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Magnetotransport of Sm₂Ir₂O₇ across the pressureinduced quantum-critical phase boundary

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ARTICLE OPEN Magnetotransport of Sm₂Ir₂O₇ across the pressure-induced

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Rare-earth pyrochlore iridates host two interlocking magnetic sublattices of corner-sharing tetrahedra and can harbour a unique combination of frustrated moments, exotic excitations and highly correlated electrons. They are also the first systems predicted to display both topological Weyl semimetal and axion insulator phases. We have measured the transport and magnetotransport properties of single-crystal $\text{Sm}_2\text{lr}_2\text{O}_7$ up to and beyond the pressure-induced quantum critical point for all-in-all-out (AIAO) Ir order at $p_c = 63$ kbar previously identified by resonant X-ray scattering and close to which Weyl semimetallic behavior has been previously predicted. Our findings overturn the accepted expectation that the suppression of AIAO order should lead to metallic conduction persisting down to zero temperature. Instead, the resistivity-minimum temperature, which tracks the decrease in the AIAO ordering temperature for pressures up to 30 kbar, begins to increase under further application of pressure, pointing to the pression of Ir magnetism, however, with a strong hysteresis observed only within the AIAO phase boundary, similar to that found for Ho₂Ir₂O₇ and attributed to plastic deformation of Ir domains. Around p_c we find the emergence of a new type of electronic phase, characterized by a negative magnetoresistance with small hysteresis at the lowest temperatures, and hysteresis-free positive magnetoresistance above approximately 5 K. The temperature dependence of our low-temperature transport data are found to be best described by a model consistent with a Weyl semimetal across the entire pressure range.

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quantum-critical phase boundary

INTRODUCTION

The magnetic pyrochlore lattice, with spins arranged into a network of corner-sharing tetrahedra, has a strong history of yielding new physical states, discoveries and insights. The seminal observation of a spin-ice state in $Ho_2Ti_2O_7$ and $Dy_2Ti_2O_7^{-1.2}$ and the associated magnetic monopole picture for its low-energy excitations^{3–6} have led to lasting interest in this family of materials. The pyrochlore iridates add to the richness of these compounds by introducing a second inter-penetrating magnetic lattice with its own order, and interactions between the two. This complex magnetic behavior, their highly correlated electronic states, and predictions of topological properties, make pyrochlore iridates fascinating systems for investigating exotic quantum phenomena.

The family of lanthanide pyrochlore iridates $Ln_2Ir_2O_7$, where Ln is a magnetic trivalent lanthanide ion, provide an ideal environment for investigating fundamental magnetism and correlatedelectron physics. They share the crystal structure shown in Fig. 1a — a cubic pyrochlore structure with space group $Fd-3m^{7-9}$. There are two interwoven sub-lattices of magnetic ions: a network of corner-sharing Ln^{3+} tetrahedra with an oxygen site in the center, and another of corner-sharing Ir^{4+} tetrahedra. Each Ir site sits in the center of an oxygen octahedron. While $Pr_2Ir_2O_7$ remains paramagnetic to low temperatures, the Ir sublattices in most (if not all) of the other members of this family undergo a transition to all-in-all-out (AIAO) antiferromagnetic long-range order⁹⁻¹³, as illustrated in Fig. 1b. In this configuration, all the spins on one Ir tetrahedron point radially outwards, and all those on its neighboring tetrahedra point inwards, giving zero total moment.

In addition to the AIAO magnetism of the Ir ions, the rare-earth sublattice in these materials may also show interesting and correlated behavior. In Dy₂Ir₂O₇ and Ho₂Ir₂O₇, the large rare-earth moments are constrained to lie along the local [111] directions and are coupled by both dipolar and exchange interactions, which are overall ferromagnetic^{14–16}. These interactions favor the two-intwo-out (2120) configuration, promoting spin-ice physics, akin to that seen in the frustrated pyrochlore titanates, and the appearance of magnetic monopole guasiparticles^{2,17}. The added effect of exchange with Ir nearest neighbors means that at sufficiently low temperatures both $Dy_2Ir_2O_7$ and $Ho_2Ir_2O_7$ are expected to undergo a transition to a fragmented monopole crystal^{14,15,18}. Experiments on Ho₂Ir₂O₇ at intermediate temperatures show that the magnetoresistance is sensitive to the density of magnetic monopoles and that the Ir AIAO domains can be manipulated, via the Ir-Ho coupling, using an applied magnetic field¹⁶. The other members of the family have considerably smaller moments on the rare-earth site and their magnetic ground state remains the subject of debate.

The materials in this series additionally all share equivalent transport behavior — metallic high-temperature resistivity and a resistivity minimum T_{min} (described in the literature as a metal-insulator transition temperature) below which the resistivity increases with cooling. This T_{min} occurs at the same temperature

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Fig. 1 Crystal structure. a Ambient-pressure crystal structure of $Sm_2Ir_2O_7$, based on structural data from refs.^{8,9}. The crystal forms a cubic pyrochlore structure, with space group *Fd*-3*m*. The Sm atoms form a network of corner-sharing tetrahedra. Between these is a second network of corner-sharing tetrahedra of Ir atoms shown in (**b**). Each Ir is surrounded by an oxygen octahedron. The black arrows in (**b**) denote an example of the all-in-all-out Ir magnetic order. The green arrow shows the direction of one of the [111] axes. The lattice parameter a = 10.311(2) Å for Sm₂Ir₂O₇ at room temperature and ambient pressure.

as the Ir AIAO ordering and has been attributed to a spin-orbitcoupling assisted Mott transition^{19,20}. The exception is Ln = Pr, in which the AIAO order is absent and which remains metallic at all temperatures. ARPES measurements have shown that in the paramagnetic phase the band structure exhibits quadratic-bandtouching points^{21,22}. Perturbations to this electronic structure, such as symmetry breaking due to magnetic order on the rareearth sites, are predicted to give rise to topologically non-trivial electronic states, including Weyl semimetals (WSMs) and accompanying Fermi arcs²¹, and evidence for such phenomena has been reported extensively^{19,23-27}.

It is possible to suppress the AIAO order in this family leading to what is reported in the literature as a quantum critical point, or sometimes a quantum metal-insulator transition, associated with significant quantum fluctuations^{12,20,24,25,28-30}. There are two principal means to tune interactions in these materials. Compressing the lattice parameter, a, will predominately strengthen the hopping term t in the Mott insulator description, while preserving local spin environments. Chemical substitution of the rare-earth site, on the other hand, mostly affects the oxygen coordinate parameter x. This parameter describes distortions away from a perfect octahedral environment (x = 0.3125) around the lr sites. towards a cubic environment around the Ln sites (x = 0.375) as its value increases (and the Ir-O-Ir bond angle decreases)¹². The reduction in x caused by decreasing Ln ionic radius as one moves from $Gd \rightarrow Eu \rightarrow Sm \rightarrow Nd$ reduces the local Ir Ising anisotropy, pushing down the AIAO ordering temperature^{12,19,31–33}. (Contrary to the Sm magnetism, where the presence of a strong single-ion anisotropy due to crystal electric field effects is robustly established, the single-ion behavior of the Ir magnetism is less well understood. By 'Ising anisotropy' of the Ir moments we refer to the fact that an AIAO order with well-defined local axes develops at low temperature and pressure. It remains to be determined whether this is purely an effect of the anisotropic direct Dzyaloshinskii-Moriya interaction combined with isotropic AFM exchange, or whether a single-ion CEF contribution plays a role). The reduction of AIAO ordering tempearture suppresses T_{min} accordingly and reduces the resistivity towards metallic behavior. The next compound in this progression, $Pr_2Ir_2O_7^{25,29,34}$, is then metallic at all temperatures and does not show any magnetic order^{12,29,31}. Reduction of either *a* or *x* has the result of suppressing the Ir AIAO order, but through distinct mechanisms. It is intriguing to ask what differences may result from the two approaches.

The strong sensitivity of the Ir ordering temperature, and the concomitant resistivity minimum, to the fine details of the crystal structure, as well as the many theoretical predictions of the presence of topological electronic states in the phase diagram³⁵, means that controlled tuning of the lattice is expected to be particularly enlightening in these materials. Applied hydrostatic pressure provides a clean, continuous and reversible method for achieving this tuning.

There have been several pressure studies reporting the evolution of resistivity in $Ln_2lr_2O_7$ samples. Most show a reduction in resistivity with pressure as they are driven towards metallization. Polycrystalline Nd₂lr₂O₇ reproduces the metallic behavior of Pr₂lr₂O₇ beyond 65 kbar of applied pressure^{19,25}. The T_{min} transition is continuously suppressed with pressure until it disappears at this point^{19,25,33}. Magnetic field has also been shown to induce metallicity in Nd₂lr₂O₇ by flipping the AIAO order to a 2-in-2-out state³⁶. In Gd₂lr₂O₇, T_{min} linearly increases with applied pressure³³. In Eu₂lr₂O₇ there are reports of a slight increase of T_{min} under pressures up to 22 kbar in polycrystalline pellets³³ or very little change up to 120 kbar in single crystals³⁷.

The differences in this behavior across the family are likely due to the interplay of the competing changes in x and a lattice parameters induced in each compound. $\text{Sm}_2\text{lr}_2\text{O}_7$, which in the *Ln* series lies between Eu and Nd and hence between these two regimes of different pressure dependence, is therefore an interesting material for investigation.

The poly- or single-crystalline nature of the samples in this family is known to be important. Polycrystalline samples of $Ln_2lr_2O_7$ can differ in transition temperature and form of resistivity to single-crystal samples^{15,24,38}. These differences likely arise from slight variations in composition, the effects of the broken symmetry at the surface or the restricted magnetic domain size in polycrystalline samples, the random orientation of individual grains, and the unknown transport properties of inter-grain boundaries present in pressed pellets. For these reasons, conclusions derived from transport measurements on polycrystalline samples should be treated with caution. Single crystals are indispensable for true clean tests of the physics in these systems, and for direction-sensitive magnetotransport measurements, but have only recently been successfully grown for many of these materials^{12,15,16,38}.

Single crystals of Sm₂Ir₂O₇ exhibit a high-temperature metallic phase (in which resistivity drops with decreasing temperature). A transition to AIAO Ir order occurs around $T_{AIAO} = 115$ K at ambient pressure, and the associated T_{min} is clearly seen in transport data

as a resistivity minimum below which resistivity rises with decreasing temperature^{31,33}. There is some evidence from muon-spin relaxation measurements that the Sm moments undergo ordering, possibly short-range, at temperatures below about 10 K³⁹, but the details remain unknown. Recently Wang et al. showed, using resonant X-ray diffraction on single crystals, that the Ir AIAO ordering temperature is reduced to a guantum critical point (QCP) at a pressure $p_c = 63 \text{ kbar}^{12}$. Above p_c a metallic state is expected once the magnetic transition responsible for driving insulating behavior is removed, similar to that seen in Pr₂Ir₂O₇ or in Nd₂Ir₂O₇ in applied fields or pressures^{19,25,33}. There are also several predictions that Sm₂lr₂O₇ hosts WSM states^{9,12,20,40,41}, most likely in proximity to the guantum-critical region close to where the magnetic order is suppressed^{12,25}. Initial transport measurements on polycrystalline samples have indeed shown that T_{min} drops gradually in Sm₂Ir₂O₇ as pressure is applied in the low pressure region up to 20 kbar³³. However, so far no measurements of electronic properties have been performed in the vicinity of the QCP, nor have any transport experiments been performed on single crystals of this material.

In this work, we report magnetotransport measurements on a high-quality single-crystal sample of Sm₂Ir₂O₇ in fields up to 15 T and pressures up to and beyond the 63 kbar quantum critical point. Our magnetoresistance data show that the AIAO Ir order is indeed suppressed at this pressure, in line with the X-ray measurements¹². However, while the resistance drops dramatically with pressure, the T_{min} resistivity minimum does not fall to zero temperature with the magnetic order as expected. Instead, for pressures above about 30 kbar the high-temperature metallic state is disrupted upon cooling by another process, which becomes more robust as pressure is increased further. Moreover, the magnetotransport changes significantly in the vicinity of the OCP, indicative of the emergence of a new low-temperature electronic state. Finally, a model consistent with Weyl semimetallic behavior is found to best describe the temperature dependence of the low-temperature resistivity across the entire pressure range. The temperature window over which the model effectively reproduces the data is largest at the lowest pressures.

RESULTS

Magnetic properties

The main panel of Fig. 2a shows the magnetic susceptibility of $Sm_2Ir_2O_7$, measured in a [111] magnetic field of 0.1 T while warming after cooling in zero field (ZFC) and after cooling in the same 0.1 T field (FC). The temperature where the FC and ZFC data bifurcate, $T_{AIAO} = 110(2)$ K, coincides with the onset of the Ir AIAO ordered state, in agreement with the results of previous magnetic measurements⁷, resonant X-ray diffraction^{9,12} and muon-spin rotation³⁹ studies. The FC susceptibility data continue to rise as temperature is reduced further. A hump is seen in the data at temperatures just below 150 K, and an additional shoulder-like feature appears in the FC data centered at about 60–70 K. There is no strong evidence in these data for long-range magnetic order on the Sm sublattice below 10 K, as suggested by ref. ³⁹ from features in the muon-spin relaxation rate. Instead, our data are consistent with the presence of paramagnetic Sm moments at temperatures down to 2 K.

The inset of Fig. 2a shows the field dependence of the magnetization M(H) of a powder sample at low temperature. Figure 2b shows the extension of these data to high magnetic field. The high-field data were measured on a powder sample using extraction magnetometry⁴² at the National High Magnetic Field Laboratory, Los Alamos, in pulsed fields up to 60 T and temperatures down to 0.9 K. As the Ir moments are ordered below 110 K, we expect that these data are dominated by the paramagnetic Sm moments.



Fig. 2 Magnetic properties. a Magnetic susceptibility data (dark and light blue symbols) of a single crystal of $Sm_2lr_2O_7$ in an applied field of 0.1 T parallel to [111], measured as a function of temperature after cooling in zero field (ZFC) and with the field applied (FC). The Ir AIAO ordering temperature where the ZFC and FC data bifurcate, T_{AIAO} , is marked with an arrow. Inset shows the magnetic-field dependence of magnetization in a powder sample at 1.8 K. **b** Magnetization data (purple symbols) against field at 0.9 K for a powder sample measured to 60 T in a pulsed magnet. In both panels the black line is the result of single-ion calculations of the Sm^{3+} moment taking into account the crystal-field environment as described in the text.

Initial crystalline electric field (CEF) calculations using a pointcharge CEF Hamiltonian and the Stevens' operator formalism (see Supplementary Information, Supplementary Note 13), show that the Sm³⁺ ground-state doublet wavefunction is approximately $|J, m_I\rangle = |5/2, \pm 3/2\rangle$. The magnetic moment of this approximate ground state is expected to be 0.43 $\mu_{\rm B}$ at T = 0. Calculations of the single-ion properties for $Sm_2Ir_2O_7$ can be improved by employing intermediate coupling and allowing mixing of higher J levels, as implemented in the SPECTRE software⁴³. This method was used in ref. ⁴⁴ to obtain a set of CEF parameters for Sm^{3+} in $Sm_2Ti_2O_7$. Assuming the parameters therein for Sm₂Ti₂O₇ are a good approximation for $Sm_2Ir_2O_7$, we calculate that at T=0 K and $\mu_0 H = 0.5$ T, the moments exhibit a strong easy-axis anisotropy along the local [111] axis with $\mu_{\parallel [111]} = 0.233 \,\mu_{\rm B}$ and $\mu_{\perp [111]} = 0.001 \ \mu_{\rm B}$ leading to four non-colinear Ising axes in each Sm tetrahedron. The moment calculated in this way is around half the value of that from the $|5/2, \pm 3/2\rangle$ doublet alone. The discrepancy is largely due to a small admixture of terms from higher J levels in the ground state, especially the J = 7/2 level⁴⁴.

The black line in the main panel of Fig. 2a is the single-ion susceptibility for *H*//[111] calculated using the intermediate-coupling wavefunctions. In spite of the simplicity of the single-



Fig. 3 Resistivity vs temperature at near-ambient pressure. The temperature of the resistivity minimum, T_{min} , is marked with an arrow. The inset shows the result of differentiating the data with respect to temperature, which reveals a subtle kink or shoulder around 60 K, T_{kink} .

ion approximation, which neglects Sm–Sm as well as Sm–Ir interactions, we see that it captures the leading behavior of the ZFC susceptibility. In particular, it shows that the low-temperature Curie-like tail can be accounted for entirely by the Sm sublattice. It is no surprise that this calculation is unable to reproduce the hump seen below 150 K, as this feature is likely caused by domains of weak ferromagnetism associated with the Ir AIAO order, which have been proposed in other pyrochlore iridates^{38,45}. The solid black line in Fig. 2b is the powder-averaged single-ion calculation for the magnetization. The calculations predict that elevated fields mix the CEF-split energy levels, causing the Sm moment to become less Ising-like and grow significantly beyond its low-field value, consistent with the experimental data.

While dipolar interactions dominate the behavior of the rareearth ions in other spin-ice materials such as holmium or dysprosium titanate, the small size of the Sm moments suggests that they can be safely disregarded in the temperature and field regime of interest in our work. The same argument however does not apply to exchange interactions, and further work beyond the scope of the present paper is needed to ascertain the effects of Sm-Sm and Sm-Ir exchange couplings to the behavior of the magnetisation in an applied field. It is reasonable to expect the Sm ions to prefer to adopt an AIAO order at sufficiently low temperature, similar to $Ho_2Ir_2O_7$ and $Dy_2Ir_2O_7$, but they may favour a 2I2O configuration, depending on the sign and relative size of the Sm–Sm exchange interaction compared to the Sm–Ir interaction^{14,15}.

Transport properties under applied pressure - temperature dependence of resistivity

Figure 3 shows the temperature dependence of the resistivity of a single crystal of $Sm_2lr_2O_7$, measured in a diamond anvil cell as discussed in the Methods section and at approximately zero applied pressure (0.3(3) kbar).

The resistivity shows metallic behavior (defined as positive $d\rho/dT$) at high temperature, but undergoes a transition to a negative $d\rho/dT$ regime below $T_{min} = 110(3)$ K, where the resistivity exhibits a minimum. This transition is referred to as a metal-insulator transition in the literature. However, as discussed below, our data indicate that the low-temperature state cannot be described as a conventional insulator characterized by a gap at the Fermi energy. To avoid confusion, we refer to this region as 'non-metallic'. The resistivity minimum occurs at the same temperature as the Ir AIAO



Fig. 4 Resistivity vs temperature at high pressures. a Resistivity vs temperature for increasing applied pressures, plotted on a logarithmic *y*-axis. The temperature position of the resistivity minimum, T_{min} , is marked with arrows. **b** Values of T_{min} plotted against applied pressure, blue open circles. Data from the work of Liu et al.³³ are overlayed as closed black circles. **c** Resistivity extrapolated (via a quadratic fit below 8 K) to T = 0 K plotted against pressure. Note that the vertical axis is logarithmic. Dotted lines are intended as guides to the eye, highlighting a change of behavior around 60 kbar, close to the Ir AIAO QCP. Error bars show the uncertainties in determining pressure (see Methods) and in the values of T_{min} and ρ_0 .

magnetic transition seen in the magnetization, in agreement with the literature. Our resistivity data are consistent with previously reported low-pressure data on polycrystals by Liu et al.³³ and on pressed pellets by Matsuhira et al.³¹. We note that the anomalous results of Graf et al.⁴⁶, also on pressed pellets, who report a much lower transition temperature are not reproduced in other studies.

The temperature derivative of the resistivity reveals a subtle change of gradient or kink, which we denote T_{kink} , at around 60–70 K in both the 0.3(3) kbar and 1(1) kbar data sets (Fig. 3 Inset). This feature is not seen or resolved in our data for $p \ge 30$ kbar, but it is seen, after differentiation, in the data of Liu et al. across their 0–22 kbar pressure range, where it increases in temperature with applied pressure (see Supplementary Fig. 12).

Figure 4a shows the resistivity of the same $Sm_2Ir_2O_7$ crystal at each pressure measured, increasing from 0.3(3) kbar at the top, blue, to 87(2) kbar at the bottom, red. Applying pressure up to 87 kbar continuously reduces the magnitude of the resistivity across all temperatures (by a factor of 40 at room temperature and

190 at 2 K). The absolute value of resistivity is initially comparatively high (~10⁶ μ Ωcm) in the metallic state, suggesting that this material may not be best described as a simple coherent metal even above T_{min} .

A transition from metallic to non-metallic behavior upon cooling is observed at all pressures. Liu et al.³³ measured the pressure dependence of the resistivity of their polycrystal sample up to 22 kbar and reported a linear decrease in T_{min} over this range and this fits well with our results. However, just above the maximum pressure achieved by Liu et al., our data show a change in behavior: the value of T_{min} begins to increase again, in a linear trend, from approximately 30 kbar, as shown in Fig. 4b. This resistivity minimum is sharp and well defined up to 30(2) kbar, but becomes markedly broader and smoother at higher pressures, possibly indicative of a change from a phase transition to a crossover. Also at this point, the rise in resistivity for $T < T_{min}$ becomes less pronounced than that observed at lower pressures.

The nature of the electronic state below T_{min} in this family remains the subject of debate — several studies have compared the form of the low-temperature resistivity to various models of localized transport to attempt to shed light on this matter, see e.g. refs. ^{33,47-51}. In the present case, the functional form of the resistivity temperature dependence in the non-metallic phase below T_{min} proves challenging to describe with common simple models. To start, a simple Arrhenius-type activated exponential behavior falls very far from describing the data, which are much shallower in slope than an exponential. It is clear from inspection of the curves in Fig. 4a that although the magnitude of the resistivity increases dramatically at low temperatures, it does not appear to diverge at zero temperature. Extrapolating the curves to T = 0 results in a finite y-axis intercept ρ_0 for each pressure. We show the result of doing just this in Fig. 4c, with a quadratic function fit to the data below 8 K and extrapolated below their lower limit. The extrapolated ρ_0 value is seen to drop off rapidly with applied pressure up to around 60 kbar, the location of the AIAO QCP, after which it decreases at a much slower rate.

As discussed by Savary et al.²⁸, a theoretical treatment of the transport properties in the pyrochlore iridates is complex and challenging; no clear predictions have been published at this time and there is as yet no experimental consensus. In polycrystalline ${\rm Sm_2 lr_2 O_7}^{33}$, and in ${\rm Gd_2 lr_2 O_7}^{50}$, the low-temperature resistivity has been described with a $\rho \propto \exp{(T_0/T)^{1/4}}$ variable-range hopping expression — an activated insulating behavior typical of systems with a high degree of local disorder^{52–56}. Telang and Singh, in their study of the effect of Bi doping in Sm₂Ir₂O₇ pressed powders⁵¹, describe the undoped sample as obeying a 1/Ttemperature dependence typifying WSM behavior. This then reportedly evolves through a $T^{1/4}$ power law dependence tentatively attributed to quantum critical effects as doping suppresses the resistivity, to a ln(T) region interpreted as signifying quadratic-band-touching. In polycrystalline $Y_2 Ir_2 O_7$, a T^{-4} model⁴⁸ and a $T^{-3/2}$ model⁴⁹ were fit to the transport data, and attributed to a WSM state with thermally screened charged impurities, and with the presence of impurities and electron-electron correlations, respectively. Fits to all these models, and more, are shown in the Supplementary Information, Supplementary Notes 1-6, to compare their relative merits with respect to our single-crystal data. With the exception of the $T^{1/4}$ function, all of these models diverge at zero temperature, so it is unsurprising that they are not able to describe our data beyond highly restricted temperature ranges. (We point out that a $\rho(T) \propto T^{1/3}$ temperature dependence, shown in Supplementary Fig. 3, was found to give a markedly better fit to our data than any of these models previously employed in the literature. However, there seems to be no clear physical basis for this functional form and we just note its ability to fit the data).

The simplest physical model that can reproduce the behavior of our resistivity data is a combination of two parallel resistance channels, one which diverges and accounts for the non-metallic nature of the material and a second term which leads to a finite resistivity at zero temperature. The latter term could be any function that varies slowly compared to the diverging term, including $\rho \propto T$ or T^2 , but for simplicity we choose a temperature-independent value, set to be equal to the ρ_0 values determined from data extrapolation (shown in Fig. 4c). This term implies a metallic channel with impurity scattering is present in the sample that could potentially arise due to surface states or conduction along magnetic domain-wall boundaries, as suggested to be present in Nd₂lr₂O₇³⁶. (The strong, monotonic pressure dependence of ρ_0 rules out that it arises from an experimental artefact, such as an electrical short).

The best fits (see Supplementary Notes 3-5) to the data at all pressures were found using the model $\rho^{-1} = \rho_0^{-1} + (A/T)^{-1}$, where A is a (pressure dependent) constant, and are shown in Fig. 5a, b. As mentioned earlier, a $\rho \propto 1/T$ divergent term has previously been predicted for WSMs⁵⁷ and it describes our data particularly well at low pressures as can be seen in the inset to Fig. 5a. This shows the 0.3(3) kbar $\rho(T)$ data lying on a straight line for 2 < T < 55K when plotted versus 1/T after the ρ_0^{-1} contribution has been subtracted. The temperature range over which the model effectively reproduces the data is largest at the two lowest pressures, farthest from the QCP. This is indicated by the arrows in Fig. 5b, which show the upper limit of the region of validity of the fit (pragmatically set here at 2% divergence between fit and data). The coefficient A of the 1/T term also varies with pressure, as shown in Fig. 5c. Similar to ρ_0 , A is dramatically suppressed with initial pressurization, by two orders of magnitude up to the vicinity of the QCP at p_{cr} where a change of behavior is seen, and further reductions are much more gradual. Indeed, the pressure dependences of ρ_0 and A are found to agree strikingly well, see Fig. 5c, suggesting that the two conduction channels may be connected in some way.

In Sm₂Ir₂O₇, the lattice constant is known to decrease linearly with pressure across the whole range measured here, while the oxygen-coordinate parameter x changes only a little¹². The swift drop on increasing pressure up to 60 kbar in the parameters ρ_0 and A that quantify the non-metallic behavior at low temperatures indicates that the degree of localization in the AIAO Ir-ordered phase is highly sensitive to reductions in the lattice constant and the concomitant enhancement of the hopping parameter, which is consistent with the ground state being underpinned by electronic correlations. By contrast, in the state that emerges above the QCP, the localization of electrons is much reduced as compared to the behavior seen under ambient conditions, and further changes in pressure have only a small effect. The relative decoupling of this state from changes in the lattice constant suggests that correlations play a smaller role in the electronic ground state in this pressure regime.

Transport properties under applied pressure - hysteresis in magnetoresistance

At each pressure, we measured the magnetic-field dependence of the resistivity at various temperatures down to 1.5 K. The field was varied between \pm 15 T in the direction perpendicular to the plane of the resistivity measurement, which is along the diamond anvil cell axis and close to the [111] direction. Figure 6 shows that the low-temperature [111] magnetoresistance exhibits a marked and reproducible hysteresis, which persists for pressures below about 61 kbar. Also shown is the first field sweep performed on cooling in zero field from high temperatures (green line) which lies between the up and down sweeps of subsequent field sweeps. The strong hysteresis remains as temperature is increased up to about 50 K, but closes gradually at higher temperatures, and before the AIAO-ordering transition is reached, see Fig. 7. As can be seen in Fig. 7 and the Supplementary Note 9 (data for all



Fig. 5 Fits of the low-temperature resistivity. a Resistivity data (points) and fits (black line) to $\rho^{-1} = \rho_0^{-1} + (A/T)^{-1}$, as described in the text. Inset shows ρ with the ρ_0 contribution removed versus 1/T for p = 0.3(3) kbar, yielding a straight line for T up to 55 K. **b** The same as (**a**) plotted on a logarithmic vertical axis, to allow the higher pressure curves to be seen more clearly. Black arrows indicate the temperatures where each fit diverges from the data by 2%. **c** Pressure dependence of the fit parameter A, the coefficient of the 1/T term (red circles). Dashed lines are guides to the eye (note that the vertical axis is logarithmic), highlighting a change of character around 60 kbar close to the QCP. The pressure dependence of ρ_0 from Fig 4c is also shown overlaid as green squares for comparison. Error bars show the uncertainties in determining pressure (see Methods) and the standard error on the fit for A.

pressures) and Supplementary Fig. 18 (an interpolated colormap plot of hysteresis magnitude across the entire temperaturepressure range) of the Supplementary Information, this trend continues at elevated pressures until about 61 kbar, at which point the size of the hysteresis reduces dramatically, disappearing close to the AIAO QCP.

This hysteretic behavior is highly reminiscent of that observed in Ho₂Ir₂O₇ (see Supplementary Fig. 17 for a comparison of Fig. 6 with Ho₂Ir₂O₇ and Nd₂Ir₂O₇ data). In that case, a magnetic field close to the [1 1 1] direction acts to align the large Ho moments in a 311O/113O monopole-crystal arrangement which, via the Ho-Ir coupling, energetically favors one Ir AIAO domain over the other



Fig. 6 Hysteresis in MR. Magnetoresistance at T = 1.45(5) K and near-ambient pressure, showing a clear hysteresis loop. The field was applied along the approximate [1 1 1] direction, perpendicular to the plane of measurement. Data are normalized to their zero-field values. The green points denote the first application of the field up to +15 T after cooling the sample in zero field, then arrows around the subsequent blue data points show the evolution of the resistivity as the field is swept to -15 T and then back to +15 T again, at a rate of 1 Tmin⁻¹.

and alters the domain ratio in the sample. On sweeping down, the domain ratio remains constant until a negative field is applied, whereupon the second domain is favored and begins to grow preferentially. In Sm₂Ir₂O₇, the Sm ions have a significantly smaller moment than the Ho ions in Ho₂Ir₂O₇ (= $10\mu_B$). If the same effect is responsible for the hysteresis in both materials, this would explain why the magnitude of the hysteresis is lower in $Sm_2Ir_2O_7$ and also why the hysteresis is seen not to close even at the highest fields reached. At these temperatures, a field of about 4 T is sufficient to close the hysteresis in Ho2lr2O7 and achieve a steady 70:30% domain ratio (a value determined by the presence of strong pinning sites). By contrast, in Sm₂Ir₂O₇, the field required to saturate the moments and fully close the hysteresis loops is far greater than the 15 T achieved in these measurements; there are no high-field regions where the resistance becomes single-valued. Indeed, from the ratio of the moment sizes in the wo cases we would expect the field required to close the hysteresis to be roughly ten times larger in Sm₂Ir₂O₇ than in Ho₂Ir₂O₇. This also explains why the hysteresis loops shown by Pearce et al. appear smoother than found here for $Sm_2Ir_2O_7$ – the data in Fig. 6 access only the low-field region of the hysteresis loop, which likely extends out towards 100 T.

The sensitivity of the magnetoresistance in Ho₂Ir₂O₇ to changes in Ir-domain ratio arises because the monopole density varies differently with field in the two Ir-domain types, which in turn affects the monopole-quasiparticle scattering rate via two coupling mechanisms linking: (i) electronic spin and magnetic monopole charges; and (ii) electronic charge and the electric dipoles associated with monopoles. The validity of the magnetic monopole description in Sm₂Ir₂O₇ has not yet been definitively established and so it is not fully clear if the hysteresis has the same origin as in Ho₂Ir₂O₇; however we can estimate that mechanism (i) should be smaller for the low-moment Sm ions. On the other hand, mechanism (ii), which is driven by Sm–Sm interactions, could be of a similar size in both cases, as we expect the Sm exchange energy to be similar to or stronger than the equivalent in Ho₂Ir₂O₇.

An alternative, related effect is proposed to account for the hysteresis observed in $Nd_2Ir_2O_7$, which also has small rare-earth moments. Ueda et al. suggest that the current is preferentially carried through the Ir-domain walls and so the resistance



Fig. 7 Evolution of MR with pressure. Magnetoresistance ratio $\rho(B)/\rho(B=0)$ at fixed temperatures for pressures of (a) 1(1) kbar, (b) 43(2) kbar (c) 61(2) kbar and (d) 75(2) kbar. The field was applied along the approximate [111] direction, perpendicular to the plane of measurement. By 61 kbar the form of the hysteresis loop is changed, and the magnetoresistance is negative at low temperatures. Green points in (b) illustrate examples of binning and symmetrizing the data to obtain effective single-valued $\rho(B)$ data, separating out the hysteresis effects, as displayed for all data in Fig. 8. The data all have value 1 at B = 0, and are offset vertically for visibility.

responds to changes in the size and distribution of these as the field is swept³⁶. Which of these mechanisms is responsible for the effects seen in $Sm_2lr_2O_7$ remains an interesting open question, but what is clear is that the large hysteresis is characteristic of the AIAO phase and disappears as the QCP is approached. For

pressures above p_{cr} , some hysteresis is still seen in the magnetoresistance at the lowest temperatures (see Fig. 7c, d), but it is significantly smaller than that observed within the AIAO phase. Furthermore, the background form of the magnetoresistance undergoes a marked change at this pressure.

Transport properties under applied pressure - background magnetoresistance

In order to highlight the background form of the magnetoresistance of the sample separately from the hysteresis effects, one can average the multivalued resistivity at the same field value, to find an effective single-valued $\rho(H)$ for each pressure and temperature.

The low-temperature magnetoresistance data at all pressures show an asymmetry between positive and negative fields (Fig. 6) -an overall linear slope. Recent work has shown a strong, but non-hysteretic asymmetry in the low-temperature magnetoresistance of Eu₂Ir₂O₇ after field-cooling, caused by the formation of single domains of AIAO Ir order which remain robust up to 9 T^{58,59}. This is contrary to the situations in Nd₂Ir₂O₇ and Ho₂Ir₂O₇ which both show switching of domains at low fields and strong hysteresis in magnetoresistance that, as mentioned above, are attributed to coupling between the Ir ions and the Ising-like rareearth ions^{16,36}. As we have shown, the Sm ions in the present material are also Ising-like and the MR is strongly hysteretic with only a weak asymmetry, and so we expect Sm₂Ir₂O₇ to behave similarly to those materials. Instead, we attribute the asymmetry to a component of the Hall voltage mixing into the measured longitudinal resistivity. The magnitude of such a Hall signal is inversely proportional to the carrier density (for a simple single band at least). Here, the strongest slope is seen at low pressures where resistivity is high. To reliably extract absolute values of carrier density, a Hall-bar geometry on a lamellar sample with accurately placed contacts would be required; the small and irregularly shaped sample in our restrictive diamond anvil cell measurements precludes this treatment. To remove the Hall asymmetry, the data were symmetrized by averaging negative and positive field traces.

Examples of the resulting $\rho(H)$ plots are shown in Fig. 7b in green and in Fig. 8 (data for all pressures are shown in Supplementary Notes 7–9). With these data we can systematically describe the form and sign of the [111] magnetoresistance separately from the large hysteresis seen at low pressures. As can be seen from Fig. 8, the low-temperature, low-pressure background magnetoresistance first drops with field before rising as field is increased. A negative component is expected in the lowtemperature magnetoresistance of materials with locally fluctuating or disordered moments, such as a paramagnet or a Kondo system. As the field aligns the spins, spin scattering is reduced and resistance drops. In this material at low pressure and temperature, only the Sm³⁺ moments are free to align with the field, as indicated by magnetometry measurements. The behavior of the magnetoresistance in this region is reminiscent of that seen in $Ho_2Ir_2O_7$ under an applied [111] field¹⁶. In that case, there is also a negative magnetoresistance at low fields, caused by the alignment of the Ho³⁺ moments, followed by change to positive magnetoresistance as the effects of enhanced monopole scattering starts to dominate. As temperature is raised the magnetoresistance flattens out in both materials in similar way.

As pressure is increased, the background magnetoresistance retains the overall shape it has at low pressures, until the critical pressure for complete suppression of the Ir AIAO is approached. Here, the low temperature magnetoresistance becomes strongly negative across the whole field range (see Fig. 8) with a small amount of hysteresis (Fig. 7). As temperature is raised, the negative magnetoresistance diminishes, becomes weakly positive and then flattens out. The narrow hysteresis closes and disappears at the same temperature the MR changes sign. This form of the



Fig. 8 Single-valued MR at high pressures. Background magnetoresistance ratio $\rho(B)/\rho(B = 0)$ at fixed temperatures for pressures of (**a**) 1(1) kbar, (**b**) 43(2) kbar (**c**) 61(2) kbar and (**d**) 75(2) kbar. By 61 kbar the MR is markedly negative at low temperature. These data are extracted from Fig. 7 by binning and symmetrizing the data to remove ρ_{xy} components, and averaging up and down sweeps to separate out the hysteresis effects. The data all have value 1 at B = 0, and are offset vertically for visibility.

magnetoresistance continues for all pressures measured above 61(2) kbar.

There are several potential causes for positive magnetoresistive behavior. Simplest is the case of orbital motion of charge carriers in non-magnetic or antiferromagnetic (semi)metals with closed Fermi surfaces. Tafti et al. attribute the positive magnetoresistance in high-pressure $Eu_2lr_2O_7$ to this effect³⁷.

DISCUSSION

Figure 9 shows the pressure-temperature phase diagram of both the magnetic and transport properties of $Sm_2Ir_2O_7$ constructed by pulling together all of the features seen in our measurements, as well as prior results in the literature. The background colormap displays the ratio of the single-valued magnetoresistance at 15 T

and zero field, to distinguish regions of positive (blue) and negative (green) MR. Regions where the MR is flat, showing no variation in field, are shown in white on this scale. (See Supplementary Fig. 16 and Supplementary Note 12). The purple stars represent the temperatures at which the hysteresis in MR closes (see Supplementary Fig. 18) and are seen to follow closely the AIAO magnetic transitions at T_{AIAO} (orange triangles) found in the resonant X-ray measurements of Wang et al.¹². The curved solid line is a guide to the eye that connects these points.

The measured transport properties are clearly consistent with the pressure-induced suppression of the AIAO phase as manifested in three specific trends in the data. First, the crossover from metallic to non-metallic conduction, marked by the resistivity minimum at $T = T_{min}$, as reported by Liu et al.³³ (closed circles) and our low-pressure data (open circles), closely tracks the decrease in T_{AIAO} at all pressures up to 30 kbar. This is consistent with the interpretation of T_{min} as a Mott transition associated with the Ir ordering at low pressures^{12,19,20}. Second, the magnitude of the resistivity continuously decreases as pressure is raised (Fig. 4), and the extrapolated finite ρ_0 values drop rapidly with pressure up to 60(4) kbar, which is where AIAO Ir order is completely guashed. As mentioned earlier, the AIAO magnetic order is driven by the combination of U/t (Coulomb repulsion/hopping strength) and local Ising anisotropy¹². As pressure is applied, the lattice parameter decreases significantly, the 5d bandwidth and hopping strength increases, and U/t is suppressed¹². The combined result of pressure then is to stymy the AIAO ordering temperature and enhance conductivity. Third, the marked hysteresis seen in the magnetoresistance for magnetic field applied close to the [111] direction is present only within the AIAO ordered phase. This makes sense if, as discussed earlier, the hysteresis has its origin in the plastic deformation of the Ir domains via coupling with the Sm³⁺ moments, equivalent to that reported in Ho₂Ir₂O₇ or Nd₂Ir₂O₇. Once the AIAO order is suppressed, the domains are no longer present and the large magnetoresistance hysteresis disappears. All these magnetotransport features affirm the existence of the same quantum critical point at $p_c = 63$ kbar suggested by Wang et al. from their X-ray data.

Despite these connections between the Ir AIAO order and the measured transport properties, an important and clear result of our experiments is that $Sm_2lr_2O_7$ remains metallic at high temperatures and non-metallic at low temperatures below a finite T_{min} for the entire pressure range measured. In particular, the transition marked by the resistivity minimum at T_{min} is not suppressed to zero temperature along with the Ir AIAO order to yield metallic behavior at all temperatures, as was naively anticipated from the coincidence of the two transitions at low pressures and from the pressure dependence of T_{min} in Nd₂Ir₂O₇^{19,25,33}.

As mentioned, T_{min} initially decreases, following the phase line of the Ir AIAO order observed by Wang et al.¹². However, above 30 kbar this trend reverses and T_{min} increases linearly up to the highest pressures measured. It seems that some additional mechanism is causing a resurgence of localization of the charge carriers once the pressure pushes the AIAO transition below a certain temperature. This new localization mechanism then strengthens with further application of pressure. Interestingly, an extrapolation of the high-pressure linear trend of $T_{\min}(p)$ back to low pressures, (dashed black line) coincides with the position of the subtle change in dp/dT observed in our sample at T_{kink} . This feature is also present in the low-pressure data of Liu et al. (closed squares, see also Supplementary Fig. 12) — albeit at a slightly lower temperature than in our data — and is seen to increase linearly with pressure at a gradient equal to the high-pressure dependence of T_{min} . We note that a similar feature is observed in the resistivity of Eu₂Ir₂O₇, where it also appears in the heat capacity³⁸. There it is shown to be most clear in the purest crystals with smallest deviations from Eu and Ir stoichiometric values,



Fig. 9 Pressure-temperature phase diagram for Sm₂Ir₂O₇. The background colormap displays interpolated values of the magnetoresistance (MR) ratio and is indicative of the size of the MR. Temperatures of the resistivity minimum T_{min} are plotted as open circles. Previous data from the literature up to 20 kbar³³ are overlaid as closed circles Above about 30 kbar T_{min} shows a linear increase with pressure (solid line), which can be extrapolated back to ambient pressure (dotted line) where it coincides with the kink feature observed in the gradient of ρ (T) at the lowest pressures. These are plotted (open squares) along with similar features found in the data of Liu et al.³³ up to 20 kbar (closed squares). Triangles denote the Ir AIAO ordered phase observed by Wang et al.¹² and star symbols represent the temperatures at which the magnitude of the hysteresis in the MR extrapolates to zero. The curved solid line is a guide to the eye through these points. The green shaded region highlights a regime around the QCP with a negative background magnetoresistance. The diamond symbols denote the temperatures where the magnetoresistance switches from negative to positive, which bound this region. Error bars show the uncertainties in determining pressure (see Methods) and estimated uncertainty for their corresponding temperature values.

suggesting in this case that it is an intrinsic effect that gets smeared out in lower quality samples. Possible origins for this kink feature include low-temperature structural and/or magnetic variations, and we note again that a shoulder-like feature appears in the field-cooled susceptibility data close to $T_{\rm kink}$ (see Fig. 2).

Whatever the cause, it is apparent that T_{min} and T_{AIAO} separate for pressures above 30 kbar, and the Ir AIAO order ceases to be the driver of the non-metallic state when the temperature scale associated with the new, as yet unidentified, mechanism begins to dominate. This explains why the material remains non-metallic once the AIAO order is suppressed at the $p_c = 63$ kbar QCP. For pressures between 30 and 63 kbar, the Ir ordering still takes place below T_{AIAO} . The main markers of the AIAO ground state in the transport in this intermediate pressure region are the large magnetoresistive hysteresis discussed earlier and the strong negative pressure dependence of the extrapolated ρ_0 value.

A new form of magnetotransport emerges around the QCP. The large hysteresis and competition between negative and positive magnetoresistance components, seen at low temperatures in the AIAO phase, now changes to a region of weakly hysteretic but strongly negative magnetoresistance that extends from just below p_c to beyond the highest pressure measured (green region in Fig. 9). The negative magnetoresistance evolves slowly to a weakly positive response as the temperature is raised above about 5 K,

disappearing altogether at elevated temperatures. At around the same temperature the weak hysteresis closes gradually (as can be seen more clearly in Supplementary Figs. 18 and 15). The open diamond symbols in Fig. 9 illustrate the temperatures where the MR is zero (see Supplementary Figs. 16 and 19). According to Wang et al., while the long-range Ir AIAO order is suppressed above p_{c} , the axial anisotropy is preserved, leaving behind weakly correlated or paramagnetic Ising-like moments¹². Any lowtemperature magnetic order on the Sm sublattice is also predicted to be lost once the Ir order is gone. Alignment of these fluctuating spins in an applied magnetic field could account for the negative magnetoresistance observed above the QCP. The positive magnetoresistance at higher temperatures is consistent with orbital effects arising from small Fermi-surface pockets, such as might be expected for a disordered semimetal and, as mentioned above, a similar origin was proposed for the high-pressure magnetoresistance seen in $Eu_2Ir_2O_7^{37}$.

The precise origin of these new electronic properties remains to be determined. Several theoretical proposals have been made for the electronic phases that might exist in the vicinity of the AIAO $QCP^{20,28,40,41,60}$. These include a WSM phase expected to be located in a relatively narrow region of Coulomb *U* between the AIAO Mott insulator and the paramagnetic-metal phase (previously believed to occur close to where AIAO order is suppressed)^{12,61,62}, and tentatively reported in the results of Telang and Singh⁵¹. However, as Wang et al. point out from their estimates of the shift in hopping *t* required to realize the QCP in $\text{Sm}_2\text{lr}_2\text{O}_7$, this theoretically predicted narrow region where the WSM exists could span the entire pressure range measured¹². Our analysis of the single-crystal transport data finds them to be well described by a 1/*T* functional form, which is indeed consistent with models of WSMs⁵⁷, albeit with the inclusion of a parallel slowly-varying (relative to 1/*T*) conduction channel, possibly arising from surface or domain-wall conduction. The temperature range over which this 1/*T* dependence is valid is found to be largest at the lowest pressure, away from the QCP.

Another theoretical prediction is that the OCP in Sm₂lr₂O₇ separates the WSM from a non-Fermi-liquid semimetal phase distinguished by quadratic-band touching (QBT)²⁸. Indeed, ARPES measurements suggest that this is the ambient pressure phase exhibited by $Pr_2Ir_2O_7$ at low temperatures²¹ and by $Nd_2Ir_2O_7$ in its metallic state at elevated temperatures²². It should also be pointed out that optical conductivity spectra for Sm₂lr₂O₇ point towards a semimetal nature⁶¹, and are distinct from the finite gap seen in these data for Eu2lr2O7. Recent work on Pr2lr2O7 has additionally reported topological Weyl semimetal states under the application of magnetic field⁶³. A recent paper reporting transport measurements on Sm₂Ir₂O₇ speculates that the WSM is present at ambient pressure and that doping more than 5% of Bi on the Sm site leads to a transformation to the QBT phase, although the evidence in favor of this interpretation is limited⁵¹. Our data also do not provide evidence for QBT behavior emerging under pressure; however, we highlight again that our experiments reveal that the paramagnetic metallic phase is not realized once AIAO order is destroyed and instead another, as yet mysterious, electronic phase emerges.

Our data share certain similarities with work done by Tafti et al. on Eu₂Ir₂O₇, neighbor to Sm₂Ir₂O₇ in the chemical series of this family³⁷. While the T_{min} in that material does not appear to shift very much at all with applied pressure, the authors do find that the resistivity drops significantly as pressure is increased. In the low-pressure region, they can fit their $\rho(T)$ data to an activated model at low temperatures (which does not work for Sm₂Ir₂O₇). The size of the insulating gap in Eu₂Ir₂O₇ drops swiftly towards zero at 80 kbar and then remains low and slowly varying upon further increase in pressure, reminiscent of the A coefficient we extract from the 1/T fits for our $Sm_2Ir_2O_7$ data. The authors describe their high-pressure, low-temperature state as a novel metal with resistivity that increases on cooling. In contrast to our results, Tafti et al. find that the magnetoresistance of Eu₂lr₂O₇ is positive down to the lowest temperatures in the high pressure region³⁷. It should be noted that the AIAO phase transition has not yet been tracked with pressure in the Eu material. Tafti et al. also find that the resistivity at high temperatures and low pressures in Eu₂lr₂O₇ increases on cooling. This is in contrast to the more metallic behavior (i.e. decreasing resistivity on cooling) observed at ambient pressure in single-crystal measurements of other pyrochlore iridates $Ln_2Ir_2O_7$, including $Ln = Pr^{29}$, Nd³⁶, Sm (our data), Ho¹⁶ and even in other studies of the same Eu material considered by Tafti et al.³⁸, as well as in some pressed-powder measurements, e.g. Pr³¹, Nd^{19,31,64} and Sm⁴⁶. We suggest that a metallic temperature dependence above \mathcal{T}_{min} is the true intrinsic behavior of the pyrochlore iridates in this region of the phase diagram.

We point out that another origin has been proposed for the metal insulator transition seen in the pyrochlore osmate $Cd_2Os_2O_7$. This material is a well-compensated, low-carrier density semimetal in its high-temperature paramagnetic phase. At 227 K, the AIAO magnetic transition on the Os sublattice is suggested to shift the electron and hole bands by a relatively small, but sufficient amount to cause a Lifshitz-type transition in this finely balanced semimetal, opening a small band gap^{65,66}. The transport

below the transition is governed by the opening gap, a number of in-gap impurity states caused by oxygen non-stoichiometry, and the presence of conduction channels in magnetic domain walls (as already mentioned above in the context of $Nd_2lr_2O_7$). It is interesting to draw parallels with our material, which also shows non-activated behavior and a non-diverging resistivity. Additional Hall effect measurements would be required to distinguish between this Lifshitz mechanism and the Mott transition more commonly attributed to the pyrochlore iridates. Such measurements are planned but require a further refinement of the synthesis to obtain larger high-quality single crystals than are currently available.

In summary, our data show a significant change in the electronic properties of Sm₂Ir₂O₇ as the AIAO Ir magnetic order is gradually destroyed by pressure. Although the metallic-to-nonmetallic crossover at $T = T_{min}$ is not suppressed in tandem with the magnetic transition, the magnitude of the resistivity, the extrapolated finite T = 0 resistivity, and the coefficient of the 1/Tresistivity component all drop swiftly as the OCP is approached. then switch over to a much reduced rate of reduction with further increasing pressure. The 1/T temperature dependence, which is predicted for a Weyl semimetal, fits the data up to 55 K at 0.3 kbar, but this range is reduced at higher pressures. While it is not currently possible from these data alone to precisely identify the electronic state that emerges beyond the QCP, we have measured both transport and magnetotransport data across the entire phase diagram in order to provide information to guide theoretical studies. Our results also act as a basis for future experimental efforts, with optical conductivity and angle-dependent magnetotransport particularly well suited to shedding light on the possible Weyl semimetal and anisotropic, guadratic-band-touching phases.

METHODS

Crystal growth

Single crystals of $Sm_2lr_2O_7$ were grown by the flux method, using KF as the flux, as described in ref. ⁶⁷. They were characterized and verified to be of good crystalline quality by X-ray diffraction.

High pressure transport measurements

Magnetotransport data were carried out in a diamond anvil cell^{68,69} with non-magnetic BeCu body and hardened BeCu gasket. The culet of the diamonds used was 1 mm and the gasket hole was drilled to 0.5 mm. The cell was pressurized at room temperature for each pressure point, then measured on a homebuilt probe loaded into a Oxford Instruments 17 T cryomagnet fitted with a variable temperature insert. A single crystal was prepared for resistivity measurements in the standard 4-wire geometry with gold wires bonded to the crystal with DuPont 6838 silver-loaded epoxy. Glycerol was chosen as the pressure transmitting medium due to its favorable compressibility and hydrostatic properties. Across the pressure range of interest, glycerol is known to perform well in comparison to other choices of media⁷⁰. Making diamond anvil cell measurements in pressures close to 100 kbar inevitably brings some risk of a degree of nonhydrostaticity; nevertheless we are confident that extrinsic effects of the pressure-transmitting medium can be ruled out as an explanation of all the main features in our data. The pressure was measured via the fluorescence spectrum of a chip of ruby placed within the pressure region⁷¹. Typical pressure uncertainties, as denoted on values given in the text are estimated from two factors. Firstly, there is a gradient or distribution of pressure within the sample volume, which we quantify via the broadness of the ruby fluorescence peak and its fitting uncertainty. Secondly, previous calibrations on identical cells have shown pressure losses upon cooling, due to thermal expansion. The resulting uncertainty from these two factors we determine as ± 1 kbar below 10 kbar

and ± 2 kbar above, noting that the pressure will always be above 0 kbar. The magnetic field was applied along the cell axis, a direction close to the crystallographic [111]. The crystal was sat on the diamond anvil faces and the current applied in the plane of the anvil faces, perpendicular to the field.

Magnetization measurements

Low-field DC magnetization measurements were carried out on a powder sample in a Quantum Design MPMS SQUID magnetometer system. The sample was then additionally measured in up to 60 T of applied field via extraction-coil magnetometry at the NHMFL, Los Alamos. As this technique does not return absolute values of the magnetization, these were obtained via calibrating the low-field slope up to 7 T with the MPMS data at 1.8 K. There is no discernible difference between the latter and the pulsed field data at 1.5 K and at 0.9 K shown in the manuscript.

DATA AVAILABILITY

Data presented in this paper are made available online at wrap.warwick.ac.uk/ 182307.

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AUTHOR CONTRIBUTIONS

P.A.G. and D.P. conceived the experiments. D.P. grew and characterised the crystal. M.J.C. prepared the pressure cell. M.J.C. and K.G. ran the magnetotransport and SQUID measurements. J.P.T. carried out a structural analysis. J.S. carried out the pulsed-field experiments. T.N.D.L.F., C.C. and A.T.B. conceived and performed the crystal-field calculations. M.J.C. analysed the data. M.J.C. and P.A.G. prepared the manuscript with input from all coauthors. P.A.G., D.P., A.T.B. and C.C. supervised the project.

COMPETING INTERESTS

The authors declare no competing interests.

ADDITIONAL INFORMATION

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