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Characterizing Nanoscale Precipitation in a Titanium Alloy by Laser-Assisted Atom Probe Tomography

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Abstract

Atom-probe tomography was performed on the metastable β -Ti alloy, Ti-5Al-5Mo-5V-3Cr wt.% (Ti-5553), aged at 300 °C for 0 to 8 h, to precipitate the embrittling isothermal ω phase. A dependency of precipitate quantification laser pulse energy is found by comparing measurements. Ultraviolet laser pulse energies above ~ 30 pJ result in significant complex molecular ion formation during field-evaporation, causing mass spectral peak overlaps that inherently complicate data analyses. Observations and accurate quantification of the ω phase under such conditions are difficult. The effect is minimized or eliminated by using smaller laser pulse energies. With a small laser pulse energy, Ti-rich and solute depleted precipitates of the isothermal ω phase with an oxygen enriched interface are observed as early as after 1 h aging time utilizing the LEAP 5000X S (77% detection efficiency). We note that these precipitates were not detected below a 2 h ageing time with the LEAP 4000X Si (58% detection efficiency). The results correct the archival literature and the discrepancies are rationalized. The Al concentration in the matrix/precipitate interfacial region increases during aging. Nucleation of the α -phase at longer aging times may be facilitated by the O and Al enrichment at the matrix/precipitate interface (both strong α -stabilisers). The kinetics and compositional trajectory of the ω -phase with aging time are quantified. facilitating direct correlation of the APT data to previously published mechanical testing.

Keywords

atom probe tomography, titanium alloys, ω -phase, precipitation, aging **Introduction**

Near α -Ti alloys, α plus β -Ti alloys and metastable β -Ti alloys are susceptible to thermally induced phase-transformations, which dramatically alter their mechanical properties. The characterisation and quantification of these phases can be inherently difficult, particularly when the precipitates are small (<10 nm) relative to the thickness of a TEM foil or when the compositional contrast between the precipitate and matrix is small, making detection by atom-probe tomography (APT) challenging. Such limitations with characterization methods have hindered our understanding of α_2 formation in near- α and α plus β -Ti alloys, which is detrimental to fracture toughness and cold dwell fatigue resistance of components in gas-turbine engines [1-6]. It has also hindered our understanding of athermal ω (ω_a), which forms on guenching, and isothermal ω (ω_i), which forms on subsequent lowtemperature aging of metastable β -Ti alloys [7-12]. The ω_a phase forms by a lattice shift to a different crystal structure without diffusion. The ω_i phase forms by subsequent diffusion for low-temperature aging at and below ~350 °C and therefore it possesses a different composition from that of the β -matrix. Typically, the ω -phase is undesirable as it embrittles the alloy [10]; it is, however, generally believed to be a precursor to the formation of stable nanoscale α -precipitates that highly strengthen the alloy [8, 9, 13, 14]. This precipitation sequence is not fully understood, and a vast amount of APT research has been published recently to help further our insight into the precipitation mechanisms of the high-strengthto-weight β-Ti alloys: Ti-5Al-5Mo-5V-3Cr wt.% (Ti-5553); and Ti-6Cr-5Mo-5V-4Al (wt.%) (Ti-6554) [9, 11-17].

Figure 1(a) illustrates the deleterious effects on mechanical properties associated with ω_i precipitation in 24 h aged Ti-24-Nb-4Zr-8Sn wt.% (Ti-2448) [10]. This allow became increasingly brittle and stiff with higher aging temperatures up to 250 °C. It is striking that an age of just 80 °C/24 h has such a profound effect on mechanical properties. Identifying such early stage precipitation of ω_i from ω_a by TEM is inherently complicated. The associated diffraction spots are weak and diffuse and diffraction spot brightness is dependent on foil-thickness. Such weak diffraction also complicates dark-field imaging. In this study of Ti-2448 the embrittlement following the 80 °C/24 h treatment could only be associated with ω_i formation from the trend in the mechanical data and observing ω_i formation in the 250 °C/24 h aged sample. Figure 1(b) shows an increase in Vickers hardness of Ti-5553 after a 300 °C/1 h aging treatment, associated with ω_i formation [10]. Previously, these precipitates were not identified in this heat-treatment by APT analyses utilising a LEAP 4000X Si [11] but were observed by TEM and small-angle neutronscattering (SANS) [10]. It is clear from this introduction that advancement in materials characterization techniques is required to characterize the earliest stages of nanoprecipitation, which can alter dramatically the mechanical properties.

The ongoing development of commercially available local-electrode atom-probe (LEAP) tomographs has recently permitted unparalleled compositional analyses of nanoscale precipitates in Ti-alloys. Characterisation of the ω -phase in Ti-5553 and the similar Ti-6554 alloy has been attempted with voltage pulsing [13, 18] and laser pulsing [9, 11,14,15,17]. Because ω -phase containing Ti-alloys are brittle, laser pulsing can assist in achieving large datasets per APT experiment prior to a specimen undergoing fracture. The

laser pulse energy can, however, induce undesirable thermal effects, such as surface diffusion or formation of complex molecular ions that make data analyses more difficult or obscure second-phase precipitates. There are clear discrepancies in the results obtained regarding the ω -phase in β -Ti alloys, specifically focusing on Ti-5553 and Ti-6554. The first APT measurements of Ti-5553 with a 2 h aging treatment at 350 °C identified the ω -precipitates as a phase with very similar composition to that of the β - matrix [18]. More recently, the same authors demonstrated that slow heating at 5 °C /min to 350 °C (a 70 min aging treatment) resulted in ω -phase precipitates, which are depleted in all solute additions [13]. This is in agreement with our current observations. Finally, it has also recently been reported that the ω -phase in Ti-6554 aged at 300 °C for 0 to 8 h is a Modepleted precipitate, with all the other solute additions equal to their bulk concentrations [14,17]. These discrepancies are of significance as the results are used to rationalize the precipitation sequence of $\beta \rightarrow \beta + \omega_a \rightarrow \beta + \omega_i \rightarrow \beta + \alpha$. Thus, a study of the characterization of the ω -phase in β -Ti alloys by APT is timely and warranted.

This article demonstrates potential characterization errors, which can arise when the laser pulse energy implemented is too high. It also notes a case where early stage precipitation is detected and characterized by the LEAP 5000X S and *not* the LEAP 4000X Si, which suggests that there may also be an instrumental dependence. Our results do not identify athermal ω -formation at Mo depleted zones in Ti-5553, in agreement with SANS measurements of the same alloy [9] and contrary to those identified by APT in Ti-6554 [14,17]. The titanium-rich and solute-depleted isothermal ω -phase is identified as early as after 1 h aging at 300 °C, in agreement with [13], and contrary to [14]. The discrepancies are rationalised by a detailed consideration of the measurement and analysis methodologies. Finally, the evolution of the isothermal ω -phase between 0 and 8 h aged at 300 °C, termed embryonic ω -phase elsewhere [14,17]), is quantified herein.

Materials and Methods

Materials and Methods: Sample Preparation

The forged billet of Ti-5553 used in this research has previously been studied and described [9,19]. All samples were prepared by initially heat-treating Ti-5553 at 900 °C for 30 min, followed by water quenching. Four samples were subsequently aged at 300 °C for 1, 2, 4, and 8 h and finally air-cooled. These aging conditions complement previously published studies of Ti-5553 by small-angle neutron-scattering (SANS), transmission electron microscopy (TEM), micro-hardness measurements and APT [9,11]. All heat-treatments were performed with the samples encapsulated within an argon atmosphere.

Following heat-treatment, APT samples of the quenched condition and aged conditions were prepared using a standard lift-out, nanotip mounting, and nanotip sharpening method [20] employing an FEI Helios NanoLab 600 Dual-Beam focused ionbeam (FIB) microscope equipped with an Omniprobe Autoprobe 200 micromanipulator. Additional aspects of the FIB lift-out procedure can be found elsewhere, for example [21].

Materials and Methods: APT Data Acquisition & Test Matrix

All APT experiments were performed with laser pulsing on Cameca's LEAP instruments, representing four different models and configurations at different research facilities. The full test matrix is displayed in Table 1, which was designed to examine the dependence of precipitate identification and quantification with laser pulse energy. The combination of the APT measurements provide accurate quantitative evaluation of the ω -phase evolution during isothermal aging at 300 °C for aging times ranging from 0 to 8 h.

A 300 °C/8 h aged APT specimen was measured with an initial 30 pJ laser pulse energy. The laser pulse energy was then decreased to collect data at 10 pJ. Similarly, a 300 °C/ 2 h APT specimen was measured with an initial 30 pJ laser pulse energy, then increased for measurements at 40 pJ and finally at 50 pJ, Table 1. These experiments were performed with the a LEAP 4000X Si (ultraviolet picosecond laser wavelength = 355 nm) at Northwestern University's Northwestern University Center for Atom-Probe Tomography (NUCAPT) core facility, Evanston, IL 60028, USA, and designed to identify any systematic dependency of precipitate quantification with laser pulse energy.

The 300 °C/8 h aged APT specimens were measured on the LEAP 3000X HR at Oxford University, the LEAP 4000X HR at Harvard University, and the LEAP 4000X Si at Northwestern University. Quenched samples and 300 °C /1 h aged samples were measured on the LEAP 4000X Si at Northwestern University and LEAP 5000X S at CAMECA instruments, Table 1. The detection efficiency of the reflectron instruments (LEAP 3000X HR and LEAP 4000X HR) is ~37 %, while the LEAP 4000X Si and LEAP 5000 XS straight flight-path instruments have ~58 and ~77 % detection efficiency, respectively [22,23].

Materials and Methods: APT Data Reconstruction & Analysis

After data collection, runs were reconstructed and examined in three-dimensions (3-D) utilising CAMECA's IVAS data analysis software program, version 3.6.12. The massto-charge-state ratio (m/z) binning size of all the mass spectra was 0.01. The mass spectra were analysed carefully to assign correctly the isotopic elemental or molecular identity and to detect potential mass spectral peak overlaps, specifically between (TiH)⁺ and Mo²⁺ ions, in addition to the known direct mass spectral peak overlaps of isotopes of different elements, such as ⁵⁰Ti⁺ and ⁵⁰Cr⁺, or ¹⁰⁰Mo²⁺ and ⁵⁰Cr⁺. The criterion for ranging each mass spectral peak was defined at 10% of the peak intensity. The presence of Ti-rich ω precipitates was identified with a combination of statistical methods and real-space visualization. Firstly, Ti concentration frequency distribution functions were calculated to verify whether statistically significant deviations from a binomial distribution were present on a scale relevant to nanometer-sized precipitate [24]. The binomial distribution represents a uniform solute distribution of an alloy with the same mean composition. Block sizes varying between N=100 and N=250 atoms were calculated using the IVAS program, corresponding to sampling volumes between approximately 3.5 and 8.8 nm³. Secondly, Ti isoconcentration surfaces were used to identify and visualize the statistically significant Tirich ω -regions. Thirdly, proximity histogram concentration profiles (proxigrams) [25-30] were then created relative to the Ti isoconcentration-surfaces for quantitative compositional analysis of the ω -precipitates. Phase compositions were measured from the plateau regions in the proxigrams. In an iterative approach, the average Ti-concentration between the Ti-rich ω-phase and Ti-depleted beta-phase was then used for establishing a

refined isoconcentration surface threshold and proxigram concentration profile analysis. Table 1 presents the total ion counts acquired in each APT measurement run for the different aging times as well as a designation of whether Ti-rich precipitates were identified for each experimental run or not. Table 2 presents the total ion counts used in each reconstruction.

Results

Results: Mass Spectra Analysis

The APT mass spectra obtained with different laser pulse energies of 10 and 30 pJ with the same nanotip are displayed in Figure 2. Figure 2(b) – (e) are enlarged sections of these mass spectra shown in Figure 2(a), to highlight individual peaks of isotope groups and their relative intensities for the elemental or molecular ions in question. Specifically, we are focusing on the mass spectral overlap of several Mo²⁺ isotopes and (TiH)⁺ peaks in Figure 2(c), and of $(Mo_2)^{3+}$ and $(TiO)^+$ peaks in (d), and Mo⁺ and $(Mo_2)^{2+}$ in 2(e). From Figure 3(c), it is clear that the two mass spectra differ in their relative peak intensities. The higher laser pulse energy produced the formation of complex molecular (TiH)⁺ ions, which overlap with the Mo²⁺ isotopic peaks at m/z = 47, 48, 49, and 50 Da. Additionally, the higher laser pulse energy promotes field-evaporation of singly-charged Cr and possibly V ions, creating additional overlaps of (TiH)⁺ with ⁵⁰Cr⁺ at m/z = 50 Da and ⁵¹V⁺ at m/z = 51 Da. As peaks are expected for Mo, V, and Cr in this region, the subtle changes of the relative peak intensities associated with (TiH)⁺ can go undetected, and errors with concentration quantification can arise from these peak overlaps. These (TiH)⁺ peaks were not immediately apparent and were only noticed on overlaying the 30 and 10 pJ mass spectra. Once overlaid, the relative change in peak intensities with laser pulse energy is then clear, which is highlighted in Figure 3(c).

(TiO)⁺ is also formed, overlapping $(Mo_2)^{3+}$ at m/z = 62, 63, 64, 65, and 66 Da, Figure 2(d). Complex molecular-ion formation is particularly apparent in Figure 2(e), where the 30 pJ laser pulse energy results in creation of $(Mo_2)^{2+}$. The $(Mo_2)^{2+}$ peaks overlap with Mo⁺ peaks between m/z = 92 - 100 Da. The formation of $(Mo_2)^{2+}$ ions in laser-pulsed APT at higher laser pulse energies was previously noted for pure Mo [31,32] and for Ni-based superalloys [33]. The higher laser pulse energies produce higher temperatures of the nanotip's surface at the time of field-evaporation, and these references provide detailed analyses of the relative intensities of peaks for Mo⁺ and $(Mo_2)^{2+}$.

Employing lower laser pulse energies reduces the probability of forming complex molecular ions. At higher laser pulse energies, the overlapping peaks make the processing of the mass spectra and quantification of compositions inherently more difficult for this alloy. The Mo²⁺ peaks at m/z = 46, 47.5, and 48.5 Da are unaffected by the isobaric overlap with (TiH)⁺ at m/z = 47, 48, 49 and 50 Da. Thus, quantitative concentrations of Mo can be determined by using the Mo²⁺ peaks at m/z = 46, 47.5, and 48.5 Da and calculating the correct number of Mo atoms using the tabulated natural abundances of the isotopes [34]. The correction process is, however, tedious. Conversely, the contribution of (TiH)⁺ can be estimated by calculating the intensity of the ⁹⁸Mo²⁺ peak from the natural isotopic abundances of Mo and subtracting from them the intensity of the peak at m/z = 49 Da, representing 73.72% of (TiH)⁺.

At lower laser pulse energies, the $(TiH)^+$ ions constitutes only a negligible fraction of the detected Ti atoms (< 1%). Their contribution may therefore be safely neglected while maintaining a relative error below 1%.

Results: Bulk Composition

The bulk compositions following mass spectra analysis are presented in Table 2. The compositions presented have not been corrected for the previously described overlapping peaks, which are particularly prevalent at higher laser pulse energies. Therefore, the apparent bulk concentration of Mo is seen to increase from 2.8 to 4.5 at.% when the laser pulse energy is increased from 30 to 50 pJ, highlighted in bold font in the table. The error is associated with an increase in complex molecular-ion formation at higher laser pulse energies, which has not been accounted for in the mass spectra analyses. The bulk concentrations deduced at lower laser pulse energies are in good agreement with those measured by inductively coupled plasma optical emission spectrometry (ICP-OES), creating confidence in the consistency of the mass spectra analyses performed.

The bulk concentrations may effectively monitor potential issues associated with detector pile-up problems when comparing instruments with different detector efficiencies. Titanium would predominantly be affected by detector pile-up as it is the majority element in this alloy (~80 at.%, Table 2) and has a dominant isotope ⁴⁸Ti (73.8 % abundance). No such dependency on detector efficiency was observed in the Ti bulk concentration, Table 2, and thus we conclude that the evaporation conditions were in a regime where detector pile-up effects were not of great influence.

Results: APT Reconstructions and Concentration Frequency Distribution Analysis

Figure 3(a) shows the effect of increasing the laser pulse energy from 30 to 50 pJ for the same nanotip with nearly constant nanotip geometry and size, in a 3-D APT reconstruction of a sample aged for 2 h at 300 °C. Superposed is an 86 at.% Ti isoconcentration surface (red) to highlight the Ti-enriched regions. The 30 and 40 pJ reconstructions look quite similar. There is, however, a clear effect at 50 pJ with visually fewer and smaller Ti-enriched regions observed to permit the determination of ω precipitate sizes, compositions and volume fractions. There also appears to be more Mo ions (blue) in the 50 pJ section of the reconstruction. This is also noted in the apparent bulk concentration results, Table 2.

The uncorrected measured Ti-concentration frequency distribution functions for the sample aged for 2 h at 300 °C are displayed in Figure 3(b) – (d), measured with (b) 30, (c) 40, and (d) 50 pJ laser pulse energies. The subtle changes in elemental concentrations with increased laser pulse energy are evident from the shift of the mean Ti concentration (vertical dashed lines). The experimentally measured distribution is shifted toward lower Ti concentration-values for the higher laser pulse energies, Figure 3(b) to 3(d). This is related to the formation of complex molecular ions at higher laser pulse energies, incorrectly being identified as Mo^{2+} ions in the mass spectra. The three concentration frequency distributions are significantly broader than their respective binomial functions, specifically with a lower maximum and with a significant tail on the Ti-rich side. These Tirich tails are indicative of the presence of Ti-rich regions associated with the isothermal ω - phase. The tails are most pronounced at lower laser pulse energies, Figure 3(b) versus 3(d), respectively. In practice, however, the laser pulse energy cannot be arbitrarily low as this results in a high dc field and a preferential loss of atoms that evaporate between laser pulses (so-called *dc-field evaporation*). Since the atoms evaporating between laser pulses increases the random noise in the mass spectrum, a lower limit for the laser pulse energy is established by ensuring that the background noise does not increase when decreasing the laser pulse energy.

Figure 4(a) shows the effect of decreasing the laser pulse energy from 30 to 10 pJ for an APT reconstruction of a sample aged for 8 h at 300 °C. The laser direction is highlighted, and an 88 at.% Ti isoconcentration surface is superposed (red) to highlight the Ti-enriched regions. This is higher than the 86 at.% Ti isoconcentration surface displayed in Figure 3. It illustrates the apparent Mo enrichment (blue)/precipitate depletion (red) on the laser side of the nanotip at the 30 pJ laser pule energy setting. This effect is clearly minimized or eliminated entirely with the lower laser pulse energy at 10 pJ. Figure 4(b) and (c) show 40 nm thick slices of the 3-D reconstruction, obtained with a 30 pJ laser pulse energy and 10 pJ laser pulse energy, respectively. Figure 4(a) – (c) are shown on the same length scale and employ the same isoconcentration surface value of 88 at.% Ti. The apparent compositional inhomogeneity throughout the specimen, induced by the higher laser pulse energy is particularly apparent in Figure 4(b) when compared to Figure 4(c). It is clear that the issues arising from complex molecular ion formation are inhomogeneous, with greater formation on the laser side of the nanotip than the shadow side.

Results: Quantifying the Initial Nucleation and Growth of the Isothermal ω Phase

It is known from SANS that ω_i precipitation occurs in under 1 h of aging at 300 °C in Ti-5553 [9]. However, characterization of initial ω_i nucleation by APT has proven elusive, and is now carefully revisited. Figures 5 (a) and (b) display 3-D APT reconstructions for the sample aged 1 h at 300 °C, collected on the LEAP4000X Si and 5000X S, respectively. Eightfive at.% Ti isoconcentration surfaces are superposed to highlight Ti-rich regions. The significance of any Ti-enriched region is analyzed in Figures 5 (c) and (d) utilizing a Ti concentration frequency distribution, for the data collected on the LEAP4000X Si and 5000 XS, respectively. From the significant tail in the distribution obtained with the LEAP 5000X S, indicated by an arrow in Figure 5(d); it is clear that Ti-rich precipitates are present. The frequency of Ti-rich bins exceeds the random binomial reference distribution. From this distribution it is apparent that the 85 at.% Ti isoconcentration threshold value is effective in identifying Ti-rich precipitates in the 3-D reconstructions displayed in Figures 5(a) and (b). There is no indication, however, of these precipitates from the Ti concentration frequency distribution obtained with the LEAP4000X Si, Figure 5(c). Thus, it is clear from Figure 5 that Ti-rich precipitates have been identified in a 300 °C/1 h aged sample measured by a LEAP 5000X Si but not in a sample measured utilizing a LEAP 4000X Si when using similar instrument parameters, Table 1. This is discussed further in the subsection titled *Discussion: Discrepancies in the Experimental Data*.

Compositional analysis of the Ti-rich precipitates is performed utilizing the proximity histogram concentration profile method. The proxigrams for samples aged 1 and 8 h at 300 °C are displayed in Figure 6, which were created from 85 at.% Ti

isoconcentration surface values. The precipitates are Ti-rich relative to the β -matrix, Figure 6(a). They are depleted in all solute additions (Al, V, Cr, Mo, and Fe), Figure 6(b). We note that the precipitate/matrix interfacial region is rich in oxygen (O), Figure 6(c), commencing just after the 1 h aging treatment. Due to the low O concentration in the alloy, a large dataset is required to identify the peak at the interface with good counting statistics. For this reason, the peak is clear in the 1 h aging treatment dataset measured on the LEAP5000X S (46 million atoms, Table 2), but somewhat obscured in the error range of the 8 h aging treatment dataset measured employing the LEAP4000X Si (8 million atoms, Table 2). An O rich ω/β heterophase-interface has been noted elsewhere [11]. The precipitates and the interfacial region become richer in Al between the 1 h and 8 h aging treatments, Figure 6(b).

Eighty-five at.% Ti isoconcentration surfaces were used to quantify volume fraction, precipitate number density, and volume equivalent mean-radius of the ω precipitates, Figure 7(a), (b), and (c), respectively. The volume fraction was calculated by the ratio of all atoms contained within all precipitates to the total of atoms in each APT 3-D reconstruction. The number density of precipitates was calculated accounting for detector efficiency and assuming the β body-centered cubic (bcc) unit cell volume determined at room temperature for Ti-5553 [19]. This is a reasonable assumption due to the low volume fraction of precipitates. The equivalent spherical volume of each precipitate was calculated from the atom counts, correcting for detector efficiency, and assuming a hexagonal ω unit cell volume of pure Ti [35]. The volume-equivalent radius of each precipitate was calculated and the dependence of the mean volume-equivalent radius with aging time is plotted in Figure 7(c). Guides to the eve have been added to each plot under the assumption that the 10 pJ laser pulse energy data sets are consistently the most accurate measurements. Approximate error bars of 5% error are indicated for these microstructural parameters as an estimate, Figure 7(a)-(c). It is assumed that the error is dominated by the counting statistics of the number of precipitates (n) encountered in a dataset, and is therefore given by $1/\sqrt{n}$.

Discussion

Discussion: Mass Spectra & Charge-State-Ratios

The presence of the $(Mo_2)^{2+}$ dimer ions between m/z = 92 - 100 Da in the mass spectra during laser pulsing at higher laser pulse energies is indicative of field-evaporation at a larger temperature increase relative to those at lower laser pulse energies, Figure 2(e). Thermally induced surface diffusion may also be promoted due to the higher temperatures of the nanotip's surface achieved with the higher laser pulse energies. Surface diffusion reduces the spatial resolution in the 3-D APT reconstruction through a smearing or blurring effect [36].

The creation of complex molecular ions during field-evaporation at higher laser pulse energies may go entirely unnoticed by the user if: (a) the peaks related to the molecular ions exclusively overlay those related to elemental peaks already assigned; and (b) specifically if the peak intensities of the molecular ions are small relative to the elemental peaks. In Ti alloys containing Mo, such as Ti-5553, peaks associated with (TiH)⁺ may not be immediately apparent as they overlie with strong Mo^{2+} and/or Cr^+ peaks, Figure 2(c). Furthermore, the contribution of both an elemental ion and a complex molecular ion to a peak at the same mass-to-charge ratio cannot always be readily deconvoluted from each other. In many cases, this deconvolution process requires significant additional analysis time and introduces additional errors through error propagation from the limited counting statistics.

In the case of Ti alloys with a Mo addition, minimization of complex molecular ion formation can be achieved during experimentation by decreasing the laser pulse energy. This decreases the surface temperature at the time of field-evaporation, thereby minimizing $(Mo_2)^{2+}$ formation. This should also limit the formation of complex molecular ions associated with TiH and TiO formation, which also depend on the partial pressures of H₂ and O₂ in the vacuum system. They are furthermore associated with the surface diffusivities of interstitial atoms [37]. Although the physical chemistry of gas-metal reactions in a high electric field is a complex topic and not the focus of this article, it is noted that a metal-hydride/oxide was formed with titanium at higher laser pulse energies. The affinity of these elements for titanium is a well-known problem in titanium production. The high price of titanium production is a result of the high reactivity of oxygen with this metal. Similarly, the strong tendency of titanium to absorb hydrogen from air moisture (hydrogen pickup) is a major processing concern as it results in the well-documented material failures caused by hydrogen embrittlement [38].

The results herein are presented in terms of the laser pulse energies of commercially available LEAP instruments manufactured by Cameca (Madison, WI USA). This is a user-defined parameter of these instruments and the discussion is relevant to the analysis of small precipitates and clusters in general. In practice, the effect of the laser pulse energy will vary, however, somewhat from sample-to-sample. In the present study of Ti-5553 we find that minimizing both formation of molecular ions and dc-field evaporation can be achieved simultaneously with laser pulse energies between approximately 10 and 20 pJ in the 5-8 kV range. This was performed while tracking the relative intensities of the peaks for different charge states (the so-called *charge-state ratios*) or molecular cluster ions, specifically minimizing or eliminating (Mo₂)²⁺ and (TiH)⁺ molecular cluster ions. Table 2 presents the intensity ratios of ${}^{46}Mo^{2+} / {}^{92}(Mo^+ + Mo_2^{2+})$ calculated from the total ions detected in each reconstruction analysis, to use a simple definition for practical use that can be monitored in real time during data acquisition (without isobaric overlap deconvolution). It is clear from the 3-D reconstruction of the 2 h aged sample displayed in Figure 3a that the change of this ratio from 2.4 to 5.2 (Table 2) between 30 - 50pJ laser pulse energies caused analysis difficulties. While it is possible to calculate an average nanotip temperature over the evaporation duration, Marguis and Gault [39] clearly illustrated discrepancies in the determined tip temperature depending on the species used in the calculation. Thus, accurate nanotip temperature remains an area of on-going research.

Discussion: APT Reconstructions

The analysis issues arising from complex ion formation at higher laser pulse energies are most significant on the side of the APT nanotip facing the incident laser beam, Figure 4(a) - (c). Figure 4(a) displays the full 3-D reconstruction at different laser pulse energies, while

Figure 4(b) and 4(c) show a 40 nm through-thickness slice at 30 pJ and 10 pJ, respectively. The 3-D reconstructions illustrate clearly that the issues associated with complex molecular ion formation are inhomogeneous. It is certainly more difficult to discern the Tirich ω -precipitates on the laser side of the nanotip, Figure 4. The effect of the higher laser pulse energy on the detected ω volume fraction is apparent visually with the applied Ti-rich isoconcentration surface values in Figure 3(a) and 4. There is a clear depletion of the precipitates in the regions with significant formation of complex molecular ions at the higher energies, Figures 3(a) and 4.

Discussion: Proximity Histogram Concentration Profiles (Proxigrams)

If the elemental ion peaks in the mass spectrum are not deconvoluted from the overlapping complex molecular ion peaks, which arise at high laser pulse energies then the resultant bulk composition is too rich in the Mo concentration (bold font, Table 2). Although not presented in this article, there are similar discrepancies observed in the proxigrams generated from the higher laser pulse energy data. Proxigrams were generated for the 30, 40, and 50 pJ laser pulse energy data using a Ti-rich isoconcentration surface value (86 at.%) for the 2 h aged sample. From these proxigrams, the apparent concentration of Mo in the matrix increases with laser pulse energy, with a corresponding decrease of the Ti concentration.

The proximity histogram concentration profiles for the 300 °C/1 h and 300 °C/8 h aging-treatments are displayed in Figure 6, which are created using the 85 at.% Ti isoconcentration surface values. The ω -phase is Ti-rich relative to the β -matrix, and depleted in all solute additions. The ω/β interface is rich in oxygen (O) after just a 1 h aging-treatment. The interfacial region becomes richer in Al between the 1 and 8 h aging-treatments. This is notable, as 0 and Al are strong α -stabilisers. Thus, this interfacial enrichment of Al and 0 may play a role in the subsequent formation of the stable α -phase by lowering the nucleation barrier. This hypothesis is strengthened by recent high-angle annular dark-field high-resolution scanning transmission electron microscopy (HAADF-HRSTEM) results displaying α -laths between ω -precipitates and the β -matrix, which are in contact with the ω -interface [13,14].

Discussion: Precipitate Nucleation and Growth

The spread in results in Figures 7 (a) – (c) with increasing laser pulse energy is associated with difficulties and statistical limitations arising from deconvoluting Ti containing complex molecular ions from the solute additions in overlapping peaks in the mass spectra. The Ti concentration determined by ICP-OES is 81.6 at.%, Table 2. For the 2 h aging-treatment, the apparent Ti bulk concentration decreased from 80.4 to 78.6 at.% as the laser pulse energy is increased from 30 to 50 pJ, Table 2. Therefore when an 85 at.% Ti isoconcentration surface value is utilized, it is clear that the higher laser pulse energy sample will yield a lower apparent volume fraction, as for the 2 h data of Figure 7a. The same logic applies to the precipitates's mean radius.

Although not displayed, an increase in the isoconcentration surface value from 85 to 87 at.% resulted in an order of magnitude decrease in the apparent precipitate number

density when measured at high laser energies, specifically the 50 pJ laser pulsed energy data used of the 2 h aged sample.

It is clear from the number density value that there is extensive nucleation of the isothermal ω -phase during the first hour of aging (Figure 7b), presumably nucleating from the athermal ω -phase that is initially present. Following the first hour the number density achieves a plateau, while the volume fraction continues to increase slightly, Figures 7(a) and (b). The increase in volume fraction is associated with the continuous growth of the mean precipitate radius, Figure 7(c). Since the volume fraction is increasing through an aging time of at least 8 h, the nucleation and growth and transition to growth and coarsening regimes are observed, but certainly a quasi-steady-state coarsening regime is not acheived. Given that the alloy is quenched, the matrix may still be supersaturated after the initial nucleation regime in the first hour of aging, leading to the simultaneous increase of the volume fraction and precipitates's mean radius.

Discussion: Comparison to Archival Literature

The first APT analysis of Ti-5553 alloys was reported in 2009 utilizing the voltage mode, with a pulse voltage that was 30% of the steady-state applied dc voltage [18]. In this research, the isothermal ω -phase was identified by rather small compositional fluctuations, and the measured Ti bulk composition appears to be smaller than anticipated. The same authors note elsewhere that in general large pulse fractions should be avoided when measuring Ti-5553 [40]. The lower dc-fields can lead to adsorption of residual gases in the vacuum system and result in field corrosion, and therefore the possibility of effects on the measured concentrations cannot be ruled out. Thus, although the artefacts induced by large pulse fractions may be different to those induced by large laser pulse energies, it is clear there were initial concerns regarding the experimental methodology when studying Ti-5553 by APT using voltage pulsing.

More recently, the isothermal ω -phase in Ti-5553 was measured at a lower pulsing voltage, 20 % of the steady-state applied dc voltage [16]. The authors illustrate employing proximity histogram concentration profiles that the ω -phase is enriched in Ti and depleted in all solute additions. This is in agreement with the results presented in this current article. Their samples had been prepared by isochronal heating with a 5 °C/ min rate to 350 °C, followed by rapid cooling.

The isothermal ω -precipitates of Ti-5553 observed herein are Ti-enriched and also depleted in all solute elements (Mo, Cr, Al, V and Fe) after aging for more than 1 h at 300 °C. The nucleation mechanism of athermal ω -precipitates that form on quenching from a solutionizing temperature in the beta-phase field [11], which is currently being debated. Although not shown, a concentration frequency distribution analysis was applied to the quenched (0 h aging) Ti-5553 alloy and it revealed no evidence that athermal ω forms with a compositional change. This is in agreement with the results of a SANS study of Ti-5553, which suggests that there is very little compositional contrast, if any, of athermal ω with the β -matrix in these alloys [11]. This is in contradiction to the APT results for a Ti-6554 alloy [14,15,17], which is very similar to the Ti-5553 alloy studied herein. In their work on Ti-6554, these authors discuss the formation of ω -embryos for aging times between 0 - 12 h at 300 °C. These embryos are depleted solely in Mo, and the concentration of all the other

solute additions (Cr, V, Al) within the precipitates is approximately equal to their bulk concntrations. Thus, the embryos must also be enriched in Ti. This is not observed in the current study of Ti-5553 with the same heat-treatments but employing lower laser pulse energies. The earlier APT analyses of Ti-6554 [14,17] were performed with laser pulse energies between 70 - 80 pJ, significantly higher than those illustrated to cause data analysis difficulties in our research (>30 - 50pJ for nanotips field-evaporating at 5-8 kV). A mass spectrum was not provided in the research performed on Ti-6554, and comments were not made on the occurrence of molecular ions or on charge-state ratios. The requirement of deconvoluting the overlapping peaks in the mass spectra, which arise from complex molecular ions at the higher laser pulse energies, is also not mentioned [14,15,17].

Our research demonstrates that after a 1 h aging treatment at 300 °C, there is clear partitioning of all solute additions between the matrix and isothermal ω -precipitates. It is highly unlikely that the small compositional difference between Ti-5553 and Ti-6554 (1 wt.% Cr and 1 wt.% Al) would change the partitioning this dramatically. At aging times longer than 12 h of Ti-6554, they observed that the isothermal ω -precipitates are depleted in all solute elements [14], similar to our results. It appears highly likely that the discrepancy between the results is a consequence of complex molecular ion formation not being taken into account when assigning peaks to elemental identities.

Discussion: Discrepancies in the Experimental Data

It is not fully understood as to why the LEAP5000X S identified the Ti rich ω -precipitates after a 1 h aging treatment at 300 °C, while the measurement performed on the LEAP4000X Si did not. The nanotips were produced from the same lift-out procedure (i.e., the APT specimens were produced from within 20 µm of each other within the same grain), and the isothermal ω -phase nucleates homogeneously [8,9]. The difference in charge state ratios of the 1 h aged samples presented in Table 2, although not dramatic, may be significant. They indicate that the sample measured using the LEAP4000X Si had an average higher effective nanotip temperature at the time of field-evaporation than that measured by the LEAP5000X S. A hypothesis is that the improved counting statistics provided by the higher detection efficiency of the LEAP 5000 XS *may* partially compensate for a loss of clear distinction when trying to delineate between phases with a small compositional amplitude or difference in composition.

Discussion: Correlation to Mechanical Properties

The increase in Ti-5553 hardness after a 300 °C/1 h aging treatment (Figure 1b) is now conclusively shown to be related to early stage nucleation of high number density (Figure 7b) Ti-rich/solute-depleted isothermal ω -precipitation (Figure 6) and not a variation of ω , which is solely enriched in Ti and depleted in Mo. With the LEAP5000X S APT and accounting for the sensitivity of materials characterization to instrument parameters, it is possible to correlate directly nanoprecipitates to mechanical properties at early nucleation times, rather than back-extrapolating the correlation from longer aged samples.

Summary and Conclusions

- It has been shown that higher laser pulse energies should be avoided when studying nanoscale precipitation in Ti-alloys. This is due to an associated high surface temperature at the time of field evaporation, which favors the formation of complex molecular ions. The associated mass spectral peak overlaps inherently complicates data analysis. It is critical to minimize these overlaps to enable accurate quantification or even discovery of nanoscale precipitates, even if they are highly concentrated like the ω -precipitate results presented in our study presented in this article. It also appears from the archival literature that high pulse-fractions in the voltage pulsing mode should also be avoided [13,40]. The discrepancies among APT measurements in the archival literature regarding the composition and solute partitioning in the isothermally formed ω -phase are addressed in detail. These discrepancies are rationalized based on an analysis of the complex molecular ions that form at higher laser pulse energies.
- We note that a measurement performed on a LEAP5000 XS detected early precipitation in Ti-5553 after a 1 h ageing treatment at 300 °C. These precipitates were not observed in a measurement performed on a LEAP 4000X Si in a separate APT specimen from the same grain.
- There is no indication from our APT analyses that athermal ω–precipitates form in Mo depleted zones in Ti-5553, in agreement with previously reported small-angle neutron-scattering (SANS) data [9].
- After a 300 °C/1 h aging-treatment Ti-rich/solute-depleted isothermal ω -precipitates are observed, with oxygen enrichment at the ω -precipitate/ β -matrix heterophase interface. After an 8 h aging-treatment, the heterophase interface is enriched with Al relative to the 1 h aging-treatment, in addition to the 0 enrichment. Both Al and 0 are α -phase stabilizers and it is suggested that the isothermal ω/β heterophase interface serves as an α nucleation site due to a reduced nucleation barrier. This is a reasonable hypothesis, based on recently published high-angle annular dark-field-high resolution scanning transmission electron microscopy (HAADF-HRSTEM) results [13,14]. With the exception of the increase of the Al concentration at the interfacial region, there is little change in element partitioning between 1 8 h of aging.
- The isothermal ω -phase is in a nucleation regime for the first hour of aging at 300 °C, and nucleation and growth regimes that last for at least 8 h. There is a rapid increase in number density in the first hour, followed by a plateau. The isothermal ω -phase volume fraction and mean precipitate radius continue to increase throughout the 8 h aging-treatment.

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LEAP instrument	Laser λ (nm)	lon detection efficiency (%)	Laser energy (pJ)	Stage temp. (K)	Aging time (h)	lon counts (M)	Precipitates identified
3000X HR	532*	38	200*	40	8	52	Υ
4000X HR	355	38	50	44	8	57	Υ
4000X Si	355	58	30–10	30	8	48	Υ
4000X Si	355	58	10	30	4	82	Υ
4000X Si	355	58	30–50	45	2	78	Υ
4000X Si	355	58	10	30	1	29	Ν
5000X S	355	77	10	30	1	56	Υ
4000X Si	355	58	10	30	0	53	Ν
5000 X S	355	77	10	30	0	14	Ν

Table and Figure Captions

Table 1. Test matrix of the four different types of LEAP instruments used in this study of isothermal ω -precipitation in Ti-5553 alloys, aged for 0 - 8 h at 300 °C. The HR tomographs have an energy-compensating reflectron lens that improves mass resolution mostly in voltage mode and reduces background noise to some degree, albeit with decreased

detection efficiency. The Si and S tomographs have a straight flight-path with higher detection efficiency. The green laser ($\lambda = 532$ nm) is identified with an asterix*, while all other tomographs are equipped with a UV laser ($\lambda = 355$ nm). The laser pulse energies and specimen stage temperatures used for each measurement are listed along with the specific tomograph, ion detection efficiency, total ion counts of each measurement run in millions (M) detected, and whether precipitates were identified by APT (Y) or not (N).

Instrument	Laser energy (pJ)	⁴⁶ Mo/ ⁹² (Mo ⁺ + Mo ₂ ²⁺)	Aging time (h)	lon counts (M)	Ti	V	Fe	AI	Мо	Cr	0
ICP-OES	-	-	-	-	81.6	4.6	0.3	8.2	2.4	2.6	0.4
3000X HR	200*	9.8	8	48	79.9	5.1	0.3	8.3	2.9	2.9	0.5
4000X HR	50	5	8	52	80.6	4.9	0.2	8.3	2.7	2.9	0.4
4000X Si	10	19.9	8	8	79.9	5.1	0.3	8.5	2.8	2.8	0.5
4000X Si	30	3.8	8	19	80.4	4.7	0.2	8.2	3.2	2.8	0.5
4000X Si	10	72.7	4	54	80.1	5.2	0.2	8.4	2.7	2.8	0.5
4000X Si	30	5.2	2	13	80.4	4.9	0.2	8.4	2.8	2.7	0.6
4000X Si	40	3.2	2	16	80	4.8	0.2	8.4	3.2	2.8	0.5
4000X Si	50	2.4	2	13	78.6	4.7	0.2	8.4	4.5	29	0.5
4000X Si	10	18.9	1	21	79.6	5.4	0.3	8.6	2.5	2.9	0.5
5000X S	10	39.9	1	46	79	5.5	0.4	8.6	2.9	3.1	0.5
4000X Si	10	39.6	0	52	80.3	5.2	0.3	8.3	2.5	2.9	0.5
5000X S	10	63.1	0	11	79.4	5.1	0.3	9.3	2.6	2.7	0.6

Table 2. Bulk compositions in atomic % of the Ti-5553 alloy, determined by inductively coupled plasma optical emission spectrometry (ICP-OES) [7] and by different atom probe tomography LEAP instruments, aged for 0 - 8 h at 300 °C. The green laser is identified with an asterix*, while all other tomographs are equipped with a UV laser. The laser pulse energies for each measurement are listed along with the total ion counts used in each data reconstruction in millions (M) detected. The intensity ratio during field evaporation of Mo to (Mo + Mo₂) at *m*/*z* 46 and 92, respectively, are shown. Measurements where the laser pulse energy was changed during the data acquisition, in order to investigate the dependence between these energies and precipitate characterization, are highlighted in the first two columns by italic font. The effect of high laser pulse energy on apparent Mo composition is highlighted in bold font. Further experimental parameters are provided in Table 1.



Figure 1. a) Effect of low temperature aging between 80 – 250 °C for 24 h on roomtemperature tensile properties of cold rolled metastable β -Ti alloy Ti-2448 (Ti-24Nb-4Zr-8Sn wt.%) [10]. b) Effect of low temperature aging on room-temperature hardness of Ti-5553 (Ti-5Al-5Mo-5V-3Cr wt.%). Samples were aged at 300 °C for up to 8 h, and a dual heat-treatment sample of 300 °C/8 h + 500 °C /2 h was also performed. Images reproduced with the permission of the publisher.



Mass-to-Charge-State Ratio m/z

Figure 2. APT mass spectra of Ti-5553 aged for 8 h at 300 °C obtained with 10 or 30 pJ laser pulse energies utilizing a LEAP 4000X Si atom-probe tomograph. The 10 and 30 pJ mass spectra are from 8 and 19 million ions, respectively, and the mass-charge-state ratio binning size was 0.01. (a) Displays the complete spectrum and (b) – (e) are magnified sections to highlight the isotopic distributions of individual ions or molecular ions. Specifically the isobaric overlap of several Mo²⁺ and (TiH)⁺ peaks in (c), and of (Mo2)³⁺ and (TiO)⁺ peaks in (d), and Mo⁺ and (Mo₂)²⁺ in (e) are highlighted.



Figure 3. (a) APT reconstruction of Ti-5553 aged for 2 h at 300 °C. Data obtained with the UV laser pulse energy stepped from 30 to 40 and 50 pJ after ~15 million ions collected, respectively, utilizing a LEAP 4000X Si atom-probe tomograph. To highlight the Ti-rich precipitates (ppts), an 86 at.% Ti isoconcentration surface is superposed in red. For clarity, only 15 % of the Mo ions are displayed (blue dots). A 10° shank angle was utilized to obtain a continuous reconstruction. (b) – (d) Ti concentration frequency distributions, red, for sections with the different laser energies, 30 pJ in (b), 40 pJ in (c), and 50 pJ in (d), respectively, for the same dataset, calculated with a block size of 250 atoms. A dashed line highlights the mean value. Binomial distributions, representing a random solid-solution with the same average composition, are shown as black lines for comparison with the

experimental data. The Ti-rich tails, representing Ti-rich precipitates, are most pronounced at lower laser pulse energies (arrows in (b) - (d)).



Figure 4. (a) APT reconstruction of Ti-5553 aged for 8 h at 300 °C. Data obtained with the UV laser pulse energy stepped from 30 pJ to 10 pJ after ~19 million ions collected, utilizing a LEAP 4000X Si tomograph. To highlight the Ti-rich precipitates, an 88 at.% Ti-rich isoconcentration surface is superposed in red. For clarity, only 35 % of the Mo ions are shown (blue dots). A 10° shank angle was utilized to obtain a continuous reconstruction. (b) - (c) View through 40 nm thickness of the APT reconstruction with Ti-rich precipitates again highlighted with an 88 at.% Ti-rich isoconcentration surface superposed in red and with 75 % Mo ions shown (blue dots), for 30 and 10 pJ laser pulse energies, respectively. All sub-figures are to the same scale, and highlight the incident laser direction.



Figure 5. APT reconstructions of Ti-5553 aged for 1 h at 300 °C obtained with (a) a LEAP 4000X Si atom-probe tomograph, and (b) a LEAP 5000 XS. An 85 at.% Ti isoconcentration surface is superposed (red). 10 % of the Mo ions detected (blue dots) are displayed to outline the shape of the reconstruction. (c) Corresponding Ti concentration frequency distributions (stepped red line) constructed with block sizes of 250 atoms for the LEAP 4000X Si data, and (d) the LEAP 5000 XS data, respectively. The binomial distributions, representing a random solid-solution with the same average composition, are displayed as black lines for comparison with the experimental data.



Figure 6. (a) – (c) Proximity histogram concentration profiles of Ti-rich precipitates with respect to an 85 at.% Ti isoconcentration surface, for Ti-5553 aged for 1 or 8 h at 300 °C. The approximate inflection points of the Ti profiles are highlighted with a dashed line, as an approximation for the center of the interfacial region between the matrix and precipitate. The larger error range in the oxygen concentration deduced in the 8 h data compared to the 1 h data is associated with the poorer counting statistics, where the 8h profile is formed from 8 million ions and the 1 h profile is formed from 46 million ions.



Figure 7. (a) Measured volume fraction, (b) number density and (c) mean volumeequivalent radius of Ti-rich precipitates versus aging time in Ti-5553 aged for 0 - 8 h at 300 °C. The microstructural parameters are calculated from an applied 85 at.% Ti isoconcentration surface value. Guides to the eye are included along with error bars of 5% error, approximated from the counting statistics of the number of precipitates encountered in the datasets