Phase Stability of Water-Quenched U-6 wt.% Nb Alloy During Low-Temperature Aging

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Abstract
TEM analyses, microhardness measurements, and tensile testing were carried out to investigate the microstructure and phase stability of a water-quenched (WQ) U-6 wt.% Nb alloy subjected to low-temperature aging experiments that include (i) 200 °C aging of as-water-quenched alloy, (ii) Ambient-temperature aging of WQ alloy for 15 years (i.e., 15-year-old WQ alloy), and (iii) Accelerated aging of 15-year-old WQ alloy at 200 °C and 212 °C. Age hardening occurred during the initial stages of 200 °C aging of as-water-quenched alloy containing mainly an α” (monotonic) martensite. The microhardness continuously increased with aging time up to 8 hours because of the onset of a spinodal decomposition to form a modulated structure with the modulation wavelength ~3 nm propagating along the [001]α” direction. A decrease in microhardness occurred when coarsening of the modulated structure took place after thermally aged for more than 8 hours. TEM observations of swirl-shape antiphase domain boundaries (APBs) formed in the 15-year-old WQ alloy, however, revealed the occurrence of a chemical-ordering reaction in the α” (monotonic) martensite. The chemical ordering reaction can be attributed to the development of a composition fluctuation with the wavelength ~0.5 nm propagating along the [001] α” direction. As a result of the chemical ordering, the 15-year-old WQ alloy revealed an aging behavior distinctly different from that of the as-water-quenched alloy during low-temperature aging. The microhardness continuously increased to much higher values after prolonged aging up to 240 hours at 200 °C and 96 hours at 212 °C. The accelerated aging of the 15-year-old WQ alloy resulted in the decomposition of the chemically ordered α” martensite into a Nb-depleted α (orthorhombic) phase and a Nb-enriched ordered phase, which resulted in precipitation hardening and gave rise to a significant ductility reduction. Aging mechanisms and transformation sequences are accordingly proposed to depict the phase stability of WQ-U6Nb alloy during low-temperature aging.

Introduction
It is known that uranium-niobium alloys, as exploited for a variety of engineering and structural applications, has a microstructure containing martensitic phases supersaturated with Nb that can be obtained by rapid quenching the alloy from γ (bcc)-field solid solution to room temperature [1,2]. U-6 wt.% Nb alloy contains approximately 14 atomic percent of niobium that is close to the monotectoid composition (13.3 at.% Nb). At temperatures...
above 647 °C, the equilibrium phase is a γ phase [body-centered cubic (bcc); a = 0.3347 nm]. A water-quenched U-6 wt.% Nb (WQ-U6Nb) from the γ (bcc)-field solid solution has a microstructure containing martensitic phases that are supersaturated with 14 at.% Nb. It is worth noting that the crystal lattice of supersaturated solid solution formed by rapid quenching is unstable and is severely distorted since the solubility of Nb in the α phase (orthorhombic; a = 0.285 nm, b = 0.5861 nm, c = 0.4947 nm) at room temperature is nearly zero in stable equilibrium [1]. In the stresses-induced martensitic transformation, a variant of the low-temperature orthorhombic (α) phase is usually formed. This phase is designated α′ martensite since its lattice parameters (orthorhombic; a = 0.285 nm, b = 0.5802 nm, c = 0.4988 nm) differ from the equilibrium α phase. Two additional variant phases, a monoclinic distortion of α′, named α″ martensite (monoclinic; a = 0.285 nm, b = 0.576 nm, c = 0.4997 nm, γ = 91.3°), or a tetragonal distortion of γ, named γ° martensite (tetragonal; a = b = 0.701 nm, c = 0.674 nm), can also form WQ-U6Nb alloy. The WQ-U6Nb alloy has improved mechanical properties (ductility and toughness) [3,4] and excellent corrosion resistance [2,5] because of the uniform distribution of niobium atoms in the solid solution that suppresses the diffusional decomposition reaction to form a two-phase (Nb-depleted α phase and Nb-enriched γ phase) cellular microstructure. It is also known that the water-quenched U6Nb alloy containing α″ martensite, which consists of many different variants of fine twins, reveals shape memory effect that results in low yield strength (~200 MPa) and high tensile ductility (~30%) [2,6,7]. Thermal aging of the α″ martensite in a temperature ranging from 250 °C to 400 °C resulted in an increase of yield strength [3,4,8,9]. Using Atom Probe Field Ion Microscopy (APFIM) method, Beverini and Edmonds [4] found an age hardening/softening phenomenon associated with nanometer length-scale of segregation reaction presumably due to spinodal decomposition [10]. According to their investigation, an age hardening/softening phenomenon associated with nanoscale solute segregation presumably caused by spinodal decomposition was detected. In the alloy samples aged at 300 °C for 16 hours, Nb content was fluctuated between 5 at.% and 30 at.%, and the wavelength of the composition fluctuation was reported to be about 3 nm. Preliminary TEM analyses that clarified the formation of fine-scale modulated microstructure due to spinodal decomposition in WQ-U6Nb thermally aged at 200 °C were reported by Zhou and Hsiung [11]. Note that spinodal decomposition is a continuous transformation mechanism in which the supersaturated solid solution separates spontaneously into solute-lean and solute-rich phases within the parent phase-domain through uphill diffusion [12]. It is a phase transient mechanism leading to aging hardening and subsequent softening when the modulated structure becomes coarsened.

The structural components made of WQ-U6Nb alloy have been in service for more than two decades. The main concerns with these components are how the microstructure and mechanical properties have changed, and how will these changes proceed in the future. To address these issues, there is a need to know how the microstructure has evolved since the alloy was fabricated and put into service decades ago. There is also a need to know how the microstructure evolution will affect the mechanical stability. In this study, aging behavior of as-water-quenched alloy thermally aged at 200 °C was investigated to compare with the age behavior of 15-year-old WQ alloy thermally aged at temperatures below 250 °C. The emphasis has been placed on the low-temperature aging mechanisms of WQ-U6Nb alloy.

**Experimental**

The U6Nb alloy used for this investigation was wrought-processed from Rocky Flats VAR VAR ingot at the BWXT/Bechtel Y-12 plant. The detailed information regarding the fabrication process can be found in Ref. [2]. As-water-quenched U6Nb alloy samples were obtained by solution-treating the wrought U6Nb alloy at 800 °C for 4 hours followed by quench in water. The WQ-U6Nb alloy samples naturally aged at ambient temperatures for 15 years (designated as 15-year-old WQ alloy hereafter) were machined directly from a 38-mm thick plate. Rod-shaped specimens of 3 mm in diameter were prepared for microstructure analyses. Aging experiments of both as-water-quenched and 15-year-old alloy samples were carried out at 200 °C and 212 °C in a vacuum furnace. Microhardness measurements were performed on all these samples using a Vickers-hardness indenter. This was conducted to monitor the change of mechanical property during aging processes. The applied load was 50 g, and the time-duration was 35 seconds for...
Experimental Results

The result of microhardness (in HV unit) measured from as-water-quenched alloy samples thermally aged at 200 °C is shown in Figure 1, in which the results of microhardness measured from accelerated heating of the 15-year-old WQ alloy samples at 200 °C and 212 °C are also shown for a comparison. Age hardening can be clearly seen from both as-water-quenched and 15-year-old WQ alloy samples because of the thermal aging. However, distinctly different aging behaviors are shown between the as-water-quenched and the 15-year-old WQ alloy samples. Unlike the as-water-quenched alloy that reached peak hardness after heating for 8 hours, the microhardness of 15-year-old WQ alloy continuously increased even after prolonged aging with the microhardness increasing to a much greater value compared to that of the as-water-quenched alloy. The different aging behaviors suggest that different aging mechanisms take place in the as-water-quenched and the 15-year-old WQ alloys, which will be further elucidated below.

Thermal aging of as-water-quenched U6Nb at 200 °C

The as-water-quenched alloy had an average microhardness (in HV unit) of 190. The microhardness value increased to 210 after thermal aging for 2 hours, increased to 237 after thermal aging for 4 hours, and increased to 254.8 after thermal aging for 8 hours. Within the first 8 hours of aging, the strengthening continued but with a decreasing trend. Further aging caused softening after the microhardness reached a maximum value at 8 hours, and the microhardness reduced to 237 at 16 hours. While no notable change of the lamellar microstructure of α′′ martensite can be found from the low-magnification images shown in Figure 2, a modulated structure of dark/bright fringes in association with satellite spots excited on either side of the main Bragg diffraction spots can be readily seen in the high-magnification phase-contrast images shown in Figure 3, which verify the occurrence of spinodal decomposition. The modulation formed because of the wave of Nb fluctuation with a wavelength of 3 nm propagating predominantly...
Figure 2: Low-magnification TEM images show the lamellar microstructure of α” martensite in the as-water-quenched (WQ) alloy and in the alloy samples after thermally aged at 200 °C for 2 hours (2 h) and 16 hours (16 h). Selected-area diffraction (SAD) patterns of the [112]_α”- and [312]_α”-zone were inserted to demonstrate the existence of α” phase in the WQ and the 16 h alloy samples.

Figure 3: Phase-contrast TEM images show the development of a modulated structure in WQ-U6Nb after thermally aged at 200 °C for 2 hours (2 h), 4 hours (4 h), 8 hours (8 h), and 16 hours (16 h). Satellite diffraction spots arising from the modulated structure with the wavelength of 3 nm are marked by arrows shown in the [010]_α”- and [110]_α”-zone diffraction patterns generated from the 2 h, the 4 h, and the 8 h samples. Coarsening of the modulated structure was observed from the 16 h sample.
along the elastically softest direction: The [001]$_{α''}$ direction. As a result, the modulated structure contains Nb-lean and Nb-enriched phase-domains of 3 nm thick accompanied with satellite reflections in electron diffraction patterns can be observed using TEM techniques [13]. The satellite diffraction spots formed in the direction of modulation because of the contribution of lattice parameter variation [14]. After the modulated structure was fully developed at 8 hours, further thermal aging led to coarsening of the modulated structure and decrease in microhardness.

**Ambient-temperature aging of WQ-U6Nb for 15 years**

Chemical ordering of the $α''$ martensite in the 15-year-old WQ alloy was clarified using TEM phase analysis and electron diffraction techniques. Like the as-water-quenched alloy, the lamellar microstructure of $α''$ martensite in the 15-year-old WQ alloy remained no change, as shown in Figure 4a. However, a unique feature observed from the 15-year-old alloy is the formation of swirl-shape antiphase domain boundaries (APBs) shown in Figure 4b and Figure 4c. As will be demonstrated later that the APBs (also known as π boundaries) can be verified based on the following visibility criteria: The contrast of APBs is visible when the phase angle $α = 2π$ is invisible when $α = 2π$ (where $g$ is the reflection vector for imaging, and $P$ is the displacement vector of APB) [13]. The observation of swirl-shape APBs in the 15-year-old alloy reveals a continuous order-disorder transition takes place in the $α''$ martensite. Since the ordering transition occurs at ambient temperatures, crystal structure of the ordered $α''$ martensitic phase likely remains monotonic as that of the disordered $α''$ martensitic phase except that the unit cell takes up a superlattice arrangement due to the occupation of lattice sites by specific atom species. An APB is produced by an antiphase vector $P$ which brings about the displacement of one atomic species from one sublattice to another. Crystal structure of the ordered $α''$ martensitic phase can therefore be identified using TEM phase-analysis method by matching the observed and simulated diffraction patterns.

To explain the formation of APBs in the 15-year-old WQ alloy, a chemically ordered $α''$ martensitic phase is accordingly proposed. The existence of ordered $α''$ phase can be verified by matching the observed diffraction pattern with the simulat-
Figure 5: Two proposed schemes of chemical ordering taken place in the 15-year-old U6Nb: a) Scheme I and b) Scheme II; c) Scheme II ordered superlattice can be regarded because of a spinodal modulation from which the Nb content fluctuates between 0 and 28 at.% and the wavelength ($\lambda$) of the composition fluctuation is equal to the lattice parameter of the ordered $\alpha''$ unit cell in the [001] direction, i.e. $\lambda = 0.497$ nm.

Figure 6: Microhardness variation: a) And microstructure evolution; b,c,d) Of the 15-year-old WQ alloy thermally aged at 200 °C up to 240 hours.
ed diffraction patterns generated according to the proposed superlattice structures. Figure 5 illustrates two possible schemes for the order-disorder transition in the supersaturated $\alpha''$ solid solution with a mean concentration $c_o = 14$ at.% Nb. Here, the probability of random site occupancy on each atomic site of the disordered $\alpha''$ unit cell (monoclinic; $a = 0.285$ nm, $b = 0.576$ nm, $c = 0.4997$ nm, $\gamma = 91.3^\circ$; 4 lattice sites/unit cell) by U and Nb is 86% and 14%, respectively. Note that for the con-

![Diffraction Patterns]

**Figure 7:** Phase analysis of the ordered phase that revealed the swirl-shaped APB in the 15-year-old WQ-U6Nb: a) Scheme I and Scheme II have the same selected-area diffraction (SAD) patterns of the [110]-, the [310]-, and the [312]-zone; b) Simulated diffraction pattern of Scheme II is a better match with the SAD pattern of the [100]-zone.
venience of viewing purpose, the unit cell is illustrated herewith the coordinate origin (0,0,0) locating at an atomic site. For the partially ordered superlattice structure of Scheme I, as illustrated in Figure 6a, the following three lattice sites (0,0,0), (0,5/6,1/2), and (1/2,1/3,1/2) are occupied solely by U atoms, and the (1/2,1/2,0) lattice site is randomly occupied by U and Nb atoms. For the partially ordered superlattice of Scheme II, as illustrated in Figure 6b, the (0,5/6,1/2) and (1/2,1/3,1/2) lattice sites are occupied solely by U atoms, the (0,0,0) and (1/2,1/2,0) lattice sites are randomly occupied by U and Nb atoms, and the probability of site occupancy by U and Nb are 72% and 28%, respectively. The systematic variations in the atomic positions result in different ordered domains separated by APBs across which the atoms have the wrong immediate neighbors. Note that Scheme II ordering transition can also be regarded as a spinodal decomposition in which the Nb content fluctuates between 0 and 28 at. % (i.e., c₀ ± δc, and δc = 14 at.%), and the wavelength (λ) of composition fluctuation is equal to the lattice parameter of the ordered α'' unit cell in the [001] direction, i.e. λ = 0.495 nm, which is an order of magnitude smaller than the wavelength of spinodal decomposition observed in new WQ alloy aged at 200 °C. This type of ordering transition without a change of composition resulting from the composition fluctuation with the wavelength equal to that of the interatomic spacing is termed as continuous ordering [15]. Like the spinodal decomposition, the continuous ordering is solely a diffusion reaction involving atoms exchanging lattice sites, and the parent and product phase share a common crystal lattice.

Selected-area diffraction (SAD) patterns of the [110]-, the [310]-, the [312]-, and the [100]-zone observed from a 15-year-old WQ alloy sample are shown in Figure 7, in which the simulated diffraction patterns of these three zones are also displayed. While the simulated diffraction patterns of the [110]-, the [310]-, and the [312]-zone (see Figure 7a) show no difference between Scheme I and Scheme II superlattices, the simulated diffraction patterns of the [001]-zone are different between Scheme I and Scheme II superlattices (see Figure 7b). The structure of ordered α'' martensitic phase is thus identified to be of Scheme II by matching the observed and simulated [100]-zone diffraction patterns. An image-contrast analysis was conducted to verify the APBs formed in the 15-year-old alloy, and the result is shown in Figure 8. Here, the APBs are visible when 001 and 021 superlattice reflections were used for imaging but become invisible when 020 fundamental reflection was used for imaging. The observation and contrast analysis of swirl-shaped antiphase domain boundaries (APBs) are demonstrated in Figure 8. The contrast of antiphase domain boundary (also known as π boundary) is visible when the phase angle α = 2πg•P = π and is invisible when α = 2π (where g is the reflection vector for imaging, and P = 1/2 <011> is the displacement vector of APB). The bright-field (BF) image shown in Figure 8 clearly displays the swirl-shape feature of APBs. The APBs are visible in the dark-field images (DF-a and DF-b) obtained using the 021 and the 001 superlattice reflections, respectively. However, the APBs become invisible in the dark-field image (DF-c) obtained using the 020 fundamental reflection. The formation of APBs reveals the occurrence of an ordering transformation, which leads to the formation of chemically ordered domains within the 15-year-old alloy. The ordered phase domain has the same crystal structure as that of the disordered phase domain, except it takes up a superlattice arrangement because of the periodic occupation of lattice sites by specific atom species.

Accelerated aging of 15-year-old WQ-U6Nb at 200 °C and 212 °C

To accelerate the ongoing chemical ordering of the 15-year-old WQ alloy, thermal aging experiments were carried out at 200 °C and 212 °C. The changes of microhardness with different aging durations are shown in Figure 1 to compare with the microhardness changes of the as-water-quenched alloy thermally aged at 200 °C. Notice that the initial microhardness of the 15-year-old alloy sample had a value of 189.4, which is about the same value as that of the as-water-quenched alloy. This suggests that the 15-year natural aging of WQ-U6Nb at ambient temperatures hadn’t caused any noticeable change in mechanical property. However, age hardening occurred, and the microhardness continuously increased to 292.5 whence the 15-year-old WQ alloy was acceleratedly aged at 200 °C for 240 hours. It took only 96 hours to reach 292.5 whence the 15-year-old alloy acceleratedly aged at 212 °C because of increasing age-hardening rate. It is worth noting that the microhardness does not de-
crease even after thermal aging at 200 °C for 240 hours, which is significantly different from thermal aging of the as-water-quenched alloy: The microhardness starts to decrease after thermal aging at 200 °C for longer than 8 hours.

Microstructure evolution and microhardness variation of the 15-year-old WQ alloy thermally aged at 200 °C up to 240 hours are shown in Figure 6. Notice that the swirl-shape APBs formed in 15-year-old WQ alloy (Figure 6a) no longer exist after thermal aging at 200 °C; age hardening caused by the increasing volume fraction of plate-like precipitates can be clearly seen in 24h (Figure 6b) and 96 h (Figure 6c) alloy samples. The occurrence of precipitation hardening in 15-year-old WQ alloy at 200 and 212 °C suggests that the sluggish on-going ordering transition at ambient temperatures kinetically accelerates whence thermal aging at 200 and 212 °C. The crystal structure of the ordered precipitates likely remains monotonic as that of the parent α” martensitic phase except that the unit cell takes up a superlattice arrangement. TEM phase analysis was accordingly conducted to identify the crystal structure of ordered precipitates. Typical selected-area electron diffraction (SAD) patterns of the [010]-, the [001]-, and the [110]-zone are presented in Figures 9a, Figures 9b and Figures 9c. The SAD patterns match very well with the simulated diffraction patterns generated from a Nb-depleted α (U) phase and the simulated diffraction patterns generated based on an ordered U3Nb (25 at. % Nb) phase in which three U atoms occupy the (0,0,0), (0,5/6,1/2), and (1/2,1/3,1/2) lattice sites; one Nb atom occupies the (1/2,1/2,0) lattice site. The U3Nb phase is likely a metastable phase since it does not exist on the equilibrium U-Nb phase diagram. A further study is required to understand its phase stability.

Figure 8: Bright-field (BF) and dark-field (DF) TEM images observed from the same area of the 15-year-old alloy sample. The swirl-shape APBs are visible in the DF images obtained using: a,b) The 0 2 1 and the 00 2 superlattice reflections but become invisible in the DF image obtained using; c) The 0 2 0 fundamental reflection.
Figure 9: TEM analyses of Nb-depleted α (U) phase and U₃Nb phase formed after accelerated aging of 15-year-old alloy at 200 °C for 96 hours. SAD patterns and simulated diffraction patterns are generated from: a) The [010]-zone; b) The [001]-zone; and c) The [110]-zone.
Stress-strain behavior of acceleratedly aged 15-year-old WQ alloy

Tensile stress-strain curves obtained under a quasi-static loading condition for the 15-year-old WQ alloy and the same alloy acceleratedly aged at 200 °C for 2 hours (2 h) and 96 hours (96 h) are shown in Figure 10. The stress-strain behavior clearly reveals the occurrence of age strengthening in the 15-year-old WQ alloy acceleratedly aged at 200 °C, which is consistent with the occurrence of age hardening in the acceleratedly aged 15-year-old WQ alloy measured by microhardness method (Figure 1). The ultimate strength continuously increases from ~121 ksi at 200 °C/0 h, to ~124 ksi at 200 °C/2 h, and to ~135 ksi at 200 °C/96 h; whereas the ductility continuously decreases from ~0.3 at 200 °C/0 h, to ~0.28 at 200 °C/2 h, and to ~0.15 at 200 °C/96 h because of the age strengthening.

Discussion

While low-temperature age hardening was observed from both as-water-quenched and 15-year-old WQ alloys (see Figure 1), a much longer aging time was required to reach hardness peak for the 15-year-old alloy. This discrepancy can be rationalized by different aging mechanisms occurred in these two alloys. The age hardening of as-water-quenched alloy is caused by spinodal decomposition: A modulated structure with a wavelength of 3 nm is developed along the [001]α″ direction. The age hardening of 15-year-old WQ alloy is caused by chemical ordering and precipitation. The occurrence of chemical ordering in the 15-year-old WQ alloy results in the formation of swirl-shaped antiphase domain boundaries (APBs) in the ordered α″ martensitic phase. Through the crystallographic simulation framework demonstrated in Figure 7 and Figure 8, it is suggested that the ordered α″ martensite can be generated from a spinodal modulation along the [001]α″ direction with the modulation wavelength (λ = 0.5 nm) which is equal to the distance between two (001) planes, as illustrated in Figure 11. Note that the modulation wavelength at ambient temperatures is approximately an order of magnitude smaller than the wavelength of spinodal modulation occurred at 200 °C. This temperature-dependence of spinodal modulation, i.e. the modulation wavelength decreases with decreasing temperature, agrees with the phenomenological theory of spinodal decomposition [12]. The formation of such an atomic scale modulation can be attributed to low thermal energy at ambient temperatures, which makes atoms difficult to migrate over a long distance. Instead, the atoms exchange their positions more effectively with their neighboring atoms. When the same species of atoms move to the same atomic planes, an atomic-scale modulation forms and the supersaturated α″ phase becomes chemically ordered. Such modulation and chemical ordering help to reduce the free energy of the supersaturated solid solution. At elevated temperatures (200 °C), higher thermal energy enables atoms to diffuse over relatively longer distances, which leads to the formation of modulated structures with a longer wavelength (λ = 3 nm). Compared to the atomic modulation formed at ambient temperatures, the modulated structure with longer wavelength has fewer interfaces. Thus, the longer wavelength modulation is an energy-favored modulation at elevated temperatures.

Two types of continuous transformations were observed to occur in WQ-U6Nb alloy at two different temperatures, i.e., continuous ordering (λ = 0.5 nm) at ambient temperature and spinodal decomposition (λ = 3 nm) at 200 °C. These low-temperature transformations reveal that the supersaturated solid solution in WQ-U6Nb alloy is thermodynamically unstable and is prone to cause composition fluctuations propagating along the [001]α″ direction, and the wavelength of the composition waves is dependent on the aging temperatures. For thermal aging of WQ-U6Nb alloy at 200 °C, the transformation rate is fast, and the transformation sequence is: Disordered solid solution α″ → spinodal decomposition → α′(Nb-lean) + α′(Nb-enriched). For ambient-temperature aging of WQ-U6Nb alloy, the transformation rate is sluggish, and the transformation sequence is disordered α″ → continuous ordering → ordered α′. For accelerated aging of ordered WQ-U6Nb alloy at 200 °C and 212 °C, the transformation sequence is ordered α″ → phase decomposition → Nb-depleted α(U) + Nb-enriched ordered phase. It can be readily seen from Figure 10 that the age strengthening associated with the strength increase and the ductility reduction occurred presumably due to the precipitation of ordered U₃Nb phase, and the ductility of the 15-year-old WQ alloy reduced by 50% from ~0.3 to ~0.15 after acceleratedly aged at 200 °C for 96 hours.

Conclusions

Low-temperature aging mechanisms of WQ-
U6Nb have been investigated using both as-water-quenched and 15-year-old WQ alloy samples. Microstructure and microhardness changes during thermal aging of the as-water-quenched WQ alloy at 200 °C and thermal aging of the 15-year-old WQ alloy at 200 °C and 212 °C have been studied. Transformation sequences were proposed to rationalize low-temperature aging mechanisms of WQ-U6Nb alloy. The age hardening behavior observed from thermally aged as-water-quenched alloy can be attributed to the occurrence of spinodal decomposition: α″ → α₁ (Nb-lean) + α₂ (Nb-enriched). A composition wave propagating along the [001]_α direction with the wavelength λ = 3 nm took place during

Figure 10: Tensile stress-strain curves obtained under a quasi-static loading condition for the 15-year-old WQ alloy acceleratedly aged at 200 °C for 0 hour (0 h), 2 hours (2 h), and 96 hours (96 h) respectively.

Figure 11: Schematic illustrations show the temperature dependence of the wavelength of spinodal modulation, in which the dark spheres represent the Nb-enriched atomic positions.
the spinodal decomposition. A continuous ordering reaction: Disordered solid solution \( \alpha'' \) \( \rightarrow \) ordered \( \alpha'' \) took place during the ambient-temperature aging of WQ alloy, which can be attributed to an atomic-scale composition wave propagating along the [001] \( \alpha'' \) direction with the wavelength \( \lambda \approx 0.5 \) nm. During accelerated aging of the 15-year-old WQ alloy at 200 °C and 212 °C, the ordered \( \alpha'' \) martensite further decomposed into a Nb-depleted \( \alpha \) phase and a Nb-enriched ordered phase which led to the increase of microhardness over an aging duration of 240 hours at 200 °C and 96 hours at 212 °C. The tensile strength increased from \( \sim 121 \) ksi to \( \sim 135 \) ksi, whereas the tensile ductility reduced from \( \sim 0.3 \) to \( \sim 0.15 \) after accelerated aging of the 15-year-old WQ alloy at 200 °C for 96 hours because of the precipitation of a Nb-enriched ordered phase.

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