Imaging Seebeck drift of excitons and trions in MoSe₂ monolayers^{*}

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Hyperspectral imaging at cryogenic temperatures is used to investigate exciton and trion propagation in MoSe₂ monolayers encapsulated with hexagonal boron nitride (hBN). Under a tightly focused, continuous-wave laser excitation, the spatial distribution of neutral excitons and charged trions strongly differ at high excitation densities. Remarkably, in this regime the trion distribution develops a halo shape, similar to that previously observed in WS_2 monolayers at room temperature and under pulsed excitation. In contrast, the exciton distribution only presents a moderate broadening without the appereance of a halo. Spatially and spectrally resolved luminescence spectra reveal the buildup of a significant temperature gradient at high excitation power, that is attributed to the energy relaxation of photoinduced hot carriers. We show, via a numerical resolution of the transport equations for excitons and trions, that the halo can be interpreted as thermal drift of trions due to a Seebeck term in the particle current. The model shows that the difference between trion and exciton profiles is simply understood in terms of the very different lifetimes of these two quasiparticles.

Introduction.— In the last decade, two-dimensional crystals of transition metal dichalcogenides (TMDC) such as MX₂ (M=Mo, W; X=S, Se, Te) have emerged as promising, atomically thin building blocks for applications in nanoelectronics and optoelectronics [1, 2], as well as ideal platforms for the study of Coulombcorrelated excitations and many-body physics [3–5]. Due to quantum confinement, weak dielectric screening and large carrier effective masses in these 2D materials, excitons are characterized by large binding energies and are stable even at room temperature, completely dominating the optical properties of these semiconductors [6–11]. In addition, the interplay between inversion symmetry breaking in monolayers (ML) and the strong spin-orbit coupling induced by the heavy transition metal atoms yields a unique spin/valley locking which is expected to provide additional functionalities in future devices [12–15]. Therefore, a better understanding of the transport of excitonic particles is required for the development of future technologies based on the excitonic properties of TMDC MLs and their heterostructures [16].

Very recently, room temperature experiments investigating the temporal and spatial dynamics of the excitonic photoluminescence (PL) under pulsed laser excitation have reported the appereance of a halo in the PL's spatial distribution at high fluence [17]. This has been attributed to the drift of excitons driven by an effective temperature gradient [18]. Indeed, a strong and focused laser excitation generates an inhomogeneous distribution of hot carriers. Their subsequent relaxation is at the origin of an inhomogeneous lattice and carrier

temperature which in turns pushes excitons out of the excitation spot via the Seebeck effect. In this work, we simultaneously measure the spatial dynamics of excitons and trions in ML MoSe₂ using hyperspectral imaging at cryogenic temperatures. Thanks to the encapsulation by hexagonal boron nitride (hBN), the linewidth of both exciton and trion peaks are close to the homogeneous limit at low temperatures [19], thus allowing for the clear separation of both excitonic complexes and the study of their propagation in the presence of a common temperature gradient. We show that the thermal drift is responsible for the appereance of a halo on the trion's spatial distribution, which is not visible under the exact same conditions for the exciton distribution due to its much shorter lifetime.

Samples and Experimental Set-up. — Van der Waals heterostructures such as the one shown in Fig.1 (a) were fabricated by mechanical exfoliation of bulk MoSe₂ crystals from 2D semiconductors. First, a 250 nm thick bottom layer of hBN was mechanically exfoliated onto a freshly cleaved SiO_2 (80 nm)/Si substrate using a viscoelastic stamp [20]. The deposition of the subsequent MoSe₂ ML and the second hBN capping layer (thickness 8 nm) is obtained by repeating this procedure. The thicknesses of all the layers were measured by Atomic Force Microscopy (AFM).

A hyperspectral micro-PL (μ PL) set-up is used to spatially monitor the exciton and trion concentration at cryogenic temperatures [21, 22]. The samples are kept inside a closed cycle He cryostat and excited either with a red He-Ne laser (1.96 eV) or with continuous wave tunable lasers. Unless otherwise stated, the wavelength of the laser is 570 nm and the sample temperature is 20

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FIG. 1. (a) Schematic drawing of the sample. The MoSe₂ monolayer is encapsulated between two thin h-BN flakes to provide high optical and transport quality. Here, t-hBN and b-hBN stands for top and bottom hBN, respectively. The whole heterostructure is deposited onto a silicon substrate with a 80 nm oxyde layer. (b) Optical microscope image of the sample. (c) Normalized photoluminescence spectrum at T = 300 K (blue) and T = 15 K (red) under a 1.96 eV cw excitation. The full width at half maximum (FWHM) of each peak is indicated.(d) Spatial map of the total PL intensity at room temperature. Despite bubbles and cracks, one can identify zones which are homogeneous over several μ m.

K. The laser beam is focused onto a diffraction-limited spot in the sample plane thanks to a vacuum (and cryogenic) compatible, apochromatic objective mounted inside the cryostat. The resulting PL spot is imaged onto the entrance slit of a 320 mm focal length spectrometer equipped with a 600 grooves/mm diffraction grating. Two methods are used to study the resulting spatial distribution of excitonic complexes: with the slit open, the 0th order specular reflection off the grating is used to image the PL spot onto a cooled Si-CCD camera. Tunable long and short-pass filters were used to selectively image the neutral exciton or the trion's luminesence. This provides, for each line, a spectrally averaged spatial resolution. The second method consists in closing the entrance slit of the spectrometer so that its image on the CCD is only one pixel wide, and an image of the first order diffraction is recorded. In this configuration, the spatial distribution of the PL is kept in the direction parallel to the slit, whereas the spectral content at a fixed position is found by taking a cross section along the direction perpendicular to the slit.

Results and Discussion.— Fig.1 (a) shows a schematic drawing of the Van der Waals heterostructure deposited onto a SiO_2/Si substrate. A microscope image of the sample studied here is shown in Fig.1(b). The PL

spectrum under a 1.96 eV He-Ne laser excitation at room and cryogenic temperatures is shown in Fig.1(c). At low temperature, the spectrum exhibits two well defined peaks, ascribed to the neutral exciton (X^0) and the charged trion (T) which lies 26.3 meV below the neutral exciton energy. The trion forms due to unintentional doping of the MoSe₂ monolayer. A spatial map of the total PL intensity at 300 K is shown in Fig.1(d), exhibiting regions with uniform PL across several μ m, which are therefore suitable for the study of exciton propagation.

We now focus on the power dependence of the PL properties. The excitation power is varied over more than 6 orders of magnitude, from tens of nW up to several tens of mW focused onto a diffraction limited spot, going well beyond the usual power densities used in μ PL experiments. Interestingly, the sample exhibits no evidence of damage after exposure to such power densities, and no irreversible changes in the PL spectrum were observed except for the very first power cycle (see supplementary material), suggesting relatively small photodoping effects which are strongly reduced compared to samples deposited onto SiO₂ thanks to the hBN encapsulation [23]. Figure 2(a) shows the spatially-averaged PL spectrum on a logarithmic scale



FIG. 2. (a) PL spectra for different excitation powers with a continuous wave laser at 570 nm. (b) Integrated PL intensity for trion and exciton peak as a function of excitation power. The lines correspond to a linear relationship between intensity and excitation power. Also shown is the temperature extracted from the low energy tail of the trion peak, here called Trion temperature.

for excitation powers ranging from 15 nW up to 23 mW. At high powers, the linewidth of both lines increases while the peak position redshifts at 20 mW by 2 meV and 1 meV for X^0 and T, respectively. These are typical signatures of local heating of the lattice induced by the optical excitation. Such a redshift of the neutral exciton would correspond to a heating of the lattice up to ~ 55 K according to the temperature dependence of the PL's peak position (see Supplementary material) However, here the laser excitation at high power may locally change by different amounts both the lattice and the carrier temperatures.

In Fig.2(b) we show the integrated intensity for both T and X^0 as a function of excitation power. The continuous lines represent a perfect linear relationship, which is therefore well verified below 100 μ W for both peaks. Remarkably, the trion's PL intensity starts to saturate above 100 μ W and achieves a plateau above 800 μ W, which we attribute to the fact that the photoexcitation density becomes larger than the equilibrium carrier density present due to unintentional doping. By taking a trion lifetime of 140 ps [24] and an absorption coefficient of 1.5×10^{-3} (see supplementary material) we can explain the saturation of the trion's PL with a resident carrier density of $\sim 5 \times 10^{10}$ cm⁻².

In contrast, the X^0 intensity only slightly becomes sublinear above 1 mW, possibly due to Auger-like exciton-exciton annihilation, which has otherwise been shown to be dramatically supressed in hBN encapsulated monolayers. By using a relatively large Auger coefficient of $\gamma \sim 0.1 \text{ cm}^2/\text{s}$ [25–27], the absorption coefficient mentioned above and neglecting diffusion (or thermal drift), we indeed expect exciton-exciton annihilation effects to become relevant only above 1 mW for trions and above 10 mW for neutral excitons, which is in agreement with the observed behaviour of the X^0 PL intensity.

Also shown in Fig.2(b) is the effective temperature of the trion gas extracted from the low energy tail of its luminescence peak, which takes into account the trion's center-of mass momentum dependence of the transition energy [28, 29]. The trion's luminescence intensity is :

$$I(h\nu) \propto e^{-(E^0 - h\nu)/\varepsilon} \Theta \left(E^0 - h\nu \right) * g(h\nu) \tag{1}$$

where $h\nu$ is the emitted photon's energy, E_0 is the emission energy of a trion with zero center-of-mass momentum, g is a Lorentzian representing the non-zero linewidth of the transition, Θ is the Heaviside function, and $\varepsilon \approx 2k_b T$ where k_B is Boltzmann's constant. The latter approximation is valid since the trion's radius is of the order of nanometers and the electron and hole effecive masses are comparable. As seen in Fig.2(b) (blue line and markers), the spatially averaged trion temperature significantly increases above 1 mW. As will be shown later, this spatial averaging strongly underestimates the increase of the effective temperature due to the peculiar spatial distribution of trions at high densities. Note that the trion temperature at low excitation, on the order of 10 K, is smaller than the lattice temperature. This trend is verified at different lattice temperatures, as shown in the Supplementary material. We cannot exclude a systematic error present in the model behind Eq.(1) whose precise determination goes beyond the scope of this work.

In order to study the effect of this local heating on excitonic transport, we performed spatially resolved microluminescence imaging. The diffraction-limited Airy disk of the laser in the sample plane can be approximated at short distances by a Gaussian profile of the form e^{-r^2/w^2} , where $w \approx 0.5 \ \mu m$. Figure 3(a) shows



FIG. 3. (a) PL's spatial distribution of excitons and trions for selected excitation powers. The images have a lateral size of 14 μ m. (b) Normalized radial profile for excitons and trions at low excitation power in a logarithmic scale. Also shown is the laser profile and fits of the 2D diffusion equation. (c) Radial profiles at the highest excitation power, normalized to one at the center in a linear scale. Also shown is the extracted trion temperature from the image of panel (d). The dashed line corresponds to the exciton profile at low power.(d) First order diffraction image at high excitation power, allowing to obtain the spectrum as a function of the distance to the excitation spot.

the spatial images of the luminescence of both excitons and trions, each properly filtered by interferential tunable filters. Remarkably, the spatial distribution of excitons and trions are practically equal at low power, but significantly differ at high excitation, due to the appearance of a halo in the trion's intensity at the center, similar to the halo observed at 300 K on WS₂ monolayers [17, 18], with the main difference here being the cryogenic temperature and a continuous excitation.

Due to the rotational symmetry of the PL images, one-dimensional profiles are obtained by averaging the PL intensity along different radial directions. The result is shown in Fig. 3(b) for an excitation power of 1 μ W, where it can be seen that the exciton and trion luminescence comes from a spatial region which extends significantly beyond the excitation spot. Suprisingly, both profiles are very similar. A fit with the analytical solution of the 2D steady-state diffusion equation [19] yields similar effective diffusion lengths, of $1.49\pm0.05 \ \mu$ m and $1.27 \pm 0.05 \ \mu m$ for X^0 and T, respectively. This result is already surprising given that the lifetime of both species differ by more than one order of magnitude. As suggested by recent experiments in encapsulated MoSe₂ monolayers [24], we interpret these low-excitation profiles as being the result of diffusion of hot excitons, which relax into bright excitons and trions on a timescale of the order of 18 ps. If the subsequent diffusion of excitons and trions is slow enough, then we would indeed expect to see the same profile for both species. Note that momentum-indirect dark excitons can also constitute a reservoir of excitons at the steady state and possibly contribute to the low excitation profile [30].

Now we discuss the high power regime, characterized by the spatial broadening of the PL profiles and the appereance of a dip at the center of the trion's spatial distribution, clearly visible in the radial profile of Fig. 3(c). We interpret this as a thermal drift induced by a temperature gradient. Indeed, the measured trion's

temperature as a function of space is also shown and exhibits a monotonic decrease from 30 K at the center to 15 K at a few microns. This was obtained by recording the 1st order diffraction image on the CCD detector, shown in Fig. 3(d). We attribute the formation of a temperature gradient as a consequence of the energy relaxation of hot carriers by emission of phonons. This gives rise to a hot spot and subsequent thermal drift of excitons and trions. This scenario is in agreement with the fact that the observed temperature gradient exhibits a similar radial profile as the low excitation power profiles observed for X^0 and T, previously attributed to the hot carrier distribution. Note that in these encapsulated monolayers, Auger recombination does not seem to play a significant role in the halo formation under non-resonant excitation, in contrast with the situation found in non encapsulated samples at room temperature [18]. Indeed, if the temperature gradient was created by Auger-like non radiative recombination, we would expect the steady-state spatial dependence of the temperature to be non monotonic and to exhibit a halo as well.

To justify this picture, we have modeled the exciton and trion's profiles by numerically solving the 2D steady state drift-diffusion equation:

$$G - \frac{n}{\tau} - \gamma n^2 = \vec{\nabla} \cdot \left(-D\vec{\nabla}n - \sigma S\vec{\nabla}T \right)$$
(2)

where n is the concentration of either X^0 and T, G is the generation rate, here representing the spatial distribution of hot carriers (low excitation profiles in Fig. 3(b)), τ is the lifetime, γ is the Auger coefficient, $D = (k_B T/q)\mu$ with μ the mobility, q the elementary charge, $\sigma = q\mu n$ is the conductivity and S the Seebeck coefficient, which is of the order of $S \sim 2k_B/q$. Details of this calculation are shown in the supplementary material. Note that the Seebeck coefficient S, which is here a universal constant independent on doping and temperature, is not the same as the phenomenological Seebeck coefficient usually measured in thermoelectric experiments which describes the proportionality between the electrochemical potential gradient (and not the electric field) and the temperature gradient [31]. In addition, in this work we suppose for the sake of simplicity that trions can be modelled by a unipolar diffusion equation. In reality, charge conservation implies that trions cannot diffuse unless the background charge density is modified accordingly. If the trion density remains small compared to the background resident carrier density, which is actually guaranteed by the saturation of the trion formation rate, we may employ a unipolar model. Finally, we also suppose that there is no coupling between the trion and exciton populations, which is in agreement to previous time-resolved measurements where totally independent dynamics has been observed for both complexes.

Figure 4 shows a comparison between the measured radial profiles and the ones predicted by the numerical solution of Eq. (2) when taking a mobility of $\mu = 100 \text{ cm}^2 V^{-1} \text{s}^{-1}$ for trions and $\mu = 150 \text{ cm}^2 V^{-1} \text{s}^{-1}$ for excitons. The Seebeck coefficients were chosen as $S = 300 \ \mu V/K$ and $S = 400 \ \mu V/K$ for trions and excitons, respectively. The trion lifetime of $\tau = 140$ ps is extracted from [24] and the exciton lifetime is estimated to be $\tau = 5$ ps by considering an intrinsic radiative lifetime (in vacuum) of 2.7 ps and cavity effects. The model reproduces well the main features of both the exciton and trion profiles with a reasonable choice of adjustable parameters. At this stage, we cannot claim a precise measurement of the Seebeck coefficient, since for example the mobility μ is unknown and we do not claim a precise determination of the absolute trion temperature. We have chosen $\mu = 100 \text{ cm}^2 V^{-1} s^{-1}$ for the trion which is low enough to ensure a low power profile dominated by hot exciton diffusion. For the excitons, we have taken into account the effective mass difference, although little is known about the momentum relaxation time for both species.

Note that the Seebeck coefficients for excitons and trions do not need to be equal. Indeed, for a classical 2D gas for which the momentum relaxation time τ depends on the kinetic energy ε as $\tau(\varepsilon) \propto \epsilon^r$, the Seebeck coefficient writes $S = (2+r)k_B/q$ and r can differ between excitons and trions.

Finally, we show that the trion's halo formation can be further enhanced with resonance excitation. This is in contrast with recent experiments performed on high purity GaAs samples, for which halo formation of excitons was attributed to a spatially dependent population balance between excitons and free electron hole plasma, and shown to disappear under resonance excitation [32]. Here, the laser energy is set to match the X^0 energy. The absorption, of around 5 %, is estimated by considering optical cavity effects. Fig. 5(a) shows the obtained first order diffraction images at different excitation powers. At high powers, the dip in the trion's luminescence becomes very pronounced, with a maximum trion intensity at distances lager than 2 μ m from the center at 20 mW, as shown in the radial profiles in Fig. 5(b). The spatial dependence of the trion's temperature is shown in Fig. 5 (c), where it can be seen that the temperature gradient is a monotonic function of the excitation power below 10 mW. The fact that a temperature gradient exists despite resonant excitation could indicate an important role played by resonant exciton-exciton Auger scattering involving an excited conduction band, which was recently proposed to explain the very efficient PL upconversion observed in MoTe₂ and MoSe₂ MLs under resonant excitation of the neutral exciton [33].

The inset of Fig. 5(c) shows that the temperature at the excitation spot starts to saturate above 10 mW and even decreases at 20 mW. Note that a saturation of the



FIG. 4. (a) Measured PL's radial profile for trions (left) and excitons (right) for selected excitation powers. (b) Calculated profiles obtained with a numerical solution of Eq.(2)

excess temperature of a hot exciton gas at high densities has been previously reported in encapsulated WSe₂ MLs and attributed to the balancing between Auger heating and phonon cooling [27]. In addition, due to the resonant excitation condition, at the highest excitation powers the redshift of the exciton transition may result in a lower absorption coefficient, counterbalancing the effect of a higher excitation on the photocreated density. Fig. 5 (d) shows the trion's peak energy as a function of space. Above 5 mW, a non-monotonic dependence is observed. with a remarkable blueshift-redshift crossover near the center. Note that such a crossover of the exciton absorption energy on ML WS_2 at 300 K has been reported and attributed to the onset of a repulsive exciton-exciton interaction at the Mott density 2×10^{12} cm⁻²[34]. Here, the crossover can be equally attributed to a repulsive interaction between hot excitons which, near the center, are then converted into trions by binding to an extra charge carrier.

Conclusions In conclusion, we have imaged the spatial distribution of trions and excitons in ML $MoSe_2$ at cryogenic temperatures as a function of excitation power. We reveal the appereance of a halo in the trion's distribution, which so far was only observed under pulsed laser excitation at room temperature. Hyperspectral imaging permits a measurement of the trion temperature as a function of space and confirms

the existence of a strong temperature gradient at high excitation. The results are consistent with the following simple picture: photons generate hot excitons out of the light cone, which then diffuse over typically one micron before relaxing their energy and convert into either bright excitons or trions. This energy relaxation creates a temperature gradient which extends across the hot carrier distribution. Excitons and trions then drift under the presence of this temperature gradient, which dominates over diffusion. The measured exciton profiles, which do not exhibit a halo but a moderate broadening, are consistent with the exciton having a much shorter Finally, we have shown that this halo can lifetime. be observed under resonant excitation of the neutral exciton, which can be explained by the fact that resonant exciton-exciton interactions creates hot excitons, thus mimicking the effect of an off-resonance excitation. This work is a step towards a more detailed understanding of the different physical phenomena that can govern excitonic transport at high excitation fluences.

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FIG. 5. (a) Spatial dependence of the trion's luminesence spectrum for selected excitation powers when exciting resonantly the neutral exciton (b) Normalized trion profiles obtained from the figures in (a) by integrating the trion's spectrum at different distances from the excitation spot. (c) Trion's temperature as a function of position. The inset shows the measured temperature at the excitation spot. (d) Trion's peak energy as a function of position.

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