The origin of anisotropy and high density of states in the electronic structure of \( \text{Cr}_2\text{GeC} \) by means of polarized soft x-ray spectroscopy and \textit{ab initio} calculations

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Abstract

The anisotropy in the electronic structure of the inherently nanolaminated ternary phase \( \text{Cr}_2\text{GeC} \) is investigated by bulk-sensitive and element selective soft x-ray absorption/emission spectroscopy. The angle-resolved absorption/emission measurements reveal differences between the in-plane and out-of-plane bonding at the (0001) interfaces of \( \text{Cr}_2\text{GeC} \). The \( \text{Cr} L_{2,3}, \text{C} K \), and \( \text{Ge} M_{1,2,3} \) emission spectra are interpreted with first-principles density-functional theory (DFT) including core-to-valence dipole transition matrix elements. For the \( \text{Ge} 4s \) states, the x-ray emission measurements reveal two orders of magnitude higher intensity at the Fermi level than DFT within the General Gradient Approximation (GGA) predicts. We provide direct evidence of anisotropy in the electronic structure and the orbital occupation that should affect the thermal expansion coefficient and transport properties. As shown in this work, hybridization and redistribution of intensity from the shallow \( 3d \) core levels to the \( 4s \) valence band explain the large \( \text{Ge} \) density of states at the Fermi level.

Keywords: MAX-phases, nanolaminates, soft x-ray spectroscopy, electronic structure, density functional theory

(Some figures may appear in colour only in the online journal)

I. Introduction

The group of ternary nanolaminated carbides and nitrides known as \( M_{n+4}AX_n \)-phases is the subject of intense research \cite{1, 2}. Three related crystal structures are classified by stoichiometry as 211 \((n = 1)\), 312 \((n = 2)\) and 413 \((n = 3)\) phases, where the letter \( M \) is an early transition metal, \( A \) is an element in the groups III–V and \( X \) is either carbon or nitrogen. The \( M_{n+4}AX_n \)-phases have a technologically important combination of properties that are metallic and ceramic \cite{3}. These properties are related to the crystal structure, the choice of the three constituent elements, as well as the strength of the chemical bonding between the nanolaminated layers.
For the 211 type of crystal structure, there are more than 50 different ternary carbides and nitrides, where Cr$_2$GeC was recognized by Jeitschko et al. already in the 1960s [4]. This multifunctional metallic and ceramic compound is particularly interesting as it exhibits a number of peculiar properties and is relatively little studied. First, Cr$_2$GeC has the highest thermal expansion coefficient $\alpha$, among all the presently known $M_{n+1}AX_n$-phases [5-7], which is largest in the M–A bonds and smallest in the M–X bonds as indicated by thermal motions of the atoms in neutron diffraction experiments [8]. There is also a discrepancy between the theoretical and experimental $\alpha$-values of 30% [9]. This could be related to a change in the orbital occupation as the temperature is raised that allows the system to show large volume variation with temperature.

Second, the magnetic ordering and electron correlations affect other properties such as thermal expansion and bulk modulus of Cr-based MAX-phases, even though they are only weakly magnetic [10-12]. Although pure Cr$_2$GeC is non-magnetic macroscopically at room temperature, the individual Cr atoms in Cr$_2$GeC have a magnetic order that could depend on the temperature [13, 14]. However, due to Cr 3$d$ electron correlation effects, the electronic origin of the unique properties is not well understood or conclusive. Transport properties cannot be interpreted in terms of ground state electronic structure calculations of Cr-based $M_{n+1}AX_n$-phase materials since the electron-phonon coupling is expected to be strong and most likely anisotropic.

Third, the calculated density of states (DOS) at $E_F$ (7.7 states/eV/unit cell for $U = 0$, and 7.9 states/eV/unit cell with $U_{eff} = 2.09$ eV ($t_{2g}$) and 2.33 eV ($e_g$) for a chosen GGA+U functional) [12], dominated by the Cr 3$d$ states, is much lower than the experimentally reported value (21 states/eV/unit cell [15] and 22 states/eV/unit cell [16]), and there is a large electron-phonon coupling [8]. For Cr$_2$GeC, the reported value of the DOS at $E_F$ is by far the highest measured among the $M_{n+1}AX_n$-phases (Cr$_2$AIC has the second highest) [15].

In this paper, we investigate the anisotropy and the orbital occupation in the electronic structure of single-crystal Cr$_2$GeC (0001) thin films. By applying bulk-sensitive and element-specific soft x-ray absorption spectroscopy (XAS), x-ray emission spectroscopy (XES) and resonant inelastic x-ray scattering (RIXS), we characterize the unoccupied and occupied bands of the containing elements, respectively [17-19]. This enables exploring the orbital occupation of the electrons buried several hundred nanometers below the surface. The energy of the x-rays is tuned to the specific core levels of Cr, Ge, and C and the incidence angle of the x-rays is changed from grazing to near normal relative to the laminate plane. This allows probing the occupation of the occupied electronic orbitals across $(d_{x^2−y^2}, p_x)$ and in $(d_{x^2−y^2}, d_{xy}, p_y)$ the laminate plane as illustrated in figure 1. We provide direct evidence of anisotropy in the electronic structure supported by ab initio band structure calculations. We further observe that the intensity ratio between the Ge 4$s$ and 3$p$ states in the valence band obtained from DFT-GGA is highly underestimated in comparison to the one measured via XES spectra.

II. Experimental and computational details

II.A. Cr$_2$GeC (0001) thin film synthesis

Cr$_2$GeC (0001) thin films were deposited by dc magnetron sputtering (base pressure $\sim 5 \times 10^{-10}$ Torr) from elemental targets of Cr, C, and Ge in an argon discharge pressure of 4 mTorr on MgO (1 1 1) substrates. Details on the synthesis process are given in [20].

II.B. X-ray emission and absorption measurements

The XAS and XES measurements were performed at room temperature (300 K) at the undulator beamline IS11-3 at MAX II (MAX-IV Laboratory, Lund, Sweden), comprising a 49-pole undulator, and a modified SX-700 plane grating monochromator [21, 22]. The measurements were made at a base pressure lower than 6.7$\times 10^{-9}$Pa. The XAS measurements were performed in total fluorescence yield (TFY) mode with 0.14 eV and 0.05 eV energy resolutions at the Cr 2$p$ and C 1$s$ absorption edges, respectively. The XAS-TFY spectra were measured at 15° (along the c-axis, near perpendicular to the basal ab plane) and 90° (normal, parallel to the basal ab-plane) incidence angles. The XAS spectra were normalized to the step before, and after the absorption edges and the self-absorption effects were estimated with the program XANDA [23].

The XES spectra were measured at 15° (in the basal ab plane) and 75° (perpendicular to the basal ab-plane) incidence angles. For comparison of the spectral profiles, the measured XES data were normalized to unity and plotted on a common photon energy scale (top) and relative to the $E_F$ (bottom). The XES spectra were recorded with 0.7, 0.2, 0.2 and 0.3 eV beamline monochromator energy resolutions at the Cr L, C K, Ge...
$M_1$, and Ge $M_{2,3}$ edges and 0.68, 0.19, 0.19 and 0.30 eV spectrometer resolutions, respectively. Self-absorption is known to affect the shape of the XES spectra on the high-energy flank of the main peak at the overlap with the XAS spectra. The observed XES intensity $I$ can be written as [24]:

$$I = I_0 \times \frac{1}{1 + \frac{\mu_{\text{out}}}{\mu_{\text{in}}} \times \frac{\sin \alpha}{\cos \beta}} \quad (1)$$

where $I_0$ is the unperturbed intensity, $\mu_{\text{in}}$ and $\mu_{\text{out}}$ are the absorption coefficients for the incident and outgoing radiation, $\alpha$ and $\beta$ are the incident and exit angles of the photon beam relative to the sample surface and surface normal, respectively. The estimated self-absorption effect is largest (5–10% at $E_\text{F}$ depending on the excitation energy) when XAS is measured at grazing incidence (unoccupied out-of-plane states) and for XES at near normal incidence (occupied out-of-plane states). On the contrary, the effect is almost negligible (0–3% at $E_\text{F}$ depending on the excitation energy) for XAS measured at normal incidence (unoccupied in-plane states) and XES measured at grazing incidence (occupied in-plane states).

II.C. First-principles calculations

All the ab initio calculations performed in this work are based on the well-known Density Functional Theory (DFT) [25]. For the computations of the XES spectra, we used the so-called final-state rule [26], where no core-hole was created at the photo-excited atom. Particularly, emission spectra were obtained by means of the single-particle transition model using the Full Potential Linearized Augmented Plane Wave (FPLAPW) method [27]. Exchange and correlation effects were taken into account through the Generalized Gradient Approximation (GGA) as parameterized by Perdew, Burke and Ernzerhof (PBE) [28] using a matrix-size of $R_{\text{MT}}^*K_{\text{max}} = 10$, where $K_{\text{max}}$ is the plane wave cut-off and $R_{\text{MT}}$ the smallest of all atomic sphere radii (1.55 for carbon atoms). The charge density and potentials were expanded up to $l = 12$ inside the atomic spheres, and the total energy was converged with respect to the Brillouin zone (BZ) integration. The electric-dipole approximation was here used to calculate XES spectra, implying that only the electronic transitions arising from core levels with orbital angular momentum $l$ to the $l \pm 1$ components of the valence bands were studied. For core-hole lifetime and broadening parameters we used the available experimental numbers. In the fitting procedure to the nonresonant $L_{2,3}$ XES spectra, we used the experimental values for the $L_3/L_2$ branching ratio of 10:1 in the basal plane and 15:1 along the c-axis. For the XAS spectra, we used the experimental values for $2p_{1/2}/2p_{3/2}$ branching ratio (1.1:1) and for the $L_{2,3}$ peak splitting (8.0 eV), which is somewhat larger than our calculated ab initio spin–orbit splitting of 7.6 eV. Absorption spectra were calculated within the same theoretical scheme used for XES, but with the addition of core-hole effects [29]. Specifically, we generated such a self-consistent-field potential using a $2 \times 2 \times 1$ hexagonal supercell of 32 atoms containing one core-hole on the investigated element. Calculated nonresonant $L_{2,3}$ XES and $2p$ XAS spectra are shown at the bottom of figure 2, while $C K$ XES and $C 1s$ XAS in figure 3 and Ge $M_1$, $M_{2,3}$ XES in figure 4.

II.D. Calculation of phonon density of states

The supercell approach was used to compute the phonon densities of states (PhDOS) within the framework of DFT and Density-Functional Perturbation Theory (DFPT) [30]. Specifically, the q-ESPRESSO software package [31] was employed to calculate real space force constants on a $2 \times 2 \times 1$ supercell system. Phonon frequencies were successfully obtained from the force constants by means of the QHA code [32]. Vanderbilt ultrasoft pseudo-potentials [33] were used for all the atomic species, while exchange-correlation effects were treated with the PBE generalized gradient approximation. Integration over the BZ was performed by using the special k-points technique [34] and a Gaussian broadening of 0.002 Ry. Electronic wave functions were plane-wave expanded with a cut-off of 70 Ry and a kinetic energy up to 700 Ry.
III. Results

III.A. Cr 2p3/2, 1/2 x-ray absorption and L2,3 emission

Figure 2 (top-right) shows XAS-TFY spectra measured at the Cr 2p absorption edge and normalized at 600 eV. The intensity is related to the unoccupied Cr states of Cr2GeC that are projected via the Cr 2p dipole selection rules. The XAS spectra are shown as measured with the polarization vector both in the ab basal-plane and along the c-axis of Cr2GeC. The main XAS peak structures correspond to the 2p3/2 and 2p1/2 core levels. The observed sub-peak splittings are most salient features for the in-plane orbitals, while the peaks across the basal plane have lower intensity. The structures agree with the DFT calculations shown at the bottom-right part of figure 2.

The branching ratios observed experimentally are different from the statistical 2:1 ratio since they are affected by both exchange as well as mixed terms between the 2p core states and the valence states [35]. At the E_F, the intensity of the unoccupied states is 35% higher in the ab basal laminate plane than along the c-axis.

The central part of figure 2 shows Cr L2,3 XES and RIXS spectra. These spectra are related to the occupied Cr 3d and 4s valence bands through the Cr 3d4s → 2p dipole selection rule of Cr2GeC. The spectra are shown both in the ab laminate plane and along the c-axis and were excited at two different energies at the 2p3/2 absorption maxima (indicated by vertical arrows in the XAS spectra), and also nonresonant at 620 eV. The main L3 peak (between −2 and −5 eV below E_F) is due to the Cr 3d–C 2p orbital overlap. As observed, the in-plane spectra have higher intensities than the out-of-plane spectra. The intensity at the E_F is 40% larger in the ab basal laminate plane compared to the intensity along the c-axis. The calculated spectra (figure 2, bottom) confirms the greater spectral intensity at the E_F for the in-plane states (30%) than for the out-of-plane states, as shown in the inset.

III.B. C 1s x-ray absorption and C K emission

Figure 3 (top right) shows measured C 1s XAS-TFY spectra of Cr2GeC. Left top-to-bottom: resonant spectra excited at 285.0 (indicated by the upward arrow) and nonresonant C K XES spectra at 310.0 eV. All spectra are plotted on a photon energy scale (top) and a relative energy scale (bottom) with respect to the top of the valence band.

Figure 3. Top-right: C 1s XAS-TFY spectra of Cr2GeC. Left top-to-bottom: resonant spectra excited at 285.0 (indicated by the upward arrow) and nonresonant C K XES spectra at 310.0 eV. All spectra are plotted on a photon energy scale (top) and a relative energy scale (bottom) with respect to the top of the valence band.

Figure 4. Ge M1 XES spectra of Cr2GeC excited at 220 eV. Bottom panel: Ge M2,3 XES spectra excited at 165 eV. A common energy scale with respect to the top of the valence band edge (vertical dotted line) is indicated between the top and bottom panels. The dashed curves below the experimental data are calculated spectra for comparison.
the difference between the XAS spectra measured across and in the \textit{ab} basal laminate plane is relatively large. At the \(E_p\), the data exhibit 67\% more empty hole-like states in the \textit{ab} basal laminate plane than along the \textit{c}-axis. This difference is consistent with our calculated spectra (74\%) at the bottom-right in figure 4. Differences between in- and out-of-plane also occur at higher energies above \(E_p\). The shoulder at 0.5 eV and the peak at 6.0 eV that occur in-plane are more pronounced in the experiment than in the calculation. This may be due to carbon contamination on the sample or on the beam line mirrors.

The C K RIXS and nonresonant XES spectra of Cr\(_2\)GeC were excited at 285.0 and 310.0 eV photon energies, respectively. Calculated nonresonant spectra are shown at the bottom of figure 3. For the in-plane excitation, a main peak and a shoulder with \(\sigma\) orbital character are clearly observed at \(-4.0\) eV and \(-6.0\) eV below \(E_p\). This intensity corresponds to the C 2\(p_{\sigma} \rightarrow 1s\) dipole transitions of Cr\(_2\)GeC. The spectra probed along the \textit{c}-axis instead exhibit a single peak at \(-4.3\) eV below \(E_p\) and are due to 2\(p_{\pi}\) states with \(\pi\) orbital character. The 2\(p\) peak structures hybridize mainly with the upper part of the Cr 3\(d\) bonding orbitals, but also with the Ge 4\(p\), 4\(s\) orbitals and the shallow 3\(d\) core levels. Contrary to the occupied Cr 3\(d\) states, the intensity at the \(E_F\) of the occupied C 2\(p\) states is similar in the \textit{ab} basal laminate plane and along the \textit{c}-axis.

\section{III.C. Ge \(M_2\) and \(M_{2,3}\) x-ray emission}

Figure 4 (top panel) shows experimental Ge \(M_1\), XES spectra (4\(p \rightarrow 3\)s transitions) of Cr\(_2\)GeC measured nonresonantly at 220 eV photon energy probing the Ge 4\(p\) states across and in the laminate \textit{ab}-plane. The spectra are plotted on a photon energy scale and relative to the \(E_F\). Calculated Ge \(M_1\) spectra are shown below the experimental spectra. Experimentally, the anisotropy of the Ge 4\(p\) states is found to be rather large with more states in the \textit{ab}-plane than along the \textit{c}-axis.

The in-plane 4\(p_{\sigma}\) states with three \(\sigma\) orbitals are spread out between 0 and \(-10\) eV below \(E_F\). On the contrary, the 4\(p_{\pi}\) states with a single bonding \(\pi\) orbital are more localized at \(-3\) eV below \(E_F\). The experimental out-of-plane Ge 4\(p_{\pi} \rightarrow \pi^*\) spectrum is broader than the sharper calculated spectrum, possibly due to electron-phonon interactions with Ge displacements. At the \(E_F\), the Ge 4\(p\) intensity is \(\sim 10\%\) higher in the basal \textit{ab}-plane than along the \textit{c}-axis, both experimentally and theoretically.

The bottom panel in figure 4 shows experimental Ge \(M_{2,3}\) XES spectra (3\(d \rightarrow 3\)p and 4\(s \rightarrow 3\)p transitions) of Cr\(_2\)GeC aligned to the \(E_F\) at the 3\(p_{1/2}\) core level. Here, we find that the isotropic 4\(s\)-states of the Cr\(_2\)GeC valence band have an exceptionally high intensity in a broad range from \(-20\) eV up to the \(E_F\) in comparison to other Ge-containing Ti\(_3\)GeC\(_2\) [37] and V\(_2\)GeC [38] compounds. Contrary to the experimental observations on Ge in Cr\(_2\)GeC, the calculated 4\(s\) valence band is more than 10 times weaker with most intensity in the range between \(-8\) to \(-12\) eV below \(E_F\). Note that at the \(E_F\), the intensity is two orders of magnitude higher in the experiment than in the calculated spectra.

The shallow Ge 3\(d\) core levels at the bottom of the valence band between \(-30\) and \(-35\) eV also participate in the Cr–Ge bonding in Cr\(_2\)GeC. The measured 3\(d_{5/2, 3/2}\) spin–orbit splitting of 3.6 \(\pm 0.1\) eV is consistent with the difference between the 3\(p_{3/2, 1/2}\) and 3\(d_{5/2, 3/2}\) spin–orbit splittings observed in XPS (4.1–0.5 eV) and in Ti\(_3\)GeC\(_2\) [37] and V\(_2\)GeC [38]. Theoretically, the spin–orbit splitting is larger (4.4 eV) and the states are also about 4.5 eV closer to the \(E_F\) than in the experiment. The enhanced Ge 4\(s\) intensity observed experimentally may be attributed to electron-correlation, many-body effects or phonon vibrations, but more extensively to hybridization and charge-transfer effects, as in the case of Ga in GaN [39].

Another interesting observation is that the intensity of the experimental Ge 3\(d\) states is higher in the basal \textit{ab}-plane than along the \textit{c}-axis. This is also the case for the calculated Ge \(M_{2,3}\) spectra (dashed curves) with the statistical 3\(p_{3/2}\) core-level ratio of 2:1. Note that the \(M_1/M_2\) branching ratio is lower (0.8:1) for measurements with polarization along the \textit{c}-axis than in the basal \textit{ab}-plane (1:1). The trend in XES branching ratios in the transition-metal compounds is a signature of the degree of ionicity in the systems [40], due to the additional Coster-Kronig process [24, 41]. The lower 3\(p_{3/2}/3p_{1/2}\) peak ratio along the \textit{c}-axis is thus an indication of higher ionicity (resistivity) than in the basal \textit{ab}-plane. The valence-to-core matrix elements are thus found to play an important role to the spectral shape of the Ge \(M_{2,3}\) XES spectra. The redistribution of intensity from the localized 3\(d\) states to the valence band may explain the large discrepancy between experiment and calculations for the DOS observed at \(E_F\).

Figure 5 shows calculated Phonon Density of States (PhDOS) for the Cr, Ge and C atoms in Cr\(_2\)GeC. The Ge atoms move preferentially within the \textit{ab}-basal plane, while both Cr and C have more preference for the \textit{c}-axis direction.

Figure 5. Calculated phonon frequency spectra of the Cr, Ge and C atoms in Cr\(_2\)GeC. The Ge atoms move preferentially within the \textit{ab}-basal plane, while both Cr and C have more preference for the \textit{c}-axis direction.
0.097 Å and a z-displacement of 0.114 Å from the phonon calculations. For Ge, the x, y-displacement is 0.128 Å and the z-displacement 0.081 Å, while for the C-atoms, the x, y-displacement is 0.085 Å and the z-displacement 0.109 Å.

Rapidly moving atoms is known to change the electronic DOS at $E_F$ [44]. However, the relatively small magnitude of these displacements alone cannot account for the large difference in DOS between experiment and theory for Cr$_2$GeC. The hybridization and redistribution of charge between the shallow 3d core-level and the 4s valence band in Ge is also a necessary prerequisite for exceptionally high DOS at $E_F$ compared to other Ge-containing compounds. However, for the Cr-atoms, the influence of phonons to the enhanced DOS at $E_F$ is not obvious, as the motions along the x, y and z directions are mixed.

IV. Discussion

The present results indicate that the peculiar transport properties in Cr$_2$GeC are more complex due to the Ge-element in comparison to other $M_{n+1}AX_n$-phase systems where they are governed by the choice of 3d transition element. The observation of the dominating contribution of the 4s states in the Ge $M_{2,3}$ XES in figure 4 explains the large difference in the DOS at $E_F$ between experiment (22 states/eV/cell) [16] and DFT calculations (7.7 states/eV/cell) [12].

In the DFT calculations, the Ge 3d states are overestimated and not enough intensity is redistributed to the 4s valence band states. The same type of deficiency in DFT calculations is known also for Ge in Ti$_3$GeC$_2$ [37] and V$_2$GeC [38] as well as for Ga in GaN [39]. We clearly observe that the calculated 4s states at the bottom of figure 4 are too low in intensity, in particular at the $E_F$. As this problem is a shortcoming in 3d/4s intensity ratio, we anticipate that this phenomenon is attributed to 3d–4s hybridization effects in combination with mixing of other states that change the relative population of the 3d and 4s states. On the other side, obtaining the DOS at $E_F$ by measuring the specific heat [16] may be sensitive to Cr$_2$Ge$_3$C and Cr$_2$C$_2$ impurities. It should also be emphasized that the samples used in [16] contained substantial amounts of Cr$_2$O$_3$ [45] and that the fitting of the specific heat was not satisfactory above 180 K [16]. This may lead to an overestimated value of the actual DOS at $E_F$. As our Cr$_2$GeC sample is phase pure with no Cr$_2$Ge$_3$C impurities [20], we only consider the very small fraction of Cr$_2$C$_2$ phase that could contribute to an increased DOS at $E_F$ [46]. However, if there is an impurity phase that causes the high DOS at $E_F$, this impurity should contain Ge and that excludes Cr$_2$C$_2$.

The Cr and Ge atoms are also known to have a correlated motion where the Cr–Ge bond length is kept constant [8]. The rattling Ge atoms thus drive the Cr atoms in the structure. The Cr 3d DOS have an intensity at $E_F$ that is sensitive to the particular in-plane and out-of-plane phonon modes of the Cr atoms. However, the change in DOS at the $E_F$ is not only due to the movement of the Cr atoms, but also to the moving Ge and C atoms where the Ge atoms move preferentially within the $ab$-basal plane, while both Cr and C have more preference for the c-axis direction.

The lowest-frequency vibrations are associated with the Ge atoms in Cr$_2$GeC, similarly to the case of Al in Cr$_2$AlC [36] and Ti$_3$AC$_2$ (A = Si, Al, and Ge), where the lowest-energy vibrations are from the A element [37, 44, 47]. This finding is consistent with the fact that the Cr–C bonding type is the stiffest. This is also in agreement with what was recently found in Cr$_2$AlC [36]. However, the Ge and Cr vibrations in Cr$_2$GeC appear to be more decoupled than in the case of Al and Cr in Cr$_2$AlC [36]. This type of tangled phonon scenario in MAX-phases will likely influence the core-level spectra at finite temperature values as well.

The electronic structure anisotropy of Cr$_2$AlC has been studied previously by electron energy loss spectroscopy (EELS) at the C-K edge and by polarized XAS at the Al-K edge [36]. A marked anisotropy at the Al-site was observed, in addition to a much smaller anisotropic character of the C-site, essentially attributed to the octahedral environment of the C-sites in the $M_{n+1}AX_n$-phase structure, which is much more isotropic than the trigonal prismatic site occupied by Al-atoms. The relatively small anisotropy of the C-K edge was exacerbated by the average of crystallographic orientations probed in EELS, as opposed to polarized x-ray beams. Here, for Cr$_2$GeC, the anisotropy of the C-K edge is present in the XAS and XES data presented in figure 3, although it may also be somewhat influenced by the presence of amorphous C, resulting from contamination of the sample. The anisotropy is consistent with the case of other $M_{n+1}AX_n$—phase systems such as V$_2$GeC [38] and Ti$_3$SiC$_2$ [44]. Although the local symmetry at each atomic site is important, the distribution of the electronic states also obeys the whole k-space topology of the $M_{n+1}AX_n$-phase unit cell, that is not highly symmetric. This is clearly observed at, for instance, the c-axial elongation of the unit cell.

Our results clearly show that the anisotropy in the electronic structure and chemical bonding of Cr$_2$GeC is substantial for the Cr 3d, C 2p and Ge 3d states. The generally greater amount of both occupied and unoccupied Cr 3d$_{x^2}$, 3d$_{y^2}$ and C 2p$_{xy}$ electronic states in the basal $ab$-plane than Cr 3d$_{z^2}$ and C 2p$_z$ states along the c-axis should contribute to higher electrical conductivity in the basal $ab$-plane for a large range of temperatures [20]. However, since the C 2p states are in minority they should not affect the conductivity as much as the Cr 3d states.

The calculations indicate that the Ge 4p$_{x+y}$σ states of the basal $ab$-plane are spread out in the region 0 to −10 eV, while the more localized single out-of-plane 4p$_{x+y}$π states along the c-axis have a peak maximum around −3 eV below $E_F$. Contrary to the case of the phonon rattling of the Si 3d states in Ti$_3$SiC$_2$ [44], the phonon modes of the isotropic Ge 4s states in Cr$_2$GeC should not affect the anisotropy of the transport properties as much as the 3d states of the transition element. The differences in intensity of the Ge 4s band explain the large difference in the DOS at $E_F$ between experiment and theory.

The strong covalent Cr 3d−C 2p bonding in Cr$_2$GeC occurs in the hybridization region between −2 to −5 eV below $E_F$ and the shallow Ge 3d core level states are located at −30 eV below the Fermi level. Intercalation of Ge monolayers into CrC with relatively weak Cr 3d−Ge 4p bonds thus implies that
the Cr–C bonds are strengthened. This also affects the Cr–Cr bonds that are strengthened, which enhances the metallic character and the density of states at the Fermi level of Cr$_2$GeC. Contrary to the elastic and stiffness properties, the transport properties depend more clearly on the states at the $E_F$.

The fact that the number of electronic states at $E_F$ are larger experimentally than those computed with DFT, is due to the additional electronic states provided to the $E_F$ by the Ge 4s states, which exhibit high intensity in the Ge 4s XES. The Ge 4s and Ge 3p states hybridize more with the Cr 3d states than DFT predicts, transferring more electron density towards the Fermi level. This effect is clearly observed in the 3p/4s intensity branching ratio in figure 4. Experimentally, the intensity ratio between the 3p $\rightarrow$ 3d and the 4s $\rightarrow$ 3d peaks is much lower than the one computed with DFT, as a consequence of the charge transfer from the shallow Ge 3p core levels to the Ge 4s states. This hybridization effect clearly manifests the Ge 4 core levels to be lower than the one computed with DFT, as a consequence of the hybridization regions in Cr$_2$GeC. The combination of hybridization of the electronic structure, chemical bonding, and hybridization regions in Cr$_2$GeC. The combination of hybridization, charge-transfer redistribution, Cr 3d electron correlation effects and electron-phonon coupling give rise to the observed transport properties. The occupied electronic DOS in the basal ab-plane is larger than those along the c-axis for the Cr 3d and Ge 4p states. The Ge 4s and shallow Ge 3p core-level states exhibit stronger hybridization with the Cr 3d states than DFT predicts in the one-electron approximation, transferring extra electron density towards the $E_F$. Thus, as shown in this work, the large intensity difference between the experimentally observed Ge 4s valence band states and the 3d states essentially explain the large difference in the experimental and calculated density of states at the Fermi level.

V. Conclusions

Polarization-dependent experimental and theoretical results provide understanding of what electronic states affect the anisotropy of the electronic structure, chemical bonding, and hybridization regions in Cr$_2$GeC. The combination of hybridization, charge-transfer redistribution, Cr 3d electron correlation effects and electron-phonon coupling give rise to the observed transport properties. The occupied electronic DOS in the basal ab-plane is larger than those along the c-axis for the Cr 3d and Ge 4p states. The Ge 4s and shallow Ge 3p core-level states exhibit stronger hybridization with the Cr 3d states than DFT predicts in the one-electron approximation, transferring extra electron density towards the $E_F$. Thus, as shown in this work, the large intensity difference between the experimentally observed Ge 4s valence band states and the 3d states essentially explain the large difference in the experimental and calculated density of states at the Fermi level.

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