Effect of Ta addition on the electrochemical behavior and functional fatigue life of metastable Ti-Zr-Nb based alloy for indwelling implant applications

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In this paper, the effects of Ta substitution for Nb, on the microstructure and corrosion fatigue behavior of metastable Ti-18Nb-14Zr biomedical alloy are investigated for the first time. For this purpose, besides previously studied ternary Ti-18Zr-14Nb alloy, new quaternary Ti-18Zr-13Nb-1Ta (at.%) biomedical alloy was also synthesized by vacuum arc remelting method and subjected to thermomechanical treatment processes. The in situ electrochemical behavior and subsequent functional fatigue life of both alloys showed a strong dependence on the composition and microstructure. The electrochemical results from the test bench demonstrated that during monotonic cycling the fatigue life of Ta-added alloy was significantly longer than that of the Ti-18Zr-14Nb alloy. Thus, the addition of Ta to the ternary Ti-18Zr-14Nb alloy was found to be very effective in increasing the resistance to fatigue degradation mainly by exhibiting excellent protective passivation tendency. Based on the electrochemical evaluation and fractographic characteristics, it is concluded that the combined effect of the stable passive film formation and higher development of a nanosubgrained structure in β phase results in the prolonged fatigue life of Ta-added alloy.

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1. Introduction

Ti-Ni based alloys are one of the most distinctive families of superelastic alloys, which provide myriad advantages in the diverse engineering domains [1,2]. At present, the superior functional fatigue properties of superelastic Ti-Ni based alloys have been commercially exploited in the manufacturing of various indwelling implants such as endovascular stents, catheter guide wire, dental braces, etc. [1–3]. Unfortunately, the risks of the hypersensitivity associated with Ti-Ni-based alloys due to the release of Ni ions from corrosion reactions against the human body have also been significantly identified [2–6].

Over the years, in order to establish greater efficacy of the biomedical implants, several Ni-free biomedical superelastic alloys had been exquisitely elaborated as a replacement for the conventional Ti-Ni based alloys [4–16]. Among them, multicomponent metastable β-type Ti-Nb- [2–4,6,9,11–13,17–20] and Ti-Zr- [4,7,10] based alloys, with different β-stabilizing element substitutions are most prevalent in the biomedical field.

Typically, the superelastic effect in metastable β-Ti alloy is derived from the reversible stress-induced martensitic transformation during the mechanical loading-unloading cycling [3,5,9,15]. Some factors affecting cyclic fatigue properties, like nanoscale concentration modulation inducing confinement of the martensitic transformation, have been described recently in Refs. [20–22].

For this reason, the evaluation of functional fatigue life under monotonous mechanical cycling is of pivotal significance [3,5,15,16,23].

Recently, it has been reported that the addition of Ta as a substitute for Nb in Ti-18Nb-14Zr alloy is a promising way to further improve the functional fatigue properties of Ti-Zr-Nb based superelastic alloys [24]. Undoubtedly, the metastable β-type ternary Ti-Zr-Nb [4,10] and/or quaternary Ti-Zr-Nb-Ta base alloys, are beginning to emerge as the latest research frontiers in
biomedical applications \cite{4,10,24}.

Yet surprisingly, according to our knowledge, none of the studies have been meticulously conducted to characterize the in situ corrosion functional fatigue performance and fracture characteristics of Ti-Zr-Nb alloy.

In this line of thought, from a practical viewpoint, it would be very enticing to synergistically elucidate and compare the effects of alloying substitutions i.e., Ta on the electrochemical behavior and the functional fatigue life of newly developed Ti-18Zr-14Nb alloy in the adverse physiological environments \cite{5,25,26}.

For this purpose, we have recently built and validated a prototype electrochemical test bench which emulates the in vitro conditions to which the wire-shaped implants will be actually subjected in vivo \cite{27,28}.

2. Materials and methods

The Ti-18Zr-13Nb-1Ta and Ti-18Zr-14Nb at. % alloys ingots (all compositions are hereafter referred in atomic percent) were prepared by the vacuum arc-melting method with a consumable electrode. In order to ensure the chemical homogeneity of the alloys, each ingot was remelted several times and flipped before each melting step. Each ingot was sliced in to 1.5 mm thick plates using electric discharge machining. Next, the rectangular shaped plates were homogenized at 1173 K for 1.8 ks under Ar atmosphere, followed by rapid quenching in water. Afterward, all the plates were subjected to thermomechanical treatments (TMT). The TMT included the following: cold rolling with a true logarithmic strain of 0.3 and post-deformation annealing at 873 K for 1.8 ks followed by water quenching. The post-deformation annealing at 873 K (1.8 ks) creates a nanosubgrained structure in the beta-phase of the Ti-Zr-Nb alloy.

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3. Results and discussion

3.1. Microstructure of Ti-18Zr-14Nb and Ti-18Zr-13Nb-1Ta alloys

The optical microscopy study was carried out for the purpose of grain size comparison of both alloys before TMT. Typical grain structures are shown in Fig. 1. It follows from Fig. 1 that the alloys possess approximately the same grain size (100–500 μm) which points to the fact that Ta for Nb substitution of 1 at.% does not affect initial grain structure of the material.

In order to discuss the effect of TMT on the microstructure of Ti-18Zr-14Nb and Ti-18Zr-13Nb-1Ta alloys, the TEM study was performed. Fig. 2 shows the TEM bright and dark field images with the corresponding selected-area diffraction patterns (shown in insets) with a beam parallel to [113]β of Ti-18Zr-14Nb and Ti-18Zr-13Nb-1Ta alloys. The corresponding dark field images for both specimens (as shown in Fig. 2b and d) were obtained using the (110) type reflection of the β-phase which is the main phase in both alloys. The bright and dark field images shown in Fig. 2 reveal that after the thermomechanical treatment both specimens consist of a polygonized dislocation substructure, with high dislocation density inside subgrains (indicated by white arrows in Fig. 2b and d). Note that this type of substructure formation has also been observed in
Ti-Ni, Ti-Nb-Zr and Ti-Nb-Ta based alloys [5,9,24]. While comparing both the images, following comparative observations may be of interest. According to the obtained diffraction contrast images, Ti-18Zr-13Nb-1Ta alloy tends to demonstrate more developed subgrained structure after TMT with somewhat smaller subgrain size compared to Ti-18Zr-14Nb (Fig. 2). Apparently, the formation of smaller subgrains in Ti-18Zr-13Nb-1Ta alloy is ascribable to lower diffusion coefficient of Ta in the $\beta$ structure which, consequently, led to the slower subgrain growth [5]. Nonetheless, the subgrain size of Ti-18Zr-13Nb-1Ta and Ti-18Zr-14Nb alloys occupies rather wide ranges of 50–500 and 100–600 nm, respectively. For that reason, more in-depth statistical analysis of large amount of TEM images is needed in order to evaluate the influence of Ta for Nb substitution on the features of subgrains structure.

3.2. Electrochemical behavior of studied materials during functional fatigue cycling

Fig. 3 displays the open circuit potential (OCP) values (mV) as a function of exposure time (s) for the Ti-18Zr-14Nb and Ti-18Zr-13Nb-1Ta alloy specimens subjected to functional fatigue cycling in Hank’s solution. The summary of the experimental results is also illustrated in Table 1. Prior to functional fatigue cycling the respective time-invariant steady states ($E_{st}$) for both the alloys was established. The $E_{st}$ value of Ti-18Zr-14Nb alloy ($-118$ mV) is slightly lower than that of Ti-18Zr-13Nb-1Ta ($-76$ mV). Once, the respective $E_{st}$ for both the specimens was achieved the functional fatigue cycling with a constant bending strain of 1.5% was imposed. Firstly, it can be identified that after the initiation of cycling the OCP for both Ti-18Zr-14Nb and Ti-18Zr-13Nb-1Ta alloy specimens shifted towards the negative values. Nonetheless, the shifts in the OCP towards negative potentials are associated to the abrupt cathodic polarizations [25–32]. Indeed, the $E_c$ values for the Ti-18Zr-14Nb and Ti-18Zr-13Nb-1Ta alloy are about $-453$ and $-588$ mV respectively.

Although, at the initial stage of fatigue cycling, the shift in $\Delta E$ towards negative values is larger in case of Ti-18Zr-13Nb-1Ta alloy specimen when compared to Ti-18Zr-14Nb alloy specimen (as shown in Fig. 3(a and b)) respectively. However, as the cycling proceeded, the $E_c$ of Ti-18Zr-13Nb-1Ta alloy specimen exhibits
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increasing tendency with the time duration for about 400 s and then keeps stable and higher OCP values during the entire cycling (as shown in Fig. 3b).

It is because the protective passive films once forms begin to grow on the Ti-18Zr-13Nb-1Ta alloy specimen surface and then forms a stable and strong passive film during the cycling. The higher protective ability of the surface film on Ti-18Zr-13Nb-1Ta alloy can be attributed to Ta\(^{5+}\) ions favourable dissolution in TiO\(_2\) matrix film as compared to Zr\(^{4+}\) ions due to closer ionic radii (0.064, 0.066 and 0.082 nm for Ti\(^{4+}\), Ta\(^{5+}\) and Zr\(^{4+}\), respectively) resulting in lower point defect concentration and, consequently, the stability is increased.

Another reason, for this stable passive film formation, might be due to the “chemomechanical effect” which is elicited at the specimen/electrolyte interface [27–29].

In other words, during the functional fatigue cycling unlike to Ti-18Zr-14 Nb alloy, the surface of the Ti-18Zr-13Nb-1Ta alloy specimen is more defant to the impact of cycling.

Accordingly, it can be deduced that the substitution of Ta for Nb in Ti-18Zr-14Nb alloy has resulted in the formation of a stable passive film, which in return can successfully mitigate the corrosion fatigue damage.

On the other hand, at the dynamic fracture stage for both the specimens a slight but distinct negative shift typically corresponding to fracture was observed which was finally followed by a rapid recovery in the positive direction in the OCP plots owing to passivation of the freshly formed surface [27,28]. Thus, the negative shift in the OCP values is essentially due to the local rupture of the protective film and/or due to the dissolution of the film prevailing over its formation [25–27].

The dynamic fracture for the specimens during functional fatigue cycling was estimated to occur around 1500 s for Ti-18Zr-14Nb and 2300 s for Ti-18Zr-13Nb-1Ta alloy specimen.

Eventually, while comparing the corresponding average functional fatigue life of Ti-18Zr-14Nb and Ti-18Zr-13Nb-1Ta alloy specimens, which in our case is defined as the total number of successful loading-unloading cycles (\(N_{\text{max}}\)) which a specimen can withstand before dynamic fracture. The \(N_{\text{max}}\) for Ti-18Zr-14Nb and Ti-18Zr-13Nb-1Ta alloy specimens in the physiological solution is calculated to be 854 ± 328 and 1806 ± 559 cycles respectively. Note that the \(N_{\text{max}}\) values obtained in low-cycle fatigue tests of analogous Ti-Nb-Zr-Sn alloys were much higher (over 50000 cycles) [17–19] at comparable maximum strain in a cycle. The reason for this difference is the difference in the experimental conditions of the present and reference investigations, including differences in surface roughness, sample size and dimensions, straining modes, thermomechanical conditions of tests, thermomechanical treatment modes and corresponding microstructures of alloys.

In order to discuss the longer functional fatigue lifetime of Ti-18Zr-13Nb-1Ta alloy in the physiological solution, the apparent surface components of the fracture surfaces of Ti-18Zr-14Nb and Ti-18Zr-13Nb-1Ta alloy specimens were systematically studied.

### 3.3. Fractography of fractured specimen after fatigue test in physiological solution

Fig. 4 shows the general fractography of Ti-18Zr-14Nb and Ti-18Zr-13Nb-1Ta alloy specimens, which was characterized by SEM. The precise comparison of the macroscopic and microscopic regions for Ti-18Zr-14Nb and Ti-18Zr-13Nb-1Ta alloy specimens, particularly the crack origin (noted by white arrows), fatigue striations region and a final unstable region which is nearby the dynamic fractured area are also illustrated in Fig. 4a and b respectively. On the other hand, for the sake of clarity, the segment of fatigue striations by the crack propagation in the stable region and dimples at the unstable region for both Ti-18Zr-14Nb and Ti-18Zr-13Nb-1Ta alloy specimens alloy are also highlighted with the white delineated rectangular boxes.

From general aspects, it is evident that the dynamic fracture in
both of the specimen was elicited mainly by trans-granular pattern [23,28,32] as the corresponding three regions shown in Fig. 4 are easily identifiable. Particularly, it is also noteworthy that in the case of Ti-18Zr-14Nb the stable region showing the fatigue crack propagation is much smaller than the final overloaded region. In fact, the enlarged unstable zone of Ti-18Zr-14Nb alloy (Fig. 4a) on one hand corroborates the lower fatigue life of Ti-18Zr-14Nb alloy. It is presumably because the local passive films breakage during cycling loading has accelerated the corrosion-assisted fatigue crack growth which collectively has led in the increase in the length of the final unstable region when compared to the fatigue striations region (Fig. 4a).

Unexpectedly, the fractography of the Ti-18Zr-13Nb-1Ta alloy specimen (Fig. 4b) clearly indicates a much larger fatigue striation region when compared to Ti-18Zr-14Nb alloy specimen (Fig. 4a). During cycling, the fatigue striations streaks on the specimen surface are assumed to be generated by the successive opening and/or closing of the surface cracks. The lengths of fatigue striation region can substantially influence the functional fatigue lifetime of an implant [28].

Additionally, the peculiarities of the reversible martensitic transformation $\beta \leftrightarrow \alpha'$ underlying the superelasticity effect in these alloys may influence the features of the fracture surfaces [33].

Thus, by correlating the microstructural results with the OCP response it is suggested that the increase in the functional fatigue lifetime of Ti-18Zr-13Nb-1Ta alloy is firstly related to the to the stable passive film formation which inhibits the local corrosive attack (as shown in Fig. 3b).

Meanwhile, while complementing these findings, with the stabilized OCP response of Ti-18Zr-13Nb-1Ta alloy specimen (Fig. 4b), it is claimed that the relatively higher mechanical strength of Ti-18Zr-13Nb-1Ta alloy which was due to the formation of nanosubgrained structure (as shown in Fig. 2b) can also be an additional and/or contributing factor for the longer functional fatigue lifetime of Ti-18Zr-13Nb-1Ta alloy. It is because during functional fatigue cycling the greater difference between dislocation and transformation yield stresses has also resulted in the prolonged functional fatigue life of Ti-Nb-based alloys [5,9,22,24].

4. Conclusions

The effects of Ta content on the microstructure, electrochemical behavior and fatigue life of Ti-Nb-Zr alloy were investigated after thermomechanical treatment in the custom-made pseudo-physiological test bench. The experimental results are summarized as follows:

1. The substitution of Ta for Nb promotes more developed subgrain structure and changes the subgrain size of thermomechanically treated Ti-18Zr-14Nb alloy towards the nanometer scale.
2. The Ti-18Zr-13Nb-1Ta alloy exhibited higher open circuit potential values during cycling and its passive oxide film is found to be more stable and adequately resistant to the impact of functional fatigue when compared to Ti-18Zr-14Nb alloy.
3. Substitution of 1% Nb by Ta increases functional fatigue life almost twice: $N_{max} = 1806$ for Ti-18Zr-13Nb-1Ta in contrast with $N_{max} = 854$ for Ti-18Zr-14Nb alloy.
4. Based on the fractographic characteristics and electrochemical evaluation, it is concluded that beside the stable passive film formation on the specimen surface, the combined effect of the unique $\beta$ phase nanosubgrained structure also contributes to the prolonged fatigue life of Ti-18Zr-13Nb-1Ta alloy.

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