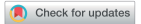
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Oxysalt based synergistic dual interfacial engineering for high performance p-i-n structured perovskite solar cells†

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Inverted perovskite solar cells (IPeSCs) have attracted extensive attention due to low-temperature processability and prospects for scale-up production. However, the power conversion efficiency (PCE) of IPeSCs lags compared to normal architecture, which is mainly due to inadequate interfaces of charge transport layers, viz electron and hole transport layers (ETL and HTL) and perovskite that suffers from carrier recombination paving the way for bulk recombination impeding the PCE and long-term stability of small devices. It is crucial to introduce an effective technique to reduce the interfacial non-radiative recombination losses at both interfaces of the perovskite layer. In this work, we demonstrated a multifunctional strategy to passivate the interface defects via the interactions and synergistic effect of anions and cations produced from sodium nitrate (NaNO₃), a π-conjugated oxysalt, at HTL/perovskite and ETL/perovskite interfaces. The synergistic effect of cation and anion bonding and interaction was confirmed by various experimental studies that decreased defect densities and improved the transfer of charges. The synergistic effect with double-side passivated devices showed a champion PCE of 21.53% as compared to pristine (19.08%), bottom (20.16%), and top (20.83%) passivated devices. The π conjugated oxysalt also improved the IPeSC stability by building a moisture-resistant interface. This work signifies the effect of chemical bonding engineering via cations and anions at dual-perovskite interfaces for synergistically enhanced PCE and long-term stability of IPeSCs.

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1. Introduction

Perovskite solar cells (PeSCs), which are the youngest generation of photovoltaic devices, have experienced an unparalleled surge in recent years due to the cost-effective and easy processing techniques, their tunable band gap, and the fact that they are made using readily available organic/inorganic materials that are compatible with industrial processing; as a result, they have arisen as the most encouraging industrial photovoltaic technology.¹⁻³

The inverted structure of PeSCs, *i.e.*, positive–intrinsic-negative (p–i–n), has attracted substantial attention due to its potential utility in flexible and tandem devices due to its low-temperature processability.⁴ Despite the immense attention that has been paid to the inverted solar cell configuration (IPeSC), the power conversion efficiency (PCE) values of IPeSCs still lag compared to those of the standard architecture, *i.e.*, the

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n-i-p structure.5 The primary reasons for the decreased performance of IPeSCs are associated with diminished charge extraction and increased non-radiative recombination losses. These issues arise from factors such as mismatched energy band alignment, the presence of interfacial defects, and the presence of trap sites, all of which collectively limit the device's fill factor and photovoltage.6 To tackle these challenges, extensive research attempts are currently in progress to devise strategies for mitigating defects, including point defects, surface defects, and bulk trap defects.7,8 The charge transport layers (CTLs) and the perovskite interface are the most crucial interfaces, as the growth of perovskite films and charge transportation depend on these interfaces. In p-i-n-structured PeSCs, NiO is most commonly used as a hole transport layer (HTL) owing to its wide band gap and deep valence band maximum (VBM) values, while [6,6]-phenyl C₆₁-butyric acid methyl ester (PCBM) has been identified as an electron transport layer (ETL) due to its outstanding charge transport abilities as well as low-temperature solution processability.9 However, the chemical species (such as water molecules and hydroxyl groups) and surface charge defects at the NiO surface can result in charge recombination, thereby reducing the open circuit voltage (V_{oc}) and fill factor (FF), while the low solubility and viscosity of PCBM can lead to a low-quality non-uniform film on

perovskite.9,10 These issues result in charge accumulation, recombination, and inefficient charge extraction, thus leaving perovskite vulnerable to degradation, which in turn limits device performance and serves as an obstacle to obtaining high PCE.

To date, several approaches have been used to modulate the interfaces of CTLs with the perovskite, such as solvent engineering, perovskite composition engineering, additive engineering, anti-solvent additive engineering, interface engineering, etc. 11-15 Among the available options, interface engineering is an efficient strategy to enhance perovskite film quality via modulating the growth and nucleation process. However, it could be highly desirable to identify a molecule that is also suitable for passivating the defects within the perovskite layer. The use of ionic compounds (cations and anions) has emerged as a novel strategy to passivate the defects and improve the interface quality. For instance, Bang et al. reported sodium bis(trifluoromethanesulfonyl)imide (Na-TFSI), a defect-tolerant dopant that improved both the performance and stability of PeSCs. 16 On a similar note, potassium nitrilotriacetate (NTAK) was used in SnO₂/perovskite interfaces, where the amino and carboxyl groups interacted with SnO2 and undercoordinated Pb²⁺ in the perovskite layer and simultaneously improved crystallization and passivated defects, accordingly, improving device performance and stability.17 Zhang et al. evaporated potassium chloride salt (KCl) at both the HTL and ETL interfaces and achieved a champion PCE of 19.2%, eventually resulting in reduced recombination at the defect sites.18 Lowcost alkali salts have demonstrated immense possibility due to the readily available cations and anions passivating both holes and electron trapping defects in the perovskite layer. 19,20 For instance, Park et al. evaluated alkali metal iodide series and concluded that K⁺ ions leads to hysteresis-free PeSCs.²¹ Moreover, Zang et al. used potassium sulphate (K2SO4), in regular PeSCs as a multifunctional layer to improve the SnO₂/perovskite interface. The SO₄²⁻ coordinated with Pb²⁺ at the bottom interface, while the K⁺ ion diffused into the perovskite layer, subsequently passivating the interfacial defects and modulating the energy level alignment. Hence, a synergistic effect of K⁺ and SO₄²⁻ was achieved in realizing high-performance PeSCs.²² Similarly, Li *et al.* utilized a π -conjugated oxysalt, *i.e.*, potassium carbonate (K2CO3), for the interfacial molecule binder and defect free layer at the SnO₂/perovskite interface in n-i-p PeSCs. The π -conjugated oxysalt interlayer offered a robust interface that enhanced the device stability and achieved impressive PCE.23

Although there have been various reports demonstrating modification at HTL/perovskite or ETL/perovskite interfaces using a suitable strategy, there is yet to be a strategy demonstrating interfacial engineering that utilizes both cations and anions from the same salt used at both HTL and ETL interfaces with perovskite in IPeSCs to improve the device performance. It is therefore urgently necessary to investigate the most suitable and multifunctional ionic compound to build a strong interface having reduced defect density, diminished charge recombination, and a synergistic effect for high performance IPeSCs.

Herein, we believe that we have demonstrated for the first time a cost-effective, simple yet effective interfacial modifier sodium nitrate (NaNO₃, ionic compound), a π -conjugated oxysalt, so that the collaborative effect of cation and anion interaction from both ends could contribute to the formation of robust interface layers. The introduction of NaNO3 at both charge transfer interfaces promoted the growth of superior perovskite films at the HTL interface along with improved charge transport properties at the ETL interface. The detailed experimental studies of both the bottom and top-side passivated devices along with double-side passivated devices revealed improved charge transfer properties with reduced defect density and recombination at the interfaces. The optimized double-side passivated device showed a champion PCE of 21.53% with enhanced V_{oc} , FF and short-circuit current density (I_{sc}) compared to the pristine, bottom, and top-side passivated devices. Moreover, the stability of the devices exhibited remarkable long-term light and nitrogen stability for unencapsulated devices.

Results and discussion

In designing the scientific approach for the two-fold optimization of the triple cation-based perovskite layer's interfaces, we use sodium nitrate ionic solution as the dual modifier for both HTL and ETL interfaces with the perovskite. Fig. 1a shows the schematics of the device architecture with the sodium nitrate surface passivator on dual sides of the photoactive layer. The HTL side (bottom-side passivated devices) and ETL side (topside passivated devices) are designated as BSP and TSP, respectively. The dual-side passivated devices are designated as DSP. Materials and experimental procedures are described in depth in the ESI documentation.†

To ascertain the distribution of NaNO₃ in the entire device, we performed time of flight secondary ion mass spectrometry (ToF-SIMS). The depth profiling images of the main elements were obtained and are shown in Fig. 1b and c. The depth profile images for DSP reveal that the sodium ions are mainly located at the interfaces of the HTL and ETL. Fig. 1c clearly shows the distribution of diffusion of Na ions at the interfaces and in the bulk for the DSP samples. The distribution along the interface reveals that the interaction mostly occurs at the interfaces leading to interfacial bonding, while the diffusion of Na ions in the bulk could also help passivate the bulk defects. 18,24

Furthermore, XRD analysis was conducted to examine the effect of alkali salt solution treatment on the structural characteristics of perovskite films. The perovskite films in pristine and DSP are shown in Fig. 1d. Both samples exhibited a nearly identical orthorhombic perovskite crystal structure, without apparent shift of the perovskite phase, implying that neither Na⁺ nor NO₃⁻ has been incorporated in the perovskite crystal lattice.11,25 The small PbI2 signals in pristine samples deposited on the NiO surface disappeared in the NiO/NaNO₃-based perovskite films, indicating that introducing NaNO3 can effectively restrict the interfacial reactions.26 There was an observable increase in the intensity of the perovskite films deposited on the NiO/NaNO3 surface, implying that the crystallinity of the

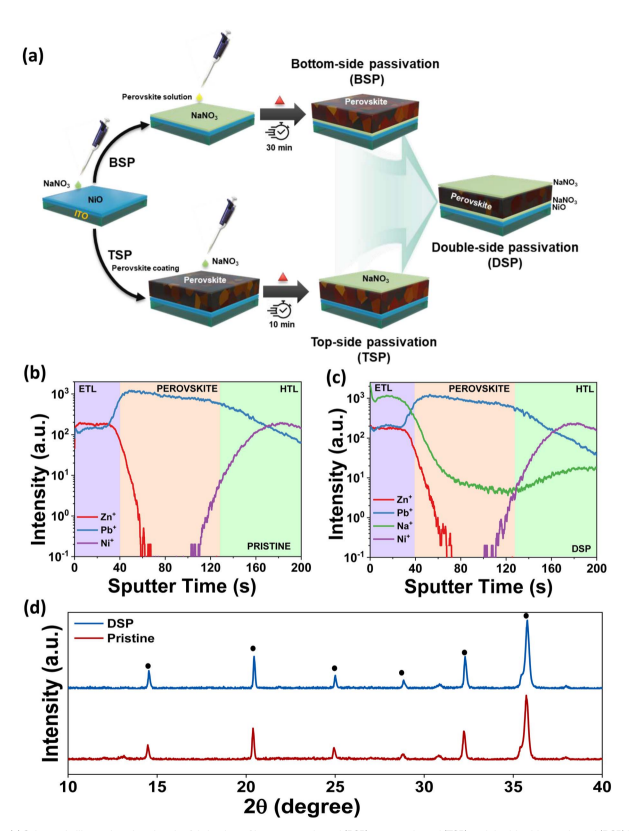


Fig. 1 (a) Schematic illustration showing the fabrication of bottom passivated (BSP), top passivated (TSP) and double side passivated (DSP) layers. TOF-SIMS depth profiles (from surface to bottom) of perovskite films: (b) pristine and (c) DSP films. (d) XRD patterns for perovskite films for pristine and DSP perovskite layers.

perovskite films was improved. This further indicates that the crystal orientation was controlled in an efficient way to decrease the distortions in lattice and micro-strains in crystals and ultimately reduce the formation of traps and defects.^{27,28} To analyze the effect of NaNO₃, XRD analysis of the sample without heating of the perovskite films was also carried out. We observed that even without heating, the XRD peak intensity in the case of DSP was significantly improved in comparison to pristine, as shown in Fig. S1.† The FWHM also decreased significantly, indicating an increase in the crystallite size. The FWHM for the perovskite layer with and without heat is shown in Table S1.† As previous studies have speculated,29,30 the dissolved salt can retard the heterogeneous nucleation process occurring at the HTL and perovskite interface, thus leading to an enlarged perovskite crystallite size. Therefore, the crystallite size was found to increase even without heating the perovskite layer.29,31 Ultimately, the dual passivation strategy enhanced crystallinity and increased grain size, which is anticipated to lower defect density and inhibit recombination.

2.1 HTL interface engineering

The HTL interface interactions were studied in detail to elucidate the effect of NaNO3 as an HTL interfacial layer. In this process, X-ray photoelectron spectroscopy (XPS) was carried out to investigate various chemical properties and more deeply understand the interaction of NaNO₃ with the NiO lattice. The XPS spectra are illustrated in Fig. 2a and b. In the figure it can be seen that the XPS spectra shifted to lower binding energy after treatment of NaNO₃ solution on the NiO surface, indicating the possibility of a chemical interaction between NiO and NaNO₃, as can be noticed in Fig. S2a and b.† The Ni 2p_{3/2} characteristic peaks at 854.06 and 855.78 eV shift to the lower binding energy values of 853.81 and 855.38 eV, respectively, after NaNO3 solution treatment. As reported previously, alkali metal ions tend to occupy the interstitial and oxygen vacancy (Ov) sites in metal oxide semiconductors. Huang et al. reported on the formation of interfacial dipoles at ITO/alkali interfaces.33 Similarly, Tai et al. reported on the formation of a Zn-O-Na/K bond by the replacement of H from the -OH group. These permanent dipole moments could alter the distribution of the electrical charges on the surface.32-34 Analogously, we speculate from previous reports that Na⁺ ions could interact with the hydroxyl groups at the NiO surface and reduce O_v. With detailed analysis of the Ni $2p_{3/2}$ and O 1s spectra, we further confirmed the possible interaction. The fitted Ni 2p_{3/2} XPS spectra showed three peaks corresponding to NiO at 853.8, Ni₂O₃ at 855.4 eV, and NiOOH at 856.5 eV. As observed in Fig. 2a, after NaNO₃ treatment, the relative composition of NiO and NiOOH decreased, whereas the amount of Ni₂O₃ increased. Similarly, in the O 1s spectrum, the XPS peak at 531 eV may be attributable to the metal deficiency of Ni³⁺ or excess oxygen of NiOOH. More explicitly, the ratio of Ni₂O₃/NiO in the case of the NaNO₃-treated layer was calculated to be 0.59, compared to the value of 0.49 calculated for the NiO layer. Meanwhile, the ratio of NiOOH was reduced from 0.43 to 0.39 for the NiO and NiO/NaNO₃ layers, respectively, suggesting that the sodium ion could help to minimize the high valence

nickel ions at the NiO surface layer. These observations are consistent with the O 1s spectra, and they further confirm that the Na ions could react with adsorbed hydroxyl groups at the NiO surface4 (Table S2†). These results show that NaNO3 treatment could lead to a reduction in defect sites and hydroxyl groups on top of the HTL surface, ultimately leading to improved charge transfer properties.35 The main reason for the reduction in hydroxyl groups could be the linking of Na with the -OOH group of NiOOH, thus leading to the formation of a Ni-O-Na bond and simultaneously reducing the defect sites and hydroxy groups.32 Furthermore, the Na 1s core-level spectra indicated the presence of a Na peak in the binding energy range of 1072–1073 eV, thereby confirming the presence of Na⁺ in the modified samples.36 Moreover, nitrate peaks were confirmed on the surface of NiO (Fig. S2c and d†).

Furthermore, FTIR studies were conducted to understand the mechanism in detail. Fig. S3† shows the FTIR spectra for the NiO, NaNO₃, and NiO/NaNO₃ films. The NiO film showed a characteristic peak at \sim 500 cm⁻¹. The NaNO₃ film showed the characteristic peak for antisymmetric stretching vibration at \sim 1383 and \sim 836 cm⁻¹.³⁷ The deposition of NaNO₃ on the NiO surface showed the characteristic peaks of both NiO and NaNO₃, indicating that NaNO₃ was well anchored on the NiO surface.38 To confirm this, we measured the water contact angles for the NiO and NaNO3 films. The water contact angle was observed to have significantly improved, increasing from 26.44° to 53.17° for control and modified films, respectively (Fig. S4†). This increase in the contact angle also confirms the reduction of surface hydroxyl groups on the NiO film.10 This further supports the XPS and FTIR results indicating that NaNO₃ can efficiently reduce the surface hydroxyl groups on the top surface of NiO and ultimately lead to the formation of a better perovskite film.38

Next, Atomic Force Microscopy (AFM) and Scanning Kelvin Probe Microscopy (SKPM) studies were conducted to empathize the impact of NaNO3 treatment further clearly on the surface properties. Some previous reports suggest that bottom passivation could amend the growth of perovskite and influence defect distribution on the perovskite surface and bulk.18 Fig. 2c shows topography images of ITO/NiO and ITO/NiO/NaNO₃. The surface roughness was reduced from 0.89 nm to 0.77 nm for the NiO/NaNO₃-modified samples. Moreover, the surface potential measured from SKPM analysis (Fig. 2d) for ITO/NiO/NaNO3 showed a low value of 27.16 mV, in comparison to the value of 34.54 mV for ITO/NiO. The lower surface potential value suggests that the utilization of NaNO₃ can help to improve the work function of NiO. These results support that the holes from the photoactive layer can be more effectively extracted and transported towards the anode.36,39,40

The uniformity of the substrate (NiO) typically represents the most important parameter influencing the film quality and morphology of spin-coated perovskite films. The energydispersive X-ray spectroscopy (EDS) mapping images demonstrated the distribution of NaNO3 on the NiO surface. The elements were identified to be evenly distributed on the HTL surface, as represented in Fig. S5.† The surface top-view FESEM analysis of BSP and pristine films was conducted to

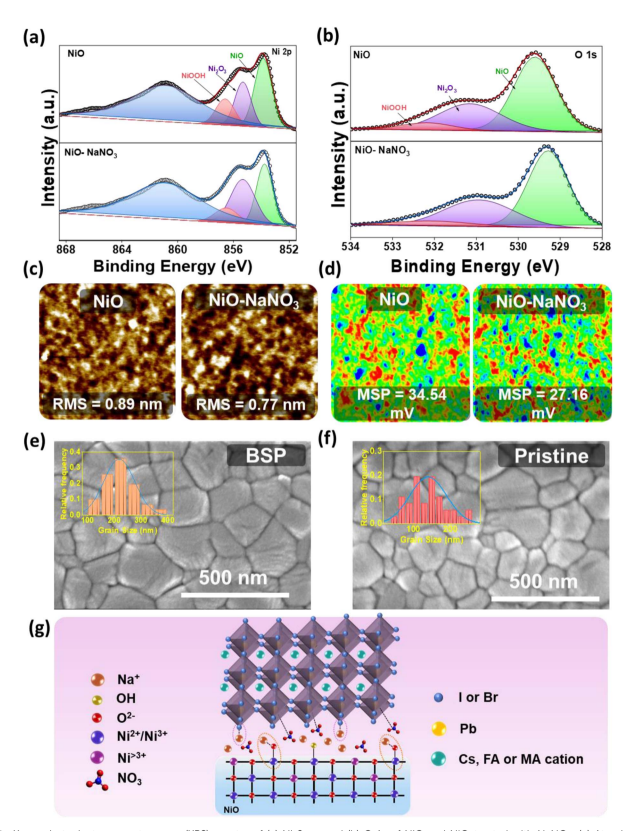


Fig. 2 X-ray photoelectron spectroscopy (XPS) spectra of (a) Ni $2p_{3/2}$ and (b) O 1s of NiO and NiO-treated with NaNO₃. (c) Atomic force microscopy (AFM) and (d) scanning Kelvin probe microscopy (SKPM) images of NiO and NiO treated with NaNO₃, (e) FESEM images of the perovskite layer for BSP (inset shows average grain size distribution) and (f) pristine (inset shows average grain size distribution). (g) Probable mechanism for interaction of NaNO₃ at the NiO/perovskite interface.

demonstrate the influence of NaNO3 treatment on the surface morphological qualities of the spin-coated perovskite film, as shown in Fig. 2e and f, respectively. Contrasted to perovskite spin-coated on pristine NiO, a smooth and uniform morphology with a greatly increased grain size, from ~150 nm to over 350 nm, was obtained for NaNO₃-treated NiO, as shown in the inset of Fig. 2e and f. This improvement in the grain size correlated well with the contact angle measurements, which revealed the hydrophobic nature of the modified layer, as depicted in Fig. S4.† The contact angle (CA) value for NiO was measured to be 26.44, which significantly improved to 53.14 for NiO/NaNO₃. This implies a reduction in nucleation sites for perovskite and promotes crystal growth. This phenomenon is attributed to the homogeneous nucleation and growth of perovskite, ultimately leading to the formation of larger grains.30,41 This suggests that NaNO3 treatment could not only passivate the NiO surface but also increase grain size, reduce trap states at the NiO/perovskite interface, and eliminate grain boundary defects, thus improving the charge carrier extraction leading to devices with improved cell parameters. 5,42

Depicting upon the preceding analysis, we propose that, considering the robust ionic bond of NaI/Br, it is reasonable to suggest that Na $^+$ ions from NaNO $_3$ could readily combine to form NaI/Br at the interface of NiO and perovskite. ^{43,44} The NaX (X = I/Br) formed in this way could readily link with PbX $_2$ and act as a nucleus for perovskite growth. ⁴³ The above results and observation are consistent with the XRD result of samples without annealing the perovskite layer, thus implying that the Na ions could facilitate perovskite crystal growth even without thermal annealing. FESEM analysis also supports this evidence, where the perovskite grain size was distinctly improved with a smooth morphology. At the same time, the Na ions could replace the hydroxyl groups in NiO with Na, thus forming a Ni–O–Na bond and aiding in the reduction of the hydroxyl groups on the NiO surface (Fig. 2g).

2.2 ETL interface engineering

After gaining insights into the HTL/perovskite interface, we studied the ETL/perovskite interface in terms of the possibility of a reaction at the surface. To this end, XPS measurements were conducted on top of perovskite films after NaNO $_3$ treatment, *i.e.*, for TSP films, to validate the chemical interactions between the photoactive perovskite material and the NaNO $_3$ small molecules. As exhibited in Fig. 3a and b, the characteristic peaks of Pb $4f_{7/2}$ and $4f_{5/2}$ and N 1s shift clearly toward lower binding energies than the pristine perovskite film, thus indicating the possibility of interactions between perovskite and passivation molecules, implying that the defects are electrostatically passivated. 6,45

Most importantly, the atomic percentage of O 1s improved from 2.64 to 4.11%, as shown in Table S3.† Furthermore, when the relative ratio of iodine and oxygen was calculated, it was observed that the oxygen/iodine ratio increased from 0.0715 (pristine) to 0.119 (TSP). The ratio of oxygen/nitrogen was also found to increase from 0.14 to 0.25 for pristine and TSP samples, respectively. This clearly implies that the relative

oxygen percentage is higher even after accounting for the inevitable bonding of oxygen in the environment, and it is evident that sodium nitrate exists on the perovskite's surface.

SKPM and UPS analyses were carried out to study the effect of NaNO₃ solution treatment on the electronic states of perovskite. Fig. 3c reveals the surface potential photo-images of pristine and TSP-based perovskite films. As observed from the images, the mean surface potential of the pristine perovskite film was 22 mV, which increased by almost 12 mV and reached 34 mV after NaNO₃ treatment. These results indicate that there is an up-shift in the quasi-Fermi level, which is consistent with previous reports. ^{14,46} This trend was in line with the workfunction change obtained by UPS analysis (Fig. 3d). The Fermi level of TSP was calculated to be -4.97 eV, whereas that of pristine devices was calculated to be -5.08 eV (inset of Fig. 3d). This up-shift by 0.11 mV implies an increase in the built-in potential that consequently leads to an enhancement in the open-circuit voltage ($V_{\rm oc}$) of devices. ^{14,47}

Similarly, the FTIR analysis of perovskite and perovskite/NaNO₃ films was conducted to understand the interactions occurring at the interfaces, as shown in Fig. 3e. The FTIR spectra were consistent with the spectra for triple cation-based perovskite films. ^{48,49} The characteristic –C=N symmetric stretch around ~1700 cm⁻¹ of perovskite shifted to a lower wavenumber along with reduced intensity, thus indicating the possibility of bonding between the lone pair of electrons available on nitrate ions with perovskite *via* coordination bonding. ⁴⁶ Furthermore, a small peak appeared around 1341.13 cm⁻¹ with a lower wavenumber shift from its original value (~1383 cm⁻¹) in the sodium nitrate-treated samples, thus indicating interaction of nitrate ions on the perovskite surface (Fig. 3f). These results correlated well with XPS analysis, further confirming the interaction of nitrate ions on the perovskite surface.

Steady-state photoluminescence (PL) spectroscopy was performed for glass/NiO/perovskite (pristine) and glass/NiO/ perovskite/NaNO3 (TSP) films to assess the impact of the passivation strategy on the ultra-fast charge transfer behaviors and the trap states. As exhibited in Fig. 4a, the TSP samples showed a significant increase in PL intensity compared to that of the pristine sample, indicating that the top-side passivated layer reduced the non-radiative recombination of the photogenerated carriers. Furthermore, to ascertain this observation, time-resolved PL (TRPL) and decay transient analyses for pristine and TSP samples were conducted as shown in Fig. 4b. Biexponential functions that correspond to two decay functions were used to fit the TRPL curves. 43 The TRPL curves were plotted and fitted with a biexponential decay fitting function, thus resulting in two constituents' system (τ_1 (short-lived) and τ_2 (long-lived)). The τ_1 indicates the monomolecular recombination time span, whereas the τ_2 shows the time span for the recombination of charges. The TRPL analysis fitting parameters are summarized in Table S4†. The average lifetime (τ_{PL}) of the pristine film was around 40.37 ns, whereas the TSP showed a lifetime of 52.12 ns. TSP perovskite showed a longer carrier lifetime (52.12 ns) than the pristine perovskite sample (40.37 ns), implying that the trap states have been efficiently eliminated for effective charge extraction towards the ETL.50

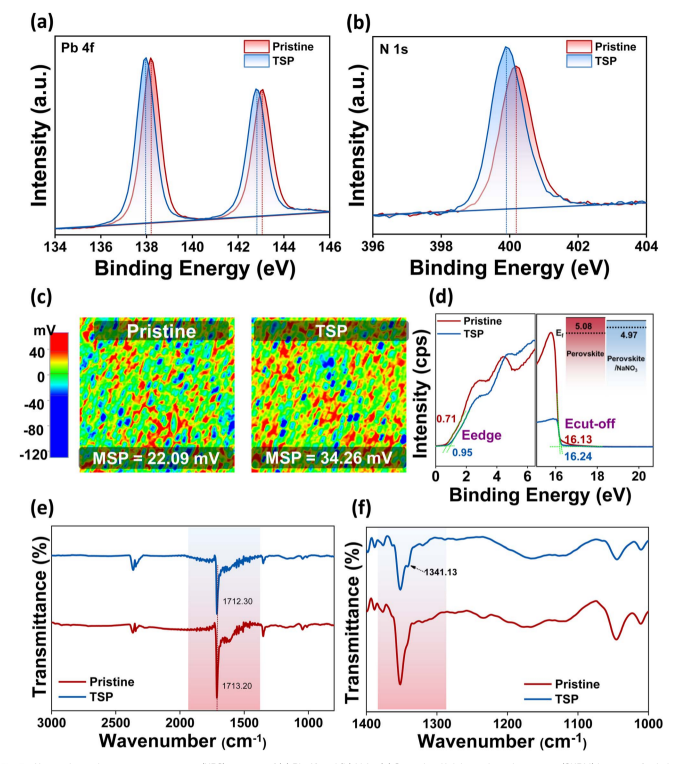


Fig. 3 X-ray photoelectron spectroscopy (XPS) spectra of (a) Pb 4f and (b) N 1s. (c) Scanning Kelvin probe microscopy (SKPM) images of pristine and TSP perovskite films. (MSP = mean surface potential). (d) UPS spectra of the perovskite and TSP films (inset shows energy level alignment). (e) FTIR spectra of pristine and TSP films. (f) FTIR spectra of pristine and TSP in the expanded fingerprint region for 1400–1000 cm⁻¹.

From the above analysis, we propose that there is a possibility that the nitrate group could interact with unreacted Pb²⁺, which leads to the formation of Pb(NO₃)₂ as shown in eqn (1).

$$Pb^{2+} + NO_3^- \to Pb(NO_3)_2$$
 (1)

The lead nitrate formed at the surface might serve as a water scavenger and is expected to improve the moisture stability of perovskite.51 Meanwhile, Na+ ions could possibly interact with I or Br, thus resulting in the formation of NaI or NaBr.43 Furthermore, Na⁺ ions could diffuse into the inside perovskite

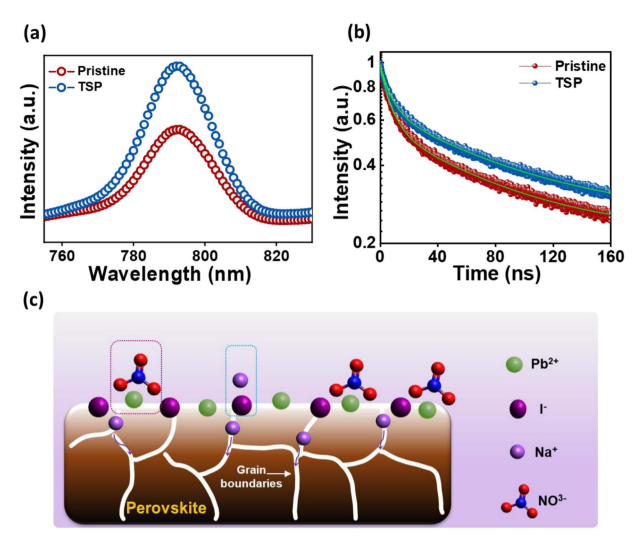


Fig. 4 (a) PL spectra for pristine and TSP films. (b) TRPL spectra for pristine and TSP films. (c) Probable mechanism for interaction of NaNO $_3$ at the perovskite/ETL interface.

layer, thus passivating the defects within the perovskite layer, as depicted in Fig. 4c.

2.3 Dual interface engineering

The photovoltaic parameters of all passivation strategies were measured at optimal concentrations for passivating the bottom surface (BSP), the top surface (TSP), and both surfaces (DSP). Fig. 5a shows the average performances of pristine, BSP, TSP, and DSP devices. The DSP devices exhibited the best performance justifying the passivation effect on both sides. All other parameters, including V_{oc} , J_{sc} , and FF, were also improved for the DSP devices compared to the BSP, TSP, and pristine devices (Fig. S6 and Table S5†). Furthermore, to verify the effect of NaNO₃, we fabricated devices without heating of the spincoated perovskite active layer, as presented in Fig. S7.† To our surprise, the devices with NaNO3 showed improved performance, and the DSP again showed the best among all devices.

The pristine devices showed PCE up to 19.08%, with a V_{oc} of 1.06 V, J_{sc} of 22.63 mA cm⁻², and FF of 75.18%. Meanwhile the DSP strategy-based fabricated device exhibited a notably

superior $V_{\rm oc}$ up to 1.12 V, $J_{\rm sc}$ of 24.20 mA cm⁻², and FF of 79.49%, thus resulting in an increased PCE of 21.53%, which is an improvement of 12.84% compared to the control devices (Fig. 5b). Moreover, V_{oc} and FF were improved more than 5%, while $J_{\rm sc}$ was improved ~7% after DSP treatment compared to the pristine devices. Furthermore, the histogram of the PCE distribution for pristine and DSP-based devices, as summarized in Fig. 5c, showed improved values for DSP devices as compared to pristine. These photovoltaic results were confirmed through external quantum efficiency (EQE), where DSP-based devices showed a better photon-to-current conversion efficiency, implying that they have better charge-collecting ability than pristine devices for the overall spectrum (Fig. 5d).

An additional significant study of the charge carrier dynamics of the IPeSCs was performed: transient photovoltage (TPV), transient photocurrent (TPC), the dependency of V_{oc} on the intensities of light, and SCLC measurements. The lifetime of photo-induced charge carriers can be directly monitored using TPV by measuring changes in solar cell photovoltage, while TPC enables the determination of operational-time

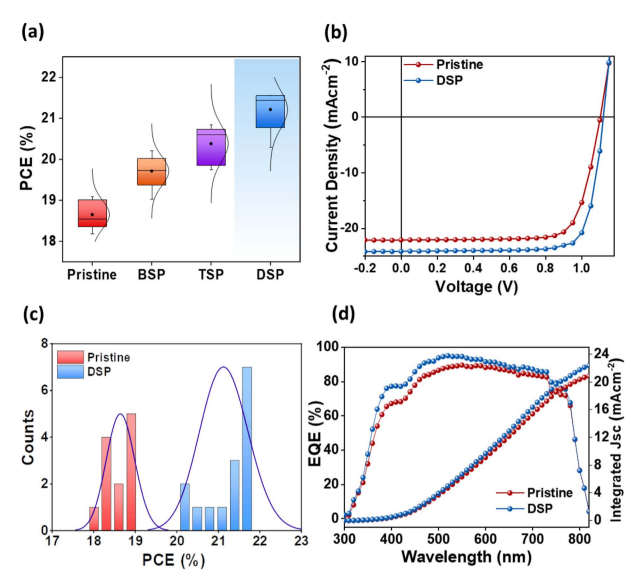


Fig. 5 (a) Statistical analysis of PCE for pristine, BSP, TSP and DSP devices. (b) J-V characteristics of pristine and DSP based devices. (c) Statistical histogram of PCE for pristine and DSP devices. (d) External quantum efficiency curves of the pristine and DSP based devices.

reliant charge extraction features of photo-induced charge carriers in IPeSCs.25,52 The TPC and TPV measurements were carried out in a real-time operating process, and they were determined through a double-exponential fitting decay function. As illustrated in Fig. 6a, the TPC decay lifetime of PeSCs with DSP was found to be 0.71 µs, while that for pristine was found to be 0.26 µs. Meanwhile, the TPV decay time of DSP increased from 0.24 ms (pristine) to 0.31 ms, indicating improved charge extraction and reduced charge recombination (Fig. 6b).25 Hence, the overall increase in PeSC performance could be ascribed to excellent charge transport abilities with efficiently diminished charge carrier recombination caused by perovskite defects.

The dependency of V_{oc} on the light intensities of the IPeScs for pristine and DSP devices is demonstrated in Fig. 6c. The degree of recombination occurring inside the perovskite solar cells was determined by using following relation:

$$V_{\rm oc} = n \frac{kT}{e} \ln(I) + \text{constant}$$
 (2)

where n, e, k, and T indicate the ideality factor, elementary charge, Boltzmann constant, and absolute temperature, respectively. If the value of "n" is equal to 1, it indicates the reduced charge carrier recombination. Hence, from this value, we may determine the inward recombination occurring inside IPeSCs from linear characteristics V_{oc} as a function of light intensities. 53,54 The pristine devices demonstrated a high slope value of 2.66 kT/q, whereas the DSP denoted a lower slope value of 1.38 kT/q. These results imply that the double-side passivation can help reduce trap-assisted recombination sites.

SCLC analysis was performed to confirm the defect densities in the photoactive layer film with and without passivation strategies, as exhibited in Fig. 6d. The value of $V_{\rm TFL}$ was estimated to be reduced for the passivated film (0.72 V) in comparison to the pristine film (0.80 V), thus indicating that the

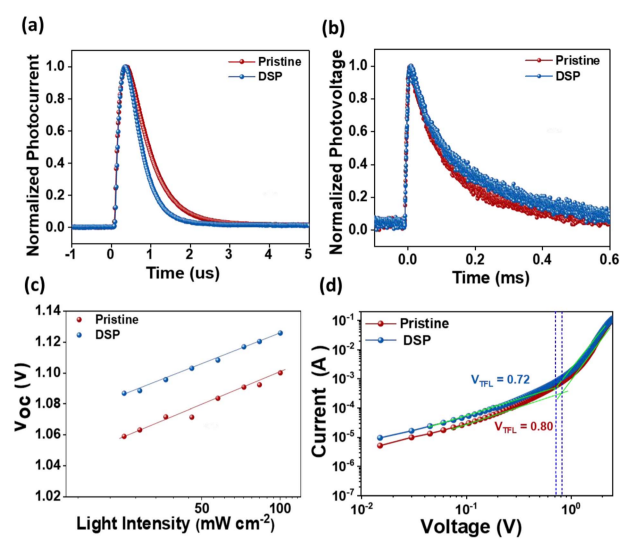


Fig. 6 (a) Transient photocurrent (TPC) and (b) transient photovoltage (TPV) of pristine and DSP devices, (c) V_{OC} versus light intensity of the devices for pristine and DSP devices and (d) SCLC measurements of hole-only devices for pristine and DSP devices.

defect densities tend to decrease with passivation. The defect densities at passivated and pristine devices were calculated to be 9.17×10^{15} and 1.02×10^{16} , respectively. These results imply that the newly employed NaNO3 can enhance perovskite filmquality via better crystallinity with large size grain and significantly decreased defect densities, thereby improving the IPeSCs' overall performance.

The interfacial instability between the CTLs is an intractable obstruction to obtaining substantially efficient and long-term stable PeSCs. It is therefore necessary to address these reactions to achieve stability in PeSCs.26 To ascertain the effect of passivation on both sides, we determined operational (MPP), light, and nitrogen long-term stability for pristine and DSP unencapsulated devices. Fig. 7a shows the devices' steady-state photocurrent density measured both under continuous illumination and at their highest power point voltage under room temperature conditions of 25 \pm 5 °C and 45 \pm 5% relative humidity. The pristine and DSP-based devices exhibited stabilized steady-state photocurrent outputs. The DSP devices attained a steady-state photocurrent of \sim 21.5 mA cm⁻², whereas the pristine devices exhibited \sim 17.9 mA cm⁻² for 500 s. Fig. 7b shows the continuous light-soaking long-term durability of the unencapsulated device when operated under ambient conditions of 25 \pm 5 °C and 45 \pm 5% relative humidity. The efficiency of DSP PeSCs degraded to 50% of their original first values in 450 h, while pristine devices showed a decreasing PCE trend of approximately 10% over the same amount of time. More specifically, we observed the T_{80} lifetime, which is defined as the time until the photovoltaic performance of the device reduces to 80% of its first value, which is 72 h and 270 h in pristine and DSP IPeSCs, respectively, thus indicating that DSP IPeSCs demonstrate excellent long-term durability. This was proven once more by the outstanding long-term stability of DSP IPeSCs in a nitrogen environment (Fig. 7c). The DSP PeSC exhibited 76% stability for more than 1800 h, while the pristine degraded to 49% for the same amount of time. Moreover, the DSP devices maintained a T_{80} of \sim 1582 h, compared to pristine, which maintained a T_{80} of 648 h. Furthermore, the T_{80} for BSP and TSP

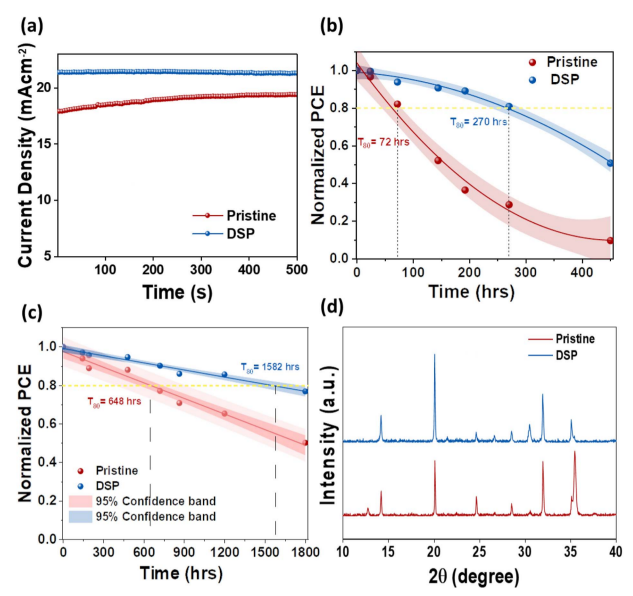


Fig. 7 (a) Steady-state photocurrent and PCE of pristine and DSP devices. (b) Light soaking stability of pristine and DSP devices. (c) Normalized PCE as a function of time for the unencapsulated devices stored in the nitrogen environment for the PSC with pristine and DSP. (d) XRD analysis for the degraded samples for pristine and DSP devices.

was measured to be 793 and 1081 h respectively (Fig. S8†). These results again signify that the DSP passivation could effectively boost the device stability. We speculate that the improvements in both light soaking and illumination stability could be associated with the improved perovskite crystallinity, with large grains, reduced grain boundary density, and an improved passivation effect from both ETL and HTL sides by DSP treatment.14 To assess the occurrence of the chemical reactions at the interfaces, we conducted XRD analysis of the aged devices after 30 days, with the results shown in Fig. 7d. The XRD peak demonstrated a PbI₂ peak at 12.8° for pristine, with a slight decreasing trend in the peak intensity of the main perovskite facets, unlike the DSP films. These results explicitly suggest that the unbound Pb2+ ions were bonded successfully, as shown in eqn (1), and that ultimately, sodium nitrate could effectively

passivate the defects from the bottom and top sides, thus increasing the DSP outcomes.

3. Conclusions

In this overview, we have successfully developed a π -conjugated oxysalt-assisted dual-interface engineering strategy using NaNO3 ionic solution, and we have also investigated the effect of each perovskite interface modification. The synergistic effect of Na⁺ and NO₃ simultaneously addressed both the surface and bulk defect issues. To elaborate, the Na⁺ ions could interact at the NiO surface, while bonding with I/Br defects from the perovskite surface. Moreover, the passivated NiO surface can promote the growth of perovskite with improved uniform large grain and crystallinity. At the perovskite/ETL interface, we speculate that Na⁺ ions diffused inside the perovskite, thus resulting in reduced trap-density, while NO₃ was coordinated with Pb²⁺, in turn leading to the formation of Pb(NO₃)₂, and eventually finetunning the alignment energy level and enlarging the device stability. These changes resulted in enhanced charge extraction and transportation at both interfaces of the perovskite. Therefore, the DSP devices showed a champion efficiency of 21.53%, surpassing those of BSP (20.16%), TSP (20.83%), and pristine (19.08%) devices. The devices also retained more than 80% stability for more than 1700 h in nitrogen for unencapsulated devices. This work paves the way to minimizing the bulk and interface non-radiative recombination losses, thus leading to simultaneous enhancement in PCE and stability through cation and anion synergistic engineering.

Author contributions

Conceptualization: P. P. and S.-N. K.; methodology: P. P.; validation: P. P. and S.-I. N.; formal analysis: P. P., S. S. and D. S. M.; investigation: P. P. and S. S.; writing - original draft: P. P.; writing - review and editing: S.-I. N.; visualization: P. P.; supervision: S.-I. N.; project administration: S.-N. K. and S.-I. N.; funding acquisition: S.-N. K. and S.-I. N.

Conflicts of interest

The authors declare no conflict of interest.

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References

- 1 G. Hodes, Science, 2013, 342, 317-318.
- 2 N. J. Jeon, J. H. Noh, W. S. Yang, Y. C. Kim, S. Ryu, J. Seo and S. I. Seok, Nature, 2015, 517, 476-480.
- 3 G. Xing, N. Mathews, S. Sun, S. S. Lim, Y. M. Lam, M. Grätzel, S. Mhaisalkar and T. C. Sum, Science, 2013, 342, 344-347.
- 4 C. Li, Y. Zhang, X. Zhang, P. Zhang, X. Yang and H. Chen, Adv. Funct. Mater., 2023, 33, 2214774.
- 5 M. Degani, Q. An, M. Albaladejo-Siguan, Y. J. Hofstetter, C. Cho, F. Paulus, G. Grancini and Y. Vaynzof, Sci. Adv., 2021, 7, eabj7930.
- 6 Y.-J. Kang, S.-N. Kwon, S.-P. Cho, Y.-H. Seo, M.-J. Choi, S.-S. Kim and S.-I. Na, ACS Energy Lett., 2020, 5, 2535-2545.
- 7 S. Gharibzadeh, P. Fassl, I. M. Hossain, P. Rohrbeck, M. Frericks, M. Schmidt, M. R. Khan, T. Abzieher, B. A. Nejand and F. Schackmar, Energy Environ. Sci., 2021, **14**, 5875-5893.
- 8 F. Li, X. Deng, F. Qi, Z. Li, D. Liu, D. Shen, M. Qin, S. Wu, F. Lin and S.-H. Jang, J. Am. Chem. Soc., 2020, 142, 20134-20142.

- 9 P. Patil, D. S. Mann, U. T. Nakate, Y.-B. Hahn, S.-N. Kwon and S.-I. Na, Chem. Eng. J., 2020, 397, 125504.
- 10 D. S. Mann, P. Patil, D.-H. Kim, S.-N. Kwon and S.-I. Na, J. Power Sources, 2020, 477, 228738.
- 11 Y.-J. Kang and S.-I. Na, Nano Energy, 2022, 97, 107193.
- 12 S.-P. Cho, H.-J. Lee, Y.-J. Kang, Y.-H. Seo and S.-I. Na, Appl. Surf. Sci., 2022, 602, 154248.
- 13 S. Thakur, S.-N. Kwon, D. S. Mann and S.-I. Na, J Mater., 2022, 8, 1165-1171.
- 14 P. Patil, D. S. Mann, S. R. Rondiya, N. Y. Dzade, S.-N. Kwon and S.-I. Na, Sol. RRL, 2021, 5, 2100547.
- 15 S. S. Mali, J. V. Patil, J. A. Steele, Y. H. Jung, M. K. Nazeeruddin and C. K. Hong, Mater. Today, 2023, 33-45
- 16 S.-M. Bang, S. S. Shin, N. J. Jeon, Y. Y. Kim, G. Kim, T.-Y. Yang and J. Seo, ACS Energy Lett., 2020, 5, 1198-1205.
- 17 M. Yang, G. Han, Y. Xiao and W. Hou, Chem. Commun., 2022, 58, 5638-5641.
- 18 S. Zhang, X. Yan, Z. Liu, H. Zhu, Z. Yang, Y. Huang, S. Liu, D. Wu, M. Pan and W. Chen, J. Energy Chem., 2021, 54, 493-500.
- 19 T. Bu, X. Liu, Y. Zhou, J. Yi, X. Huang, L. Luo, J. Xiao, Z. Ku, Y. Peng and F. Huang, Energy Environ. Sci., 2017, 10, 2509-
- 20 H. Si, C. Xu, Y. Ou, G. Zhang, W. Fan, Z. Xiong, A. Kausar, Q. Liao, Z. Zhang and A. Sattar, Nano Energy, 2020, 68, 104320.
- 21 D.-Y. Son, S.-G. Kim, J.-Y. Seo, S.-H. Lee, H. Shin, D. Lee and N.-G. Park, J. Am. Chem. Soc., 2018, 140, 1358-1364.
- 22 C. Zhang, H. Wang, H. Li, Q. Zhuang, C. Gong, X. Hu, W. Cai, S. Zhao, J. Chen and Z. Zang, J. Energy Chem., 2021, 63, 452-460.
- 23 W. Fan, Y. Shen, K. Deng, Q. Chen, Y. Bai and L. Li, Nano Energy, 2022, 100, 107518.
- 24 W. Chen, Y. Zhou, G. Chen, Y. Wu, B. Tu, F. Z. Liu, L. Huang, A. M. C. Ng, A. B. Djurišić and Z. He, Adv. Energy Mater., 2019, 9, 1803872.
- 25 Y. Guo, H. Lei, L. Xiong, B. Li and G. Fang, J. Mater. Chem. A, 2018, 6, 2157-2165.
- 26 J. Zhang, J. Yang, R. Dai, W. Sheng, Y. Su, Y. Zhong, X. Li, L. Tan and Y. Chen, Adv. Energy Mater., 2022, 12, 2103674.
- 27 S. Zhang, S. Wu, R. Chen, W. Chen, Y. Huang, H. Zhu, Z. Yang and W. Chen, J. Phys. Chem. Lett., 2019, 10, 2898-2903.
- 28 G. Zheng, C. Zhu, J. Ma, X. Zhang, G. Tang, R. Li, Y. Chen, L. Li, J. Hu and J. Hong, Nat. Commun., 2018, 9, 2793.
- 29 Y. Tidhar, E. Edri, H. Weissman, D. Zohar, G. Hodes, D. Cahen, B. Rybtchinski and S. Kirmayer, J. Am. Chem. Soc., 2014, 136, 13249-13256.
- 30 X. Xu, Y. Sun, D. He, Z. Liang, G. Liu, S. Xu, Z. Li, L. Zhu and X. Pan, J. Mater. Chem. C, 2021, 9, 208-213.
- 31 S. Zhang, S. Wu, R. Chen, W. Chen, Y. Huang, Z. Yang and W. Chen, J. Mater. Chem. C, 2020, 8, 1642-1648.
- 32 D. Zhang, X. Zhang, T. Guo, J. Zou, Y. Zhou, J. Jin, Z. Zhu, Q. Cao, J. Zhang and Q. Tai, Small, 2023, 19, 2205604.
- 33 J. Huang, G. Li and Y. Yang, Adv. Mater., 2008, 20, 415-419.

- 34 J. Xia, J. Luo, H. Yang, F. Zhao, Z. Wan, H. A. Malik, Y. Shi, K. Han, X. Yao and C. Jia, Adv. Funct. Mater., 2020, 30, 2001418.
- 35 D. S. Mann, S.-N. Kwon, P. Patil and S.-I. Na, *Nano Energy*, 2023, **106**, 108062.
- 36 D. Di Girolamo, N. Phung, M. Jošt, A. Al-Ashouri, G. Chistiakova, J. Li, J. A. Márquez, T. Unold, L. Korte and S. Albrecht, *Adv. Mater. Interfaces*, 2019, 6, 1900789.
- 37 J. Ciro, D. Ramírez, M. A. Mejia Escobar, J. F. Montoya, S. Mesa, R. Betancur and F. Jaramillo, ACS Appl. Mater. Interfaces, 2017, 9, 12348–12354.
- 38 T. Guo, Z. Fang, Z. Zhang, Z. Deng, R. Zhao, J. Zhang, M. Shang, X. Liu, Z. Hu and Y. Zhu, *J. Energy Chem.*, 2022, 69, 211–220.
- 39 S. Ahmad, R. Ma, J. Zheng, C. K. Gary Kwok, Q. Zhou, Z. Ren, J. Kim, X. He, X. Zhang and K. M. Yu, *Small Methods*, 2022, 6, 2200787.
- 40 C. Bi, Q. Wang, Y. Shao, Y. Yuan, Z. Xiao and J. Huang, *Nat. Commun.*, 2015, **6**, 1–7.
- 41 C. Bi, Q. Wang, Y. Shao, Y. Yuan, Z. Xiao and J. Huang, *Nat. Commun.*, 2015, 6, 7747.
- 42 K. M. Reza, A. Gurung, B. Bahrami, S. Mabrouk, H. Elbohy, R. Pathak, K. Chen, A. H. Chowdhury, M. T. Rahman and S. Letourneau, *J. Energy Chem.*, 2020, 44, 41–50.
- 43 T. Bu, J. Li, F. Zheng, W. Chen, X. Wen, Z. Ku, Y. Peng, J. Zhong, Y.-B. Cheng and F. Huang, *Nat. Commun.*, 2018, 9, 4609.

- 44 D. J. Kubicki, D. Prochowicz, A. Hofstetter, S. M. Zakeeruddin, M. Grätzel and L. Emsley, *J. Am. Chem. Soc.*, 2018, 140, 7232–7238.
- 45 M.-J. Choi, Y.-S. Lee, I. H. Cho, S. S. Kim, D.-H. Kim, S.-N. Kwon and S.-I. Na, *Nano Energy*, 2020, **71**, 104639.
- 46 P. Patil, A. Maibam, S. S. Sangale, D. S. Mann, H.-J. Lee, S. Krishnamurty, S.-N. Kwon and S.-I. Na, ACS Appl. Mater. Interfaces, 2023, 29597–29608.
- 47 S. Ding, S. Li, Q. Sun, Y. Wu, Y. Liu, Z. Li, Y. Cui, H. Wang, Y. Hao and Y. Wu, *J. Mater. Chem. C*, 2019, 7, 5686–5694.
- 48 J. Yang, Q. Hong, Z. Yuan, R. Xu, X. Guo, S. Xiong, X. Liu, S. Braun, Y. Li and J. Tang, *Adv. Opt. Mater.*, 2018, **6**, 1800262.
- 49 K. Hills-Kimball, Y. Nagaoka, C. Cao, E. Chaykovsky and O. Chen, *J. Mater. Chem. C*, 2017, **5**, 5680–5684.
- 50 M. A. Mahmud, T. Duong, Y. Yin, H. T. Pham, D. Walter, J. Peng, Y. Wu, L. Li, H. Shen and N. Wu, *Adv. Funct. Mater.*, 2020, 30, 1907962.
- 51 D. V. Shinde, L. Pyeon, M. Pei, G.-W. Kim, H. Yang and T. Park, ACS Appl. Mater. Interfaces, 2017, 9, 14023–14030.
- 52 C. R. McNeill, I. Hwang and N. C. Greenham, *J. Appl. Phys.*, 2009, **106**, 024507.
- 53 Y. Yang, H. Peng, C. Liu, Z. Arain, Y. Ding, S. Ma, X. Liu, T. Hayat, A. Alsaedi and S. Dai, *J. Mater. Chem. A*, 2019, 7, 6450–6458.
- 54 Y. Huang, S. Wu, R. Chen, S. Fang, S. Zhang, G. Wang and W. Chen, ACS Appl. Mater. Interfaces, 2019, 11, 18415–18422.