Intrinsic insulating ground state in transition metal dichalcogenide TiSe₂

Daniel J. Campbell,^{1,*} Chris Eckberg,¹ Peter Y. Zavalij,² Hsiang-Hsi Kung,^{3,4} Elia Razzoli,^{3,4} Matteo Michiardi,^{3,4,5}

Chris Jozwiak,⁶ Aaron Bostwick,⁶ Eli Rotenberg,⁶ Andrea Damascelli,^{3,4} and Johnpierre Paglione^{1,7,†}

¹Center for Nanophysics and Advanced Materials, Department of Physics, University of Maryland, College Park, Maryland 20742, USA

²Department of Chemistry, University of Maryland, College Park, Maryland 20742, USA

³Quantum Matter Institute, University of British Columbia, Vancouver, British Columbia V6T 1Z4, Canada

⁴Department of Physics & Astronomy, University of British Columbia, Vancouver, British Columbia V6T 1Z1, Canada

⁵Max Planck Institute for Chemical Physics of Solids, 01187 Dresden, Germany

⁶Advanced Light Source, Lawrence Berkeley National Laboratory, Berkeley, California 94720, USA

⁷Canadian Institute for Advanced Research, Toronto, Ontario M5G 1Z8, Canada

(Received 25 September 2018; published 14 May 2019)

The transition metal dichalcogenide $TiSe_2$ has received significant research attention over the past four decades. Different studies have presented ways to suppress the 200 K charge-density-wave transition, vary low-temperature resistivity by several orders of magnitude, and stabilize magnetism or superconductivity. Here we give the results of a synthesis technique whereby samples were grown in a high-pressure environment with up to 180 bar of argon gas. Above 100 K, properties are nearly unchanged from previous reports, but a distinct hysteretic resistance region begins around 80 K, accompanied by insulating low-temperature behavior. An accompanying decrease in carrier concentration is seen in Hall effect measurements, and photoemission data show a removal of an electron pocket from the Fermi surface in an insulating sample. We conclude that high inert gas pressure synthesis accesses an underlying nonmetallic ground state in a material long speculated to be an excitonic insulator.

DOI: 10.1103/PhysRevMaterials.3.053402

I. INTRODUCTION

Titanium diselenide is one of the most studied members of the transition metal dichalcogenide (TMD) family. As with other TMDs, weak van der Waals bonding along the caxis of its hexagonal structure means that relatively minor tweaks to unit cell size, stoichiometry, or interlayer dynamics can have dramatic effects on physical properties. In TiSe2, these changes are most evident in investigation of the chargedensity wave (CDW) that emerges at 200 K under normal circumstances [1]. The TiSe₂ Fermi surface is not susceptible to nesting, a typical driver of charge ordering in TMDs [2], so other explanations have been proposed: different variations of the Jahn-Teller effect [3–5] or an excitonic insulator state resulting from a small indirect band gap or overlap [6]. Recent experiments have given backing to the latter scenario [7,8]. Despite this, TiSe₂ single crystals show metallic low-temperature behavior, with an overall resistivity decrease from room temperature.

Both the application of high pressure and the intercalation or substitution of new atoms to $TiSe_2$ have been used to change the character of the CDW, generally suppressing it and in some cases leading to superconductivity or magnetic ordering [9–16]. In this paper, we present a way of stabilizing new properties in $TiSe_2$ at ambient pressure without the use of additional atoms. By applying up to 180 bar of pressure with argon gas during growth, we have synthesized both single and polycrystals that, below 100 K, exhibit a first-order transition together with a large increase in the resistance and magnitude of the Hall coefficient. We find pressure growth to be fundamentally distinct from substitution, as samples show very similar transport, magnetic, and structural properties to typical TiSe₂ at higher temperatures. Instead, the presence of high inert gas pressure reduces selenium vacancy formation, counteracting an extrinsic metallic component in the resistance and allowing for observation of a previously obscured insulating ground state, traces of which have been seen in other studies. Photoemission measurements give evidence for the disappearance of an electron pocket in an insulating sample that is present in semiconducting and metallic ones, in line with this assertion.

II. EXPERIMENTAL METHODS

Typically, TiSe₂ crystals have been grown by chemical vapor transport (CVT) with excess Se or, more popularly, I₂ as the transport agent [1,12,17–19]. In contrast, for this study, samples were grown at elevated pressure using argon gas in a Morris HPS-3210 furnace [Fig. 1(a)]. This furnace can reach pressures up to 200 bar at 1000 °C by introducing Ar into a stainless steel growth chamber. The pressure in the chamber varies in a consistent manner with temperature; the values reported here correspond to the maximum observed pressure for each growth, which (depending on maximum temperature) was about 60–70% greater than at room temperature.

^{*}Corresponding author: djcampbe@umd.edu

[†]Corresponding author: paglione@umd.edu



FIG. 1. (a) The Morris HPS-3210 furnace used for growth. A quartz ampule, open at one end, was inserted with its closed end on the right-hand side near the heating element. The entire chamber was then sealed and the furnace lid closed. Tilting the furnace is suspected to help nucleate crystals. (b) A typical ampule after growth. The dark area on the closed right end is polycrystalline $TiSe_2$, while the region in the middle is elemental Se. (c) A small, fragile, pressure-grown $TiSe_2$ crystal. (d) Larger pressure-grown $TiSe_2$ single crystals (on $1 \times 1 \text{ mm}^2$ scale paper).

Traditional CVT is not possible in the pressure furnace, both because of the large amount of Ar gas present and the fact that iodine vapor would damage the chamber, so the actual process was closer to a "Se flux" growth. Se shot (99.999+%, Alfa Aesar) and crushed Ti slugs (99.98%, Alfa Aesar) were mixed together at the bottom of a quartz ampule with a 40-cm length and 0.75-cm inner diameter. Growths were attempted with Ti:Se ratios ranging from 1:2 to 1:9, and several different temperature sequences. The most frequently used profile was a 1:9 ratio and heating the furnace at 48 °C/hour to 700 °C, where it stayed 24-48 h. The sample space was then slowly cooled at 4.8 °C/h to 400 °C, after which it was passively cooled to room temperature; only then was the chamber returned to ambient pressure. For comparison, we also grew single crystals via CVT with I₂ and a flux technique [20] using excess Se (in a 9:1 ratio with Ti) in alumina crucibles in about 1/3 atm of Ar gas inside a sealed quartz tube.

All pressure furnace growths produced a large number of polycrystalline chunks of TiSe₂, but only about half also resulted in single crystals large enough for transport measurements. There was no identifiable correspondence between growth pressure, temperature profile, or Ti:Se ratio and the successful production of large crystals. Empirically, it seemed that propping up the end of the furnace opposite from where the reactants were located at an angle helped to form large single crystals. Doing this may concentrate Se at the end of the ampule where the Ti slugs are located, since the excess Se often condensed further up the length of the tube [Fig. 1(b)]. It could also help amplify any natural temperature gradient in the long, narrow furnace and approximate the conditions for vapor transport.

Resultant single crystals varied in appearance between growths, as seen in Figs. 1(c) and 1(d). Some were small, whispy, and flexible, less than half a millimeter in length and 15–60 μ m in thickness. Others were larger and sturdier, over 1 mm wide and 200 μ m thick. In either case, single crystal x-ray diffraction (XRD) confirmed that the platelike crystals always grew with the *c* axis out of plane, as would be expected for hexagonal, layered TiSe₂. Throughout this paper, single crystals are labeled by the maximum pressure reached during growth, with those from the same batch distinguished by lettering.

Synchrotron powder XRD data were obtained through the 11-BM beamline rapid access mail-in program at the Advanced Photon Source at Argonne National Laboratory, and refinements were made with the GSAS-II software package [21]. Single-crystal XRD measurements were made on a Bruker APEX2 Diffractometer with Mo K_{α} radiation. The integral intensities were corrected for absorption with the SAD-ABS software [22], using the integration method. The structure was solved with the SHELXS-2015 program and refined with the SHELXL-2015 program and least-square minimization using the SHELX software package [23]. Angle-resolved photoemission spectroscopy (ARPES) measurements were performed at the MAESTRO beamline 7.0.2.1 of the Advanced Light Source in the Lawrence Berkeley National Laboratory, where samples were cleaved and measured at 20 K with a base pressure of 2×10^{-11} Torr. Photoelectrons were detected by a Scienta R4000 analyzer equipped with a deflector, where the energy and angular resolution were better than 20 meV and 0.2° , respectively. Electrical transport measurements were carried out in 9 T and 14 T Quantum Design Physical Properties Measurement Systems, and a 14 T Quantum Design DynaCool. The 14 T PPMS and DynaCool were also used for heat capacity measurements. Magnetization was measured using the DynaCool vibrating sample magnetometer as well as two versions of the 7 T Quantum Design Magnetic Properties Measurement System, the MPMS XL and MPMS3.

III. PHYSICAL PROPERTIES

A. Structural characterization

Synchrotron diffraction patterns were taken at 295 K of ground polycrystals from 13 growths: three separate CVT growths with temperatures of either 550 or 575 °C at the hot end of the ampule (thought to be optimal for CVT) [1], a Se flux growth, and nine pressure growths with maximum pressures in the range 56–173 bar. Figure 2 shows (a) the lattice parameters, (b) volumes, and (c) Se site occupancies for all data sets, where vapor transport is denoted as "0 bar" of growth pressure, Se flux as "1 bar," and growth methods are further distinguished by symbols. Values for all samples



FIG. 2. TiSe₂ structural data obtained from refinement of powder synchrotron data of CVT (squares, "0 bar"), Se flux (triangles, "1 bar"), and pressure (circles) growths. (a) shows the *a* (left, blue) and *c* (right, red) axis lengths and (b) the volumes, with error bars magnified ten times from those given by GSAS-II. Note that data points for two of the CVT growths nearly completely overlap. (c) is the refined Se occupancy of the same data, with error bars, where a CVT and Se flux growth overlap at 0.98.

are very close to those reported previously [11,19]. Elemental Se was present in some cases, unsurprising given the use of excess in the growth. There is a very small but nevertheless consistent difference in unit cell size and composition between the synthesis techniques. Both Se flux and pressure growths have smaller lattice parameters, perhaps attributable to the absence of larger iodine atoms that can substitute on the Se site. This shift is much smaller than that brought about by small amounts of chemical dopants or intercalants [10,11,16]. Among pressure-grown samples, there is again no trend in lattice size with the amount of Ar pressure applied. This

TABLE I. Parameters obtained from single crystal XRD at 150 K. Here and in subsequent figures, single crystals from the same batch (and therefore with the same P_{Growth}) are distinguished by lettering.

P _{Growth} (bar)	<i>a</i> (Å)	<i>c</i> (Å)	wR_2
101A	3.5415(11)	6.0198(19)	0.0494
101B	3.5355(9)	6.0112(16)	0.0556
114A	3.5332(15)	6.007(3)	0.0553
138	3.5432(10)	6.0195(17)	0.0615
140	3.5275(11)	5.9969(19)	0.0509

is not surprising; via the Clausius-Clapeyron equation, the boiling point of a material varies logarithmically with external pressure. A factor of about 4 in the high pressure growths will have much less of an impact of Se volatility than the differences of several orders of magnitude between pressure growth, flux, and CVT.

There is also a clear difference in Se occupancy between pressure growth and the other two methods. The presence of Se vacancies is known to be highly sensitive to growth method and temperature, and can impact transport behavior [1,24]. Vapor transport and Se flux result in vacancies of 2-4%, while those for pressure growth are all within 1% of full occupation. An explanation for this is that the presence of a significant amount of inert gas suppresses the high vapor pressure of Se and, as a result, reduces vacancy formation. In contrast, vapor transport relies on reaction in the gas phase with a third element, iodine, making it more susceptible to reduced Se content or site substitution. Se flux avoids this issue, but with a low argon pressure that does not sufficiently combat the volatility of Se. Comparable temperatures and amounts of excess Se were used in flux and pressure growths, thus it is evident that higher pressure is the key factor to reduced vacancies. Additionally, vacancy formation is not directly dependent on lattice volume, since the values for Se flux and high pressure growth are very similar. Significant differences in the properties of pressure-grown samples compared to Se flux or I₂ CVT that we will show further on are then attributable to Se occupancy, rather than a slightly smaller lattice.

Five single crystal samples grown in the range 101-140 bar, including two from the same batch, were also selected for XRD at 150 K [Table I]. These data are not directly comparable to room temperature values as they are determined by a combination of thermal contraction and CDW-related lattice distortion. Nevertheless, they demonstrate that T_{CDW} is above 150 K for these samples, in spite of different growth conditions and low-temperature resistivities 2-5 times higher than the 300 K value. Using the reported TiSe₂ lattice parameters [19] and thermal expansion coefficients [25] we can estimate 150 K lattice parameter values, before accounting for the effect of the CDW on the lattice, to be a = 3.530 Å and c = 5.993 Å for typical TiSe₂. With one exception, we see 150 K lattice parameters that are similar to or larger than room-temperature values. The reason for this is charge ordering, which by 150 K can result in a distortion of more than 10^{-2} Å in



FIG. 3. Resistance (scaled to 300 K value) as a function of temperature for pressure-grown TiSe₂ single crystals listed in Table I, with maximum growth pressures noted. Note that even the two samples from the same batch exhibit very different behavior. Hysteresis in the 30–80 K region is marked by arrows for the black curve to show the difference in warming and cooling, which is the same for unmarked samples. In some cases, there is also hysteresis around the CDW-related upturn from 150–200 K.

typical TiSe₂ [1,26]. While applied pressure is known to suppress the CDW [12], pressure growth evidently does not, since at 150 K samples have experienced a lattice expansion.

B. Electrical transport

The temperature-dependent resistance of the single crystals from Table I is shown in Fig. 3, with resistances scaled to 300 K values. The behavior shown is representative of what is seen in a larger number of samples. Above 100 K, the behavior does not differ from previous reports on TiSe₂. Following convention, we used the kink in the derivative (a peak in the second derivative) to identify the onset of charge ordering [1]. The values we see for all samples are just above 200 K, the same as standard TiSe₂, and concordant with the conclusion from 150 K single crystal XRD. All samples have a rise upon cooling at 200 K regardless of behavior at lower temperatures. On the other hand, the peak in resistance, normally centered around 165 K, comes at a lower temperature in most of our samples. A similar effect has been reported in samples grown without iodine [27], or at higher temperature [1,24]. The ratio R_{peak}/R (300 K) for our samples can be as high as six, larger than has been achieved with CVT crystal growth [24,27]. An increased peak height (relative to 300 K resistivity) has previously been interpreted as signifying fewer Se vacancies and correspondingly higher crystal quality [1,17,28]. Another recent paper argued that Se deficiency is actually beneficial to charge ordering, a conclusion based purely on the size of the resistance increase in the 150–200 K region [24]. In comparing multiple growth methods, pressure-grown crystals show a taller peak, but at a lower temperature, and without Se deficiency. From this, we believe that there is no correspondence between peak height, the temperature at which it occurs, and crystal quality. The broadness of the peak, and the fact



FIG. 4. $\Delta R \equiv R_{\text{warming}} - R_{\text{cooling}}$, scaled to room temperature resistance, as a function of temperature for seven crystals from the same 101 bar growth. Hysteresis is most evident around 80 K but also manifests at the higher temperature CDW resistance peak. $\Delta R/R$ reaches a maximum of 5.0, 6.7, and 6.9 for the red, purple, and blue curves, respectively. Inset: $\rho(T)$ of each sample, with matching colors. Stars indicate samples for which data are also presented in other figures: 101B (Table I and Fig. 3, here green), 101C (Fig. 5, pink), and 101D (Fig. 7, purple).

that it comes 30-50 K lower than T_{CDW} , means that it likely represents simply a change in dominant scattering mechanism within the charge ordered phase, and that its specifics are not as significant. This is supported by the fact that our own samples with similar peak heights show differing behavior upon further cooling.

The more significant departure from previous observations comes at lower temperatures, where many pressure-grown samples show a large increase in resistivity. Additionally, temperature hysteresis often opens around 80 K in a similar range to where insulating character emerges, before closing near 30 K. Visible in Fig. 3, this is emphasized in Fig. 4, which shows ΔR , defined as the difference in resistance value between warming and cooling, scaled to its room temperature value, for five samples from a single 101 bar growth. Except for a single nonhysteretic sample (the green curve), the greatest difference occurs between 30-100 K, with samples also showing appreciable ΔR near and above the higher temperature resistance peak. This effect is still present when temperature is swept slowly or stabilized at each data point, and so is not a result of temperature lag or sample heating. In some cases, there is a noticeable kink at the hysteresis opening, further confirmation that the transition is more than just measurement error. Application of fields up to 140 kOe does not affect overall behavior. Polycrystalline samples were always insulating and hysteretic, often to a far greater degree, but we present only single-crystal resistance data throughout this paper to demonstrate that the effect is inherent to pressure-grown TiSe₂ and not a result of insulating impurities, grain boundaries, or other effects that make polycrystalline transport measurements less reliable.

Not all single crystals show the 80 K transition; some behave like typical, metallic, CVT-grown TiSe₂, and behavior below 100 K varied even for samples from the same growth



FIG. 5. (a) Temperature-dependent resistivity for $TiSe_2$ single crystals grown by I_2 chemical vapor transport, excess Se flux, and 101 bar of Ar gas pressure. At high temperatures, the behavior of all three samples is similar, but differences emerge below 150 K. No hysteresis is observed in the CVT or flux crystals. (b) The derivative of the cooling data from (a) in a narrower temperature region. The inset is the second derivative of the same data, where the peak is identified as the onset of the CDW.

[Fig. 4, inset; note the logarithmic y axis]. The height of the CDW peak is also inconsistent and its temperature can vary by as much as 10 K. The lowest growth pressure to produce single crystals was 56 bar, and the highest was 140 bar, but polycrystals were grown at maximum pressures of 10–180 bar and were more universally insulating. As with lattice parameters, there is no clear link between a specific growth pressure and resistivity behavior (or any other measurable quantity). Instead, it seems that pressure synthesis can stimulate this behavior, but does not guarantee it. The difference between 50 and 200 bar of pressure is likely insignificant, since in any case the pressure is much higher than during CVT or flux growth.

A comparison of $\rho(T)$ for crystals grown by I₂ CVT, Se flux, and Ar pressure [Fig. 5(a)] makes clearer the differences in resultant single crystals. All show the CDW-associated rise in ρ at the same temperature. The CVT crystal has a peak at 165 K and $\rho(1.8 \text{ K}) < \rho(300 \text{ K})$. The Se flux sample has a larger peak, suppressed in temperature to 150 K, and slightly higher resistivity at 1.8 K than room temperature. A pressuregrown crystal has a local maximum at even lower temperature (140 K) with a comparable size [relative to $\rho(300 \text{ K})$] and a hysteretic, insulating transition. T_{CDW} has typically been identified as the beginning of the flat minimum region in the first derivative [Fig. 5(b)], which corresponds to the onset of charge ordering in neutron measurements [1]. However, the minima for Se flux and pressure-grown samples are influenced by the suppression of T_{peak} . We therefore used the peak in the second derivative, equivalent to the kink in the first derivative, to define T_{CDW} . In the inset to Fig. 5(b), it is clear that this occurs at the same temperature for all three samples, and in fact the CVT and pressure-grown samples have nearly identical first and second derivatives above 200 K. While both Se flux and pressure-grown crystals have an elevated low-temperature resistance compared to CVT samples, pressure growth is further distinguished by the more significant insulating behavior and temperature hysteresis.

As with longitudinal resistivity, the Hall resistance [Fig. 6(a)] in pressure-grown crystals above 200 K is generally similar to typical TiSe₂ [1]. The Hall coefficient R_H is initially positive and crosses zero at 150–170 K [Fig. 6(a), inset], which, like the $\rho(T)$ peak, is slightly lower than our own CVT crystals and previous reports [1,27]. Elemental substitution leads to a more substantial temperature suppression [29,30]. Some pressure-grown crystals showed a slight increase in R_H just before the zero crossing, not seen with vapor transport. At low temperatures, R_H can reach large negative values up to two orders of magnitude larger than those of typical $TiSe_2$ [1] and an order of magnitude above those measured even for insulating $Ti_{1-x}M_xSe_2$ (M = As, Sc, Nb, Ni, Re, or Y) [30], indicating that a reduced carrier concentration, rather than impurity scattering, is the reason for increased resistance. Generally, samples with more insulating low-temperature behavior had a larger $|R_{\rm H}|$. The Hall signal becomes more linear with decreasing temperature below the R_H sign change, indicating transport dominated by a single electron band despite the presence of multiple carrier types at higher temperatures. Overall, the Hall data support the idea of a change in band structure at low temperatures between pressure and CVTgrown TiSe₂. Another thing we noticed in measurements was an 80 K maximum in $|R_H|$ for CVT crystals. Extrema in R_H at a similar temperature have been seen in other intrinsic [1] and metal-doped [30] samples. Given that this is the same temperature as the beginning of hysteresis, it suggests that pressure growth emphasizes or strengthens some phenomenon already present in other forms of TiSe₂.

C. Heat capacity

Heat capacity measurements were taken from 300 K to 2 K on a polycrystalline chunk of TiSe₂ grown at 160 bar [Fig. 6(b)], which had shown an insulating transition in transport measurements. We observe no features in the corresponding temperature range, and the shape and values of the data are very similar to what has been measured before [28]. The lack of a feature in the hysteretic region is not wholly surprising, as even that corresponding to the higher temperature CDW is subtle. Low-temperature measurements on the same polycrystal and two others grown at different pressures are shown in the inset of Fig. 6(b). All data fit well to the standard specific heat equation $C/T = \gamma + \beta T^2$. In this equation, γ is the Sommerfeld coefficient and β can be used to calculate



FIG. 6. (a) The temperature dependence of the Hall coefficient of $TiSe_2$ single crystals. Note that the CVT sample's temperature dependence, including a zero-crossing, is too small to see on this scale. The inset shows the zero-crossing region for selected samples from the main plot. (b) Specific heat of a polycrystalline $TiSe_2$ sample with a maximum growth pressure of 160 bar as a function of temperature. Inset: Low temperature data for the same sample and two other polycrystals, with maximum growth pressures noted. Lines are fits to the Debye low-temperature specific heat model. (c) Paramagnetic susceptibility of polycrystalline $TiSe_2$ chunks taken in constant applied fields of 20 (116 bar), 50 (80, 127, and 156 bar), or 70 kOe (CVT and 114 bar) due to the small intrinsic moment. (d) The hysteretic area (see Discussion for details) versus resistance increase for pressure-grown single crystals, on a log-log scale. Inset: The base temperature Hall coefficient as a function of the same quantity. More insulating samples generally showed a larger $|R_H|$. Error bars come from uncertainty in the measurement of sample thickness and in some cases are smaller than the symbol size.

the Debye temperature $\theta_D = (\frac{12\pi^4 N_A k_B n}{5\beta})^{\frac{1}{3}}$, where N_A is the Avogadro number, k_B Boltzmann's constant, and n = 3 the number of atoms per formula unit. Results were similar for all three samples. The computed γ values are small: 0.14, 0.16, and 0.19 $\frac{\text{mJ}}{\text{mol K}^2}$, respectively, for the 98, 130, and 160 bar samples, reflecting the small low-temperature density of states at the Fermi energy. θ_D is 220, 244, and 209 K for the same data. The reference values for TiSe₂ powder [28] are $\gamma = 0.19$ $\frac{\text{mJ}}{\text{mol K}^2}$ and $\theta_D = 251$ K, both similar to but slightly higher than those derived from pressure-grown samples.

D. Magnetic susceptibility

Like heat capacity, magnetization measurements for multiple growths differed little from vapor transport-grown TiSe₂. The total magnetic susceptibility is small, on the order of 10^{-6} emu (Oe mol Ti)⁻¹, since the paramagnetic and diamagnetic components are comparable in magnitude [1,11,14]. Figure 6(c) shows the paramagnetic susceptibility χ_P after subtraction of the core diamagnetic contribution [31]. The shape matches results from our own CVT-grown samples

and what has been presented in the past [1], and in fact has a similar shape to $d\rho/dT$. We attribute the rise at lowest temperatures in some curves to paramagnetic impurities. Data were taken under large fields (≥ 20 kOe) to enhance the weak signal, but curves had the same appearance over the range 0–140 kOe.

E. ARPES

To understand the origin of the insulating ground state in some crystals, we performed comparative ARPES measurements on two crystals, one semimetallic [114B in Fig. 7(a)] and another more insulating (101D). Measurements were made at 20 K, below the onset of hysteretic resistance. Despite discussions of the semimetallic versus semiconducting nature of TiSe₂ above the CDW transition [32], the previous consensus was that an electron pocket crosses the Fermi level at the *L* point (Se-4*p*) in the CDW phase [33,34]. However, we find that the more insulating sample presents neither an electron nor hole pocket anywhere in *k*-space at the Fermi energy. Figure 7(b) compares the constant binding energy contours measured in the *A-L-H* plane ($k_z = \pi/c$), where



FIG. 7. (a) Resistivity as a function of temperature for the two pressure-grown ARPES samples, with a schematic drawing of the Brillouin zone. 114B shows a small amount of hysteresis that is less obvious with the logarithmic scale. (b) Comparison of the constant energy contour measured at $k_z = \pi/c$ (the *A*-*L*-*H* plane) at E_F for 114B (left) and the *L* valence band maximum (60 meV below E_F) for 101D (right) at T = 20 K, plotted in the $k_x < 0$ and $k_x > 0$ half-plane, respectively. (c) and (d) ARPES map measured along the *A*-*L* cut [the dashed red line in (a)] for 114B and 101D, respectively, with the same photon energy of 122 eV for both. To the right of each cut are the EDCs at the *L* point (vertical red solid line), which were fit by Voigt functions multiplied by the Fermi-Dirac distribution (blue curves).

data for 114B (at E_F) and 101D (60 meV below E_F), at the maximum of the L-point valence bands) are shown in the left and right half-planes, respectively. The Fermi surface of 114B is composed of electron pockets at the L point, whereas in the more insulating 101D the states closest to the Fermi level are the hole bands at the Γ and L points at a larger binding energy. This is most evident from the measured band dispersion along the A-L direction (the red dotted line in the Fig. 7(a) inset). For 114B [Fig. 7(c)], a small electron pocket crosses the Fermi level at $k_y \approx 1.0 \text{ Å}^{-1}$. The energy distribution curve (EDC) at this momentum (marked by solid red line) is shown in the right panel of Fig. 7(c), where it is fit with two bands, consistent with previous reports [34]. This is distinct from 101D, shown in Fig. 7(d), where no band crosses the Fermi level and the EDC can be fit with just a single Voigt distribution multiplied by the Fermi-Dirac function. The lack of any quasiparticle band crossing the Fermi energy would naturally explain the insulating behavior of 101D and should also result in a reduced carrier concentration. Scans throughout the entire k_z dispersion confirmed the absence of any intensity at E_F at the Brillouin zone edge.

IV. DISCUSSION

Above 100 K, there is little to distinguish TiSe₂ grown with iodine vapor transport or at high argon pressure. The difference between the two of the sudden decrease in $\chi(T)$ and the maximum in $d^2\rho/dT^2$ (the $d\rho/dT$ inflection point), both associated with the onset of charge ordering [1,29], is 5 K or less. XRD data show that the lattice of pressuregrown samples is about 0.1% smaller than for CVT. However, pressure growth results in no appreciable selenium vacancies, compared to about 2-4% reduced stoichiometry in CVT or flux samples. While small, these changes are very consistent. The most noteworthy aspect of pressure growth is that it can lead to insulating and hysteretic low temperature behavior. And although TiSe₂ becomes more insulating with $\leq 5\%$ V doping [9,29] or the intercalation of Cr, Fe, and Co [10], those samples do not display temperature hysteresis. Furthermore, they show CDW temperature suppression [13], antiferromagnetic or Curie-Weiss behavior [10], and a more significant change in room-temperature lattice parameters [10,16], distinctly different effects than pressure growth.

The introduction of Pd and Pt, in contrast, do have similarities to our findings. In $Pd_x TiSe_2$ with $x \leq 0.03$, a second inflection point in $d\rho/dT$ occurs near 80 K and transmission electron microscopy shows a strengthened CDW at lower temperature [14]. However, the *a* and *c* axes are both larger than in unintercalated samples, and further Pd intercalation leads to metallic behavior and superconductivity. Up to x = 0.13, resistivity increases by eight orders of magnitude at low temperatures in $Ti_{1-x}Pt_xSe_2$, while 300 K lattice constants and T_{CDW} hardly change [15]. Transport measurements and density-functional theory calculations on Pt-doped samples attribute insulating behavior to an increased energy gap. Similarly, our ARPES results demonstrate that pressure growth can lead to the disappearance of an electron pocket. However, hysteresis is not noted with either Pd or Pt.

 $TiSe_2$ has shown inconsistent transport behavior, with I_2 CVT-grown single crystals having an overall decrease in resistivity from room temperature[1,24] and polycrystals being more semimetallic [11,14,15]. Our own CVT or Se flux crystals can similarly differ in low-temperature properties [Fig. 5(a)]. Pressure-grown samples have a decrease in lowtemperature carrier concentration that is more dramatic in more insulating samples [Fig. 6(d), inset]. This also seems to be connected to the hysteretic region. In Fig. 6(d), we plot the "hysteretic area" of single-crystal samples against their scaled resistance increase, R(1.8 K)/R(300 K). This quantity is defined as the area under the $\Delta R/R(300 \text{ K})$ versus T curve (like those shown in Fig. 4) between 30 and 80 K. The correlation spans several orders of magnitude between more insulating behavior and more pronounced hysteresis, even after scaling the raw resistance.

ARPES results give more insight, showing that a very insulating crystal does not have the electron pocket at the Fermi surface seen in a semiconducting one and previous reports on CVT samples. As ARPES is unable to probe above E_F , the exact change to the conduction band cannot be determined: there may be a downward shift of the chemical potential, an increase in the conduction-valence band gap at L, orbitaldependent distortion, or a combination of all three. However, we note that Se vacancies would contribute electron carriers to the system. Reducing those vacancies via pressure growth would be equivalent to hole doping. We speculate that reduced vacancy formation lowers the chemical potential, as has also been shown to occur with growth of Bi₂Se₃ in the same furnace [35]. It may also change the size of the gap, similar to the suspected effect of Pt doping. This naturally results in a decreased carrier concentration and increased resistivity at low temperatures.

The fragility of TiSe₂'s structure in both real and momentum space means that subtle changes, such as a slight change in E_F or the gap size, will be magnified when transposed onto other properties. The small gap and the variability of a Fermi energy change would also explain the spread of semiconducting-insulating behavior in different samples, emphasized in Fig. 6(d), where even samples from the same batch show different transport behavior. The two ARPES samples have different Fermi surfaces, after all, but were grown under similar pressures. Variation could come from temperature gradients in the furnace, which could approximate the conditions for vapor transport, or other factors more difficult to observe and control during the growth process. Given the consistency of diffraction and transport measurements on polycrystals, it seems likely that the majority of pressure-grown material has reduced vacancies, with fluctuations among individual crystals. Pressure growth of Bi_2Se_3 had similar variability in resistive behavior and carrier concentration [35].

Anomalies at 80 K have appeared in previous studies of TiSe₂. The 200 K commensurate CDW (CCDW) can be suppressed with Cu intercalation or applied pressure, the CDWs in these samples have been shown by x-ray scattering and scanning tunneling microscopy to be incommensurate (ICDWs) in some regions of the phase diagram above the induced superconducting transition [36-38]. The change in ordering vector is first seen in the 65-80 K range, very close to where hysteresis first emerges in pressure-grown samples, and under pressure the ICDW-CCDW transition is weakly first order. The usual transition of TiSe₂ directly to a CCDW is actually atypical for a TMD. TaS₂, for example, has three progressively more insulating CDW transitions, with the two at lower temperature being hysteretic [39,40]. They correspond to the onset of (with decreasing temperature) incommensurate, nearly commensurate, and fully commensurate charge order. The consistent onset temperature of the ICDW in TiSe₂ in comparison to the continuous suppression of the CCDW with pressure or Cu intercalation has led to speculation that there is an inherent mechanism for lower temperature charge ordering that is "boosted" to 200 K by excitonic interactions [37]. A transition in a similar temperature region was suggested for Pd_xTiSe_2 with $x \leq 0.03$, where samples are insulating and T_{CDW} unchanged [14]. Even the |R_H| maximum at 80 K for CVT samples supports the notion of an underlying feature at that temperature. The two transitions we see in pressure-grown TiSe₂ crystals may signify that pressure growth allows for the observation of both the "natural" and "boosted" CDWs, perhaps with differing wave vectors, where the former is otherwise obscured by the effects of nonstoichiometry.

V. CONCLUSION

We have shown that TiSe₂ crystals grown under argon gas pressures of 10-180 bar can have a much larger resistance and reduced carrier concentration at low temperatures. Synchrotron data show a smaller lattice and reduced Se vacancy formation compared to vapor transport growth, and there is evidence for the elimination of an electron pocket at the Fermi level from photoemission. Prior examples in which the introduction of new atoms caused a low temperature resistance increase lack the hysteretic behavior that we have observed starting around 80 K. We suspect this new behavior stems from an enhancement of charge ordering and suppression of the metallic behavior, attributable to selenium vacancies, that dominates transport behavior in crystals grown by vapor transport. This first-order transition may be a signature of a true charge-ordered, excitonic insulator ground state in TiSe₂, that has been hinted at in work with applied pressure [38], Cu intercalation [36,37], or Pd doping [14].

The association between changes to charge ordering in $TiSe_2$ and superconductivity, demonstrated by the observation of CDW incommensuration near a quantum critical point in pressurized or Cu-intercalated samples, is a reason to further

explore the possibilities of high-pressure crystal growth. Due to their weak interlayer bonding, applied pressure or chemical substitution can significantly impact the behavior of TMDs. Pressure growth also presents a method of manipulating band gaps in bulk materials without the introduction of extrinsic atoms. This and other aspects of high-pressure synthesis can alter observed properties and lead to further discoveries related to TMDs or the many other materials with unstable lattice configurations.

ACKNOWLEDGMENTS

This work was supported by Air Force Office of Scientific Research Award No. FA9550-14-1-0332, National Science Foundation Division of Materials Research Award No. DMR-1610349, and the Gordon and Betty Moore Foundation's EPiQS Initiative through Grant No. GBMF4419. D.J.C. was funded in part by the U.S. Department of Energy (DOE), Office of Science, Office of Workforce Development for

- F. J. Di Salvo, D. Moncton, and J. Waszczak, Phys. Rev. B 14, 4321 (1976).
- [2] K. Rossnagel, J. Phys.: Condens. Matter 23, 213001 (2011).
- [3] H. Hughes, J. Phys. C 10, L319 (1977).
- [4] M. H. Whangbo and E. Canadell, J. Am. Chem. Soc. 114, 9587 (1992).
- [5] J. van Wezel, P. Nahai-Williamson, and S. S. Saxena, Phys. Rev. B 81, 165109 (2010).
- [6] D. Jérome, T. M. Rice, and W. Kohn, Phys. Rev. 158, 462 (1967).
- [7] H. Cercellier, C. Monney, F. Clerc, C. Battaglia, L. Despont, M. G. Garnier, H. Beck, P. Aebi, L. Patthey, H. Berger, and L. Forró, Phys. Rev. Lett. 99, 146403 (2007).
- [8] A. Kogar, M. S. Rak, S. Vig, A. A. Husain, F. Flicker, Y. I. Joe, L. Venema, G. J. MacDougall, T. C. Chiang, E. Fradkin, J. van Wezel, and P. Abbamonte, Science 358, 1314 (2017).
- [9] F. J. Di Salvo and J. V. Waszczak, Phys. Rev. B 17, 3801 (1978).
- [10] A. Kuranov, V. Pleshchev, A. Titov, N. Baranov, and L. Krasavin, Phys. Solid State 42, 2089 (2000).
- [11] E. Morosan, H. Zandbergen, B. Dennis, J. Bos, Y. Onose, T. Klimczuk, A. Ramirez, N. Ong, and R. Cava, Nat. Phys. 2, 544 (2006).
- [12] A. F. Kusmartseva, B. Sipos, H. Berger, L. Forro, and E. Tutiš, Phys. Rev. Lett. **103**, 236401 (2009).
- [13] M. Sasaki, A. Ohnishi, T. Kikuchi, M. Kitaura, K. Shimada, and H.-J. Kim, J. Low Temp. Phys. 161, 375 (2010).
- [14] E. Morosan, K. E. Wagner, L. L. Zhao, Y. Hor, A. J. Williams, J. Tao, Y. Zhu, and R. J. Cava, Phys. Rev. B 81, 094524 (2010).
- [15] J. S. Chen, J. K. Wang, S. V. Carr, S. C. Vogel, O. Gourdon, P. Dai, and E. Morosan, Phys. Rev. B 91, 045125 (2015).
- [16] H. Luo, J. W. Krizan, E. M. Seibel, W. Xie, G. S. Sahasrabudhe, S. L. Bergman, B. F. Phelan, J. Tao, Z. Wang, J. Zhang, and R. J. Cava, Chem. Mater. 27, 6810 (2015).
- [17] B. Hildebrand, C. Didiot, A. M. Novello, G. Monney, A. Scarfato, A. Ubaldini, H. Berger, D. R. Bowler, C. Renner, and P. Aebi, Phys. Rev. Lett. **112**, 197001 (2014).

Teachers and Scientists, Office of Science Graduate Student Research program, administered by the Oak Ridge Institute for Science and Education for the DOE under Contract No. DESC0014664. The work at the University of British Columbia was undertaken thanks in part to funding from the Max Planck-UBC-UTokyo Centre for Quantum Materials and the Canada First Research Excellence Fund, Quantum Materials and Future Technologies Program. It was also supported by the Killam, Alfred P. Sloan, and Natural Sciences and Engineering Research Council of Canada's (NSERC's) Steacie Memorial Fellowships (A.D.), the Alexander von Humboldt Fellowship (A.D.), the Canada Research Chairs Program (A.D.), NSERC, Canada Foundation for Innovation (CFI), and the CIFAR Quantum Materials Program. Use of the Advanced Photon Source at Argonne National Laboratory was supported by the DOE, Office of Science, Office of Basic Energy Sciences, under Contract No. DE-AC02-06CH11357. This research also used resources of the Advanced Light Source, which is a DOE Office of Science User Facility under Contract No. DE-AC02-05CH11231.

- [18] D. B. Lioi, D. J. Gosztola, G. P. Wiederrecht, and G. Karapetrov, Appl. Phys. Lett. 110, 081901 (2017).
- [19] C. Riekel, J. Solid State Chem. 17, 389 (1976).
- [20] P. C. Canfield and Z. Fisk, Philos. Mag. B 65, 1117 (1992).
- [21] B. H. Toby and R. B. Von Dreele, J. Appl. Crystallogr. 46, 544 (2013).
- [22] L. Krause, R. Herbst-Irmer, G. M. Sheldrick, and D. Stalke, J. Appl. Crystallogr. 48, 3 (2015).
- [23] G. M. Sheldrick, Acta Crystallogr. C 71, 3 (2015).
- [24] S. H. Huang, G. J. Shu, W. W. Pai, H. L. Liu, and F. C. Chou, Phys. Rev. B 95, 045310 (2017).
- [25] G. Wiegers, Physica B + C 99, 151 (1980).
- [26] C. Monney, C. Battaglia, H. Cercellier, P. Aebi, and H. Beck, Phys. Rev. Lett. 106, 106404 (2011).
- [27] I. Taguchi, M. Asai, Y. Watanabe, and M. Oka, Physica B + C 105, 146 (1981).
- [28] R. Craven, F. Di Salvo, and F. Hsu, Solid State Commun. 25, 39 (1978).
- [29] F. Levy, J. Phys. C 12, 3725 (1979).
- [30] F. Levy, J. Phys. C 13, 2901 (1980).
- [31] G. A. Bain and J. F. Berry, J. Chem. Educ. 85, 532 (2008).
- [32] J. C. E. Rasch, T. Stemmler, B. Müller, L. Dudy, and R. Manzke, Phys. Rev. Lett. 101, 237602 (2008).
- [33] Z. Vydrova, E. F. Schwier, G. Monney, T. Jaouen, E. Razzoli, C. Monney, B. Hildebrand, C. Didiot, H. Berger, T. Schmitt, V. N. Strocov, F. Vanini, and P. Aebi, Phys. Rev. B 91, 235129 (2015).
- [34] P. Chen, Y.-H. Chan, X.-Y. Fang, Y. Zhang, M.-Y. Chou, S.-K. Mo, Z. Hussain, A.-V. Fedorov, and T.-C. Chiang, Nat. Commun. 6, 8943 (2015).
- [35] P. Syers and J. Paglione, Phys. Rev. B **95**, 045123 (2017).
- [36] S. Yan, D. Iaia, E. Morosan, E. Fradkin, P. Abbamonte, and V. Madhavan, Phys. Rev. Lett. 118, 106405 (2017).

- [37] A. Kogar, G. A. de la Pena, S. Lee, Y. Fang, S. X.-L. Sun, D. B. Lioi, G. Karapetrov, K. D. Finkelstein, J. P. C. Ruff, P. Abbamonte, and S. Rosenkranz, Phys. Rev. Lett. 118, 027002 (2017).
- [38] Y. I. Joe, X. Chen, P. Ghaemi, K. Finkelstein, G. de La Peña, Y. Gan, J. Lee, S. Yuan, J. Geck, G. MacDougall, T. Chiang,

S. Cooper, E. Fradkin, and P. Abbamonte, Nat. Phys. 10, 421 (2014).

- [39] A. Thompson, R. Gamble, and J. Revelli, Solid State Commun.9, 981 (1971).
- [40] B. Sipos, A. F. Kusmartseva, A. Akrap, H. Berger, L. Forró, and E. Tutiš, Nat. Mater. 7, 960 (2008).