Introduction

Absorbable metallic alloys have an outstanding potential for biomedical applications when their biocompatibility is adequate and its release sub products have toxicity to host tissues [1]. Although most of the biodegradable materials developed are based on polymers and ceramics, these materials have limitations regarding mechanical strength and brittleness, respectively [1]. The advantages of magnesium alloys as implantable devices for bone fracture stabilization rely on its adequate mechanical strength, toughness, and a variable elastic modulus most similar to human bone tissue. The basal slip or twinning strain from the hexagonal closed packed structure ensure improvements into mechanical strength since not all the five independent slip systems become operative at room temperature [2,3]. The potential wear debris and metal ions release toxicity expected depends on the friendly interactions of the metallic elements with human body [1]. In this perspective, magnesium has several advantages, being one of metals most committed in human metabolism, in a content just overcome by potassium element [1].

Magnesium has participation in different mechanisms reaction in human body like bone turnover process and maintenance of inorganic content of mineralized structures as teeth [1]. When associated with calcium, its potential enhances, encompassing cell membranes and intracellular organelles, systemic triggers glands and its hormones release, neural signal transduction, muscles and blood clotting [4,5]. Calcium also improves friendly behavior of magnesium in host body, as recognized by its co-factor for many enzymes and with stimulatory influence on growth of new mineralized tissues [4,5]. The performance of an absorbable alloy under physiologic environment is gambled on interactions between corrosion degradation and maintenance of its mechanical integrity [6]. In cases of device for bone fractures stabilization the degradation must occur at a range in synergism with the ability of human tissue to heal itself, otherwise failure should take over before bone fractures were consolidated [7,8].

Rare earth (RE) elements addiction is one of the strategies for control the corrosion rate dissolution of magnesium alloys under physiological environment. Besides degradation control, RE elements can also enhance mechanical and physical properties at once, allowing slower alloy deterioration, which prevent significant amounts of hydrogen evolution during corrosion process [9]. As fast as bio-absorbable magnesium alloys are degraded, higher is the possibility of painful gas pockets emergence interspersed to...
local tissues around implantation sites, serious local alkalization and hemolysis [4,6]. The objective of this work was to compare the corrosion resistance of Mg-Ca based alloys with and without RE alloy under simulated physiological environment.

Materials and Methods

Mg-Ca-Gd alloy was produced by casting process from pure Mg (99.9%), Mg-Ca (30 wt. %) and Mg-Gd (30 wt. %) master alloys. Mg-Ca was casted from pure Mg and Ca elements. Melting process was executed in an induction crucible furnace at a melting temperature of 720°C, under a protective atmosphere of purged SF6 gas. Stirring (at 720°C) was performed for 5 minutes and molten metals were casted in cylinders ingots 80 mm in diameter. Cast ingots were annealed at 400°C for 12h and hot extruded in rods with 15 mm in diameter. For electrochemical test cylindrical samples with 1 mm in thick were prepared superficially in SiC paper up to grit of 1200 and included into a sample holder with an exposure area of 0.1257 cm². The electrochemical test was performed using a three- electrode flat cell with physiological solution (0.9mol L⁻¹ NaCl) as electrolyte. Corrosion cell was filled with 500 mL of electrolyte and maintained at 37°C under a pH=7.4. Saturated calomel electrode was used as reference and platinum was addressed as counter electrode, both connected to a potentiostat (PG-3901, OMNIMETRA INSTRUMENTS). Potential dynamic measurements were acquired ± 1.5 V of the open circuit potential of each sample at scan rates of 1, 0.1 and 0.01 Vs⁻¹.

In order to compare results, immersion corrosion tests were also carried out and mass loss was calculated. The samples were cut with close dimensions, sanded in 320 grit sandpaper, weighed in the analytical balance and they had their surface areas (initial) calculated. Each sample was placed in a conical tube with a capacity of 50 mL. Physiological solution (0.9mol L⁻¹ NaCl) was added; the samples were removed, cleaned with distilled water, weighed again and the electrolyte replaced into oven. This procedure was repeated with 48 and 72 hours of immersion. The tests were conducted in triplicate sample. The alloys had their chemical compositions determined by x-ray fluorescence spectrometry (Epsilon 3 UL PANalytical). Theoretical density and corrosion rates were calculated. The roughness measurement was performed by the Zygo three- dimensional rugosimeter model New View 7100 before and after the polarizations.

Results

Potentiodynamic polarization curves from Mg-40Ca and Mg-Ca-Gd alloys were shown in Figure 1. Experimental results were plotted in terms of I (current) vs. potential, being current the dependent variable when the potentiostat is initiated to overcome the free energy barrier by changes into potential intensity. Specific points considered for data acquirement included onset point, extrapolations and additional calculations Tafel constants, polarization resistance and corrosion rate. Table 1 shows the chemical composition of the alloys and theoretical densities. A density of approximately 1.80 g/cm³ was found for the MgCaGd alloy and 1.65 g/cm³ for the MgCa alloy. Both values are close to the density of pure magnesium which is 1.74 g/cm³. Open circuit potential (Table 2) of Mg-Ca and Mg-Ca-Gd alloys was -1.760 V and -1.550, respectively. OCP values identified on alloy with RE contents showed that these samples were chemically more stable than binary Mg-Ca alloy. Corrosion potential was equal to -1.715V for Mg-40Ca alloy and -1.550 V for Mg-Ca-Gd. From Tafel intersections and data calculations (Table 2), for tests conducted at the rate of 1 Vs⁻¹, corrosion current density was measured as 4.35E-03 A cm⁻² on Mg-40Ca alloy and 4.31E-03 A cm⁻² was measured from Mg-Ca-Gd. Tafel constants calculated from cathodic curves revealed a gradient of βc= -0.028 V dec⁻¹ for Mg-40Ca curve and βc= -0.033 V dec⁻¹ from Mg-Ca-Gd alloy. Polarization resistance was calculated as 4.35E-03 A cm⁻² on Mg-40Ca alloy and 4.31E-03 A cm⁻² was measured from Mg-Ca-Gd. Tafel constants calculated from anodic curves revealed a gradient of βc= -0.28 V dec⁻¹ for Mg-40Ca curve and βc= -0.033 V dec⁻¹ from Mg-Ca-Gd alloy. The corrosion rate was extrapolated as 104.637 mm year⁻¹ from Mg-40Ca sample and 95.620 mm year⁻¹ from Mg-Ca-Gd.
Table 1: Results of the chemical analyzes of the alloys and their respective calculated densities.

| Chemical Analysis (wt %) | Alloy       | Mg         | Ca         | Gd         | Calculated density (g/cm³) |
|--------------------------|-------------|------------|------------|------------|---------------------------|
|                          | MgCaGd      | 0.9132     | 0.03407    | 0.04727    | 1.80                      |
|                          | Mg40Ca      | 0.6084     | 0.38744    | --         | 1.65                      |

Table 2: Open circuit potential (OCP), corrosion current density ($i_{corr}$), corrosion potential ($E_{corr}$), Tafel coefficient at cathode ($\beta_c$), polarization resistance and corrosion rate from different samples. Rate was settled at 1, 0.1 and 0.01 V/s.

| Alloy      | OCP (V) | Rate (V/s) | $i_{corr}$ (A/cm²) | $E_{corr}$ (V) | $\beta_c$ | Polarization resistance (ohm/cm²) | Corrosion rate (mm/year) |
|------------|---------|------------|-------------------|----------------|-----------|----------------------------------|--------------------------|
| MgCaGd     | -1.550  | 0.1        | 1.23E-03          | -1.292         | -0.040    | 353.987                          | 27.228                   |
|            |         | 0.01       | 6.30E-04          | -1.132         | -0.040    | 689.599                          | 13.977                   |
| Mg40Ca     | -1.760  | 0.1        | 3.01E-03          | -1.537         | -0.037    | 144.246                          | 72.489                   |
|            |         | 0.01       | 1.82E-03          | -1.425         | -0.030    | 238.139                          | 43.908                   |

During electrochemical corrosion test with a rate of 0.1 Vs⁻¹, the corrosion current density was measured as 3.01E-03 Acm⁻² for Mg-40Ca alloy and 1.23E-03 Acm⁻² for Mg-Ca-Gd. Tafel constants calculated from cathodic curves revealed a gradient of $\beta_c = -0.037 \text{ Vdec}^{-1}$ for Mg-40Ca curve and $\beta_c = -0.040 \text{ Vdec}^{-1}$ from Mg-Ca-Gd alloy. Polarization resistance was calculated as 144.246 Ω/cm² on Mg-40Ca alloy and 353.987 Ω/cm² from Mg-Ca-Gd. The corrosion rate was extrapolated as 72.489 mm year⁻¹ for Mg-40Ca sample and 27.228 mm year⁻¹ for Mg-Ca-Gd. Finally for the electrochemical corrosion test with a rate of 0.01 Vs⁻¹, corrosion current density was measured as 6.30E-04 Acm⁻² for Mg-40Ca alloy and 1.82E-03 Acm⁻² was measured for Mg-Ca-Gd. Tafel constants calculated from cathodic curves revealed a gradient of $\beta_c = -0.030 \text{ Vdec}^{-1}$ for Mg-40Ca curve and $\beta_c = -0.040 \text{ Vdec}^{-1}$ for Mg-Ca-Gd alloy. Polarization resistance was calculated as 238.139 Ω/cm² on Mg-40Ca alloy and 689.599 Ω/cm² for Mg-Ca-Gd. The corrosion rate was extrapolated as 43.908 mm year⁻¹ from Mg-40Ca sample and 13.977 mm year⁻¹ from Mg-Ca-Gd.

Figure 2: Visual comparison of samples Mg-40Ca and Mg-Ca-Gd at 24, 48 and 72 hours of immersion in NaCl at 36°C.

Figure 2 shows the samples of the two alloys after immersion during in physiological solution at 36°C for 24 hours, 48 hours and 72 hours. In Figure 3 it is possible to compare graphically the evolution of mass loss over time. With the data obtained in the mass loss tests during immersion in physiological solution at 36°C, the corrosion rates were calculated (Table 3) and the hydrogen evolution for each alloy was estimated. The measurement of the roughness of the alloys was made by means of six different measurements of the same two parameters (Ra and PV) in three different regions of the sample. Table 4 shows the general averages and the standard deviation of these results for each parameter, all in micrometers.
Table 3: Corrosion rates (mm/y) for the MgCaGd and MgCa alloys calculated from the mass loss results and their respective hydrogen evolution values (mg/ml/d).

| Alloy    | Corrosion rate (mm/y) | Hydrogen evolution rate (ml/cm²/d) |
|----------|------------------------|-----------------------------------|
| MgCaGd   | 153.11                 | 69.41                             |
| Mg40Ca   | 284.49                 | 118.99                            |

Discussion

Comparing curves of Mg-40Ca and Mg-Ca-Gd alloys (Figure 1), differences in slopes curves can be identified and reflect straightly in Tafel constants. Despite the initial overlap of cathodic curves from alloys with RE content, as potential is raised by potentiostat, curves assumed different slopes being slower the slope of Mg-40Ca sample than Mg-5Ca-5Gd. A more vertical anodic curve was identified on Mg-Ca alloy with Gd content which indicates that passivation film is easily formed; being also possible that passive film might be more stable than observed into Mg-40Ca during anodic reaction. The alloy with Gd addiction showed a lower value of current density and a nobler corrosion potential when compared to Mg-40Ca. Similarities into OCP value from alloys with RE content brought too little in indication of differences in to chemical stability of both samples, although the sample without RE addiction and higher Ca content demonstrated the least chemical stability with the lowest OCP value (-1.760V). The influence of calcium addiction in the alloys corrosion behavior was previously discussed [5,10]. This information is in line with our results, where besides the influence of Gd, the least corrosion rate was observed in Mg-Ca-Gd alloy. From our perspective, the higher calcium content of binary Mg-Ca alloy should be the major reason of the higher corrosion susceptibility of this alloy when compared to the others ternary ones. Most of this susceptibility might come from the higher volume of Mg₂Ca phase normally expected in binary alloys with higher calcium content.

Another two parameters showed in Table 2 are the corrosion rate (Rᵣ) and the polarization resistance (Rₚ). Both factors need to be calculated from different equations and are referred to the rate of alloy corrosion degradation, as a consequence of current density. Polarization resistance is referred to the material refusal in allowing current passage. Rᵣ and Rₚ were calculated from Tafel equation, where after rearrangements, Rᵣ was related directly to the corrosion current through the equation:

\[ Rᵣ = \frac{0.13 I_{corr} m}{d} \]

where \( I_{corr} \) = corrosion current density; \( m \)= equivalent weight of corroding species [11] and \( d \)=density of samples.

Polarization resistance (Rₚ) was calculated as an observance of the anodic and cathodic reactions from Tafel equations. The Tafel equation was rearranged in terms of Butler-Volmer equation for consideration of mutually influence of anodic and cathodic reactions, and then simplified to get the Stern-Geary equation [12]. The rearrangement was presented below. \( β_a \) is the anodic \( β \) Tafel constant and \( β_c \) is the cathodic one.
The calculated average corrosion rate related to mass loss were conducted according to technical standard ASTM G31:

\[
\text{Corrosion rate} = \frac{(K \times W)}{(A \times t \times d)}
\]

where \( K \) = a constant; \( W \) = mass loss in g; \( A \) = area in cm²; \( t \) = time of exposure in hours and \( d \) = sample corrosion density.

The results of Mg-5Ca-5Gd alloy presented lower values of corrosion rates than Mg-40Ca. This finding is coherent with the discussion above related to calcium content and the positive effects of RE elements effects into polarization resistance, since Mg-Ca-Gd showed higher polarization resistance (in all three cases) than Mg-40Ca. Previous works showed [13-15] that it is possible relate the hydrogen evolution rate (\( V_H \)) with mass loss (\( \Delta W \)) using:

\[
V_H = \frac{\Delta W}{1.085}
\]

Where the results were coherent since higher is the mass loss, higher will be the hydrogen evolution.

Analyzing Table 4 it was found that the Ra mean values were close, for the two alloys. However, the standard deviation of the Mg-Ca alloy (Ra and PV) was higher indicating that there was considerable variation in the measured points. It means that the corrosion on the surface was more aggressive. This can be proved by the tridimensional images (Figure 4).

Table 4: The general averages and the standard deviation of the measurement of the roughness of the alloys.

| Alloy    | Ra (µm) Before | Ra (µm) After | PV (µm) Before | PV (µm) After |
|----------|---------------|--------------|---------------|--------------|
| MgCaGd   | 0.254±0.008   | 1.562±0.094  | 2.353±0.136   | 7.933±0.548  |
| Mg40Ca   | 0.397±0.036   | 1.542±1.146  | 4.083±0.503   | 18.140±8.180 |

Figure 4: Tridimensional representation of surface morphology after corrosion of Mg40Ca alloy (left) and Mg-Ca-Gd (right).

Conclusion

a) Magnesium based alloys with Ca and Gd has higher corrosion resistance than the binary alloy Mg-40Ca alloy;

b) The corrosion rate was adjusted with adequate RE elements and Ca content;

c) The most adequate results were achieved in ternary alloys with Gd and limited Ca addition.

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