Energy level tuned indium arsenide colloidal quantum dot films for efficient photovoltaics

Jung Hoon Song¹, Hyekyoung Choi¹, Hien Thu Pham¹ & Sohee Jeong¹,²

We introduce indium arsenide colloidal quantum dot films for photovoltaic devices, fabricated by two-step surface modification. Native ligands and unwanted oxides on the surface are peeled off followed by passivating with incoming atomic or short ligands. The near-infrared-absorbing n-type indium arsenide colloidal quantum dot films can be tuned in energy-level positions up to 0.4 eV depending on the surface chemistry, and consequently, they boost collection efficiency when used in various emerging solar cells. As an example, we demonstrate p–n junction between n-type indium arsenide and p-type lead sulfide colloidal quantum dot layers, which leads to a favorable electronic band alignment and charge extraction from both colloidal quantum dot layers. A certified power conversion efficiency of 7.92% is achieved without additionally supporting carrier transport layers. This study provides richer materials to explore for high-efficiency emerging photovoltaics and will broaden research interest for various optoelectronic applications using the n-type covalent nanocrystal arrays.

¹ Nanomechanical Systems Research Division, Korea Institute of Machinery and Materials, Daejeon 34103, Korea. ² Department of Nanomechatronics, Korea University of Science and Technology (UST), Daejeon 34113, Korea. These authors contributed equally: Jung Hoon Song, Hyekyoung Choi, Hien Thu Pham. Correspondence and requests for materials should be addressed to S.J. (email: sjong@kimm.re.kr) or (email: sohee.jeong@gmail.com)
Colloidal quantum dots (CQDs) are attractive materials for next-generation photovoltaics (PVs) owing to their solution processability and size-dependent optical bandgaps that enable efficient absorption across a broad range of the solar spectrum. Since the first report of CQD solar cells, research interest has been focused on p-type CQD materials mainly including PbSe and PbS CQDs, and power conversion efficiency (PCE) has reached 12% as a result of proper device architecture and surface control of the CQDs. Most highly efficient CQD solar cells are based on a rectifying junction between a p-type CQD film and an n-type metal oxide semiconductor with a wide bandgap. These wide-bandgap metal oxide semiconductors absorb only a limited amount of solar energy and thus do not significantly contribute to solar energy harvesting.

So far, n-type CQD materials have been mainly achieved by chemically post-processing as-synthesized lead chalcogenides, for example, by hydrazine treatment, radical anion chemically post-processing as-synthesized lead chalcogenides, for example, by hydrazine treatment, radical anion chemically post-processing as-synthesized lead chalcogenides, for example, by hydrazine treatment, and optoelectronic devices. InAs has been shown to have inherent n-absorbing enhanced ambient stability. Furthermore, the near-infrared (NIR) solar cells. Built-in potentials and efficiency among devices using thin-film n-type InAs CQDs instead of wide-bandgap metal oxides.

Results and discussion
Two-step surface modification. Recently, we have reported a synthetic method of highly monodispersed, size-tunable InAs CQDs via growing InAs seeds through a continuous supply of amorphous pre-nucleation InAs clusters. In this study, synthesis of colloidal InAs CQDs was carried out following an improved version of a procedure from our previous report. We added diocyltiane (DOAm) as a co-surfactant as shown in Supplementary Fig. 1. Amines have been reported to significantly alter the reaction kinetics of III–V CQDs. Specifically, amines inhibit the decomposition of precursor (TMSI), and allow a wider tunability in sizes when preparing InP CQDs. Similarly, we found that InAs CQDs prepared with the addition of DOAm had a narrower size distribution (standard deviation less than 3.7%), measured by transmission electron microscopy (TEM) analysis and half width at half maximum (HWWM, 57 meV) for CQDs with 1.1 eV bandgap. Furthermore, the prepared InAs CQDs also offered a wider spectral range with an optical transition in the NIR between the first excited states from 700 to 1500 nm, compared with our previous results (Supplementary Figs. 2, 3).

As-synthesized InAs CQDs are usually passivated by insulating long-chain native ligands (NLs), which need to be removed for conductive CQD film formation. In addition, in situ formation of oxidizing layers on the surface of InAs CQDs at high temperature above 250 °C could decrease the film conductivity, which is a huge obstacle to high-performance optoelectronic devices. For efficient surface modification at oxidized surface layers in InAs CQDs, we established a two-step approach as shown in Figure 1. First, as-synthesized oleate-InAs CQDs were treated with NOBF4 to remove NLs, as well as oxides on the surface through solution-phase transfer (Fig. 2a); then, the CQD surface was reconstructed using five different ILs, including halide (Br–, Cl–, and I–) and short thiol ligands (3-mercaptopropionic acid (MPA) and 1,2-ethanedithiol (EDT)).

After the first step of peeling off NLs, the cubic zinc blend crystal structure of bulk InAs remained unchanged, as confirmed by the X-ray diffraction (XRD) patterns of InAs CQDs (Supplementary Fig. 4). On the basis of our observations, pristine oleate-capped InAs CQDs exhibit As2O3 formation, as evidenced by high-resolution As 3d X-ray photoelectron spectroscopy (XPS) spectra. After NOBF4 treatment, As2O3 was removed as confirmed by the disappearance of the shoulder peak at around 44 eV corresponding to As2O3, as shown in Fig. 2b. According to Dong A. and co-workers, NOBF4 could immediately etch the surfaces of metal oxide nanocrystals. The InAs CQDs obtained after the first step are termed naked CQDs. After NOBF4 treatment, the naked InAs CQDs were purified using toluene and well re-dispersed in N,N-dimethylformamide (DMF). The UV–vis–NIR spectra in Fig. 2c indicated that the InAs CQD peaks were slightly blue-shifted after NOBF4 treatment. Transmission electron microscopy (TEM) images of naked InAs CQDs with a diameter of 5.24 ± 0.35 nm confirmed the preservation of the particle size and size distribution after the first step (Supplementary Fig. 5a). Higher concentrations of NOBF4 solution, additionally, etch InAs CQD surfaces, as confirmed by blue-shifted absorption spectra (Supplementary Fig. 6).

BF4−, a very weak nucleophile, cannot bind to the CQD surface but instead forms an electrical double layer around the InAs CQD as a counter-ion for a stable colloid.
InAs CQDs from hexane to DMF, and Step-2 occurs in the same phase of the DMF solvent passivation of a naked surface by incoming ligands (ILs), including thiolate, carboxylate, and halides. In this process, Step-1 leads to the phase transfer of corresponding to the aliphatic $C–C$–$H$ stretching vibration at 2800–3000 cm$^{-1}$  and vinyl $=C–C–H$ stretching vibration at 3005 cm$^{-1}$ ascribed to the original oleate ligands before NOBF$_4$ treatment. No signal attributable to B (boron) was present, and a tiny amount of F (fluorine) remained from the BF$_4$ in the XPS spectra (Supplementary Table 1).

After peeling off the NLs, the CQD surfaces were reconstructed using a library of small and compact ligands, as shown in Fig. 3a. The reconstructed CQDs could be well re-dispersed in various hydrophilic solvents such as DMF and even highly polar formamide after purification with toluene. In order to analyze the surface compositions of reconstructed InAs CQDs, we performed XPS measurements by depositing InAs CQDs on indium tin oxide (ITO)/glass substrates. The XPS spectra of As 3d core level (Supplementary Fig. 7a) confirmed the absence of $\text{As}_2\text{O}_3$ on CQD surfaces after surface reconstruction with various ligands under ambient conditions. Supplementary Fig. 7b–f show the XPS spectra of Cl 2$p$, Br 3$p$, I 3$p$, and S 2$p$ for Cl-treated, Br-treated, I-treated, MPA-treated, and EDT-treated InAs CQDs, respectively. Depending on the binding affinities of the incoming ligands, the peaks in the In 3d spectra slightly shifted toward a higher binding energy compared with naked InAs CQDs (from 433.9 to 444.5 eV), as shown in Supplementary Fig. 8, which suggests that Cl, Br, I, and thiol ligands were bound directly to InAs CQD surfaces. Notably, no fluorine was detected in the XPS spectra after the second step.

Furthermore, we observed evidence of interactions of ligands with CQD surfaces through photoluminescence (PL) analysis, as shown in Fig. 3b. After peeling off NLs, the PL was strongly quenched owing to imperfect passivation, but the PL intensity was improved significantly after surface reconstruction. These results suggest that the surface defect density of InAs CQDs was reduced by surface passivation with the incoming ligands. After MPA treatment, the first excitonic peak in the absorbance spectrum is preserved (Supplementary Fig. 9). $\zeta$-potential measurements showed that, after the first step, InAs CQDs dispersed in DMF were positively charged, whereas InAs CQDs after surface reconstruction with MPA in DMF were negatively charged, providing further evidence for the interaction of ILs with naked CQD surfaces (Fig. 3c).

Figure 3d, e shows the UPS spectra in the secondary cutoff region (d) and valence band region (e) for InAs CQD films (100 nm thick). UPS is a viable tool for extracting the electronic states of materials and thus provides the Fermi level ($E_F$) and valence band maximum ($E_{V\text{B}}$) positions estimated by adding the optical bandgap of 1.1 eV determined from absorption spectra. Figure 3f shows the energy levels of InAs CQDs capped with different ligands. The $E_{V\text{B}}$ levels of all InAs CQD films are located close to their $E_C$ levels, and thus, the InAs CQD films exhibit an $n$-type behavior regardless of the ligands.
The energy levels of InAs CQDs shifted by up to 0.4 eV depending on the chemical ligands, which can be attributed to the contribution of the total sum of ligand/CQD interface dipole moment and the intrinsic dipole moment of the ligand30. Halide-capped InAs CQDs showed deeper band energy levels because halide ligands cause large interface dipole moments to form between the surface atom of the CQD and the binding group of the ligand, without the ligand having an intrinsic dipole moment; this observation is similar to that for halide-capped PbS CQDs27. On the other hand, thiol-capped InAs CQDs showed a small shift in their energy levels possibly due to the offset direction of the ligand/CQD interface dipole moment and the intrinsic dipole moment of the ligand because of thiol ligands with their own dipole moment. Among the halide-capped InAs CQDs, the CQDs capped with I− showed the deepest electronic energy-level position. We consider that energy levels can also be influenced by the degree of passivation because trap filling can affect the increase in n-type doping, which leads to a shift of $E_F$ in InAs CQD films31. Indeed, according to the XPS quantitative analysis (Supplementary Table 1) of atomic ratios of halide to In, the ratio of I to As is higher than those of Br and Cl.

**Charge transport in InAs CQDs.** Through two-step surface modification, bulky organic ligands were replaced with small and compact ligands such as Br and MPA, which facilitate the electronic communication among individual InAs CQDs. The electron mobility of InAs CQD arrays was conveniently measured from the transfer characteristics of a FET. Briefly, a densely packed 50-nm-thick InAs CQD film was prepared on pre-patterned interdigitated electrodes (Fig. 4a) on a 300-nm-thick silicon dioxide (SiO$_2$/silicon (Si) substrate by spin coating without annealing (please refer to the Methods section for details). Figure 4b depicts the representative transfer curves, i.e., drain current ($I_{DS}$) versus gate voltage ($V_G$), of the FET comprising MPA-passivated InAs CQDs, obtained by sweeping $V_G$ from −40 to +40 V at $V_{DS}$ of 40 V, which clearly exhibit an n-type behavior. A positive gate bias causes the depth of the accumulation mode to be exceeded since the concentration of the electron majority carriers contributes to the increase in $I_{DS}$. Such n-type transport behavior of InAs CQDs is in good agreement with our UPS results (Fig. 3f).

As shown in Supplementary Fig. 10, naked InAs CQDs before the second-step treatment exhibited an electron mobility ($\mu_{lin}$) corresponding to the linear regime of FET operation (measured at $V_{DS} = 20$ V), which was determined to be $1.2 \times 10^{-4}$ cm$^2$ V$^{-1}$ s$^{-1}$. The saturation electron mobility ($\mu_{sat}$) of $1.3 \times 10^{-4}$ cm$^2$ V$^{-1}$ s$^{-1}$ measured at $V_{DS} = 40$ V was slightly higher than $\mu_{lin}$. On the other hand, FETs assembled from InAs CQD-capped Br and MPA ligands (denoted by InAs CQD–Br and InAs CQD–MPA) showed $\mu_{lin}$ of $1.4 \times 10^{-3}$ and $2.0 \times 10^{-3}$ cm$^2$ V$^{-1}$ s$^{-1}$ at $V_{DS} = 10$ V and $\mu_{sat}$ of $1.7 \times 10^{-3}$ and $2.5 \times 10^{-3}$ cm$^2$ V$^{-1}$ s$^{-1}$, respectively, as summarized in Table 1. Compared with naked InAs CQDs, the channel currents for InAs CQD–Br and InAs CQD–MPA were one order of magnitude higher. We consider that the lower conductivity of naked InAs CQDs originates from the presence of surface trap states created by unpassivated metal cation sites after
ligand removal by NOBF₄. The trap states located in the bandgap often behave as a local electrostatic barrier around the CQD. After surface reconstruction with small and compact ligands such as Br and MPA, the surface traps were suppressed, as evidenced by PL analysis (Fig. 3b), and resulted in higher carrier mobility. We also obtained smaller subthreshold swing values of 12.9 and 9.7 V decade⁻¹ for InAs CQD–Br and InAs CQD–MPA, which is a critical metric describing how well a device turns on and off. Furthermore, we extracted the carrier concentrations of the InAs CQDs that depend on the ligands by performing the FET device analysis as shown in Table 1. The InAs–MPA CQD film contains more electron carriers than that contained in case of the InAs–Br.

As shown in Fig. 4c, MPA-capped InAs CQD FETs demonstrated excellent air stability, maintaining the original n-type characteristic even after air exposure over 22 days. It is noted that n-type InAs CQD films are resistant to atmospheric exposure at the device level because air stability is strongly affected by adsorption of molecular oxygen species. In n-type CdSe nanocrystals using a radical anion as the n-type dopant, the decay of the 1Sₓ–1Pₓ transition occurs within 30 min to 24 h at room temperature. Also, the doping polarity of n-type lead chalcogenides tends to switch from n-type to p-type upon air exposure because molecular oxygen acts as the p-type dopant. Unlike those unstable n-type CQDs reported so far, our n-type InAs CQD films, which are resistant to air exposure, can be used for various applications without being restricted by processing environments.

Additionally, we extracted the trap density, effective lifetime, and diffusion length using the PL-based interparticle coupling method that was originally developed by Zhitomirsky et al. and successfully implemented in PbS CQD layers with varied coupling efficiency. The interparticle coupling among the assembled InAs CQDs was monitored as follows (please refer to Supplementary Note 1 for details). Two InAs CQDs with 1.5 eV (donor CQDs) and 1.1 eV (acceptor CQDs) bandgaps were prepared, and the pristine oleate ligands were replaced with Br and MPA, respectively, using the two-step surface modification method that has been proposed in this paper, as depicted in Supplementary Fig. 11. Then, we prepared InAs CQD mixture solutions with two sizes of InAs CQDs in various composition ratios (0%, 0.1%, 1%, 10%, and 100%). We measured the PL after the deposition of the InAs CQD mixture on glass, as depicted in Supplementary Fig. 12. The ratio of integrated PL emissions from the large bandgap (donor) CQDs to the small bandgap (acceptor) CQDs was expected to decrease upon enhanced interparticle coupling (Supplementary Fig. 13). The characteristics of the InAs CQD layers obtained using the 3D diffusion method are described in Supplementary Table 2. The mobility of the InAs CQD matched well with that obtained from the FET analysis; InAs–MPA exhibited a long diffusion length because of their low trap density.

**Physical parameters of the InAs CQD layer.** The InAs CQDs show n-type doping polarity and thus we search for p-type materials for the p–n junction solar cell devices. We consider the PbS CQDs to be the p-type layer. Due to the InAs CQDs with high n-type doping concentration from 2.77 × 10¹⁶ to 2.8 × 10¹⁷ cm⁻³ (Table 2), we need to use lightly doped p-type layers for an increase in depletion width in the device. The PbS–I CQD layer in this paper exhibits n-type properties based on the FET characteristics that have been measured in an N₂-filled glove box, as depicted in Supplementary Fig. 14 and Supplementary Table 3. However, the doping polarity switches from n-type to p-type when this device is exposed to air for an hour because oxygen can act as a p-type dopant. We used the air-exposed PbS–I CQD layer as a lightly p-type doped material with a carrier concentration of approximately 10¹⁵ cm⁻³. We can individually control optical bandgaps and the energy-level positions in the CQD–CQD p–n junction solar cells as shown in Fig. 5.

Based on the carrier concentrations of InAs–MPA (1.28 × 10¹⁷ cm⁻³) and PbS–I (8.11 × 10¹⁵ cm⁻³) CQDs, the device structure was designed such that the PbS CQDs (1.3 eV...
bandgap as shown in Supplementary Fig. 15) serve as the main absorber for a high efficiency of CQD solar cell devices according to the Shockley–Queisser limit\(^6\). We further used InAs CQDs (1.1 eV bandgap) with an aim to extend the absorption range beyond the bandgap of PbS CQDs. Among the large-sized InAs CQDs having a bandgap of less than 1.3 eV, which is the PbS CQD bandgap, the 1.1 eV InAs CQDs are observed to be the most reproducibly synthesized and exhibit a narrow size distribution (standard deviation less than 4%, measured using TEM analysis). Furthermore, the InAs CQDs that are too large lead to significant \(V_{OC}\) losses, as apprehended.

Representatively, we fabricated the Br-capped and MPA-capped InAs CQD films of 50 nm in thickness (Fig. 5b) under the same conditions as those for the film preparation for FET devices. Then we deposited the PbS CQD films (PbS–I and PbS–EDT) of 200 nm in thickness on the InAs–Br and InAs–MPA CQD films (Supplementary Fig. 16)\(^{11,13,34}\). All film deposition procedures were carried out under atmospheric conditions. Furthermore, we expect no contribution from the space charge region that is formed between the PbS CQDs (PbS–I and PbS–EDT). Therefore, it is necessary to confirm whether the carrier is extracted by the space charge region formed by PbS–I and PbS–EDT. To clarify this assumption, No-InAs devices were fabricated.

In a planar \(p–n\) junction CQD solar cell, the photo-induced carriers are transported by the drift in the space charge region and by the diffusion in the quasi-neutral region. Therefore, we performed capacitance–voltage (\(C–V\)) measurements to study the carrier concentration and depletion width in the InAs/PbS CQD solar cell devices. Supplementary Fig. 17 depicts the Mott–Schottky (\(C–2–V\))\(^{40}\) curves of the InAs/PbS CQD solar cell. Two distinct slopes can be observed, among which one corresponds to the InAs CQD layer (in high reverse bias) and the other corresponds to a PbS–I layer (in low forward bias)\(^{41}\) because the InAs CQDs exhibit a much higher carrier concentration than that exhibited by the PbS CQDs, as confirmed by FET analysis. The built-in potentials (\(V_{bi}\)) were determined from the curves and revealed that a higher \(V_{bi}\) was obtained for the InAs–MPA-based CQD solar cell as compared to that obtained in case of the InAs–Br-based CQD solar cell. A large \(V_{bi}\) in the solar cell will cause a strong built-in electric field, which will facilitate charge carrier extraction (Supplementary Table 4). The depletion width (\(W_{D}\)) within the InAs/PbS CQD solar cells was calculated using Mott–Schottky analysis, which is commonly used in CQD devices\(^{42}\), and the details are presented in the experimental section. Supplementary Fig. 17b and 17c depict the calculated \(W_{D}\) of the InAs CQD and PbS CQD layers over the applied voltage in the InAs–Br/PbS and InAs–MPA/CQD solar cell devices. Under the short-circuit condition (voltage = 0 V), the \(W_{D}\) for the InAs–Br and InAs–MPA CQD layers was calculated to be around 40 and 60 nm, whereas the \(W_{D}\) were around 350 and 450 nm for the PbS CQD layers that were deposited on InAs–Br and InAs–MPA, respectively.

| Materials     | Majority carrier | Mobility (cm\(^2\) V\(^{-1}\) s\(^{-1}\)) | Carrier concentration (cm\(^{-2}\)) | Trap density (cm\(^{-2}\)) | Diffusion length (nm) |
|---------------|------------------|-----------------------------------------|------------------------------------|----------------------------|------------------------|
| InAs–Br       | Electron         | 1.55 × 10\(^{-3}\) (FET)\(^a\) 1.34 × 10\(^{-3}\) (3D PL)\(^b\) | 2.77 × 10\(^{10}\) (FET)\(^c\) 4.65 × 10\(^{15}\) (C–V)\(^{-2}\) | 3.32 × 10\(^{17}\) (3D PL)\(^d\) | 30 (3D PL)\(^e\)      |
| InAs–MPA      | Electron         | 2.25 × 10\(^{-3}\) (FET)\(^a\) 1.78 × 10\(^{-3}\) (3D PL)\(^b\) | 8.11 × 10\(^{15}\) (FET)\(^c\) 1.27 × 10\(^{17}\) (C–V)\(^{-2}\) | 1.24 × 10\(^{17}\) (3D PL)\(^d\) | 60 (3D PL)\(^e\)      |
| PbS–I         | Hole\(^f\)       | 2.74 × 10\(^{-3}\) (FET)\(^a\) (air) | 5.30 × 10\(^{15}\) (C–V)\(^{-2}\) (InAs–Br) | -                          | -                      |
|               |                  |                                        | 5.18 × 10\(^{15}\) (C–V)\(^{-2}\) (InAs–MPA) | -                          | -                      |

\(^{a}\)Supplementary Fig. 14 and Supplementary Table 3

\(^{b}\)Mean value of linear and saturation field-effect mobility of Table 1

\(^{c}\)Supplementary Figs. 11, 12, 13, and Supplementary Table 2

\(^{d}\)Supplementary Fig. 17 and Supplementary Table 4

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Table 3 The best PV performance of the various CQD solar cells

| Device         | \(V_{OC}\) (V) | PCE (%) | FF (%) | \(J_{SC}\) (mA cm\(^{-2}\)) | \(J_{SC}\# EQE\) (mA cm\(^{-2}\)) |
|----------------|----------------|---------|--------|---------------------------|----------------------------------|
| No-InAs        | 0.21           | 1.11    | 39.57  | 13.13                      | 8.39                             |
| InAs–Br        | 0.46           | 4.61    | 56.54  | 17.78                      | 18.61                            |
| InAs–MPA       | 0.54           | 7.23    | 63.73  | 20.92                      | 20.73                            |

Figure 5c shows the current–voltage (\(I–V\)) characteristics of the PVs measured under simulated AM 1.5 G illumination (please refer to Supplementary Figs. 18–20 and Supplementary Table 5 for detailed measurement conditions and various reliability tests). PV performance metrics such as open-circuit voltage (\(V_{OC}\)), short-circuit current density (\(J_{SC}\)), fill factor (FF), and PCE are summarized in Table 3. Interestingly, the PbS CQD PV device functioned even without an \(n\)-layer such as ZnO, which might be caused by the homojunction formation between PbS–I CQDs and PbS–EDT CQDs. However, the PCE was lower than 2% because of a low built-in potential, which led to a low \(V_{OC}\) of 0.21 V. All PV parameters were significantly improved after InAs CQDs were incorporated (Table 3). The PCEs of CQD PVs fabricated using InAs–Br and InAs–MPA CQDs were 4.61% and 7.23%, respectively. In the InAs–MPA-based PVs, the higher \(V_{OC}\) is attributed to an increased built-in potential (Supplementary Fig. 17 and Supplementary Table 4) due to an upward shift of the energy bands and an increase of carrier concentration (Table 1 and Supplementary Table 4). Simultaneously, the higher \(J_{SC}\) and FF values originate from the enhanced charge carrier transport in InAs–MPA CQDs, as demonstrated by the FET (Fig. 4) and PL-based interparticle coupling measurements (Supplementary Table 2).

In addition, the external quantum efficiency (EQE) of InAs–MPA-based PVs mainly increased at short wavelength with a short penetration length compared with InAs–Br-based PVs, as shown in Fig. 5d. These results are in good agreement with the InAs CQD film thickness dependence analyses (Supplementary Figs. 21, 22 and Supplementary Table 6), the depletion width (Supplementary Fig. 17 and Supplementary Table 4), and the diffusion length (Supplementary Figs. 11–13, and Supplementary Table 2). The carrier extraction efficiency of InAs–MPA CQD films decreases at a relatively thick thickness than InAs–Br CQD films (Supplementary Fig. 21 and Supplementary Table 6). The PbS CQD PVs without the InAs CQD layers exhibited a significantly low EQE over the entire wavelength because the formation of a depletion width to extract the carriers was considerably weak, which indicated that carrier extraction in the CQD solar cells occurred by the drift mechanism. The depletion width of the InAs/PbS CQD solar cell device was nearly obtained in the PbS CQD films, which limited the increase of the thickness of the InAs CQD film. The best method to overcome this limitation is to increase the...
of CQD solar cells, especially device structure in both forward (\(J_{\text{InAs}}\)) and backward (\(J_{\text{Br}}\)), which corresponds to the EQE of this solar cell obviously increased near 1000 nm (Supplementary Fig. 23c). The EQE peak at 710 nm in the absorption spectrum, which is distinguishable from that of InAs CQDs (Supplementary Fig. 23a, b). The EQE of this solar cell obviously increased near 1000 nm (Supplementary Fig. 23c), which corresponds to the first excitonic peak in the absorption spectrum of InAs CQDs. This result suggests that the NIR photoactive \(n\)-type InAs CQDs can boost the collection efficiency of CQD solar cells, especially \(J_{\text{SC}}\), by extending the absorption range beyond the PbS CQD bandgap.

A statistical analysis of PCE was carried out for InAs-MPA, InAs-Br, and no InAs CQD PV devices, with six devices for each device structure in both forward (\(J_{\text{SC}} \rightarrow V_{\text{OC}}\)) and backward (\(V_{\text{OC}} \rightarrow J_{\text{SC}}\)) directions, as shown in Fig. 5e. The average PCE and its standard deviation were recorded as PCE of 6.78 ± 0.27% for InAs–MPA-based CQD PV devices. The stability of CQD solar cell performance (Fig. 5f) was monitored while storing the solar cell in air without any encapsulation. Similar to the FET stability, we found that the solar cell device performance was stable for over 13 days. To support our results, we sent our best device to Newport Corporation for independent certification. Figure 5g, h shows Newport certification results for a certified PCE of 7.92% without additional supporting transport layers, which is the record efficiency among devices using thin-film \(n\)-type InAs CQDs instead of wide-bandgap metal oxides. The accreditation certificate is shown in Supplementary Fig. 24. Further efficiency increases are expected by enhancing the conductivity in the \(n\)-layer, optimizing band alignment via size and surface modification of both \(n\)-type and \(p\)-type CQDs, and introducing additional supporting transport layers or blocking layers.

In conclusion, we report the use of InAs CQD thin films in photovoltaics by overcoming their limitations with respect to optoelectronic devices through fabrication of defect-controlled conductive InAs CQD assemblies. Specifically, we developed a facile and efficient surface modification method via two-step solution processing. First, InAs CQD surfaces were treated with NOBF4 to remove native oleate ligands, as well as metal oxide formed on the surface during or after the synthesis before being subjected to surface reconstruction by various ligands in the second step. Regardless of the capping ligands, the InAs CQD films showed \(n\)-type characteristics with from \(10^{16}\) to \(10^{17}\) cm\(^{-3}\) doping concentration and also exhibited noticeably improved air stability in FET devices. Most importantly, the electronic energy diffusion length by eliminating the deep-level trap states or by controlling the doping concentration of the InAs CQD layers.

Additionally, the EQE onsets were redshifted with increasing InAs CQD film thickness. In order to monitor the contribution from InAs CQDs to the solar cells, we used PbS CQDs with the first excitonic peak at 710 nm in the absorption spectrum, which is distinguishable from that of InAs CQDs (Supplementary Fig. 23a, b). The EQE of this solar cell obviously increased near 1000 nm (Supplementary Fig. 23c), which corresponds to the first excitonic peak in the absorption spectrum of InAs CQDs. This result suggests that the NIR photoactive \(n\)-type InAs CQDs can boost the collection efficiency of CQD solar cells, especially \(J_{\text{SC}}\), by extending the absorption range beyond the PbS CQD bandgap.

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levels were tuned by over 0.4 eV depending on the surface ligands. Accordingly, the n-type InAs QDs formed rectifying junctions with p-type PbS QDs, thus forming p-n junction devices. Our best device showed a certified PCE of 7.92%. These energy-level-tuned InAs QDs represent an emerging class of stable n-type materials that enable simultaneous near-infrared absorption and energy-level tuning.

**Methods**

**CQD synthesis.** Indium acetate (InOAc, 99.99%), oleic acid (OA, ≥ 99%), dioc-tanoyl (DOA, ≥ 97%), 1-octadecene (ODE, 90%), nitrosyl tetrafluoroborate (NOBF₄, ≥ 98%), 3-mercaptopropionic acid (MPA, ≥ 99%), dimethylformamide (DMF), toluene (99%), ammonium iodide (NH₄I, ≥ 99%), ammonium chloride (NH₄Cl, 99.99%), and ammonium bromide (NH₄Br, ≥ 99%) are purchased from Sigma-Aldrich Chemical Co. and used without further purification. Triethylsilanesulfurine (TMSi), 99% is purchased from JSI Silicone and is distilled before use. All solvents, including hexane, butanol, anhydrous n-dimethylformamide (DMF), formamide (FA), anhydrous octane, tolouene, and tetrachloroethylene (TCE) are purchased from Sigma-Aldrich Chemical Co.

Collodial InAs CQD via continuous injection synthesis: Indium arsenide quantum dots (InAs CQDs) have been prepared following those that have been described earlier with some modification. Typically, InAs CQDs have been synthesized by using continuous injection of cluster solution into the seed solutions, and the details are as follows:

1. Synthesis of InAs seeds. The InAs solution: In total, 0.29 g (1 mmol) of InOAc is mixed with 0.85 g (3 mmol) of oleic acid in 5 mL of ODE in a 50 mL round-bottom flask degassed at 120 °C for 2 h under vacuum. The As solution: In total, 0.14 g (0.5 mmol) of TMSiAs, As is mixed with 0.36 g (1.5 mmol) of DOA degassed in 1 mL of degassed ODE solution, and the solution is kept at 80 °C for 1 h until the no-color transparent solution turns to brown, in a glove box. The As solution is rapidly injected into InOAc solution at 300 °C. The temperature is dropped to 287 °C and continued for 20 min.

2. Synthesis of amorphous InAs nanoclusters. The InOAc solution: In total, 1.74 g (6 mmol) of InOAc is mixed with 5.10 g (18 mmol) of oleic acid in 30 mL of ODE in a 50 mL round-bottom flask degassed at 120 °C for 2 h and heated at 120 °C in vacuum. The As solution: In total, 0.84 g (3 mmol) of TMSiAs, As is mixed with 2.17 g (9 mmol) of DOA dissolved in 6 mL of degassed ODE solution, and the solution is kept at 60 °C for 1 h, in a glove box. The As solution is injected into InOAc solution at room temperature with constant stirring for 10 min.

3. Synthesis of InAs quantum dots. InAs nanoclusters solution prepared has been loaded in a syringe (diameter 20 mm) and injected into InAs QCD seed solution at growth temperature (300 °C) at desired rates (0.05 mmol indium/min, arsenic equivalent). Final sizes are depending on the injection rate, amount of precursor added, and also InAs seeds (size and size distribution).

4. Purification. Synthesized InAs CQD solution is divided into smaller volume fractions of 10 mL each. To each fraction, 40 mL of butanol is added and centrifuged at 6000g rpm for 5 min. The precipitate is dispersed in 10 mL of hexane, and subsequently, 15 mL of butanol is added and centrifuged at 4000g rpm for 5 min. The precipitate is left of the supernatant is collected in a fresh tube, and 20 mL of butanol was added and centrifuged at 6000g for 5 min. This step effectively removes any by-product (In₂O₃) formed. The precipitate is dispersed in 10 mL of hexane. An aliquot of 35 mL of butanol is added and centrifuged at 6000g for 5 min and this step is repeated twice more. Finally, the precipitate is dried under vacuum for overnight before it disperses in octane for further steps.

Step 1: preparation of naked InAs CQDs–solution treatment with NOBF₄:

Typically, a total of 200 mg of InAs CQDs are dispersed in 4 mL of octane followed by the addition of a 10 mL solution of NOBF₄ (0.02 M in DMF). CQDs completely precipitated in octane phase within 2 min. The colorless octane phase is discarded by the addition of a 10-mL solution of NOBF₄ (0.02 M in DMF). CQDs completely precipitated in octane phase within 2 min. The precipitate is washed by the addition of a 10-mL solution of octane phase and centrifuged at 6000g rpm for 4 min. The final product is dried in vacuum for overnight before being dispersed in DMF for further use.

Step 2: surface reconstruction in InAs CQD: For halide ligand reconstruction, the halide salts such as ammonium chloride (NH₄Cl), ammonium bromide (NH₄Br), or ammonium iodide (NH₄I) are purchased from Sigma-Aldrich and used as received without further purification. The following stock solution: 0.1% halide salts in anhydrous MeOH are tested in this work. In brief, 10 mL of naked InAs CQD (10 mg mL⁻¹) in DMF is mixed with 5 mL of halide salts in MeOH. After stirring for less than 3 min, we obtain a stable CQD solution in mixture solvent. Tolouene is employed to precipitate the CQD, which is followed by centrifugation at 6000g rpm for 4 min. The final product is dried in vacuum for 2 h before being recovered in DMF without affecting the colloidal stability.

For thiol ligand reconstruction, 0.01 vol% of MPA in anhydrous DMF is prepared. In brief, 10 mL of naked InAs CQD (10 mg mL⁻¹) in DMF is mixed with 5 mL of MPA solvent. After stirring for less than 3 min, toluene is used to precipitate the QCD, which is followed by centrifugation at 6000g rpm for 4 min. The final product is dried in vacuum for 2 h before being recovered in DMF without affecting the colloidal stability. InAs CQD prepared with MPA are stable in FA for 1 h.

**FET fabrication.** InAs CQD FET is fabricated by spin-coating a layer of InAs CQD ink onto pre-patterned interdigital electrodes (IDEs) on SiO₂/Si substrates. Commercial silicon wafers (a highly doped p type using boron) with 300 nm of thermally oxidized SiO₂ are used as substrates and gate dielectrics. Au and Cr are used as the source and drain electrodes. The IDE arrays had a 300 μm wide and 5 μm spacing between the electrodes. Typically, 150 mg mL⁻¹ InAs CQD ink in DMF is deposited on the substrate (spread: 4000 rpm, 30 s) and the film thickness is around 35 nm. After deposition, the QCD films are dried in vacuum for 1 h.

**Solar cell fabrication.** Heterojunction CQD solar cells (ITO/InAs/PbS/Au) are fabricated, and the details are as follows. PbS–InCQD ink with around 3 nm in diameter is fabricated following a published recipe. InAs CQD layers and PbS CQD layers. The capacitance of the InAs/PbS CQD solar cells was determined using the impedance analysis to determine the carrier concentration and depletion width of InAs CQD layers and PbS CQD layers. The capacitance of the InAs/PbS CQD solar cells was measured by a spectral measurement system (IQE-200, Newport).

**Characterizations.** For X-ray diffraction (XRD) analyses, we used PanAnalytical X'Pert PRO diffractrometer (Cu source). Transmission electron microscopy (TEM) images were acquired using Tecnai F30 Super-Twin TEM microscopy operated at 300 keV. The InAs QCDs dispersed in hexane and InAs CQD ink was dispersed in DMF and drop casted on a carbon-coated Cu grid. Absorption spectra were collected using an UV-visible spectrometer (Shimadzu, UV3600). X-ray photoemission spectroscopy (XPS) and ultraviolet photoelectron spectroscopy (UPS) were performed using Sigma Probe Model (Thermo VG Scientific). Samples were prepared via spin coating on ITO glass substrate as a conducting substrate to avoid the charging effect. The thickness of the QCD film was over 100 nm and kept in an inert condition until loading to the analysis chamber. The Fermi level was determined as used as an energy reference point, which was calibrated through clean Au (111). Cross-sectional field-emission scanning electron microscopy (FE-SEM) was conducted on a JSM-7500F microscope (JEOL, Akishima, Tokyo, Japan) to measure the thickness of the QCD layer.

The transfer characteristics of QCD FET devices were conducted on an Agilent B2902A (4 ½ Digits) (Hewlett-Packard) (HP) and Agilent B2902A (4 ½ Digits) (Hewlett-Packard) (HP) (Hewlett-Packard) (HP). The transfer characteristics were measured by scanning gate voltage (V₉) from −40 to 40 V and drain–source voltage (V DS) of 40 V. We performed the impedance analysis to determine the carrier concentration and depletion width of InAs CQD layers and PbS CQD layers. The capacitance of the InAs/PbS CQD solar cells was obtained by varying the applied bias. We used an impedance analyzer (Agilent, 4291A) with the signal of 1000 Hz frequency and 50 mV amplitude for C–V measurement under the C–Rₒ model. The current–voltage was extracted from the C–V curve and the depletion width was calculated using diode approximation analysis. Current density–voltage (J–V) characteristics were measured by using a solar simulator (SolAStar Class-94043A, Newport) at 100 mW cm⁻² from a 450-W Xe short arc lamp filtered by an AM 1.5 G filter. Voltage swept from 0.7 to −0.1 V (backward sweep direction) and from −0.1 to 0.7 V (forward sweep direction) with the speed of 0.01 V per second and a dwell time of 500 ms (0.02 s V⁻¹). A mask of 0.047 cm² was attached to the device to clearly define the active area. The measurement of J–V characteristics was taken after light soaking for 2 min under 1 sun. The EQE was measured by a spectral measurement system (QE-200, Newport).

**Data availability**

The data that support the plots within this paper and other findings of this study are available from the corresponding author upon request.
References

1. Carey, G. H. et al. Colloidal quantum dot solar cells. Chem. Rev. 115, 12732–12763 (2015).
2. Yuan, M., Liu, M., Stovergert, E. H. Colloidal quantum dot solids for solution-processed solar cells. Nat. Energy 1, 16016 (2016).
3. Bi, Y. et al. Infrared solution-processed quantum dot solar cells reaching external quantum efficiency of 80% at 1.35 μm and 6% in excess of 34 mA cm⁻². Adv. Mater. 30, 1704928 (2018).
4. Fan, Z. J. et al. Halide re-shelled quantum dot inks for infrared photovoltaics. ACS Appl. Mater. Interfaces. 9, 37356–37354 (2017).
5. Choi, H., Ko, J.-H., Kim, Y.-H. & Jeong, S. Steric hindrance-driven shape transition in PbS quantum dots: understanding size-dependent stability. J. Am. Chem. Soc. 135, 5278–5281 (2013).
6. Mcdonald, S. A. et al. Solution-processed PbS quantum dot infrared photodetectors and photovoltaics. Nat. Mater. 4, 138–142 (2005).
7. Pattanyus-Abraham, A. G. et al. Depleted-heterojunction colloidal quantum dot solar cells. ACS Nano 4, 3374–3380 (2010).
8. Talapin, D. V. & Murray, C. B. PbSe nanocrystal solids for p- and n-channel thin film field-effect transistors. Science 310, 86–89 (2005).
9. Shim, M. & Guyot-Sionnest, P. N-type colloidal semiconductor nanocrystals. Nature 407, 981–983 (2000).
10. Woo, J. Y. et al. Ultrastable PbSe nanocrystal quantum dots via in situ formation of atomically thin halide adlayers on PbSe (100). J. Am. Chem. Soc. 136, 8883–8889 (2014).
11. Zhitomirsky, D. et al. N-type colloidal-quantum-dot solids for photovoltaics. ACS Nano 6, 1700301 (2017).
12. Balazs, D. M. et al. Lead-chalcogenide colloidal-quantum-dot solids: novel assembly methods, electronic structure control, and application prospects. Adv. Mater. 30, 1800802 (2018).
13. Kroupa, D. M. et al. Tuning colloidal quantum dot band edge positions through solution-phase surface chemistry modification. Nat. Commun. 8, 15257 (2017).
14. Yaacobi-Gross, N. et al. Combining ligand-induced quantum-confinement stark effect with type II heterojunction bilayer structure in CdTe and CdSe nanocrystal-based solar cells. ACS Nano 6, 3128–3133 (2012).
15. Shockley, W. & Queisser, H. J. Detailed balance limit of efficiency of p-n junction solar cells. J. Appl. Phys. 32, 510–519 (1961).
16. Rydberg, E. H. & Williams, R. H. in Metal-Semiconductor Contacts. (Clarendon, Oxford, 1988).
17. Ning, Z. et al. Graded doping for enhanced colloidal quantum dot photovoltaics. Adv. Mater. 25, 1719–1723 (2013).
18. Willis, S. M., Cheng, C., Assender, H. E. & Watt, A. A. R. The transitional heterojunction behavior of PbS/ZnO colloidal quantum dot solar cells. Nano Lett. 12, 1522–1526 (2012).

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Author contributions

J.H.S., H.C. and H.T.P. contributed equally to this work. H.C. and H.T.P. synthesized and characterized their photovoltaic performance, and physical properties of InAs nanocrystals. J.H.S., H.C., H.T.P. and S.J. discussed the results and co-wrote the manuscript.

Additional information

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