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Controlled interlayer exciton ionization in an electrostatic trap in atomically thin heterostructures

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Atomically thin semiconductor heterostructures provide a two-dimensional (2D) device platform for creating high densities of cold, controllable excitons. Interlayer excitons (IEs), bound electrons and holes localized to separate 2D quantum well layers, have permanent out-of-plane dipole moments and long lifetimes, allowing their spatial distribution to be tuned on demand. Here, we employ electrostatic gates to trap IEs and control their density. By electrically modulating the IE Stark shift, electron-hole pair concentrations above $2 \times 10^{12} \, \mathrm{cm^{-2}}$ can be achieved. At this high IE density, we observe an exponentially increasing linewidth broadening indicative of an IE ionization transition, independent of the trap depth. This runaway threshold remains constant at low temperatures, but increases above 20 K, consistent with the quantum dissociation of a degenerate IE gas. Our demonstration of the IE ionization in a tunable electrostatic trap represents an important step towards the realization of dipolar exciton condensates in solid-state optoelectronic devices.

Semiconductor van der Waals (vdW) heterostructures offer a unique platform where strong light-matter interactions enable novel device capabilities that expand our ability to study fundamental mesoscopic phenomena. As their building blocks, monolayer transition metal dichalcogenides (TMDs) provide a synergy of electronic and photonic degrees of freedom that manifest as excitons, optically generated bound electron-hole pairs, with large binding energies¹ and spin-valley locking properties^{2,3}. In type-II TMD heterostructures, such as MoSe₂/WSe₂, electrons and holes are localized on separate 2D quantum well layers. This band alignment leads to the formation of IEs with

permanent out-of-plane dipole moments⁴⁻⁷, extended lifetimes^{4,8,9}, and long interaction-driven diffusion lengths^{4,7,10}, while maintaining strong binding energies¹¹⁻¹³. These properties also make IEs ideal for the design of excitonic devices with engineered spatial emission profiles and controllable IE density with electrostatic gates.

The generation and control of large IE concentrations is desirable to explore the rich physics predicted by the excitonic phase diagram. In GaAs double quantum wells, indirect excitons have been shown to condense into a degenerate state ¹⁴⁻¹⁶ that is expected to be a superfluid, providing a route to dissipationless optoelectronic devices.

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While increasing exciton density is generally favorable for condensation, increasing wavefunction overlap eventually leads to complete screening of the IE binding energy at the Mott density¹⁷⁻¹⁹. Even below this critical condition, a fraction of the IEs can be thermally ionized at a finite temperature, enhancing the screening effect. Previous theoretical studies showed that this density dependent screening leads to a runway IE dissociation process near the ionization threshold²⁰⁻²⁵. In TMD heterostructures, the orders of magnitude larger binding energies allow the system to sustain larger IE populations before ionization effects become relevant⁹. This property opens the door to the realization of electrically tunable high-temperature condensates^{13,26,27}.

Electrostatics traps are a useful tool for exploring the IE phase diagram. They allow us to deterministically localize IEs and increase their density, bypassing the need for high laser powers that lead to heating or nonlinear Auger recombination^{28,29}, and potentially even damage to the sample. Furthermore, the control over their diffusion dynamics opens the avenue for reliably designing excitonic devices, as other trapping approaches like moiré localization depend sensitively on sample homogeneity^{30–33}. Separate control of the IE density and temperature paves the way towards constructing tunable coherent optoelectronic devices and studying correlated phenomena emerging from them.

Results

Electrostatic tunability of interlayer exciton emission

In this article, we design a multi-gated structure to create electrostatic traps that allow us to generate high densities of IEs, control their

spatial distribution, and determine their critical dissociation density as a function of temperature. All the data shown is from device D1 unless otherwise stated. Figure 1a shows a schematic of the device design. A high quality, h-BN encapsulated MoSe₂/WSe₂ heterostructure is fabricated with matching top and bottom metal gates (Methods). A matching set of grounded ~250 nm-wide stripe gates maintains a constant IE energy in the trap region. On either side of the stripe are two sets of top and bottom gates separated by ~500 nm that control the IE energy in the outside region. Applying voltages to the outer gates creates a neighboring electric field, $E_{og} = \frac{\varepsilon_{h-BN}}{\varepsilon_{TMD}} \frac{V_{tg} - V_{bg}}{t_{total}}$, where $\varepsilon_{h\text{-}BN}$ = 3.9, ε_{TMD} = 7.2 are the h-BN and TMD dielectric constants, V_{tg} and $V_{h\sigma}$ are the top and bottom gate voltages, and t_{total} is the total h-BN thickness. Applying positive E_{og} raises the IE energy in the outer region due to the linear Stark effect4. We note that the intralayer exciton energies are not affected in this gating configuration since their dipole moments are in the TMD plane⁴. The electric field for the center stripe gates is approximately zero since these are grounded. The relative field difference between the stripe gate region and the outer area creates a confinement potential along the former so that a high IE density can accumulate in the trap without diffusing away (red curve in Fig. 1b). Figure 1c shows an optical image of the device detailing the spatial region for the experiment.

We demonstrate our gate modulated trap by performing scanning photoluminescence (PL) spectroscopy at various E_{og} across the stripe gates (Methods). All PL measurements are performed at T = 4 K, unless otherwise stated, with an excitation wavelength of 660 nm (1.87 eV), and a diffraction limited spot. In Fig. 1d, we measure the PL spectra as a

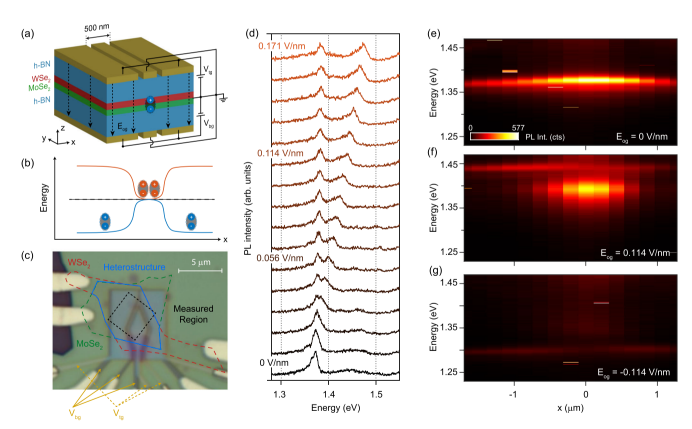


Fig. 1 | **Modulating the interlayer exciton energy with electric fields. a** Device schematic showing the stripe gate and large outer gates on either side. V_{tg} and V_{bg} are applied to create electric fields in the outer gate (E_{og}) regions while the stripe gates are grounded. **b** IE energy schematic as a function of x-position perpendicular to the stripe gate. The red (black dashed) curves represent the electric field profile felt by the IEs with (without) E_{og} in a trapping configuration. The blue curve represents the anti-trap regime. **c** Optical image of the device. Red and green dashed lines outline the WSe₂ and MoSe₂ regions, respectively. Blue line outlines

the heterostructure region. Gold arrows identify the relevant top gate and bottom gate structures. The black dashed square indicates the approximate region for the PL diffusion measurements. **d** IE spectra as a function of $E_{\rm og}$. The linearly shifting peak at higher energy is from the outer gate region while the lower energy peak is from within the stripe gate region. **e**-**g** Scanning PL spectroscopy, where the laser excites at x = 0 µm and the spectra is collected across the stripe gate at $E_{\rm og} = 0$ and ± 0.114 V/nm, showing spatially modulated IE energy.

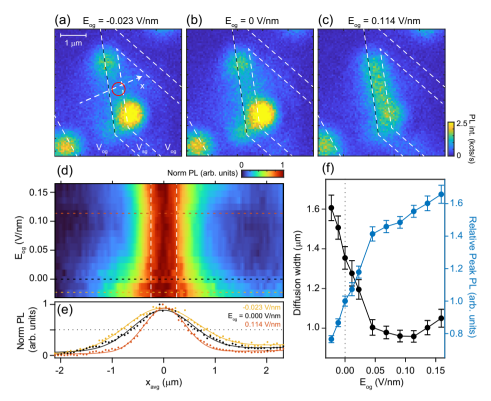


Fig. 2 | **Controlling the diffusion behavior.** \mathbf{a} - \mathbf{c} Spatial mapping of the diffused IE PL emission at $E_{\rm og}$ = -0.023, 0, and 0.114 V/nm. The outer gate and the stripe gate areas are labeled $V_{\rm og}$ and $V_{\rm sg}$, respectively. White dashed arrow indicates positive x direction. Red circle is the excitation position. Bright spots that naturally occur on the sample are reduced with the trapping potential. \mathbf{d} Normalized PL along the x direction, averaged over the uniform y-cut trap region, as a function of $E_{\rm og}$. The dotted lines correspond to the $E_{\rm og}$ line cuts in (\mathbf{e}). The white dashed lines indicate

the expected trap width of ~ 500 nm. **e** Line cuts (scatter) and Gaussian fittings (lines) at $E_{\rm og}$ = -0.023, 0, and 0.114 V/nm showing narrowing (broadening) of the diffusion width when forming a trap (anti-trap). **f** Extracted diffusion width and relative peak intensity (PL_{peak}(E_{\rm og})/PL_{peak}(E_{\rm og}=0)) as a function of E_{\rm og}. We observe the trap saturates near $E_{\rm og} \approx 0.04$ V/nm. Error bars come from the fitting of the spatial PL distribution. Note that "a.u." is the abbreviation for "arbitrary units".

function of E_{og} at a power (P) of 50 μ W at the edge of the stripe gates. As E_{og} increases, a second peak splits off, with only the higher energy peak shifting linearly with the external field. We extract a dipole separation of $d \approx 0.6$ nm from the slope of the energy shift $\Delta E =$ edE_{og} (e is the elementary charge), corresponding to the expected interlayer separation between the electron and hole wavefunctions localized on the TMD layers. In Fig. 1e-g, we excite the heterostructure with $P = 500 \,\mu\text{W}$ at the center of the stripe gate and measure the PL spectra at a transverse distance x away from the center of the stripe gate for three representative fields, $E_{og} = 0 \text{ V/nm}$ and $\pm 0.114 \text{ V/nm}$. When $E_{og} = 0 \text{ V/nm}$, the PL energy is constant at ~1.38 eV across the spatial linecut, consistent with the expected electrostatic profile. When $E_{og} = 0.114 \text{ V/nm}$, the PL energy in the outer gate region rises to ~1.44 eV, consistent with the gate dependent spectra, while the stripe gate region emits lower energy. Thus, we conclude that the higher energy peak originates from IEs outside the trap, and we identify the lower energy peak as the trapped IEs. Regardless of the trapping potential, we observe that the higher-energy free IE emission extends nearly 3 µm away from the center of the stripes. The observed long diffusion lengths of IEs in TMDs is caused by dipolar repulsion⁴. However, we note that the high energy emission is darker when the confinement potential is active, hindering IE diffusion perpendicular to the stripe, while favoring localization in the trap. We also demonstrate an anti-trap potential when $E_{og} = -0.114 \text{ V/nm}$ (Fig. 1g), where the overall emission is darker because the electrostatic profile helps IEs diffuse away from the excitation spot (see also Supplementary Fig. S1).

Spatial control over interlayer exciton dynamics

To understand the diffusive behavior under confinement, we take spectrally filtered PL intensity maps at the IE energy. Figure 2a-c shows

the spatially dependent IE PL emission for anti-trap, flat, and trap potentials when the excitation spot is fixed near the edge of the stripe gate for $P = 550 \,\mu\text{W}$ (red circle in Fig. 2a). For $E_{og} = -0.023$ and 0 V/nm, we observe emission around the laser spot and at uncontrolled disorder-related bright spots. In the absence of the externally applied trapping potential, IEs diffuse and localize in these naturally occurring traps (Supplementary Fig. S2b, c). However, when $E_{og} = 0.114 \text{ V/nm}$, we observe uniform and elongated emission along the stripe, implying that the IE diffusion is constrained along it. In Fig. 2d, we average the emission along the stripe (y-direction) and plot the normalized intensity as a function of x perpendicular to the stripe for various E_{og} . We observe a narrowing of the spatial IE cloud width when the trap is active and a broadening when an anti-trap is created. In Fig. 2e, we show three characteristic spatial PL profiles obtained along the linecuts shown in Fig. 2d, at constant E_{og} corresponding to anti-trap (yellow), no-trap (black), and trap (red) configurations. These profiles exhibit a spatially narrowed (broadened) distribution for $E_{og} = 0.114 \text{ V/}$ nm (-0.023 V/nm) compared to 0 V/nm. We perform a similar analysis along the stripe gate and find that the emission is uniform above $E_{og} \approx 0.09 \text{ V/nm}$ (Supplementary Fig. S3), corresponding to a trap depth of ~55 meV. This estimated natural trap energy scale is consistent with reports of defect or strain-trapped IEs in these heterostructures34,35.

The emission profile across the stripe gate can be described by the balance between the diffusion of laser-generated IEs near the edge of the stripe, and the electrostatic trap (anti-trap) that hinders (favors) exciton transport. We obtain the spatial full-width half-maximum (FWHM) centered at the trap using a Gaussian fit. Figure 2f shows the FWHM as a function of E_{og} . We observe that the trap diffusion width saturates near 0.95 μ m at $E_{og} \approx$ 0.04 V/nm. The FWHM is wider than the

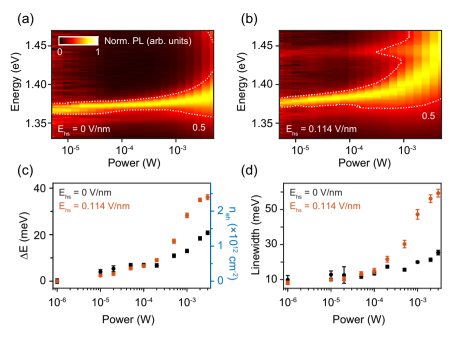


Fig. 3 | **Tuning the interlayer exciton density. a** Normalized power dependent PL emission from the stripe gate region at $E_{og} = 0$ V/nm, when the excitation laser is slightly outside the region. The dotted white line shows a contour of 0.5 for the intensity of the trapped IEs. **b** Same as (**a**) for $E_{og} = 0.114$ V/nm. **c** The blueshift

energy, ΔE , for $E_{\rm og}$ = 0 and 0.114 V/nm as a function of power. Right axis (blue) shows the corresponding calculated electron-hole pair density, $n_{\rm eh}$. **d** Fitted trapped PL linewidth for $E_{\rm og}$ = 0 and 0.114 V/nm as a function of power. The error bars in (**c**) and (**d**) are from the fitting of the spectral peaks.

expected trap width (-500 nm), which could be explained by imperfect alignment of the top and bottom gates or by convolution with the point spread function of the system (Supplementary Information S4). We note that the FWHM increases slightly for $E_{og} > 0.12$ V/nm, probably due to a shorter IE lifetime and greater emission outside the trap for larger electric fields⁴. The relative peak PL (normalized to the emission at $E_{og} = 0$ V/nm) also increases rapidly until E_{og} reaches -0.04 V/nm, before increasing at a slower rate. We observe the longest IE lifetime at the center stripe region when a trapping potential is applied, which increases by up to 20% at the largest E_{og} (Supplementary Fig. S5). These signatures indicate stronger confinement at higher IE densities.

Generation of a high-density trapped electron-hole ensemble

The PL peak energy shifts as a function of electron-hole pair density due to interactions. In the excitonic limit, strong dipolar repulsion leads to a blueshift for increasing IE density^{4,5,9,18,19,36-38}. Similarly, an increasing density of unbound electrons and holes on opposite layers generates an electric field leading to a blueshift^{18,39,40}. Figure 3a shows the PL spectra as a function of the excitation power P at the center of the trap (see Fig. 2). Even without trap activation (i.e., $E_{og} = 0$), the PL peak exhibits a strong blueshift as P increases (Fig. 3a). This increase of the IE emission energy ΔE can be described by a balance between the diffusion, generation, and emission of IEs⁴. With increasing P in the excitonic regime, the electron-hole pair density n_{eh} under the stripe gates increases as excitons diffuse from the excitation site, leading to an increase in ΔE . The pair density can be further increased by deepening the confinement potential: a larger E_{og} hinders diffusion out of the trap as in Fig. 2f, leading to a larger blueshift (Fig. 3b) for the same power range compared to the flat potential case.

For a more quantitative estimate of n_{eh} , we employ a formula obtained by considering dipolar excitonic interactions using a mean-field approach³⁹: $\Delta E = \frac{(n_{eh} - n_0)e^2d}{\varepsilon_{TMD}\varepsilon_0}$, where n_0 is the initial density from which we reference ΔE , and ε_0 is the vacuum permittivity. Although this linear relationship tends to underestimate the pair density⁴⁰, we treat ΔE as an indication of increasing electron-hole concentration in

the trap. In Fig. 3c, we plot the measured ΔE (left axis) and the estimated n_{eh} (right axis) at the center of the stripe gate region as a function of P with (orange symbols) and without (black symbols) the trap potential activated. We estimate a small initial density n_0 for the lowest measured excitation power (P=1μW) (Supplementary Information S6). Since n_0 and the blueshift at low powers are small, this estimation contributes only a small error at higher densities. At low powers ($P < 10^{-4}$ W), we find that n_{eh} behaves similarly with and without a trap potential, increasing slowly with increasing P. This behavior suggests that IE diffusion is confined to the trap region, independent of the trap depth, and therefore n_{eh} is limited by the IE generation and recombination rates. However, at high powers ($P > 10^{-4}$ W), the maximum n_{eh} with the trap on is nearly twice as large as with the trap potential off, indicating that the electrostatic profile confines electronhole pairs that would otherwise diffuse away at the same generation rate. Without a trap, we obtain a maximum $n_{eh} = 1.38 \times 10^{12} \, \text{cm}^{-2}$, in agreement with previous measurements⁴. With the trapping potential, we reach almost twice this density: $n_{eh} = 2.40 \times 10^{12} \, \text{cm}^{-2}$. We note that the enhanced blueshift occurs only for trapped IEs and not for IEs outside of the trap even at E_{og} = 0.114 V/nm (Supplementary Fig. S6). This confirms that the blueshift enhancement is a consequence of increasing interactions from higher local electron-hole densities.

In addition to the blueshifts, we observe a broadening of the linewidth, shown by the contour lines of half maximum normalized intensity (Fig. 3a, b). Figure 3d shows the fitted linewidth as a function of power for $E_{og} = 0$ and 0.114 V/nm. At lower excitation powers, we have a linewidth of ~10 meV in both cases. At higher E_{og} , we observe an increase in the linewidth with lower power, similar to the deviations observed in n_{eh} as discussed above. We rule out heating as the main mechanism causing this trend, since a temperature increase due to laser heating would lead to linewidth broadening and peak redshifting regardless of the trap potential set by E_{og} (see S.11).

Interlayer exciton ionization phase diagram

The rapid increase of the spectral linewidth at high power invites a more detailed investigation. Figure 4a shows the density dependent

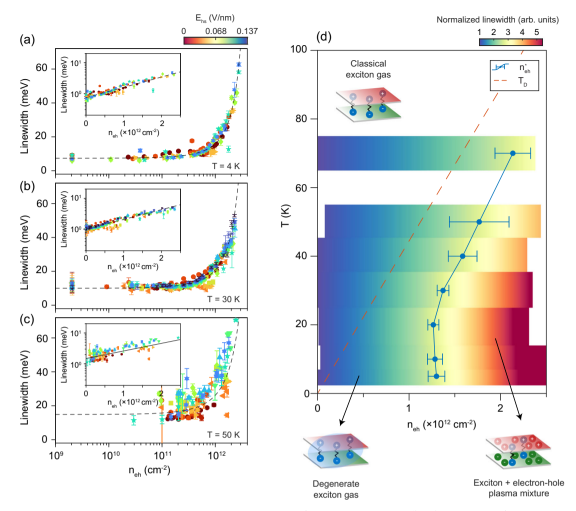


Fig. 4 | **Interlayer exciton phase diagram.** a Scatter plot of linewidth vs. electronhole density for various trap depths at T = 4 K. The error bars indicate the uncertainty of the fittings. The black dashed line shows the exponential fit giving a universal critical density $n_{eh}^*(4\text{ K}) = 1.30(\pm 0.09) \times 10^{12} \text{ cm}^{-2}$. Inset: Same data but where linewidth is on a log scale and density on linear scaling, showing the exponential fit is shown as a linear curve. **b**, **c** Same as (**a**) but at 30 K with

 $n_{eh}^*(30~\text{K}) = 1.37(\pm 0.06) \times 10^{12} \, \text{cm}^{-2}$ and 50 K with $n_{eh}^*(50~\text{K}) = 1.77(\pm 0.32) \times 10^{12} \, \text{cm}^{-2}$, respectively. **d** Plot of the fitted normalized linewidth over the collected data range as a function of pair density and measured temperature. The blue data shows the extracted $n_{eh}^*(T)$ at different temperatures with the error bars showing the uncertainty of the fits. The red dashed line is the degeneracy temperature, T_D as a function of exciton density per flavor.

linewidth $w(n_{eh})$ at different E_{og} and P. As discussed in Fig. 3, n_{eh} can vary over a wide range (10^9 to 2×10^{12} cm⁻²). Interestingly, we find that $w(n_{eh})$ follows a universal curve determined solely by n_{eh} , independent of E_{og} and P. We find that the linewidth increases exponentially with n_{eh} (inset of Fig. 4a), suggesting that we can fit the behavior with the functional form $w(n_{eh}) = w_0 \exp\left(\frac{n_{eh}}{n^2}\right)$, where w_0 is an intrinsic linewidth at low pair density and n_{eh}^* is a characteristic density for the exponential upturn. We consider a rapid increase in the spectral linewidth for $n_{eh} > n_{eh}^*$ as a signature of the ionization threshold of IEs, equivalent to the Mott transition 9,17,20-24,37,41. At base temperature, we estimate $n_{eh}^*(4K) = 1.30 \ (\pm 0.09) \times 10^{12} \ cm^{-2}$ (Fig. 4a). We observed similar behavior exponential behavior in two additional devices, D2 and D3, with different stacking orientations and interlayer h-BN spacing (see Fig. S11 and Supplementary Section S10 for further discussion). Interestingly, all heterostructures exhibit an $n_{eh}^* \sim 1 - 2.5 \times 10^{12} \, \text{cm}^{-2}$. For D1, we repeated similar measurements at different temperatures T ranging from 4 K to 70 K, where $n_{ab}^*(T)$ can be similarly obtained from the exponentially increasing linewidth (see Fig. 4b, c for representative data at higher temperatures and more data is available in Supplementary Fig. S8). Figure 4d shows the

normalized linewidth as a function of n_{eh} and T. In this plot we also show $n_{eh}^*(T)$ obtained from the exponential fit. For $T \le 20$ K, we find that $n_{eh}^*(T)$ is constant within our fitting uncertainty, while above 20 K, $n_{eh}^*(T)$ increases with increasing temperature.

The broadening of the spectral linewidth as a function of n_{eh} has been previously observed for indirect excitons in GaAs double quantum wells^{17,18,38,41} and for IEs in MoSe₂/WSe₂ heterostructures^{9,37}, and is attributed to the dissociation of the IEs. Similarly, we attribute n_{eh}^* to the density driven ionization threshold for IEs. As n_{eh} increases, Coulomb screening becomes more pronounced, reducing the binding energy and gradually increasing the fraction of ionized carriers²¹. At n_{eh}^* , the mixed IE-plasma system reaches a threshold and ionization becomes a runaway process^{20–25}, which can explain the rapid increase of the PL linewidth, although a complete theoretical model of the exponential dependence is beyond the scope of our work. Following this scenario, the temperature dependent behavior of $n_{eh}^*(T)$ shown in Fig. 4d is understood by considering the competing effects of screening and bosonic degeneracy. Here, the three different length scales that characterize this competition are the excitonic Bohr radius a_B , the mean interpair distance $D = n_{eh}^{-1/2}$, and the excitonic thermal de

Broglie wavelength $\lambda_T = \left(\frac{2\pi h^2}{m_{lr}k_BT}\right)^{-\frac{1}{2}}$, where m_{lE} is the total IE mass. In the low temperature and high density regime where $D < \lambda_T$, the quantum statistical nature of the bosonic IE becomes appreciable. The condition $\lambda_T = D$ defines the degeneracy temperature per flavor $T_D = \frac{2\pi h^2}{k_B m_{JE} n_{eh}}$ (red dashed line in Fig. 4d), which marks the crossover between the classical (non-degenerate) and quantum (degenerate) IEs. Experimentally, we find that the temperature-dependent ionization threshold $n_{ch}^*(T)$ occurs well below this line for all measured temperatures in our experiment, indicating that the transition we observe is related to the quantum dissociation of the IEs. Below 20 K, we find that $n_{oh}^*(T)$ remains at a nearly constant value of ~1.3 × 10¹² cm⁻². This n_{eh}^* behavior can be understood in terms of the charge-neutral nature of IEs leading to a self-stabilization effect^{20,21} and is consistent with claims of a temperature-independent Mott transition at the lowest temperatures¹³. It is worth noting that the Mott density n_M at which the binding energy vanishes is larger than n_{lE}^* , but of the same order²¹. Theoretically, the excitonic Mott density is expected to be given by $n_M a_B^2 \sim 0.1$ in our experiment^{13,42,43} (see Supplementary Information S9). Using $a_B \approx 1.9$ nm¹¹, we estimate $n_M \sim 2.8 \times 10^{12}$ cm⁻², providing a consistent upper bound for our measured n_{eh}^* below 20 K. At higher temperatures, we find that $n_{eh}^*(T)$ increases with increasing T with a similar slope to the degeneracy limit defined by $\lambda_T = D$. Here, as T increases, λ_T becomes smaller and a higher IE density is required for IE wavefunction overlap to become significant enough to reach the ionization threshold^{20-22,24}.

Discussion

We note that in similar systems with an h-BN spacer layer, an opposite trend for the IE phase diagram has been observed $^{44-47}$. However, due to the increased electron-hole separation, the IEs in those systems have a binding energy that is an order of magnitude smaller than in our study. For weakly bound IEs, temperature might be relevant in promoting IE dissociation, resulting in a reduction of the Mott density. In a strongly bound IE system, as studied here, the temperature is always below 10% of the binding energy and ionization physics is dominated by the quantum ionization process described above. Our arguments are supported by a recent exciton drag experiment, which observed a similar $n_{eh}^*(T)$ trend as in our work at the lowest temperatures, before transitioning to a thermal ionization dominated regime 45 .

Using a gate-tunable trap architecture, we have demonstrated a novel feature of critical IE dissociation dynamics: an exponential linewidth broadening. This reveals a crucial piece of the IE phase diagram, and we hope future theoretical studies will help address its microscopic origins. Combining these strong many-body effects with highly gate tunable devices paves the way for novel optoelectronic devices such as exciton transistors^{10,48,49} or high population density inverted cavity-less lasers⁵⁰. Furthermore, depth-tunable traps are an important step towards funneling a controlled density of cold dipolar excitons away from the excitation spot, thereby eliminating any laser-induced heating or coherence effects that could prevent unambiguous identification of a condensate.

Methods

Device fabrication

Figure 1c shows our h-BN encapsulated $MoSe_2/WSe_2$ heterostructure device D1 with electrical contacts to each layer. The $MoSe_2$, WSe_2 , and h-BN layers are prepared via mechanical exfoliation on 285 nm SiO_2 substrates. The $MoSe_2$ and WSe_2 bulk crystals are grown by the flux method. The h-BN thicknesses are 36 nm and 59 nm for the top and bottom h-BN, respectively. The stack is assembled by picking up each layer starting with the top h-BN using a standard dry transfer method. The final stack is placed on ultra-flat Cr/PdAu alloy (1 nm/9 nm) bottom

gates written on a 285 nm SiO_2 substrate. The stripe gate width (\sim 250 nm) and outer gate separations (\sim 500 nm) are measured via scanning electron microscopy images (Supplementary Fig. S13). The same gates are evaporated on top of the completed heterostructure. Edge contacts were fabricated by reactive ion etching parts of the monolayer region with O_2 /CHF $_3$ /Ar gas mixture and then evaporating Cr / Au leads (5 nm/120 nm). It should be noted these contacts are not sufficient for electronic transport but can be used for electrostatic doping measurements.

Measurement details

Optical measurements were performed using a 4f confocal microscope system with a 0.75 NA 100x objective in a Montana Instruments, closedloop optical 4 K cryostat. Measurements were performed at T=4 K, except for temperature dependent measurements. PL measurements were performed with a Thorlabs continuous wave diode laser with excitation wavelength of 660 nm and collected into a spectrometer with a PIXIS:256BR or BLAZE:400HR camera (both Princeton Instruments) unless otherwise noted. Scanning images were taken by fiber coupling into an avalanche photodiode (Excelitas SPCM-900-14-FC). Time resolved PL measurements are performed with a supercontinuum laser from NKT with an 80 MHz repetition rate and through a tunable bandpass filter (SuperK VARIA). The laser is synchronized to a single photon counting module (Excelitas Technologies) and a picosecond event timer (Picoharp 300, PicoQuant). Scanning confocal photoluminescence spectra are taken by using two separate optical paths to excite and collect from separate positions. The optical path is split using a 50:50 pellicle beamsplitter, each with a set of controllable two-axis galvanometer mirrors. The excitation channel has a co-aligned collection line that can be used to choose the excitation position based on PL map. With the excitation fixed, the collection channel is scanned to measure PL emission counts or spectra from the various parts of the sample. All PL measurements presented in the manuscript are measured this way unless stated otherwise. The PL map in Fig. S2b is taken where the excitation and collection are co-aligned. All electrostatic gates are controlled via Keithley 2400 sourcemeters.

Data availability

All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Information. All raw data generated during the current study are available from the corresponding author upon request.

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Author contributions

A.Y.J., L.A.J., and P.K. conceived the study. A.Y.J., A.M.M.V., L.A.J., K.P., D.D., Y.Z., G.S., K.D.G. and A.S. performed the experiments. A.Y.J., L.A.J., and K.P. fabricated the device. B.K. and J.H. performed the TMD crystal growth. T.T. and K.W. performed the h-BN crystal growth. M.D.L., H.P., and P.K. supervised the experiments. A.Y.J., A.M.M.V., and P.K. analyzed the data and wrote the manuscript with extensive input from all authors.

Competing interests

The authors declare no competing interests.

Additional information

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