Finding and Fixing Traps in II–VI and III–V Colloidal Quantum Dots: The Importance of Z-Type Ligand Passivation

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§ Supporting Information

ABSTRACT: Energy levels in the band gap arising from surface states can dominate the optical and electronic properties of semiconductor nanocrystal quantum dots (QDs). Recent theoretical work has predicted that such trap states in II–VI and III–V QDs arise only from two-coordinated anions on the QD surface, offering the hypothesis that Lewis acid (Z-type) ligands should be able to completely passivate these anionic trap states. In this work, we provide experimental support for this hypothesis by demonstrating that Z-type ligation is the primary cause of PL QY increase when passivating undercoordinated CdTe QDs with various metal salts. Optimized treatments with InCl₃ or CdCl₂ afford a near-unity (>90%) photoluminescence quantum yield (PL QY), whereas other metal halogen or carboxylate salts provide a smaller increase in PL QY as a result of weaker binding or steric repulsion. The addition of non-Lewis acidic ligands (amines, alkylammonium chlorides) systematically gives a much smaller but non-negligible increase in the PL QY. We discuss possible reasons for this result, which points toward a more complex and dynamic QD surface. Finally we show that Z-type metal halide ligand treatments also lead to a strong increase in the PL QY of CdSe, CdS, and InP QDs and can increase the efficiency of sintered CdTe solar cells. These results show that surface anions are the dominant source of trap states in II–VI and III–V QDs and that passivation with Lewis acidic Z-type ligands is a general strategy to fix these traps. Our work also provides a method to tune the PL QY of QD samples from nearly zero up to near-unity values, without the need to grow epitaxial shells.

INTRODUCTION

Colloidal quantum dots (QDs) have been the focus of a substantial body of research due to their size-dependent optoelectronic properties. States in the band gap that lead to localization of charge carriers (trapping) can strongly influence optoelectronic properties in these QDs and are typically attributed to the presence of undercoordinated surface atoms. Consequently, control over these trap states remains an outstanding challenge in the field. Until recently theoretical understanding of the exact chemical nature of surface traps was limited, and as a result, efforts to add or remove trap states in QDs have been largely empirical in nature. In this work, we experimentally test the latest theoretical models of trap formation in II–VI and III–V QDs. In doing so we verify the chemical nature of the traps affecting the photoluminescence quantum yield (PL QY) and develop a general strategy to reduce their density by ligand passivation. This approach affords core-only QDs with near-complete passivation and near-unity PL QYs.

Existing experimental studies on the control of trap states have focused on the effect of adding¹⁻¹⁰ or removing¹⁰⁻¹² organic ligands on the photoluminescence (PL) of II–VI metal chalcogenide QDs, which is highly sensitive to the presence of traps. Unfortunately, these and other studies in the literature do not offer a consistent description of the relationship between trap states and the chemistry of the bond formed between ligands and surface atoms. For example, it is known that amines increase PL QYs of CdSe QDs up to 15–20% at saturation coverage¹ and that Se-rich CdSe QDs exhibit PL QYs up to 50% with addition of phosphine ligands.¹ These results are usually rationalized by considering a simplified molecular orbital analysis of a Cd–Se bond, which predicts that lone-pair electrons on Se or empty orbitals on Cd (“dangling bonds”) will leave trap states in the band gap. While this predicts amines or phosphines can act as a Lewis base to fill the dangling bond on a Cd site, it is not clear why maximum PL QY values are higher for the Se-rich QDs with phosphine passivation compared to Cd-rich QDs with amine passivation.¹ Furthermore, a general understanding of the...
requirements for complete trap passivation is not available. Reports of core-only QDs with near 100% PL QY exist, e.g., by treatment of QDs with several ligands at once or via highly optimized syntheses, but the precise ligands and surface sites responsible for the lack of trap states remain ambiguous, and it is not clear if there is any generality between different QD materials or crystal structures. As a result, a general experimental strategy to tune trap densities in colloidal QDs is lacking.

Density functional theory (DFT) calculations have led to an improved understanding of the origin of surface traps in QDs. Recently, we have demonstrated that in undoped and charge-balanced nanocrystals only dicoordinated surface chalcogenide atoms should contribute to an electronic state within the band gap for zincblende II–VI nanocrystals. This argument is ultimately based on orbital symmetry and can be expanded to other tetracoordinated (e.g., wurtzite) semiconductor QDs such as the III–V family and invites the prediction that the only requirement to achieve a trap-free QD should be a surface free of dicoordinated anions.

The other important factor in understanding QD passivation is an accurate, atomistic description of the bond formed between a ligand and the QD surface. Some progress has been made by adoption of the covalent bond classification (CBC) method proposed by Green for metal–organic complexes. This formalism categorizes ligands into three classes based on the nature of the resulting bond formed to the QD surface site. Z-type ligands bind as a neutral two-electron acceptor (a Lewis acid) to an occupied lone pair on a surface anion site (a Lewis base). L-type ligands bind as neutral two-electron donors (Lewis bases) to unoccupied surface metal orbitals (Lewis acids). Finally, X-type ligands formally share one electron with a singly occupied orbital on a surface site (covalent bond). In the QD literature, anionic ligands (e.g., Cl⁻) bound to QD surfaces are often called X-type by considering a Cd²⁺–Cl⁻ bond where each atom contributes one electron. However, this assignment is not very intuitive and highlights that complications arise when applying the CBC method to bonds with significant ionic character.

Combined with the prediction from DFT calculations that only lone pairs on dicoordinated anion sites should lead to trap states, the CBC classification of ligands offers the hypothesis that Z-type ligands are the only class of ligand necessary to completely passivate trap states in QDs. Consequently only Z-type ligands should influence the PL QY, whereas X- and L-type ligands should not have any direct effect. There is already evidence for this in the literature in studies showing that the removal of Z-type ligands decreases the PL of QDs. Most notably Sanieyapour et al. recently concluded that the PL of CdSe QDs is much more dependent on certain Z-type binding sites than others, which supports the notion that only a fraction of surface chalcogenide sites (i.e., two-coordinated Se) are responsible for trap states. Furthermore, studies utilizing charged X-type ligands to transfer QDs from nonpolar to polar media do not result in increases of PL QY. However, it is not clear if there is any generality between different QD materials or crystal structures.

In this work we investigate the effects on PL of treatment of colloidal QDs with a wide range of Z-type ligands as well as X-type (chloride anions) and L-type (amines) ligands. We chose CdTe as a model system because it is a well-studied QD material in which we have earlier identified Te surface atoms as a source of traps, and it has a high potential for application in solution-processable photovoltaic devices. However, we show that our results can be generalized to other II–VI and III–V nanocrystals. We present a set of experiments demonstrating that trap states in CdTe nanocrystals can be almost completely removed by the binding of a wide range of metal halide or metal carboxylate Z-type ligands. Our results point toward the presence of a single binding energy between Z-type ligands and PL QY-active surface sites, hence suggesting that a single Te trap site is responsible for the PL QY. However, we find that ligands outside the Z-type classification such as amines and alkylammonium chlorides can also result in small but significant PL increases. This shows that the QD surface is complex and dynamic, and we discuss the implications of this result for the hypothesis that only dicoordinated anions can form trap states.

## METHODS

**Materials.** All anhydrous materials and QDs were stored and handled in a nitrogen-filled glovebox atmosphere and/or using air-free Schlenk line techniques. Cadmium(II) oxide (CdO, anhydrous, 99.5%), tellurium (Te powder, 99.8%), tetradecapephosphonic acid (TDA, 97%), palmitic acid (≥99%), trioctylphosphine (TOP, anhydrous, 97%), butylamine (99.5%), octylamine (99%) sodium caprylate (≥99%), sodium palmitate (≥98.5%), cadmium chloride (CdCl₂, anhydrous, 99.99%), indium chloride (InCl₃, anhydrous, 99.99%), zinc chloride (ZnCl₂, anhydrous, 99.99%), zinc bromide (ZnBr₂, anhydrous, 99.99%), zinc iodide (ZnI₂, anhydrous, 99.99%), magnesium chloride (MgCl₂, anhydrous, 99.99%), lithium chloride (LiCl, anhydrous, ≥99%), gallium(III) chloride (GaCl₃, anhydrous, 99.99%), aluminum chloride (AlCl₃, anhydrous, 99.99%), zinc acetate (Zn(OAc)₂, 99.99%), indium acetate (In(OAc)₃, anhydrous, 99.99%), cadmium acetate (Cd(OAc)₂, anhydrous, 99.99%), lead chloride (PbCl₂, anhydrous, 99.99%), gold chloride (AuCl₃, 99%), platinum chloride (PtCl₄, ≥99.9%), and palladium chloride (PdCl₂, ≥99.9%) were purchased from Sigma-Aldrich and used without further purification. Oleic acid (OA, ≥95%, Sigma-Aldrich), octadecene (ODE, 90%, Sigma-Aldrich), and oleylamine (OAM, 90%, Acros Organics) were degassed at 100 °C for 1 h before storage in a nitrogen glovebox. All solvents (toluene, methanol, hexane, acetone, and methyl acetate) were purchased anhydrous from Sigma-Aldrich and used without further puriﬁcation. Rhodamine 6G and rhodamine 101 reference dyes were obtained from Lambda Physik GmbH.

**Preparation of TOP-Te Precursor.** A 1 M solution of TOP–Te in TOP was prepared by heating 2.553 g of Te in 20 mL of TOP to ∼150–200 °C in a N₂-filled glovebox until a clear yellow liquid was obtained. In a separate vial, 0.2 mL of this 1 M TOP–Te solution was diluted by addition of 0.8 mL of TOP and 1.0 mL of ODE, giving a final Te concentration of 0.1 M.

**Synthesis of CdTe NCs.** Cadmium telluride nanocrystals were synthesized according to a protocol described by Wang et al. All synthesis and washing steps were carried out under an inert atmosphere and with anhydrous solvents. Briefly, CdO (25.6 mg, 0.2 mmol), tetradecapephosphonic acid (140 mg, 0.5 mmol), and ODE (4 mL) were loaded into a three-neck round-bottom flask and attached to a Schlenk line. Water and oxygen were removed by heating the flask to 100 °C under vacuum (<1 mbar) for 1 h. The solution was heated to 315 °C until a clear Cd–TDA complex formed, then cooled to 290 °C, at which point 2 mL of a 0.1 M solution of TOP–Te in TOP and ODE was swiftly injected. After a growth time of 2–15 min depending on the desired QD size, the
CdCl₂ and InCl₃ (630 L or as number of ligands added per nm² of total CdTe surface area) ligand concentration. Ligand concentrations can be expressed as mol/L or as number of ligands added per nm² of total CdTe surface area. Preheated aluminum heating block on a hot plate set to 95 °C in toluene. In samples with higher MX⁻ ligand concentrations, a white precipitate appeared during cooling due to the insolubility of the excess MX⁻ ligand in toluene. Once cooled, the samples were passed through a 0.2 μm PTFE syringe filter to remove these precipitates.

Optical Characterization. Optical characterization was conducted using gastight cuvettes loaded in a N₂-filled glovebox. Absorbance measurements were acquired using a PerkinElmer Lambda 1050 or Lambda 40 absorbance spectrometer. Fluorescence spectra were acquired using an Edinburgh Instruments FLS980 spectrometer. Photoluminescence quantum yields were collected on an Edinburgh Instruments Lifespec TCSPC setup with a 400 nm pulsed laser diode excitation (<1 ns instrument response time).

Electron Microscopy and Elemental Analysis. Transmission electron microscopy (TEM) images, electron diffractionograms, and energy-dispersive X-ray (EDX) spectra were acquired using a JEOL JEM1400 transmission electron microscope operating at 120 keV with a built-in EDX detector. Prior to sample deposition onto grids for TEM and EDX measurements, QDs treated with various MX⁻—amine complexes were washed twice with methyl acetate and resuspended in toluene to remove any traces of unreacted ligand. Reported EDX values are averages of 3–6 measurements taken on different areas of each sample.

RESULTS AND DISCUSSION

Treatment of CdTe QDs with MX⁻, Lewis Acids. CdTe QDs capped with tetradecylphosphonic acid were synthesized and washed as described by Wang et al. Absorption spectra, PL spectra, and PL transients of the washed QDs are shown in Figure 1 (gray traces). X-ray and electron diffraction determined the QDs to have a zincblende crystal structure (see SI, Figure S1). After washing three times the PL QY was 8%, which indicates very incomplete surface passivation.

In order to test the hypothesis that Z-type ligands can remove all trap states by passivating dicoordinated Te (Te-2c) surface sites, the washed QDs were then treated with a Lewis acidic MX⁺ complex (where M is a Lewis acidic metal site and X⁻ is a singly negative anion) using a modification of the protocol developed by Page et al. for CdCl₂ passivation of CdTe. Each MX⁺ ligand was first dissolved at 95 °C in toluene with 9 equiv of OAM to form a clear MX⁺—OAM acid—base adduct. The OAM confers solubility in nonpolar solvents, which dissolve QDs coated in hydrophobic ligands and impede the formation of free X⁻ anions, which can bind to Cd sites. The MX⁺—OAM complex, for brevity referred to hereafter as MXO, was then added to the CdTe QDs in toluene to give a desired ligand concentration. In this work, we typically express the ligand concentration as ligands added per nm² of total CdTe surface area, as this facilitates comparison of results between QDs of different sizes (see supplementary methods for details of this calculation), but we stress this is not the bound ligand density, which is expected to be at most around 3 MX⁻ per nm². The mixture of CdTe QDs and MXO ligand was heated at 95 °C for 15 min in a nitrogen-filled glovebox and afterward allowed to cool to room temperature and filtered to remove excess ligand. We find that this procedure results in a highly reproducible increase of the PL QY of up to 90% for InCl₃ treatment (sample-to-sample standard deviation of 3.7% and estimated 5% systematic error).

Figure 1. Optical characterization of CdTe QDs (3.8 nm diameter, 0.456 μmol/L in toluene) before and after treatment at 95 °C for 15 min with CdCl₂ and InCl₃ (630 μmol/L, equivalent to 30 ligands added per nm² CdTe surface area) with 9 equiv of OAM (5.67 mmol/L). An OAM control experiment (6.3 mmol/L, 300 ligands/nm²) is also shown; it overlaps with the original spectrum. (A) Absorption spectra and (B) PL spectra of CdTe QDs before and after treatment with each ligand. PL QY values determined via reference dye method are shown in brackets next to the PL trace of each sample. Inset of (B) shows photographs of CdTe QD solutions under UV illumination before and after treatment with CdCl₂ and InCl₃. (C) Time-resolved PL traces of the same samples (60 s acquisition time).
of 0.456 μmol/L. A red-shift of the absorbance and PL is observed. The treatment does not alter the width of the first absorption peak of the QDs, indicating no significant change in QD polydispersity occurred. In addition, we observe changes in the absorption spectrum for higher transitions and a small increase of absorption at the first exciton peak, all consistent with a slight increase in the effective nanocrystal diameter due to the Z-type binding of MXn to undercoordinated Te sites on the nanocrystal surface.

Treatment of the CdTe QDs with InCl3 and CdCl2 resulted in a PL QY increase from 8% to 90% and 73%, respectively, clearly evident from the photo taken under UV light in the inset of Figure 1B. The PL lifetimes become significantly longer after treatment and can be fitted with a single-exponential function (lifetime 25.0 ns; see SI). This is the result of the loss of fast nonradiative decay components, indicating reduced trapping at surface defects.

In order to investigate the effects from the 9 equiv of OAM (i.e., equivalent to 270 lig/nm²) used to solubilize the MXn ligands, we performed a control treatment with only oleylamine at 300 added ligands/nm² (black traces in Figure 1A, B, and C). There was no appreciable change in the QD absorbance, but a marginal increase of the PL QY to 15% was observed and the PL lifetime became slightly longer. Although this result rules out OAM as the cause of near-unity PL QY for CdCl2 and InCl3, oleylamine is nonetheless able to afford a modest PL increase, despite being unable to bind to undercoordinated Te.

Figure 2. PL QY of CdTe QDs (0.456 μmol/L) after treatment with various MXn-type ligands for 15 min. (A) Treatment with metal chloride ligands. Each data set utilizes different QD diameters and equivalents of OAM (per MXn) as shown in the legend. Treatment temperature and added ligand concentration: 80 °C, 60 lig/nm² (sets 1 and 3); 95 °C, 30 lig/nm² (set 2); 95 °C, 14 lig/nm² (set 4). (B) Effect of varying anion of MXn ligands: X = iodide (I⁻), bromide (Br⁻), chloride (Cl⁻), acetate (H₃C₂O₂⁻), caprylate (H₁₅C₈O₂⁻), and palmitate (H₃₁C₁₆O₂⁻). Treatment conditions: 4.1 nm QDs, 60 lig/nm² (20 lig/nm² for InCl3), 70–85 °C, 9 equiv of OAM, 15 min. (C) PL QY of samples treated with various MXn ligands after each of two washing steps (precipitation with methyl acetate and resuspension in toluene). Lower PL QY values in panels B and C are due to a smaller treatment volume with suboptimal temperature control (75–80 °C).

Optimal Treatment Conditions for Near-Unity PL QY.
In general, we observed higher PL QYS after treatment with MXn at higher temperatures (SI, Figure S2A,B), although above 100 °C more significant changes in the QD absorbance indicative of Ostwald ripening were observed. Treatment with CdCl2 at 120 °C for 15 min (in ODE) gave 89% PL QY, whereas at room temperature the PL QY increased from 8% to 15% over a period of 1 h (SI, Figure S2C). This suggests that an activation barrier must be overcome for the MXn ligand to attach to the nanocrystal surface. We tentatively attribute this to the need to dissociate the MXn−OAM Lewis acid/base adduct to form a new Te−MXn adduct or the rearrangement of bulky Cd−TDPA ligands from the synthesis to achieve full surface coverage. From our experiments we concluded a treatment temperature of 95 °C offers the best balance between increasing the PL QY while avoiding ripening or agglomeration of QDs, but note that each ligand has slightly different optimal conditions. True optimization was only performed for CdCl2 treatment of CdTe QDs, and hence we believe treatments with other ligands and of other QDs discussed below may be optimized further.

Treatment with Other MXn Ligands. We found that many other metal chloride treatments result in a significant increase of the PL QY of CdTe QDs under appropriate conditions (Figure 2A). We achieved the highest PL QY of 90% with InCl3, and ZnCl2 gave comparable results to CdCl2. Other divalent (MgCl2), trivalent (GaCl3, AlCl3), and univalent (LiCl) metal halides gave significant PL QY increases up to 60%. An increase in PL QY was also observed for...
treatment with metal bromides, iodides, and carboxylates, with a general trend toward higher PL QY with shorter carboxylate carbon chains or smaller halide anions (Figure 2B). Noble metal chlorides (e.g., AgCl, AuCl, PdCl) were not compatible with this reaction system, as they were reduced by amines or the CdTe QDs themselves to give metal flakes or metal nanocrystals (SI, Figure S3C–H). Addition of lead chloride (PbCl2) induced cation exchange to give PbTe QDs (SI, Figure S3B and Zhang et al.45).

We found that the amount of amine used to dissolve the $\text{MX}_n$ ligand can be varied between 3 and 9 equiv with no change in the PL QY observed for $\text{MX}_n$ treatment (Figure 2A and SIFigure S9). In addition CdTe QDs of varying diameters between 3.6 and 4.6 nm were used (Figure 2A) with very similar results, showing that the Z-type passivation is a general surface effect that can result in near-unity PL QYs independent of the QD size.

These results show clearly that the PL increase is not particular for CdCl2, which was already known to enhance the PL QY of CdTe QDs and is also indispensable in the fabrication of high efficiency in CdTe solar cells, but indeed is general across all $\text{MX}_n$-type Lewis acids. In fact, we observe that under identical conditions the addition of InCl3 increases the PL QY further than CdCl2, and small metal carboxylates such as Cd, Zn, and In acetate are almost as effective as metal chlorides.

The trends in the PL QY data in Figure 2B can be explained through steric hindrance, with smaller $\text{MX}_n$ complexes affording greater surface coverage and longer ligand chains potentially having limited accessibility to some surface sites. However, the PL QY will also be influenced by the equilibrium between bound and unbound ligand, which will be determined by the solubility of the ligand–OAM complex in the reaction solvent (toluene) and the binding affinity of the ligand. Indeed the trend toward lower PL QY with longer metal carboxylate ligands and larger, less ionic halogen salts can be understood in terms of increased solubility of the $\text{MX}_n$ complex,46 pushing the equilibrium toward ligand in solution. $\text{MX}_n$–butylamine complexes also gave rise to greater PL QY values than $\text{MX}_n$–OAM ones (see SI, Figure S4), consistent with the steric hindrance and solubility arguments. We explore the effect of the ligand binding affinity and equilibrium constant in more detail below.

To test the possibility that X-type binding of anions might account for these observed increases the PL QY of CdTe, we treated the CdTe QDs with tetrabutylammonium chloride (TBACl) dissolved in toluene with 9 equiv of OAM. TBACl has been reported as a source of X-type chloride ligands,27 and the bulky alkylammonium cation lacks a Lewis acidic site to bind as a Z-type ligand to surface Te. Interestingly, TBACl treatment resulted in a reproducible PL QY increase to 30% at 14−30 lig/nm2 irrespective of whether 3 or 9 equiv of OAM was employed (Figure 2A; for full optical characterization of

Table 1. EDX Elemental Analysis and Cd/Te Ratios for Samples Treated with Various Ligand–OAM Complexes

| element (line) | atom % (average) ($^a$) |
|---------------|------------------------|
|               | untreated OAM CdCl2 ZnCl2 InCl3 Zn(acetate)2 TBACl |
| Cd ($\alpha_1$) | 56.0 55.6 59.6 32.3 33.7 46.4 50.6 |
| Te ($\alpha_1$) | 44.0 44.4 29.3 25.6 28.1 37.7 40.4 |
| Zn ($\alpha_1$) | – – – 30.7 – 15.9 – |
| In ($\alpha_1$) | – – – 11.2 11.4 38.2 – 8.9 |
| Cl ($\alpha_1$) | – – – – – 11.2 11.4 38.2 – 8.9 |
| total (%) | 100.0 100.0 100.1 100.0 100.0 100.0 100.0 100.0 |
| Cd/Te | 1.27 1.25 2.04 1.26 1.20 1.23 1.23 1.25 |

$^a$Values are averages from 3 to 6 measurements. $^*\text{Indium content was difficult to quantify due to overlap with Te L} \alpha_1$ and Cd L$\beta_1$ signals so was omitted from quantitative analysis. $^\sim$ indicates element not detected and so not included in the analysis.

Figure 3. (A) TEM micrographs of CdTe QDs treated with various ligands. (B) Histograms of QD diameters taken from TEM micrographs.
TBACl treatment see SI, Figure S5). This PL increase is significantly greater than that of OAM-only-treated samples even at high OAM concentrations (300 lig/nm$^2$). This somewhat surprising observation will be discussed further below.

**QD Morphology and Composition.** We investigated changes in QD morphology and composition during the treatment, as it is known that chloride salts facilitate grain growth of CdTe crystals.$^{35,36,47}$ Samples were washed two times after ligand treatment to remove excess unbound ligands and studied using TEM and EDX. We note that the PL of CdCl$_2$- and InCl$_3$-treated samples was unaltered after each washing step (see Figure 2C), suggesting that these ligands were tightly bound to the surface. ZnCl$_2$-, Zn(acetate)$_2$-, and TBACl-treated samples exhibited PL decreases during washing, suggesting loss of ligand in these samples.

The results in Figure 3 show that the QDs do not undergo any observable change in shape or diameter after treatment with a variety of ligands. EDX results in Table 1 show that all but one sample gave a Cd/Te ratio of about 1.25, which falls into a typical range for EDX measurements on II–VI QDs giving cation/anion ratios of 1.2–1.5.$^{7,48,49}$ The only exception is the CdCl$_2$-treated sample with a Cd/Te ratio of 2.04. As expected, zinc was observed in samples treated with ZnCl$_2$ and Zn(acetate)$_2$. Indium was also observed in samples treated with InCl$_3$ (Table 1), although quantification was not possible due to overlap with Te L$_4$ and Cd L$_{β1}$ spectral lines. Chlorine was measured in all samples treated with a chloride salt. We note that the Zn/Cl ratio does not match the expected 2:1, and the amount of Zn or extra Cd present in the ZnCl$_2$- or CdCl$_2$-treated samples (respectively) is much higher than would be expected for a monolayer on the QD surface. These discrepancies may reflect the lack of sensitivity toward lighter elements in EDX measurements or excess MX$_n$ ligand loosely attached to the QDs. XPS measurements corroborate the presence of metal and chloride ions in each sample (see SI, Figure S6).

These TEM and EDX results show that the QDs are Cd-rich prior to treatment, probably due to Cd-phosphonate ligands on the surface, and the addition of MX$_n$ ligands does not affect this ratio unless CdX$_2$ is used. The increase in Cd, Zn, or In content for CdX$_2$, ZnX$_2$, and InX$_3$ addition, respectively, confirms the addition of entire MX$_n$ complexes onto the QD surface, rather than just the X$^-$ anion. Comparing Zn(acetate)$_2$ and ZnCl$_2$, the higher PL QY of ZnCl$_2$-treated sample (see Figure 2B) correlates with a higher Zn content measured with EDX (Table 1), suggesting that higher PL QYs are correlated with higher Z-type ligand surface coverage.

**Effect of Ligand Concentration.** To further investigate the binding of each type of ligand to QD surfaces, we performed treatments with a series of increasing added ligand concentrations for CdCl$_2$, InCl$_3$, TBACl, and OAM. To avoid QD dilution each data point is taken from a separate reaction with a fixed QD concentration of 0.456 μmol/L and varying ligand concentration. In Figure 4A the PL QY is plotted as a function of added ligand concentration per nm$^2$ of total CdTe surface area. Top axis shows the final ligand concentration in the solution. Lines are fits using eq 7 (see text). (B) Photographs under ambient room light (top) and UV light (bottom) showing effect of CdCl$_2$ treatment at increasing CdCl$_2$ concentrations (in nm$^{-2}$) on QD PL. (C) PL spectra of CdCl$_2$-treated QDs at increasing amounts of ligand added from 0.1 to 100 lig/nm$^2$. Dashed gray line shows position of PL peak before treatment. (D) PL decay curves of CdCl$_2$-treated QDs as a function of ligand concentration.

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**Figure 4.** Effect of ligand concentration on CdTe QDs (3.8 nm, 0.456 μmol/L in toluene) after treatment at 95 °C for 15 min with CdCl$_2$, InCl$_3$, TBACl (9 equiv of OAM), and OAM only ligand. (A) PL QY of CdTe QDs after treatment with ligands as a function of concentration of added ligand per nm$^2$ of total CdTe surface area. Top axis shows the final ligand concentration in the solution. Lines are fits using eq 7 (see text). (B) Photographs under ambient room light (top) and UV light (bottom) showing effect of CdCl$_2$ treatment at increasing CdCl$_2$ concentrations (in nm$^{-2}$) on QD PL. (C) PL spectra of CdCl$_2$-treated QDs at increasing amounts of ligand added from 0.1 to 100 lig/nm$^2$. Dashed gray line shows position of PL peak before treatment. (D) PL decay curves of CdCl$_2$-treated QDs as a function of ligand concentration.
Figure 4A clearly shows that the PL QY increases with concentration for all MX$_2$, TBACI, and OAM ligands, but that the highest obtained PL QY (at very high ligand concentrations), as well as the functional dependence on the ligand concentration, is markedly different. The CdCl$_2$- and InCl$_3$-treated samples can be tuned from very low PL QY (under 3% increase more rapidly than [MX$_2$]$_m$ concentration, is markedly different. The equilibrium for ligands and takes into account the possibility that some amine (l equivalents) is not freed into solution after MX$_2$, binds to the surface. The value of m, the number of amines coordinated to the Lewis acidic ligand in solution, has been shown to be 2 for primary amines bound to Cd-carboxylate Lewis acids.$^{10}$ The equilibrium constant $K$ for the ligand binding can be expressed in terms of the fractional ligand surface coverage, $\theta$:

$$K = \frac{[MX,L_m][L]^n}{[MX,L_m][S]} = \frac{[L]^{n-1} \theta}{[MX,L_m] (1 - \theta)}$$

(2)

If more ligand is added, $[MX,L_m]$ increases, but due to the excess of amine utilized in these experiments (9 equiv in Figure 4A), [L] will also increase. If $m - l > 1$, then $[L]^{n-1}$ will increase more rapidly than $[MX,L_m]$, so $\theta$ should decrease in eq 2 to maintain constant $K$. This situation corresponds to a net removal of Z-type ligands upon addition of MX$_2$. Our data however clearly show that the PL QY (and therefore $\theta$) increases with added ligand, meaning that $m - l < 1$. In fact, we found that the PLQY vs concentration series for CdCl$_2$ treatment in Figure 4A is identical with 3 or 9 equiv of amine (SI, Figure S9). Therefore, we conclude that the concentration of amine does not affect the binding of MX$_2$ ligand to CdTe, i.e., $m - l = 0$. Effectively oleylamine is removed from the equilibrium, either through L-type binding to surface Cd or remaining bound to the newly added MX$_2$ ligand. Additionally, amines can hydrogen bond to phosphonate ligands on CdTe QD surfaces,$^{16}$ and indeed $^{31}$P and $^1$H NMR experiments we conducted show that oleylamine forms adducts with native Cd-phosphonate ligands after treatment (see SI, Figure S10 and Supplementary Note 1).

We note that this expression only accounts for the effects of a single ligand on the PL QY. In general, the rate of PL QY will be governed by the rate equation

$$\frac{dN_{eh}}{dt} = -c_{trap} N_{trap} N_{eh} - k_{rad} N_{eh}$$

(4)

where $N_{trap}$ is the concentration of traps and $k_{rad}$ is the radiative recombination rate. Taking the trapping rate and concentration of traps to be time-independent quantities, $c_{trap}$ is a constant and eq 4 becomes a quasi-first-order rate equation.

$$\frac{dN_{eh}}{dt} = -k_{trap} N_{trap} N_{eh} = k_{rad} N_{eh}$$

(5)

We note that this expression offers an explanation for the deviation of a QD ensemble PL lifetime from a single exponential, in that the surface coverage and hence observed radiative rate may vary between QDs. After combining the constants $c_{trap}$ and $N_{trap}$ into a single constant $c_{trap}$ the PL QY will then be given by

$$\Phi = \frac{k_{rad}}{c_{trap}(1 - B\theta) + k_{rad}} = \frac{1}{c_{trap}(1 - B\theta) + 1}$$

(6)

Finally we combine eqs 3 and 6 to obtain an expression for the PL QY in terms of the concentration of added ligand $[MX,L_m]$:

$$\Phi = \frac{c_{trap}}{c_{trap}(1 - B\theta)} + 1$$

(7)

Table 2. Fit Values Obtained by Fitting Eq 7 to the Data in Figure 4A

| ligand   | ligand type (CBC) | $K$ (L mol$^{-1}$) | $B$ | $\Phi_{\text{max}}$ | $\Delta G$ (kJ/mol) |
|----------|-------------------|-------------------|-----|---------------------|---------------------|
| CdCl$_2$ | Z                 | $[(7.78 \pm 0.37) \times 10^6$ | 0.99967 & 0.00024 | 0.83 | $\pm 48.5 \pm 0.2$ |
| InCl$_3$ | Z                 | $[(7.97 \pm 1.06) \times 10^6$ | 1.0 & 0.000075 | 1.0 | $\pm 49.2 \pm 0.3$ |
| TBACI    | X                 | $[(1.24 \pm 0.08) \times 10^7$ | 0.9961 & 0.00013 | 0.29 | $\pm 50.0 \pm 0.2$ |
| OAM      | L                 | $[(6.18 \pm 1.04) \times 10^6$ | 0.9944 & 0.00045 | 0.22 | $\pm 33.8 \pm 0.5$ |

**Saturation PL QY predicted by the fit.**
We used eq 7 to simulate the dependence of the PL QY on added ligand and found reasonable behavior for physically relevant parameters (see Supplementary Note 2 and Figure S11). We were able to measure the value of \( \frac{c_{\text{app}}}{k_{\text{rad}}} \) by measuring the PL lifetime of a CdTe QD sample with near-unity PL QY (\( \theta \approx 1; k_{\text{rad}} \approx k_{\text{sen}} = 25 \text{ ns} \)) and the average trapping lifetime of a sample with near-zero PL QY using ultrafast transient absorption spectroscopy (\( \theta \approx 0; c_{\text{app}} \approx k_{\text{sen}} = 39.4 \text{ ps} \)), giving a ratio of \( \frac{c_{\text{app}}}{k_{\text{rad}}} \approx 634.5 \) (see Supplementary Note 3 and Figure S12). Holding this ratio constant, we fit eq 7 to the data in Figure 4A (solid lines) to obtain values for fit parameters \( B \) and \( K \) (in \( \text{L mol}^{-1} \)) reported in Table 2. The free energy difference \( \Delta G \) between bound and unbound ligand was also computed from the relation \( \Delta G = -RT \ln K \).

CdCl\(_2\) and InCl\(_3\) give similar \( \Delta G \) values, suggesting similar binding strengths, and TBACl is moderately higher but similar in magnitude. OAM exhibits far lower binding affinity, in line with earlier reports of weak binding of amines to II–VI QDs.\(^{51}\) Our reported \( \Delta G \) values suggest a very high affinity of MCI\(_3\) ligands to the QD surface, but are not unprecedented in the literature. Munro et al.\(^ {31} \) report \( \Delta G \) values of approximately \(-50 \text{ kJ/mol for octadecanethiol binding to CdSe using a fitting model that relates } \theta \text{ to PL QY via the number of binding sites, but they note that their value of } K \text{ is not unique, as they must also fit the number of binding sites. Bullen et al.}\(^ {6} \) assumed a linear relationship between \( \theta \) and PL QY and report much lower \( \Delta G \) values of \(-24 \text{ kJ/mol (decylamine) and } -26 \text{ kJ/mol (octanethiol).} \) Clearly the model used to relate \( \theta \) and PL QY has a large influence on the estimation of \( \Delta G \) values. Other studies have utilized quantitative NMR to more directly measure \( \theta \) when stripping Cd–carboxylate ligands from CdSe using amines and report \( \Delta G \) values of cadmium carboxylate ligand binding from the amine complex in solution between \(+13.5 \text{ and } -17 \text{ kJ/mol depending on the NC size, binding site, and the amine used.}^{10,11} \) It is instructive to compare these and our binding affinities with those reported for dissolved MX\(_2\) binding to Lewis bases in solution, which are directly measured using NMR or absorbance spectroscopy; for example, for ZnCl\(_2\) in diethyl ether \( \Delta G \) ranges from \(+4.2 \text{ kJ/mol (binding to } 2\text{-methyl-4-nitroaniline)} \text{ to } -20.3 \text{ kJ/mol (4-methoxybenzamide).}^{42} \) The most relevant interaction to compare these molecular \( \Delta G \) values to is likely the QD–oleylamine binding affinity, for which we obtain a \( \Delta G \) of \(-33.8 \text{ kJ/mol.}\)

From these comparisons we tentatively conclude that our model is overestimating the binding affinity of ligands, which can be due to a number of factors. First of all, our assumed proportionality between ligand coverage and passivated trap densities (eq S) is a continuum model that reports only an average ligand coverage for the ensemble; the high value of \( \frac{c_{\text{app}}}{k_{\text{rad}}} \) means that the PL QY is very sensitive to the ligand coverage around \( \theta = 1 \). Thus, only a small drop in \( B \) (the maximum fraction of passivated traps) from 1 to 0.995 is required to lower the saturation PL QY value to 20% (see SI, Figure S11). We expect that a discretized binding model that accounts for the distribution of bound and unbound ligands would better model the system, especially at high ligand concentrations, and provide more accurate binding affinities. We also note that our values are sensitive to the measured \( \frac{c_{\text{app}}}{k_{\text{rad}}} \) value, so improved the understanding of these rates will also afford greater accuracy. Finally, the validity of assuming Langmuir binding, especially with a single binding energy, is potentially questionable for QDs.\(^ {52} \) Nonetheless, the model presented here is based on physical assumptions and measurable recombination rates and gives unique fits, which will assist with comparing results across QD samples with different nonradiative and radiative rates.

We also note that our data fits well to a single binding site model. This is particularly important in light of recent studies that have shown that there are at least two distinct ligand binding affinities for Z-type ligands to CdSe QDs arising from the heterogeneity in QD surface sites.\(^ {10,11} \) Saniepay et al.\(^ {11} \) further argued that the binding site with the highest affinity for ligands (largest \(- \Delta G \) value) dominated the changes in PL when removing ligands. Because we measure the PL QY to infer the ligand binding equilibrium, we only observe the equilibrium linked to PL-sensitive surface sites. The good fits we obtain with a model assuming only one binding affinity therefore support the notion that the large majority of traps responsible for PL QY losses are associated with a single Lewis basic ligand binding site. On the other hand, the ability of non-Lewis acidic ligands (OAM, TBACl) to increase PL shows that a holistic explanation of the PL QY is not necessarily achieved through a single binding site model.

**Relationship between Binding Mechanism and PL.** We have demonstrated that MX\(_2\)-type Lewis acids significantly increase the PL QY of CdTe QDs up to near unity values without any observable physical change to the QDs and have shown that both the metals and anions are attached to the resulting QDs. This does not however constitute direct evidence of Z-type binding; for instance, the presence of the metal in elemental analysis can be explained as X-type binding of the anion to Cd with the metal cation providing charge compensation, i.e., a bound ion pair.\(^ {24,53} \) We can, however, present several counter-arguments against X-type binding being the cause of near-unity PL QY.

First, the large difference in maximum PL QY between MX\(_n\) (Lewis acid) and TBACl (no Lewis acidity) ligands supports the notion that the ability of a ligand to bind as a Lewis acid (Z-type) is most important in achieving near-unity PL QYs. Indeed, we found that samples treated with both CdCl\(_2\) and TBACl gave the same PL QY as a CdCl\(_2\)-only-treated sample. This highlights that MX\(_n\) ligands are sufficient for achieving complete trap passivation. Second, we observed a higher PL QY for ZnX\(_2\) ligands with a higher measured Zn content in the EDX data (Figure 2B and Table 1), so PL QY correlates with the presence of Lewis acidic metal sites. Finally, for ZnCl\(_2\)-treated CdTe we observed a weak signal in the nonresonant Raman spectrum at 275 cm\(^{-1}\), which may be attributable to a Zn–Cl vibration (250 cm\(^{-1}\) in ZnCl\(_2\) powder) arising from ZnCl\(_2\) bound to Te (see Supplementary Note 4 and Figure S13). Therefore, we conclude that our results provide a strong argument that the majority of the PL QY increase achieved by various MX\(_n\) ligands tested is indeed due to the metal binding as a Z-type Lewis acid to undercoordinated Te sites.

However, the observations that OAM and TBACl can increase the PLQY to \( \sim 20–30\% \) while exhibiting markedly different affinities for the QD surface than CdCl\(_2\) or InCl\(_3\) need to be addressed. If we assume that TBACl can only bind to Cd as an X-type ligand and OAM as an L-type, then these results can be considered evidence against the hypothesis that only two-coordinated Te can contribute trap states to a QD. We propose four possibilities: (i) There are surface Cd sites that create trap states not predicted by our recent DFT calculations.\(^ {20} \) While we did not identify such sites, this does not disqualify their existence. One candidate is Cd–Cd dimers: it has been shown for PbS NCs that free conduction band
electrons can become trapped by populating the bonding orbital of a dynamically formed Pb–Pb dimer, which lies within the band gap. A similar process might be possible for CdTe. Alternatively, for relatively large NCs facet-specific Cd surface sites, such as edge sites or adatoms, could present states in the band gap. These would not easily be found in DFT calculations on moderately sized QD model systems. However, our results show that if Cd traps exist, they must contribute less to nonradiative recombination than Te traps and are removed by the addition of MX ligands. (ii) Some surface reconstruction follows binding of Cl or other ligands, resulting in a reduction of two-coordinated Te sites. We observed no sign of morphological changes in our treatment experiments, but this does not rule out reconstruction of the QD surface. (iii) TBACl and OAM are able to passivate Te sites in a manner not well described by the covalent bond classification method. One can imagine that the tetrabutyrammonium ion complexes to Te-2c surface sites. However, we have performed DFT calculations of this situation that show the TBA cation is too bulky to influence the Te trap state enough to remove it from the band gap (see SI, Figure S14). Finally, (iv) TBACl and OAM could bind to the QD and prevent diffusion of Z-type ligands across the QD surface. We have observed in molecular dynamics simulations that Z-type ligands diffuse over the QD surface. This implies that Te traps are effectively passivated only part of the time. If the density of ligands on the surface increases, the diffusion of Z-type ligands is hindered, leading to an increased time-averaged surface coverage of two-coordinated Te sites.

Which scenario, or combination of scenarios, is responsible for the increase of the PL QY with increased concentration of amines or TBACl remains unclear. What is clear is that the QD surface is a complex and dynamic system. Overall we conclude that Lewis acidic (Z-type) ligand passivation is able to remove the vast majority of surface traps responsible for the low PL QY and that our hypothesis that Te-2c surface sites are the only source of traps for CdTe QDs is mostly verified. At the same time, secondary effects are still present on the QD surface that remain not completely understood and that are not easily captured in a simple picture of passivating a single type of trap with suitable coordinating ligands.

**Generalizability of Results to Other QDs and Devices.**

To test how general the results discussed in this work are, we performed the same treatment with CdCl₂ on other types of II–VI QDs and III–V QDs. Figure 5 shows the results of this treatment for CdSe, CdS, and In(Zn)P QDs. We observed a PL QY increase from 0.4% (before treatment) to 11.5% (after treatment) for CdSe QDs, whereas a control experiment with OAM only increased the PL to 0.9% (Figure 5B). A PL QY enhancement was also observed for InP and InZnP QDs (Figure 5A), most notably from 0.4% to 18% for InZnP QDs. For the InZnP QDs, ligand treatment with X-type ligand (TBACl) and L-type ligand (OAM) gave no increase in the PL QY. The band-edge PL of CdS was also increased by a factor of 4 (Figure 5A), although a broad sub-band-gap PL was also enhanced (SI, Figure S15). We also observed large PL increases when treating CdTe tetrapods featuring a predominantly wurtzite crystal structure (SI, Figure S16). This demonstrates that the that MX₄ treatment also works for passivating undercoordinated anions on wurtzite surfaces. Achieving unity PL QY may require optimization of ligand treatment conditions for each QD material and in some cases could be hindered by the presence of defects not related to the surface. However, these results suggest that Lewis acidic (Z-type) ligands are in general far more efficient than non-Lewis acidic ligands at passivating surface traps in II–VI and III–V QDs and therefore at increasing the PL QY.

Finally we stress that understanding the surface of semiconductor nanocrystals is not only of use to enhance the PL QY. Surface state passivation in general is an important scientific and technological challenge with particular relevance for semiconductor devices such as LEDs, lasers, and solar cells. To show the relation between surface state passivation on QDs and the passivation in bulk semiconductor devices, we have constructed sintered CdTe/ZnO heterojunction solar cells from CdTe QDs that were “pretreated” with CdCl₂–octylamine (>100 lig/nm², 95 °C, 9 equiv of amine) after synthesis (Figure 5C, blue solid lines) and from control QDs treated with octylamine only (Figure 5C, blue dashed lines). The pretreatment of QDs with CdCl₂ increased the device efficiency from 0.01% to 0.8%. We also “post-treated” some devices with CdCl₂ by dipping the as-deposited CdTe QD films into a saturated solution of CdCl₂ in methanol before sintering while these devices showed overall higher efficiencies (red lines in Figure 5C), pretreated QDs still provided higher device efficiencies (3.3%) compared to control...
QDs (2.3%). We observed similar results for CdTe QD devices pretreated with InCl₃, which will be the focus of an upcoming publication. This highlights again the generality of passivating the undercoordinated surface anions with Lewis acidic ligands. These results demonstrate that surface passivation of QDs in solution by MCl₃–amine complexes is a viable method to improve the performance of QD-based devices.

**CONCLUSION**

In conclusion, we have shown that CdTe QDs can be passivated with a wide range of MX₃ Lewis acidic ligands, affording tunable PL QY increases up to near-unity values for optimized conditions. The PL QY of these CdTe QDs depends on the equilibrium between the MX₃ ligands that are added and the surface, which is well described by a model considering only a single binding energy to the surface. These results show that the passivation of Te surface sites by Lewis acidic (Z-type) ligands is the most important factor for eliminating traps in these materials and support our recent theoretical study that two-coordinated anions are the origin of trap states in II–VI QDs. We have also found that the addition of ligands that we do not expect to bind to the two-coordinated Te trap state may increase the PL QY, albeit to significantly lower values and with a smaller binding energy. This result highlights that the QD surface is a complex and dynamic system that is not yet completely understood.

In addition we have shown that the same MX₃ ligand passivation is also effective on other II–VI and III–V QDs, demonstrating that the passivation of undercoordinated anions is in general key to achieve high PL QY values in QDs. Finally, we have demonstrated that the relevance of ligand binding model simulations; and DFT calculations to TSCPC and transient absorbance data; and manipulation of CdSe nanocrystal surface stoichiometry. J. Am. Chem. Soc. 2007, 129 (10), 2841–2848.

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**ASSOCIATED CONTENT**

Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.8b07783.

Supplementary methods (metal carboxylate ligand synthesis, surface area calculations, PV device fabrication); extended QD characterization (XRD, XPS, NMR, Raman spectroscopy, time-resolved spectroscopy); extended data sets for CdTe and other QD materials; calculation of radiative recombination and trapping rates from fits to TSCPC and transient absorbance data; ligand binding model simulations; and DFT calculations (PDF)

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Notes
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**ACKNOWLEDGMENTS**

A.J.H. acknowledges support from the European Research Council Horizon 2020 ERC Grant Agreement No. 678004 (Doping on Demand). This research is supported by the Dutch Technology Foundation STW (project No. 13903, Stable and Non-Toxic Nanocrystal Solar Cells, and project No. 12734, Cadmium-free All-Inorganic Quantum Dots as Down-Conversion LED Phosphors), which is part of The Netherlands Organization for Scientific Research (NWO) (Vidi grant, No. 723.013.002), and which is partly funded by Ministry of Economic Affairs.

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