Extraordinary phase coherence length in epitaxial halide perovskites

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Highlights
Epitaxial halide perovskites with extraordinary quantum phase coherence
Quantum transport properties with weak antilocalization observed in tetragonal CsSnI₃
Demonstration of quasi-2d charge carrier behavior with spin-orbit coupling
Epitaxial halide perovskites emerging materials for quantum electronic applications

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Extraordinary phase coherence length in epitaxial halide perovskites

Kostyantyn Nasyedkin,1,4,5 Isaac King,2,5 Liangji Zhang,1,5 Pei Chen,2 Lili Wang,2 Richard J. Staples,3 Richard R. Lunt,1,2,* and Johannes Pollanen1,6,*

SUMMARY
Inorganic halide perovskites have emerged as a promising platform in a wide range of applications from solar energy harvesting to computing and light emission. The recent advent of epitaxial thin film growth of halide perovskites has made it possible to investigate low-dimensional quantum electronic devices based on this class of materials. This study leverages advances in vapor-phase epitaxy of halide perovskites to perform low-temperature magnetotransport measurements on single-domain cesium tin iodide (CsSnI3) epitaxial thin films. The low-field magnetoresistance carries signatures of coherent quantum interference effects and spin-orbit coupling. These weak anti-localization measurements reveal a micron-scale low-temperature phase coherence length for charge carriers in this system. The results indicate that epitaxial halide perovskite heterostructures are a promising platform for investigating long coherent quantum electronic effects and potential applications in spintronics and spin-orbitronics.

INTRODUCTION
Halide perovskite semiconductors are being developed worldwide in numerous thin-film electronic applications due to their exceptional properties. Initially these materials were investigated for light applications (Hattori et al., 1996) and subsequently for their photovoltaic properties (Lee et al., 2012b; Nie et al., 2015; Yang et al., 2017), long carrier diffusion lengths (Saidaminov et al., 2020), and exceptional emission properties (Gao et al., 2020; Xu et al., 2019). More recently, thin films of CsSnX3 (X = Cl, Br, I) have also been used for lasers (Wang et al., 2015) and light-emitting diodes (LEDs) (Kazim et al., 2014; Yuan et al., 2016). Given their high-quality and flexible crystal structure, it is natural to ask if halide perovskites might serve as a novel platform for creating quantum two-dimensional electron systems (2DESs). However, despite being considered as “defect-tolerant” films when grown from solution, there is still a need to transition these materials from polycrystalline layers to epitaxial single-crystal layers to enable the study of delicate and emergent quantum many-electron phenomena. By reducing the defect density in the perovskite layer, controlling strain and crystal phase, and improving crystalline order, electronic properties can be enhanced and quantum devices can be developed. Indeed, the ability to exploit epitaxy and epitaxial strain in silicon (Si) and III-V semiconductors revolutionized modern computing and optoelectronics and led to improved photovoltaics (Bertness et al., 1994; Dai et al., 2015; King et al., 2007; Lee et al., 2012a; Takamoto et al., 1997), LEDs (Ghosh et al., 1986; Huang et al., 1992; Kuo et al., 2010; Peng and Wu, 2004), as well as 2DESs with ultra-high carrier mobility (Dingle et al., 1978; Hatke et al., 2017; Manfra, 2014; Pfeiffer et al., 1989; Umansky et al., 2009). For example, strained Si is now commonly found in every MOSFET (Sun et al., 2007) due to the increased interatomic distance in the silicon layer that nearly doubles the mobility. Additionally, molecular beam epitaxy (MBE) can now produce GaAs/AlGaAs heterostructures having charge carrier mobilities \( \mu > 3 \times 10^6 \text{cm}^2/\text{V} \cdot \text{s} \) (Umansky et al., 2009) hosting a wide variety of exotic collective phases of quantum electronic matter (Eisenstein and Stormer, 1990). Advances in epitaxial growth have also benefitted a wide range of materials beyond low-bandgap semiconductors including graphene, diamond, SiC, and oxide perovskites just to name several.

The ability to precisely grow epitaxial halide perovskite thin films, as well as halide perovskite quantum wells has recently been demonstrated (Chen et al., 2017, 2019; Oksenberg et al., 2020; Wang et al., 2017a, 2017b, 2019). The vapor phase epitaxy of our CsSnI3 samples was achieved on lattice-matched metal halide crystals with congruent ionic bonding (see Figure 1A) (Hu et al., 2017; Wang et al., 2017b, 2019). The heteroepitaxy of these perovskite films allows for precise film thicknesses, controllable phase and
orientation, integration into quantum wells, and opens the door to quantum transport devices. In fact, perovskite films have been shown to exhibit numerous quantum phenomena at low temperatures, such as superconductivity (Reyren et al., 2007), ferroelectricity (Cohen, 1992), and quantum coherent transport (Caviglia et al., 2010; Herranz et al., 2004; Keshavarz et al., 2017; Wang et al., 2020; Zhang et al., 2021), the last of which we show in this paper for epitaxial CsSnI3 thin films. In particular, we exploit the growth of epitaxial halide perovskite devices to perform low-temperature quantum transport measurements on thin films composed of CsSnI3. These measurements reveal quantum coherent transport of charge carriers in this halide perovskite 2DES, as well as the presence of a spin-orbit crystal field.

In low-dimensional electronic systems quantum interference of charge carriers leads to a variety of coherent phenomena, including Aharonov-Bohm oscillations (Washburn and Webb, 1986), universal conductance fluctuations (Lee et al., 1987), as well quantum interference induced localization of charges (Altshuler et al., 1980; Bergmann, 1982; den dies et al., 1981; Pierre et al., 2003). Of particular interest are the effects of weak localization (WL) (Altshuler et al., 1980) and weak anti-localization (WAL) (Bergmann, 1982) as they can be used as tools to determine the phase coherence length for charge carriers in low-dimensional devices by performing magnetoresistance experiments at low temperature. Given a sufficiently long phase coherence length, the diffusive trajectory of a charge carrier in a low-dimensional system can interfere with its time-reversed partner. When the interference is constructive, it produces WL, which manifests as a reduction in the electrical conductivity near zero magnetic field. In contrast, the presence of sufficiently strong spin-orbit coupling produces an additional rotation of the charge carrier spin and leads destructive interference, i.e. WAL. In this work, we have observed clear signatures of phase coherent transport and WAL in the low-field magnetotransport measurements of epitaxial CsSnI3. These results allow us to extract a phase coherence length of charge carriers in the device, which significantly exceeds the thin film thickness and indicates the low-dimensional character of the system.

RESULTS AND DISCUSSION
The epitaxial CsSnI3 crystalline film samples were grown in a custom Angstrom Engineering thermal evaporator by co-evaporation from two tungsten boats containing precursor materials (CsI and SnI2), with each
source having an independently calibrated quartz-crystal-microbalance rate monitor and source shutter. A 50 nm thick film was deposited stoichiometrically at a rate of 0.007 nm/s on a cleaved [100] surface of a potassium chloride (KCl) single crystal substrate. The growth was performed at pressures less than 3×10⁻⁶ torr and a temperature of 22 °C. In situ crystal analysis was performed in real-time using reflective high-energy electron diffraction (RHEED) to determine structure and film quality. The RHEED scans were performed at 30.0 kV and an emission current of less than 50 nA to reduce damage and charging on the perovskite film during growth. These measurement conditions no damage was observed over typical deposition times of up to 1-2 hr. Gold contacts (50 nm thick) were deposited on the perovskite layers using electron-beam evaporation at a rate of 0.02 nm/s through a shadow mask in the same deposition chamber (with the same temperature and pressure).

The resulting CsSnI₃ epitaxial film is pseudomorphic to the KCl substrate (a = 0.629 nm) as shown in Figure 1A and confirmed by rotation-dependent RHEED patterns (Figure 1B), as well as X-ray pole figure analysis (Nagai, 1974) and TEM (see Figures S1–S3). The streaky patterns observed are indicative of smooth growth compared with spotty patterns seen in rougher films (see Figures S4 and S5). From the cleaved metal halide crystals, we observe atomic terrace lengths of 1-5 microns. Since the length of the streaks for the pseudomorphic epitaxial layer is similar to the streak lengths from the substrate, this indicates that the terrace length is simply limited by that of substrate. The patterns we observe are representative of high quality epitaxial and heteroepitaxial oxide perovskite layers (Choi et al., 2012; Soukiassian et al., 2008). We note that it is possible that a small amount of oxygen and water impurities could be incorporated into the epitaxial thin film despite the overall low pressure during growth. In fact, future investigations systematically varying the growth pressure could be conducted to assess the impact of the base pressure on the resulting transport properties of the epitaxial thin film.

The RHEED data show a clear change in the symmetry from face-centered-cubic (FCC) of the substrate to a primitive perovskite cell by the emergence of (01) streaks between the (02) streaks of the substrate. The crystal structure of epitaxial CsSnI₃ is similar to the high temperature cubic phase ([Wang et al., 2017b]) but tetragonally distorted due to the in-plane tensile strain resulting in c = 0.612±0.002 nm (see Figure S6 and Table S1). Thus, this epitaxial phase and band structure are unique from the commonly observed orthorhombic phase at room temperature ([Wang et al., 2017b]).

Two 2.0 mm × 5.0 mm gold pads spaced by 50 μm on the CsSnI₃ layer enabled standard low-frequency (10 Hz) ac electrical transport measurements as shown in Figure 1C. While the epitaxial perovskite film is more stable than the bulk orthorhombic phase in air, it too degrades when exposed to air after many hours, so protocols were developed to keep the devices in a high vacuum or dry-nitrogen filled environment, similar to a glovebox, during sample fabrication, transferring, and measurement. Additionally we have characterized the air sensitivity of epitaxial CsSnI₃ using X-ray, optical, and electrical transport techniques (see Figure S7). We note that it is conceivable that for particular future applications the material could be encapsulated in epoxy or edge sealed in an inert atmosphere after growth to prevent degradation in air. Alternatively, the material could be kept under vacuum, which is the typical environment for cryogenic quantum devices at milliKelvin temperatures. Devices for the quantum transport measurements reported here were mounted in a custom indium o-ring sealed copper sample holder (see Figure S8), to minimize exposure to air, and thermally anchored to the mixing chamber of a dilution refrigerator having a lowest temperature of T ~ 10 mK. A variable magnetic field B perpendicular to the plane of the CsSnI₃ epitaxial film was supplied by a superconducting solenoid and enabled the measurement of the magnetoconductivity of the device up to B = ±13.5 T.

Before investigating the magnetotransport, we first characterized the temperature dependent electrical properties of the device in the absence of a magnetic field (for comparison we present measurements on additional CsSnI₃ devices which show consistent behavior as discussed below. Similar measurements on epitaxial CsSnBr₃ devices were also performed. See Figure S9). In Figure 2 we show the conductivity, σ, of a CsSnI₃ epitaxial film as the sample was cooled from room temperature down to T = 16 mK. The resistance was measured using standard lock-in techniques and the conductivity was calculated from the measured device resistance by taking into account the geometric factor between the gold pads. The low overall measured resistance indicates that the Au makes ohmic contact to the CsSnI₃, consistent with previous electrical measurements on bulk CsSnI₃ ([Han et al., 2019]). Since the total measured resistance is given by R_{tot} = 2R + R, where R is the intrinsic sample resistance, these measurements place an upper
bound of \( R_c \leq 10 \, \Omega \) on the contact resistance between the gold pads and the thin film and indicate good electrical contact between the two. The conductivity of the device increases with decreasing temperature indicating the presence of mobile charge carriers that do not freeze out upon cooling to low temperature. This increase in the conductivity is consistent with a reduction in phonon density upon lowering the temperature of the device. The existence of charge carriers in the material is likely the result of intrinsic doping in the epitaxial thin film, which is a direct bandgap semiconductor (Peedikakkandy and Bhargava, 2016) (see Figure S10 and Table S2). In fact, previous electrical conductivity measurements on orthorhombic CsSnI\(_3\) have also reported metal-like conduction attributable to hole-doping associated with Sn vacancies (Chung et al., 2012). We note that it is possible the magnitude of the low temperature conductivity shown in Figure 2 is limited by the contact resistance at the Au/CsSnI\(_3\) interface even though there is clearly good ohmic contact. This would impact the shape of the temperature dependence of \( \sigma \), as well as its maximum value but would not change the conclusions we draw regarding phase coherent transport nor the existence of spin-orbit coupling described below.

At low-temperatures, the application of a magnetic field normal to the plane of the CsSnI\(_3\) layer (see Figure 1C) strongly modifies the transport of charge carriers. In Figure 3A we present the conductivity of the CsSnI\(_3\) epitaxial film device as a function of magnetic field up to \( \pm 13.5 \, T \). Above several Tesla we observe a non-saturating magnetoconductivity that is approximately linear in the applied field. This apparent reduction in the conductivity with increasing magnetic field likely arises from a relatively mundane source, i.e. from a convolution of the longitudinal conductivity measurement by the Hall voltage across the sample due to the geometry of our device. In fact, this type of effect is common in magnetotransport measurements such as ours and has been observed in 2DESs in semiconductors (Fang and Stiles, 1983; Russell et al., 1990), as well as graphene (Skachko et al., 2010) and is associated with the formation of so-called “hot-spots” near corners of device contacts (Klaβ et al., 1991; Russell et al., 1990; Skachko et al., 2010).

More importantly, the low-field magnetoconductivity can be used to reveal signatures of coherent quantum phenomena. In the vicinity of \( B=0 \) we observe clear signs of quantum interference. The inset of Figure 3A shows the change in the magnetoconductance \( \sigma(B) - \sigma(0) \) near zero magnetic field and \( T = 16 \, mK \). These data exhibit a cusp-like maximum that broadens and weakens as the temperature is increased as shown in Figure 3B. As we describe below, these features are the characteristic signatures of WAL, i.e. phase coherent diffusive magnetotransport in the presence of spin-orbit coupling (Bergmann, 1982; Lee and Ramakrishnan, 1985).

The Hikami-Larkin-Nagaoka (HLN) theory of WAL provides a description of the magnetoconductivity behavior of quasi-2D systems in the quantum diffusive transport regime (Hikami et al., 1980). We utilize
a simplified empirical HLN formula to analyze our data and to extract a phase coherence length \( L_F \) for the charge carriers. This model contains two fit parameters \( L_F \) and \( a \), which accurately capture the physics of WAL in our data. In this analysis, the change in magnetoconductivity away from its zero field value is given by,

\[
\sigma(B) - \sigma(0) = \frac{a}{\pi} \left[ \Psi\left(\frac{1}{2} + \frac{\hbar}{4eBL_F^2}\right) - \ln\left(\frac{\hbar}{4eBL_F^2}\right) \right] \tag{Equation 1}
\]

where \( e \) is the electron charge, \( \hbar \) is the reduced Planck’s constant and \( \Psi(x) \) is the digamma function. The sign of the coefficient \( a \) indicates the type of localization (He et al., 2011; Kurzman et al., 2011). Specifically, \( a > 0 \) is associated with WL while \( a < 0 \) indicates WAL. For the entire range over which we observe phase coherent transport (\( T < 15K \)) we find a negative value of \( a \), which demonstrates the presence of spin-orbit coupling in the epitaxial CsSnI\(_3\) film. Representative fits of the data to Equation 1 at different temperatures are shown in Figure 3B. The magnitude of spin-orbit coupling in a material is dictated by the mass of the constituent atoms in the crystal and whether the crystal lacks inversion symmetry. Inversion symmetry is broken if a compound is intrinsically non-centrosymmetric and results in Dresselhaus spin-orbit coupling (Dresselhaus, 1955). It is not known whether the epitaxial phase of CsSnI\(_3\) intrinsically lacks an inversion center as it is nearly impossible to perform precise atomic refinement on an epitaxial thin film. Regardless, the presence of the interface between the KCl substrate and the CsSnI\(_3\) epitaxial film breaks inversion symmetry and could lead to a Rashba spin-orbit field (Bychkov and Rashba, 1984) along the growth direction of the heterostructure. In fact, it has been shown that the large Rashba splitting observed in tetragonal MAPbI\(_3\) is likely not a bulk- but rather a surface-induced phenomenon (Frohna et al., 2018). Alternatively, it is possible that the spin-orbit coupling in CsSnI\(_3\) that we observe arises from the relatively large mass of the constituent atoms in the crystal. We note that our quasi-two-terminal measurements do not provide a measure of the absolute sample conductivity, which could be affected by the contact resistance. Therefore in our analysis the magnitude of \( a \) serves only as an overall scale factor needed to accurately fit the data and to extract the phase coherence length. However, as described above, the sign of \( a \) does carry intrinsic physical information, i.e. that spin-orbit coupling is present in the system. Additionally we emphasize that in this analysis.
\( \alpha \) and \( \ell_F \) are not meaningfully dependent on one another. This can be seen in Equation 1 where \( \alpha \) is an overall scaling pre-factor that cannot simply be subsumed into the digamma and log functions to compensate for a change in \( \ell_F \).

The temperature dependence of the phase coherence length extracted from these measurements is shown in Figure 4 along with value of \( \ell_F \) for other low-dimensional quantum electronic materials in the quantum diffusive transport regime. For our data on epitaxial CsSnI\(_3\) each value of \( \ell_F \) corresponds to the average value obtained from multiple measurements at each temperature and the error bars are the standard deviation. As the temperature is increased we observe a reduction in \( \ell_F \) due to increased inelastic scattering of charge carriers. Several important inelastic scattering mechanisms can result in the loss of phase coherence including electron-electron scattering, electron-phonon scattering, and scattering from magnetic impurities (Lin and Bird, 2002). In the quantum diffusive regime \( \ell_F = \sqrt{D} \tau_\phi \), where \( D \) is the charge carrier diffusivity and \( \tau_\phi \) is the rate of dephasing scattering events. The temperature dependence of \( \tau_\phi \) carries information regarding which of the scattering mechanisms dominates; however multiple competing mechanisms often lead to mixed temperature dependences that are difficult to disentangle (Lin and Bird, 2002). We find that \( \tau_\phi \) scales as \( 1/T^p \) with \( p \approx 2 \), which is consistent with phonon scattering as the dominant contributor to dephasing with increasing temperature. In fact, previous studies on semiconductors and metals have reported electron-phonon scattering as leading to \( p \approx 2 \) – 4 (Lin and Bird, 2002). Regarding the role of magnetic scattering, while we do not purposefully introduce magnetic defects/impurities (e.g. Fe, Mn, etc.), magnetic scattering is not impossible in our devices. Despite the purity of the source materials being high (CsI 99.9% (Sigma-Aldrich), SnI\(_2\) 99+% (Alfa Aesar), SnI\(_2\) 99% (Strem)), it is possible for a residual amount of magnetic impurities to be incorporated during growth from the source materials. Additionally, it has recently been demonstrated that vacancy-induced magnetism can arise in solution processed lead halide perovskites (Sun et al., 2020). Future studies could be conducted on other epitaxial halide perovskites, such as CsSnI\(_3\), to systematically investigate the role of magnetism on quantum transport devices based on these materials. Regardless of the underlying inelastic scattering mechanism, \( \ell_F \) increases with decreasing temperature and reaches \( \approx 5 \mu m \) below roughly 350 mK, below which it saturates, possibly due to thermal decoupling of the charge carriers at low temperatures. We note that recent work on three-dimensional quasi-epitaxial layers of CsPbBr\(_3\) have reported WL signatures in photoelectric transport with \( \ell_F \approx 50 \text{ nm at } 3.5 \text{ K} \) (Wang et al., 2020), likely limited by the quasi-epitaxial domain size.

**Figure 4. Phase coherence length**

Temperature dependence of the charge carrier phase coherence length \( \ell_F \) in the epitaxial CsSnI\(_3\) thin film. Each value of \( \ell_F \) corresponds to the average value obtained from multiple measurements at a given temperature and the error bars are the standard deviation. For comparison, we also show results from the literature for other electronic materials.
The value of the charge carrier phase coherence length we find in epitaxial CsSnI$_3$ can be compared to other high-quality low-dimensional electronic materials in the quantum diffusive regime where localization measurements are possible such as GaAs (Beenakker and van Houten, 1991; Mailly, 1987), oxide thin films (Yun et al., 2017), graphene (Baker et al., 2012; Elias and Henriksen, 2017; Ki et al., 2008; Tikhonenko et al., 2008), as well as silicon (Beenakker and van Houten, 1991). We note that longer phase coherence exists in extremely high-mobility GaAs 2DESs and graphene but are not reported due to the absence of localization effects in sufficiently disorder free samples. In fact, in graphene ballistic transport has been observed over longer than 15 microns (Wang et al., 2013).

In summary, our results on phase coherent transport in the presence of spin-orbit coupling show that micron-scale phase coherence lengths can be achieved in epitaxial CsSnI$_3$. These findings attest to the fact that epitaxial halide perovskites are emerging as a novel material for future quantum coherent devices with potential applications in spin-orbitronics where gate-control could enable manipulation of spin via the Rashba-effect (Kepenekian and Even, 2017; Manchon et al., 2015).

Limitations of the study
As described in the main text of the manuscript, transport measurements were performed using a two-terminal configuration. These measurements reveal, via the manifestation of WAL, spin-orbit coupling, and long phase coherent transport of charge carriers. Future work using a four-terminal van der Pauw geometry would allow for measurements of the zero-field mobility needed to characterize the elastic mean-free path of charge carriers in this material and four-terminal measurements of the Hall resistance as a function of magnetic field would allow characterization of the type and density of charge carriers. These measurements could be performed to extensively map out the dependence of the mobility and density on various epitaxial growth parameters, doping level, disorder, etc. Such future measurements would be ideal for optimizing sample growth with an eye toward quantum electronic device applications as they could be used to understand how to develop samples with decreasing levels of impurity scattering. Additionally, such measurements would allow for a quantitative measure of the strength of the spin-orbit coupling revealed by our two-terminal measurements.

STAR METHODS
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SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.isci.2021.102912.

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AUTHOR CONTRIBUTIONS
K.N. and L.Z. performed the low-temperature magneto-transport experiments and I.K. and P.C. grew the halide perovskite films and devices. I.K., P.C., L.W., and R.J.S. performed X-ray characterization of the samples. J.P. and R.R.L. conceived of the experiments and supervised the project. All authors contributed to data analysis and writing the manuscript.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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STAR METHODS

KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Software            | Mathematica [https://www.wolfram.com/mathematica/](https://www.wolfram.com/mathematica/) | version 12.3 |
|                     | Materials Studio [https://www.3ds.com/products-services/biovia/products/molecular-modeling-simulation/biovia-materials-studio/](https://www.3ds.com/products-services/biovia/products/molecular-modeling-simulation/biovia-materials-studio/) | version 7.0 |

RESOURCE AVAILABILITY

Lead contact
Further information and request for resources should be directed to the lead contact, Johannes Pollanen (pollanen@msu.edu).

Materials availability
This study did not generate new unique reagents.

Data and code availability
The data are available upon reasonable request by contacting the lead contact. No new code was generated during the course of this study.

METHOD DETAILS

CsSnI3 sample growth
The epitaxial CsSnI3 crystalline film samples were grown in a custom Angstrom Engineering thermal evaporator by co-evaporation from two tungsten boats containing precursor materials (CsI and SnI2), with each source having an independently calibrated quartz-crystal-microbalance rate monitor and source shutter. A 50 nm thick film was deposited stoichiometrically at a rate of 0.007 nm/s on a cleaved [100] surface of a potassium chloride (KCl) single crystal substrate. The growth was performed at pressures less than $3 \times 10^{-6}$ torr and a temperature of $22 \pm 4 \degree C$. In-situ crystal analysis was performed in real-time using reflective high-energy electron diffraction (RHEED) to determine structure and film quality. The RHEED scans were performed at 30.0 kV and an emission current of less than 50 nA to reduce damage and charging on the perovskite film during growth. With these measurement conditions no damage was observed over typical deposition times of up to 1-2 hours. Gold contacts (50 nm thick) were deposited on the perovskite layers using electron-beam evaporation at a rate of 0.02 nm/s through a shadow mask in the same deposition chamber (with the same temperature and pressure).

Characterization of CsSnI3 air sensitivity and device handling
To understand the effects of exposure to air, we have characterized the air sensitivity of additional epitaxial CsSnI3 devices using x-ray, optical and electrical transport, as shown in Figure S7. In Figure S7C we show how the resistance of a CsSnI3 epitaxial layer evolves as a function of time when the device was exposed in air at room temperature. For the first 15-20 minutes, the device resistance does not significantly change. With continued exposure we clearly observe an increase in the device resistance with increasing time, which continues until it is > 50 M$\Omega$. We attribute this behavior to the known transformation of CsSnI3 into Cs$_2$SnI$_6$ upon exposure to air (even for the epitaxial phase). Importantly for the results reported in this manuscript no such changes in device resistance were observed for any of the devices that were handled and sealed in a dry-nitrogen environment before being measured at low temperature. Moreover, repeated transport measurements performed at low temperature did not show changes in the overall resistance of the device nor changes in the observed magnetotransport phenomena.

Although the epitaxial films were structurally stable in air for hours, care was taken to mitigate this as a convoluting effect. Thus the growth, handling and characterization of the devices in this article (CsSnI3...
and CsSnBr3) were performed, from start to finish, in a dry, oxygen-free environment. After the fabrication, the devices were sealed in containers filled with dry nitrogen gas and transported from the growth laboratory to the low temperature quantum measurement laboratory. These devices were wired up in a nitrogen environment and loaded into a custom hermetically sealed sample cell containing an 18-pin chip carrier inside (see Figure S8). The cell was sealed with a conventional indium O-ring compatible with cryogenic measurements. Then the cell was transferred to dilution refrigerator for low temperature (~10 mK) and high magnetic field (~14 T) transport measurements. At no point during this entire process (from growth to low-temperature measurement) were the devices exposed to air.

**Low-temperature magneto-transport measurements**

After growth the devices were wired up using 0.02” diameter gold wire and InSn solder in a nitrogen environment and loaded into a custom hermetically sealed sample cell containing an 18-pin chip carrier inside. This experimental cell was sealed with a conventional indium O-ring compatible with cryogenic measurements. Then the cell was transferred to cryogen-free dilution refrigerator for low temperature (~10 mK) and high magnetic field (~14 T) transport measurements which were performed using low-frequency (10 Hz) lock-in techniques. At no point during this entire process (from growth to low-temperature measurement) were the devices exposed to ambient condition.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Each value of $L_F$ shown in Figure 4 of the main manuscript corresponds to the average value obtained from multiple weak-anti-localization measurements at a given temperature and the error bars are the standard deviation.