



Article Crystal Chemistry and Electronic Properties of the Al-Rich Compounds, Al₂Cu, ω-Al₇Cu₂Fe and θ-Al₁₃Fe₄ with Cu Solution

Changming Fang *[®], Maaouia Souissi, Zhongping Que and Zhongyun Fan

Brunel Centre for Advanced Solidification Technology (BCAST), Brunel University London, Uxbridge UB8 3PH, UK; maaouia.souissi@brunel.ac.uk (M.S.); zhongping.que@brunel.ac.uk (Z.Q.); zhongyun.fan@brunel.ac.uk (Z.F.)

* Correspondence: changming.fang@brunel.ac.uk

Abstract: In this work, we investigate Cu solution in θ -Al13Fe4 and related Al-rich ω -Al7Cu2Fe and Al2Cu phases in the Al-Cu-Fe system using the first-principles density functional theory (DFT) with on-site Coulomb interaction correction. The results show preference of Cu at Al7, forming a ternary θ -Al76Cu2Fe24 at ambient conditions, and both Al7 and Al9 sites (in Grin's note), forming θ -(Al76–xCu2+x)Fe24 at a high temperature. The relative stability of the Al-rich compounds and their crystal and electronic properties are investigated. We show the importance of the Hubbard U correction to the standard DFT functionals for Cu-containing metallic materials. This study helps characterize the intermetallic compounds in Cu-containing Al alloys, and helps further control Fe-containing intermetallic compounds in the solidification of Al-based alloys.

Keywords: Cu solution in θ -Al13Fe4; Cu-containing compounds; crystal chemistry; on-site coulomb correction; density functional theory

1. Introduction

Fe is the most common impurity in commercial Al metals [1,2]. Due to its low solubility in Al, Fe exists in the form of Fe-containing intermetallic compounds (Fe-IMCs), including the primary θ -Al₁₃Fe₄. These Fe-IMCs deteriorate the mechanical performance of cast Al-based parts [1–5]. Al-Cu alloys, including the 2000- and 800-series, have attractive advantages: being light-weight, high strength, and age-hardening. Thus, they have been widely utilized in aerospace and automobile industries [1,5–7]. Information about the Fe-IMCs, including θ -Al₁₃Fe₄ and the AlCu precipitates, Al₂Cu and ω -Al₇Cu₂Fe, in the Al-rich part of the Al-Fe-Cu system is of importance to achieve the cast parts of fine microstructure and desirable properties. Moreover, the recycling economy requires that harmful Fe-IMCs, including θ -Al₁₃Fe₄, be minimized or at least controlled in the products during the casting of Al alloys, especially Al scrap, which contains various impurities, including Fe [8–10]. In this respect, knowledge about the related compounds/precipitates and Cu solutions in θ -Al₁₃Fe₄ is of crucial importance.

Experimental and theoretical efforts have been made on the phase relations in the Al-related compounds and the structures of Fe-IMCs, including θ -Al₁₃Fe₄ [1,2,11–33]. Grin and co-workers determined the crystal structure of θ -Al₁₃Fe₄ [22] and showed that this compound exhibits rich crystal chemistry. It has a monoclinic lattice (space group C2/m, nr. 12) with lattice parameters, *a* = 15.492 Å, *b* = 8.078 Å, and *c* = 12.471 Å, β = 107.69° [22]. This cell contains 15 Al and 5 Fe species, and 102 atoms in total (78 Al and 24 Fe) (Grin's note) [22]. There is a family of this type of structure, θ -Al₁₃M₄ (M = Fe, Co, Ni, and Ru) [23,24]. Moreover, this structure is chemically flexible, and Fe/Al can be partially replaced to form ternary compounds [23,25,26]. As an example, the Cu solution in θ -Al₁₃Fe₄ has been under discussion [16–18,25,26]. Freiburg and Grushko investigated the chemical



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Copyright: © 2022 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). composition and structural model of θ -Al₁₃Fe₄ in the Al-Fe-Cu system and proposed Cu substitution at the Al7 and Al9 sites [25]. Genba and co-workers prepared Al-Fe alloys under Ar protection in a quartz (SiO₂) vessel using pure Al and Fe raw materials [26]. They first heated the sample at 850 °C and then cooled it to 750 °C by quenching it in water. They observed that many needle-like crystals appeared on the surface of the SiO₂ glass tube. Using the X-ray diffraction patterns analysis and electron probe microanalysis, they found Cu at the Fe1, Fe2, Fe3, Al9, Al12, and Al14 sites in θ -Al₁₃Fe₄ [26]. First-principles density functional approaches have been applied to investigate the structural, electronic, and magnetic properties of pure θ -Al₁₃Fe₄ [23,24,31,32]; its surfaces for catalysis [33,34]; as well as the intrinsic defects in this binary compound [35,36]. Recently, we investigated Si substitution in θ -Al₁₃Fe₄ [36,37]. Here, we investigate the Cu substitution in θ -Al₁₃Fe₄ using a density functional theory (DFT) method with on-site Coulomb interaction correction to Cu 3d electrons. We reveal the preference of Cu at special Al sites, forming Al₇₆Cu₂Fe₂₄ at ambient conditions.

The rest of this manuscript is arranged as follows. We first introduce the computational technique and settings in Section 2. Then, we discuss the effect of on-site Coulomb interaction correction on the cohesive properties of the elemental solid Cu and related solids in Section 3.1. This helps us to choose the density functional for future work. The energetics, phase relations, and structural and electronic properties of the Al-rich compounds, θ -Al₁₃Fe₄, θ - and θ' -Al₂Cu, and ω -Al₇Cu₂Fe are addressed in Section 3.2. Cu substitution at the atomic sites in θ -Al₁₃Fe₄ is described in Section 3.3. We present the crystal structure of the stable θ -Al₇₆Cu₂Fe₂₄ in Section 3.4. We compare the theoretical calculations with the experimental observations in Section 4. The results are summarized in Section 5.

2. Methods

We utilized the first-principles' Vienna Ab initio Simulation Package (VASP) [38,39]. This code employs the density functional theory (DFT) within the Projector Augmented Wave (PAW) approach [40]. The spin-polarized Generalized Gradient Approximation (GGA-PBE) [41] was applied for the exchange and correlation energy terms because the GGA approximations describe the 3d metals such as iron, Cu, and related compounds better than the Local Density Approximation (LDA) [42]. We employed a cut-off energy of 550 eV for the wave functions and 700 eV for the augmented wave functions, which better describe the localized 3d electrons of the transition metals. These values are higher than the default values ($E_{MMAX}/E_{AUG} = 240.3/291.1$ eV for Al, 267.9/511.4 eV for Fe, and 295.4/587.0 eV for Cu). The electronic wave functions were sampled with a dense, $4 \times 8 \times 6$ grid (70–100 *k*-points) in the irreducible Brillouin zone (BZ) of θ -Al₁₃Fe₄ and the related compositions depending on the symmetry, using the Monkhorst-Pack method [43]. First-principles structural optimizations were performed for both lattice parameters and the coordinates of the atoms. Different *k*-meshes and cut-off energies for the wave functions and augmentation wave functions were tested, which showed suitable convergence (<1 meV per atom).

3. Results

First, we report the calculations for the bulk elemental solids, body-centered cubic (BCC) α -Fe and face-centered cubic (FCC) Al [44], using the DFT-GGA approach with the settings mentioned above. The calculations produced a lattice parameter a = 2.831 Å for ferro-magnetic Fe, and 4.040 Å for Al. The calculated lattice parameters are in agreement with the available experimental values at 0 K, a = 2.8607 Å for Fe and 4.0325 Å for Al [45], as shown in Table 1. The calculations produced a magnetic moment of 2.18 $\mu_{\rm B}$ /Fe, which reproduced the previous results [36,37] and is close to the experimental value 2.10 $\mu_{\rm B}$ /Fe [36,46].

3.1. Effect of on-Site Coulomb Repulsion Correction on Cohesive Properties of Cu

The electron correlation in narrow energy bands was recognized by Hubbard [47]. Based on a Green function technique, Hubbard proposed a simplified model to correct this effect for localized orbitals (Hubbard U) [47]. This approach has been successfully applied to describe, e.g., compounds containing 3d transition metals [48,49]. Recently, we [50] performed calculations using the DFT + U approach and revealed U = 4 eV based on matching the calculated energy position of Cu 3d peak to the photoemission measurements [50,51]. This U value for the occupied Cu 3d electrons is the same as that used in the oxides containing Cu [49] and the systematic study for 3d transition metals using a constrained random-phase approximation [52].

Here, we introduce in detail the cohesive properties of Cu using the standard DFT functionals (LDA and GGA) with/without the Hubbard U (=4 eV) correction. The cohesive energy of an elemental solid is defined as:

$$E_{\rm coh.} = E({\rm solid}) - E({\rm atom}) \tag{1}$$

Here, *E*(solid) and *E*(atom) represent the calculated energies for the solid and isolated atom, respectively. The calculated results are listed in Table 1 compared with selected data from previous calculations and experiments.

One issue to obtain cohesive energy of an elemental solid is the ground state of the individual atom, E(atom) in Equation (1). We performed calculations for the individual atoms with different spin-polarization values in a cube with an axis length of 21 Å. The analyses show that all elemental atoms are spin-polarized and have an integral number of moments: $1 \mu_B/\text{atom}$ for Al and Cu, corresponding to their one unpaired 3 s and 4 s electron, respectively (Table 1). Fe atom has a complex electronic configuration with partially occupied 3d orbitals. We performed the atomic energy with different spin-polarization values, including fractional occupations. The results are plotted in Figure 1. An isolated iron atom has an integral spin-polarization number of $4 \mu_B/\text{Fe}$. Based on the calculations, cohesive energies of the elemental solids are obtained according to Equation (1). The available experimental values are included in Table 1 for comparison.



Figure 1. (a) (Color on line) The calculated energy on magnetic moment for an individual Fe atoms using the standard density functional, LDA and GGA. (b) The effect of on-site Coulomb interaction (U = 4 eV) on the Cu 3d states using both GGA (top) and LDA (bottom).

Table 1 shows the calculated results using different density functionals. The DFT-GGA approximation predicts the lattice parameters of the elemental solids, Al and Fe, with deviations within 1%. Meanwhile, DFT-LDA underestimates the lattice parameters of the elemental solids, especially for iron. Moreover, the estimated cohesive energies of the elemental solids from the DFT-GGA are in much better agreement with the available

experimental values in the literature than those from the DFT-LDA, which agrees with the previous work that DFT-GGA works better for solids, especially for transition metals and their compounds [41,42].

Table 1. Calculated cohesive properties (lattice parameter, cohesive energy) for elemental solids Al, Fe, and Cu using both DFT and DFT+ U methods (U = 4 eV) with comparison with available experimental values and previous calculations in the literature.

Property	LDA	LDA + U	GGA	GGA + U	Exper. (0 K)	
Al: Face-Centered Cubic (FCC) lattice. Electronic configuration, [Ne] 3s ² 3p ¹						
<i>a</i> (Å)	3.984 (-1.2%)	-	4.039 (+0.2%)	-	4.0325 [45]	
$E_{\rm coh}$ (eV/Al)	-4.020 (+15.7%)			-	-3.39 [46,53]	
α -Fe: Body-Centered Cubic (BCC) lattice. Electronic configuration, [Ar] 3d ⁶ 4s ²						
<i>a</i> (Å)	2.747 (-4.0%)	-	2.831 (-1.0%)	-	2.8607 [45]	
$E_{\rm coh}$ (eV/Fe)	-6.480 (+51.4%)	-	-4.933 (+15.3%)	-	-4.28 [46]	
Cu: Face-Centered Cubic (FCC) lattice. Electronic configuration, [Ar] 3d ¹⁰ 4s ¹						
a (Å)	3.523 (-2.2%) 3.553 (-1.4%) [46]	3.496 (-2.9%)	3.635 (+0.9%) 3.630 (+0.7%) [30] 3.676 (+2.0%) [46]	3.622 (+0.6%)	3.6032 [45]	
$E_{\rm coh}$ (eV/Cu)	-4.676 (+34.4%) -4.353 [46]	-4.288 (+23.2%)	-3.501 (+0.6) -3.48 [30]	-3.079 (-11.5%)	-3.48 [30] -3.524 [54]	
<i>E</i> (3d) (eV)	-1.78	-2.49	-1.65	-2.34	-2.4 (UPS) [51] -3.2 (XPS) [51]	

Calculations using both LDA and LDA + U approaches markedly underestimate the lattice parameter of Cu (over 2%) (Table 1). Moreover, the calculated cohesive energies are notably higher than the experimental values. The DFT-GGA calculations produced lattice parameters in agreement with the experimental value (deviation of 0.9%), and the GGA + U calculations resulted in an even better agreement with the experiment (0.6%). The value of the cohesive energy from GGA using the current settings is slightly higher than the experimental values (~4%). The GGA + U calculations produced cohesive energy of about -3.078 eV/Cu in Al, which is notably smaller than the GGA calculations. The experimental values are in between the GGA and the GGA + U calculations. However, this GGA + U value is close to the value -3.046 eV/Cu from the advanced hybrid functional approach [30,55,56]. This is understandable, that the GGA + U approximation corrects the unphysical interaction between the 3d electrons and the neighboring Cu atoms, similar to that in the advanced hybrid functionals [55,56]. This result also indicates that Cu 3d electrons are better treated as a semi-core level.

The Hubbard U correction puts the Cu 3d states about 0.7 eV lower in energy for both LDA(+U) and GGA(+U) methods (Table 1 and Figure 1b), in agreement with the recent study [50]. The peak of the Cu 3d states is about 2.3 eV below the Fermi level, close to the experimental value from ultra-violet photoelectron spectrum (UPS) observations (2.4 eV) [51].

Overall, the DFT-GGA predicts the cohesive properties of the elemental solids, especially the 3d transition metals, notably better than the DFT-LDA. The Hubbard U approach corrects the unphysical interaction of the Cu 3d electrons with the neighboring atoms and thus produces less cohesive energy. In the rest of this work, we only use the DFT + GGA with the Hubbard U correction for the Cu-containing compounds. 3.2. The Al-Rich Compounds θ -Al₁₃Fe₄, Al₂Cu, and ω -Al₇Cu₂Fe in the Al-Fe-Cu System

The formation energy of, e.g., a ternary compound $A_l B_m C_n$ with respect to the elemental solids A, B, and C is defined as,

$$\Delta E(\mathbf{A}_l \mathbf{B}_m \mathbf{C}_n) = E(\mathbf{A}_l \mathbf{B}_m \mathbf{C}_n) - [l \ E(\mathbf{A}) + m \ E(\mathbf{B}) + n \ E(\mathbf{C})]$$
(2)

where $E(A_lB_mC_n)$ and E(A), E(B), and E(C) are the calculated total-electron energies for $A_lB_mC_n$ and the elemental solids A, B, and C, respectively.

The computed results (lattice parameters and formation energies) for the Al-rich compounds in the Al-Cu-Fe system are listed in Table 2. Available experimental results are included for the sake of comparison. The schematical structures and related coordination of the transition metals are shown in Figure 2.

The calculated lattice parameters for the compounds are in agreement with the available experimental data within 1%, except for the a-axis of θ' -Al₂Cu, the calculated value of which is about 1.3% greater than the experimental data.

Table 2. Calculated results (lattice parameters, formation energies according to Equation (2)) for the stable structure in the binary and ternary systems at ambient conditions. Experimental data in the literature are included for comparison.

Compounds	Calculated Latt. Para. (Å)/Form. Energy		Fyneriments		
Compounds	GGA	GGA + U			
θ-Al ₁₃ Fe ₄ , Mon. C2/m, (nr. 12)	a = 15.426 b = 8.022 c = 12.425 $\beta = 107.68 (^{\circ})$ $V = 1464.92 (Å^{3}/cell)$		a = 15.492 b = 8.078; c = 12.471 $\beta = 107.69(^{\circ});$ $V = 1486.88(Å^3/cell) [22]$		
θ-Al ₂ Cu, Tetr. I4/mcm (Nr.140)	$\Delta E_{form} = -0.329 \text{ eV} / \text{atom}$ a = 6.066 c = 4.874 V = 179.36 $\Delta E_{form} = -0.157$	6.064 4.864 178.88(Å ³ /cell) -0.125(eV/atom)	$\Delta E_{form} = -0.225 \text{ to } -0.310 \text{ eV}/\text{ atom } [38-40]$ a = 6.054 [57]; 6.063 [58]; 6.067 [59,60] c = 4.864 [57]; 4.872 [58]; 4.877 [59,60] $V = 178.27 (\text{Å}^3/\text{cell})$ [57], 179.09 [58], 179.52 [59,60]		
θ'-Al ₂ Cu, Tetr. I-4m2 (Nr.119)	a = 4.093 c = 5.786 V = 96.94 $\Delta E_{form} = -0.176$	4.092 5.782 96.81(Å ³ /cell) -0.124 (eV/atom)	a = 4.04 [60] c = 5.80 [60] V = 94.67 (Å ³ /cell) [60]		
ω-Al ₇ Cu ₂ Fe, Tetr. I-4m2 (Nr.119)	a = 6.333 c = 14.763 V = 592.14 $\Delta E_{form} = -0.253$	328 14.773 591.49 (Å ³ /cell) -0.236 (eV/atom)	a = 6.336 [61] 6.338 [62] c = 14.870 [61] 14.832 [62] V = 597.0 (Å ³ /cell) [61] 595.73 [62]		

Table 2 shows that the calculated formation energy for θ -Al₁₃Fe₄ is more prominent than/close to the scattered experimental data. The GGA calculations provide lower formation energy for θ' -Al₂Cu than that for θ -Al₂Cu. This is against the experimental observations that the latter is the ground state [17–19]. Meanwhile, the GGA + U approximation produces a significant difference of correction effect on the formation energies of the Al₂Cu phases [50] so that θ -Al₂Cu has a lower energy than θ' -phase. Such a difference of the on-site Coulomb interaction correction originates from the local coordination of Cu in the two phases (Figure 2). The Cu atoms in θ -Al₂Cu have eight Al and two Cu neighbors, whereas the Cu atoms in θ' -Al₂Cu have only eight Al neighbors (Figure 2c,f).

To gain insight into the physics behind these results, we also performed electronic structure calculations for the three Cu-containing compounds using the GGA + U approach (Figure 3).



Figure 2. (Color on line) Schematic structure of ω -Al₇Cu₂Fe along its [100] direction (**a**) and related coordination of Fe (**b**) and Cu (**e**); of θ -Al₂Cu (**d**) and the coordination of Cu (**c**); and of θ' -Al₂Cu (**g**) and the coordination of Cu (**f**). The silvery spheres represent Al, the blue spheres represent Cu, and the goldish spheres represent Fe.



Figure 3. (Color on line) Partial and total density of states for the Al-rich Cu-containing compounds in Al-Fe-Cu system using the DFT-GGA with Hubbard U correction for Cu 3d electrons. Subfugures a) to c) for the partial density of states (pDOS) of atoms in and total density of states (tDOS) of θ -Al₂Cu, d)-f) θ' -Al₂Cu in (**a**) and h)-m) for ω -Al₇Cu₂Fe in (**b**). For partial density of states, the green curves represent s-characters, the blue curves represent p-characters, and the red curves represent d-characters. The black curves represent the total DOS. The unit for pDOS is states/eV per atom and for tDOS is states/eV per formular unit.

As shown in Figure 3a, the frames of the DOS curves for θ' - and θ -Al₂Cu phases are similar. The DOS curves for both phases are dominated by Cu 3d states. However, there are some subtle differences. (1) The DOS of θ -Al₂Cu starts at -10.7 eV, whereas for θ' -Al₂Cu, it starts at -10.1 eV. (2) At the DOS curve of θ -Al₂Cu, the Cu 3d states range from -6.2 eV to -3.2 eV with a width of 3.0 eV, whereas for θ' -Al₂Cu, the Cu 3d states range from -6.0 eV to -3.9 eV with a width of 2.1 eV. This indicates a more localized character of the Cu 3d in θ' -Al₂Cu. The broader 3d band of θ -Al₂Cu is due to the Cu-Cu interaction, as shown in Figure 2. The more localized character of the Cu 3d in θ' -Al₂Cu is the cause of the more significant on-site Coulomb interaction correction effect on the formation energy.

Figure 3b shows that the tDOS curve for ω -Al₇Cu₂Fe is primarily determined by the Cu and Fe 3d states. The tDOS starts at -10.7 eV, similar to that of θ -Al₂Cu. The Cu 3d states range from -6.2 eV to -3.1 eV with a width of 3.1 eV, which is also close to that of θ -Al₂Cu. This is understandable, since the Cu atoms have eight Al and two Cu neighbors in both structures, as shown in Figure 2. There is little direct interaction between Cu 3d electrons and those at the Fe 3d orbitals in ω -Al₇Cu₂Fe (Figure 2). This agrees with the study that Cu 3d electrons behave as a semi-core level (Section 3.1). The Fe 3d states dominate the upper part of the valence band, ranging from -3.5 eV to -0.2 eV and extend to the upper part of the valence band and the lower part of the conduction band. This relates to the more delocalized character of the unfilled Fe 3d states in such metallic systems.

3.3. Cu Substitution at the Atomic Sites in θ -Al₁₃Fe₄

We first define the energy of a substituted Cu at the Al and Fe sites in θ -Al₇₈Fe₂₄ with respect to θ -Al₁₃Fe₄ and elemental Cu, Al, and Fe as,

$$\Delta E_{subs}(\theta - Al_{77}CuFe_{24}) = E(\theta - Al_{77}CuFe_{24}) - [E(\theta - Al_{78}Fe_{24}) + (E(Cu) - E(Al)]$$
(3)

$$\Delta E_{\text{subs}}(\theta - \text{Al}_{78}\text{CuFe}_{23}) = E(\theta - \text{Al}_{78}\text{CuFe}_{23}) - [E(\theta - \text{Al}_{78}\text{Fe}_{24}) + (E(\text{Cu}) - E(\text{Fe})]$$
(4)

Equation (3) is for Cu substitution of an Al, and Equation (4) is for Cu substitution of an iron in θ -Al₁₃Fe₄. The calculated results, including the coordination of Cu and the substitution energy, together with the local coordination in pure θ -Al₁₃Fe₄ as a comparison, are listed in Table 3.

The explorations reveal that Cu substitutions of Al generally do not change the local coordination. However, in the local chemical bonds for Cu at the Al sites, the Cu-Fe bond-lengths are slightly larger than the corresponding Al-Fe ones, whereas the Cu-Al bond-lengths are slightly shorter than the corresponding Al-Al bonds. Meanwhile, Cu substitutions of Fe cause longer Cu-Al bonds and shorter Cu-Fe bonds than the corresponding Fe-Al and Fe-Fe ones, respectively.

Table 3 shows that substituting one Fe with one Cu costs high energy (>0.45 eV), being unlikely. The Cu substitution of an Al7 atom is favored with negative formation energy (-0.123 eV/Cu), whereas the rest of the Cu substitutions of Al atoms cost energy. The lowest cost is one Cu atom at Al9 sites with a formation energy of 0.246 eV/Cu (Table 3 and Figure 4). Interestingly, at Al7 sites, each Cu has only two iron neighbors, compared with other sites, e.g., Al4, Al8, and Al9, where each substitution Cu has four Fe neighbors. This indicates that the number of Fe neighbors plays no significant role, corresponding to the weak interaction between the semi-core Cu 3d electrons and Fe 3d states. Meanwhile, the Al7 sites have the highest local symmetry (2/m), indicating that symmetry may play a role.

We performed structural optimization and total energy calculations for more Cu substitutions at the Al sites in θ -Al₁₃Fe₄. The configurations with low substitution energies are shown in Figure 4.

Table 3. Results (local coordination and substitution energy according to Equations (3) and (4)) for one Cu substitution at the Al and Fe sites in θ -Al₁₃Fe₄ from the GGA + U (U = 4 eV for Cu 3d) approach. Coordination of the related Cu-M (M = Al or Fe) with 3.0 Å in the substituted structures and Al/Fe-M in pure θ -Al₁₃Fe₄ is included.

Spe., Site (sym)	Al/Fe-M in Pure θ-Al ₁₃ Fe ₄	Cu-M in Doped θ -Al ₁₃ Fe ₄	ΔE_{sub} (eV/Cell)	
Al1, 4i (m)	Al1-7Al(2.54–2.89)Cu-7Al(2.44–2.84)-3Fe: 2.45, 2.53, 2.62-3Fe: 2.49, 2.66, 2.79		0.637	
Al2, 4i (m)	Al2-4Al(2.92–2.97) -2Fe: 2.36(×2)	Cu-4Al(2.76–2.88) -2Fe: 2.49(×2)	0.801	
Al3, 4i (m)	Al3-10Al(2.71–2.96) -2Fe: 2.44, 2.52	Cu-10Al(2.64–2.85) -2Fe: 2.53, 2.62	0.448	
Al4, 4i (m)	Al4-7Al(2.53–2.75) -4Fe: 2.51, 2.61(×2), 2.63	Cu-7Al(2.41–2.68) -4Fe:2.62, 2.65(×2), 2.74	0.453	
Al5, 4i (m)	Al5-8Al(2.67–2.86) -2Fe: 2.41, 2.44	Cu-10Al(2.62–2.84) -2Fe: 2.50, 2.52	0.422	
Al6, 4i (m)	Al6-7Al(2.56–2.86)Cu-7Al(2.48–2.79)-3Fe: 2.47, 2.48, 2.55-3Fe: 2.52, 2.58, 2.70		0.805	
Al7, 2d (2/m)	Al7-8Al(2.69–2.80) -2Fe: 2.47(×2)	Cu-8Al(2.58–2.69) -2Fe: 2.59(×2)	-0.123	
Al8, 4i (m)	Al8-7Al(2.56–2.81) -4Fe: 2.47(×2), 2.61, 2.70	Cu-7Al(2.41–2.68) -4Fe:2.55(×2), 2.69, 2.77	0.646	
Al9, 4i (m)	Al9-7Al(2.53–2.67) -4Fe: 2.46, 2.49(×2), 2.86	Cu-7Al(2.39–2.57) -4Fe: 2.57, 2.58(×2), 2.89	0.246	
Al10, 8j (1)	Al10-9Al(2.67–2.86) -2Fe: 2.48, 2.52	Cu-9Al(2.59–2.84) -3Fe: 2.58, 2.62, 2.84	0.453	
Al11, 8j (1)	Al11-9Al(2.63–2.97) -3Fe: 2.46, 2.51, 2.77	Cu-9Al(2.54–2.82) -3Fe: 2.56, 2.62, 2.87	0.387	
Al12, 8j (1)	Al12-8Al(2.64–2.96) -3Fe: 2.46, 2.56, 2.64	Cu-9Al(2.59–2.96) -3Fe: 2.55, 2.62, 2.85	0.586	
Al13, 8j (1)	Al13-7Al(2.67–2.88) -3Fe: 2.44, 2.55, 2.58	Cu-8Al(2.60–2.97) -3Fe: 2.52, 2.59, 2.75	0.661	
Al14, 8j (1)	8j (1) Al14-8Al(2.64–2.92) -3Fe: 2.45, 2.46, 2.81 Cu-9Al(2.3		2.074	
Al15, 8j (2)	Al15-8Al(2.78–2.89) Cu- 15, 8j (2) -4Fe: 2.46(×4) -4		0.879	
Fe1, 4i (m)	Fe1-11Al(2.46–2.86)	Cu-11Al(2.51–2.93)	0.548	
Fe2, 4i (m)	Fe2-10Al(2.44-2.70)	Cu-10Al(2.49–2.85)	0.548	
Fe3, 4i (m)	Fe3-10Al(2.44–2.77) -1Fe: 2.91	Cu-10Al(2.50–2.88) -1Fe: 2.86	0.638	
Fe4, 4i (m)	Fe4-10Al(2.41–2.72) -1Fe: 2.93	Cu-10Al(2.47–2.77) -1Fe: 2.91	0.672	
Fe5, 8j (1)	Al1-9Al(2.36–2.61)	Cu-9Al(2.37–2.70)	1.004	

The calculations reveal that Cu substitution at the Al sites is favorable and the formation energy decreases with increasing Cu concentration, reaching the minimum when Cu fully occupies the Al7 sites. This produces a stable structural model, θ -Al₇₆Cu₂Fe₂₄. As shown in Figure 4, three Cu substitutions at two Al7 sites and one Al9 site has a small formation energy of +0.030 eV. This minor energy cost can be easily compensated by configuration entropy contribution at elevated temperature. Partial Si substitution of Al7 and Al9 sites produces extra freedom (number of independent configurations): w = 2 for one Si atom at Al7 and w = 1 for two Si occupying the Al7 sites; $w = 2 \times 4 = 8$ for one Si at Al7 and one at Al9. For the full occupation of the Al7 sites, w = 4 for one extra Si, 6 for two Si, 4 for 3 Si, and 1 for four Si atoms occupying the Al9 sites. The Gibbs free energy expression is given as $\Delta G = \Delta H - T \Delta S_{conf}$, where $\Delta S_{conf} = kB \ln w$. Here, kB is the Boltzmann's constant, and w represents the extra freedom. We calculated the Gibbs free energy for the configurations, with Si substitutions at the Al7 and Al9 sites. The obtained results are shown in Figure 4. At 1000 K, we found that the Gibbs free energy of the configuration, where two Si are at Al7 sites and one Si is at the Al4 site, is negative due to the configurational entropy contribution. Next, we focus on the crystal chemistry of θ -Al₇₆Cu₂Fe₂₄.



Figure 4. (Color on line) Dependence of substitution energy on Cu concentration for configurations of high stability at 0 K and related free energy at 1000 K.

3.4. Crystal and Electronic Properties of θ -Al₇₆Cu₂Fe₂₄

We present the results of the stable θ -Al₇₆Cu₂Fe₂₄ phase. The first-principles GGA + U calculations produce the lattice parameters and formation energy with respect to the elemental solids according to Equation (2). The results for θ -Al₇₆Cu₂Fe₂₄ and the parent θ -Al₇₈Fe₂₄ are shown and compared in Table 4. Figure 5 shows the schematic structure of θ -Al₇₆Cu₂Fe₂₄ projected along its [10] direction and the local coordination of a Cu atom.

The Cu substitution reduces the lengths of *b*-axis but increases the lengths of the *a*-axis, the *c*-axis, and the angle, β . The volume of the unit cell of the Cu substituted crystal becomes smaller, as well. With respect to the elemental solids, the ternary θ -Al₇₆Cu₂Fe₂₄ phase has a lower formation energy as compared to the parent θ -Al₇₈Fe₂₄ and the elemental Al and Cu solids.

The atomic coordinates in the ternary θ -Al₇₆Cu₂Fe₂₄ phase follow the same trend in that they differ slightly from the corresponding ones in the binary θ -Al₇₈Fe₂₄, as shown clearly in Table 4. To better compare the ternary θ -Al₇₆Cu₂Fe₂₄ phase with the parent θ -Al₇₈Fe₂₄, we performed electronic band structures for the two compositions. The partial density of states (pDOS) for the 3d transition metals and total density of states (tDOS) in the two compounds and the dispersion curves are compared in Figures 6 and 7, respectively.

For both compounds, the DOS curves have one band, starting at -10.9 eV. The upper part of the valence band from -3.0 eV to -0.2 eV is dominated by Fe 3d states, which form rather flat bands, as shown in Figure 7. The Cu 3d states form a band ranging from -5.6 eV to -2.5 eV with a peak at -4.3 eV. There are tails of the Cu 3d states at about -1.0 eV. Overall, the Cu 3d orbitals are fully occupied and contribute little to the chemical bonding in the crystal. The frame of the dispersion curves near the Fermi level is similar (Figure 7) since they are determined by the Fe-Al interaction. Figure 7 also shows that the states along the Γ -X line (about 0.6 eV) are slightly more dispersive than those at the Γ -Y and Γ -Z lines (typically ~0.4 eV). Considering the length ratios, there is no strong anisotropy in both θ -Al₇₈Fe₂₄ and θ -Al₇₆Cu₂Fe₂₄ crystals, which is different from other Fe-IMCs, e.g., β -AlFeSi [28].



Figure 5. (Color on line) Schematic structure of the ternary θ -Al₇₆Cu₂Fe₂₄ phase projected along its [010] orientation and the related coordination of Cu by Fe and Al (**b**). The meaning of the spheres is the same as in Figure 2. The red lines in (**a**) represent the axis.

Table 4. The calculated lattice parameters, formation energy, and coordinates of atoms in θ -Al₇₆Cu₂Fe₂₄. The related results for the parent θ -Al₇₈Fe₂₄ are included for sake of comparison. * The Al7 sites are occupied by Cu.

Compound		θ-Al ₇₈ Fe ₂₄		θ -Al ₇₆ Cu ₂ Fe ₂₄			
Lattice Parameters (Å)		a = 15.426, b = 8.022, c = 12.425;		a = 15.504, b = 7.930, c = 12.459;			
		B = 107.68(°), V = 1464.92(Å ³)		B = 108.16 (°), V = 1455.62(Å ³)			
ΔE_{form} (eV/atom)		-0.329		-0.332			
Species	site	х	у	Z	х	У	Z
Al1	4i	0.065	0	0.173	0.066	0	0.1727
Al2	4i	0.323	0	0.2812	0.3214	0	0.2786
Al3	4i	0.2371	0	0.5354	0.2334	0	0.5362
Al4	4i	0.0737	0	0.5803	0.0738	0	0.5794
Al5	4i	0.2409	0	0.9613	0.2391	0	0.9557
Al6	4i	0.4788	0	0.83	0.4778	0	0.8312
Al7	2d	0.5	0	0.5	0.5	0	0.5000 *
Al8	4i	0.3049	0	0.7726	0.305	0	0.7715
Al9	4i	0.0874	0	0.7883	0.0826	0	0.7845
Al10	8j	0.1854	0.2175	0.1107	0.1859	0.2166	0.1108
Al11	8j	0.368	0.2117	0.1097	0.3682	0.2111	0.1102
Al12	8j	0.1782	0.2193	0.3345	0.1787	0.2187	0.3366
Al13	8j	0.4922	0.2333	0.33	0.4901	0.2222	0.3352
Al14	8j	0.3631	0.2186	0.4781	0.367	0.2064	0.4768
Al15	4g	0	0.2503	0	0	0.2501	0
Fe1	4i	0.0853	0	0.383	0.0836	0	0.3812
Fe2	4i	0.4015	0	0.6237	0.396	0	0.6298
Fe3	4i	0.0908	0	0.9878	0.0904	0	0.9872
Fe4	4i	0.402	0	0.9851	0.4022	0	0.9861
Fe5	8j	0.3198	0.2935	0.2779	0.3205	0.2942	0.2774



Figure 6. (Color on line) The pDOS of the selected atoms and tDOS curves for the binary θ -Al₇₈Fe₂₄ (**a**) and the θ -Al₇₆Cu₂Fe₂₄ phase (**b**). For the pDOS, **a**) to f) and h) to m), the green curves represent contributions from the s-character states, blue the p-character, red the d-character and black the sum of the s, p and d states of the atoms.



Figure 7. (Color on line) Dispersion curves of states around the Fermi level along the main axis for θ -Al₇₈Fe₂₄ (**a**) and for θ -Al₇₆Cu₂Fe₂₄ phase (**b**). The red lines represent the Fermi level (at 0 eV).

4. Discussion

First, we discuss the importance of the on-site Coulomb correction for metallic compounds containing narrow bands, e.g., Cu 3d. The DFT + U correction has been used to describe Cu-containing inorganic compounds, e.g., oxides [48,49]. Recently, our work has shown that it is also essential for metallic compounds containing Cu [50]. In this paper, we further describe this effect in detail. The Cu 3d states are fully occupied in the element Cu and Cu-containing metallic compounds. However, the standard DFT approximations have the Cu 3d states too close to the Fermi level, thus producing a physically meaningless interaction between Cu 3d and other metals. The Hubbard U correction pushes the Cu 3d orbitals to lower energy and thus avoids the unphysical interaction. Our calculations also showed a narrower band width of Cu 3d in θ' -Al₂Cu (2.1 eV) as compared to that in θ -Al₂Cu (3.0 eV) and in the pure Cu (~3.6 eV). Consequently, the Hubbard U correction is more pronounced for θ' -Al₂Cu, which results in the correct prediction of stability of the Al₂Cu phases.

The calculations predicted the stable Al-rich compounds in the Al-Fe-Cu system: θ -Al₁₃Fe₄ in the binary Al-Fe line, θ -Al₇₆Cu₂Fe₂₄ in Cu-poor conditions, ω -Al₇Cu₂Fe in Cu-rich conditions, and θ -Al₂Cu in the Al-Cu line. This is in line with the thermodynamics assessments in the Al-rich phase diagrams [11–13,17–19]. In the Al-Fe-Cu system, the stable Al-rich compounds (with Cu concentrations larger than 0) are θ -Al₇₆Cu₂Fe₂₄, ω -Al₇Cu₂Fe, and θ -Al₂Cu. We can also find the composition relation: ω -Al₇Cu₂Fe = 1/24(Al₇₆Cu₂Fe₂₄) + (11.5/3)(θ -Al₂Cu). Using the calculated total valence electron energies, we obtained the energy difference $\Delta E = E(\omega$ -Al₇Cu₂Fe) – [(1/24) $E(Al_{76}Cu_2Fe_{24})$ + (11.5/3) $E(\theta$ -Al₂Cu)] = -0.234 eV/f.u. (formula unit), and ω -Al₇Cu₂Fe is favored, in agreement with the experimental observations [17–19].

The calculations showed that θ -Al₇₆Cu₂Fe₂₄ is the ground state compound at low temperature. In this compound, the Cu content is 1.96 at %, in line with the phase diagrams [17–19]. However, additional Cu substitutions at the Al9 sites cost a moderate amount of energy. Partial occupation of the Al sites induces extra freedom (configurational entropy contribution in thermodynamics). At 1000 K, an additional Cu at Al9 sites is favored, forming θ -Al₇₅Cu₃Fe₂₄ with about 2.94 at % Cu. At 1360 K, two additional Cu can be doped into the Al9 sites, forming the θ -Al₇₄Cu₄Fe₂₄ structure with 3.92 at % Cu. This is close to the experimental observation by Freiburg and Grushko, who found Cu at both Al7 and Al9 sites [25], and the thermodynamics studies [17–19]. However, our calculations are different from the experimental observation of quenched samples in [26]. We believe that the quenched samples in [26] may contain impurities such as Si from the vessel, as our previous work showed that Si atoms can be doped into θ -Al₁₃Fe₄ [37], which in turn increases Cu solution content. This topic deserves further investigation.

5. Summary

Using the first-principles DFT approach with on-site Coulomb interaction correction, we investigated the stability, phase relations, structural properties, and electronic properties of the Al-rich compounds, θ -Al₁₃Fe₄, and Al₂Cu phases, ω -Al₇Cu₂Fe, as well as Cu substitution in θ -Al₁₃Fe₄. Due to the more localized nature of the Cu 3d electrons in θ' -Al₂Cu, the current approach predicted the θ -phase being more stable. We also found that Cu substitution at the Fe sites is too costly and thus unlikely. At ambient conditions, Cu prefers the Al7 sites, forming θ -Al₇₆Cu₂Fe₂₄. At a high temperature, Cu occupies the Al7 sites and partial Al9 sites, forming θ -Al_{76-x}Cu_{2+x}Fe₂₄. This agrees with the experimental observations. Electronic structure calculations also showed moderate contributions of the Cu 3d states to the electronic properties and chemical bonding in the Cu-containing metallic compounds.

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