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Coexisting Charge-Ordered States with Distinct Driving Mechanisms in Monolayer VSe₂

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ABSTRACT: Thinning crystalline materials to two dimensions (2D) creates a rich playground for electronic phases, including charge, spin, superconducting, and topological order. Bulk materials hosting charge density waves (CDWs), when reduced to ultrathin films, have shown CDW enhancement and tunability. However, charge order confined to only 2D remains elusive. Here we report a distinct charge ordered state emerging in the monolayer limit of 1T-VSe₂. Systematic scanning tunneling microscopy experiments reveal that bilayer VSe₂ largely retains the bulk electronic structure, hosting a tridirectional CDW. However, monolayer VSe₂ —consistently across distinct substrates—exhibits a dimensional crossover, hosting two CDWs with distinct wavelengths and transition temperatures. Electronic structure calculations reveal that while one CDW is bulk-like and arises from the well-known Peierls mechanism, the other is decidedly unconventional. The observed CDW-lattice decoupling and the emergence of a flat band suggest that the second CDW could arise from enhanced electron–electron interactions in the 2D limit. These findings establish monolayer-VSe₂ as a host of coexisting charge orders with distinct origins, and enable the tailoring of electronic phenomena via emergent interactions in 2D materials.

KEYWORDS: two-dimensional materials, transition metal dichalcogenides, monolayer, VSe₂, charge density waves, scanning tunneling microscopy, band structure

INTRODUCTION

Charge order in crystalline materials typically manifests as a static modulation of electron density, known as a charge-density wave (CDW), accompanied by periodic modulations of the atomic lattice.1 The prototypical CDW arises in (quasi)one-dimensional (1D) systems from the “nesting” of parallel Fermi surface (FS) regions connected by the CDW propagation vector $Q_{\text{CDW}}$. As real materials do not exhibit perfect nesting, CDW formation is supported by either electron–phonon coupling (EPC), other collective excitations, or electron–electron interactions.2−7 In layered materials, CDWs often exist in proximity to other ordered phases, for example, superconductivity and magnetism,8 due to a precarious balance between competing interactions. Approaching the two-dimensional (2D) limit enhances the potential for such interplay,9 while providing new knobs to tune electronic phases, such as electric fields and strain.10−13 Notably, electron–electron interactions in the 2D limit are expected to induce competition among different CDW driving mechanisms as well as other ordered states.14−16 In practice, however, a crossover toward electronic charge order driven by dimensional reduction remains to be discovered.

Transition metal dichalcogenides (TMDCs) are well-studied hosts of conventional and unconventional CDWs.4−6,9,17 The tunability of CDWs in the ultrathin limit of several TMDCs is particularly relevant to practical electronic applications.11−13,18,19 1T-VSe₂ is a prototypical metallic TMDC with layered hexagonal crystal structure (Figure 1a). Bulk 1T-VSe₂ is paramagnetic, with a three-dimensional (3D) FS. Below temperature $T_{\text{bulk}}^{\text{cdw}} \approx 110$ K, it hosts a triple-Q (triangular) CDW with 3D character. The CDW periodicity

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\(\lambda_{\text{CDW}} \approx 4a \times 4a \times 3c + \delta\) is commensurate with the in-plane lattice constant \(a\), but incommensurate with the interlayer distance \(c\), and corresponds to a weakly nested FS region, supported by a structured EPC.\(^7\),\(^{20}\) For thicknesses below 20 nm, the FS of 1T-VSe\(_2\) transitions to 2D character, while maintaining triple-Q, 4\(a\) CDW order.\(^{21}\)

Meanwhile, monolayer (ML)-VSe\(_2\) grown epitaxially in several recent works, purportedly hosts a ground state with concomitant charge and spin orders, the nature of which is controversial.\(^{22}\)\(^{−}\)\(^{25}\) First, while some claim 4\(a\) CDW to be absent even at low temperatures,\(^{22}\)\(^{24}\)\(^{25}\) others indicate its persistence to well above room temperature.\(^{23}\) Second, several works report incommensurate superstructures with varying periodicities, viz. \(\sqrt{3}a \times 2a\), \(\sqrt{3}a \times \sqrt{7}a\), and \(\sim 2a \times 3a\),\(^{22}\)\(^{−}\)\(^{25}\) whose purported origins vary from structural distortions to nested CDWs. The relation of all these superstructures—identified via electronic density distributions over small real space regions—to any long-ranged charge order remains unclear. Finally, magnetism is suggested to emerge in ML-VSe\(_2\) despite its absence in the bulk,\(^{22}\)\(^{26}\) but both its existence and interplay with charge order are actively debated.\(^{27}\)\(^{28}\)

Disentangling these apparently conflicting observations is paramount to revealing the true nature of charge order in ML-VSe\(_2\), its driving mechanism, and its ramifications on other phases. This requires a controlled and systematic study of the CDW under varying thermodynamic conditions.

Here we report a comprehensive experimental and theoretical investigation of charge order in ultrathin epitaxial 1T-VSe\(_2\). Scanning tunnelling microscopy (STM) experiments show that while the CDW in BL-VSe\(_2\) is closely related to that in bulk, charge order in ML-VSe\(_2\) is qualitatively different. By systematically varying substrates, film thickness, and temperature, we find that ML-VSe\(_2\) consistently hosts two unidirectional (single-Q) CDWs with periods 4\(a\) and 2.8\(a\), with strikingly distinct phenomenologies. Band structure calculations elucidate that while the 4\(a\) CDW is stabilized by conventional FS nesting and EPC, the 2.8\(a\) CDW cannot be explained by such mechanisms. Instead, we find the 2.8\(a\) instability to originate from a flat band region, wherein electron–electron interactions are expected to be strongly enhanced. Our results establish ML-VSe\(_2\) as a host of coexisting CDWs with distinct driving mechanisms, demonstrating the potential of correlations for tuning electronic phases in the 2D limit.

RESULTS AND DISCUSSION

STM Imaging Experiments. Thin films of VSe\(_2\) were grown using molecular beam epitaxy (MBE) on highly oriented pyrolytic graphite (HOPG) and MoS\(_2\) substrates under ultrahigh vacuum conditions (see Methods). Both substrates are known to stabilize the 1T polymorph of VSe\(_2\) whose crystal structure is shown in Figure 1a. The films were characterized in situ using STM over temperatures of 77−200 K (see Methods). As shown in Figure 1b, controlled growth of an average thickness of 1.5 layers resulted in the formation of both ML- and BL-VSe\(_2\) regions (on HOPG) within fields-of-view accessible to STM imaging. Topographic characterization of a terraced region at 78 K (Figure 1b inset) reveals step heights of 0.9 and 0.6 nm for the first and second VSe\(_2\) layers respectively, in line with values reported previously.\(^{23}\)

Figure 1c,d displays atomic resolution topographs obtained in the BL and ML regions, respectively. As expected, both cases show the expected hexagonal arrangement of atoms with lattice constant, \(a \approx 0.34\) nm.\(^{22}\)\(^{23}\) Meanwhile, the atomic-scale superstructures seen on ML- and BL-VSe\(_2\) appear starkly different. For BL-VSe\(_2\) (Figure 1c), the superstructure is tridirectional, that is, it manifests along all three lattice directions with a single length scale. The overall phenomenology is very similar to that of the triple-Q CDW reported in bulk and thinned 1T-VSe\(_2\) crystals.\(^{21}\) In contrast, for ML-VSe\(_2\) (Figure 1d), the superstructure appears unidirectional, and has multiple length scales, consistent with recent results reported by other groups.\(^{22}\)\(^{23}\)

Crucially, complementary imaging of the ML using noncontact atomic force microscopy under similar conditions shows no corrugations beyond those of the atomic lattice (see Supporting Information (SI) Section S2), which
rules out structural distortions. Therefore, we conclude that the superstructures observed in STM imaging of ML-VSe₂ must be of electronic origin, and putatively regard them as CDWs.

In light of conflicting reports on the CDW phenomenology in ultrathin VSe₂, we systematically examine in Figure 2 the Fourier space modulations from larger STM topographs obtained for both BL and ML cases. For BL-VSe₂, Figure 2b shows the Fourier transform (FT) of a typical STM topograph. Here we find prominent peaks at $Q_1 \approx 0.25a^*$ (green circles), where $a^*$ is the magnitude of the reciprocal lattice vector, with $C_6$ symmetry, that is, along all three Bragg directions. Meanwhile, the anisotropy of Bragg peak intensities may indicate either local uniaxial strain within the sample, or asymmetry in the tip shape. Regardless, these observations are consistent with the triple-Q, 4a CDW reported in bulk and thinned 1T - VSe₂ crystals. In contrast, the FT for ML-VSe₂ shown in Figure 2d appears more complex, with only $C_2$ symmetry present. First, c.f. the BL, the ML shows the persistence of the $Q_1 \approx 0.25a^*$ peak (green circle) along a single Bragg direction, corresponding to a single-Q, 4a CDW. Second, the most prominent Fourier peak for the ML is seen at $Q_2 \approx 0.36a^*$ at an angle $\theta_1 \sim 30^\circ$ relative to the Bragg direction (magenta circle). As shown in Figure 2f, a careful inspection of the FT for the ML suggests that all remaining Fourier peaks can be assigned to higher harmonics or Bragg reflections of $Q_1$ and $Q_2$ including previously reported multiplet superstructures. While such superstructures may, in principle, be identified with several distinct wave-lengths over small topographic regions, such identifications are not consistent over length scales above 5 nm in any of the reported data. Instead, we propose that these apparent supercells are merely the result of superposing two single-Q CDWs, one of which is aligned away from a high-symmetry direction and also incommensurate with the atomic lattice.

**Temperature Dependence.** To further establish the character of CDW(s), we studied the evolution of CDW peaks in BL- and ML-VSe₂ with temperature, across both substrates. Notably, the FT of ML-VSe₂ recorded at higher temperatures (Figure 2e) reveal only a single Fourier modulation with magnitude $Q_2$, as well as its harmonics and reflections. This further evidence the presence of only two principal CDWs—$Q_1$ and $Q_2$—and suggests that they may have independent origins. At the same time, the slight thermal variation in the direction of $Q_1$ with respect to the lattice shows that the Q₁ CDW is not strongly coupled to the lattice. It also suggests a potential interplay between the two CDWs, which may lower the energetic cost of the charge ordered state when harmonics and reflections of $Q_1$ are connected by $Q_2$ (Figure 2f). Meanwhile, both BL- and ML-VSe₂ grown on MoSe₂ substrate (see SI Section S1) exhibit identical CDW phenomenology to their counterparts grown on HOPG (Figure 2), limiting the potential role of substrate-induced strain effects in driving CDW formation.

The thermal evolution of the CDW intensity in STM topographs is an established thermodynamic marker of the CDW transition.$^{29,30}$ In Figure 3, we show representative STM topographs for ML-VSe₂ on HOPG for different temperatures (extended data set in SI Section S3). While the data were recorded over varying fields-of-view, we emphasize that, within our experiments, none of the CDWs exhibit any macroscopic spatial variation across atomically smooth regions. For ease of comparison, the CDW peak intensities plotted in Figure 3 are normalized to the corresponding Bragg peak intensities for each STM topograph. Consistently across BL- and ML-VSe₂, we find that the intensity of $Q_1$ (4a CDW) drops sharply at $\sim 110$ K to a negligible magnitude, consistent with the thermal evolution of its bulk counterpart.$^{21}$ The small, finite magnitude

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**Figure 2.** Comparison of CDWs in BL and ML-VSe₂. (a–d) STM topographs (a,c: 10 × 10 nm², $V_{tip} = -0.2$ V, $I_{tip} = 200$ pA) and their respective Fourier Transforms (FTs: b,d) acquired at 78 K on BL- (a,b) and ML- (c,d) VSe₂ from adjacent terraces with no observable grain boundary. Dashed color-coded lines in (a,c) represent the real space CDW wavefronts, and corresponding circles in (b,d) denote the respective CDW wavevectors $Q_1$ (b,d: green) and $Q_2$ (d: magenta), whose magnitudes are indicated in reciprocal lattice units (rlu). Red circles denote atomic Bragg peaks in all FT images. (e) FT of STM topograph acquired on ML-VSe₂ at 116 K. Magenta circle denotes the $Q_2$ peak, while the $Q_1$ peak is absent. Dashed lines indicate the orientation of $Q_1$ and its harmonics with respect to the Bragg peak (red circle). (f) Annotated FT of ML-VSe₂ at 78 K (c.f. data in d). Green ($Q_1$) and magenta ($Q_2$) circles identify the primary CDW peaks. Color-coded arrows indicate the positions of harmonics with respect to primary and Bragg peaks. All peaks can be accounted for this way.
Figure 3. Temperature Dependence of CDW Intensities in ultrathin VSe$_2$. Thermal evolution of intensities of the CDW peaks, normalized to the averaged intensities of the six primary Bragg peaks at that temperature, as measured from FTs of 10 $\times$ 10 nm$^2$ STM topographs acquired on (a) BL- and (b) ML-VSe$_2$, respectively (data set in SI Section S3, Figure S3). Error bars show the standard deviation, incorporating the variance in Bragg and CDW peak intensities at each temperature. Insets show STM topographs at selected temperatures for BL (left) and ML (right), respectively.

of Q$_1$ in BL-VSe$_2$ at higher temperatures likely arises from small CDW pockets near defects, similar to defect-pinned CDWs at $T \gg T_{CDW}$ reported in other TMDCs.$^{29,30}$ Meanwhile, for ML-VSe$_2$, the intensity of Q$_2$—in sharp contrast to Q$_1$—remains sizable well above ~110 K, and drops to nearly zero at ~140 K. Finally, no CDW signatures are observed in the 204 K topographs (Figure 3c,g), precluding the persistence of either CDW to room temperature.$^{23}$

Overall, our systematic analysis sheds much-needed light on the presence, character, and robustness of putative charge order in ML-VSe$_2$ in view of conflicting reports in literature.$^{23,24,31-33}$ First, our AFM-STM comparison confirms the purely electronic (CDW) origin of all observed superstructures on ML- and BL-VSe$_2$ (c.f. ref 23). Second, T-dependent experiments conclusively establish the presence of two, and only two, independent single-Q CDWs in ML-VSe$_2$—Q$_1$ $\approx$ 0.25$a_*$ (i.e., $\lambda_1$ $\approx$ 4$a_*$) and Q$_2$ $\approx$ 0.36$a_*$ (i.e., $\lambda_2$ $\approx$ 2.8$a_*$), respectively. The Q$_1$ CDW is identical in magnitude, orientation, and transition temperature to the triple-Q CDW observed in BL-VSe$_2$, and to (the in-plane projection of) the CDW reported in bulk crystals. Meanwhile, the Q$_2$ CDW persists at temperatures well beyond Q$_1$ and exhibits thermal variations in its orientation with respect to the atomic lattice. Finally, the observed consistency of Q$_1$ and Q$_2$ across distinct substrates (c.f. refs 32 and 33), and of BL-VSe$_2$ with bulk (c.f. ref 31), strongly constrain the potential influence of substrate-induced strain effects on the CDW characteristics reported here. To understand the origin of this observed dichotomy in CDW characteristics within the same material, we conduct a detailed examination of the electronic structure of ultrathin VSe$_2$.

**Band Structure Calculations.** Density functional theory (DFT) calculations were performed to investigate the atomic and electronic structure of ultrathin 1T-VSe$_2$ using the Vienna *Ab Initio* Simulation Package (VASP, see Methods).$^{35}$ ML-VSe$_2$ was simulated by requiring the interlayer distance to be 25 Å, and relaxing a $4 \times 4$ atomic supercell structure, both with and without the symmetry constraints of the underlying P$\bar{3}$m1 space group.$^{36}$ In both cases, the resulting lattice is purely hexagonal, and free of any structural distortions (c.f. ref 23). This further points to the electronic origin of superstructures observed in ML-VSe$_2$, in line with our experimental findings. Subsequently, the electronic structure was computed, both with and without including spin polarization. The resulting energies are nearly equal for both cases. This suggests, in conjunction with the absence of spin splitting in angle-resolved photoemission spectroscopy (ARPES) results,$^{7,5,24,26,27,34,37,38}$ that magnetic order, even if present in ML-VSe$_2$, is unlikely to play a significant role in the energetics of charge ordered states.

The DFT band structure (Figure 4a) is broadly in agreement with the ARPES spectral function measured for ML-VSe$_2$.$^{32}$ The data in ref 34 provides a valuable benchmark given its high quality, large momentum range, and qualitative agreement with other ARPES reports, including data acquired on our samples (SI Section S4, Figure S4).$^{26}$ Both techniques find a single band of predominantly d-orbital character crossing the Fermi energy $E_F$. Previous works have emphasized the importance of the nesting of the sides of the FS lobes at the BZ edge.$^{23,24,39}$ The DFT electronic structure, however, underestimates $k_F$ along M$-$K and suggests a “nesting vector” along $a_*$ of length 0.21 rlu. This falls short of the vector extracted from ARPES data (0.54 $\pm$ 0.04 Å$^{-1}$), which corresponds to 0.25 $\pm$ 0.02 rlu.$^{7,37}$ The DFT band along $\Gamma$–M also appears more dispersive than that in ARPES, while along $\Gamma$–K the DFT band is higher (50–200 meV) than the magnitude expected from the high photoelectron count around $\Gamma$.$^{23,24,26,27,34,37}$ These discrepancies are likely due to the inability to duly account for electronic correlations.$^{38}$ As a result, our *ab initio* calculations may not capture the electronic structure near $E_F$ with sufficient quantitative accuracy to describe CDW energetics.

We therefore complement the DFT calculation with a tight-binding (TB) fit to the ARPES data in ref 34 (see Methods), the results of which are compared to the DFT in Figure 4. In agreement with reported ARPES spectra, the TB fit shows a flat band region around the $\Gamma$-point, an indicator of strong...
Figure 4. Electronic Structure of ML-VSe$_2$. (a) Electronic band structure of ML-VSe$_2$ obtained from DFT calculations (red line), compared to published ARPES measurements of the spectral function for epitaxially grown ML-VSe$_2$ at $T = 170$ K (shaded yellow, crosses: peak positions, lines: full width at half-maximum). The line width of the experimental data greatly exceeds the experimental resolution. Blue line is a tight-binding (TB) fit to the ARPES-measured, near-$E_F$ band structure, where $E_F$ is the Fermi energy. (b) Fermi surface (FS) of ML-VSe$_2$, obtained from the TB fit in (a) by plotting states within $± 1$ meV (dark blue) and $± 10$ meV ($± k_BT$ for $T = 100$ K, light blue) of $E_F$. Dotted red line shows the DFT FS, which qualitatively deviates from the TB fit. Hexagon shows the Brillouin zone (BZ), and the arrows indicate FS regions visually appearing to be nested by the experimentally determined CDW wavevectors.

correlations. The difference in topology between the DFT and TB FS (Figure 4b) is due to the proximity of a van Hove singularity to $E_F$.4,34 Overlaying the CDW vectors extracted from our STM data onto the FS visually suggests that $Q_1$ corresponds to nesting between the sides of neighboring triangular FS pockets at the BZ edge, while $Q_2$ connects the flat-band region around $\Gamma$ to the pocket corners around K.

**Nesting and Correlated Instabilities.** A conventional CDW instability at wavevector $Q_{CDW}$ results from a maximum in its electronic susceptibility $\chi(Q)$ for $Q = Q_{CDW}$.2,44 In the weak electron–phonon coupling (EPC) limit (see Methods), $D_2(Q)$ can be expressed as

$$D_2(Q) = - \sum_{k \in BZ} \bar{\epsilon}_{k+q}^2 \frac{f(E_k) - f(E_{k+q})}{E_k - E_{k+q} + i\delta}$$

(1)

Here, $f(E)$ is the Fermi–Dirac function, $E_k$ is the bare (nonrenormalized) electronic dispersion, and $\delta$ is a small regularizer (0.1 meV in this work). The EPC matrix elements, $\bar{\epsilon}_{k+q}^2$, are often approximated to unity, resulting in the Lindhard, or bare susceptibility, $\chi(Q)$. However, for TMDCs whose near-$E_F$ behavior is governed by $d$-band(s), several works have established a more realistic approximation to $\chi(Q)$ via the electronic band structure.41,42 (see Methods). Here, we use the TB fit to calculate the bare ($\chi(Q)$) and structured ($D_2(Q)$) electronic susceptibilities, which are shown in Figure 5a,b, respectively.

The green circles in Figure 5a,b indicates the maximum of the bare (structured) susceptibility, which lies at $Q = (0,0.28) \approx Q_1$. Its proximity to a commensurate value suggests that the corresponding CDW will lock to 0.25 $rlu$ ($\lambda = 4\alpha$) due to CDW-lattice interactions. Although its periodicity is the same as that of the CDW observed in bulk and BL-VSe$_2$, the FS for the ML is strictly 2D, and the parts of the FS involved in CDW formation may differ. To elucidate the role of the FS in the observed CDWs, we plot in Figure 5c,d the k-resolved contributions to $\chi(Q)$ and $D_2(Q)$ for $Q = Q_1$. As anticipated in Figure 4b, the dominant contributions to $\chi(Q_1)$ arise from the parallel edges of the K-centered pockets, while the $\Gamma$-centered FS region plays a negligible role. The well-nested K-pocket edges with opposite group velocities are therefore inherently unstable to a Peierls-like CDW. The EPC matrix elements further enhance the contribution of these Q$_1$-connected regions to $D_2(Q_1)$, thereby confirming the conventional origin of the Q$_1$ CDW in ML-VSe$_2$.

In contrast, the phenomenology for $Q = Q_2$ does not fit the conventional CDW framework. As highlighted by the magenta circles in Figure 5a,b, this wavevector lies in the middle of a susceptibility plateau, and lacks a well-defined maximum. The dominant contribution to the bare susceptibility at $Q = Q_2$ comes from the $\Gamma$-centered flat band region, with smaller contributions from the K-centered pockets (see Figure 5e). However, the corresponding $D_2(k,Q_2)$ in Figure 5f shows that the EPC matrix elements strongly suppress the intensity in these regions, and the remaining contributions are insufficient to drive the Q$_2$ CDW according to an EPC-assisted Peierls scenario. While the perturbative expansion used for the structured susceptibility calculations$^{40,42}$ may not fully capture EPC in flat bands, that the origin of the Q$_2$ CDW lies beyond the Peierls description of CDWs is consistent with its empirical characteristics, viz. varying orientation with respect to the lattice, absence in BLs (and beyond), and the lack of a discernible peak in $\chi(Q)$. In the 2D limit of layered TMDCs like 1T-VSe$_2$, the screening of Coulomb interactions between electrons is much reduced.43 The relative importance of the unscreened interactions is further enhanced within flat bands associated with a van Hove singularity, such as at the near-$\Gamma$ region in ML-VSe$_2$ (Figure 4).23,34,37,38 Indeed, the measured line width, or self-energy, of the band near $E_F$ is much larger than the experimental resolution,43 supporting the presence of strong electronic correlations.44 Such interactions can considerably renormalize electron and phonon properties, and enable CDW order at momenta that do not correspond to peaks in the conventional susceptibility ($\chi(Q)$ or $D_2(Q)$). Indeed, such correlation-driven CDWs have been predicted to exist in TMDCs$^8$ including in ML-VSe$_2$,39 and are consistent with the unusual characteristics of the Q$_2$ CDW. Crucially, a correlation-driven mechanism for the Q$_2$ CDW offers the only viable explanation of its prevalence over a well-nested
counterpart (Q1), and the complete gapping of the FS,23,34,37,38 despite the absence of any associated feature in susceptibility calculations based on models of noninteracting electrons. Further, we conjecture that the single-q character of the Q2 CDW, which breaks the 3-fold rotational symmetry of the lattice, makes it energetically favorable for the Q1 CDW (nominally triple-q) to also order in a single-q configuration.

The interplay of these CDWs could be examined in future theoretical works by iteratively incorporating the resulting lattice distortions.

CONCLUSIONS

In summary, our systematic experimental and theoretical efforts elucidate that 1T-VSe2 undergoes a dimensional crossover as its thickness is reduced to a single layer. While BL-VSe2, akin to bulk, hosts a conventional triple-Q CDW, ML-VSe2 hosts two distinct single-Q CDWs with contrasting characteristics. One, with λ1 ≃ 4a, behaves similarly to its BL/bulk counterpart, and arises from a weak-coupling Peierls mechanism utilizing nested FS regions. In contrast, the dominant CDW, with λ2 ≃ 2.8 a, cannot be explained within the conventional EPC-assisted Peierls framework. Instead, the observed thermal evolution and the calculated susceptibility suggest that this CDW—unique to the ML—arises from a flat region of the electronic band structure, where interactions and correlation effects are expected to dominate.

Monolayer VSe2 stands apart in hosting two coexisting charge orders with distinct physical origins. Conventional electronic materials are typecast by the mechanisms and phenomena they host. Our work suggests that ML-VSe2 transcends such labeling, and hosts coexisting ordered states originating from contrasting coupling mechanisms. The prospect of such emergent electron correlations and ensuing ordered states presenting themselves in 2D TMDCs more generally is particularly promising given their prominence in the plethora of proposed designs for heterogeneous layered materials.10,45,46 Their potential for tunability and their interplay with conventional charge and spin orders in the ultrathin limit is promising for realizing exotic ordered states on one the hand, and for applications in multifunctional electronics on the other.
temperature. Electrochemically etched tungsten tips were used with bias voltage applied to the tip, while the sample holder was grounded. STM images were acquired using constant current mode. For nc-AFM imaging, the constant-height mode with an oscillation amplitude of 10 nm was used to record the frequency shift (Δf) of the qPlus resonator (sensor frequency f_s ≈ 24 kHz, Q ≈ 8000). A lock-in technique was used to measure dI/dV spectra, with a modulation of 625 Hz and 30 mV.

**DFT Calculations.** 1T-VSe_2 belongs to the space group P6_3m1, with the lattice parameters for the monolayer crystal being a = b = 3.33 Å, c = 12.0 Å. First-principle atomic and electronic structure calculations were performed within the density functional theory (DFT) framework as implemented in the Vienna Ab Initio Simulation Package (VASP) with a plane-wave basis up to a cutoff of 500 eV. To simulate the monolayer, we artificially set the distance between two layers of VSe_2 to 25 Å. The Perdew–Burke–Ernzerhof (PBE) functional form was used for the exchange-correlation functional. The Γ-centered k-mesh was set to be 25 × 25 × 1 in the Brillouin zone for the self-consistent calculation. The relaxation of atomic structure was done in two ways. First, a 4 × 4 supercell was relaxed under the symmetry constraints of the space group. This process was then repeated without any symmetry constraints applied.

**Tight-Binding Calculations.** A tight-binding fit was performed for the single d-orbital band crossing the Fermi level in the available ARPES data of ref 34. To obtain the best fit, we used an expansion of the dispersion E(k) in functions respecting the lattice symmetries. Including terms to fifth order, the fit can be expressed as

\[
E(k) = t_{1} + t_{2}(2 \cos(\xi) \cos(\eta) + \cos(2\zeta)) + t_{3}(2 \cos(\xi) \cos(\eta) + \cos(2\phi)) + t_{4}(2 \cos(2\xi) \cos(2\eta) + \cos(4\zeta)) + t_{5}(2 \cos(\xi) \cos(3\eta) + \cos(3\phi)) + t_{6}(2 \cos(3\xi) \cos(2\eta) + \cos(6\phi)) \tag{2}
\]

where \( \xi = \frac{k_x}{a} \) and \( \eta = \frac{k_y}{b} \), while \( k_x, k_y \) are given in units of \( \frac{\pi}{a} \) and \( \frac{\pi}{b} \) respectively. The best fit to ARPES data based on this form of the dispersion is shown in Figure 4.

**Susceptibility Calculations.** In the limit of weak electron–phonon coupling, the electronic susceptibility can be derived from a perturbative expansion of the phonon propagator, using the random phase approximation (RPA). Neglecting vertex corrections, which should be small,\(^{10}\) the renormalized phonon propagator is described by \( D_{\text{RPA}} = (D_{\text{bare}} - D_{\text{bare}}^{-1})^{-1} \), with bare phonon propagator \( D_{\text{bare}} \) and electronic susceptibility \( D_{\text{bare}} \) given by\(^{10}\)

\[
D_{2}(q) = -\sum_{k \in \text{BZ}} g_{k+q}^{2} f(E_{k}) f(E_{k+q}) \tag{3}
\]

Here, \( f(E) \) is the Fermi–Dirac distribution function, \( E_{k} \) is the nonrenormalized electronic dispersion and we use a small regulator \( \delta = 0.1 \text{ meV} \). If the system has an intrinsic or electron–phonon driven CDW instability within the weak-coupling limit, the susceptibility will exhibit a maximum at the CDW wave vector \( Q_{L} \). Generally, the electron–phonon coupling (EPC) matrix elements \( g_{k+q}^{2} \) are difficult to compute exactly. However, for this reason, it is common to set them to unity, resulting in the Lindhard function:

\[
\chi(q) = \sum_{k \in \text{BZ}} f(E_{k}) f(E_{k+q}) \tag{4}
\]

In previous works, it has been shown that the EPC matrix elements can be approximated based purely on the electronic dispersion. This approximation has been well-tested for transition metal compounds with d-orbital character at \( E_{\text{F}} \). In the case of a single band crossing \( E_{\text{F}} \), the expression becomes

The orientation of \( g_{k+q} \) indicates the direction of phonon polarization. We consider longitudinal CDWs, such that the relevant component of the EPC tensor is parallel to the in-plane phonon momentum: \( g_{k+q} = g_{k+q_0} \cdot |q_0|/|q_0| \).

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.nano.1c08304.

Supporting data on MoS_2 substrate; comparison of STM- and AFM-imaging of ML-VSe_2; full data set for temperature-dependence presented in manuscript; and comparison of tight-binding fit to ARPES data (PDF)

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**Author Contributions**

R.C. and J.H. contributed equally to this work.

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R.C., Y.L.H., J.G., and X.H. performed the experiments and analyzed the data. S.S. and T.D. performed the ab initio calculations.
calculations. J.H. performed the tight-binding and susceptibility calculations. J.v.W., A.S., and A.T.S.W. coordinated and supervised the work. All authors discussed the results and provided inputs to the manuscript.

Notes
The authors declare no competing financial interest.

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