

Highly tunable ground and excited state excitonic dipoles in multilayer 2H-MoSe₂

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The fundamental properties of an exciton are determined by the spin, valley, energy, and spatial wavefunctions of the Coulomb-bound electron and hole. In van der Waals materials, these attributes can be widely engineered through layer stacking configuration to create highly tunable interlayer excitons with static out-of-plane electric dipoles, at the expense of the strength of the oscillating in-plane dipole responsible for light-matter coupling. Here we show that interlayer excitons in bi- and tri-layer 2H-MoSe₂ crystals exhibit electric-field-driven coupling with the ground (1s) and excited states (2s) of the intralayer A excitons. We demonstrate that the hybrid states of these distinct exciton species provide strong oscillator strength, large permanent dipoles (up to 0.73 ± 0.01 nm), high energy tunability (up to -200 meV), and full control of the spin and valley characteristics such that the exciton g-factor can be manipulated over a large range (from -4 to $+14$). Further, we observe the bi- and tri-layer excited state (2s) interlayer excitons and their coupling with the intralayer excitons states (1s and 2s). Our results, in good agreement with a coupled oscillator model with spin (layer)-selectivity and beyond standard density functional theory calculations, promote multilayer 2H-MoSe₂ as a highly tunable platform to explore exciton-exciton interactions with strong light-matter interactions.

A range of exotic collective effects are predicted to arise from dipolar interactions^{1,2}, which have a quadratic dependence on the magnitude of the static electric dipoles (p). For example, strong dipolar interactions may result in exciton crystals, which exhibit ordering due to a balance between exciton kinetic energy and many-body Coulomb interactions³⁻⁷, or lead to nonlinear exciton switches which can reach the quantum limit when the strength of the interaction is larger than the exciton's radiative linewidth⁸⁻¹⁰. Hence, in the solid-state, much emphasis has been placed on engineering interlayer excitons with large p in pioneering III-V heterostructures¹¹ and more recently in

transition metal dichalcogenide (TMD) heterostructures¹²⁻¹⁴ which host excitons with huge binding energies and thus small Bohr radii^{15,16} that enable high exciton densities^{17,18}. In TMD heterostructure devices, tunable interlayer excitons with large p have been realised in homobilayers¹⁹⁻²⁹ and heterobilayers³⁰⁻³³, even at the single exciton level³⁴⁻³⁶. However, many of the exotic collective effects underpinned by strong dipolar interactions remain to be observed, motivating further exploration of interlayer excitons and ways to manipulate their spin and optical properties. For example, it is desirable to increase the electron-hole spatial separation beyond the interlayer distance, but

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not at the cost of vanishing oscillator strength. This goal is intrinsically difficult for bare interlayer exciton states (e.g. in TMD heterobilayers with type II interfaces). Recent experimental and theoretical efforts^{21,23,37} suggest that each ‘bare exciton’ is actually composed of a mixture of other exciton wave functions, and their mixture can be tuned by detuning energy between transitions. Here we use the term bare exciton to denote the majority of each transition for simplicity. While previous studies focused on hybrid interlayer excitons formed in adjacent ‘natural’ homobilayers^{21–23}, an open question remains if larger hybrid interlayer dipoles, with the electron and hole highly confined in separate layers, can be generally obtained in multilayer TMD platforms. Further, due to the decreasing oscillator strengths of excited Rydberg-like states^{15,16,38}, the observation of excited states of interlayer excitons has proven elusive to date. Characterisation of the excited state spectrum of interlayer excitons provides additional information about their basic properties and a potential means to further engineer dipolar interactions by taking advantage of their larger Bohr radii, which has been crucial to realise optical nonlinearity³⁹ and excitonic analogues of spatially ordered structures in ultracold atomic gases⁴⁰. To address these issues, we take advantage of hybridised excitons in bilayer (2L) and trilayer (3L) 2H-MoSe₂ composed of the bare exciton states of the interlayer excitons (both 1s and 2s) and the ground (1s) and excited (2s) states of the intralayer A-excitons. We simultaneously demonstrate strong oscillator strength and wide tunability of the fundamental properties (spin, valley, energy, and spatial wavefunction) of the hybrid exciton species in multilayer MoSe₂.

Results and discussion

Device structure and introduction to excitons in multilayer 2H-MoSe₂

Figure 1a shows a sketch of our device, which consists of a terraced 2H-MoSe₂ flake with 2L and 3L regions encapsulated by hexagonal boron nitride (h-BN) layers with nearly identical thicknesses (~18 nm). Graphene layers act as electrical contacts for the MoSe₂ crystal and the top and bottom gates (see method section for further details of the device fabrication). In our experiments, the MoSe₂ contact is grounded while we apply voltages to the top and bottom gates, labelled as V_T and V_B , respectively. This configuration allows us to apply a vertical electric field with a magnitude V_E (where $V_E = V_B = -V_T$) while keeping the carrier concentration in the MoSe₂ sample constant at charge neutrality. Similar to other TMDs, each layer in our terraced 2H-MoSe₂ flake hosts tightly bound intralayer excitons with both ground (1s) and excited (2s, 3s, etc.) states¹⁵. Ground and excited intralayer exciton states present

the same exact spin-valley configuration. In the case of the lowest energy excitons in MoSe₂ (the so-called A excitons, X^A), both the ground and excited exciton states originate from Coulomb-bound electron-hole pairs in the lower- (higher-) lying conduction (valence) band at $\pm K$ (see Fig. 1b), respectively, which endows them with the same optical selection rules: excitons at $\pm K$ couple to σ^\pm -polarised light⁴¹. In addition to X^A , there are intralayer B excitons (X^B) composed of the electron (hole) at the top (bottom) of the conduction (valence) band at $\pm K$, respectively, with a considerable energy difference and opposite spin index compared to X^A .

Beyond intralayer exciton states, multilayer TMDs also host interlayer excitons, in which the layer-localised electron wave functions can bind to holes with wave functions confined predominantly within adjacent layers or spread across several layers, giving rise to excitons with spatially extended wave functions^{19,21–25}. Interlayer and intralayer excitons have been shown to coexist in multilayer²⁰ and 2L 2H-MoSe₂²⁴. Figure 1a shows a sketch of the possible intra- and interlayer exciton spatial configurations in our terraced 2L/3L 2H-MoSe₂, where we have assumed that, despite the possible spatial spread of the carrier wave functions, the carriers are predominantly confined to a single layer. This assumption leads to three different bare exciton species: (i) intralayer excitons within each individual layer (i.e. ground and excited states of the A and B exciton series in MoSe₂⁴²); (ii) interlayer excitons with the carriers residing in adjacent layers (IX_{2L}), in which the electron and hole occupy the upper spin-orbit-split conduction band and the topmost valence band of each layer, respectively (see Fig. 1b); and (iii) interlayer excitons in which the carriers reside in the outermost layers of the 3L MoSe₂ region (IX_{3L}), where the electron (hole) occupies the lowermost (topmost) conduction (valence) band (see Fig. 1b). In the 3L MoSe₂ region, IX_{2L} can be formed with carriers from the middle MoSe₂ layer (L_2) and carriers from either the bottom (L_1) or top (L_3) MoSe₂ layers. Note that, despite the spatially indirect character of IX_{2L} and IX_{3L} , these interlayer exciton species exhibit momentum-direct (intravalley) optical transitions within $\pm K$. Moreover, the vertical displacement of the electron and hole wave functions of the bare interlayer excitons results in an out-of-plane static electric dipole (negligible for intralayer excitons), with a dipole polarity and magnitude that depend on the positions and the spatial separation of the electron and hole in the multilayer, respectively (see Fig. 1a). Such out-of-plane permanent electric dipoles of interlayer excitons in other TMD multilayers, homostructures, and heterostructures have been shown to lead to large shifts of the exciton transition energies via the quantum confined Stark effect^{21–23,32,35,43}. Finally, the natural 2H

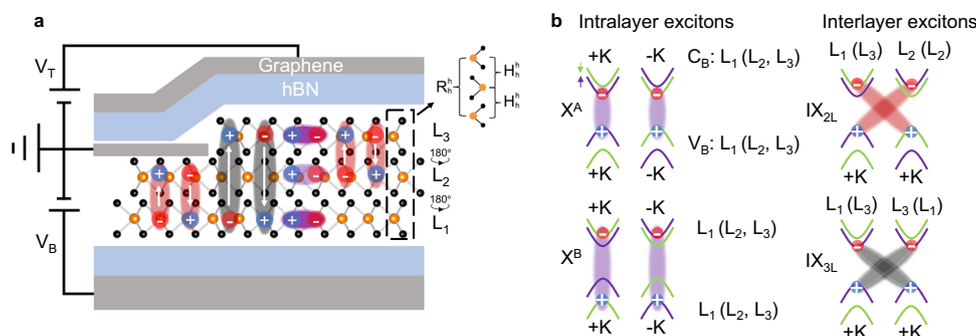


Fig. 1 | Intralayer and interlayer excitons in a terraced 2L/3L 2H-MoSe₂ sample.

a Sketch of the sample and device used in this work: a 2H-MoSe₂ crystal with 2L- and 3L-thick terraces in a dual-gated device configuration. Electrons and holes (red and blue circles, respectively) can be localised either in the bottom (L_1), middle (L_2), or top (L_3) MoSe₂ layers, giving rise to different species of strongly bound intralayer and interlayer excitons. The vertical white arrows indicate the direction of the permanent electric dipoles for the different interlayer excitons. The interlayer stackings are highlighted as R_h^h between L_1 and L_3 and H_h^h between L_1 (L_2) and L_2 (L_3),

where h denotes the hexagon centre in the crystal lattice of each layer. **b** Spin-valley configurations of the intralayer and interlayer excitons depicted in panel a: intralayer A and B excitons (purple shaded ovals), bilayer and trilayer interlayer excitons (red and grey shaded ovals, respectively). The layer labels (L_1 – L_3) on each panel indicate the layer origin of the corresponding electronic states in the conduction (C_B) and valence (V_B) bands. The purple (green) bands correspond to spin up (down).

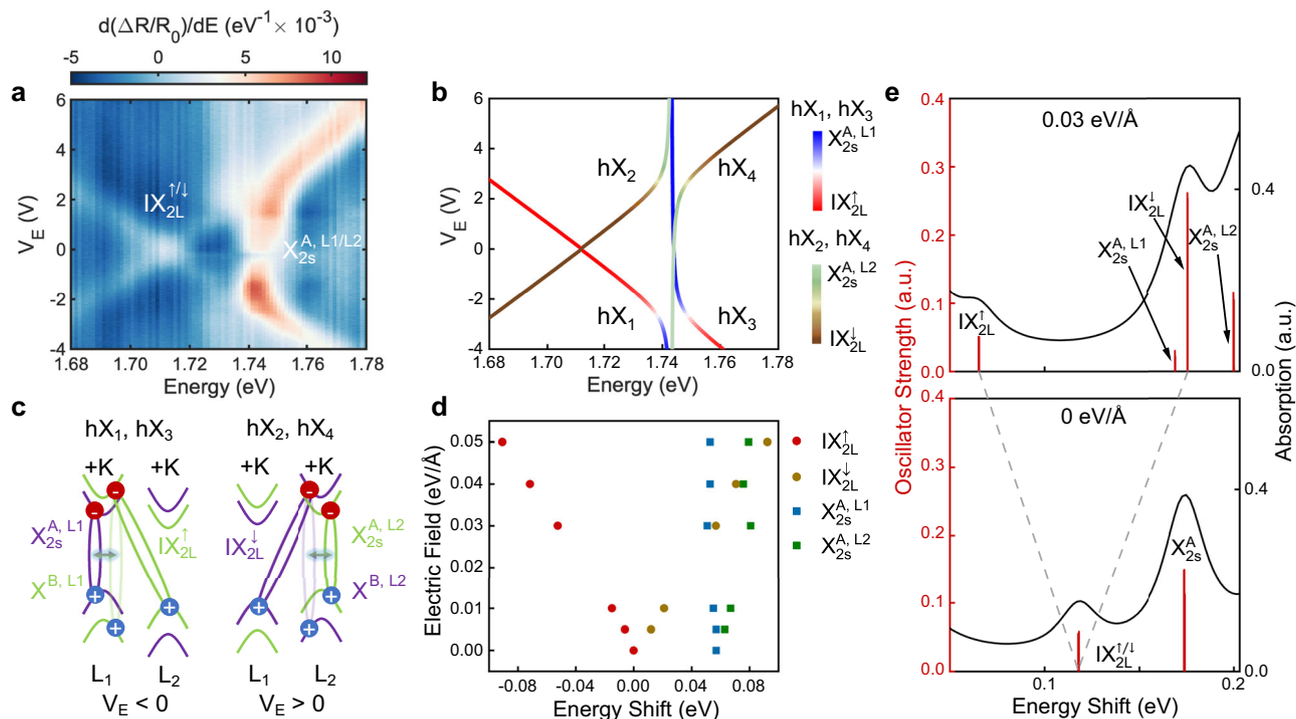


Fig. 2 | Layer hybridised excitons in 2L 2H-MoSe₂. **a** V_E dependence of the first derivative of the reflectance contrast spectra with respect to photon energy ($d(\Delta R/R_0)/dE$) in our 2L 2H-MoSe₂ in the spectral range 1.68–1.78 eV. **b** Calculated energies of the different hybrid $IX_{2L}^i-X_{2s}^A$ exciton states as a function of V_E , which we label as hX_1 , hX_2 , hX_3 , and hX_4 from low to high energy at $V_E > 0$ V, respectively. The colour of the solid lines denotes the contribution of the different bare exciton states to each hybrid exciton. **c** Schematics of the spin, valley, and layer configuration of the exciton states responsible for the exciton hybridisation shown in panel a for negative and positive applied V_E (left and right panels, respectively). The

exciton hybridisation is attributed to a second-order effective coupling between IX_{2L} and the intralayer A exciton facilitated via the A and B exciton admixture (depicted by the glowing double arrows). **d** The energy position of IX_{2L} and X_{2s}^A as a function of the applied electric field, as obtained from GW + BSE calculations. The labels identify the simplified bare exciton states. **e** Normalised theoretical oscillator strengths (red vertical line) and absorption spectra (black line) at 0 eV/Å and 0.03 eV/Å focusing on the energy range of IX_{2L} and X_{2s}^A . The numerical precision of our calculations is estimated to be of the order of ± 5 meV, see computational details in Supplementary Note S2.

stacking of our terraced 2L/3L MoSe₂ flake results in a different relative stacking configuration between the different layers. While IX_{2L} involves MoSe₂ layers with a relative 2H stacking, IX_{3L} originates from MoSe₂ layers with a relative R_h^i -type stacking, where h denotes the hexagon centre of the crystal lattice in each layer (see Fig. 1a). Hence, each IX species is endowed with distinct spin-layer-valley configurations (see Fig. 1b) that can be optically probed. Finally, we note that the relative R_h^i -stacking between L_1 and L_3 is distinct from 3R-stacked homobilayers^{27,44}. For 3R-type bilayers the two layers are laterally shifted so that the hexagon centres in each layer are not vertically aligned, rendering a different symmetry and resulting in a forbidden hole tunnelling⁴⁵.

Electric field-dependent excitonic transitions in 2L 2H-MoSe₂

To investigate the layer-dependent and intra-/interlayer nature of the different exciton species in the terraced 2H-MoSe₂ flake, we perform differential reflection contrast ($\Delta R/R_0$) spectroscopy at cryogenic temperature (4 K) as a function of V_E at charge neutrality, where $\Delta R = R_s - R_0$, and R_s (R_0) is the intensity of the light reflected by the flake (substrate). We focus first on the 2L region of our MoSe₂ sample. Supplementary Fig. S1 shows a representative reflection contrast spectrum of our 2L 2H-MoSe₂ at $V_E = 0$ V, where several excitonic resonances with different linewidths and absorption strengths are observed (see Supplementary Fig. S2 for the extracted exciton linewidths). The strongest exciton resonances at low (-1.63 eV) and high energy (-1.86 eV) correspond to the ground states of intralayer A (X_{1s}^A) and B (X_{1s}^B) neutral excitons in 2L MoSe₂, respectively. Two additional weaker exciton resonances, separated only by ~25 meV, are also observed in the energy range between the A and B intralayer 1s exciton

states, which can be attributed to IX_{2L} (-1.715 eV) and the first excited exciton state X_{2s}^A of the neutral A exciton (-1.74 eV), in agreement with previous experimental and theoretical results on bulk²⁰ and 2L 2H-MoSe₂²⁴. More precisely, we perform GW + BSE calculations (see Supplementary Note 2 for computational details) and obtain IX_{2L} and X_{2s}^A peaks 0.11 and 0.17 eV above the X_{1s}^A one, respectively (see Supplementary Fig. S3), which qualitatively agrees with our experiment in which IX_{2L} and X_{2s}^A are 0.07 and 0.11 eV above X_{1s}^A . To corroborate the intra-/interlayer character of the IX_{2L} and X_{2s}^A states and unravel their layer-dependent properties, we show in Fig. 2a a density plot of the V_E -dependence of the first derivative of the reflectance contrast spectra with respect to photon energy ($d(\Delta R/R_0)/dE$) in the spectral range 1.68–1.78 eV, which helps to visualise these exciton transitions (see Supplementary Fig. S5 for comparison with bare $\Delta R/R_0$ spectra). At $V_E = 0$ V, we observe the two excitonic resonances attributed to IX_{2L} and X_{2s}^A . The application of a V_E lead to a stark contrast in the behaviour of the two resonance peaks. For small positive applied V_E , the resonance energy of the high energy peak remains almost constant, while the low energy peak splits into two exciton branches which shift symmetrically towards lower and higher energies with a linear dependence with V_E . This phenomenon can be interpreted in terms of the DC Stark energy (ΔU) tuning experienced by interlayer excitons under applied out-of-plane electric fields: $\Delta U = -pE$, with E the strength of the vertical electric field and $p = ed$ the out-of-plane electric dipole moment (where e represents the electron charge and d the electron-hole distance). The absolute values of the energy shifts of the two IX_{2L} branches allow us to estimate an average electron-hole spatial separation of $d = 0.34 \pm 0.01$ nm for IX_{2L} , which is comparable to the reported values of 0.39–0.47 nm for IXs in other TMD homostructures

such as 2L MoS₂^{22,23}, and 0.63 nm (0.26 nm) for momentum direct (indirect) IXs in twisted 2L MoSe₂²⁸. This result unambiguously demonstrates the presence of a sizeable static electric dipole in the out-of-plane direction and corroborates the interlayer exciton nature of IX_{2L}^{20,24}. Moreover, the symmetrical but opposite energy shifts of the two IX_{2L} branches reveal the presence of IX_{2L} excitons with different polarities (i.e. with static electric dipoles aligned parallel and anti-parallel to the applied electric field), as sketched in Fig. 1a. For positive V_E , IX_{2L} excitons shifting to lower (higher) energies originate from Coulomb-bound electron-hole pairs in which the hole is spatially located in the top (bottom) layer of the 2L MoSe₂, i.e. with static electric dipoles pointing up (IX_{2L}[↑]) and down (IX_{2L}[↓]), respectively. Note that for negative V_E , the behaviour of IX_{2L}^{↑(↓)} is reversed. Therefore, the Stark shifts of IX_{2L}^{↑(↓)} allow us to unravel their layer configuration.

Further, the large DC Stark tuning of IX_{2L} allows us to explore the possible hybridisation between IX_{2L} and the energetically close X₂₅^A by reducing their energy detuning via the applied V_E . For $|V_E| \approx 2$ V, IX_{2L} and X₂₅^A show an energy anti-crossing characteristic of coupled systems, suggesting the hybridisation of the two exciton species. We note that this observation is reproduced across the 2L 2H-MoSe₂ sample. Supplementary Fig. S6 shows the results for a different spatial location in the 2L 2H-MoSe₂.

In order to estimate the magnitude of the coupling between IX_{2L} and X₂₅^A, we employ a phenomenological model in which the hybridisation between the exciton states is treated as a coupling between oscillators with resonance energies corresponding to the bare exciton states (see Supplementary Section SI for details). In our model, we take into account the spin, valley, and layer degrees of freedom of each exciton species, which leads to eight exciton resonances with different spin, valley, and layer properties: two IX_{2L} with opposite polarities (IX_{2L}^{↑(↓)}) and momentum-direct transitions at $\pm K$ each, and X₂₅^A localised in the top (X₂₅^{A,L1}) and bottom (X₂₅^{A,L2}) MoSe₂ layers with momentum-direct optical transitions at $\pm K$ each. Regarding the IX_{2L}-X₂₅^A hybridisation, we include it in our model as a phenomenological interlayer coupling of holes at $\pm K$, with a magnitude which we assume to be independent of V_E . In our calculations, the energies of the exciton states at $V_E = 0$ V and the slope of the DC Stark shift for IX_{2L}^{↑(↓)} are set to match the corresponding experimental values, while the value of the coupling strength between the exciton states is left as a free parameter that can be tuned to fit our experimental data. Figure 2b shows the calculated energies of the resulting hybrid exciton states as a function of V_E , which we label as hX₁, hX₂, hX₃, and hX₄ from low to high energy at $V_E > 0$ V, respectively. The colour of the solid lines in Fig. 2b denotes the contribution of the different bare exciton states to each hybrid exciton. The phenomenological model captures well the hybridisation-induced renormalisation of the exciton resonance energies with increasing electric field, allowing us to estimate an IX_{2L}-X₂₅^A coupling strength $\kappa_{2L-25} \approx 5.2$ meV, which is slightly smaller but of the same order of magnitude as the linewidths of the exciton states at $V_E = 0$ V (see Supplementary Fig. S2).

The physical origin of the observed hybridisation between an interlayer exciton and the first excited state of an intralayer exciton is intriguing, and to the best of our knowledge has not been reported in any other homobilayer TMD system. We discard the possibility of spin-conserving electron hopping between the electron states involved in IX_{2L} and X₂₅^A, since such interlayer electron hopping is forbidden in 2L 2H-MoSe₂ due to the C₃ symmetry of the d_{z^2} orbitals of the conduction band states at $\pm K$ ^{45,46}. However, recent theoretical work has shown that the application of a vertical electric field can lead to the hybridisation of IX_{2L} and X₁₅^B in 2L 2H-MoSe₂ via spin-conserving interlayer hole tunnelling⁴⁷, in agreement with previous experimental and theoretical results for 2L 2H-MoSe₂^{23,48}. Moreover, similar to 2L 2H-MoSe₂⁴⁹, the theoretical results in Ref. 47 also suggest a weak admixture between the A and B excitons in 2L 2H-MoSe₂, which leads to a non-vanishing second-order effective hybridisation of IX_{2L} and the intralayer X₁₅^A

exciton. Since X₁₅^A and X₂₅^A have the same nature (i.e. same exact spin and valley configuration), we tentatively attribute the observed IX_{2L}-X₂₅^A hybridisation to a second-order effective coupling between IX_{2L} and the intralayer A exciton facilitated by through intravalley exchange interaction between the A (spin up) and B (spin down) excitons (see Fig. 2c)⁴⁷⁻⁴⁹. This result agrees well conceptually with the theoretical modelling in ref. 47, and leads to a layer selective coupling between IX_{2L} and the intralayer A exciton, as also captured by our phenomenological model (see Fig. 2b). Our estimated value of the coupling strength $\kappa_{2L-25} \approx 5.2$ meV is very similar to the coupling strength reported for IX_{2L} and X₁₅^A excitons in 2H-MoSe₂, and significantly smaller than the direct coupling between the IX_{2L} and X₁₅^B excitons in the same 2H-MoSe₂^{23,48} and 2H-MoSe₂ sample⁵⁰, which supports our hypothesis.

To further corroborate our hypothesis, we calculate GW + BSE-based estimates of the excitonic transition energies and their corresponding oscillator strengths at various external electric field values (see Fig. 2d, e). For vertical applied positive electric fields, the IX_{2L} excitons experience a significant Stark shift, which supports their interlayer nature. The X₂₅^{A,L1} remains relatively unchanged, while the X₂₅^{A,L2} exhibits an energy shift with the applied electric field, indicative of the layer-selective mixing with the IX_{2L} state. In fact, our calculations reveal a sizeable mixing between IX_{2L} and X₂₅^{A,L2} even at zero applied electric field. Table 1 in Supplementary Section S2 summarises the oscillator strengths of the different bare optical transitions contributing to each excitonic resonance in Fig. 2d, e, corresponding to 0 V/Å and 0.03 V/Å applied electric fields, respectively. The conversion from V_E to electric field is shown in Fig. S4.

Electric field-dependent excitonic transitions in 3L 2H-MoSe₂

To explore the potential for multilayer TMDs to host IX with dipole moments even larger than IX_{2L} and with greater tunability, we optically probe the 3L region of our terraced 2H-MoSe₂ sample at charge neutrality as a function of V_E . Figure 3a shows the V_E -dependence of $d(\Delta R/R_0)/dE$ in the spectral range 1.58–1.78 eV (with the values in the energy range 1.66–1.78 eV multiplied by a factor 4 for visualisation purposes). The reflectance spectrum at $V_E = 0$ V is markedly different to the one observed in the 2L region (see Fig. 2a). We observe two exciton transitions in the energy range corresponding to X₁₅^A with an energy splitting of -11 meV, and two exciton resonances in the energy range of IX_{2L} with an energy splitting of -14.5 meV. The resonances at low energy can be attributed to X₁₅^A excitons localised in the different layers of our sample, in which the lower average permittivity environment of L₁ and L₃ (h-BN/L₁₍₃₎/MoSe₂) compared to L₂ (MoSe₂/L₂/MoSe₂) results in a dielectric-induced energy blue shift for X₁₅^A excitons in L₁ and L₃ (X₁₅^{A,L1(L3)}) compared to X₁₅^A excitons in L₂ (X₁₅^{A,L2})⁵¹, similar to what has been observed for 3L MoS₂²³.

To unravel the nature of the excitons in the energy range corresponding to IX_{2L}, we focus on their behaviour as a function of V_E . As shown in Fig. 3a, the application of a vertical electric field gives rise to a Stark-effect-induced splitting of each resonance into two distinct exciton branches which shift symmetrically towards lower and higher energies with a linear dependence with V_E (see the comparison between $\Delta R/R_0$ and $d(\Delta R/R_0)/dE$ in Supplementary Fig. S7). For the lower exciton resonance, we estimate an average electron-hole spatial separation of $d = 0.36 \pm 0.01$ nm, which agrees very well with the value found for IX in the 2L region of our sample, confirming its IX_{2L} nature. The energy splitting of the two branches belonging to the higher energy resonance yields an average electron-hole separation of $d = 0.73 \pm 0.01$ nm, which exceeds the interlayer distance of TMD homobilayers⁵² and is approximately twice the IX_{2L} dipole size. We therefore identify this resonance as IX_{3L}. Note that the deviation between the effective dipole moment and physical interlayer distance (-0.6 nm and 1.4 nm for 2L and 3L, respectively) further indicates that these hybrid interlayer excitons are not pure 'bare excitons' as in the

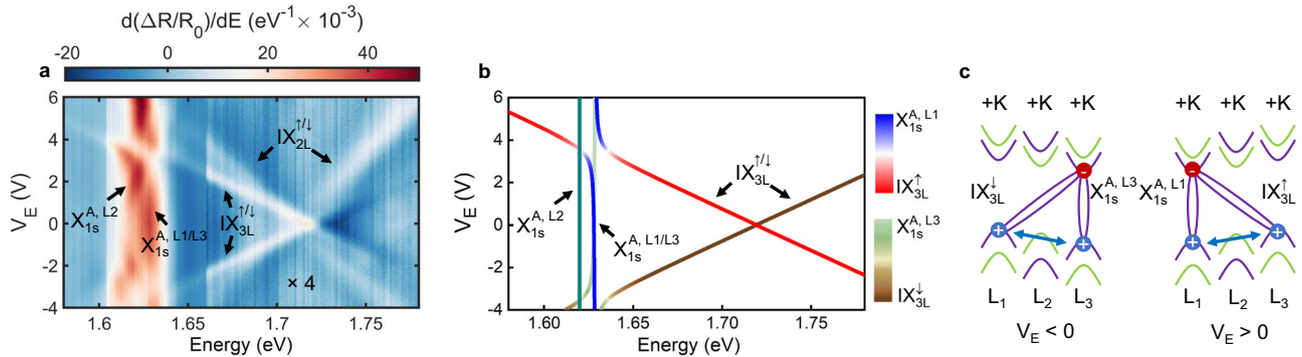


Fig. 3 | Layer hybridised excitons in 3L MoSe₂. **a** V_E dependence of $d(\Delta R/R_0)/dE$ in the 3L MoSe₂ region of our sample. **b** Energies of the different hybrid $IX_{3L}^A - X_{1s}^A$ exciton states as a function of V_E , where the colour of the solid lines denotes the contribution of the different bare exciton states to each hybrid exciton.

c Schematics of the spin, valley, and layer configuration of the exciton states responsible for the exciton hybridisation shown in panel (a) for negative and positive applied V_E (left and right panels, respectively). The exciton hybridisation is attributed to direct spin-conserving interlayer hole tunnelling between L_1 and L_3 .

simple cartoon picture, rather, they are composed of a mixture of intra/interlayer exciton wave functions. The wavefunction admixture leads to intra/interlayer contributions to the effective dipole moments of the hybrid IX_{2L} and IX_{3L} . Using IX_{2L} as an example, it consists of the bare intralayer exciton with negligible vertical dipole moment and the bare interlayer exciton with dipole moment -0.62 nm. In real space, this is in line with a qualitative description that the electron of the IX is relatively layer localised while the holes are relatively delocalised in both layers³⁷, which yields an effective dipole size that is roughly 50% of the physical interlayer distance. The DFT calculations also suggest similar exciton admixtures at high field i.e. 52% of intralayer X_{2s} and 38% IX_{2L} (Table I in Supplementary Information, $X_{2s}^{A,L2}$ list in lower panel), qualitatively supporting the experimental results. As a result of the complex wave function mixing, the hybrid K-valley IX may have material-dependent effective dipole moments. In addition, the choice of the dielectric constants differs in hybrid IX studies⁵³, which also introduces discrepancies in the extracted dipole moments.

Furthermore, the larger Stark shift of IX_{3L} allows tuning its energy into resonance with X_{1s}^A (see Fig. 3a). For $|V_E| = 4$ V we observe a clear avoided crossing between IX_{3L} and $X_{1s}^{A,L1/L3}$, indicative of the hybridisation between the exciton species with carriers hosted in the outermost layers of the 3L MoSe₂ sample. Supplementary Fig. S8 shows the results for a different spatial location in the 3L 2H-MoSe₂. We note that contrary to the results for the 2L region, we observe negligible oscillator strength of X_{2s}^A in the 3L MoSe₂ region, which prevents clear observation of coupling between this exciton state and the two IX species. Therefore, we focus on the clear $IX_{3L} - X_{1s}^A$ hybridisation.

Similar to the 2L case, we simulate the V_E -dependent energy dispersion of the hybridised excitons in the 3L MoSe₂ system using a phenomenological model of coupled oscillators, in which the exciton coupling is both spin- and layer-selective (see Fig. 3b). In this case, we include three different X_{1s}^A excitons (i.e. one in each layer) and two IX_{3L} with opposite polarities ($IX_{3L}^{\uparrow(\downarrow)}$) and momentum-direct transitions at $\pm K$ each. Spin-conserving hole tunnelling results in a layer-selective coupling between IX_{3L} and X_{1s}^A ; the polarity of IX_{3L} is locked to the layer degree of freedom of X_{1s}^A (e.g. $IX_{3L}^{\uparrow(\downarrow)}$ only couples to $X_{1s}^{A,L1(L3)}$). However, the energy degeneracy of $X_{1s}^{A,L1}$ and $X_{1s}^{A,L3}$ prevents clear observation of such layer-selective coupling experimentally. Figure 3b shows the results of the best fit of the model to our experimental data. The colour of the solid lines denotes the contribution of the different bare exciton states to each hybrid exciton. Overall, the phenomenological model captures well the hybridisation-induced renormalisation of the exciton resonance energies with increasing electric field, allowing us to estimate an $IX_{3L} - X_{1s}^A$ coupling strength $\kappa_{3L-1s} \approx 7.5$ meV, which is slightly larger but of the same order of magnitude as κ_{2L-2s} in the 2L region. Furthermore, it is worth noting that the R-type relative stacking of the

layers L_1 and L_3 leads to some differences between the $IX_{3L} - X_{1s}^A$ and $IX_{2L} - X_{2s}^A$ couplings. The R-type stacking between L_1 and L_3 results in IX_{3L} with electron-hole pairs with the same spin-valley configurations as X^A (see Fig. 1a), which allows the hybridisation of the two exciton species via direct spin-conserving interlayer hole tunnelling in the presence of an additional layer between L_1 and L_3 ^{29,54}. For the specific stacking registry of R_r^h between L_1 and L_3 , a recent work calculated an 11 meV tunnel splitting of the valence band (with monolayer BN as L_2)²⁹, supporting our interpretation. On the contrary, due to the 2H relative stacking between adjacent layers, the direct hole tunnelling between the valence band states in IX_{2L} and X^A has to compete with a sizable detuning equal to the spin splitting at the valence band edges (150 meV in MoSe₂⁵⁴), and is thus only facilitated by the admixture of X^A and X^B . Finally, we note that the energy detuning between IX_{2L} and the energy degenerate $X_{1s}^{A,L1}$ and $X_{1s}^{A,L3}$ exciton resonances is slightly smaller in the 3L region as compared to the 2L one. As a consequence, we are able to observe experimental signatures of the coupling between these two exciton species at $V_E = 6$ V (see Fig. 3a and Supplementary Information S4), which corroborate our results for the 2L 2H-MoSe₂. We summarise the rules for determining which excitons can hybridise based on our new insight and other references (see Supplementary Information S5).

Electric-field-dependent magneto-optical properties of hybrid excitons in MoSe₂

Beyond tuning of the exciton resonance energies, oscillator strengths, and effective permanent electric dipoles, here we explore if the electric-field-dependent control of the exciton nature enables precision tuning of the effective Landé g -factors. The application of a vertical magnetic field B results in the Zeeman splitting of the optical transitions of each exciton at $\pm K$, with an energy splitting $\Delta E(B) = g\mu_0 B$, with μ_0 the Bohr magneton. Optical transitions at $\pm K$ can be independently probed by σ^\pm -polarised light, respectively, which enables us to perform circularly polarised reflectance contrast measurements to estimate the experimental Zeeman splitting $\Delta E = E^{\sigma^+} - E^{\sigma^-}$, with E^{σ^\pm} the energy of the σ^\pm -polarised transition. We focus first on the hybrid $IX_{2L} - X_{2s}^A$ exciton states in our 2L region. Figure 4a shows the measured Zeeman splittings (blue dots) for hX_4 at three different applied V_E values. Supplementary Fig. S9 shows the linecuts for the σ^\pm -resolved $d(\Delta R/R_0)/dE$ at $V_E = 0$ V for 5 T. The blue solid lines represent linear fits of the experimental data, from which we estimate the effective g -factor at each applied V_E . We observe the effective g -factor of hX_4 is tuned from a negative value (-1.9 ± 0.5) to a relatively large positive value (11.8 ± 0.4). To explore this effect in more detail, we employ the same experimental procedure and extract the effective g -factors of hX_3 and hX_4 in the range of applied V_E in which the oscillator strength of each

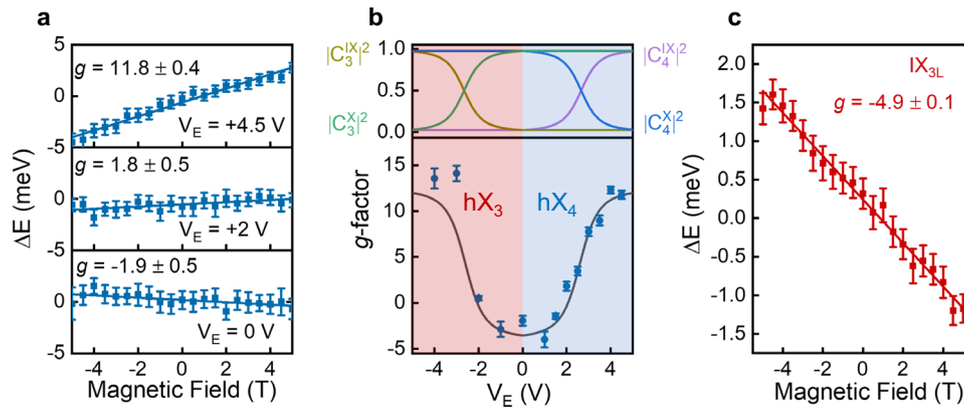


Fig. 4 | Magneto-optical properties of layer-hybridised excitons in 2L MoSe₂ and 3L MoSe₂. **a** Zeeman splitting of hX₄ at three different applied V_E . The blue dots represent the experimental values, while the blue solid lines show linear fits of the experimental data, from which we are able to estimate the effective g -factor of this hybrid exciton at each applied V_E . **b** V_E -driven evolution of the g -factor of the hybrid

excitons hX₃ (−5 V to 0 V, red shaded area) and hX₄ (0 V to 5 V, blue shaded area) in bilayer MoSe₂ (bottom panel). The top panel shows the V_E -dependent contributions of each bare exciton state $|C_{3,4}^{IX(X)}|^2$ to the corresponding hybrid excitons. **c** Zeeman splitting of IX_{3L} at measured at $V_E = 0$ V. Error bars are from uncertainties in Lorentz peak fit and linear fit, respectively.

transition and their energy detuning with respect to other transitions enable a reliable estimate of $E^{\sigma\pm}$, i.e. $V_E < 0$ for hX₃ and $V_E > 0$ for hX₄ (as indicated by the red and blue shaded areas in Fig. 4b). The results highlight a continuous and smooth transition of the g -factor of the two hybrid exciton states from -1.9 ± 0.5 to 11.8 ± 0.4 by changing V_E from 0 V to ± 4.5 V. Such evolution of the effective g -factor arises from the V_E -dependent hybridisation between IX_{2L} and X_{2s}^A, and can be quantitatively explained by our phenomenological model of coupled oscillators. In this model, the wave function of each hybrid exciton is expressed as a superposition of the bare intralayer and interlayer exciton wave functions (see Supplementary Section S1). In the case of hX₃ and hX₄, these exciton wave functions are expressed as follows:

$$|hX_3(V_E)\rangle = C_3^{IX}(V_E)|IX_{2L}^\uparrow\rangle + C_3^X(V_E)|X_{2s}^{A,L_1}\rangle \quad (1)$$

and

$$|hX_4(V_E)\rangle = C_4^{IX(X)}(V_E)|IX_{2L}^\downarrow\rangle + C_4^X(V_E)|X_{2s}^{A,L_2}\rangle, \quad (2)$$

where $C_i^{IX(X)}(V_E)$ represents the V_E -dependent amplitude of the bare interlayer (intralayer) exciton state in hybrid exciton state $|hX_i\rangle$, with $|C_i^{IX}(V_E)|^2 + |C_i^X(V_E)|^2 = 1$. Therefore, the effective g -factor of each hybrid exciton state (g_{hX_i}) can be expressed as

$$g_{hX_i}(V_E) = |C_i^{IX}(V_E)|^2 g_{IX_{2L}} + |C_i^X(V_E)|^2 g_{X_{2s}}, \quad (3)$$

where $g_{IX_{2L}(X_{2s})}$ represents the g -factor of the bare IX_{2L} (X_{2s}) state. The solid line in Fig. 4b represents a fit of the experimental data to Eq. (3), in which we have used the $C_i^{IX(X)}(V_E)$ values shown in the top panel of Fig. 4b (obtained from the fits in Fig. 2b), and $g_{IX_{2L}}$ and $g_{X_{2s}}$ have been left as free fitting parameters. Our model captures well the hybridisation-induced evolution of the g -factors and allows us to estimate of the effective g -factors of the bare states: $g_{X_{2s}} \approx -4$ and $g_{IX_{2L}} \approx 12.5$. We note that, although to the best of our knowledge $g_{X_{2s}}$ has not been previously reported for 2L MoSe₂, our estimated value of $g_{X_{2s}}$ is in very good agreement with reported experimental (-3.6 ± 0.1)⁵⁵ and theoretical (-3.7)⁵⁶ values for this excited exciton state in monolayer MoSe₂ and other TMD systems such as bulk WSe₂ (-3.3 ± 0.6)²⁰. Regarding $g_{IX_{2L}}$, the estimated sign and value are also in good agreement with the value predicted by a simplistic “atomic picture”, in which the g -factor of the bands hosting the electron-hole pairs are assumed to be equal to the sum of their spin, orbital, and valley magnetic moments⁵⁷. Within this model, the spin-valley configuration of IX_{2L} results in a $g_{IX_{2L}} = 2(\frac{m_0}{m_c} + \frac{m_0}{m_v}) + 4 \approx 9.7$, with m_0

the free electron mass, and m_c^* and m_v^* the experimentally reported electron and hole effective masses for the bottom conduction band and top valence band in monolayer MoSe₂, respectively⁵⁸. The relatively large and positive value of the g -factor estimated with this atomic picture model provides additional confirmation of the spin-valley configuration and interlayer nature of IX_{2L}. Also, we note that the small discrepancy between the experimental and the calculated value for $g_{IX_{2L}}$ might arise from a combination of several factors, including the limitation of this simple model to estimate accurately the g -factor of the relevant bands⁵⁹ and the absence of experimental values for the electron effective mass of the top conduction band in 2L MoSe₂, which theoretical calculations predict to be slightly smaller than for the bottom conduction band in Mo-based TMDs such as 2H-MoSe₂⁶⁰ and 2H-MoS₂⁶¹. Nevertheless, we note that the extracted g -factors of the bare IX_{2L} and X_{2s} states allow us to include the effects of the Zeeman splitting in the calculated V_E -dependent evolution of the hybrid IX_{2L}-X_{2s} states, for which we find a very good agreement with the experimental results (see Supplementary Note S3 and Supplementary Fig. S14).

Finally, we investigate the magneto-optical properties of IX_{3L} in the 3L region of the sample. Figure 4c shows the experimental Zeeman splitting for this exciton species at $V_E = 0$ V, from which we estimate a g -factor $g_{IX_{3L}} = -4.9 \pm 0.1$. The extracted value for $g_{IX_{3L}}$ is very similar to the experimentally reported g -factor of X_{1s}^A in 3L MoSe₂⁵⁹, which confirms the identical spin-valley configuration of these two exciton species and corroborates that the giant dipole IX_{3L} indeed originates from electron and hole in L₁ and L₃ (or vice versa), respectively. The identical spin-valley configuration of X_{1s}^A and IX_{3L} has an important consequence on the magneto-optical properties of the hybrid X_{1s}^A-IX_{3L} states: contrary to the hybrid X_{2s}^A-IX_{2L} states in 2L MoSe₂, hybrid X_{1s}^A-IX_{3L} do not feature a V_E -tunable g -factor (see $V_E = 3$ V Zeeman splitting in Fig. S10), since the two bare exciton states already exhibit a similar g -factor.

Excited state interlayer excitons in multilayer 2H-MoSe₂

Although excited (Rydberg) exciton states have weaker oscillator strength than their corresponding ground states¹⁵, in this section, we show that the hybridisation of interlayer excitons with intralayer transitions leads to clear spectroscopic signatures of excited Rydberg states for both IX_{2L} and IX_{3L}. Figure 5a,b show $d^2(\Delta R/R_0)/dE^2$ spectra as a function of V_E for 2L and 3L 2H-MoSe₂ regions, respectively, corresponding to spatial locations where the interlayer excited exciton states present appreciable oscillator strengths. The data shown in the spectral ranges 1.797–1.830 eV and 1.665–1.800 eV in Fig. 5a, b have

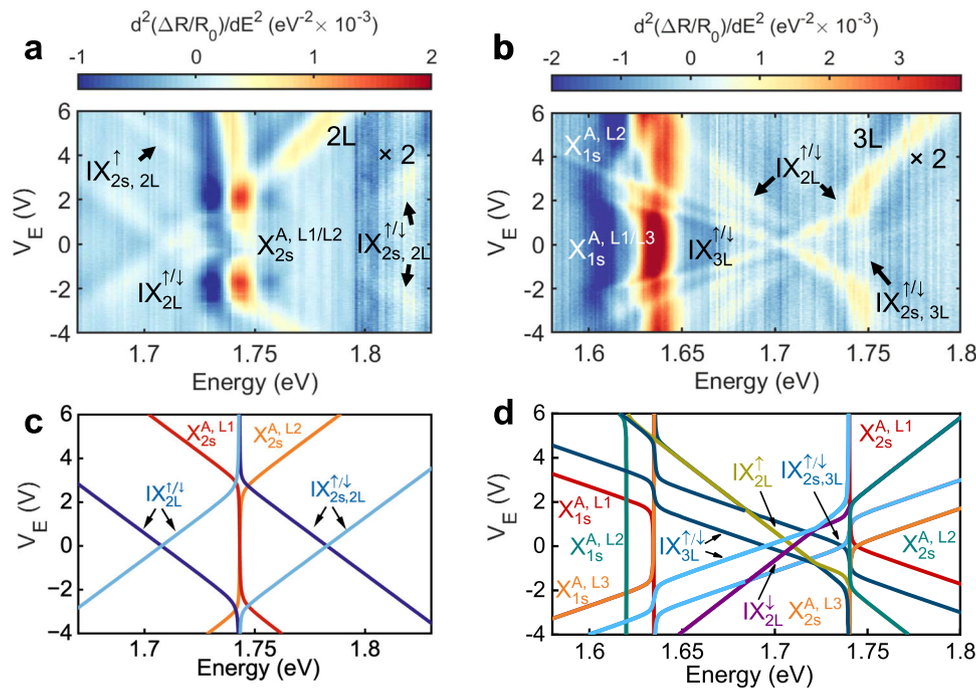


Fig. 5 | Observation of excited states of IX_{2L} and IX_{3L} . **a** V_E dependence of $d^2(\Delta R/R_0)/dE^2$ in a second location of the 2L 2H-MoSe₂ sample in the spectral range 1.67–1.83 eV. **b** $d^2(\Delta R/R_0)/dE^2$ in another location of the 3L 2H-MoSe₂ sample in the spectral range 1.58–1.8 eV. **c** Calculated energies of the different exciton states of

2L 2H-MoSe₂ including $IX_{2s,2L}$ with the same spin and layer configuration as IX_{2L} . **d** Calculated energies of the different exciton states in 3L 2H-MoSe₂ including $IX_{2s,3L}$ with the same spin and layer configuration as IX_{3L} .

been multiplied by a factor 2 for visualisation purposes. In the 2L region (Fig. 5a), for $4 \lesssim V_E \lesssim 6$ V we resolve an additional exciton transition in the energy range ~ 1.7 eV with a linear Stark shift parallel to the IX_{2L}^\uparrow . Extrapolation of the linear energy shift to $V_E = 0$ V gives an estimated energy of ~ 1.77 eV for this excitonic transition with clear interlayer nature. We note that the relative energy position of this excitonic peak with respect to both IX_{2L} and X_{2s}^A agrees well with the first excited Rydberg state of IX_{2L} predicted by our GW + BSE results (see Fig. 2e and Supplementary Fig. S3). This, together with the dipole moment of 0.30 ± 0.01 nm estimated from the linear Stark shift (almost identical to the one for IX_{2L}), allows us to attribute the observed resonance to the first excited Rydberg state of IX_{2L}^\uparrow (i.e. $IX_{2s,2L}^\uparrow$). Notably, for $|V_E| \sim 3$ V and energies ~ 1.82 eV, we observe two additional resonances with opposite Stark shift slopes that also extrapolate to an energy of ~ 1.77 eV at $V_E = 0$ V, which we attribute to $IX_{2s,2L}^\downarrow$ ($V_E > 0$ V) and $IX_{2s,2L}^\uparrow$ ($V_E < 0$ V), corroborating our peak assignment. We note that the $IX_{2s,2L}$ state is a general feature rather than a location-dependent feature of our sample, as we also observe $IX_{2s,2L}$ in the main locations of 2L and 3L MoSe₂ shown in Figs. 2–4, as depicted in Supplementary Fig. S11. Further, we observe a very weak transition near 1.72 eV for $4 \lesssim V_E \lesssim 6$ V with a Stark shift parallel to $IX_{2s,2L}$, which we tentatively ascribe as the 3s Rydberg state of IX_{2L} . Extrapolation to $V_E = 0$ V gives an estimated energy for the 3s state of ~ 20 meV higher than the 2s at zero applied electric field (Supplementary Fig. S12).

Similar to Fig. 5a, the results in Fig. 5b also display clear excited Rydberg states in the 3L 2H-MoSe₂ sample. In addition to the coexisting interlayer exciton species IX_{2L} and IX_{3L} already shown in Fig. 3, we observe two additional transitions with Stark-induced linear energy shifts parallel to the IX_{3L} , which we attribute to the two opposite polarities of the 2s excited Rydberg state of IX_{3L} (i.e. $IX_{2s,3L}^\uparrow$). Extrapolation of the measured linear energy shifts of $IX_{2s,3L}^\uparrow$ to $V_E = 0$ V gives an estimated energy ~ 40 meV above the IX_{3L} ground state (near the X_{2s}^A transition).

We simulate the V_E -dependent energy dispersions of the hybridised ground and excited exciton Rydberg states using the phenomenological coupled oscillator model previously described. We build on

the models used for Figs. 2b and 3b and add the the observed excited Rydberg states, which we assume to have identical Stark shifts and the same spin- and layer-selective couplings as their corresponding ground states. Figure 5c, d shows the simulated energy dispersions corresponding to the results in Fig. 5a, b, respectively. As can be observed in these figures, the simulated energy dispersions capture well the V_E -induced hybridisation and energy dispersion of the different excitonic transitions (see Supplementary Information Fig. S17 for the simulated absorption spectrum as a function of V_E corresponding to Fig. 5b using this model and Supplementary Information section S1 for simulation details), supporting our interpretation of their different intra-/interlayer origin, ground/excited state character, and spin and valley configurations. On top of the electric field sweeps, we performed additional characterisations on the excited state IXs, which reveal the doping-dependent energy dispersion and magnetic field-dependent diamagnetic shift of $IX_{2s,2L}$ and higher order excited state of IX_{3L} (Supplementary Information S7 and 8).

Finally, we note that the relative energy order of IX_{2L} and IX_{3L} at 0 V varies across the sample. Here we summarise our experimental findings and provide a qualitative explanation. In the primary 3L location (Fig. 3), the IX_{2L} (lower energy) and IX_{3L} (higher energy) are separated by 12.6 meV; in the secondary 3L location (Fig. 5), the IX_{2L} and IX_{3L} energetic ordering is reversed with a 15 meV difference (opposite sign to the primary location); in a tertiary location (Fig. S8b), we observe similar trend as the first location, where IX_{2L} can be extrapolated to be around 17 meV lower in energy than IX_{3L} at 0 V. The energy order is interpreted as a combined effect of interband transition energies (determined by the band gap and binding energy), dielectric screening, local strain, and possible exciton state mixing (see Supplementary Information S6 for a more detailed discussion)

Discussion

Our work reports the observation of giant Stark splitting of the interlayer excitons in 2L and 3L 2H-MoSe₂. First, we observe hybridisation between IX_{2L} and X_{2s}^A . In addition to their spectral evolution, a

hybridisation-driven g-factor evolution of the coupled excitons is resolved and understood via a unified coupled oscillator model. The ability to drive the exciton Zeeman splitting from negative to positive through zero has potential in electrically tunable valleytronics and spin-dependent exciton-exciton interactions. Next, we report the giant excitonic trilayer dipole IX_{3L} , which exhibits distinct spin-layer selection rules for hybridisation with X_{15}^A . A salient feature of the large IX_{3L} dipole is that it can be Stark tuned to become the ground state (e.g. lower energy than X_{15}^A), promising for applications in exciton transport. Finally, by harnessing the exciton hybridisation effects, we successfully probe the excited state IX for the first time in a TMD system. Future theory and experimental effort is encouraged to better understand the excited IX states, including comparisons with the hydrogen model and the magnetic field and doping dependence to reveal the g-factor, effective mass⁵⁸ and Roton-like⁶² properties of the interlayer Rydberg excitons. In future work, an external cavity could be used to enhance the light-matter interaction strength⁶³ and take advantage of the large energy tunability. The excited state IX with large spatial extension, as well as dipolar nature, may further facilitate polariton blockade in real space that may even lead to quantum nonlinearity^{64,65}. Altogether, these results promote a new TMD homostructure candidate for applications with enhanced exciton-exciton interactions with strong light-matter coupling. Beyond 3L 2H-MoSe₂, a strategy of further engineering IX dipoles by tuning the layer number, including thicker multilayer (>3L) 2H-TMDs or heterostructures with multilayer TMD components and hBN spacers, is encouraged.

Following the submission of our manuscript, a related work on interlayer excitons (similar transition as IX_{3L} in our work) in multilayer 2H-WSe₂ has been published⁵³.

Methods

Sample fabrication

Bulk MoSe₂ crystal was exfoliated onto polydimethylsiloxane (PDMS) stamps and a flake with a terraced 2L and 3L region was identified using optical contrast. Few-layer graphene and hBN layers were also prepared and identified on PDMS. The flakes were then stacked sequentially onto pre-patterned Au electrodes on SiO₂/Si substrates using the all-dry viscoelastic transfer technique in an Ar-filled glove box⁶⁶.

Optical measurements

The sample was held in a closed-cycle cryostat (Attodry 1000) at 4 K, where a magnetic field can be applied out-of-plane to the sample (Faraday configuration). For the reflectance measurements, the broadband spectrum from a power-stabilised tungsten lamp was collected by a multimode fibre. The light was collimated by a 20× objective and focused on the sample with an achromatic objective with a 0.82 numerical aperture. The reflected light was collected with the same objective and then focused onto a single-mode fibre and detected using a liquid nitrogen-cooled CCD spectrometer. The setup is confocal in collection due to the small diameter of the core of the collection fibre. The incident and collected polarisation of the light was controlled using a series of linear polarisers, quarter-wave and half-wave plates.

Data availability

The dataset generated and analysed during the current study is available at <https://doi.org/10.17861/5a2ae35f-9787-47bc-bb0e-a0395f064509>. Source data are provided with this paper.

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Author contributions

S.F. and A.J.C. contributed equally to this work. S.F. and A.J.C. performed the optical measurements. H.B. fabricated the sample. D.A.-P. performed transfer matrix analysis. S.F. and M.B.-G. performed coupled oscillator model simulation. S.F., A.J.C., M.B.-G., B.U., and B.D.G. analysed the data. I.C.G. performed the theoretical calculations. T.T. and K.W. grew the hBN crystals. B.D.G. and M.B.-G. conceived and supervised the project. S.F., A.C., M.B.-G., and B.D.G. wrote the paper with input from all authors.

Competing interests

The authors declare no competing interests.

Additional information

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