

Temperature-dependent local structure and lattice dynamics of 1T-TiSe₂ and 1T-VSe₂ probed by X-ray absorption spectroscopy

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Abstract

The local atomic structure and lattice dynamics of two isostructural layered transition metal dichalcogenides (TMDs), 1T-TiSe₂ and 1T-VSe₂, were studied using temperature-dependent X-ray absorption spectroscopy at the Ti, V, and Se K-edges. Analysis of the extended X-ray absorption fine structure (EXAFS) spectra, employing reverse Monte Carlo (RMC) simulations, enabled tracking of the temperature evolution of the local environment in the range of 10-300 K. The atomic coordinates derived from the final atomic configurations obtained using the RMC method were used to calculate the partial radial distribution functions (RDFs) and the mean-square relative displacement (MSRD) factors for the first ten coordination shells around the absorbing atoms. Characteristic Einstein frequencies and effective force con-

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stants were determined for Ti–Se, Ti–Ti, V–Se, V–V, and Se–Se atom pairs from the temperature dependencies of MSRDs. The obtained results reveal differences in the temperature evolution of lattice dynamics and the strengths of intralayer and interlayer interactions in TiSe_2 and VSe_2 .

Keywords: Transition metal dichalcogenides; Interlayer and intralayer coupling; Extended X-ray absorption fine structure; Reverse Monte Carlo simulations; Effective force constants

1. Introduction

Transition metal dichalcogenides (TMDs) have garnered significant attention in the scientific community due to their layered structure, resulting in remarkable electronic, optical, mechanical, tribological, catalytic, and magnetic properties as well as numerous possible technological applications [1, 2, 3, 4]. Among all, the charge density wave (CDW) behavior observed for a number of TMDs MX_2 ($\text{M} = \text{Ti}, \text{V}, \text{Ta}$; $\text{X} = \text{S}, \text{Se}$) is particularly fascinating [5, 6, 7]. Indeed, TMDs represent the first layered materials in which the presence of CDWs was detected [8]. The CDW state involves the periodic modulation of the electron density within the material coupled with lattice distortion, leading to changes in the material electronic and optical properties [9, 10]. Models describing CDW mechanisms are based solely on crystal structure distortion/superlattice formation, exciton–phonon interactions, or electron–phonon interactions [11, 12]. Due to the minute nature of distortion or atomic shifts in the CDW state ($\sim 0.1\text{--}0.15 \text{ \AA}$), the resulting superstructure might generate only relatively faint secondary peaks within the X-ray diffraction patterns [10].

In this paper, two isostructural TMDs, namely titanium selenide (TiSe_2) and vanadium selenides (VSe_2), were selected to evaluate the effect of the CDW ordering on their local environments. The choice of TMDs was dictated by the feasibility of measuring high-quality X-ray absorption spectra at both the metal and chalcogenide K absorption edges.

Under ambient conditions, both TiSe_2 and VSe_2 crystallize in the 1T polytype with a space group of $P\bar{3}m1$ (164) (Fig. 1) [13, 14]. In this structure, the Ti(V) atoms are octahedrally coordinated to six covalently bonded

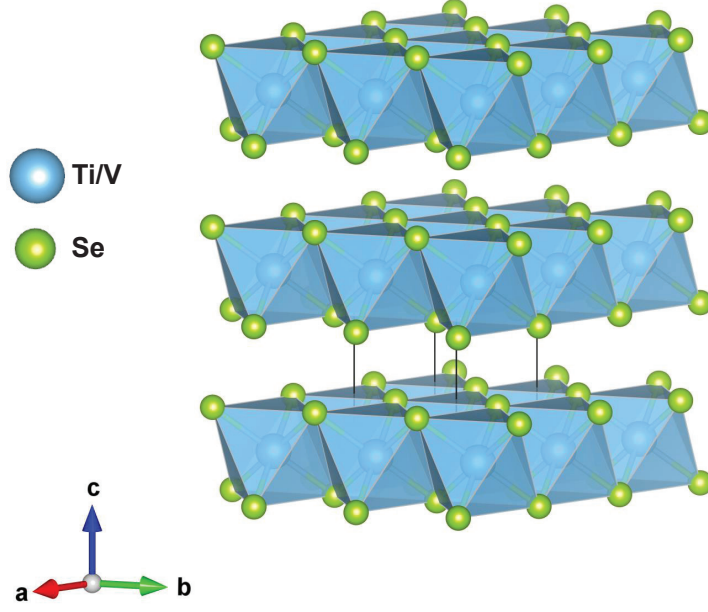


Figure 1: Crystallographic structure of trigonal (space group $P\bar{3}m1$ (164)) TiSe_2 (VSe_2) [13, 14].

Se atoms, and $[\text{Ti(V)}\text{Se}_6]$ octahedra are connected by edges into layers, joined by weak van der Waals (vdW) interactions. The interactions between neighbouring layers play an important role in assembling and exfoliation of vdW (hetero)structures and, thus, allow engineering of their electronic, optical, and mechanical properties [15, 16, 17]. While the importance of weak inter-layer coupling is recognized, its accurate experimental quantification remains challenging [16, 18].

A commensurate CDW phase in bulk 1T- TiSe_2 forms below $T_{\text{CDW}} \approx 202$ K with a $2a \times 2a \times 2c$ superlattice [19, 11]. The neutron diffraction study proposed the low-temperature (at 77 K) lattice distortion due to displacements of about 0.085 Å for Ti and 0.028 Å for Se parallel to the plane of the layer [19].

However, more recent synchrotron X-ray scattering experiments pointed to a zone-boundary phonon softening mechanism of CDW in 1T-TiSe₂ [20, 21].

1T-VSe₂ undergoes an incommensurate CDW transition at $T_{\text{CDW}} \approx 110$ K and a commensurate CDW transition at ≈ 80 K, forming a $4a \times 4a \times 3c$ superlattice [22, 23]. The origin of the CDW in VSe₂ was attributed to the electron-phonon interaction, based on inelastic X-ray scattering and first-principles calculations [24]. Besides, an important role of vdW forces in the CDW melting was proposed, likely caused by the out-of-plane nature of the CDW, which modulates interlayer distance [24]. Recently, a pressure-induced CDW state was observed in VSe₂ at room temperature within the pressure range of 10-15 GPa by Raman spectroscopy [25] and X-ray diffraction (XRD) [10]. The Se K-edge X-ray absorption spectroscopy (XAS) was also used in [10] to probe local distortions but only qualitative analysis was reported.

In this study, we employed XAS combined with the advanced data analysis methodology of extended X-ray absorption fine structure (EXAFS), based on reverse Monte Carlo (RMC) simulations coupled with an evolutionary algorithm (EA) approach [26, 27], to investigate the local atomic structure and lattice dynamics in bulk 1T-TiSe₂ and 1T-VSe₂.

The RMC method was originally proposed more than 30 years ago by McGreevy and Pusztai [28] to determine a three-dimensional model of the atomic structure of disordered systems that is consistent with the experimentally measured structure factor or the obtained radial distribution function. Two years later, it was first used for the analysis of EXAFS spectra by Gorman and McGreevy [29]. In recent years, the RMC method has gradually gained popularity due to its ability to properly incorporate thermal and static

disorder effects in multiple-scattering EXAFS calculations, easily introduce different constraints provided by other experimental techniques, and account for EXAFS spectra measured at multiple absorption edges [30, 31, 32, 33]. Further details on the strengths and weaknesses of the method can be found in a recent review [34].

We have demonstrated recently [18] that employing the RMC method for $2\text{H}_c\text{-MoS}_2$ enables the retrieval of structural data from distant coordination shells and, thus, provides valuable information on both interlayer and intralayer coupling. Here we took advantage of high-quality EXAFS spectra measured at two (Ti/V and Se) K absorption edges and performed their simultaneous analysis to reconstruct the temperature evolution of the local environment in the corresponding TMDs in the range of 10-300 K. This allowed us to determine the amplitudes of thermal vibrations for atoms located in the first ten coordination shells around absorbing atoms, which were further used to evaluate the strengths of interlayer and intralayer interactions. We showed that despite the isostructural nature of TiSe_2 and VSe_2 , the temperature-dependent evolution of their lattice dynamics has some peculiarities.

2. Materials and methods

2.1. Synthesis procedure

TiSe_2 and VSe_2 were prepared using the chemical vapor transport technique with an I_2 transport agent [35, 36]. The starting materials titanium (or vanadium) and selenium in the form of powder were weighed out with a molar ratio of $\text{M:Se} = 1:2$ ($\text{M} = \text{Ti}$ or V) and loaded into a quartz am-

poule together with the iodine (molar ratio $I_2:M = 0.05$). The filled ampoule was evacuated to pressure 10^{-5} torr and sealed at a length of approximately 12-14 cm using the oxygen-methane flame. For $TiSe_2$ synthesis the ampoule was heated in a two-zone furnace with $T_{hot} = 700$ °C at the hot zone and $T_{cold} = 600$ °C at the cold zone for 24 h, then naturally cooled down (for VSe_2 $T_{hot} = 820$ °C, $T_{coldt} = 650$ °C, heating time 72 h).

2.2. X-ray diffraction

The phase purity of the TMD samples was confirmed using X-ray powder diffraction (XRD). Diffraction patterns were collected at room temperature using a benchtop Rigaku MiniFlex 600 diffractometer with Bragg-Brentano θ - 2θ geometry. An X-ray tube with copper anode (Cu $K\alpha$ radiation, $\lambda = 1.5418$ Å), operated at $U = 40$ kV and $I = 15$ mA, was used as a source. The obtained X-ray diffraction patterns of $TiSe_2$ and VSe_2 are compared in Fig. 2 with the reference ones. All Bragg peaks were indexed to pure $TiSe_2$ (PDF Card 04-003-1758) and VSe_2 (PDF Card 01-074-1411) phases possessing $P\bar{3}m1$ (164) space group. Slight variations in peak intensities are noticeable, likely attributed to the favored alignment of layered structures that could arise during the process of sample preparation. No impurity phases were detected, demonstrating the single-phase composition of the synthesized samples.

2.3. X-ray absorption experiments

The temperature-dependent (10-300 K) X-ray absorption spectra of bulk 1T- $TiSe_2$ and 1T- VSe_2 were recorded at Ti (4966 eV), V (5465 eV) and Se (12658 eV) K-edges in transmission mode at the DESY PETRA III P65

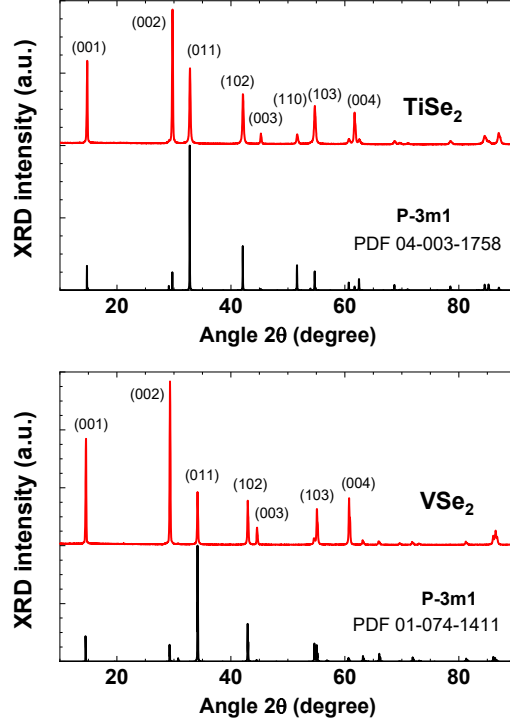


Figure 2: Powder X-ray diffraction patterns of synthesized 1T-TiSe₂ and 1T-VSe₂. Main Bragg peaks are indexed to the $P\bar{3}m1$ (164) space group. Reference patterns corresponding to TiSe₂ (PDF Card 04-003-1758) and VSe₂ (PDF Card 01-074-1411) phases are shown for comparison.

Applied XAFS undulator beamline [37]. The storage ring operated in top-up 480 bunch mode at the energy $E = 6.08$ GeV and current $I = 100$ mA. The synchrotron radiation was monochromatized using a Si(111) double-crystal monochromator, and its intensity before and after the sample was measured by two ionization chambers. The harmonic rejection was achieved by the uncoated silicon plane mirror.

Experimental Ti, V and Se K-edge EXAFS spectra were extracted following the conventional procedure [38] using the XAESA code [39]. The EXAFS

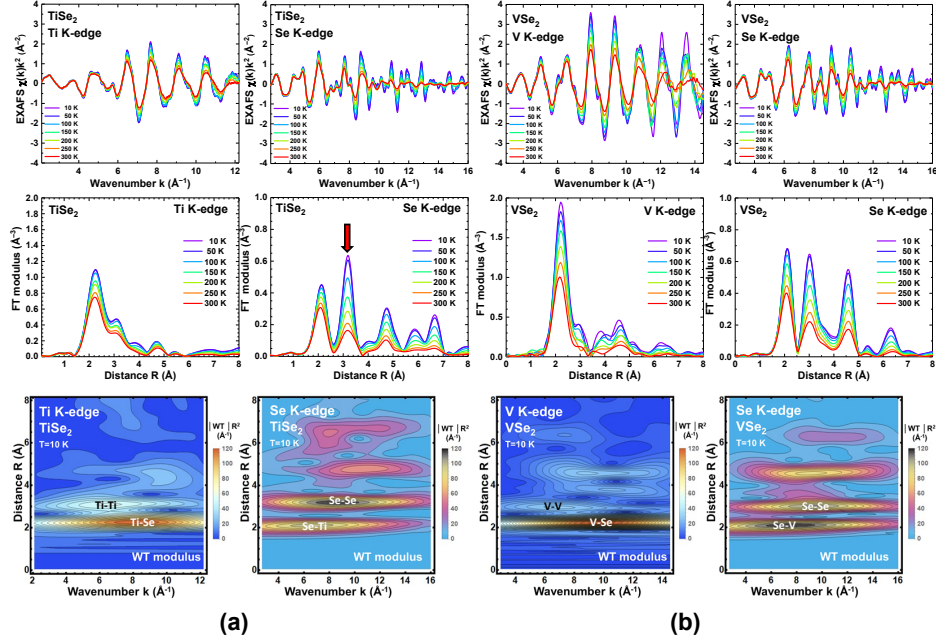


Figure 3: Temperature-dependent EXAFS spectra $\chi(k)k^2$ (top panels), their Fourier transforms (only modulus is shown) (middle panels) and wavelet transforms at 10 K (bottom panels) for TiSe₂ (a) and VSe₂ (b).

spectra $\chi(k)k^2$ and their Fourier transforms (FTs) are shown at selected temperatures in Fig. 3. Note that the FTs were not adjusted to compensate for the backscattering phase shift of atoms, therefore, the positions of all peaks appear shifted towards shorter distances compared to their crystallographic values.

2.4. Reverse Monte Carlo simulations

The structural information encoded in the experimental EXAFS spectra was extracted using the reverse Monte Carlo (RMC) method based on an evolutionary algorithm (EA) approach implemented in the EvAX code [26, 27]. The RMC/EA-EXAFS technique aims to minimize the discrepancy

between the experimental and calculated EXAFS spectra. As a result, it enables the reconstruction of the three-dimensional structural model of the material by introducing random atomic displacements within specific geometric constraints [26, 27].

In this study, the initial structural models for the RMC/EA calculations were constructed based on diffraction data [13, 14]. TiSe_2 and VSe_2 crystallize in the space group $P\bar{3}m1$ (164) with the lattice parameters $a = b = 3.540 \text{ \AA}$, $c = 6.008 \text{ \AA}$ for TiSe_2 [13] and $a = b = 3.355 \text{ \AA}$, $c = 6.134 \text{ \AA}$ for VSe_2 [14]. A supercell with a size of $6a \times 6b \times 4c$, including 432 atoms, and periodic boundary conditions was constructed for both compounds. During the RMC calculations, the size and shape of the supercell were fixed, and the number of atoms in the supercell was constant. At each iteration, all atoms in the supercell were displaced with the maximum allowed displacement set to $0.4 \text{ \AA}$. As a result of such small displacements, the coordination numbers were held constant throughout the calculation. 32 atomic configurations were employed simultaneously in the EA method [27].

The configuration-averaged EXAFS spectra at the Ti/V and Se K-edges were calculated using the ab initio self-consistent real-space multiple-scattering FEFF8.5L code [40, 41] taking into account multiple-scattering contributions up to the 5^{th} order. The complex energy-dependent exchange-correlation Hedin-Lundqvist potential was employed to account for inelastic effects [42]. The values of the non-structural parameters, such as E_0 and S_0^2 [38], were determined by performing simulations for the highest-quality experimental data (the EXAFS spectra measured at 10 K) to achieve optimal alignment and minimize residuals in the wavelet transform space. Subsequently, the

amplitude scaling parameter S_0^2 was set to 0.9 (for Ti K-edge) or 1.0 (for V and Se K-edges) and remained fixed for the entire series of experimental data, albeit potentially introducing some systematic errors. Multiple RMC calculations were conducted at each temperature using different sequences of pseudo-random numbers to estimate statistical dispersion. The mean-square relative displacement (MSRD) factors σ^2 for the first ten coordination shells around the absorbing atoms were calculated using the median absolute deviation (MAD) method [44, 45]. The deviation between the smallest and largest MSRD values obtained from these calculations is comparable to the effect resulting from adjusting S_0^2 by ± 0.1 .

The structural model was adjusted in each RMC/EA calculation by minimizing the difference between the Morlet wavelet transforms (WTs) [43] of the experimental and calculated EXAFS spectra at two absorption edges (Ti and Se for TiSe_2 or V and Se for VSe_2) simultaneously. As a result, good agreement between the experimental and configuration-averaged EXAFS spectra was achieved in both the direct (R) and reciprocal (k) space.

An example of fits at the selected temperatures is shown in Fig. 4. The convergence of each RMC/EA simulation was attained after 3000 iterations. To improve statistics, seven independent RMC/EA calculations were carried out for each experimental data set, employing distinct sequences of pseudo-random numbers. The agreement between the configuration-averaged EXAFS spectra and the experimental data at all temperatures affirms the reliability of the obtained structural models. The calculated partial contributions of single-scattering and combined multiple-scattering paths to the total EXAFS spectra $\chi(k)k^2$ at 10 K are shown in Fig. 5. Note that the total number

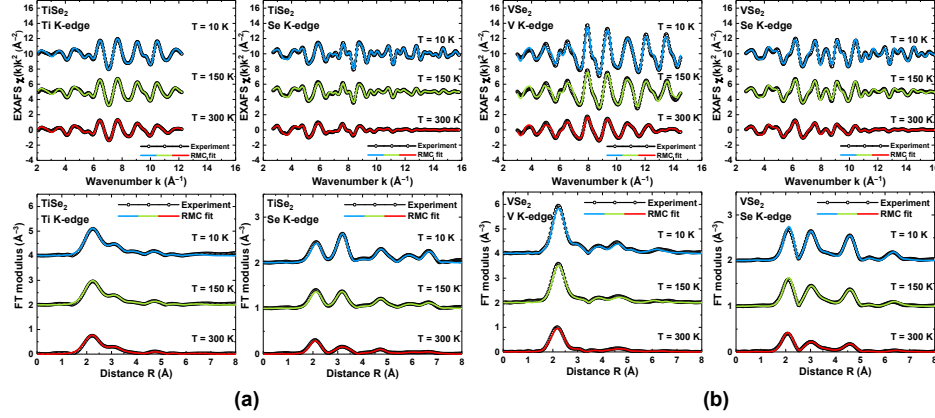


Figure 4: Results of the RMC/EA calculations for the Ti and Se K-edges in TiSe_2 (a) and V and Se K-edges in VSe_2 (b) at selected (10, 150 and 300 K) temperatures. The EXAFS spectra $\chi(k)k^2$ are displayed within their respective fitting ranges. The fitting R -space ranges were 1.5–5.9 Å (Ti K-edge) and 1.2–7.1 Å (Se K-edge) for TiSe_2 and 1.0–7.0 Å (V K-edge) and 1.2–7.5 Å (Se K-edge) for VSe_2 .

of single- and multiple-scattering paths included in the analysis was up to 1200, with those having a distance difference of less than 0.5 Å grouped together in the RMC calculations.

The atomic coordinates derived from the final RMC/EA configurations were utilized to calculate the partial radial distribution functions (RDFs) and to obtain relevant structural parameters. The MSR factors σ^2 for Ti–Se, Ti–Ti, V–Se, V–V, and Se–Se atom pairs, which account for the thermal and static disorder, were calculated using the MAD method for the first ten coordination shells [44, 45].

The temperature dependencies of the obtained MSR factors, $\sigma^2(T)$, were approximated using the correlated Einstein model [46]

$$\sigma^2(T) = \sigma_{\text{st}}^2 + (\hbar/2\mu\omega_E) \coth(\hbar\omega_E/2k_B T) \quad (1)$$

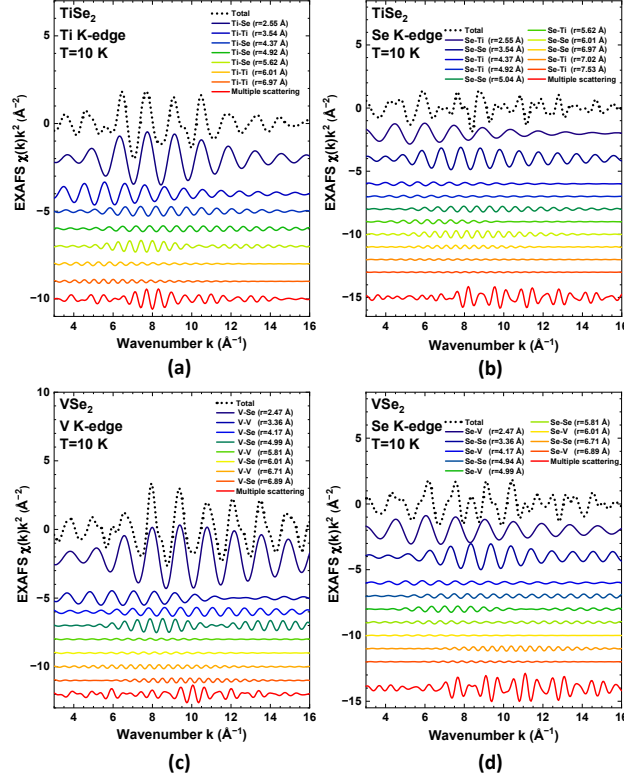


Figure 5: Example of the calculated partial contributions (solid curves) of single-scattering and combined multiple-scattering paths to the total (dotted curves) EXAFS spectra at the Ti(Se) K-edges in TiSe_2 (a, b) and V(Se) K-edges in VSe_2 (c, d) at 10 K. The effective distances of the single-scattering paths are given in brackets.

where σ_{st}^2 is the static contribution, μ is the reduced mass of the atomic pair, ω_E is the characteristic Einstein frequency, and k_B is the Boltzmann constant. The Einstein frequency ω_E can be used to determine the effective force constant $\kappa = \mu\omega_E^2$ [47], which is a useful parameter for estimating and comparing the strength of interatomic interactions. The values of the characteristic Einstein frequencies ω_E and the effective force constants κ for the relevant atomic pairs are reported in Tables 1 and 2. The uncertainties

reported in the tables correspond to standard errors of the fits.

3. Results and discussion

The crystallographic structure of TiSe_2 (VSe_2) (Fig. 1) consists of a titanium (vanadium) atomic layer sandwiched between two selenium atomic layers [13, 14]. These layers are weakly bonded to each other along the c -axis through vdW forces. The interlayer gap between Se atoms is about 3.12 Å in TiSe_2 and 3.16 Å in VSe_2 . Ti(V) atoms are covalently bonded to six selenium atoms, forming regular octahedra. The values of Ti–Se, Ti–Ti, and Se–Se interatomic distances for the first ten coordination shells in TiSe_2 are reported in Table 1, while the values of V–Se, V–V, and Se–Se interatomic distances in VSe_2 are summarized in Table 2.

Despite the similarity between the two crystallographic structures, a notable difference can be observed as well. In the case of VSe_2 , the interatomic distances between selenium atoms along the c -axis direction within the same layer and across adjacent layers coincide ($r_{\text{Se}_0-\text{Se}_{3/4}^{**}} = 3.63$ Å), adding complexity to the analysis. In contrast, in TiSe_2 , the Se_3^* atoms in the adjacent layer are approximately 0.1 Å closer to the absorbing Se_0 than the Se_4 atoms within the same layer, so the two interatomic distances are different $r_{\text{Se}_0-\text{Se}_3^*} = 3.58$ Å and $r_{\text{Se}_0-\text{Se}_4} = 3.68$ Å.

The Ti K-edge EXAFS spectra of TiSe_2 and their FTs are dominated by a contribution from the first two coordination shells at all studied temperatures in the range of 10-300 K (Fig. 3(a)). These coordination shells consist of six selenium and six titanium atoms, respectively. The complex pattern of WTs observed at longer distances is attributed to outer shells and

multiple-scattering effects. Note that heavier Se atoms with an atomic mass of 78.971 amu produce a greater impact on the EXAFS spectra at larger k values, whereas lighter Ti elements with an atomic mass of 47.867 amu contribute at lower k -values.

The Se K-edge EXAFS spectra of TiSe_2 contain contributions from scattering paths that extend beyond ~ 7 Å as evident in the FTs and WTs in Fig. 3(a). Their analysis can yield novel insights into the interactions between layers [18]. Indeed, the second peak at about 3.2 Å in FTs (indicated with a red arrow in Fig. 3(a)) contains contributions from intralayer ($r_{\text{Se}_0-\text{Se}_2} = 3.54$ Å, $r_{\text{Se}_0-\text{Se}_4} = 3.68$ Å) and interlayer ($r_{\text{Se}_0-\text{Se}_3^*} = 3.58$ Å) selenium atoms, and its amplitude becomes significantly suppressed upon sample heating. Note that the amplitude of other peaks up to ~ 7 Å is also reduced upon increasing thermal disorder but the peaks remain distinguishable even at 300 K.

While the reliable analysis of distant coordination shells is challenging within the conventional EXAFS methodology [38], it can be performed using the RMC/EA method [18, 27, 48]. Examples of the RMC/EA simulations of the Ti and Se K-edge EXAFS spectra for TiSe_2 at three selected temperatures (10, 150 and 300 K) are shown in Fig. 4 in k - and R -space. Good agreement between experimental and calculated EXAFS spectra is observed, enabling a comprehensive analysis of thermal disorder effects within the first ten coordination shells up to about 6 Å.

Different partial contributions due to single-scattering and combined multiple-scattering paths to the total EXAFS spectra $\chi(k)k^2$ of TiSe_2 and VSe_2 at 10 K are compared in Fig. 5. As expected, the strongest contributions come from the nearest shells. At the Ti and V K-edges, the first shell dominates,

while the next four shells around Ti and three shells around V also exhibit large amplitudes. In the case of Se, the first and second shells dominate, while the amplitudes of other shells are relatively smaller. Note that the combined multiple-scattering contributions are important at all three absorption edges.

The coordinates of atoms derived from the RMC/EA simulations were employed to calculate the partial RDFs $g(r)$ (Fig. 6) and the MSDs σ^2 (Fig. 7) for Ti–Ti, Ti–Se(Se–Ti), and Se–Se atom pairs at each temperature. The temperature dependencies of the obtained MSDs $\sigma^2(T)$ in the range of 10–300 K were further approximated using the correlated Einstein model [46]. As a result, the characteristic Einstein frequencies ω_E and the effective force constants κ were obtained for all coordination shells and are reported in Table 1.

The effective force constants κ for the first two coordination shells of titanium (Ti₀–Se₁ and Ti₀–Ti₂) are about 52 N/m and 39 N/m, respectively, suggesting strong interaction between the nearest atoms. The interactions between absorbing Ti₀ and the next Se atoms located in the same (Se₃) or in the adjacent layer (Se₄^{*}) are close, as indicated by similar values of $\kappa \approx 27$ N/m. At the same time, the intralayer Ti–Ti interactions (Ti₀–Ti₂, Ti₀–Ti₈) are slightly stronger ($\kappa \approx 39$ –42 N/m) than the interlayer Ti–Ti interactions (Ti₀–Ti₆^{*}, Ti₀–Ti₉^{*}) with $\kappa \approx 26$ –35 N/m. Among all Se–Se atom pairs, the interaction between the nearest selenium atoms is the weakest ($\kappa_{\text{Se}_0-\text{Se}_2} \approx 28$ N/m), while the interaction between two selenium atoms located at opposite sides of the same layer is more than twice stronger ($\kappa_{\text{Se}_0-\text{Se}_4} \approx 66$ N/m). At the same time, the interaction between selenium atoms in adjacent layers across the vdW gap is rather strong with $\kappa_{\text{Se}_0-\text{Se}_3^*} \approx$

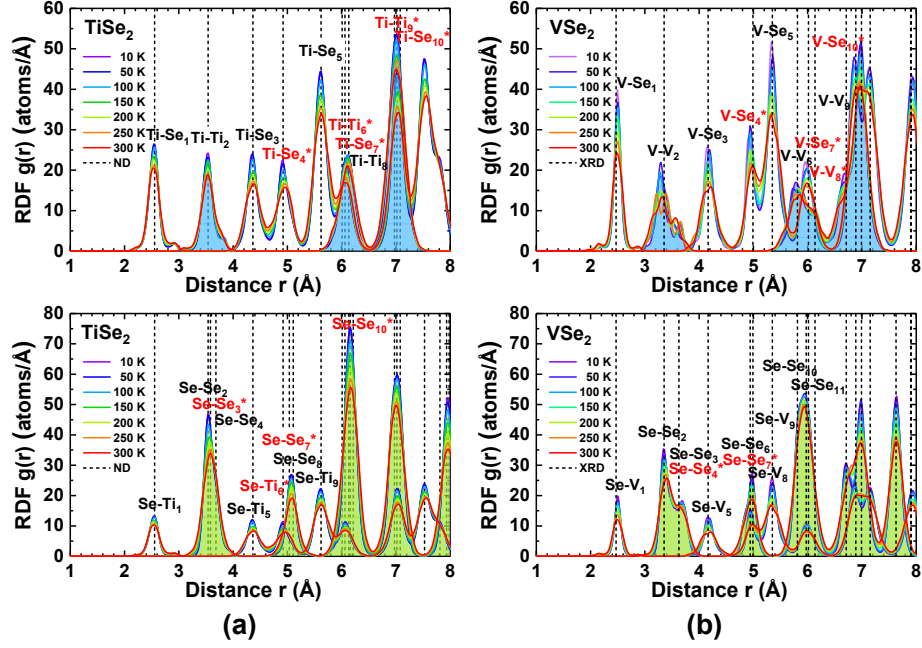


Figure 6: Partial radial distribution functions (RDFs) around Ti and Se atoms in TiSe_2 (a) and V and Se atoms in VSe_2 (b) as a function of temperature. Open and filled peaks correspond to different partial RDFs. Vertical dashed lines are used to show the interatomic distances estimated from diffraction at room temperature [13, 14].

52 N/m. To conclude, the temperature dependencies of the MSRD factors in TiSe_2 do not show any unusual behaviour.

In spite of the similarities in the crystallographic structures of TiSe_2 and VSe_2 , the EXAFS spectra $\chi(k)k^2$ of VSe_2 have stronger amplitude and, as a result, more intense peaks in FTs (Fig. 3). The V K-edge EXAFS spectra of VSe_2 , their FTs and WT (Fig. 3(b)) are dominated by a contribution from the first coordination shell composed of six selenium atoms. Contributions from distant coordination shells are also present up to 7 Å but they are relatively weaker. This fact contrasts with the Se K-edge EXAFS spectra

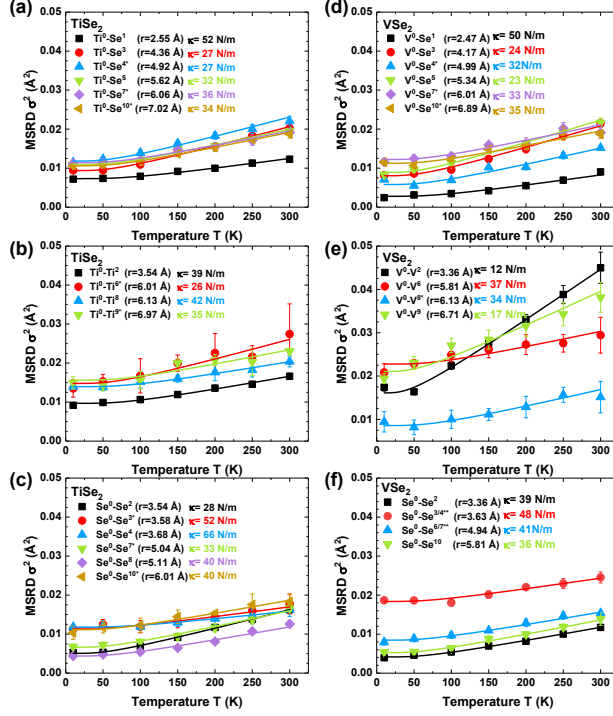


Figure 7: Temperature dependence of the mean-square relative displacement (MSRD) factors σ^2 for Ti(V)–Se (a,d), Ti–Ti(V–V) (b,e), and Se–Se (c,f) atom pairs in TiSe_2 and VSe_2 . Solid lines correspond to the fits using the correlated Einstein model [46]. Calculated values of the effective force constants κ are also given.

where strong contributions from distant shells are observed in FTs up to 5 Å but appreciable signals are visible even at longer distances.

The results of the RMC/EA fits for VSe_2 are in good agreement with the experimental data at both V and Se K-edges (Fig. 4(b)). The partial RDFs for V–V, V–Se(Se–V), and Se–Se atom pairs are shown in Fig. 6(b), whereas the characteristic Einstein frequencies ω_E and the effective force constants κ for all coordination shells are reported in Table 2.

The RDFs obtained for both TMDs (Fig. 6) exhibit an expected temper-

ature dependence: they become broadened at higher temperatures. At the same time, no peak splitting is observed for the nearest shells at low temperatures in the CDW state. This suggests that the expected lattice distortions due to atom displacements [19, 10] are relatively small compared to thermal disorder effects. Nevertheless, upon comparing the partial RDFs for TiSe_2 and VSe_2 , a notable difference in the temperature dependence between Ti-Ti and V-V RDFs is observed within the range of the second coordination shell. Upon increasing temperature, the $\text{V}_0\text{-V}_2$ RDF peak at 3.36 Å in VSe_2 becomes significantly more broadened compared to the $\text{Ti}_0\text{-Ti}_2$ RDF peak at 3.54 Å in TiSe_2 . This difference is well reflected by the temperature dependence of respective MSRDs in Fig. 7(b) and (e). Indeed, the MSRD for the $\text{V}_0\text{-V}_2$ atom pair exhibits a rapid increase above 50 K, displaying anomalous behavior compared to MSRDs for other atom pairs. Note that the value of the force constant $\kappa_{\text{V}_0\text{-V}_2} \approx 12$ N/m is the smallest one (Table 2).

There is also a prominent difference in the splitting of the peak around 3.6 Å in the Se-Se RDFs between the two TMDs. A single peak at 3.55 Å is present in TiSe_2 , whereas a double peak at about 3.35 Å and 3.68 Å exists in VSe_2 . This difference is due to that the $\text{Se}_0\text{-Se}_2$ distance in VSe_2 is significantly shorter by 0.18 Å than in TiSe_2 (see Tables 1 and 2).

4. Conclusions

Highly crystalline bulk 1T- TiSe_2 and 1T- VSe_2 were synthesized using the chemical vapor transport technique, and their local atomic structure and lattice dynamics were studied by temperature-dependent (10-300 K) X-ray absorption spectroscopy at the Ti, V, and Se K-edges.

The extended X-ray absorption fine structure (EXAFS) spectra were analysed using the reverse Monte Carlo (RMC) simulations [26, 27] by a simultaneous fitting of metal (Ti/V) and selenium K-edge spectra (Fig. 4). Such an approach allowed us to reconstruct the local environment in both selenides in terms of the partial radial distribution functions (Fig. 6) and to gain reliable information on disorder effects in the nearest and distant coordination shells. The mean-square relative displacements σ^2 for atom pairs located in the first ten coordination shells were determined, and their temperature dependencies were approximated using the correlated Einstein model. As a result, the characteristic Einstein frequencies ω_E and the effective force constants κ were obtained (Tables 1 and 2) for all atomic pairs and used in the assessment of the strengths of intralayer and interlayer interactions.

The partial RDFs obtained for both TMDs (Fig. 6) demonstrate the expected temperature dependence and do not show any evidence of the peak splitting for the nearest shells at low temperatures due to the CDW state. This indicates that CDW-induced atom displacements [19, 10] are relatively small compared to thermal disorder effects. Nevertheless, a comparison of the RDFs for Ti–Ti and V–V atom pairs located within the same layer shows their different temperature variations. In particular, the thermal disorder significantly affects the V_0 – V_2 atom pair as can be seen from the temperature dependence of its MSD $\sigma_{V_0-V_2}^2$ (Fig. 7) and small value of the force constant $\kappa_{V_0-V_2} \approx 12$ N/m (Table 2).

Thus, the RMC/EA-EXAFS technique shows great potential for studying the local atomic structure and disorder effects in TMDs and other layered materials.

CRedit authorship contribution statement

Inga Pudza: Investigation, Visualization, Writing – original draft, Writing – review & editing. **Boris Polyakov:** Investigation, Writing – original draft, Writing – review & editing. **Kaspars Pudzs:** Investigation. **Edmund Welter:** Resources. **Alexei Kuzmin:** Conceptualization, Investigation, Methodology, Writing – original draft, Writing – review & editing.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Data availability statement

Data will be made available on request.

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Table 1: Values of coordination numbers (N) and Ti–Se, Ti–Ti, and Se–Se interatomic distances r for the first ten coordination shells of Ti and Se calculated from the crystallographic structure of TiSe_2 [13]. Interlayer distances are marked with “*”. The values of characteristic Einstein frequencies (ω_E) and the effective force constants (κ) obtained from the RMC/EA analysis are also given for each atom pair.

Atom pair	N	Distance r (Å)	ω_E (THz)	κ (N/m)
Ti ₀ –Se ₁	6	2.55	32.3±0.4	51.61±0.01
Ti ₀ –Ti ₂	6	3.54	31.3±0.6	39.02±0.02
Ti ₀ –Se ₃	6	4.37	23.3±0.4	26.92±0.01
Ti ₀ –Se ₄ *	6	4.92	23.3±0.6	26.94±0.02
Ti ₀ –Se ₅	12	5.62	25.6±0.9	32.43±0.04
Ti ₀ –Ti ₆ *	2	6.01	25.6±1.7	25.99±0.12
Ti ₀ –Se ₇ *	6	6.06	26.8±1.2	35.65±0.07
Ti ₀ –Ti ₈	6	6.13	32.4±0.6	41.69±0.01
Ti ₀ –Ti ₉ *	12	6.97	29.8±1.9	35.41±0.14
Ti ₀ –Se ₁₀ *	12	7.02	26.2±0.2	33.96±0.01
Se ₀ –Ti ₁	3	2.55	32.3±0.4	51.61±0.01
Se ₀ –Se ₂	6	3.54	20.7±0.2	28.12±0.01
Se ₀ –Se ₃ *	3	3.58	28.1±1.2	51.95±0.10
Se ₀ –Se ₄	3	3.68	31.7±1.3	65.81±0.12
Se ₀ –Ti ₅	3	4.37	23.3±0.4	26.92±0.01
Se ₀ –Ti ₆ *	3	4.92	23.3±0.6	26.94±0.02
Se ₀ –Se ₇ *	3	5.04	22.4±0.5	32.89±0.02
Se ₀ –Se ₈	3	5.11	24.7±0.6	39.96±0.03
Se ₀ –Ti ₉	6	5.62	25.6±0.9	32.43±0.04
Se ₀ –Se ₁₀ *	2	6.01	24.6±0.8	39.53±0.04

Table 2: Values of coordination numbers (N) and V–Se, V–V, and Se–Se interatomic distances r for the first ten coordination shells of V and Se calculated from the crystallographic structure of VSe₂ [14]. Interlayer distances are marked with “*”. Coordination shells composed of both intralayer and interlayer atomic pairs are marked with “**”. The values of characteristic Einstein frequencies (ω_E) and the effective force constants (κ) obtained from the RMC/EA analysis are also reported for each atom pair.

Atom pair	N	Distance r (Å)	ω_E (THz)	κ (N/m)
V ₀ –Se ₁	6	2.47	31.1±0.8	49.69±0.04
V ₀ –V ₂	6	3.36	16.6±0.4	11.67±0.01
V ₀ –Se ₃	6	4.17	21.7±0.4	24.25±0.01
V ₀ –Se ₄ [*]	6	4.99	24.9±1.0	31.84±0.06
V ₀ –Se ₅	12	5.35	21.3±0.6	23.23±0.02
V ₀ –V ₆	6	5.81	29.5±2.3	36.81±0.22
V ₀ –Se ₇ [*]	6	6.01	25.4±1.6	33.13±0.13
V ₀ –V ₈ [*]	2	6.13	28.4±1.3	34.03±0.07
V ₀ –V ₉	6	6.71	20.1±1.2	17.16±0.06
V ₀ –Se ₁₀ [*]	12	6.89	26.2±1.1	35.22±0.06
Se ₀ –V ₁	3	2.47	31.1±0.8	49.69±0.04
Se ₀ –Se ₂	6	3.36	24.5±0.4	39.32±0.01
Se ₀ –Se _{3/4} ^{**}	3+3	3.63	27.1±1.4	48.18±0.12
Se ₀ –V ₅	3	4.17	21.7±0.4	24.25±0.01
Se ₀ –Se _{6/7} ^{**}	3+3	4.94	25.1±0.6	41.19±0.02
Se ₀ –V ₈	3	4.99	24.9±1.0	31.84±0.06
Se ₀ –V ₉	6	5.35	21.3±0.6	23.23±0.02
Se ₀ –Se ₁₀	6	5.81	23.5±0.4	36.16±0.01