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Charge Carrier Cooling Bottleneck Opens Up Nonexcitonic Gain Mechanisms in Colloidal CdSe Quantum Wells

Renu Tomar,†‡ Aditya Kulkarni,§ Kai Chen,∥§ Shalini Singh,†‡ Dries van Thourhout,†‡∥ Laurens D.A. Siebbeles,§ Zeger Hens,†‡⊥ and Pieter Geiregat†‡¶

†Physics and Chemistry of Nanostructures and §Photonics Research Group, Department of Information Technology, Ghent University, Gent 9000, Belgium
‡Center for Nano and Biophotonics, Gent 9000, Belgium
§Optoelectronic Materials Section, Department of Chemical Engineering, Delft University of Technology, Van der Maasweg 9, Delft 2629HZ, The Netherlands
∥The MacDiarmid Institute for Advanced Materials and Nanotechnology, Wellington 6012, New Zealand
⊥School of Chemical and Physical Sciences, Victoria University of Wellington, Wellington 6012, New Zealand
¶The Dodd-Walls Centre for Photonic and Quantum Technologies, Wellington 6012, New Zealand

ABSTRACT: Ultrathin two-dimensional (2D) materials have received much attention in the past years for a wide variety of photonic applications because of their pronounced room-temperature excitonic features, leading to unique properties in terms of light–matter interaction. However, only a few studies focus on light amplification and the complex photophysics at high excitation density. The beneficial nature of strong excitonic effects on optical gain remain hence unquantified, and despite the increased binding energies of the excitonic species, it remains unclear what the involvement of 2D excitons would be in optical gain. Here, we use colloidal CdSe nanoplatelets as a model system for colloidal 2D materials and show, using a quantitative and combinatory approach to ultrafast spectroscopy, that several excitation density-dependent optical gain regimes exist. At low density, optical gain originates from excitonic molecules delivering large material gains up to 20 000 cm⁻¹ with an Auger limited lifetime of a few hundred picoseconds. At increasing pair density, we observe a persistence of this excitonic gain regime and the unexpected coexistence of blue-shifted and significantly enhanced optical gain up to 10⁵ cm⁻¹. We show that this peculiar situation originates from a carrier cooling bottleneck at high density that limits further exciton formation from unbound charge carriers. The insulating (multi-)exciton gas is found to coexist with the conductive phase, indicating the absence of a full Mott transition. Our results shed a new light on the photophysics of excitons in strongly excited 2D materials.

1. INTRODUCTION

Two-dimensional (2D) materials are considered as prime candidates for realizing low-cost and tunable optoelectronic devices.¹,² Applications in fields as diverse as light emission, photodetection, and quantum optics can make use of the distinct and high-oscillator strength excitonic transitions and/or the tunable, narrow photoluminescence (PL) found in such materials at room temperature.³⁻⁵ Recent years saw an increase in the number of available inorganic 2D materials such as transition-metal dichalcogenide monolayers (MLs),¹ layered perovskites,⁶ and II–VI nanoplatelets.⁷ The strong Coulomb correlations leading to stable and strong excitonic features might be very beneficial for increasing the light–matter interactions required for large and/or low-threshold optical gain. Several reports indicated, for example, low-threshold lasing using 2D materials.⁸⁻¹⁰ However, because net modal gain develops only at very high carrier densities, the possible role of those exciton(s) (complexes) in the process remains unclear.¹⁰⁻¹ⁱ Indeed, within a mass-action model, excitons—and higher complexes such as excitonic molecules—form only when the associated binding energy and pair density is sufficiently large. On the other hand, it is well known that 2D excitons dissociate more readily to form unbound carriers at high excitation density. This effect is due to the density-induced screening of the attractive Coulomb interactions which reduces the exciton binding energy, leading to so-called...
Optical gain in epitaxially grown quantum wells, for example, is mostly mediated by an electron–hole plasma, even at cryogenic temperatures. Given the exceptionally large binding energy of excitons in colloidal 2D materials, in large due to reduced screening, it is not clear to what extent these concepts still hold. As one of the few quantitative reports on the optical gain mechanism in 2D materials, Chernikov et al. observed a strong band gap renormalization and transition to an unbound electron–hole plasma in WS₂ at high carrier density associated with the development of gain. This observation is remarkable, given the large binding energies (>300 meV) for the 2D excitons.

It is clear that quantitative studies on optical gain are therefore much needed to assess the relation between stimulated emission (SE) and the complex mixture of carrier types that exist in strongly excitonic 2D materials at high density, such as 2D excitons, excitonic molecules, and free charges.

CdSe 2D nanoplatelets, also called colloidal quantum wells, provide an excellent test-bed for this purpose, as they can be obtained as scatter-free colloidal dispersions with high luminescence quantum yield and excellent colloidal stability. Several reports have demonstrated excellent light amplification properties using these colloidal quantum wells, such as large modal gains up to 500 cm⁻¹ and low lasing thresholds down to tens of μJ/cm², pointing toward a seemingly huge potential of these materials. Apart from core only CdSe platelets, also more complex architectures have been studied such as core/shell and core/crown systems. The development of optical gain in CdSe-based platelets was however studied mainly through thin-film measurements, where either amplified spontaneous emission (ASE) or lasing was employed. Unfortunately, such approaches fail to elucidate the fundamentals of the gain mechanism, that is, which type of interactions and excitations (free charge carriers, excitons, ...) are required to achieve net optical gain. Nor do they provide any prospect on the full potential of the material as the measured properties are inherently linked to the development of net SE dictated by the film and/or cavity losses. A case in point here is the alleged threshold for stimulated emission (SE), where energy densities (typically in μJ/cm²) are reported instead of generated excitation densities. Although energy densities have a high practical value, the density of carriers, that is, the number of absorbed photons, is the most relevant parameter for a proper mechanistic understanding of optical gain in this unique class of materials. Nanoplatelets can be deceptive in this respect, as they offer larger intrinsic absorption coefficients at short wavelengths because of a reduced in-plane screening.

Herein, we report on an extensive quantitative study of optical gain development and the associated high excitation density photophysics of colloidal 2D CdSe quantum wells. By combining a comprehensive set of femtosecond transient absorption (TA), photoluminescence (PL), and terahertz spectroscopy, we were able to identify different optical gain regimes and a remarkable absence of a clear exciton Mott transition. At low electron–hole pair densities, a gain band appears at the low-energy side of the heavy-hole (HH) exciton absorption that was assigned to SE by biexcitonic molecules. Here, we show that this exciton-mediated gain regime persist at pair densities well above the expected Mott density, where a transition of the exciton–molecule gas into an electron–hole plasma is expected. At ~20,000 cm⁻¹, the saturated material gain in this regime is significantly larger than for comparable zero-dimensional materials, such as state-of-the-art CdSe/CdS or perovskite nanocrystals, and explains the observations of large modal gains in the literature. Apart from the exciton-mediated regime, a charge carrier cooling bottleneck manifests itself, which results in the peculiar coexistence of neutral excitonic species with unbound, hot electrons and holes. This combination leads to large, up to 10⁵ cm⁻¹, and broadband material gain and luminescence, a second regime. At the same high excitation densities, we inferred a remarkable preservation of the exciton polarizability, as evidenced by optical pump–terahertz probe experiments, and HH and light-hole (LH) exciton oscillator strength, both of which indicates that a pure Mott-type transition does not occur. Our results show that even though strong excitonic features are not beneficial for the development of broadband gain, they do significantly increase the achievable material gain magnitude. Moreover, the extreme stability of the exciton gas in colloidal 2D materials will be beneficial for applications relying on the scattering of excitons at high density, such as exciton-polariton lasers.

2. EXPERIMENTAL PROCEDURES

2.1. Synthesis of 4.5 ML Platelets. The synthesis of 4.5 ML CdSe nanoplatelets was carried out by degassing cadmium myristate (0.34 g), Se (24 mg) and 1-octadecene (ODE) (25 mL) in a three-necked flask in vacuum and backfilling with N₂. To this solution, 0.08 g of cadmium acetate was added at 205 °C when the solution turned yellow. Next, the solution was heated at 240 °C for 10 min, and 1.6 g of cadmium oleate in ODE was added at end. Samples were washed with a hexane/ethanol mixture. For spectroscopy, the CdSe platelets were dispersed in a transparent solvent (hexane) to achieve optical densities of 0.1 at the first HH exciton transition (510 nm). The PL quantum yield was determined using an integrating sphere after CW photoexcitation at 400 nm.

2.2. Quantification of Absorbed Photons per Platelet. The average number of absorbed photons (or photogenerated electron–hole pairs) at time zero, here labeled \(\langle N \rangle\), generated by a pump pulse at wavelength \(\lambda\) was calculated as

\[
\langle N \rangle = J_{ph} \times \sigma_s
\]

Here, \(J_{ph}\) is the photon fluence in photons/cm², \(\sigma_s\) is the absorption cross section of the platelets, and \(J_s\) is the sample absorbance at the pump wavelength. The beam size used for calculating \(J_{ph}\) is measured using a Thorlabs CCD Camera Beam profiler. The additional factor corrects for variation of the pump fluence along the pump beam path length. The cross section is determined starting from the intrinsic absorption coefficient (see Supporting Information S2).

2.3. Ultrafast TA Spectroscopy. Samples were pumped using 110 fs pulses (1 kHz) at varying wavelengths, created from the 800 nm fundamental (Spitfire Ace, Spectra-Physics) through nonlinear frequency mixing in an optical parametric amplifier (OPA, TOPAS, Light Conversion). Broadband probe pulses were generated in a CaF₂ crystal using the 800 nm fundamental. The probe pulses were delayed using a delay stage with maximum delay of 3.3 ns (Newport TAS). Noise levels of 0.1 mOD (rms) are achieved by averaging out over 5000 shots. The probe spectrum covers the VIS-NIR window.
from 350 up to 700 nm. Using 2 mm path length cuvettes, the samples were stirred during pump-probe measurements to avoid effects of photocharging and sample degradation. Only at extreme densities ( ⟨N⟩ > 150) was sample degradation observed. No air-free sample handling was required as CdSe platelets are, under our conditions, insensitive to oxidation as is evidenced by a lack of PL peak shift or change in PL quantum yield when exposing samples to ambient conditions.

2.4. Ultrafast PL Spectroscopy. For (ultrafast) PL spectroscopy, samples were measured using a cuvette with a 1 mm optical path length and an optical density of 0.1 at the HH absorption peak to avoid strong re-absorption. The quantum yield was determined using an integrating sphere with excitation at 400 nm. For the ultrafast experiments, samples were translated along 1 axis to avoid photocharging. The detection of the broadband PL on femtosecond timescales was made possible by using a newly developed transient grating technique by Chen et al.37 A Ti:sapphire amplifier system (Spectra-Physics Spitfire Ace) operating at 1 kHz generating 100 fs pulses was split into two parts. One part was frequency doubled using a BBO crystal and focused to a 70 µm spot on the sample. The PL is collimated using an off-axis parabolic mirror and refocused on a polished slice of fused silica. The second part of the 800 nm output was split using a 50:50 beam splitter creating two gate beams that are focused on the sample. The PL is collimated using an off-axis parabolic mirror and refocused on a polished slice of fused silica. The instantaneous grating generated by the interfering gate beams creates an instantaneous gate which is used to temporally resolve the decay over a broad wavelength range. The scatter of the pump beam was suppressed using a 430 nm long-pass filter, and the pump was set a magic angle relative to the PL collection. Data are averaged over 15 000 shots for every time delay.

2.5. Terahertz Spectroscopy. Samples were photoexcited by optical pump pulses of wavelength 480 nm and the photoconductivity due to mobile charge carriers, and the polarizability of excitons was probed by single cycle THz pulses. The real part of the THz conductivity is due to in-phase charge velocity in the probing THz field and leads to a reduction of the amplitude of the THz field.38,39 The imaginary THz conductivity is due to the out-of-phase velocity of charges and the polarizability of excitons, leading to a phase retardation of the THz field.39–45 Single cycle terahertz pulses were generated in a ZnTe nonlinear crystal by optical rectification of 800 nm optical pulses of duration 45 fs, delivered from a chirped pulse amplified femtosecond laser system (Mira, Coherent), with a frequency of 1.4 kHz. The generated THz pulses were detected in a ZnTe crystal by the electro-optic effect with a chirped optical detection pulse, which encodes and decodes the entire THz pulse in a single laser shot. Optical pulses of wavelength 480 nm were generated in an OPA (TOPAS, Coherent). The THz conductivity due to charge carriers and excitons at time t after the laser pump pulse, that generates charges and excitons, was obtained from the differential THz field given by

\[
\Delta E(t_p,t) = E_{\text{excited}}(t_p,t) - E_{\text{unexcited}}(t_p),
\]

where \(E_{\text{excited}}(t_p,t)\) and \(E_{\text{unexcited}}(t_p)\) are the transmitted THz field in the presence and absence of optical pumping, respectively. \(t_p\) is the delay between the THz generation and detection pulse. The real THz conductivity was obtained from the ratio \(\Delta E(t_p,t)/E_0(t_p)\), where \(E_0(t_p)\) is the time at which the amplitude of the THz field is maximum, according to44–46

\[
\Delta \sigma(t) = \frac{2n_0c\epsilon_0}{L} \frac{\Delta E(t_p, t)}{E_0(t_p)}
\]

with \(n_0 = 1.42\), the refractive index of n-hexane,47 \(\epsilon\) as the speed of light in vacuum, \(\epsilon\) as the permittivity of free space, and \(L\) as the sample thickness. The imaginary THz conductivity was obtained from the differential THz field with \(t^\text{zero}\) the time at which the THz field is zero, according to48,42,45

\[
\Delta \sigma(t) = \frac{2n_0c\epsilon_0}{L} \frac{\Delta E(t_p, t)}{E_0(t_p)}
\]

2.6. Material Overview. CdSe platelets with a thickness of 1.37 nm, that is, 4.5 MLs, were synthesized according to Ithurria et al.7,20 and their lateral dimensions were determined using brightfield transmission electron microscopy (TEM). While the measurements were repeated on multiple samples, all the data shown in the manuscript were recorded on a sample measuring 34 × 10 nm², see Figure 1a and Supporting Information S1. These dimensions are well above the electron Bohr diameter of the exciton, see further, leading to overall weak in-plane 2D confinement. Figure 1a shows the intrinsic absorption and spontaneous emission spectrum of a dispersion of CdSe platelets in n-hexane. As is common for direct gap II–VI semiconductor quantum wells, two excitonic features are observed, attributed to the HH (510 nm) and LH (480 nm) exciton transitions.7,49 The spontaneous PL has a high
Figure 2. Optical gain at high carrier density after excitation with 400 nm. (a) 2D time—wavelength map of the intrinsic absorption coefficient $\mu_i$ (cm$^{-1}$) for $\langle N \rangle = 84$ where we limit the color scale to negative values showing net gain: $g_i = -\mu_i > 0$. The vertical line indicates the position of the HH exciton resonance. (b) 2D time—wavelength map of the PL taken at the same excitation density as (a). Black trace is the spontaneous emission for $\langle N \rangle \ll 1$. (c) Intrinsic absorption coefficients taken 2.5 ps after photoexcitation with a 400 nm pump pulse for densities ranging from 4.6 to 112 excitations per platelet. (d) Material gain spectra $g_i$ for the same carrier densities as (c). (e) Required $\langle N \rangle$ to achieve transparency, $\mu_i (2.5 \text{ ps}) = 0$, as a function of wavelength. Optical gain is achieved at 41 and 35 in the red- and blue-shifted gain regimes respectively. (f) PL at 2.5 ps for increasing pair density from 4.7 to 78. The inset shows the fwhm wavelengths of the PL spectra.

quantum yield of >50% (see Experimental Procedures) and shows no Stokes shift, indicating the absence of strong phonon-coupling\(^{25}\) fine-structure relaxation,\(^{51}\) or spurious effects such as charging, exciton localization, or trap-emission.\(^{49}\) We represent the absorbance spectrum as an intrinsic absorption coefficient $\mu_{i,0}$ (cm$^{-1}$), a quantity that is independent of the concentration of platelets in the dispersion analyzed. $\mu_{i,0}$ can be calculated through theory as $\mu_{i,0} = 5.9 \times 10^5$ cm$^{-1}$, see Supporting Information S2, as validated experimentally by Achstein et al.\(^{50}\) Besides providing a means of normalizing linear and transient spectra, $\mu_{i,0}$ enables us to calculate the absorption cross section $\sigma$ of a single platelet from the relation $\sigma(\lambda) = \mu_{i,0}(\lambda) \times V$, with $V$ as the platelet volume.\(^{48}\) This brings us to a cross section of 8 $\times$ 10$^{-14}$ cm$^2$ at 400 nm for the platelets used here. Following the well-established methodology of epitaxially grown quantum wells, Figure 1b shows a decomposition of the absorbance spectrum into (C) step-like features representing 2D-continuum transitions and (X) narrow resonances representing the excitonic transitions.\(^{22,49}\) This decomposition shows that the sharp exciton features are actually superimposed on a broadened step-profile, typical for the 2D density of states, which is due to free carrier transitions between the first of the 2D HH states to the first 2D conduction-band states $C_{\text{HH}}$, see also the inset of Figure 2b. At higher energy, a similar step associated with the LH valence band is observed, denoted as $C_{\text{LH}}$. The fit also enables us to estimate the exciton binding energies $E_{b,XX}$ as 193 ± 5 meV (HH exciton) and 277 ± 13 meV (LH exciton). Such figures are in excellent agreement with the literature,\(^{22,53}\) where the large values for the exciton binding energies are confidently ascribed to the reduced screening of the Coulomb interaction compared with bulk, given the low permittivity environment consisting of ligands and solvent.\(^{15,50}\) Using the exciton binding energy, we can calculate the Bohr radius of the 2D HH exciton using $m_i = 0.085$ a$_{\text{HH}} = \frac{h}{\sqrt{2m_{i}E_{b,\text{HH}}}} = 1.52$ nm.\(^{54}\)

Given the area $S$ of the nanoplatelets, we can define $\langle N \rangle = \langle n a_{\text{HH}}^2 \rangle^{-1} \times S$ as the number of excitons required to physically occupy all of the nanoplatelets surface area, that is, the average interexciton distance is the Bohr diameter. For our sample, this amounts to 45 excitations per platelet, a number that sets a scale for $\langle N \rangle$ in this particular system.

3. RESULTS

We analyzed the development of optical gain in dispersed CdSe platelets by a combination of femtosecond pump-probe TA, ultrafast PL spectroscopy, and optical pump—terahertz probe spectroscopy; see Experimental Procedures section for technical details. All techniques involve the excitation of an ensemble of dispersed platelets using a 110 fs laser pulse at 400 nm, unless mentioned otherwise. In TA, photoexcitation is followed by probing the change in absorbance $\Delta A(\lambda, t) = A(\lambda, t) - A_0(\lambda)$, where $A$ is the absorbance after photo-excitation and $A_0$ the absorbance in absence of pump pulse—as a function of wavelength and pump-probe time delay. Note that the occurrence of optical gain, or net SE, at a given wavelength is most clearly evidenced by a negative nonlinear absorbance $A = \Delta A + A_0 < 0$.

Starting from this nonlinear absorbance $A(\lambda, t)$, the time- and wavelength-dependent intrinsic absorption coefficient of the nanocrystal (NC) under study can be calculated as

$$\mu_i(\lambda, t) = A(\lambda, t)/A_0(300)h_i(300)$$

(4)

see also Supporting Information S2.\(^{56}\) When $\mu_i$ turns negative, we have net SE for which $-\mu_i = g_i$ can be identified as the material gain. This quantity is independent of the measurement conditions and can be translated to any device or thin-film context through a rescaling with the appropriate modal confinement and volume fraction.\(^{13}\)

Figure 1c shows a 2D time—wavelength map of $\Delta \mu_i$ recorded on a dispersion of nanoplatelets using excitation at
400 nm. As outlined in the Experimental Procedures section and Supporting Information S2, the average excitation number \( \langle N \rangle \) such a fluence brings about per platelet can be calculated from the product \( f_{\text{exc}} \times \sigma \) of the photon flux and the absorption cross section, an exercise yielding \( \langle N \rangle = 84 \) for the data represented in Figure 1c. At these pumping levels, pronounced, long-lived bleach features arise at the HH and LH exciton positions because of state-filling. Importantly, the absorbance at the pump wavelength remains nearly unaffected, implying that no corrections are needed for the absorbed number of photons. At early times, a short-lived photoinduced feature occurs near 530 nm. This photoinduced absorption gets overtaken by a bleach signal at later times and was assigned to exciton–molecule transitions, opposed to the common notion of spectral shifts.\(^{31}\) Comparing the \( \Delta \mu \) signal to \( \mu_{\text{no}} \), we can readily identify regions of the 2D map where net optical gain occurs. These are indicated by the white dashed contour line in Figure 1c and remarkably show up both at the low- and high-energy side of the HH resonance.

To evaluate optical gain more precisely, we map out the regions of the spectrum where the material gain \( g_i = -\mu_i \) turns positive. Figure 2a shows such a false color map for a higher excitation density corresponding again to an average photoexcitation density of \( \langle N \rangle = 84 \). We observe two distinct regimes of optical gain, separated from each other by the persisting HH absorption at 510 nm. First, a \( \approx 60 \) nm (250 meV) wide gain band red-shifted from the HH exciton peak is observed which persists for ca. 100 ps. This gain window coincides perfectly with reports on ASE and lasing experiments on 4.5 ML CdSe nanoplatelets and recent theoretical work, see also Supporting Information S3.\(^{22,31}\) At these high pump powers, an unexpected second window of optical gain develops at the high-energy side of the HH resonance. It offers significantly larger gain than the low-energy band, builds up instantaneously but is sustained only until \( \approx 10 \) ps after photoexcitation. Figure 2b shows the map of the PL under similar conditions of optical pumping as in Figure 2a. The emission spectrum is clearly broadened to lower and higher photon energy compared with the spontaneous emission of a single exciton. Note that the spectrum decays asymmetrically, showing a shorter \( \approx 10 \) ps lifetime for the high-energy luminescence and a longer \( \approx 100 \) ps lifetime for the red-shifted luminescence, overall very comparable to the material gain dynamics shown in Figure 2a.

### 3.1. Gain Bands

To deepen our understanding of both gain regimes, we sweep the photogenerated density—or average number of excitations per platelet \( \langle N \rangle \)—over 2 orders of magnitude, from nearly single excitations to close to 112 charge pairs per platelet. Note that the highest flux corresponds to a 2D pair density of \( 3.4 \times 10^{13} \) cm\(^{-2} \). Figure 2c shows the intrinsic absorbance \( \mu_i \) at 2.5 ps for these increasing densities. We observe that gain initially builds up on the low-energy (long wavelength) side of the HH resonance. This first gain window is sustained throughout the entire density range. The gain in the blue-shifted regime is observed only when the excitation exceeds ca. 40. A number of intriguing observations can already be made at this point: the gain at the high-energy side is disruptively larger than the low-energy gain; both the HH and LH exciton absorption persist, showing only slight saturation and effectively counteract broadband gain development; and at high density, a reduction of the absorbance plateau at shorter wavelengths (\(<480 \) nm) is observed.

Zooming in on the net material gain \( g_i = -\mu_i \), Figure 2d shows that for the low-energy gain band, the intrinsic gain coefficient in cm\(^{-1} \) reaches up to 18 000 cm\(^{-1} \) and eventually saturates close to 84 excitations per platelet. Quantitative variable stripe length experiments verify that such large gain coefficients are indeed achievable, see Supporting Information S3, and line up with the report of Guzel turk et al.\(^{25}\)

The gain at the blue side shows no signs of saturation and is also disruptively larger with values close to \( 10^3 \) cm\(^{-1} \). A more detailed analysis of the gain threshold is shown in Figure 2e, see also Supporting Information S4, where we observe that the low-energy gain band has a threshold of 4.1 \( \pm \) 0.5 and the high-energy gain band of 35 \( \pm \) 2, see also Supporting Information S4.

Finally, Figure 2f shows a similar fluence dependence as Figure 2c,d for the PL after 2.5 ps. Also here, a transition occurs at ca. \( \langle N \rangle = 35 \) which is perhaps more clearly observed looking at the wavelengths (energies) for which the luminescence has its half-maximum value, as is shown in the inset of Figure 2f. A gradual increase at the low-energy side clearly contrasts the disruptive jump on the high-energy side near \( \langle N \rangle = 35 \). These results show that not only are nanoplatelets able to amplify incoming probe light in the high-energy gain window, they also exhibit spontaneous emission in the same spectral region, with a similar density-dependent onset. This is crucial for their use in lasers because the material’s ability to amplify its own spontaneous emission is what sets it apart from optical amplifiers.

As shown elsewhere, we assign the low-energy gain band to SE from excitonic molecules.\(^{31}\) Such a biexcitonic molecule is a bound two-exciton state with a binding energy \( E_{\text{BM}} \approx 45 \) meV for 4.5 ML CdSe platelets, outlasting thermal energy at room temperature and typical optical phonon energies of CdSe.\(^{22,31,49}\) This complex can recombine radiatively to a single exciton state, giving rise to SE and net amplification in a narrow band on the low-energy side of the HH exciton absorption. The quantified material gain coefficients of the excitonic molecule gain mechanism shown here are close to 1 order of magnitude larger than observed for state-of-the-art CdSe/CdS or perovskite quantum dots (QDs).\(^{33–35}\) which rationalizes recent reports on extremely small footprint lasers operating under low excitation power.\(^{22–24}\) Supporting Information S5 shows that across the entire density range, exciton–exciton annihilation is the main culprit for the vanishing carrier density and hence the overall optical gain lifetime of ca. 100 ps, as confirmed by other reports.\(^{33–37}\)

The exciton-mediated red-shifted gain window observed at low density persists through the entire density range and shows a saturation close to 100 excitations, see Figure 2d. The latter already indicates that excitonic species can indeed persist up to such high carrier densities. In particular for the biexcitonic molecules, which show a much smaller stabilization energy compared with excitons, this result is remarkable. An initial confirmation is found in the absence of a full collapse of the exciton oscillator strength as is observed via the preservation of the exciton luminescence, gain, and absorption features, see Figure 2. However, before we analyze the limited exciton saturation in more detail, we want to unambiguously confirm the existence of excitons at high density.\(^{38}\)

### 3.2. Excitonic Stability at High Density

To this end, we employ optical pump–THz probe spectroscopy as similar to previous studies on 2D InSe nanosheets,\(^{39}\) one can use these measurements to characterize the nature of photoexcited
Figure 3. THz spectroscopy (a) real (positive traces) and imaginary (negative traces) components of the THz conductivity as a function of time after photoexcitation with $\langle N \rangle$ equal to 6 and 53. The optical bleach at the HH exciton transition at comparable densities is shown for comparison with the imaginary THz conductivity. (b) Measured real (red circles) and imaginary (blue squares) THz conductivity averaged between 6 and 10 ps after photoexcitation as a function of $\langle N \rangle$, together with fitted results from the Saha model (solid lines). Inset below shows a schematic depiction of the interexciton distance for this density range. (c) Reduction of normalized HH oscillator strength (red circles) and HH spectral shift $\Delta E$ (blue triangles), normalized to the HH binding energy, for increasing carrier density.

species. The real component of the THz conductivity is due to free mobile charges moving with velocity in-phase with the THz field, while the imaginary conductivity is due to the out-of-phase velocity of charges (resulting from scattering) and the polarizability of excitonic species (see Experimental Procedures).59,60

Figure 3a shows the THz conductivity as a function of time after photoexcitation for $\langle N \rangle = 6$ and 53. The real conductivity is much smaller than the imaginary component for both densities, which is a strong qualitative indication for a significant contribution from excitons. To get more insights into the contribution of excitons, we plot the kinetics of the optical bleach at the HH transition due to excitons on top of the imaginary THz conductivity for comparable excitation densities. The decay kinetics of the bleach and imaginary THz conductivity are similar at lower and higher densities, indicating that the imaginary THz conductivity is mainly due to excitons.

The initial real THz conductivity $\Delta \sigma_R$ is given by

$$\Delta \sigma_R = \frac{eN_0 \phi(n) \sum_{\mu R} \mu R}{L} \tag{5}$$

where $\sum_{\mu R}$ is the sum of the real components of the electron and hole mobility, $\phi$ is the quantum yield of charges, $e$ is the elementary charge, and $N_0$ is the number of absorbed photons per unit area in the sample solution with cuvette length $L$. As denoted, $\phi$ depends on the average pair density $n = \langle N \rangle / S$ where $S$ is the average platelet area. The imaginary THz conductivity $\Delta \sigma_I$ is given by the sum of the contribution of (localized) charges and excitons, according to

$$\Delta \sigma_I = \frac{eN_0}{L} \times \left[ \sum_{\mu I} \mu_I \phi(n) + \frac{a_0 \omega}{e} (1 - \phi(n)) \right] \tag{6}$$

where $\sum_{\mu I}$ is the sum of the imaginary electron and hole mobility, $a_0$ is the polarization of excitons, and $\omega$ is the radian frequency of the THz electric field. The imaginary THz conductivity in Figure 3b increases close to linearly with $\langle N \rangle$ (or $n$). The sublinear increase of the contribution of free charges observed in the real conductivity indicates $\phi(n)$ increases strongly sublinearly. Given eq 2 and the observed linear increase with $n$, this implies a significant contribution of the second term in eq 2, that is of excitons.

The fraction $\phi$ of free charges is dictated by the Saha equilibrium $e + h \rightleftharpoons X$, which assumes a thermal equilibrium between excitons and free charge carriers. In Supporting Information S6, this equilibrium is worked out formally. Using the Saha model at a temperature of 293 K, and fitting that to the experimental data with $\sum_{\mu R}$ and the exciton polarizability $a$ as adjustable parameters, yields the drawn curves in Figure 3b. The real conductivity is reproduced with $\sum_{\mu R} = 54 \pm 12$ cm$^2$/V s. For the imaginary conductivity, we found $\sum_{\mu I} = 7 \pm 5$ cm$^2$/V s and $a = 3.1 \pm 0.2 \times 10^{-16}$ cm$^3$/V. The imaginary charge mobility and exciton polarizability for the 4.5 ML CdSe platelets studied in the current work are close to those reported earlier for similar CdSe platelets.53

We remark that the Saha equation can fit the data reasonably well beyond the broadband gain onset of $\langle N \rangle = 35$, see Figure 3b. We should however proceed with caution in this regime for two reasons. The temperature of the system is not well defined beyond this density and most likely above room temperature, as is evidenced by the detailed analysis of the PL (see further), and (b) full occupation of the platelet occurs around the same number, $\langle N \rangle_L = 45$, as derived earlier. The first observation complicates the validity of the Saha fitting procedure as thermal equilibrium is not guaranteed, whereas the second observation indicates possibly stronger carrier scattering limiting the mobility and hence possibly saturating the transient real conductivity.

In any case, the THz results unambiguously show that across the entire density range probed in this work, excitons are present with no significant change in the polarizability compared with the isolated, low density regime. This agrees with the observation that the red-shifted gain window, assigned to excitonic species, can persist up to the highest densities used here. For the highest carrier density employed in this work (112), the average interexciton distance is only $d_i = 2 \sqrt{\frac{3}{\pi \times \langle N \rangle}} = 1.9$ nm, fairly close to the Bohr radius estimated earlier. Though exciton-carrier phase diagrams can be very complex, a general understanding is that a so-called Mott-type transition should occur at these separations. Closely tied to the transition of an exciton gas into a conductive phase is the saturation and spectral shifting of the exciton absorption line itself. It is well known that the oscillator strength of the exciton transitions can diminish and vanish because of state-filling and screening-induced renormalization effects.61 In concert with this, the exciton line will also undergo spectral shifts for increasing density because of a balance between renormalization of the band gap (red shift) and reduction in exciton binding energy (blue shift).60–62 Figure 3c shows the evolution of the normalized oscillator strength of the HH-exciton line, extracted from the integrated HH-peak area (see.
Supporting Information S2), and the HH spectral shift $\Delta E_{HH}$ for increasing excitation density. A qualitative fit, $f = \frac{1}{1 + \langle N \rangle / \langle N \rangle_s}$, to the oscillator strength indicates a characteristic saturation density $\langle N \rangle_s = 12.7$. A full Mott transition is however not observed as the oscillator strength persists up to $\langle N \rangle = 112 \approx 10\langle N \rangle_s$, with a lingering strength of ca. 0.25$f_0$. This observation indicates that new types of carriers, or interactions between them, manifest themselves at higher density, a point we will elaborate further in the following section.

Finally, the exciton line shows a clear red shift, indicating that band gap renormalization (BGR) overtakes the blue shift that would originate from a binding energy reduction through increased screening of the exciton. Our quantitative observations on CdSe are in-line with recent observations on other high binding energy 2D materials such as WS$_2$, where similarly a red shift and no full saturation of the exciton line was observed until densities of $5 \times 10^{13}$ cm$^{-3}$, equivalent to $\langle N \rangle = 16$ for our experiments, yet at slightly higher densities, Chernikov et al. showed that the exciton line does get fully saturated and a transition to free carrier-mediated gain takes over.

3.3. Gain via Unbound Charge Carriers. Having established the excitonic nature of the universal gain regime at low photon energy, it remains an open question what causes the short-lived broadband gain and luminescence at high photon energy just after strong photoexcitation. To deepen our understanding, we first conceptualize how charge carrier thermalize and excitons are formed in direct gap polar semiconductors. Upon photoexcitation with 400 nm (3.1 eV), free electrons and holes with significant excess energy are created. These “hot” carriers have to cool down by optical phonon emission and finally condense into a HH exciton, see Figure 4a. This is quite a remarkable process as the stabilization of the exciton alone, that is, its binding energy, requires the dissipation of close to 200 meV. Given typical phonon energies of $E_{LO,CdSe} \approx 26$ meV, this implies that multiple optical phonons need to be emitted. Several reports in the literature indicated that this coupling to phonons is relatively weak for single excitons and gets even more slowed down with increasing excitation density, hence slowing down exciton formation after nonresonant excitation. Nonetheless, we observe a sub-picosecond, near resolution-limited, buildup of the band-edge exciton population and the associated gain at low exciton densities.

To understand how a cooling bottleneck can still manifest with such fast single exciton formation, we should consider the accumulative nature of the phonon emission process, see also Figure 4a. At early times, carriers cool down extremely fast to the band edge by rapid phonon emission, denoted as stage (1) in Figure 4a, giving rise to fast buildup of exciton PL and band-edge bleach, stage (2). However, as more carriers cool down, the phonon population increases up to a point where the temperature (occupation) of (a subset of) the phonon modes, generalized as $T_q$, exceeds that of the electronic system, generalized as $T_e$. At this point, net energy transfer to the phonon bath will substantially reduce, giving rise to a cooling bottleneck for the remaining carriers. Overall, this would result in the formation of a hot electron–hole gas which cannot cool down effectively to form excitons. This will eventually lead to spontaneous and also SE from unbound charge carriers from higher energy free carrier states.
Key elements to validate the proposed model are (a) the (prolonged) occupation of high-energy levels corresponding to unbound charges and (b) an increase of the overall carrier temperature, that is, a modified distribution of charge carriers over those high-energy states. The TA and luminescence data in Figure 2c,f already show that we can indeed attain a significant amount of carriers occupying energy levels blue-shifted from the HH line because we achieve spontaneous emission and net amplification of incoming probe light in this spectral window. As the TA data are congested because of the overlap of several spectral features (bleach, induced absorption, shifts, ...), we focus here on the PL data to extract additional information on the unbound charge carrier distribution, that is, their temperature and absolute number density.

Figure 4b shows the high-energy tail of the normalized PL spectrum 2.5 ps after photoexcitation with 400 nm, for increasing excitation density from $\langle N \rangle = 1.9 - 78$. We clearly observe both an increased emission at higher energy and a change in the slope of the high-energy tail. The latter indicates that the distribution of the carriers over the high-energy levels is changing, not just the occupation of the levels themselves. Fitting the decaying tail with a Boltzmann distribution,

$$I(h\nu) \propto e^{-E/h\nu/T_e},$$

we are able to extract an effective carrier temperature $T_e$ which starts off at 300 K for low excitation density and reaches up to 1200 K for $\langle N \rangle$ close to 80. A Boltzmann fit is a good approximation if we restrain the fit to high energies ($>2.5 \ eV$; 496 nm) effectively matching the expected energy range of free carrier related transitions, shown by the dotted black curve in Figure 4b. Supporting Information S7 shows moreover that quasi-Fermi level shifts are limited. The lower density limit corresponds well to the data of Sippel et al., who found similar carrier temperatures at low densities.

Figure 4b shows the full dynamics of the extracted temperature for three initial excitation densities: 15, 50, and 78. A single component fit of the lowest density trace shows that the temperature decay is initially very fast with a characteristic lifetime of $\tau_1 = 355 \pm 35$ fs. Fixing this fast lifetime, we can also do a restricted two-component fit on the high density kinetic trace and find a second component of 10 ± 1.4 ps. The latter matches well with the observed gain lifetime in the high-energy region, see Figure 2a. This two-stage cooling corresponds well to the notion of initial fast LO phonon emission followed by overheating of the phonon bath.

Having established that there is a population of high-energy unbound carriers with elevated temperature, we can now rationalize their role in optical gain development. Looking closer at the TA data of Figure 2c, we observe that at high enough density (Figure 2c), the high-energy plateau (<495 nm; 2.50 eV) of the absorbance starts to collapse. Looking back at Figure 1b, this plateau is formed by the staircase absorption profile of a 2D electron gas, in particular from the quantized HH valence band to the conduction band. Filling up these energy levels will indeed lead to a bleach of the plateau $C_{\text{HH}}$ and eventually optical gain, as is observed for densities above ca. 35 electron–hole pairs. To gain more insights, we calculated the intrinsic absorption coefficient for varying carrier densities, considering only the HH continuum $C_{\text{HH}}$ of Figure 1c, see also Supporting Information S7. In Figure 4d, we obtain both the experimentally observed collapse of the absorbance and a clear transition from net absorption to optical gain. A detailed zoom around the gain threshold is shown in Supporting Information Figure S6 and reveals a gain threshold close to 20 excitations. This number is substantially larger than the molecule-mediated gain threshold (4.1) but slightly lower than the observed $\langle N \rangle = 35$, see Figure 2e. This is to be expected because not all excitations are unbound charge carriers and competing LH/HH absorption will counteract the gain buildup. The correspondence of the experimental high-energy gain data with the classical quantum well gain model is excellent as also the gain magnitude, for example, $5 \times 10^4 \ \text{cm}^{-1}$ for $\langle N \rangle = 84$ is retrieved. However, the gain from free carriers is at slightly higher energy than observed experimentally. It is however well established that strongly photoexcited (2D) semiconductors show strong BGR. Estimates based on universal models indicate shifts of 2 excitonic Rydbergs

$$R_y^{2D} = \frac{2\hbar^2}{5e^2a_0^{2D}}$$

up to a density of $\langle N \rangle = (45)$ and reaching even up to 4 Ry2D for higher densities. For CdSe, this would amount to shifts of 60 meV at low density and up to 240 meV at higher density. Figure 4c shows that such a moderate 150 meV renormalization for the highest density indeed pushes the optical gain window in between the two exciton resonances, exactly where optical gain is observed.

4. DISCUSSION

The increased formation of free charge carriers at elevated carrier temperatures and densities over excitons explains the observations made in Figure 3c on the lack of exciton saturation and blue shift. Indeed, exciton saturation through state-filling will be ineffective because carriers do not thermalize to occupy the band-edge levels and additionally, a gas of unbound 2D charge carriers is considered less effective at screening Coulomb interactions than cold excitons. Also the persistent exciton red shift, as was observed in Figure 3c, can be rationalized this way because the position of the exciton transition is a balance between band gap renormalization (BGR) and screening-induced blue shifts. The latter are too weak to overcome the substantial BGR and the overall line red shifts. The remarkable absence of a full Mott-type transition merits further theoretical investigation. Several estimates of this Mott density exist, based on dynamic or static screening arguments. Static screening would put the Mott transition at $n_{aq}^M = 0.1$, which for our system would amount to a mere $\langle N \rangle \approx 14$, clearly a huge underestimation. A possible explanation for the high-energy gain window could be the emission/gain from multie excitons, similar to what is observed in colloidal 0D CdSe-based QDs. It is however important to stress that the photophysics of excitons in 2D materials is fundamentally different from that found in zero-dimensional systems. The example of bie exciton formation is illustrative for this purpose. Whereas in a 0D system, excitons have due to spatial localization no choice than to form the bie exciton state, either repulsive or attractive, this is not valid for a 2D system with lateral extent (326 nm2) much larger than the exciton size (7.2 nm3). Indeed, if the interactions between excitons are repulsive in a 2D system, the excitons simply do not stabilize and retain their single exciton characteristics. As such, repulsive exciton interactions cannot explain the blue-shifted gain window. Another "0D"-argument could be that we are just observing state-filling where optical gain will manifest itself first on the low-energy side and then span out to higher transitions, such as P-type transitions found in QDs, upon increased population. First off, this contradicts our observation that the HH exciton line persists in absorption,
see Figures 2c and 3c. If state-filling was applicable, it should quench similar to the absorption line in 0D QDs which saturates (and inverts) when the population reaches (half of) the band-edge degeneracy and then span out to provide gain of excited exciton states. Moreover, fluorescence and gain of nonthermalized excitons holds no ground because only zero momentum ($K = 0$) transitions are allowed for 2D excitons. Finally, the density of states at the band edge is enormous in a 2D system. Even modest estimates of the number of available states $N_0$ in the 50 meV window taken up by the HH line, puts the number at ca. 230, see Supporting Information S8. An additional proof that state-filling plays no role in the blue-shifted gain band is found by exciting the system at 480 nm. Significantly less energy, ca. 500 meV, needs to be dissipated in this case, leading to much more efficient thermalization. Using the same exciton density, we observe only limited blue-shifted gain under these conditions with a much narrower bandwidth, see Supporting Information S8, again evident that exciton state-filling cannot explain the blue-shifted gain window.

5. CONCLUSION

We have quantified and modeled the distinctive mechanisms leading to room-temperature optical gain in 2D CdSe nanoplatelets, a model system for colloidal 2D materials. At high charge carrier density, a phonon bottleneck slows down hot carrier cooling leading to a unique situation of strong light amplification by unbound charge carriers, up to $10^5$ cm$^{-1}$, and excitonic molecules simultaneously. The remarkable retention of excitonic features until extreme densities indicated an absence of a full Mott-type collapse of the exciton gas, as confirmed by THz spectroscopy. Although it shows from our results that this remarkable property is not directly beneficial for development of very broadband SE due to lingering exciton absorption, it does indicate gain at high exciton densities is feasible with concomitantly high gain coefficients. Moreover, it could render other applications possible such as high brightness LEDs or more complex devices relying on high density multie exciton phenomena such as Bose–Einstein condensation. Indeed, the latter are often hampered by exciton instability at high carrier densities, a problem which is absent in these high exciton binding energy quantum wells. Clearly, (colloidal) 2D materials hold great promise to realize small footprint, low-threshold microlasers. Hence, it would be of great interest to see how these observations translate to other 2D material systems with different types of carrier–carrier and carrier–phonon interactions such as transition-metal dichalcogenides or layered perovskites.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcc.9b02085.

TEM analysis, absorption cross section and gain threshold calculation, more detailed analysis of recombination kinetics, Saha fitting and the quantum well gain model (PDF)

AUTHOR INFORMATION

Corresponding Author

*E-mail: Pieter.Geiregat@Ugent.be.

ORCID

Aditya Kulkarni: 0000-0002-5840-8768

Shalini Singh: 0000-0001-8607-8383
Justin M. Hodgkiss: 0000-0002-9629-8213
Laurens D.A. Siebbeles: 0000-0002-4812-7495
Zeger Hens: 0000-0002-7041-3375
Pieter Geiregat: 0000-0001-7217-8738

Notes

The authors declare no competing financial interest.

REFERENCES


(49) Pelant, I.; Valenta, J. Luminescence Spectroscopy of Semiconductors; Oxford University Press, 2012.


