Quantum-assisted photoelectric gain effects in perovskite solar cells

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Abstract

Further boosting the power conversion efficiencies (PCEs) of perovskite solar cells (PSCs) without excessively increasing production expenses is critical for practical applications. Here, we introduce silicon quantum dots (SiQDs) to enable perovskites to harvest additional sunlight without changing PSC processes. These SiQDs can convert shorter wavelength excitation light (300–530 nm) into visible region light and reflect longer wavelength perovskite-unabsorbed visible light (550–800 nm), leading to broadband visible light absorption enhancement in PSCs. As a result, the SiQD-based photocurrent gain can improve the external quantum efficiencies of PSCs over a wide wavelength range of 360–760 nm, yielding relatively enhanced short-circuit current density (+1.66 mA/cm²) and PCE (+1.4%). Surprisingly, even the PSC with a low-purity perovskite layer shows an ultrahigh PCE improvement of 5.6%. Our findings demonstrate QD-assisted effects based on earth-abundant and environmentally friendly silicon, leading to effective optical management that remarkably promotes the performance of PSCs and enables the balance of costs to be substantially addressed.

Introduction

The past decade has witnessed amazing advances in organic–inorganic perovskite solar cells (PSCs), with the power conversion efficiency (PCE) drastically increasing from 3.8% to more than 25%1–10. The extraordinary PCEs have already exceeded those of multocrystalline Si solar cells and are not far from the single crystal values10. Even so, crystalline Si modules still dominate the current photovoltaic industry because their mature technology has led to a continuous decrease in costs11. To date, the major costs for solar panel manufacturing are area-related balance-of-system costs; thus, increasing the PCE is a key direction for reducing the levelized costs of solar energy11–15. Unfortunately, the PCEs of single-junction solar cells are limited to the Shockley-Queisser limit due to the thermal relaxation of photocarriers with energies in excess of the bandgap energies (known as thermalization loss)11–17. Multijunction solar cells offer an effective solution to broaden the sunlight-sensitive region by utilizing multiple light harvesters with various absorption bands. A recent study on the perovskite (wide bandgap)/Si (narrow bandgap) tandem solar cell realized a certified PCE of 28.0%10, which once again places perovskite photovoltaics on the list of commercial candidates. However, to fabricate efficient perovskite/Si tandem photovoltaics on the list of commercial candidates, the upper perovskite crystals need to be well tailored to produce ultratransparent properties at long wavelengths while achieving excellent stability18. Therefore, the development of a simple method through which sunlight can be maximally absorbed to boost the performance of PSCs without changing the crystallization
of the perovskite and without excessive material costs remains a challenge.

Silicon quantum dots (SiQDs) are fascinating materials for photovoltaics because of their superior properties, for example, strongly quantum-confined photoluminescence (PL) and color-tunable emission, as well as their nontoxic and earth-abundant nature. Recently, we successfully synthesized colloidal SiQDs through pulsed ultraviolet (UV) laser illumination of porous Si powders. With the introduction of hydrogen fluoride (HF), the dissolution of the surface oxide is more efficient, promoting surface passivation provided by hydrogen-terminated Si surfaces. Therefore, after hydrosilylation reactions, the PL quantum efficiencies of colloidal SiQDs with visible luminescence are increased.

To date, state-of-the-art PSCs require well-tailored high-quality crystals or custom-designed device structures to achieve high performance. In this work, multifunctional SiQDs synergistically boost the efficiencies of PSCs by fully capturing sunlight without changing the original PSC process. This quantum dot-assisted photovoltaic strategy uses facilely synthesized SiQDs to harvest the unabsorbed sunlight in PSCs, which has the cost-effective production advantages of an earth-abundant element, a solution process and unchanged perovskite crystallization. The broadband spectral response of PSCs can be enhanced by harvesting the visible luminescence generated by SiQDs excited by short wavelength light between 300 and 530 nm and reflecting the perovskite-unabsorbed visible light in the 500–800 nm region, resulting in minimization of the thermalization loss. As a result, the SiQD-based PSC yields a promising PCE of 19.6%, compared with 18.2% for the bare PSC device. The best-performing PSC reaches a short-circuit current density ($J_{SC}$) of $\gtrsim 23$ mA/cm$^2$. Most impressively, using the SiQD-assisted photon management strategy, a low-quality PSC can exhibit an increase in the PCE from 7.3 to 12.9%. This approach provides a simple pathway to improve the photocurrent generation in PSCs, giving an innovation method for realizing PSCs on the path toward commercialization.

**Materials and methods**

**SiQD synthesis procedures**

As shown in Supplementary Fig. 1, colloidal SiQDs were prepared by pulsed UV laser irradiation of porous Si (obtained from Quantum14, Ltd.) in a cosolvent (including 1-decene and HF). Porous Si powder was prepared by standard electrochemical etching in a HF/C$_2$H$_5$OH aqueous solution followed by drying under ambient conditions. To prepare the colloidal SiQDs, the porous Si powder was first dispersed in an organic solvent (such as 1-decene) in a quartz cuvette. Then, a small amount of aqueous HF solution was added to the organic solution. Finally, 266 nm UV laser pulses from a Q-switched Nd:YAG laser (continuum) irradiated the sample with a pulse duration of 5 ns and a repetition rate of 15 Hz for 2 h. The laser fluence was $\sim 0.2$ J/cm$^2$. During laser irradiation, the porous Si powder dispersed in the HF/1-decene solution was constantly stirred by a magnetic stirrer. After laser irradiation, the supernatant of the liquid was centrifuged at 13,000 rpm for 20 min using a membrane filter with a pore size of $\sim 200$ nm to remove the HF and residual powder. This process enabled a colloidal SiQD solution in toluene to be obtained.

**Solar cell fabrication**

A SnO$_2$ solution (Alfa Aesar, tin(IV) oxide, 15% in a H$_2$O colloidal dispersion) was diluted with water to 2.67%. The solution was then spin-coated on clean ITO glass at 3000 rpm for 30 s. The coated ITO substrate was annealed at 150 °C for 30 min in air and then transferred to a glovebox. Solutions of C$_{60}$ pyrroline tris-acid (CPTA) (Sigma-Aldrich, 97%) in dimethylformamide (DMF) at a concentration of 2 mg/ml were prepared. The ITO/SnO$_2$ substrates were spin-coated with the CPTA solutions at 4000 rpm for 30 s and then annealed at 140 °C for 10 min to facilitate esterification between the carboxyl groups of CPTA and the hydroxyl groups of SnO$_2$. Next, 1 ml of a perovskite solution [1.25 M in dimethylformamide (DMF) with a 1:1 molar ratio of lead iodide (PbI$_2$, Alfa Aesar, 99.9985%) to methylammonium iodide (MAI with high purity, Xi’an Polymer Light Technology Corp, $\gtrsim 99.5$, MAI with low purity, Sigma-Aldrich, 98%)] was mixed with 5 mg of urea (Alfa Aesar, 98%+) and then stirred at 60 °C for 8 h in a glovebox. Afterwards, the perovskite precursor solution was deposited on the ITO/SnO$_2$/CPTA template in a glovebox by spin-coating at 5000 rpm for 30 s, with chlorobenzene (CB) treatment 8 s after the spin-coating began. Then, the film was annealed at 100 °C for 10 min. A hole transport solution was prepared by combining 1 ml of a 72.3 mg/ml solution of 2,2’,7,7’-tetrakis-(N,N-di-p-methoxyphenylamine)9,9’-Spirobifluorene (Spiro-OMeTAD) (Boron Chemical, 99.7%) in CB, 28.8 µl of 4-tert-butylpyridine (Sigma-Aldrich, 96%), and 17.5 µl of a 520 mg/ml lithium bis(trifuoromethylsulfonyl)imide (Sigma-Aldrich, 99.9%) stock solution in acetonitrile. A hole transport layer was spin-coated from the Spiro-OMeTAD solution at 2000 rpm for 30 s, followed by deposition of a 75 nm patterned Au top electrode by thermal evaporation. All of the abovementioned solutions were filtered through a 0.45 µm polytetrafluoroethylene filter prior to use.

**Characterization and measurements**

High-resolution transmission electron microscopy (HR-TEM) images and selected area electron diffraction (SAED) patterns were obtained using a Philips Tecnai G2...
(FEI-TEM) microscope operating at 200 kV. Fourier transform infrared (FTIR) chemical analyses were performed with a NICOLET 6700 (Thermo Fisher Scientific, USA) instrument. X-ray photoelectron spectroscopy (XPS) measurements were recorded using a Kratos AXIS ULTRA DALD XPS/UPS system. XPS survey scans were collected to identify the overall surface composition using a monochromatic Al Kα X-ray source (1486.6 eV). The measurement procedures for the photocurrent density–voltage (J–V) curves and the external quantum efficiency (EQE) spectra were as follows. The cell parameters were obtained under incident light with an intensity of 100 mW/cm² (as measured using a thermopile probe; Oriel 71964) generated by a 300 W solar simulator (Oriel Sol3A Class AAA Solar Simulator 9043A, Newport) and passed through an AM 1.5 filter (Oriel 74110). The light intensity was further calibrated using an Oriel reference solar cell (Oriel 91150) and adjusted to 1 sun. The monochromatic quantum efficiency was recorded using a monochromator (Oriel 74100) under short-circuit conditions. PL spectra were obtained using a fluorescence spectrophotometer (F-4600, Hitachi Ltd., Tokyo, Japan) with a 150 W Xe lamp as the excitation source at room temperature. UV/Visible transmission spectra were obtained using a Lambda 750 UV/VIS/NIR spectrophotometer (PerkinElmer). AFM (Hitachi AFM500II, tapping mode) was used to observe the surfaces of the perovskite and perovskite/SiQD films. Size distribution analysis via dynamic light scattering was performed with a Nanotrac Wave (MicrotracBEL Corp., Japan) instrument. Ultraviolet photoelectron spectroscopy (UPS) measurements were conducted using a Shimadzu spectrometer (AXIS ULTRA DLD) with a Hel (21.2 eV) discharge lamp.

Results

Characterization of SiQDs

SiQDs were synthesized mainly using a pulsed Nd:YAG laser for 266 nm irradiation of a porous Si suspension [consisting of porous Si powder, long unsaturated alkyl ligands (i.e., 1-decane) and HF] with minor modifications according to our previous results28,29 (see Supplementary Fig. 1 and the “Materials and methods” section). After laser fragmentation, the supernatant fluid was centrifuged at 13,000 rpm for 20 min and then filtered through a membrane mesh (pore size circa 200 nm). Finally, the filtered SiQDs were redispersed into toluene at a concentration of 10 mg/ml to form the target colloidal solution. Figure 1a shows an HR-TEM image of the as-prepared colloidal SiQDs with diameters between approximately 4 and 7.5 nm. These SiQDs were formed by fragmentation of porous Si powder, and thus, the size distribution of the SiQDs is similar to the size distribution of the Si cores constituting porous Si29. Since the porous Si cores have a relatively wide size distribution, the size distribution of the SiQDs is also wide (that is, >5% of the average diameter). A dynamic light scattering measurement was performed, showing a colloidal hydrodynamic size distribution of 10.9 ± 1.4 nm (see Supplementary Fig.
2). The wide size distribution will benefit PSCs, which will be discussed later. Furthermore, in Fig. 1a, the visible crystal fringes of three SiQDs with two types of interplanar lattice distances of 1.9 and 2.3 Å match the Si (220)[29] and SiOx (111)[30] crystal planes, respectively, indicating the existence of oxide regions in the SiQDs. As presented in Fig. 1b, the SAED patterns indicate reflection bands for the (111), (220), and (311) index facets of the SiQDs, corresponding to the face-centered cubic (FCC) silicon phase with a lattice parameter of 5.43 Å[30]. In addition, the [200] planes in the SAED pattern are attributed to the crystal plane of FCC silicon with a smaller lattice parameter of 3.90 Å[30] due to the presence of silicon oxide, implying that the SiQDs are partially oxidized in air.

To introduce SiQDs into PSC devices, toluene-diluted SiQD suspensions with various concentrations (0.001–1 mg/ml) were spin-coated onto perovskite layers at 4000 rpm for 30 s. After deposition of the SiQDs, the chemical structure of the SiQD films could be analyzed using the FTIR technique, as depicted in Fig. 1c. The FTIR spectrum of the as-synthesized film is characterized by absorption bands attributable to the surface functionalities of the SiQDs after hydroxylation, namely, the Si–O asymmetric stretching mode [v asym(Si–O)] in the region from 1010 to 1080 cm−1 (refs. 28,31), v sym(Si–C) at 864 cm−1 (ref. 32), and the Si–H symmetric stretching mode [v sym(Si–H)] at 2100 cm−1 (ref. 31). The as-synthesized SiQDs exhibit Si–O vibrations in Fig. 1c due to oxidation of the silicon surface exposed to air, thus corroborating the HR-TEM and SAED results. The presence of the Si–C band indicates that 1-decene molecules are directly attached to the SiQD surfaces via the hydrosilylation reaction. Herein, the C–H bending modes [δ(C–H)] and C–H stretching vibrations [ν(C–H)] located in regions 1200–1500 cm−1 (refs. 28,31) and 2800–3000 cm−1 (ref. 31), respectively, are from this long alkyl ligand. Figure 1d plots a survey XPS spectrum to identify the oxygen content in the SiQDs. The XPS scan yielded an O 1s/Si 2p ratio of 1.07, indicating that the SiQDs have a large oxidized proportion, in agreement with the abovementioned FTIR results.

**Photovoltaic performance**

The SiQD-containing PSC with the architecture of indium tin oxide (ITO) glass anode/SnO2 electron transporter/CPTA electron transporter/CH3NH3PbI3 perovskite/SiQDs/Spiro-OMeTAD hole transporter/Au cathode is schematically shown in Fig. 2a, wherein the SiQDs are used as photocurrent enhancers. Supplementary Fig. 3a shows a cross-sectional scanning electron microscopy (SEM) image of the pristine PSC device (without SiQDs); the perovskite absorber remained conformally flat over a wide range, with a film thickness of 320 nm. Figure 2b shows J–V curves of PSCs without and with various concentrations of SiQDs (from 0.001 to 1 mg/ml) under simulated AM 1.5G illumination (100 mW/cm²). The detailed J–V values are summarized in Table 1. These PSCs exhibit an open-circuit voltage (V OC) of 1.07 V and similar fill factors (FFs) of ~79% up to the addition of excess SiQDs (1 mg/ml) to the PSC device. The champion PSC incorporating the photocurrent-enhancing SiQDs at a concentration of 0.01 mg/ml yields a PCE of 19.6% with a short-circuit current density (J SC) of 23.10 mA/cm². These two parameters are comparable to those of the control PSC without any SiQD coating, which shows a PCE of 18.2% and a J SC of 21.44 mA/cm², indicating that the SiQDs can increase the efficiency of the PSC device through the superior photocurrent gain effect.

A small J–V hysteresis is observed in the PSCs constructed with SiQDs in different scan directions, as shown in Fig. 2c. Fortunately, a stable photocurrent output is observed in Fig. 2d. After 1500 s of illumination, the SiQD-based PSC device still presents a high photocurrent of 21.20 mA/cm² with a stabilized PCE of 19.3% at the maximum power output (at 0.91 V). Statistical J–V results were collected by examining 15 PSCs from five batches, as depicted in Fig. 2e and given in Table 1. The same trend is observed, in which the PCE and J SC values of these PSCs are maximized when the SiQD concentration reaches 0.01 mg/ml. Moreover, the increase in the PCEs is mainly due to the high photocurrent gain contributed by the SiQDs. In general, high-efficiency PSCs can be achieved by using high crystallinity perovskite layers. However, to fabricate these well-tailored perovskites, multistep sublimation of precursors is required, which greatly increases the costs in the PSC industry. Since SiQDs can harvest photon energy more effectively in PSCs, we applied this method to low-efficiency but inexpensive PSCs. SiQDs were added to a PSC device with a low-purity perovskite layer and remarkably suppressed the defect-induced photocarrier transport barrier, resulting in a PCE of 12.9% with a J SC of 22.96 mA/cm² and an FF of 52.4% (see Supplementary Fig. 4). Compared with the bare PSC, the SiQD-based PSC exhibits a surprising increase in the PCE by 5.6% (from 7.3 to 12.9%), indicating that SiQDs can provide another low-cost and effective way to promote PSCs in the commercial market.

**Photocurrent analysis**

From the above results, the efficiency improvement of the PSCs mainly comes from the additional photocurrent boost, and thus, the photocurrent gain can be further analyzed using the EQE. The EQE spectra of PSCs with various concentrations of SiQDs (0.001–1 mg/ml) and without SiQDs are depicted in Fig. 3a. The best-performing PSC with 0.01 mg/ml SiQDs shows a broad EQE spectrum and a significantly high value of >75% from 360 to 760 nm, giving an integrated photocurrent of
Table 1  Photovoltaic characteristics of PSCs incorporating SiQDs at various concentrations measured in the forward bias (FB) to short-circuit (SC) direction or in the SC to FB direction under AM 1.5G illumination (100 mW/cm²).

| SiQD (mg/ml) | V_OC (V) | J_SC (mA/cm²) | FF (%) | PCE (%) |
|--------------|----------|---------------|--------|---------|
| Without (FB → SC) | 1.07 (1.05 ± 0.01) | 21.44 (21.59 ± 0.13) | 79.4 (78.1 ± 0.9) | 18.2 (17.7 ± 0.2) |
| Without (SC → FB) | 1.06 | 21.21 | 78.3 | 17.6 |
| 0.001 (FB → SC) | 1.07 (1.05 ± 0.02) | 22.18 (22.45 ± 0.35) | 79.5 (78.3 ± 1.1) | 18.8 (18.5 ± 0.3) |
| 0.01 (FB → SC) | 1.07 (1.06 ± 0.01) | 23.10 (22.80 ± 0.32) | 79.7 (79.2 ± 0.7) | 19.6 (19.2 ± 0.4) |
| 0.01 (SC → FB) | 1.06 | 23.16 | 75.3 | 18.5 |
| 0.1 (FB → SC) | 1.07 (1.06 ± 0.02) | 22.72 (22.62 ± 0.46) | 79.3 (78.5 ± 0.8) | 19.2 (18.8 ± 0.3) |
| 1 (FB → SC) | 1.06 (1.06 ± 0.01) | 21.82 (22.09 ± 0.54) | 77.5 (74.6 ± 2.5) | 17.9 (17.4 ± 0.3) |

The average results in brackets represent the standard deviation of 15 devices from five different batches.
21.98 mA/cm². This value is consistent with the $J_{SC}$ parameter measured from the $J$–$V$ characteristics under simulated AM 1.5G irradiation (see Fig. 2b). Furthermore, the corresponding EQE enhancements were extracted using the EQEs of SiQD-based PSCs divided by those from the EQE spectrum of the reference PSC (without SiQDs) as shown in Fig. 3b. The enhancement in the EQE ($\Delta EQE$) demonstrates that the deposition of 0.01 mg/ml SiQDs can remarkably improve the photocurrent generation efficiency, with EQE enhancement peaks of more than 10% at 360, 410, 630, and 758 nm. We also provide two other sets of EQE and $\Delta EQE$ data, as shown in Supplementary Figs. 5 and 6, respectively. The results show that PSCs with 0.01 mg/ml SiQDs still have the highest EQE gain effect. Once the PSC is covered by SiQDs, its EQE improvement effect at short wavelengths is obvious. At the same time, when the concentration of SiQDs spin-coated on the PSC device exceeds 0.01 mg/ml, the EQE improvement in the visible band also increases. The results show that SiQD-based PSC devices exhibit photocurrent gain effects over a wide range from UV to visible light.

Discussion

Photon management in SiQD-based PSCs

To gain insight into the photon management mechanism, the PL signal from SiQDs was studied in detail using fluorescence mapping spectra, as demonstrated in Fig. 4a. Two emission bands of SiQDs are displayed in the visible wavelength range from 380 to 775 nm under excitation at UV-visible regions (300–530 nm). The long wavelength luminescence can be attributed to quantum confinement effects, while the short wavelength luminescence is caused by oxidized SiQDs. Placing SiQDs in air to measure the PL signal may cause some of them to be oxidized. Fortunately, the oxidation of SiQDs only occurs on surfaces without ligands. Once these ligand-free surfaces are oxidized, the oxidation of SiQD stops. The PL signal of SiQD shown in Fig. 4a is the data after the oxidation has stopped. The luminescence of these two bands (see Fig. 4a) can be reabsorbed by perovskite crystals and converted into electrical energy. Figure 4b clearly shows that SiQDs absorb photons from 300 to 530 nm and emit visible photons (380–775 nm) after the SiQDs are coated onto the perovskite [CH$_3$NH$_3$PbI$_3$ (MAPbI$_3$)] surfaces. Then, the visible emission can be reabsorbed by MAPbI$_3$ light harvesters. PSC photovoltaics can only effectively convert sunlight in the 400–750 nm spectral range into electricity. Fortunately, the SiQDs show an excellent UV-visible absorption capacity and can convert short wavelength light to a perovskite-operating visible band. In addition, the size distribution of the SiQDs is relatively wide (see Supplementary Fig. 2). Therefore, a broad excitation peak is achieved between 300 and 530 nm due to quantum size effects. As a result, the photoelectric conversion efficiency across this broad UV-visible region will be more efficient in PSCs. Therefore, the EQE responses of all SiQD-based PSCs are enhanced at short wavelengths, as shown in Fig. 3b.

Moreover, the EQEs of SiQD-containing PSCs in the 550–770 nm wavelength region are also higher than those of the PSC without SiQDs. The optical properties were further analyzed using UV-visible reflection spectra. After coating the MAPbI$_3$ layer with SiQDs, the reflection response was significantly improved at wavelengths between 550 and 800 nm, as shown in Fig. 4c. The strong reflection caused by SiQDs is beneficial for reabsorbing the visible light that the perovskite does not absorb. In addition, the reflection intensity in the 550–800 nm band...
is enhanced as the concentration of SiQDs coated on the perovskite increases. This result is consistent with the ΔEQE signals (see Fig. 3b). When the SiQD concentration of the SiQD-based PSC samples is increased from 0.001 to 0.1 mg/ml, the corresponding ΔEQE in the 550–800 nm range is improved. However, the EQE response requires not only the capability of the material to absorb light but also the ability to convert light into electricity. Although the sample with a SiQD concentration of 1 mg/ml has a stronger reflection spectrum than the 0.01 mg/ml sample, its ΔEQE response at 550–800 nm is decreased. This shows that the photoelectric conversion ability has a great impact after deposition of a high concentration of SiQDs. Thus, enhancement of the EQE in the long wavelength visible region (>550 nm) can be obtained by covering the perovskite crystals with SiQDs (see Fig. 4b). Because of these two SiQD-assisted photon management effects, namely, converting short wavelength light into visible light and reflecting long wavelength visible light, the EQE response of the best-performing PSC device (0.01 mg/ml) can be enhanced across a 400 nm ultrawide region (~350–750 nm) (see Fig. 4b). In addition, the traditional passivation layer deposited after the perovskite is used to promote the carrier transport of defective perovskite crystals (including defects, grain boundaries and incomplete crystals). In this work, SiQDs are mainly used to capture the UV-visible light unabsorbed by the perovskite and then recycle this light through reabsorption and reflection. Since SiQD-based PSCs capture more light than the pristine device (without SiQDs), a higher photocurrent will be generated. In these PSCs, the upper SiQD clusters do not change or passivate the bottom perovskite crystals. Therefore, the SiQDs are a photocurrent enhancer, not a passivator (as confirmed in Supplementary Fig. 8).

Morphology of SiQDs and the photoelectric impacts on PSCs

To further explore the reasons for the reduction in the photoelectric responses (including the J–V curves and EQE characteristics) after adding excess SiQDs (concentration >1 mg/ml), we systematically studied the morphology evolution of SiQDs deposited onto MAPbI₃ perovskites with increasing concentration. Figure 5a presents atomic force microscopy (AFM) images of the pristine MAPbI₃ perovskite (without SiQDs) and of SiQDs spin-coated on perovskite crystals at concentrations of 0.01 and 1 mg/ml. Compared with the pristine perovskite crystals that exhibit clear lattice fringes, the MAPbI₃ coated with 0.01 mg/ml SiQDs appears to have different size particles on the perovskite crystals. As shown in Supplementary Fig. 3b, c, the AFM depth profiles show that the thicknesses of the 0.01 mg/ml SiQDs and 1 mg/ml SiQDs are 12.7 and 15.2 nm, respectively. The SiQD clusters are approximately tens of nanometers in diameter, which are much larger than the as-prepared SiQDs (see Fig. 1a). This phenomenon occurs because the surface of the as-synthesized SiQDs has a long carbon chain, and once exposed to the hydrophilic perovskite surface, these SiQDs will aggregate into clusters to reduce the interface energy. The survey XPS spectrum demonstrates a Si 2p/Pb 4f value of ~13.5% (see Supplementary Fig. 9); hence, the amount of SiQDs in the perovskite/SiQD (0.01 mg/ml) sample can be quantified. After a large amount of SiQDs was deposited onto the perovskite (1 mg/ml), the perovskite surface was covered by SiQD clusters, resulting in the disappearance of the perovskite grains. These samples (without SiQDs, 0.01 and 1 mg/ml) are plotted as...
corresponding side schematics to explain the influence on the photocarrier dynamics of the SiQD morphologies, as shown in Fig. 5b. In the pristine perovskite (without SiQDs) and the low SiQD concentration sample (0.01 mg/ml), perovskite-generated holes can be directly transported to the hole transport layer [Spiro-OMeTAD] because most perovskite grains are exposed. Although the valence band level of the SiQDs (see Supplementary Fig. 10) is lower than that of MAPbI3, hole transport in the SiQD-based PSC is still easy. However, as the concentration of SiQDs increases to 1 mg/ml, the holes will be blocked due to the lower valence band level of SiQDs with high coverage. This will cause hole accumulation at the perovskite/SiQD interface, leading to poor performance of PSCs with many SiQDs. These results illustrate that appropriate adjustment of the SiQD amount can improve the PCEs, FFs, and $J_{SC}$ of PSCs without excessively increasing the cost.

**Conclusion**

In summary, we have successfully introduced SiQDs into PSC devices to achieve a PCE enhancement up to 1.4% (increased from 18.2 to 19.6%). SiQDs with a relatively wide size distribution were synthesized using pulsed UV laser irradiation, yielding a broad excitation peak from 300 to 530 nm. After excitation at broad UV-visible regions, the SiQDs converted the short wavelength photons into perovskite-absorbable visible bands. At the same time, reflection occurred at the perovskite/SiQD interface, resulting in 550–800 nm perovskite-unabsorbed light being reflected and then reabsorbed by the perovskite crystals. Through these sunlight harvesting methods, UV light and unabsorbed visible light can be effectively utilized by the PSCs, achieving a high $J_{SC}$ of 23.10 mA/cm², and these phenomena are economically beneficial to low-purity perovskite-based PSCs, with an impressive PCE boost of up to 5.6% (increased from
7.3 to 12.9%). In general, low-purity perovskites lose too much light due to the poor crystallinity. Once SiQDs are coated onto the surface of low-quality perovskites, a large amount of unabsorbed light will be reused. For high-purity perovskites, since most of the light is absorbed by these perovskites, the ratio of recovered light is relatively low. Therefore, the SiQDs are more effective for low-purity PSCs than for high-purity PSCs. Generally, perovskite crystals require high-purity precursors and strict process control to achieve high-crystallinity characteristics. If SiQDs are introduced into inexpensive low-purity perovskite solar cells, this will provide a new way to commercialize PSCs. Therefore, the photocurrent management strategy has tremendous implications for the real application of low-cost, high-efficiency PSCs.

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Author contributions

Y.-C.W., C.-W.C., and K.T. conceived and supervised this project. Y.-C.W. carried out the final device optimization. T.N., N.K., and T.S. synthesized the colloidal SiQDs. Y.-C.W., S.-K.H., T.N., Y.-T.K., C.-H.C., D.-Y.W., Y.J.C., and S.L. performed the data collection, interpretation, and analysis. Y.-C.W., C.-W.C., and K.T. wrote and revised the paper. All authors commented on and agreed to its content.

Conflict of interest

The authors declare that they have no conflict of interest.

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