High-Efficiency Optical Gain in Type-II Semiconductor Nanocrystals of Alloyed Colloidal Quantum Wells

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Abstract: Colloidal nanocrystals having controlled size, tailored shape, and tuned composition have been explored for optical gain and lasing. Among these, nanocrystals having Type-II electronic structure have been introduced toward low-threshold gain. However, to date, their performance has remained severely limited due to diminishing oscillator strength and modest absorption cross-section. Overcoming these problems, here we realize highly efficient optical gain in Type-II nanocrystals by using alloyed colloidal quantum wells. With composition-tuned core/alloyed-crown CdSe/CdSe$_x$Te$_{1-x}$ quantum wells, we achieved amplified spontaneous emission thresholds as low as 26 μJ/cm$^2$, long optical gain lifetimes ($\tau_{\text{gain}} \approx 400$ ps), and high modal gain coefficients ($g_{\text{modal}} \approx 930$ cm$^{-1}$). We uncover that the optical gain in these Type-II quantum wells arises from the excitations localized to the alloyed-crown region that are electronically coupled to the charge-transfer state. These alloyed heteronanostructures exhibiting remarkable optical gain performance are expected to be highly appealing for future display and lighting technologies.

Colloidal semiconductor nanocrystals hold great promise for full-color lasers$^1$ as a potential game changer for future solid-state lighting and innovative display systems.$^6,9$ To date, numerous structures of nanocrystals with their shape tailored, composition tuned, and size controlled have been extensively investigated to aim for efficient optical gain performance.$^3,4,5,10,12$ Yet their optical gain has been limited. This is a fundamental limitation arising due to the fact that single-exciton state does not provide net gain in common II–VI nanocrystals. To achieve net optical gain, biexcitons had to be generated by pumping these materials harder.$^3$ However, biexcitons in conventional nanocrystals are very short-lived because they are rapidly annihilated via fast Auger recombination processes ($\sim$10–50 ps),$^{13,14}$ immediately depleting the available optical gain and substantially shortening gain lifetime ($\tau_{\text{gain}} \ll 100$ ps).

Previously, heterostructured nanocrystals having Type-II electronic structure$^{6,17}$ or soft-confinement potential$^1,2$ have been proposed to circumvent Auger losses by either introducing single exciton gain or minimizing the Auger recombination process. However, optical gain performance has been commonly hampered in these nanocrystals due to reduced oscillator strength and small absorption cross-section, leading to overall modest stimulated emission cross sections and hence limited gain coefficients. Recently, colloidal quantum wells (CQWs), also known as colloidal nanoplatelets,$^{14,29}$ have been shown to achieve superior optical gain performance among colloidal nanocrystals, resulting in low gain thresholds$^{27,28}$ and high gain coefficients.$^{27}$ In these aforementioned works, only CQWs having either Type-I or quasi-Type-II electronic structure have been investigated. More recently, CdSe/CdTe (core/crown) CQWs having pure Type-II electronic structure have been synthesized; these CQWs combine large Stokes-shifted emission with long fluorescence lifetimes,$^{25,29}$ making them potentially interesting for amplified spontaneous emission (ASE) and lasing, which have remained elusive to date.

Here we demonstrate that Type-II CQWs that have composition-tuned alloyed-crowns exhibit remarkable optical gain performance with ASE threshold as low as 26 μJ/cm$^2$, accompanied by large net modal gain coefficients up to $\sim$930 cm$^{-1}$ and long gain lifetimes ($\tau_{\text{gain}} \approx 400$ ps). The performance...
of the Type-II CQWs studied here represents more than an order of magnitude improvement over the previous best reports in Type-II colloidal nanocrystals. We corroborate the underpinning mechanism of this efficient optical gain by ultrafast transient absorption (TA) spectroscopy and show that the excitations localized to the alloyed-crown region, which are electronically coupled to the charge transfer (CT) excitations at the Type-II interface, are responsible for the gain. The electronic coupling between the crown excitations and the CT state results in strong energetic shift, allowing for the minimization of the reabsorption losses, which is otherwise detrimental for optical gain in conventional quantum dots, and enable four-level-like gain system.

CdSe core-only CQWs having a vertical thickness of \( \sim 1.2 \text{ nm} \) (corresponding to four monolayers of CdSe with an additional Cd layer) and a lateral size of \( 14.0 \pm 1.6 \text{ nm} \) by \( 12.7 \pm 1.5 \text{ nm} \) were synthesized. (See Figure S1 for the TEM image.) Using this core-only CQW as the seed, we synthesized core/alloyed-crown CQWs of CdSe/CdSe\(_{1-x}\)Te\(_{x}\) (\( x = 0, 0.25, 0.50, 0.75 \)), where the crown layer grows laterally and the vertical thickness is kept fixed (see Methods).\(^{26,30}\) The lateral size of the core/alloyed-crown CQW was analyzed using high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM). Figure 1a,b shows the STEM images of the CdSe/CdSe\(_{1-x}\)Te\(_{x}\) CQWs with \( x = 0 \) and 0.50, respectively. (See Figure S2 for the other CQW samples.) After the crown growth, the lateral extent of the CQWs was uniform and the lateral size reached, on average, 29.3 nm by 20.6 nm. (See Table S1 for the lateral sizes of the CQWs with different crown composition.)

To inspect the elemental composition of the alloyed-crown CQWs, we performed X-ray photoelectron (XPS) and energy-dispersive X-ray (EDS) spectroscopies. Table S2 compares the elemental compositions of the alloyed-crows, estimated from the injected crown growth precursors and also analyzed via XPS measurements, showing a good agreement. Figure 1c depicts the line-probe EDS analysis of a single CdSe/CdSe\(_{0.50}\)Te\(_{0.50}\) CQW. (See Figure S3 for the CdSe/CdTe CQW.) The CdSe core and the CdSe\(_{0.50}\)Te\(_{0.50}\) alloyed-crown regions can be clearly distinguished in the green and orange shaded areas, respectively. These observations strongly indicate the homogeneous lateral growth of the alloyed-crown with a well-estimated stoichiometry owing to the similar reactivity of the Se and Te precursors.

Figure 1d shows the UV–vis absorbance and steady-state photoluminescence (PL) spectra of the core-only and core/alloyed-crown CQWs. Absorbance of the core-only CdSe CQW exhibits sharp excitonic features arising from the electron/ heavy-hole (e/hh) and electron/ light-hole (e/lh) transitions of CdSe at 512 and 480 nm, respectively.\(^{21,31}\) This seed CQW is quite pure in terms of vertical thickness population, as we do not observe any emission or absorption feature belonging to another CQW population having a different vertical thickness. As alloyed-crown layers are laterally grown, new absorption features emerge. (See Figure 1d.) For example, the absorbance peak at 560 nm in the CdSe/CdTe CQW is due to the e/hh transition in the CdTe crown.\(^{25}\) The absorbance features arising from the alloyed-crowns red shift as more Se is incorporated into the crown (see Figure 1d) due to the band gap bowing effect, which was previously observed in the alloyed CdSe\(_{1-x}\)Te\(_{x}\) quantum dots.\(^{22}\)

The steady-state PL emission of the CdSe/CdTe CQW peaks at 659 nm with a full width at half-maximum (fwhm) of 49 nm, arising from the spontaneous emission of the spatially indirect CT state across the Type-II interface.\(^{25,26,33}\) In the alloyed-crown CQWs, the PL emission peak blue-shifts as the fraction of Se in the crown is increased because the valence band edge energy is increased with more Se incorporated into the alloyed-crown. (See Figure S4 for the energy band alignments estimated from the absorptions and emission of the CQWs.) PL quantum yields are also noted in Figure 1d, which increase from 20 to 51% as the fraction of Se in the alloyed-crown is increased.

We investigated ASE in the thin films of the alloyed Type-II CQWs. For that, we tested both drop-casted and spin-coated films on quartz substrates. We found that spin-coated films (thickness \( \sim 150 \pm 20 \text{ nm} \)) achieve lower ASE thresholds, possibly due to better film-packing and also smaller optical
waveguide losses. Therefore, we only studied the spin-coated films for the further investigations. To minimize sample degradation during ASE measurements, we encapsulated the samples in a nitrogen-filled glovebox using coverslips and applied epoxy to prevent contact with oxygen and moisture. (See Figure S5.) As the pump source, we employed frequency-doubled output (400 nm, using a BBO crystal) of a femtosecond Ti:sapphire regenerative amplifier (Spitfire Pro, Spectra Physics) having a 120 fs pulse duration with a 1 kHz repetition rate.

Figure 2a shows the PL spectrum of the CQW samples as the pump fluence is gradually increased, showing the transition from spontaneous emission into the ASE. The ASE peaks are observed to be blue-shifted with respect to their corresponding spontaneous emission peaks. The amount of blue shift changes with the alloyed-crown composition. For example, the ASE peak in the CdSe/CdSe0.50Te0.50 CQW emerges at 609 nm, which is ~19 nm blue-shifted with respect to the spontaneous PL emission peak. In Figure 2b, luminance intensity versus pump fluence is plotted. In the CQW having pristine CdTe crown, the ASE threshold is the largest (~119 μJ/cm²). We checked over 10 different CdSe/CdTe CQW samples having different core and crown sizes, where the ASE thresholds were in the range of 100−300 μJ/cm² with an average threshold of ~200 μJ/cm² (see Figure S6), although the ASE peak was spectrally at the same position (~3 nm). Therefore, it was not possible to further reduce the ASE thresholds in the pristine crown Type-II CQW. The alloyed-crown CQWs, on the contrary, exhibited remarkably improved ASE performance with much lower ASE thresholds. The alloyed-crown CQW in the form of CdSe/CdSe0.50Te0.50 achieved the lowest ASE threshold (IASE = 26 μJ/cm²). Also, we checked three different batches of CdSe/CdSe0.50Te0.50 CQWs, which exhibited almost the ASE threshold (25−30 μJ/cm²). The other alloyed-crown CQWs exhibited ASE thresholds of 48 and 66 μJ/cm² for Se fractions of 25 and 75%, respectively. Therefore, alloyed-crown Type-II CQWs achieve substantially lower ASE thresholds than those of the nonalloyed-crown versions. Furthermore, slope-efficiency of the alloyed-crown CQWs is better than that of the pristine-crown CQW, suggesting effective utilization of the excitation for optical gain and hence higher gain coefficients.

Previously, the lowest ASE threshold in Type-II core/shell quantum dots was reported to be on the order of 2 mJ/cm².16,34 Therefore, the alloyed-crown Type-II CQWs here exhibit more than an order of magnitude improvement among Type-II nanocrystals. As a side note, ASE was not realized using Type-II CdSe/CdTe quantum dots before, possibly due to their strongly reduced oscillator strength in their core/shell quantum dot form. Therefore, Type-II nanostructures in the form of core/crown quantum wells are superior to their spherical core/shell counterparts. Here the best ASE threshold is comparable or better than the recently reported highly performing heterostructure-engineered nanocrystal systems including giant-shell CdSe/CdS quantum dots (IASE ~ 29 μJ/cm²),3,35 CdZnS/ZnS quantum dots (IASE ~ 50 μJ/cm²),3,35 and multishell CdSe/CdZnS/ZnS quantum dots (IASE ~ 90 μJ/cm²).3 Also, our ASE threshold is on par with the average ASE threshold reported using Type-I (or quasi-Type-II) core/shell CdSe/CdS CQWs,3,35 where statistically averaged ASE threshold was ~24 μJ/cm², whereas their hero sample was reported to achieve 6 μJ/cm². Similarly, giant-shell quantum dots have shown ASE thresholds down to 6 μJ/cm².
Previously, in Type-II quantum dots (e.g., core/shell CdS/ZnS), optical gain has been shown to arise from both single excitons and also spatially indirect biexcitons. A blue-shifted ASE was observed as attributed to the spatially indirect biexciton states. A blue-shifted ASE was observed as attributed to the spatially indirect biexciton states.16,36 A blue-shifted ASE was observed as attributed to the spatially indirect biexciton states, which possess positive binding energy \( E_{\text{BX}} - 2E_{\text{XX}} = E_{\text{B}} + E_{\text{R}} > 0 \) due to repulsive Coulomb interaction. In our case, if the ASE were to arise from spatially indirect biexcitons, then it would suggest the existence of a giant repulsive biexciton binding energy in these Type-II CQWs. For example, in the CdSe/CdTe CQW the blue shift between the ASE and the spontaneous emission peaks is 190 meV. (See Figure 2a.) Therefore, \( \Delta E_{\text{BX}} \) had to be on the order of 200 meV. However, Coulombic binding energy for an exciton in a CdSe CQW was theoretically calculated to be on the order of 150–200 meV37 and was experimentally determined to be 180 meV.38

Considering the large lateral extent of the CQWs, which should reduce the exciton-exciton interactions, one would expect much lower \( E_{\text{BX}} \). Therefore, giant biexciton binding energy in Type-II CQWs does not seem likely to explain the origin of the gain, and hence the gain mechanism in the Type-II CQWs is different than what has been reported for the conventional Type-II quantum dots.

In Figure 2c, we show the spectral positions of the ASE peak (red triangles), spontaneous emission peak (black circles), and the e/hh transition of the crown (blue squares) in all CQW samples. Although energy difference between the ASE and the spontaneous emission peaks monotonically decreases, surprisingly, the ASE peak is found to spectrally follow the bandgap (e/hh transition) of the alloyed-crown. This observation indicates that optical gain is associated with the excitations that are localized to the alloyed-crown region.

To further investigate the presence of the excitations in the alloyed-crown, we performed ultrafast TA measurements. Previously, internal charge transfer within CdSe/CdTe CQW has been studied and CT state formation has been observed on a picosecond time scale. However, in this aforementioned study the excitation fluence was intentionally kept low to prevent the generation of more than one exciton per CQW. Therefore, the nature of the excitations and their spatial localization throughout the CQW under elevated pump conditions have not been understood. Here we carried out pump-probe TA spectroscopy in the alloyed-crown Type-II CQWs as a function of the pump fluence. Experimental details of the TA system are summarized in the Supporting Information. Figure 3a shows the TA spectra of the CdSe/CdSe\(_{0.50}\)Te\(_{0.50}\) CQW (see Figure S7 for the other samples) (excited @400 nm with a fluence of 79 \( \mu \)J/cm\(^2\)) at three different pump-probe delay times: just after the excitation (t = 0–0.2 ps), at early time (t = 1–2 ps), and at later time (t = 800–1000 ps). Just after the excitation, we observe the bleach signals at 580 and 512 nm arising from the e/h transitions of the CdSe\(_{0.50}\)Te\(_{0.50}\) alloyed-crown (T1) and the CdSe core (T2), respectively, indicating the photogeneration of excitons in both the core and alloyed-crown regions. Also, a photoinduced absorption (PIA) feature is observed at 597 nm for early time, which marks the position of the biexciton state in the alloyed-crown. After a short time delay (t = 1–2 ps), the CT state is formed (appearing as bleach at 606 nm) via electron (hole) transfer from the alloyed-crown (core) into the core (alloyed-crown). Interestingly, at long pump-probe time delay (t = 800–1000 ps), both T1 and CT are found to coexist with persisting bleach signals at this fluence level (79 \( \mu \)J/cm\(^2\)), which is expected to generate more than one exciton per CQW. T1...
transition persists for long time delays (t > 800 ps) even after the formation of the CT state. This indicates that excitations could persistently remain in the alloyed-crown even after initial CT across the Type-II interface.

In Figure 3b, we present the decay (or bleach recovery) kinetics of the T1 (solid) and CT (dotted) states under three different pump fluences. (See Figure S8 for the other CQW samples.) At a low pump fluence level (e.g., 23 μJ/cm²), T1 has a fast decay component (∼1 ps) that leads to the formation of the CT state, indicating an efficient CT state formation. On the contrary, T1 decay is observed to persist after a few picoseconds, suggesting that there seem to remain charges in the crown at long time delay. This becomes clearer at elevated pump fluences. At 79 μJ/cm², T1 partially decays. While the fast decaying part forms the CT state, the remaining excitations in the crown decay slowly (∼100 ps or longer). More importantly, as shown in Figure 3b, the decay kinetics of the remaining excitations in T1 and the CT state are found to be quite similar to each other. This indicates that both the CT state and the remaining excitation in the alloyed-crown are coupled via a shared electronic state (i.e., hole in the alloyed-crown). This coupling is expected to shift the energy levels of the excitations involved for the optical gain, and hence it can explain observed large spectral shift of the ASE peak with respect to the ground-state absorption (i.e., T1) and emission (i.e., CT state) in these Type-II CQWs.

In Figure 3c, we show the normalized absorption (ΔA(t)/A₀) calculated at the ASE peak for each corresponding sample for a pump–probe delay of t = 2.5 ps. The normalized absorption term being larger than 1 indicates the presence of net optical gain in the system. Optical gain lasts for only ∼10 ps in the CdSe/CdTe CQW, which explains the large ASE thresholds because gain active species are rapidly depleted before they could contribute to gain. The alloyed-crown CQWs, on the contrary, exhibit substantially longer optical gain lifetimes. Also, the gain amplitude is considerably larger in the alloyed-crown CQWs.

Next, we investigate the net modal gain coefficients in the alloyed-crown CQWs using variable stripe length (VSL) method as a function of the pump fluence. (See the Supporting Information for the experimental details.) Emission spectrum was measured as a function of the excitation stripe length, while the width of the stripe was kept fixed at ∼120 μm.

In Figure 3d, we depict the gain lifetimes for each sample. The best performing sample (CdSe/CdSe₀.5Te₀.5 CQW) attained a very long gain lifetime τ_gain ≈ 400 ps, almost three times larger as compared with the previous best reported measurements using heterostructured colloidal nanocrystals (i.e., core/shell CQWs and CdSe/CdS nanorods).

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WINVERSION can be expressed as ineq 3, where the population inversion. The density of the population peak modal gain coefficient, and \( \alpha(\lambda) \) is the waveguide loss coefficient. The term \( G(\lambda) - \alpha(\lambda) \) is the net modal gain coefficient. Equation 1 provides an excellent fit to the VSL data. (See Figure 4a.) In Figure 4b we present the spectra of the net modal gain coefficients at increasing fluence levels, which could be fitted with a Gaussian function having a width of \( \sim 9 \) nm (similar to the fwhm of the ASE peak). As the pump fluence is increased, the net modal gain coefficients grow and also slightly blue-shift, possibly due to the state-filling effect and multiexciton Coulomb interactions.

Figure 4c plots the net peak modal gain coefficients as a function of the pump fluence. At low pump fluence, net modal gain coefficient increases linearly, then a gradual saturation is observed at elevated excitation levels. To fit the net peak modal gain coefficients as a function of the pump fluence, we consider the following

\[
I(\lambda, l) = \frac{I_s (e^{G(\lambda) - \alpha(\lambda)} - 1)}{G(\lambda) - \alpha(\lambda)}
\]

(1)

Here \( I(\lambda, l) \) is the measured intensity from the sample, \( l \) is the stripe length, \( I_s \) is the spontaneous emission proportionality constant, \( G(\lambda) \) is the modal gain coefficient, and \( \alpha(\lambda) \) is the waveguide loss coefficient. The term \( G(\lambda) - \alpha(\lambda) \) is the net modal gain coefficient. Equation 1 provides an excellent fit to the VSL data. (See Figure 4a.) In Figure 4b we present the spectra of the net modal gain coefficients at increasing fluence levels, which could be fitted with a Gaussian function having a width of \( \sim 9 \) nm (similar to the fwhm of the ASE peak). As the pump fluence is increased, the net modal gain coefficients grow and also slightly blue-shift, possibly due to the state-filling effect and multiexciton Coulomb interactions.

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\[
\text{net modal gain} = G - \alpha = \Gamma \times N_{\text{pop-inv}} \times \sigma_s
\]

(2)

\[
N_{\text{pop-inv}} = \frac{W_p}{1 + \frac{W_p}{W_c}}
\]

(3)

\[
G - \alpha = \Gamma \times \sigma_s \times \frac{W_p}{1 + \frac{W_p}{W_c}}
\]

(4)

where \( \Gamma \) is the optical mode confinement factor, \( \sigma_s \) is the stimulated emission cross-section, and \( N_{\text{pop-inv}} \) is the density of the population inversion. The density of the population inversion can be expressed as in eq 3, where \( W_p \) is the pump intensity and \( W_c \) is the saturation pump intensity. From the fit, we find \( W_c \) to be 1 mJ/cm² and \( \alpha \) to be \( \sim 50 \) cm⁻¹. The net peak modal gain coefficient is as high as 929 cm⁻¹. Thus achievement of very large gain coefficients from the alloyed-crown Type-II CQWs makes them exciting not only for colloidal lasers but also for high-speed applications including optical switches.

To find the modal gain coefficients \( G \), we measure the waveguide loss coefficient in these CQW thin films. For this, we measured the ASE intensity as a function of the distance from the edge of the sample. (See the inset of Figure 4d and Supporting Information for the experimental details.) We fit the data by using the relation: \( I_{\text{ASE}} = I_0 e^{-\alpha l} \), where \( I_0 \) is the maximum ASE intensity and \( l \) is the distance from the edge of the sample and \( \alpha \) is the waveguide loss coefficient. Here \( \alpha \) is found to be 46.0 ± 1.2 cm⁻¹, which is in good agreement with the fitting by eq 4. Relatively high waveguide loss coefficient in the CQW film could be attributed to their needle-like face-to-face assemblies, likely to increase the scattering losses. (See Figure S12.) We identify \( \sim 22 \) mJ/cm² as the transparency excitation density. Furthermore, the extrapolation of the modal gain coefficients below the transparency point suggests that the modal gain coefficient starts increasing without a clear threshold (shown by the dotted line in Figure 4d extrapolated to cross at the origin). This indicates that the Type-II CQWs could accomplish significantly lower gain thresholds if the waveguide loss coefficient (i.e., system loss) would be further reduced. Moreover, this gain behavior in the alloyed-crown CQWs, which resembles a four-level-like gain system, is expected to make them promising for ultralow threshold lasers based on low loss microcavities.

In conclusion, we explored optical gain in the CQWs having a Type-II electronic structure. Tuning the composition of the crown via alloying allowed us to realize exceptional optical gain performance in these colloidal nanocrystals. Also, we elaborated that the gain arises from the excitations in the crown that are coupled to the CT state at the Type-II interface. A deeper understanding of the crown’s excitation-CT state coupling will be important for designing new colloidal heterostructures as ultraefficient optical gain media. These alloyed-crown quantum-well structures are expected to inspire novel colloidal gain media that will open up exciting revenues toward practical nanocrystal lasers.

**METHODS**

**Synthesis of the 4 ML CdSe CQWs.** CdSe nanoplatelets having four-monolayer (4 ML) thickness were synthesized with the modified recipe from the literature. For a typical synthesis, 340 mg of cadmium myristate, 24 mg of Se, and 30 mL of octodecene (ODE) were loaded into a three-necked flask. The solution was degassed at room temperature for half an hour and heated to 240 °C under an inert atmosphere. When the temperature reached 195 °C, 120–130 mg of cadmium acetate dihydrate was injected into the reaction solution having bright yellowish color. After 10 min of the growth at 240 °C, the temperature of the solution was decreased to room temperature with the injection of 1 mL of oleic acid (OA). With the purification of as-synthesized CdSe CQWs and the removal of other side-products including CQWs having different thicknesses and quantum dots, 4 ML thick CdSe CQWs can be achieved having an emission peak around 513 nm with a moderately high quantum yield (30–50%). Finally, 4 ML thick CdSe CQWs were dissolved in hexane and stored for next steps.

**Preparation of Anisotropic Growth Mixture for CdSe, Te₇₋ₓ Alloyed-Crown Region.** Anisotropic growth mixture was prepared according to a slightly modified recipe. For a typical cadmium precursor preparation, 480 mg of cadmium acetate dihydrate, 340 µL of OA, and 2 mL of ODE were mixed in a three-necked flask. The solution was heated to 150 °C under ambient conditions with rigorous stirring while also regularly sonicating. After the formation of a homogeneous gel with whitish color, the cadmium precursor was cooled to room temperature and stored for the crown coating. Te-TOP-ODE (0.03 M) solution was used for the tellurium (Te) precursors. It was obtained by the dilution of Te-TOP (1 M) solution with ODE and prepared inside the glovebox. Se-TOP-ODE (0.03 M) solution was used for the selenium (Se) precursors. It was obtained by the dilution of Se-TOP (1 M) solution with ODE and prepared inside the glovebox.

**Synthesis of 4 ML Thick Core/Alloyed-Crown CdSe/CdSe₆Te₁₋ₓ CQWs.** For typical crown coating, 2 mL of 4 ML thick CdSe CQWs dissolved in hexane (having an optical density of 0.5 at 350 nm), 5 mL of ODE, 40 µL of OA, and 0.39 mL of cadmium precursors were put into a three-necked flask. First, the solution was evacuated at room temperature for the complete
removal of hexane. Then, the solution was further evacuated around 100 °C for the removal of other residuals including water and any other organic solvents. Following that, the temperature of the solution was increased to 240 °C under an argon flow for the coating of the crown region. When the temperature reached 240 °C, a desired amount of tellurium precursor (Te-TOP-ODE (0.03 M)) started to be injected at the rate of 8 mL/h. After the complete injection of Te precursor, the synthesis was further annealed for 5 min at 240 °C. Finally, the reaction was stopped with the injection of 1 mL of OA and cooled to room temperature. In addition, depending on the injection amount of Te precursor (between 1.0−2.0 mL), different sizes of the CdTe crown region can be grown.

For the synthesis of CdSeTe1−x alloyed-crown region, we also followed the same recipe with the synthesis of pure CdTe crown region. Simply changing the ratio of Se to Te in the injection precursor, we performed the synthesis with the same amount of injection, which is used for the growth of pure CdTe crown region. For the samples used in these studies, 1.5 mL of injection precursors having different ratios (Te to Se) was used. For the further characterization of CdSe/CdTe core/crown and core/alloyed crown QWJs, they were purified with successive precipitation and finally dispersed in hexane.

PL QE measurement: We compared the emission intensity to Rho6G having 95% PL QE.

■ ASSOCIATED CONTENT
5 Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcl.7b02367.

TEM of the core-only QWJs, XPS analysis of the alloyed-crown composition, photograph of the spin-coated and encapsulated samples, statistical distribution of the ASE thresholds, and photoluminescence lifetime components and their contributions. (PDF)

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■ ACKNOWLEDGMENTS
We acknowledge the financial support partly from National Research Foundation, Prime Minister’s Office, Singapore under its NRF Investigatorship Award (NRF-NRFI2016-08) and A*STAR SERC PHAROS Program (152 73 00025) of Singapore and partly from EU-FP7 Nanophotonics4Energy NoE and TUBITAK 114F236 and 114E449. H.V.D. acknowledges additional support from ESF-EURYI and TUBA, and Y.K. acknowledges support from TUBITAK-BİDEB. T.L. also gratefully acknowledges the financial support from the U.S. National Science Foundation (CHE-1309817). We thank Prof. Alexander L. Efros for the discussions on the repulsive exciton−exciton interactions.

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