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Evaluation of design and device parameters for lead-free Sr₃PBr₃/Sr₃NCl₃ duel-layer perovskite photovoltaic device technology

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The study looks into how Sr₃PBr₃ and Sr₃NCl₃ double perovskite materials can be used as absorbers in perovskite solar cells (PSCs). Computational Sr₃PBr₃ and Sr₃NCl₃ simulations were employed to assess the performance of each absorber together with electron transport layers (ETL), with a particular emphasis on optimizing ETL thickness to improve charge transport and synchronize current outputs. The simulations yielded valuable insights into the electronic and optical characteristics of the individual absorbers. Subsequently, a tandem simulation was performed to adjust each layer's thickness, ensuring that both devices' current outputs were aligned for maximum system efficiency. The findings revealed that the tandem configuration of Sr₃PBr₃ and Sr₃NCl₃ surpassed the performance of the individual absorber setups, attributed to the optimized ETL thicknesses that enhanced charge transport and facilitated effective current matching. This study makes a significant contribution to the design and optimization of tandem PSCs utilizing Sr₃PBr₃ and Sr₃NCl₃ absorbers, paving the way for improved overall device efficiency. We investigated three device configurations to find the optimum structure. FTO/SnS₂/Sr₃PBr₃/Ni, FTO/SnS₂/Sr₃NCl₃/Ni, and FTO/SnS₂/Sr₃PBr₃/Sr₃NCl₃/Ni are considered as Device-I, II, and III. In Device-I, the execution parameters are power conversion efficiency (PCE) of 24.26%, an open-circuit voltage (V_{OC}) of 1.23 V, a short-circuit current density (J_{SC}) of 24.65 mA cm⁻², and a fill factor (FF) of 87.42%. For Device-II, PCE, FF, V_{OC} , and J_{SC} are correspondingly 20.35%, 87.91%, 1.28 V, and 18.07 mA cm⁻². The further refined tandem configuration achieved a PCE of 30.32%, with a V_{OC} of 1.27 V, an FF of 90.14%, and a J_{SC} of 26.44 mA cm⁻², demonstrating the potential of this methodology in enhancing PSC performance.

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1. Introduction

The need for energy worldwide is steadily increasing, driven by an expanding population, which raises concerns about the exhaustion of nonrenewable supplies and the associated ecological consequences. This situation has catalyzed a shift towards renewable energy sources, with solar energy standing out as a prominent choice due to its plentiful, clean, and limitless characteristics.¹ The adoption of solar energy not only mitigates environmental challenges but also fosters sustainable

energy solutions, such as solar architecture,² solar heating,³ photovoltaics,4 and photocatalytic water-splitting technologies.^{5,6} Among these, photovoltaics has attracted considerable interest because of their ability to use the PV effect to turn sunlight into energy. Solar cells (SCs), which are essential components of photovoltaics, capture sunlight to generate electricity and can be divided into three generations. The first generation is based on wafer technology; the second generation utilizes thin films; and the third generation features organic materials. Although first- and second-generation SCs have historically led the market, their high production costs, intricate manufacturing processes, and environmental concerns have limited their widespread use. Researchers are diligently investigating new materials for solar cells (SCs) that prioritize both cost-effectiveness and environmental sustainability. A variety of SC types have been created, including polycrystalline and single-crystalline silicon SCs,7 copper indium gallium selenide (CIGS) SCs,8 cadmium telluride (CdTe) SCs,9 quantum dot SCs,10 organic SCs,11 and perovskite solar cells (PSCs),12 each presenting unique benefits. In the commercialization process of

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SCs, PCE and material costs are critical considerations. Currently, third-generation silicon-based SCs dominate the market, achieving impressive PCEs of approximately 25-26%.13 However, innovative third-generation PSCs are emerging as formidable alternatives to conventional silicon SCs, boasting remarkable PCEs of up to 22.1%.14,15 To further improve efficiency, researchers are utilizing various strategies, including structural and compositional engineering, optimizing ETLs and HTLs, and investigating double and graded absorber configurations in PSCs.¹⁶⁻¹⁹ The Sr₃PBr₃ and Sr₃NCl₃ structure, characterized by the substitution of P (phosphorus) and N (nitrogen) for lead, is recognized as one of the most encouraging configurations due to its superior crystalline nature, extended carrier duration, and enhanced stability in comparison to lead-based perovskites.^{20,21} Double perovskite compounds present considerable advantages over their single perovskite counterparts across various applications. Notable benefits of double perovskites include increased stability, as the incorporation of two different cations in the A-site of the crystal lattice bolsters structural integrity and resistance to degradation, making them more suitable for prolonged device operation.²² Furthermore, double perovskites facilitate property tuning through the modification of cation combinations at the A-site, allowing for precise control over bandgap, electronic structure, and photosensitivity, thus enabling customization to fulfill specific application requirements.23 Double perovskite materials present a wider absorption spectrum compared to single perovskites, primarily due to the presence of multiple cations. This diversity can lead to an increased bandgap, enabling the absorption of a broader range of solar photons. As a result, the enhanced absorption improves the utilization of sunlight, which in turn boosts photocurrent generation and elevates power conversion efficiency. Furthermore, double perovskites typically demonstrate reduced carrier recombination rates. The incorporation of various cations helps to optimize energy-level alignments, thereby minimizing nonradiative recombination and prolonging the lifetimes of charge carriers.24-26 To optimize the performance of double-layer perovskite materials like Sr₃PBr₃ and Sr₃NCl₃ in solar cells, composition tuning and improving crystallinity through advanced deposition methods are crucial. Selecting a suitable electron transport layer (ETL) with high electron mobility is essential for efficient charge extraction, and optimizing the ETL thickness can help balance electron collection and recombination losses. Implementing a double-layer architecture can enhance charge separation by minimizing energy barriers at the interfaces. Incorporating light-trapping techniques and optical coatings can significantly boost light absorption and overall efficiency. Effective encapsulation protects the perovskite layer from moisture, improving device longevity. Exploring tandem configurations that combine perovskite layers with other materials can maximize energy harvesting across a broader spectrum. Finally, using computer simulations to model device performance can guide material and structural optimizations for improved efficiency. These materials also exhibit excellent charge transport characteristics, attributed to their unique crystal structures and the deliberate inclusion of different cations, which enhance charge

carrier mobility and promote effective charge extraction. These features render double perovskites particularly appealing for application in diodes that produce light, solar cells, and photosensors, as they provide improved stability, adjustable propabsorption erties, expanded capabilities, decreased recombination rates, and enhanced charge transport, all of which contribute to the advancement of more efficient and stable optoelectronic devices. The ETL, characterized as n-type, is essential for influencing the $J_{\rm SC}$ and $V_{\rm OC}$ in photovoltaic devices. The V_{OC} is notably affected by the alignment of the band at the junction of the absorber and backstop layers. This alignment, named the conduction band offset (CBO), can be classified into three types: cliff, plane, or spike. A spike at the interface is beneficial as it minimizes recombination losses, while a cliff alignment decreases in V_{OC} . To ensure efficient transmission of electrons between the ETL and the absorber, thereby maintaining the fill factor and density of current, the height of the spike should be kept to a minimum. Tin sulfide (SnS₂) is chosen for the ETL in this solar cell because of its advantageous electrical properties, including a band gap of approximately 2.24 eV, which positions it as an effective ETL material. SnS₂ is recognized for its low absorption losses, high electron mobility, and good visibility spectrum transparency. The way its conduction band aligns with absorber materials like Sr₃PBr₃ facilitates efficient electron extraction and inhibits hole back-transfer. Furthermore, SnS₂ establishes stable interfaces with the FTO foundation; this reduces the losses from interface recombination and increases adhesion. Its chemical stability and ease of fabrication further confirm its appropriateness for solar energy applications. In the structure, the absence of a hole transport layer (HTL) could be a deliberate design decision based on the favorable alignment of energy levels between the materials. This alignment allows for efficient hole extraction directly to the Ni electrode, eliminating the need for a dedicated HTL. Such an approach simplifies the device architecture, reducing fabrication complexity and minimizing potential interface losses that might occur with an additional layer. If the electron transport layer (SnS₂) effectively facilitates electron extraction and the absorber materials (Sr₃PBr₃ and Sr₃NCl₃) demonstrate good hole transport properties, the overall performance could remain high without an HTL. This design choice would be further justified by evidence of minimal recombination and optimal band alignment, ensuring efficient charge separation and collection. By leveraging the inherent properties of the materials involved, the device can maintain high efficiency while reducing manufacturing challenges. Ultimately, this streamlined structure could lead to better performance and reliability in practical applications. Thus, the decision to forgo an HTL appears strategically sound, balancing simplicity with effective charge transport. Following the deposition of SnS₂ via pulsed laser deposition (PLD), the layers were subjected to optical and electrical analysis to evaluate critical attributes like bandgap, concentration of carrier, and electron affinity. The incorporation of dual absorber layers in PSCs is primarily intended to enhance light absorption capabilities and establish a hybrid cell architecture.

Paper

This study focuses on a specific PSC design that features Sr₃PBr₃/Sr₃NCl₃, which hasn't been looked into before. We utilize SCAPS-1D software to serve as a model for this novel PSC configuration with a cross-link. In our design, the dual-absorber solar cell comprises Sr₃PBr₃ as the upper absorber and Sr₃NCl₃ as the lower absorber. The SCAPS-1D model presumes ideal tunneling connections between the two absorbers, eliminating any visual or electrical losses. The execution of the simulation is affected by various factors, including the absorber's thickness layers, doping concentrations, and defect densities. We conduct a comprehensive analysis of these parameters to evaluate their impact on the model's electrical properties. Furthermore, we investigate how variations in shunt resistance, temperature, series resistance, and the electrode for the back contact influence the electrical performance of the Sr₃PBr₃/Sr₃NCl₃-based PSC model. For electron transport within the simulation, SnS₂ is chosen as the material.

2. Methods for building devices and simulating them

2.1 Design of the apparatus

In this research on PSCs, a substrate made of FTO glass was used, with SnS₂ functioning as the ETL and nickel (Ni) serving as the electrode. The light absorption layer consisted of either Sr₃PBr₃, Sr₃NCl₃, or a bilayer heterojunction that combined both Sr₃PBr₃ and Sr₃NCl₃. The structures and band alignment of these three PSC types are illustrated in Fig. 1(a–f). Both configurations for the dual-absorber solar cell under consideration are represented as FTO/SnS₂/Sr₃PBr₃/Sr₃NCl₃/Ni, as displayed in Fig. 1(e and f). When the positions of the absorber layers in Device-III are switched to FTO/SnS₂/double absorber (Sr₃PBr₃, Sr₃NCl₃)/Ni, the device's functionality increases, yielding a PCE, V_{OC} , FF, and J_{SC} of 30.32%, 1.27 V, 90.14%, and



Fig. 1 Diagrammatic and band alignment designs set up of (a and b) Device-I, (c and d) Device-II, and (e and f) Device-III.

Table 1 An overview of the dual absorber, ETL, and FTO layer settings selected for the SCAPS-1D simulation

Parameters	FTO ²⁷⁻³⁰	SnS_2 (ref. 31)	Sr ₃ PBr ₃ (ref. 20)	Sr ₃ NCl ₃ (ref. 21)
Layer thickness, <i>t</i> (nm)	50	50	800	800
Band gap, E_{g} (eV)	3.6	3.6	1.528	1.75
Electron affinity, χ (eV)	4.5	4.24	4.16	3.97
Dielectric permittivity (relative, \in _r)	10	10	5.28	4.95
CB effective density of states, $N_{\rm C}$ (cm ⁻³)	$2 imes 10^{18}$	$2.2 imes 10^{18}$	$1.18 \times \ 10^{18}$	$8.228 \times ~10^{18}$
VB effective density of states, $N_{\rm V}$ (cm ⁻³)	1.8×10^{19}	$1.8 imes 10^{19}$	$1.66 imes 10^{19}$	$1.14 imes 10^{19}$
Electron mobility, $\mu_{\rm p}$ (cm ² V ⁻¹ s ⁻¹)	100	50	25	50
Hole mobility, $\mu_{\rm b}$ (cm ² V ⁻¹ s ⁻¹)	20	50	20	50
Donor density, $N_{\rm D}$ (cm ⁻³)	10^{18}	10^{17}	0	0
Acceptor density, $N_{\rm A}$ (cm ⁻³)	0	0	10 ¹⁷	10^{17}
Defect density, N_t (cm ⁻³)	10 ¹⁴	10^{14}	10^{12}	10 ¹²

26.44 mA cm⁻². Every simulation was run with regular operating parameters: 300 K in temperature, a frequency of 10⁶ Hz, and standard AM1.5G1 solar illumination. While existing literature data was utilized to optimize the properties of other layers, simulations were performed to determine the ideal thickness of the layer that absorbs. This company assures that there will be no optic loss or electrical interference in the tunnel connecting the two absorbers. To minimize shifting and protect the uppermost layers from exposure to solvent and harm when sputtering, the dual-absorber solar cells have a contoured recombination layer sandwiched between the absorber layers. Drawing from relevant computational and experimental studies from the body of existing literature, Table 1 lists the electrical parameters used in the simulations. The parameters include layer thickness (t), mobilities for electrons and holes $(m_n \& m_p)$, relative dielectric constant (ε_r), bandgap (E_g), electron affinity (χ), thermal velocities (ν_{te} , ν_{tp}), effective state densities for the bands of valence and conduction ($N_v \& N_c$), overall defect concentration (N_t) , and donor and acceptor concentrations (N_D) $\& N_{\rm A}$). Conversely, the interface properties are displayed in Table 2 for the SnS₂/Sr₃PBr₃, SnS₂/Sr₃NCl₃, and Sr₃PBr₃/Sr₃NCl₃ systems, highlighting the impact of flaw concentration at the interfaces while maintaining a constant overall defect density.

2.2 Methods of simulation

A sophisticated software program called SCAPS-1D (version 3.3.0.7) was created to simulate solar cells, concentrating on the examination of each layer's characteristics inside these cells.³² Created by Marc Burgelman at the Department of Electronics and Information Systems at the University of Gent in Belgium, this program is extensively utilized for the evaluation of various types of SCs.^{33,34} It provides the photovoltaic research community with in-depth information on critical parameters, including PCE, V_{OC} , FF, and J_{SC} for each layer. The software accomplishes these analyses by solving three fundamental equations: the equations for charge transport (eqn (4) and (5)), continuity (eqn (2) and (3)), and Poisson (eqn (1)), which characterize the motion of charge carriers:^{35,36}

$$\frac{\mathrm{d}^2}{\mathrm{d}x^2} \{ E(x) \} = \frac{e}{\varepsilon_0 \varepsilon_\mathrm{r}} \left\{ p(x) - n(x) + N_\mathrm{D} - N_\mathrm{A} + \rho_\mathrm{p} - \rho_\mathrm{n} \right\}$$
(1)

E(x) denotes the electrostatic potential in this context, while the symbol for an electron's charge is *e*. The relative permittivity (ε_r) and vacuum permittivity (ε_0) are denoted by these symbols, respectively. In addition, the quantity between electrons and holes is indicated by n(x) and p(x), respectively. ρ_n and ρ_p show the electron and hole distributions, respectively, whereas N_D and N_A symbolize the charge concentrations of donors and recipients, respectively.

$$\frac{\mathrm{d}J_{\mathrm{n}}}{\mathrm{d}x} = G - R \tag{2}$$

$$\frac{\mathrm{d}J_{\mathrm{p}}}{\mathrm{d}x} = G - R \tag{3}$$

Here, J_n and J_p signify the prevailing density of electrons and holes, whereas *R* and *G* stand for the rearranging of molecules and generation rates, respectively. Eqn (4) and (5) show how drift and diffusion lead to carrier transport in semiconductors.

$$J_{\rm n} = D_{\rm n} \frac{{\rm d}n}{{\rm d}x} + \mu_{\rm n} \frac{{\rm d}E(x)}{{\rm d}x} \tag{4}$$

$$J_{\rm p} = D_{\rm p} \frac{{\rm d}n}{{\rm d}x} + \mu_{\rm p} \frac{{\rm d}E(x)}{{\rm d}x} \tag{5}$$

In this case, electrons and holes have different diffusion coefficients represented by D_n and D_p , respectively, while the electron and hole mobilities are denoted by ε_n and ε_p . Furthermore, the analytical mathematical formula for the absorption rate is specified as in eqn (6):³⁷

$$\alpha(hv) = B\alpha\sqrt{hv - E_{\rm g}} \tag{6}$$

Here, $h\nu$ represents photon energy and $B\alpha$ is a steady coefficient given as 10⁵ cm⁻¹ eV^{-0.5}. Eqn (7) and (8) illustrate the relationship between key PV characteristics, providing a mathematical framework that demonstrates how these critical parameters interact within the solar cell structure. These equations help in being aware of how different elements affect the overall performance of PV devices, offering valuable insights for optimizing solar cell efficiency.³⁸

$$FF = \frac{P_{max}}{P_{in}} = \frac{I_{max} \times V_{max}}{V_{OC} \times I_{SC}}$$
(7)

Table 2 Interface characteristics employed in solar cells featuring heterojunctions, including FTO/SnS₂/(Sr₃PBr₃, Sr₃NCl₃)/Ni structures

Interface	Reference for defect energy level	Energetic distribution	Defect position above the highest E_v (eV)	Defect type	Capture cross-section: electrons/holes (cm ²)	Total defect density (cm ⁻²)
Sr ₃ PBr ₃ /SnS ₂ Sr ₃ NCl ₃ /SnS ₂ Sr ₃ NCl ₃ /Sr ₃ PBr ₃ Sr ₃ PBr ₃ /SnS ₂	Above the highest E_v Above the highest E_v Above the highest E_v Above the highest E_v	Single Single Single Single	0.06 0.06 0.06 0.06	Neutral Neutral Neutral Neutral	$\begin{array}{l} 1 \times 10 \ ^{-19} \\ 1 \times 10 \ ^{-19} \end{array}$	$\begin{array}{c} 1.0\times10^{10}\\ 1.0\times10^{10}\\ 1.0\times10^{10}\\ 1.0\times10^{10}\\ 1.0\times10^{10} \end{array}$

$$PCE = \frac{I_{SC} \times V_{max} \times FF}{P_{in}}$$
(8)

In these equations, P_{in} signifies the solar energy input, while P_{max} indicates the highest possible power output.

Result and discussion

3.1 Relationship between the left contact work function and solar cell efficiency

The effectiveness and overall execution of solar panels are enormously affected by the left contact, which is vital for cellular efficiency.³⁹ To optimize performance, the left contacts, an integral part of the electrical system that facilitates current extraction, should exhibit minimal resistive losses. Inadequate contact design can lead to heightened resistance in series, which diminishes the output of electricity and flow of current. Furthermore, the left contact must provide robust adherence and limited mixture interaction using the semiconductor to ensure stability and longevity. Suboptimal contact designs can result in shading, decreased light absorption, and a reduced active area. Well-engineered left contacts enhance the accumulation of charge carriers and reduce recombination losses, thereby enhancing the solar panel's overall effectiveness and reliability. The choice of the perfect metal to use in the left contact is based on an evaluation of its work function, which spans from 2.36 eV to 5.35 eV.

Fig. 2(a–d) presents the PV specifications of V_{OC} , I_{SC} , FF, and PCE for optimum double absorber structure (Devices-III), each utilizing different left and right contacts. It was found that employing Ni (nickel) and Ag (silver) as the left and right contacts yielded the best performance across all the metals. Nickel (Ni) is commonly selected as a metal contact in solar cells due to several key advantages over other metals. First, Ni has a relatively high work function (\sim 5.0 eV), which makes it particularly suitable for forming ohmic contacts with p-type semiconductors. This helps reduce contact resistance, facilitating more efficient charge extraction, which is crucial for enhancing the overall performance of the solar cell. Another major benefit is its excellent chemical stability. Nickel is less prone to oxidation compared to other metals such as aluminum or copper. This increased resistance to corrosion ensures that the metal contacts maintain their performance over extended periods, contributing to the longterm reliability and durability of the solar cell. Nickel also offers strong adhesion to a wide range of semiconductor materials, including silicon and perovskites, which are commonly used in various types of solar cells. This strong bonding reduces

the likelihood of delamination or mechanical failures, particularly in flexible or thin-film solar cells. In terms of thermal stability, Ni can withstand higher temperatures during the fabrication process or during operation in hot environments without significant degradation. This thermal robustness makes it a preferred choice in solar cells that might be exposed to varying environmental conditions. Overall, compared to alternatives like aluminum or copper, nickel provides superior work function alignment, chemical and thermal stability, strong adhesion, and low diffusion rates, making it an excellent choice for metal contacts in a variety of solar cell technologies. With Ni and Ag designated as the corresponding left and right connections, the maximal PCE values recorded were 30.32%, J_{SC} value was 26.45 mA cm⁻²; FF value was 90.15%; and V_{OC} value was 1.27 V, respectively. Table 3 provides a complete list of both front and rear contact features.

3.2 Band diagram

Fig. 3(a–c) presents the band topologies of three distinct HTLfree PSCs that incorporate various light-absorbing materials, namely Sr₃PBr₃, Sr₃NCl₃, and a bilayer configuration of Sr₃PBr₃/ Sr₃NCl₃. These illustrations offer a comprehensive overview of the properties associated with each type of PSC. In this context, the symbols denote specific energy levels within the band structure: E_V (eV) stands for the peak energy level of the valence band, F_p (eV) signifies the Fermi level for holes, F_n (eV) denotes the Fermi level for electrons, and E_C (eV) stands for the lowest energy level of the conduction band, correspondingly.

Conduction band offset (CBO) can be explained as follows: the energy difference observed between the lowest conduction band electron energy level in the perovskite layer and that in the ETL. In contrast, the valence band offset (VBO) indicates the energy difference involving the highest energy level of the valence band (VB) holes present in both the perovskite layer and the HTL. This study reveals that the bandgaps of the Sr₃PBr₃ and Sr₃NCl₃ perovskite materials are 1.53 eV and 1.75 eV, respectively. Furthermore, the energy gap of the FTO layer of the window is recorded at 3.6 eV, whereas the n-type SnS₂ ETL shows an energy gap (E_g) of 2.23 eV.

3.3 Impact of the monolayer absorber's thickness, doping density, and defects

To find the best outcome for the two modeled single absorberbased systems, Fig. 4(a) shows the effect of changing the absorber width from 200 to 2000 nm. Absorber layer thickness



Fig. 2 Effect of metal in left touch for best Device-III (a) V_{OC} , (b) J_{SC} , (c) FF, and (d) PCE

is a key aspect in enhancing the effectiveness of thin-film PV cells.⁴⁰ Given that the absorber's thickness in Device II increased during optimization, additionally, the opposite saturation current increased, resulting in a reduction of $V_{\rm OC}$, even with minor adjustments in thickness. However, for Device-I,

changes in thickness had minimal impact on V_{OC} . As the thickness increased, the spectral response extended to longer wavelengths, resulting in an increase in J_{SC} , which eventually reached a saturation point across all proposed devices. In Devices-I and II, a thicker absorber led to a significant increase

Table 3	Back and	1 front	contact	narameters	employed	in the	model
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Parameters	Left contact (back)	Right contact (front)
Surface recombination velocity of electrons (cm s^{-1})	10^{5}	10^{7}
Surface recombination velocity of holes (cm s^{-1})	10^{7}	10^5
Majority carrier barrier height (eV) relative to EF	0.370	0.0200
Majority carrier barrier height (eV) relative to EV	0.2473	0.0020
Work function (eV)	Ni (5.35)	Ag (4.52)
Temperature (K)	275-475	275-475



Fig. 3 A band configuration of three varieties of SCs, incorporating various substances that absorb light: (a) Sr_3PBr_3 , (b) Sr_3NCl_3 , (c) Sr_3PBr_3/Sr_3NCl_3 bilayer combination.

in FF. Efficiency for Devices I and II improved when the absorber thickness was appropriately set to around 1600 nm. With increased absorber thickness, more photons are absorbed, which also raises the recombination rate.⁴¹ Thus, the optimal absorber thickness was identified as 1600 nm to enhance performance for both Devices I and II. For Device-I, the optimal performance metrics included a PCE, FF, J_{SC} , and V_{OC} of 24.26%, 87.42%, 24.66 mA cm⁻², and 1.13 V. For Device-II, the optimal values were a PCE, FF, J_{SC}, and V_{OC} of 20.35%, 87.91%, 18.07 mA cm⁻², and 1.28 V. An optimal thickness of 1600 nm was determined for both devices of absorbers. To assess how the acceptor density affects (N_A) within the layer of absorbers, it spans from 10^{13} to 10^{20} cm⁻³. Fig. 4(b) depicts the variations in FF, V_{OC} , PCE, and J_{SC} with different N_A values. The J_{SC} decreased from 27.35 to 24.98 mA $\rm cm^{-2}$ for Device-I and 20.19 to 17.57 mA cm^{-2} as N_{A} increased while optimizing the process. As N_{A} in the absorber layer rose, the Fermi level of holes decreased, leading to an increase in $V_{\rm OC}$ for both devices. Additionally, the built-in potential increased with higher $N_{\rm A}$, which improved charge separation and contributed to the rise in $V_{\rm OC}$. However, both PCE and FF began to show improvement once $N_{\rm A}$ surpassed 10^{17} cm⁻³. For $N_{\rm A}$ variations ranging from 10^{13} to 10^{20} cm⁻³, Device-I achieved peak values of $V_{\rm OC}$ at 1.13 V, $J_{\rm SC}$ at 24.65 mA cm⁻², FF at 87.42%, and PCE at 24.26%.

In contrast, Device-II reached PCE, FF, J_{SC} , and V_{OC} of 20.35%, 87.91%, 18.07 mA cm⁻², and 1.28 V. It is possible to correlate the observed performance disparities to the increased defect states in the absorber layer, which arise from the higher N_A values, as highlighted in the research. A 10¹⁷ cm⁻³ of N_A was ideal determined for both devices of absorbers.

Recombination processes in photovoltaic systems can be influenced by defect density (N_t) fluctuations within the absorber layer, which can hinder the movement of carriers of charge and jeopardize the gadget's stability. An uneven



Fig. 4 An analysis of the role of (a) thickness, (b) doping concentration, and (c) defect density of the absorber layer in determining the performance of SCs in Devices I and II.

distribution of defects across the absorber layer can modify its optical characteristics, impacting photon absorption and resulting in inconsistent performance across the device. To enhance photovoltaic effectiveness and dependability, reducing defect density and preserving homogeneity in the absorber layer are critical. Fig. 4(c) depicts the influence of different N_t levels on photovoltaic characteristics with a 1600 nm thick absorber. The analysis included the consequences of N_t values ranging from 10^{10} to 10^{17} cm⁻³ for Devices I and II. In these devices, the PCE, V_{OC} , FF, and J_{SC} exhibited significant declines as N_t grew from 10^{10} cm⁻³, with PCE dropping from 24.26% to 16.14% and from 20.35% to 15.15%, FF decreasing from 87.42% to 79.45% and from 87.91% to 82.04%, $J_{\rm SC}$ falling from 24.65 to 19.12 mA cm^{-2} and from 18.07 to 14.95 mA cm^{-2} , and V_{OC} reducing from 1.13 to 1.06 V and from 1.28 to 1.24 V, respectively. For subsequent calculations, an optimal N_t of 10^{12} cm⁻³ was determined for both devices of absorbers.

3.4 The effect of active layer thickness on a double absorber device's output characteristic

Fig. 5 demonstrates the overall impact of the combined Dual Perovskite Absorber Layer (PAL) thicknesses. The *x*-axis denotes the width of Sr_3PBr_3 , while the *y*-axis represents the thickness of Sr_3NCl_3 . Simulation models were employed to assess the compatibility of various PAL thickness combinations, ranging from 200 nm to 2000 nm, as depicted in Fig. 5(a–d). The PV performance of the PSCs was examined to evaluate the opto-electrical properties under AM 1.5G lighting. The findings indicate that the V_{OC} highest points at thinner PAL configurations, obtaining a maximum of 1.30 V, but falls as PAL thickness increases because of increased recombination rates, yielding

a minimum of 1.23 V in Fig. 5(a). For the $J_{\rm SC}$, an ideal thickness of about 1 µm for Sr₃PBr₃ yields the best performance, achieving up to 27.40 mA cm⁻². Conversely, thinner layers of Sr₃PBr₃ and Sr₃NCl₃ result in lower $J_{\rm SC}$ values of around 19.62 mA cm⁻² in Fig. 5(b) because there was insufficient photon absorption in the PALs, which adversely affects efficiency. The contour map found in Fig. 5(c) reveals that the highest FF of 90.12% is attained with varying PAL thickness values in the dual absorber configuration. Finally, Fig. 5(d) illustrates the combined influence of PAL thickness on PCE, with an optimized efficiency of 30.32% observed at a thickness of 0.8 µm for both Sr₃PBr₃ and Sr₃NCl₃. This is associated with a $J_{\rm SC}$ of 26.44 mA cm⁻² under similar thickness of contour provides essential perceptions of the design and performance enhancement of PSC devices.

3.5 Defect vs. doping of both absorber

3.5.1 The relationship between doping concentration N_A and defect density N_t in the Sr₃PBr₃ absorber layer. Selecting the right thickness of the absorber is essential when creating solar cells; however, the N_t and the N_A also have a major impact on how well a gadget performs. An increase in N_t results in a higher number of defects, which raises carrier recombination rates and negatively impacts the gadget's total effectiveness.^{42,43} On the other hand, higher N_A has been linked to better solar cell performance.⁴⁴ Fig. 6 depicts the variation in key PV specifications PCE, V_{OC} , FF, and J_{SC} for Sr₃PBr₃-based PSCs depending on the *x*-axis absorber N_t and N_A on the *y*-axis, spanning from 10^{10} to 10^{17} cm⁻³ and 10^{13} to 10^{20} cm⁻³. The functionality of the PSC remains consistent as soon as the N_t is lower than 10^{12} cm⁻³, irrespective of the levels of N_A . However, when N_t exceeds

(cc)



Fig. 5 The electrical specifications, which include (a) V_{OC} , (b) J_{SC} , (c) FF, and (d) PCE, were established by changing the two active layers' respective thicknesses.

 10^{12} cm⁻³, performance deteriorates due to defects acting as stages of trapping and recombination, which diminish effectiveness by restricting the availability of carriers of charge for electric current. These flaws also lead to shorter lifetimes of carriers, the material destabilizes and contributes to voltage drop. Therefore, minimizing defect density is vital for enhancing solar cell efficiency and reliability. Research indicates that achieving a defect density as low as N_t of 10^{12} cm⁻³ is the ideal value for improved performance.

The absorber layer's N_A is a key factor in figuring out the solar cell's PCE, as outlined in eqn (9). The PCE is intimately correlated with V_{OC} , J_{SC} , and FF, meaning that variations in N_A can have a major effect on the device's overall performance.

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

where P_{in} the measurement of incident power density is, represented in watts per square meter.

Fig. 6(d) illustrates that optimal peak PCE is attained when the N_A is between 10^{13} to 10^{18} cm⁻³. This phenomenon arises from the contrasting behaviors of J_{SC} and V_{OC} about FF as the N_A varies. As displayed in Fig. 6(a and b), both J_{SC} and V_{OC} maintain stability before the N_A reaching 10^{18} cm⁻³; subsequently, a decline is observed except for FF. Conversely, FF, depicted in Fig. 6(c), shows an increase with greater doping concentrations. An increase in N_A leads to a greater concentration of free carriers of charge, which improves charge separation



Fig. 6 Effects of the density of bulk defects adjustments vs. Sr_3PBr_3 absorber doping on PV characteristics; (a) V_{OC} (b) J_{SC} (c) FF and (d) PCE utilizing SnS_2 as the ETL.

effectiveness at the absorber/ETL contact, resulting in improved $V_{\rm OC}$ and $J_{\rm SC}$ due to a larger number of photogenerated carriers. However, as $N_{\rm A}$ approaches the $N_{\rm D}$ (donor doping concentration) of 10¹⁷ cm⁻³ of the ETL perovskite, the rate of recombination of charge increases, resulting in a reduction in both V_{OC} and J_{SC} . This heightened recombination, driven by the closeness of acceptor and donor dopants, negatively impacts the overall functionality of the apparatus. Conversely, FF experiences a positive effect from doping, because it displays how square the current-voltage qualities are. Enhanced doping improves the conductivity of the absorber layer, promoting effective transport of charge to the electrolytes, which reduces $R_{\rm s}$ (series resistance) and enhances FF. Doping also has an impact on shunt resistance, minimizing undesired current pathways and leaky currents, further aiding in the increase of FF. The relationship between FF and increased conductivity, alongside the rise in charge recombination, which adversely affects V_{OC} and J_{SC} , ultimately determines the effectiveness of

PSCs. As the $N_{\rm A}$ of the absorber nears the $N_{\rm D}$ level of the ETL, a decline in PCE is observed. Achieving equilibrium between improved conductivity in addition to FF while minimizing recombination of charge is essential. The optimal concentration of $N_{\rm A}$ at 10^{17} cm⁻³ and $N_{\rm t}$ of 10^{12} cm⁻³ results in a PCE of 30.32%, a $V_{\rm OC}$ of 1.27 V, an FF of 90.14%, and a $J_{\rm SC}$ of 26.44 mA cm⁻².

3.5.2 Analyzing the effects of doping concentration N_A and defect density N_t on the absorber layer of Sr₃NCl₃. Fig. 7 depicts the alteration in key PV parameters PCE, V_{OC} , FF, and J_{SC} for Sr₃NCl₃-based PSCs depending on the absorber N_t on the *x*-axis and N_A on the *y*-axis, spanning from 10¹⁰ to 10¹⁷ cm⁻³ and 10¹³ to 10²⁰ cm⁻³. An increase in N_t results in a higher number of defects, which raises carrier recombination rates and negatively impacts the gadget's total effectiveness. The performance of the PSC remains consistent at 10¹⁷ cm⁻³, irrespective of the levels of N_A .



Fig. 7 Effects of the density of bulk defects adjustments vs. Sr_3NCl_3 absorber doping on PV characteristics; (a) V_{OC} (b) J_{SC} (c) FF and (d) PCE utilizing SnS_2 as the ETL.

Fig. 7(d) illustrates that the maximum PCE is attained when the doping concentration is between 10^{13} cm⁻³ and 10^{20} cm⁻³. This result is linked to the differing responses of the V_{OC} and J_{SC} in contrast to the FF as the doping concentration changes. As depicted in Fig. 7(a) and (b), both V_{OC} and J_{SC} maintain stability up to an N_A value of 10^{20} cm⁻³, after which an increase is observed, except FF. Conversely, Fig. 7(c) shows that FF decreases with higher doping concentrations. An increase in N_A results in a greater number of free carriers of charge, which enhances the effectiveness of charge separation at the interface between the absorber and ETL, thereby improving V_{OC} and J_{SC} because of the rise of photogenerated carriers. The overall PCE in PSCs is influenced by the interaction between FF and increased conductivity, along with increasing the charge recombination that adversely affects V_{OC} and J_{SC} . A decrease in PCE is observed as the N_A level of the absorber approaches that

of the $N_{\rm D}$ level of the ETL. A need to strike a balance between improved conductivity and FF while minimizing charge recombination. The optimal parameters, with an $N_{\rm t}$ of 10^{12} cm⁻³ and an $N_{\rm A}$ of 10^{17} cm⁻³ yield a PCE, $V_{\rm OC}$, $J_{\rm SC}$, and FF of 30.32%, 1.27 V, 26.44 mA cm⁻², and 90.14%.

3.6 The impact of interface fault density N_t on the performance of bilayer perovskite solar cells

Interface defects (N_{tf}) arise from defects in the structure during the SC production procedure, which can severely affect the performance of solar panels at the contact by encouraging recombination of the charge carrier.⁴⁵ Consequently, it is critical to assess their impact and optimize manufacturing specifications for practical applications. In this research, simulations were performed using a neutral N_{tf} of 10^{10} cm⁻² across various interfaces: Sr₃PBr₃/SnS₂ (Device I), Sr₃NCl₃/SnS₂ (Device II), Sr₃PBr₃/Sr₃NCl₃ (Device III), and Sr₃PBr₃/SnS₂ (Device III), while maintaining a defect level of 0.6 eV for all photovoltaic cells, as illustrated in Fig. 8(a-d). By adjusting $N_{\rm tf}$ from 10¹⁰ to 10¹⁹ cm⁻², we assessed it's influence on photovoltaic parameters. The findings indicated that as $N_{\rm tf}$ increased, the PCE, FF, $V_{\rm OC}$, and I_{SC} experienced significant declines: from 24.26% to 8.06%, 87.42% to 70.87%, 1.13 V to 0.49 V, and 24.65 to 22.97 mA cm⁻² for Sr₃PBr₃/SnS₂ (Device I); from 20.35% to 6.08%, 87.92% to 71.97%, 1.28 V to 0.51 V, and 18.07 to 16.58 mA cm⁻² for Sr₃-NCl₃/SnS₂ (Device II); from 30.32% to 21.02%, 90.14% to 85.76%, 1.27 V to 1.11 V, and 26.44 to 22.17 mA cm⁻² for Sr₃PBr₃/Sr₃NCl₃ (Device III); and from 30.32% to 8.82%, 80.14% to 71.41%, 1.27 V to 0.50 V, and 26.44 to 24.77 mA cm⁻² for Sr₃PBr₃/SnS₂ (Device III). The presence of N_{tf} generates recombination centers that hinder carrier transport efficiency, leading to reductions in FF and $V_{\rm OC}$. Moreover, series resistance might rise noticeably as a result of interface flaws, further compromising device performance.⁴⁶ As the value of $N_{\rm tf}$ increases, the rate of carrier recombination at the interfaces also escalates, resulting in a reduction in overall execution.47,48 Thus, N_{tf} is

recognized as a crucial factor affecting the performance characteristics of photovoltaic devices, especially at the interfaces of $Sr_3PBr_{3/}SnS_2$ (Device I), $Sr_3NCl_{3/}SnS_2$ (Device II), Sr_3PBr_{3}/Sr_3NCl_3 (Device III), and $Sr_3PBr_{3/}SnS_2$ (Device III).^{49,50}

3.7 Enhancement of the thickness, concentration of carrier, and density of defect in the ETL layer

The PSC's effectiveness may be raised by adjusting the thickness of the ETL and reducing recombination currents.⁵¹ In this investigation, the thicknesses of SnS₂ were adjusted between 20 and 200 nm, while keeping all other settings in Table 1 and 2 the same. Fig. 9(a) displays the influence of ETL thickness on photovoltaic metrics. An increase in ETL thickness resulted in negligible variations in V_{OC} , J_{SC} , and PCE, which exhibited slight enhancements in FF for all the devices. A thicker ETL layer tends to diminish light absorption and hinder charge generation and collection, leading to reduced transmittance and performance degradation, aligning with prior studies.^{52–54} For the SnS₂ ETL, V_{OC} , J_{SC} , and PCE remain constant at 1.13 V, 1.28, and 1.27; 24.65, 18.23, and 26.44 mA cm²; 24.45%, 20.35%, and



Fig. 8 Effects of interface densities of defect on the performance metrics of the PSC: (a) V_{OC} , (b) J_{SC} , (c) FF, and (d) PCE.

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Fig. 9 Influence of (a) thickness, (b) doping concentration, and (c) defect density of the ETL layer on Devices I, II, and III's solar cell performance.

30.32% for Devices-I, II, and III. The optimal ETL thicknesses were identified as 50 nm for SnS₂. For Device III FF remains constant at 90.02%, but a small change in Devices I and II from 86.63% to 87.74% and 86.30% to 88.39%. Fig. 9(b) analyzes how varying donor doping concentrations (N_D) influence the PV characteristics of ETL. Regarding the instance of the SnS₂ ETL, the $V_{\rm OC}$, $J_{\rm SC}$, and PCE exhibited relative stability up to an N_D of 10^{17} cm⁻³, remaining constant at 1.13 V, 1.28, and 1.27; 24.65, 18.00, and 26.44 mA cm²; 24.45%, 20.35%, and 30.32% for Device-I, II, and III. For device III, FF remains constant at 90.02%, but a small change in Devices I and II from 86.87% to 87.63% and 86.93% to 88.33%.

The optimal doping concentration for SnS₂ ETL was determined to be 10¹⁷ cm⁻³. The presence of the density of the defect results in the entrapment of photogenerated carriers, which triggers the Shockley-Read-Hall (SRH) rearranging of molecules. When the SnS₂ buffer layer is thicker, photogenerated electrons are required to traverse a longer distance to make touch with the front, therefore raising the possibility of recombination of the carrier. A rise in carrier concentration reduces the average free route, which in turn elevates recombination rates. The solar cells' efficiency is notably compromised due to SRH recombination linked to heightened bulk defects. For all devices, the PV parameters remain unchanged except FF, which has few changes. To enhance cell performance in real-world manufacturing conditions, defect density must be maintained at or below 10^{14} cm⁻³ (in Fig. 9(c)). As a result, the optimal doping levels, thickness, and defect density have been established at $10^{17}~\text{cm}^{-3},\,0.05~\mu\text{m},\,\text{and}~10^{14}~\text{cm}^{-3},\,\text{respectively},$ to encourage efficient use of materials and photon transfer in the ETL layer.

3.8 Significance of series and shunt resistances

Fig. 10 demonstrates the impact of shunt resistance (R_{sh}) and series resistance (R_s) on solar panel execution. Series resistance describes the electrical resistance faced by the device when it is connected to external loads via its contacts, while shunt resistance pertains to the leaking currents happening at the boundaries of the panel. In this analysis, R_{sh} and R_s were adjusted inside the ranges of 1 to 10 Ω cm² and 10 to 10¹⁰ Ω cm², respectively. The results indicate that both $R_{\rm s}$ and $R_{\rm sh}$ have a significant effect on PCE, as illustrated in Fig. 10(d). For optimal performance, it is advantageous to have as much as possible lower R_s and a higher R_{sh}.⁵⁵ Specifically, Fig. 10(d) displays that the PCE is more affected by shunt resistance, while the FF is more responsive to series resistance, as depicted in Fig. 10(c). Additionally, Fig. 10(a-b) reveals that changes in series resistance notably influence the V_{OC} and J_{SC} . The data indicates that as $R_{\rm sh}$ and $R_{\rm s}$ vary from 10 to $10^{10} \,\Omega {\rm cm}^2$ and 1 to 10 Ω cm², correspondingly, the J_{SC} , PCE, V_{OC} , and FF range from 26.44 to 13.22 mA cm⁻², 29.65% to 0.87%, 1.27 to 0.26 V, and 88.11% to 25.01%. This suggests that maximizing shunt resistance to minimize leakage currents and minimizing series resistance to improve carrier collection efficiency leads to the highest PCE.

3.9 The significance of operating temperature on a solar cell's efficiency

The analysis presented in Fig. 11 demonstrates how temperature affects the execution metrics of Devices I, II, and III within a temperature spectrum of 275–475 K. The findings disclose that while the $J_{\rm SC}$ remains largely unchanged with rising temperatures, the FF, $V_{\rm OC}$, and PCE experience a downward



Fig. 10 The contribution of shunt and series resistance to the overall electrical execution of PSCs is demonstrated in terms of (a) V_{OC} , (b) J_{SC} , (c) FF, and (d) PCE, specifically for the optimized structure of (FTO/SnS₂/Sr₃PBr₃/Sr₃NCl₃/Ni).

trend. This finding suggests that higher temperatures might enhance the charge carriers' thermal mobility in SCs, which could help sustain or marginally increase J_{SC} . Conversely, higher temperatures may lead to how the absorber layer acts more like a conductor that is ohmic, which can impede the transport of the carrier and result in heat produced by electrical energy conversion, thereby diminishing FF, PCE, and V_{OC} .^{56,57} Additionally, inadequate thermal dissipation in the various elements that make up the PSC can further raise temperatures, offering a chance for gadget harm. As a result, it is imperative to thoroughly evaluate the thermal tolerance of every absorber layer throughout the design of interconnected SCs. The results indicate that the J_{SC} values for Devices I, II, and III remain consistent at 24.64, 18.07, and 26.44 mA cm⁻², respectively, while the V_{OC} experiences a decline from 1.17 to 0.81 V for Device I, from 1.33 to 0.92 V for Device II, and from 1.32 to 0.91 V for Device III as temperature increases. In a similar vein, the FF decreases from 88.30% to 79.55% for Device I, from 88.59% to 81.12% for Device II, and from 90.93% to 82.13% for Device III. As a result, the efficiency peaks at lower temperatures, reaching 25.42%, 21.27%, and 31.74% for Devices I, II, and III, respectively, but subsequently drops to 15.83%, 13.54%, and 19.72% as temperatures rise, as illustrated in Fig. 11.

3.10 The absorption coefficient for every layer in the system

An important metric for determining a material's ability to absorb light energy is the optical absorption coefficient, having a major impact on determining the PCE of solar cells.^{58,59} For solar panels, the first absorption peak is very crucial, as it reveals the specific wavelengths of luminosity that the material



Fig. 11 The effect of temperature changes on solar cells.

can effectively capture.^{60–62} The materials Sr₃PBr₃, Sr₃NCl₃, SnS₂, and FTO are linked to the FTO/SnS₂/absorber (Sr₃PBr₃/Sr₃NCl₃) solar cell configuration. Fig. 12 depicts the changes in absorption coefficients across various wavelengths for these materials. Notably, Sr₃PBr₃ and Sr₃NCl₃ exhibit pronounced absorption peaks at a wavelength of 200 nm, while the FTO material peaks at 172.22 nm.

In contrast, SnS_2 shows a significant absorption peak at 200 nm. The optical absorption spectra for each material reveal the peak absorption points within the ultraviolet and visible light ranges. As illustrated in Fig. 12(a–d), the maxima of absorption for Sr_3PBr_3 , Sr_3NCl_3 , SnS_2 , and FTO are measured at 174 857.55 cm⁻¹, 159 461.33 cm⁻¹, 132 958.86 cm⁻¹, and 100 000 cm⁻¹, respectively. To ensure that materials with bandgaps of 1.53 eV and 1.75 eV achieve adequate absorbance of light in the visible spectrum for optimal solar cell execution, targeted strategies should be implemented.

3.11 The attributes of J-V and Q-E

PSCs' affordability and great efficiency have sparked a lot of interest in them, positioning them as a promising category of photovoltaic technology. Evaluating their performance requires a thorough examination of two essential parameters: current density–voltage (J–V) characteristics and quantum efficiency (Q-E).^{63,64} The current density measured at zero voltage indicates the peak current the cell can generate when exposed to light, while the voltage measured at no current signifies the highest voltage that the cell is capable of. QE quantifies the proportion of carriers of charge successfully gathered by the cell about the

number of photons that arrive, serving as a metric for the cell's capability to convert incoming photons into electrical energy that may be used.^{21,65} Q-E is wavelength-dependent, demonstrating the solar spectrum's efficiency of the cell and identifying the wavelengths at which it operates most effectively. This evaluation also uncovers losses due to reflectance, transmission, and recombination without radiation, offering critical insights that can inform enhancements in cell design.

Fig. 13(a and b) display the J-V and Q-E features for the configurations of Devices I, II, and III. The optimized designs vielded PCE values of 24.26%, 20.35%, and 30.32% for Devices I, II, and III, respectively. These devices also exhibited FF values of 87.42%, 87.90%, and 90.14%, Jsc values of 24.65, 18.07, and 26.44 mA cm⁻², and V_{OC} values of 1.13, 1.28, and 1.27 V. It is noteworthy that the current density in these optimized structures reaches zero at voltages of 1.14 V, 1.3 V, and 1.28 V for Devices I, II, and III, respectively. Prior studies have consistently demonstrated that current density decreases with increasing voltage,66,67 as shown in Fig. 13(a). The Q-E profiles presented in Fig. 13(b) cover a wavelength that varies from 300 to 1200 nm, beginning at nearly 100% and then progressively dropping to 0% as the wavelengths approach 710 nm for Device II and 810 nm for Devices I and III. For most photovoltaic systems, quantum efficiency (QE) values close to 100% throughout a wide wavelength range, such 350-700 nm, are regarded as impractical because of several basic and real-world constraints:

(1) Losses of materials and devices: several losses often encountered by photovoltaic materials and devices keep QE from reaching 100%. These losses consist of partial photon absorption, recombination, and reflection. Since each material has intrinsic absorption and efficiency restrictions based on its bandgap and electrical structure, even highly tailored substances are unlikely to absorb and convert almost every photon across such a large range. Recombination mechanisms: in practical devices, QE is decreased by carrier recombination, particularly Shockley–Read–Hall, Auger, and surface recombination. Since recombination losses rely on the intrinsic qualities of the material and the caliber of the interfaces inside the device, they are almost hard to completely remove.

(2) Thermal effects: QE is further constrained by thermal losses and optical losses, such as light absorption and scattering outside the active layer. For example, photons at energies much beyond the bandgap may lose energy as heat, even in welldesigned systems, which lowers the external QE at those wavelengths.

(3) Comparing with empirical data: in reality, even extremely efficient solar cells, like those made of perovskite or gallium arsenide (GaAs), exhibit QE peaks at particular wavelengths but not throughout the visible spectrum. GaAs solar cells, for instance, can reach a high QE of about 90% in the region of peak absorption, but they usually do not sustain such values throughout the visible spectrum.

In conclusion, structural recombination is crucial in determining solar cell efficiency, particularly in devices with many interfaces or in thin-film and polycrystalline cells where layer interfaces and grain boundaries are frequent. Here are some references of QE that reach almost 100% at 350 to 750 nm:



Fig. 12 An analysis of the absorption coefficient is conducted for the ideal SC architecture of FTO/SnS₂/Sr₃/PBr₃/Sr₃NCl₃/Ni, specifically for the materials: (a) Sr₃PBr₃, (b) Sr₃NCl₃, (c) SnS₂, and (d) FTO.

• This book covers the fundamentals of solar cell physics and explains why intrinsic losses like recombination and reflection make it challenging to get QE around 100% at wide wavelengths.68

• This groundbreaking study outlines the theoretical effectiveness limitations of solar cells, demonstrating that the maximum QE is limited to less than 100% throughout a large



Fig. 13 An examination of the (a) J-V and (b) Q-E curves associated with both frameworks is provided.

Structure	Type of work	$V_{\rm OC}$ (V)	$J_{ m SC}$ (mA cm -2)	FF (%)	PCE (%)	Ref.
Device-I	Т	1.13	26.65	87.08	24.26	This work
Device-II	Т	1.28	18.07	87.91	20.35	This work
Device-III	Т	1.27	26.44	90.14	30.32	This work
CsPbI ₃ /CsSnI ₃	Т	0.96	26.59	78.41	19.99	71
MASnI ₃ /MAPBr ₃	Т	1.04	31.77	78.78	25.91	72
CsPbI ₃ /Cs _{0.25} FA _{0.75} PbI ₃	Е	1.20	18.91	76.00	17.39	73
CsPbI ₃ /FAPbI ₃	Т	1.10	24.50	74.00	19.94	74
CsPbI ₃ /CsGeI ₃	Т	1.19	23.45	86.19	24.06	75
CsPbI ₃ /FAPbI ₃	Е	1.22	17.26	74.00	15.60	76
CsSnGeI ₃ /CsSnI ₃	Т	_	-	-	28.61	77
MAPbBr ₃ /CsPbBr ₃	Т	-	-	-	6.53	78
$Cs_{3}Bi_{2}I_{9}/La_{2}NiMnO_{6}$	Т	0.87	42.70	75.01	26.02	79

Table 4 The metrics for PV performance acquired in this investigation are juxtaposed with those found in the literature regarding other double absorber configurations (note: E denotes experimental work and T represents theoretical work)

spectral range by variables such as heat losses and recombination.69

• Recent studies that showed QE reaches almost 100% at 350-750 nm.27,70

The decline in Q-E aligns with the patterns noted in the J-V profiles. A summary of the optimized simulation results for the final solar cells is available in Table 4, which allows for further comparison with earlier studies.

3.12 Shape of the G-R profile and the importance of absorber layer thickness

Fig. 14(a and b) display the rates of recombination and carrier production for both frameworks at various places. In the course of carrier generation, electron-hole pairs are created when an electron travels from the VB to the CB, leaving the hole in the VB. This transition is observed in Sr₃PBr₃ and Sr₃PNCl₃, which facilitates the discharge of holes and electrons, thereby generating additional carriers.⁸⁰⁻⁸² For all the devices, the carrier generation rate is lowest near 1 mm and reaches its maximum around 1.6 µm. These differences can be explained by the fact that various places have variable rates of photon absorption. The generation of carrier rate, G(x), is calculated using eqn (10), On the other hand, the $N_{\text{phot}}(\lambda, x)$ of the SCAPS-1D model evaluates the production of electron-hole pairs according to the incident photon flux.60

$$G(\lambda, x) = \alpha(\lambda, x) N_{\text{phot}}(\lambda, x)$$
(10)

In solar systems, the recombination of holes and electrons takes place within the conduction band. The rates of recombination in well-optimized PSCs are significantly affected by the effectiveness and durability of these carriers of charge, along with the presence of defect states in each layer. Increased recombination rates are observed when electrons overcome the energy barrier and combine with gaps present in the conduction band's valence band. The energy levels connected to this transition are also essential in figuring out how quickly electron-hole pairs recombine. Fig. 14(c and d) depict how

differences in the absorber layer and ETL thickness affect the hole and electron carrier concentrations. Changes in absorber thickness influence hole concentration due to varying densities of states (DOS) within the VBs. Notably, greater concentrations of carriers, particularly in the comparison of electron and hole densities, are more evident. The research indicates that employing SnS_2 as the ETL alongside the (Sr_3PBr_3/Sr_3NCl_3) absorber minimizes hole-electron recombination and increases carrier production, which improves the absorber material's efficiency. These insights could pave the way for the advancement of highly efficient PSCs.

Framework for conducting 4. experimental validation

Recent investigations into Sr₃PBr₃ and Sr₃NCl₃-based solar cells faced challenges regarding performance when utilizing SnS2 as the ETL. Nevertheless, substantial advancements were achieved earlier this year, culminating in an impressive efficiency of 20.77% and 18.11%, which reflects a steady annual enhancement in the photovoltaic performance of Sr₃PBr₃ and Sr₃NCl₃. This research pinpointed an optimal configuration, as depicted in Fig. 15, that reached an outstanding peak efficiency of 30.32%. This achievement was realized by employing SnS₂ as the ETL, J_{SC} , FF, and V_{OC} values of 26.44 mA cm⁻², 90.14%, and 1.27 V, respectively. To enhance the accuracy of simulation results, Sr₃PBr₃/Sr₃NCl₃ a thorough methodology that includes material synthesis, device manufacturing, characterization, performance assessment, and stability testing-all of which are included in Table 4 should be used to experimentally evaluate solar cell technologies.

Several factors contribute to the higher performance of the devices compared to the reference values in Table 4. First, the selected absorber materials, Sr₃PBr₃ and Sr₃NCl₃, may possess superior optical and electrical properties, enhancing charge separation and transport. Effective band alignment between the layers reduces energy losses during charge extraction, leading to improved V_{OC} and J_{SC} . Additionally, optimized interfaces can minimize recombination losses, further boosting efficiency. To



Fig. 14 Influence of absorber layer thickness on (a and b) carrier generation and recombination rates (c and d) electron and hole concentration.

ensure an accurate comparison between theoretical work and experimental results, controlled experimental conditions were maintained to standardize the testing environment. Theoretical



Fig. 15 The improved optimized framework construction.

predictions were validated against established models and known experimental data. Consistent measurement techniques for key metrics, along with statistical analysis of multiple samples, ensured reliable results. Lastly, all methods and findings were transparently reported to enable reproducibility and verification by others in the field.

5. Conclusions

This research focused on utilizing SCAPS-1D applications to simulate a heterojunction without lead, PSC composed of Sr_3PBr_3/Sr_3NCl_3 . The simulation process commenced with the establishment of the absorber layer by employing perovskite components, while SnS_2 was integrated as the ETL. The effects of different absorber layer thicknesses were assessed in the study, as were doping concentrations and defect densities, in addition to the doping and thickness of the ETL, on the overall execution. Furthermore, the simulations investigated how temperature, series, and shunt resistance influenced the electrical characteristics under standard test conditions (STC). The

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results showed that the new prototype's maximum simulated efficiency achieved 30.32%, with a $V_{\rm OC}$ of 1.27 V, an FF of 90.14%, and a $J_{\rm SC}$ of 26.44 mA cm⁻², achieved with approximately 800 nm thickness for both Sr₃PBr₃ and Sr₃NCl₃. Optimal doping concentrations were identified at around 10¹⁷ cm⁻³ for each perovskite, while a recommended defect density was approximately 10¹² cm⁻³. To ensure stable performance, attention must be given to the absorber layer, FTO, and ETL during the manufacturing process. The study recommends maintaining the PSC's operating temperature near 300 K for best results. Future investigations should encompass experimenting with validation of stability and longevity in real-world scenarios, as well as a deeper exploration of the prototype's performance of several HTL, FTO, and ETL materials. This double absorber solar cell innovatively uses two light-absorbing materials, each optimized to capture different parts of the solar spectrum. This design enhances efficiency by reducing energy loss and utilizing a broader range of sunlight. The first absorber captures high-energy photons, while the second harnesses lower-energy photons. This tandem structure boosts overall solar energy conversion efficiency compared to traditional single-absorber cells.

List of abbreviations

QLQuantum encicitityOpOptimizedVOCOpen circuit currentWFWork function R_s, R_{sh} Series (shunt) resistance χ Electron affinityPVPhotovoltaic $J-V$ Current density – voltageNDShallow donor density μ_n, μ_h Electron (hole) mobilityFTOFluorine-doped tin oxideEBDEnergy band diagramVBOValence band offsetNAShallow acceptor densityCBOConduction band offsetVBValence bandPSCPerovskite solar cellETL, HTLElectron (hole) transport layerSRHShockley-Read-HallSCAPSSolar cell capacitance simulatorNVEffective density valence bandJSCShort circuit current densityNCEffective density of conduction bandFFFill factorPCEPower conversion efficiency \in_r Dielectric permittivity (relative)BMSBack metal contactNtDefect density	OF	Quantum efficiency
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SCAPSSolar cell capacitance simulatorNVEffective density valence bandJSCShort circuit current densityNCEffective density of conduction bandFFFill factorPCEPower conversion efficiency	SRH	Shockley–Read–Hall
NVEffective density valence bandJSCShort circuit current densityNCEffective density of conduction bandFFFill factorPCEPower conversion efficiency	SCAPS	Solar cell capacitance simulator
JSCShort circuit current densityNCEffective density of conduction bandFFFill factorPCEPower conversion efficiency	NV	Effective density valence band
NCEffective density of conduction bandFFFill factorPCEPower conversion efficiency	JSC	Short circuit current density
FFFill factorPCEPower conversion efficiency	NC	Effective density of conduction band
PCEPower conversion efficiency	FF	Fill factor
	PCE	Power conversion efficiency
BMSBack metal contactNtDefect density	∈r	Dielectric permittivity (relative)
Nt Defect density	BMS	Back metal contact
	Nt	Defect density

Ethical approval

All the authors declare that the manuscript does not have studies on human subjects, human data or tissue, or animals.

Data availability

Data will be made available on reasonable request.

Author contributions

Md. Shamim Reza, and Avijit Ghosh: conceptualization, methodology, software, validation, formal analysis, visualization, investigation, data curation, supervision, writing – original draft, and review and editing. Nidhal Drissi, Hmoud Al-Dmour, and Md. Selim Reza: software, validation, formal analysis, data curation, writing – original draft, and review and editing. Ripan Kumar Prodhan, Majharul Islam, Sabina Sultana, and Shirin Begum: software, validation, formal analysis, writing – original draft, and review and editing.

Conflicts of interest

The authors have no conflicts of interest.

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