Experimental and DFT investigation of (Cr,Ti)₃AlC₂ MAX phases stability

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ABSTRACT
Using a synergistic combination of experimental and computational methods, we shed light on the unusual solubility of (Cr,Ti)₃AlC₂ MAX phase, showing that it may accommodate Cr only at very low concentrations (< 2 at%) or at the exact Cr/(Cr + Ti) ratio of 2/3, even when the ratio of reactants is far from this stoichiometry (1/2 ≤ Cr/(Cr + Ti) ≤ 5/6). In both phases, Cr exclusively occupies the 4f sites, bridging carbide layers with the Al layer. Despite this, the peculiar stability of (Cr₂/₃Ti₁/₃)₃AlC₂ is attributed to the formation of strong, spin-polarized Cr–C bonds, which result in volume reduction and a marked increase in c/a ratio.

IMPACT STATEMENT
Solubility of Cr and Ti in (Cr,Ti)₃AlC₂ was investigated using experimental and DFT techniques. It was also determined that (Cr₂/₃Ti₁/₃)₃AlC₂ owe its remarkable stability to the formation strong Cr–C bonds.

1. Introduction
Mₙ₊₁AXₙ compounds crystallizing with the P6₃/mmc space group (where M is an early transition metal, A is a group 13–16 element, X is carbon or nitrogen and n is an integer) constitute a material family called MAX phases. The MAX phases are known for having a particular combination of properties owing to the duality of their structure. Indeed, a MAX phase structure consists of the stacking of n ‘ceramic’ M-X planes interposed with a ‘metallic’ A layer. As a consequence, MAX phases exhibit a favorable combination of metallic properties (high thermal shock resistance, high thermal and electrical conductivities and good machinability) and ceramic properties (high decomposition/melting temperature and elastic stiffness) [1].

Over 70 ternary MAX phases have been synthesized and a fast-growing number of quaternary MAX phases [2,3], that is, containing either two M, two A or two X elements. The interest in quaternary MAX phases is multiple:

- Firstly, they can be used to tailor a certain property to get a value in-between or even better than that of the ternary end-members [4–9]. As an example, Cabioch et al. [9] showed that when substituting 25% of the Al atoms for Ge in Cr₂AlC, its thermal expansion becomes isotropic, presumably diminishing thermal stresses in the material.
- Secondly the addition of a fourth element may trigger new properties not observed in the ternary host system. For instance, incorporating Mn in Cr₂GeC [10–12] or Cr₂GaC [4,13–16] was found to render these ternary MAX phases ferromagnetic.
- Thirdly, it allows integrating elements that are not able to form a bulk MAX phase by themselves. In a recent publication, we showed that the bismuth containing...
Zr$_2$(Al$_{0.42}$Bi$_{0.58}$)C MAX phase can be synthesized despite no M$_{n+1}$Bi$_X$ MAX phase being known [17]. Similarly, the recently reported Zr$_2$(Sb$_{0.7}$Al$_{0.3}$)C represents the firstly reported antimony-based MAX phase [3].

- Fourthly, it is also interesting on a fundamental basis to help understand the reasons driving the stability (or non-stability) of MAX phases, as this matter is still open to debate [1,18,19].
- Lastly, it allows synthesis of MAX phases with a composition as close as possible to a non-stable ternary one. For example, it can be conservatively said that Cr$_3$AlC$_2$ cannot be formed as a bulk phase, possibly because Cr$_2$AlC is more stable [18,20,21]. However, replacing only one-third of the chromium atoms for titanium atoms leads to the remarkable MAX phase (Cr$_{2/3}$Ti$_{1/3}$)$_3$AlC$_2$ [22,23], which may be stabilized by the ordering of Cr and Ti [24] (Cr atoms exclusively on the 4f Wyckoff site, Ti exclusively on the 2a sites, as illustrated in Figure 1).

Continuing on this last example, we decided to investigate the stability of quaternary 312-MAX phases across the compositional range from Ti$_3$AlC$_2$ to Cr$_3$AlC$_2$. This was done through a combination of computational methods and experimental synthesis attempts. First the solubility of Cr in Ti$_3$AlC$_2$ is considered at increasing levels of Cr, and the relevant accommodation mechanism is revealed. Then the driving force for ordering in (Cr$_{2/3}$Ti$_{1/3}$)$_3$AlC$_2$ is investigated, together with its ability to accommodate non-stoichiometry. The source of the peculiar stability of (Cr$_{2/3}$Ti$_{1/3}$)$_3$AlC$_2$ is then discussed in detail.

2. Methodology

2.1. Experimental details

Full details of the experimental procedure employed for ceramic synthesis and characterization are found in our previous publication [23]. In brief, TiH$_2$, Cr, Al and graphite powders are employed. All syntheses stages are done under argon atmosphere (except for the introduction at room temperature of the blended powders in the furnace) to limit oxygen contamination. First the elemental powders are mixed by ball–milling (Nanjing University Instrument Plant) according to the ratio Ti + Cr/Al/C = 3/1.1/1. Then the powder mixes are poured in graphite crucibles and placed in a hot-press (FCT Systeme HP W/25/1, Rauenstein, Germany) used as a regular furnace (pressureless mode). The furnace is heated up to 1450°C at a ∼ 20°C min$^{-1}$ rate and held at the target temperature for 1 h. The furnace is allowed to cool to room temperature before opening it to ambient atmosphere. Sintering of the reacted powder was then performed by spark plasma sintering (FCT Systeme HP D/25/1 equipment) at 1220°C and 35 MPa. X-ray diffraction (XRD) characterization was performed with a Bruker D2 Phaser SSD160 (Karlsruhe, Germany) and the treatment of results was done with Fullprof Suite program and Xpert High Score Plus software (PANalytical, Almelo, the Netherlands) using the implemented ICDD (International Center for Diffraction Data) database. SEM (scanning electron microscopy) observations were realized using a JEOL JSM-6400 (Tokyo, Japan), operated at 20 keV. The same microscope was used for EDX (energy dispersive X-ray spectrometry) analyses using an INCA ultra-thin polymer window detector (Oxford Instruments, Oxford, UK). The samples for SEM/EDX were prepared by embedding pieces of the sintered pellets in a resin, then grinding by successive SiC papers down to 5 μm and finally polishing with 3 μm and then with 1 μm diamond paste. The as-prepared resin blocks are finally gold-coated (∼ 10 nm thick) before the analyses. Further details on the characterization are found in [23]. When reporting statistical uncertainties of EDX measurements, we use the symbol ‘±’ to indicate maximum deviation from average and the Greek letter σ to indicate standard deviation.
2.2. Computational methodology

All density functional theory (DFT) simulations were carried out using the Castep code [25,26] by employing the PBE exchange-correlation functional [27] and ultra-soft pseudo potentials [28] with a consistent plane-wave energy cut-off of 450 eV. Point defects were investigated in a $3 \times 3 \times 1$ supercell (108 atoms) of the conventional unit cell of M$_3$AX$_2$, in conjunction with the aneto finite size correction [29]. Defect clusters were simulated in a $4 \times 4 \times 1$ supercell (192 atoms) instead. To test the accuracy of these supercells, selected defects were simulated in both supercells: the difference in energy was consistently below 0.02 eV for point defects, whilst it was as high as 0.10 eV for defect clusters (details provided in Section 3). $k$-point spacing was kept as close as possible to 0.03 Å$^{-1}$ throughout, which equates to Monkhorst-Pack [30] grids of $3 \times 3 \times 2, 4 \times 4 \times 2$ and $10 \times 10 \times 2$ for the 192-atom supercell, 108-atom supercell and the unit cell, respectively. No symmetry operations were enforced when calculating point defects and all simulations were spin-polarized. Since these systems are metallic, density mixing and cold smearing of bands (broadening width $= 0.1$ eV) were used. Electronic densities of states (DOS) were analyzed with the aid of the Optados program [31] with the adaptive broadening scheme [32].

3. Dilute Cr incorporation in Ti$_3$AlC$_2$

The incorporation of Cr in the Ti$_3$AlC$_2$ MAX phase [1,33–35] was tried with 5 and 33% substitution of Cr for Ti (target compositions (Cr$_{0.05}$Ti$_{0.95}$)$_3$AlC$_2$ and (Cr$_{1/3}$Ti$_{2/3}$)$_3$AlC$_2$, respectively). The samples are labeled as Cr$_5$Ti$_{95}$ and Cr$_{1/3}$Ti$_{2/3}$, respectively. According to XRD and after matching with the ICDD database, the Cr$_5$Ti$_{95}$ sample contained mainly a $312$ MAX phase in the $\alpha$ polymorph structure ($\sim 85$ vol%) accompanied with TiC ($\sim 15$ vol%) [23], the relative volume fraction being determined by Rietveld refinement. The unit cell parameters associated with the obtained MAX phase ($a = 3.0744(9)$ Å; $c = 18.567(7)$ Å) are in the range of literature values for Ti$_3$AlC$_2$ ($3.065 \leq a \leq 3.0786$ Å and $18.487 \leq c \leq 18.73$ Å) [34–38]. To investigate if significant amounts of Cr entered the $\alpha$-Ti$_3$AlC$_2$ structure, EDX measurements were performed on the sintered Cr$_5$Ti$_{95}$ sample. Global analyses ($\sim 100 \times 100$ μm square area) confirmed the expected Ti/Cr/Al ratios, while C was not measured (because of the low atomic mass of C, which precludes precise quantification by EDX and because of the probable contamination by SiC and diamond during polishing). Point measurements were then done on individual grains; the corresponding results are presented in Figure 2(a). The micrograph taken in back-scattered electron (BSE) mode shows three different phases:

- The darker phase (i.e. the phase containing the lightest elements by average atomic number and thereby likely to contain significant amount of C) was found to contain primarily Ti, possibly with some Al and/or Cr; however, the Al and Cr signal may be due to signal contamination by the surrounding grains as the grain size of this dark phase is of the same order as the EDX

![Figure 2](image-url)
analysis interaction volume (~1 μm³). As this phase is lighter than the other in the sample and as XRD determined that Cr₅Ti₉₅ contains ~15 vol% of TiC, it is safe to assume this first phase to be TiC.

- The main gray phase is expected to be the 312 MAX phase due to the XRD results and the lamellar shape of the grains. The seven-point analyses done on different grains of this phase in the area presented in Figure 2(a) have sensibly the same Al/(Al + Cr + Ti) ratio of 24 at% (with a maximum deviation of the results from the average value of ±1 at%), which agrees with the expected ratio for a 312 MAX phase. The Cr/(Cr + Ti) ratio is of 1.7 ± 0.3 at%. When combined with the 15-point analysis performed in other areas of the SEM sample, the average value for Cr/(Cr + Ti) becomes 2 ± 1 at%. According to EDX analyses and in conformity with the unit cell parameters of the MAX phase found to be equivalent to that usually reported for Ti₃AlC₂ [34–38], the Cr incorporation in Ti₃AlC₂ appears to be limited to a maximum of 2 at% of replacement of Ti by Cr, thereby forming (Cr₀.₀₂Ti₀.₉₈)₃AlC₂.

- The presence of a brighter Cr-rich third and last phase corroborates the above-mentioned findings. This phase is observed by SEM-BSE imaging and associated EDX but not by XRD. Excluding C, which is not detected, the composition of these white grains was found to be close to that of AlCr₂ with some Cr deficiency and small amounts of Ti (5–14 at%). As the phase has very bright contrast compared to the other phases in BSE imaging and is thus ‘heavier’, it is very likely that the phase does not contain much carbon and it is thus labeled as a metallic AlCr₁.₈₃Ti₀₂. XRD was not able to detect this phase either because it constituted too small a volume fraction (< ~ 5 vol%) or because it is amorphous.

The synthesis results for Cr₁/₃Ti₂/₃ are presented in [23]. Knowing at this stage the synthesis results of the previous compound, it was anticipated Cr and Ti would split into two separate MAX phases (e.g. Cr₂AlC + Ti₃AlC₂ or (Cr₂/₃Ti₁/₃)₃AlC₂ + Ti₃AlC₂); however, not a single MAX phase was found by XRD analysis and the Cr₁/₃Ti₂/₃ only consists of ~65 vol% TiC + ~35 vol% Al–Cr based alloys. This shows that when increased to a certain level, Cr not only does not enter the Ti₃AlC₂ structure but also precludes its formation, leading to this TiC + Al₃Cr c composite.

Accommodation of Cr in Ti₃AlC₂ was investigated at the atomic scale with DFT simulations, with the assumption that Cr would exclusively reside on the metallic M sites. This choice was made given prior DFT studies on this and similar systems [3,19] but also neutron diffraction studies such as that of Caspi et al. [39] which all agree, the usual M, A and X elements are only present on their respective crystallographic sites in the absence of radiation damage. The solution energies associated with the substitution of Cr for Ti were calculated for the two distinct Wyckoff sites:

\[
\begin{align*}
\text{Cr}(s) + \text{Ti}(4f) & \rightarrow \text{CrTi}(4f) + \text{Ti}(s), \quad (1) \\
\text{Cr}(s) + \text{Ti}(2a) & \rightarrow \text{CrTi}(2a) + \text{Ti}(s), \quad (2)
\end{align*}
\]

where Kröger–Vink notation is used to denote the substitution defect [40]. In both cases, the solution energy is positive (unfavorable), suggesting that Cr should exhibit only limited solubility in Ti₃AlC₂, in agreement with the experimental findings presented earlier. Notably, substitution onto the 4f site is 0.63 eV less unfavorable compared to substitution on the 2a site. Consequently, if small (dilute) amounts of Cr were added to Ti₃AlC₂, according to the Boltzmann distribution, 98.5% of the total Cr content should reside on the 4f site at the synthesis temperature of 1450°C, with the ratio increasing to 99.2% at the sintering temperature of 1220°C.

The relaxation volume associated with the incorporation of a single Cr atom, as defined in [41], was calculated as −6.508 Å³, in a 3 × 3 × 1 supercell containing 108 atoms (total supercell volume = 1378.15 Å³). This equates to a change in volume of −0.51% for every addition of 1 at% Cr. Equivalently, at the solubility limit of 2 at% Cr, as measured by EDX above, the predicted volume change from Ti₃AlC₂ to (Cr₀.₀₂Ti₀.₉₈)₃AlC₂ is only 1.02 vol%. This relatively small change confirms that standard powder XRD is unable to prove or disprove Cr presence in Ti₃AlC₂ at such low substitution rate, given the wider range of existing unit cell parameter values for Ti₃AlC₂ [34–38].

To investigate whether the Cr solubility was dependent on Cr concentration, defect clusters containing two Cr atoms were considered, in which at least one Cr atom occupies the more favorable 4f site. Figure 2(b) shows the binding energy associated with the formation of a [Cr₇₁: Cr₄₁] cluster, following Equation (3):

\[
E_b = E_{2\text{(Cr₇₁)}} - 2 E_{\text{Cr₄₁}},
\]

where \(E_b\) is the formation energy of the defect. A negative binding energy indicates increased stability against dilute defects. Clusters where both Cr atoms occupy 4f sites show small but negative binding energies. On the other hand, the clusters where one Cr atom occupies the 2a site are unstable by 0.6–0.7 eV, which is similar to the energy difference between the accommodation of Cr in 2a vs 4f sites in the dilute cases. This indicates that Cr exhibits limited solubility in Ti₃AlC₂ even when defect clusters...
are considered. It also indicate that Cr prefers to occupy the 4f site at low and higher concentrations alike, and will therefore tend toward the formation of a layered structure where all the Cr sites allowed to enter Ti3AlC2 resides on 4f sites, in line with the experimental observation of ordered (Cr2/3Ti1/3)3AlC2 [24].

4. Ordering and non-stoichiometry of (Cr2/3Ti1/3)3AlC2

(Cr2/3Ti1/3)3AlC2 was first obtained and reported by Liu et al. in 2014 [22,24] and in parallel synthesized by our research group and reported a year later [23]. We agree that the stable polymorph at room temperature of this MAX phase exhibits the α structure. After a thorough investigation by Raman spectroscopy, TEM, XRD and especially neutron diffraction (ND), Liu et al. [24] further found experimentally that α-(Cr2/3Ti1/3)3AlC2 has an ordered structure since the Cr atoms are, in the limit of sensitivity of ND, all located on the 4f sites, while the Ti atoms are only present on the 2a sites. This was corroborated by Dahlqvist and Rosen [19], who calculated by DFT the theoretical temperature of transition from an ordered to a disordered state to be over 3300°C, thereby inaccessible with conventional powder processing techniques. An interesting point also found in previous studies [22,23] concerns synthesis attempts of compositions close to (Cr2/3Ti1/3)3AlC2, which are (Cr3/4Ti1/4)2AlC2, (Cr1/2Ti1/2)3AlC2, (Cr4/7Ti3/7)3AlC2 and (Cr5/6Ti1/6)3AlC2: in all cases a 312 MAX phase with distortion and lattice parameters in line with that of ‘pure’ (Cr2/3Ti1/3)3AlC2 samples was formed along with either TiC (when (Cr1/2Ti1/2)3AlC2 and (Cr4/7Ti3/7)3AlC2 are originally targeted) or CrC + Cr2AlC (for (Cr5/6Ti1/6)3AlC2). This thus signifies that (Cr2/3Ti1/3)3AlC2 prefers to segregate excess Cr or Ti in secondary phases rather than to deviate from its nominal 2/3 Cr + 1/3 Ti ratio. The unit cell parameters determined for the 312 MAX phases in ‘(Cr3/4Ti1/4)2AlC’, ‘(Cr1/2Ti1/2)3AlC2’ and ‘(Cr2/3Ti1/3)3AlC2’ samples, given in Ref. [23] are very close to one another—typically within associated errors. This indicates that if a deviation of stoichiometry in (Cr2/3Ti1/3)3AlC2 occurs when the reactants contain an inadequate ratio of Cr/Ti/Al/C, the deviation is low enough that the resulting variation in lattice parameters measured by XRD falls within the range of reported XRD values for the lattice parameters of (Cr2/3Ti1/3)3AlC2 (2.921 ≤ a ≤ 2.935, 17.878 ≤ c ≤ 17.894) [22,23]. Similarly, the neutron diffraction refinement carried out by Liu et al. [24] did not show any sign of systematic non-stoichiometry in (Cr2/3Ti1/3)3AlC2 (e.g. through the presence of vacancies). To try further investigating possible non-stoichiometry in (Cr2/3Ti1/3)3AlC2, EDX was performed on the samples that we previously obtained when aiming at synthesizing (Cr3/4Ti1/4)2AlC, (Cr1/2Ti1/2)3AlC2 and (Cr2/3Ti1/3)3AlC2 thereafter labeled 211–Cr3/4Ti1/4, 312–Cr1/2Ti1/2 and 312–Cr2/3Ti1/3, respectively. Some characteristics of these samples syntheses are given in Table 1 for clarity.

An example of EDX results is given in Figure 3 for 211–Cr3/4Ti1/4. This SEM sample was analyzed in three separate occasions (8 months span time) and as detailed hereafter, the average quantifications from one session to another were found to be less than 1% different, suggesting the absence of a drifting of the EDX apparatus with time. In the example presented in Figure 3, we first note that the average Al/(Al + Ti + Cr) ratio for 6 measurements in the (Cr2/3Ti1/3)3AlC2 grains is 24.7 ± 0.7 at%. The value increases to 25.0 ± 2.1 at% (standard deviation σ = 1.0%) when averaged over 24 measurements taken during the three sessions and over a number of different locations. These values compare very closely with the EDX measurements performed on the 312–Cr1/2Ti1/2 (25.4 ± 1.1 at%) and the 312–Cr2/3Ti1/3 (24.6 ± 1.5 at%) samples. The total average for the (Cr2/3Ti1/3)3AlC2 phase across all samples remains 25.0 ± 2.1 at% (σ = 0.9%). Amongst MAX phases, one can easily find works where non-stoichiometries were measured [1,42–47] or suggested according to DFT calculations [20,47–51]. As in most if not all cases an X or A elemental deficiency was reported (e.g. Ti4AlN2.9 [44,45], Ti3Ge0.8C2 [42]), it appears reasonable to make the assumption that M sites in (Cr2/3Ti1/3)3AlC2 are fully occupied and that this can serve as the necessary reference for stoichiometry determination. Thereby, as the Al/(Al + Ti + Cr) ratio stays very close to 25 at% throughout the measurement sessions and the samples, it can be concluded that EDX, in the limit of its sensitivity, was not able to evidence any deficiency (or excess) in Al and the stoichiometry of Al in (Cr2/3Ti1/3)3AlC2 is therefore expected to be exactly 1, regardless of the initial reactants stoichiometries.

Table 1. (Cr2/3Ti1/3)3AlC2-containing samples considered for EDX characterization.

| Sample name | Targeted compound | Powder batch composition | Obtained phases (and Cr2/3Ti1/3AlC2 volume ratio) |
|-------------|------------------|--------------------------|--------------------------------------------------|
| 211–Cr3/4Ti1/4 | (Cr0.35Ti0.65)2AlC | 1.5 Cr + 0.5 TiH2 + 1.05 Al + 0.95 C | Cr2/3Ti1/3AlC2 (40%) + TiC + Al2Cr5 + Al2Cr3 |
| 312–Cr1/2Ti1/2 | (Cr0.35Ti0.65)3AlC2 | 1.5 Cr + 1.5 TiH2 + 1.05 Al + 1.9 C | Cr2/3Ti1/3AlC2 (50%) + TiC + Al2Cr5 + Al2Cr3 |
| 312–Cr2/3Ti1/3 | (Cr0.35Ti0.65)3AlC2 | 2 Cr + 1 TiH2 + 1.05 Al + 1.9 C | Cr2/3Ti1/3AlC2 (95%) + TiC + Al2Cr5 |
Regarding the Cr/(Cr + Ti) ratio in $^{\text{211}}$–Cr$_{\text{3/4}}$Ti$_{\text{1/4}}$, the results are presented following the same data treatment and considering the same EDX points selected for the previous analysis on Al content. The Cr/(Cr + Ti) ratio for the 6 data points given in Figure 3 is 65.4 ± 2.4%. The average over 24 points is 65.7 ± 2.7% ($\sigma = 1.6\%$), respectively; this leads to an average value of 65.7 ± 3.3% ($\sigma = 1.5\%$) over all three samples. Once again the measured ratio does not vary much from a measurement session to another, proving the stability over time of the SEM/EDX apparatus, but more importantly Cr/(Cr + Ti) does not vary from one sample to another, despite differences in initial powders stoichiometries (excess Ti in the $^{3\text{12}}$–Cr$_{\text{1/2}}$Ti$_{\text{1/2}}$ and $^{3\text{12}}$–Cr$_{\text{2/3}}$Ti$_{\text{1/3}}$ samples, and excess Cr in the $^{\text{211}}$–Cr$_{\text{3/4}}$Ti$_{\text{1/4}}$ sample). Therefore it can be inferred that, within the limits of precision of our EDX characterization, the Cr/(Cr + Ti) ratio in (Cr$_{2/3}$Ti$_{1/3}$)$_3$AlC$_2$ appears independent of the reactants stoichiometries. Besides one can also note that all samples show a Cr/(Cr + Ti) about 1% lesser than the theoretical and expected 2/3 (66.7%) value. Of course the scattering of data and the associated standard deviation are generally greater than 1%, but the fact that the deviation from the 2/3 theoretical value is for the 3 samples always falling on the same side indicates that if a real deviation exists, this EDX characterization suggests it be in favor of a Ti excess over Cr, i.e. through the substitution of Ti for Cr onto 4f Wyckoff sites, as it is calculated hereafter by DFT.

For DFT investigations, the $\alpha$–(Cr$_{2/3}$Ti$_{1/3}$)$_3$AlC$_2$ structure was first simulated by ordering the Cr atoms on the 4f sites and the Ti atoms on the 2a sites (Figure 1), in line with DFT results from Dahlqvist and Rosen [7], experimental findings of Liu et al. [24] and observations of such ordering in similar MAX phases [39,52]. The formation of energy of (Cr$_{2/3}$Ti$_{1/3}$)$_3$AlC$_2$ and its ternary end-members are presented, in Equations 3–5:

\begin{align}
3\text{Ti}_\text{(hcp)} + \text{Al}_\text{(fcc)} + \text{C}_\text{(2h)} & \rightarrow -4.68\text{eV} \rightarrow \text{Ti}_3\text{AlC}_2, \quad (3a) \\
\frac{1}{2}\text{Ti}_2\text{AlC} + \frac{1}{2}\text{Ti}_4\text{AlC}_3 & \rightarrow -0.078\text{eV} \rightarrow \text{Ti}_3\text{AlC}_2, \quad (3b) \\
2\text{Cr}_\text{(bcc)} + \frac{1}{2}\text{Ti}_\text{(hcp)} + \text{Al}_\text{(fcc)} + \text{C}_\text{(2h)} & \rightarrow -2.31\text{eV} \rightarrow \text{Cr}_2\text{TiAlC}_2, \quad (4a) \\
\text{Cr}_2\text{AlC} + \text{TiC} & \rightarrow -0.081\text{eV} \rightarrow \text{Cr}_2\text{TiAlC}_2, \quad (4b) \\
3\text{Cr}_\text{(bcc)} + \text{Al}_\text{(fcc)} + \text{C}_\text{(2h)} & \rightarrow -0.26\text{eV} \rightarrow \text{Cr}_3\text{AlC}_2, \quad (5a) \\
\text{Cr}_2\text{AlC} + \frac{1}{3}\text{Cr}_3\text{C}_2 + \frac{1}{3}\text{C}_\text{(2h)} & \rightarrow +0.41\text{eV} \rightarrow \text{Cr}_3\text{AlC}_2. \quad (5b)
\end{align}

Here we consider both the formation energy from standard state (Equations 3a, 4a, 5a), and the reaction energy from the most competing phases, as identified by Dahlqvist and Rosen [19,20] (Equations 3b, 4b and 5b). (Cr$_{2/3}$Ti$_{1/3}$)$_3$AlC$_2$ exhibits the lowest reaction energy when considering the most competing phases. It has been shown that (Cr$_{2/3}$Ti$_{1/3}$)$_3$AlC$_2$ exhibits multiple metastable spin states [19,53]; therefore all previously reported spin states were considered and the reaction energies of equation 4b are presented in Table 2. In agreement with Dahlqvist and Rosen [19], we find that the in-plane antiferromagnetic ordering type XX (following the nomenclature of Wang et al. [53]) is the most favorable spin state, closely followed by simple ferromagnetic ordering with a difference of only 1 meV. Note that this energy difference is much below the expected accuracy of point defect calculations within DFT (see methodology section, and the effect of on-site Coulomb interaction on states and energies).
Table 2. Relative stability of (Cr$_{2}/3$Ti$_{1}/3$)$_3$AlC$_2$ under different spin configurations.

| Spin order | Supercell size | Equation 4b (meV) |
|------------|----------------|------------------|
| NM         | Unit cell      | 40               |
| FM         | Unit cell      | $-62$            |
| AFM-X      | Unit cell      | 21               |
| AFM-1      | Unit cell      | 0                |
| AFM-A      | Unit cell      | 33               |
| AFM-FF     | $2 \times 1 \times 1$ | $-62$            |
| AFM-XX     | $2 \times 1 \times 1$ | $-81$            |
| AFM-11     | $2 \times 1 \times 1$ | 36               |

correction discussed by Wang et al. [53]); therefore all defect calculations were performed with the ferromagnetic ordering only, thus allowing greater flexibility on the size of supercell adopted.

As discussed in the previous section, if the phase forms by progressive additions of Cr into the Ti$_3$AlC$_2$ structure, then it is expected to retain the ordered, layered structure of (Cr$_{2}/3$Ti$_{1}/3$)$_3$AlC$_2$. However, to investigate the driving force for ordering of (Cr$_{2}/3$Ti$_{1}/3$)$_3$AlC$_2$, the formation energy of anti-site defects (i.e. swapping the positions of a Cr and a Ti atom to form a (Ti$_{Cr}$,Cr$_{Ti}$) defect pair) is considered, following the method outlined in [54]. Six anti-site defect configurations were considered: three configurations where the swapping atoms reside in neighboring planes (part of the same carbide layer) and three configurations where the defects are separated by the Al layer. In all cases, the formation energy was unfavorable at 0.82 ± 0.06 eV, confirming that there is a significant preference for ordered layering of Cr and Ti atoms at this composition.

Interestingly, similarly to the [Cr$_{Ti}$/Cr$_{Ti}$] defects clusters in Ti$_3$AlC$_2$ discussed earlier, the formation energies of the anti-site clusters were found to be largely insensitive to the distance or location of the defect pair (standard deviation of only 0.06 eV). Furthermore, the formation energy for dilute non-interacting defects (i.e. Ti$_{Cr}$ + Cr$_{Ti}$ in separate simulations) was calculated at 0.80 eV, which is within the uncertainty of the formation energy of the bound defects. This suggests that there is no driving force for either clustering or separation of anti-site defects if these were forcefully introduced in the system (e.g. through radiation damage).

The ability of the structure to accommodate more (or less) Cr than exactly 2/3 of the total M sites can be calculated from the formation energy of individual Cr$_{Ti}$ (or Ti$_{Cr}$) defects in the ordered (Cr$_{2}/3$Ti$_{1}/3$)$_3$AlC$_2$ structure. The formation energy of the defect is sensitive to the choice of reference phase, that is, what is the chemical form of the local reservoirs of Ti and Cr atoms. Here we consider six possible scenarios where the reference phases were taken (Equation 6) from standard state (i.e. metallic Ti and Cr), (Equation 7) from the parent 312–MAX phases, and (Equation 8) from the lower order 211–MAX phases.

\[
\text{Ti}_{(s)} + \text{Cr}_{(s)} \rightarrow \text{Ti}_{Cr} + \text{Cr}_{Ti}, \quad (6a)
\]

\[
\text{Cr}_{(s)} + \text{Ti}_{(s)} \rightarrow \text{Cr}_{Ti} + \text{Ti}_{(s)}, \quad (6b)
\]

\[
\frac{1}{3}\text{Cr}_3\text{Al}_2\text{C}_{2(s)} + \text{Cr}_{Cr} \rightarrow \text{Ti}_{Cr} + \frac{1}{3}\text{Cr}_3\text{Al}_2\text{C}_{2(s)}, \quad (7a)
\]

\[
\frac{1}{3}\text{Cr}_3\text{Al}_2\text{C}_{2(s)} + \text{Ti}_{Ti} \rightarrow \text{Cr}_{Ti} + \frac{1}{3}\text{Ti}_3\text{Al}_2\text{C}_{2(s)}, \quad (7b)
\]

\[
\frac{1}{2}\text{Ti}_2\text{Al}_2\text{C}_{2(s)} + \text{Cr}_{Cr} \rightarrow \text{Ti}_{Cr} + \frac{1}{2}\text{Cr}_2\text{Al}_2\text{C}_{2(s)}, \quad (8a)
\]

\[
\frac{1}{2}\text{Cr}_2\text{Al}_2\text{C}_{2(s)} + \text{Ti}_{Ti} \rightarrow \text{Cr}_{Ti} + \frac{1}{2}\text{Ti}_2\text{Al}_2\text{C}_{2(s)}. \quad (8b)
\]

The values reported do not consider all possible phases that may form, only the simplest, most intuitive, cases where reactants and products are the constituent metals or ternary MAX phases or carbides. A comprehensive analysis of all the possible binary, ternary and quaternary phases that may form is clearly beyond the scope of this investigation, and conclusions may be inferred without such considerations. For instance, since the formation energy for accommodating excess Cr (Equations 6b, 7b and 8b) is unfavorable in all scenarios, then it is clear that considering additional (more stable) reactants may only render Cr$_{Ti}$ accommodation even more unfavorable, thus (Cr$_{2}/3$Ti$_{1}/3$)$_3$AlC$_2$ is expected to show no meaningful deviation of stoichiometry in the case of Cr excess. These findings are in agreement with the results of Dahlqvist and Rosen [19], where they simulated the compositional range Ti$_{3-x}$Cr$_x$AlC$_2$, where $x = \{0,0.5,1,1.5,2,2.5,3\}$ (equivalent to concentration steps of 8.33 at%), whereas the current work considered dilute deviations from stoichiometry of $\sim$0.925 at% (1 atom replaced in 108-atom supercells). Regarding Ti excess (Equations 6a, 7a and 8a), on the other hand, the reaction energy varies from highly favorable ($\sim$1.10 eV) in the case of metal reactants (unlikely in the presence of C) to highly unfavorable (0.37 eV) in the case of a hypothetical MAX phase Cr$_3$AlC$_2$ product. Since Cr$_3$AlC$_2$ is known not to form, any binary or ternary compound that forms in its place must have lower formation energy, thus reducing the reaction energy for accommodation of Ti$_{Cr}$. Nevertheless, irrespective of reactants and products, the accommodation of excess Ti is consistently more favorable than the accommodation of excess Cr, thus suggesting that if any non-stoichiometry may be accommodated in (Cr$_{2}/3$Ti$_{1}/3$)$_3$AlC$_2$, it will necessarily be toward Ti excess.
5. Factors contributing to the stability of 
(Cr$_{2/3}$Ti$_{1/3}$)$_3$AlC$_2$

Calculated lattice parameters of the (Cr$_x$Ti$_{1-x}$)$_3$AlC$_2$ series are presented in Figure 4, including, for the purpose of comparison, hypothetical ordered phases (Cr$_{1/3}$Ti$_{2/3}$)$_3$AlC$_2$ and Cr$_3$AlC$_2$ that are known not to form experimentally. Incidentally those phases that are thought to be unstable also proved difficult to converge at the computational level, suggesting that perhaps at the root of their instability is the electron distribution. Only the lowest energy magnetic configuration is reported for each composition in Figure 4 as these best represent real-life materials. However, changes in magnetic ordering are known to affect the lattice parameters of these materials significantly, as thoroughly investigated by Wang et al. [53] Notably, the change in lattice parameters between ferromagnetic and non-magnetic Cr$_3$AlC$_2$ was of the same order as the change in lattice parameter between Ti$_3$AlC$_2$ and Cr$_3$AlC$_2$. The calculated lattice parameter of Ti$_3$AlC$_2$ ($a = 3.078$ Å, $c = 18.654$ Å) fall within the range of experimental values from literature ($3.065$ Å $\leq a \leq 3.076$ Å and $18.487$ Å $\leq c \leq 18.73$ Å [34–38]).

For (Cr$_{2/3}$Ti$_{1/3}$)$_3$AlC$_2$ the calculated lattice parameters vary significantly depending on the spin ordering; however, for the two most stable spin states the lattice parameters were $a_{FM} = 2.930$ Å, $c_{FM} = 17.830$ Å and $a_{AFM-XX} = 2.921$ Å, $c_{AFM-XX} = 17.874$ Å. These are also within or very close to the range of experimental values found in the literature: $2.921$ Å $\leq a \leq 2.935$ Å, $17.878$ Å $\leq c \leq 17.894$ Å [23,24]. It is well established that the choice of GGA exchange-correlation functional may lead to an overestimation or underestimation of the lattice parameter for most material systems. However, it is also acknowledged that the error is generally consistent within a system series; therefore the prediction of lattice parameter change due to a change in composition is typically highly accurate with DFT-GGA.

(Cr$_{2/3}$Ti$_{1/3}$)$_3$AlC$_2$ exhibits a significant deviation from Vegard’s law, resulting in an overall reduction in volume compared to the linear trend and, surprisingly, an exceptional increase in $c/a$ ratio above that of any other ternary or quaternary 312–MAX phase in the series. This is a strong indication that the nature of the in-plane and out-of-plane bonds is different at this specific composition. To investigate this further, the electronic DOS were calculated for the four phases and are presented in Figure 5.

Addition of Cr creates sharp peaks in the energy region near the Fermi level (Figure 5), which are typically indicative of localized states. Furthermore, addition of Ti or Cr beyond the 2/3:1/3 ratio causes spin polarization of the structure as indicated by the asymmetric shape of the DOS of (Cr$_{1/3}$Ti$_{2/3}$)$_3$AlC$_2$ and Cr$_3$AlC$_2$. The DOS of the stable phases, (Cr$_{2/3}$Ti$_{1/3}$)$_3$AlC$_2$ and Ti$_3$AlC$_2$, further expanded into the individual orbital contributions in Figure 6. It is evident that spin polarization of the quaternary MAX phase is not limited to the Cr atoms (where it is most pronounced), but it is also observed in the projected DOS of all other atomic species. Furthermore Al and C exhibit changes in densities around the Fermi level, with new narrow bands forming in the spin-up component of the DOS at the same energies as the most pronounced Cr spin-up bands, suggesting that there are (partly) covalent Al–Cr and C–Cr bonds in (Cr$_{2/3}$Ti$_{1/3}$)$_3$AlC$_2$. To further investigate this, Mulliken population analysis was performed (Table 3) and the charge density difference (with respect to the sum of individual atomic charge densities) was extracted from

![Figure 4](image-url)
the relaxed lattices (Figure 7). The results are shown for the ferromagnetic configuration only, but the AFM–XX structure yielded population analysis results within 2% of the FM values.

The 4f site—occupied by Cr in (Cr2/3Ti1/3)3AlC2—links the carbide layer with the aluminum layer, whilst the 2c site—occupied only by Ti—is surrounded exclusively by C atoms (see Figure 1). Consequently, it seems sensible to attribute the increase in stability of the (Cr2/3Ti1/3)3AlC2 phase to Al-Cr interactions. However, the population analysis shows that the electron density of the Al–Cr bond in (Cr2/3Ti1/3)3AlC2 is reduced by \( \sim 20 \) vol% compared to the Al–Ti bond in Ti3AlC2. And whilst the Al–Cr bond is 19.3 pm shorter than the Al–Ti bond, which typically is an indication of a stronger bond, the difference in bond length is entirely accounted for by the smaller atomic radius of Cr compared to Ti (124.9 pm vs. 144.8 pm) [55].

On the other hand, a higher localization of electron is observed around the 4f Cr atom compared to the 4f Ti atoms in Ti3AlC2 (Figure 7). A corresponding increase in electron density of \( \sim 15 \) vol% is found around the C–Cr bond compared to the same sites in Ti3AlC2. At the same time, the C–Ti and Al–Cr bonds are markedly depopulated. This suggests that the increase in stability of the (Cr2/3Ti1/3)3AlC2 is largely due to the C–Cr interaction, despite the preferential location of Cr atoms (bridging Al layers and carbide layers). As a consequence of the redistribution of electrons, the local electron density in and around the Al layer is increased (\( \sim 36 \) vol%),

**Figure 5.** Electronic density of states, aligned with respect to the Fermi level, of the (Cr,Ti)3AlC2 series.
which may also play a role in the stability of layered (Cr$_{2/3}$Ti$_{1/3}$)$_3$AlC$_2$. Dahlqvist and Rosen [19] suggested that the addition of M elements with higher electronegativity than Ti (such as Mo in their paper and Cr in this paper) near the Al layer causes the Al atoms to become more positively charged, thus reducing the number of electrons available for the Al–Al anti-bond. This is in contrast to our finding that the local (relative) density around Al atoms increases. Nevertheless, Figure 6 shows that the DOS of the Al $p$ orbital in (Cr$_{2/3}$Ti$_{1/3}$)$_3$AlC$_2$ is shifted toward lower energies compared to Ti$_3$AlC$_2$, suggesting an energetically more favorable bond, in line with the findings of Dahlqvist and Rosen [19]. This is corroborated by a subtle change in the electron density surrounding the Al atom: the in-plane density maximum between Al atoms in Ti$_3$AlC$_2$ splits into two (stronger) maxima around Al atoms.
in \((\text{Cr}_{2/3}\text{Ti}_{1/3})_3\text{AlC}_2\), one above and one below the Al layer. Finally, it is observed that the in-plane M–M interactions also exhibit a significant charge transfer when Cr is included. In \(\text{Ti}_3\text{AlC}_2\), the basal plane \((2a–2a)\) Ti–Ti bonds are nearly identically populated to the in-plane \((4f–4f)\) Ti–Ti bonds. However, in \((\text{Cr}_{2/3}\text{Ti}_{1/3})_3\text{AlC}_2\) the in-plane Cr–Cr bond \((4f–4f)\) is markedly depopulated in favor of the Ti–Ti basal bond \((2a–2a)\). This may be a further source of increased stability for the Ti carbide layers in the quaternary Cr–Ti–Al–C MAX phase.

It is believed that similar redistribution of bond charges might be at the root of stability of other \((M, M')_{n+1}\text{AX}_n\) MAX phases recently reported. ANasori et al. [52] reported a quaternary \((\text{Mo,Ti})_3\text{AlC}_2\) with the same layered ordering and \(M/(M + Ti)\) ratio as \((\text{Cr}_{2/3}\text{Ti}_{1/3})_3\text{AlC}_2\). The analogy with the Cr-Ti system continues since, like \(\text{Cr}_3\text{AlC}_2\), \(\text{Mo}_3\text{AlC}_2\) is known not to form. Interestingly Zhou et al. [56] and Caspi et al. [39] reported stable \((\text{Cr}_{0.5}\text{V}_{0.5})_{n+1}\text{AlC}_n\) MAX phases with \(n = 1, 2, 3\). This behavior is distinctly different from that of the Cr-Ti-Al-C and Mo-Ti-Al-C systems, where stable phases are observed only at the specific \(M/(M + Ti)\) ratios that allow layered ordering of M and Ti atoms \((2/3\) in the case of \(312\)-MAX phases). In fact, the lack of a stable solid solution \((\text{Cr},\text{Ti})_2\text{AlC} [23]\) may be explained in terms of the inability of \(211\)-MAX phases (i.e. \(n = 1\)) to achieve such layer ordering at any \(\text{Cr}/(\text{Cr} + \text{Ti})\) ratio. The fact that \((\text{Cr}_{0.5}\text{V}_{0.5})_{n+1}\text{AlC}_n\) may be achieved even for \(n = 1\), suggests that the layered ordering of M atoms is less important to the phase stability in the V–Cr–Al–C
system. This is further corroborated by the recent results of Hamlin et al., [57] where (Cr_{0.5},V_{0.5})_{n+1}AlC_n MAX phases (n = 1,2,3) were synthesized with Cr/(Cr+V) ratios of 1/4, 1/2 and 3/4 (the latter being unstable only for n = 3). Nevertheless, a strong tendency toward ordering is observed for n = 2 and n = 3, as reported by the neutron diffraction data [39]. The results of the current work, together with those of Caspi et al., [39] suggest that if the stoichiometry of the compounds is adjusted to reflect the multiplicity of the different Wyckoff sites (i.e. Cr/(Cr+V) = 2/3 for n = 2), then a V–Cr–Al–C 312-MAX phase with layered ordering of Cr and V atoms may be obtained, and this may have improved stability. Nevertheless, the ability of (Cr,V)_{n+1}AlC_n to accommodate disorder on the M site suggests that perhaps it may be more radiation tolerant than MAX phases such as (Cr_{2/3}Tl_{1/3})_3AlC_2, where ordering is a pre-requisite for stability.

6. Conclusions

Through a combination of DFT simulations and experimental synthesis, it was found that:

- Cr exhibits limited solubility in Ti_3AlC_2 (~ 2 at%) and that all the Cr is accommodated on the 4f Wyckoff site, that is, bridging the Al layer to the carbide layer.
- Larger amounts of Cr (Cr/(Cr + Ti) ratio ≥ 1/2) enable formation of (Cr_{2/3}Tl_{1/3})_3AlC_2, which is confirmed to be a fully ordered structure with Cr atoms on the 4f sites and Ti atoms on the 2a site.
- Deviation from stoichiometry of (Cr_{2/3}Tl_{1/3})_3AlC_2 is predicted and measured to be negligible in the case of Cr-excess but the results are inconclusive regarding Ti-excess, where if deviation occurs it is expected to be mediated by TiC substitutions. EDX also found that the ratio Al/(Al + Cr + Ti) in (Cr_{2/3}Tl_{1/3})_3AlC_2 does not deviate significantly from the expected 1/4 value suggesting that the A site is fully occupied by Al.
- (Cr_{2/3}Tl_{1/3})_3AlC_2 has a smaller volume (132.6 Å³ simulated, 133.2 ± 0.2 Å³ measured) and surprisingly larger c/a ratio (6.084 simulated, 6.101 ± 0.003 measured) than the other 312-MAX phases in the series.
- Increase in stability is primarily attributed to the formation of favorable Cr–C bond despite the preferential location of Cr (bridging Al and carbide layers), though magnetization may also play an important role, as may the redistribution of charge in the Al layer.
- Other (M,M')_{n+1}AX_n MAX phases with layered ordering of M on 2a sites and M' on 4f sites are expected to exhibit increased stability due to the same factors (point above).

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