Enhanced power conversion efficiency via hybrid ligand exchange treatment of p-type PbS quantum dots

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Abstract

PbS quantum dot solar cells (QDSCs) have emerged as a promising low-cost solution processable solar energy harvesting device and demonstrated good air stability and potential for large-scale commercial implementation. PbS QDSC achieved a record certified efficiency of 12% in 2018 by utilizing an n+-n-p device structure. However, the p-type layer has generally suffered from low carrier mobility due to the organic ligand 1,2-ethanedithiol (EDT) that is used to modify the quantum dot (QD) surface. The low carrier mobility of EDT naturally limits the device thickness as the carrier diffusion length is limited by the low mobility. Herein, we improve the properties of the p-type layer through a 2-step hybrid organic ligand treatment. By treating the p-type layer with two types of ligands, 3mercaptopropionic acid (MPA) and EDT, the PbS QD surface was passivated by a combination of the two ligands, resulting in an overall improvement in open circuit voltage, fill factor and current density, leading to an improvement in cell efficiency from 7.0% to a champion cell efficiency of 10.4%. This achievement was a result of improved QD passivation and a reduction in interdot distance, improving charge transport through the ptype PbS quantum dot film.

Keywords: PbS quantum dot, quantum dot solar cell, p-type layer, ligand exchange, interdot distance

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Graphical abstract



1. Introduction

The earliest colloidal quantum dot (CQD) implementation in solar cells achieved an efficiency of 1-3% based on Schottky junctions ^{1,2}. Since then, the structure of QDSCs has evolved from a simple metal-PbS-metal Schottky junction, to the current n⁺-n-p structure ³. This, along with improvements in the passivation and fabrication of the absorber layer via solution ligand exchange with halide ligands, has allowed the performance of QDSCs to improve to a PCE of 12% ^{4,5}. However, challenges remain with the p-type PbS hole transport layer (HTL), which is a vital component of QDSC devices.

The p-type quantum dot layer based on 1,2-ethanedithiol (EDT) treatment was first demonstrated by Luther et al. in 2008⁶. This discovery enabled the formation of p-type solids after oxidation in air and formed a film with high electronic coupling that drastically improved electronic properties over PbS QDs capped with oleic acid. This makes EDT treated PbS QD film (PbS-EDT) a suitable HTL that can be implemented into high performance n⁺-n-p solar cell structures. Various optimizations have been performed on the EDT treatment with adjustments to the ligand concentration and treatment duration and the recipe differs for different publications ^{5,7,8}. The current highest efficiency cells published have a power conversion efficiency (PCE) of 12.48% and use PbS-EDT as the HTL ⁵. However, these recent advancements have all been due to progress in improving the passivation of the absorber layer.

To date, there is a lack of progress in improving the p-type PbS-EDT HTL for PbS QDSC. The passivation of PbS QDs with EDT has not fundamentally changed since it was first demonstrated in 2008^{6,9}. The key to a good HTL is to have sufficiently high p-doping concentration in order to increase the charge carrier concentration and form a wide depletion region when interfaced with an n-type absorber layer. Wider depletion regions can enable thicker devices to increase light harvesting that will boost the current output of the solar cell. One main drawback of PbS-EDT is the low carrier mobility. PbS-EDT thin films have a carrier mobility of around 10^{-4} cm² V⁻¹ s⁻¹, which is one order of magnitude lower than PbS treated with 3-mercaptopropionic acid (MPA) and two orders of magnitude lower than halide treated PbS QD films¹⁰. As such, typical methods to increase doping concentration such as the introduction of impurity atoms have limited improvement as the mobility will be drastically reduced due to the impurity atoms. Hu et al. has demonstrated

that a 1.0% Ag-doping in PbS-EDT can improve the performance of QDSCs from a PCE of 9.1% to 10.6% by improving the hole carrier density and p-type character¹¹.

While higher Ag doping concentrations further increases hole carrier density, the device performance peaked at 1.0% doping concentration due to a decrease in carrier mobility and diffusion length that negates the gain from improved charge carrier density. Furthermore, the improved hole mobility is still lower than that of the n-type QD layer. This is an inherent issue with EDT treatment as it has a low carrier mobility due the larger interdot distance. There is an urgent need to find an alternative p-type PbS material with improved charge transport such that the doping limit can be raised to further improve the charge carrier concentration and reduce carrier recombination velocity at the junction.

An alternative ligand to EDT that contributes a higher carrier mobility is MPA. MPA differs from EDT in that one of the thiol group has been replaced by a carboxylic acid. QD passivated with carboxylic acid ligands have better charge transport compared to thiols ^{12,13}. Devices using MPA treated PbS quantum dot film (PbS-MPA) as HTL have shown mixed results. PbS-MPA heterojunction devices have demonstrated to have a higher J_{SC} over PbS-EDT devices ¹⁴. However, the gain in V_{OC} of PbS-MPA over PbS-EDT did not seem to be reproducible in another study comparing MPA and EDT treatments across various device structures ¹⁵. The lower V_{OC} of PbS-MPA over PbS-EDT devices were also observed in this work, although PbS-MPA still demonstrated improvements in J_{SC} over PbS-EDT. Likewise, majority of PbS QDSC publications prefer EDT over MPA as a p-type ligand due to the superior V_{OC} of PbS-EDT based devices, while the J_{SC} shortfall is tackled through passivation and thickness improvements in the PbS-PbI₂ absorber layer ^{4,5,16-19}, including the world record 12.01% PCE device⁵.

In this paper, we introduce a novel 2-step MPA and EDT hybrid ligand treatment method to capitalize on the high J_{SC} characteristic in PbS-MPA devices and high V_{OC} characteristic of PbS-EDT devices. The ligand exchange procedure is shown in Figure 1. The presence of the carboxylic group in MPA from the first treatment would help improve device performance through improvements in the conductivity and mobility by one order magnitude when compared to the thiols ^{10,13,14}. The two-step treatment also introduces EDT as the primary passivating ligand to achieve a higher V_{OC}, while still retaining MPA ligands for improved conductivity over pure PbS-EDT films. This results in an improvement in PCE from 7.0% for PbS-EDT devices and 8.4% for PbS-MPA devices to 10.4% for PbS-Hybrid devices.



Figure 1: Procedure for 2-step MPA and EDT hybrid ligand treatment. The conventional ligand exchange method for the p-type layer involves only one ligand treatment step conducted on an oleic acid capped PbS QD film. The method described in this work demonstrates that a solid layer can also be further treated with a 2nd step to alter and improve the film's properties.

2. Experimental section

PbS CQD synthesis: The materials used were lead (II) oxide (PbO) (Sigma-Aldrich 99.999% trace metals basis), oleic acid (OA) (Sigma-Aldrich 90% technical grade), 1-octadecene (ODE) (Sigma-Aldrich 90% technical grade), bis(trimethylsilyl)sulfide (TMS) (Sigma-Aldrich synthesis grade), hexane (Chem-Supply \geq 95%) and acetone (Ajax Finechem \geq 99.5%). The synthesis of PbS QDs was based on a previous published recipe ²⁰. The reactants 0.45 g PbO, 1.62 ml OA and 20 ml ODE were added to a 100 ml three-neck flask and degassed and stirred at 60 °C for 2 hours and then 100 °C for 2 hours to form colorless Pb-oleate precursor solution. Once the solution is completely transparent and colorless, 140 μ l TMS dissolved in 5 ml ODE was injected into the Pb-oleate solution at 80 °C and left heating for 10 s for QD nucleation and growth. After 10 s, the heating mantle was removed and turned off and the solution was allowed to naturally cool to room temperature. The PbS CQD solution was washed and precipitated with hexane and acetone with a volume ratio of 1:3. The precipitation was repeated twice and the PbS CQD was dried under vacuum for 30 minutes.

MgZnO synthesis: The materials used were Zinc acetate dihydrate (Ajax Finechem \geq 99.5%), magnesium acetate tetrahydrate (Sigma-Aldrich \geq 99% ReagentPlus®), anhydrous 2-methoxyethanol (Sigma-Aldrich \geq 99.8%) and monoethanolamine (Sigma-Aldrich ACS reagent \geq 99.0%). 1g of zinc acetate dihydrate and 0.05g of magnesium acetate tetrahydrate were added to 10 ml of 2-methoxyethanol solvent and 280 µl of monoethanolamine complexant. The solution was stirred at 75 °C for 12 hours until all the precursors were dissolved into a clear homogenous sol-gel.

Device fabrication: The materials used were lead (II) iodide (PbI₂) (Sigma-Aldrich 99%), dimethylformamide (Analar NORMAPUR 99.9%), 3-mercaptopropionic acid (Sigma-Aldrich 99%), methanol (RCI Labscan Ltd \geq 99.9%), 1,2-ethanedithiol (TCI >99.0%), and acetonitrile (RCI Labscan Ltd \geq 99.8%). The QDSC device fabrication was based on a previous published recipe¹¹. ITO glass was cleaned with detergent, deionized water and acetone with ultrasonication. The MgZnO sol-gel was spin-coated on ITO glass substrates at 3000 RPM for 30 s then dried at 100 °C for 10 minutes and annealed at 360 °C for 20 minutes. The MgZnO film fabrication was repeated 3 times until ~130nm thickness was achieved. For absorber layer, a colloid containing 80 µl of PbS CQDs suspended in hexane at 30 mg/ml concentration was spin-coated on the MgZnO film at 3000 RPM for 30s, treated then with 10 mM PbI₂ in DMF for 50s via dip coating, and rinsed with acetonitrile and dried under N₂ gas flow. This process was repeated 6 times until iodide capped QD (PbS-iodide) absorber layer with ~150 nm thickness was fabricated.

For the HTL layers, 80 μ l of PbS CQDs with a concentration of 15 mg/ml suspended in hexane was spin-coated at 3000 RPM for 30s on the substrate. Two ligands were used for this study, MPA and EDT. For PbS-MPA layer, the PbS-OA QD film was treated with 10% v/v MPA in methanol solution for 10 s followed by a methanol rinse. For PbS-EDT layer, the PbS-OA film was treated with 0.02% v/v EDT in acetonitrile solution for 10 s followed by rinsing with acetonitrile. For the hybrid ligand treated layer (PbS-Hybrid), the same PbS-OA film was treated in 2-steps, i. e. by MPA treatment followed by EDT treatment using the same recipe described above. The fabrication was carried out in air. The PbS QDSC devices were annealed at 75 °C in air for 15 minutes to oxidize the HTL and help form a p-type film. The devices were stored in a desiccator box in dry air overnight to further facilitate oxidation and improve p-type properties of the HTL in order to achieve the best efficiencies. 100 nm of gold electrodes were then fabricated on top of the HTL via thermal evaporation. The absorber area of the device was 0.0314 cm².

Characterization: The optical absorption spectra of the PbS colloid were measured with the Perkin Elmer Lambda1050 UV/Vis/NIR spectrophotometer in ambient condition. The steadystate photoluminescence (PL) spectra of PbS CQDs dispersed in hexane were measured by QuantaMaster 500 (Horiba) spectrofluorometer with an excitation from a xenon arc lamp at 600 nm and detected by a InGaAs detector. The Fourier transform infrared spectroscopy (FTIR) measurements were carried out with a PerkinElmer FTIR Spectrometer in reflection mode. The Transmission electron microscopic (TEM) imaging was done on dilute OA-capped PbS QDs dropped onto a copper TEM grid. For treated TEM samples, MPA and EDT ligands with concentrations described in the previous section were dropped onto the copper grid that contains PbS QDs. There samples were then put in an oven for drying to remove any remaining solvents. TEM imaging was carried out on a Philips CM200 field emission TEM. The scanning electron microscopic (SEM) images of the cross section and surface morphology were taken with the FEI Nova NanoSEM 230 Field Emission SEM. The photoemission yield spectroscopy (PYS) was measured using a StelaNet SL3 Deuterium lamp as a light source and JobinYvon HR 250 monochromator with stepper motor to control the wavelength of the incident light, and the amplified photoemission current is measured by grounding the sample stage through a Keithley 617 Electrometer.

The light current density–voltage (J-V) curves of different devices were obtained using a Keithley 2400 (I-V) digital source meter under simulated AM 1.5G solar irradiation at 100 mW/cm² (Newport, AAA solar simulator, 94023A-U). Dark J-V measurements were also carried out in the same system using a dark enclosure. The mobility of the HTL was measured with a field effect transistor (FET) using an Agilent B1500A semiconductor characterization system.

3. Results and discussion

3.1 IV performance of solar cell device



Figure 2: (a) Schematic of the QDSCs . The HTL is the only variable being investigated in this work. (b) The current density-voltage (J-V) curves of champion devices for different HTL ligand treatments. The device with MPA and EDT hybrid ligand treated HTL achieves both high V_{OC} and J_{SC} , while the device with EDT treated HTL has a high V_{OC} and MPA treated HTL has a high J_{SC} .



Figure 3: (a) PCE, (b) J_{SC} , (c) V_{OC} and (d) FF variation between different solar cell devices with different HTLs. The hybrid ligand treatment produces higher performance and reproducibility of the V_{OC} and FF.

	J _{sc}	Voc	FF	R _{SH}	Rs	PCE
Treatment	(mA cm ⁻²)	(mV)	(%)	(Ω cm²)	(mΩ cm²)	(%)
	24.1 ± 0.2	557 ± 5	57.9 ± 1	3814 ± 529	32 ± 2	7.8 ± 0.2
IVIPA	(25.1)	(573)	(61.1)	(7004)	(23)	(8.4)
FDT	21.6 ± 0.2	617 ± 4	47.5 ± 0.6	2654 ± 157	51 ± 2	6.3 ± 0.1
EDI	(21.9)	(635)	(51.0)	(3698)	(35)	(7.0)
Hybrid	23.5 ± 0.2	617 ± 3	65.4 ± 0.2	11771 ± 439	34 ± 2	9.5 ± 0.1
	(25.3)	(633)	(66.8)	(14064)	(22)	(10.4)

Table 1: Average J_{SC}, V_{OC}, FF, R_{SH}, R_S and PCE of solar cell device with different HTL as described. The values in brackets are the results from the best performing devices.

For device J-V characteristic comparison, two reference cells with PbS-MPA and PbS-EDT thin films as HTLs were used as a benchmark against the PbS-Hybrid devices. All other layers of the device are identical. The cross-sectional SEM image of the solar cell device is show in **Error! Reference source not found.**.

From the J-V curves shown in Figure 2b, it is clear that the PbS-EDT device has a superior V_{oC} at 635 mV compared to PbS-MPA, which is also reflected in other comparison studies between MPA and EDT ^{15,21}. This is likely due to the higher proportion of thiol groups in EDT that can be easily oxidized to achieve a higher doping concentration during air annealing to form a p-type film. The carboxylic group is inherently more resistant to oxidation and MPA devices have been shown to be more resistant to performance degradation compared to EDT devices when exposed to air [14]. Apart from doping via oxidation, direct addition of thiol ligands to carboxylic acid ligands was also demonstrated to significantly improve the V_{OC} of a QD heterojunction solar cell by passivating exposed sulfur atoms on the QD surface ²². PbS-EDT has also shown to have a longer lifetime than PbS-MPA¹⁴, possibly reducing carrier recombination.

Note that EDT passivation is effective at a much lower concentration of 0.02% v/v, compared to MPA which requires 2% v/v for complete passivation. Based on hard and soft acid base theory (HSAB), thiolate is a soft base and should have a much stronger affinity with the borderline soft acid Pb²⁺ compared to carboxylate, which is a hard acid ²³. This makes thiols a more potent passivating ligand compared to carboxylic acid.

The shunt resistance for the PbS-EDT device is noticeable lower than the other two devices. From the SEM images as shown in Figure S6, some pin holes can be observed for PbS-EDT but not for PbS-MPA or PbS-Hybrid, which might be the cause of the small R_{sh} for PbS-EDT device.

On the other hand, the PbS-MPA device has a better J_{SC} at 25.1 mA cm⁻² and a lower R_S of 23 m Ω cm² compared to PbS-EDT. This improvement is consistent with other studies on both heterojunction and n⁺-n-p device^{14,15,21}. The higher J_{SC} is due to the better charge transport of PbS-MPA over PbS-EDT¹⁴, and higher conductivity has been demonstrated to improve J_{SC} and R_S ¹¹.

The PbS-Hybrid device successfully combine the characteristics of higher J_{SC} and lower R_S of PbS-MPA and higher V_{OC} of PbS-EDT as shown in Figure 3 and Table 1. The champion device achieves a PCE of 10.4% with a V_{OC} of 633 mV, J_{SC} of 25.3 mA cm⁻² and FF of 66.8%. The EDT treatment after MPA improves the V_{OC} to EDT levels, indicating improved passivation of the HTL.

Despite the EDT treatment after the MPA treatment, the J_{SC} and R_S of PbS-Hybrid are relatively unchanged compared to PbS-MPA, which indicates that PbS-Hybrid device has a similar conductivity, charge generation and collection as PbS-MPA device. External quantum efficiency measurements as shown in Figure S8 indicates an overall improvement in charge collection for PbS-MPA and PbS-Hybrid devices over PbS-EDT. The high conductivity of PbS-Hybrid can be attributed to the presence of a small amount of MPA ligands remaining on the PbS QD surface and the shorter interdot distance shown discussed in section 3.3.



3.2 FTIR spectroscopic study on PbS QD surface passivation

Figure 4: FTIR spectra of PbS QD thin films treated with various ligands. The spectra show the replacement of long chain ligands by short chain linking ligands by tracking the reduction in C-H and C-O stretch peaks which corresponds to the long carbon chain and carboxylic acid group respectively on OA.

	C-O asymmetric stretch					
Ligand	Peak	Peak	MPA %			
	position	height				
MPA	1533	0.0116	100%			
Hybrid	1532	0.0020	18%			
EDT 1533		0.0004	0%			

Table 2: Percentage of MPA ligand coverage based on FTIR peak position and height of C-O asymmetric peak

To understand the nature of the hybrid ligand treatment, PbS-MPA, PbS-EDT and PbS-Hybrid thin films were characterized with FTIR spectroscopy. FTIR spectroscopy reveals the presence of organic ligands passivating the PbS QD surface for a dry PbS QD thin film absent of solvents and precursors. As seen from the FTIR spectra in Figure 4, OA molecules are efficiently removed by all ligand treatments. This is evidenced by the significant reduction of C-H stretch peaks at 2928 cm⁻¹ and 2856 cm⁻¹ whereby the long carbon chain OA ligands have been replaced by short carbon chain ligands. There is also a clear reduction of the C-O asymmetric and symmetric stretch peaks at 1533 cm⁻¹ and 1414 cm⁻¹ after treatment with MPA, EDT and hybrid ligand treatment. The MPA and hybrid ligand treatment both retain some C-O peak signal due to the presence of carboxylates from the MPA ligand. For PbS-MPA, there are no peaks at 1700 and 1300 cm⁻¹ position, each belonging to the C=O stretch and C-O stretch peak, indicating that the carboxylic acid is fully deprotonated and the ligand bonds to the QD surface in carboxylate form. For EDT, there is no S-H peak at around 2550 cm⁻¹ which indicates that the ligand is bound in its thiolate form to the QD surface²⁴.

Comparing the FTIR spectra of PbS-MPA and PbS-Hybrid, it is clear that hybrid ligand treatment results in the removal of MPA ligands, as evidenced by the significant reduction in both asymmetric and symmetric C-O stretch peaks; the addition of a 2nd step of EDT treatment in the hybrid ligand treatment removes majority of MPA ligands from the QD surface. The ability of EDT to remove and replace MPA ligands is consistent with the HSAB theory discussed previously. This demonstrates that the EDT treatment is effective even for a solid QD thin film linked with short chain MPA ligands.

According to the Beer-Lambert law, absorbance is directly proportional to the concentration of the absorbing species. The percentage of MPA ligands that were replaced by EDT can be estimated by taking the ratio of the C-O stretch peak before and after EDT treatment. The C-O asymmetric stretch peak at 1531 cm⁻¹ was chosen for the calculation as there is no overlap with EDT peaks in that region. The C-O symmetric stretch peak cannot be used as there is a wide peak at 1434 cm⁻¹ for EDT which is very close to the 1388 cm⁻¹ C-O symmetric peak for MPA. As seen in

Table 2, based on the C-O asymmetric stretch peak height of PbS-Hybrid, it is estimated that only 18% of the MPA ligands from the 1st treatment step remains on the surface of the PbS QDs after the 2nd step of EDT treatment, indicating 82% of the MPA ligands have been replaced by EDT.

The replacement MPA ligands with EDT justifies the improved V_{OC} of the PbS-Hybrid device. The built-in voltage of a PbS-EDT device can be improved through surface oxidation by air annealing dithiol treated PbS⁹. The replacement of MPA ligands with EDT increases the abundance of air sensitive thiol groups for oxidation and this increased oxidation improves the hole doping concentration responsible for the higher V_{OC} of the device.

Considering that most of the ligands in the PbS-Hybrid film are EDT, the reduced presence of MPA ligands may not be sufficient to justify the high J_{SC} and R_S characteristics of the PbS-Hybrid device, which match PbS-MPA device performances. This suggests that there are other contributors that can improve the interdot coupling that are independent of the presence of carboxylates such as interdot distance. In summary, the FTIR spectra show a clear replacement of MPA ligands by EDT in the hybrid ligand treatment, demonstrating that

EDT is the main ligand passivating the PbS QD surface, while about 18% of MPA ligands remain on the QD surface.

3.3 TEM study on interdot distance

Interdot distance has been shown to play a key role in determining the mobility and conductivity of QD thin films. The impact of ligand type and interdot distance to photovoltaic performance has been discussed comprehensively in Wang and co-workers' review paper ²⁵. Study of different lengths of dithiol ligands on PbSe QDs have shown an exponential increase in mobility as ligand length is reduced ²⁶. Investigation on Cu_{2-x}Se QDs functionalized by dithiol ligands has also shown that the conductivity of Cu_{2-x}Se thin films increases as the carbon backbone length is reduced ¹³. Under the dithiol family of ligands, EDT is the shortest with only 2 carbon backbone and is unsurprisingly the most commonly used ligand for the HTL in order to maximize film conductivity and mobility.

However, the dithiol ligand has an inherent problem of forming longer chain dimers when exposed to air which reduces mobility and conductivity. Weidman et. al. has observed an unusual trend whereby EDT treated QD film has a larger interparticle spacing of 1.2 nm than that of longer chain 1,4-butane dithiol treated film with an interparticle spacing of 0.8 nm ²⁷. This goes against the otherwise linear decreasing trend of interdot distance with decreasing carbon chain length of the ligand^{27,28}. This anomaly in the trend was similarly observed by Lynch et. al. who also observed a detrimental effect on conductivity due to the dimerization of EDT ¹³, which is illustrated in

Figure 5a. They proposed that performing the ligand exchange in air will result in the oxidation of the thiol functional group and the formation of sulfur-sulfur bond between two thiols on different QDs, effectively doubling the ligand length, i.e. dimerization. Performing the ligand exchange in an inert environment does not completely eliminate this effect, showing the high sensitivity of EDT even under low oxygen concentration environment¹³.



Figure 5: Ligand bonding between QDs for different treatment types and their molecular structures. (a) EDT treatment forms an EDT dimer with a disulfide linkage. (b) MPA treatment forms two overlapping MPA ligands. (c) Hybrid ligand treatment forming the proposed EDT monomer bond. The short interdot distance for the hybrid ligand is likely due to bridging by only one EDT monomer as opposed to the EDT dimer for EDT treated PbS QDs.

To understand the impact of hybrid ligand treatment to the interdot distance using TEM imaging, slightly larger QDs i.e. 4.5 nm in diameter were synthesized and treated with the appropriate ligands and observed under the TEM. The QDs used were larger than those synthesized for QDSC device in order to facilitate clear imaging. The larger QDs were synthesized by increasing the OA:PbO mole ratio from 2.5:1 to 14:1, which yields a QD size of 4.5 nm with a narrow size distribution as shown in **Error! Reference source not found.**. After the images were taken as shown in

Figure 6, the QD diameter and QD centre-to-centre distance were measured using an algorithm written in python and the OpenCV software packages. The interdot distance is the difference between the average centre-to-centre distance and the average diameter as shown in **Error! Reference source not found.**



Figure 6: TEM images of 4.5 ± 0.6 nm PbS CQDs functionalized by the ligands of (a) OA, (b) EDT, (c) MPA, and (d) and (e) hybrid ligands. The measured interdot distance is 2.9 ± 0.4 nm for PbS-OA, 1.2 ± 0.1 nm for PbS-EDT, 1.1 ± 0.2 nm for PbS-MPA, and 0.6 ± 0.2 nm for PbS-Hybrid. (e) shows PbS QDs multilayered superlattice matrix after hybrid ligand treatment with highly ordered self-assembled regions, which dominates most of the TEM imaging of the hybrid ligand samples. Such dense regions were not found in the PbS-EDT and PbS-MPA samples. (f) shows the interdot distance between PbS QDs treated with EDT, MPA and hybrid ligands. The PbS-Hybrid QDs show the shortest interdot distance as the QDs are only bridge by one EDT monomer as oppossed to an EDT dimer for PbS-EDT sample.

From

Figure 6f**Error! Reference source not found.**, PbS-EDT QDs show the largest interdot distance of 1.2 ± 0.1 nm after exchanging the original ligands of OA by EDT, which matches literature ²⁷. This value is double the molecular length of EDT, suggesting EDT molecules are dimerized as shown in Figure 5a. The interdot distance of PbS-MPA sample is 1.1 ± 0.2 nm, consistent with literature ²⁷. The molecule length of MPA was calculated to be 6.7 Å from software simulations using the MMFF94Fs model for energy minimization ²⁹. This implies that the QDs in PbS-MPA film are separated by two MPA molecules as shown in Figure 5b.

As can be seen in

Figure 6f, the hybrid treatment significantly reduces the interdot distance to 0.6 ± 0.2 nm. Since the hybrid treatment is a 2-step treatment, i.e. an MPA treatment followed by an EDT treatment, the interdot distance is firstly reduced from 2.9 nm for PbS-OA to 1.1 nm with the 1st MPA ligand treatment. At this point, the QDs are separated by two MPA molecular lengths. The 2nd step of the treatment with EDT introduces a slightly shorter

ligand with a length of 6.1 Å to further reduce the interdot distance by displacing MPA. Unlike the direct treatment of EDT on PbS-OA, treating PbS-MPA QDs with EDT does not lead to EDT dimerization that is responsible for the large interdot distance, allowing the QDs to directly couple by just one EDT ligand length as shown in Figure 5c. It is likely that the more densely packed PbS-MPA layer prevents the diffusion of excessive amounts of EDT ligands between QDs that drives the oxidation of thiols to disulfides. The reduction in interdot distance for PbS-Hybrid helps enhance interdot coupling and improves the mobility and conductivity of hybrid ligands over longer interdot distance such as PbS-EDT. This mobility and conductivity gains translate into the J_{SC} and R_S improvements of the QDSC devices using PbS-Hybrid as the HTL over PbS-EDT which will be elaborated in a later section.

The hybrid ligand treatment also yields a densely packed and highly self-assembled 3D structures shown in

Figure 6e that are not seen in the TEM images of MPA or EDT treated PbS QD film. This demonstrates that the reduced interdot distance by the hybrid ligand treatment allows for high degree of ordering. Single ligand treatment of PbS QDs tend to yield localize short range ordering of QDs separated by empty space, resembling a web of QDs such as the ones shown in

Figure 6b and c, which drastically reduces charge transport throughout the PbS QD film.

3.4 Conductivity and mobility

To observe the changes in the conductivity of PbS QD film with different treatment, a Schottky junction with an ITO/PbS/Au device structure was fabricated and the dark J-V was measured. The resistance was determined from the slope of the J-V curve at current = 0. The thickness of the device was estimated to be 70 nm and the device area is 0.0314 cm². The conductivity was then calculated from the following equation:

$$\frac{1}{\sigma} = \rho = R \frac{A}{l}$$

where σ is conductivity, ρ is resistivity, R is resistance, A is area of the device, and I is thickness of the device.



Figure 7: Conductivity of PbS QDs treated with MPA, EDT and Hybrid ligands

The conductivity of the PbS QD film treated by hybrid ligands lies between EDT and MPA-treated samples, as shown in Figure 7. This is a marked improvement over EDT despite the PbS-Hybrid containing 82% EDT ligands as inferred from the FTIR spectra. This confirms that the reduction in interdot distance resulted in an improvement to the conductivity of the film. However, the increase in conductivity falls short of PbS-MPA, indicating that the functional group on the QD surface plays an equally significant role in determining the conductivity. The conductivity trends well with the J_{SC} performance of the three devices, further confirming the impact of conductivity towards J_{SC}.

Mobility					
Treatment	(cm² V ⁻¹ s ⁻¹)	Reference			
MPA	0.00510	14			
EDT	0.00018	this work			
Hybrid	0.00120	this work			

Table 3: Mobility of PbS QDs treated with MPA, EDT and Hybrid ligands

To further identify the contribution towards improved conductivity, the mobility of PbS-Hybrid and PbS-EDT films were measured by fabricating a PbS thin film field effect transistor and the results are shown in Table 3. Similar to the conductivity trend, the mobility of PbS-Hybrid is in between PbS-MPA and PbS-EDT, suggesting that the mobility is likely the main contributor to the improvement in conductivity of the film.

There are possibly two contributions towards the improvement of conductivity and mobility of PbS-Hybrid sample. First is the reduction in the interdot distance compared to PbS-EDT sample due to the lack of dimerization of EDT. The shorter interdot distance can increase carrier tunnelling probability between QDs hence improving film's mobility and conductivity^{26,28}. Secondly, the mobility is also enhanced due to the presence of small amounts of the carboxylate functional group from the remaining MPA ligands as shown by the FTIR results. The mobility and conductivity of QDs capped with carboxylic acids ligands with similar ligand lengths have been previously reported to be superior to those capped by thiols ^{12,13}.

3.5 Ionisation energy

The improved J_{SC} of PbS-Hybrid can be further interpreted from the changes to the valence band edge of the p-type PbS films. The valence band edge of PbS-EDT, PbS-MPA and PbS-Hybrid is determined from the ionization energy of the film. The ionization energy was measured using photoelectron yield spectroscopy (PYS), which is suitable for quantum dots³⁰ and ultra-thin buffer layers³¹. This method is especially suitable for ligand exchanged p-type quantum dots with low mobility and thickness.

	lonisation energy
Sample	(eV)
PbS-EDT	4.94
PbS-MPA	4.87
PbS-Hybrid	4.85

Table 4: Ionisation energy measured using PYS

For the PbS-Hybrid film, the ionisation energy closely matches with that of MPA as shown in Table 4. This contrasts with the large reduction in MPA ligands shown in the FTIR results. The PbS-Hybrid film retains the ionisation energy of MPA with the presence of just a minority amount of MPA. When there are two species of ligands on the surface, likely additional shallower valence states from the MPA ligand contribute to the overall shallower ionisation energy of the PbS-Hybrid sample. The shallower ionisation energy of PbS-MPA and PbS-Hybrid prevents the backflow of holes into the absorber layer, thus improving the hole transport properties and reducing recombination. With a lower recombination, both PbS-MPA and PbS-Hybrid devices show enhanced J_{SC} over PbS-EDT.

Note that the ionisation energy of all species is about 0.2-0.3 eV shallower compared to the reported value measured by UV photoelectron spectroscopy (UPS) ³², perhaps due to the inherent charging problem for samples measured with UPS. However, both the measurements here and in literature show a similar trend of MPA having a shallower ionization potential³². Comparison between oleic acid and octadecyl thiol also show a similar 0.05 eV reduction in ionization energy³⁰.

4. Conclusion

This work demonstrates a novel method to produce closely packed p-type QD thin films with a short interdot distance of ~0.6 nm using the 2-step hybrid ligand treatment. This is the first observation of a 0.6 nm interdot distance with EDT as a dominant ligand, as confirmed by TEM imaging. EDT ligands have previously been only observed in its dimer form with an interdot distance of 1.2 nm. The 2-step hybrid ligand exchange works by using MPA as an intermediate ligand to prevent dimerization of the EDT ligands during the EDT ligand exchange step. This yields a film that has improved interdot coupling and conductivity. As shown in the J-V results, the devices based on PbS-Hybrid HTL have demonstrated significant improvement in J_{SC} and FF compared to that of the PbS-EDT device, while the V_{oc} and FF have improved compared to that of PbS-MPA devices. The best performing PbS-Hybrid device has achieved a PCE of 10.4% which demonstrates an improvement by 23% over PbS-MPA device (PCE of 8.4%) and 48% compared to PbS-EDT device (PCE of 7%).

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