Sensitization of p-GaP (100) Photoelectrodes With CdSe Quantum Dots

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Abstract

This thesis contains two sections of my undergraduate research. Chapter one is dedicated to demonstrate a light-stimulated hole injection process by using p-GaP(100) photoelectrodes and CdSe quantum dots. The valence band edge of CdSe is higher than the valence band edge of p-GaP. After excitation of light, the photoinduced hole in the valence band of CdSe is thermodynamically possible to be injected into the valence band of GaP. A series of systematic electrochemical measurements including photoresponse, time-resolved photoluminescence decays, and steady-state current-potential responses have indicated the occurrence of such quantum dots sensitized hole injection processes.

The excitonic peak of the wet-chemically synthesized CdSe quantum dots locates around 560 nm; however, the photoresponse position of p-GaP photoelectrode is also locating around that wavelength. Namely, this similarity makes it challenging to deconvolute the spectral signatures of the GaP substrate and the CdSe sensitizer. Chapter two demonstrates my work focusing on surface modification of CdSe quantum dots in order to shift the excitonic dynamics. When the size of quantum dots is too small, hole injection processes cannot be demonstrated with photoresponse measurement because the spectral profile of GaP and CdSe will be convoluted in external quantum yield spectra. With ligand exchanging procedures inspired from literatures, the CdSe sensitization peaks have shown a bathochromic shift, well out of the spectral signatures of the GaP substrate. UV-Vis spectroscopy, fluorescence spectroscopy, and NMR spectroscopy have been used to show the redshifted exciton peak and characterize the synthesized surface ligands. Photoresponse measurement is used to demonstrate the same hole injection process in chapter one.
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CHAPTER 1

Sensitization of p-GaP (100) photoelectrode with CdSe Quantum Dots:
Light stimulated hole injection

1.1 Introduction

Photosensitization of semiconductor photoelectrodes has been a main method to utilize solar energy among current photoelectrochemical studies\(^1\)\(^-\)\(^5\). In such heterostructure systems, a light-absorbing material, which generates an electron-hole pair, is deposited onto a semiconductor photoelectrode. The photo-induced electron and hole quickly separate into two different phases, and these charge carriers are transported via a redox couple and collected by respective electrodes\(^3\). One example of such photovoltaic devices is dye-sensitized solar cell, where dye molecules are the light absorbers deposited onto semiconductor photoelectrodes (mainly TiO\(_2\))\(^2\),\(^6\)\(^-\)\(^{12}\). Nevertheless, the relatively low energy conversion efficiency and absorption coefficients among dye-sensitized solar cells have stimulated the need to seek new photo-absorber materials\(^{13}\)\(^-\)\(^{14}\).

Semiconductor nanocrystals (Quantum Dots) have attracted much attention due to their large extinction coefficients, tunable band gap, and well-established synthetic strategies\(^3\),\(^5\),\(^5\)\(^-\)\(^6\)\(^6\)-\(^{16}\). Due to the quantum confinement effect, intrinsic optical band gaps of these semiconductor materials can be tuned by changing the size of synthesized particles in order to maximum solar photon absorption\(^{17}\)\(^-\)\(^{18}\). Due to these benefits of QDs, instead of dye sensitization, QDs sensitization of semiconductor photoelectrodes has been explored by many research groups\(^{19}\)\(^-\)\(^{22}\).

\(^1\) Portions of this chapter were published in Wang, Z.; Shakya, A.; Gu, J.; Lian, S.; Maldonado, S. J. Am. Chem. Soc. 2013, 135 (25), 9275-9278
Several studies have demonstrated sensitization of a semiconductor photoelectrode by using the light-stimulated electron transfer from the conduction of a quantum dot to the conduction band of metal oxides\textsuperscript{1,20,23}. Nevertheless, photoelectrochemical studies involving quantum dots sensitization on the hole injection part are still poorly explored, which can be attributed to the difficulty in finding viable combination of materials that is thermodynamically possible to inject light-excited hole from the valence band of a quantum dot to the valence band of semiconductor photoelectrode. Inspired by several literatures regarding dye sensitization based on hole injection processes of planar p-type GaP photoelectrodes, the possibility of using p-GaP and quantum dots to demonstrate hole injection have been explored in this study\textsuperscript{2,7}.

The valence band edge potential of GaP has been reported to be +1.0 V vs Ag/AgCl, which is more negative than that of CdSe\textsuperscript{2,7,24-25}. This feature suggests that hole injection from a CdSe quantum dot into p-GaP should be thermodynamically possible.

In this study, wet-chemically synthesized CdSe quantum dots were surface-absorbed onto p-GaP (100) photoelectrodes. The native surface ligands of colloidal quantum dots (triocylphosphine oxide and oleate species) were exchanged with more conductive organic ligands such as ethylenediamine in order to increase the external quantum yield signals. A combination of optical characterization, photoresponse measurement, time-resolved photoluminescence decay measurement, and stead-state current-potential responses techniques has been utilized to demonstrate the light-stimulated hole injection process.

A control experiment was also performed in order to comprehensively study the proposed hole injection mechanism. Type I CdSe/ZnSe core/shell quantum dots were
synthesized and used to sensitize the p-GaP (100) photoelectrodes. In a Type I core/shell quantum dots, the valence and conduction band edge of the shell material is respectively higher and lower than the valence and conduction band edge of the core material. After shining light on p-GaP/CdSe/ZnSe pair, hypothetically, all the photoinduced excitons should be confined within CdSe core. In external quantum yield spectra, no additional photoresponse signals should be observed apart from the GaP’s intrinsic profile.
1.2 Materials and methods

**Materials and Chemicals**

Selenium powder (Se, ≥99.5%), 1-octadecene (ODE, ≥99.5%), cadmium acetate dehydrate (Cd(CH$_3$CO$_2$)$_2$, ≥98%), oleic acid (OA, ≥99.0%), trioctylphosphine oxide (TOPO, 90%), zinc stearate (technical grade), trioctylphosphine (TOP, 90%), ethylenediamine (EDA, ≥99.5%), potassium chloride (KCl, ≥99%), (3-aminopropyl)triethoxysilane (≥98%), heptanes (99%), isopropanol (≥99.7%), toluene (99.8%), dodecylamine (99%) and ammonium sulfide ((NH$_4$)$_2$S, 20% in H$_2$O) were used as received from Sigma Aldrich. Europium (III) chloride (EuCl$_3$, 99.9%) was obtained from Strem Chemicals. Methanol (99.8%) and acetone (99.5%) were obtained from VWR International. Hexanes (98.5%) and 40% v/v NH$_4$F (aq) were purchased from MARCRON Chemicals and Transene, respectively. Chloroform (99.9%) and concentrated HCl (aq) were obtained from Fisher Scientific. Silver print was from GC Electronics. H$_2$O was purified (>18 MΩ cm) with a Barnstead Nanopure III purifier and used throughout. N$_2$ (g), Ar (g) and forming gas (5% H$_2$ in N$_2$) were obtained from Metro Welding. GaP photoelectrodes were prepared from a 500 μm thick, single-crystalline, single-side polished p-GaP(100) wafer doped with Zn at $5.0 \times 10^{17}$ cm$^{-3}$. GaP wafers were obtained and used as received from ITME. Epoxy was purchased from Hysol C.

**Synthesis of CdSe quantum dots**

CdSe quantum dots were prepared using a modified approach reported by Peng et al.\textsuperscript{26-27} 0.047g Se powder and 7.092 g ODE were put in a 50 ml three-necked round-bottom flask and purged under Ar flow with constant stirring for 30 min. A 25 ml
Erlenmeyer flask containing 0.1596 g Cd(CH\textsubscript{3}CO\textsubscript{2})\textsubscript{2} and 3 ml OA was purged under N\textsubscript{2} flow with constant stirring. The Se/ODE mixture was then heated to 280 °C using a heating mantle and the temperature was monitored by a thermocouple until the Se powder was completely dissolved and a pale yellow solution was generated. The Cd(CH\textsubscript{3}CO\textsubscript{2})\textsubscript{2} and OA mixture was heated on a hot plate until ~120 °C and injected into the Se/ODE mixture. Within 2-3 min of injection, the solution turned orange indicating the nucleation of quantum dots and the color became deeper with increasing reaction time. After cooling to room temperature, the quantum dots were purified with a combination of methanol and acetone three times respectively. The final quantum dots were dispersed in hexanes for characterization.

*Synthesis of CdSe/ZnSe core/shell quantum dots*

CdSe/ZnSe quantum dots were synthesized using procedures reported by Bard et al.\textsuperscript{28} 6 g TOPO and 0.05 g Cd(CH\textsubscript{3}CO\textsubscript{2})\textsubscript{2} were put in a three-necked round-bottom flask and heated to 140 °C with continuous purging with Ar flow. The temperature was kept at 140 °C for 1 hr and then increased to 330 °C. 2.4 mL of 1M Se in TOP solution was injected into the hot cadmium solution. The instant color change suggested the nucleation of CdSe quantum dots. Quantum dots were purified with the same procedure mentioned above and finally dispersed in heptanes. 3 g TOPO and 2 g dodecylamine were placed in a three-necked flask, degassed with Ar and heated to 150 °C. After 1 hr, 6 mL of CdSe in heptanes solution was added into the flask with slowly injected Zn precursor solution (0.2 g Zn stearate in 2.5 mL toluene and 0.03 g Se in 2.5 mL TOP). The core/shell quantum
dots were purified using the same procedure as described above and dispersed in chloroform for optical and photoelectrochemistry measurements.

**Preparation of p-GaP electrodes**

One side polished 500 µm thick single crystal p-GaP(100) wafer with doping density of $5 \times 10^{17}$ cm$^{-3}$ was diced into 0.5 cm × 0.5 cm squares. Ohmic contact was prepared by etching the backside of wafer briefly (30 s) with concentrated NH$_4$F (aq), rinsing with water, soldering a thin, even film of pure indium on the back, and then annealing in forming gas for 10 min at 400 °C. Electrodes were prepared by attaching the GaP section (using silver print) onto a copper wire coil threaded through a glass tube and sealing with inert epoxy. Electrode areas were defined by the edge of the epoxy and were nominally 0.2 cm$^2$.

**Preparation of CdSe-sensitized and CdSe/ZnSe-sensitized p-GaP electrode**

The p-GaP electrode was first etched using concentrated HCl for 30 s and then merged in HCl-neutralized (NH$_4$)$_2$S for 6 hrs, rinsed with water, and dried with N$_2$. The electrode was merged in CdSe quantum dots hexanes solution (or CdSe/ZnSe chloroform solution) for 20 min. The loosely attached quantum dots were removed by rinsing with hexanes (CdSe/ZnSe using chloroform) and soaking in hexanes (CdSe/ZnSe using chloroform) for 20 min. Finally, the quantum dots sensitized p-GaP electrode was soaked in 10% v/v EDA methanol solution for 10 min for ligand exchange before being used for photoelectrochemical measurements.
Photoelectrochemical and spectroscopic measurements

Photoelectrochemical measurements were conducted in an airtight quartz cell with an optically flat bottom. A Pt counter electrode and an Ag/AgCl reference electrode were used. Aqueous solution containing 0.1 M KCl & 0.002 M EuCl$_3$ were adopted as the electrolyte. The electrolyte was purged with N$_2$ flow when measurements were conducted. External quantum yield was measured with an Oriel 150 W Xe arc lamp (Newport) and a quarter-turn single-grating monochromator (Newport). Sample measurements were recorded with chopped illumination (20 Hz), and a quartz beam splitter was used to simultaneously record the light output intensity with a separate Si photodiode (Newport) to adjust the fluctuations in lamp intensity. The potential of the working photoelectrode was set to -0.6 V vs Ag/AgCl, and the absolute photocurrents were measured by a digital PAR 273 potentiostat. The output current signal was connected to a Stanford Instruments SR830 lock-in amplifier, and the output signals from the lock-in amplifier and the reference Si photodiode were fed into a computer controlled by custom-written LabVIEW software. Current-potential curves were measured using the digital PAR 273 potentiostat under the unchopped illumination from the monochromator.

Absorption spectra were obtained with a Varian Cary 5000 ultraviolet–visible–near-infrared (UV–Vis–NIR) photospectrometer. Luminescence spectra were measured using Fluoromax-2 Fluorimeter. Luminescence lifetime spectroscopy was measured using ALBA microscope system from ISS (Illinois) combining fluorescence lifetime imaging (FLIM, FRET) with fluorescence fluctuation spectroscopy (FCS, FCCS, FCH) capabilities. A monolayer of CdSe quantum dots on glass was prepared following the
same procedure with that on p-GaP, the glass was pretreated by soaking in 10% v/v (3-aminopropyl)triethoxysilane isopropanol solution overnight.

### 1.3 Results and discussions

Figure 1.1 d illustrates a schematic depiction of sensitized hole injection process form a CdSe QD to p-GaP. Theoretically, the valence band position of CdSe is lower than the valence band position of p-GaP. After shining light on the p-GaP/CdSe pair, remaining holes in the valence band of CdSe should be injected into the valence band of GaP. Figure 1.1 a shows two representative time-resolved photoluminescence decays after pulsed excitation at 561 nm. When deposited onto insulating glass, CdSe QDs capped with ethylenediamine exhibited an average decay time of 8.8 ns; when surface absorbed onto p-GaP wafer, CdSe QDs showed an average decay time of 1.4 ns, with nearly an 4-fold decrease. The decreased PL lifetime indicates that a new fluorescence-quenching pathway has been generated. Exciton relaxation can be accelerated due to the hole injection processes happened at the interface between CdSe and GaP.

Figure 1.1 b contains representative steady-state photoelectrochemical response for CdSd/p-GaP pairs. Under illumination with sub-bandgap nomochromatic light (605 nm), an enhanced photocurrent has been observed compared with the bare p-GaP photoelectrodes under illumination.

Figure 1.1 c is the wavelength-dependent external quantum yield for photoelectrodes fabricated with CdSe-EDA QDs. The inset figure is the absorption spectrum for the QDs that are absorbed onto the surface of GaP. After sensitization (green curve), an increased external quantum yield values were observed with a qualitatively matched wavelength
range with the absorption profile of CdSe. Compared with bare p-GaP photoelectrodes, the observed additional signals beyond 550 nm should be attributed to the injected photoinduced hole coming from CdSe QDs.

The combination of these data strongly suggest that CdSe QDs are able to inject light stimulated hole in to p-GaP photoelectrodes.

Figure 1.1. a) Time-resolved photoluminescence decays measured from adsorbed CdSe quantum dots (diameter = 4.5 nm) on (red line) glass and (green line) p-GaP(100), respectively. Pulsed excitation wavelength = 561 nm. b) Steady-state current-potential responses for a (black lines) mirror-polished, bare p-GaP(100) photoelectrode and (green lines) a mirror-polished p-GaP(100) photoelectrode with adsorbed CdSe quantum dots (diameter = 4.5 nm). Responses both (dashed lines) in absence of any illumination and (solid lines) under monochromatic illumination at 605 nm at 0.33 mW cm$^{-2}$ are shown. c) External quantum yield spectra for (black) a bare p-GaP(100) photoelectrode and (green) a p-GaP(100) photoelectrode coated with a monolayer of CdSe quantum dots. Inset: Absorption spectrum for the same CdSe quantum dots dispersed in hexane. d) Schematic depiction of sensitized hole injection from a CdSe quantum dot into p-GaP.

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In order to comprehensively study the hole injection mechanism, a control experiment was also performed with type I core/shell CdSe/ZnSe quantum dots. In a Type I core/shell system, the conduction and valence band edge energies for ZnSe extend both above and below the conduction and valence band-edge energies for CdSe respectively. After excitation of light, hypothetically all the photoinduced excitons will be confined within the CdSe core based on fundamental thermodynamic theories. Hence, the hole injection process should be forbidden. Figure 3 b shows that the sensitization signals from p-GaP/CdSe decreases with the increasing thicknesses of ZnSe shells, which is consistent with the proposed hypothesis. The ZnSe shell here works as an additional barrier, which blocks charge carriers from being transferred between the CdSe core and GaP. Compared with previous studies, in the p-Gap/CdSe system, the photoinduced hole should be able to transfer as the proposed mechanism (from the valence band of CdSe into the valence band of GaP).

Figure 1.2. a) External quantum yield spectra for a textured p-GaP(100) photoelectrode sensitized with CdSe quantum dots (diameter = 4.5 nm) after various quantum dot surface treatments. b) External quantum yield for a textured p-GaP(100) photoelectrodes sensitized with CdSe quantum dot cores (diameter = 3.7 nm) coated with ZnSe. The shell thicknesses were controlled through variation in the ZnSe shell growth time.

1.4 Conclusion

A systematic collection of optical and electrochemical measurements have demonstrated that p-GaP and CdSe QDs is a viable combination of materials to demonstrate hole injection processes from the valence band of CdSe into the valence band of p-GaP. Before CdSe being surface absorbed onto GaP photoelectrode, apart from its own photoresponse signature, there was no observable photoresponse beyond 550 nm; after CdSe sensitization, external quantum yield measurements indicated that additional charge carriers have been collected beyond the GaP’s optical band range, which should be attributed to the photoinduced hole injected from CdSe. Based on the optical studies of CdSe, the excitonic dynamic profile is also consistent between the absorption spectra and external quantum yield measurements.
CHAPTER 2

Tuning sensitization photoresponse of p-GaP/CdSe pair by phenyldithiocarbamate surface ligand

2.1 Introduction

Due to the nature of synthetic systems, colloidal semiconductor nanocrystals (Quantum Dots) usually are passivated with long-chain aliphatic compounds (stearic acid and oleic acid)\(^{26, 29-31}\). The most established method is the so-called hot-injection technique\(^{32-33}\). It involves the rapid injection of metal precursors into a hot solvent dissolved with another inorganic precursor. The subsequent temperature drop controls the nucleation and growth of QDs\(^{34}\). Typical organic solvents for this system are oleic acid, n-octadecyphosphonic acid, alkylphosphines and alkylphosphine oxides\(^{35}\). Such organic molecules will adhere to the surface of growing crystals to facilitate the kinetics of nucleation and growth, prevent the semiconductor nanocrystal core from oxidation, and control the shape of QDs. Bawendi et al. has demonstrated a systematic method to synthesize uniformly monodispersed nanocrystals by using a combination of long-chain alkylphosphines, alkylphosphine oxides, and alkyamines organic compounds\(^{32}\). Nevertheless, the non-conductive long carbon chains in such systems can also compromise charge transfer processes in the system of interest because they separate individual nanocrystals to prevent efficient charge transport, induce charge trapping, and complicate the surface chemistry of QDs\(^{30-31, 36}\). Hence, numerous researchers have attempted to study how these surfactant organic molecules can impact the morphology, electronic properties, and chemical functionalities of QDs so that they can chemically functionalize these nanostructured materials\(^{1, 28, 30, 36-38}\). Weiss has reported several roles
of organic molecules in terms of controlling the growth, surface structure, and redox activity of colloidal semiconductor nanocrystals. According to her account, in the “hot-injection” reaction mixture, using long-carbon-chain organic molecules can control the shape of metal-chalcogenide QDs. Bulky alkylphosphines such as trioctylphosphine can confine the growth of QDs in three dimension in order to prevent crystal defects. Interestingly, organic surfactant molecules can impact not only the shape and composition of final products, but also the absorption and photoluminescence of QDs. When the highest occupied molecular orbital (HOMO) of the organic surface ligand couples with the HOMO of QDs, the anti-bonding orbital of this coupling, along with the lowest occupied molecular orbital (LUMO) of QDs will determine the new optical band gap of the QD’s material.

Previously, we have demonstrated a light stimulated hole injection process from a CdSe quantum dot to a p-GaP photoelectrode. In that study, the excitonic peak of CdSe is centered at 560 nm, which is similar to the native absorption of the planar GaP substrate. Namely, this similarity makes it challenging to deconvolute the spectral signatures of the GaP substrate and the CdSe sensitizer. Weiss et al. has demonstrated a ligand exchange procedure, which uses phenylthiocarbamate (PTC) ligand to control hole delocalization in QDs through coupling effect between the HOMO of PTC and the valence band edge of CdSe QDs. In that study, a bathochromic shift up to 220 meV in the optical bandgap of CdSe induced by PTC has been observed. Leveraging this ligand exchange procedure, a photoelectrochemical study of the p-GaP/CdSe pair, along with the surface modification of colloidal CdSe QDs has been performed in this chapter. The native ligands (TOPO) of wet-chemically synthesized CdSe QDs have been exchanged
with PTC molecules and para-substituted derivatives of PTC. Subsequently, the surface modified CdSe QDs have been further used to sensitize p-GaP (100) photoelectrode. As what figure 2.1 depicting, theoretically, the HOMO of PTC can be coupled with the valence band edge of CdSe. The newly formed band gap of CdSe-PTC is the energy difference between hybrid anti-bonding orbital and the conduction band edge of CdSe. If the PTC molecule is para-substituted with an electron-withdrawing group, the inductive effect can lower the HOMO energy level of PTC. Hypothetically, the coupling effect should be stronger so that the bathochromic shift should be more severe. If the PTC molecule is para-substituted with an electron-donating group, the HOMO energy level of PTC should be increased. The coupling effect should lead to a less obvious bathochromic shift.

In this study, PTC, PTC-OMeth, and PTC-F (shown in table 1) have been synthesized and exchanged with the native TOPO ligand of CdSe quantum dots. UV-Vis spectroscopy was utilized to study the excition dynamics. Furthermore, the surface modified CdSe QDs were used to sensitize p-GaP (100) photoelectrodes. Quantum yield and photoluminescence decay measurements were used to demonstrate hole injection processes from the valence band of CdSe QDs to the valence band of GaP.
Schematic 1. Synthesized PTC molecular structures and PTC molecule-CdSe quantum dots linkage demonstration.

Figure 2.1. Schematic demonstration of coupling effect between the HOMO of PTC (PTC-electron withdrawing group and PTC-electron donating group) and the valence band edge of CdSe QDs.
2.2 Materials and methods

*Synthesis of CdSe quantum dots*

CdSe quantum dots were prepared using a modified method reported by Weiss et al.\(^{40}\). A 50 ml three-necked round-bottom flask containing 7.76 g of trioctylphosphine oxide, 7.76 g of hexadecylamine, and cadmium acetate (0.660 mmol) was purged under Ar flow with constant stirring for 60 min in 150 °C. One arm of the flask was left open to remove any remaining water in the system. After that, the flask was sealed and heated up to 330 °C. After the Cadmium precursor was completely dissolved in the reaction mixture, trioctylphosphine selenide solution (4mL of 1 M solution) was rapidly injected and the reaction mixture was immediately cooled to room temperature. The resulting products were purified by methanol and re-dispersed in hexanes.

*Synthesis of dithiocarbamate ligands*

NH\(_4\)X-PTC ligands were synthesized based on a method reported by Weiss et al.\(^{39}\). Two equivalents of carbon-disulfide to one equivalent of the appropriate para-substituted aniline (Aniline, OMeth-Aniline, and F-Aniline) were added into excess concentrated ammonium hydroxide at 0 °C and stirred overnight. The resulting powder were washed with chloroform and dried under vacuum and characterized by NMR.

*Film preparation and ligand exchange*

QDs were drop-casted onto borosilicate glass microscope slip covers. Stock solutions of X-PTC in methanol were prepared at 10mM, 15mM, and 20mM. The glass
slip covers coated with QDs were put into the solution overnight for ligand exchange. Uv-Vis spectroscopy was used for exciton dynamic studies.

**Preparation of CdSe-sensitized p-GaP electrode**

The p-GaP electrode was first etched using concentrated HCl for 30 s and then soaked in HCl-neutralized (NH₄)₂S for 6 hrs. Rinsed with water thoroughly and dried with N₂ flow, the electrode was immersed in CdSe quantum dots hexanes solution for 20 min. In order to form a monolayer of quantum dots on GaP surface, the loosely attached quantum dots were removed by thoroughly rinsing with hexanes. Subsequently, the quantum dots sensitized p-GaP electrode was soaked in 10% v/v EDA methanol solution for 10 min for ligand exchange before being used for photoelectrochemical measurements. After obtaining the CdSe-EDA/p-GaP photoresponse spectra, the CdSe sensitized GaP electrodes were further merged in the appropriate PTC Methanol solution [15mM] overnight for ligand exchange. External quantum yield measurements were performed subsequently.

**Preparation of p-GaP electrodes and Photoelectrochemical and spectroscopic measurements**

Preparation of p-GaP electrodes and the conditions of photoelectrochemical and spectroscopic measurements in this section are same as they are in chapter one.
2.3 Results and discussions

A combination of optical and photoelectrochemical studies has shown that exchanging the native surfactant ligands (TOPO) of CdSe QDs with PTC ligands can shift the excitonic dynamic and further enable the hole injection from CdSe to GaP. Figure 2.2 a shows that after treatments of PTC in methanol solution, CdSe-PTC QDs have demonstrated a red shift up to 20 nm. Equation 1 describes an empirical fitting function of the diameter of CdSe QDs in terms of their excitonic position\textsuperscript{44}. Based on calculation, the original size of CdSe-TOPO was 5.29 nm radius. After ligand exchange, the calculated radius was up to 5.60 nm. Figure 2.2 b illustrates a set of photoelectrochemical studies in terms of external quantum yield measurement. All three measurements (Black: bare p-GaP, Red: p-GaP/CdSe/EDA, and Blue: p-GaP/CdSe/PTC) were performed by using the same electrode, same calibration curve, and same conditions. Due to the instrumental filter switch, there was an artificial peak at 570 nm. However, between the red(p-GaP/CdSe-EDA) and black(bare p-GaP) curve, the 560-580 nm range did not show photoresponse, which means that, theoretically, the delocalized holes were not injected from the valence band of CdSe to the valence band of GaP OR the hole injection process was not observed by external quantum yield measurement due to the convolution between the photoreponse profile of CdSe and GaP. Blue curve shows an increased area within 560-580 nm. Hypothetically, these increased signals should be attributed to the additional charge carriers coming from CdSe. Besides, after 610 nm, all three curves coincide with each other and go towards zero, which further indicates that the PTC treatments have shifted the excitionic peaks of CdSe from the original 570 nm to near 600 nm and enabled hole injection processes within this specific range. Additionally,
the increased photoresponse appeared within a range that is matching with the exciton
dynamic of the surface modified CdSe depicted in UV-Vis spectrum (Figure 2.1 a).
Figure 2.2 c more clearly shows an increased signal at 580 nm and 600 nm between the
before-ligand-exchange CdSe sensitization and the after-ligand-exchange CdSe
sensitization. In Figure 2.1 a, the wavelength range of excitation states are also consistent
with such observations.

Equation 1\textsuperscript{44}:

\[ D(\text{nm}) = (-6.6521 \times 10^{-8})\lambda^3 + (1.9557 \times 10^{-4})\lambda^2 - (9.2352 \times 10^{-2})\lambda + 13.29 \]

**Figure 2.2.** a) UV-Vis absorption spectra of CdSe quantum dots with TOPO native ligand (black line) and
with PTC ligands in different concentrations. b) External quantum yield spectra for (black) a p-GaP (100)
photoelectrode coated with a layer of CdSe with ethyldiamine ligand (black) and phenyldithiocarbamate
ligand (red). c) External quantum yield difference between p-GaP/CdSe-EDA and p-GaP/CdSe-PTC.
Figure 2.3 shows a set of Time-resolved photoluminescence decay measurements. Based on equation 2, when deposited onto insulating glass, CdSe QDs capped with TOPO has shown an average decay time of 2.407 ns. When deposited on GaP, CdSe-TOPO has displayed an average decay time of 2.653 ns. Nearly same photoluminescence decay times demonstrate that the same exciton relaxation pathway may be operative for these two samples. However, after the treatment of ligand exchange, the CdSe-PTC on glass has demonstrated an average decay time of 9.599 ns, while on GaP, an average decay time of 8.110 ns has been observed. Based on calculation, a 1.489 ns difference shows that a new exciton-relaxing pathway may have been generated. Theoretically, if light stimulated hole of CdSe on GaP has been injected into GaP, the photoluminescence decay for CdSe should be faster than the CdSe on Glass samples due to the transfer mechanism of charge carriers. Based on observations, the experimental data is consistent with this hypothesis.

Equation 2: 
\[ \frac{\alpha_1 \tau_1^2 + \alpha_2 \tau_2^2 + \alpha_3 \tau_3^2}{\alpha_1 + \alpha_2 \tau_2 + \alpha_3 \tau_3} \]

Figure 2.3. a) Time-resolved photoluminescence decays measured from absorbed CdSe quantum dots with TOPO native ligands on (red line) GaP and on (black line) glass. b,c,e,f) Fluorescence image of CdSe on glass, GaP, CdSe-PTC on glass, and GaP. d) Time-resolved photoluminescence decays measured from absorbed CdSe quantum dots with PTC ligands on (red line) glass and on (black line) GaP.
As what discussed in introduction part, the coupling effect between the HOMO of PTC and the valence band edge of QDs can result in the excitonic shift of QDs. If the PTC molecule is para-substituted with an electron-withdrawing group, the inductive effect can lower the HOMO energy level of PTC. Hypothetically, the coupling effect should be stronger so that the bathochromic shift should be more severe. If the PTC molecule is para-substituted with an electron-donating group, the HOMO energy level of PTC should be increased. The coupling effect should lead to a less obvious bathochromic shift. Figure 2.4 a shows a set of UV-Vis absorption spectra of CdSe after treatment of OMeth-PTC ligand exchange. Methoxy group is an electron-donating group. The HOMO energy level of this para-OMeth-PTC ligand should be higher than that of PTC ligand. Theoretically, the week coupling effect between p-OMeth-PTC and CdSe QDs should not generate severe bathochromic shift in UV-Vis spectra. Figure 2.4 a is consistent with this hypothesis. If the exciton dynamic of QDs is not altered, in photoresponse measurements, both before-ligand-exchange and after-ligand-exchange CdSe/p-GaP pairs should not show hole injections because the originally synthesized QDs are too small and the hole injection is thermodynamically forbidden. Figure 2.4 b is also consistent with this hypothesis. Both of such observations are consistent with literatures\textsuperscript{39}. 
Figure 2.4. a) UV-Vis absorption spectra of CdSe quantum dots with TOPO native ligand (black line) and with PTC ligands in different concentrations. b) External quantum yield spectra for (black) a bare p-GaP (100) photoelectrode (black line) and a p-GaP (100) photoelectrode coated with a layer of CdSe with ethyldiamine ligand (red) and PTC ligand (blue).

Figure 2.5 shows a collection of UV-Vis spectra of CdSe-FPTC. After ligand exchange, a red shift up to 40 nm has been observed, which is higher than the observed shift in CdSe-PTC. Based on Equation 1, the increased radius after ligand exchange is 1.32 nm. The CdSe-PTC samples have 0.31 nm of increased radius. These observations are consistent with the coupling theory previously stated and depicted in literatures\textsuperscript{38, 42}.

Figure 2.5 b shows the external quantum yield comparison. Black line is the p-GaP/CdSe sensitization with TOPO surface ligand. There is no observable sensitization signal within 550-600 nm range due to the small sizes of synthesized quantum dots. After surface modification, according to the UV-Vis spectra, quantum dots have shown excitonic peaks shifted to higher wavelength. Accordingly, in quantum yield spectra, additional signals have been observed within this range. Using same electrodes, calibrations, and physical conditions to eliminate instrumental interferences performed both of the measurements.

In this entire study, all observed quantum yield are relatively lower than that in p-GaP CdSe sensitization with EDA ligand in chapter one. This can be attributed to the
nature of long molecular length and bulky molecular structure of phenyldithiocarbamate ligand. Several studies have shown that organic surfactants can separate individual quantum dots and prevent efficient charge transport at the interface. The relatively shorter molecular length of ethylenediamine molecules and better conductivity of amine group may have provided a better charge transfer pathway compared with the phenyldithiocarbamate molecule at the interface between GaP and CdSe.

Figure 2.5. a) UV-Vis absorption spectra of CdSe quantum dots with TOPO native ligand (black line) and with F-PTC ligands in different concentrations. b) External quantum yield spectra for (black) a p-GaP (100) CdSe-TOPO photoelectrode and a p-GaP (100) photoelectrode coated with a layer of CdSe with FPTC (red).

2.4 Conclusions

A systematic collection of optical and electrochemical measurements have demonstrated that exchanging the native surface ligand of CdSe QDs with phenyldithiocarbamate will red shift the excitonic peak of QDs. Before ligand exchange, the small CdSe QDs, which have exciton peak at 550 nm did not show increased signals in photoresponse measurement due to the convolution between their exciton dynamic and the GaP’s photoresponse signature. After ligand exchange, the increased area within the sub-band range should indicate the occurrence of hole injection from CdSe to GaP. Sensitization range is also consistent with the exciton position of the surface modified QDs.
References


35. Nozik, A. J.; Beard, M. C.; Luther, J. M.; Law, M.; Ellingson, R. J.; Johnson, J. C., Semiconductor quantum dots and quantum dot arrays and applications of multiple


Appendix

C-13 NMR

- N-CS₂

C-OMe

Aromatic

DMSO
Figure 3.1. (a), (b) and (c) Absorption, luminescence and x ray diffraction spectra of CdSe with different ZnSe shell thickness. Samples for XRD measurement were prepared by drop-casting CdSe hexanes solution on glass substrate.