ARTICLE

Exciton-polarons in the presence of strongly correlated electronic states in a MoSe₂/WSe₂ moiré superlattice

Aidan J. Campbell, Mauro Brotons-Gisbert, Hyeonjun Baek, Valerio Vitale, Takashi Taniguchi, Kenji Watanabe, Johannes Lischner and Brian D. Gerardot

Two-dimensional (2D) materials have emerged as a new playground to investigate strongly correlated electronic phenomena. For example, due to a direct bandgap, huge exciton binding energies, and straightforward control of carrier concentration, monolayer transition metal dichalcogenides (TMDs) provide a platform to probe the interaction of an exciton with a Fermi sea (2D electron or hole gas) described by the Fermi-polaron model. With increasing Fermi energy, a neutral exciton evolves into two branches due to both attractive and repulsive interactions with charge carriers. By extension, TMD moiré heterostructures provide access to a highly tunable many-body physical system consisting of an exciton dressed by a Fermi sea which forms a series of charge-ordered (Mott insulating and generalised Wigner crystal) electronic states as the carrier concentration is tuned. Stacking two monolayer TMDs with either a lattice mismatch and/or relative twist angle forms a moiré superlattice with a periodicity that far exceeds the inter-atomic spacing of the constituent crystals. Itinerant electrons in a Fermi sea can be spatially localised by the moiré superlattice of a molybdenum diselenide (MoSe₂)/tungsten diselenide (WSe₂) twisted hetero-bilayer. At a multitude of fractional fillings of the moiré lattice, we observe ordering of both electrons and holes into stable correlated electronic states. Magneto-optical measurements reveal extraordinary Zeeman splittings of the exciton-polarons due to exchange interactions in the correlated hole phases, with a maximum close to the correlated state at one hole per site. The temperature dependence of the Zeeman splitting reveals antiferromagnetic ordering of the correlated holes across a wide range of fractional fillings. Our results illustrate the nature of exciton-polarons in the presence of strongly correlated electronic states and reveal the rich potential of the MoSe₂/WSe₂ platform for investigations of Fermi–Hubbard and Bose–Hubbard physics.

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INTRODUCTION

Two-dimensional (2D) materials have emerged as a new playground to investigate many-body interactions and strongly correlated electronic phenomena. For example, due to a direct bandgap, huge exciton binding energies, and straightforward control of carrier concentration, monolayer transition metal dichalcogenides (TMDs) provide a platform to probe the interaction of an exciton with a Fermi sea (2D electron or hole gas) described by the Fermi-polaron model. With increasing Fermi energy, a neutral exciton evolves into two branches due to both attractive (lower energy) and repulsive (higher energy) interactions with charge carriers. By extension, TMD moiré heterostructures provide access to a highly tunable many-body physical system consisting of an exciton dressed by a Fermi sea which forms a series of charge-ordered (Mott insulating and generalised Wigner crystal) electronic states as the carrier concentration is tuned. Stacking two monolayer TMDs with either a lattice mismatch and/or relative twist angle forms a moiré superlattice with a periodicity that far exceeds the inter-atomic spacing of the constituent crystals. Itinerant electrons in a Fermi sea can be spatially localised by the moiré superlattice of a molybdenum diselenide (MoSe₂)/tungsten diselenide (WSe₂) twisted hetero-bilayer. At a multitude of fractional fillings of the moiré lattice, we observe ordering of both electrons and holes into stable correlated electronic states. Magneto-optical measurements reveal extraordinary Zeeman splittings of the exciton-polarons due to exchange interactions in the correlated hole phases, with a maximum close to the correlated state at one hole per site. The temperature dependence of the Zeeman splitting reveals antiferromagnetic ordering of the correlated holes across a wide range of fractional fillings. Our results illustrate the nature of exciton-polarons in the presence of strongly correlated electronic states and reveal the rich potential of the MoSe₂/WSe₂ platform for investigations of Fermi–Hubbard and Bose–Hubbard physics.
generalised Wigner crystals. At \( v = -1 \) we observe that the repulsive WSe\(_2\) exciton-polaron gains oscillator strength from the attractive, due to the reduced screening of the exciton by carriers in the insulating state. After \( v = -1 \), the oscillator strength is then abruptly transferred to the attractive polaron. Furthermore, we observe filling-factor-dependent \( g \)-factors of these positively-charged attractive and repulsive polarons, with a maximum at \(-1\) hole per site. Under the assumption that the \( g \)-factors are proportional to the magnetic susceptibility of the correlated phases induced by exchange interactions, temperature dependent measurements for \( v = -0.7 \) to \(-1.3\) reveal an antiferromagnetic spin coupling of the moiré pinned Fermi-hole sea. The experimental magnetic behaviour is theoretically explored using a model that solves the Heisenberg Hamiltonian for charged-ordered hole states with antiferromagnetic next-neighbour interactions. Our results highlight the behaviour of excitons dressed by a Fermi sea which is spatially ordered in a series of correlated states. This has importance for optical studies of correlated electronic phenomena in 2D materials and demonstrates the potential of MoSe\(_2\)/WSe\(_2\) heterostructures for future investigations and implementations of highly tunable 2D Fermi–Hubbard or Bose–Hubbard models.

RESULTS

Device structure and Fermi-polarons

Figure 1a shows a sketch of our dual-gated hetero-bilayer device, consisting of a monolayer MoSe\(_2\) and a monolayer WSe\(_2\) vertically stacked with a twist angle (\( \Delta \theta \)) of \(-57^\circ\). The relative twist angle from perfect H stacking (i.e., \( \Delta \theta = 60^\circ \)), estimated from the optical micrograph of the hetero-bilayer (see Methods) and confirmed by our gate dependent measurements (described later), is beyond the proposed critical angle for lattice reconstruction. The hetero-bilayer was encapsulated by hexagonal boron nitride (hBN) layers with nearly identical thicknesses (~18 nm). Graphene layers act as electrical contacts for the top, bottom and hetero-bilayer gates (see ref. 34 for more details). Moreover, the combination of the layer twist and the lattice mismatch between MoSe\(_2\) and WSe\(_2\) results in the formation of a triangular moiré superlattice in our device (see sketch in Fig. 1b) with a period of ~6 nm. This causes a periodic variation in the interlayer hopping that results in a flattening of the conduction and valence bands in the type-II band structure characteristic of TMD hetero-bilayers (see Fig. 1c) [26,43].

The bare intralayer excitons of the constituent TMDs can be probed via absorption spectroscopy. The moiré lattice carrier concentration is tuned via the application of a gate voltage \( (V_g) \) between the top/bottom graphene contacts and the hetero-bilayer. As depicted by the schematic in Fig. 1b, these carriers are spatially ordered in a series of correlated states by the moiré superlattice, whilst also dressing the photo-excited intralayer exciton to form attractive and repulsive exciton-polaron complexes (as shown in Fig. 1c).

Correlated electronic states

To investigate doping-dependent phenomena, we perform differential-reflection contrast \((\Delta R/R_0)\) spectroscopy as a function of \( V_g \), where \( \Delta R = R_s - R_0 \) and \( R_0 \) is the intensity of the light reflected by the hetero-bilayer (substrate). Figure 2a shows the \( V_g \) dependence of the first derivative of the reflectance spectra with respect to photon energy \((d(\Delta R/R_0)/dE)\). The doping dependence of the intralayer exciton-polarons in the MoSe\(_2\)/WSe\(_2\) hetero-bilayer is markedly different to that observed for individual monolayers (see Supplementary Fig. 1). At charge neutrality, we observe three excitonic resonances: \( X_{0}^{W} \) at high energy, and two resonances separated by 36 meV in the spectral range corresponding to \( X_{0}^{Mo} \) which we label as \( X_{0,1}^{Mo} \) (low energy) and \( X_{0,2}^{Mo} \) (high energy). We assign the two MoSe\(_2\) peaks to be a consequence of the formation of moiré minibands, arising from the band folding at the edges of the reduced Brillouin zone. Furthermore, with increasing electron (hole) doping \( X_{0}^{W} \) \( (X_{0}^{Mo}) \) dominate the spectrum, as expected for a type-II band alignment.

We employ the parallel plate capacitor model to estimate the dependence of the nominal carrier concentration \( n \) on the applied \( V_g \). Using the density of moiré sites \( n_0 \) corresponding to \( \Delta \theta = 57^\circ \), we estimate the \( V_g \)-dependent nominal fractional filling \( v = n/n_0 \) of the moiré lattice (see Methods). The excitonic features shown in Fig. 2a exhibit strong modulations in their transition energy, linewidth and oscillator strengths for applied voltages close to the nominal \( V_g \) values corresponding to \( v = 0 \) and \pm 1\). Similar modulations of the excitonic transitions, observed in WSe\(_2\)/WS\(_2\) hetero-bilayers [17,20,21,46], have been attributed to the suppressed charge screening originating from the formation of correlated insulator phases at different fractional fillings of the moiré superlattice. In order to corroborate the presence of a robust moiré lattice at the same spatial position in our sample where we observe strongly correlated states, we measure the low-temperature (4 K) photoluminescence (PL) spectrum at charge neutrality using confocal spectroscopy, revealing a series of...
discrete peaks with narrow line-widths (<100 μeV) that demonstrate the existence of an underlying moiré lattice responsible for the interlayer exciton trapping32–34 (see Supplementary Fig. 2).

Moreover, Fig. 2a also reveals that each monolayer in the WSe2/MoSe2 hetero-bilayer is capable of sensing the doping-induced changes in their dielectric environment originating from the fractional filling of the other layer, similar to the effects observed using a WSe2 sensor layer in proximity to a WSe2/WS2 heterostructure21. Figure 2b shows an example of the sensing capabilities of the MoSe2 layer for hole doping of the WSe2 layer: the transition energies of \( \chi^0_{\text{Mo},1} \) and \( \chi^0_{\text{Mo},2} \) blue-shift and peak at \( \Delta V_g = -1.34 \) V, consistent with a decrease in the permittivity of the heterostructure arising from the formation of a correlated insulating state at 1 hole per moiré site in the WSe2 layer17,20. In addition to the modulation in the transition energy, the linewidth of \( \chi^0_{\text{Mo},1} \) also presents a clear minimum at \( v = -1 \) (see Fig. 2c), which can be understood as the result of reduced charge disorder originating from a correlated insulating state19. These results demonstrate the potential of intralayer excitons as sensors that can probe the formation of correlated states in the adjacent layer (see sketch in Fig. 2d) and corroborate the calibration of \( v = \pm 1 \) in our device. To estimate the \( V_g \) values corresponding to other fractional fillings of the moiré lattice, we assume a linear dependence of \( v \) with \( V_g \) and extrapolate from the experimental \( V_g \) values determined for one hole/electron per site, as shown in the right panel of Fig. 2e. To increase the sensitivity to doping-induced modulations of the reflectance signal, we plot the first derivative of \( \Delta R/R_0 \) as a function of \( V_g \) (see left panel of Fig. 2e). The d(\( \Delta R/R_0 \))/d\( V_g \) spectrum highlights a series of abrupt changes in the reflected signal at \( v = 0, \pm 1/3, \pm 1/2, \pm 2/3, \pm 1, -5/4 \) (as indicated by the horizontal lines in Fig. 2e), suggesting the formation of correlated states at these fractional fillings of the triangular lattice. These results reveal symmetric loading of carriers, with an identical \( \Delta V_g = \pm 1.34 \) V required to fill the moiré superlattice with either one electron (\( V_g = 1.34 \) V) or one hole (\( V_g = -1.34 \) V) per site, respectively. We
assign the stable phases at $\nu = \pm 1$ to be either Mott $^{17,18,21}$ or charge-transfer $^{16}$ insulator states and the remaining states to be generalised Wigner crystals $^{13,18,20,21,27}$.

To gain deeper insight into the strength of the electronic correlations in our system, we investigate the melting temperature of the different correlated states. Supplementary Fig. 9 shows the dependence of $d(\Delta R/R_0)/dV_g$ on $V_g$ for temperatures ranging from 4 K to 90 K. With increasing temperature, the abrupt changes in the $d(\Delta R/R_0)/dV_g$ spectrum (indicative of correlated state formation in both the electron and hole doping regimes, see Fig. 2e) progressively smooth out until they can no longer be observed at 90 K. We quantitatively estimate a melting temperature of $\sim 55$ K for the correlated state at one hole per moiré site (see Supplementary Fig. 10).

Exciton-polaron behaviour at one hole per site

We now investigate the behaviour of the WSe$_2$ exciton-polarons as the hole fractional filling of the moiré superlattice is tuned. Figure 3a, b shows the $\sigma^-$ and $\sigma^+$-helicity-resolved evolution of the $d(\Delta R/R_0)/dE$ spectrum, respectively, for negative $V_g$ under an applied magnetic field $B$ of 5 T in Faraday configuration. The intensity colour scale in these figures is saturated to improve the visibility of the WSe$_2$ intralayer repulsive and attractive exciton-polarons (labelled RP$_W^+$ and AP$_W^+$, respectively), while the application of a magnetic field breaks the energy degeneracy between the exciton transitions at $\pm K$, helping to disentangle the behaviour of each excitonic species. As for the monolayer case, at $V_g = 0$ V only the neutral exciton resonance which becomes RP$_W^+$ is present. As the hole fractional filling increases, an additional resonance gains oscillator strength at $\sim 10$ meV lower energy, in agreement with the formation of AP$_W^+$. $^{47-49}$

The helicity-resolved results in Fig. 3a, b reveal additional features of the RP$_W^+$ and AP$_W^+$ complexes for fractional hole filling. First, hole doping results in a larger blue-shift of RP$_W^+$ compared to AP$_W^+$, as also observed for excitons interacting with a 2D fermionic sea in ML TMDs $^{4,6,50}$. Second, the oscillator strength of the RP$_W^+$ resonance shows a non-monotonic behaviour: for small gate voltages it decreases with increasing hole doping, which can be understood as a progressive transfer of oscillator strength from the neutral exciton to the positive trion-like state as the Fermi energy moves deeper into the valence band. However, the RP$_W^+$ resonance regains oscillator strength for hole doping levels corresponding to $\nu = -1$. This can be attributed to the suppressed charge screening in the correlated insulating phase forming in the moiré lattice. For further hole doping ($\nu < -1$), RP$_W^+$ abruptly quenches as the oscillator strength transfers to AP$_W^+$. Third, upon hole doping the $\sigma^-$-polarised transitions of both RP$_W^+$ and AP$_W^+$ appear at a higher energy than their respective $\sigma^+$-polarised transitions, indicative of a positive Zeeman splitting $\Delta E$ (according to the convention based on $\Delta E = E^{\sigma^+} - E^{\sigma^-}$, with $E^{\sigma^\pm}$ the energy of the transition with $\sigma^\pm$ polarisation). This behaviour contrasts with the negative $g$-factor of exciton-polarons in ML WSe$_2$ based on their spin and valley configurations $^{51}$.

To gain insight into the origin of the positive Zeeman splitting, we investigate the energies of RP$_W^+$ and AP$_W^+$ as a function of the applied $B$ field for $\nu = -1$. Figure 3c, d shows the $\sigma^-$- and $\sigma^+$-helicity-resolved evolution of the $d(\Delta R/R_0)/dE$ spectrum for applied $B$ fields...
between −2.5 and 2.5 T at ν = −1. Figure 3e shows linecuts of the bare ∆R/RO spectrum at different B fields for σ− (red dots) and σ+ polarised (blue dots) collection while the solid lines represent fits from which we estimate the energy, linewidth, and oscillator strength of both RPW and APW as function of the applied magnetic field (see the Supplementary text for a detailed description of the fitting procedure). The B-field-dependent estimated energies of RPW and APW extracted from the fits are overlayed in Fig. 3c, d (purple and black dots, respectively). A large positive Zeeman splitting is clearly observed, which suggests an interaction-enhanced magnetic response of the correlated hole state at ν = −1. Finally, we note that Fig. 3a–e reveal a large spin polarisation of RPW and APW under applied B fields. Such spin polarisation originates from the different effective hole doping in the ±K valleys induced by the large Zeeman splitting. APW is an intervalley exciton complex in which the momentum-direct electron-hole pairs at ±K are dressed by holes in the opposite valley52,53. On application of a positive magnetic field of 5 T, the valence band edge at +K is shifted to higher energy relative to −K. Exchange interactions favour single valley occupancy of carriers, so nearly all the holes are doped in the +K valley, rather than the −K, leading to the observed spin polarisation6. When we probe the exciton in −K using σ− light, there is a large population of holes available to form APW (see Fig. 3a). In contrast, there is a smaller population of holes at −K available to form APW when we probe the +K exciton using σ+, leading to a higher relative intensity of RPW (see Fig. 3b). When we sweep the magnetic field from positive to negative, we shift the band maximum from +K to −K, leading to the observed transfer in oscillator strength between RPW and APW (see Fig. 3c–e).

Magnetic interactions probed by exciton-polarons
Next, we investigate the fractional-filling-dependence of the Zeeman splitting of the RPW and APW resonances. Figure 4a

![Fig. 4 Exciton-polaron magnetic interactions at different hole fractional fillings. a σ− (blue) and σ+ resolved (red) ∆R/RO spectra at representative hole ν values under B = 1 T. b B-field-dependent Zeeman splitting of APW from −1 to 1 T at representative hole ν values. c Evolution of the g-factor of APW as a function of the hole ν extracted from linear fits of ∆E in the range |B| ≤ 1 T (solid lines in panel b). d Valley Zeeman splitting of the APW resonance at ν = −1 for different temperatures. e Evolution of the measured g-factor of APW as a function of temperature for ν = −1 in the temperature range for which the oscillator strength and linewidth of APW are sufficient to enable a reliable estimate of the Zeeman splitting. The red solid line represents a fit of the experimental data (black dots) to a Curie–Weiss law from which the Weiss constant is estimated to be θ = −4.6 ± 0.9 K. The negative sign of the extracted Weiss constant reveals antiferromagnetic ordering of neighbouring hole spins. f Theoretical prediction of the ν dependence of the g-factor based on a model that solves the Heisenberg Hamiltonian for charge-ordered hole states with antiferromagnetic exchange interactions between nearest neighbour spins. For all panels the error bars represent 68% confidence intervals.](image-url)
shows $\sigma^-$ (blue) and $\sigma^+$-resolved (red) $\Delta R/R_o$ spectra at representative hole $\nu$ values at $B = 1$ T. The dots represent experimental data while the solid lines are fits of the experimental $\Delta R/R_o$ to the model described in the Supplementary text, from which we estimate the energy of the resonances. The spectra in Fig. 4a reveal a clear positive $\Delta \nu$ for all hole doping levels, although with a magnitude that depends strongly on $\nu$. Figure 4b shows the $B$-field-dependent Zeeman splitting of $AP^\nu_1$ from $-1$ to $1$ T at representative hole $\nu$ values. The estimated Zeeman splitting exhibits a linear dependence with $B$ at small fields (i.e., $|B| < 1$ T). We note that $RP^\nu_1$ shows a similar positive linear dependence with $B$ at small fields, although it saturates at larger $B$ (see Supplementary Fig. 6). The linear evolution of $\Delta \nu$ at small $B$ can be associated to an effective exciton valley $g$-factor according to $\Delta \nu(B) = g\mu_B B$, where $\mu_B$ is the Bohr magneton. Figure 4c shows the evolution of the $g$-factor of $AP^\nu_1$ as a function of the hole $\nu$ extracted from linear fits of $\Delta \nu$ in the range $|B| < 1$ T (solid lines in Fig. 4b). As already inferred from the results in Fig. 4b, the $g$-factor of $AP^\nu_1$ shown in Fig. 4c exhibits a strong dependence on $\nu$, peaking around $\nu = -1$, where it reaches a maximum value of $g = 145$.

Figure 4d shows the Zeeman splitting of $AP^\nu_1$ measured for $|B| \leq 1$ T at $\nu = -1$ for different temperatures, where we observe the slope of the Zeeman splitting (and therefore the $g$-factor) decreases with increasing temperature. Figure 4e shows the evolution of the measured $g$-factor of $AP^\nu_1$ as a function of temperature for $\nu = -1$ in the temperature range in which the oscillator strength and linewidth of $AP^\nu_1$ enable a reliable estimate. We observe that the $g$-factor decreases by a factor $\sim 5$ when the temperature increases from 4 to 39 K. We assume that the interaction-induced enhancement of the attractive polaron $g$-factor is proportional to the magnetic susceptibility of the correlated states (e.g., $\chi \propto g$) and observe that the decrease of $g$-factor with increasing temperature follows a Curie–Weiss law $\chi^{-1} \propto T - \theta$ (red solid line in Fig. 4e), with $T$ being the temperature and $\theta$ the Weiss constant. From the fit in Fig. 4e we estimate a Weiss constant of $\theta = -4.6 \pm 0.9$ K, which suggests an antiferromagnetic behaviour of the interactions between the localised hole moments for $\nu = -1$. Supplementary Fig. 11 shows the temperature dependence of the $AP^\nu_1$ $g$-factor for a range of hole filling factors from $\nu = -0.7$ to $\nu = -1.37$. The Weiss constants extracted from the Curie–Weiss fits are negative for all the explored hole filling factors, suggesting an antiferromagnetic phase for all correlated hole states. We note we observe no magnetic hysteresis in the Zeeman splitting at $\nu = -1$ when the magnetic field is swept from negative to positive values followed by a subsequent positive to negative sweep (see Supplementary Fig. 7). In contrast to the large polaron $g$-factor enhancement observed under hole doping, we only observe a modest $g$-factor enhancement in the electron doping regime (see Supplementary Fig. 8).

DISCUSSION

The extraordinary $g$-factors observed under hole doping can be understood by considering the effect of a magnetic field on a localised hole in the triangular moiré superlattice. As a result of exchange interactions with other holes in its environment, such a hole experiences an effective magnetic field which is the sum of the externally applied field and the field induced by the other holes which in turn is proportional to the magnetisation $M$ of the localised hole gas (i.e., $\propto \lambda_0 \nu M$ with $\lambda_0$ being a coupling constant).

To calculate the induced field for a given fractional filling, we first determine the configuration of localised carriers that minimises the electrostatic repulsion energy using a simulated annealing technique (see Supplementary text for details). To describe the spin response of this arrangement of charges, we consider a Heisenberg Hamiltonian with distance-dependent antiferromagnetic isotropic exchange interactions $J(r) = J_0 \exp(-r/r_0)$ with $J_0$ denoting the magnitude of the exchange coupling at the characteristic length scale $r_0$. For this Hamiltonian, the induced magnetic field and the corresponding $g$-factor enhancement, $g' = \frac{g}{g_0}$, of a localised hole are calculated within mean-field theory (see Supplementary text for a detailed description). To obtain the effective $g$-factor of the attractive exciton-polaron which is probed in our experiments, we assume that its $g$-factor enhancement due to exchange interactions with localised holes is the same as that of a single hole, but that the 'non-enhanced' $g$-factor $g$ can be different. The experimental value of this non-enhanced $g$ is unknown since the $g$-factors of exciton-polarons are dependent on paramagnetic interactions and phase-space filling effects as carrier concentration is changed, even for monolayer TMDs. Therefore, we treat $g$ as an adjustable parameter choosing its value such that the calculated and experimentally measured $g$-factors agree at $\nu = -1$.

Figure 4f shows that the filling dependence of the calculated $g$-factor is in good qualitative agreement with the experimental results. Specifically, it reaches a maximum at $\nu = -1$, where the average number of occupied moiré sites around the localised hole is largest and the strong exchange interactions between neighbouring spins give rise to a large effective magnetic field. The model also captures the plateau-like feature between $\nu = -1/3$ and $-2/3$. In contrast to the experimental findings, however, the calculated $g$-factor is symmetric around $\nu = -1$. Potential reasons for this discrepancy include (i) band structure effects and (ii) the doping-induced reduction of frustration. Regarding (i), Tang et al. pointed out that the maximum $g$-factor occurs at a filling factor that corresponds to a van Hove singularity of the density of states. Regarding (ii), Zhang et al. carried out exact diagonalisation studies of a triangular t-J model and found that at finite temperatures the maximum spin susceptibility occurs at a hole filling higher than $\nu = -1$. This is in agreement with a study by Koresutse et al. which employs a high-temperature expansion of the spin response function. The increase in the spin susceptibility was explained in the work by the "release" of frustration by doping. Further theoretical work is required to fully understand the detailed behaviour of the $g$-factor.

Finally, we estimate $U/t$ in our device. The antiferromagnetic coupling between neighbouring spins due to the kinetic exchange mechanism can be estimated as $J = -t^2/\sqrt{U}$, where $t$ is the hopping amplitude between neighbouring moiré lattice sites. Using the value of $\theta$ at $\nu = -1$ we estimate $J = -0.4$ meV. By combining $J$ with the estimated melting temperature of the correlated state at $\nu = -1$ ($\sim 55$ K) we obtain $U/t = 3.5 \pm 0.4$. This experimental value agrees well with predicted values for MoSe$_2$/WSe$_2$ heterostructures with stacking angles $\sim 3^\circ$ and $\sim 57^\circ$.

Our results illustrate the properties of exciton-polarons in the presence of correlated states in moiré heterostructures. Using the changes in energy, oscillator strength, and linewidth of intralayer excitons dressed by itinerant carriers occupying narrow electronic moiré bands in a MoSe$_2$/WSe$_2$ hetero-bilayer, we observe the formation of correlated electron and hole states at a multitude of fractional fillings of the moiré lattice. Upon hole doping, the WSe$_2$ attractive polaron transfers oscillator strength back to the repulsive polaron branch at 1 carrier per site, demonstrating the reduced screening of the exciton by the free charges in the presence of the insulating state. In addition, we observe the magnetic interactions within the correlated hole states via both the attractive and repulsive WSe$_2$ exciton-polarons, which exhibit enhanced Zeeman splittings due to exchange interactions with the moiré pinned carriers. Through temperature dependent measurements, the magnetic ordering of the correlated holes is shown to be antiferromagnetic in the range $\nu = -0.7$ to $\nu = -1.37$, and the $U/t$ ratio of our device is estimated to be $\sim -3.5$. Further investigations could exploit the small lattice mismatch between MoSe$_2$/WSe$_2$, which enables a highly tunable moiré period, to simulate condensed matter phase diagrams over a large range of $U/t$ ratios. Our observation of the formation of flat electronic
bands compliments recent reports of moiré trapped interlayer excitons in MoSe2/WS2 hetero-bilayers\textsuperscript{12–38} and highlights the exciting prospects to investigate Fermi–Hubbard and Bose–Hubbard physics in this system.

**METHODS**

**Sample fabrication**

The sample was fabricated using the all-dry viscoelastic transfer technique\textsuperscript{55}. Bulk crystal was exfoliated onto PDMS stamps and monolayer flakes were identified using an optical microscope. As shown in a previous work for the same sample studied in this work, the zigzag edges of the TMD monolayers were identified using optical images in order to achieve a nearly angle-aligned heterostructure (0° or 60°). Further details can be found in\textsuperscript{34}. The interface between the two TMD layers is kept pristine (free of polymer contamination) by a transfer of one flake from its PDMS stamp onto the other flake on its PDMS stamp before subsequent transfer onto the bottom hBN layer of the device. The twist angle is estimated to be −57° from the optical images. Gold contacts to the graphene layers were fabricated using standard lithography techniques.

**Optical measurements**

The sample is held in a closed-cycle cryostat at 4 K unless otherwise specified (for temperature dependent measurements). For differential reflectivity measurements, light from a power stabilised tungsten lamp is collected by a multi-mode fibre. The light is collimated by a 20x objective and focused on the sample with an achromatic objective (0.82 numerical aperture). The reflected light is 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 is controlled using a series of linear polarisers, quarter-wave and half-wave plates.

**Filling factor calibration**

The carrier concentration \( n \) in the heterostructure can be calculated using the parallel plate capacitance model, \( n = \frac{\epsilon_\text{hBN} \Delta V_0}{\epsilon_\text{hBN} \lambda_\text{m} + \epsilon_\text{hBN} \Delta V_0} \), where \( \epsilon \) is the permittivity of hBN, \( \Delta V_0 \) is the voltage offset between both the top and bottom gates and the heterostructure gate, and \( \lambda_\text{m} \) (\( d_\text{hBN} \)) is the thickness of the top (bottom) hBN layer measured to be 17.4 ± 0.2 nm (18.2 ± 0.3 nm) using nulling ellipsometry. For a small angular difference between two stacked layers the moiré periodicity can be estimated using \( \lambda_\text{m} = \frac{d_\text{e2}}{\sqrt{\theta^2 + \delta^2}} \), where \( d_\text{e2} \) is the lattice constant of WS\textsubscript{2}, \( \delta \) is the fractional lattice mismatch between the two layers and \( \theta \) is the twist angle in radians. For a triangular moiré pattern, the number of carriers required for one hole per site is given by \( n_0 = \frac{\sqrt{3}}{2\pi} \lambda_\text{m} \). Using the lattice constants of 0.3280 and 0.3288 nm for WS\textsubscript{2} and MoSe\textsubscript{2}, respectively\textsuperscript{38} and a permittivity of 3.8 for the hBN\textsuperscript{56} and \( \Delta V_0 = 1.34 \text{ V} \) for \( \nu = \pm 1 \) as determined in the main text, we calculate the twist angle in the main measurement location to be 56.9°. This agrees well with the angle of −57° estimated from the optical micrograph.

**DATA AVAILABILITY**

Data described in this paper are available online at https://researchportal.hw.ac.uk/en/persons/brian-d-gerardot/datasets/.
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COMPETING INTERESTS
The authors declare no competing interests.