The Polythermal Section of Ti-22Al-xNb (30-78 at.% Ti) in Ti–Al–Nb System

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Abstract: The polythermal section of Ti-22Al-xNb (30–78 at.% Ti) in the Ti-Al-Nb system was studied using X-ray diffraction analysis (XRD), differential thermal analysis (DSC), and electron probe micro-analysis (EPMA). No new ternary compounds were found in this work. The polythermal section has five three-phase regions, nine two-phase regions, and three single-phase regions. The O phase transition is confirmed to occur below 1000 °C. A four-phase invariant reaction \( \beta + \sigma \rightarrow O + \delta \) was found at 931 °C.

Keywords: Ti-Al-Nb system; phase diagram; polythermal section

1. Introduction

Because of their light weight, promising mechanical properties and good oxidation resistance [1], titanium–aluminum intermetallic alloys have become very important for the development of high-performance engines for airplanes [2]. The addition of an appropriate amount of niobium can significantly improve the oxidation resistance, the thermal deformation and hot creep properties of these alloys [3]. TiAlNb alloys are typical representatives of them [4,5]. The general composition of these alloys is usually Ti-(18–30 at.%) Al-(12.5–30 at.%) Nb. Many research works have been undertaken to fully understand the relationship between the composition, process, microstructure, mechanical properties, and performance of these alloys [6,7]. Among them, Ti-22Al-25Nb and Ti-22Al-27Nb have shown excellent potential for aeronautical applications [8,9]. To further improve the properties of these alloys, it is necessary to have a deeper understanding of the microstructure evolution during the thermomechanical processing. Therefore, the phase equilibrium information of the Ti-Al-Nb system is indispensable to model and to monitor the microstructure evolution of these alloys by accurately selecting the process temperatures.

Previous studies of Ti-Al-Nb systems have mostly focused on isothermal sections or the phase transformation of specific alloys [10–16]. Subsequently, calculations of the phase diagram (Calphad method) [17,18] were carried out by several researchers. In 1989, Kaltenbach et al. [10] obtained the first isothermal section of Ti-Al-Nb at 1200 °C experimentally. In 1992, Kattner and Boettinger [19] optimized the ternary system for the first time based on the existing information of the binary system. Due to the lack of thermodynamics data and phase equilibria, their results are questionable. In 1995,
Miracle et al. [20] obtained a partial polythermal section of Ti-22Al, but the Ti-Al boundary and the O phase region of the polythermal section are quite different from those reported later [21,22]. For example, Muraleedharan et al. [23] found that the upper temperature limit for the stability of the O phase is about 1000 °C. Above 1000 °C, the O phase disappears. However, in Miracle’s research [20], the O phase did not disappear until 1050 °C. In 2005, Raghavan [15] reviewed the phase relationships in the ternary system reported in the previous literature, and experimentally obtained a liquid surface projection diagram and the 1200, 1100, 1000 °C isothermal sections. In 2009, Witusiewicz et al. [17,18] re-evaluated the binary systems in the Ti-Al-Nb ternary system and re-optimized the ternary parameters of this ternary system. The calculated phase diagrams are consistent with the experimental results at that time. The same year, Cupid et al. [24] also independently completed the optimization of the Ti-Al-Nb ternary system. The calculated isothermal sections were in good agreement with the experimental results obtained at 1100 °C and 1200 °C, but differed from those of Witusiewicz’s work [18] in the β, the α, and the O phase regions. Raghavan [25] summarized the recent results again in 2010. In 2018, Li et al. [21] updated the isothermal sections at 1000 °C and 1150 °C again, and studied the phase relationship at 1100°C completely. They verified that the O phase was only stable below 1000 °C and the β0 phase was found to be stable at 1000 °C. However, the γ1 phase reported by Ding et al. [26] and Chen et al. [12] was not confirmed at 1000 °C. Table 1 lists the accepted phase names and crystal structures of the Ti-Al-Nb ternary system [18].

Although significant progress has been made in the phase diagram research of the Ti-Al-Nb ternary system, the existing phase diagram is not completely correct. For example, according to the polythermal section calculated by Witusiewicz et al. [18], the α phase does not appear in the Ti-22Al-23Nb alloy. However, after solution treatment at 1050, 1100 and 1150 °C for 2 h, and aging at 815 °C for 8 h, Rollett et al. [27] found α phase in this alloy. The composition range of the existing phase regions in the polythermal section are inconsistent with each other [18,20,24,28]. The purpose of this work is to determine the polythermal section of Ti-22Al-xNb (30-78 at.% Ti). The results will be useful for determining the heat treatment or hot working process of Ti-22Al-xNb alloys and to provide more accurate data for the optimization of the Ti-Al-Nb ternary system.

**Table 1. Crystal structure data of different phases in Ti-Al-Nb system.**

| Phase                  | Pearson Symbol | Space Group | Strukturbericht Designation | Prototype |
|------------------------|----------------|-------------|-----------------------------|-----------|
| (Al)(αAl), fcc_A1      | cF4            | Fm-3m       | A1                          | Cu        |
| α, (αTi), hcp_A3       | hp2            | P63/mmc     | A3                          | Mg        |
| α0, TiAl               | hP8            | P63/mmc     | D019                        | NiSn      |
| β, (βTi), bcc_A2       | cl2            | Im-3m       | A2                          | W         |
| β0, bcc_B2             | cl2            | Pm-3m       | B2                          | CsCl      |
| γ1, γTiAl, TiAl        | tP4            | P4/mmc      | L10                         | AuCu      |
| δ, Nb3Al               | cP8            | Pm-3n       | A15                         | CeSi      |
| ξ, (Ti3Nb3)Alβ         | tI8            | I4/mmm      | D022                        | TiAl(h)   |
| TiAl(h), TiAl3(h)      | tI32           | I4/mmm      | —                           | TiAl(l)   |
| ξ, Ti3Al2(x)           | tP28           | P4/mmc      | —                           | TiAlβ     |
| η, TiAl5               | tI24           | I4/mmm      | —                           | HiGa2     |
| θ, Nb3Al               | tP30           | P4/mmc      | D8β                         | αCrFe     |
| TaAl5                  | tP32           | P4/mmc      | —                           | TiAlβ     |
| O1, O, O1(h), Ti3NbAl  | oC16           | Cmcm        | —                           | NaHg      |
| O2, O2(r), Ti3NbAl     | oC16           | Cmcm        | —                           | NaHg      |
| τ, Ti3Nb2Al5           | hP6            | P63/mmc     | B82                         | Ni3In     |
| γ1-Ti3Nb2Alβ           | tP16           | P4/mmc      | —                           | γ1        |

This table lists the crystal structure data of different phases in the Ti-Al-Nb system. The Pearson symbols and space groups are provided for each phase, along with the Strukturbericht designations and prototypes. The table includes phases such as (Al)(αAl), fcc_A1, α, (αTi), hcp_A3, α0, TiAl, β, (βTi), bcc_A2, β0, bcc_B2, γ1, γTiAl, TiAl, δ, Nb3Al, ξ, (Ti3Nb3)Alβ, TiAl(h), and O1, O, O1(h), Ti3NbAl, among others. The table is a comprehensive resource for understanding the crystal structures of these phases in the Ti-Al-Nb system.
2. Materials and Method

Nine alloy components (listed in Table 2), 27 samples (listed in Table 3), each with a mass of approximately 12 g, were prepared by arc melting on a water-cooled copper crucible under high purity argon (99.99%). Single metals, Ti (purity of 99.99 at.%), Al (purity of 99.99 at.%), and Nb (purity of 99.99 at.%), were supplied by China JinYu Materials Technology Co. All samples were weighed to a precision of 0.001 g and formulated to contain 22 at.% Al. At the same time, to ensure the uniformity of the alloys, the sample buttons were melted, turned over and re-melted at least six times. The mass loss of each sample was less than 1%. Then, the obtained samples were evenly divided into four parts, three of which were sealed into different evacuated quartz capsules back-filled with high purity argon (99.99%) together with Ti chips as oxygen getters. Then, one of three parts was annealed at 1000 °C for 720 h. Another part was annealed at 900 °C for 1440 h and the last part was annealed at 800 °C for 2160 h in laboratory muffle furnaces. After annealing, samples were quenched in ice water to freeze the microstructure obtained during the annealing at the considered temperatures.

Table 2. The designed alloy compositions in the present work.

| No. | Ti | Al | Nb |
|-----|----|----|----|
| 1   | 73 | 22 | 5  |
| 2   | 68 | 22 | 10 |
| 3   | 63 | 22 | 15 |
| 4   | 58 | 22 | 20 |
| 5   | 53 | 22 | 25 |
| 6   | 48 | 22 | 30 |
| 7   | 43 | 22 | 35 |
| 8   | 38 | 22 | 40 |
| 9   | 33 | 22 | 45 |

The annealed samples were mechanically polished to remove contaminated surface layers. Electron Probe Microanalysis (EPMA) (JEOL JXA-8530F, Japan Electron Optics Laboratory CO., Ltd, Tokyo, Japan) was used to observe their microstructure and to determine the chemical compositions of single phases. Based on the EPMA composition results, the existing phases in the alloys were preliminarily identified.

To further confirm the phases in each sample, X-ray diffraction analysis (XRD) was carried out on powder and bulk materials using an X-ray diffractometer (Rigaku D-MAX/2550 VB, Japan Rigaku Co., Tokyo, Japan) with Cu Kα radiation (40 kV and 250 mA). The scanning range was 10–90° with a speed of 8°/min. The XRD results were analyzed using the program Jade 6.0 (Materials Data Inc., California Grown, CA, USA).

Differential scanning calorimetric (DSC) analysis was conducted either on a Labsys EVO (Setaram Inc., Lyon, France) or a Netzsch 449 F3 (Netzsch-Gerätebau GmbH, Selb, Germany) to determine the temperature of phase transformations. The experiments were performed under high purity argon (99.99%), using Al2O3 crucibles and platinum crucibles. The heat rate of 20 °C/min was selected as it ensures the best phase transition initiation during thermal analysis, as the driving force is higher in a shorter time. The choice of the crucibles was discussed by Kostov et al. [29], who showed that the crucible stability depends on the titanium activity and temperature. As the activity of Al and Ti in the alloy is lower than one, the oxidation pressure is then higher compared to those of the Al/Al2O3 and Ti/TiO2 equilibria, respectively [30]. Therefore, the alloy does not react with the crucible. The DSC curve remains stable throughout the heating process. Before the measurements, the apparatus was calibrated using high purity calibration standards including low-melting pure metals (Sn (213.9 °C), Zn (419.53 °C), etc.), medium-melting metals (Al (660 °C), Ag (961.78 °C), etc.) and high-melting metals (Au (1064.18 °C), Cu (1083.4 °C), etc.). Since the Ti-Al-Nb system was studied in this work, pure Ti metal was also used during the calibration process. Unlike other pure metals, with pure Ti metal, careful attention was paid to check whether it could have been contaminated by an
unwanted reaction with the crucible materials. It was proven that pure Ti and crucibles remained stable during the experiments, and no reaction occurred.

3. Results

3.1. Analysis of DSC Results

Prior to DSC analysis, all samples were annealed at 1000 °C for 720 h, 10–20 mg thin slices were cut from the ingots, and placed with forceps in the DSC instrument and it was subsequently flushed with argon. The DSC heating curves are shown in Figure 1. The temperature that corresponds to the intersection of the horizontal baseline and the tangent line of the endothermal peak is marked as the phase transformation temperature.

As shown in Figure 1, only one endothermal effect was found in the temperature heating curves of alloy #5. According to the results of Miracle et al. [20], this was ascribed to the phase transformation O + β → β. In alloys #6, #7, #8, and #9, an endothermic reaction was observed at the same temperature (931 °C). According to the experiment and calculation results of Witusiewicz et al., it is preliminarily speculated that this reaction is a four-phase reaction of β + σ ↔ O + δ. The subsequent EPMA and XRD experimental results also support this inference.
3.2. Analysis of Typical Alloys in Polythermal Section

To accurately analyze the phase transformation which occurred at each temperature peak observed in the DSC curves, alloys were annealed at 800, 900, and 1000 °C for 720, 1440, and 2160 h, respectively. The corresponding microstructures and phase compositions were investigated by means of EPMA, and the phases were identified by XRD analysis.

Based on the experimental data obtained by EPMA and XRD, Table 3 summarizes the phase constitution and corresponding phase compositions in all equilibrium alloys at 800, 900, 1000 °C, respectively. Meanwhile, we used imagetool to roughly calculate the phase percentage in the samples. These are also listed in Table 3. The phase relationships and transformations of several key alloys are discussed below.

Table 3. Equilibrium compositions and phases determined in the present work.

| No. | Heat Treatment | Phase Constitution | Phase Composition, at.% | Phase Fraction |
|-----|----------------|---------------------|--------------------------|---------------|
|     |                | Phase 1/2/3         | Phase 1 | Phase 2 | Phase 3 | Phase1:2:3 |
|     |                | α/β                 | Al   | Ti     | Al   | Ti     |            |
| 1   | 1000 °C 720 h  | β/αζ                | 16.7 | 75.9   | 23.7 | 74.3   | 0.53:0.47  |
| 2   | 900 °C 1440 h  | αζ                  | 21.6 | 73.7   | -    | -      | -           |
|     | 800 °C 2160 h  | αζ                  | 21.6 | 75.8   | -    | -      | -           |
| 3   | 1000 °C 720 h  | β/αζ                | 18.4 | 68.3   | 23.3 | 68.3   | 0.58:0.42  |
|     | 1440 h         | β/αζ                | 14.3 | 67.9   | 23.4 | 67.9   | 0.59:0.41  |
|     | 2160 h         | αζ/αζ               | 22.5 | 68.9   | 18.0 | 68.9   | 0.42:0.58  |
| 4   | 1000 °C 720 h  | β/αζ                | 20.4 | 61.9   | 24.6 | 65.3   | 0.60:0.40  |
|     | 1440 h         | β/αζ                | 15.7 | 61.3   | 23.4 | 64.7   | 0.62:0.38  |
|     | 2160 h         | αζ/αζ               | 22.7 | 64.9   | 15.3 | 61.2   | 0.43:0.57  |
| 5   | 1000 °C 720 h  | β/αζ                | 20.5 | 53.5   | 24.5 | 61.2   | 0.63:0.37  |
|     | 1440 h         | β/αζ                | 15.6 | 56.8   | 23.5 | 62.9   | 0.60:0.33  |
|     | 2160 h         | β/αζ                | 18.3 | 52.1   | 23.0 | 59.4   | 0.58:0.02  |
| 6   | 1000 °C 720 h  | β/αζ                | 22.5 | 53.2   | -    | -      | 1           |
|     | 1440 h         | β/αζ                | 23.9 | 53.9   | 15.7 | 52.3   | 0.57:0.43  |
|     | 2160 h         | β/αζ                | 23.9 | 54.5   | 19.4 | 52.8   | 0.60:0.40  |
| 7   | 1000 °C 720 h  | β/αζ                | 22.3 | 47.0   | 26.8 | 57.8   | 0.64:0.36  |
|     | 1440 h         | β/αζ                | 23.1 | 50.0   | 15.2 | 47.0   | 0.55:0.45  |
|     | 2160 h         | β/αζ                | 24.8 | 49.5   | 18.6 | 46.7   | 0.61:0.39  |
| 8   | 1000 °C 720 h  | β/αζ                | 21.5 | 44.8   | 27.2 | 56.8   | 0.67:0.33  |
|     | 1440 h         | β/αζ                | 23.3 | 52.1   | 14.6 | 47.7   | 0.04:0.40  |
|     | 2160 h         | β/αζ                | 22.6 | 45.1   | 17.7 | 42.1   | 0.58:0.42  |
| 9   | 1000 °C 720 h  | β/αζ                | 19.5 | 46.7   | 25.0 | 59.1   | 0.11:0.37  |
|     | 1440 h         | β/αζ                | 14.7 | 46.9   | 23.6 | 52.8   | 0.02:0.41  |
|     | 2160 h         | β/αζ                | 23.7 | 35.3   | 24.8 | 42.5   | 0.01:0.41  |
| 10  | 1000 °C 720 h  | β/αζ                | 19.3 | 42.4   | 22.6 | 54.3   | 0.52:0.09  |
|     | 1440 h         | β/αζ                | 22.8 | 50.1   | 20.3 | 32.5   | 0.41:0.59  |
|     | 2160 h         | β/αζ                | 25.8 | 43.3   | 19.3 | 33.2   | 0.40:0.60  |
The Nb concentration in each phase can be calculated as 100 (Al concentration in at.% + Ti concentration in at.%).

Alloy #1 was used to verify the transformation process of $\alpha \leftrightarrow \alpha + \beta$. According to the DSC measurements (Figure 1 #1), the phase transformation should occur at 950 °C. The microstructure of annealed alloy #1 and the corresponding XRD analysis results after heat treating at 900 and 1000 °C are shown in Figure 2a,b and Figure 2c,d, respectively. If we compare (a)/(c) and (b)/(d) in Figure 2, the $\alpha_2$ phase transforms into the $\alpha_2 + \beta$ phase between 900 and 1000 °C. Therefore, it can be concluded that the phase transformation of $\alpha_2$ to $\alpha_2 + \beta$ occurs at 950 °C.

![Figure 2](image-url)

**Figure 2.** Electron probe micro-analysis (EPMA) images and XRD results of alloy #1 after annealing at 1000 °C, 900 °C. (a) The microstructure of alloy #1 at 1000 °C; (b) the XRD result of alloy #1 at 1000 °C. (c) The microstructure of alloy #1 at 900 °C; (d) the XRD result of alloy #1 at 900 °C.

Alloy #4 was designed to confirm the transformation of the ternary $\beta + \alpha + O$ phase region. The DSC results (see Figure 1, #4) show that the alloy phase transformations took place at 821, 932, and 976 °C. Figure 3 shows the corresponding microstructures and XRD patterns of this alloy after annealing at 800, 900 and 1000 °C. The alloy is composed of three phases: the dark $\alpha_2$ phase, the gray O phase and the white $\beta$ phase at 800 and 900 °C. When the temperature reaches 1000 °C, the gray O phase disappears, whereas the dark $\alpha_2$ and the white $\beta$ phases remain.
Figure 3. EPMA images and XRD results of alloy #4 after annealing at 1000, 900, 800 °C. (a) The microstructure of alloy #4 at 1000 °C; (b) the XRD result of alloy #4 at 1000 °C. (c) The microstructure of alloy #4 at 900 °C; (d) the XRD result of alloy #4 at 900 °C. (e) The microstructure of alloy #4 at 800 °C; (f) the XRD result of alloy #4 at 800 °C.

At 800 °C, the α2 and the O phases are formed as a small litter of ellipsoids and fine needles, respectively, and are homogeneously dispersed across the β matrix. They exhibit the classic microstructures of α2-based titanium alloys [31]. After annealing at 900 °C, the α2 phase gathers in a lamellar shape at the grain boundaries. At the same time, the O phase was changed into short rods and distributed throughout the β matrix. It can be seen that the decomposition of O phase into β + α2 occurs at 900 and 1000 °C.

The annealed microstructure of alloy #5 at 800 and 900 °C (Figure 4) exhibits the β and O phases. Moreover, when the annealing temperature was raised from 800 to 900 °C, the volume fraction of the β phase increased significantly and the O phase gradually changed into a short, rod-like shape. However, at 1000 °C (Figure 4), only the β single phase was observed in the annealed microstructure. This means that the phase transformation occurred in the 900–1000 °C temperature range. Combining the DSC results, the phase transformation in alloy #5, β + O ⇌ β, was confirmed. The phase transformation temperature was found to be close to 966 °C.
According to the DSC results, alloys #6, #7, #8, and #9 experienced the same phase transformation at 931 °C. The annealed microstructure of alloy #7 is shown in Figure 5. The annealed microstructure at 800 °C contains only the β and O phases, in which the two phases are interweaved and the β phase is the matrix phase. A three-phase equilibrium composed of the gray β phase, dark O phase, and the white δ phase was found in the annealed structure at 900 °C, where the O phase was distributed at the grain boundary or in the β phase. Equally, the annealed structure at 1000 °C is made of black β and white δ phases and the black β phase is dispersed in the white δ phase matrix. It can be ascertained that the four-phase reaction $\beta + \sigma \rightarrow O + \delta$ takes place between 900 and 1000 °C.
°C. Xu [22] et al. measured the isothermal cross sections at 980 and 900 °C. Through their analysis of the isothermal sections, they also found the four-phase reaction of \( \beta + \sigma \leftrightarrow O + \delta \), which is in agreement with the current experimental results. At the same time, since the alloy undergoes a four-phase reaction of \( \beta + \sigma \leftrightarrow O + \delta \) at 931 °C and the \( \sigma \) phase is clearly seen at 1000 °C, the phase transition peak at 986 °C should include the transformation of the \( \beta + \sigma \) and \( \beta + \sigma + O \) peaks. The \( \beta + \sigma/\beta \) transition peak should be at a higher temperature and \( \beta + O \leftrightarrow \delta \) should occur at 878 °C.

Figure 5. EPMA images and XRD results of alloy #7 after annealing at 1000 °C, 900 °C, 800 °C. (a) The microstructure of alloy #7 at 1000 °C; (b) the XRD result of alloy #7 at 1000 °C. (c) The microstructure of alloy #7 at 900 °C; (d) the XRD result of alloy #7 at 900 °C. (e) The microstructure of alloy #7 at 800 °C; (f) the XRD result of alloy #7 at 800 °C.
Figure 6 show the annealed microstructures and XRD diffraction analysis patterns of alloy #8. The annealed structures of alloy #8 more directly prove the existence of the $\beta + \sigma \leftrightarrow O + \delta$ four-phase reaction. At 900 °C, the alloy is in the gray $\beta$ phase, dark black $O$ phase and white $\delta$ phase; while, at 1000 °C, the alloy structure has changed to the black $\beta$ phase, dark gray $O$ phase and white $\sigma$ phase. The changes in the two structures indicate that there is a phase transformation. Combined with the changes in alloy #7, it can be confirmed that there is a $\beta + \sigma \leftrightarrow O + \delta$ four-phase reaction at 931 °C.

**Figure 6.** EPMA images and XRD results of alloy #8 after annealing at 1000 °C, 900 °C, 800 °C. (a) The microstructure of alloy #8 at 1000 °C; (b) the XRD result of alloy #8 at 1000 °C. (c) The microstructure of alloy #8 at 900 °C; (d) the XRD result of alloy #8 at 900 °C. (e) The microstructure of alloy #8 at 800 °C; (f) the XRD result of alloy #8 at 800 °C.
3.3. Polythermal Section

Based on the above experimental results, the polythermal section in the Ti-Al-Nb ternary system covering the composition range of 30–78 at.% Ti was established and is illustrated in Figure 7. The experimental data in this work are highlighted in red and compared with some reported ones from the literature [17,21,22] (in blue). The phase-transformation temperatures in our experiments were derived from the DSC and symbolled using red dots. The experimental alloy compositions are indicated by red dotted lines. The BCC$_A2$-BCC$_B2$ phase transition could not distinguished in the present work and is therefore not drawn in the diagram. The polythermal section consists of five three-phase regions: $\alpha + \beta + \alpha_2$, $\alpha + \beta + O$, $\beta + O + \delta$, $\beta + O + \sigma$, and $\delta + \beta + \sigma$; nine two-phase regions: $\alpha + \beta$, $\alpha + \alpha_2$, $\alpha + \beta$, $O + \alpha_2$, $\beta + O$, $\beta + \sigma$, $O + \delta$ and $\delta + \beta$; three single-phase regions: $\alpha$, $\alpha_2$ and $\beta$.

![Figure 7. Overview of the polythermal section of the Ti-Al-Nb ternary system using data from this work and from [18,21,22].](image-url)

4. Discussion

Combining the microstructure and XRD results, the phase transformation at different temperatures were ascribed. The results are summarized in Table 4.
The boundary between the two sections and the temperature at which the α phase from the β phase is 1120 °C. The experimental results of this work exhibit agreement with the research results of Witusiewicz et al. [18]. The experimental results of this work exhibit good agreement with the results of Miracle et al. [20], when the Nb content is zero (the binary boundary conditions of Ti-Al), the precipitation temperature of the α phase from the β phase is 1120 °C. The temperature at which β transforms into α and α + β is 1075 °C whereas the α phase is converted into α at 1000 °C. However, this is quite different from the recent research results of Witusiewicz et al. [18]. The experimental results of this work exhibit good agreement with those of Witusiewicz [18]. Explicitly, this means that the α single-phase region appears in the section and the temperature at which the α phase drops to 900 °C. Thus, the position of the three-phase region of α + β + α2 is closer to the binary boundary. The boundary between the two-phase region of β + α2 and the three-phase region is steeper.

| No. | Composition       | Crucible Type | Mass (mg) | Rate (°C/min) | Phase Transformation | Temperature (°C) |
|-----|-------------------|---------------|-----------|---------------|----------------------|-----------------|
| 1   | Ti74.3Al21.4Nb3.6 | Al2O3         | 13.56     | 20            | \(\alpha2 \leftrightarrow \alpha2 + \beta\) \(\alpha2 + \beta \leftrightarrow \beta\) | 950(±10)        |
| 2   | Ti80.3Al22.2Nb9.5| Al2O3         | 17.75     | 20            | \(\alpha2 \leftrightarrow \alpha2 + \beta\) \(\alpha2 + \beta \leftrightarrow \beta\) | 1110(±10)       |
| 3   | Ti83.2Al22Nb4.8   | Al2O3         | 14.45     | 20            | \(\beta + O \leftrightarrow \alpha2 + \beta + O\) \(\alpha2 + \beta + O \leftrightarrow \alpha2 + \beta\) | 748(±10)        |
| 4   | Ti86.4Al22Nb21.6  | Al2O3         | 18.73     | 20            | \(\alpha2 + \beta + O \leftrightarrow \alpha2 + \beta\) \(\alpha2 + \beta \leftrightarrow \beta\) | 882(±10)        |
| 5   | Ti83.2Al22.8Nb0.8 | Al2O3         | 26.67     | 20            | \(\beta + O \leftrightarrow \beta\) \(\beta + \sigma \leftrightarrow \beta + \sigma\) | 966(±10)        |
| 6   | Ti85Al22.4Nb29.8  | Al2O3         | 22.23     | 20            | \(\beta + O \leftrightarrow \beta + O + \sigma\) \(\beta + \sigma \leftrightarrow \beta + \sigma\) | 931(±10)        |
| 7   | Ti45.5Al21.4Nb33.1| Pt            | 14.78     | 20            | \(\beta + O \leftrightarrow \beta + O + \delta\) \(\beta + \delta \leftrightarrow \beta + \delta\) | 878(±3)         |
| 8   | Ti38.7Al21.2Nb9.1 | Pt            | 15.67     | 20            | \(\beta + O + \sigma \leftrightarrow \beta + \sigma + O\) \(\beta + \sigma \leftrightarrow \beta + \sigma\) | 931(±3)         |
| 9   | Ti39.9Al21.8Nb5   | Al2O3         | 11.96     | 20            | \(\beta + \sigma \leftrightarrow \beta + \sigma\) \(\delta + \sigma \leftrightarrow \delta + \sigma\) | 1191(±10)       |

Figure 8 shows a comparison of the experimental results of the Ti-22Al polythermal section of Miracle et al. [14] with the results of this work. As shown in the results of Miracle et al. [20], when the Nb content is zero (the binary boundary conditions of Ti-Al), the precipitation temperature of the α phase from the β phase is 1120 °C. The temperature at which β transforms into α and α2 is 1075 °C whereas the α phase is converted into α at 1000 °C. However, this is quite different from the recent research results of Witusiewicz et al. [18]. The experimental results of this work exhibit good agreement with those of Witusiewicz [18]. Explicitly, this means that the α single-phase region appears in the section and the temperature at which the α phase drops to 900 °C. Thus, the position of the three-phase region of α + β + α2 is closer to the binary boundary. The boundary between the two-phase region of β + α2 and the three-phase region is steeper.
The O phase was not observed at 1000 °C in this work. The phase region containing the O phase will not appear at this temperature, that is, the β + O, α2 + β + O phase region should be shifted to lower temperatures. This result is consistent with the experimental results of Chen et al. [12], Ding et al. [26], and Li et al. [21] obtained at 1000 °C and Xu et al. [22] measured at 980 °C.

Figure 9 shows the calculated polythermal section from Witusiewicz’s work [18] compared with the present results. However, the calculated results differ from the current experimental results in terms of the four-phase reaction temperature, the size of the δ + β + σ three-phase region and the location of the α2 + β + O three-phase region. The four-phase reaction temperature calculated by Witusiewicz et al. [18] is 954 °C, whereas the temperature we experimentally measured is 931 °C. Compared with the Ti-22Al-48Nb boundary calculated by Witusiewicz et al. [18], it was found that the phase region above 921 °C is not only the δ + β phase region, but also the δ + β + σ and the δ + σ phase regions. According to the experimental data of Li et al. [21], Xu et al. [22], the Ti-22Al-48Nb alloy is located in the δ + σ phase region at 1000 °C, in the δ + β + σ phase region for temperatures between 1100 and 1300 °C. When the temperature is raised to 1400 °C, it is converted to the single β phase. This work is consistent with their results. In addition, it was also determined in this work that the α2 phase does not disappear until the Nb content in the alloy is increased up to 24 at.%, instead of the 21 at.% Nb in Witusiewicz’s calculations [18]. At the same time, Witusiewicz et al. described the O phase as the O1 and O2 phases and optimized the two phases independently during the optimization process. However, in their experiments, they failed to distinguish the two O phases. In this work, the two phases cannot be distinguished, so the O phase is also used as a unified representation.
Figure 9. Experimentally determined the polythermal section of the Ti-Al-Nb ternary system compared with the calculated results of Witusiewicz et al. [18].

5. Conclusions

The polythermal section of the Ti-22Al-Nb (30–78 at.% Ti) in the Ti-Al-Nb ternary system was determined using electron probe microanalysis (EPMA), X-ray diffraction (XRD) and differential scanning calorimetric (DSC).

The polythermal section has five three-phase regions, nine two-phase regions, and three single-phase regions. Compared with the literature results, the $\alpha_2 + \beta + O$ three-phase region moves to a lower temperature and the $\alpha + \beta + \alpha_2$ three-phase region is shifted to the Ti-Al binary boundary. As a result, all the phase boundaries change. The O phase is confirmed to occur below 1000 °C. There are differences between the present experimental work and Witusiewicz's calculated results. A four-phase invariant reaction was found at 931 °C: $\beta + \sigma \rightarrow O + \delta$. This polythermal section will help in the thermodynamic evaluation of the Ti-Al-Nb system and the practical application of O-phase alloys.

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