

Proximity-induced chiral quantum light generation in strain-engineered WSe₂/NiPS₃ heterostructures

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Quantum light emitters capable of generating single photons with circular polarization and non-classical statistics could enable non-reciprocal single-photon devices and deterministic spin–photon interfaces for quantum networks. To date, the emission of such chiral quantum light relies on the application of intense external magnetic fields, electrical/optical injection of spin-polarized carriers/excitons or coupling with complex photonic metastructures. Here we report the creation of free-space chiral quantum light emitters via the nanoindentation of monolayer WSe₂/NiPS₃ heterostructures at zero external magnetic field. These quantum light emitters emit with a high degree of circular polarization (0.89) and single-photon purity (95%), independent of pump laser polarization. Scanning diamond nitrogen-vacancy microscopy and temperature-dependent magneto-photoluminescence studies reveal that the chiral quantum light emission arises from magnetic proximity interactions between localized excitons in the WSe₂ monolayer and the out-of-plane magnetization of defects in the antiferromagnetic order of NiPS₃, both of which are co-localized by strain fields associated with the nanoscale indentations.

Defined as the process by which an atomically thin material acquires properties from adjacent materials via quantum mechanical interactions, proximity effects have recently emerged as a means of inducing desirable magnetic, topological, transport and optical properties in two-dimensional (2D) systems^{1–3}. Strong enhancements in valley Zeeman splitting^{4,5} and spin-dependent charge transfer^{6–9} have been observed through the coupling of semiconducting transition metal dichalcogenide (TMD) monolayers to magnetic thin films (for example, EuS)⁴ and van der Waals magnets (for example, CrI₃ (refs. 8–10), CrBr₃ (refs. 6,7), Cr₂Ge₂Te₂ (ref. 5) and Fe₃GeTe₂ (ref. 11)). The inverse

proximity effect where TMD monolayer altering 2D magnetic materials through spin–orbit coupling has also been reported¹². However, few studies have focused on exploiting proximity effects to manipulate the chirality of quantum emitters (QEs) in monolayer TMDs^{5,8,11}. To date, such manipulations demand the application of external magnetic fields^{13,14}, injection of spin-polarized carriers/excitons^{2,15–17} or coupling with complex photonic metastructures^{18,19}.

In parallel, the studies of transition metal phosphorous trichalcogenide TMPX₃ (TM = Mn, Ni, Fe, Co; X = S, Se) van der Waals antiferromagnetic (AFM) crystals have recently opened new frontiers for

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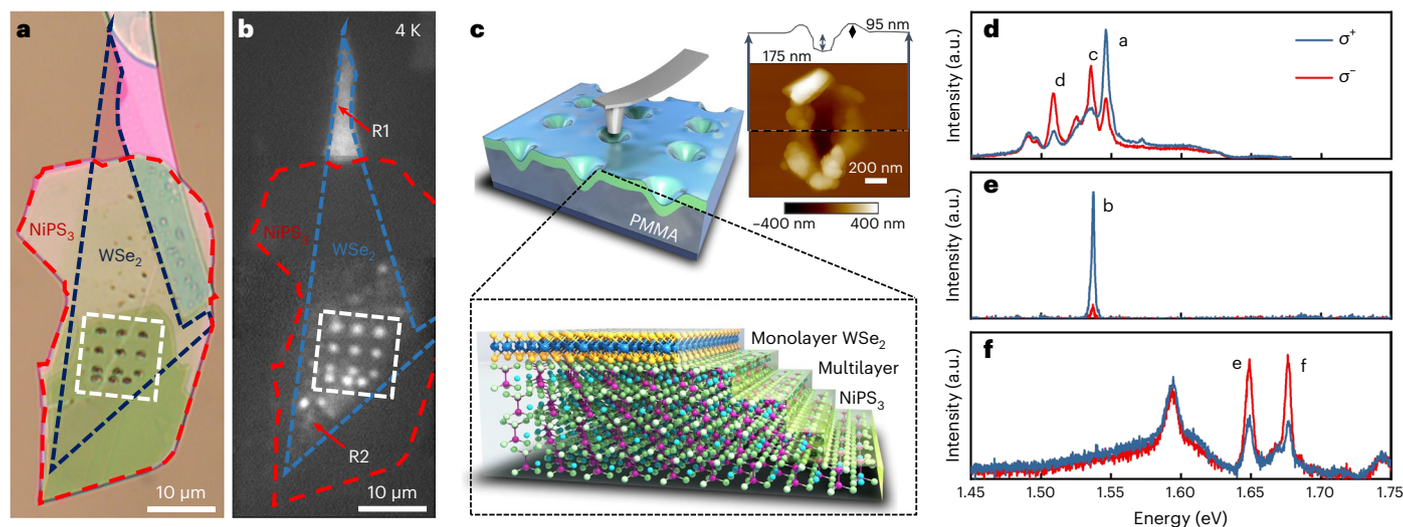


Fig. 1 | Strain-engineered WSe₂/NiPS₃ heterostructures host QEs displaying sharp, localized PL peaks with a strong degree of spontaneous circular polarization. **a, b**, Optical (**a**) and PL (**b**) images of a WSe₂/NiPS₃ heterostructure. The portion of the WSe₂ monolayer that does not overlap with NiPS₃ emits bright PL (region R1), whereas PL is quenched when the WSe₂ is coupled to the NiPS₃ (region R2). Strong PL was restored by indentations marked by the white square. **c**, Schematic of the sample structure and an atomic force microscopy topography image and cross section of a representative indentation (inset).

d–f, Representative σ^+ -resolved (blue) and σ^- -resolved (red) PL spectra from different individual nanoindentations marked under linearly polarized laser excitation at 4 K. Although the peaks marked as **a** and **b** are σ^+ polarized with DCP values of 0.36 and 0.89, peaks marked **c–f** are σ^- polarized with DCP values of -0.40 , -0.29 , -0.33 and -0.37 , respectively. Some of the localized emission peaks ride on top of a broad PL background that displays little polarization, suggesting that the DCP of localized emission peaks could even be higher if the broad PL background is subtracted.

exploring 2D magnetism with applications in spintronic and quantum information technologies²⁰. Recent breakthroughs include the discovery of charge–spin correlation²¹, the spin-induced linear polarization of excitons^{22–24} and the switching of the Néel vector under strain²⁵. The potential of these materials to support magnons, which are the collective excitation of local magnetic moments²⁶, as a means of coupling QEs represents an additional opportunity for the realization of microwave–magnon–spin quantum interfaces²⁷. Still, the study of whether TMPX₃ materials are capable of modifying the properties of free and localized excitons in 2D TMD materials via the proximity effect remains in its nascent stage²⁸. Here we show that WSe₂/NiPS₃ heterostructures locally strained via nanoindentations²⁹ can support QEs capable of emitting single photons with strong circular polarization at zero magnetic field and independent of excitation polarization. This observation is surprising because NiPS₃ displays AFM order²⁶, and is therefore not expected to yield a net out-of-plane magnetization capable of enhancing the valley Zeeman effect and thus induce circularly polarized photoluminescence (PL). This finding could be exploited in the development of non-reciprocal single-photon devices and deterministic spin–photon interfaces^{18,19} for quantum networks.

Circularly polarized PL at zero magnetic field

Figure 1a–c displays a representative heterostructure consisting of an ~ 50 -nm-thick NiPS₃ flake and a monolayer-thick WSe₂ flake stacked atop a Si wafer coated with a 300-nm-thick polymer layer. Wide-field PL images acquired at cryogenic temperatures ($T = 4$ K) under 514 nm continuous-wave laser excitation (Fig. 1b) show that PL from the WSe₂ monolayer is strongly quenched when coupled to NiPS₃. The PL spectra acquired from unstrained regions R1 and R2 showed no degree of circular polarization (DCP), indicating that electron–hole pairs preferentially pumped non-resonantly into the K^+ valley relax equally to both valleys before recombination (Extended Data Fig. 2). Next, localized QEs on the heterostructure were created using strain engineering²⁹ by pressing a blunt atomic force microscopy probe (~ 100 nm radius of curvature) into the heterostructure to form an array of nanoindentations measuring ~ 250 nm in diameter, 175 nm in depth and surrounded

by an ~ 100 -nm-high circular ridge (Fig. 1c). Bright localized PL emission with spectrally narrow peaks is observed at indentation locations (Fig. 1d–f)²⁹. In contrast to the unpolarized PL emission of narrow peaks reported on WSe₂ monolayer on polymethyl methacrylate (PMMA)²⁹ and hexagonal boron nitride (Extended Data Fig. 3a), we observed clear, circularly polarized PL from some of the sharp emission peaks under linearly polarized laser excitation at zero magnetic field (Fig. 1d–f). This circularly polarized emission behaviour was also observed to remain unchanged when the excitation laser polarization is changed to σ^+ , σ^- or linear polarization of different orientations (Extended Data Fig. 3b,c). Both polarity and DCP ($(I_{\sigma^+} - I_{\sigma^-}) / (I_{\sigma^+} + I_{\sigma^-})$) vary widely among the localized PL emission peaks. Some indented spots display multiple localized peaks with opposite circular polarizations (Fig. 1d). An analysis of the spectral image of sharp PL peaks from an individual indentation (Extended Data Fig. 4) reveals that these sharp PL peaks originate from independent localized exciton states separated by 50 to a few hundred nanometres. Also, σ^+ and σ^- peaks of all the localized emitters were found to occur at identical energies. Localized emitters also exhibit weak linear polarization (Extended Data Fig. 5a,b) with zero energy splitting between the two components. Based on these observations, we attribute the σ^+ and σ^- PL peaks of localized emitters to the emission of a single-exciton state exhibiting a strong but incomplete DCP. The chiral emission peaks are observed in the same 1.5–1.7 eV range as in the case of strain-engineered WSe₂ monolayers³⁰ with linewidths of 3.6 ± 1.3 meV (Extended Data Fig. 5c), broader than the sharpest linewidths reported³⁰, but comparable with those formed via nanoindentations²⁹ and those induced via strain on the edge of SiN photonic waveguides³¹.

Chiral quantum light emission

To demonstrate the capability of chiral emission sites to act as QEs, we performed Hanbury Brown–Twiss experiments. Figure 2a,b shows circular-polarization-resolved PL spectra recorded from an indentation excited with either σ^+ and σ^- polarization, respectively. Spectrally filtering the lowest-energy peak ($E = 1.629$ eV, DCP = -0.3) showing strong σ^- polarization, we next acquired the intensity versus time trace,

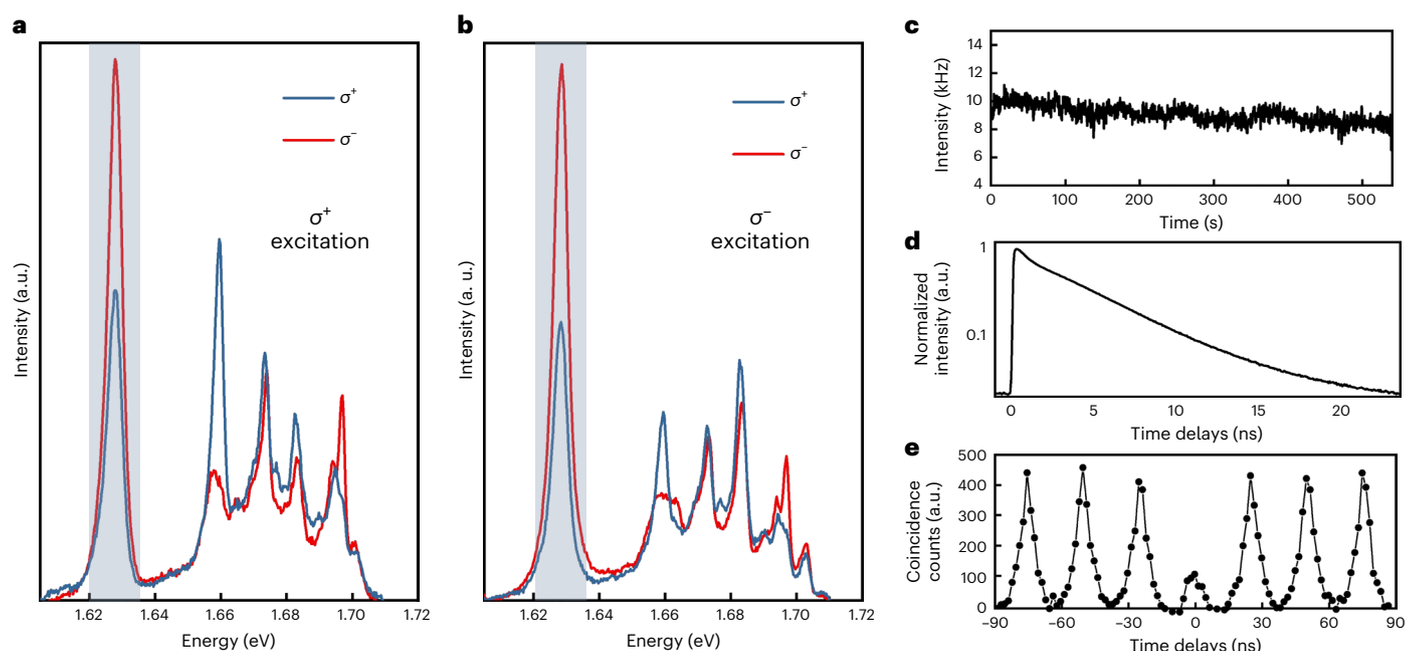


Fig. 2 | Demonstration of quantum light emission from strain-engineered WSe₂/NiPS₃ heterostructures. **a, b**, σ^+ -polarized (blue) and σ^- -polarized (red) PL spectra of an indented WSe₂/NiPS₃ heterostructure acquired under σ^+ -polarized (a) and σ^- -polarized (b) laser excitation. **c–e**, PL intensity time trace (c), PL decay

curve (d) and second-order photon correlation trace (e) measured from the PL peak marked by the blue column in a and b acquired under 514 nm, 40 MHz pulsed laser excitation.

decay curve and second-order photon correlation function ($g^{(2)}(\tau)$) of the PL (Methods). The PL intensity versus time trace (Fig. 2c) shows blinking-free emission, whereas the decay curve (Fig. 2d) can be fitted to a single exponential with a lifetime of 4.35 ± 0.01 ns. Ultimately, the $g^{(2)}$ trace (Fig. 2e) provides clear evidence of quantum light emission such that for more than 80% of the time, the QE emits a single photon per excitation–emission cycle (meaning single-photon purity of 80%) comparable with other reported WSe₂ QEs³⁰. We attribute the residual multiphoton emission probability ($g^{(2)}(0)/g^{(2)}(T)$) of 0.2 to the broadband emission tail extending from the group of higher-energy PL peaks. The dataset for another chiral QE emitting more than -0.7 DCP and having greater than 95% single-photon purity is shown in Extended Data Fig. 6.

Together, these findings provide clear evidence that the local strain engineering of WSe₂/NiPS₃ heterostructures can form QEs capable of generating chiral single photons without the need for an external magnetic field and independent of laser excitation polarization (Figs. 1 and 2 and Extended Data Fig. 3b,c). This observation is fundamentally different from PL polarization ($\rho(B)$) reported to arise from spin-dependent charge transfer in WSe₂/CrI₃ heterostructures^{9,10}. These studies compare σ^+ -polarized PL detected under near-resonant (1.96 eV) σ^+ laser excitation (I_{σ^+/σ^+}) with σ^- -polarized PL detected under σ^- laser excitation (I_{σ^-/σ^-}). A non-zero value of $\rho(B) = (I_{\sigma^-/\sigma^-} - I_{\sigma^+/\sigma^+}) / (I_{\sigma^-/\sigma^-} + I_{\sigma^+/\sigma^+})$, therefore, does not indicate the emission of chiral light. In fact, studies of the spin-dependent charge transfer effect in MoSe₂/CrBr₃ heterostructures^{6,7} that measured DCP independent of excitation laser polarization⁷ or under linearly polarized laser excitation⁷ report no detectable DCP at zero magnetic field. These studies indicate that a simple TMD/MX₃ material system cannot emit chiral classical light. Since these works^{6,7,9,10} primarily focused on the 2D exciton, quantum light (single-photon) emission is not possible. Proximity interaction between the strain-engineered QEs of WSe₂ and 2D magnets such as Cr₂Ge₃Te₆ (ref. 5), Fe₃GeTe₂ (ref. 11) and CrI₃ (ref. 8) were recently investigated. No chiral light emission was reported when QEs were coupled to Cr₂Ge₃Te₆ (ref. 5) and Fe₃GeTe₂ (ref. 11) (only enhancement in Zeeman

splitting⁵ and some increase in zero field splitting of the fine-structure states of QEs¹¹ were reported, respectively). A small value of DCP ≈ 0.15 was observed at zero magnetic field for the localized excitons of WSe₂ coupled to CrI₃ but no photon antibunching was demonstrated¹⁸. Our demonstration of quantum light emission with DCP up to 0.89 in the absence of an external B field and spin-polarized pumping, therefore, clearly stands in contrast with these prior works. Our observations demonstrate the ability of strain-engineered NiPS₃ to modify the optical characteristics of QEs via magnetic proximity interactions. Besides immediate relevance for chiral quantum photonics, these results may also be relevant to the long-standing pursuit of spin lasers^{15,32} by providing a means to ensure the robust spin polarization of electrons and holes via magnetic proximity effects.

Magnetic proximity interaction evidenced by scanning NV microscopy

Because NiPS₃ exhibits zig-zag AFM order with spins aligned parallel or antiparallel to the in-plane a axis compensating each other at the lattice constant (Fig. 3a)^{22–24,26}, the net magnetization needed to modify the optical properties of nearby materials through proximity effects is practically averaged out to zero. However, studies have revealed that localized ferromagnetic (FM) moments can emerge in AFM materials due to flexomagnetism (the emergence of net magnetic moments due to bending and/or strain gradients)^{33,34} or due to spatial variations in the AFM order parameter, as they occur, for example, in domain walls³⁵. Based on these findings, we hypothesize that the magnetic moments of Ni atoms in the strained region around the indentation rotate to point out of plane to form local FM moments due to severe deformations of the NiPS₃ lattice and/or changes in the magnetic anisotropy energy. Since the indentation could create localized exciton states in the WSe₂ layer in close vicinity of the local FM moments of NiPS₃, the proximity-induced exchange interaction between the FM moments and localized exciton further leads to the time-reversal symmetry breaking necessary for the emission of chiral single photons. Such proximity interaction has been well established on both experimental and theoretical grounds^{1,2,36}.

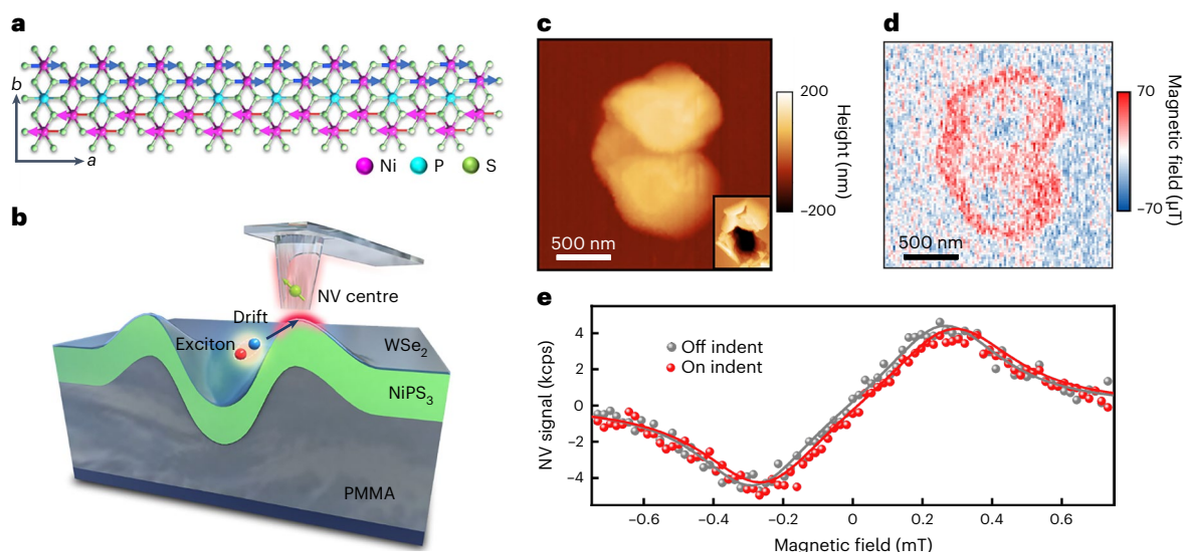


Fig. 3 | Correlation of chiral quantum light emission with local magnetization. **a**, Zig-zag AFM order overlaid with the atomic NiPS₃ structure where Ni electronic spins are aligned parallel (blue arrows) and antiparallel (red arrows) to the crystallographic *a* axis. **b**, Schematic of an indentation region cross section where the local strain creates exciton-capture centres in WSe₂. **c,d**, Scanning NV-centre mapping of the surface magnetization associated with the morphology of a nanoindentation (**c**) is strongly correlated with the magnetic

field measured in the indentation region (**d**). **e**, NV spectra taken both off (grey) and on (red) the indented region show a shift in the detected magnetic signal of $B_{\text{indent}} = 26 \pm 2 \mu\text{T}$. A high-resolution atomic force microscopy image of the nanoindentation is provided (inset of **c**) as the NV scanning probes have reduced spatial resolution. The scanning NV centre has a standoff distance of $70 \pm 20 \text{ nm}$, meaning that the measured magnetic field is reduced by spatial averaging and inverse distance scaling. kcps, kilo counts per second.

To confirm this hypothesis, we imaged the magnetic signal from an indent in a bare NiPS₃ flake using a nitrogen-vacancy (NV) centre³⁷ in a diamond atomic force microscope at $T \approx 4.4 \text{ K}$ (a strong PL background from the WSe₂ monolayer prevented a similar experiment on WSe₂/NiPS₃ heterostructures). Topography and magnetic-field images of an indent (Fig. 3c,d) show a magnetic signal of $B_{\text{mean}} = 22 \pm 9 \mu\text{T}$ from the ridge region surrounding the indentation. The magnetic signal was confirmed by obtaining the full NV lock-in spectra (Methods) both on and off the indent, which indicates a magnetic-field strength of $B_{\text{indent}} = 26 \pm 2 \mu\text{T}$ (Fig. 3e). A corresponding control experiment that was conducted at room temperature on the same nanoindentation showed no measurable stray magnetic fields on the sample (Extended Data Fig. 7). This magnetic signal is one order of magnitude weaker compared with that measured on a CrI₃ monolayer³⁸ and comparable with that arising due to flexomagnetism in AFM Cr₂O₃ films³³. Because the NV centre is separated from the sample surface by $70 \pm 20 \text{ nm}$ (standoff distance), the measured field is reduced by inverse vertical-distance scaling and the measurement only yields an average response of the $\sim 70\text{-nm}$ -diameter region directly beneath the NV centre (Fig. 3b). The measured field value, therefore, reflects only a fraction of the actual magnetization on the surface and loss of local fluctuations occurring at length scales of less than 70 nm .

This measurement of the stray magnetic field provides evidence for the emergence of local out-of-plane ferromagnetism in the indented NiPS₃ layer below the Néel transition temperature. However, it is crucial to note that the chiral light emission we observed does not arise from the interaction between the stray magnetic field and the QE. Instead, it originates from a proximity-induced magnetic-exchange interaction that occurs when the wavefunction of the localized exciton in WSe₂ overlaps with the electronic wavefunction of Ni ions, forming the local ferromagnetic moment. Since each Ni ion can carry the magnetic moment of the order of $1\mu_B$, the effective magnetic field inducing the chiral emission could be substantially stronger than the measured stray field of local magnetization by a few orders of magnitude. Since the interaction is highly localized and the irregular strain field of nanoindentation could create multiple local magnetizations of varying

directionalities and strengths, the QEs from the same indentations display varying directionalities and degrees of circular polarization.

Magnetic proximity interaction evidenced by magneto-PL

To provide further evidence linking our chiral light emission to localized ferromagnetic proximity effects, we performed an additional magneto-optical characterization of the PL emission from nanoindentations in WSe₂/NiPS₃ heterostructures. First, we analysed the 1.47 eV ultrasharp exciton emission peak of NiPS₃ (refs. 22–24), revealing that although the emission is strongly linearly polarized (degree of linear polarization, DLP = 0.76) perpendicular to the Néel vector of NiPS₃ on the unstrained region, the DLP of this peak from an indented region drastically reduced to 0.24 and became circularly polarized with DCP = 0.4 (Extended Data Fig. 5). This result indicates the existence of multiple domains with reoriented Néel vectors within an indented region—a prerequisite for the emergence of localized magnetization and ferromagnetic proximity effects.

Studies of NiPS₃ have shown that external fields greater than 15 T are required to reorient the Néel vector to the direction of the magnetic field at cryogenic temperatures below the Néel transition temperature ($T_N \approx 150 \text{ K}$)²³. Strain-induced localized magnetization associated with AFM defects is, therefore, expected to be stable against external B fields of lesser strength. To test this hypothesis, polarization-resolved PL spectra were acquired from an indentation under external B fields ranging from -6 to $+6 \text{ T}$ in the Faraday geometry (Fig. 4a,b and Extended Data Fig. 8). These PL spectra reveal that a strongly circularly polarized PL peak at -1.59 eV with a DCP of -0.6 exhibits no notable change in DCP as a function of B field (Fig. 4b). Consistent with the previous conclusion that the σ^+ and σ^- peaks of the chiral emitter originated from a single electronic state exhibiting incomplete circular polarization, energy splitting between the σ^+ and σ^- peak remains essentially zero for all the B fields (Fig. 4b). The spin doublets of localized excitons are known to display B -field-dependent, incomplete circular polarization behaviour when strain-induced, anisotropic exchange splitting competes with Zeeman splitting resulting from the external B field^{30,39} (Supplementary Information). We could interpret

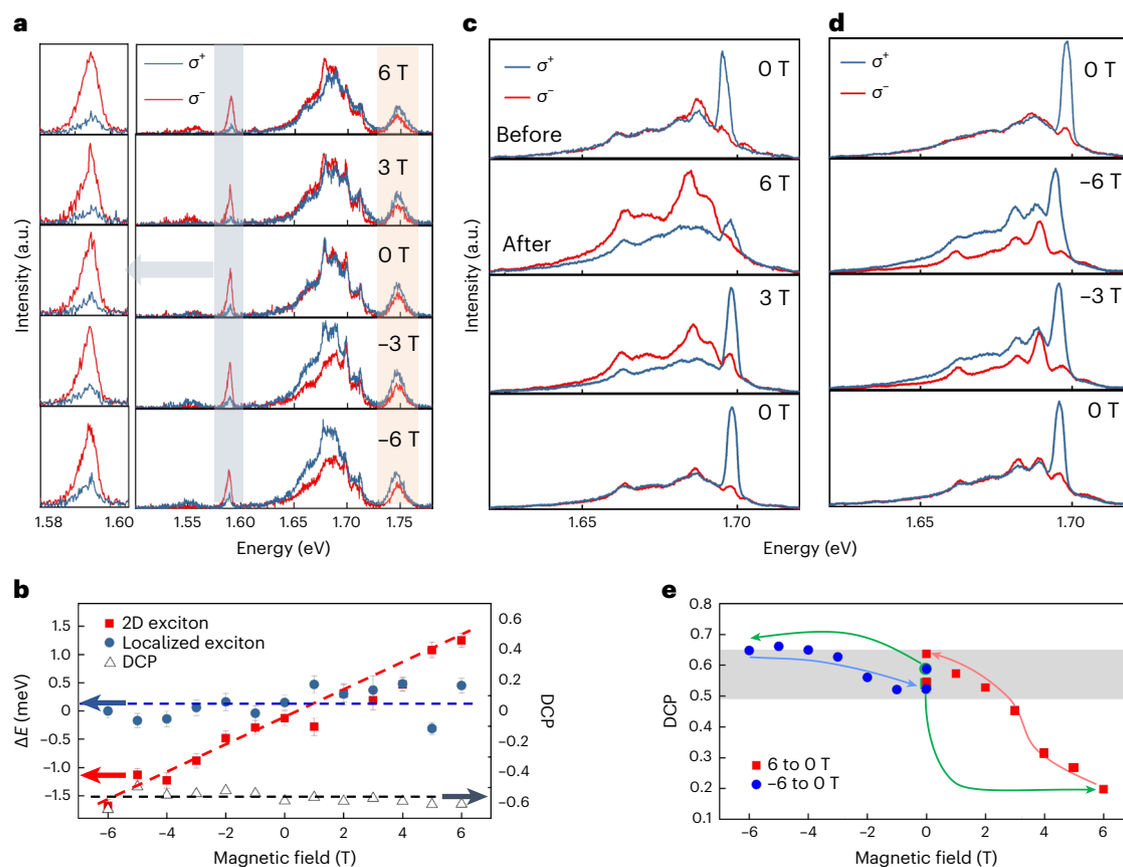


Fig. 4 | Magneto-PL studies of chiral QEs. a, Low-temperature, polarization-resolved PL spectra acquired as a function of external B field applied perpendicular to the sample plane (Faraday geometry). An expanded view of a localized PL peak (1.59 eV; grey bar) displays a strong DCP of -0.6 (left). **b**, Energy splitting (left axis) between the σ^+ and σ^- PL peaks of a localized exciton at 1.59 eV (blue data points) and a 2D exciton at 1.75 eV (red data points). The error bars represent the uncertainty in the peak positions determined by the Gaussian fit. The DCP (right axis) of the 1.59 eV peak versus B field is plotted as black triangles. The blue, red and black dashed lines are guides to the eye. **c**, Low-temperature, polarization-resolved PL spectra of an indented region with a PL peak at -1.70 eV, exhibiting a strong DCP of -0.7 before the temperature- and field-dependent measurement (top row). Low-temperature, polarization-resolved PL spectra of the same location acquired after heating the sample to 180 K, raising the field to 6 T and cooling the sample

back down to 4 K while maintaining B at 6 T (second row). Low-temperature, polarization-resolved PL spectra acquired as the field is reduced back to 0 T (two bottom rows). **d**, The experiment in **c** was repeated by applying an external field of -6 T at 180 K. **e**, DCP of the 1.69 eV peak plotted against an external B field. The green arrows originate from the DCP value originally observed in the top-most panels of **c** and **d** and indicate the process involving heating the sample to 180 K and cooling back down to 4 K while maintaining the B field at 6 T (-6 T). The red (blue) arrows and data points represent the process and DCP observed during the reduction in B from 6 T (-6 T) to 0 T at 4 K immediately after cooling down. Since the DCP values are calculated from the peak intensities of the σ^{\pm} -polarized spectra, they are contributed not only by the QE emission but also by the broad PL background that increases as the field is ramped from $+6$ to -6 T. The grey-shaded region represents variations in the DCP measured at 0 T.

the chiral PL peak as the lower-energy state of such a spin doublet. The B -field invariance in both PL emission energy and DCP suggests that the magnetic-exchange interaction with FM moments within NiPS₃ is dominating over the effect of the external B field in inducing valley splitting and chiral quantum light emission. In marked contrast with the localized QE emission, the 2D WSe₂ exciton peak at 1.75 eV displays normal Zeeman splitting (-1.68 meV at ± 6 T) similar to uncoupled monolayers, suggesting that proximity interactions average out to zero for delocalized 2D excitons and that chiral PL emission demands the co-localization of exciton and local moments arising from AFM defects.

The defect-induced local magnetic moment in NiPS₃ (to which localized excitons in WSe₂ couple), can, however, be influenced by the application of large magnetic fields, if applied at elevated temperatures. These studies are shown in Fig. 4c,d. First, a QE emitting strongly circularly polarized PL at 4 K and $B = 0$ T was identified (1.7 eV PL peak; Fig. 4c). We then raised the temperature to 180 K (above T_N) and applied a $+6$ T magnetic field along the sample normal. The sample was then cooled back to 4 K and the applied field was maintained at $+6$ T. The PL spectra acquired after cooling to 4 K at 6 T (Fig. 4c, second row) showed a dramatic decrease in the DCP of QE to -0.2 from the original value of -0.7 , indicating

that the local magnetization responsible for the QE's chiral PL emission is reduced, probably as a consequence of the applied B field opposing the local moment's preferred orientation direction. When the external B field is decreased, the DCP was found to increase until it was fully restored to its original value at 0 T (Fig. 4c,e and Extended Data Fig. 9), suggesting that the local strain distribution and underlying AFM order restores the original local magnetization in the absence of an external field. In contrast, when a B field in the opposite direction (-6 T) is applied at 180 K, no notable change in the DCP of QE is observed since B aligns parallel to the preferred local magnetization, making it more stable (Fig. 4d,e). Importantly, in both cases, the chiral emission is restored at $B = 0$ T, and sweeping the fields from $+6$ to -6 T after the restoration of chiral emission at 0 T (Extended Data Fig. 9e,f) reproduces the B -field-independent behaviour of chiral QE (Fig. 4a). This restoration of chiral emission and thermal hysteresis behaviour indicate that the underlying AFM order is unaffected by ± 6 T applied fields, as expected. This experiment provides strong evidence that our chiral QEs form due to proximity interactions with localized out-of-plane magnetizations that are intimately tied to FM moments in NiPS₃ and AFM order of the surrounding NiPS₃ stabilized the orientation and strength of the FM moments against the external B field.

In summary, our observations reveal that local strain engineering can be utilized not only to create QEs but also to localize ferromagnetic proximity effects required for the creation of chiral single-photon emitters in $\text{WSe}_2/\text{NiPS}_3$ heterostructures. Interestingly, a similar chiral localized excitonic emission was also observed in our more recent experiment performed on $\text{WSe}_2/\text{MnPS}_3$ and $\text{WSe}_2/\text{FePS}_3$ heterostructures with nanoindentations (Extended Data Fig. 10). These discoveries establish TMD/TMPX₃ AFM insulators as an exciting material platform for the further exploration of novel emergent phenomena and the realization of solid-state quantum transduction and sensing technologies.

Online content

Any methods, additional references, Nature Portfolio reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at <https://doi.org/10.1038/s41563-023-01645-7>.

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Methods

Sample preparation

A PMMA solution was spin coated on a Si substrate with a 285 nm SiO₂ layer to achieve a 300-nm-thick uniform membrane. NiPS₃ flakes were exfoliated from bulk crystals (2D Semiconductors) using Scotch tape and deposited onto the PMMA layer. Monolayers of WSe₂ (HQ Graphene) were mechanically exfoliated onto a silicone Gel-Film stamp. The layer thickness was identified by optical microscopy contrast and room-temperature PL. The WSe₂ monolayers were then transferred onto the selected NiPS₃ flakes using a 2D material transfer system from HQ Graphene. NiPS₃ and WSe₂ exfoliations as well as layer stacking were performed in an argon-filled glove box to minimize degradation. Once the heterostructure was fabricated, nanoindentations were deterministically created by using a Bruker Dimension Icon instrument using blunt atomic force microscopy probes with a nominal tip radius of 100 nm and a spring constant of 80 N m⁻¹. Indentations were fabricated by performing approach curves with a maximum force setpoint between 50 and 60 μN. This combination of probe radius and force setpoint was found to consistently produce chiral QEs within the WSe₂/NiPS₃ heterostructures. Topography imaging of the surface following nanoindentation was performed using tapping-mode imaging with a second, higher-resolution topography probe (tip radius, ~10 nm).

Low-temperature polarization-resolved PL spectroscopy and Hanbury Brown–Twiss experiments

The schematic of the experimental setup for optical measurements is illustrated in Extended Data Fig. 1. The sample was held at 4 K in a continuous-flow cryostat (Oxford Instruments, MicrostatHires). The heterostructures were excited with a 514 nm continuous-wave laser (Coherent OBIS) and the PL signal was collected through a ×50, 0.55-numerical-aperture Olympus microscope objective. A quarter-wave plate (QWP) was placed immediately after the objective to circularly polarize the laser excitation and converted the σ^+ - and σ^- -polarized PL signal into *S* and *P* linear polarization. A Wollaston prism was then used to split the signal into channels, which were then projected onto a liquid-nitrogen-cooled charge-coupled device detector through a 500 mm spectrometer (Acton SpectraPro). This arrangement allowed the simultaneous collection of the spectral images of σ^+ and σ^- PL signals on two different portions of the charge-coupled device camera (Extended Data Fig. 1). The σ^+ and σ^- PL spectra and the extracted DCP are, therefore, free of artifacts that could result from random intensity fluctuations and spectral wandering. For linear polarization analysis, the QWP was removed, and two half-wave plates (HWP1 and HWP2) were inserted in the excitation and collection channels before the Wollaston prism. The PL spectra were acquired as a function of the rotation angles of the half-wave plates to determine the degree and orientation of PL linear polarization. HWP2 was replaced with a QWP to acquire σ^+ - and σ^- -polarized PL spectra under linearly polarized excitation (Extended Data Fig. 3). For the time-tagged time-correlated PL measurements, we pumped the sample with a tunable visible picosecond laser (TOPTICA Photonics, 40 MHz repetition rate, 3.5 ps pulse width) filtered by a 10 nm band-pass filter centred at 514 nm. The emission was then filtered by using a tunable band-pass filter (SLI) and coupled into a Hanbury Brown–Twiss spectrometer composed of a 50/50 beamsplitter and two avalanche photodiodes (MPD PDM Series) with a timing accuracy of 35 ps. The macro- and microtimes of each photon detection event were recorded with a HydraHarp 400 time-correlated single-photon counting module and the PL intensity time traces, PL lifetime and second-order photon correlation ($g^{(2)}(\tau)$) were extracted from the recorded photon stream. Here $g^{(2)}(\tau)$ depicts the probability of registering two successive photons as a function of the time delay (τ) between the detection events. Under pulsed laser excitation, the $g^{(2)}$ trace displays a periodic series of peaks separated by the delay time between the laser excitation pulses. These peaks indicate the detection of two successive photons that result from

excitation by different laser pulses. Since a single QE can only produce at most one photon per pulsed excitation, the peak at zero delay time ($g^{(2)}(0)$) should be completely absent. The $g^{(2)}$ trace in Fig. 2e shows the $g^{(2)}(0)$ peak with an area of 20% compared with the side peak. This fact indicates that our chiral QE produces single photons 80% of the time.

Scanning diamond NV-centre magnetometry imaging

To perform local magnetometry on the NiPS₃, a new WSe₂/NiPS₃ heterostructure on PMMA was prepared with nanoindents on both pure NiPS₃ and WSe₂/NiPS₃ heterostructures. The sample was mounted inside a combined optical and atomic force microscope housed in a cryostat (attoLIQUID1000) where the atomic force microscope sensor consisted of a diamond and hosted the sensing NV at its tip (Qnami, Quantilever MX)³⁸. All the measurements were performed at an approximate temperature of $T \approx 4.4$ K. The NV centre was excited with a 532 nm laser to optically polarize the NV spin and to read out the spin state via a spin-state-dependent PL, whereas the electron spin was driven with a frequency-tunable microwave source to record the electron spin resonance spectra. To reduce the background light from the sample, a 600 nm long-pass filter and a 700 nm short-pass filter (Thorlabs) were used before the avalanche photodiode detector. At each position, a lock-in measurement is performed to attain the Larmor precession frequency of the electron spin, which is determined by the magnetic-field projection along the NV axis (inclined at 60° with respect to the sample normal) through the Zeeman splitting of the NV spin's energy levels⁴⁰. The lock-in measurement works by introducing a frequency modulation to the microwave driving field that rapidly samples two frequencies that are offset by the measured linewidth of the NV spectrum. The difference in these two frequencies forms the feedback signal that linearly depends on the stray magnetic field, within a certain dynamic range (Fig. 3e). The NV-containing diamond probe has an ~500 nm tip diameter⁴¹ and the NV centre is separated from the sample surface by a 70 ± 20 nm standoff distance. Although the height image can, therefore, not fully resolve the indentation, the key features in the ridge area can be identified via a direct comparison with the high-resolution AFM image taken on a different system. The standoff distance leads to an inverse-distance-scale-down measured *B* field⁴² and limits the lateral resolution to 70 ± 20 nm with an additional blurring of the edge features by 250 nm assuming that the NV centre is located close to the centre of the 500-nm-diameter tip. The measurement of the *B* field surrounding the indents of the WSe₂/NiPS₃ heterostructures was not feasible due to the strong optical background associated with the PL emission from the WSe₂ monolayer.

Temperature-dependent magneto-PL experiment

To perform the magneto-PL experiments, the sample was mounted in the variable-temperature insert of a 7 T magneto-optical cryostat (Oxford Instruments Spectramag). An in situ aspheric lens (numerical aperture = 0.68), mounted on piezoelectric nanopositioners (attocube), focused the incident light. The sample was excited using a continuous-wave 632.8 nm HeNe laser. The polarization of the laser was controlled using a QWP and a linear polarizer. Emission from the sample in the Faraday geometry was collected by the same aspheric lens, and its polarization was analysed using a QWP and linear polarizers. The emission was dispersed in a 500 mm spectrometer (Acton) and detected by a liquid-nitrogen-cooled charge-coupled device detector. For the temperature-dependent magneto-PL measurements, we first set the sample temperature to 180 K (above the Néel temperature of the NiPS₃) and then ramped the magnetic field to either +6 or -6 T. The sample was then cooled back to 4 K to repeat the field-ramping experiments.

Data availability

Source data are provided with this paper. All other data supporting the plots within this paper and other findings of this study are available

from the Supplementary Information or from the corresponding author upon reasonable request.

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Author contributions

H.H. conceived and led the experiment. X.L., under the supervision of H.H., initiated the idea of using the nanoindentation method for enhancing the magnetic proximity effect along with the deterministic creation of localized QEs, prepared the samples and conducted the experiments. A.C.J. guided X.L. in creating nanoindentations in the WSe₂/NiPS₃ heterostructure and scanning probe microscopy studies. S.A.C. contributed the key idea of the temperature-dependent magneto-PL experiment and X.L. and J.C. conducted the experiment. H.Z., M.T.P. and V.C. assisted X.L. in a variety of optical spectroscopy experiments. N.S. and A.P. provided the theoretical model. D.A.B., M.A.T. and P.R. performed the NV magnetometry measurements under the supervision of P.M. H.H. and X.L. wrote the manuscript with the assistance of all the authors.

Competing interests

The authors declare no competing interests.

Additional information

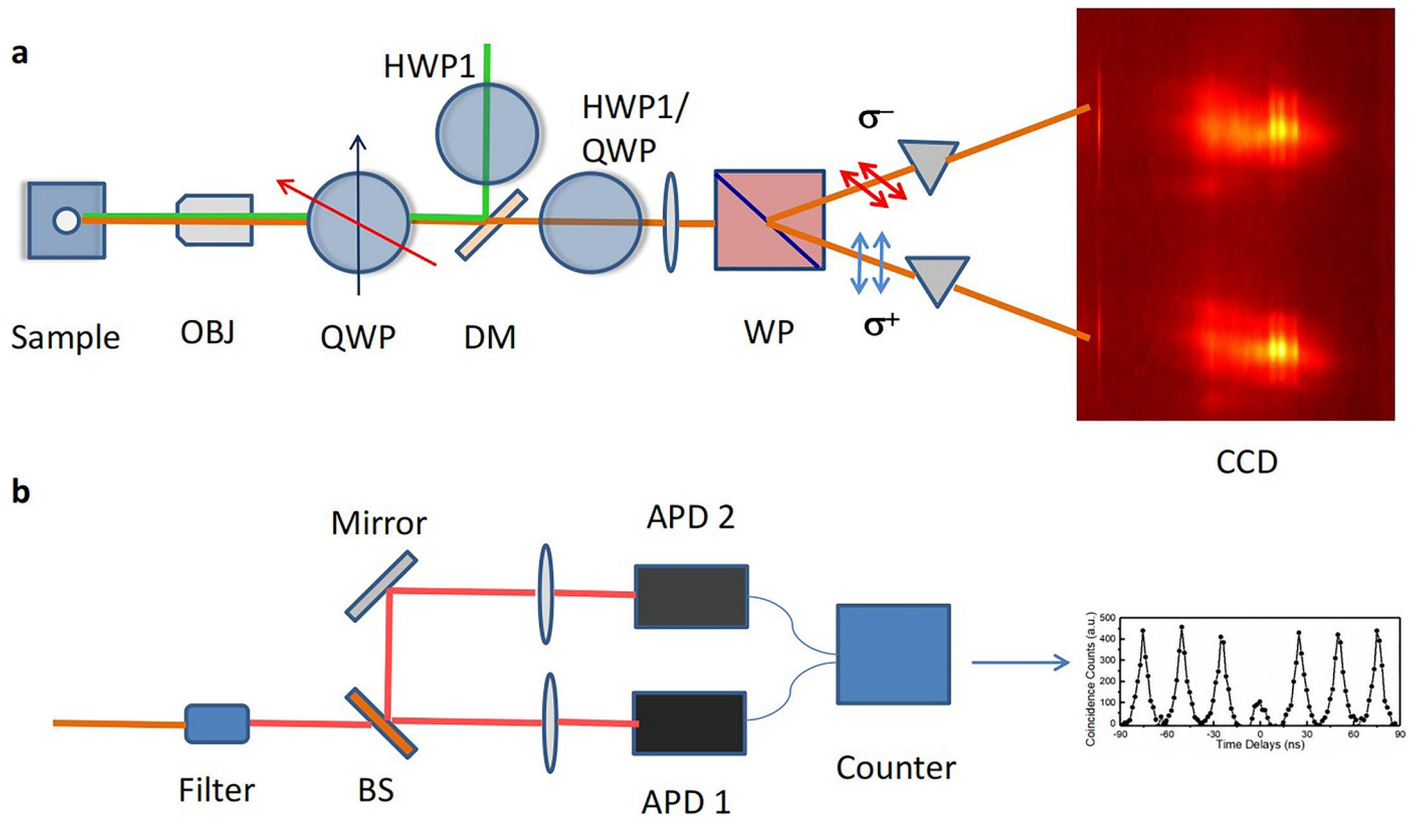
Extended data is available for this paper at <https://doi.org/10.1038/s41563-023-01645-7>.

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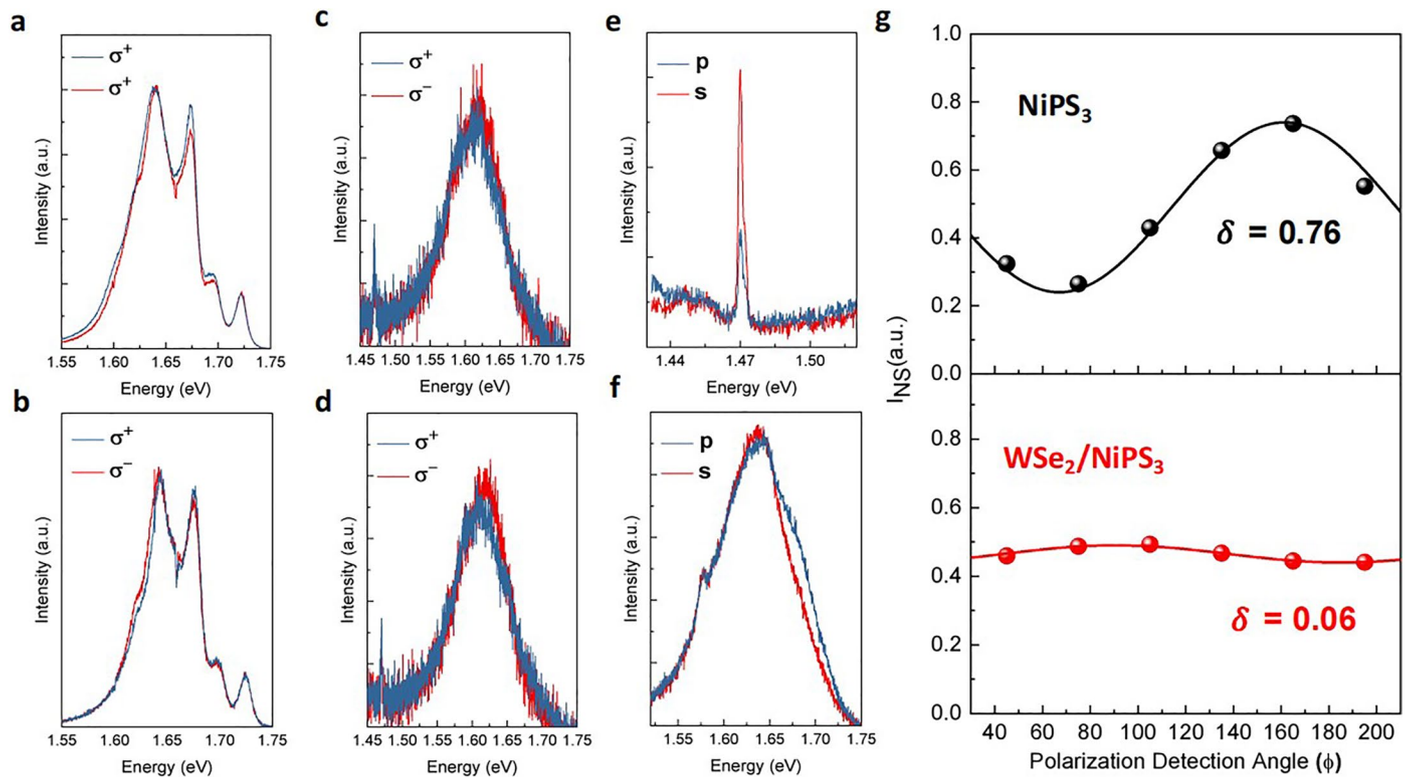
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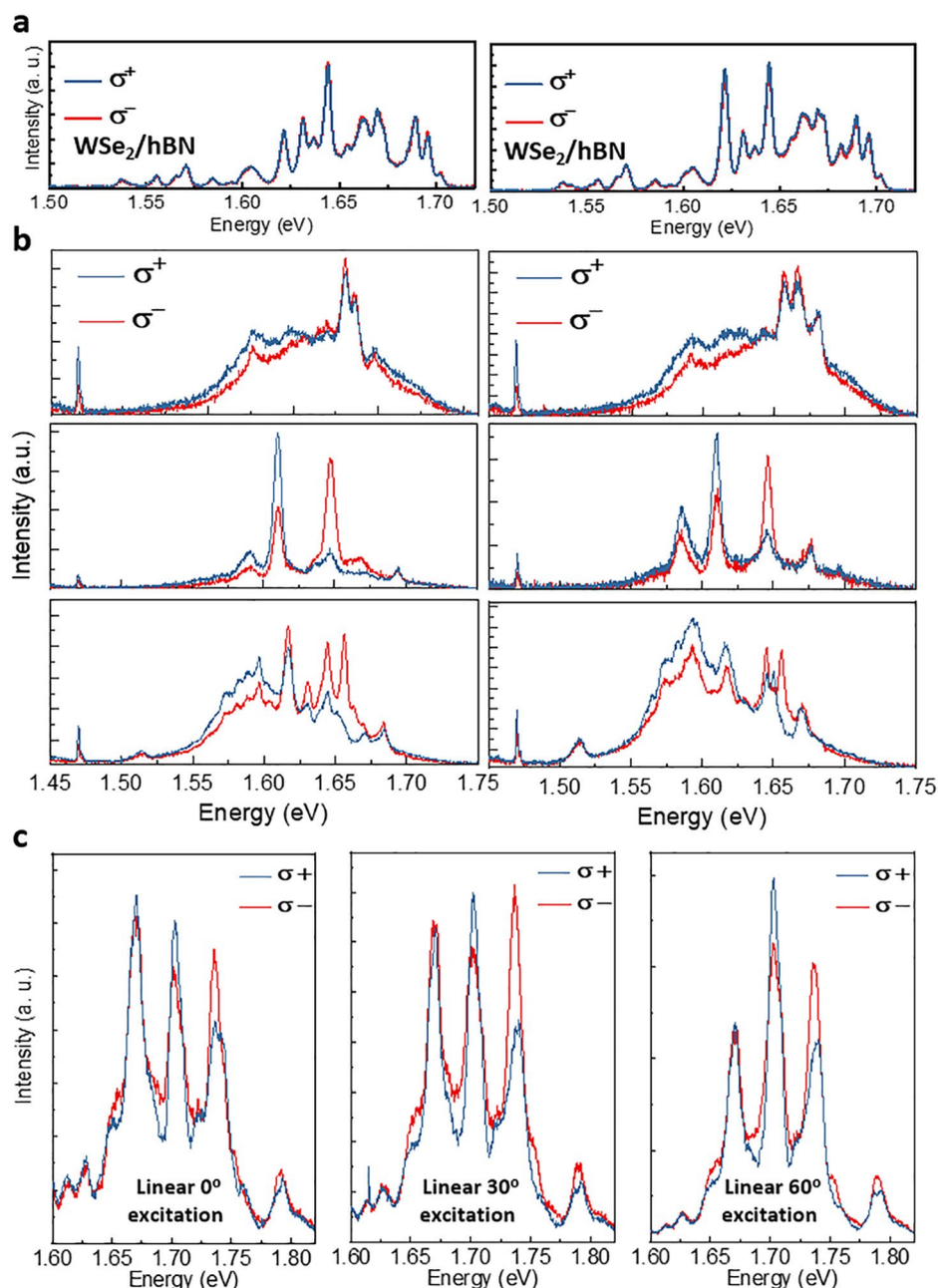
Extended Data Fig. 1 | Experimental setup for low-temperature polarization resolved PL spectroscopy. a. OBJ: Objective, DM: Dichroic mirror, QWP: quarter-wave plate, WP: Wollaston prism. For the linear polarization measurements, QWP is removed and two half-wave plates (HWP1 and HWP2) are inserted in excitation

and collection channels. **b.** Schematic of the HBT experiment. BS: 50/50 beam splitter, APD1, APD2: Avalanche photo diodes, Counter: Hydra400 TCSPC module.



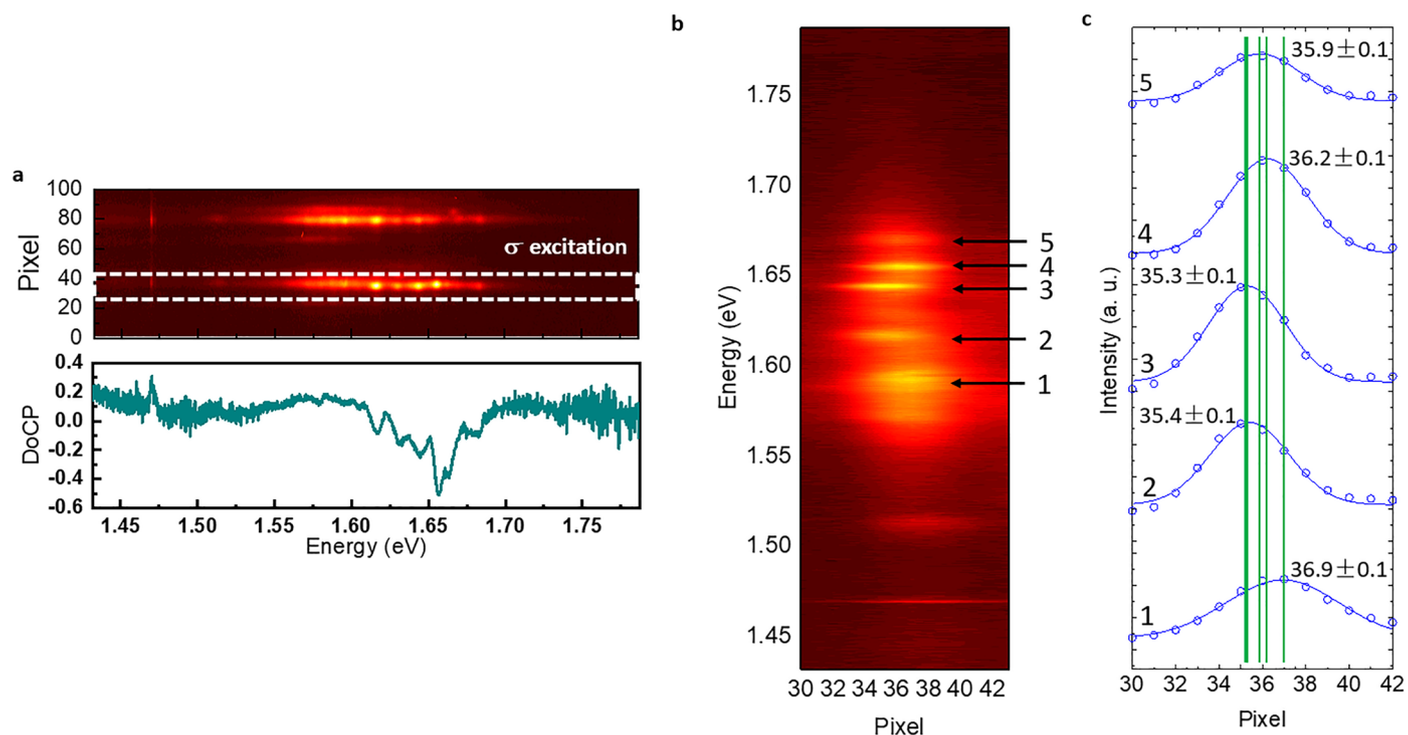
Extended Data Fig. 2 | Control experiments. a-d. Low temperature, σ^{\pm} polarized PL spectra of WSe₂ monolayer, region R1 of Fig. 1b (a, b) and un-indented WSe₂/NiPS₃ heterostructure, region R2 of Fig. 1b (c, d) excited by σ^+ (a, c) and σ^- (b, d) 514 nm laser. The data show no detectable DCP in the emission of both monolayer and un-indented WSe₂/NiPS₃ heterostructure indicating that spin-polarized excitons injected at 514 nm were depolarized completely upon relaxation to the band-edge. e, f. Low temperature, linearly (S, P) polarized PL

spectra (left panel) and plot of DLP as the function of polarization detection angle (right panel) for the sharp emission peak at 1.47 eV (e) and WSe₂ exciton emission (f). (g) **Upper Panel.** Strong DLP (0.76) observed for 1.47 eV sharp emission peak is consistent with prior studies that attributed the emission to highly anisotropic excitons reflecting the emergence of AFM order in NiPS₃. **Lower Panel.** PL of unstrained WSe₂/NiPS₃ heterostructure shows a weak degree of linear polarization consistent with the behavior of WSe₂ monolayer.



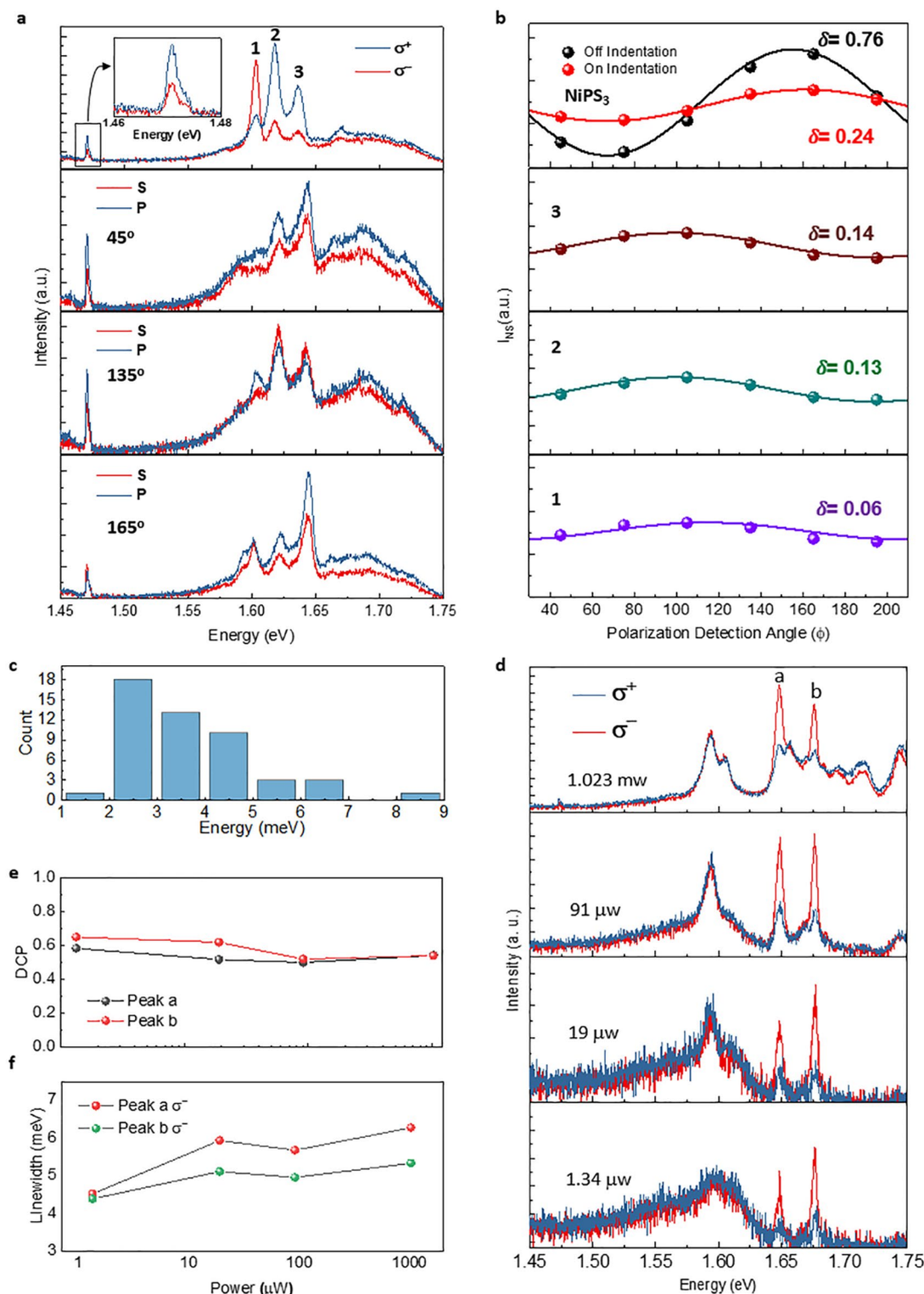
Extended Data Fig. 3 | Control experiment and invariance of circularly polarized PL emission under laser excitation of different polarization. a. σ^+ / σ^- polarization resolved PL spectra from WSe₂/hBN/PMMA structures with similar nano-indentations of Fig. 1, acquired under σ^+ (left) and σ^- (right) polarized laser excitation, respectively. While localized exciton emission peaks of similar line shapes were created, no circularly polarized emission were detected. **b.** σ^+ / σ^- polarization resolved PL spectra from 3 nano-indentations of Fig. 1a, b acquired under σ^+ (left) and σ^- (right) polarized laser excitation, respectively. While most

nano-indentations show circularly polarized localized exciton emission (bottom two panels), some show no circular polarization (top panels). When the excitation laser polarization is switched from σ^+ to σ^- , the polarities of the circularly polarized emission peak remain the same (peak marked *a* and *b*) but some quantitative change in DCPs as well as relative intensities of the localized PL peaks occur. **c.** Circularly polarized PL emission also remain qualitatively the same when the orientation of linearly polarized PL excitation is changed from 0 – 60 degree.



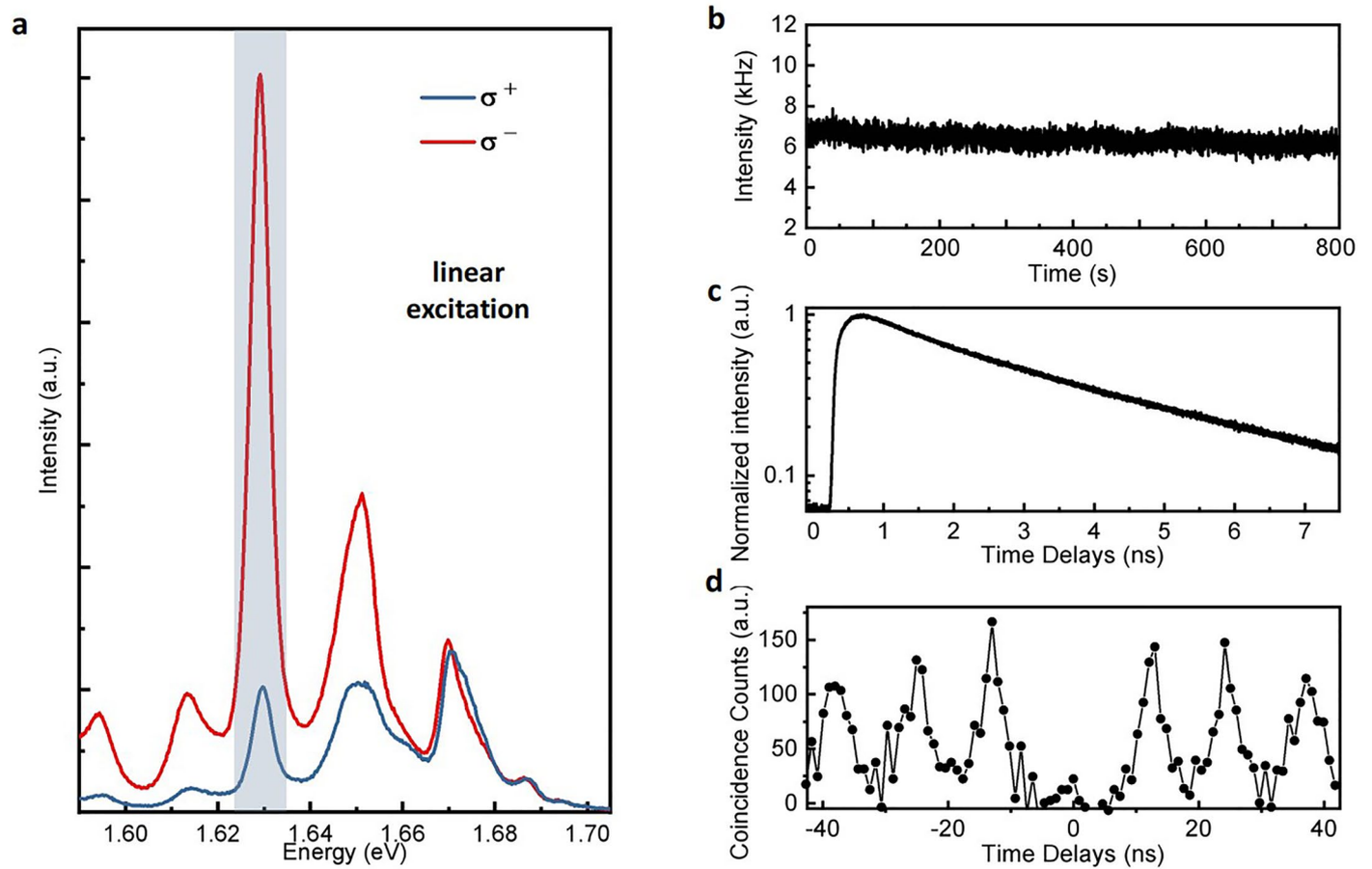
Extended Data Fig. 4 | Spectral image and analysis of the spatial distribution of PL intensity. **a.** PL spectral image of a nanoindentation in Fig. 1 a,b. σ^+ and σ^- PL emissions are simultaneously recorded on two different regions of the CCD array (**upper**). Degree of circular polarization (DoCP) spectrum calculated from integrated σ^+ and σ^- PL spectra (**lower**). **b.** Magnified spectra image of the region marked in (a). **c.** Blue data points: Intensity distribution of the localized emission peaks marked 1-5 in (b). By fitting the PL intensity distribution with Gaussian

function (black fitted curves), we can determine the centroid of the emission peak with ± 0.1 pixel precision. The position of each peak and fitting error is listed next to the peak. Based on the magnification factor of the microscope objective (50X) and pixel dimensions of the CCD (20 μm), we can estimate that all peaks, marked 1-5 originated from slightly different locations around the nano-indent. The separation between Peak 2 and 3 is ~ 50 nm.

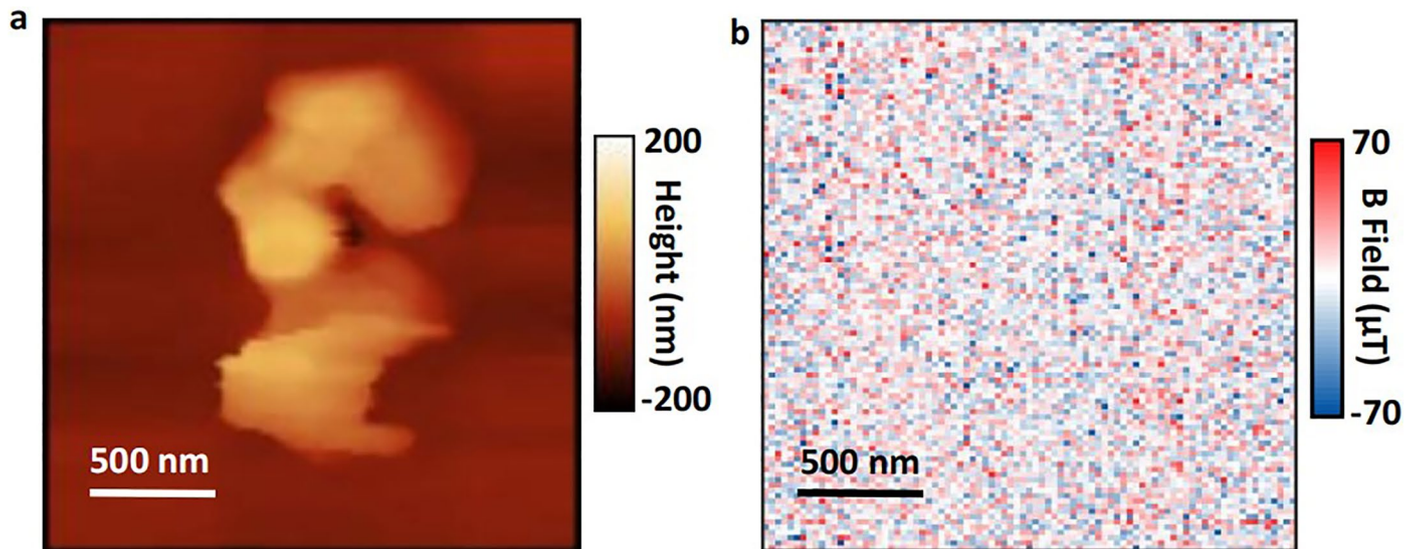


Extended Data Fig. 5 | Analyzing linear polarization of strain-engineered WSe₂/NiPS₃ heterostructure and distribution of linewidth of chiral quantum emitters and pump-dependent measurements. a, top panel. σ^+ / σ^- polarized PL spectra acquired under σ^+ excitation. **Inset**. Zoom-in view of the NiPS₃ anisotropic exciton peak displaying peak DCP of 0.4. **lower panels**. S and P linearly polarized PL spectra acquired at different polarization detection angles indicated in the panels. All three spectra are excited by σ^+ polarized laser in the same way as the top panel. **b**. DLP of NiPS₃ anisotropic exciton (red data points and trace of **top panel**) and PL peak 1-3 (**lower panels**) as the function of polarization detection angle. Black data points and trace plot DLP of NiPS₃

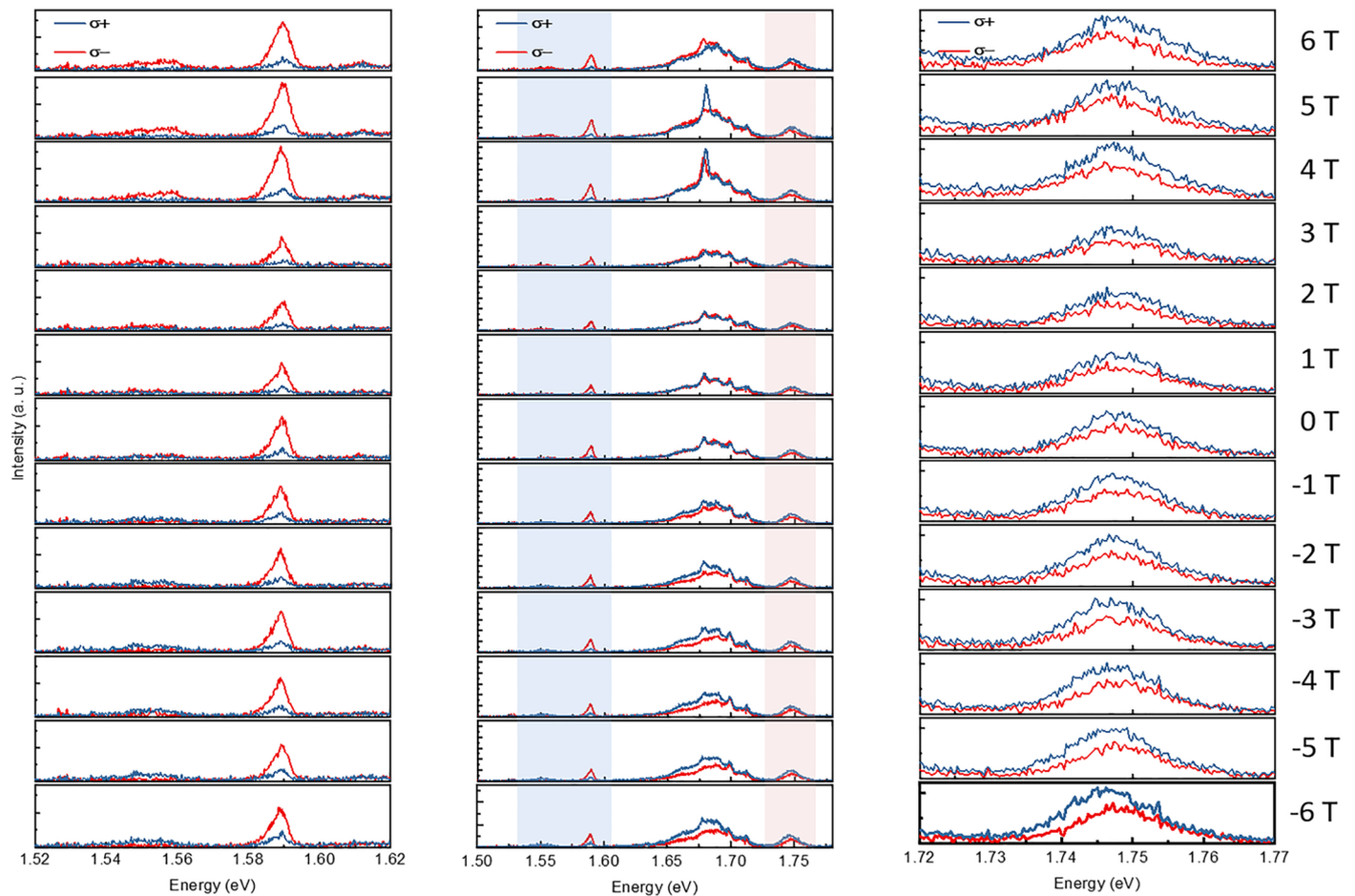
anisotropic excitons in unstrained NiPS₃. **c**. Histogram of the linewidths of chiral quantum light emitters yielding 3.6 ± 1.3 meV average and standard deviation, respectively. ~40% of the QEs display linewidth narrower than 3 meV. We believe the linewidth can be further minimized via encapsulation in hBN. Both linewidth and chirality show no notable dependence on laser excitation power. **d**. Pump-dependent study of a nanoindentation with two circularly polarized PL peaks. **e, f**. The plot of DCP and linewidth as the function of the pump displays a little change over 3 orders of magnitude change in pump power. The DCP of peaks a & b is calculated by taking the rise of the un-polarized PL background into account.



Extended Data Fig. 6 | Another Chiral Quantum Light Emitter showing -0.7 DCP and 96% single photon purity. **a.** Chiral PL spectra (σ^+ (blue) and σ^- (red)) of another indentation spot under linearly polarized laser excitation showing >70% of DCP. **b.** PL intensity time trace, **c.** PL decay curve, **d.** 2nd order photon correlation trace collected at 10 K showing a $g^{(2)}(0) = 0.04$.

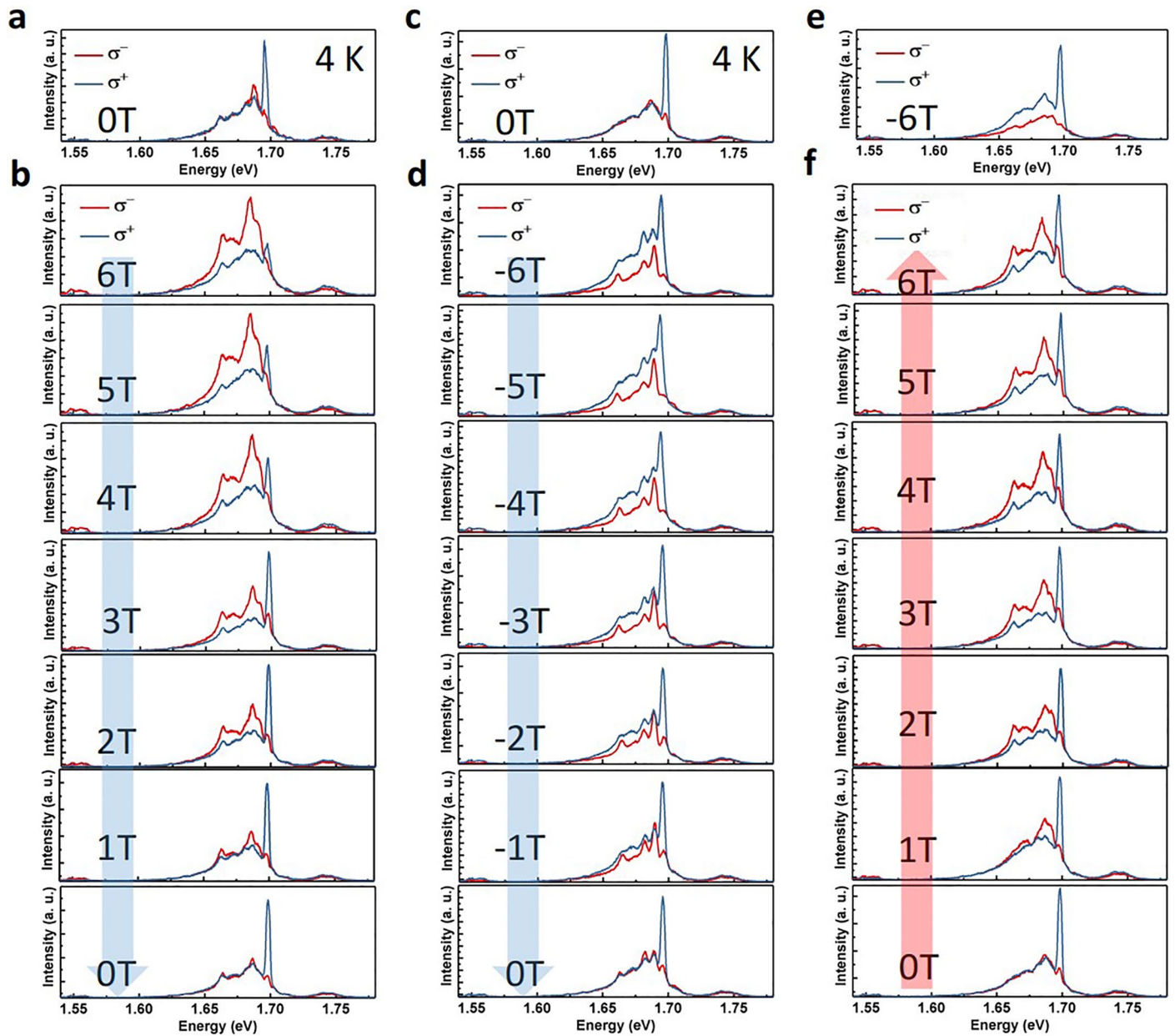


Extended Data Fig. 7 | Room temperature (RT) scanning diamond NV microscopy study of indented NiPS. **a.** Topographical image of the indentation shown in Fig. 3c. The RT scanning NV system utilized a slightly sharper NV probe yielding a higher resolution topographical image. **b.** Magnetic field image of the same area showing only the noise.

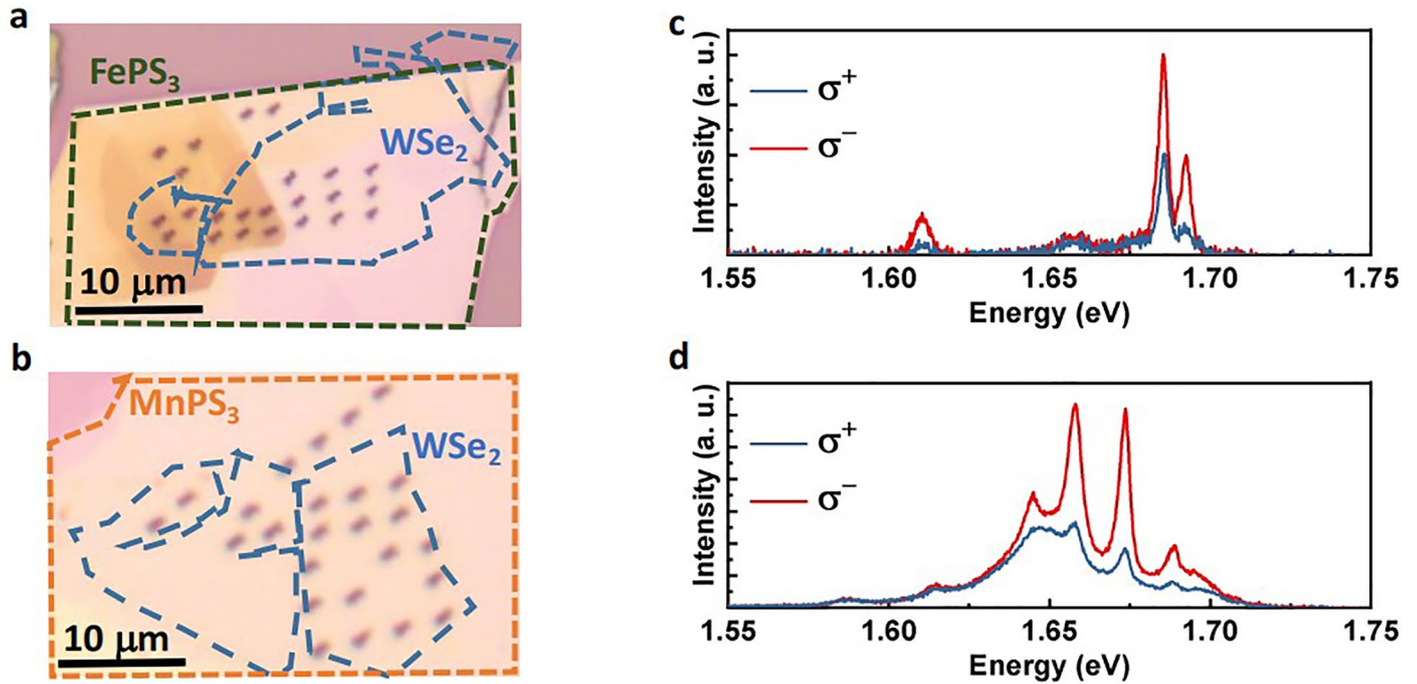


Extended Data Fig. 8 | Complete data set for Fig. 4a. Center panel. Low-temperature σ^+ and σ^- polarized PL spectra acquired under -6.0 T to +6.0 T external magnetic field applied perpendicular to the sample plane. **Left and right panels.** Zoom-in view of the gray and red spectra ranges of the center

panels. Because magneto-PL spectra were acquired under 633 nm σ^+ laser excitation, a weak DCP is observed at zero B field. A weak magnetic field-dependent change in DCP likely resulting from field-induced polarization of localized excitons is also observed in the 1.65 to 1.70 eV spectral range.



Extended Data Fig. 9 | Complete data set for Figs. 4c and 4d. a-b, c-d. Complete B field dependent σ^{\pm} polarized PL spectra shown in the bottom 3 panels of Fig. 4c and Fig. 4d. e-f. After chiral emission is fully restored at 0 T in b, PL spectra of the same location were collected by sweeping the field from 0 T to 6 T and then 6 T to -6 T.



Extended Data Fig. 10 | Observation of Chiral Localized Exciton Emission from WSe₂/FePS₃ and WSe₂/MnPS₃ heterostructures with nano-indents.
a-b. Optical images of the WSe₂/FePS₃ (a) and WSe₂/MnPS₃ (b) heterostructures

with nano-indents. c-d. σ⁺ and σ⁻ polarized PL spectra acquired under linearly polarized laser excitation at 4 K from two different nano-indents of WSe₂/FePS₃ (c) and WSe₂/MnPS₃ (d), respectively.