Comparative study of fluoride conversion coatings formed on biodegradable powder metallurgy Mg: The effect of chlorides at physiological level

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Abstract

The development of a biodegradable metallic implant demands a precisely defined degradation profile and adequate mechanical properties. Mg has been proposed for this purpose but it has an excessively high corrosion rate and insufficient yield strength. In the present work pure Mg mechanically reinforced by a powder metallurgy (Mg(PM)) route and treated with KF was used. The effect of chlorides, at the physiological level, on four fluoride conversion coatings (F-CC) formed on Mg(PM) was evaluated comparatively. The behavior of Mg(PM) during fluoride treatments (0.01 M-0.3 M fluoride-containing solutions) before and after the addition of chlorides (8 g L⁻¹ NaCl) was investigated by conventional corrosion techniques and by scanning electrochemical microscopy (SECM) complemented with SEM observations and EDX analysis. Results showed that the composition and the microstructural characteristics of the F-CCs as well as their corrosion behavior change with KF concentration and immersion time. Treatments in the 0.01 M-0.1M KF range prove to be effective to protect Mg(PM) against corrosion in the absence of chlorides while higher KF solution concentration (0.3 M) adversely affects the corrosion resistance of this metal. In the presence of chloride ions the F-CCs progressively lose their fluoride content and their corrosion resistance at a rate that depends on the treatment conditions. Such temporary corrosion protection is appropriate for biodegradable implants.

1. Introduction

Magnesium and its alloys are considered the lightest structural materials with potential applications in different areas such as automotive, aircraft, electronics and aerospace [1–4]. In the last few years they have also been proposed as promissory biodegradable biomaterials. These biomaterials have several advantages in relation to fixed implants, since they can avoid some of the harmful effects of non-degradable implants and prevent the second surgery necessary to remove the

traditional implants. Additionally, their mechanical properties are closer to those of bone and they are able to stimulate growth of new bone tissue and reduce stress shielding [5–9]. Their degradation properties are also useful in the case of cardiovascular implants [10,11], particularly for preterm babies [12]. From the biological point of view, the use of pure Mg is more advisable than Mg alloys because it prevents the release of toxic ions.

However, there are negative issues that should be solved before using Mg-based materials. Pure magnesium typically has a yield strength of 55 MPa, lower than natural bone (e.g. femur \approx 110 MPa) and shows excessively high degradation rates in body fluids [13–20]. Powder metallurgical (PM) route was used to prepare Mg/HA biocomposite with cytotoxicity and corrosion resistance dependent on the HA content [21,22]. PM route was also used to upgrade the mechanical properties of pure Mg [23–25]. Mg(PM) benefits from a large grain size refinement in comparison with cast magnesium, and the development of a considerable crystallographic texture and, additionally, the presence of the dispersion of magnesium oxide particles produces mechanical reinforcement. As a result of these microstructural improvements the material obtained in the present work has a 2% ductility, offset stress (at 0.2% strain) of 280 MPa and a tensile stress of 320 MPa close to values obtained in the widely used AZ31 magnesium alloy [24].

To improve the corrosion resistance of Mg(PM), among different inhibition treatments proposed [26,27], the treatment with KF solution seems to be promissory since it is low-cost, non-toxic simple and easily handled for biomedical applications. Besides, fluoride-treated biomaterials have shown lower cytotoxicity level and better biocompatibility than controls [28,29]. The effect of fluoride treatments on Mg alloys has been recently described [30]. The authors showed that the behavior of AZ91D Mg alloy is very complex when chlorides are also present in the aqueous solution. Since Mg is potentially useful as degradable implant and considering that chloride is very abundant in body fluids, this condition deserves a particular investigation in the case of Mg(PM).

The techniques so far used to study corrosion problems (potentiostatic and potentiodynamic polarization curves, transient currents, electrochemical impedance spectroscopy, etc.) give valuable surface-average information on the system.

On the other hand, scanning electrochemical microscopy (SECM) is a powerful technique which can be applied to monitor both insulating and conducting surfaces [31]. This microelectrochemical technique consists in the movement of an ultramicroelectrode tip (UME) parallel to the surface of the sample inside an electrochemical cell. The UME can be positioned close to the interface with submicron precision and can probe the topography, reactivity or permeability of that interface with high spatial resolution [32,33].

Due to the fact that some electrochemical mediators employed in SECM can interfere with the electrochemical reactions that take place in the corrosion process, it is necessary to use as electrochemical mediators species already present in the system under study. For example, the ionic species resulting from localized corrosion processes can be detected at the tip [34] and the local variations in the extent of metal corrosion can be measured [35]. It is also possible to use oxygen dissolved in the electrolyte as mediator, since it is an electroactive molecule. Thus, the need for the addition of a redox mediator is avoided [35,36].

This technique has been applied to several corrosion problems, such as pitting corrosion [37,38], mapping of the surface conductivity of aluminum alloys [39], imaging of cathodic protection of aluminum [40], and recently to evaluate a Mg surface treated with alternating voltage [41]. In the present work we report a novel use of this electrochemical technique that involves monitoring changes in surface reactivity over time during passivating treatments.

The aim of this work was to evaluate the protective characteristics of different fluoride conversion coatings (F-CC) formed on Mg(PM) surfaces when they are in contact with chloride ions at the physiological levels in an attempt to contribute to the elucidation of the chloride-assisted F-CC degradation process without the interference of other ions. With this purpose the

electrochemical behavior of Mg(PM) during fluoride treatments before and after the addition of chloride ions was investigated. Conventional corrosion techniques (open circuit potential measurements (E_{ocp}), transient currents, and electrochemical impedance spectroscopy (EIS)) together with SECM were used.

2. Materials and methods

2.1. Materials

Magnesium powder (99.8%, 325 mesh) was supplied by Alfa Aesar. The powders were coldpressed and the resulting cylindrical compacts of 40 mm in diameter were extruded in the form of bars at 420 °C, employing an extrusion ratio of 16:1 [24,25].

Cylindrical samples (1 cm diameter) were cut from Mg(PM) bars. The lateral surfaces were covered by epoxy resin leaving a circular exposed area of 0.785 cm^2 . The samples were ground and dry polished with SiC papers of successively finer grit down to 1500 grit. Once polished, the samples were rinsed in ethanol and dried with nitrogen.

The sample surface appearance was carefully examined by optical microscopy before and after the experiments so as to check color and texture uniformity of the conversion coating. SEM observations were also made (EI Quanta 200 in high vacuum mode). The equipment used for the characterization of magnesium samples was a Philips XL30 Scanning Electron Microscope, coupled to the EDX DE4i analyser. The phases present in the coating were determined by X-ray diffractometry (X'Pert PRO XRD, Panalytical).

2.2. Reagents, solutions and pretreatments of the samples for electrochemical measurements

All the solutions were prepared with ultrapure water by means of a Millipore Milli-Q system (18.2 $M\Omega^{-1}$). The compositions of the solutions employed in the experiments were: 0.01 M KF, 0.03 M KF, 0.1MKF, 0.3M KF and 8gL⁻¹ NaCl. Analytical grade reagents were used in all cases.

Prior to some of the electrochemical experiments the Mg(PM) electrodes were immersed in 8 g L^{-1} NaCl solutions for 1 h to induce surface attack (Mg(PM)SA).

Electrochemical measurements were performed after different treatments and under different experimental conditions in order to investigate the corrosion resistance of the Mg(PM) and Mg(PM)SA treated with KF solutions in the absence and in the presence of chloride ions: a) transient currents and SECM were recorded with Mg (PM)SA during the passivating treatments with KF solutions, b) EIS experiments were performed in 8 g L⁻¹ NaCl solution with Mg(PM) passivated with 0.01 M, 0.03 M, 0.1 M and 0.3 M KF solutions; and c) transient currents and E_{ocp} with Mg(PM)SA were recorded in 0.1 M KF solution to which NaCl was added after different periods up to physiological level (8 g L⁻¹).

2.3. SECM characterization

The surface topography was studied before and during fluoride conversion coating formation (F-CC) using a SECM (Sensolytics) equipped with a 10 μ m platinum tip (ultramicroelectrode, UME) as probe, Ag/AgCl/KCl reference electrode, and a platinum wire as counter electrode, all placed within a polytetrafluoroethene cell. Mg (PM)SA specimens were mounted horizontally facing upwards. Open circuit potential (E_{ocp}) experiments were performed in 0.1 M KF solution as background electrolyte. The scan rate of the microscope tip over the specimen surface was 10 μ ms⁻¹ in unidirectional lines, with 5 μ m separation between line scans. The maps were generated with the tip located c.a. 10 μ m from the Mg(PM)SA surface.

In order to study the passivating action of fluoride ions on Mg(PM) samples they were previously corroded in 8 g L⁻¹ NaCl for 1 h (Mg (PM)SA) before dipping into the SECM electrolytic cell with 0.1 M KF aerated solution (treatment (a)). The chemical activity was measured in a feedback mode using O2 dissolved in solution as electrochemical mediator. The oxygen related current was recorded directly by the SECM-tip set at -0.70 V. At this potential, O2 reduction occurs under diffusion control and the current magnitude at the microelectrode is dependent on the O2 concentration.

The UME was scanned over a 2000 μ m×600 μ m area of the sample. The localized differences in the concentration of oxygen lead to variations in the faradic current which passes through the tip.

2.4. Transient currents and E_{ocp} measurements

An electrochemical cell with a three-electrode setup was used. A platinum wire (99.99% purity) served as auxiliary electrode. All potentials are quoted with respect to the Ag/AgCl/KCl reference electrode. The exposed area of the working electrode (Mg(PM)SA disk) was 0.785 cm² for all the experiments.

A TEQ-02 potentiostat was used for electrochemical measurements. Transient currents were recorded using Mg(PM)SA as working electrode at fixed potentials (E_{ocp} +50 mV, vs. Ag/AgCl/KCl) in 0.01 M, 0.03 M, 0.1 M and 0.3 M KF solutions (treatment (a)).

 E_{ocp} and transient currents recorded at Eocp +50 mV,vs.Ag/AgCl/KCl with Mg(PM)SA in 0.1 M KF with NaCl additions up to physiological level, (8 g L⁻¹) (treatment (c)) after different times (0.5 min and 1 min) were also measured to analyze the effect of chloride ions on the protective characteristics of the film.

2.5. Electrochemical impedance spectroscopy measurements

The EIS were performed with an Autolab PGSTAT 30 potentiostat from Eco-Chemie. The Mg(PM) samples were pre-treated in aereated 0.01 M, 0.03 M, 0.1 M KF and 0.3 M KF solutions for 1 h. A sinusoidal potential modulation of ± 10 mV amplitude in the 10⁵ Hz–10⁻² Hz frequency range, spaced logarithmically (120 per 8 decades), was superimposed on the E_{ocp} of F⁻-coating Mg(PM) in 8 g L⁻¹ NaCl.

A control assay for the electrodes without pre-treatment could not be included because of the high scattering of the results due to the unstable behavior of the Mg(PM)SA bare surface in NaCl solutions [42].

3. Results

3.1. F-CC formation on corroded surfaces

3.1.1. Transient currents

The change in the average activity of the electrode surface during KF treatments was followed by transient current measurements and was used to identify the more appropriate protective treatments. Prior to each electrochemical measurement, and in order to enhance the detection of the protective effect of F⁻ ions on active surfaces, the Mg (PM) sample was immersed into 8 g L⁻¹ NaCl for 1 h. The microscopic observation of Mg(PM) with surface attack (Mg(PM)SA) (Fig. 1a and b) shows regions with and without localized attack and others covered by corrosion products. After this pretreatment the samples were immersed in KF of different concentrations to identify the best passivating process. Fig. 2 shows that the transient currents recorded at 50 mV were more anodic than the instantaneous E_{ocp}. Initially, the best performance was depicted in 0.1 M KF solutions,

which showed the lower and more stable current values. In solutions with lower F⁻ concentrations (0.01 M and 0.03 M KF) the initial current values were high and very unstable. Interestingly, these current values decreased sharply and reached cathodic values after some seconds. Although 0.3 M KF assays showed low current values initially, these values increased later, revealing the adverse effect of high KF concentrations.

3.1.2. SECM characterization

The local current, related to the in situ conductivity was mapped by the SECM revealing the hillor valley-like current fluxion in the vertical Z direction. Fig. 3 shows the surface scanning before and during the formation of the conversion coating in 0.1 M KF solution. The effect of F⁻ during the immersion of the corroded sample in a 0.1 M KF solution followed by this electrochemical imaging technique allows in situ mapping of the chemical reactivity of surfaces and visualization of spatially-confined variations in conductivity. When the tip moves over the areas covered by a hydroxide film, the tip current decreases, because the diffusion of O2 is physically hindering towards the probe (negative feedback, i.e. negative current decreases). However, when it moves over the active areas (localized corrosion micropits), the tip-sample distance increases, decreasing the O2 concentration that reaches the tip, causing the decrease of the reduction current. The successive curves plotted in the X–Z plane represent the map of the active areas at each X position while those of Y–Z plane represent the change in the reactivity with time. Due the protective effect of the background electrolyte (0.1 M KF) which gradually covers the active areas, lower variation in the current values was progressively detected during the successive scans, giving rise to c.a. constant current values at the end of the experiment, associated with the flat image of a passive surface

Unfortunately, due to the formation of hydrogen bubbles, SECM measurements in the presence of chloride ions cannot be made.

3.2. F-CC formation on non-corroded surfaces

Fig. 4a–c shows that after 1 h exposure in solutions of 0.01, 0.03 and 0.1 M KF a compact film is formed with some needle shape crystals on the surface. EDX analysis showed that F-at.% increases with time (from 7% (1 h) to 20 at.% (22 h) in the case of 0.1 M KF).

A dramatic change is detected after 0.3 M KF treatments (Fig. 4d) due to the formation of small (c.a. size =100 nm wide) cubic crystals. Importantly, crevices can be observed on the surface.

With the aim of comparison, results of EDX analysis after 1 h immersion period in the 0.01–0.3 M KF concentration range are shown in Fig. 5. It can be noticed that the oxygen at.% decreases when the concentration of KF solution increases while the amount of fluoride increases. Interestingly, a significant increase of K at.% [from 0.05 (0.1 M KF) to 1.39 (0.3 M)] was detected. After a long exposure (7 days) of Mg(PM) to 0.1 M KF solutions the microstructural characteristics of the layer coating change and KMgF₃ was identified by XRD analysis (not shown).

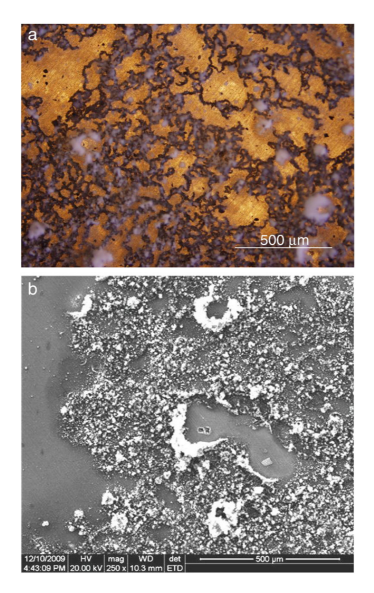


Fig. 1. (a) Optical microscope and (b) SEM images of the Mg(PM) after being immersed in 8 g L^{-1} NaCl for 1 h (Mg(PM)SA).

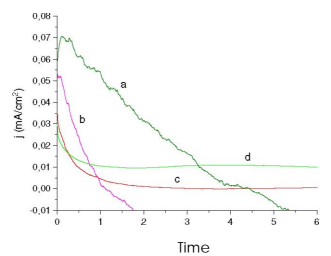
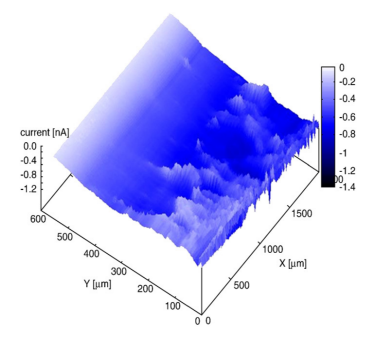
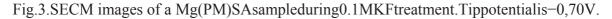


Fig. 2. Transient current corresponding to Mg(PM)SA recorded at 50 mV more anodic than the open circuit potential in different KF solutions: (a) 0.01 M, (b) 0.03 M, (c) 0.1 M and (d) 0.3 M.





3.3. The effect of Cl⁻-containing solutions on F-CC

3.3.1. Transient current and E_{ocp} measurements

To test the change of the passivating characteristics of F-CC by chloride ions (the most aggressive among the anions of biological fluids) transient currents were recorded at $E_{ocp}+50$ mV with Mg (PM)SA. Firstly, very thin F-CCs were formed in chloride-free solution during 0.5 and 1 min respectively and subsequently chloride ions were added to the F-containing medium. If the addition of chlorides is made after 1 min it can be noticed (Fig. 6) that the current initially decreases due to the passivating action of fluorides, then increases slowly and finally remains nearly constant. However, when the addition was made after 0.5 min the current increased sharply.

Similarly, E_{ocp} variation with time also showed different responses depending on the time at which the NaCl addition was made. Results obtained during 0.1 M KF treatments before and after the addition of NaCl up to physiological level (8 g L⁻¹) are shown in Fig. 7. Initially, the E_{ocp} value increased sharply up to c.a. -1.17 V and then remained nearly constant without change when the addition of NaCl was made after 1 min. Conversely, E_{ocp} decreased abruptly and very unstable values were recorded when the addition was made 0.5 min after the beginning of the experiment.

3.3.2. Electrochemical impedance spectroscopy (EIS)

EIS experiments were performed with Mg(PM) electrodes previously treated in KF solutions (0.01–03 M concentration range) to make a comparative analysis of the protective characteristic of the F-CCs. EIS was performed under E_{ocp} condition in 8 g L⁻¹ NaCl. The impedance results are shown in both the Nyquist and the Bode plots in Fig. 8. The Nyquist diagrams (Fig. 8a, imaginary part Z'' vs. real part Z') show a semicircle, corresponding to the charge transfer resistance limiting process related to F-CC/electrolyte interface (R_{CT}). Its intersection with the real axis, at high frequencies, corresponds to electrolyte resistance (R_e). At low frequency an inductive behavior also appears.

The capacitance was calculated from experimental data as C=1/(2 $\pi f_i R_{TC}$). The frequency f_i was

selected to obtain a maximum value of Z_i , where Z_i is the imaginary part of the impedance (Table 1).

The impedance modulus Bode diagrams (Fig. 8b and c) revealed a plateau at high frequencies. The impedance value at this region, with $\varphi = 0$ (Fig. 8c), gives the electrolyte resistance in each case. At decreasing frequencies (10³ Hz–10 Hz) the slope can be attributed to one capacitor in parallel with a resistance corresponding to the F-CC/ electrolyte interface with a phase angle whose value is shown in Table 1. In the lower frequency range, the phase angle is close to 0°, corresponding to a plateau in the impedance modulus vs. frequency plot. Thus, the total resistance of the circuit (R_e+R_{CT}) can be obtained. The experimental results are shown in Table 1.

3.3.3. EDX analysis and SEM of the F-CCs after immersion in NaCl solutions

Fig. 5 shows that the F-content of the F-CC drastically decreases during the immersion in NaCl solution (compare F atomic percent before and after the immersion in the chloride solution ((F) and (F (Cl sol)) respectively)). Additionally an important increase in the oxygen content can also be observed (Table 2).

The F-CCs are more vulnerable to chloride attack when the fluorides are absent in the solution (i.e. biological solutions). After 1 h exposure periods in KF followed by immersion in a fluoride-free NaCl solution for 1 h, the Mg(PM) surface showed areas with corrosion products and localized attack (Fig. 9a) and other regions that remain protected (Fig. 9b). Fe and Si impurities detected by EDX seem to favor localized attack.

4. Discussion

The protective characteristics of F-CCs formed on Mg(PM) with KF solutions in the 0.01–03 M concentration range were evaluated comparatively. Mg(PM) samples with F-CCs were subsequently immersedina8 g L^{-1} NaCl solution to elucidate the effect of chloride on the degradation of F-CCs without the interference of other ions. Useful information to select an appropriate treatment to provide temporary protection to Mg(PM) in biological media could be obtained from the results.

Several corrosion studies of magnesium and magnesium alloy dissolution have been performed for industrial and medical applications. The overall corrosion mechanism accepted for neutral and alkaline media is [1–3]:

$$Mg + 2H_2O \leftrightarrow Mg(OH)_2 + H_2$$

(1)

With the following partial reactions:

$$Mg \leftrightarrow Mg^{2+} + 2e^{-}$$
$$2H_2O + 2e^{-} \leftrightarrow H_2 + 2OH^{-}$$

or

 $2H^+ + 2e^- \leftrightarrow H_2$.

(3b)

(3a)

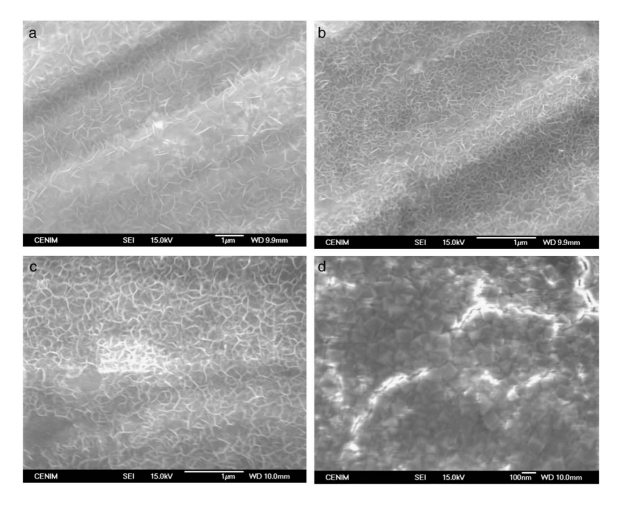


Fig. 4. SEM microphotographs of Mg(PM) after 1 h immersion in (a) 0.01 M; (b) 0.03 M; (c) 0.1 M and (d) 0.3 M KF. Cubic crystals and crevices can be distinguished in (d).

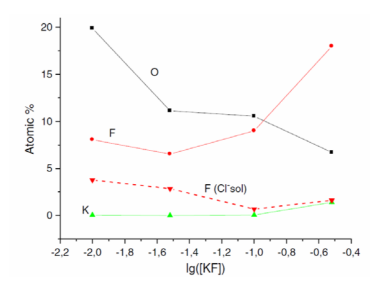


Fig. 5. Atomic percent of the elements (O, F, and K) obtained by EDX analysis of F-CC formed with different KF concentrations (0.01 M, 0.03 M, 0.1 M, and 0.3 M) (—). F(Cl⁻ sol) corresponds to the atomic percent of F after the F-CC-Mg(PM) immersion in 8 g L⁻¹ NaCl (- -).

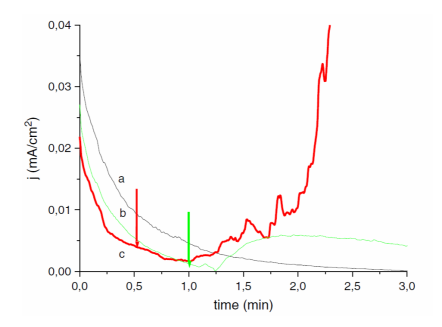


Fig. 6. Transient currents corresponding to Mg(PM)SA, recorded at 50 mV more anodic than the open circuit potential in 0.1 M KF. (a) Control, (b) NaCl up to 8 g L^{-1} is added after 1 min, and (c) after 0.5 min. The addition of NaCl is indicated with the arrows.

Alternately, a one electron step followed by a chemical step was also proposed,

$$\begin{split} Mg &\leftrightarrow Mg^{1+} + 1e^{-} \\ 2Mg^{+} + 2H_2O &\leftrightarrow 2Mg^{2+} + 2OH^{-} + H_2 (\text{chemical reaction}). \end{split}$$

Followed by

 $Mg^{2+} + 2OH^{-} \leftrightarrow Mg(OH)_2(s).$

The overall reaction consumes H^+ or produces OH^- (Eqs. (3a) and (3b)), and consequently the pH increases and favors the formation of Mg(OH)₂.

As Mg(OH)₂ is a poor electron conductor this suggests that the H₂ reaction may be occurring mainly in the pores at points at which are the shortest distance to the metal [43]. In the case of Mg(PM) the microstructure includes MgO particles that reinforce mechanical properties. These particles may also affect the corrosion behavior of Mg(PM) since electrochemical measurements show more anodic E_{ocp} and lower current densities for pure Mg with MgO coating than for bare Mg [44]. SECM proved to be a sensitive localized technique able to evaluate the changes of the chemical activity of the Mg(PM)SA surface with time. It was revealed that the activity of spatially-confined sites can be sensed and that this reactivity decreases when the immersion time in KF solution increases. SECM results showed that the Mg(PM)SA surface, initially very active and showing scattering current values, is progressively passivated in the 0.1 M KF solution, reaching low and constant current values that indicate the passive state of the sample. It seems that in

aqueous fluoride-containing solutions the $Mg(OH)_2$ film (slightly stable in water) reacts with fluorides and forms F⁻-containing Mg compounds according to [45]:

$$Mg(OH)_2 + xF^- \rightarrow Mg(OH)_{2-x}F_x + xOH^-.$$

If
$$x = 2$$

$$Mg(OH)_2 + 2F^- \rightarrow MgF_2 + 2OH^-$$
.

Additionally in the presence of K cations, and depending on the fluoride concentration, MgKF₃ could also be obtained according to:

$$Mg(OH)_2 + K^+ + 3F^- \rightarrow MgKF_3 + 2OH^-$$

or

 $MgF_2 + KF \rightarrow MgKF_3$

The decrease of oxygen at.% at the surface and the increase of fluor at.% when the concentration of the KF solution increases as well as the increase in K content of the layer (Fig. 5) and the identification of KMgF3 for long immersion times are in agreement with this sequence of reactions.

According to SEM observations (Fig. 4d) F-CC formed with 0.3 M KF revealed crevices where corrosion seems to be facilitated. These results are in agreement with those reported for AZ91D alloys [30].

The protective F-CCs can also be formed on corroded samples (Mg (PM)SA). Transient currents revealed that in the case of 0.3 M KF solution a less protective layer is formed.

The impedance diagrams of Mg(PM), in 8 g L⁻¹ NaCl solution, after 1 h immersion pre-treatment in 0.03 M, 0.01 M, 0.1 M and 0.3 M KF solutions, exhibit one capacitive loop. For simplicity one might take the real impedance at which the imaginary part vanishes for the capacitive part to be the charge transfer resistance RTC and relate it to the corrosion resistance. Therefore, the corrosion resistance RCT decreases from 1032.29 Ω cm² to 144.18 Ω cm² (close to that obtained in the absence of F-treatment [26]) when the concentration of the KF solution increases from 0.01 M to 0.3 M. This behavior seems to reflect that, in the presence of Cl⁻, the charge transfer resistance, associated with the inhibitory effect of F⁻, decreases when the concentration of the KF in the pretreatment increases.

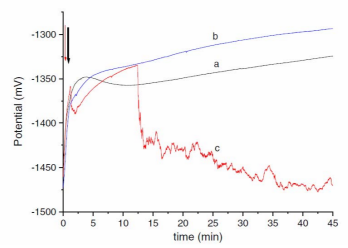


Fig. 7. E_{ocp} vs. time plot corresponding to Mg(PM)SA in 0.1 M KF. (a) Control, (b) NaCl up to 8 g L^{-1} is added after 1 min, and (c) after 0.5 min. The addition of NaCl is indicated with the arrows.

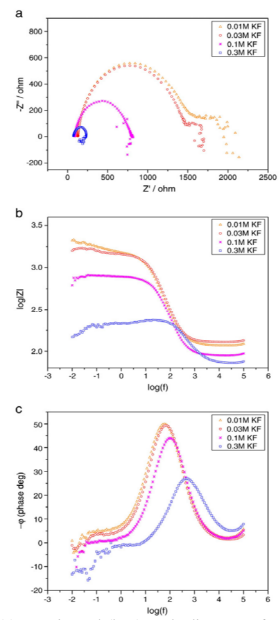


Fig. 8. EIS spectra of the (a) Nyquist and (b, c) Bode diagrams of Mg(PM) immersed in 8 g L^{-1} NaCl after the following treatments for 1 h: 0.01 M, 0.03 M, 0.1 M and 0.3 M KF.

At low frequencies, an inductive behavior appears, more defined in the case of 0.01 M and 0.03 M solutions, usually associated to adsorption–desorption of ions in a corrosion process. Our results and those of Chiu [24,26] revealed that the composition and structural characteristics of the F-CC affect the degradation profile of this layer. SEM and EDX analysis revealed (Figs. 4 and 5) that the microstructural characteristics and composition of the F-CC change with the concentration of the KF solutions used. Crystallographic KMgF₃ or more amorphousMgF₂, of different protective characteristics, can be obtained by varying the conditions of the pre-treatment [24].

Transient currents and E_{ocp} measurements (Figs. 6 and 7) also revealed interesting results in relation to the competitive action of chloride ions. These results indicate that the F-CC progressively achieves its protective characteristics in F⁻-containing solutions both in corroded and

non corroded samples, in the absence of chloride ions. However, after the addition of chloride ions at the physiological level they show a very aggressive behavior competing with the fluoride ions of the solution and of the coating for the adsorption sites. Thus, the very labile F-CC formed after 0.5 min in 0.1MKF solution can be easily attacked by chloride ions. After 1 min of immersion in the KF solution a more resistive behavior was shown, revealing the importance of the immersion time of fluoride treatments. Importantly, in fluoride-free NaCl solutions (like biological fluids) the F-content decreases drastically while oxygen-content increases. Consequently, the protective characteristics are progressively lost (Table 2) and localized attack can be observed after the 1 h immersion period (Fig. 9a)

The complex behavior of F-CC in chloride solutions could be explained considering: (a) the local change of pH and the high electron affinity of fluoride in the absence of chlorides and (b) the competition for the adsorption sites in the presence of chlorides [30,43,45-47]. Thus, there must be two successive processes occurring at the anodic interface. Firstly, the hydrolysis of the metal ions coming from the corrosion reaction and then the formation of $H_i F_{i+1}$ adsorbates due to the presence of F⁻ (highly electronegative ions with strong binding energy with hydrogen ions) that protects the Mg(PM) surface. When chloride is present in the electrolyte media both, chloride and fluoride of the F-CC compete for the adsorption sites. Results showed that chloride ions induce the dissolution of F-containing products, since the F-content of F-CC drastically decreases after the immersion in chloride solutions (Table 2, Fig. 5). Thus, the surface layer becomes more labile to the corrosion attack and oxygen-containing corrosion products are formed on the surface according to the increase of the O atomic percent of the layer detected by EDX (Table 2). In agreement, results also showed that chloride aggressive action is more effective when a thin, weak F-CC is formed, i.e., when a treatment with a short immersion period in the KF solution is applied (Fig. 6). Consequently, the comparative analysis of the data reveals that the composition, the structural characteristics and the corrosion resistance of the F-CC depend on the fluoride concentration and the immersion time of the solution. After the immersion in chloride solutions and in the absence of fluorides (like biological fluids) the corrosion resistance decreases due to the reduction of the Fcontent of the F-CC layer. This temporary protection of F-CC is appropriated for their application as biodegradable implants.

Accordingly, Mg(PM) with F-CC shows low corrosion rates initially and could originally preserve their mechanical integrity while the bone is healing. Subsequently a chloride assisted corrosion process could complete the degradation of the biomaterial. Importantly, it should be emphasized that an advantage of Mg(PM) with respect to Mg alloys is that its dissolution is not accompanied by the release of other less biocompatible metals.

Overall, results showed that SECM is a sensitive technique which enabled the evaluation of the progressive passivating action of fluoride on Mg(PM)SA in the absence of chloride ions. These results are in concordance with transient current measurements that revealed that the current decreases with time (except for 0.3 M KF treatment where crevices are formed). On the other hand, the variable protective behavior of F-CC in the presence of chloride ions at physiological level could be followed by EIS, E_{ocp} and transient current records. It was demonstrated that the F-CC progressively loses its corrosion resistance. The dissolution rate depends both on the immersion period and on the concentration of KF solution of the pre-treatment. High KF concentrations (0.3 M) or short immersion times adversely affect the corrosion resistance of the layer in chloride solutions. Thus, the degradation profile of F-CC on Mg(PM) could be controlled to satisfy the implant requirements by the adequate selection of the concentration of the KF solution and of the immersion period of the inhibition treatment.

KF	$E_{oc}(V)$	$R_e(\Omega)$	$R_{CT} (\Omega cm^2)$	$C(F cm^{-2})$	b	φ
0.3 M	-1.620	72.55	144.18	4.8910^{-6}	-0.33	-27.03
0.1 M	-1.550	89.51	586.86	6.9910^{-6}	-0.52	-41.41
0.03 M	-1.570	128.53	1018.96	9.0810^{-6}	-0.63	-49.31
0.01 M	-1.560	116.69	1032.29	10.2610^{-6}	-0.64	-49.83

Table 1 Experimental parameters obtained from EIS plots of Mg(PM) in 8 g L^{-1} NaCl with pretreatments in KF solutions of different concentrations.

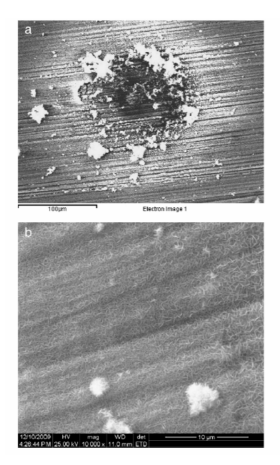


Fig. 9. SEM image of a Mg(PM) sample after immersion in 0.1 M KF solution for 1 h and then in a fluoride-free 8 g L^{-1} NaCl solution for 1 h. (a) Localized attack and corrosion products can be observed. (b) Region without attack

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