Effects of Al on Precipitation Behavior of Ti-Nb-Ta-Zr Refractory High Entropy Alloys

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Abstract: Addition of Al can decrease density and improve oxidation resistance of refractory high entropy alloys (RHEAs), but may cause complicated precipitation and further affect mechanical properties. The present work studied the microstructural evolution of Al-contained RHEAs at elevated temperatures. The effects of Al on precipitation behavior were discussed. Results show that, TiNbTa0.5ZrAlx alloys (x ≤ 0.5) have single BCC (Body Centered Cubic) structure, but the primary BCC phase is supersaturated. Precipitation of BCC2(Nb,Ta)-rich solid solution phase, HCP(Zr,Al)-rich intermetallic phase, and ordered B2 phase can occur during heat treatment at 600~1200 °C. The precipitation of BCC2 phase mainly exists in RHEAs with low content of Al, while HCP (Hexagonal Close Packed) precipitates prefer to form in RHEAs with high content of Al. Interestingly, ordered B2 precipitates with fine and basket-weave structure can form in TiNbTa0.5ZrAl0.5 alloy after annealing at 800 °C, producing significant precipitation hardening effect.

Keywords: high entropy alloy; refractory metal; precipitation behavior; microstructure

1. Introduction

High entropy alloys (HEAs) are a new class of metallic material that consist of multi-principal components but typically exhibit simple phase structure [1–3]. HEAs have many superior properties, such as high strength and toughness, good corrosion, and radiation resistance, mainly due to their flexible chemical compositions [4–6]. Refractory high entropy alloys (RHEAs) are a specific category of HEAs, which are primarily made up of high-melting-point elements, typically from group IV–VI in the periodic table [7–9]. RHEAs usually have single BCC structure because of the BCC structured components (elements from group V–VI are BCC structure and elements in group IV are also BCC structure at high temperatures). Most RHEAs possess high strength at high temperatures, making them a candidate for aerospace engineering, and have attracted extensive attention [10]. For example, the well-known WMoTaNb and WMoTaNbV RHEAs have yield strength above 400 MPa up to 1600 °C, which were considered as potential substitution for nickel based superalloys [11]. But the heavy elements result in high density (usually > 10 g/cm³), limiting their engineering application. Besides, some refractory metals, such as V and Mo, have poor high-temperature-oxidation resistance, making RHEAs difficult to work in atmosphere at high temperatures [12,13]. Addition of Al to RHEAs can effectively reduce density and improve oxidation resistance at the same time. For instance, Lin et al. [14] introduced Al into HfNbTaTiZr alloy and reduced its density from 9.96 to 8.84 g/cm³. Bronislava et al. [13] introduced Al into TaMoCrTi alloy and found that the oxidation resistance improves significantly. Thus, alloying with Al may be a feasible approach for RHEAs to apply in aerospace engineering.
Nevertheless, alloying with Al could lead to complex precipitation behavior in RHEAs. Yurchenko et al. [15] reported phase transformation in Al\textsubscript{x}NbTiVZr RHEAs and found that the addition of Al leads to the formation of C14 Laves and Zr\textsubscript{2}Al-type phases. Furthermore, the phase evolution in Al\textsubscript{x}NbTiVZr RHEAs is sensitive to the content of Al. Besides, they also found the addition of Al causes precipitation of another (Ti, Al)-rich orthorhombic phase in a Ti\textsubscript{40}Nb\textsubscript{30}Hf\textsubscript{15} alloy [16]. Schuh et al. [17] introduced Al into TiVNb alloy and observed precipitation of Nb\textsubscript{2}Al and Ti\textsubscript{3}Al intermetallic phases. Ge et al. [18] studied phase equilibrium in Al-contained MoNbTaTiV RHEAs through CALPHAD method and presented that the Al-contained RHEAs have BCC structure at high temperatures but decomposed into multiphase structure (including BCC1, BCC2, HCP, Sigma, and Ti\textsubscript{3}Al) at intermediate temperatures. Apparently, addition of Al can induce formation of various second phases in RHEAs, and the precipitation behavior is relevant to the content of Al as well as the composition of the matrix. In our previous work [19], we have designed Ti-Nb-Ta-Zr based RHEAs with good mechanical properties. In addition, we have demonstrated that the addition of Al can significantly enhance the high-temperature-oxidation resistance of the RHEAs [20]. However, the microstructural evolution of the Ti-Nb-Ta-Zr RHEAs with the addition of Al still remains unclear, which has become an urgent issue.

In this work, we fabricated a series of Al-contained Ti-Nb-Ta-Zr RHEAs and studied the microstructural evolution at elevated temperatures. The effects of Al on precipitation behavior were analyzed, and the detailed structures of the precipitates were characterized. The RHEAs were prepared through powder metallurgy method rather than conventional casting, so as to avoid severe chemical-segregation and make the analysis on precipitation behavior more precise.

2. Experimental

Elemental Ti, Nb, Ta, Zr, and Al powders with purities above 99.5% and average particle sizes of 18–42 \( \mu \)m were used to prepare the Ti\textsubscript{Ni-Ta\textsubscript{0.5}ZrAl\textsubscript{x}} (X = 0, 0.2, 0.5, 1.0) alloys through powder metallurgy method. The alloys were named “RHEA-Al0,” “RHEA-Al0.2,” “RHEA-Al0.5,” and “RHEA-Al1.0,” respectively. The powders were blended under argon atmosphere for 10 h in a V-type mixture machine (Zgsujia V-20, Yupan, Nanjing, China). The powder mixtures were compressed via cold isostatic pressing (CIP) at a pressure of 200 MPa. Then, the compressed compacts were sintered at 1400 \( ^\circ \)C for 10 h in a vacuum of 1 \( \times \) 10\(-3\) Pa followed by furnace cooling. To investigate precipitation behavior of the RHEAs, the as-sintered samples were cut into pieces and subjected to annealing at different temperatures. The pieces were encapsulated in quartz tubes and annealed at 600–1200 \( ^\circ \)C for 2 h, and then quenched in water to retain the microstructural characteristics.

The phase constitution, microstructures, and micro-hardness were studied to investigate the precipitation behavior of the RHEAs. The phase analysis was performed using X-ray diffraction (XRD, Advance D8, RigakuD, Tokyo, Japan) with CuK\( \alpha \) radiation. Microstructures were observed using a scanning electron microscope (SEM, Helios NanoLab G3 UC, FEI, Hillsboro, OR, USA) equipped with a backscatter electron (BSE) detector and an energy-dispersive X-ray spectroscopy (EDS). The chemical composition was analyzed by EDS. The details of microstructures were studied using a transmission electron microscope (TEM, Titan G260–300, FEI, Hillsboro, OR, USA). Specimens for TEM test were prepared using focused ion beam (FIB) apparatus (FEI, Hillsboro, OR, USA). The micro-hardness was measured using an indentation tester (BUEHLER5104, BUEHLER, Lake Bluff, IL, USA).

3. Results

The XRD patterns and microstructures of the as-sintered RHEAs are shown in Figure 1. RHEA-Al0, RHEA-Al0.2, and RHEA-Al0.5 all exhibit single BCC phase structure, while RHEA-Al1.0 shows BCC + HCP dual-phase structure. The HCP phase can be identified as Zr\textsubscript{5}Al\textsubscript{4} phase according to its diffraction peaks. Figure 1b–e presents the microstructures of as-sintered RHEAs with different Al content. Obviously, RHEA-Al0, RHEA-Al0.2, and RHEA-Al0.5 are single phase, and the microstructures are homogeneous. The contrasts of
The BSE images are uniform, indicating that the chemical components distribute uniformly in the alloys. RHEA-Al1.0 has a BCC matrix and HCP precipitates. The HCP phase is quasi-continuous at grain boundaries but forms needlelike morphology inside grains. The microstructures indicate that the addition of Al can lead to formation of an HCP structured Zr5Al4 intermetallic phase. Figure 2 shows detailed phase structure of the as-sintered RHEAs obtained by TEM. As a representative, RHEA-Al0.2 was characterized to analyze the BCC phase, since the three (RHEA-Al0, RHEA-Al0.2, and RHEA-Al0.5) have similar phase structures. The bright field (BF) image presents the BCC phase without any precipitate, and the inserted selected area electron diffraction (SAED) patterns further verify the single BCC structure. The high-resolution TEM (HRTEM) image shows that the BCC phase is disordered solid solution phase, and the interplanar spacing is measured as \( d = 0.242 \) nm for the crystal plane of (110). Figure 2c shows the BF image of RHEA-Al1.0 and the chemical composition of the HCP phase. The interface between the BCC matrix and the HCP precipitate is clear without any visible second phase. The chemical composition shows that the HCP phase is not a conventional Zr5Al4 intermetallic compound, instead, it also contains a small amount of Ti, Nb, and Ta, indicating a multiple Zr5Al4 type intermetallic phase. The SAED patterns of the intermetallic phase further verify its HCP structure.

**Figure 1.** XRD patterns and microstructures of the as-sintered refractory high entropy alloys (RHEAs): (a) XRD patterns of the as-sintered RHEAs, (b) BSE image of RHEA-Al0, (c) BSE image of RHEA-Al0.2, (d) BSE image of RHEA-Al0.5, and (e) BSE image of RHEA-Al1.0.

The phase constitutions of the RHEAs after annealing are shown in Figure 3. Apparently, addition of Al has strong impact on phase transformation of the annealed RHEAs. As for RHEA-Al0, annealing at temperatures above 1000 °C can maintain single BCC structure, while annealing at temperatures below 800 °C leads to phase decomposition and forms a BCC2 phase. The diffraction peaks of the BCC2 phase are very close to those of the primary BCC phase, indicating their lattice parameters should also be similar. RHEA-Al0.2 has analogous precipitation behavior with RHEA-Al0, as shown in Figure 3b, but the temperature for maintaining single BCC structure is higher (1200 °C) than that for RHEA-Al0, which means addition of Al can stimulate phase decomposition in the RHEAs. As for RHEA-Al0.5, the precipitation behavior becomes more complicated. Annealing at 800 °C could also lead to phase decomposition and forms the BCC2 phase, but annealing at 1000–1200 °C results in precipitation of an HCP phase, which was identified as the Zr5Al4 type intermetallic phase. The precipitation behavior of RHEA-Al1.0 is rather simplex. Additionally, the results show same phase constitutions at 600–1200 °C. The annealed RHEA-Al1.0 consists of BCC and HCP phases, and the intensity of the diffraction peaks of the HCP phase increases with the increasing annealing temperature, indicating a further precipitation of the HCP Zr5Al4 type intermetallic phase.
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Figure 2. Detailed phase structure obtained by TEM: (a) bright field (BF) image of BCC phase in RHEA-Al0.2 (inserted image gives the corresponding selected area electron diffraction (SAED) patterns), (b) high-resolution TEM (HRTEM) image of BCC phase in RHEA-Al0.2 (inserted image is the magnified view of the marked region), (c) BF image of RHEA-Al1.0 (inserted table gives the composition of the HCP phase), and (d) SAED patterns of the HCP phase in RHEA-Al1.0.

Figure 3. XRD patterns of the RHEAs annealed at 600–1200 °C for 2 h: (a) RHEA-Al0, (b) RHEA-Al0.2, (c) RHEA-Al0.5, and (d) RHEA-Al1.0.
Figure 4 presents microstructures of the RHEAs annealed at 600–1200 °C, which gives a direct comparison of the precipitation characteristics. As for RHEA-Al0 and RHEA-Al0.2, annealing at 600–1000 °C causes phase decomposition and forms the BCC2 phase. The BCC2 phase primarily distributes around grain boundaries in RHEA-Al0, but forms both around grain boundaries and inside grains in RHEA-Al0.2. The BCC2 phase was identified as (Nb, Ta)-rich precipitates, while the BCC phase was identified as Zr-rich matrix. The (Nb, Ta)-rich precipitates become coarser with annealing temperature increasing. As for RHEA-Al0.5, annealing at 800 °C results in a fine, basket-weave, nano-lamellar structure inside the grain, which can be regarded as BCC and B2 phases. Similar structure has also been observed by Senkov et al. [21] in a AlMo0.5NbTa0.5TiZr RHEA. The B2 precipitates are too fine to analyze using SEM, which are carefully investigated using TEM below. Annealing at 1000–1200 °C leads to a formation of needle-like (Zr, Al)-rich HCP precipitates. The precipitates are fine and distribute homogeneously when RHEA-Al0.5 annealed at 1000 °C. With annealing temperature increasing to 1200 °C, the precipitates become coarser. The precipitation behavior of RHEA-Al1.0 is much more simple compared with others. Annealing at 600–1200 °C results in further precipitation of (Zr, Al)-rich HCP precipitates, while the phase transformation of BCC2 is restrained. The microstructural characteristics of the annealed RHEAs are in good accordance with the XRD results. Figure 5 shows detailed structures of the B2 precipitates in RHEA-Al0.5 after annealing at 800 °C. The B2 precipitates have needle-like morphology with length of 10–50 nm and width below 10 nm. The SAED patterns show a crystal orientation of [001]B2 // [001]BCC. The (001)_{B2} super-lattice reflections indicate that the B2 phase has an ordered crystal structure. The HRTEM image reveals that the B2 phase is coherent with the BCC phase. The inserted images give magnified view of the atoms, which present an ordered arrangement for B2 phase (note the brightness variation), and the corresponding Fast Fourier Transform (FFT) images also indicate ordered structure for B2 phase (note the diffraction spot of (010) plane, which should not exist in disordered BCC phase), further demonstrating that the B2 phase is ordered, agreeing with the observations by Senkov et al. [21,22].

Figure 4. Microstructures of the RHEAs annealed at 600–1200 °C for 2 h. (a) RHEA-Al0, (b) RHEA-Al0.2, (c) RHEA-Al0.5, and (d) RHEA-Al1.0.
Precipitation could remarkably affect mechanical properties of RHEAs and typically causes hardening effect. The hardness of the current Al-contained RHEAs at different annealing status is shown in Figure 6. Apparently, the RHEAs become harder with the Al content increasing, mainly due to solid solution hardening. As for RHEA-Al0 and RHEA-Al0.2, the precipitation of BCC2 phase (annealing at 600–800 °C) causes increase in hardness. But, annealing above 1000 °C makes the RHEAs transform back to single BCC structure, resulting in decreasing in hardness. As for RHEA-Al0.5 and RHEA-Al1.0, the precipitation of HCP phase also causes hardening. In addition, it can be noted that the hardness reaches the maximum values when the RHEAs are annealed at 1000 °C, mainly because of the fine and dispersive HCP precipitates. Annealing at 1200 °C causes coarsening and dissolution of the HCP phase, leading to the decrease in hardness. It can also be noted that the hardness of RHEA-Al0.5 increases remarkably when annealing at 800 °C, indicating that the ordered B2 phase has significant precipitation hardening effect.

![Figure 5](image_url)

**Figure 5.** Detailed structures of the precipitates in RHEA-Al0.5 annealed at 800 °C for 2 h: (a) BF image of the precipitates, (b) SAED patterns of the RHEA-Al0.5, (c) DF image of the precipitates, (d) HRTEM image of the precipitates (inserted images are the magnified view of the atoms and corresponding Fast Fourier Transform (FFT) patterns).

![Figure 6](image_url)

**Figure 6.** Hardness of the Al-contained RHEAs at different annealing status.
4. Discussion

RHEAs normally form solid solution phase with single BCC structure when being prepared by casting method, because the solid solution phase can be stabilized by high entropy effect at high temperatures and possible phase transformation or precipitation could be kinetically hindered during rapid solidification process [8,23]. Similarly, RHEAs synthesized by high-temperature sintering also have single BCC structure. When being cooled down to room temperature, the cooling rate is too fast for RHEAs to accomplish possible phase transformation, therefore, forming supersaturated solid solution phase. The supersaturated BCC phase may decompose or transform to various second phases when subjected to annealing at intermediate temperatures and, consequently, affect the mechanical properties of RHEAs. Phase equilibrium of RHEAs can be described by several empirical parameters, such as the atomic size mismatch (δ) [24], the enthalpy of mixing (ΔH_{mix}) [25], the parameter Ω [26,27], and the valence electron concentration (VEC) [26,27]. The δ and ΔH_{mix} can be expressed as [24,25]:

\[ \delta = \sqrt{\sum_{i=1}^{n} c_i \left(1 - r_i / \bar{r}\right)^2} \]  

(1)  

\[ \Delta H_{mix} = 4 \sum_{i=1, i \neq j}^{n} \Delta H_{ij}^{mix} c_i c_j \]  

(2)

where \( c_i \) is the molar ratio of \( i \)th element, \( r_i \) is the atomic radius of \( i \)th element, \( \bar{r} \) is the average radius, given by \( \bar{r} = \sum_{i=1}^{n} c_i r_i \), and \( \Delta H_{ij}^{mix} \) is the enthalpy of mixing between \( i \)th and \( j \)th elements. The parameter δ describes the comprehensive effect of the atomic-size difference in the multicomponent alloy, while ΔH_{mix} reflects the tendency of forming stable intermetallic compounds. It was proposed that \( \delta \leq 6.6\% \) and \(-15 \leq \Delta H_{mix} \leq 5 \text{kJ/mol} \) can be used as criterion for forming solid solution phase of HEAs [28]. The parameter Ω and VEC can be expressed as [26,27]:

\[ \Omega = \frac{T_m \Delta S_{mix}}{|\Delta H_{mix}|} \]  

(3)  

\[ VEC = \sum_{i=1}^{n} c_i (VEC)_i \]  

(4)

where \( T_m \) is the average melting temperature, given by \( T_m = \sum_{i=1}^{n} c_i (T_m)_i \) and \( \Delta S_{mix} \) is the entropy of mixing of the alloy system, given by \( \Delta S_{mix} = -R \sum_{i=1}^{n} c_i \ln c_i \), \( R \) is the gas constant (\( R = 8.314 \text{ J/K/mol} \)), and \( (VEC)_i \) is the VEC of \( i \)th element. The parameter Ω reflects the impact of entropy effect and enthalpy effect, i.e., a high value of Ω indicates a dominant impact of entropy effect, while a low value of Ω means a dominant impact of enthalpy effect. Solid solution phase is suggested existing at \( \Omega > 1.1 \) [28]. The VEC act as an efficient parameter to predict phase structure of most of HEAs, and it is suggested that single BCC phase exists at \( VEC < 6.87 \) [27].

The above empirical parameters for the current TiNbTa0.5ZrAl4 RHEAs were calculated and shown in Figure 7. The δ increases with increasing Al content and reaches the maximum value at \( x = 0.75 \). Noted that all the values of δ are below 6.6%, indicating the atomic size mismatch has limited effect on the precipitation behavior. The ΔH_{mix} decreases with increasing Al content and reaches the threshold value for intermetallic phase at \( x = 0.75 \), which means that an excess addition of Al can lead to the formation of intermetallic phase. The phase equilibrium based on the calculation of enthalpy of mixing agrees well with the experimental results, demonstrating that the precipitation of HCP Zr5Al4 type intermetallic phase may be primarily induced by enthalpy effect. The values of ΔH_{mix} for atomic pairs between elements in Ti-Nb-Ta-Zr-Al system are given in Table 1 [14]. Obviously, the ΔH_{mix} between refractory elements is positive, but Al has large negative ΔH_{mix} with other refractory elements, especially with Zr, therefore, leading to the formation of Zr-Al intermetallic phase. Figure 7c exhibits the trend of parameter Ω with Al content increasing. When the Al content is low, the value of Ω is
match.

Table 1. Calculation of empirical parameters for the Al-contained RHEAs: (a) atomic size mismatch ($\delta$), (b) mixing of enthalpy ($\Delta H_{\text{mix}}$), (c) parameter $\Omega$, and (d) valence electron concentration (VEC).

Figure 7. Calculation of empirical parameters for the Al-contained RHEAs: (a) atomic size mismatch ($\delta$), (b) mixing of enthalpy ($\Delta H_{\text{mix}}$), (c) parameter $\Omega$, and (d) valence electron concentration (VEC).
Table 1. The values of $\Delta H_{\text{mix}}$ (kJ/mol) for atomic pairs between elements in Ti-Nb-Ta-Zr-Al system [14].

| Element | Ti | Nb | Ta | Zr | Al |
|---------|----|----|----|----|----|
| Ti      | -  | -  | -  | -  | -  |
| Nb      | 2  | -  | -  | -  | -  |
| Ta      | 1  | 0  | -  | -  | -  |
| Zr      | 0  | 4  | 3  | -  | -  |
| Al      | -30| -18| -19| -44| -  |

5. Conclusions

Precipitation behavior of the TiNbTa$_{0.5}$ZrAl$_x$ RHEAs was investigated. Accordingly, the following conclusions can be drawn:

(1) The powder metallurgy TiNbTa$_{0.5}$Zr RHEA has single BCC structure. Addition of Al may lead to the formation of Zr$_5$Al$_4$ type intermetallic phase.

(2) The TiNbTa$_{0.5}$ZrAl$_x$ RHEAs exhibit different precipitation behavior as a function of Al contents: for $x \leq 0.2$, the RHEAs form (Nb, Ta)-rich BCC2 precipitates and for $x \geq 0.5$, the RHEAs mainly form (Zr, Al)-rich HCP precipitates.

(3) Annealing at 800 $^\circ$C causes precipitation of an ordered B2 phase in TiNbTa$_{0.5}$ZrAl$_{0.5}$ RHEA, and the precipitation behavior is associated with spinodal decomposition. The ordered B2 precipitates have significant precipitation hardening effect on the TiNbTa$_{0.5}$ZrAl$_{0.5}$ RHEA.

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