Spin-correlated exciton-polaritons in a van der Waals magnet

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Strong coupling between light and elementary excitations is emerging as a powerful tool to engineer the properties of solid-state systems. Spin-correlated excitations that couple strongly to optical cavities promise control over collective quantum phenomena such as magnetic phase transitions, but their suitable electronic resonances have yet to be found. Here we report strong light-matter coupling in NiPS3, a van der Waals antiferromagnet with highly correlated electronic degrees of freedom. A previously unobserved class of polaritonic quasiparticles emerges from the strong coupling between its spin-correlated excitons and the photons inside a microcavity. Detailed spectroscopic analysis in conjunction with a microscopic theory provides unique insights into the origin and interactions of these exotic magnetically coupled excitations. Our work introduces van der Waals magnets to the field of strong light-matter physics and provides a path towards the design and control of correlated electron systems via cavity quantum electrodynamics.

Strong light-matter coupling has recently emerged as an attractive platform to engineer quantum materials. When coherent optical feedback drives material excitations into a strong coupling regime, new hybrid quasiparticles known as polaritons determine the energy scales of the coupled system. Dressing specific collective modes, such as excitons or phonons with tailored optical fields is expected to cause selective modifications of material properties [1, 2]. Pioneering experiments using excitons in semiconductors embedded in optical cavities realized Bose–Einstein-like condensates, demonstrated fundamental phenomena, such as superfluidity, and spawned device concepts like polaritonic interferometers and Hamiltonian simulators [3, 4]. More recently, the prospects of strong light-matter coupling for manipulating collective phenomena like magnetism, superconductivity, and ferroelectricity in correlated materials has garnered much attention [5–7].

Among different classes of quantum materials, magnetic van der Waals (vdW) crystals provide access to a large variety of electronic and magnetic phases. Their strong electronic correlations, exotic magnetic orders, and potential for pressure-induced superconductivity currently motivates immense research efforts [8]. In addition to magnetic ordering, experiments demonstrated strong internal coupling of phonons and magnons [9]. Such intertwined degrees of freedom make these materials an ideal platform for exploring new aspects of tailored optical control. Particularly the recent discovery of optically active excitons with coupling to the antiferromagnetic order [1, 10, 11, 13] opens an extraordinary opportunity in this endeavor.

Here we demonstrate strong coupling between an optical microcavity mode and spin-correlated excitons hosted in the vdW magnetic insulator NiPS3. Hybridization results in a previously unobserved class of polaritons with unique signatures of excitons, photons, and spins. These newly formed quasiparticles are utilized as a probe to study the nature and interactions of spin-correlated excitons. We find that long-range excitonic interactions are severely suppressed owing to the tightly bound and highly localized nature of the excitons. By modeling excitonic transitions in strongly correlated insulators, we capture the key experimental signatures and demonstrate that spin-correlated excitons in NiPS3 have an origin that is distinct from that of excitons in conventional band semiconductors.

Within the class of vdW magnets, the family of transition metal thiophosphates (MPX3, with M being a transition metal and X a chalcogen) realizes different types of antiferromagnetism, such as Néel, stripy, and zig-zag phases [14]. NiPS3 is a correlated insulator that hosts zig-zag chains of ferromagnetically oriented Ni spins, which align antiferromagnetically inside each layer below the Néel temperature $T_N = 155$ K (see Fig. 1a) [15].

Our low-temperature photoluminescence (PL) measurements of NiPS3 crystals exfoliated onto standard SiO2/Si substrates (see Fig. 1b) show a narrow exciton emission peak at 1.476 eV, in agreement with recent studies [10, 11]. Due to their exceptional coherence, these excitons exhibit spectral widths as low as 350 µeV in high-resolution PL experiments [11]. Our theoretical analysis of strongly correlated electronic states in NiPS3 based on an extended Hubbard model (cf. Supplementary Section 1A) reveals excitonic transitions between S $p$-orbitals with zero net-magnetization and spin-polarized, long-range ordered Ni $d$-orbitals (cf. red ellipses in Fig. 1a). As a result, these excitons intrinsically couple to the magnetic order of Ni spins. Their highly anisotropic dipole moment predicted by our model (Supplementary Section 1C) is in excellent agreement with the large degree of linearly polarized PL emission (>80%) observed in our experiments (cf. inset of Fig. 1b).

The optical absorption spectrum derived from the lin-
ear reflectance contrast in Fig. 1c shows three distinct resonances close to the excitonic transition observed in PL spectra. The first resonance, $X_1$, coincides with the PL emission at 1.476 eV and is therefore ascribed to the absorption of these spin-correlated excitons [10]. Another absorption peak, $X_2$, at 1.479 eV, as well as a much broader feature, $X_3$, at 1.498 eV, were also reported in ref. [10]. Their line-shape and temperature dependence indicate that, like $X_1$, they are coupled to the long-range magnetic order and may be related to magnon modes [10, 16, 17]. Indeed, photo-excitation of these resonances with ultrashort laser pulses has recently led to the generation of coherent magnon oscillations [13].

As the temperature approaches $T_N$, all the excitonic features discussed above vanish (see Fig. S3, S6 and S10). This strong correlation between excitons and antiferromagnetism is also apparent in an external in-plane magnetic field, which simultaneously rotates the magnetic axis and the emitted photon polarization [1]. In its antiferromagnetic state, NiPS$_3$ thus presents an excellent opportunity for exploring strong light-matter coupling with spin-correlated excitons.

To realize strong coupling between these exotic quasi-particles and microcavity photons, we transfer NiPS$_3$ crystals onto dielectric bottom mirrors prior to evaporating a 35 nm-thin top silver mirror (see Methods and Fig. 2a). The thickness of each individual crystal directly determines the overlap with the photon cavity field. About one out of four NiPS$_3$ crystals exhibits a cavity resonance suitable for the observation of strong coupling in our experiments (cf. Fig. S4). Figure 2b displays an optical image of a NiPS$_3$ microcavity flake with lateral dimensions around 15 µm and a thickness of 170 nm determined by atomic force microscopy.

Figure 2c shows an angle-resolved optical reflectance contrast map measured at $T=4$ K. The photonic mode has multiple anti-crossing features that are characteristic of polariton formation via strong coupling of cavity photons and excitons. As indicated by the splittings, each absorption resonance ($X_1, X_2, X_3$) strongly couples to the optical cavity mode. We fitted the new modes using a coupled oscillator model (cf. Supplementary Section 6) and determined Rabi splitting energies $\hbar \Omega$ comparable to the values observed in standard band semiconductors [18]. However, the anti-crossing signature of strong coupling vanishes upon approaching $T_N$ (cf. Fig. 2d and Fig. S10), indicating a relation between the polaritons and long-range magnetic order.

First, we study the cavity PL emission obtained under non-resonant laser excitation at $T=4$ K. Angle-integrated spectra comprise a main peak at 1.476 eV, a pronounced high-energy shoulder at 1.478 eV and a broad emission
FIG. 2. Strong light-matter coupling in NiPS₃ microcavities. a NiPS₃ crystals enclosed by a dielectric bottom mirror and a 35 nm-thin top silver layer form a microcavity. Scale bar is 5 µm. b Optical microscope image of a single NiPS₃ crystal inside the cavity. c Angle-resolved optical reflection contrast map at 4K of the same crystal plotted together with a coupled oscillator model of a single cavity mode and absorption resonances X₁, X₂ and X₃ for negative angles (cf. also reflectance map and profile analysis in Fig. S7). Anti-crossings at each intersection between the cavity mode (white dashed line) and an excitonic resonance (black dash-dotted lines) are reproduced with good agreement between the model and the experimental data. The resulting multiple branches of the polariton dispersion are depicted as blue solid lines. The Rabi splittings indicated by gray circles are ℏΩ₁ = 4 meV, ℏΩ₂ = 2 meV, and ℏΩ₃ = 10 meV, as obtained from fits. d Anti-crossing features are almost absent in the reflectance contrast map recorded at 120K. e Integrated PL intensity of X₁ as a function of analyzer angle. f Angle-integrated PL emission shows strong linear polarization of more than 50% and pronounced peaks at the X₁ and X₂ absorption resonances. Blue and grey spectra represent 90 deg and 0 deg analyzer angles, respectively.

band around 1.46 eV (cf. Fig. 2f). The linear polarization observed in Fig. 2e shows that the antiferromagnetic order of NiPS₃ is preserved in our strongly coupled cavity sample. Like in the bare flakes on SiO₂/Si, the degree of polarization varies from sample to sample.

We gain valuable insight into the nature of the resulting exciton-polaritons from angle-resolved emission maps. Unlike in most exciton-polariton systems [3, 19], the strongest PL signal of our NiPS₃ cavity is observed at energies close to the X₁ and X₂ exciton resonances (see Fig. 3a,b) and only a small fraction of the total polariton emission occurs from the lowest energy states of the polariton dispersion. This behavior indicates the presence of a pronounced polariton relaxation bottleneck towards the bottom of the dispersion at 1.46 eV, and thus weak polariton scattering processes and interactions. Despite matching with phonon energies found in recent Raman studies (ΔE ≈ 10–20 meV) [20], exciton scattering rates are not sufficient to populate these low-energy polariton states during the short polariton lifetime [21].

Even in the absence of efficient polariton relaxation via phonons, long-range exciton-exciton exchange interactions typically overcome the relaxation bottleneck and facilitate majority population of low-lying polariton states [3]. However, the observed weak polariton relaxation indicates that these interactions are greatly suppressed in NiPS₃. As shown in Fig. 3c,d, only the enhanced phonon scattering at elevated temperatures is able to reduce the relaxation bottleneck and thus increases the fraction of PL observed from low-energy polariton states around 1.46 eV.

We also perform polariton excitation spectroscopy – a
FIG. 3. Pronounced bottleneck of polariton relaxation. a Angle-resolved PL emission recorded at 4K superimposed by the coupled oscillator model shown in Fig. 2c. b Schematic illustration of polariton relaxation and the bottleneck effect: Excitons from the reservoir scatter with phonons to weakly populate the lowest polariton branch, while efficient scattering due to long-range exciton-exciton interactions is suppressed. c Cavity PL emission at 60K. Intensity-scale is indicated in a by the number given in brackets. d Comparison of angle-integrated normalized PL spectra at 4K and 60K (intensity spikes were removed).

FIG. 4. Polariton non-linearities under increasing density. a Polariton dispersion for the lowest ($P_0$) and highest ($\approx 360P_0$) excitation power of the incident broadband laser pulses. Inset shows a magnified view of the fitted oscillator model for both powers. b Density-dependent exciton energies directly determined from absorption minima in line-cuts at 0° ($X_1$ and $X_2$) and at 20° ($X_3$) with high exciton fraction. c-e Reduction of Rabi splitting $\Omega(n_p)$ relative to the Rabi splitting $\Omega_0$ obtained at the lowest excitation power. Red solid lines represent a fit based on Eq. (1) to determine the saturation density.
powerful technique highly susceptible to non-linear optical phenomena [2, 3, 22] – to further study the interactions of spin-correlated excitons (see Methods). As shown in Fig. 4a, tuning the power density of the incident broadband laser pulses allows us to measure the microcavity dispersion under increasing excitation density. Since the excitonic fraction is dispersive, exciton energies can be obtained from absorption minima in line-cuts taken at angles at which the exciton fraction is close to unity (see Fig. S9). Additionally, Rabi splitting energies can be determined from fits to the dispersion in the anti-crossing region.

Due to repulsive long-range interactions and the Pauli exclusion principle, polaritons formed by delocalized excitons in band semiconductors typically respond to increasing density by shifting towards higher energies [2, 22]. However, for the three exciton resonances plotted in Fig. 4b, no significant shifts are noticeable despite the high polariton densities ($n_{P} \approx 5 \times 10^{12} \text{cm}^{-2}$) reached in our experiment. Very similar behavior has recently been observed for localized moiré excitons in 2D vdW heterostructures [2]. In excellent agreement with our microscopic model presented in Fig. 1a and the conclusions drawn from Fig. 3, excitons in NiPS$_3$ are thus characterized by the suppressed long-range interactions of strongly localized excitons.

Despite the absence of intrinsic dispersive exciton nonlinearities, strong coupling introduces a collective nonlinear response in the electronic dispersion of our NiPS$_3$ microcavity. Saturation of optical absorption modifies the dispersion due to a gradual reduction of Rabi splitting energies $\Omega(n_{P})$ under increasing polariton density $n_{P}$ (cf. Fig. 4c-e). While the negatively detuned cavity analyzed in Fig. 4 indicates no noticeable saturation for $\Omega_3$, higher total polariton density reached in a near zero-detuned cavity reveals its saturation at high powers (cf. Figs. S12 and S13). The large saturation density ($n_{sa}$) of $X_1$ and $X_2$ exceeding $10^{13} \text{cm}^{-2}$ indicates that excitons in NiPS$_3$ are strongly localized and tightly bound by attractive Coulomb forces, despite the pronounced electrostatic screening effects inherent to bulk crystals. As expected from our microscopic model presented in Fig. 1a, the average Bohr radius of $X_1$ and $X_2$ excitons in NiPS$_3$ estimated from their optical saturation ($a_B = (6 \pm 1) \text{Å}$, see Supplementary Section 10) is comparable to the order of the size of the unit cell [27]. This conclusion is also in excellent agreement with recent first-principles calculations [24] and the large effective masses predicted in ref. [25] resulting in Bohr radii well below 1 nm. The apparent inconsistency between the large exciton binding energies (several hundred meV) and the fact that spin-correlated excitons vanish around $T = T_N$ highlights the distinct origin of these magnetically coupled excitations in NiPS$_3$ from excitons in conventional band semiconductors. Despite their exceptionally small Bohr radii, the modest Rabi splitting energies observed in our microcavities further demonstrate the presence of a relatively small exciton oscillator strength in NiPS$_3$ (see Supplementary Section S9). Finally, we emphasize that pronounced electronic correlations in NiPS$_3$ [15] are responsible for the strong exciton localization supported by the results in Figs. 3 and 4.

In summary, we demonstrate strong coupling between microcavity photons and spin-correlated excitons in the vdW magnetic insulator NiPS$_3$. Hybridization leads to a new type of quasiparticle which inherits signatures of excitons, photons, and spins. A detailed experimental and theoretical analysis of these polaritons demonstrates the absence of typical long-range exciton-exciton interactions to result from the strong localization of tightly bound excitons. Future studies could explore the potential of spin-correlated polaritons and their coupling to magnons for optical control of the high-speed information processing demonstrated in antiferromagnets [26]. Furthermore, with the ability of spin-correlated excitons to strongly couple to a microcavity photon field emerges the unique opportunity of realizing polariton Bose-Einstein condensates interacting with magnetically ordered spin systems. Finally, our experiments on correlated vdW antiferromagnets represent a first step towards the cavity-induced optical engineering of strongly correlated quantum materials.

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Data availability: The data sets generated during and/or analysed during the current study are available from the corresponding authors on reasonable request.

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**METHODS**

Sample fabrication and characterization

For microcavity fabrication, highly reflective Bragg mirrors (from 1.39 eV to 1.62 eV) were grown by plasma-enhanced chemical vapor deposition of 30 pairs of Silicon nitride/Silicon dioxide layers on Silicon wafers. Commercial NiPS3 bulk crystals (acquired from 2D Semiconductors) were thinned down in multiple cycles of stick-and-release on blue tape (PVC tape 224PR, Nitto) and transferred onto a polydimethylsiloxane (PDMS, AD se-ries, Gel-Pak) film. After moderately pressing the PDMS film onto a Bragg mirror (or a SiO2/Si substrate, oxide thickness 285 nm) and swiftly releasing it, the majority of crystals that were transferred showed a thickness of a few tens to hundreds of nanometers. Subsequent to NiPS3 transfer, a 35 nm-thin layer of silver was deposited by electron-beam physical vapor deposition to form a top mirror, resulting in measured quality factors of around 100. Within 24 hours after transfer, the samples were cooled down to T=4 K and kept inside the high-vacuum...
chamber of a dry-flow cryostat (Montana Systems).

**Optical spectroscopy**

Reflectance spectra and maps were obtained from the attenuated output of a spectrally broadband tungsten-halogen lamp (250W LSH-T250, Horiba), spatially filtered, and focused to a spot size of 2.0 µm by the 100 × microscope objective (NA=0.66) mounted inside the cryostat. The reflected signal was analyzed by a linear polarizer along the axis of maximum PL intensity. Angle-integrated and angle-resolved measurements were recorded by respectively focusing either real-space images or the back-focal plane of the objective onto a spectrometer connected to a charge-coupled device (CCD). Polariton dispersions were fitted by varying the vacuum Rabi splitting and the exciton–photon detuning while keeping the exciton energy and the effective refractive index of NiPS$_3$ constant.

For the PL experiments, unless specified otherwise, a continuous-wave laser with 2.33 eV energy and $P = 120$ kW/cm$^2$ average output power was focused onto the sample to a spot size of 1.0 µm by the same objective used for reflectance spectroscopy. The collected PL signal was directed towards the spectrometer and spectrally filtered to remove the laser emission. For polarization-resolved measurements, stationary linearly polarized laser excitation was used to obtain PL emission, which was subsequently passed through a rotating half-waveplate and a polarizer fixed along the direction parallel to the entrance slit of the spectrometer. The degree of linear polarization is thus defined as $\rho = (I_\perp - I_\parallel)/(I_\perp + I_\parallel)$, where $I_\perp$ ($I_\parallel$) denotes the PL intensity when the emission polarization is analyzed perpendicular (parallel) to the antiferromagnetic spin orientation [1].

**Nonlinear polariton spectroscopy**

For nonlinear spectroscopy, spectrally broad pulses from a super-continuum laser (NKT Photonics, SuperK Extreme) with a repetition rate of 78 MHz and a pulse width of 20 ps were filtered by a bandpass filter from 1.41 eV to 1.52 eV to minimize absorption at higher energies and the resulting heating effects. The time-averaged laser power of the spectrally filtered pulses was increased from 0.07 mW to 25.3 mW, while the reflected signal was attenuated by optical density filters before entering the detector. Polariton branches were monitored for different input powers by mapping the numerically determined reflectance minima at different angles. The resulting dispersion plots were then fitted in a small angle range around the anti-crossing region for each laser power with a coupled oscillator model to determine the Rabi splitting energies.

Under increasing laser-induced polariton density $n_p$, saturation of optical absorption modifies the dispersion due to a gradual reduction of Rabi splitting energies $\Omega(n_p)$. The decrease with respect to the Rabi splitting obtained at the lowest density, $\Omega_0$, is given by [2]

$$\frac{\Omega(n_p)}{\Omega_0} = \frac{1}{\sqrt{1 + \frac{n_p}{n_s}}}.$$  

Fitting Eq. (1) to the experimental data yields a saturation density $n_s$ for each set of polariton branches (see Supplementary Section S9).

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