Evidence for a spinon Kondo effect in cobalt atoms on single-layer 1T-TaSe$_2$

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Quantum spin liquids are highly entangled, disordered magnetic states that are expected to arise in frustrated Mott insulators and to exhibit exotic fractional excitations such as spinons and chargons. Despite being electrical insulators, some quantum spin liquids are predicted to exhibit gapless itinerant spinons that yield metallic behaviour in the charge-neutral spin channel. We deposited isolated magnetic atoms onto single-layer 1T-TaSe$_2$, a candidate gapless spin liquid, to probe how itinerant spinons couple to impurity spin centres. Using scanning tunnelling spectroscopy, we observe the emergence of new, impurity-induced resonance peaks at the 1T-TaSe$_2$ Hubbard band edges when cobalt adatoms are positioned to have maximal spatial overlap with the local charge distribution. These resonance peaks disappear when the spatial overlap is reduced or when the magnetic impurities are replaced with nonmagnetic impurities. Theoretical simulations of a modified Anderson impurity model show that the observed peaks are consistent with a Kondo resonance induced by spinons combined with spin-charge binding effects that arise due to fluctuations of an emergent gauge field.

Gapless quantum spin liquids (QSLs) are predicted to act like ‘neutral metals’ that exhibit metallic behaviour in the spin channel despite being Mott insulators$^{1-4}$. Evidence for this exotic metal-like behaviour has been observed using thermal and magnetic measurements for several gapless QSL candidates$^{5-11}$. These bulk probes, however, average over impurities and disorder, leading to inconsistent results for different samples$^{12-16}$ and/or alternative explanations for the experimental signatures$^{17}$. Local probe measurements, on the other hand, allow direct comparison of defect-free regions of a material with regions containing well-defined impurities$^{14,16}$. Local magnetic moments are a particularly interesting type of impurity because their direct interactions with spinons allow them to potentially probe the effects of spinon itinerancy$^{15}$. In a conventional metal, the impurity moment is collectively screened by itinerant conduction electrons via the Kondo effect$^{15}$. For a gapless QSL with a spinon Fermi surface, it has been predicted that itinerant spinons might also perform Kondo screening of magnetic impurities, yielding Kondo temperatures that scale similar to the conventional metallic case$^{16}$. Some experimental evidence for spinon Kondo screening has been obtained from muon spin rotation studies of the QSL candidate Zn-brochantite (ZnCu$_x$(OH)$_{1-x}$SO$_4$) that show reduced magnetic moments arising from Cu–Zn inter-site disorder$^{17}$. Local measurements of individual Kondo impurities in a QSL, however, have not yet been reported.

Here, we demonstrate the use of scanning tunnelling microscopy/spectroscopy (STM/STS) to probe the Kondo response of individual magnetic impurities in a gapless QSL. This measurement is complicated by the fact that QSLs are typically good insulators whereas STM requires a nonzero electrical conductivity to function. We have overcome this challenge by using a new two-dimensional gapless QSL candidate, single-layer (SL) 1T-TaSe$_2$, that is supported by a sheet of conductive graphene and so allows injected charge to be drained away$^{12-14}$. STM imaging allows us to identify defect-free patches of SL 1T-TaSe$_2$ where we have previously discovered long-wavelength spatial modulations indicative of a half-filled spinon Fermi surface$^{14}$. In this work, we introduce individual magnetic impurities onto the clean 1T-TaSe$_2$ surface by vacuum deposition of a low coverage of Co atoms. By using STS at $T=5\,\text{K}$, we observe that Co atoms having maximal spatial overlap with the 1T-TaSe$_2$ d-orbital charge distribution exhibit new resonance peaks at both the upper and lower Hubbard band edges in the vicinity of Co impurities. Nonmagnetic Au atoms deposited onto the 1T-TaSe$_2$ surface do not show this unusual behaviour. To understand why Co atoms on SL 1T-TaSe$_2$ generate new electron–hole band-edge resonances, we simulated our magnetic impurity system utilizing a modified Anderson model. Strong interaction effects were accounted for both in the substrate and in the impurity through a slave-rotor mean-field theory approach employing a U(1) gapless

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The main experimental result reported here was obtained for individual magnetic Co atoms positioned at the centre of the 1T-TaSe$_2$ charge density wave (CDW) unit cell. Magnetism in Co adatoms for this configuration is supported by density functional theory (DFT) simulations (Methods and Supplementary Fig. 1). As sketched in Fig. 1a, in the CDW phase of 1T-TaSe$_2$, the Ta atom lattice distorts (due to electron–phonon interactions$^{25}$) to form a triangular super-lattice of star-of-David (SOD) clusters, each composed of 13 Ta atoms. An STM image of a single Co atom sitting at the centre of a SOD cluster can be seen in the STM topograph of Fig. 1b (obtained at a sample bias voltage of $V_b = -0.5$ V). At this bias voltage, the SOD CDW superlattice can clearly be seen, and the Co adatom creates a bright protrusion $\sim$1.4 Å high. Imaging the same region at a high positive bias of $V_b = 1.5$ V reduces the CDW contrast, allowing the position of the Co atom to be determined more clearly (Fig. 1c).

An STS $dI/dV$ spectrum obtained for a single Co atom on 1T-TaSe$_2$ is shown in Fig. 1d. Here, the blue curve shows the spectrum obtained when the STM tip is held over the centre of a pristine SOD cluster with no Co adatom (Fig. 1b,c, blue dot). Two peaks are seen, marking the lower Hubbard band (LHB) at $V_t = -0.34$ V and the upper Hubbard band (UHB$_1$) at $V_t = 0.62$ V for pristine 1T-TaSe$_2$, in agreement with previous results.$^{21}$ The upper Hubbard band is marked ‘UHB,’ because another Hubbard band feature (UHB$_2$) occurs at lower energy ($V_t = 0.17$ V) but with little weight (that is, reduced local density of states (LDOS)) at the SOD centre (Fig. 2b and ref. $^{23}$). The red curve shows the $dI/dV$ spectrum measured while placing the STM tip above the Co atom shown in Fig. 1b,c (red dot). Two outer peaks are seen, corresponding to the LHB and UHB$_2$ features of the pristine surface. Strikingly, however, two new peaks are observed at $V_t = -0.15$ V (labelled P$_0$, just below the LHB band edge) and at $V_t = 0.53$ V (labelled P$_e$, just above the UHB$_2$ band edge) (Supplementary Fig. 3). These two peaks are the main experimental finding of this paper. The P$_0$ and P$_e$ peaks are relatively narrow with a full-width at half-maximum of 106 and 90 meV, respectively.

The Co electronic structure is sensitive to where on the surface the adatom sits (Fig. 2). Here, local electronic structure variation can be seen in both our pristine surface spectroscopy and our ‘on-Co’ spectroscopy, depending on atomic position. In general there are three types of positions that Co atoms can occupy (Fig. 2a, sketch): (1) on-centre, (2) off-centre and (3) midway (with respect to the centre of a SOD unit cell). Co atoms occupying these three different positions can be seen in the topographs of Fig. 2c,d. One of the main differences between the three different positions is the strength of the pristine SL 1T-TaSe$_2$ Hubbard band LDOS (Fig. 2b). As shown previously$^{21}$, both the LHB and UHB$_2$ features exhibit high LDOS at the on-centre position but decrease monotonically as one moves outwards. The UHB$_2$ feature, in contrast, peaks at locations away from the SOD centre. This behaviour is reflected in the pristine (that is, off-Co) $dI/dV$ spectra, which show the UHB$_2$ feature absent at the on-centre position but appearing more strongly for the off-centre or midway position (Fig. 2e, blue curves).

The on-Co $dI/dV$ spectra in Fig. 2e (red curves) differ substantially when the Co atom sits at different positions. While the P$_0$ and P$_e$ features appear strongly for the on-centre position, they are completely absent when Co atoms occupy either the off-centre or midway position. At the off-centre position, Co atoms exhibit features that correspond to the LHB, UHB$_2$ and UHB$_1$ features, but the UHB$_1$ peak observed at $V_t = 0.25$ V is slightly offset from the pristine UHB$_1$ feature at $V_t = 0.17$ V. At the midway position, Co atoms exhibit features corresponding to LHB and UHB$_2$, but show no features corresponding to UHB$_1$. All of the spectroscopic features shown in Fig. 2c–e were reproducibly observed when we shifted individual Co atoms from one position to another for all three positions using atom manipulation (Supplementary Figs. 4, 5 and Note 3). This helps to clarify that the spectroscopic features highlighted above arise from individual Co atoms and are not due to other types of surface defects.
As a control measure, we performed these same types of measurements using a nonmagnetic impurity, Au. Au atoms were deposited onto SL 1T-TaSe₂ under the same general conditions that we used for depositing Co atoms. An STM topograph of a single Au atom on SL 1T-TaSe₂ can be seen in the inset to Fig. 3. Au atoms show a preference for binding at the centre of SOD clusters. Occasionally, Au atoms were found at other positions on the surface, but in those cases the atoms were unstable under STM spectroscopic conditions (the only stable position for Au atoms being the on-centre position).

The results of STM dI/dV spectroscopy performed with the STM tip held over a Au atom in the on-centre position can be seen in Fig. 3 (red curve). The Au atom spectrum shows no new resonances compared with the reference spectrum (blue curve, using the same tip) and notably does not show either the P₁ or the P₂ peak that was observed for Co atoms. Features in the Au atom spectrum corresponding to the pristine LHB and UHB resonances can be seen, although the LHB feature appears to be shifted to slightly lower voltage for the on-Au spectrum (Vₛ = -0.46 V) compared with the pristine SL 1T-TaSe₂ spectrum (Vₛ = -0.31 V).

The appearance of two new resonances symmetrically aligned adjacent to the Hubbard band edges, as seen for Co atoms in the on-centre position, is an unusual type of defect behaviour. Inelastic tunnelling channels associated with impurity spin or vibrational excitations can induce dI/dV steps or replicate existing lower-energy elastic dI/dV peaks but cannot create new dI/dV peaks. Impurity doping can induce polaronic states dressed by phonons or spins, but they occur at only one side of the gap and/or require a static magnetic background (which is not present here). Impurities coupled to a CDW can produce a pair of bound states, but they are typically located at the CDW gap edges rather than the Hubbard band edges. For conventional semiconductors, band-edge resonances do occur for shallow donor and acceptor impurities, but at only one band edge. Donor states reside below the conduction band edge while acceptor states reside above the valence band edge, but it is abnormal for a single type of defect to exhibit a resonance at both energies simultaneously. Defect behaviour in Mott insulators is more poorly understood than in conventional semiconductors, but the behaviour we observe is unusual even for Mott insulators. In 'conventional' Mott insulators such as cuprates and iridates, for example, impurities have generally been found to induce broad in-gap states that fill up the Mott–Hubbard gap and metallize the system through a doping mechanism. In bulk 1T-TaS₂, which has many similarities to SL 1T-TaSe₂, the Mott insulating state has
been shown to partially or fully collapse upon the introduction of defects or K adatoms, or the substitution of S with Se. The appearance of symmetrically placed resonances near the Hubbard band edges, however, has not previously been seen.

We are able to explain the observed behaviour of Co adatoms at the surface of 1T-TaSe\textsubscript{2} by the appearance of the Kondo effect for a magnetic impurity in contact with a gapless QSL. For the ‘conventional’ Fermi liquid Kondo effect, scattering between itinerant conduction electrons and a localized impurity spin leads to a many-body singlet state whose signature is a narrow ‘Kondo resonance’ at the Fermi level ($E_F$). Such features have been observed by STM for individual magnetic impurities on metal surfaces and exhibit a Fano line shape in the electronic LDOS. For a gapless QSL the Kondo behaviour is predicted to be strikingly similar to the Fermi liquid case if one considers only the spin channel (since these are the elementary excitations of QSLs). The problem lies in how to understand the experimental detection of a Kondo resonance that exists only in the spin channel of a Mott insulator. The solution is to take into account both the spin and charge channels simultaneously. Since conventional STMs can only tunnel electrons into a material, these electrons must decompose into both spinons and chargons once they are inside a QSL (since these are the elementary excitations of QSLs). The STM electron tunnel current into a QSL can thus be expressed as a convolution of the spinon and chargon density of states. If one wants to use STM to detect a Kondo resonance in the spin channel of a QSL, one must therefore consider how it will appear when the spinon density of states is appropriately convoluted with the simultaneously occurring chargon density of states.

Such reasoning led us to calculate both the spinon and chargon density of states for a magnetic impurity in contact with a QSL. To accomplish this, we utilized a modified Anderson model incorporating a U(1) QSL having a spinon Fermi surface (the model parameters are mostly determined by experimental spectra; Methods, Supplementary Note 1 and ref. 45). The model was solved using a slave-rotor mean-field technique whereby electrons are decomposed into auxiliary fermions and charged bosons (gauge fluctuations are ignored at this level). The resulting theoretical DOS spectra of impurity auxiliary fermions (a) and QSL auxiliary fermions (‘spinons’) (b) in the spin channel. Impurity auxiliary fermions show a sharp resonance at the Fermi level due to Kondo screening from itinerant QSL spinons. QSL spinon DOS shows a dip at $E_F$ due to level repulsion from the Kondo resonance.

The model was solved using a slave-rotor mean-field theory where electrons are decomposed into spin-1/2 auxiliary fermions (the spin channel) and charged spinless bosons (the charge channel)\textsuperscript{44,46}. The auxiliary fermions and charged bosons within the QSL (not including the impurity) are referred to as spinons and chargons, respectively. The solution of the Anderson model in the spin channel yields a Kondo resonance peak in the density of states (DOS) of the impurity auxiliary fermion at the Fermi level of the QSL spinon band (Fig. 4a). In the QSL spinon spectrum, the Kondo resonance manifests as a dip at the Fermi level (Fig. 4b), which can be rationalized as level repulsion in a standard Fano-type scenario (that is, involving a discrete state in resonance with a continuum)\textsuperscript{45,46}.

In the charge channel, the impurity charged boson hybridizes with QSL chargon states, resulting in an intermixing of their DOS spectra (Fig. 4c,f). For example, in addition to having the usual bare impurity levels ($E_F$ and $E_F + U$), the impurity exhibits new spectral features that appear at the QSL Hubbard band energies (Fig. 4c, dashed lines) which are inherited from the QSL chargon spectrum. Similarly, the QSL chargon DOS spectrum exhibits weak peaks at $E_F$ and $E_F + U$ (Fig. 4d, dashed lines) that originate from the bare impurity levels.

To obtain the final predicted tunnelling DOS for electrons, the spin and charge channels must be combined via convolution (Methods and refs. 24,44,45) into a total electron channel. The result of this convolution can be seen in Fig. 4 for both the impurity (Fig. 4c) and the QSL continuum (Fig. 4f). The resulting impurity electronic DOS spectrum (Fig. 4e) exhibits a step function at the Hubbard band edges (that is, at the energies $\pm \Delta_{\text{Mott}}$) that is more abrupt than the linear dispersion seen in the absence of impurity (Supplementary Fig. 6c). However, there are no peaks at the band edges that resemble the experimental $P_1$ and $P_2$ features of Fig. 1d.

Explaining the experimentally observed side-peaks in the on-Co STS requires the incorporation of more realistic effects into our model, namely the inclusion of longitudinal gauge-field fluctuations (Methods, Supplementary Note 2 and ref. 45). The reason for this is that, in a U(1) QSL with a spinon Fermi surface, both spinons and chargons couple to an emergent U(1) gauge field\textsuperscript{46}. Since spinons...
and chargons have opposite gauge charge\(^{34,35}\), the gauge field produces an effective binding interaction \(U_b\) (which is short ranged here owing to the screening of the gauge field by the spinon Fermi sea). Around a magnetic impurity, the localized spinon Kondo screening cloud therefore attracts chargons in both the holon and doublon branches (Fig. 5a), forming composite spinon–chargon states at energies both near the bottom of the doublon branch and near the top of the holon branch (Fig. 5b), in analogy to the donor and acceptor states of a semiconductor.

Inclusion of this gauge binding effect into our model causes a pile-up of spectral weight near the band edges of the impurity electronic DOS. The pile-up appears for small binding interaction \(U_b = 0.08\) eV and increases smoothly with increasing \(U_b\) in the range shown in Fig. 5c. The value \(U_b = 0.19\) eV results in a pair of resonance peaks at the Hubbard band edges of the impurity electronic DOS that best fits our data. The \(P_1\) and \(P_2\) side-peaks in our experimental STM spectrum (Fig. 1d) can thus be understood to reflect composite spinon–chargon states induced by the spinon Kondo cloud around a magnetic impurity, with peak widths determined by hybridization with QSL electrons. When \(U_b\) is increased beyond a critical value of \(\sim 0.35\) eV (Fig. 5d and Supplementary Fig. 7), the band-edge resonances move inside the gap, forming narrow in-gap bound states that have no intrinsic width and require an extrinsic broadening to be better visualized (Supplementary Note 2). We find that varying model parameters such as the Coulomb repulsion, \(U_c\), and hybridization, \(V\), shifts the positions of the LHB and UHB peaks in this overall scenario. Because the SOD unit cell is composed of 13 Ta atoms (each containing one \(d\) electron\(^{23}\)), the fundamental building block of the SL \(1T\)-TaSe\(_2\) Mott bands is more akin to a molecular orbital than to an individual atomic state. This is manifest, for example, in the position dependence of UHB\(_2\) for Co in the on-centre position is qualitatively consistent with the experimental data (Fig. 3).

In conclusion, we deposited isolated magnetic adsatoms onto the single-layer QSL candidate \(1T\)-TaSe\(_2\). Magnetic impurities having large hybridization with the underlying Hubbard band LDOS develop a pair of resonance peaks at the Hubbard band edges, consistent with the impurity electronic DOS obtained from a modified

![Image](https://www.nature.com/naturephysics)

Fig. 5 | Gauge-field fluctuations induce electronic band-edge resonances and bound states. **a**, Sketch showing the inclusion of gauge-field fluctuations generating an effective attraction \(U_b\) that binds a QSL chargon to the spinon Kondo cloud. FS, Fermi surface. **b**, Energy diagram illustrating the formation of spinon–chargon band-edge resonance states due to the attraction between chargons and the spinon Kondo cloud. **c**, With increasing binding interaction \(U_b\), the impurity electronic DOS spectrum exhibits growing resonance peaks at the Hubbard band edges. At \(U_b = 0.19\) eV, the theoretical electronic resonances (green line) best reproduce the experimental observation in Fig. 1d. At even larger binding interaction \(U_b\), the spinon–chargon states move inside the Mott gap and form very narrow bound states that do not match the experiment (an extrinsic broadening of 0.01\(\text{eV}\) was applied in **d** to better visualize the bound-state peaks; Supplementary Note 2).
Anderson model incorporating a U(1) gapless QSL. This provides evidence supporting the existence of a spinon-based Kondo effect in a gapless QSL. Future experiments that more directly probe the magnetic moments of impurities could provide further insight into spinon-induced Kondo screening. Our ability to perform atomically precise impurity manipulation and spectroscopy with the STM tip might be applied to other layered QSL systems to gain microscopic control over other exotic excitations such as non-abelian anyons.

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Methods
Sample growth. Single-layer 1T-TaSe₂ films were grown on epilaxial bilayer graphene-terminating 6H-SiC(001) substrates in a molecular beam epitaxy chamber operating at ultrahigh vacuum (base pressure 2 × 10⁻¹⁰ Torr) at the HERBS endstation of Beamline 10.0.1, Advanced Light Source, Lawrence Berkeley National Laboratory. High-purity Ta (99.9%) and Se (99.999%) were evaporated from an electron beam evaporator and a standard Knudsen cell, respectively, with a Ta-Se flux ratio set between 1:10 and 1:20 and a substrate temperature of 660 °C. The growth process was monitored by reflection high-energy electron diffraction. Before taking the films out of vacuum for STM/STS measurements, Se capping layers with ~10 nm thickness were deposited onto the samples for passivation. These were later removed by ultrahigh-vacuum annealing at ~180 °C for 1 h.

STM/STS measurements. STM/STS measurements were performed using a commercial CreaTec STM/AFM system at T ≈ 5 K under ultrahigh-vacuum conditions. To avoid tip artefacts, STM tips were calibrated on a Au(111) surface by measuring its herringbone surface reconstruction and Schottky surface state before all STM/STS measurements. STS dI/dV spectra were obtained using standard lock-in techniques with a small bias modulation at 401 Hz. High-purity Cu wire (99.99%) was used to support Co atoms from an electron beam evaporator onto cold samples kept in the STM head (to avoid adatom clustering). Co evaporation was first calibrated on Au(111) substrates where the existence of isolated Co atoms was confirmed through their known Kondo resonance features. Single-layer 1T-TaSe₂/bilayer graphene on SiC was then transferred into the STM and brought to thermal equilibrium before performing another Co evaporation using identical deposition parameters. High-purity Au wire (99.99%) was used to evaporate Au atoms via direct heating onto cold samples kept in the STM head (that is, by wrapping the Au wire around a W filament heater).

Anderson impurity model on QSL. The following Hamiltonian was used to describe an atomic impurity with on-site repulsion U embedded in a QSL: $H = \sum_{\sigma} E_{\sigma} d_{\sigma}^\dagger d_{\sigma} + U d_{\uparrow}^\dagger d_{\uparrow} d_{\downarrow}^\dagger d_{\downarrow} + V \sum_{\sigma} \left( \epsilon_{\uparrow} d_{\uparrow}^\dagger d_{\downarrow} + \text{h.c.} \right) + H_b$. Here $E_{\sigma}$ is the on-site energy, $\sigma = \uparrow, \downarrow$ is the spin index, $V$ is the coupling between the impurity and the QSL electronic states at the nearest-neighbour site $R$ and $H_b = J \sum_{\sigma,\sigma'} c_{\sigma R}^\dagger c_{\sigma' R} + U_b \sum_{\sigma} c_{\sigma R}^\dagger c_{\sigma R} - \Delta_{\text{Mott}}$. The half-filled Hubbard model for the QSL has a slave-rotor mean-field solution, which fractionalizes the electrons in the QSL as $c_{\sigma R} = f_{\sigma R} X_{\sigma}$, where $f_{\sigma R}$ and $X_{\sigma}^{(1)}$ being the annihilation (creation) operators of the spinons and chargons, respectively. The spinon and chargon band dispersions in the QSL are taken to be

$$\delta_\sigma = 2t \cos k^x a \cos \frac{\sqrt{3}}{2} k^y a + \cos k^y a - \mu_f,$$

$$\epsilon_\sigma = 2t \cos k^x a \cos \frac{\sqrt{3}}{2} k^y a + \cos k^y a - 3 - \Delta_{\text{Mott}},$$

where $(k^x, k^y)$ denotes the quasimomentum of the Bloch states in a triangular lattice and $a$ is the lattice constant. Here $\Delta_{\text{Mott}} = 0.25 eV$ is fixed by the experimentally observed Mott gap size (which coincides with $U_b/2$ in the atomic limit).

The hopping parameters $t = 0.03 eV$ and $t_c = 0.03 eV$ are determined by the experimentally observed band width of pristine SL 1T-TaSe₂, while the half-filling requirement of the spinon band fixes the chemical potential to be $\mu_f = -0.024 eV$.

In the slave-rotor mean-field formalism, the electron operators of the impurity are rewritten as $d_{\sigma}^\dagger = f_{\sigma R} X_{\sigma}^\dagger$, $d_{\sigma} = f_{\sigma R} X_{\sigma}$ for the impurity electron, $\sigma = \uparrow, \downarrow$ annihilates (creates) a spin-1/2 auxiliary fermion that represents the spin degree of freedom of the impurity electron and $X_{\sigma}^{(1)}$ annihilates (creates) a spinless charged boson that corresponds to its charge degree of freedom. The coupling between the Anderson impurity and the QSL was solved using the slave-rotor mean-field so that the Green functions for the spin-1/2 auxiliary fermion and the charged boson at the impurity could be obtained. Here, the bare Co Coulomb energy $U$ and the hopping $V$ between the Co atom and the QSL are taken to be $U = 2eV$ and $V = 0.23 eV$, which yield a Kondo temperature of $T_K = 19 K$. Variation of $U$ and $V$ changes the Kondo temperature but does not significantly affect our main results (that is, the appearance of band-edge resonance peaks in the Kondo regime with $T < T_K$). More details regarding the Green function constructed from the slave-rotor mean-field theory can be found in Supplementary Note 1 and ref. 1.

Spectral function calculations. The electronic spectral function (that is, the LDOS) at the impurity and the QSL is the imaginary part of the retarded Green's function. For the impurity electron, the thermal Green's function is given by the correlation function $G_\sigma^{\text{R}}(i\omega_n) = -\left\langle \left( d_{\sigma}^\dagger(0,i\omega_n) d_{\sigma} (t,i\omega_n) \right) \right\rangle = -\frac{1}{\pi} \int_0^\infty \left\langle \left( d_{\sigma}^\dagger(0,i\omega_n + i\Delta) d_{\sigma} (t,i\omega_n + i\Delta) \right) \right\rangle \Delta^2 d\Delta$, where $\omega_n = \frac{n\pi eV}{k_B T}$, with $\Delta$ being the holon and doublon annihilation (creation) operators, respectively, which together compose the chargon creation and annihilation operators, $X_{\sigma}^{(1)} = a_{\sigma}^\dagger + a_{\sigma}$ and $X_{\sigma} = a_{\sigma} + a_{\sigma}^\dagger$. Incorporating the spinon–chargon binding interaction into the Anderson impurity Hamiltonian results in a modified correlation function that yields the final impurity electron thermal Green's function:

$$G_\sigma^{\text{R}}(i\omega_n) = \left\langle \left( d_{\sigma}^\dagger(0,i\omega_n) d_{\sigma} (t,i\omega_n) \right) \right\rangle$$

Similarly, the thermal Green's function for the QSL electron at $R$ is $G_{\sigma R}^{\text{R}}(i\omega_n) = \left\langle \left( c_{\sigma R}^\dagger(0,i\omega_n) c_{\sigma R} (t,i\omega_n) \right) \right\rangle$. The half-filled Hubbard model for the QSL exhibits a pair of resonance peaks at the Hubbard band edges as shown in Fig. 5. The detailed form of $G_{\sigma R}^{\text{R}}(i\omega_n)$ can be found in Supplementary Note 2.

DFT simulation of Co on single-layer 1T-TaSe₂. First-principles calculations of Co on the CDW phase of single-layer 1T-TaSe₂ were performed using DFT as implemented in the Quantum ESPRESSO package. A slab model with a 16 Å vacuum layer was adopted to avoid interactions between periodic images. We employed optimized norm-conserving Vanderbilt pseudopotentials including Ta 5s and 5p semicore states (with a plane-wave energy cut-off of 90 Ry) as well as the Perdew–Burke–Ernzerhof exchange–correlation functional in the generalized gradient approximation. Every CDW unit cell was assumed to host a Co atom above its central Ta atom, and the structure was fully relaxed at the DFT Perdew–Burke–Ernzerhof level until the force on each atom was less than 0.02 eV Å⁻¹. On-site Hubbard interactions were added through the simplified rotationally invariant approach using $U = 2eV$ for each Ta atom and $U = 3eV$ for each Co atom. DFT simulation results show that the Co atom on single-layer 1T-TaSe₂ slightly $r$-dopes the system and hosts a magnetic moment of 1.86 $\mu_B$, where $\mu_B$ is the Bohr magneton. The spin density associated with Co on single-layer 1T-TaSe₂ is shown in Supplementary Fig. 1.

Data availability
The data represented in Figs. 1d, 2e, 3, 4 and 5 are available as Source Data files. All other data that support the plots within this paper and other findings of this study are available upon request. Source data are provided with this publication.

Code availability
The codes used in the calculations shown in Figs. 4, 5c, 5d, S5a–c, S7, S8 and S9 are available as Supplementary Data Files 1–8.

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**Author contributions**

Y.C., W.R., P.A.L. and M.F.C. initiated and conceived this project. Y.C., W.R., R.L.L., T.Z. and C.Z. carried out STM/STS measurements under the supervision of M.F.C. J.H., S.T. and H.R. performed sample growth under the supervision of Z.-X.S. and S.-K.M. W.Y.H. performed slave-rotor calculations and theoretical analysis under the supervision of P.A.L. M.W. performed DFT calculations under the supervision of S.G.L. Y.C., W.Y.H., W.R., P.A.L. and M.F.C. wrote the manuscript with the help from all authors. All authors contributed to the scientific discussion.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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