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Atomic-scale study of low-temperature equilibria in iron-rich Al-C-Fe

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The capability of the thermodynamic approach based on the independent point defect approximation to describe low-temperature phase equilibria is investigated and applied to the Al-C-Fe system. The method gives a reasonable description of the multicomponent and multisublattice Fe-rich corner and evidences numerous peculiarities concerning the ordered phases as well as the density-functional-theory (DFT) energy models. The study of Fe3Al(C), revealing strong defect-induced instabilities, rules out the LDA, SLDA and GGA schemes and leaves (spin-polarized) SGGA as the only valid one. C stabilizes L12 Fe3Al with respect to D03, which justifies the fcc-type structure of the κ Fe3AlC compound. The present work also helps in justifying the experimentally observed depletion of C in the κ phase. Finally, a correct description of both Fe3C and κ requires inclusion of interstitial carbon at low temperature, emphasizing the unexpected importance of interstitial defects in ordered phases.

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I. INTRODUCTION

The Al-C-Fe system is currently used to create multiphase alloys with controlled mechanical properties at high temperatures. During the thermomechanical elaboration treatments, transient phases appear, among which the κ Fe3AlC carbide with E21 structure shows especially intriguing properties. At high temperature (>1000 K), this compound is experimentally known1 to display a pronounced off-stoichiometry, since it appears in the equilibrium phase diagram only for low C contents (0.10<xC<0.15), which indicates the presence of high amounts of point defects with possibly intricate degrees of order. Besides this experimental study, the Al-C-Fe phase diagram was also calculated2 through an analysis combining phenomenological thermodynamics and ab initio calculations, including evaluations of the energies of selected structures with various compositions. Although these calculations show qualitative agreement with experiment (presence of off-stoichiometry), the calculated composition extent of the κ phase is significantly shrunk to a line compound, a signature of sharp free-energy curves.

The κ compound is thus expected to play a central role in multiphase Al-C-Fe systems, and it would certainly be useful to characterize more accurately its properties, a task that can adequately be performed using atomic-scale approaches. However, the only previous work that tackled this issue, with ab initio calculations,3 was concerned merely with the behavior of the C vacancy in κ. This work showed the strong relaxation of the iron atoms around the defect, inducing a switch from the local environment of L12 Fe3Al to that of an iron vacancy in B2 FeAl. From these calculations, the authors were able to propose a structural path for the formation of the κ phase from L12 Fe3Al. However, due to the free boundary conditions employed, this result could not be used to infer the bulk thermodynamic properties of κ. Apart from this analysis of a particular point defect, no attempt has been made so far toward atomic-scale theoretical investigations of the properties of κ Fe3AlC.

In order to approach the low-temperature thermodynamic behavior of complex alloys such as Fe3AlC, it is necessary to have an exhaustive picture of their point defect properties, since the latter are directly connected to the chemical potentials of the species, the knowledge of which is required for realistic modeling. To achieve this, the most general approach would involve the elaboration of a cluster description of the energetics of the compound, suitable for use in either semianalytical (variational method) or numerical (Monte Carlo simulations) equilibrium calculations. The previously mentioned pieces of experimental evidence require the consideration of all kinds of point defects, including vacancies, which implies handling a cumbersome pseudoquaternary Al-C-Fe-vacancy system. Whereas the vast majority of the available cluster analyses have been concerned with binary systems, only a few approaches have tried to extend the procedure to [binary+vacancies].4,5 Alternatively, an independent point defect analysis constitutes a convenient way to estimate the behavior of ordered compounds. In spite of its limitations (low amounts of defects; hence moderate temperature and off-stoichiometry), better theoretical knowledge of the point defect properties of Fe3AlC would eventually help in: (i) estimating the trends of the Al-C-Fe system in the iron-rich composition range and (ii) understanding the kinetic paths followed by the system during the elaboration processes and, therefore, the final microstructures.

With the κ compound appearing experimentally at equilibrium with other alloys, a realistic modeling should also encompass the surrounding phases, which justifies investigations extending over the whole Fe-rich corner of the Al-C-Fe system. In this context, the present work is devoted to a detailed atomic-scale analysis of the T=0 K energetics of the iron-rich Al-C-Fe system, with special attention paid to κ Fe3AlC, by means of ab initio density-functional calculations coupled with a thermodynamic treatment based on point defects.
As tackling the properties of ternary Al-C-Fe also requires the knowledge of the underlying binary systems, it is worth recalling the amount of work already performed on Al-Fe and C-Fe in the relevant composition domain (iron-rich corner). It is a well-known fact that serious difficulties are encountered when using density-functional theory (DFT) to assess the properties of iron-containing alloys (Fe-Al dilute solid solution; B2, D03, and L12 ordered compounds). Indeed, some improvement was achieved recently for the D03 phase through the LDA+U method, but the latter remains confined to specific studies and its inherently higher degree of parameterization makes it uneasy to generalize. As more specifically regards the D03 and B2 compounds, the predicted point defect properties have been shown to depend strongly on the DFT formalism used, with additional intricacy due to the importance of including magnetism. In particular, vacancies in B2 FeAl are seemingly better described by spin-polarized local-density approximation (SLDA). However, as will be emphasized in the present work, SLDA can by no means be considered as intrinsically better than spin-polarized generalized gradient approximation (SGGA) in any situation. Further investigation is greatly required to estimate the (S)LDA’s and (S)GGA’s respective merits for iron alloys. (For brevity, we will use the notation (S)LDA (respectively (S)GGA) to refer to both the LDA and SLDA (respectively GGA and SGGA) frameworks.) Regarding the low-temperature D03 Fe3Al phase, very few theoretical results are available when compared to those pertaining to B2 FeAl. Valuable (in reasonable agreement with experiments) semi-quantitative information concerning the point defect formation energies were however provided by DFT calculations without magnetism, which justifies studies (such as the present one) relying on classical DFT approaches without the U term.

Among the identified Fe-C compounds, namely, the solid solution, the D011 Fe14C cementite, and the Fe60C and Fe2C phases, only the first two ones were retained in our investigations, since the other carbides show a degree of metastability sufficient for their influence to be neglected in an equilibrium study. Conversely, the metastability of cementite being much lower, it was not reasonable to rule it out a priori, all the more since it may be strongly stabilized by ternary additions such as Cr. A DFT (projector-augmented wave PAW–SGGA) analysis of complex point defects in the Fe-C solid solution at moderate temperature (430 K), including defect free energies, showed a considerable binding between C and vacancies. It is also worth mentioning the DFT studies of carbon dissolution and diffusion in ferrite and austenite. To our knowledge, however, no atomic-scale study of ordered C-Fe compounds has been performed.

This paper is thus concerned with the low-temperature equilibrium properties of the Al-C-Fe system in the composition domain limited by the following phases: (i) the Fe(-Al,C) solid solution, (ii) the Fe2Al compound with low amounts of C, (iii) the Al-doped Fe14C cementite, and (iv) the Fe3AlC ordered ternary carbide. Although it is the best way to perform this investigation, the exhaustive determination of the convex hull of the system in the whole \[0 < (x_{Al}, x_C) < 20\%\] composition range (as ordinarily done in \(T=0\) K cluster studies) would be a task exceedingly long (requiring the determination of an optimal cluster expansion for the pseudoquaternary fcc-based Al-C-Fe-vacancy system) and inherently incomplete, due to the variety of underlying crystallographic structures (\(L12, D03, D011,\) and \(\kappa\)). Our purpose here is therefore to obtain, at low temperature \((T=0\) K), relevant information on this pseudoquaternary system via a more tractable point defect analysis [enabling examination of the relative positions of the \(H(\alpha_{Al}, \alpha_C)\) surfaces], since each such surface around stoichiometry is approximately a linear function of the point defect energies.

The present study is thus intended to help in estimating the relative phase stabilities, with particular attention paid to the effect of C and to the possible reasons for the experimental off-stoichiometry of the \(\kappa\) phase. It also provides additional information about the respective merits of the DFT approximations currently used to describe the properties of iron aluminides, these alloys being well known as sources of difficulties in atomic-scale simulations. To this aim, after outlining the relevant methods (Sec. II), a point defect analysis is carried out for each phase in its composition range (Sec. III), yielding the chemical potentials around stoichiometry. The latter are then employed to assess the enthalpies as a function of composition, which eventually allows us to discuss the main low-temperature trends (Sec. IV).

## II. METHODS

The \(\kappa\) Fe3AlC phase with \(E2_1\) perovskite structure (CaTiO3,cP5), of chief interest here, has a unit cell (Fig. 1) that can be viewed as an \(L12\) Fe3Al (AuCu3,cP4) compound with one supplementary C atom occupying the central site. Its three sublattices thus give rise to nine types of simple point defects (six antisites and three vacancies). In addition, because of the practical importance of interstitial occupancy for carbon in iron-based alloys, the present study takes into account interstitial C in either tetrahedral \((111)\) or octahedral \((000)\) position, the latter labeled as \(O_{Fe-Al}\) below, due to its mixed (four Fe and two Al) neighbor environment. From considerations of atomic size, the possibility of interstitial Al or Fe was neglected. Concerning the \(D03\) Fe3Al compound (BiF3,cF16), we respectively labeled as Fe2 and Fe3 the mixed Al-Fe and pure Fe cubic sublattices and as \(O_{Fe-Al}\) and \(O_{Fe}\) the interstitial octahedral sites located at the centers of

![FIG. 1. Unit cell of the Fe3AlC \(\kappa\) phase, showing both types of interstitial sites \(O_{Fe}\) and \(O_{Fe-Al}\) of the \(L12\) "parent" structure (the former site being fully occupied by C in \(\kappa\)).](image-url)
TABLE I. Reference (eV/atom) and GC energies (eV) of point defects in $D_0_3$ and $L_1_2$, Fe$_3$Al(-C), calculated in the (S)LDA and (S)GGA frameworks with $1 \times 1 \times 1$ (16 atoms) and $2 \times 2 \times 2$ (32 atoms) cubic supercells for $D_0_3$ and $L_1_2$, respectively. In order to further check the convergence and some surprising trends, noted for both compounds with all but the SGGGA DFT formalisms, values for antisites pertaining to larger supercells ($2 \times 2 \times 2$, 32 atoms, and $3 \times 3 \times 3$, 108 atoms, for $D_0_3$ and $L_1_2$, respectively) are also provided in parentheses (see text).

<table>
<thead>
<tr>
<th>$D_0_3$</th>
<th>Ref.</th>
<th>$V_{Al}$</th>
<th>$Fe_{Al}$</th>
<th>$V_{Fe_2}$</th>
<th>$V_{Fe_3}$</th>
<th>$Al_{Fe_2}$</th>
<th>$Al_{Fe_3}$</th>
<th>$C_{Al}$</th>
<th>$C_{Fe_2}$</th>
<th>$C_{Fe_3}$</th>
<th>$C_{oct_{Fe}}$</th>
<th>$C_{oct_{Fe-Al}}$</th>
</tr>
</thead>
</table>

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<tr>
<th>$L_1_2$</th>
<th>Ref.</th>
<th>$V_{Al}$</th>
<th>$Fe_{Al}$</th>
<th>$V_{Fe}$</th>
<th>$Al_{Fe}$</th>
<th>$Al_{Fe_2}$</th>
<th>$Al_{Fe_3}$</th>
<th>$C_{Al}$</th>
<th>$C_{Fe}$</th>
<th>$C_{oct_{Fe}}$</th>
<th>$C_{oct_{Fe-Al}}$</th>
</tr>
</thead>
</table>

The properties of point defects are conveniently described in terms of the so-called grand canonical (GC) quantities, measuring the excess of the property (with respect to the reference undefected system) in a volume surrounding the defect.23,24 The GC parameters are essentially the thermodynamic extensive variables (namely, the energy, the volume, and the magnetic moment); under the common conditions of zero pressure and magnetic field, the defect-dependent equilibrium properties of a phase depend merely on the GC energies. Therefore, the ground-state point defect structure can be obtained by minimizing the $T=0$ K enthalpy of the system with respect to the point defect numbers at constant composition,25 the key quantities in this approach being the GC energies of point defects, identical to GC enthalpies under zero pressure, and noted as $h_{d}$ for a defect $d$ on sublattice $s$:

$$h_{d} = H(N_{d}^{s} = 1, N_{d' \neq d}^{s} = 0) - H_{0}$$

($H_{0}$ being the enthalpy of the undefected crystal). Under the assumption of independent defects, the $T=0$ K enthalpy of the system containing $M$ unit cells and amounts $\{N_{d}\}$ of defects of various types $d$ is thus written as

$$H(\{N_{d}\}) = Mh_{0} + \sum_{d} N_{d}h_{d},$$

with $h_{0}$ the enthalpy per unit cell for the perfect (undefected) crystal.

### III. POINT DEFECTS

#### A. Defect-induced instabilities in Fe$_3$Al(-C)

Table I displays the GC point defect energies of Fe$_3$Al(-C), as obtained in LDA or GGA calculations possibly including spin polarization, together with the reference energies of the undefected supercells. At perfect stoichiometry (no point defect), in agreement with previous calculations,6 all DFT formalisms predict the stable phase for Fe$_3$Al to be $D_0_3$ except SGGGA, for which both structures have almost identical stabilities (within the 1 meV/atom uncertainty of the computational method). The situation is however modi-

the mixed Fe-Al and pure Fe faces. In the $L_1_2$ phase, C additions were also supposed to be located only at octahedral neighbors, site fully occupied in the $L_2$ phase, the C with atomic positions $C_0$ and $4_{d}$ Wyckoff positions were used in $D_0_3$ compounds such as NiAl$_3$ (Ref. 22). The latter (4a and 4b Wyckoff positions) were thus included in our study, this choice being found legitimate a posteriori (see below). Finally, for the Fe(-AlC) solid solution, a $3 \times 3 \times 3$ supercell (54 sites) with $5 \times 5 \times 5$ k grid was used. With a plane-wave cut-off energy equal to 500 eV throughout, the previous parameters ensured a 1 meV/atom convergence for all energy values. In order to get a zero local pressure around the defects, all calculations included atomic relaxations together with energy minimization with respect to the supercell size and shape. In magnetic calculations, the ground-state search also allowed the relaxation of spins (including inversion).
fied significantly by point defects, since the dispersion between the DFT schemes appears to be quite large, a fact that will be found to have important consequences in terms of stability.

The first information that can be inferred from Table I is the point defect structure of each compound around stoichiometry. However, in the L12 one, achieving these calculations was surprisingly found to be impossible except for the SGGA framework, which solely could yield reasonable results. In all other formalisms, trying to minimize $H$ [relation (2)] led to no solution, this function being unbounded with the numerical values in Table I. The only acceptable $T=0$ K defect structure of $L1_2$ Fe$_3$Al, obtained from SGGA calculations, is therefore that depicted in Fig. 2(a). The case of $D0_3$ similarly provided no solution in SLDA. Whereas the other three formalisms led to $T=0$ K results according to Fig. 2(b), both the LDA and GGA nonmagnetic calculations for this compound were ruled out by temperature effects (not shown for brevity). These unexpected trends were checked for this compound were ruled out by temperature effects.

The only acceptable $T=0$ K defect structure of $L1_2$ results according to Table I, which indicated: (i) in $D0_3$ no significant amendment and (ii) for $L1_2$ a strong sensitivity to the supercell size for Al antisites in the nonmagnetic formalisms, which however led to the same unphysical (S)LDA and GGA behavior. The net conclusion is thus, with the inadequacy of the (S)LDA and GGA formalisms for both $D0_3$ and $L1_2$ Fe$_3$Al, only the SGGA calculations provide a reasonable picture of these compounds around stoichiometry. Note that this conclusion could be reached only by examination of the point defect properties, which proves the importance of including point defects in atomic-scale investigations of intermetallic compounds.

From the underlying linear minimization scheme, it can easily be demonstrated that there exists a relation between these (S)LDA and GGA defect-induced instabilities and the negative (or close to zero) sum of GC enthalpies $h_{Al}^{Fe}+h_{Fe}^{Al}$ for antisite pairs in these DFT approaches (see Table I). More precisely, the independent point defect framework imposes that $h_{Al}^{Fe}+h_{Fe}^{Al}$ be positive in order to avoid unphysical behaviors such as spontaneous formation of exceedingly large amounts of antisite defects. In $L1_2$, the trend is already pronounced $<−1$ eV) in both nonmagnetic schemes. In $D0_3$, although these values are also slightly positive for nonmagnetic LDA and GGA (considering the favored Fe$_5$ sublattice), the instability actually arises at any temperature, as confirmed by low-temperature calculations (not displayed for.
brevity) showing unrealistic populations of thermal defects. The situation in SLDA is close to that obtained for $L1_2$. Therefore, the SLDA and GGA frameworks equally yield incorrect GC energies for the $D0_3$ phase. These instabilities were suggested, although not pointed out clearly, in a previous study using a nonmagnetic pseudopotential DFT approach with a mixed basis, with the latter work however providing no GC energies but rather formation energies. More recently, a similar situation was found in $B2$ FeCo, with spontaneous formation of antisites at stoichiometry if magnetism is neglected. Such a behavior was also detected in $Ni_2Al_3$: When modeling this compound with an embedded-atom potential, unphysical amounts of interstitial point defects were obtained, whereas correct values were provided by DFT calculations. $L1_2$ Fe$_3$Al however seems to be more critical, since its defect structure is dramatically sensitive to the DFT approach chosen. Finally, in contrast with previous results concerning $B2$ FeAl, we conclude that point defect calculations for Fe$_3$Al should therefore be performed in SGGA.

This failure of the (S)LDA and GGA approximations for Fe$_3$Al is a somewhat surprising result, revealed in the course of the (free-)energy minimization scheme of the compound with respect to its point defect populations. However, one can also get convinced of its validity in a simpler way, merely by considering the stoichiometric case at $T=0$ K. As long as the point defects can be regarded as roughly independent (this is probably true until a few percent of the sites are affected), its enthalpy can be uniformly reduced by creating pairs of antisites (hence at constant composition). This increase in the long-range disorder clearly violates the stability of the ordered structure. Of course, this unphysical process may not lead to complete disorder, since it should be modified when the defects begin to interact. Nevertheless, it is sufficient to demonstrate the inconsistency of the (S)LDA and GGA schemes for the $L1_2$ (and $D0_3$) compound. It should be noted that this unrealistic behavior concerns the spontaneous creation of disorder at constant composition, and it is therefore an intrinsic property (not depending on the overall equilibrium of the compound with the surrounding phases). Such a criterion of long-range-order stability against generation of composition-conserving point defects is quite general and is therefore widely used as a test of the validity of empirical potentials designed for ordered phases (see, for instance, Ref. 30). Its use up to now remains however scarce in the increasing area of ab initio calculations about intermetallics. However, such tests would probably be useful, at least for the most common intermetallics (among which are the iron and nickel aluminides).

Finally, the unphysical behavior of antisites in $L1_2$ Fe$_3$Al obtained in (S)LDA and GGA can be related to a strong tetragonal deformation of the defect supercell around Al$_{Fe}$, the latter being itself a hint of a structural instability in this compound. Such an instability indeed occurs, as illustrated in Fig. 3 in the GGA case, with the tetragonal Fe$_3$Al ($c/a \approx 0.688$) found to be more stable than the cubic $L1_2$ one by more than 0.05 eV/atom, the two variants being separated by a very low-energy barrier ($<0.01$ eV). On the contrary, in SGGA a tetragonal Fe$_3$Al unit cell recovers the cubic symmetry, showing that no such transition exists in this frame-

![FIG. 3. Energy path from $L1_2$ cubic to tetragonal Fe$_3$Al (nonmagnetic GGA calculations).](image1)

![FIG. 4. Stability domains of $L1_2$ and $D0_3$ Fe$_3$Al(-C) at $T = 0$ K according to SGGA calculations.](image2)
should be improved in priority on the Al-rich side of stoichiometry. Finally, when C is added, the stability domain of $D_0_3$ shrinks, showing the stabilizing effect of this element for $L_1_2$. This suggests a C-induced stabilization of the fcc underlying structure and corroborates the preference for such a structure when the C content is increased up to the $E2_1$ phase.

**B. Fe$_3$C(-Al) cementite and Fe(-Al,C) solid solution**

The other binary ordered compound of interest in our work is the $D_0_1$ Fe$_3$C(-Al) cementite. This phase, known as slightly metastable, must nevertheless be taken into account in practice, due to its easy formation in the various thermo-mechanical processes. For the present purpose, in a way in practice, due to its easy formation in the various thermo-mechanical processes. For the present purpose, in a way similar to that obtained for C in the $L_1_2/D_0_3$ competition in Fe$_3$Al, we are interested in a possibly stabilizing influence of Al on this phase [guided by the existence of such a influence for Cr (Ref. 9)]. Following the same procedure as the previous one, the point defect structure of Fe$_3$C(-Al) was therefore determined, and the results are shown in Fig. 5 and Table II. The trends are quite clear, since all DFT formalisms yield similar behaviors (only SGGA is shown). On the whole, Fe$_3$C(-Al) appears to be a vacancy-type compound on the Fe-rich side of stoichiometry, while C excess is accommodated by 4$a$ interstitials. Al preferentially occupies Fe$_2$ sites, with no influence of the composition ($x_C=1/4$ isoplet). When the temperature rises significantly (for example, $T=1000$ K in Fig. 5), the secondary defects become important for C-rich compositions, whereas the Fe-rich defect structure remains simple, and Fe$_3$C(-Al) therefore exhibits a dissymetric point defect behavior.

Before turning to the $\kappa$ compound, let us briefly estimate the properties of the solid solution of Al and C in bcc iron, as the latter will also be used in Sec. IV. Consistency with the previous studies of ordered compounds implies treatment of this system within the independent point defect approximation, and in this framework, only the iron vacancy $V_{Fe}$, aluminum antisite $Al_{Fe}$, and octahedral carbon $C_{octa}$ were taken into account, with respective GC enthalpies of 10.460, 3.820, and $−8.557$ eV (the reference value being $h_0/2 = −8.309$ eV/atom). Whatever the composition (low amounts of Al and C), the structural point defects were necessarily $Al_{Fe}$ and $C_{octa}$; the corresponding $T=0$ K chemical potentials read therefore as

$$\mu_{Al} = \frac{h_0}{2} + \frac{h_{Fe}}{Al}, \quad \mu_{C} = h_{C_{octa}}, \quad \mu_{Fe} = \frac{h_0}{2}. \quad (3)$$

**C. Interstitials in the $\kappa$ Fe$_3$AlC compound**

Completing the point defect study, the reference and GC point defect energies for Fe$_3$AlC, obtained with LDA or GGA calculations including or not including spin polarization, are displayed in Table III. Although all frameworks show qualitative agreement, significant discrepancies occur, mainly for interstitial carbon, vacancies, and Al antisites. As before, no direct conclusion on the stable point defects can be drawn from the GC energies, since the defects are associated with local composition changes that need to be in-

**Table II.** Reference (eV/atom) and GC energies (eV) of point defects in $D_0_1$ Fe$_3$C(-Al) cementite, calculated in the (S)LDA and (S)GGA frameworks (32-atom $2\times2\times2$ supercell).

<table>
<thead>
<tr>
<th>Ref.</th>
<th>$V_{Fe_1}$</th>
<th>$C_{Fe_1}$</th>
<th>$V_{Fe_2}$</th>
<th>$C_{Fe_2}$</th>
<th>$V_{Fe}$</th>
<th>$C_{FeC}$</th>
<th>$Al_{Fe_1}$</th>
<th>$Al_{Fe_2}$</th>
<th>$Al_{C}$</th>
<th>$C_{4a}$</th>
<th>$C_{4b}$</th>
</tr>
</thead>
</table>
TABLE III. Reference (eV/atom) and point defect GC energies (eV) in $E_2$, $Fe_3AlC$, calculated in the (S)LDA and (S)GGA frameworks (40-atom $2 \times 2 \times 2$ supercell).

<table>
<thead>
<tr>
<th>Ref.</th>
<th>$V_{Al}$</th>
<th>$Fe_{Al}$</th>
<th>$C_{Al}$</th>
<th>$V_{Fe}$</th>
<th>$Al_{Fe}$</th>
<th>$C_{Fe}$</th>
<th>$V_{C}$</th>
<th>$Al_{C}$</th>
<th>$Fe_{C}$</th>
<th>$C_{octa}$</th>
<th>$C_{intra}$</th>
</tr>
</thead>
</table>

included globally in the analytic treatment [Eq. (2)]. Table III nevertheless shows the marked preference of C for octahedral sites, since $E_{GC}^{Al} (C_{octa}) - E_{GC}^{Al} (C_{tetra}) = -2$ eV whatever the DFT framework used. As regards the $T = 0$ K point defect structure, the minimization procedure naturally yields zone limits of two types: $x_i = 1/5$ or $3/5$ and $x_i = 3x_j$ (with $i$, $j$ = Al, Fe, or C), corresponding to sharp changes in the point defect nature, a well-known effect in ordered compounds, possibly inducing drastic changes in the properties for slight composition changes. This effect, already noted for additions in binary ordered systems, is also present in intrinsically ternary ones. The role of interstitial C in the compound is not easy to infer intuitively. From a crystallographic point of view, both the central and the octahedral interstitial positions in $Fe_3AlC$ are octahedral sites, the difference merely lying in the chemical nature of the neighboring atoms [pure Fe and mixed (Fe,Al), respectively]. As pointed out in Ref. 1, systematic studies of $F_4MC$ compounds (with $T$ = transition metal and $M = metal \neq T$) indicated that C occupies only pure Fe octahedra, its insertion into a mixed octahedron leading to an energetically unfavorable distortion. Such a hypothesis, conceiving the perovskite as a binary Fe-Al alloy with a posteriori additions of C, provides an estimation of the C solubility limit, related to the number of available Fe octahedra in the Fe-Al underlying structure. It cannot however be rigorously justified, since the equilibrium phase structure results from a unique energy minimization for the ternary compound (in equilibrium with other candidate phases) with respect to its internal variables. On the whole, there exists no convincing argument to justify interstitials being neglected a priori in the $\kappa$ phase nor as regards the respective roles of octahedral and tetrahedral sites.

However, in order to evaluate explicitly the sensitivity of the results to interstitial C, it is instructive to perform at first an “interstitial-free” calculation of the $T = 0$ K point defect structure. Figures 6(a) and 6(b) show the conclusions of such a calculation: Neglecting interstitial C leads to somewhat dispersed results, the SGGA here again behaving differently from the other frameworks. The ambiguity occurs essentially in the composition domain corresponding to $x_{Al} < 1/5$ and $x_C > 1/5$, with SGGA yielding Al vacancies and a $x_C = 3x_C$ zone limit, whereas all other calculations clearly point out C defects with a $x_C = 3/5$ limit. Thus, although the C-depleted off-stoichiometry is, whatever the DFT formalism, coherently predicted to be accommodated by $Al_{C}$, $Fe_{C}$, and $V_{C}$ (with changes at $x_{Fe} = 3x_{Al}$, $x_{Al} = 1/5$, and $x_{C} = 1/5$), disagreement between the DFT formalisms occurs in the C-rich part of the composition diagram.

Including interstitial C (in octahedral sites) dramatically clarifies the situation [Fig. 6(c)], the $T = 0$ K point defect structure being depicted then more coherently: Only four composition domains appear, delimited by the $x_{Fe} = 3x_{Al}$ and $x_C = 1/5$ isoplets, and the off-stoichiometry leads to $Al_{C}$, $Fe_{C}$, $V_{C}$, and $C_{octa}$ point defects. This clearly shows the importance of including interstitial C (in octahedral sites) to ensure a proper description of $Fe_3AlC$. At moderate temperature, the latter should thus be modeled reasonably using a four-defect model with the formula $(Fe,Al)_1(Al,Fe)C(C_{vacancy})_{octa}$, refining the currently used three-sublattice description of $\kappa$. In the present case, due to the relative simplicity of the cubic underlying crystallographic structure, the role of interstitial C could perhaps have been expected from intuitive arguments. However, other cases (for example, $Fe_2C$) may be encountered for which no such heuristic guidelines exist, and a correct initial defect identification would then be critical.

The knowledge of the $T = 0$ K point defects in each phase and each composition domain eventually makes it possible to estimate analytically the low-temperature chemical potentials of the three elements, which show quite different behaviors (Table IV). In both $Fe_2Al$-(C) structures, the C chemical potential is independent of the composition (being simply equal to the GC energy of the interstitial defect), whereas this parameter has an influence on the potentials of both other (intrinsical) elements. In cementite, a similar behavior is noticed for the intrinsic chemical potentials, together with that of Al, which is sensitive to the composition (due to its substitutional occupancy). In $Fe_3AlC$, the C chemical potential is

![FIG. 6. $T = 0$ K domains of stability of point defects in $Fe_3AlC$ as a function of the DFT formalism used, with (a) and (b) neglecting and (c) including the octahedral occupancy for C.](image-url)
sensitive only to the atomic fraction of this element, while those of both other elements show a more complex behavior and vary when crossing each zone limit. These analytic expressions for the chemical potentials will be used in Sec. IV, enabling numerical estimations of the low-temperature enthalpy per atom \( H_e = \sum_i x_i \mu_i \) of the phases in each composition domain.

Finally, since the GC energies and chemical potentials together give access to the defect formation energies (more readily comparable with experiments) at \( T=0 \) K, the latter quantities are also provided in Table V (in the SGGA scheme) for the ordered compounds considered in the present work [namely, \( D0_3 \) and \( L1_2 \) Fe₅Al-C, \( D0_{11} \) Fe₅C-Al, and \( E2_1 \) Fe₅AlC]. As expected, the structural defects in each relevant composition domain have zero formation energies, and those with low positive values should be formed easily by thermal activation. Also note the large values occurring in Fe₅AlC, which recall those obtained in nonmetallic systems (oxides, etc.).

### IV. LOW-TEMPERATURE ANALYSIS OF THE Al-C-Fe SYSTEM

Relying on the previous point defect studies, we then proceed to an analysis of the possible low-temperature equilibrium between these phases. First of all, it should be recalled that the previous conclusions exclude the use of any formalism except SGGA. The latter will consequently constitute the only framework of the following discussion, but this does not dismiss the other formalisms which may perform better than the SGGA in some cases (for instance, SLDA in perfect Fe₅Al). In fact, it might also be conceivable to devise mixed (S)LDA/(S)GGA schemes, using distinct DFT calculations for the different phases. However, we did not retain such a procedure, by reason of its significant level of arbitrariness and also because it obviously precludes any assessment of the merits of a single DFT framework.

In Ref. 2, the \( L1_2 \) and \( E2_1 \) compounds were simultaneously treated as a continuous solution of \( C \), and the switch-

### TABLE IV. Chemical potentials at \( T=0 \) K in Fe₅Al(-C) (\( D0_3 \) and \( L1_2 \)), Fe₅C(-Al) and Fe₅AlC, calculated in the SGGA framework.

<table>
<thead>
<tr>
<th>Phase</th>
<th>Composition range</th>
<th>( \mu_{Al} )</th>
<th>( \mu_{C} )</th>
<th>( \mu_{Fe} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( D0_3 ) Fe₅Al-C</td>
<td>( x_{Al} &lt; x_{Fe}/3 )</td>
<td>( h_0/16 - 3/4 h_{Al}^{Fe} )</td>
<td>( h_{C}^{Fe} )</td>
<td>( h_0/16 + h_{Al}^{Fe}/4 )</td>
</tr>
<tr>
<td></td>
<td>( x_{Al} &gt; x_{Fe}/3 )</td>
<td>( h_0/16 + 3/4 h_{Al}^{Fe} )</td>
<td>( h_{C}^{Fe} )</td>
<td>( h_0/16 - h_{Al}^{Fe}/4 )</td>
</tr>
<tr>
<td>( L1_2 ) Fe₅Al-C</td>
<td>( x_{Al} &lt; x_{Fe}/3 )</td>
<td>( (h_0 - 3h_{Al}^{Fe})/4 )</td>
<td>( h_{C}^{Fe} )</td>
<td>( (h_0 + h_{Al}^{Fe})/4 )</td>
</tr>
<tr>
<td></td>
<td>( x_{Al} &gt; x_{Fe}/3 )</td>
<td>( (h_0 + 3h_{Al}^{Fe})/4 )</td>
<td>( h_{C}^{Fe} )</td>
<td>( (h_0 - h_{Al}^{Fe})/4 )</td>
</tr>
<tr>
<td>( D0_{11} ) Fe₅C-Al</td>
<td>( x_{C} &gt; 1/4 )</td>
<td>( h_0/12 - h_{C}^{Fe}/3 + h_{C}^{Fe}/4 )</td>
<td>( h_{C}^{Fe} )</td>
<td>( h_0/12 - h_{C}^{Fe}/3 )</td>
</tr>
<tr>
<td></td>
<td>( x_{C} &lt; 1/4 )</td>
<td>( h_0/12 + h_{C}^{Fe}/3 + h_{C}^{Fe}/4 )</td>
<td>( -h_{C}^{Fe} )</td>
<td>( h_0/12 + h_{C}^{Fe}/3 )</td>
</tr>
<tr>
<td>( E2_1 ) Fe₅AlC</td>
<td>( x_{C} &gt; 1/5; x_{Al} &lt; x_{Fe}/3 )</td>
<td>( (h_0 - 3h_{Al}^{Fe} - h_{C})/4 )</td>
<td>( h_{C}^{Fe} )</td>
<td>( (h_0 + h_{Al}^{Fe} - h_{C})/4 )</td>
</tr>
<tr>
<td></td>
<td>( x_{C} &lt; 1/5; x_{Al} &lt; x_{Fe}/3 )</td>
<td>( (h_0 + 3h_{Al}^{Fe} + h_{C})/4 )</td>
<td>( -h_{C}^{Fe} )</td>
<td>( (h_0 - h_{Al}^{Fe} + h_{C})/4 )</td>
</tr>
</tbody>
</table>

### TABLE V. \( T=0 \) K point defect formation energies (eV) in the ordered compounds encountered in this work (SGGA framework throughout).

<table>
<thead>
<tr>
<th>Phase</th>
<th>Composition range</th>
<th>( \mu_{Al} )</th>
<th>( \mu_{Fe} )</th>
<th>( \mu_{C} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( D0_3 ) Fe₅Al-C</td>
<td>( x_{Al} &lt; x_{Fe}/3 )</td>
<td>1.904</td>
<td>4.002</td>
<td>0.971</td>
</tr>
<tr>
<td></td>
<td>( x_{Al} &gt; x_{Fe}/3 )</td>
<td>2.489</td>
<td>3.806</td>
<td>0.190</td>
</tr>
<tr>
<td>( L1_2 ) Fe₅Al-C</td>
<td>( x_{Al} &lt; x_{Fe}/3 )</td>
<td>1.769</td>
<td>1.201</td>
<td>1.201</td>
</tr>
<tr>
<td></td>
<td>( x_{Al} &gt; x_{Fe}/3 )</td>
<td>2.670</td>
<td>1.201</td>
<td>1.604</td>
</tr>
<tr>
<td>( D0_{11} ) Fe₅C-Al</td>
<td>( x_{C} &gt; 1/4 )</td>
<td>1.467</td>
<td>2.297</td>
<td>1.936</td>
</tr>
<tr>
<td></td>
<td>( x_{C} &lt; 1/4 )</td>
<td>1.913</td>
<td>4.082</td>
<td>1.732</td>
</tr>
<tr>
<td>( E2_1 ) Fe₅AlC</td>
<td>( x_{C} &lt; 1/5; x_{Al} &lt; x_{Fe}/3 )</td>
<td>1.191</td>
<td>7.003</td>
<td>1.944</td>
</tr>
<tr>
<td></td>
<td>( x_{C} &lt; 1/5; x_{Al} &gt; x_{Fe}/3 )</td>
<td>3.519</td>
<td>9.331</td>
<td>3.103</td>
</tr>
<tr>
<td></td>
<td>( x_{C} &gt; 1/5; x_{Al} &lt; x_{Fe}/3 )</td>
<td>0.306</td>
<td>2.580</td>
<td>1.060</td>
</tr>
<tr>
<td></td>
<td>( x_{C} &gt; 1/5; x_{Al} &gt; x_{Fe}/3 )</td>
<td>2.634</td>
<td>3.103</td>
<td>4.907</td>
</tr>
</tbody>
</table>
Fe₃AlC. It may however help to understand a surprising fea-
ture in the behavior of C when going from Fe₃Al
ichiometry, cannot therefore give information about the evo-

duced by the possibility of
diagrams are presented in Fig. 7, showing only three-

The effect of off-stoichiometry is examined in the second
step, by superposing onto \( H_{\text{stochio}} \) the \( (H_{\text{st}}=x_{\text{C}}\mu_{\text{C}}) \) surfaces of the
phases obtained via the previously described point defect
analysis, which requires the chemical potentials of
Fe₃Al(C), Fe₃C(-Al), and Fe₃AlC (Table IV). Figure 8
displays the convex hull of the three-phase stoichiometric
mixtures (exhibiting an almost flat shape), together with the
superposition of these nonstoichiometric compounds. The

Using the present methodology, assessing the effect of
off-stoichiometry cannot be undertaken directly, due to the
nonderviability of the enthalpies (as a function of composition) at \( T=0 \) K, which prevents application of
the usual rule of common tangent planes. This difficulty can be overcome
by means of a two-step analysis, the first of which consists in
determining the “stoichiometric convex hull” associated with
the phases considered, namely, the Fe₃AlC -C
-Fe₃AlC_{1-x}+C_{\text{graphite}}ardo the experimental maximum C solubility of 15% at 1000 K.1

As regards graphite C, the value \( h_{\text{C}}=0 \) eV/atom (obtained from SGGa calculations) was used. As regards graphi-
te, owing to the doubtful value provided by DFT, its energy
was calculated by adding the experimental difference
\( \Delta E = \Delta E_{\text{expt}} - \Delta E_{\text{DFT}} \) to the DFT values
obtained for diamond, yielding \( \Delta E = -9.112 \) eV/atom for (S)GGA. The ideal stoichiometric sta-

FIG. 7. (Color online) Stability domains of three-phase systems
involving stoichiometric compounds at \( T=0 \) K (magnetic GGA calculations).

FIG. 8. \( T=0 \) K enthalpy-composition diagram for the phases
considered in this work (bcc, \( D_{01}, D_{011}, \) and \( E_{21} \)), together with
the enthalpy of the mixtures of the three stoichiometric phases
(stoichiometric convex hull) corresponding to Fig. 7 (magnetic
GGA calculations).

FIG. 9. (Color online) Cut of the \( T=0 \) K enthalpy-composition
diagram in Fig. 8 along the \( x_{\text{Fe}}=3x_{\text{Al}} \) line, for comparison of the
stoichiometric mixtures and \( E_{21} \) Fe₃AlC.
mains when $T$ increases, and cementite shows an intermediate behavior, with similar slopes for C depletion and excess. Our results also provide a rough interpretation of the experimental symmetric behavior of $\kappa$ (according to whether $x_C < 0.2$ or not) by considering the influence of the C content on the relative positions of $H_{\text{stoich}}$ and $H_e$. Figure 9, which corresponds to a cut of Fig. 8 along the $x_C=0.3x_{\text{Al}}$ line, thus indicates that $H_e-H_{\text{stoich}}$ is larger for $x_{\text{Fe}}<0.6$ (that is, for $x_C>0.2$), which is coherent with a poorer accommodation of off-stoichiometry and, thus, an increased trend to unmixing in this composition domain. In agreement with experimental conclusions, $\kappa$ therefore appears as preferentially understoichiometric in C, even at low temperature.

Finally, the question arises as to whether this phenomenon is related to interstitial C. As already noted in Fig. 6, the role of interstitials is limited to the C-rich domain, which implies that for $x_C<1/5$ the $\kappa$ enthalpies with and without interstitial C are identical. Moreover, for $x_C>1/5$ the interstitial-free curve is necessarily above that with interstitial (since the latter structure is the most stable one). It follows that neglecting interstitial C should lead to the same conclusion as regarding phase separation. This may partly explain why three-sublattice (hence ignoring interstitial C) approaches may succeed in predicting such a trend.2

V. CONCLUSION

The thermodynamic properties of complex alloys and their equilibria strongly depend on their point defects, which constitute the elementary low-temperature excitations governing the free energy. The energetics of these defects can be conveniently obtained using the available \textit{ab initio} methods, which now offer the possibility of tackling multiconstituent and multistructured mixtures provided a tractable thermodynamic framework is employed. The independent point defect approximation used here seems to be well designed for this task, since the present study has demonstrated its capability of yielding realistic trends for Al-C-Fe; several major results obtained this way are worth emphasizing.

First, only the SGGA point defect calculations give correct results for $L1_2/D0_3$, Fe$_3$Al, since all other formalisms entail an unstable $L1_2$ phase with respect to antisite defect formation. This behavior is probably related to the strong structural relaxation around $A_{\text{Fe}}$ that can be traced back to the existence of a tetragonal phase predicted to be more stable than the cubic one. Carbon stabilizes the $L1_2$ structure, suggesting the existence of a continuous connection between the $L1_2$ Fe$_3$Al(-C) solid solution and the $\kappa$ phase; this may help to understand why Fe$_3$AlC has an $L1_2$ rather than a $D0_3$ related structure.

The study of relative stability between $D0_3$ and $L1_2$ Fe$_3$Al(-C) points out the intrinsic difficulties of the DFT methods in describing the Fe-Al system. These particular phases and their point defect properties may constitute a benchmark to check possible improvements in electronic structure calculations. The point defect study of the various phases investigated also evidences the importance of taking into account interstitial carbon in Fe$_3$C and Fe$_3$Al in describing carbon-rich phases, although such phases may not appear in the equilibrium phase diagram.

By constructing $H_{\text{stoich}}$ the stoichiometric convex hull associated with the considered phases allows us to draw a $T=0$ K phase diagram made of three-phase coexistence domains. A cut of the enthalpy surfaces displays trends that indicate that the Fe$_3$AlC kappa compound should be carbon depleted at equilibrium, as experimentally observed. The shapes of the enthalpy-composition surfaces are quite different: flat for Fe$_3$Al(-C) and Fe(-Al,C), intermediate for Fe$_3$C(-Al), and sharp for Fe$_3$AlC.

To summarize, beyond the particular case presented here, we more generally believe that the thermodynamic analysis relying on the independent point defect approximation is a powerful tool in spite of its simplicity. It allows handling multicomponent phases with general crystallographic structures. It therefore constitutes a soundly based and convenient approach to address thermodynamics at the atomic scale, before possibly resorting to more powerful but much heavier methods.