In the 1950s, Pomeranchuk predicted that, counterintuitively, liquid He may solidify on heating. This effect arises owing to high excess nuclear spin entropy in the solid phase, where the atoms are spatially localized. Here we find that an analogous effect occurs in magic-angle twisted bilayer graphene (MATBG). Using both local and global electronic entropy measurements, we show that near a filling of one electron per moiré unit cell, there is a marked increase in the electronic entropy to about 1 $k_B$ per unit cell ($k_B$ is the Boltzmann constant). This large excess entropy is quenched by an in-plane magnetic field, pointing to its magnetic origin. A sharp drop in the compressibility as a function of the electron density, associated with a reset of the Fermi level back to the vicinity of the Dirac point, marks a clear boundary between two phases. We map this jump as a function of electron density, temperature and magnetic field. This reveals a phase diagram that is consistent with a Pomeranchuk-like temperature- and field-driven transition from a low-entropy electronic liquid to a high-entropy correlated state with nearly free magnetic moments. The correlated state features an unusual combination of seemingly contradictory properties, some associated with itinerant electrons—such as the absence of a thermodynamic gap, metallicity and a Dirac-like compressibility—and others associated with localized moments, such as a large entropy and its disappearance under a magnetic field. Moreover, the energy scales characterizing these two sets of properties are very different: whereas the compressibility jump has an onset at a temperature of about 30 kelvin, the bandwidth of magnetic excitations is about 3 kelvin or smaller. The hybrid nature of the present correlated state and the large separation of energy scales have implications for the thermodynamic and transport properties of the correlated states in twisted bilayer graphene.

Systems of strongly interacting fermions exhibit competition between localization, which minimizes the potential energy, and itineracy, which minimizes the kinetic energy. The advent of two-dimensional moiré systems, such as magic-angle twisted bilayer graphene (MATBG), allows this physics to be studied by controlling the ratio between the electronic interactions and bandwidth in a highly tunable way. When this ratio is large, electrons tend to localize and form Mott insulators. When the bandwidth dominates, a Fermi liquid state is formed in which electrons are itinerant. MATBG is at the boundary between these two extremes, showing a host of fascinating electronic phases, including correlated insulators, Chern insulators, superconductors and ferromagnets. Scanning tunnelling spectroscopy and electronic compressibility measurements indicate that in this system the strengths of the Coulomb interaction and the kinetic energy are indeed comparable. In this regime, there is an inherent tension between localized and itinerant descriptions of the physics. Moreover, the topological character of the nearly-flat bands in MATBG implies that a simple 'atomic' description, in which electrons are localized to individual moiré lattice sites, may not be appropriate. Instead, a picture analogous to that of quantum Hall ferromagnetism has been proposed. Understanding this interplay between itinerancy and localization, and the new physics that emerges from it, remains a major challenge.

In this work we find that, surprisingly, the correlated state in MATBG above a filling of one electron per moiré site has a hybrid nature, with some properties resembling those of an itinerant system, and others resembling those of localized electrons. At temperatures of a few kelvin we measure unusually large excess entropy, which is rapidly suppressed by a moderate in-plane magnetic field. This suggests that even at such low temperatures, there are strongly fluctuating magnetic moments in the system, a behaviour typically associated with local moments. On the other hand, our measurements find that this state is metallic and has no thermodynamic gap, naturally fitting an itinerant picture.
The presence of fluctuating moments at temperatures much below the electronic bandwidth indicates the existence of a new, anomalously small energy scale associated with the bandwidth of magnetic excitations, which is an order of magnitude smaller than the scale where a small energy scale associated with the bandwidth of magnetic excitations indicates the existence of a new, anomalously high nuclear spin entropy of the atoms in the solid. Similarly, we note that another domain with a small difference in twist angle dominates the averaged result over domains with different twist angle. Therefore, the resistance maxima are slightly shifted from the usual integer $v$ values, probably because another domain with a small difference in twist angle dominates the transport characteristics globally.

Our data are measured using two independent techniques on two conceptually different devices. The bulk of the results are obtained from local measurements of the electronic entropy and compressibility using a scanning nanotube single-electron transistor (SET) that probes a hexagonal boron nitride (hBN)-encapsulated MATBG device (device 1, Fig. 1a). We focus on a large (5 μm × 4 μm) region with an extremely homogenous twist angle that is close to the theoretical magic angle $\theta = 1.13^\circ \pm 0.005^\circ$. Similar results are obtained from global entropy measurements using a monolayer graphene sensor (device 2; see section 'Global measurements of the entropy'). Both methods have been described elsewhere.

**Electrical compressibility and transport**

The inverse compressibility, $d\mu/dn$, measured in device 1 at $T = 15$ K as a function of the filling factor, $\nu = n/(\pi/4)$ (where $\mu$ is the chemical potential, $n$ is the carrier density, and $n_\nu$ corresponds to four electrons per moiré superlattice unit cell), is shown in Fig. 1b. As reported previously, sharp jumps in $d\mu/dn$ are observed close to integer values of $\nu$, reflecting Fermi surface reconstructions. These were termed Dirac revivals as they were interpreted as resets of partially filled energy bands back to near the Dirac point, leading to the decreased compressibility. The cascade of revivals is already very prominent at this relatively high temperature. Measurements of the two probe resistance, $R$, versus $\nu$ at various temperatures $T$ (Fig. 1c) show insulating behaviour at $\nu = 2, 3$ and semi-metallic behaviour at $\nu = 0$. As previously noted, $R$ shows a step-like increase across $\nu = 1$, which gradually disappears with decreasing temperature, very different to the behaviour at other integer values of $\nu$.

The unusual physics near $\nu = 1$ is revealed by the dependence of $d\mu/dn$ on $T$ and parallel magnetic field, $B_\parallel$. At low temperature and $B_\parallel = 0$ T (Fig. 2a), the jump in $d\mu/dn$ occurs at $\nu$ slightly larger than 1. Increasing the temperature moves the jump towards lower $\nu$, and surprisingly, increases the magnitude of the jump rather than smearing it. Similar measurements with $B_\parallel = 12$ T at low $T$ (Fig. 2b) show a much larger jump, which is also closer to $\nu = 1$. With increasing temperature, this jump remains close to $\nu = 1$, but in contrast to the $B_\parallel = 0$ T case, its amplitude is reduced and its width increased.

**Local measurements of electronic entropy**

The chemical potential, $\mu(\nu, T)$ (measured relative to that at charge neutrality), can be obtained by integrating $d\mu/dn$ over carrier density (Fig. 2c, d). Visibly, $\mu$ depends strongly on $T$ for a range of values of $\nu$. This is clearly seen when we plot $\mu$ versus $T$ at two representative $\nu$ values (Fig. 2c, inset). At $\nu = 0.2$, $\mu$ is practically independent of $T$ (blue). In contrast, at $\nu = 0.9$ (red) $\mu$ is nearly constant until $T = 4$ K, and then decreases approximately linearly with $T$. At $\nu > 1.15$, $\mu$ is again nearly temperature independent. Comparison of $\mu$ at $B_\parallel = 0$ T (Fig. 2c) with that at $B_\parallel = 12$ T (Fig. 2d) reveals a clear contrast: whereas for $B_\parallel = 0$ T, $\mu$ is a decreasing function of temperature for $0.4 < \nu < 1.15$, for $B_\parallel = 12$ T, $\mu$ decreases with $T$ for $\nu < 0.9$ and increases for $\nu > 0.9$.
These measurements allow us to directly determine the entropy of the system, by integrating Maxwell’s relation, \( \frac{\partial s}{\partial v} = -\frac{1}{T} \frac{\partial \mu}{\partial T} \), to obtain \( s(v, T) \) (where \( s \) is the entropy per moiré unit cell). For more details on this procedure, see Supplementary Information section 1. Figure 2e shows \( s(v) \) at \( T = 10 \, \text{K} \) (obtained from the slope of \( \mu \) versus \( T \) in the range \( T = 4.5–15 \, \text{K} \), for \( B_0 = 0, 4, 8, 12 \, \text{T} \)). At \( B_0 = 0 \), the entropy is small at low \( v \), climbs close to \( v = 1 \), remains roughly constant between \( v = 1 \) and 2, at \( s = 1.2k_B \), drops rapidly near \( v = 2 \), and decreases towards zero after \( v = 3 \). Clearly, the \( v \) dependence of the entropy is qualitatively different from that of the compressibility: whereas the latter drops sharply near \( v = 1 \) (Fig. 2a), the former remains at a high value.

An important insight into the origin of this large entropy is given by its magnetic field dependence. As seen in Fig. 2e, the entropy above \( v = 1 \) depends strongly on \( B_0 \). In particular, at \( B_0 = 12 \, \text{T} \), most of the entropy between \( v = 1 \) and 2 is quenched. The inset shows that \( s(B_0 = 0 \, \text{T}) - s(B_0 = 12 \, \text{T}) \) versus \( v \) (the purple shading indicates error bars; see Supplementary Information section 1). The entropy difference increases sharply near \( v = 1 \), reaching a maximum of \((0.85 \pm 0.1)k_B \) between \( v = 1 \) and 2. To appreciate the importance of this value, we recall that an entropy of \( k_B \ln 2 \approx 0.7k_B \) corresponds to two degenerate states on each moiré unit cell. Moreover, in a Fermi liquid, we would expect a much weaker change in the entropy with \( B_0 \) (see Supplementary Information section 4), of the order of \( k_B \) times the ratio of the Zeeman energy (about 1\, meV at \( B_0 = 12 \, \text{T} \)) to the bandwidth, estimated to be \( W = 30 \, \text{meV} \) (see below). Finally, we observe that at \( B_0 = 12 \, \text{T} \) the entropy shows a cascade of drops following each integer \( v \). These drops are similar to the revival drops observed in the compressibility (Supplementary Information section 5) and are reproduced by mean-field calculations (Supplementary Information section 3). The strong quenching of entropy by moderate \( B_0 \) strongly suggests a magnetic origin.

Global measurements of the entropy
To test the robustness of our results, we measured the entropy in a completely different setup, in which a sheet of monolayer graphene senses the chemical potential of MATBG, averaged over the entire device\textsuperscript{2} (Fig. 3a). Figure 3b shows the entropy extracted in three different temperature ranges. We see (inset) that the globally measured entropy for \( T = 4–16 \, \text{K} \) is in good agreement with the locally measured one over a similar range of temperatures, both in the overall shape, the magnitude of \( s(v) \), and the detailed features. At elevated temperatures, the minimum in the entropy at \( v = 0 \) gradually fills in, evolving from a double-dome structure at low \( T \) (corresponding to the valence and conduction flat bands) to a single dome at high \( T \). This dependence is qualitatively reproduced by a naive calculation for a system of non-interacting electrons, whose density of states rises linearly from the charge neutrality point until the band edges (Fig. 3c). The merging of the domes in \( s(v) \) occurs when the temperature exceeds a fraction of the bandwidth. Calibrating the bandwidth using the measured entropy at \( T = 55 \, \text{K} \) gives \( W = 30 \, \text{meV} \), where \( W \) is the full bandwidth— from valence band bottom to conduction band top, in rough agreement with scanning tunnelling microscopy\textsuperscript{15–17} and compressibility\textsuperscript{18} experiments. This free-electron picture is of course invalid at low temperatures, where interactions are important. The measured \( s(v) \) in the valence band is approximately a mirror image of \( s(v) \) in the conduction band (Fig. 3b), although it is smaller and has less pronounced features. This is consistent with the weaker \( d\mu/dn \) revivals observed in the valence band relative to the conduction band\textsuperscript{21,26} (Supplementary Information section 9).

Mapping the phase diagram
So far, we have shown the change in the magnetic entropy and compressibility near \( v = 1 \). This change may be due to a continuous build-up
of electronic correlations. Alternatively, it could be interpreted as an underlying first-order phase transition between two distinct phases. Naively, one would then expect a discontinuous jump in thermodynamic properties and hysteretic behaviour across the transition, which are not observed. However, we note that a true first-order phase transition is forbidden in two dimensions in the presence of disorder or long-range Coulomb interactions\(^3\), as these broaden the transition into a mesoscopic coexistence region (Supplementary Information Section 10). Experimentally, although the revival transition is very sharp and may be consistent with a Coulomb- and/or disorder- smearing first-order transition, we cannot rule out a sharp crossover or a higher-order phase transition. Nevertheless, the sharpness of the rise of \(d\mu/d\nu\) at the revival transition allows us to precisely track its filling factor, \(\nu = \nu_0\) (Fig. 4a, and map a phase diagram, which is naturally explained when this feature is interpreted as a proxy for a first-order transition.

The measured \(\nu_0\) versus \(B_{||}\) and \(T\) forms a surface in \((\nu, B_{||}, T)\) space (Fig. 4b), whose projections onto the \((\nu, B_{||})\) and \((\nu, T)\) planes are shown in Fig. 4c, d. At \(T = 2.8\) K and at low \(B_{||}\), \(\nu_0\) depends weakly on \(B_{||}\), but decreases linearly above \(B_{||} = 4\) T (Fig. 4c, blue). A similar crossover is observed at higher temperatures, but with a crossover \(B_{||}\) that increases with temperature. The \(T\) dependence of \(\nu_0\) at \(B_{||} = 0\) T (Fig. 4d) is linear at low temperatures and curves up at higher temperatures. As \(B_{||}\) increases, the curve shifts towards smaller values of \(\nu\), and simultaneously its slope at low temperatures changes sign. At \(B_{||} = 12\) T, \(\nu_0\) first increases with \(T\), reaches a maximum at \(T = 9\) K, and then decreases.

The phenomenology seen in Fig. 4b–d can be understood in terms of a first-order phase transition at \(\nu = \nu_0\) between a Fermi liquid phase below \(\nu_0\), and a ‘free moment’ phase above it. The latter has a high concentration of free moments (about one per moiré site), coexisting with a low density of itinerant electrons. Within this framework, the shift of \(\nu_0\) as a function of \(B_{||}\) and \(T\) reflects the magnetization and entropy differences between the two neighbouring phases.

At \(B_{||} = 0\) T, the free moment phase has a higher entropy than the Fermi liquid, owing to thermal fluctuations of the moments. Hence, the former becomes entropically favourable at high temperatures. This explains the observed decrease of \(\nu_0\) with increasing \(T\) at low fields (Fig. 4d). Raising the temperature at fixed \(\nu\) may therefore drive a transition from the Fermi liquid to the free moments phase, an electronic analogue of the Pomeranchuk effect. As \(B_{||}\) increases and the Zeeman energy exceeds the temperature, the moments become nearly fully polarized and their entropy is quenched (as is observed directly in Fig. 2e). Consequently, at low temperatures and sufficiently high fields, the Fermi liquid phase is favoured by raising the temperature. The trend reverses once the temperature exceeds the Zeeman energy. This explains the non-monotonic behaviour of \(\nu_0\) as a function of \(T\), seen at \(B_{||} = 12\) T in Fig. 4d. The main features of the phase boundary are qualitatively reproduced in a thermodynamic model of the two phases (Supplementary Information Section 7, and insets of Fig. 4b–d). Note that the experiment probes moments that couple to an in-plane field. This includes Zeeman-coupled spins and may also include the valleys if their in-plane orbital moment is non-zero.

**Discussion**

The observation of free magnetic moments at surprisingly low temperatures has profound implications for the physics of MATBG. Low-energy magnetic fluctuations are destructive for superconductivity, and may be the limiting factor for the superconducting transition temperature. Moreover, increased scattering from fluctuating moments can account for the ‘strange metal’ behaviour reported over a broad range of temperatures\(^3,\)\(^4\).

An important question raised by our observations concerns the origin of the free moments. Soft collective modes have been predicted in insulating states of MATBG\(^3\)\(^5\), but our experiments show metallic behaviour near \(\nu = 1\). Moreover, the energy scale associated with the appearance of free moments is strikingly low (3 K or less), much below the microscopic energy scales in the system. Understanding the state near \(\nu = 1\), which combines behaviours associated with electron localization and itinerancy, and its surprisingly low onset temperature, poses an important challenge for the theory of MATBG.
Online content

Any methods, additional references, Nature Research 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/s41586-021-03319-3.

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Fig. 4 | Experimental phase diagram. a. The inverse compressibility, dµ/dn, measured as a function of ν near ν = 1, at several values of parallel magnetic field, B∥. We track the filling factor that corresponds to the centre of the jump in dµ/dn (labelled ν₀). The application of B∥ is seen to push ν₀ to lower values. b. Left panel, measured ν₀ as a function of B∥ and T, plotted as dots in (ν₀, B∥, T) space (the dots are coloured by their temperature as in c; the dashed lines are polynomial fits to the dots at constant B∥ or constant T). Right panel, the same surface calculated from a simple model that assumes a transition between a Fermi liquid and a metallic phase that contains one free moment per moiré site (see text). c. Projection of the data in b onto the (ν₀, B∥) plane, showing the dependence of ν₀ on B∥ for various temperatures. At low fields, ν₀ is independent of field but becomes linear in B∥ at high fields, a behaviour expected from the field polarization of free moments (see text). Inset, curves calculated from the model. d. Projection onto the (ν₀, T) plane, showing the dependence of ν₀ on T for various magnetic fields. At B∥ = 0 T, ν₀ is linear in T at small values of T and then curves up at higher values of T. At high magnetic field, the dependence of ν₀ on T becomes non-monotonic. Inset, curves calculated from the model.
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**Data availability**

The data in the main text are available at https://github.com/uzondi/MA_Pomeranchuk.

**Code availability**

The code used in this work is available at https://github.com/erezberg/pomeranchuk_tbg_theory.

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**Author contributions**

A.R., J.M.P., U.Z., Y.C., P.J.-H. and S.I. designed the experiment. A.R. and U.Z. performed the scanning SET experiments, and J.M.P. and Y.C. performed the monolayer graphene sensing experiments. D.R.-L. and Y.C. fabricated the twisted bilayer graphene devices. A.R., J.M.P., U.Z., Y.C., P.J.-H. and S.I. analysed the data. E.B., Y.O. and A.S. developed the theoretical model. K.W. and T.T. supplied the hBN crystals. A.R., J.M.P., U.Z., Y.C., A.S., E.B., P.J.-H. and S.I. wrote the manuscript.

**Competing interests**

The authors declare no competing interests.

**Additional information**

**Supplementary information**

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