Hydrogen absorption and hydride formation in pure titanium T40 (grade 2) and TA6V ELI (grade 23) under cathodic polarization in artificial seawater

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Abstract

Different kinetics of hydrogen absorption in T40 (grade 2) and TA6V ELI (grade 23) under cathodic polarization in artificial seawater have been highlighted. These polarizations were made by applying potentials from -0.8 to -1.8V/SCE in artificial seawater and NaCl solution. Four stages were identified and related in term of hydrogen ingress, hydrides formation and calcareous deposit growth. The formation of γ and δ-hydrides have been observed, localized and characterized using several techniques. On T40, hydrides form as a layer that increases the surface roughness and clusters form in the bulk after first moments of hydrogen absorption. Whereas in TA6V ELI, hydrogen is absorbed by β-phase leading to a volume expansion of this phase. Then after reaching the hydrogen solubility limit of β-phase, hydrides form on interfaces α/β or α/γ and in γ grains. For long durations, the hydrogen ingress is limited by the subsurface hydrides and the stabilized calcareous deposit. These different steps are time dependent processes which need to be tackled into account to improve knowledge of hydrogen embrittlement in titanium alloys.

1. Introduction

Titanium alloys are chosen for the manufacturing of components used in a marine environment. These alloys efficiently protect themselves from any damage due to this aggressive environment by forming a stable passive film of titanium oxide TiO₂. When they are used in multi-materials systems under cathodic protection, these conditions could favour the hydrogen evolution reaction (HER), which may lead to the material embrittlement by hydrides formation and/or solute/plasticity interactions [1, 2]. Consequently, characterise the hydrogen state is a key feature to understand the hydrogen embrittlement sensitivity of titanium alloys. A commercially pure titanium single phase (α) T40 (grade 2) and a two-phase titanium alloy (α / β) TA6V ELI (grade 23) are studied in order to assess the impact of the nature of each phase on the sensitivity to hydrogen absorption in artificial seawater and on hydrogen and hydrides distribution.

2. Materials and experiments

Two materials were studied, the commercially pure grade 2 and the grade 23 titanium alloy. Grade 2 was received in the form of 3 mm-thick rolled sheet from TIMET and grade 23 in the form of 6 mm-thick rolled sheet from AUBERT & DUVAL. Grade 2, Figure 1, presents a single α-phase equiaxed structure with grains measuring 46 ± 10 μm and some intergranular iron inclusions (<3% weight). The texture analysis shows a preferential crystallographic orientation of the basal (0002) pole. The axes are perpendicular to the rolling direction (RD) with the basal pole rotated about 35° toward the transverse direction (TD). Grains show strong misorientations (>15°). Additionally, it has been observed by TEM only few dislocations in α-grains. Grade 23, Figure 1, presents a α/β bimodal structure with a volume fraction of β-phase about 6%. Primary α grains measure 22 ± 5 μm, secondary α grains measure 9 ± 5 μm and β grains measure approximately 1 μm. Macro-zones of 100 to 150 μm-thick are stacked all along the normal direction (ND). The dislocation density in the α-phase is on the order of 9.4x10¹³ m⁻² and TEM observations highlight dislocation sub-boundaries.

Hydrogen charging conditions of samples by cathodic polarization were mainly performed in artificial seawater (ASTM-D1141) at pH 8 and 30°C with potentials between -0.8V/SCE and -1.8V/SCE. Hydrogen charging were also made with these potentials and temperature in NaCl solution (35g/L) pH 7 and in H₂SO₄ (0.1M) solution at -15mA/cm² and 25°C to verify some assumptions. All the samples were charged on the ND face. Hydrogen concentration was measured by thermo-desorption spectroscopy (TDS Horiba EMGA Serie) on 10x4x0.5 mm samples after specific charging steps on the side of 104x4 mm (2400-SiC polished before charging). Others hydrogen charging conditions were made on 10x10x2mm OPS polished samples. XRD (Bruker advance AXS D8 with Cu source) and AFM were performed on samples charged at different times corresponding to each regime of the -1.8V/SCE artificial seawater absorption curve. Then, transversal surfaces were polished until OPS Master-Met vibro-polishing and observed by SEM and EBSD (FEI Quanta 200 ESEM/FEG with EBSD TSL/EDAX). TEM samples were taken parallelly to the surface exposed to the cathodic charging and electro-polished with a solution composed by 70% methanol, 20% 2-butoxyethanol and 10% perchloric acid at 20V until the drill by Struers TenuPol-5. TEM observations were performed with a JEOL JEM 2011 at 200 kV.

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3. Results

Hydrogen absorption

Figure 2 shows the evolution of hydrogen concentration in time for grade 2 and grade 23 with different cathodic potentials in artificial seawater. It is first observed that there is a significant absorption of hydrogen in these durations only with potentials equal to -1.6V/SCE or more negatives. On -1.8V/SCE curves, we can observe four regimes of absorption depending on time and hydrogen concentration. Hydrogen picking linearly rises with time until 30h (regime I) then a quasi-plateau is observed for both alloys (regime II) at 250 wppm for grade 2 and 300 wppm for grade 23. After 60h for grade 2 and 72h for grade 23, hydrogen uptake rises again (regime III) until the last step at 1000 wppm after 96h for grade 2 and 2000 wppm after 144h for grade 23 (regime IV). Charging performed in H2SO4 for 96h on grade 2 and in NaCl for 144 h on grade 23 show the same kind of behaviour but nothing can be assumed for the last step, which was not clearly reached.

Hydrides formation

XRD analyses in Figure 3 were performed on samples charged at -1.8V/SCE in artificial seawater for each regime of the absorption curve (Figure 2). We see the formation of γ-hydrides TiH (FCT) and δ-hydrides TiHx (1.5). Some bumps are observed on the surface and their dimensions could not allow measuring a Ra by AFM. A small evolution of the roughness of grade 23, Figure 3 is observed on regime II and an important uptake is seen on regime III. In parallel to these analyses, SEM revealed the formation of hydrides in the two alloys with the same evolution as analysed by XRD. Figure 6 shows the formation of a hydride layer on the surface of grade 2. A 2 µm-thick inhomogeneous layer is formed at regime I, transforming to a homogeneous 4 µm-thick layer at regime II and to a 16 µm-thick one at regime III. Hydrides have been detected around iron inclusions from regime I even in the bulk. From regime II, the formation of hydrides in all the bulk is noticed. SEM analyses performed on grade 23 (Figure 7) revealed the formation of hydrides from regime III. An evolution of the localisation of hydrides from the surface to the bulk is noticed. From the surface up to 100 µm in depth, intergranular and intragranular hydrides are observed, whereas only intergranular hydrides are present in the region between 100 µm and 800 µm in depth. From 800 µm in depth to the rest of the bulk, no hydrides are identified.
Figure 3: XRD analyses (a) symmetric on grade 2 and (b) asymmetric 5° on grade 23 after cathodic polarization at -1.8V/SCE in artificial seawater.

Figure 4: Optical and AFM analyses on the exposed surface of grade 2 after cathodic polarization at -1.8V/SCE in artificial seawater.

Figure 5: AFM analyses on the exposed surface of grade 23 after cathodic polarization at -1.8V/SCE in artificial seawater.
Figure 6: SEM analyses on grade 2 without charging and charged 10h, 30h and 72h in artificial seawater at -1.8V/SCE.

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Figure 7: SEM analyses on grade 23 charged until 50h, 96h and 144h in artificial seawater at -1.8V/SCE. Measures represents the depth from the surface to the bulk until this kind of structures are seen.

Regimes I-II / until 50h | Regime III / 96h | Regime IV / 144h | Hydrides identification

20 µm | 20 µm | 20 µm | 10 µm
100 µm | 150 µm | α Hydrides β
800 µm | 550 µm

TEM analyses revealed further details (Figure 8, Figure 9). On grade 2, Figure 8, the δ-hydride layer grown on regime II is nanostructured and becomes homogeneous on regime III. Different orientations relationships between γ-hydrides, δ-hydrides and the substrate were identified [4]. γ-hydrides in OR2 were observed in regime I. In regime II, incoherent γ-hydride and δ-hydrides in OR2 and OR3 are present. The regime III revealed mainly δ-hydrides in OR2 in the bulk. In grade 23, Figure 9, three kinds of hydrides were identified in regime III. First, δ-hydride in OR1 grows from α/β interface toward the bulk of α-phase. Then, δ-hydride OR2 formed at the interface of α-grain and β grain in epitaxy [5]. Finally, γ-hydrides formed parallel to each other in OR2 with α-grains.
Figure 8: TEM analyses on hydrides formed at 50 μm depth in titanium grade 2 after 10h (regime I), 30h (regime II) and 72h (regime III) of hydrogen charging by cathodic polarization at -1.8V/SCE in artificial seawater.

Figure 9: TEM analyses on hydrides formed at 50 μm-depth in titanium grade 23 after 96h (regime III) of hydrogen charging by cathodic polarization at -1.8V/SCE in artificial seawater.
4. Discussion

Despite a large difference between both microstructures of the alloys studied, similar behaviour were observed in term of hydrogen absorption (Figure 7). The absorption rate of hydrogen at applied potentials ranging from -0.8 to -1.5V/SCE in artificial seawater is low in the range of our considered times (<200h) with hydrogen concentration less than 100 wppm. This result is in good agreement with previous works. Absorptions of hydrogen up to 120 wppm were identified at -1.05 mV/Ag/AgCl after 2 years of polarizations of grade 2 [6] and 100 wppm after 1 year of polarizations on grade 23 (stabilized at 90 wppm measured after 2 years) [7]. In opposite, for -1.8 V/SCE, a largest increase of hydrogen has been identified. Four regimes were observed (Figure 2). Stage IV corresponds to a pseudo saturation equal to 1000 wppm for grade 2 and 2000 wppm for grade 23. Polarization curves of grade 2 and grade 23 in artificial seawater presents three domains, A from -0.5 V/SCE to -1 V/SCE then B up to -1.5V/ECs and C from this value to more negatives potentials. Domains A and B restrict the hydrogen absorption because it is in competition with oxygen reduction and formation of Aragonite calcareous deposit on the surface. In domain C, there is the formation of inhomogeneous Brucite and Calcite. The peculiar kinetic modifications seen with the -1.8V/SCE curve suggests surface modifications on samples. The calcareous deposit formation could be an explanation for the quasi-plateau of regime II. But after performing hydrogen charging in NaCl and in H2SO4, which don’t produce calcareous deposits, this step is still present. It could also be caused by a modification of the behaviour of the titanium oxide film, as seen by Ohtsuka [8] and Vezvaie [9], which chemically change from TiO2 to TiO2H on cathodic polarization in neutral solutions. Therefore, Ohtsuka [8] also determined that on cathodic polarization in acidic solutions, the oxide TiO2 is reduce. Consequently, regimes I and II are only associated with hydrogen activity.

On the grade 2, a homogeneous nanostructures hydride layer in sub-surface was observed for 30h of polarization at -1.8V/SCE, Figure 6. This layer becomes massive at 72h. The quasi-plateau (regime II) could take his origins on the slow-down of hydrogen diffusion during the homogenisation of the hydride layer which present a two decades lower diffusion coefficient (10^-16m2/s in hydride at 30°C [10,11]) than in α-phase (10^-14m2/s in a 30°C [12]). Hydride formation leads to volume expansion (12, 5-14% for γ-hydrides and 21-29% for δ-hydrides [1]) which explain the evolution of roughness over time of grade 2, Figure 4. Even if a hydride layer forms, hydrides clusters were observed in the bulk. (Figure 6 and Figure 8). The structure of this grade 2 has a particularity, his grains are mostly with strong misorientations, which means a large fraction of random grain boundaries with high excess volume. This could form diffusion short-circuits and allow a part of hydrogen atoms to penetrate in all the bulk and form a large fraction of hydrides in stage III. To resume, four processes are in competition during the hydrogen charging: the growing of a hydride phase from the surface and three processes of the hydrogen diffusion (a first in the α-phase, a second other in the hydride phase and third along the grains boundaries).

In the grade 23, first hydrides formation were identified on regime III, Figure 3, Figure 7, Figure 9. Therefore, AFM analyses, Figure 5, show a small evolution of the roughness even before these formations. Hydrogen solubility limits are from 20 to 100 wppm in α-phase at 25°C depending on sources [1, 13, 14, 15] and is much higher in β-phase. Regarding solubility limits of β-alloys and phase-diagrams of titanium, the solubility of hydrogen could reach around 5 000 to 10 000 wppm at 25°C in β-phase [13, 14, 16, 17, 18]. Hydrogen diffuses also faster in β-alloys [18] and is a β-stabilizer [19, 20, 21]. For these reasons, we can assume that hydrogen diffuse in β until its solubility limit are reached (regime II). Before this point, β growth and increases the roughness of the surface, then hydrogen cannot be absorbed by β anymore (only 6% of the volume fraction of phases), which can cause the quasi-plateau of the regime II. In regime III, hydrides form first on interfaces and when they are saturated, hydrogen penetrates in α-grains, quickly reach the low solubility limit of this phase and forms hydrides (Figure 7, Figure 9). The quantity of hydrogen being high in sub-surface allow the formation of intergranular and intragranular hydrides. Below, there is not enough hydrogen for the formation of intragranular and just intergranular are present. Deeper, no more hydrides are present because of the fewer hydrogen concentration. Hydrides and β-phase volume expansion explains the strong roughness’s rise on regime III, Figure 5. This expansion cannot be accommodated in the bulk and induces long-range internal stresses.

Finally, the calcareous deposit becomes more compact with time, stabilized with the help of the roughness increase (this phenomenon has been studied to improve the stability of implants and prosthesis for biomedical purposes [22, 23, 24]), and lead to a decrease of the current intensity by adding resistance on the surface and could cause the stabilization of regime IV. The saturation of the sub-surface by hydrides also strongly decrease the absorption rate for both alloys.

5. Conclusion

Different kinetics of hydrogen absorption in grade 2 and grade 23 during hydrogen charging by cathodic polarization in artificial seawater have been highlighted. For our charging durations (until 200 hours), only cathodic polarizations of -1.6V/SCE and more negatives lead to a noticed hydrogen uptake. Kinetic variations of hydrogen absorption during polarizations at -1.8V/SCE have been attributed first to a competition between the diffusion processes and the formation of hydrides for grade 2 and to the ingress of hydrogen in β-phase reaching the solubility limit for grade 23, then to the growth of a large fraction of hydrides in the bulk. On longer durations the diminution of the current intensity is caused by the presence of a calcareous deposit and the saturation of sub-surface by hydrides. The formation of γ-hydrides have been localized in grade 2 as a layer that increases the surface roughness, then γ-hydrides in OR2 and δ-hydrides in OR2 and OR3 have been localized as clusters in the bulk. In grade 23, hydrogen is absorbed by β-phase leading to a volume dilatation of this phase affecting the surface roughness. Then after reaching the hydrogen solubility limit of β-phase, first δ-hydrides in OR1 and OR2 forms on interfaces α/β and α/a and with more hydrogen, γ-hydrides in OR2 forms in α grains. All these kinetics can affect the damage processes, which will be discussed in a further works.

References


