Single-Exciton Gain and Stimulated Emission Across the Infrared Telecom Band from Robust Heavily Doped PbS Colloidal Quantum Dots

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ABSTRACT: Materials with optical gain in the infrared are of paramount importance for optical communications, medical diagnostics, and silicon photonics. The current technology is based either on costly III−V semiconductors that are not monolithic to silicon CMOS technology or Er-doped fiber technology that does not make use of the full fiber transparency window. Colloidal quantum dots (CQDs) offer a unique opportunity as an optical gain medium in view of their tunable bandgap, solution processability, and CMOS compatibility. The 8-fold degeneracy of infrared CQDs based on Pb-chalcogenides has hindered the demonstration of low-threshold optical gain and lasing, at room temperature. We demonstrate room-temperature, infrared, size-tunable, band-edge stimulated emission with a line width of \( \sim 14 \) meV. Leveraging robust electronic doping and charge-exciton interactions in PbS CQD thin films, we reach a gain threshold at the single exciton regime representing a 4-fold reduction from the theoretical limit of an 8-fold degenerate system, with a net modal gain in excess of \( 100 \) cm\(^{-1} \).

KEYWORDS: Optical gain, colloidal quantum dots, stimulated emission, infrared, doping

The potential for narrower line widths\(^1\) and the lower-than-bulk degeneracy\(^2\) has led to dramatic progress toward successful demonstration of optical gain, stimulated emission,\(^4\) and lasing\(^4\)−\(^7\) in the visible part of spectrum utilizing CdSe-based colloidal quantum dots (CQDs). Therefore, low threshold, band-edge amplified spontaneous emission (ASE) in CQDs has been at the center of intensive research over recent years as a prerequisite toward the demonstration of CQD lasing.\(^2\)−\(^9\) Engineered CQDs with suppressed Auger\(^4\) and photodoping\(^10\) have allowed the realization of low-threshold ASE and lasing\(^1\) at the single exciton regime, in the visible, for CdSe CQD systems that possess a 2-fold degeneracy value.\(^12,13\) Nevertheless, low-threshold band-edge ASE in the near-infrared (NIR), based on CQDs, has remained a challenge due to the high degeneracy of Pb-chalcogenide CQDs. PbS CQDs are the material of choice for solution processed infrared optoelectronics with successful demonstrations in LEDs,\(^14\) solar cells,\(^15,16\) and photodetectors.\(^17,18\) The 8-fold degeneracy of PbS(e) CQDs, however, has hindered the demonstration of low-threshold optical gain and ASE, at room temperature, in the infrared across the telecommunications wavelength band.\(^19,20\)

We posited that a CQD film, robustly n-doped in the heavy doping regime, can address this challenge by utilizing the doping electrons present in the first excited state of the CQDs (conduction band) to reach the population inversion condition at reduced pumping fluence. Previously, the use of charged CQDs has resulted in lowering the ASE threshold in CdSe-based systems employing photodoping,\(^10,11\) an approach that requires inert conditions and that is reversible in time, preventing thus the demonstration of robust, permanently doped CQDs and devices thereof. Therefore, we sought to develop a robust electronic n-doping method for PbS CQDs in the heavy doping regime based on atomic substitutional doping. Over the years, various doping strategies have been employed to CQDs to tune their electronic properties.\(^21\) Prior reports of heavily doped PbS(e) QDs have relied on aliovalent cation substitution: Ag\(^+\) substitution of Pb\(^{2+}\) induces p-type-doped PbSe\(^{22}\) CQDs, while the substitution of Pb\(^{2+}\) by In\(^{3+}\) leads to n-doped PbSe\(^{23}\) CQDs. Remote transfer of electrons from cobaltocene molecules as well as photochemical doping have also been reported as doping mechanisms for n-type PbSe CQDs.\(^24,25\) However, none of the above approaches have led to robust permanent doping, preventing thus their use in...
devices. We instead posited that aliovalent anionic substitution may lead to a more robust doping scheme. Density functional theory (DFT) calculations have theoretically predicted n-type doping upon iodide substitution, yet not experimentally observed.27 Here we sought to use PbS (100) surfaces as a platform to facilitate heavy doping through iodide substitution (Figure 1a). Our selection has been guided by the DFT calculations predicted n-type doping upon iodide substitution, yet not experimentally observed.28,29 However, bare iodide-exchanged samples did not show any absorption bleaching as the DFT calculations predicted (see Supporting Information Section S2.1). The absence of heavy doping, evidenced by the lack of bleaching in the absorption measurements, is ascribed to the presence of oxygen and water in the film upon exposure to ambient conditions. Oxygen and water are efficient oxidants and have been reported as effective p-type dopants in lead chalcogenides.30,31 In order to preserve heavy n-doping in our films under ambient conditions, we submitted our samples to atomic layer deposition (ALD) of alumina with a 2-fold purpose: impeding oxygen to further incorporate in the film, and preventing oxidation caused by oxygen/water adsorbates pre-existent in the films upon their formation. Infilling with alumina (supported by XPS results in Supporting Information Section S3), hence, is crucial to preserving robust n-type doping provided by iodide ligand-exchange procedure. We have observed that only the samples that have undergone iodide doping and ALD encapsulation demonstrate optical bleaching and, herein in this study, are assigned as doped PbS CQD and the samples only with iodine substitution as undoped samples. Control samples of thiol-based ligand exchange chemistry did not show any signatures of heavy doping upon ALD encapsulation, further corroborating the role of iodine as the n-type dopant (see Supporting Information Section S2.1). Absorption spectra at shorter wavelengths show that the higher energy states of the CQDs remain unaffected by the doping process (see Supporting Information Section S2.2).

Figure 1d shows an exciton peak bleach after alumina deposition, indicating that the ALD process was successful in maintaining heavy doping at ambient conditions. We have been able to quantify the doping (number of electrons in the CB per dot) in our samples optically, by analyzing the bleach in the absorption (see Experimental Section in the Supporting Information), while we further confirm the doping level electrically via field effect transistor measurements (see Supporting Information Section S4), as shown in Figure 1e. Small dots have an octahedral shape with Pb-rich (111) facets, while, as the dot diameter increases, their morphology evolves progressively to a cuboctahedron that has six sulfur-rich (100) facets.30 Because (100) surfaces are progressively exposed with increasing PbS CQD size, the doping efficacy of the process increases with the size of the dots. Figure 1e shows that particles smaller than 4 nm in diameter do not undergo this doping process due to the lack of (100) exposed facets, whereas in particles with an exciton peak of more than 1800 nm (solid black lines). After doping of the CQD films the absorption bleaches (red dash lines). (e) The number of electrons in the CB, upon doping, depends on the size of the CQD due to the degree of (100) surface presence that enables doping. Red dots represent the experimentally extracted number of electrons from measuring the bleaching of the absorption of the CQDs, while blue dots represent the ZnI2/MPA hybrid ligand treated CQD films, before and after doping. The CQD sizes are 5.5 and 6.1 nm with respective exciton peaks at 1480 and 1580 nm (solid black lines). After doping of the CQD films the absorption bleaches (red dash lines). (e) The number of electrons in the CB, upon doping, depends on the size of the CQD due to the degree of (100) surface presence that enables doping. Red dots represent the experimentally extracted number of electrons from measuring the bleaching of the absorption of the films at the exciton peak with the use of 1-ethyl-3-methylimidazolium iodide (EMII) for ligand exchange, while blue dots represent the ZnI2/MPA hybrid ligand treated films that are used for ASE and gain measurements. Yellow dots indicate the extracted doping values from electrical FET measurements of transistors based on EMII-treated CQDs. Both ligand treatments are equally effective in the doping of the CQDs. The red area represents the doping range variation based on size distribution of the CQDs, indicating even the doping distribution across the films is ±0.5 electrons. The data have been fitted with a sigmoidal function (black dash line) as a guide to the eye.

Figure 1. (a) Schematic representation of the S2− substitution to I− in (100) surface in large, cuboctahedral-shaped PbS CQDs. (b, c) Calculated density of states of the (100) surface before and after I− substitution showing that the Fermi level, EF, is shifted to the CB. (d) Absorption spectra of two representative PbS CQD films, before and after doping. The CQD sizes are 5.5 and 6.1 nm with respective exciton peaks at 1480 and 1580 nm (solid black lines). After doping of the CQD films the absorption bleaches (red dash lines). (e) The number of electrons in the CB, upon doping, depends on the size of the CQD due to the degree of (100) surface presence that enables doping. Red dots represent the experimentally extracted number of electrons from measuring the bleaching of the absorption of the films at the exciton peak with the use of 1-ethyl-3-methylimidazolium iodide (EMII) for ligand exchange, while blue dots represent the ZnI2/MPA hybrid ligand treated films that are used for ASE and gain measurements. Yellow dots indicate the extracted doping values from electrical FET measurements of transistors based on EMII-treated CQDs. Both ligand treatments are equally effective in the doping of the CQDs. The red area represents the doping range variation based on size distribution of the CQDs, indicating even the doping distribution across the films is ±0.5 electrons. The data have been fitted with a sigmoidal function (black dash line) as a guide to the eye.

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Henceforth, we have employed a hybrid ligand exchange treatment based on ZnI₂/MPA that caters for equally effective doping (as shown in Figure 1e) as well as serves better passivation of the CQDs and higher photoluminescence, as previously reported. 14 The undoped PbS CQD films demonstrate optical gain threshold $\langle N \rangle_{\text{thr}}$ expressed in excitons per dot of four, as expected from their 8-fold degeneracy (Figure 2a). Upon doping, the $\langle N \rangle_{\text{thr}}$ reduces, and for the case of initial doping $\langle N \rangle_D$ of 5.4, the $\langle N \rangle_{\text{thr}}$ reaches a value of 0.9 excitons per dot (Figure 2b). This 4-fold reduction of the gain threshold upon doping outperforms the 2-fold reduction reported in CdSe based CQD systems. 14 By varying the initial doping of the CQD films, according to the size-doping dependence shown in Figure 1c, we have measured $\langle N \rangle_{\text{thr}}$ values of 4.3, 2.1, 1, and 0.9 excitons per dot for QD sizes with diameter (initial doping) of: 5.0 nm ($\langle N_D \rangle = 1.4$), 5.6 nm ($\langle N_D \rangle = 3.4$), 5.9 nm ($\langle N_D \rangle = 4.4$), and 6.2 nm ($\langle N_D \rangle = 5.4$) respectively (Figure 2c). This finding further corroborates our hypothesis of the effect of doping on the optical gain threshold. It is noteworthy that while the gain threshold measured in the undoped films is in good agreement with the value predicted by simple theoretical and Poisson-statistical models (see Supporting Information Section S8), the experimentally determined gain threshold for the doped QDs is distinctly lower that the one predicted by those models. However, it comes in good agreement when taking into account charged-exciton and exciton-exciton interactions, 15 which lower the gain threshold in heavily doped PbS CQDs (see Supporting Information Section S8).

The corresponding TA traces of both the undoped and doped films (Figure 2d,e) have been collected at the gain threshold. The calculated Auger rates have been calculated using the experimental biexciton Auger lifetime of 210 ps (measured for 5.5 nm PbS CQDs) at the exciton occupancy that reaches the gain threshold depending on the initial doping occupancy of the dots.
reproduce the Auger scaling dependence (Figure S9.1). We
note that in the gain regime, the TAS dynamics do not
follow the Auger scaling. It is noteworthy that in the
exciton recombination (see Supporting Information Section
S9). TAS analysis in the gain regime (Supporting Information
Section S9, Figure S9.2) discloses the presence of an
excitation-independent recombination pathway that does not
follow the Auger scaling, similar to the undoped films, and that
we assign as the gain lifetime of the system. To further
elucidate the dynamics of the system, we have both calculated
and extracted experimentally the Auger lifetime for a series of
doped CQDs at their corresponding gain threshold. Figure 2f
plots the experimentally measured excitation-independent
wavelength. We notice that already below the gain threshold,
the undoped sample shows multi-exponential dependence,
while above the gain, both doped and undoped samples are
characterized by a recombination channel that saturates with
−Δα/α_{PbS} and whose lifetime component is excitation
independent. In order to elucidate the multie exciton and charge
exciton dynamics, we used as a case study ~5.7 nm PbS QDs,
and we collected the TAS for the undoped and doped (⟨N⟩_PbS = 3.6) films probing the relaxation dynamics up to 1 ns (shown
in Supporting Information Section S9). In our model, we have
considered that the photogenerated exciton occupancy
distribution follows Poisson statistics. By fitting the lifetime
traces of the undoped samples at low exciton occupancy (⟨N⟩ < 1),
we extract the single exciton lifetime on the order of 1.5 ns as well as the Auger lifetime that follows statistical scaling
with a measured biexciton lifetime of ~200 ps. Therefore,
below the gain threshold, the relaxation mechanism is due to
Auger recombination as well as singe exciton relaxation.53
Above the gain threshold, there is a rise of an excitation
independent component with a lifetime of ~25–30 ps for large
CQDs (6.2 nm) and 15 ps for smaller sized dots (5.0 nm) that
saturates following the −Δt/t_{Φ_{PbS}} which we attribute as
the gain lifetime, as it does not follow the Auger scaling. It is
noteworthy that in the gain regime, the TAS dynamics do not
reproduce the Auger scaling dependence (Figure S9.1). We
instead measure a strong relaxation component of 25–35 ps
assigned as the gain lifetime) despite the fact that at those
conditions (excitation of 4 excitons per dot), Auger scaling,
taking into account Poisson statistics, predicts much faster
Auger lifetime components for the population of dots that are
excited with more than 4 excitons/dot (see table in Supporting
Information Section S9). This suggests that the stimulated
emission channel is favorable over non-radiative Auger
recombination for carriers when in the gain regime. The
mechanism behind this intriguing observation is still unclear
and merits further investigation.

In the doped CQD films, at fl uences below the gain regime,
Auger recombination takes place even at the single exciton
occupancy excitation as a result of the dopant electrons at the
CB. Our experimental results however point out the existence
of two recombination channels, one following the Auger scaling
and a longer one that we attribute to single charged-exciton
recombination (see Supporting Information Section S9). TAS analysis in the gain regime (Supporting Information
Section S9, Figure S9.2) discloses the presence of an
excitation-independent recombination pathway that does not
follow the Auger scaling, similar to the undoped films, and that
we assign as the gain lifetime of the system. To further
elucidate the dynamics of the system, we have both calculated
and extracted experimentally the Auger lifetime for a series of
doped CQDs at their corresponding gain threshold. Figure 2f
plots the experimentally measured excitation-independent
component assigned as the gain lifetime along with the experimentally extracted Auger lifetime at gain threshold. The experimentally extracted Auger lifetime matches well with the calculated Auger lifetime, when we consider only the nominal exciton occupancy of the dots at their corresponding thresholds (i.e., when ignoring the faster Auger components from the dots with larger exciton occupancy due to Poisson statistics). Interestingly, we note that doped QDs at threshold possess longer Auger lifetimes than gain lifetimes thanks to reaching the gain regime at lower exciton occupancies. The Auger lifetime on threshold is found to increase for a doping occupancy up to 3 electrons per dot. Beyond this point, Auger lifetime accelerates due to increasing number of free carriers in the dots. Throughout the experimental conditions of this work, the gain lifetimes remain in the range of 20–30 ps, thus faster compared to the experimentally determined Auger lifetimes, which supports that the stimulated emission is the main de-excitation channel.

Optical gain is a prerequisite for stimulated emission. Having achieved this, next we performed amplified spontaneous emission (ASE) measurements of our thin films. In line with the TA measurements, we observed stimulated emission from both doped and undoped samples (see Supporting Information Section S11). We calculated the stimulated emission occupancy threshold \( \langle N \rangle_{\text{th}} \) by fitting the integrated photoluminescence (PL) spectra (see Supporting Information Section S12) from the power-dependent \( S \) curves of all the samples. In Figure 3a,b we plot the integrated ASE peak as a function of exciton occupancy and the respective power dependence measurement of two representative PbS sizes of 5.4 nm (\( \langle N \rangle = 2.7 \)) and 5.8 nm (\( \langle N \rangle = 4.0 \)) (Figure 3c–f). All the undoped samples have an \( \langle N \rangle \) of 4, in agreement with the TAS measurements of Figure 2, and the ASE signal is saturated when 8 electrons have fully populated the CB. Power dependence measurement of those samples are shown in Figure 3c–f positioning the stimulating emission peak in wavelengths above 1500 nm. The sharp ASE peak has an fwhm of 14 meV, characteristic of a stimulated emission process and comparable with reported values in the visible from CdSe-based systems.10 To our knowledge, this is the first report of CQD ASE in the infrared characterized by ASE saturation and narrow line width, essential features of ASE that had remained elusive hitherto.34,35 The stimulated emission threshold occupancy is summarized in Figure 3g. The undoped samples preserve a constant value of \( \langle N \rangle_{\text{th}} \) of 4 independent of their size, whereas in the case of doped CQDs increasing their size, and thereby the initial doping occupancy, the stimulated emission threshold decreases to a minimum value of 1.3 excitons, in good agreement with the TA measurements.

A figure of merit of paramount importance for applications in optical amplification and lasing is the net modal gain of the material. We have experimentally measured the net modal gain \( g_{\text{modal}} \) using the variable stripe length (VSL) technique (see Supporting Information Section S13). We report an average \( g_{\text{modal}} \) of 30 cm\(^{-1}\) nearly constant for all the undoped samples (Figure 3h). Upon doping, \( g_{\text{modal}} \) increases up to a value of 114 cm\(^{-1}\) for an \( \langle N \rangle \) of 4, when the CB is half-filled. This modal gain value outperforms Er-doped fiber systems with values in the range of 0.01–0.1 cm\(^{-1}\) and compares favorably to costly epitaxial III–V multiquantum well and quantum dot systems.36 We demonstrate tunable coverage across the optical fiber communication spectrum (Figure 3i) from a solution processed material, extending beyond the spectral coverage of Er-doped fiber systems. The tunable spectral coverage across the infrared taken together with the high gain values and the low threshold represent a significant advance toward the development of infrared CQD solution processed lasers. Further improvements in doping efficacy may lead in the future toward threshold-less lasing in such doped CQD films. The latter taken together with future developments of engineered lead chalcogenide CQDs with suppressed Auger16 and the appropriate selection of thermal dissipation schemes9 can pave the way toward CW lasing (predictive calculations on this are provided in Supporting Information Section S14).

- **ASSOCIATED CONTENT**

- **Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.nanolett.0c01859.

Experimental methods, characterization methods, density of states calculations, control experiments, additional absorption spectra, XPS and UPS spectra and analysis, FET experiments and analysis, doping stability, ASE threshold calculations, exciton dynamics models, Auger analysis, additional TA traces, additional power dependence spectra and analysis, VSL measurements, CW lasing calculations, and exciton occupancy calculations (PDF)

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