. Young's modulus (E) reflects stiffness and resistance to elongation under uniaxial stress. Fig. 11(c) shows that In₂CeCl₆ and Tl₂CeBr₆ have the highest E values, consistent with their B and G moduli, making them ideal for rigid supports, optical mounts, and stiff structural components. Conversely, Ga₂CeCl₆ (5.34 GPa) exhibits enhanced elasticity, suitable for flexible electronics, wearable devices, and straintolerant applications. Overall, the variation of B, G, and Eacross cation-anion substitutions demonstrate the tunable mechanical behavior of D2CeX6 perovskites, enabling design for either rigid, high-strength applications (In- and Tl-Br based) or flexible, damping-oriented applications (Ga-based). The ductile or brittle nature of a material can be assessed using Poisson's ratio (ν) and Pugh's ratio (B/G). In general, materials with ν

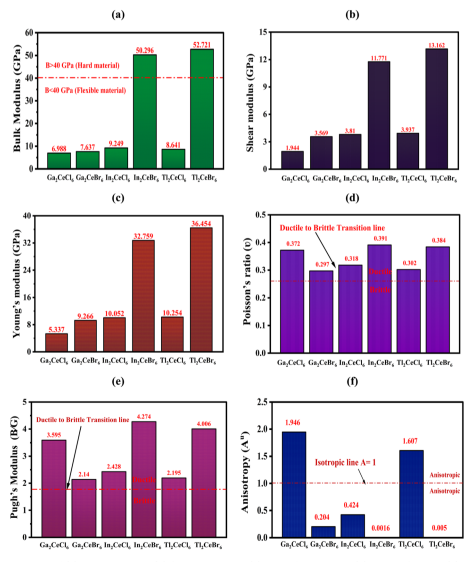


Fig. 11 Mechanical parameters of (a) Bulk modulus, (b) Shear modulus, (c) Young's modulus, (d) Poisson's ratio, (e) Pugh's modulus, and (f) anisotropy of D_2CeX_6 double perovskite materials.

exceeding 0.26 and B/G greater than 1.75 are classified as ductile.²⁹

$$v = \frac{3B - 2G}{2(3B + G)} \tag{25}$$

Poisson's ratio (ν), illustrated in Fig. 11(d), describes the ratio of lateral to axial strain under applied stress, reflecting both bonding characteristics and ductility. For the D₂CeX₆ perovskites studied, the calculated ν values are 0.372, 0.297, 0.318, 0.391, 0.302, and 0.384 for Ga₂CeCl₆, Ga₂CeBr₆, In₂CeCl₆, In₂CeBr₆, Tl₂CeCl₆, and Tl₂CeBr₆, respectively. Since all values exceed the critical threshold of 0.26, these compounds exhibit ductile behavior, indicating their ability to undergo plastic deformation and their suitability for flexible applications such as strain-tolerant devices, vibration-damping layers, and ductile electrodes. Likewise, Pugh's ratio (B/G), shown in Fig. 11(e), provides a complementary measure of ductility, where values above 1.75 confirm ductile behavior. All investigated D₂CeX₆ perovskites surpass this criterion, reinforcing the conclusion from Poisson's ratio that these materials are inherently ductile.

The Universal Anisotropy Index (A^{U}) in Fig. 11(f) quantifies the directional dependence of elastic properties, where zero denotes perfect isotropy.⁸⁷

$$A^{\rm U} = \frac{B_{\rm V}}{B_{\rm r}} + 5\frac{G_{\rm V}}{B_{\rm R}} - 6 \ge 0 \tag{26}$$

 ${\rm In_2CeBr_6}$ ($A^{\rm U}=0.0016$) and ${\rm Tl_2CeBr_6}$ ($A^{\rm U}=0.005$) exhibit near-isotropic behavior, suitable for applications where mechanical stability is critical, for example, in sensors, microelectronic substrates, and integrated circuits. On the other hand, ${\rm Ga_2CeCl_6}$ displays significant anisotropy ($A^{\rm U}=1.946$), implying strongly direction-dependent mechanical responses, which could be leveraged in anisotropic optical or piezoelectric devices where orientation-dependent properties are desired.

3.8. Anisotropy

Anisotropy refers to the variation of material properties with direction, and its evaluation is crucial in understanding the mechanical behavior of D₂CeX₆ double perovskites.

From the graphical assessments, Fig. 12–14, Ga_2CeCl_6 displays the highest mechanical anisotropy with Young's modulus ranging between \sim 2.5 and 9.0 GPa, indicating its stiffness varies significantly with direction. In contrast, Tl_2 -CeBr₆ shows the highest and most isotropic stiffness (\sim 35 to 37 GPa), followed by In_2CeBr_6 (\sim 32 to 33 GPa), highlighting their superior mechanical robustness and uniformity. Similarly, in

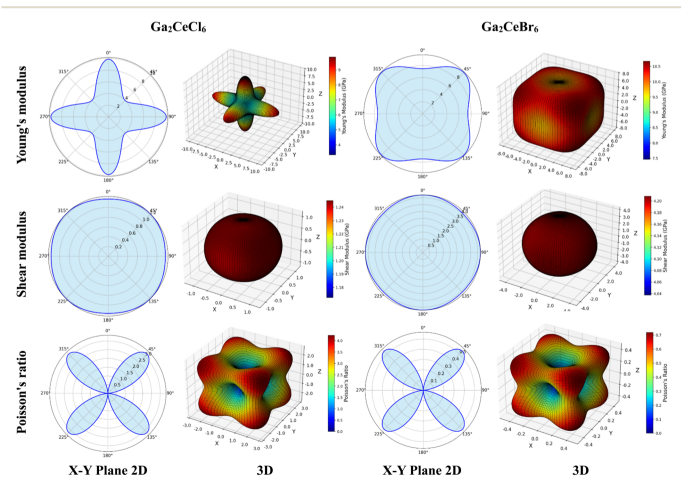


Fig. 12 Graphical assessment of Ga₂CeCl₆ and Ga₂CeBr₆ double perovskites anisotropy showing 2D and 3D figures.

shear modulus, Tl_2CeBr_6 peaks (\sim 13.5 GPa), while Ga_2CeCl_6 registers the lowest (\sim 1.17 to 1.24 GPa).

For Poisson's ratio, Tl_2CeBr_6 and In_2CeBr_6 exhibit broader, more spherical distributions (\sim 0.2 to 1.0), suggesting low anisotropy, whereas Ga_2CeCl_6 again demonstrates high directionality with values dipping to \sim 0.05.

Comparing the compounds, bromide-based materials are generally stiffer and more isotropic than their chloride counterparts, and mechanical performance improves from $Ga \to In \to Tl$ due to increasing atomic size and enhanced bonding. S8,89 These insights are vital for tailoring materials for specific directional mechanical responses or ensuring mechanical stability in multifunctional devices. Accurate anisotropy assessment helps optimize materials for real-world applications, guiding experimental synthesis and structural design.

3.9. Molecular dynamics (MD) analysis

Molecular Dynamics (MD) is a computational technique that simulates the motion of atoms and molecules over time by solving Newton's equations of motion, providing a dynamic perspective on material behavior under different energy components and temperatures.⁹⁰ In this study, simulations were performed at 300 K, 600 K, and 900 K to monitor the potential energy (P.E.), kinetic energy (K.E.), and total energy

(T.E.) of the systems. MD analysis offers valuable insights into thermal and energetic stability, atomic vibrations, diffusion mechanisms, and structural evolution under varying temperature or pressure conditions.91 This approach is particularly important for predicting material performance in real-world environments, including stability, phase transitions, and mechanical or electronic responses under thermal stress.92 Therefore, MD simulations are essential for assessing and optimizing the structural robustness and functional reliability of materials in practical applications. Fig. 15 illustrates the variation of energy components and temperature with simulation time for Ga₂CeCl₆ at different temperatures (300 K, 600 K, and 900 K). In Fig. 15(a)-(c), the potential energy (P.E.), kinetic energy (K.E.), and total energy (T.E.) are plotted against time for 300 K, 600 K, and 900 K, respectively. At 300 K (Fig. 15(a)), the system shows relatively stable energy fluctuations, with total energy oscillating mildly around equilibrium, indicating thermal stability at lower temperatures. As the temperature increases to 600 K (Fig. 15(b)), both kinetic and total energies rise significantly, accompanied by slightly larger oscillations, reflecting enhanced atomic motion. At 900 K (Fig. 15(c)), the kinetic and total energies reach their highest values, exhibiting pronounced fluctuations, which suggests strong thermal vibrations and a possible onset of lattice instability.

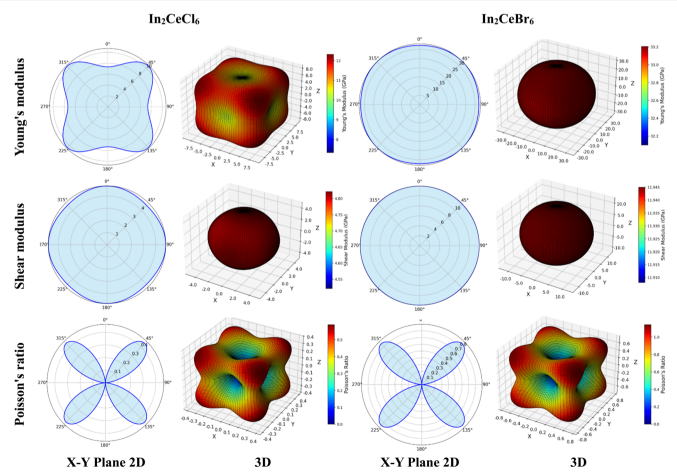


Fig. 13 Graphical assessment of In₂CeCl₆ and In₂CeBr₆ double perovskites anisotropy showing 2D and 3D figures.

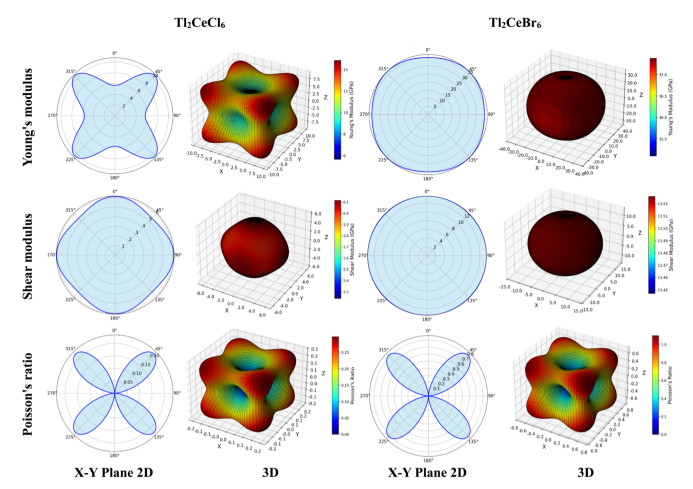


Fig. 14 Graphical assessment of Tl_2CeCl_6 and Tl_2CeBr_6 double perovskites anisotropy showing 2D and 3D figures.

The potential energy remains comparatively lower and negative in all cases, consistent with the system's bound state. Fig. 15(d) presents the temperature variation over time, showing steady temperature maintenance around the target values (300 K, 600 K, and 900 K), with moderate oscillations attributed to thermal equilibrium fluctuations.

Fig. 16 depicts the variation of kinetic energy (K.E.), potential energy (P.E.), total energy (T.E.), and temperature (*T*) of Ga₂-CeBr₆ as a function of time at different temperatures (300 K, 600 K, and 900 K) during molecular dynamics (MD) simulations. Fig. 16(a)–(c) shows that as temperature increases, the amplitude and average values of K.E. and T.E. rise, reflecting enhanced atomic vibrations and thermal activity. At 300 K, the system exhibits relatively small oscillations and stable energy values, indicating a well-equilibrated and structurally stable phase. At 600 K, both K.E. and T.E increase moderately, with more frequent fluctuations due to higher thermal motion.

At 900 K, the oscillations become much larger, and the total energy reaches its maximum level, signifying strong lattice vibrations and possibly anharmonic effects. P.E remains negative across all temperatures but shows a gradual upward shift with increasing temperature, suggesting partial weakening of interatomic interactions. Fig. 16(d) compares the temperature variation over time, showing that each system stabilizes near its

target temperature (300 K, 600 K, and 900 K) with expected thermal fluctuations.

Fig. 17 illustrates the variation of kinetic energy (K.E.), potential energy (P.E.), total energy (T.E.), and temperature (T) of In_2CeCl_6 during molecular dynamics (MD) simulations at 300 K, 600 K, and 900 K over a 50 ps timescale. Fig. 17(a)–(c), the energy profiles show that as temperature increases, both K.E. and T.E rise significantly, while P.E remains negative but slightly increases, indicating stronger atomic vibrations and enhanced atomic displacement at elevated temperatures.

At 300 K, the energy fluctuations are relatively small, implying structural stability with limited atomic motion. At 600 K, oscillations in K.E. and T.E become more pronounced, showing moderate thermal agitation. At 900 K, energy oscillations are much larger, signifying intense atomic vibration and possible lattice distortion. Fig. 17(d) compares the temperature profiles, showing stable fluctuations around the target temperatures (300 K, 600 K, and 900 K), confirming proper thermal equilibration during the MD simulations.

Fig. 18 presents the variation of kinetic energy (K.E.), potential energy (P.E.), total energy (T.E), and temperature (T) of In₂CeBr₆ over 50 ps at three different temperatures, 300 K, 600 K, and 900 K, obtained from molecular dynamics simulations. Fig. 18(a)–(c), the system exhibits increasing energy magnitudes

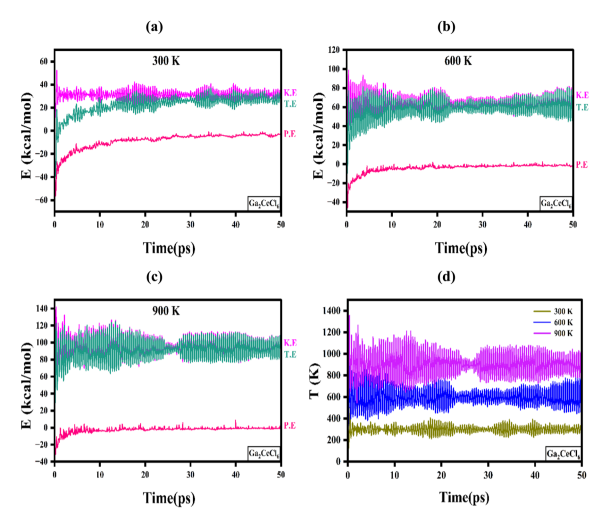


Fig. 15 Time evolution of potential, kinetic, and total energy for Ga_2CeCl_6 double perovskite at (a) 300 K, (b) 600 K, (c) 900 K, and (d) temperature variation over time.

with rising temperature, indicating stronger atomic vibrations and enhanced lattice dynamics. At 300 K, K.E. and T.E. remain relatively stable with small oscillations, reflecting a thermally equilibrated and structurally stable configuration. As the temperature increases to 600 K, both K.E. and T.E rise moderately, accompanied by more pronounced oscillations due to intensified thermal motion. At 900 K, large fluctuations in K.E. and T.E are observed, implying vigorous atomic vibrations and possible anharmonic effects, while P.E remains negative but shifts slightly upward, suggesting reduced interatomic binding strength. Panel (d) compares temperature variations over time, showing consistent stabilization near the target values of 300 K, 600 K, and 900 K with periodic fluctuations characteristic of dynamic equilibrium.

Fig. 19 presents the variation of energy and temperature with simulation time for Tl_2CeCl_6 at three different temperatures: 300 K, 600 K, and 900 K. Fig. 19(a to c) show the evolution of kinetic energy (K.E), potential energy (P.E), and total energy (T.E), while Fig. 19(d) depicts the temperature fluctuations over time. At 300 K, Fig. 19(a), the system displays stable energy oscillations with minor fluctuations, indicating good thermal equilibrium and structural stability. As the temperature rises to

600 K, Fig. 19(b), both kinetic and total energies increase noticeably, accompanied by slightly larger oscillations, suggesting enhanced atomic motion. At 900 K, Fig. 19(c), the total and kinetic energies reach their highest average values, and the oscillations become more pronounced, reflecting greater thermal agitation and the possible onset of lattice instability. The potential energy becomes less negative with increasing temperature, implying partial weakening of interatomic interactions. The temperature profile in Fig. 19(d) confirms consistent thermal behavior, where higher set temperatures produce broader fluctuations but maintain equilibrium around their respective mean values.

Fig. 20 illustrates the variation of total energy (T.E), kinetic energy (K.E), potential energy (P.E), and temperature (T) with time for Tl₂CeBr₆ at different temperatures, 300 K, 600 K, and 900 K, over a simulation period of 50 ps. Fig. 20(a)–(c) shows the energy profiles, while panel Fig. 17(d) presents the temperature fluctuations. At 300 K, Fig. 20(a), the system exhibits relatively stable energy oscillations with moderate kinetic and total energy values and a potential energy of around -40 kcal mol⁻¹, indicating structural stability. As the temperature increases to 600 K, Fig. 20(b), both K.E. and T.E rise, showing higher

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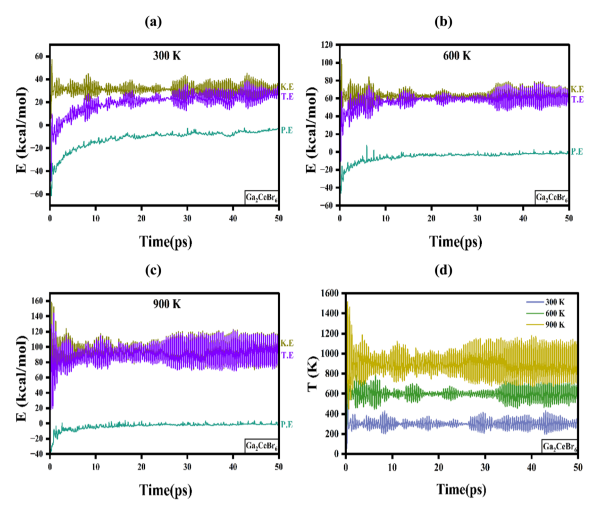


Fig. 16 Time evolution of potential, kinetic, and total energy for Ga₂CeBr₆ double perovskite at (a) 300 K, (b) 600 K, (c) 900 K, and (d) temperature variation over time.

fluctuation amplitudes, whereas P.E slightly shifts toward less negative values, reflecting increased atomic vibrations.

At 900 K, Fig. 17(c), kinetic and total energies reach their highest values, and fluctuations become more pronounced, suggesting enhanced atomic motion and thermal agitation. The temperature νs . time plot Fig. 17(d) confirms that Tl_2CeBr_6 maintains near-equilibrium average temperatures around 300 K, 600 K, and 900 K, respectively, though oscillations widen with rising temperature.

3.10. Phonon analysis

Phonon dispersion analysis is a key tool to investigate the vibrational and dynamical stability of materials.⁹³

The phonon spectra indicate whether a crystal structure is stable: the absence of imaginary (negative) frequencies confirms dynamical stability, while the presence of such frequencies may signal lattice instabilities or soft phonon modes. ⁹⁴ Fig. 21 presents the phonon dispersion curves for six Ce-based halide double perovskites plotted along the high-symmetry K-points (X-R-M- Γ -R) with frequency (THz) as a function of wave vector: (a) Ga₂CeCl₆, (b) Ga₂CeBr₆, (c) In₂-CeCl₆, (d) In₂CeBr₆, (e) Tl₂CeCl₆, and (f) Tl₂CeBr₆ double

perovskite materials. From the phonon spectra, Ga₂CeCl₆, In₂-CeCl₆, and Tl₂CeCl₆ exhibit no negative frequencies across the entire Brillouin zone, confirming their excellent dynamical stability and structurally robust behavior under equilibrium conditions. In contrast, Ga2CeBr6, In2CeBr6, and Tl2CeBr6 display very small negative frequencies near the Γ -point, suggesting the presence of slight lattice instabilities or soft phonon modes. These modes may become stabilized under finite temperature or pressure, or through minor structural relaxation. Comparing the compounds, chloride-based perovskites possess a more rigid lattice and stronger bonding interactions due to the lighter Cl atoms, resulting in higher phonon frequencies and enhanced dynamic stability. Bromide-based compounds, on the other hand, display softer vibrational modes due to the heavier Br atoms, leading to slightly reduced structural robustness.

3.11. Population analysis

Population analysis is a fundamental computational technique in solid-state physics and quantum chemistry, employed to quantify the distribution of electronic charge among atoms in a compound. It provides crucial insight into bonding character,

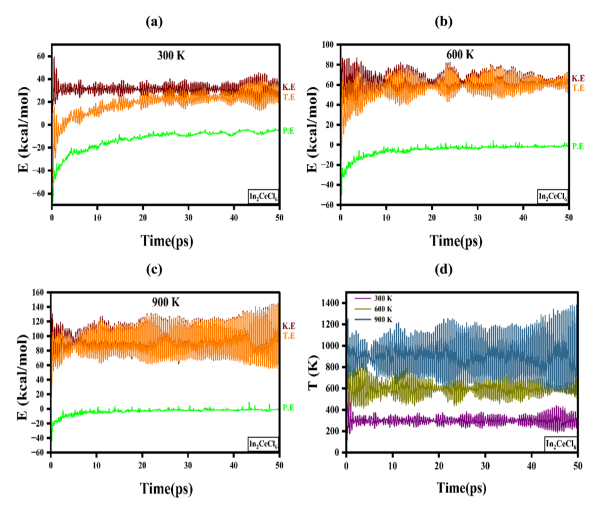


Fig. 17 Time evolution of potential, kinetic, and total energy for In_2CeCl_6 double perovskite at (a) 300 K, (b) 600 K, (c) 900 K, and (d) temperature variation over time.

charge transfer, and the electronic environment of individual species within complex materials. Among the established schemes, Mulliken and Hirshfeld population analyses are widely used. 95,96 Mulliken analysis partitions the electron density based on basis function overlaps, while Hirshfeld analysis assigns electron density relative to reference atomic densities in Table 5. These methods are particularly important for analyzing halide double perovskites such as D₂CeX₆, where mixed ionic-covalent bonding and intricate charge distributions significantly influence optoelectronic and structural properties.

In the present study, population analysis was carried out for a series of D₂CeX₆ compounds. The Mulliken atomic populations, categorized into s, p, d, and f orbitals, reveal substantial trends in Table 5. Notably, Tl atoms exhibit the highest total Mulliken population at 20.24 (in Tl₂CeCl₆), owing to their large atomic size and heavy electron contribution from the 6 s and 6p orbitals, whereas halide anions (Cl and Br) consistently show the lowest, with Br in Tl₂CeBr₆ at just 7.09. Among the cations, Ga atoms in Ga₂CeCl₆ possess the lowest total Mulliken population of 12.25, reflecting the lighter, less electron-rich nature of Ga compared to In and Tl. Cerium (Ce) atoms display significant f-orbital occupancy (1.10 to 1.15),

underlining the role of 4f electrons in determining the material's optical and electronic properties. Further, the Mulliken charge change values indicate the extent of electron gain or loss relative to neutral atoms.

The highest positive charge transfer occurs for Tl in Tl₂CeCl₆ (0.76), while the largest electron acceptance is seen in Cl in the same compound (-0.36), reaffirming the ionic nature of halides as electron acceptors. Hirshfeld charge changes, though lower in magnitude, mirror these trends, with Tl showing a maximum of 0.46 and Cl a minimum of -0.19 in Tl₂CeCl₆. Comparing across halide substitutions, replacing Cl with Br generally results in reduced total atomic populations and charge transfers, consistent with Br's lower electronegativity. Similarly, moving from Ga to heavier In and Tl systematically increases total Mulliken populations and charge changes, reflecting the influence of cationic size and electronic configuration on charge distribution. The significance of such population analysis extends to practical applications, as the charge distribution in these double perovskites directly impacts their optical absorption behavior, defect tolerance, and charge carrier dynamics. These factors are critical in determining the performance of perovskite-based materials in applications such as

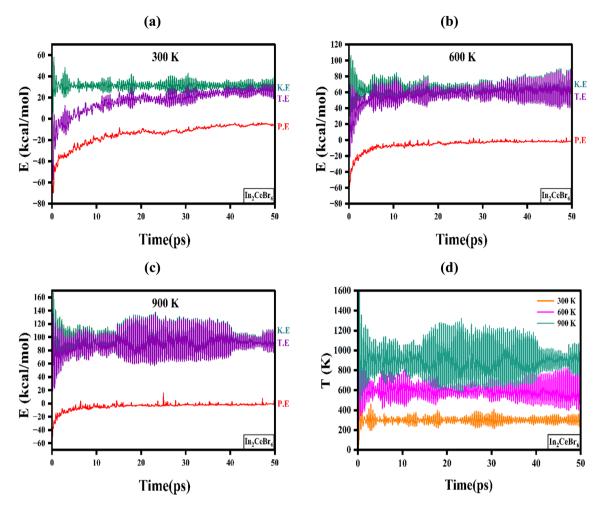


Fig. 18 Time evolution of potential, kinetic, and total energy for In_2CeBr_6 double perovskite at (a) 300 K, (b) 600 K, (c) 900 K, and (d) temperature variation over time.

photovoltaics, X-ray scintillators, and optoelectronic devices. Accurate knowledge of charge transfer tendencies aids in tuning material properties, enabling the rational design of compounds with optimized band gaps, enhanced stability, and superior optoelectronic performance. This comprehensive population analysis thus not only elucidates the fundamental electronic structure of $D_2 CeX_6$ double perovskites but also guides the engineering of advanced materials for future energy and photonic technologies.

4 Device architecture and simulation parameters

In the present work, a series of photovoltaic multilayer structures was systematically designed with the configuration Al/FTO/WS $_2$ /D $_2$ CeX $_6$ /Ni, where D = Ga, In, Tl, and X = Cl, Br. Six distinct device architectures were considered, namely Al/FTO/WS $_2$ /Ga $_2$ CeeCl $_6$ /Ni, Al/FTO/WS $_2$ /Ga $_2$ CeBr $_6$ /Ni, Al/FTO/WS $_2$ /In $_2$ CeCl $_6$ /Ni, Al/FTO/WS $_2$ /In $_2$ CeBr $_6$ /Ni, Al/FTO/WS $_2$ /Tl $_2$ CeCl $_6$ /Ni, and Al/FTO/WS $_2$ /Tl $_2$ CeBr $_6$ /Ni. These structures were modeled using SCAPS-1D (Version 3.3.09), which numerically solves the fundamental semiconductor equations. including Poisson's

equation and the electron-hole continuity equations under steady-state conditions. This ensures an accurate representation of charge generation, recombination, and transport processes within the device. The schematic layout of the proposed heterostructures is depicted in Fig. 2(b). Each solar cell consists of several functional layers engineered to optimize light absorption and charge carrier extraction. The aluminum (Al) front contact (work function: 4.20 eV) serves as the transparent conductive electrode, while the fluorine-doped tin oxide (FTO) layer acts as an additional transparent conductor, enabling efficient light transmission into the active layers. An ntype WS₂ electron transport layer (ETL) is introduced to facilitate selective electron extraction and minimize interfacial recombination losses. The D₂CeX₆ perovskite absorber layer functions as the primary photoactive medium, where incident photons are absorbed to generate electron-hole pairs. Finally, the nickel (Ni) back contact (work function: 5.22 eV) provides efficient hole collection and establishes an ohmic interface with the absorber. Nickel was deliberately selected as the back electrode because of its favorable combination of properties, including high electrical conductivity, excellent interfacial adhesion, chemical stability, and significantly lower cost

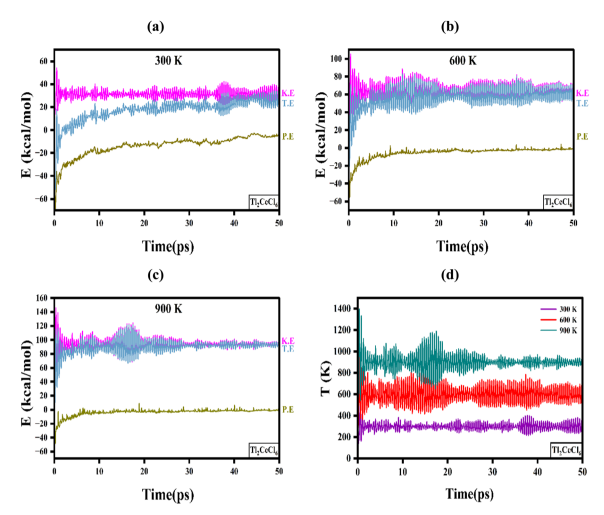


Fig. 19 Time evolution of potential, kinetic, and total energy for Tl_2CeCl_6 double perovskite at (a) 300 K, (b) 600 K, (c) 900 K, and (d) temperature variation over time.

compared to noble metals such as gold or platinum. These attributes make Ni a practical and economically viable choice for scalable perovskite solar cell fabrication.

Table 6 lists the material and structural parameters employed to configure the SCAPS-1D simulation architecture for the studied D_2CeX_6 (D = Ga, In, Tl; X = Cl, Br) double perovskite solar cells. The layer thickness is set at 50 nm for the transparent conducting oxide (FTO) and electron transport layer (WS₂), while the absorber layer is 800 nm, providing adequate optical absorption while minimizing recombination losses. The band gap (E_g) values, ranging from 0.859 eV (In_2CeBr_6) to 3.6 eV (FTO), define the spectral absorption range, with the perovskites covering a broad portion of the visible-near-infrared region for effective photon harvesting. Relative dielectric permittivity (ε_r) values between 3.36 and 13.60 influence the electrostatic screening of charge carriers, directly impacting exciton dissociation and defect tolerance. The electron affinity (χ), varying from 3.950 eV (WS₂) to 7.660 eV (In₂CeBr₆), determines band alignment at interfaces, which is crucial for efficient carrier transport and reduced energy barriers.

The effective density of states (N_V , N_C) for the valence and conduction bands, in the range of 1.429 \times 10¹⁹–1.492 \times 10¹⁹

 ${\rm cm}^{-3}$ (N_V) and 2.460 × 10¹⁹-2.980 × 10¹⁹ cm⁻³ (N_C) for perovskites, influences carrier concentration and recombination dynamics. Doping concentrations are assigned such that the FTO and WS₂ layers have a donor density (N_D) of 1×10^{18} cm⁻³ to ensure high conductivity, while the absorber layers are lightly p-type doped ($N_{\rm A}=1\times10^{17}~{\rm cm}^{-3}$) to facilitate charge separation. Hole mobility (μ_h) values for the absorbers range from 60 to 69 cm² V⁻¹ s⁻¹, and electron mobility (μ_p) from 35 to 46 cm² V^{-1} s⁻¹, supporting balanced charge transport and minimizing space-charge buildup. The defect density (N_t) , fixed at 1×10^{14} cm⁻³ for all layers, represents a moderate trap concentration to reflect realistic fabrication conditions while avoiding excessive recombination losses. Collectively, these parameters ensure that the simulated device reflects realistic material properties, enabling accurate prediction of optoelectronic performance and providing insight into how halide (Cl, Br) and A-site cation (Ga, In, Tl) substitutions influence solar cell efficiency.

This interface defect analysis in Table 7 is significant because the WS₂/absorber junction plays a critical role in determining overall solar cell efficiency, particularly in double perovskite-based devices like D₂CeX₆. Interface defects can act as recombination centers, trapping photogenerated carriers

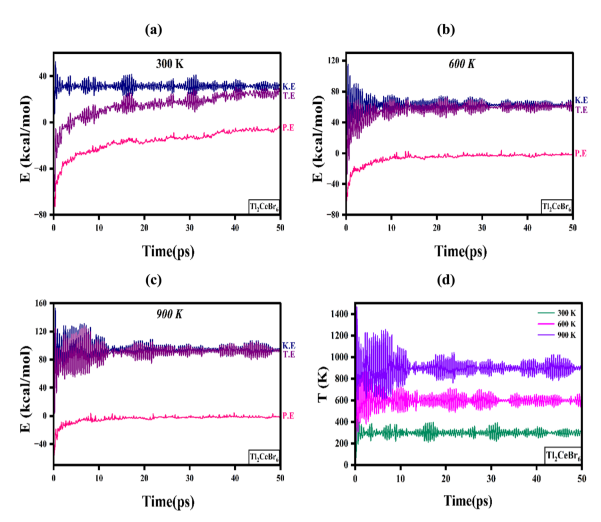


Fig. 20 Time evolution of potential, kinetic, and total energy for In_2CeBr_6 double perovskite at (a) 300 K, (b) 600 K, (c) 900 K, and (d) temperature variation over time.

before they contribute to the photocurrent, which reduces the open-circuit voltage ($V_{\rm OC}$), short-circuit current density ($J_{\rm SC}$), and ultimately the power conversion efficiency (PCE).⁹⁷ By quantifying parameters such as capture cross-section, defect type, and defect density, the simulation can realistically model the carrier recombination behavior at the heterojunction.

Table 7 summarizes the interface defect parameters used in the SCAPS-1D simulations for the heterojunctions between the electron transport layer (WS₂) and the absorber layers Ga₂CeCl₆, Ga₂CeBr₆, In₂CeCl₆, In₂CeBr₆, Tl₂CeCl₆, and Tl₂CeBr₆. For all interfaces, the capture cross-section for both electrons and holes is set at 1 imes 10⁻¹⁹ cm², representing a relatively low probability of carrier trapping, which helps to minimize interface recombination losses. The defect type is assigned as neutral, indicating that these traps do not possess a fixed positive or negative charge state, thereby reducing the likelihood of strong coulombic scattering. The total defect density is maintained at 1 × 10¹¹ cm⁻² for all combinations, corresponding to a moderate but realistic level of interface states that could arise from lattice mismatch or imperfections at the WS₂/ absorber junction. Such parameter uniformity across the Ga, In, and Tl-based D₂CeX₆ double perovskite absorbers ensures

a controlled comparison of their intrinsic optoelectronic performance, allowing any variations in simulated device efficiency to be attributed primarily to the absorber material properties rather than interface quality differences.

4.1. Band diagram

This analysis presents a comprehensive study of the energy band alignment for six perovskite absorber layers, Ga₂CeCl₆ (Fig. 22(a)), Ga₂CeBr₆ (Fig. 22(b)), In₂CeCl₆ (Fig. 22(c)), In₂CeBr₆ (Fig. 22(d)), Tl₂CeCl₆ (Fig. 22(e)), and Tl₂CeBr₆ (Fig. 15(f)) within the simulated solar cell architectures. Understanding the relative positions of the conduction band minimum (CBM), valence band maximum (VBM), and quasi-Fermi levels is essential for evaluating charge carrier generation, separation, and transport efficiency. By mapping these energy levels across the device thickness, the study reveals how the intrinsic bandgap and band offsets of each material dictate the overall photovoltaic response.98 Fig. 22 illustrates the comparative energy band diagrams of the six absorber materials; each modeled with a thickness of 0.80 μm. In each subfigure, the conduction band edge (E_C), valence band edge (E_V), electron quasi-Fermi level $(F_{\rm n})$, and hole quasi-Fermi level $(F_{\rm p})$ are plotted as functions of

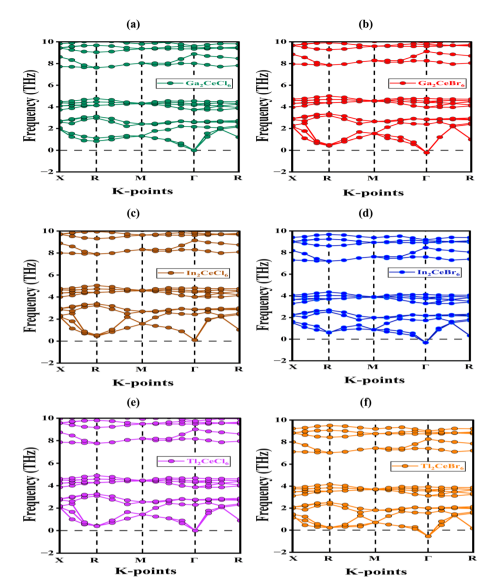


Fig. 21 Phonon dispersion curves of Ce-based (a) Ga_2CeCl_6 , (b) Ga_2CeBr_6 , (c) In_2CeCl_6 , (d) In_2CeBr_6 , (e) Tl_2CeCl_6 , and (f) Tl_2CeBr_6 double perovskites materials.

depth, providing a clear visualization of the bandgap and alignment characteristics. The results indicate systematic variations, where Ga₂CeCl₆ and Tl₂CeCl₆ exhibit the widest band gaps of 1.778 eV and 1.735 eV, respectively, while their bromide counterparts show reduced values of 1.276 eV and 1.304 eV. Similarly, In₂CeCl₆ possesses a bandgap of 1.550 eV, which decreases to 0.855 eV in In₂CeBr₆. These trends demonstrate that chloride-containing compounds generally feature wider band gaps with sharper conduction–valence band separations, leading to higher photon energy thresholds and making them favorable for achieving high open-circuit voltages. In contrast, bromide-based analogues exhibit narrower band gaps, enabling stronger absorption in the longer-wavelength region of the solar spectrum but also raising the possibility of enhanced recombination losses.

While the transport layers (FTO/WS₂) maintain consistent band offsets across all structures, the internal energy level positioning within the absorbers varies considerably, which

directly affects charge separation, extraction, and recombination dynamics. Overall, Ga_2CeCl_6 and Tl_2CeCl_6 are promising candidates for high- V_{OC} applications due to their wider band gaps and reduced thermalization losses, whereas In_2CeBr_6 , with the narrowest band gap, offers strong potential for nearinfrared light harvesting but requires careful optimization to suppress recombination and enhance device performance.

4.2. Optimizing absorber thickness for device performance

Analyzing the thickness-dependent behavior of Ce-based halide perovskite solar cells is vital for optimizing absorber layer design. In this study, the influence of absorber thickness on photovoltaic parameters PCE, FF, J_{SC} , and V_{OC} was systematically evaluated for Ga₂CeCl₆, Ga₂CeBr₆, In₂CeCl₆, In₂CeBr₆, Tl₂CeCl₆, and Tl₂CeBr₆. The results show that optimizing absorber thickness improves photon utilization across the solar spectrum while mitigating carrier recombination losses.⁹⁹

Table 5 Mulliken and Hirshfeld charge analysis of different atoms of D_2CeX_6 (where D=Ga, In, TI, and X=CI, Br) double perovskites

		Species	Mulliken atomic populations						
Compound	Change spilling		s	p	d	f	Total	Mulliken change	Hirshfeld change
Ga₂CeCl ₆	0.14%	Ga	1.93	0.32	10.00	0.00	12.25	0.75	0.48
		Ce	2.30	6.34	1.63	1.10	11.38	0.62	0.20
		Cl	1.95	5.40	0.00	0.00	7.35	-0.35	-0.19
Ga ₂ CeBr ₆	0.10%	Ga	2.11	0.39	10.00	0.00	12.50	0.50	0.45
		Ce	2.45	6.63	1.73	1.13	11.94	0.06	0.12
		Br	1.85	5.33	0.00	0.00	7.18	-0.18	-0.17
In ₂ CeCl ₆	0.14%	In	1.93	0.32	10.00	0.00	12.25	0.75	0.45
		Ce	2.30	6.34	1.63	1.09	11.37	0.63	0.21
		Cl	1.95	5.41	0.00	0.00	7.36	-0.36	-0.19
In₂CeBr ₆	0.16%	In	2.23	0.59	10.00	0.00	12.82	0.18	0.29
		Ce	2.43	6.47	1.89	1.15	11.94	0.06	0.07
		Br	1.81	5.26	0.00	0.00	7.07	-0.07	-0.11
Tl_2CeCl_6	0.11%	Tl	3.93	6.31	10.00	0.00	20.24	0.76	0.46
		Ce	2.30	6.34	1.63	1.09	11.36	0.64	0.21
		Cl	1.95	5.41	0.00	0.00	7.36	-0.36	-0.19
Tl₂CeBr ₆	0.13%	Tl	4.16	6.59	10.00	0.00	20.75	0.25	0.30
		Ce	2.44	6.47	1.89	1.15	11.95	0.05	0.07
		Br	1.82	5.27	0.00	0.00	7.09	-0.09	-0.11

Excessive thickness, however, restricts carrier diffusion and enhances bulk recombination,100 whereas insufficient thickness reduces light absorption and lowers photocurrent generation.¹⁰¹ These findings reveal material-specific trade-offs among absorption, carrier transport, and recombination, offering critical insights for designing efficient Ce-based halide perovskite absorbers. As illustrated in Fig. 23(a and b), increasing the absorber thickness from 0.2 µm to 0.8 µm leads to a pronounced improvement in both PCE and J_{SC} . This enhancement originates from more efficient photon absorption within the active layer, which increases carrier generation, while the diffusion length remains sufficient to suppress significant recombination losses. Within this range, the PCE and J_{SC} increase from 13.7% to 20.63% and 13.67 to 20.05 mA cm⁻² for Ga_2CeCl_6 , 17.18% to 24.34% and 17.74 to 25.35 mA cm⁻² for $In_2CeCl_6,\,14.10\%$ to 20.67% and 13.21 to 19.33 mA cm^{-2} for Tl_2CeCl_6 , 19.44% to 25.29% and 25.70 to 35.07 mA cm⁻² for Ga₂CeBr₆, 15.39% to 16.43% and 41.60 to 51.25 mA cm⁻² for In_2CeBr_6 , and 17.21% to 24.13% and 23.11 to 32.37 mA cm⁻² for Tl₂CeBr₆. When the thickness is further increased from 0.8 μm to 1.4 µm, the performance shows only limited gains. At this

stage, nearly all incident photons within the absorber's absorption range are harvested, so PCE and $J_{\rm SC}$ improvements primarily reflect marginal reductions in surface recombination. The highest efficiencies at 1.4 μm are 21.51% and 20.90 mA cm⁻² (Ga₂CeCl₆), 24.92% and 26.17 mA cm⁻² (In₂CeCl₆), 21.47% and 20.09 mA cm⁻² (Tl₂CeCl₆), 25.36% and 35.90 mA cm⁻² (Ga₂CeBr₆), 24.82% and 33.39 mA cm⁻² (Tl₂CeBr₆), and 15.41% and 51.67 mA cm⁻² (In₂CeBr₆). Beyond 1.4 μm , further thickness increases up to 2.1 μm yield negligible improvements, as optical absorption saturates and longer carrier transport pathways enhance bulk recombination.

The $V_{\rm OC}$ remains nearly constant across thickness variations, emphasizing that it is primarily dictated by the intrinsic bandgap and defect-related recombination of the material rather than absorber thickness. The simulated $V_{\rm OC}$ values are 1.259 V (Ga₂CeCl₆), 1.163 V (In₂CeCl₆), 1.297 V (Tl₂CeCl₆), 0.913 V (Ga₂CeBr₆), 0.497 V (In₂CeBr₆), and 0.969 V (Tl₂CeBr₆). These values confirm that Cl-based absorbers favor higher $V_{\rm OC}$, whereas Br-based compounds favor higher $J_{\rm SC}$, consistent with bandgap-dependent photovoltaic theory. Regarding the FF, increasing thickness from 0.2 μ m to 0.8 μ m causes slight

Table 6 Configurational parameters for the simulated device architecture

Parameters	FTO	WS_2	${\rm Ga_2CeCl_6}$	Ga ₂ CeBr ₆	In_2CeCl_6	In ₂ CeBr ₆	${\rm Tl_2CeCl_6}$	${\rm Tl_2CeBr_6}$
Thickness (nm)	50	50	800	800	800	800	800	800
Band gap, $E_{\rm g}$ (eV)	3.6	2.2	1.733	1.273	1.550	0.859	1.755	1.364
Dielectric permittivity, \in_{r}	10	13.60	3.520	3.750	3.650	3.360	3.540	3.36
Electron affinity, χ (eV)	4.5	3.950	4.472	5.410	4.560	7.660	4.280	7.160
$V_{\rm B}$ effective density of states, $N_{\rm V}$ (cm ⁻³)	1.8×10^{19}	2.2×10^{18}	1.450×10^{19}	1.492×10^{19}	1.489×10^{19}	1.440×10^{19}	1.478×10^{19}	1.429×10^{19}
$C_{\rm B}$ effective density of states, $N_{\rm C}$ (cm ⁻³)	2×10^{18}	1.8×10^{19}	2.460×10^{19}	2.476×10^{19}	2.980×10^{19}	2.930×10^{19}	2.504×10^{19}	2.483×10^{19}
Donor density, $N_{\rm D}$ (cm ⁻³)	1×10^{18}	1×10^{18}	0	0	0	0	0	0
Acceptor density, N _A (cm ⁻³)	0	0	1×10^{17}	1×10^{17}	1×10^{17}	1×10^{17}	1×10^{17}	1×10^{17}
Hole mobility, μ_h (cm ² V ⁻¹ s ⁻¹)	20	100	65	60	67	65	69	69
Electron mobility, $\mu_{\rm n}$ (cm ² V ⁻¹ s ⁻¹)	100	100	40	35	37	42	46	46
Total defect density, $N_{\rm t}$ (cm ⁻³)	1×10^{14}	1×10^{14}	1×10^{14}	1×10^{14}	1×10^{14}	1×10^{14}	1×10^{14}	1×10^{14}

Capture cross section: Electrons/holes (cm²) Defect type

Table 7 The simulated design of WS₂/D₂CeX₆ solar cells' specific interface input parameters

Total interface defect density (cm⁻²) Interfaces 1×10^{-19} 1×10^{11} WS2/Ga2CeCl6 Neutral 1×10^{-19} WS2/Ga2CeBr6 Neutral 1×10^{11} 1×10^{-19} 1×10^{11} WS2/In2CeCl6 Neutral 1×10^{-19} 1×10^{11} WS2/In2CeBr6 Neutral 1×10^{-19} 1×10^{11} WS2/Tl2CeCl6 Neutral 1×10^{-19} $1\times 10^{11}\,$ WS2/Tl2CeBr6 Neutral

decreases for most materials, reflecting the interplay between increased series resistance and enhanced absorption. FF decreases from 82.67% to 81.75% (Ga₂CeCl₆), 82.39% to 82.55% (In_2CeCl_6) , 83.26% to 82.44% (Tl_2CeCl_6) , 80.13% to 79.03% (Ga₂CeBr₆), 71.35% to 64.56% (In₂CeBr₆), and 77.03% to 76.91% (Tl₂CeBr₆). When the thickness is further increased to 2.3 μm, FF stabilizes for most absorbers, except In₂CeBr₆, which decreases from 64.56% to 58.82% due to increased bulk recombination and resistive losses. Overall, these results demonstrate that both cation choice (Ga, In, Tl) and anion substitution (Cl vs. Br) significantly influence thicknessdependent performance: Br-based perovskites achieve higher $J_{\rm SC}$ owing to narrower band gaps, while Cl-based perovskites retain higher V_{OC} , and the optimal thickness for balancing light absorption, carrier transport, and recombination generally lies at 0.8 µm for all materials.

4.3. Optimizing total defect density for device performance

As depicted in Fig. 24(a and b), varying the effect of varying defect density from 1011 to 1018 cm-3 on the photovoltaic parameters PCE, FF, JSC, and VOC of Ga2CeCl6, In2CeCl6, Tl2-CeCl₆, Ga₂CeBr₆, In₂CeBr₆, and Tl₂CeBr₆-based solar cells. This is commonly referred to as defect tolerance analysis, which quantifies how device efficiency responds to intrinsic material imperfections. Defect-tolerant materials exhibit minimal performance degradation even at higher defect concentrations, an essential attribute for the fabrication of scalable, costeffective, and durable perovskite solar technologies. Our results reveal a progressive degradation in all photovoltaic parameters with increasing defect density. At low defect levels (10¹¹ to 10¹⁴ cm⁻³), performance remains nearly unchanged, with only slight reductions due to enhanced Shockley-Read-Hall (SRH) recombination and trap-assisted non-radiative processes that slightly shorten carrier lifetimes. For instance, when the acceptor concentration increases, only slight variations are observed in the photovoltaic parameters. The PCE, FF, $J_{\rm SC}$, and $V_{\rm OC}$ values remain almost stable for all $D_2{\rm CeX}_6$ compounds, showing minimal fluctuations within a narrow range. For example, Ga₂CeCl₆ exhibits a minor change from 19.9% to 20.6% (PCE) and 1.26 to 1.26 V (V_{OC}), while In₂CeCl₆ and Tl2CeCl6 show similarly small differences. A comparable trend is seen for the bromide systems, where Ga₂CeBr₆, In₂-CeBr₆, and Tl₂CeBr₆ display only slight declines or adjustments in efficiency, current, and voltage.

However, when the defect density exceeds 10¹⁴ cm⁻³, a noticeable performance degradation occurs. At a higher defect density of around 10¹⁸ cm⁻³, this decline becomes severe, with significant reductions observed in all key photovoltaic parameters. The PCE, FF, J_{SC} , and V_{OC} values drop sharply for each material, indicating increased non-radiative recombination and hindered charge transport. These steep declines stem from the fact that higher defect concentrations introduce deep-level trap states, which increase carrier recombination rates, lower carrier diffusion lengths, and reduce quasi-Fermi level splitting, thereby strongly suppressing both $V_{\rm OC}$ and $J_{\rm SC}$. Overall, the analysis highlights that cation (Ga \rightarrow In \rightarrow Tl) and anion (Cl \rightarrow Br) substitutions significantly affect defect tolerance, with Clbased compounds generally showing better stability against defect-induced performance loss compared to their Br counterparts. This insight provides critical guidance for material selection in designing defect-tolerant halide double perovskite solar cells.

4.4. Effect of shallow acceptor density of the absorber layer

Fig. 26(a and b) demonstrates how changes in the shallow acceptor density (N_{Δ}) from 10^{13} to 10^{20} cm⁻³ influence key photovoltaic metrics PCE, FF, J_{SC} , and V_{OC} . This trend is observed in Ga₂CeCl₆, In₂CeCl₆, Tl₂CeCl₆, Ga₂CeBr₆, In₂CeBr₆, and Tl₂CeBr₆ solar cells. The shallow acceptor density denotes the number of shallow-level acceptor impurities per unit volume in a semiconductor, which governs its p-type (D₂CeX₆) conductivity by generating holes as charge carriers. While higher N_A values enhance hole concentration and initially improve device performance, excessive doping leads to stronger recombination losses and reduced efficiency. 102 An optimal range of 10¹⁵-10¹⁷ cm⁻³ achieves the best trade-off between carrier generation and recombination, making shallow acceptor density a key factor in the design of high-efficiency photovoltaic and electronic devices. 103 As shown in Fig. 26(a and c), increasing the acceptor concentration (N_A) from 10^{13} to 10^{17} cm⁻³ leads to a steady improvement in both V_{OC} and FF across all D₂CeX₆-based solar cells. Within this range, each compound exhibits a noticeable rise in voltage and fill factor, indicating enhanced charge separation and reduced recombination. Further increasing N_A up to around 1020 cm⁻³ continues to improve these parameters, reaching their maximum values for all materials. However, an excessively high N_A may also promote additional recombination losses. Generally, a moderate doping

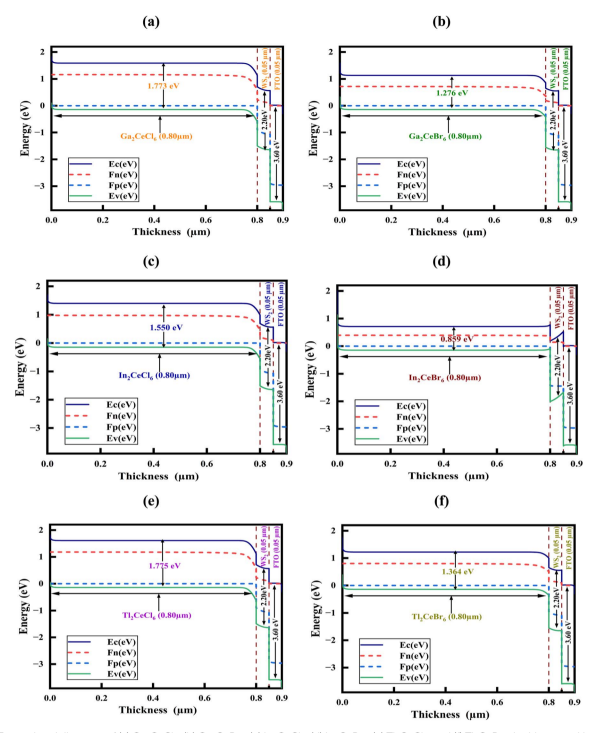


Fig. 22 Energy band diagrams of (a) Ga_2CeCl_6 , (b) Ga_2CeBr_6 , (c) In_2CeCl_6 , (d) In_2CeBr_6 , (e) Tl_2CeCl_6 , and (f) Tl_2CeBr_6 double perovskite absorber layers in the simulated solar cell structures.

level of about 10¹⁷ cm⁻³ offers the most balanced performance, ensuring optimal carrier transport and device efficiency.

As illustrated in Fig. 25(b), increasing the $N_{\rm A}$ from 10^{13} to 10^{17} cm⁻³ results in nearly constant $J_{\rm SC}$ values for most of the investigated D₂CeX₆ solar cells, except for In₂CeBr₆. The $J_{\rm SC}$ remains approximately 20.05 mA cm⁻² (Ga₂CeCl₆), 25.36 mA cm⁻² (In₂CeCl₆), 19.33 mA cm⁻² (Tl₂CeCl₆), 35.07 mA cm⁻² (Ga₂CeBr₆), and 32.37 mA cm⁻² (Tl₂CeBr₆), while In₂CeBr₆

shows a slight increase from 50.35 to 51.25 mA cm $^{-2}$ over the same $N_{\rm A}$ range. With further doping up to 10^{20} cm $^{-3}$, $J_{\rm SC}$ values largely remain constant, except for slight reductions observed in ${\rm Ga_2CeCl_6}$ (18.70 mA cm $^{-2}$) and ${\rm Tl_2CeCl_6}$ (16.53 mA cm $^{-2}$). This indicates that $J_{\rm SC}$ in these perovskite systems is less sensitive to shallow acceptor density variations compared to $V_{\rm OC}$ and FF, suggesting that carrier collection efficiency remains largely

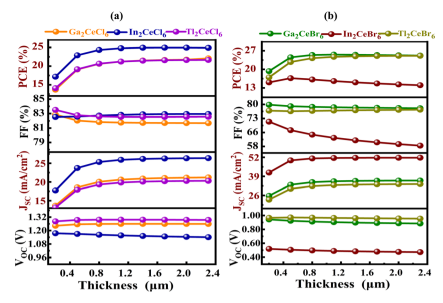


Fig. 23 Optimization of absorber layer thickness on PCE, FF, J_{SC}, and V_{OC} for (a) D₂CeCl₆ and (b) D₂CeBr₆ double perovskite materials.

unaffected by excess doping, except in specific cases where recombination effects begin to dominate.

As illustrated in Fig. 25(d), the variation of PCE exhibits a similar trend to that of $V_{\rm OC}$ and FF. When the acceptor concentration ($N_{\rm A}$) increases to around 10^{17} cm⁻³, the PCE values show a notable improvement for both chloride- and bromide-based devices. Upon further increasing $N_{\rm A}$ to approximately 10^{20} cm⁻³, the efficiencies rise even higher across all configurations, with $\rm Ga_2CeBr_6$ and $\rm Tl_2CeBr_6$ achieving the best performance. This overall enhancement with increasing $N_{\rm A}$ suggests improved charge transport and reduced recombination losses, leading to more efficient device operation. These results indicate that moderate $N_{\rm A}$ of 10^{17} cm⁻³ significantly enhances device performance by improving carrier

concentration and transport, while excessive doping continues to raise PCE but may approach saturation due to increased recombination losses. The optimized $N_{\rm A}$ for $V_{\rm OC}$, $J_{\rm SC}$, FF, and PCE was found to be 10^{17} cm⁻³ for all the studied D₂CeX₆ materials. At this $N_{\rm A}$ value, the devices achieve a balanced enhancement of carrier concentration and transport, resulting in maximum overall photovoltaic performance before recombination losses from excessive doping become significant.

4.5. Impact on temperature-dependent photovoltaic performance

This temperature-dependent photovoltaic performance analysis is important because it reveals how different D-site cations (Ga,

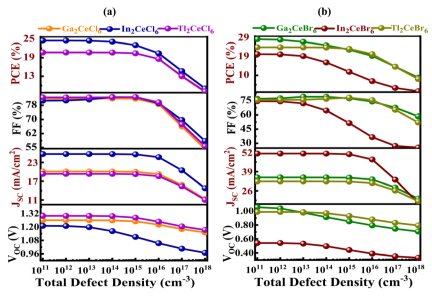


Fig. 24 Optimization of total defect density on PCE, FF, J_{SC} , and V_{OC} for (a) D_2CeCl_6 and (b) D_2CeBr_6 double perovskite materials.

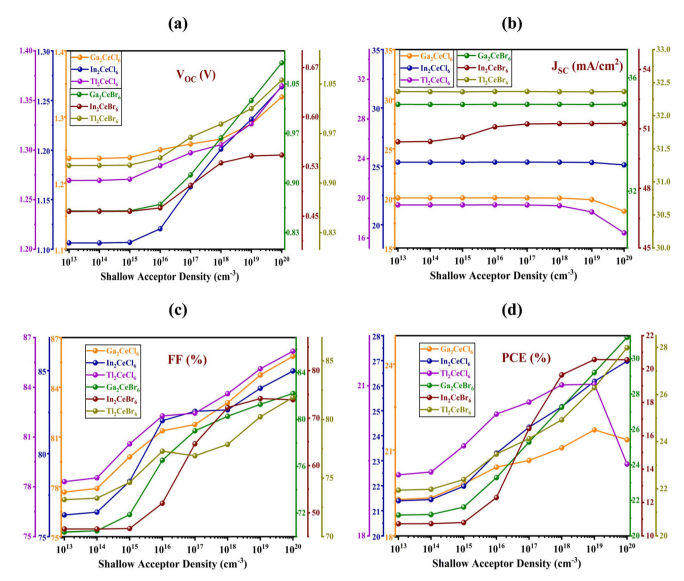


Fig. 25 Optimization of shallow acceptor density on (a) V_{OC} , (b) J_{SC} , (c) FF, and (d) PCE for D_2CeX_6 double perovskite materials.

In, Tl) and halide anions (Cl, Br) influence the thermal stability and efficiency retention of double perovskite solar cells under varying operating conditions. By examining the trends in $V_{\rm OC}$, J_{SC} , FF, and PCE across a broad temperature range, it becomes possible to identify materials that maintain high output and stability despite thermal stress, an essential requirement for real-world solar energy deployment where temperature fluctuations are inevitable. Such insights not only highlight the tradeoffs between voltage, current generation, and charge transport in different compositions but also pinpoint the most promising candidates, like Tl-based compounds, for achieving high efficiency and durability, guiding the design of next-generation, thermally robust photovoltaic devices. Fig. 26 presents the temperature-dependent photovoltaic performance parameters of six double perovskite halides: Ga₂CeCl₆, In₂CeCl₆, Tl₂CeCl₆, Ga₂CeBr₆, In₂CeBr₆, and Tl₂CeBr₆, displayed across four figures: Fig. 26(a). V_{OC}, Fig. 26(b). J_{SC}, Fig. 26(c). FF, and Fig. 26(d). PCE. Fig. 26(a), $V_{\rm OC}$ decreases consistently with increasing temperature for all materials, with Ga₂CeCl₆ and Tl₂CeCl₆ showing the

highest initial values at low temperatures, while bromide-based compounds generally exhibit lower $V_{\rm OC}$ due to their narrower band gaps. In Fig. 26(b), $J_{\rm SC}$ slightly increases with temperature for bromides (notably ${\rm Tl_2CeBr_6}$) but remains relatively stable for chlorides, reflecting enhanced carrier generation in narrower-bandgap materials at higher temperatures. Fig. 26(c) shows FF values slightly decreasing with temperature, with Tl-based compounds maintaining higher FF across the range, indicating better charge extraction stability.

Finally, Fig. 26(d) reveals that PCE declines with rising temperature for all materials, with Tl₂CeCl₆ and Tl₂CeBr₆ retaining superior efficiencies compared to Ga- and In-based counterparts, suggesting that Tl-containing perovskites offer a better balance between voltage, current, and charge transport. This analysis is crucial because it reveals how intrinsic material properties and halide composition govern the thermal stability and operational performance of perovskite solar cells, aiding in identifying the most thermally robust candidates for high-temperature photovoltaic applications.

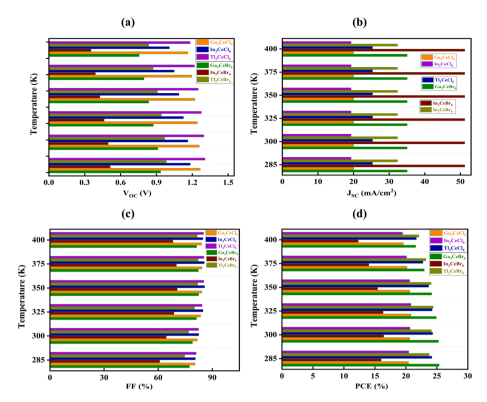


Fig. 26 Temperature-dependent photovoltaic parameters of (a) $V_{\rm OC}$, (b) $J_{\rm SC}$, (c) FF, and (d) PCE of D₂CeX₆-based solar cells.

4.6. JV and QE properties

A thorough examination of the quantum efficiency (QE) and current density–voltage (*J–V*) characteristics is indispensable for elucidating the optoelectronic behavior of photovoltaic devices. These parameters function as critical diagnostic indicators, providing quantitative insights into the efficiency of photon absorption, carrier photogeneration, and charge transport dynamics.¹⁰⁴ The QE spectrum delineates the proportion of incident photons converted into collected charge carriers, thereby serving as a direct measure of spectral responsivity. In parallel, the *J–V* profile describes the current density as a function of applied bias, enabling a rigorous evaluation of key device performance metrics.¹⁰⁵

Fig. 27(a) and (b) depict the J-V and QE characteristics of D_2CeX_6 -based double perovskite solar cells. The optimized Al/FTO/WS₂/D₂CeX₆/Au configuration was obtained using carefully tuned parameters: an absorber thickness of 800 nm, a total defect density of 10^{14} cm⁻³, and a shallow acceptor density of 10^{17} cm⁻³. As shown in Fig. 27(a), the simulated J-V curves yield J_{SC} of 20.05, 25.35, 19.33, 35.07, 51.25, and 32.37 mA cm⁻² for Ga_2CeCl_6 , In_2CeCl_6 , Tl_2CeCl_6 , Ga_2CeBr_6 , In_2CeBr_6 , and Tl_2-CeBr_6 , respectively, under zero voltage. With increasing voltage, J_{SC} initially remains nearly constant but then decreases sharply to zero at the corresponding V_{OC} of 1.259, 1.163, 1.297, 0.913, 0.497, and 0.969 V. This reduction in photocurrent at higher voltage is primarily attributed to carrier saturation and recombination effects. As shown in Fig. 27(b), the simulated QE

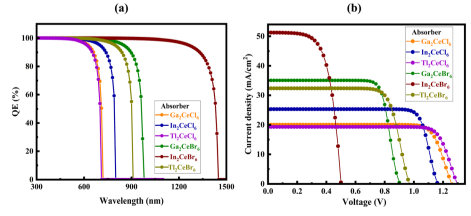


Fig. 27 (a) J-V characteristics and (b) QE spectra of D_2CeX_6 -based double perovskite solar cells.

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spectra exhibit an initial maximum value of nearly 100% for all compositions at short wavelengths, indicating excellent photonto-carrier conversion efficiency under strong absorption conditions. With increasing wavelength, QE remains relatively constant over the visible region but decreases sharply to zero at the respective cutoff wavelengths of 720, 800, 710, 980, 1450, and 910 nm for Ga₂CeCl₆, In₂CeCl₆, Tl₂CeCl₆, Ga₂CeBr₆, In₂-CeBr₆, and Tl₂CeBr₆, respectively. These cutoff positions are strongly correlated with the optical band gaps, where larger band gaps (e.g., Ga₂CeCl₆, Tl₂CeCl₆) correspond to shorter cutoff wavelengths, while smaller band gaps (e.g., In₂CeBr₆) extend absorption into the near-infrared region. The observed trends highlight the role of cationic substitution (Ga \rightarrow In \rightarrow Tl) and halide exchange (Cl → Br) in tailoring the optical response. Replacing Cl with Br reduces the band gap, redshifting the absorption edge and enhancing long-wavelength OE, thereby improving light-harvesting capability. Similarly, the choice of the trivalent cation influences orbital hybridization and band dispersion, which governs carrier generation and transport. These theoretical insights confirm that compositional engineering in D₂CeX₆ double perovskites offers a viable pathway to optimize the spectral response and photovoltaic performance of the devices.

4.7. Device overall performance

Optimization of the device parameters was carried out with an absorber thickness of 800 nm, a total defect density of 10¹⁴ cm⁻³, and a shallow acceptor density of 10¹⁷ cm⁻³. Under these optimized conditions, the photovoltaic parameters were measured as follows: Al/FTO/WS $_2$ /Ga $_2$ CeCl $_6$ /Au delivered a $V_{\rm OC}$ of 1.259 V, a J_{SC} of 20.05 mA cm⁻², an FF of 81.75%, and a PCE of 20.64%. Al/FTO/WS₂/In₂CeCl₆/Au exhibited a V_{OC} of 1.163 V, a $J_{\rm SC}$ of 25.35 mA cm $^{-2}$, an FF of 82.55%, and a PCE of 24.34%. As shown in Fig. 28, the Al/FTO/WS₂/Tl₂CeCl₆/Au device achieved the highest $V_{\rm OC}$ (~1.29 V) but moderate $J_{\rm SC}$ (~19 mA cm⁻²), resulting in a PCE of about 20.6%. In contrast, the bromide-based cells showed lower $V_{\rm OC}$ but higher $J_{\rm SC}$ values. Among them, Al/FTO/WS2/Ga2CeBr6/Au reached the best efficiency (\sim 25.3%) with a balanced $V_{\rm OC}$ (\sim 0.91 V) and $J_{\rm SC}$ (\sim 35 mA cm⁻²). The In-based device exhibited the highest J_{SC} (~51 mA cm⁻²) but lowest $V_{\rm OC}$ (~0.50 V), giving a moderate PCE (~16.4%), while Al/FTO/WS₂/Tl₂CeBr₆/Au showed a wellbalanced performance (~24.1% PCE). Overall, chloride systems favor higher voltage, whereas bromide systems enhance current generation.

A comparative evaluation highlights distinct trends associated with cationic and anionic substitutions. The chloride-based systems (Ga_2CeCl_6 , In_2CeCl_6 , Tl_2CeCl_6) generally provide higher V_{OC} owing to their wider band gaps, although at the cost of reduced photocurrent. Among them, Ga_2CeBr_6 achieves the highest efficiency of 25.29%, reflecting a favorable balance between voltage and current density. The bromide-based systems (Ga_2CeBr_6 , In_2CeBr_6 , Tl_2CeBr_6), in contrast, show red-shifted absorption edges that enhance photocurrent but simultaneously lower the voltage due to narrower band gaps. In_2CeCl_6 demonstrates the best overall performance with

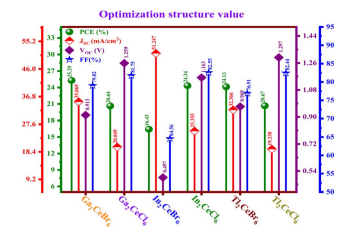


Fig. 28 Comparison of $V_{\rm OC}$, $J_{\rm SC}$, FF, and PCE for Ga₂CeBr₆, Ga₂CeCl₆, In₂CeBr₆, In₂CeCl₆, Tl₂CeBr₆, and Tl₂CeCl₆ absorber-based solar cells.

an efficiency of 24.34%, arising from a well-balanced combination of $V_{\rm OC}$ and $J_{\rm SC}$. $\rm In_2CeBr_6$, despite delivering the highest photocurrent of 51.25 mA cm⁻², suffers from severe voltage loss, restricting its efficiency to only 16.43%. $\rm Tl_2CeBr_6$ exhibits intermediate behavior, with an efficiency of 24.13%, reflecting the role of cation substitution in fine-tuning electronic structure and carrier transport. These findings underscore that chloride-based compounds are favorable for higher voltage stability, while bromide substitution enhances photocurrent generation. Among the studied materials, $\rm Ga_2CeBr_6$ emerges as the most promising candidate, demonstrating that halide exchange combined with careful cation selection can effectively optimize photovoltaic performance in $\rm D_2CeX_6$ double perovskite solar cells.

4.8. Limitations of the study

This study has several limitations. First, the initial DOS calculations employed relatively large Gaussian smearing values, which increased numerical broadening and led to inconsistencies between the DOS and band structure bandgap. In future work, this issue can be explored in greater depth to further improve the reliability of the results. Nevertheless, selecting an appropriate smearing parameter remains essential for accurately determining the electronic structure.

Second, in the earlier version, the convergence criteria used for the phonon dispersion calculations were not sufficiently strict, and relatively small supercells were employed. As a result, the acoustic phonon frequencies at the Γ -point did not appear as zero, and slight imaginary modes were observed in some bromine-containing compounds. Although these issues have been significantly corrected in the revised calculations, very small negative or near-zero frequencies still appear near the Γ -point for Ga_2CeBr_6 , In_2CeBr_6 , and Tl_2CeBr_6 . These features indicate possible soft phonon modes or minor lattice instabilities. In the future, the stability of these Br-based compounds may be enhanced through pressure, temperature effects, or doping. Finally, the phonon calculations were performed within the harmonic approximation at 0 K. Anharmonic effects, thermal stabilization, and pressure-dependent behavior were

not considered and may influence the dynamical stability of the bromine-containing compounds.

5 Conclusion

Based on the structural, electronic, optical, and molecular dynamics analyses, the overall findings highlight the promising stability and functionality of the investigated materials. This study presented a comprehensive first-principles and devicelevel investigation of lead-free, vacancy-ordered halide double perovskites D_2CeX_6 (D = Ga, In, Tl; X = Cl, Br) for solar energy applications. All compounds crystallize in a stable cubic $Fm\bar{3}m$ phase, with negative formation enthalpies and tolerance factors confirming thermodynamic and structural stability. Halide substitution (Cl \rightarrow Br) enlarged lattice constants, narrowed band gaps, and improved visible-light absorption, while D-site cation variation (Ga → In → Tl) tuned bonding strength, charge distribution, and mechanical robustness. Electronic structure analysis revealed tunable direct and indirect band gaps (0.859 to 3.632 eV), with Ce-4f and halogen-p orbitals dominating near the band edges, supporting efficient chargetransfer transitions. Optical evaluations confirmed strong UVvisible absorption coefficients ($\alpha > 10^4 \text{ cm}^{-1}$), high refractive indices, and large dielectric constants, especially in In₂CeBr₆ and Tl₂CeBr₆, making them promising for optoelectronics and photodetectors. Mechanical analysis showed ductile behavior across the series. Tl₂CeBr₆ had the highest bulk, shear, and Young's moduli, while In₂CeBr₆ exhibited excellent isotropy and flexibility suitable for stable thin-film or flexible solar cells. Phonon dispersion spectra confirm full dynamical stability for Ga₂CeCl₆, In₂CeCl₆, and Tl₂CeCl₆, while minor soft modes in bromides are expected to be stabilized at finite temperatures. Importantly, molecular dynamics simulations performed at 300 K, 600 K, and 900 K under NPT conditions demonstrate that all studied materials maintain thermal stability across a wide temperature range, highlighting their robustness for practical applications. Device simulations using SCAPS-1D demonstrated high photovoltaic potential. The device parameters were optimized with an absorber thickness of 800 nm, a total defect density of 1014 cm-3, and a shallow acceptor density of 1017 cm⁻³. In₂CeCl₆ reached an efficiency of 24.34%, while Ga₂CeBr₆ achieved the highest PCE of 25.29% due to a favorable trade-off between voltage and current. Bromide compounds enhanced photocurrent through band-gap narrowing, whereas chloride compounds delivered higher voltages. D2CeX6 family exhibits stability, tunable optoelectronic response, and strong photovoltaic performance, with In₂CeCl₆, Tl₂CeBr₆, and Ga₂CeBr₆ emerging as the most promising absorbers. These findings provide valuable guidance for experimental synthesis and compositional engineering of sustainable, lead-free perovskites for next-generation solar cells and optoelectronic technologies.

Ethical approval

The manuscript's authors agree that there are no research involving human participants, human data or tissue, or animal subjects.

Author contributions

Rifat Rafiu, Md. Sakib Hasan Saikot, Md. Azizur Rahman, Imtiaz Ahamed Apon, Imtiaz Ahmed Talukder: methodology, validation, software, conceptualization, investigation, formal analysis, data curation, visualization, writing – original draft, and review and editing. Karim KRIAA, N. Sfina, Mohd Taukeer Khan, S. AlFaify, Noureddine Elboughdiri: investigation, validation, software, formal analysis, data curation, writing – original draft, and review and editing.

Conflicts of interest

The authors have no conflicts of interest.

Data availability

Data will be made available on reasonable request.

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