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Introduction

Colloidal quantum dots (QDs) have emerged as promising candidates for optoelectronic devices, such as photodetectors, owing to their unique optoelectronic properties, low-temperature solution processing, easy device integration, etc.¹ Among various types of QDs, alloyed chalcogenide QDs have garnered significant attention due to their exceptional performance characteristics, including broad spectral tunability, low defect state density, smooth charge transfer channels, and excellent photostability.^{2,3} However, it is challenging to obtain multielement chalcogenide alloyed QDs that have significant amounts of traps during the growth periods due to the chemical reactivity difference, easy oxidation of elements, etc.4,5 The localized arrangement of traps tends to form charge combination centers hindering the charge transfer process, which deteriorate the optical properties of QDs and resultant photo-

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Defect passivation engineering of chalcogenide quantum dots via in situ fluorination treatment*

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Efficient carrier transport through reducing the traps in chalcogenide quantum dots (QDs) is crucial for their application in optoelectronic devices. This study introduces an innovative in situ fluorination treatment to remove the whole-body traps of metal chalcogenide QDs. further accelerating the carrier transport process. The selected benzene carbonyl fluoride (BF) molecular additive can efficiently peel off the caused oxide traps and dangling bonds of chalcogenide QDs in real-time through the continuous release of the HF gas of BF decomposition. Experimental results revealed that the obtained chalcogenide QDs with in situ fluorination treatment can accelerate the charge extraction and hinder the charge recombination. Finally, two types of photo-electrical conversion devices, consisting of photodetectors and sensitized solar cells, are fabricated to reveal the advantages of in situ fluorination treatment of QDs. Our findings highlight in situ fluorination treatment towards chalcogenide QDs as a viable approach to reduce the traps of QDs and improve the performance in optoelectronic technologies, offering hope for the practical application of this technology in the near future.

> electrical conversion applications.^{6,7} Therefore, focusing on the removal of trap states and surface passivation towards chalcogenide QDs is crucial to improving the carrier dynamics transport for constructing high-performance optoelectronic devices.5,8

> Owing to the high surface-to-volume ratio, individual QDs can present a relatively large number of defects that trap optical or electrically injected carriers and adsorb water and oxygen molecules, resulting in lowered optical properties and stability.9,10 In particular, for the alloyed chalcogenide QDs, the highly reactive Te atoms are usually unstable and easily oxidized in the whole-body crystals in the nucleation and growth process. Although many efforts have been made to devise surface chemistry approaches to eliminating defects (oxides and dangling bonds) on the surface of QDs, it is commonly impossible to remove the inner body defects caused during the nucleation and growth periods, such as that caused by the precursor oxide, yet this should not be ignored.^{11,12} Therefore, paying attention to the whole-body defects of QDs is crucial to enhancing the charge transport and improving the performance of optoelectronic devices.13 Fluorination treatment strategies have been widely used to improve optoelectronic device efficiency and stability from surface trap removal and passivation of various nanocrystals.^{12,14-16} The fluorine anion (F⁻) is well recognized as one of the most efficient oxygen defect quenchers owing to its unique ionic characteristics, notably its high electronegativity value of 3.98. Meanwhile, the remarkable similarity in ionic radii between



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fluorine and oxygen, at 1.36 and 1.40, respectively, combined with their electrically passive behavior as a heavy dopant, suggests that fluorine could be easily integrated into the whole-body nanocrystal to modify its structural and optical properties, as well as its surface morphology.^{12,14-17} However, in previous studies, most fluorination treatments were usually used for the photoluminescence enhancement of nanocrystals, such as perovskite and InP QDs, for light-emitting applications. This simple fluorination treatment for photo-to-electrical conversion applications of QDs, such as photodetectors and solar cells, is rare. Moreover, most fluorination treatments are post-processed after synthesizing the QDs, which is relatively restricted to modulating the defect states of the whole nanocrystals. Therefore, developing a simple and efficient fluorination treatment strategy for chalcogenide QDs from the whole-body defect passivation and revealing the charge transport process for photo-electrical conversion devices are still crucial.

Here, we present a feasible and efficient in situ fluorination treatment approach for peeling off the whole-body traps and enhancing the charge transport of chalcogenide QDs. By the in situ use of a benzene carbonyl fluoride (BF) molecule additive in the one-pot synthesis of CdSeTe QDs, the oxide defects and dangling bonds caused during the growth process can be efficiently peeled off in real-time through the continuous release of the HF gas from BF decomposition. Experimental results revealed that the obtained chalcogenide QDs with in situ fluorination treatment facilitate effective surface defect passivation and significantly enhance charge carrier mobility. Furthermore, fabricating the photodetectors and sensitized solar cells needs evaluation of the photo-electrical conversion performance of the target CdSeTe QDs with in situ fluorination treatment. As a result, the self-powered photodetector devices exhibited a responsivity and specific detectivity up to 280 mA W^{-1} and 10^8 Jones under the broad spectral range to nearinfrared region illumination, respectively. Meanwhile, the



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Qingdao University. His main research interests include semiconductor quantum dot synthesis and their optoelectronic device applications, such as solar cells and photodetectors. current density of the target QD-based photodetectors with *in situ* fluorination treatment was two orders of magnitude higher than that of the control ones. In addition, the target QD-based sensitized solar cells have improved the photovoltaic parameters. Our results underscore the potential of *in situ* fluorination treatment as a promising strategy for unlocking the full potential of chalcogenide QDs in optoelectronic technologies.

Results and discussion

Generally, CdSeTe chalcogenide QDs are synthesized through the one-pot non-injection method according to our previous literature.^{11,18} The reaction mixture composed of oleate-Cd, TOP-Se, and TOP-Te in paraffin media is directly nucleated and grows at high temperatures (the detailed synthetic procedures are described in the ESI[†]). High-angle annular darkfield scanning transmission electron microscopy (HAADF-STEM) images of the control and target CdSeTe QDs without and with in situ fluorination treatment were obtained, as shown in Fig. 1a-d. As depicted in Fig. 1a and c, the size of the target CdSeTe QDs with in situ fluorination treatment is a little smaller than that of the control one, indicating the surface etching of the F⁻ ions. The STEM energy dispersive X-ray spectroscopy (EDS) mapping result exhibited the Se and Te element distribution of CdSeTe ODs, which are shown in Fig. 1b and d. Both the target and control QDs exhibited similar Se/Te element distribution with the Te-rich core and Se-rich shell of a graded alloy structure. The structure and morphology of QDs remained unchanged by the in situ fluorination treatment method. Meanwhile, the graded alloy structure with low interface lattice strain and defects is required for efficient charge transport for photo-electrical conversion applications, especially photodetectors and solar cells.¹⁹ To determine the elemental composition of the as-prepared QDs, inductively coupled plasma atomic emission spectrometry (ICP-AES) was performed and the results are shown in Fig. 1e. Three separate ICP-AES analyses were performed for each sample. Compared to the control one, the ratio of three elements in the target QDs is slightly changed, further indicating that introducing the BF molecule slightly influences the nucleation and growth process of CdSeTe QDs.²⁰ The contents of the Cd and Se elements of the target QDs are slightly higher, while that of the Te element is lower than those of the control. This result is mainly because the BF molecules mixed with the precursors can influence the nucleation and growth rate of CdSeTe QDs via the random collision probabilities between ions to form the Cd-Te and Cd-Se bonds.

The crystalline structure of the as-obtained CdSeTe QDs was measured *via* X-ray diffraction (XRD) (Fig. 1f). The XRD patterns of the CdSeTe QDs matched well with each other, indicating that the crystal structure of the target QDs remained unchanged. Both samples exhibit three diffraction peaks corresponding to the (111), (200), and (311) planes of the zinc blende (ZB) structure (JCPDS 00-041-1324). The sharpness of



Fig. 1 (a and c) HAADF-STEM image, (b and d) STEM-EDS mapping of the target and control CdSeTe QDs with and without *in situ* fluorination treatment, respectively. The scale bar is 10 nm. (e) ICP element analysis, (f) XRD, and (g) Te 3d XPS results of the target and control CdSeTe QDs. (h) The possible reaction mechanism of the target CdSeTe QDs with *in situ* fluorination treatment.

each peak in the target QDs showed that the introduction of the BF molecule additive can improve the crystallinity of CdSeTe QDs. X-ray photoelectron spectroscopy (XPS) measurement was carried out, and the results are shown in Fig. 1g and Fig. S1.† High-resolution Te 3d XPS spectra show that the clear peaks at 569.3 eV and 579.6 eV can be detected from the TeO₂ species in the control QDs. However, these peaks corresponding to TeO₂ were removed in the target QDs. This result confirms that BF molecules can effectively etch the surface metal oxides. As for the Cd and Se ions, the peak position did not change after the BF additive, which revealed that the Cd and Se ions are much more stable than Te.¹¹ Finally, the possible in situ fluorination mechanism towards CdSeTe QDs is demonstrated in Fig. 1h. The BF molecule is expected to react with the oleic acid solvent and safely form the F-related intermediates and activated oxygen-negative ions. The F-related intermediates can release HF gas and remove the surface defects of QDs during the nucleation and growth periods. To validate the proposed reaction mechanism with the introduction of the BF additive, ¹⁹F nuclear magnetic resonance (NMR) measurements were performed, and the results are shown in Fig. S2.[†] When mixed with Cd(Ac)₂ oleic acid solution, the BF molecule was quickly transformed into three ¹⁹F resonances at room temperature at 18, -171, and -185 ppm, respectively. The ¹⁹F resonance at 18 ppm was similar to that of the BF molecule mixed with oleic acid, consistent with the ketal intermediate formation.¹⁷ Upon heating the mixture to 320 °C, the ¹⁹F resonance located at 18 ppm disappeared, indicating that the balance of the intermediate state was disturbed, accompanied by the generation of HF gas from BF molecular decomposition.¹⁷

To investigate the nucleation and growth process of the target and control CdSeTe QDs with and without *in situ* fluorination treatment, UV-Vis absorption and photoluminescence (PL) emission characterization is carried out at an interval of one minute as a function of reaction time.^{21,22} The time-resolved absorption spectra of the control and target CdSeTe QDs are depicted in Fig. S3.† Compared to the control one, an absorption feature at the target CdSeTe QDs appeared at the beginning due to the formation of CdSe clusters. With the reaction of Te ions, the absorption feature of the target CdSeTe QDs gradually disappeared and redshifted to the NIR region.²¹ During the whole reaction process of the control CdSeTe QDs, the absorption spectra were always featureless, which indicated the initial Te reaction with Cd ions to form the CdTe clusters.²¹

the control and target CdSeTe QDs also revealed the formation of CdSeTe nanocrystals. The UV-vis absorption and PL spectra of the control and target QDs are shown in Fig. 2a. The absorption edge of CdSeTe QDs extended to the NIR region at about ~850 nm. After introducing BF additives, the absorption behavior is much higher in the whole area, and the PL emission peak position is slightly red-shifted from 792 nm to 820 nm.²³ Besides, the much narrower full width at half-maximum (FWHM) further reveals that the alloyed QDs do not contain a heterogeneous population of amorphous clusters but are highly crystalline in structure and monodisperse in size.¹¹ In Fig. 2b, it can be observed that the target CdSeTe OD films exhibit slightly higher absorption coefficient α values in the whole photon energy range compared to the control film, indicating the higher sunlight capture ability with the removal and passivation of the whole-body crystal traps. In addition, the plot of $\ln \alpha vs$. energy was used to extract the Urbach energy $(E_{\rm u})$ of the target and control CdSeTe QD films (Fig. 2c). The $E_{\rm u}$ value represents dislocations, strain, and divergence from ideal stoichiometry, as well as dynamic phonon disorder and static structural disorder caused by crystal lattice point defects.²⁴ Typically, an exponential component known as the Urbach tail appears along the absorption coefficient curve and at the optical band edge.²⁵ The $E_{\rm u}$ value also represents the tail of localized states in the band gap. The $E_{\rm u}$ value of the control film was 0.16 eV, whereas it decreased to 0.14 eV with the target film with in situ fluorination treatment derived from the lower band edge disorder and trap density.23 Transient absorption (TA) spectroscopy was further utilized to study the influ-

ence of *in situ* fluorination treatment of CdSeTe QDs. The measured carrier dynamic curves of the target and control CdSeTe QDs are shown in Fig. 2d and e. Photoexcited QD materials exhibit transient absorption spectra with two negative absorption bands peaking at about 500 and 620 nm and a large positive absorption band in the 650–680 nm area. Both bleach peaks arise at 500 and 620 nm, identical to the excitonic peaks in CdSeTe QD steady-state absorption. The TA kinetic traces of the control and target CdSeTe QDs with global analysis are shown in Fig. 2f. The double exponential fitting results of the TA kinetics were analyzed. The excited state lifetime of the CdSeTe QDs subjected to *in situ* fluorination is longer than that of the samples due to trap removal and passivation.

The corresponding E_g values of the control and target QDs were estimated by extrapolating the linear portion of the plot of $(ah\nu)^2 vs. h\nu$ with the results shown in Fig. 3a, in which the *x*-axis intercept of an extrapolated line from the linear regime of the curve represented the E_g value. It is clear that the E_g value slightly reduced from 1.84 eV to 1.78 eV after introducing the BF additive. Ultraviolet photoelectron spectroscopy (UPS) characterization was used to reveal the conduction band (CB) and valence band (VB) edge positions of typical control and target CdSeTe QDs with results shown in Fig. 3b and Table S1.[†] The ionization potentials (equivalent to the VB energy level) of control and target CdSeTe QDs were calculated to be -5.29 and -5.41 eV, respectively, by subtracting the excitation energy (21.22 eV) from the width of UPS images. The CB energy level is thus estimated to be at -3.45 and -3.63 eV. The



Fig. 2 (a) The extracted UV-vis and PL emission of the final control and target CdSeTe QDs. (b) Absorption coefficient (α) and (c) plot of ln α vs. energy used to extract the Urbach energy of the target and control CdSeTe QD films. Ultrafast time-resolved TA spectra of (d) the target and (e) control CdSeTe QDs in toluene. (f) Transient absorption kinetics of the target and control CdSeTe QDs under identical conditions.



Fig. 3 (a) Tauc plots of $(\alpha h\nu)^2$ as a function of the photon energy $(h\nu)$, (b) high and low binding energy cut-off, (c) the resulting energy level of the target and control CdSeTe QDs with and without *in situ* fluorination treatment, (d) Tr-PL spectra of the control and target CdSeTe QD deposition on the TiO₂ electrodes, (e and f) time-of-flight measurements of TiO₂/QDs/Ag devices. (g and h) Schematic illustration of the trap state distribution and carrier localization for the control and target QD films and the corresponding charge transfer process diagram.

higher CB level of the target CdSeTe QDs is due to the composition change of the elemental ratio, as discussed above. The schematic diagrams of the corresponding energy level distribution are presented in Fig. 3c. The suitable CB level of TiO₂ of the control QDs promotes the driving force for electron injection from QDs into TiO₂, consequently improving the electron injection rate and photovoltaic performance. A series of transient measurements were performed to investigate further the influence of fluorination treatment on the photogenerated carrier dynamics of QDs. The transient PL decay curves for the control and target QDs are displayed in Fig. 3d, revealing a reduced average PL lifetime from 12.68 to 9.89 ns. Both the PL quenching and shortened PL lifetime of the QD films with in situ fluorination treatment demonstrate the photoexcited electron transfer from QDs to the TiO₂ electron transparent layer. The time of flight (TOF) method was also used to measure carrier mobility.^{26,27} A pulse laser of 375 nm was irradiated on the device, and the transit times under different biases were measured with an oscilloscope (Fig. 3e and f). The estimated transit time is proportional to the reciprocal of the applied bias, and the carrier mobility can be calculated from the slope divided by the thickness square. The

carrier mobility of the target CdSeTe QDs is much higher than that of the control one.²⁸ Therefore, as shown in Fig. 3g and h, *in situ* fluorination treatment effectively removes the oxide and dangling bond defects of CdSeTe QDs, further hinders the charge recombination, and accelerates the charge transport process.

To investigate the influence of the *in situ* fluorination treatment of CdSeTe QDs towards the carrier extraction and transfer process, we continued to construct self-powered photodetectors with a device configuration of ITO/TiO2/CdSeTe QDs/ PF2/Ag (Fig. 4a). The detailed fabrication of the self-powered photodetectors are demonstrated in the ESI.† The cross-sectional scanning electron microscopy (SEM) image of the QDbased photodetector device is shown in Fig. S4a;† the thickness of the TiO₂/QD active layer is approximately 0.4 nm. Here, the polymer 2-(4-[(3S,5S)-5-[(3,3-difluoropyrrolidin-1-yl)carbonyl]pyrrolidin-3-yl]piperazin-1-yl)pyrimidine (PF2) was first selected as the hole transport material. The energy level of the PF2 polymer is well-matchable with CdSeTe QDs. Meanwhile, as previously reported, the PF2 can form a much more uniform and smooth film than other hole-transport materials, reducing the interface charge recombination and accelerating



Fig. 4 (a) Schematic structure of the self-powered broadband photodetector, (b) energy band diagram and the selected polymer hole transport materials of the photodetector. The molecular structure of the PF2 hole-transport materials. (c) Current–voltage (I-V) curves of the control and target CdSeTe QD-based photodetectors in the dark and under standard sunlight (1 sun, 100 mW cm⁻²) illumination conditions. (d) Time-dependent photocurrent curves (I-t) of photodetectors of the control and target CdSeTe QD photodetectors under the standard sunlight at 0 V bias. (e) EQE spectra of CdSeTe QD photodetectors. (f) Responsivity (*R*) and specific detectivity (*D**) spectra of the control and target CdSeTe QD photodetectors as a function of the incident light wavelength. Time-dependent photocurrent curves of the control (g) and the target (h) CdSeTe QD photodetectors under a 606 nm laser with different illumination power densities at 0 V bias.

the charge transfer.²⁹ Fig. 4b shows the corresponding band energy level diagram of the photodetector and proposed charge transfer pathways. Under illumination conditions, the excitons generated from the CdSeTe QD absorbers then diffuse to the interface between the donor and acceptor, where they can dissociate into free charge carriers, followed by the charge carriers diffusing to appropriate electrodes through the charge transfer layers under the built-in electric field.³⁰ Therefore, the photodetector device can effectively operate without an external bias and realizes the self-powered operation mode. The molecular structure of PF2 is also shown in Fig. 4b. The current–voltage (I-V) measurements of the as-fabricated control and target CdSeTe QD devices under standard sunlight irradiation were performed and the results are shown in Fig. 4c. It is found that a certain photocurrent can be generated at an applied bias of 0 V, suggesting the unique selfpowered capability of these photodetectors.³¹ The target CdSeTe QD photodetectors exhibited higher photocurrent densities and lower dark-current densities than the control device at 0 V bias. These results can be attributed to the higher concentration of photogenerated carriers and lower trap density, thus resulting in more efficient charge transport in the target CdSeTe QDs. This further reinforces the conclusion that the *in situ* fluorination treatment of the target QDs is a more effective photoinduced charge dynamic process than the control one, benefiting from the removal of the whole-body defects. The time-dependent photocurrent of the photodetectors was measured at a 0 V bias under standard sunlight illumination (Fig. 4d). The target CdSeTe QD-based photodetector exhibited two orders of magnitude higher photo-

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current density compared with the control devices (0.21 mA cm^{-2} vs. 0.11 mA cm⁻²). The on-off responsibility of the target photodetectors at 0 V bias was three orders higher than that of the control one. Moreover, as depicted in Fig. 4e, the external quantum efficiency (EQE) spectra of the target CdSeTe QDbased photodetectors are nearly twice those of the control device. The photodetectors exhibited a broad visible spectral photoresponse range, consistent with the wide absorption spectra of CdSeTe QDs. The photocurrent responsivities (R) and detectivity (D^*) of the control and target photodetectors under different wavelengths are demonstrated in Fig. 4f. Since R is proportional to the EQE value, the target photodetector displayed a higher R value with a peak value of 280 mA W^{-1} at 510 nm. The specific D^* value can also be calculated according to the equation $D^* = R/\sqrt{2qI_{\text{dark}}/S}$, where q is the electron charge, and Idark is the dark current, which is considered as the major noise in the detector.³² S is the effective area of the detector, and R is the photoresponsivity at the corresponding wavelength. The D^* of the target photodetector is remarkably higher than that of the control device. Table S2[†] presents a comparative analysis of the photoelectrical response performance of Cd-based photodetectors as reported in the literature. In addition, the time-dependent photocurrent of the photodetectors with the target and control CdSeTe QDs at 0 V bias was studied under single wavelength 606 nm laser illumination with three different light intensities (Fig. 4g and h). As the

laser light intensity increased from 24.6 to 60.8 mW cm⁻², the photocurrent of the control and target CdSeTe QD-based photodetectors significantly increased. However, the on-off responsibility ratio of the target QD-based photodetectors was much higher than that of the control one under all three light intensities. The high sensitivity under a negative bias indicates the increase of photoexcited carriers, which leads to the monotonic increase in the photocurrent at higher light intensity. We continued to measure the stability of the photodetector during continuous operation under standard sunlight illumination. As shown in Fig. S5a,[†] the change trends of J_{sc} in the CdSeTe QD-based photodetector with in situ fluorination treatment exhibit a relatively sluggish behavior, in contrast to that of the control device, which demonstrates a comparably quick degradation. This disparity is attributed to the CdSeTe QDs under in situ fluorination treatment to expedite charge extraction and mitigate charge recombination, ultimately enhancing the stability of the device. These results highlight the promising potential of the target QDs with enhanced charge transfer rates for optoelectronic applications. This high response ability at single visible wavelengths gives the photodetectors good potential for further optical communication and other optoelectronic applications.33

In addition, we continued to fabricate the CdSeTe QD-sensitized solar cells to validate the influence of *in situ* fluorination treatment further. The device structure of the high-efficiency QD-sensitized solar cell is shown in Fig. 5a. The immobilizing



Fig. 5 (a) Schematic illustration of the QD-based solar cell. (b) J-V curves, (c) corresponding EQE spectra, and integrated J_{sc} of the champion solar cells with control and target CdSeTe QD-based solar cells under the standard conditions (AM 1.5 G, 100 mW cm⁻²). The corresponding (d) recombination resistance (R_{rec}) on the applied bias and (e) Nyquist curves under a bias voltage of 0.80 V recovered from the EIS measurement of solar cells under dark conditions. (f) The open-circuit voltage decay curves of the control and target CdSeTe QD-based solar cells.

MPA-capped water-soluble CdSeTe QDs were obtained via the ligand-exchange process and then deposited on TiO₂ film electrodes via the capping ligand-induced self-assembly approach.^{11,34,35} The cross-section SEM images of the QD-sensitized TiO₂ photoanode utilized in solar cells, presented in Fig. S4b,[†] demonstrate a thickness of 10.3 µm. The *I*-V characteristics of the solar cells are plotted in Fig. 5b. The control CdSeTe OD-based devices produce a power conversion efficiency (PCE) of 11.12%, with photovoltaic parameters of a $V_{\rm oc}$ of 0.794 V, a $J_{\rm sc}$ of 20.53 mA cm⁻², and a fill factor (FF) of 0.682. Encouragingly, the target CdSeTe QD-based devices obtained a PCE of 12.79%, which resulted from the simultaneously enhanced PV parameters with a $V_{\rm oc}$ of 0.869 V, a $J_{\rm sc}$ of 20.97 mA cm⁻², and an FF of 0.702. Table S3[†] presents a comparative analysis of the photovoltaic performance of CdSeTe QD-sensitized solar cells as reported in recent literature. The results indicate that our device exhibits a higher $V_{\rm oc}$ and PCE than those reported before. The enhanced photovoltaic parameters, especially $V_{\rm oc}$, of the target QD-based devices are ascribed to the suppressed charge recombination, which results from the improved charge extraction and increased charge recombination of the QDs with in situ fluorination treatment. The external quantum efficiency (EQE) spectra and integrated J_{sc} presented in Fig. 5c confirm the creation of a more significant photocurrent in the target QD devices. The integrated J_{sc} value of the champion solar cell with the target QDs is 19.54 mA cm^{-2} , while that of the control QD-based device is 17.68 mA cm⁻², similar to that obtained from the *J*-*V* curves. Electrochemical impedance spectroscopy (EIS) measurements were performed to explore the recombination kinetics behavior of solar cells and understand the increase in performance better.³⁶ The value of charge recombination resistance (R_{rec}) is directly connected to the charge recombination process, and a bigger $R_{\rm rec}$ signifies a slower charge recombination rate. As demonstrated in Fig. 5d, the $R_{\rm rec}$ value of the samples with the target QDs is much higher than that of the control sample under different voltage biases. It can be shown that the recombination impedance of the device with the target CdSeTe QDs is substantially greater than that of the control CdSeTe QD-based devices, indicating that in situ fluorination treatment may effectively suppress charge recombination and accelerate charge extraction.³⁷ The Nyquist charts for several passivation procedures at a forward bias of -0.80 V are shown in Fig. 5e, and the fitting parameters are presented in Table S4.[†] It can be found that the extracted $R_{\rm rec}$ value of the target QD-based solar cells (6.9 Ω cm²) is higher than that of the control QD-based cells (5.1 Ω cm²). The notably higher $R_{\rm rec}$ implies that the charge recombination rate of the target QDbased device is significantly suppressed. The suppressed recombination will enhance photovoltaic performance, especially for $V_{\rm oc}$. Furthermore, to explore the influence of different passivation processes on the dynamics of photo-generated electrons, the open-circuit voltage decay (OCVD) was measured.³⁶ The V_{oc} transient attenuation of the solar cell was recorded after the light was turned off. Fig. 5f shows the decay curves of the solar cells. The devices with the target CdSeTe

QDs show a much slower V_{oc} decay rate compared with the sample with the control CdSeTe QDs.³⁷ Fig. S5b† displays the time-dependent PCE curve for a CdSeTe-based solar cell. Upon exposure to continuous standard sunlight for 200 min, the CdSeTe QD-based solar cells subjected to *in situ* fluorination retain 80% of their initial PCE. In contrast, the control device demonstrates a significant deterioration, maintaining only 68% of its initial intensity; therefore, the stability of solar cells retained with BF passivated CdSeTe QDs improved apparently. These results prove that the *in situ* fluorination treatment would harden the charge transport process by removing and passing whole-body defects toward chalcogenide QDs.

Conclusion

In summary, we have developed a simple and efficient in situ fluorination treatment to passivate the surface defects and thus enhance the charge transport process of the alloyed chalcogenide QDs. The BF molecule additive can decompose and release the F⁻ ions to remove oxide traps and dangling bonds of the whole-body nanocrystals and improve the stability of QDs. As a result, the fabricated self-powered photodetector devices demonstrated high photocurrent density, two orders of magnitude higher than those of the control ones. The responsivity and specific detectivity of the target self-powered photodetectors reached up to 280 mAW⁻¹ and 10⁸ Jones under illumination with a broad spectral range to near-infrared region, respectively. Furthermore, the target QD-based sensitized solar cells have improved PCE with three simultaneously enhanced photovoltaic parameters. This study provided a facile method to achieve the removal and passivation of defects under the in situ and real-time states to promote the charge transport process for QD-related optoelectronic technologies.

Data availability

All data required to understand the conclusions in the paper are presented in the main text and ESI.[†] Additional data related to this paper are available from the corresponding author upon request.

Conflicts of interest

The authors declare that they have no conflict of interest.

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