Atomistic Modeling of RuAl and (RuNi)Al alloys

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\textbf{Abstract}
Atomistic modeling of RuAl and RuAlNi alloys, using the BFS method for alloys is performed. The lattice parameter and energy of formation of B2 RuAl as a function of stoichiometry and the lattice parameter of (Ru\textsubscript{50-x}Ni\textsubscript{x})Al\textsubscript{50} alloys as a function of Ni concentration are computed. BFS-based Monte Carlo simulations indicate that compositions close to Ru\textsubscript{25}Ni\textsubscript{25}Al\textsubscript{50} are single phase with no obvious evidence of a miscibility gap and separation of the individual B2 phases.

\textbf{Introduction}

In comparison with nickel or cobalt aluminides, B2 RuAl has appreciable room-temperature toughness and plasticity and maintains considerable strength at high temperatures [1-2]. These properties, in combination with excellent oxidation resistance [3], make this alloy a potential candidate for aerospace applications though cost and high density are a significant concern. In an effort to drive down both cost and weight and improve upon its other properties, several studies [4-6] have looked at alloying schemes for replacing Ru or Al with other elements that generally form an isostructural B2 phase such as Co and Fe for Ru and Ti for Al. But by far, the most widely studied ternary alloying addition has been Ni [7-13]. Even then, there is disagreement as to the structure of ternary Ni-Ru-Al alloys that exist between the NiAl and RuAl B2-phase fields.

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Chakravorty and West [9-10] have reported a miscibility gap centered between the two binary phases resulting in a region consisting of two distinct B2 compounds. In apparent agreement, Sabariz and Taylor [11] have also observed a two-phase alloy at the composition Ni_{25}Ru_{25}Al_{50}. While Horner et al. [13] found results similar to Chakravorty and West [10] in that many, but not all, of the ternary compounds seemed to exhibit two distinct components, the evidence seemed to overwhelmingly suggest coring as opposed to actual formation of two distinct B2 phases.

Furthermore, a sample within the miscibility gap claimed by Chakravorty and West was heavily milled and annealed so that diffusion distances would be much smaller and a better opportunity for obtaining a near equilibrium structure would exist. In this case, only a single B2 phase was observed. Furthermore, Liu et al. [12] observed only a single B2 phase across the (NiRu)Al system in mechanically alloyed samples when Ni was varied between 10 and 25 at.%, indicating complete mutual solubility between NiAl and RuAl.

Given the lack of agreement for even the structure of Ni-Ru-Al alloys and the generally scarce data for other ternary or higher order systems, it is useful to consider the development of a modeling effort to supplement the ongoing experimental work on RuAl alloys. In this area, we provide the necessary tools to perform such modeling, and use them to study the fundamental properties of RuAl and (Ru,Ni)Al alloys. The modeling is performed within the framework of the BFS method for alloys, a quantum approximate method for the description of the energetics of complex systems at the atomic level, and is validated by comparison to experimental data.

The BFS Method

The BFS method for alloys [14] has been proven to be highly effective for the study of multicomponent systems. With the proper parameterization, it allows for an extremely economical, computationally simple, and physically sound description of the energetics of large collections of atoms. The BFS method is based on the assumption that the heat of formation, \( \Delta H \), of a given collection of atoms is the sum of the contributions of each atom in the sample, \( \varepsilon_i \). Each contribution \( \varepsilon_i \) consists of two terms: a strain energy \( (\varepsilon_i)^S \) which accounts for the change in geometry with respect to a single monoatomic crystal of the reference atom, and a chemical energy \( (\varepsilon_i)^C \), linked by a coupling function \( (g_i) \) so that \( \varepsilon_i = (\varepsilon_i)^S + g_i(\varepsilon_i)^C \). Three parameters, equilibrium lattice parameter, cohesive energy, and bulk modulus, for each of the constituent
atoms are needed in the general derivative structure of the compound being studied (in this case, bcc) in order to calculate these terms. The chemical energy accounts for the corresponding change in composition, considered as a defect in an otherwise pure crystal. The chemical ‘defect’ deals with pure and mixed bonds, therefore, two additional perturbative parameters ($\Delta_{AB}$ and $\Delta_{BA}$ where A, B = Ru, Ni, Al) are needed to describe these interactions. All the parameters (listed in Table 1) are determined with the Linearized-Augmented Plane Wave method (LAPW) [15]. An alternative set of parameters, obtained with the Linear-Muffin Tin Orbital method [16] can be found in Ref. 17. We refer the reader to Ref. 14 for a detailed description of the BFS method and its parameterization, and Ref. 18 for details on Monte Carlo-Metropolis simulations using BFS.

**Results and Discussion**

A general comparison between the LAPW values and other calculated and experimental values of the lattice parameter, cohesive energy, and bulk modulus for RuAl [19-33] is shown in Table 2. The LAPW calculations correctly reproduce the value of the lattice parameter $a_0$ for stoichiometric RuAl. Although Ru and Al do not exist in the bcc phase, the lattice parameter of the computed bcc-Ru and bcc-Al are both larger than the equilibrium lattice parameter of RuAl (a 1.4% difference with respect to bcc-Ru, and a 7.8% difference for bcc-Al). The predicted value of $a_0$, 3.005 Å, compares well with the experimental values, 2.95 Å [32] to 3.03 Å [20]. Considering the typical deviations in first-principles values for the energy of formation, the LAPW prediction compares well with the experimental value of 1.28 eV/atom [21]. The LAPW input, validated in Table 2, is used for the determination of the BFS perturbative parameters $\Delta_{AB}$ and $\Delta_{BA}$, listed in Table 1 for all possible pair combinations of Ru, Al, and Ni.

We first apply the parameters listed in Table 1 to the analysis of the B2 RuAl phase field, by considering a large number of computational cells (defined previously in Ref. 14) including all possible defect structures for off-stoichiometric compositions. Whether structural vacancies are included or not in this set of possible configurations, it is seen that the RuAl phase is quite unstable with respect to changes in stoichiometry, in the sense that there is a noticeable change in energy of formation off stoichiometry. Furthermore, this change is more pronounced for Al-rich alloys, as can be seen in Fig. 1. For Ru-rich alloys, the lowest energy configurations correspond to substitutional alloys. The analysis suggests a trend towards phase separation, as configurations
showing clustering of excess Ru atoms are energetically favored. This can be explained in terms of the individual BFS contributions $e_i$ ($i = \text{Ru, Al}$) to the energy of formation of the computational cell. For B2 RuAl, $e_{\text{Ru}} = -0.3383 \text{ eV/atom}$ and $e_{\text{Al}} = -1.6516 \text{ eV/atom}$, indicating that both atoms contribute to the formation of the alloy (negative contributions to $\Delta H$). The 'weak' contribution of Ru atoms (compared to that of Al atoms) and the similarity between the atomic volume per atom in the B2 RuAl alloy and the atomic volume in a pure Ru crystal, lead to the phase separation observed for Ru-rich alloys [6]. To complete the analysis of B2 RuAl, Monte Carlo simulations using the NN approximation [18] were used to determine the melting temperature. It was found that the structure is highly ordered and stable up to 2300 K, in excellent agreement with the experimental value of 2330 K [1].

Given the possibility of 1) reducing the cost and weight of RuAl alloys through substitution of Ru with Ni, 2) the potential to use RuAl as a reinforcing phase in other intermetallic systems, possibly even NiAl [7], and 3) given the controversy surrounding the phase structure of ternary Ru-Ni-Al alloys in the region between the two binary B2 phases NiAl and RuAl [9-13], Ru-Ni-Al would appear to be an ideal system for further theoretical analysis. While the following BFS analysis is applied to Ni additions to RuAl, it should be noted that similar calculations can be easily performed for other ternary or higher order additions using, for example, the BFS parameters listed in Ref. 17.

For the ternary case, we first determine the preferred site occupancy for dilute additions of Ni in RuAl. Let $A(B)$ denote an A atom occupying a site in the B sublattice, and $A(B)'B(C)$ the same, but with the displaced B atom moving to a site in the C sublattice. For $(\text{Ru}_{50-x}\text{Ni}_x)\text{Al}_{50}$ alloys, $\text{Ni}$(Ru) = -0.98217 $e\text{V/atom}$ and $\text{Ni}$(Al)$(\text{Ru}) = -0.8780 e\text{V/atom}$, indicating that Ni prefers available Ru sites. However, for $\text{Ru}_{50}(\text{Al}_{50-x}\text{Ni}_x)$ alloys $\text{Ni}$(Al) = -0.9600 $e\text{V/atom}$ and $\text{Ni}$(Ru)$\text{Ru}(\text{Al}) = -0.9513 e\text{V/atom}$, indicating a slight preference for Al sites, although the difference in energy is so small that it is difficult to conclude any obvious preference for either site under these conditions. The strong preference of Ni for Ru sites when Ni+Ru $< 50$ at.% is not surprising, as it allows for the formation of strong Ni-Al bonds.

In terms of the phase structure of the ternary alloy(s) bridging the two binary B2 phases NiAl and RuAl, there are reports that suggest either a miscibility gap and phase separation for compositions near $\text{Ni}_{25}\text{Ru}_{25}\text{Al}_{50}$ [9-11] or a single ternary B2 phase throughout [12-13]. While BFS Monte Carlo NN simulations, as described in Ref. 18, clearly show the formation of the B2
RuAl phase, as can be seen in Fig. 2.a, the results are just as clear for Ni_{25}Ru_{25}Al_{50}, as shown in Fig. 2.b. The simulations show that both Ru and Ni share the same sublattice. The degree and type of ordering can be estimated from the coordination matrices $a$ and $b$, for nearest-neighbors (NN) and next-nearest-neighbors (NNN), respectively, where the matrix element $a_{ij}$ ($b_{ij}$) denotes the probability that an atom of species $i$ has an atom $j$ as a NN (NNN). If there is phase separation, it would be expected that atoms in the Ru/Ni sublattice will have, mostly, atoms of their same species as NNN's. If there was a ternary ordered phase other than B2, the NNN coordination of Ru or Ni atoms would reflect this fact with an increasing number of NNN's of the other species. For example, the NNN coordination matrix for a Heusler L2$_1$ ternary phase would maximize the value of $b_{NiRu}$ ($= b_{RuNi} = 1$). Phase separation, on the other hand, would lead to a very small value of $b_{NiRu}$ (and $b_{RuNi}$), maximizing in turn, $b_{NiNi}$ (and $b_{RuRu}$). Instead, the results of the simulations reflect the existence of a (Ru,Ni)Al B2 phase where Ru and Ni are randomly located throughout their own sublattice, in agreement with Liu et al. [12] and Homer et al. [13].

The BFS values of the lattice parameter for (Ru$_{50-x}$Ni$_x$)Al$_{50}$ as a function of Ni concentration (in at.%), $a(x)$, show a slight deviation with respect to the the average values obtained using Vegard's law [34], as shown in Fig. 3. Reported measurements of $a(x)$ [11,12] are also considered as having a linear behavior. The linear fit of the experimental values in Ref. 11 (normalized to the equilibrium B2 value $a_0$) is $a(x)/a_0 = 1 - 0.0006949x$. In spite of the slight positive deviation from linearity, the best linear fit of the BFS prediction is almost identical, $a(x)/a_0 = 1 - 0.0007266x$.

To further understanding of the behavior of the ternary alloys, it is possible to define within the framework of BFS, the role that each atom plays in (Ru$_{50-x}$Ni$_x$)Al$_{50}$ as a function of Ni concentration, by separately computing the BFS energy contributions. Fig. 4 shows the strain, chemical and total energy contributions of Ru, Ni, and Al atoms to ternary RuNiAl alloys, computed as the average over all atoms of similar species. Ru atoms provide a favorable (i.e., $\varepsilon^C < 0$, favoring alloying) chemical contribution to the total energy of formation, diminished by increasing strain as the Ni concentration increases. As a result, the role of Ru in favoring compound formation is diminished with increasing Ni content (Fig. 4.a). Ni atoms display the opposite behavior: as the concentration of Ni increases, the strain energy becomes substantially smaller as the average volume per atom in the alloy becomes closer to that of Ni. The chemical contribution, which does not favor alloying, increases, but at a slower rate than the decrease in strain, resulting in a total contribution that does not favor the stability of the alloy (Fig. 4.b). Al
atoms display a mild increase in strain with increasing Ni content, easily compensated by the growing (much more favorable than Ru) chemical contribution, leading to a net decrease of the total energy (Fig. 4.c), which along with the chemical contribution from Ru ultimately favors the formation of ternary B2 compounds.

Summary

The BFS method for alloys has been used to model the structure of RuAl and the extension of the B2 phase fields from NiAl to RuAl in the Ru-Ni-Al system. The BFS analysis correctly predicts the melting temperature of RuAl and indicates a strong preference for Ni in Ru sites in (Ru\textsubscript{50-x}Ni\textsubscript{x})Al\textsubscript{50} alloys, and an almost equal probability for either site in Al-deficient alloys. The results indicate that the two phases are isostructural with a single B2 phase of varying Ru:Ni ratio bridging the binary compounds. The calculated lattice parameter for (Ru\textsubscript{50-x}Ni\textsubscript{x})Al\textsubscript{50} alloys, as a function of Ni contents, displays slight deviations from the average values, but in close agreement with experiment.

Acknowledgments

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References

34. L. Vegard, Z. Phys. 5 (1921) 17.
TABLE CAPTIONS

Table 1: LAPW results for the lattice parameter, cohesive energy, and bulk modulus for the bcc phases of Ru, Al and Ni, the resulting equivalent crystal theory (ECT) parameters p, \( \alpha \), l and \( \lambda \) and the BFS perturbative parameters \( \Delta_{AB} \) and \( \Delta_{BA} \).

Table 2: Calculated (LAPW) properties compared to measured and other theoretical values for NiAl and RuAl.

Table 3: Nearest-neighbor (NN) and next-nearest-neighbor (NNN) coordination matrices for the Ru-Ni-Al simulation shown in Fig. 2.b. The matrix element in row \( i \) and column \( j \), denotes the probability that an atom of species \( i \) has a NN (top matrix) or a NNN (bottom matrix) of species \( j \), where \( i \) (or \( j \)) = 1, 2, 3 corresponds to Ru, Al, Ni, respectively.
FIGURE CAPTIONS

Fig. 1: Energy of formation (in eV/atom) vs. Ru concentration. The data points shown correspond to the lowest energy configuration found at each composition.

Fig. 2: Final state (room temperature) of Monte Carlo MC-NN simulations for (a) RuAl and (b) Ni25Ru25Al50. Ru, Ni, and Al atoms are denoted with white, dark, and light grey spheres, respectively.

Fig. 3: Comparison of experimental results and BFS predictions for the lattice parameter of (Ru50-xNi_x)Al50 alloys as a function of Ni concentration. Experimental results are from Ref. 11. The dashed line indicates the results of Vegard’s law.

Fig. 4: Individual average BFS contributions for a) Ru, b) Ni and c) Al atoms in a (Ru50-xNi_x)Al50 alloy, as a function of Ni concentration. In each case, the strain (Triangle up), chemical (triangle down) and total (solid disk) energies are shown.
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Table 1
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Table 2
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Table 3
Fig. 1
Fig. 2
Fig. 3
Fig. 4