Electron Stability and Negative-Tetron Luminescence in Free-Standing Colloidal n-Type CdSe/CdS Quantum Dots
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Abstract. We examine the effects of CdS shell growth on photochemical reduction of colloidal CdSe quantum dots (QDs) and describe the spectroscopic properties of the resulting n-type CdSe/CdS QDs. CdS shell growth greatly slows electron trapping. Because of this improvement complete two-electron occupancy of the 1Se conduction-band orbital is achieved in CdSe/CdS QDs and found to be much more stable than in past experiments. Simultaneous photoluminescence at two different energies is now observed from QDs possessing two excess conduction-band electrons, reflecting competing recombination of discretized 1Sr and 1Pe conduction-band electrons within photogenerated four-carrier negative tetrons (three electrons and one hole). Stable occupancy of the 1Pe level is not achievable under these conditions, and possible reasons are discussed. The stability and accessibility of these multielectron configurations, and the facile spectroscopic detection of negative tetrons, both make photodoped core/shell QDs attractive for exploring the physical properties of free-standing heavily n-doped colloidal CdSe-based QDs.

Introduction
A fundamental understanding of the electronic structures of colloidal semiconductor quantum dots (QDs) is vital to the application of this class of materials in future technologies. Attractive due to their size-tunable physical properties and solution processability, free-standing colloidal QDs suffer from surface defects that have presented longstanding barriers to many practical applications by limiting luminescence quantum yields, affecting Fermi levels, and ultimately impacting performance in various QD-based devices. Surface modification is commonly employed to mitigate the influence of surface traps, most often in the form of passivating small-molecule surface ligation or shell overgrowth. Through judicious selection of the core and shell parameters, the latter can also be used to strategically tune the material’s electronic structure for enhancement of target functional properties. For example, the CdSe/CdS core/shell
heterostructure has long attracted interest because of its quasi-type II band alignment, in which conduction-band electrons ($e_{\text{CB}}^-$) delocalize throughout the heterostructure but valence-band holes ($h_{\text{VB}}^+$) localize at the CdSe core.\textsuperscript{1–11} This spatial separation of charge carriers prolongs exciton radiative lifetimes, slows Auger recombination, enhances photocatalysis, suppresses luminescence intermittency (“blinking”), and eliminates some nonradiative surface recombination pathways.

Electronic doping is a powerful approach for studying the electronic structures of semiconductor nanocrystals.\textsuperscript{12} Among the various methods for introducing excess delocalized charge carriers, photodoping has proven particularly useful for achieving reversible and quantifiable tuning of electron occupancies in free-standing oxide and chalcogenide QDs.\textsuperscript{12–22} Here, we use photodoping to incorporate excess $e_{\text{CB}}^-$ into colloidal CdSe/CdS core/shell QDs with various shell thicknesses. We show that the excess conduction-band electrons become increasingly stable at room temperature with increasing CdS shell thickness. This stability allows detailed chemical and spectroscopic characterization of the resulting n-doped core/shell QDs in the absence of electrodes, excess reductant, or other potential perturbations. In QDs containing two extra $e_{\text{CB}}^-$, we observe separate photoluminescence (PL) bands from recombination of discretized $1S_e$ and $1P_e$ electrons as well as corresponding dynamic signatures of four-carrier negative tetrons (three $e_{\text{CB}}^-$ and one $h_{\text{VB}}^+$). Discrete $1S_e$ and $1P_e$ PL bands have been observed in other CdSe-based QDs under electrochemical bias,\textsuperscript{23,24} and the dynamics of negative tetrons have been described for CdSe/CdS QDs.\textsuperscript{22,25} Here, we quantify these two phenomena together in conjunction with chemical titration analysis of the extent of n-doping to demonstrate the relationship between them. The findings presented here improve our understanding of electronic materials relevant to various QD optoelectronic technologies and point to the use of photodoped n-type CdSe/CdS core/shell QDs for future fundamental electronic-structure studies of multielectron QDs.

Results and analysis

**Photodoping, Electron Stability, and Electron Titration.** Figure 1 shows TEM images of a representative series of CdSe/xCdS core/shell QDs prepared from the same CdSe cores but with different CdS shell thicknesses of $x = 2$, 4, and 7 monolayers. The average QD diameter increases with increasing shell thickness, as does the nanocrystal faceting, and a small increase in the size distribution is also observed.
Core/shell QDs were reduced photochemically following the approach described in detail previously,\textsuperscript{14,19} in which Li-[Et\textsubscript{3}BH] pre-reduces the QD surfaces, followed by photoexcitation and hole capture. Figure 2 plots normalized absorption and continuous-wave photoluminescence (CW PL) spectra of as-prepared (solid) and maximally photodoped (dashed) CdSe/xCdS QDs with increasing shell thickness (top to bottom). The combination of large valence-band offset and small conduction-band offset between CdSe and CdS leaves holes largely confined to the CdSe volumes but permits delocalization of electrons throughout the entire heterostructure volume,\textsuperscript{27} resulting in a redshift in absorption and PL energies of the as-prepared QDs with increasing CdS shell thickness. The PL quantum yield (QY) of the as-prepared QDs also increases, from \(\sim 3\%\) at \(x = 0\) to \(\sim 73\%\) at \(x = 7\). Resonant photoexcitation of the CdSe feature still leads to photodoping (see Figure S1), demonstrating hole capture at the surface despite confinement to the CdSe core.
Figure 2. (A) Visible and infrared (1S_e→1P_e) electronic absorption and (B) visible CW PL spectra of as-prepared (solid) and maximally photodoped (dashed) CdSe/CdS QDs with x = 0, 2, and 7 monolayers of CdS shell (from top to bottom) deposited on d = 4.3 nm CdSe QD cores. Major solvent peaks have been removed from the IR spectra for clarity. With increasing shell thickness, ⟨n_max⟩ increases. CW PL intensity is still observed at ⟨n_max⟩ for the shelled samples.

Concomitant with these changes, the maximum extent of photodoping that can be achieved also increases with increasing shell thickness, as gauged by the bleach of the first excitonic absorption peak (1S_h→1S_e). In the core CdSe QDs (x = 0), this absorbance is reduced by only ~35% before further photoexcitation no longer generates additional 1S_h→1S_e bleach, whereas in the largest CdSe/CdS QDs (x = 7), the first exciton is bleached by ~90% at maximum photodoping. The average number of excess e_sp CB per QD ⟨n⟩ can be quantified by analyzing the extent of this bleach using the relationship ⟨n⟩ = (2ΔA/A0). At maximum photodoping, we define ⟨n⟩ = ⟨n_max⟩. The CdSe QD cores without any shell reach only ⟨n_max⟩ ~ 0.7, but depositing just two monolayers of CdS shell increases ⟨n_max⟩ to 1.6, and at 7 shell monolayers ⟨n_max⟩ = 1.8. We have previously shown that ⟨n_max⟩ in CdSe QDs reflects the steady-state balance between photoreduction and electron trapping and that increases in ⟨n_max⟩ correlate with stabilization of the conduction-band-edge potential to reduce the number of midgap electron traps. We have also shown that even though midgap electron traps below the Fermi level are filled by photodoping, new midgap traps appear spontaneously on long time scales because of trap-state fluctuations. Following these observations, the trend of increasing ⟨n_max⟩ with CdS shell growth is attributed to (1) reduction of the number of midgap surface
electron traps through physical passivation, and (2) shifting of the conduction-band edge to more positive redox potentials relative to the electron-trap-state distribution through relaxation of electron quantum confinement.\textsuperscript{21} With these two effects, conduction-band electrons have access to fewer midgap surface traps after CdS shell growth, leading to lower electron-trapping rates (\textit{vide infra}) and hence greater values of \( \langle n_{\text{max}} \rangle \).

In the unshelled \((x = 0)\) CdSe QDs, the CW PL is almost completely quenched at \( \langle n_{\text{max}} \rangle \) due to efficient Auger recombination. Statistically, each thin-shelled \((x = 2)\) CdSe/CdS QD contains at least one \( e_{\text{CB}}^- \) at \( \langle n_{\text{max}} \rangle \sim 1.6 \), and yet considerable PL intensity remains. The thick-shelled \((x = 7)\) QDs also still luminesce even at \( \langle n_{\text{max}} \rangle \sim 1.8 \). The remaining CW PL intensity is attributed to the reduced nonradiative Auger decay rate of negative trions in these QDs, due to the combination of relaxed quantum confinement\textsuperscript{15} and increased spatial separation between charge carriers\textsuperscript{8} with increasing CdS shell thickness.

The fluctuations described above cause new midgap electron traps to appear spontaneously over very long time scales, causing slow electron trapping even under rigorously anaerobic conditions.\textsuperscript{19,21} This slow trapping is manifested as an extremely slow recovery of excitonic absorption after photodoping.\textsuperscript{21} Detailed studies have demonstrated that this trapping follows uninomolecular reaction kinetics and hence cannot be attributed to the trivial scenario of diffusive oxidants in solution.\textsuperscript{21} Figure 3 summarizes the slow anaerobic absorption recovery observed for photodoped CdSe/xCdS QDs \((x = 2, 4, 7)\). The data are plotted as the normalized first-excitonic absorption bleach \(( -\Delta A/A_0 )\) vs time. Fitting these data yields the electron decay time constants \( (\tau_D) \) plotted in the inset of Figure 3. These data show that increasing the CdS shell thickness reduces the rate of electron trapping markedly. For example, \( \tau_D \sim 15 \) min when \( x = 2 \), but increases to several hours for \( x = 7 \). Photodoped CdSe/7CdS QDs are thus remarkably stable as free-standing colloidal n-type nanocrystals.

\begin{figure}[h]
\centering
\includegraphics[width=0.8\textwidth]{figure3.png}
\caption{Anaerobic \( e_{\text{CB}}^- \) decay kinetics for CdSe/xCdS QDs with \( x = 2 \) (blue open triangle), 4 (green open square), and 7 (red open circle) monolayers, measured in solution at room temperature. The data points represent the normalized absorption at the maximum of the first excitonic absorption transition \((1S_h^- - 1S_e)\) for each sample. The solid lines show exponential fits to the data, which yield the electron decay time constants \( (\tau_D) \) shown.}
\end{figure}
Because the history of the QDs can affect the electron trapping rates, these samples were prepared and handled under identical conditions for these measurements (see Experimental Methods). The small differences in $\langle n \rangle$ at $t = 0$ reflect sample handling and do not impact the primary conclusions drawn from these data. Inset: Plot of $\tau_D$ vs CdS shell thickness.

The high stability of these excess conduction-band electrons enables the use of chemical titration as an independent measure of $\langle n \rangle$. Previous attempts to titrate large ($d = 7.0$ nm), maximally photodoped CdSe QDs were complicated by the presence of additional trapped electrons, presumably at surface sites. Figure 4 shows electronic absorption spectra of maximally photodoped CdSe/7CdS QDs collected during titration with the outer-sphere chemical oxidant, [FeCp*$_2$][BArF$_4$]. With each aliquot of oxidant, the absorption recovers partially. The inset plots $-\Delta A/A_0$ vs added equivalents of [FeCp*$_2$]$^+$ and reveals a linear dependence of the absorption recovery on added oxidant. The intercept of these data with the x axis indicates $\langle n \rangle \sim 2$. This result is consistent with the spectroscopic estimate of $\langle n \rangle = 1.8$ for the same sample. The linearity of these titration data shows that all titratable electrons contribute equally to the absorption bleach, within experimental uncertainty. These data show that few if any electrons reside on the surfaces of these CdSe/7CdS QDs, consistent with nearly complete elimination of midgap electron traps as discussed above.

Figure 4. Absorption spectra of maximally photodoped CdSe/7CdS QDs collected during titration with [FeCp*$_2$][BArF$_4$]. Inset: Normalized absorption bleach at the first excitonic maximum vs added [FeCp*$_2$]$^+$ equivalents. The titration demonstrates that the maximally photodoped QDs contain an average of $\langle n \rangle \sim 2$.

Negative-Tetron Luminescence. Figure 5 shows a more detailed view of the absorption and PL changes that accompany photodoping of one representative sample of CdSe/2CdS QDs. Spectra are plotted for these QDs photodoped to $\langle n \rangle = 0.0$, 0.4, and 1.6. In addition to the increasing bleach of the $1S_h - 1S_e$ absorption feature with increasing $\langle n \rangle$, the main PL peak (ascribed to radiative $1S_h - 1S_e$ recombination) red-shifts with photodoping, for example, by $\sim 5$ meV at $\langle n \rangle = 0.4$, consistent with emission contributions from negative trions. At a value of $\langle n \rangle = 1.6$, this main feature has shifted by $\sim 25$ meV, and a second higher-energy PL band appears. This additional PL feature has been observed in CdSe and CdSe/CdS QD films held
under an applied potential and was assigned to emission involving higher-energy $1P_e$ electrons.\textsuperscript{23,24}

Figure 5. (A) Absorption of CdSe/2CdS QDs measured at $\langle n \rangle = 0$ (thick solid), $\langle n \rangle = 0.4$ (dashed) and $\langle n_{\text{max}} \rangle = 1.6$ (thin solid). (B) PL spectra of the samples from panel A. At $\langle n \rangle > 1$, a second higher energy PL peak is observed.

Figure 6A plots PL spectra for a series of CdSe/xCdS QDs ($x = 0, 2, 7$) collected after photodoping to $\langle n_{\text{max}} \rangle$ in each case. All three spectra show evidence of an upper PL band. $\langle n_{\text{max}} \rangle$ is only 0.7 for the $x = 0$ sample, so the $1P_e$ orbital is essentially unoccupied at steady state (see Figure S7). Moreover, these unshelled CdSe QDs exhibit negligible CW PL at $\langle n_{\text{max}} \rangle$ (Figure 2). Nonetheless, their PL spectrum can be obtained by integrating just the first 800 ps of their time-resolved PL, and a small higher-energy PL band is then observed. PL spectra of the CdSe/2CdS QDs were discussed in Figure 5 and show a well-resolved higher-energy band at high photodoping levels. For the CdSe/7CdS QDs, the PL is much stronger, and the upper PL band is again observed. In this sample, the two bands are closer in energy because of relaxed quantum confinement, and they partially overlap.

Figure 6B plots the energy difference between the two PL peaks in Figure 6A ($\Delta E_{P-S(\text{PL})}$) vs the energy of the first excitonic absorption maximum ($E_{1\text{Sh}-1\text{Se}}$) for these three samples, along with the energy difference between $1S_e$ and $1P_e$ one-electron states within the conduction band, as probed by intraband absorption in the infrared ($\Delta E_{P-S(\text{IR})}$, Figure 2A). Both energies decrease with decreasing electron confinement upon increasing the CdS shell thickness, but $\Delta E_{P-S(\text{PL})}$ is consistently smaller than $\Delta E_{P-S(\text{IR})}$. The difference between
\( \Delta E_{P-S}(PL) \) and \( \Delta E_{P-S}(IR) \) reflects the additional Coulomb interactions present in the initial and final states of the luminescence transitions that are not present in the IR excited states. This correlation thus supports the assignment of this upper PL band to recombination involving 1P\( e \) electrons in QDs containing multiple excess conduction-band electrons. We note that although the trend in Figure 3 would suggest even greater \( \bar{e}C_B \) stability with increasing CdS shell thickness, the data in Figure 6 show that the upper PL band becomes increasingly obscured by the lower PL band with increasing CdS shell thickness. For this reason, CdSe/xCdS QDs with even thicker CdS shells than \( x = 7 \) were not explored in this study.

![Image](image-url)

**Figure 6.** (A) PL spectra for maximally photodoped CdSe/xCdS QDs, where \( x = 0, 2, \) and 7 monolayers. Values of \( \langle n \rangle \) are indicated on each spectrum. The dotted lines indicate the 1S\( e \) and 1P\( e \) emission energies in each spectrum. (B) \( \Delta E_{P-S} \) values determined from IR intraband absorption (Figure 2A) and visible PL (panel A of this figure), plotted vs the energy of the first excitonic absorption feature (E_{1S_e-1S_e}). \( \Delta E_{P-S}(IR) \) and \( \Delta E_{P-S}(PL) \) both decrease with decreasing quantum confinement, and the difference between the two reflects the additional Coulomb interactions probed by the PL measurement. Error bars on \( \Delta E_{P-S}(PL) \) values reflect estimated uncertainties from the peak fitting. \( \Delta E_{P-S}(IR) \) error bars are smaller than the data points themselves.

Figure 7A shows a streak-camera image describing the PL decay dynamics of the CdSe/2CdS QDs from Figure 5 possessing \( \langle n \rangle = \langle n_{\text{max}} \rangle = 1.6 \). The integrated 1S\( e \) and 1P\( e \) PL intensities are plotted vs time in Figure 7B. Although similar, the decay dynamics for the two features do differ slightly. This difference is attributable to the presence of discrete subsets of QDs with n
= 1 and n = 2 within this $\langle n \rangle = 1.6$ ensemble. Nonetheless, these results confirm that both luminescence features seen in Figure 5B ($\langle n \rangle = 1.6$) derive from the same excited state.

To explore the emission of these heavily n-doped QDs more systematically, we measured room-temperature PL decay dynamics of the CdSe/2CdS QDs at various values of $\langle n \rangle$. Figure 8 plots absorption spectra and PL decay curves of these CdSe/2CdS QDs poised at $\langle n \rangle = 0$, 0.5, and 1.8. The as-prepared QDs ($\langle n \rangle = 0$) exhibit purely excitonic emission, and their PL decay can be fit to a single exponential with a time constant $\tau_X = 35$ ns (see Figure S4 for complete fit). When n-type carriers are introduced, PL is enhanced at short times due to the greater radiative rate of the negative trion. Faster decay of the PL is also observed, consistent with a combination of this increased radiative rate and the introduction of nonradiative trion Auger recombination. At $\langle n \rangle = 0.5$, the PL decay is biexponential, reflecting a superposition of negative trion recombination ($\tau_{X^-} = 2.6$ ns) and neutral exciton recombination ($\tau_X$) from the presence of both n-doped and undoped QDs in this ensemble. The value of $\tau_{X^-} = 2.6$ ns is consistent with that predicted from the reported size dependence of $\tau_{X^-}$ in photodoped n-type CdSe QDs using the effective QD diameter, as determined by the energy of the first excitonic absorption feature (see Figure S5). The PL decay is also biexponential at $\langle n \rangle = 1.8$. Here, neutral exciton emission is completely absent because every QD possesses at least one excess $e_{CB}$. Negative trion decay (with $\tau_{X^-}$) is still observed, and in addition a faster process is
observed with a decay constant of $\tau \approx 600$ ps, or roughly $1/4 \tau_{X^-}$. This faster component is interpreted as reflecting PL from QDs with $\langle n \rangle = 2$ that have been photoexcited, thus containing four bandlike charge carriers (three $e_{CB}$ and one hole $h_{VB}$) in their luminescent excited state, that is, a negative tetron ($X^{2-}$). The inset to Figure 8 illustrates carrier configurations relevant to each of the traces in panels A and B. Trace 1 includes only neutral excitons (X), trace 2 includes X and negative trions (X$^-$), and trace 3 includes X$^-$ and X$^{2-}$. Upon reoxidation by air, both the absorption spectrum and PL dynamics return to their original values, being indistinguishable from those of the as-prepared QDs prior to photodoping. A small ($\sim 19\%$ relative) enhancement in overall PL quantum yield is observed after reoxidation, attributed to surface restructuring induced by this reduction/oxidation cycle.

**Figure 8.** (A) Absorption spectra collected at three stages of photodoping for CdSe/7CdS QDs: (1) $\langle n \rangle = 0$, (2) $\langle n \rangle = 0.5$, and (3) $\langle n \rangle = 1.8$. The black dotted trace shows the spectrum collected after QD reoxidation to $\langle n \rangle = 0$ and is indistinguishable from the initial trace, demonstrating complete sample recovery. (B) PL decay curves corresponding to (1), (2), and (3) from panel A ($\lambda_{exc} = 405$ nm, 3.06 eV). The amplitude of the reoxidized curve (dotted) has been reduced ($x = 0.84$) for clarity to account for mild sample brightening upon reoxidation. Thin solid traces illustrate biexponential fits for $\langle n \rangle > 0$ as described in the main text. Inset: Schematics illustrating the three relevant multicharge configurations observed in panels A and B: neutral exciton, negative trion, and negative tetron (from left to right).

Figure 9 summarizes these findings. Figure 9A plots the fractional contributions from each of the carrier configurations to the PL dynamics vs $\langle n \rangle$ for the same CdSe/7CdS QDs, deduced from global fitting of a more extensive series of PL decay curves (see Figure S6 and Experimental Methods). As described above, at values of $\langle n \rangle = 0$, the PL is entirely attributable
to neutral X. In the regime $0 < \langle n \rangle < 1$, the decay is biexponential with components corresponding to $\tau_X$ and $\tau_{X^-}$. At $\langle n \rangle > 1$, a third component ($\tau_{X^2^-}$) is introduced.

Similarly, Figure 9B plots the fractional $1S_e$ and $1P_e$ emission intensities vs $\langle n \rangle$ extracted from PL spectral fitting (see Figure S2). At $\langle n \rangle < 1$, the $1P_e$ emission is absent within experimental uncertainty, but when $\langle n \rangle > 1$, the $1P_e$ emission appears and increases in intensity. The relative intensity from the $1S_e$ emission is inversely correlated with that from the $1P_e$ emission, decreasing only at $\langle n \rangle > 1$. Overall, the PL QY decreases with increasing $\langle n \rangle$ (see Figure S6). The $1S_e$ data plotted in Figure 9B compare well to the sum of X and $X^-$ decay amplitudes plotted in Figure 9A, whereas the $1P_e$ data in Figure 9B show the same trend as seen for the $X^2^-$ decay amplitudes in Figure 9A.

![Figure 9](image)

**Figure 9.** (A) Relative amplitudes of PL decay components arising from neutral excitons (X), negative trions ($X^-$), and negative tetrions ($X^{2^-}$) in n-doped CdSe/CdS QDs plotted vs $\langle n \rangle$, obtained from fitting of PL decay curves collected for each value of $\langle n \rangle$. (B) Intensities of PL peaks assigned to recombination of discrete $1S_e$ and $1P_e$ electrons in n-doped CdSe/CdS QDs plotted vs $\langle n \rangle$, determined by analysis of PL spectra collected for each value of $\langle n \rangle$ (see Experimental Methods and Figure S2 for details).

The data in Figure 9 show that tetron luminescence, which can only appear when $n = 2$ in a given QD prior to photoexcitation, turns on rather abruptly at $\langle n \rangle > 1$. Our previous studies have demonstrated diffusion-limited inter-QD electron transfer following such photodoping.\textsuperscript{17,30,31} Because of this rapid electron transfer, there is Fermi level equilibration within the ensemble of QDs such that the electron distributions among QDs are not determined by Poissonian photoexcitation statistics but rather by $\langle n \rangle$ and the ensemble’s distributions in QD band-edge potentials and multielectron configurational energies. The abrupt appearance of tetron luminescence at $\langle n \rangle > 1$ in Figure 9 thus reflects the tendency of the added electrons to
distribute rather evenly across the ensemble of QDs, regardless of where they were initially generated.

From these PL decay dynamics, it is also possible to estimate radiative rate constants \(k_r\), nonradiative Auger rate constants \(k_A\), and PL QYs for the \(X\), \(X^-\), and \(X^{2-}\) electronic configurations of these QDs. Following prior analysis of multistate blinking in CdSe/CdS QDs, these parameters can be quantified using eqs 1−3.\(^8\)\(^,\)\(^25\) Here, \(\beta\) is a carrier overlap factor that relates the radiative recombination rate of a charged exciton to that of the neutral exciton, and \(i\) denotes the carrier configuration. Table 1 summarizes the values of these quantities determined from the dynamics measured for the CdSe/7CdS QDs (Figures 8 and S6). Statistically, \(k_A^{X^{2-}}\) should be 3x larger than \(k_A^{X^-}\) because of the increased number of possible recombination pairings, consistent with the factor of \(~4\) measured experimentally. Different radiative and nonradiative transition probabilities can be anticipated for recombination involving \(1S_e\) and \(1P_e\) electrons, which will cause deviations from simple statistics. The values in Table 1 were obtained by using the integrated PL decay curves to estimate QYs at different values of \(\langle n \rangle\) relative to the QY at \(\langle n \rangle = 0\), which was measured independently using an integrating sphere (see Figure S6).

\[
k_i^i = \beta^i k^X
\]
\[
k_i = k_i^i + k_A^i = 1/\tau_i
\]
\[
QY_i = \frac{k_i^i}{k_i^i + k_A^i}
\]

**Table 1.** Recombination Parameters Determined for Various Neutral and Charged Excitonic Excited States of CdSe/7CdS QDs

<table>
<thead>
<tr>
<th>(i)</th>
<th>QY</th>
<th>(\tau_i) (ns)</th>
<th>(1/k_r^i) (ns)</th>
<th>(1/k_A^i) (ns)</th>
<th>(\beta^i)</th>
</tr>
</thead>
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<td>(X)</td>
<td>0.73(^a)</td>
<td>35</td>
<td>35</td>
<td>–</td>
<td>1</td>
</tr>
<tr>
<td>(X^-)</td>
<td>0.11</td>
<td>2.6</td>
<td>23.6</td>
<td>2.9</td>
<td>1.48</td>
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<tr>
<td>(X^{2-})</td>
<td>0.06</td>
<td>0.6</td>
<td>10</td>
<td>0.6</td>
<td>3.5</td>
</tr>
</tbody>
</table>

\(^a\) Measured independently using an integrating sphere.

The \(\beta\) values determined here yield information about the Coulomb forces acting between carriers. The experimental \(\beta^{X^-}\) value is smaller than the statistical expectation (\(\beta^{X^-} = 2\)), and it is also smaller than the value observed in CdSe QDs (\(\beta^{X^-} = 2.2\)),\(^32\) but it agrees well with the value measured for CdSe/CdS QDs overcoated with silica (\(\beta^{X^-} = 1.49\)). This reduced value of
\( \beta^X \) is consistent with diminished carrier overlap in these core/shell QDs because of proper core/shell band alignment for electron but not hole delocalization.\(^25\) The value of \( \beta^X \) in Table 1 is comparable to but slightly larger than that expected purely statistically (\( \beta^X = 3 \)).

**Discussion**

The results and analysis presented above show that growth of CdS shells around colloidal CdSe QDs markedly improves the stability of excess conduction-band electrons added via photodoping. The resulting colloidal n-type QDs can be stabilized for hours in solution at room temperature and are thus amenable to investigation by various chemical and spectroscopic methods. Similar to unshelled CdSe QDs,\(^21\) we find that the maximum number of conduction-band electrons that can be added to these core/shell QDs via photodoping, \( \langle n_{\text{max}} \rangle \), increases with increasing QD diameter across a well-controlled series of related QDs. For the present QDs, we can add as many as two excess electrons per QD via photodoping, but not more. This maximum is confirmed by direct chemical titration of the excess electrons. This value of \( \langle n_{\text{max}} \rangle \) is somewhat larger than we found for unshelled CdSe QDs (\( \langle n_{\text{max}} \rangle < \sim 1.6 \)),\(^18,21\) but not dramatically so. The greatest contrast between shelled and unshelled CdSe QDs is in the long-term stability of these excess electrons. An interesting question pertains to the precise origin of \( \langle n_{\text{max}} \rangle \), which remains unclear. We have previously demonstrated\(^18,21\) that \( \langle n_{\text{max}} \rangle \) for (unshelled) CdSe QDs is not related to the specific orbital occupancies of the QDs in any way, because in some cases \( \langle n_{\text{max}} \rangle \) occurs at values well below 1, meaning many QDs within the ensemble actually possess no conduction-band electrons at maximum photodoping. Such data suggest an important role of surface traps in determining the most-negative Fermi level achievable during photodoping.\(^18,21\) For the present CdSe/CdS QDs, \( \langle n_{\text{max}} \rangle \) reaches nearly 2 with the thickest of our CdS shells (\( x = 7 \)), but we see no evidence for steady-state occupancy of the 1P\(_e\) levels in either the PL spectra or the PL decay data. We hypothesize that such 1P\(_e\) electrons, which would be at least \( \sim 150–300 \) meV (variable with quantum confinement, Figure 2) less stable than the 1S\(_e\) electrons, trap at much faster rates than the 1S\(_e\) electrons even in these core/shell heterostructures because of a greater trap density of states at these potentials, and this rapid trapping precludes stable 1P\(_e\) occupancy. Maximum carrier densities are thus limited to \( \langle N \rangle \sim 10^{19} \) cm\(^{-3}\) in these CdSe/CdS QDs, compared to an order of magnitude larger carrier densities found for photodoped metal-oxide (ZnO, In\(_2\)O\(_3\), and related) nanocrystals.\(^13,33\)

Previously, remote chemical dopants were also found to introduce no more than two excess electrons per QD into CdSe QDs.\(^14\) Interestingly, even when electrons are introduced
electrochemically, CdSe QD films still allow only partial filling of the 1P\textsubscript{e} shell.\textsuperscript{34} It is likely that both of these observations also reflect the accessibility of surface electron traps at the fairly negative potentials of the 1P\textsubscript{e} electrons. During preparation of this manuscript, another study of the same n-type photodoping of CdSe/CdS QDs using Li[Et\textsubscript{3}BH] was published in which values as large as \langle n \rangle = 6 are reported.\textsuperscript{22} It is conceivable that in these QDs, the density of surface traps was diminished significantly compared to the other samples discussed above, allowing accumulation of many more electrons per QD before reaching the potentials of these traps. We note, however, that \langle n \rangle values in ref 22 were determined by analysis of the first exciton’s absorption bleach under the assumption that distributions of excess electrons among QDs reflect the Poissonian QD photoexcitation statistics, without consideration of subsequent inter-QD electron transfer. Under this assumption, QDs with very large values of \( n \) coexist with QDs having no excess electrons, and much larger values of \langle n \rangle are thus required to fully bleach the first excitonic absorption feature. Only the 1S\textsubscript{e} electrons contribute to this bleach. Such a broad Poissonian distribution of electrons among QDs appears inconsistent with our previous observations of rapid inter-QD electron transfer,\textsuperscript{17,30,31} the abrupt appearance of tetron PL when increasing from \langle n \rangle < 1 to \langle n \rangle > 1 (Figure 9), and the confirmation by direct chemical titration that a full bleach of the first excitonic absorption feature is achieved at \langle n \rangle \sim 2 (Figure 4).

More generally, the capability to incorporate multiple, robust n-type carriers into high-optical-quality colloidal CdSe QDs via photochemical reduction described here presents exciting opportunities to investigate the various physical properties of these unusual materials. For example, long-lived spin coherence has been demonstrated in a host of singly reduced semiconductor QDs generated as films on electrode surfaces.\textsuperscript{35} In the case of the tetrons described here, the spin configuration reduces to that of a neutral exciton because the excess carriers form a paired singlet with zero spin. These negative tetrons may therefore prove of fundamental interest from the standpoint that the one unpaired electron occupies the 1P\textsubscript{e} orbital rather than the 1Se orbital, and spin-sensitive techniques such as time-resolved Faraday rotation (TRFR) or magnetoluminescence could thus be used to selectively probe specifically the 1P\textsubscript{e} electron. TRFR experiments on electrochemically reduced CdSe QDs do demonstrate a convergence of two \( g \) values in neutral QDs to a single resonance under negative bias,\textsuperscript{36} attributed to suppression of excitonic precession and demonstrating feasibility of such measurements on n-doped QDs. Stable photodoping may also help advance the understanding of dopant-carrier magnetic-exchange interactions involving impurity spin centers in colloidal CdSe QDs. For example, photodoping of Zn\textsubscript{1-x}Mn\textsubscript{x}O QDs was shown to induce long-range...
carrier-mediated Mn$^{2+}$-Mn$^{2+}$ exchange coupling and modify Mn$^{2+}$ spin-relaxation rates. Similar experiments in Cd$_{1-x}$Mn$_x$Se QDs could take advantage of the ability to retain excitonic PL in these doped QDs, allowing use of PL and magneto-PL to probe such exchange interactions. All of these experiments will benefit from the ability to generate and stabilize excess charge carriers in high-optical-quality n-doped CdSe-based QDs.

**Conclusion**

Colloidal CdSe/CdS core/shell QDs were photochemically reduced, incorporating up to two conduction-band electrons per QD. The maximum level of n-doping was shown to increase as a function of shell thickness, reflecting suppression of surface electron trapping. CdS shelling vastly enhances carrier stability in colloidal CdSe QDs, with excess electrons stable on time scales of hours in solution at room temperature, which in turn enables accurate quantification of the number of excess charge carriers per QD via direct chemical titration. Discrete emission bands involving recombination of 1S$_e$ and 1P$_e$ electrons are observed at values of $\langle n \rangle > 1$, both decaying with similar rate constants, indicating formation of four-carrier negative tetrons. Analysis of the PL decay dynamics confirms this assignment. These results highlight the exceptional control afforded by photodoping for incorporating excess conductionband electrons into freestanding colloidal CdSe/CdS QDs and suggest opportunities for development of novel nanomaterials with tailored electronic configurations relevant to future applications in optoelectronics, photoredox chemistry, and other technologies.

**Experimental Methods**

**General Considerations.** Unless stated otherwise, all syntheses and sample preparations were performed using standard Schlenk line technique under a dinitrogen atmosphere, or in a nitrogen-filled glovebox with anhydrous solvents.

**Chemicals.** Cadmium oxide (CdO; 99.99+%), 1-octadecene (ODE; 90%), oleic acid (OA; 90%), hexadecylamine (HDA; 90%), trioctylphosphine oxide (TOPO; 99%), selenium metal (Se; 99.99+%), and 1-octanethiol (OT; >98.5%) were purchased from Sigma-Aldrich. Trioctylphosphine (TOP; 97%) was purchased from Strem. All chemicals were used as received.

**Quantum Dot Synthesis.** Colloidal CdSe QDs were synthesized using a hot-injection procedure adapted from ref 41. In a 100-mL 3-neck round bottom flask, a stirred solution of
150 mg of CdO, 12g of ODE, and 1g of OA was degassed under vacuum at 110°C for 1 hour. The solution was then heated to ~280°C under N$_2$ until the red solution turned clear and colorless. The solution was allowed to cool to 100°C and was then placed under vacuum for 1 hour to remove O$_2$ produced by forming the Cd(oleate)$_2$. 2g of HDA and 2g of TOPO were then added to the flask under an overpressure of N$_2$, followed by 30 minutes of vacuum and 3 purge cycles with N$_2$. After purging the system, the flask was heated to 300°C under N$_2$. In a separate 25-mL flask, 190 mg of Se was dissolved in 4 mL of TOP and then rapidly injected into the hot Cd(oleate)$_2$ solution. After 5 minutes, the heating mantle was removed and the cooled solution of QDs was purified by flocculation with ethanol and methanol followed by resuspension in toluene.

CdS shells were grown onto the CdSe cores using a procedure adapted from ref 7. For a typical shell growth, a 0.16 M Cd(oleate)$_2$ solution in ODE and a 0.20 M solution of OT in ODE were separately injected at 2.5 mL/hour to a degassed solution containing 200 nmol of CdSe cores in 4g of ODE and 2g of HDA at 200°C. Once the injection was started, the temperature of the solution was raised to 300°C and held there until the injection was complete. The total volume of each solution injected was varied depending on the desired shell thickness, while the volumes for the two shell precursor solutions were kept equal. After the injection was complete, the solution was cooled to room temperature and the QDs purified by flocculation with ethanol and methanol followed by resuspension in toluene.

**Physical Characterization.** Transmission electron microscope (TEM) images were obtained using a FEI Tecnai G2 F20 operating at 200 kV. Quantum yields were measured on colloidal QD suspensions by exciting samples with a Xe lamp coupled to a monochromator ($\lambda_{\text{exc}} = 450$ nm) and detecting in an external quantum efficiency integrating sphere (Hamamatsu C9920-12). All spectroscopic measurements were performed at room temperature on suspensions of colloidal QDs in toluene. Absorption spectra were measured in a Cary 500 (Varian) spectrophotometer. QD concentrations were determined using the absorption intensity at the first excitonic peak and a calibration curve for CdSe QDs.$^{42}$ For infrared (IR) absorption measurements, QDs were mixed with excess LiBE$_3$H in the dark in a nitrogen-filled glovebox, and the solution loaded into an air-free IR cell with either CaF$_2$ (CdSe/0CdS and CdSe/2CdS in toluene) or KBr (CdSe/7CdS in cyclohexane) windows separated by a 100 µm Teflon spacer. Measurements were collected on an FTIR spectrometer (ThermoScientific Nicolet-8700) with a HgCdTe detector. Continuous-wave PL spectra were measured by exciting the sample with a 5 mW 405 nm laser pointer and detecting on an OceanOptics 2000+ spectrometer. Time-
resolved PL measurements were performed using the frequency-doubled output of a Ti:sapphire laser at a 200-kHz repetition rate (405 nm, 150 fs pulse, 1.2–1.5 nJ/pulse). Decay curves were measured using a monochromator and streak camera with an instrument response function of ~15 ps.

**Photodoping.** Photodoping was performed following the method introduced in ref 14. QDs were dried and resuspended in anhydrous toluene in a nitrogen-filled glovebox prior to photodoping experiments. Samples were diluted to the desired concentration (generally ~100 nM for PL experiments), and 50–1000 equivalents of Li[BEt$_3$H] were added. Samples were irradiated with a 5 mW 405 nm laser pointer to achieve desired level of photodoping. To achieve maximal photodoping, LiBEt$_3$H was often added in small increments to mitigate sample scattering and etching.

**Determination of $\langle n \rangle$.** A solution of CdSe/7CdS QDs in toluene was prepared in a nitrogen-filled glovebox and measured in a 1 cm pathlength quartz cuvette. Spectra were collected using an OceanOptics 2000+ spectrophotometer with an absorption attachment. ~200 equiv of LiBEt$_3$H was added, and photodoping was performed using the OceanOptics light source for photoexcitation until no additional bleach was observed over the course of 30 s. These excess electrons were then quantified via titration by addition of 10-µL aliquots of 9 µM [FeCp$^*$]$_2$[BArF$_4$] in THF. The light source was switched off during aliquot additions to avoid *in situ* photodoping between spectral collections. The titration took <10 min to complete, and thus anaerobic recovery for this thick-shelled sample was considered negligible over the course of the experiment.

**Anaerobic Recovery.** A series of QD samples was photodoped with careful attention paid to minimize differences in sample handling. The samples were prepared with the same optical density at their respective first absorption peak maxima in a solution of 30 µM Li[Et$_3$BH] in toluene. Each was prepared anaerobically in a quartz cuvette equipped with a magnetic stir bar and sealed with a Teflon stopper. A photolysis lamp paired with a monochromator was tuned to selectively photoexcite each sample at its respective first excitonic absorption feature, and the light was focused onto the cuvette, which was set to stir inside the Cary spectrophotometer sample cavity. The power of the photolysis light was the same for all samples (0.5 mW), and each sample was irradiated for a total of 30 min. After 30 min, the lamp was switched off, and absorption scans were intermittently collected to monitor the rate of anaerobic recovery.
**Spectral Analysis.** Absorption spectra at different values of $\langle n \rangle$ were fit to a sum of four Gaussians using a global fitting procedure. Widths were fixed, but energies and intensities were allowed to float to reflect the extent of photodoping and electron accumulation. For analysis of the PL spectra, the spectrum of neutral ($\langle n \rangle = 0$) CdSe/7CdS QDs was fitted. Upon photodoping and with increasing $\langle n \rangle$, this band was allowed to shift to lower energy as a new fixed-width Gaussian feature (ascribed to emission involving recombination of 1P$_e$ electrons) gained intensity at 2.01 eV. For PL decay analysis, global triexponential fitting of the PL decay curves collected for the CdSe/7CdS QD sample at various photodoping levels yielded the fitted values of $\tau$ corresponding to $X$, $X^-$, and $X^{2-}$ listed in Table 1, and the fitted $X$, $X^-$, and $X^{2-}$ amplitudes plotted vs $\langle n \rangle$ in Figure 9A.

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