Degradation of Titanium Sintered with Magnesium: Effect of Hydrogen Uptake

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Abstract: Multifunctional materials based on a combination of permanent and degradable metals open new perspectives for medical implants combining osseoconductivity and drug-delivery functions which can significantly decrease the number of implants’ revision. In this work, hybrid magnesium-titanium materials were produced via sintering, and the properties of the permanent titanium component before and after the degradation of the temporary magnesium part were evaluated. The changes of chemical composition and mechanical parameters were determined. Loading of hydrogen into the titanium part at room temperature was observed, which deteriorated the mechanical characteristics but could also simultaneously improve the biocompatibility of the permanent titanium implant. The control of degradation of the magnesium part and the modification of the titanium part are required for the development of partly degradable hybrid implants.

Keywords: titanium; magnesium degradation; hydrogen uptake; titanium hydrides; mechanical properties; hydrogen embrittlement

1. Introduction

Metallic implants are generally applied in orthopedic and craniofacial surgery, dental practice and as vascular stents [1]. Titanium and titanium alloys, due to their high strength, low density, high corrosion resistance, inertness to body environment and excellent integrity to the bone are widely used from hip to wrist applications and as bone fixation materials [2]. In vitro studies of titanium foams and human osteoblasts showed that cells differentiate into mature bone cells with osteoinductive properties, which makes titanium unique among metals [3]. Magnesium, as a biodegradable metallic material, offers additional opportunities due to its biocompatibility, resorbability and antibacterial properties [4]. Controlled degradation of magnesium implants is already used in modern medicine (confirmed by recent meta-analysis studies of clinical trials [5]), with growth potential as the role of Mg in bone-healing becomes clearer [6].

It is quite natural that there are significant efforts to combine the best properties of both materials (Mg and Ti), aiming to overcome well-known limitations of metallic implants, such as the stress shielding effect [7] and the susceptibility to infections [8]. The hybrid implant material, which consists of permanent Ti and temporary Mg parts (bimetallic couples), can be used in the cases when high strength of titanium and bone stimulation by degradation of magnesium are required and/or only partial bone remodelling is expected, as in the case of revision surgery [9]. These hybrid materials enable also the design of drug-delivery devices, i.e., loading the permanent and/or the temporary parts with drugs.

Application of magnesium films on porous titanium and the local release of magnesium ions from the implant surfaces were reported to enhance implant retention, osteogen-
esis and angiogenesis [10–12]. The development of a hybrid fixation system which includes the combination of Mg and Ti screws instead of only Ti screws has been announced [13,14]. It was possible not only to reduce the number of screw removals after healing but also to improve bone regeneration, suppress bone resorption and induce blood vessel formation combined with an antibacterial effect. In a rabbit model, the failure load has increased by 30% after 12 weeks of application [14].

There are special requirements for the connection combining Ti and Mg parts: it has to be strong enough for implant positioning in the body and, in case of titanium alloys, the possible diffusion of alloying elements from the Ti part into the Mg part should be negligible. The low solubility between Ti and Mg is a good presumption for the latter demand but may hinder a tight connection on an atomic scale [15]. However, studies on porous Ti/Mg composites where Mg plays a role of the degradable anodic part, which gives space for the ingrowing cells, show that the cohesion between Mg and Ti is sufficient for proper mechanical characteristics [16].

Another strong concern on simultaneous use of Ti and Mg is the large difference in electrochemical potential of Mg and Ti, which could lead to accelerated degradation of Mg. In vivo and in vitro studies demonstrated that the distance of 5 mm between Mg and Ti parts is critical for the degradation of Mg. Most probably, blood vessels form a good electrical connection which enhances degradation of Mg [17].

To overcome increased corrosion, it is possible to use a polymeric layer between Mg and Ti parts as it has been applied in a hybrid fixation system [13]. Another possibility is the selection of suitable Mg alloys like aluminum- and zinc-containing AZ alloys that still show an acceptable degradation rate in contact with titanium alloys [18,19]. However, the presence of aluminum limits their application as implant material.

In the above-mentioned investigations, it was postulated that the Ti-material is unaffected by Mg degradation. To investigate whether this assumption is justifiable, in the present study we have constructed Ti-Mg-alloy and Ti64-Mg-alloy (here: Ti64 defined as Ti-6Al-4V alloy) hybrid specimens via a powder metallurgy approach. Powder metallurgy technologies based on sintering, like metal injection molding (MIM) are frequently used to produce specimens with variable shape and mechanical characteristics and are also applied for Ti and its alloys [20,21]. The sintering of Mg powder is more challenging [22,23] due to higher melting temperature of Mg oxide layer than bulk Mg. However, the addition of a small amount of Ca helps destroy the surface oxide layer of the Mg particles, which otherwise hinders sintering. Principally, sintering gives a quite good mechanical connection between Ti and Mg and could be used for the production of implants [24]. However, the effect of degradation on the Ti-material has not yet been investigated. In the present work, the degradation of the Mg part of the hybrid specimens in saline solution was performed, and related possible changes in composition, phase structure and mechanical properties of Ti part were studied. Significant changes of the titanium parts have been observed, which should be taken into account in further simultaneous applications of Mg and Ti implants and can be used for optimizing the properties of future implants.

2. Materials and Methods

2.1. Preparation of Specimens

A rod of Titanium Grade 2 (ASTM B348) was obtained from Ankuro Int. (Rostock, Germany). This rod (with diameter 10 mm) was cut into discs with the thickness of 1.5 mm by using a diamond saw. Mg-0.6Ca alloy was produced by casting and subsequent extrusion with a speed of 4.4 m/s. Mg-0.6wt.%Ca discs of 9 mm diameter and 1.5 mm thickness were machined from the rod. Ti and Mg discs were cleaned and put pairwise into the sintering furnace (RRO 350-900, MUT Advanced Heating, Jena, Germany) for 8 h at 640 °C under Ar atmosphere.

For tensile test measurements, dog-bone shaped samples of Ti64 were manufactured via the metal injection molding (MIM) method and sintered (see the Supplementary Materials). The middle part of the obtained Ti64 dog-bone samples was covered with Mg-0.6Ca
also using MIM (Figure 1) and sintered (see the Supplementary Materials). The thickness of the Mg-0.6Ca feedstock layer in the middle part before sintering was 5 mm.

![Figure 1](image-url)  
**Figure 1.** Dog-bone-shape tensile test samples of hybrid Ti64/Mg-0.6Ca after metal injection molding (MIM) production (a), sintering (b), corrosion (c) of Mg and tensile test (d).

### 2.2. Characterisation

Corrosion experiments were performed in 0.9 wt.% NaCl solution at room temperature, and the corrosion rate was followed by the hydrogen evolution method. Corrosion products were removed from Mg samples by washing them in a 200 g·L\(^{-1}\) solution of chromic acid for 10 min. High-performance micro-X-ray fluorescence spectrometer (M4 TORNADO, Bruker Nano, Berlin, Germany) was used to verify the full removal of Mg products from the Ti surface.

Determination of chemical composition (contents of O, N and H) in the titanium part before and after degradation of Mg was done via inert gas fusion technique using a LECO apparatus (ONH836, LECO, St Joseph, MI, USA). For this, Ti discs and Ti64 dog-bone-shape samples were cut into pieces of approximately 50 mg each by using a diamond saw and finally cleaned. Results were obtained as average value of three pieces, and the measurement was repeated for three different Ti discs.

Phase identification was performed by grazing incidence X-ray diffraction (GIXRD) (see the Supplementary Materials) using a Bruker D8 Advance X-ray diffractometer (Bruker AXS, Karlsruhe, Germany).

Tensile tests were performed according to DIN EN ISO 6892-1:2009 B on a universal materials-testing machine (ZwickRoell, Ulm, Germany). Images of the Ti cross section were recorded using scanning electron microscopy (VEGA3 TESCAN-15kV, TESCAN, Kohoutovice, Czech Republic, see the Supplementary Materials).
3. Results

After sintering heat treatment of the Mg-0.6Ca/Ti discs at 640 °C for 8 h a tight connection was obtained between the two materials (Figure 2) which was expected from previous investigations [24]. For the corrosion experiments, Mg-0.6Ca/Ti couples were placed into saline solution (0.9 wt.% NaCl). The degradation of the Mg-alloy was very fast (Figure 2). In contrast to the uncoupled specimen, the Mg-alloy part was completely dissolved after only two days of immersion. Afterwards, there was no additional H$_2$ evolved. This indicates that Mg was fully dissolved based on the following reaction:

$$\text{Mg} + 2\text{H}_2\text{O} \rightarrow \text{Mg(OH)}_2 + \text{H}_2 \uparrow$$ (1)

This is in agreement with recent studies of Mg(-alloy)/Ti64 composites where pure Mg and WE alloys showed very high degradation rates [18] with complete degradation in 1–2 days. The amount of released H$_2$ (~800 mL) corresponded to the total mass of Mg (0.819 g), according to Equation (1).

![Figure 2](image2.png)

**Figure 2.** Photo of Mg-0.6Ca and Ti plates after heat treatment at 640 °C for 8 h (left) and the evaluation of H$_2$ for Mg-0.6Ca/Ti couple in saline solutions (0.9 wt.% NaCl) and, in comparison, Mg-0.6Ca alone (right).

After corrosion experiments, the Ti plates were cleaned by chromic acid from corrosion products of Mg. There were no traces of Mg on the Ti plates detectable by X-ray fluorescence analysis. On the other hand, the Ti plate partially changed the color (Figure 3) which can typically be connected to a variation of the oxide layer thickness. The comparison of the weight of the titanium plates before and after corrosion of the Mg-part pointed to a weight gain (Figure 4). Here, five Ti discs have been analyzed. It meant that titanium absorbed some elements during corrosion of Mg. However, the error bars were rather high, so the effect appeared to be small. In Figure 4, a comparison is shown to Ti samples which have been treated in a similar way (exposition in saline solution and chromic acid treatment), but without connection to Mg.

![Figure 3](image3.png)

**Figure 3.** Photos of Ti plates before (left) and after (right) corrosion of Mg.
Figure 4. The change of weight of the Ti plate after exposition to saline solution and subsequent chromic acid treatment. Left bar (Ti): without coupled Mg; right bar (Ti after Mg corrosion): with coupled Mg. Numbers are average values of five discs.

For chemical analysis, the samples after degradation of Mg were cut into smaller pieces (~50 mg) and compared to uncoupled Ti after the same treatment. The chemical analysis showed that there was a small variation of concentration in O and N and a significant increase in the content of H (Figure 5).

Figure 5. Content of H, N and O in titanium part: initial Ti, after degradation of Mg alloy sintered with Ti and after additional heat treatment at 1100 °C in a vacuum.

It should be pointed out that such a significant increase of H content from below 50 µg/g to above 1000 µg/g took place at room temperature and was the direct result of Mg corrosion. Additional tests have been performed to check out other possible sources (chromic acid treatment, simultaneous heating of Ti and Mg) of H loading within the applied process. However, the results were negative and those factors could not have caused hydrogen absorption by Ti. By application of high temperatures (1100 °C) and
vacuum for 2 h, the loaded hydrogen was released again, so the concentration decreased below 1 µg/g (Figure 5). Thus, it was proven that indeed hydrogen was loaded into titanium at room temperature, which is typically not expected because of the inherent dense and protective oxide layer on Ti.

It is quite important to know how the hydrogen was distributed in Ti. SEM images showed only some dark regions on the surface of the Ti plates which could be the location of H loading (see the Supplementary Materials, Figure S1) and EDX is not sensitive to low weight elements such as H. To overcome this problem, we have performed grazing incidence X-ray diffraction. Figure 6 presents the comparison of GIXRD patterns of initial Ti and Ti coupled with Mg after degradation of Mg. The initial titanium sample showed almost only peaks (100), (002), (101), (102), (110), (103), (112), (201) of crystalline α-Ti in hexagonal phase P63/mmc a = 2.9504 Å, c = 4.6933 Å (PDF 03-065-3362, Data 2015). In contrast, the coupled Ti sample after magnesium degradation showed peaks of Ti (similar to initial Ti) and additional peaks (111), (200), (220), (311), (222) from TiH₂ cubic phase (fcc) a = 4.42 Å (PDF 00-009-0371, Data 2015). The results suggest that hydrogen emerging during Mg degradation saturates the Ti-phase and leads to the formation of hydrides. Again, it should be mentioned that the transformation from pure hexagonal Ti phase to a mixture with the hydride phase takes place at room temperature. Industrially, a hydrogenation process is applied for some purposes. However, process temperatures in interval from 600 °C to 700 °C are usually required.

![Figure 6. Grazing incidence X-ray diffraction (GIXRD) data of titanium plate and titanium plate sintered with Mg alloy after degradation of Mg alloy.](image-url)

It is well-known that loading of a high amount of H in Ti and the formation of titanium hydrides change the mechanical properties of titanium parts rather drastically [25,26]. This might significantly affect the implant properties depending on the application. To check the evolution of mechanical properties with hydrogen uptake, we used the dog-bone samples produced by MIM consisting of Ti64 and Mg-0.6Ca (Figure 1). Even if the corrosion properties of Ti and Ti64 are not absolutely identical [27] and Ti64 shows higher corrosion resistance in Ringer’s artificial simulated body fluids, the basic results related to hydrogen uptake are not supposed to be affected in the frame of this study. After the same procedure of sintering heat treatment for the plates, tensile tests were performed after corrosion experiments in 0.9 wt.% NaCl (Figure 1). They were compared with tensile tests on uncoupled sintered Ti64 specimens. As it can be seen in Figure 7, the uncoupled Ti64 sample showed elastic and plastic features with elongation to fracture up to 12%. However,
all hybrid samples after corrosion of Mg have been broken early within the elastic area. The location of fracture for all hybrid samples are not in the center, which is shown in Figure 1d. In the case of initial Ti64, the fracture was always in the center of the sample. Thus, by uptake of hydrogen, the ductility was completely lost. The average content of hydrogen in the titanium part of the dog-bone samples after degradation of Mg was approximately two times higher ($2130 \pm 98 \mu\text{g/g}$) than in the plate samples. This was probably related to the fact that the Mg part covered the Ti alloy part almost completely.

![Stress-Strain Curve](image1)

**Figure 7.** Results of tensile tests for initial Ti64 (solid line) and Ti64 sintered with Mg alloy (three samples) after Mg degradation (broken lines). Insert shows the elastic region of measurements. Measurements of initial Ti64 were repeated three times which showed the similar behavior in elastic and plastic regions. One curve is presented to keep the figure clear.

The locations of the fracture induced by the tensile test have been examined using SEM. For the initial sample (Figure 8, left) white and dark regions were visible very clearly, corresponding to $\beta$ (white)—high solubility of hydrogen, and $\alpha$ (dark)—low solubility of hydrogen phases. After corrosion of Mg (Figure 8, right) it appeared that the structure became finer, and the comparison with the pseudo-binary phase diagram of Ti64-H [28] indicated the transformation to $\alpha_2$ (TiAl$_3$) and $\delta$ (highest solubility of hydrogen, TiH$_2$) phases with visible boundaries of $\alpha/\beta$. This fully matches the results from the diffraction experiments.

![SEM Micrographs](image2)

**Figure 8.** Scanning electron microscopy (SEM) micrographs of the fracture area of the initial Ti64 sample (left) and Ti64 sintered with Mg alloy after Mg degradation (right).
4. Discussion

Any unintended changes observed in permanent implant materials are undesired. The variation of color and weight of the titanium part after degradation of magnesium should be taken with significant attention. The observed weight gain was connected with adsorption of hydrogen and the change of thickness of the titanium oxide layer, which causes the change of color.

Hydrogen is known for prompting embrittlement in most metallic materials. Pure titanium is less susceptible and shows good ductility for up 25 at.% hydrogen [29]. In the present study, the estimated concentration of hydrogen (around 5–10 at.%) should be below the limit mentioned above but infringing ASTM standards (<150 ppm) for common Ti materials [30]. The tensile tests suggested that the actual amount of adsorbed hydrogen was higher than the plasticity limit for Ti64. Hydrogen forms mainly the hydride TiH$_2$ (face centered cubic) which can be attributed to δ phase and usually it is observed for hydrogen loading into titanium at high temperatures (~750 °C [31]) or after cathodic hydrogenation at 80 °C followed by thermal diffusion at 225 °C [32]. According to the phase diagram of Ti-H [33,34] the formation of δ phase takes place in the concentration range for H between 51.2 and 66.7 at.%. As we did not perform homogenization of Ti and Ti64 parts, it is possible that at the surface of Ti, the concentration of H was much higher than the average value obtained from chemical analysis.

The observed loading of a high amount of hydrogen into titanium takes place at room temperature under the immersion conditions and in presence of a high concentration of saline solution. The potentially important parameter is increased local pressure, it is known that by fast degradation of Mg in Mg/Ti-alloys composite, the gas pressure of formed hydrogen can be so high that it breaks the pre-sintered scaffold of titanium [18].

The other factor that could promote hydrogen absorption is a loss in stability of the oxide layer by presence and degradation of Mg. It is possible that a part of oxygen migrates from the TiO$_2$ layer to Mg because of the extremely high affinity of Mg to oxygen. This could also be a reason for the astonishingly good bonding by sintering between two metals with very low mutual solubility [24]. The destabilization of the oxide layer makes the surface of Ti easily accessible for H, which is formed during degradation of Mg. The most suitable place is the line where Mg, Ti and the solvent are in contact. We should note that fracture of all samples after degradation of Mg took place in this part of sample (Figure 1d) and not in the middle as in the initial samples. Figure 9 shows that the profile of hydrogen absorption was not constant and showed two maxima (~0 and ~25 mm) at the lines of Mg, Ti and solvent contact. The maximum at 0 mm was significantly higher (here, fracture takes place). The reasons for higher maximum at 0 mm could be the higher local residual porosity of Ti64 and/or slightly higher amount of Mg.

The observed hydrogen loading can be explained by a metallic reduction reaction (MRR) which is widely used in material science [35]. According to the MRR mechanism, TiO$_2$ is reduced to Ti by Mg that, in turn, oxidizes to MgO:

$$2Mg + TiO_2 \rightarrow 2MgO + Ti$$ (2)

As a result, there is a direct contact between Mg and Ti, which leads to transfer of electrons from Mg (→MgO) to TiO$_2$ (→Ti) due to difference in electrochemical potential. The overall reaction of magnesium corrosion (1) consists of anodic magnesium oxidation Equation (3) and cathodic water reduction Equations (4) and (5)

$$Mg - 2e^- \leftrightarrow Mg^{2+}$$ (3)

First proposed by Höche et al. [36] and then proved by Density Functional Theory (DFT) calculations, it was shown that the Volmer–Heyrovsky reaction sequence is the predominant and most favorable, from the standpoint of energy barrier, pathway for hydrogen evolution during degradation of magnesium [37]. It starts with the water
reduction step, according to Volmer, accompanied by adsorption of atomic hydrogen on magnesium surface:

$$H_2O + e^- \leftrightarrow OH^- + H_{ad} \quad \text{(Volmer)}$$  \hspace{1cm} (4)$$

Adsorbed hydrogen reacts with a partially positive hydrogen atom of the water molecule in the Heyrovsky step:

$$H_2O + e^- + H_{ad} \leftrightarrow OH^- + H_2 \quad \text{(Heyrovsky)} \hspace{1cm} (5)$$

The next step is hydroxide formation:

$$Mg^{2+} + 2OH^- \rightarrow Mg(OH)_2 \hspace{1cm} (6)$$

Figure 9. Concentration of absorbed H along the length of Ti64 dog-bone sample after degradation of Mg alloy.

The formed Mg(OH)$_2$ is not fully passivating. It does not fully block the access of water and Cl-species to MgO/Mg interface. Thus, corrosion of Mg continues even under the bulky layer of Mg(OH)$_2$.

The $H_{ad}$ or $H_2$ can be adsorbed by titanium given a high concentration of hydrogen, that is enough for the formation of hydrides following Equations (7) and (8).

$$Ti + H_2 \leftrightarrow TiH_2 \hspace{1cm} (7)$$

$$Ti + 2H_{ad} \leftrightarrow TiH_2 \hspace{1cm} (8)$$

The temperature of 640 °C, at which Ti and Mg parts were kept to get good bonding, is quite important for the MRR process and most probably without bonding, the loading of H into Ti will be not so high.

From the other side, room temperature Mg reduction of TiO$_2$ has been described recently [38]. It was shown that the contact between Mg and the oxide of Ti induces the direct transfer of electrons, which facilities the diffusion of oxygen from interfacial TiO$_2$ and collecting H from the HCl solution. Theoretical calculations have shown that due to the connection of Mg and TiO$_2$ the energy of reduction of TiO$_2$ decreases by about $-1.27$ eV, and this negative value reveals that the process of hydrogenation is exothermic [38]. In
addition, the energy of formation of oxygen vacancies is more than 1 eV lower for a Mg-
TiO$_2$ connection in comparison with TiO$_2$ standalone \[38\]. A similar process was possible
in the present system (Figure 10). There is electron transfer from Mg to TiO$_2$ which leads to
enhanced hydrogenation. The pressure due to formation of H$_2$ gas increases and plays an
important role in hydrogenation, as it is shown in \[38\].

![Figure 10. Possible scheme of loading of hydrogen (H$_{ad}$ and H$_2$) into the titanium part.](image)

Formation of hydrides at the surface of titanium implants can be an important factor
for improving biocompatibility and usually requires the etching in acids or application
of electrical discharging \[39\]. It has been shown that treatment by concentrated sulfuric
acid leads to the production of titanium hydrides with simultaneous increase of surface
roughness and weight loss of titanium samples \[40\]. In the present study, the hydrides
were obtained without such aggressive treatment. We propose that the degradation of
Mg not only leads to improved bone growth due to the release of magnesium ions, but
also potentially can improve biocompatibility of titanium implants via hydrides formation.
Identified formation of hydrides provides additional explanation of enhanced biocompa-
tibility of titanium implants after loading with magnesium, which was recently observed
in many cases \[10–12,41\]. However, this aspect could not be investigated in the frame of
this work.

5. Conclusions

In a saline solution, fast degradation of a Mg-part sintered to a Ti-part was promoted
by galvanic coupling and accompanied by enhanced hydrogen evolution with further
penetration of hydrogen into Ti at room temperature. High hydrogen content initiated the
formation of titanium hydrides and a phase transformation from $\alpha$ and $\beta$ to $\alpha_2$ and $\delta$
phases. Mechanical characteristics of Ti were strongly affected, i.e., the material became
very brittle. Plastic elongation decreased from over 12% to below the detection limit. The
formation of titanium hydrides (hydrogenation of titanium) by degradation of magnesium
could substitute such complicate treatment as etching of titanium in acids that is applied
to improve biocompatibility of Ti. Subsequent change of mechanical properties has to be
carefully considered.

Supplementary Materials: The following are available online at https://www.mdpi.com/2075-470
11/4/527/s1, Figure S1: Photos and SEM images of titanium plates. Details on sample preparation.

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