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Exciton Condensation in an Atomically-thin MoS\textsubscript{2} Semiconductor

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Condensation of a dilute Bose gas of excitons (coupled electron-hole pairs) in a direct-bandgap semiconductor was first theoretically predicted in 1968\textsuperscript{1}. This exotic state of matter is expected to exhibit spectacular non-linear properties, such as superradiance and superfluidity. However, direct experimental observation of condensation of optically-active excitons in conventional semiconductors has been hindered by their short lifetimes and weak collective excitonic interactions. Here, we have experimentally realized the condensation of short-lived excitons in a direct-bandgap, atomically-thin MoS\textsubscript{2} semiconductor. The signature is the anomalous transport of the fast-expanding exciton density, originating from a thermalized dilute gas generated under the laser spot. Below the critical temperature $T_c \sim 150$ K, the exciton liquid propagates over ultra-long distances (at least 60
micrometers) with record speed in a solid-state system of $\sim 1.8 \times 10^7$ m/s ($\sim 6\%$ the speed of light), fuelled by the unconventionally strong repulsions among excitons. The condensation is controlled by many-body interactions in the gas mixture of excitons (bosons) and free-carriers (fermions) via an electrical backgate. Our results demonstrate electrostatic doping as a simple approach for the investigation of correlated states of matter at high-temperatures, excitonic circuitry and spin-valley Hall devices mediated by exciton superfluids in semiconducting monolayers.

Similar to ultracold atomic gases $^{2-3}$, a dilute gas of excitons optically injected in a semiconductor below a critical temperature ($T_c$) is well-described as a weakly interacting Bose gas that can spontaneously condense (in momentum space) into a liquid, via many-body repulsive interactions among electron-hole pairs $^{1,4}$. Despite the efforts over the last five decades, experimental observation of condensation of optically-active (short-lived) excitons in direct-bandgap semiconductors has not been realized. To stabilize the condensate in these systems, the repulsive interactions among excitons must compensate the fast decay of the population. Furthermore, the formation of other complexes via attractive forces, such as charged states (trions), molecules (biexcitons) or even electron-hole droplets $^5$, are competing effects against exciton condensation. To overcome the limitations in direct-bandgap semiconductors with weakly-interacting excitons, top-down approaches have been taken. For instance, by using microcavities, polaritons with ultra-light effective masses are obtained, allowing for metastable Bose condensation at high densities $^6$. Alternatively, quantum fluids have been reported in double quantum-wells (DQWs) by using optically-forbidden (long-lived) dipolar excitons, whose electron and holes resides in opposite QW layers (spatially-indirect) $^7$-$^{13}$. In DQWs, the long lifetimes and the tailored repulsive interactions among indirect dipolar excitons drive the condensation.

Excitons in direct-bandgap two-dimensional semiconductors, such as monolayer transi-
tion metal dichalcogenide (TMD) MoS$_2$, have short lifetimes and stable exciton complexes
\textsuperscript{14–16}, which a priori may preclude the formation of exciton liquids. Here, by using spatially-, time- and energy-resolved photoluminescence (PL) and reflectivity measurements, we have experimentally realized the condensation of a dilute exciton gas in MoS$_2$ monolayer until $T_c \sim 150$ K, evidenced by the anomalous transport over long distances of at least 60 $\mu$m (limited by the sample size) with record speed in a solid-state system of $\sim1.8 \times 10^7$ m/s ($\sim$6% the speed of light). The condensation is driven by unconventionally strong repulsive interactions among excitons, directly revealed by $\sim$3 meV blueshift of the exciton energy. We also present the phase diagram by using an electrical backgate and temperature as relevant thermodynamic parameters that control the interplay between attractive (exciton-electron) and repulsive (exciton-exciton) many-body interactions present in the Bose-Fermi gas mixture in the monolayer semiconductor.

In our PL imaging experiments, we used a laser as excitation source with a well-defined Gaussian spatial distribution with full width at half-maximum (fwhm) of $\sigma_{\text{Laser}} \sim 1.5$ $\mu$m. By using near-resonant excitation (Fig. 1(A)), a low density gas of excitons (up to $n_X \sim 1.1 \times 10^{10}$ cm$^{-2}$, well below the Mott density $n_{\text{Mott}} \sim 10^{13}$ cm$^{-2}$) is photogenerated with minimal momentum ($\hbar k_X \sim 0$), kinetic energy ($E_X(k_X) = E_g + \hbar k_X^2/2m^*_X$) and group velocity ($v_X(k_X) = \partial E_X/\partial k_X$), where $\hbar$ is reduced Planck constant and $m^*_X$ is exciton effective mass. In this way, the exciton gas reaches thermal equilibrium with the semiconductor lattice right after photoexcitation ($E_X(k_X)-E_X(0) \sim k_B T$, where $k_B$ is the Boltzmann constant and $T$ is temperature), persisting within the radiative lifetime of the state.

Monolayer MoS$_2$ has large intrinsic concentration of electrons that markedly affects the optical response of the two-dimensional semiconductor. We control the electron density ($n_e$) and enhance the excitonic response in our samples by incorporating a backgate ($V_g$). At large positive $V_g$ (i.e., high $n_e$), the reflectance contrast in Fig. 1(B) is dominated by a single reso-
nance, denoted as $X^-$, that corresponds to the so-called charged exciton or trion state, or also described as attractive polaron. In this metallic regime ($V_g > 20$ V), the neutral exciton, also referred to as repulsive polaron, is absent. Decreasing $V_g$ (lower $n_e$), the excitonic resonance $X$ starts appearing at 38 meV above $X^-$. A further decrease in $V_g$ leads to a strong increase in the exciton absorption and a monotonic shift to lower energies. The $X^-$ feature disappears below $V_g = -30$ V. At this insulating regime ($V_g < -30$ V), only the neutral exciton resonance ($E_X(0) \sim 1.950$ eV) is resolved.

The change in the character from metallic to insulating of the MoS$_2$ monolayer drives an instability in the excitonic system that results in an anomalous excitonic transport and nonlinear optical effects. Therefore, to accurately study this, we have produced high-quality large area ($A \sim 1200 \mu$m$^2$) MoS$_2$ monolayers. The longest dimensions of our samples of 40-60 $\mu$m (Fig. 1(C) and Figs. S4-S6) are 30-40 times larger than $\sigma_{\text{Laser}}$ that allows us to monitor the long-range propagation and distribution of the exciton population in space via PL imaging as a function of $V_g$, power and temperature.

The low-temperature ($T = 20$ K) PL emission at $V_g = 0$ V is localized around the laser spot, positioned at the center of the sample (Fig. 1(D)). Remarkably, maintaining the same excitation conditions but setting $V_g = -60$ V, we observe light-emission with homogeneous intensity distribution over the entire crystal (Fig. 1(E)) outside the laser spot. Such PL profile suggests that excitons have homogeneously spread over large distances away from the excitation source (see Fig. S5 for reproducibility in other devices). Given the negligible kinetic energy and low mobility of the exciton gas $^{18-24}$, such a long-range transport is striking and has not been theoretically predicted for TMDs.

For a fixed power ($P = 1000 \mu$W) and temperature ($T = 20$ K), the sudden changes in the spatial profile on the light-emission with varying $V_g$ can be seen in the Supporting Video,
while PL image cross sections are shown in Fig. 2(A). When $V_g$ decreases from 0 to -20 V, the PL intensity profile remains unchanged and exhibits a bimodal distribution. Most of the PL intensity ($\sim 75\%$) follows a Gaussian distribution centred at the excitation spot ($x_{\text{Laser}} = 0 \ \mu m$). The Gaussian profile (region-I) indicates diffusive transport of the exciton gas away from the generation source. The fwhm at region-I $\sigma_I = (1.6\pm0.2) \ \mu m$ is comparable to $\sigma_{\text{Laser}}$ and manifest the low mobility of a non-interacting exciton gas in TMDs $^{18-24}$. The shape of this distribution remains unaffected upon variations of $V_g$, $P$ and $T$ (Fig. 2(A)-(C)).

The PL profiles exhibit a tail that extends further into the sample beyond region-I, which suggests a weakly interacting exciton gas at $V_g = 0$ V. Its intensity follows an exponential distribution (region-II). Intriguingly, when decreasing $V_g$, the amplitude and decay length ($L_X$) of the exponential distribution increase non-linearly (Figs. S7-S8). In particular, at $V_g = 0$ V, region-II represents the $\sim 25\%$ of the PL intensity that ultimately increases to $\sim 90\%$ at $V_g = -60$ V. The acquired quasi-homogeneous intensity of the PL profile at large negative voltages (limited by the finite size of the crystal), typically observed in condensed polaritons, indicates ballistic-like transport over micrometer distances of the short-lived excitons in monolayer MoS$_2$.

Moreover, the PL pattern at region-II also varies non-linearly with excitation power as shown by Fig. 2(B) at $V_g = -60$ V and $T = 20$ K. With increasing power, the exponential decay becomes longer and asymptotically approaches a step-like shape at the highest injected exciton density. Remarkably, keeping $P = 1000 \ \mu W$ and $V_g = -60$ V, the step-like profile is observed until exceptionally high lattice temperatures of $\sim 150$ K (Fig. 2(C)). Further increase of $T$ leads to a rapid decay of the exponential component of the PL profile that becomes negligible ($<10\%$) above 180 K (Fig. S6).

Such striking instabilities of the excitonic density in region-II are attributed to a thermodynamic phase transition, where a fraction of the dilute gas of excitons has spontaneously
condensed into a liquid. The formation of the exciton liquid is revealed by the anomalous transport behaviour imprinted in the exponential PL distribution. Therefore, we use the exponential decay constant ($L_X$), defining the transport length of the exciton fluid, as a characteristic macroscopic parameter to closely monitor the phase transition, which originates from microscopic many-body interactions, as a function of $V_g$, $P$ and $T$.

The phase diagram of $L_X (V_g, T)$ is shown in Figure 2(D) for $P = 1000 \, \mu W$. The liquid phase domain (left-side of the dashed guideline) shows the largest transport length (up to $L_X = (64 \pm 3) \, \mu m$) that extends until $V_g = -30 \, V$ and high-temperature of $\sim 150 \, K$, abruptly shortening by one-order-of magnitude ($L_X = (5 \pm 4) \, \mu m$) when entering the gas phase (right-side of the dashed guideline). Intriguingly, exciton condensation occurs even with low excitation power provided that the temperature and the electron density are low enough (Fig. S9). Unlike in conventional phase diagrams for indirect excitons$^{25}$, here in the case of direct excitons in MoS$_2$ monolayer, $V_g$ is an important parameter, because the fraction of the exciton gas that condenses depends critically on the electron density present in the two-dimensional semiconductor. The liquid phase is observed only when monolayer MoS$_2$ approaches the insulating regime (Fig. 1(B)).

Microscopically, what distinguishes the exciton liquid from the gas is the competition between repulsive (exciton-exciton) and attractive (exciton-electron) interactions within the mixture of excitons (boson) and electrons (fermion) (illustration Fig. 2(D)). In the gas phase, the short transport length suggests that excitons are spatially localized by the strong attractive interactions induced by the dense electron gas. In this metallic-like regime, the many-body repulsive exciton-exciton interactions required for their condensation are screened, regardless of the exciton density, which implies attractive forces are dominant to the leading order $n_e/n_X > 1$.

Nominally, interactions between electrons and excitons remain attractive for all $n_e^{26}$, 

which prevents for the formation of the exciton liquid. The short-range attractive interactions reduce when gradually decreasing $n_e$ to become smaller, or of order of $n_X (n_e/n_X \leq 1)$. In the insulating regime, the low electron density (i.e., negligible screening) allows for the repulsive exciton-exciton interactions to take over, giving rise to the spontaneous condensation of the dilute exciton gas.

Moreover, the fraction of condensed excitons is also markedly affected by temperature. Given the large binding energy of MoS$_2$ excitons, we estimate that the density of thermally excited electrons surpasses $n_X$ at $T = (160 \pm 40)$ K (Fig. S12). This range of temperatures depends on the intrinsic carrier concentration and agrees well with the gas-to-liquid phase transition. The critical exciton condensation temperature ($T_c \sim 150$ K) observed here for direct-bandgap MoS$_2$ monolayer is much higher than that for superfluid helium (2.17 K), indirect excitons in III-V DQWs ($< 10$ K), III-V and II-VI polaritons ($< 10$ K) and Cu$_2$O excitons ($< 1$ K) and is similar to the estimated temperature for tightly-bound indirect excitons in TMD DQWs. The remarkably high-$T_c$ suggests unusually strong repulsions between excitons occupying the lowest-energy state and sets a record for condensation and achievement of exciton superfluidity in direct-bandgap semiconductors.

The macroscopic expansion of the exciton liquid in the atomically-thin MoS$_2$ surface causes the up to three-orders-of magnitude increment of the integrated PL intensity with decreasing $V_g$ (Fig S10-11) that explains for the near-unity PL quantum yield reported in neutral MoS$_2$ samples. Such giant PL enhancement, or superradiance, indicates that light-emission of the liquid originates from optically-active (bright) excitons. Therefore, the long-range propagation of excitons within their ultra-short radiative lifetime, typically $\tau_X \sim 1-20$ ps in TMDs, strongly suggests the unconventionally high-speed of the exciton fluid because $L_X(T, V_g) = v_{X,L}(T, V_g) \cdot \tau_X$.

We demonstrate the ultrafast transport by performing spatially- and energy-resolved pump-
probe reflectance measurements with 150-femtosecond time-resolution. A broadband-energy source (probe) illuminates the entire MoS$_2$ crystal, while the single-frequency beam (pump) excites the center of the sample ($x = 0 \mu m$) with a reduced beam size of $\sigma_{\text{Laser}} \approx 2 \mu m$. The pump, tuned near the exciton resonance, generates an exciton gas with estimated density $n_X \sim 1.7 \times 10^{12}$ cm$^{-2}$ under the laser spot. After gas thermalization, as excitons condense via repulsive interactions, they are propelled away from the generation source, altering the spatial-dependence of the intensity and energy distribution of the reflectance profile through the monolayer.

Figure 3 shows the differential reflectance signal $DR(x, E)$ at $V_g = -60$ V for representative delay times ($\Delta t$) between pump and probe. At $\Delta t = 0$ ps (Fig. 3(A)) defined as the time frame at which the photogenerated density under the laser spot is thermalized at the exciton resonance energy, a weak positive $DR$ signal (photobleaching) arise from the exciton population at $E_X(0) = 1.950$ eV. Increasing $\Delta t$, the photobleaching feature becomes pronounced and extends through space (Fig. 3(B)-(D)), reaching 20 $\mu m$ in just 1.2 ps. On the other hand, other states at energies below $X$ do not propagate proving that the long-range transport observed in Figures 1 and 2 is entirely determined by excitons, which is further supported by the suppression of the propagation when $n_e$ is high at $V_g = 0$ V (Fig. S15). These observations have been reproduced in multiple devices (Fig. S16).

Figure 4(A) shows the time-dependence of the mean-square displacement (MSD) $\sigma_{\text{II}}^2 - \sigma_{\text{II,0}}^2$, where $\sigma_{\text{II,0}}^2$ is the fwhm at $\Delta t = 0$ ps. Similarly to ballistic transport, the MSD in the liquid phase (circles) scales super-linearly with increasing time, which is in clear contrast to the linear trend at the gas phase (squares) that is characteristic of diffusive transport. Analysis of the MSD with an parabolic function (solid-line) provides the velocity at which the exciton liquid expands.
over the atomically-thin semiconducting layer, which is $v_{X,L} = (1.8 \pm 0.3) \times 10^7 \text{ m/s}.$

The obtained velocity is exceptionally high, being only 6% the speed of light ($3 \times 10^8 \text{ m/s}$). Remarkably, $v_{X,L}$ is one-order-of magnitude larger than condensed polaritons in ultra-high quality micro-cavities in III-V \cite{6,32}, ZnO \cite{33}, than the Fermi velocity of Dirac electrons in graphene \cite{34,35} and in metals ($v_F \approx 10^6 \text{ m/s}$) and than ballistic excitons in perovskites semiconductors \cite{36,37}, three orders of magnitude larger than excitons fluids in III-V single \cite{38} and DQWs \cite{7}, in Cu$_2$O \cite{39} and II-VI \cite{40}. Therefore, the unconventionally high speed makes condensed excitons in direct-bandgap monolayer MoS$_2$ the fastest excitation in electronic matter observed in a solid-state system.

Within Keldysh’s framework \cite{1,4}, the group velocity of condensed excitons follows the hydrodynamic relation $v_{X,L} \propto \sqrt{f}$, where $f$ characterizes the many-body interactions within the Bose-Fermi gas mixture. To stabilize the condensate against disorder and the formation of charged excitons, biexcitons and electron-hole droplets, $f$ takes real positive values. In monolayer MoS$_2$, $f = f(T, V_g)$ depends on temperature and $V_g$, and defines the phase diagram in Fig. 2(D). With increasing $f$, the exciton liquid expands through the two-dimensional crystal with remarkably large speed $v_{X,L}$, suggesting a frictionless, or superfluid flow of condensed excitons.

The unconventionally strong repulsions among excitons causes a significant blue-shift of the exciton feature in the pump-on with respect to the pump-off raw reflectance spectra (Figs. S17-S18). We estimate the net repulsive energy $\Delta E_{X-X}$ arising from exciton-exciton interactions, which is shown in Fig. 4(B) for $\Delta t = 2.4 \text{ ps}$. In the range 0-5 $\mu$m, $\Delta E_{X-X}$ has the largest value of $\sim 3 \text{ meV}$, comparing well to that of dipolar indirect excitons \cite{11} and polaritons \cite{32} condensates. $\Delta E_{X-X}$ follows the decay of the exciton population given by $DR(x, E_X(0))$
(solid-pink curve) that decreases slowly to $\sim 1$ meV far from the laser spot.

In conclusion, we have realized the condensation of ultrashort-lived excitons in a direct-bandgap, atomically-thin MoS$_2$ semiconductor. The condensation resulting from many-body interactions within the mixture of Bose (excitons) and Fermi (free-carriers) gases, controlled by an electrical backgate, takes place when exciton-exciton repulsions dominate over exciton-electron attractions and remains stable until $T_c \sim 150$ K. The exciton liquid propagates over long-distances (at least 60 micrometers) with record velocity of $\sim 1.8 \times 10^7$ m/s ($\sim 6\%$ the speed of light). Our findings have implications in high-temperature superconductivity, excitonic circuitry and spin-valley Hall devices mediated by excitonic superfluids for the next-generation quantum-technologies by using semiconducting monolayers.
Fig. 1: Exciton condensation driven by an electrical-backgate in an atomically-thin MoS$_2$ semiconductor. (A) Parabolic exciton dispersion $E_X(k_X)$ (black curve). A dilute exciton gas is generated using near-resonance excitation (red wavy-arrow) with energy $\hbar \omega_{\text{Exc}}$, propagates with small group velocity $v_X(k_X) = \hbar^{-1} \partial E_X(k_X)/\partial k_X$ (blue-arrow) and finally radiatively recombine within the light cone (dashed-line) with energy $\hbar \omega_{\text{PL}}$ (gray solid-line). (B) Reflectance contrast showing the charged ($X^-$) and neutral ($X$) exciton resonance energy as a function of photon energy and backgate $V_g$. (C) Optical image of an electrically contacted monolayer with large area of $\sim$1200 $\mu$m$^2$. The sample contour is highlighted by the dashed-line. Right-bottom inset: Device schematic. Low-temperature (20 K) spatially-resolved photoluminescence images at (D) $V_g = 0$ V and (E) -60 V. All images have been taken using the same magnification.
Fig. 2: **Exciton phase diagram.** Spatial cross-section profiles in a semilogarithmic scale of the photoluminescence images as a function of (A) backgate, where $V_g$ varies from 0 V (bottom curve) to -60 V (top curve) with a step of -10 V, taken at fix $P = 1000 \mu W$ and $T = 20 K$; (B) power, where $P = 10 \mu W$, 50 $\mu W$, 100 $\mu W$ and 1000 $\mu W$ from bottom to top, taken at fix $V_g = -60 V$ and $T = 20 K$; (C) temperature, where $T = 200 K$, 180 K, 150 K, 80 K and 20 K from bottom to top, taken at fix $P = 1000 \mu W$ and $V_g = -60 V$. The sharp drop of the PL intensity in (A), (B) and (C) corresponds to the monolayer edge. All spectra have been normalized to the maximum intensity of the exponential distribution for easy comparison among experiments. (D) Phase diagram of the fluid transport length $L_X(V_g, T)$. The liquid (gas) phase domain, where the electron density is low (high), occupies the left-side (right-side) of the dashed guideline. The scale bar is given. Illustration: At low (high) electron (solid-dark spheres) density, excitons (transparent-light spheres) experience strong (weak) collective repulsions causing exciton condensation and propagation over long distances.
Fig. 3: Transport of condensed excitons. Spatially- (horizontal-axis) and energy-resolved (vertical-axis) differential reflectance \( (DR(x, E)) \) at delay times \( (\Delta t) \) of (A) 0 ps, (B) 0.2 ps, (C) 0.6 ps and (D) 1.2 ps, measured at fixed conditions \( T = 20 \) K, \( V_g = -60 \) V and \( n_X \sim 1.7 \times 10^{12} \) cm\(^{-2}\). \( DR(x, E) \) is defined as \( \frac{R_{On}(x, E) - R_{Off}(x, E)}{R_{Off}(x, E)} \), where \( R_{On}(x, E) \) and \( R_{Off}(x, E) \) are the reflectance profiles with the pump on and off, respectively. The positive (negative) \( DR(x, E) \) signal indicates strong (weak) reflectance, corresponding to pump-induced photobleaching (pump-induced absorption) of a state at energy \( E \) at position \( x \). The spot of the pump laser is located at the center of the monolayer \( (x = 0 \, \mu m) \). The photobleaching at the exciton resonance propagates 20 \( \mu m \) after a pump-probe delay of \( \Delta t = 1.2 \) ps, clearly demonstrating the unconventionally high propagation speed of condensed excitons of the order of \( 10^7 \) m/s.
**Fig. 4: Non-linear transport and energy blueshift (repulsive interaction) of condensed excitons.**

(A) Temporal-evolution of the mean-square-displacement (MSD) $\sigma^2_{II} - \sigma^2_{II,0}$ as a function of $\Delta t$ for excitons at the liquid (blue circles, $V_g = -60$ V) and gas (purple squares, $V_g = 0$ V) phase. The quadratic behaviour of the MSD is reproduced (red curve) by using a power law $\sigma^2_{II} - \sigma^2_{II,0} = v_{X,L}^2 t^\alpha$ with $\alpha = 2$, which demonstrates the ballistic-like propagation of the exciton liquid. The liquid speed $v_{X,L} = (1.8 \pm 0.3) \times 10^7$ m/s. Dashed-lines represent upper and lower limits of the MSD fitting, which provides the uncertainty of $v_{X,L}$. 

(B) Energy change of excitons $\Delta E_{X-X}$ at the liquid (blue circles, $V_g = -60$ V) and gas (purple squares, $V_g = 0$ V) phase as a function of position $x$. Large blueshifted energy (positive $\Delta E_{X-X}$) is detected in the liquid phase, due to the strong repulsive exciton-exciton interaction, while no net repulsion is observed at the gas phase ($\Delta E_{X-X} \sim 0$). $\Delta E_{X-X}$ decreases as exciton liquid propagates in space following the decrease in the density (solid-pink curve, right axis).
Methods

Device Fabrication

Large MoS$_2$ monolayers with the typical long side of 40 to 60 $\mu$m and area of $\sim$1200 $\mu$m$^2$ were mechanically exfoliated from natural bulk MoS$_2$ crystals onto a polydimethylsiloxane (PDMS) stamp. The MoS$_2$ monolayers were then carefully transferred from the PDMS stamp onto clean 285 nm SiO$_2$/Si p-type dry oxide substrates, ensuring that smooth, uncracked large area monolayers are obtained upon transfer. The successful transfers were verified under an optical microscope by optical contrast. A poly(methyl methacrylate) (PMMA) mask was spin coated on the substrates and patterned electrical contact areas were defined on the mask by standard electron beam lithography. 5 nm Cr/50 nm Au electrical contacts were subsequently deposited by thermal evaporation. The final devices were obtained after an acetone lift-off procedure. To control the electron concentration of the MoS$_2$ devices in our optical experiments, a backgate voltage was applied to the Si substrate and the MoS$_2$ monolayers grounded via the electrical contacts.

Optical Imaging and Experimental Setup

The fabricated devices were mounted on a cold finger cryostat and cooled down using a continuous flow of liquid helium. The cryostat station was mounted on the photoluminescence (PL) and transient reflectance (TR) imaging setup that operated in two different imaging configurations (see SI for details).

In the PL imaging configuration, the samples were excited with a He-Ne (632.8 nm) continuous wave (cw) laser cleaned by an interference filter resonant with the A-exciton of the MoS$_2$ monolayer. Using a 50× objective lens (NA = 0.55), the laser was focused to locally excite a small spot with diameter of 1.5 $\mu$m on the large sample. The objective then collected the PL emission. An ultra-steep-edge long pass filter placed in the detection path before the
optical detector removes reflected laser light, allowing only PL image signal to be measured. A
set of lenses were added in the detection path to change the image magnification and effective
field of view on the optical detector (sCMOS sensor). This enables imaging to be optimized
for samples of different sizes. The sCMOS chip has an area of $2300 \times 2300$ pixels, 6.5 $\mu$m
each. Our detection system provides a high spatial resolution of 40 nm per pixel in either axial
direction.

For TR imaging, a femtosecond pulsed pump laser (615 nm) is used to excite the sam-
ple. A pulsed broadband white light beam given a time delay $\Delta t$ with respect to the pump is
used to probe the changes in the reflectance of the sample in the visible (625 – 700 nm) region.
The size of the probe beam illuminating the sample surface is controlled by a series of lenses,
which focuses the probe at the back focal plane of the objective lens to maximize the probe
and cover the entire sample. The reflected probe beam was collected by the same objective and
dispersed by a single grating (600 gr/mm) monochromator with 550 mm of focal length and
the signal was detected by a scientific CCD detector cooled with liquid-nitrogen. Energy-space
reflectance spectrum map images are measured for each time delay under pump on/off condi-
tions. As the technique relays in the changes in the optical reflectance/absorption, the dynamics
of the exciton population within the light-cone, which includes the spatial propagation, radiative
recombination and relaxation into other states, is monitored in great detail.

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**Data availability**

All methods and data generated and/or analyzed supporting the findings of this study are included in this paper and its Supplementary Information file and are available from the corresponding authors upon reasonable request.
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Author contributions

A.G.D.A. and Q.X. started the project. A.G.D.A. conceived and designed the experiments. Y.R.W and X.L. designed and fabricated the large area devices. A.G.D.A, Y.R.W, X.L. and A.F. performed the experiments. Q.X., M.B., Y.R.W. and T.T.H.D. intensively discussed the results and the manuscript during its preparation. A.G.D.A analyzed and interpreted the data and wrote the manuscript with the help of all co-authors. A.G.D.A and Y.R.W contributed equally to this work.

Competing interests

The authors declare no competing interests

Supplementary Information

A supporting information file with information regarding sample fabrication, experimental details, analysis, fitting procedures, theoretical section and additional data supporting the main findings of the main manuscript file is available.
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