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Electropolishing Fe-based biodegradable metals for vascular applications: impact on surface properties, corrosion and cell viability

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Biodegradable metals constitute a new class of materials for medical application. By breaking the paradigm that a metallic biomaterial to be implanted in the body must be corrosion resistant, biodegradable metals advance surgery allowing clinicians to dispose of temporary devices. Among them, Fe–Mn–C steel has emerged due to its outstanding mechanical properties, while its degradation rate must be carefully controlled. For this purpose, especially for medical devices, the surface finishing plays a pivotal role and influences both the corrosion behavior and biological response of these materials. Therefore, this research investigated the impact of electropolishing (EP) processes on the Fe–Mn–C alloy surface finishing in terms of composition, morphology, topography, and wettability. Three electrolytes were carefully selected and used in this study: EP1 (ethanol, perchloric acid, and glycerol), EP2 (perchloric acid, acetic acid, and glycerol), and an ionic liquid EP3 (choline chloride and ethylene glycol). Corrosion behavior and cell viability were investigated and compared with those obtained on mechanically polished (MP) samples. The results displayed that electropolishing was governed by two mechanisms: 1) controlled mass transport for EP1 and EP2, and 2) an adsorption mechanism for EP3. Among the tested conditions, EP2 emerged as a promising overall EP process. It promoted the smoothest and most hydrophilic passivated surface ($R_a \sim 10$ nm, WCA = 53°, respectively) and the highest ratio of metal oxides to metallic elements. In addition, EP2 exhibited appropriate corrosion behavior suitable for biodegradable metal devices by inducing the formation of a protective oxide layer. Furthermore, cell viability with EP2 was comparable to that observed with MP. These findings emphasize the potential of electropolishing for enhancing the properties of Fe–Mn–C alloys, paving their applicability, especially in cardiovascular devices.

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1. Introduction

Biodegradable metals represent a significant advance in cardiovascular applications,¹ for example for temporary stents, where complete degradation is expected to occur between 6 and 24 months.² Despite several candidates such as Fe, Mg, and Zn alloys, only one Mg-based stent is currently available on the market.³ However, its low mechanical properties impose a larger structure design, much larger than the gold standard conventional stents made of Co–Cr alloys, or AISI 316 L.⁴ Fe–Mn–C has therefore emerged as a promising solution due to its exceptional mechanical properties, notably its high ductility (strain at break, $A \approx 20\%$) and strength (yield strength, $\sigma_Y \approx 620$

MPa), much closer than those of Co–Cr alloys, thus enabling thin stents, thereby reducing the risks of complications.^{5,6} Furthermore, *in vitro* assays with endothelial cells have exhibited satisfactory cytocompatibility levels, depending on the Mn content within Fe–Mn–C substrates.^{7,8} Metallic bare metal devices, if used as stents for vascular applications, must have a very smooth surface finish, in the range of a few nanometers.^{9,10} While this requirement is easily addressed with an electropolishing step in Cr–Co or SS316L corrosion-resistant metals, it represents a major challenge for biodegradable metals, mainly because it has the potential to initiate corrosion, thus leading to degradation. Moreover, smoother surfaces are less prone to cellular hyperplasia and inflammation when in contact with the artery wall,¹¹ and hydrophilic surfaces reduce platelet adhesion, promoting early vascular healing, and minimize late tissue response.¹² Regarding corrosion behavior, for instance, increasing surface roughness induces a higher degradation rate.¹³

In the light of these results, researchers in academia and industry agree that there is an urgent need to develop an

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efficient surface finishing process for Fe–Mn–C alloys susceptible to be used in vascular applications. A stable engineered surface that compromises corrosion behavior and improves biological response in the early stages of alloy–tissue interactions is required. To achieve this goal, recent studies have focused on the development of new surface modification processes, such as thermo-mechanical treatments,^{14,15} acid pickling,^{16,17} anodic oxidation,¹⁸ sandblasting¹⁹ and laser texturing,²⁰ which have been explored to treat Fe-based and Fe–Mn–C-based alloys.

Mechanical polishing (MP) is a common step performed before preliminary surface finishing tests on metallic materials.^{21–25} This process uses abrasives to remove microscopic chips from metallic surfaces; the process is composed of several steps: the first ones foresee the use of coarse grain abrasives, whose size is progressively reduced until the final stages, which are executed with micron-size or less abrasives, buffing to achieve a mirror-like surface.²⁶ This considerably hinders its application on devices with a complex geometry and reduced thickness, such as those found in stents.¹⁶ In addition, residual stress after MP can smear the surface or even reduce the corrosion resistance of the substrate.^{27,28}

To overcome these drawbacks, electropolishing (EP) has been adopted as a standard process for surface finishing of corrosion-resistant metallic implants with complex shapes.²⁹ This technique is employed to promote material smoothening and to increase its corrosion resistance by tuning, respectively, the surface roughness and chemical composition.³⁰ EP is a surface finishing process wherein the anodic dissolution of the workpiece (anode) occurs within a suitable electrolyte in an electrochemical cell.^{31–33} Controlling the current density and voltage, the outermost material from the workpiece surface is removed.³⁴ Subsequently, mass transport phenomena promote surface passivation through the formation of a nanometric oxide layer on the metallic surface. This technique can be used for the precise control of surface features, finally imparting a bright appearance to the material.³⁵

Several EP protocols were developed for corrosion-resistant alloys for biomedical practice, for example stainless steel,^{36–41} Co–Cr,^{42,43} and nitinol,⁴⁴ generally based on strong acids, such as sulfuric and *ortho*-phosphoric acids as electrolytes.⁴⁵ However, as far as EP processes for biodegradable metals are concerned, to our knowledge, only a few have been described in the literature, but none for Fe–Mn–C alloys. In fact, electrolytes composed of C_2H_5OH , C_4H_9OH , $AlCl_3 \cdot 6H_2O$, $ZnCl_2$ and distilled water held at 21–23 °C for 90 s were used for an EP pure Zn stent, resulting in the formation of a chemically homogeneous and smooth surface.⁴⁶ Another EP process was applied to a commonly studied magnesium alloy, WE43, featuring an electrolytic mixture of HNO_3 and $C_2H_5NO_3$.⁴⁷ The EP process effectively smoothens the surface and maintains high cell viability, comparable to that of untreated surfaces.

In parallel, recent research has been focusing on the viability of employing organic acids or ionic liquids (ILs) as alternatives to water-based solutions.⁴⁸ IL solvents are a large

class of environmentally-conductive solvents which have the advantages of not being based on a water solution and of presenting high current efficiencies with negligible gas evolution at the anode/solution interface.⁴⁹ ILs were successfully applied to the EP of stainless steel,⁵⁰ pure silver,⁵¹ and aluminum- and nickel-based alloys.⁵² The most commonly used electrolyte, composed of choline chloride (ChCl) and ethylene glycol (EG), features a unique EP mechanism, distinct from traditional aqueous acid solutions – the ChCl:EG mixture shows slower and more potential-dependent oxide removal, shifting to a mass transport-limited process after the oxide layer is removed.⁵³ In all the cases, the use of this electrolyte results in bright surfaces and a noticeable reduction in surface roughness. Furthermore, while electropolishing with water-acid solutions lasts for 1 to 15 minutes at a temperature of 30 to 90 °C, IL solutions take 10 to 120 minutes, depending on their chemical composition.³⁶ Despite this longer duration, electropolishing with IL solutions is carried out at a lower temperature, ranging from room temperature to 50 °C.⁵⁴

Therefore, the aim of this research was to develop an electropolishing process as surface finishing for Fe–Mn–C biodegradable alloys in order to achieve effective passivation, homogeneous chemical composition and surface roughness. The effect on surface features of EP process parameters among three different solutions controlling the potential (V), current density ($A\ cm^{-2}$), time (min) and electrolyte bath temperature (°C) within the electrochemical cell was investigated. In addition, the relationship between these surface characteristics and their influence on corrosion behavior was evaluated by comparing electropolished and mechanically polished samples. Finally, the impact of the surface finish on vascular cells was assessed in order to validate the potential of these treatments for vascular applications.

2. Materials and methods

2.1. Sample preparation

Square samples (10 mm × 10 mm × 1 mm) of an Fe–13Mn–1.2C alloy (ISO GX120Mn13), commercially known as Hadfield steel, were supplied by Polstar Metals Inc. (Canada). The amount of Fe and Mn in both alloys was determined by atomic absorption spectroscopy (AAS). The carbon content was analyzed by employing a pyrolytic carbon/sulfur detector (LECO CS200, USA). The experimental composition of the alloy is reported in Table 1.

The sample surface was subjected to two surface finishings, that is, MP and EP, as detailed in Table 2. First, as a pre-treatment, all samples were mechanically polished using 80–240 grit SiC paper (AS-REC condition). The MP surface finishing was obtained by applying complete mechanical polishing until a mirror-like surface finishing was achieved for the pretreated samples. Following the MP treatment, a thorough cleaning step with an industrial non-acidic cleanser (DiverseyTM, Canada) and deionized water was performed to remove any accumulated impurities on the specimen surface. Finally, samples were



Table 1 Experimental composition of the Fe–13Mn–1.2C alloy

Element	C	Mn	Si	S	P	Fe
Composition (wt%)	0.99 ± 0.02	12.80 ± 0.50	<0.4	<0.02	<0.035	Bal.

Table 2 Details of the experimental procedures for mechanical polishing and electropolishing, along with the names of the samples

Treatment	Conditions	SiC abrasive paper (grit)	Cleaning after each stage	Diamond suspension (μm)		
1. Grinding (pre-treatment)	All samples	80–240	Sonication in acetone and methanol + final compressed air drying	–		
2. Mechanical polishing	MP	320–800	Sonication in acetone and methanol + final compressed air drying	6, 3, 1		
Treatment	Conditions	Electrolyte composition (* % vol/vol † % w/w)	Potential (V)	Current density (A cm ^{−2})		
3. Electropolishing	EP1	*70% ethanol 20% HClO ₄ 10% glycerol *94.8% CH ₃ COOH 4.7% HClO ₄ 0.5% glycerol †69% choline chloride *31% ethylene glycol	60–90	0.5	2.5	24 ± 0.5
	EP2		65–100	0.5	2.5	24 ± 0.5
	EP3		6	0.25	10	40 ± 0.6

immersed in an ultrasonic bath containing 10 mL of pure acetone for 5 minutes, then extracted and dried with filtered compressed air. Subsequently, they were sonicated in a similar way with methanol before drying with filtered compressed air. This exact cleaning step was also employed prior to EP treatment.

For the EP finishing, three electrolytes were selected: two acid solutions⁵⁵ and one ionic liquid solution described by Abbott *et al.*⁵⁴ The chemical composition of electrolytes and operating parameters are described in Table 2. All the acids (CH₃COOH, 99.5 vol% and HClO₄, 80 vol%) used in the present work were commercial ACS grade solutions (Lab Mat, Canada). A volume of 150 mL of freshly prepared solution was used for each test. Square samples with the same composition as the substrate alloy were used as the cathode. The distance between the electrodes was 6 cm. The current density in this electrochemical cell setup was controlled using a Sorenson DC power supply (Ametek, USA). After the electropolishing procedure, all the samples were sonicated in acetone and methanol, as described above, and then dried with a jet of filtered compressed air.

2.2. Surface characterization

2.2.1. Scanning electron microscopy. Surface morphology changes induced by EP procedures were assessed using an FEI Quanta 250 scanning electron microscope (SEM), equipped with a W filament, from Thermo-Fisher (USA). SEM images were acquired with an acceleration voltage of 20 kV in secondary electron mode.

2.2.2. Atomic force microscopy. The topography and roughness of surfaces were investigated using a Dimension

TM3100 atomic force microscope (AFM) from Veeco (USA) in tapping mode with an etched silicon tip (model NCHV, tip radius $R = 10$ nm, Bruker, USA). The surface roughness of five randomly chosen areas per condition was evaluated over 20 $\mu\text{m} \times 20 \mu\text{m}$ images and 2 $\mu\text{m} \times 2 \mu\text{m}$. The average roughness $\left(R_a = \frac{1}{MN} \sum_{j=1}^N \sum_{i=1}^M |z(x_i, y_j)| \right)$ and root mean roughness $\left(\text{RMS} = \sqrt{\frac{1}{MN} \sum_{j=1}^N \sum_{i=1}^M |z(x_i, y_j)|^2} \right)$ were calculated using WSxM software,⁵⁶ where R_a is the arithmetic average of profile height deviations from the mean line and RMS is the quadratic mean or root mean square average of profile height deviations from the mean line.⁵⁷ In addition, a first derivative was applied to the 20 $\mu\text{m} \times 20 \mu\text{m}$ images to enhance the contrast of features such as grain boundary steps.

2.2.3. Sessile drop contact angle. The wettability of the surfaces was determined by static water contact angle (WCA) measurements, according to ASTM-D7334,⁵⁸ using a camera system AST Products Inc. VCA 2500 XE (Billerica, USA). Six measurements per sample were carried out at room temperature using 1 μL of nanopure water ($R = 18.2 \text{ M}\Omega$) droplets within 20 seconds after depositing the drop.

2.2.4. X-ray photoelectron spectroscopy. X-ray photoelectron spectroscopy (XPS) measurements were carried out with a PHI 5600-ci spectrometer (Physical Electronics, USA). An achromatic aluminum K_α X-ray source (1486.6 eV) was used to record survey and high-resolution spectra of C 1s and O 1s regions without charge compensation. Detection was carried out at a 45° angle with respect to the normal surface; the analyzed area was 0.5 mm^2 . A curve fitting procedure was performed by means of a least-squares minimization procedure employing Gaussian-



Lorentzian functions, followed by the subtraction of a Shirley-type background. The peaks were referenced to 284.8 eV (C-C and C-H). All tests were carried out in triplicate on randomly selected areas for each condition.

2.3. Electrochemical surface characterization

2.3.1. Linear scanning voltammetry. EP parameters, including potential and current density, were evaluated through LSV to determine the passivation region of current density–potential curves for three electrolytes. Subsequently, the required potential range for the polishing process was determined. Time, temperature, and chemical composition were considered on the basis of the literature.⁵⁵ Voltammetry was carried out using a VersaSTAT3 potentiostat and a Versa Studio software data analyzer (Princeton Applied Research, USA). A three-electrode cell system was used, with a graphite rod as the counter electrode, a saturated calomel reference electrode, and 0.13 cm² of sample area as the working electrode. The surface preparation process for all the conditions was already described in the previous section. For LSV, the voltage was swept from 0 to 2 V with a scan rate of 5, 20 and 100 mV s⁻¹.

2.3.2. Potentiodynamic tests and electrochemical impedance spectroscopy. Open circuit potential (OCP), potentiodynamic polarization (PDP), and electrochemical impedance spectroscopy (EIS) tests were carried out with a three-electrode cell electrochemical setup in the same way as that for the already described LSV tests. To achieve the stabilization potential, OCP monitoring was settled for a period of 3600 s; a scan rate of 0.166 mV s⁻¹, with an applied potential range of 1 V starting from -0.5 V vs. OCP, was used. The experiments were carried out in an aerated environment at 37 ± 1 °C. Five specimens were tested for each condition. For the PDP technique, tests were set at 0.166 mV s⁻¹ with a 0.5 mV step. The corrosion rate was determined based on ASTM G102-89,⁵⁹ according to eqn (1):

$$CR = K \cdot \frac{i_{corr}}{\rho} \cdot EW \quad (1)$$

where CR is the corrosion rate in µm per year, $K = 3.27 \times 10^{-3}$ (mm g µA⁻¹ cm⁻¹ per year), i_{corr} is the corrosion current density (µA cm⁻²) obtained from polarization measurements, ρ is the

density of the material (g cm⁻³) and EW is the equivalent weight calculated to be 25.44 g.

The impedance spectra were recorded using an AC amplitude with a root mean square (RMS) of ±10 mV using a frequency range from 100 kHz to 10 mHz. The electrochemical behavior of samples was studied in modified Hanks' solution⁶⁰ and M199 culture medium (Gibco, Invitrogen Corporation, Canada). Table 3 presents the ionic concentration of both solutions compared to human plasma.

2.4. Biological tests

2.4.1. Cell culture. Human umbilical vein endothelial cells (HUVECs) were selected and considered as the most representative of the vascular environment. They were isolated from human umbilical cord samples obtained from normal term pregnancies as previously described by Loy *et al.*⁶³ HUVECs were maintained in culture at 37 °C in a saturated atmosphere of 5% CO₂ in a supplemented M199 culture medium (Gibco, Invitrogen Corporation, Canada); the supplements are presented in Table 4. When 85–90% confluence was reached, the cells were enzymatically detached from the plate (0.05% trypsin, Gibco, Invitrogen Corporation, Canada) and then reseeded at a ratio of 1:3 or used for experiments. For the experiment reported here, HUVECs have been used at passage 5.

2.4.2. Indirect viability assay. The indirect cytotoxicity test was performed following the ISO 10993-5:2009 (ref. 64) procedure. Stainless steel (SS316L) was used as a reference material and prepared as described elsewhere by Campelo and coworkers.⁶⁵ Before the test, all samples were submitted to the cleaning step mentioned in the previous section “Sample preparation”. All samples were also sterilized by UV irradiation. Briefly, each side of the samples underwent two 15 minute cycles of UV irradiation (256 nm). After that, the samples were stored in a sterile 24 multi-well plate until use prior to the cytotoxicity test. Briefly, 1 cm² treated samples were immersed in 660 µL of M199 culture medium, supplemented with 1% penicillin–streptomycin for 1, 3 and 7 days. At each time point, the medium was collected from the samples and subsequently used for the viability test. Viability tests were performed using different concentrations of the extracted media: 100%, 10%, and 1% concentrations. Before putting them in contact with cells, extracted media were supplemented with the same

Table 3 Ionic composition (mg L) of modified Hanks' and M199 solutions compared to human plasma

Ions	Modified Hanks ⁶⁰	M199 (ref. 61)	Human plasma ⁶²
Cl ⁻	3542	2549	3360–3900
Na ⁺	2795	2102	3000–3400
Ca ²⁺	35	80	84–110
Mg ²⁺	14	24	15–30
HCO ₃ ⁻	1654	1342	1110–2400
HPO ₄ ²⁻ /H ₂ PO ₄ ⁻	48	133	270–450
SO ₄ ²⁻	78	94	5–150
K ⁺	172	156	130–210
D-Glucose	714	1000	600–1200
Albumin	-	-	28 000–56 000



Table 4 List of supplements for cell culture¹ and indirect viability² media

Supplement	HUVEC medium
Fetal bovine serum (FBS) ^a	5%; ¹ 10% ²
Penicillin-streptomycin (P/S) ^a	1%
Human fibroblast growth factor-basic (FGF) ^b	2 ng mL ⁻¹
Human epidermal growth factor (EGF) ^b	0.5 ng mL ⁻¹
Ascorbic acid ^c	1 µg mL ⁻¹
Heparin ^c	90 µg mL ⁻¹
Hydrocortisone ^c	1 µg mL ⁻¹

^a Gibco, Invitrogen Corporation, Canada. ^b Life Sciences, USA.

^c Sigma Aldrich, Canada.

supplements described in Table 4. The CTRL was obtained by measuring the HUVECS in cell culture solution. For the 10% and 1%, extracted media were diluted with complete M199 medium (M199 medium supplemented with the reagents as mentioned earlier). One day prior to contact with the extract, HUVECs were seeded in the well of a 96 multi-well plate at a density of 20 000 cells per cm² and incubated at 37 °C and in 5 vol% CO₂ for 24 hours in 100 µL per well of complete M199. The day after, the medium was removed, and 100 µL of the extracts were added to the well containing the HUVECs and

incubated for 24 h. The extracts were then removed, and 100 µL of 1% solution of resazurin sodium salt in complete M199 medium was added to the cells and incubated for 4 hours at 37 °C and in 5 vol% CO₂. After the incubation, the solutions containing the now reduced resorufin product were collected and the fluorescence intensity at a 545 nm_{ex}/590 nm_{em} wavelength was measured with a SpectraMax i3x multi-mode plate reader (Molecular Devices, USA). Fluorescence intensity is proportional to cell viability.

2.5. Statistical analysis

All numerical data were analyzed by ANOVA and followed by Tukey's test for specific comparisons between mean values. Results were expressed as the mean value ± standard deviation (SD). A statistically significant difference was considered, if the probability values for a set of data were found to be <0.05 ($p < 0.05$).

3. Results

3.1. Diffusion-limited current plateau

Three scan rates were applied to determine the key parameters for EP, specifically the applied voltage and current density range. Additionally, the corresponding voltage–current curves

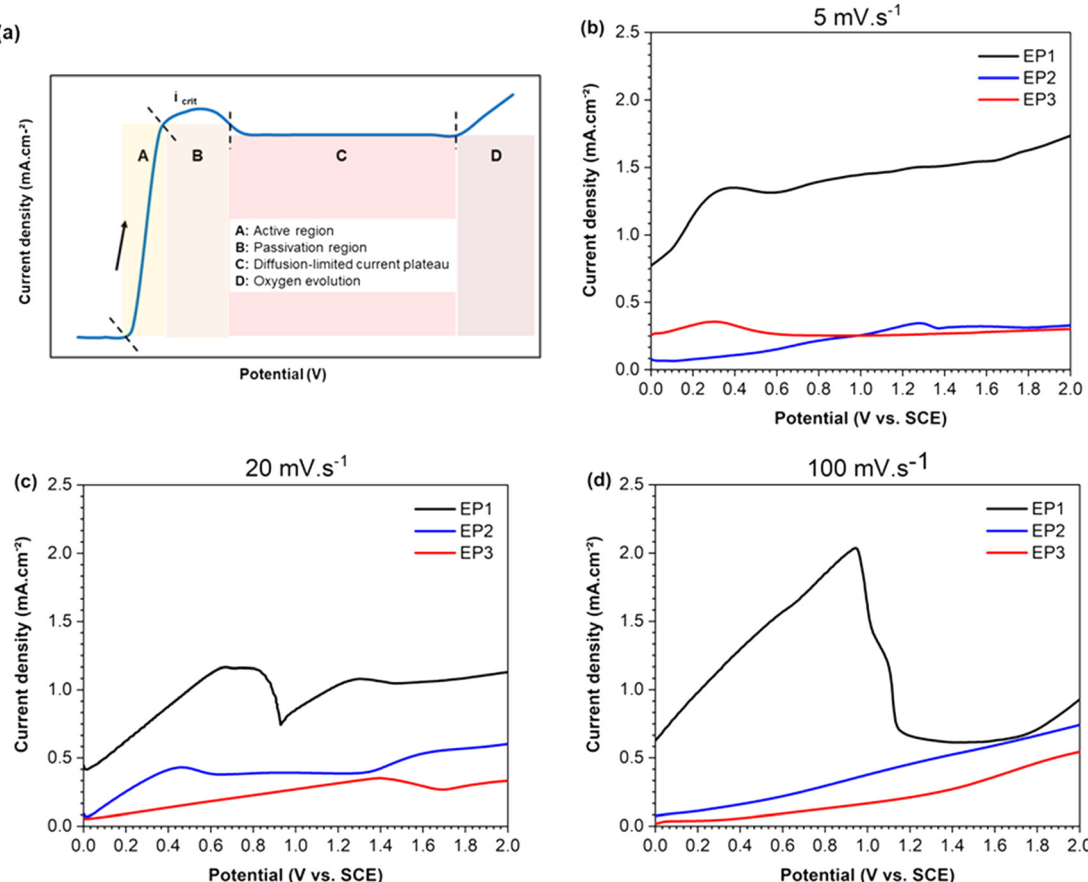


Fig. 1 (a) Schematic illustration of a reference LSV plot, and curves for Hadfield samples in EP1 (perchloric acid, ethanol and glycerol), EP2 (acetic acid, perchloric acid and glycerol) and EP3 (choline chloride and ethylene glycol) electrolytes. All experiments were conducted at 21.0 ± 0.5 °C with scan rates of (b) 5 mV s⁻¹, (c) 20 mV s⁻¹ and (d) 100 mV s⁻¹.



were used to evaluate the mass transport mechanism occurring at the solution–sample interface for each electrolyte. The schematic LSV curve in Fig. 1a shows the four characteristic regions (A, B, C and D). In brief, region A corresponds to the anodic dissolution of the metal, and therefore to the highest corrosion rate at critical current density (I_{crit}). Region B is associated with the formation of a passive layer, followed by region C, where mass transport controls both dissolution and diffusion of anions through the passive layer. Finally, region D represents the transpassive region where oxygen gas is produced, leading eventually to pitting.

Fig. 1b–d show the linear sweep voltammograms, acquired from 0 to 2 V, measuring the anodic behavior of the Hadfield alloy immersed in the three electrolyte compositions at different scan rates. When a scan rate of 5 mV s^{-1} was adopted, samples tested in EP1 solution exhibited a more positive anodic current (black line of Fig. 1b) compared to those obtained for samples tested in EP2 and EP3 solutions (blue and red lines, respectively). However, EP1 conditions displayed a continuous progressive curve within the studied potential range without reaching a maximum current peak. This observation may be correlated to the absence of a passive layer formation on the sample surface. For EP2 and EP3 electrolytes, a tiny maximum peak current ($I_{\text{crit}} \sim 0.35\text{--}0.36 \text{ mA}$) was reached, followed by a plateau of diffusion for potentials higher than 0.6 V. The rising of a low current density for samples treated with EP2 and EP3 electrolytes showed a limited mass transport of the

dissolving ions to the electrolyte–sample interface, which could be associated with the slow scan rate used (5 mV s^{-1}).

When the intermediate scan rate of 20 mV s^{-1} (Fig. 1c) was applied, EP1 conditions still exhibited the highest positive anodic current above all conditions, with a maximum current peak at $I_{\text{crit}} \sim 0.68 \text{ mA}$. The active dissolution region of EP1 was not immediately followed by a steady passive region plateau (represented by the C section of Fig. 1a). Still, the current density increased immediately until 1.5 V followed by a slight but not significant increase. In contrast, samples tested in EP2 solution at 20 mV s^{-1} , showed an $I_{\text{crit}} \sim 0.46 \text{ mA}$ (for low potentials, <0.6 V) and a passive region from 0.6 to 1.35 V. After this potential range, the current still increased, indicating oxygen evolution. The EP3 voltammogram displayed an increasing linear trend reaching a value of 0.4 mA at 1.4 V, followed by a small decrease until 1.7 V. Then a continuous slight rise until the final studied potential (2 V) was registered.

At the highest scan rate of 100 mV s^{-1} (Fig. 1d), LSV curves showed an I_{crit} peak followed by a plateau region only for samples tested in EP1 solution, suggesting that no passive film occurs on sample surfaces.

Based on these results, the current density was settled at 0.5 A cm^{-2} for EP1 and EP2 electrolytes while 0.25 A cm^{-2} for EP3. The potential range for each electrolyte was calculated based on the exposed area of the samples used in the following experiments, as indicated in Table 2.

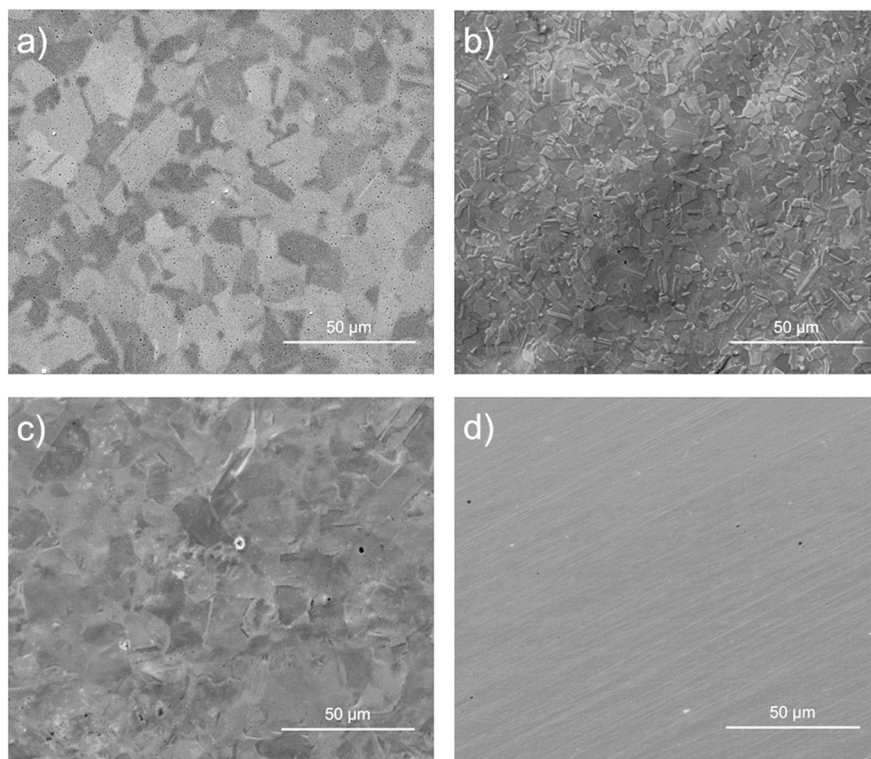


Fig. 2 Secondary electron mode SEM images for EP1 (a), EP2 (b), EP3 (c), and MP (d) samples.



3.2. Microstructure and morphology

The morphology, microstructure, and detailed patterns of the EP sample surfaces were analyzed by means of SEM, and compared to the MP surface (Fig. 2).

After EP and MP processes, samples exhibited a mirror-like surface; no noticeable features were observed with the naked eye. For EP1 conditions (Fig. 2a), several pits (small black points) across the surface were noticed, while for EP2, negligible pitting corrosion was observed (Fig. 2b). Additionally, the formation of a thin brown layer was observed during the EP1 and EP2 electropolishing processes. This layer was easily removed with methanol during the rinsing step. It is likely associated with a viscous layer that forms at the metal-electrolyte interface due to anodic dissolution facilitated by the diffusion-adsorption process during the EP process.⁶⁶

From Fig. 2b, the microstructure observed is associated with austenite (γ) and martensite (α' or ε) phases which are commonly observed in the type of steel used in the present study.⁶⁷ The EP3 conditions showed a heterogeneous surface (Fig. 2c); the microstructure was evidenced, but the grain boundaries were not distinctive as for EP1 and EP2 conditions. Different phases and grains exhibited distinct crystallographic orientations, resulting in uneven dissolution rates as observed in the AFM images of the EP2 conditions (Fig. 3). In Fig. 3a, a line was drawn through two regions (A and B) touching

prominent boundaries to assess if height profilometry could qualitatively depict the dissolution phenomena. As depicted in Fig. 3b, a height difference of nearly 20 nm was observed between regions A and B, with region B showing smaller height peaks. Additionally, Fig. 3c indicates that the phase in region B is predominantly formed of small particles (less than 10 nm), contributing to a more pronounced surface smoothening compared to region A. Moreover, the amplitude mode from the same image delineated distinct grain boundaries. In fact, Landolt³¹ explained this phenomenon as the anodization of polycrystals with random orientations which leads to the formation of crystallographic facets, revealing different patterns on crystal planes. These observations indicate that the two main phenomena observed during EP, *i.e.* macro- and micro-smoothening, are influenced by the microstructure itself and the electrolyte chemical composition.⁶⁶ However, anhydrous solutions used in EP treatments may lead to the formation of complex oxides on the surface, which could be attributed to the adsorption of organic products originating from the viscous organic solution of choline chloride and ethylene glycol, as well as from perchloric reactions.³⁶

Finally, MP samples (Fig. 2d) showed parallel lines associated with the grooves formed during the mechanical polishing process. No other features, such as grain boundaries, were observed for these conditions contrary to the tested electropolishing conditions.

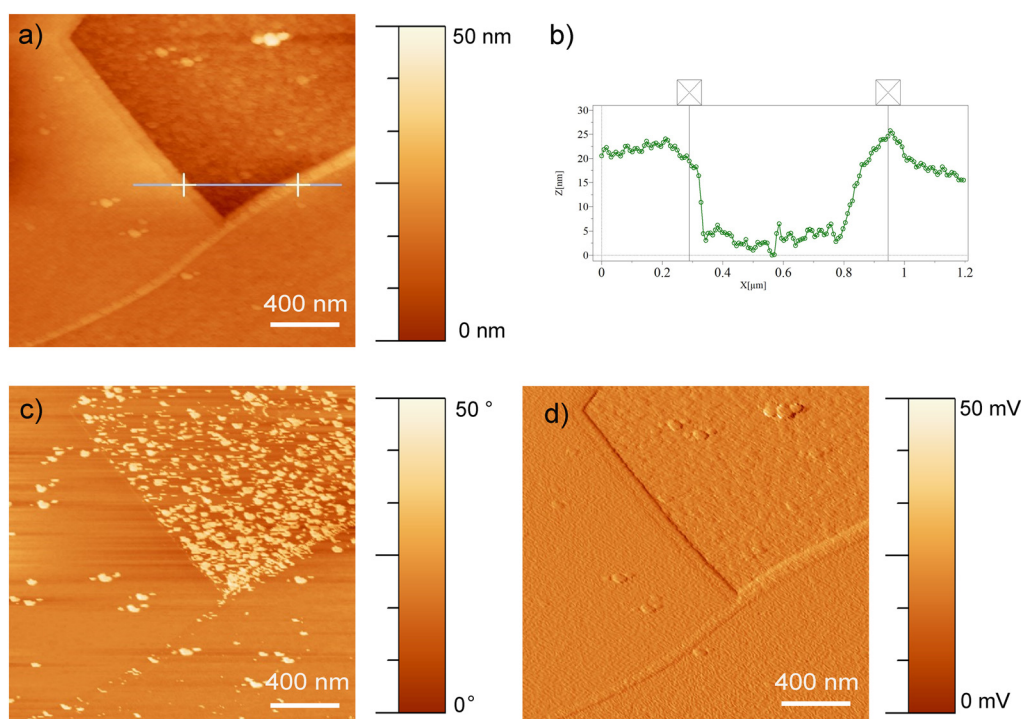


Fig. 3 AFM measurements of EP1 conditions show the a) height image and the b) height profile of the line passing through grain boundaries. The c) phase, and d) amplitude images are also presented.



3.3. Surface roughness, topography, and wettability

The RMS and R_a roughness values of EP1, EP2, EP3, and MP samples were evaluated for an area of $20 \mu\text{m} \times 20 \mu\text{m}$ (Fig. 4a) and $2 \mu\text{m} \times 2 \mu\text{m}$ (Fig. 4b).

For the large scan area (images of $20 \mu\text{m} \times 20 \mu\text{m}$), EP1 and MP treatments appeared to produce a smoother surface ($\text{RMS}_{\text{EP1}} = 5.3 \pm 0.8 \text{ nm}$ and $\text{RMS}_{\text{MP}} = 4.4 \pm 0.3 \text{ nm}$; $R_a,_{\text{EP1}} = 3.6 \pm 0.9 \text{ nm}$ and $R_a,_{\text{MP}} = 3.4 \pm 0.2 \text{ nm}$, respectively), whereas EP2 and EP3 samples exhibited a rougher surface ($\text{RMS}_{\text{EP2}} = 23.2 \pm 2.0 \text{ nm}$ and $\text{RMS}_{\text{EP3}} = 23.1 \pm 1.6 \text{ nm}$; $R_a,_{\text{EP2}} = 10.0 \pm 1.1 \text{ nm}$ and $R_a,_{\text{EP3}} = 16.9 \pm 1.3 \text{ nm}$, respectively). When the analyses were carried out on a smaller area ($2 \mu\text{m} \times 2 \mu\text{m}$), surface roughness decreased, as expected, with no significant differences between EP1, EP2, and MP conditions. However, EP3 exhibited the highest calculated surface roughness ($4.6 \pm 0.7 \mu\text{m}$) which displayed a significant difference in both RMS

and R_a calculations when compared to the others. As suggested by Sautebin *et al.*,⁶⁸ it is possible to achieve microsmoothing without macrosmoothing. Both these definitions are linked to local current densities on a rough surface and the limitation of mass transport, respectively.^{66,69} Furthermore, the first derivative images (Fig. 4g–j), revealed that grain boundaries form steps along two adjacent grains, a phenomenon primarily correlated to the dissolution process during electropolishing. For EP1, the average height step was about 20 nm. EP2 conditions displayed a similar average height of 9 nm (Fig. 4h), and EP3 conditions exhibited a maximum average height step of $\sim 32 \text{ nm}$, with the highest variation among the three EP conditions. In the case of MP, the fine grooves across the surface showed an approximate average of $\sim 16 \text{ nm}$ from the edge to the substrate.

Finally, the surface wettability for all the studied conditions was assessed by static contact angle (WCA) measurements. All

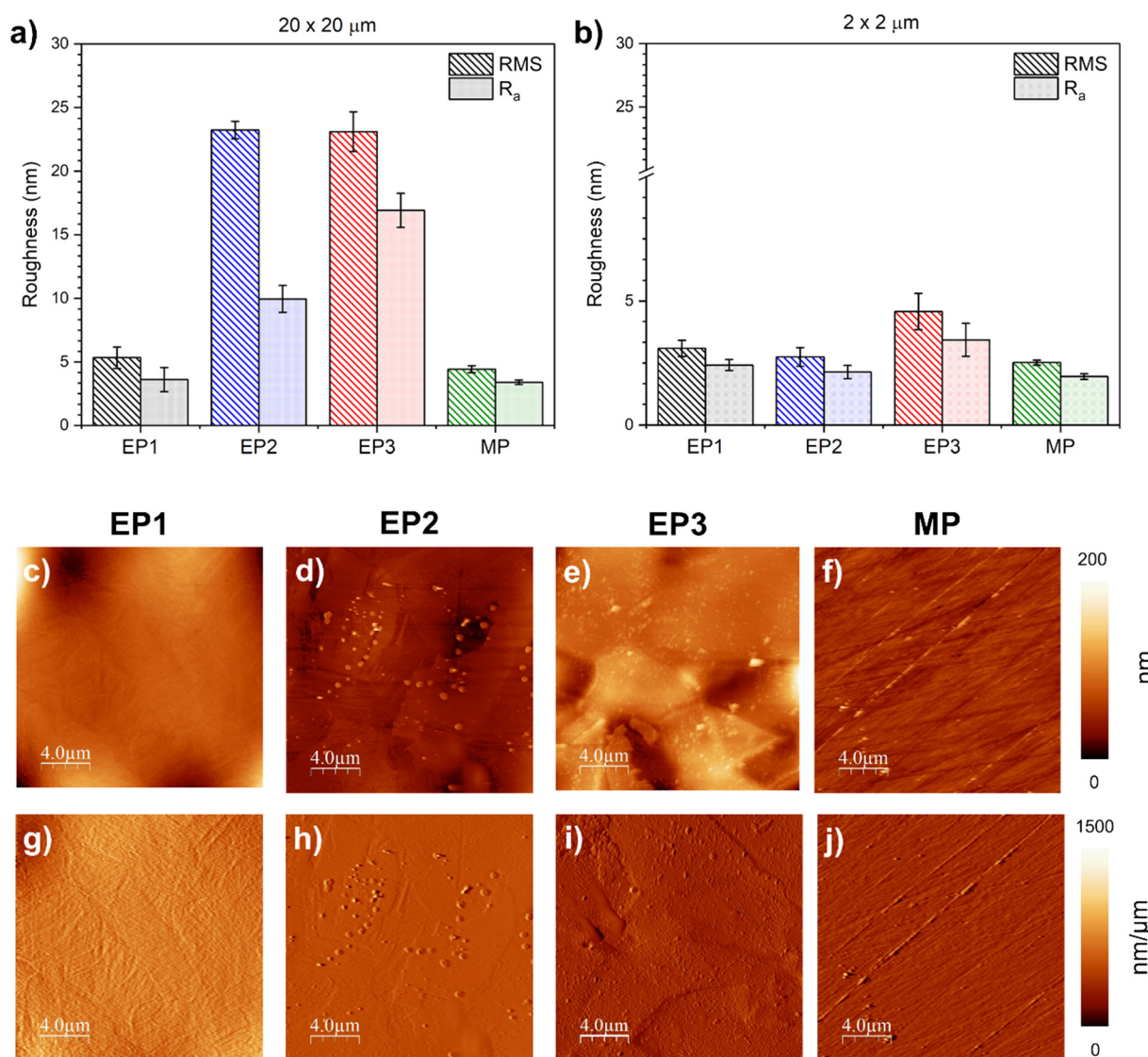


Fig. 4 Surface roughness values (R_a and RMS) were measured for two scan sizes, (a) $20 \mu\text{m} \times 20 \mu\text{m}$ and (b) $2 \mu\text{m} \times 2 \mu\text{m}$. Figures (c–f) present AFM images and their respective first derivative images (g–j) for EP1, EP2, EP3 and MP conditions.

of them showed a hydrophilic behavior; even though EP1 and MP showed the lowest roughness values compared to EP2 and EP3, the surface wettability did not follow the same trends. In fact, the EP2 surface displayed the lowest WCA_{EP2} value, $53 \pm 2^\circ$, among all treatments. EP1 showed an intermediate behavior (WCA_{EP1} = $64 \pm 2^\circ$), whereas EP3 and MP presented no significantly different CA values, WCA_{EP3} = $72 \pm 5^\circ$ and WCA_{MP} = $76 \pm 3^\circ$, respectively, compared to the other conditions.

3.4. Chemical composition after EP and MP

The chemical surface composition (at%) for all the conditions was evaluated by XPS survey analysis (Table 5).

The chemical composition of each condition appeared to be homogeneous all over the surface, with a standard deviation of less than 2% for all elements detected. The Fe amount for EP1 and EP2 conditions was similar to those mechanically polished (~ 12 at%), while EP3 conditions exhibited the highest amount of Fe concentration (16.1 ± 1.3 at%). EP1 samples presented the highest Mn content (2.9 ± 0.1 at%) followed by EP3 (2.5 at%), MP (1.8 ± 0.7 at%), and finally, EP2 (1.2 ± 0.4 at%). The overall metallic percentage (Fe + Mn) at% found on each sample surface was calculated (Table 5). EP3 and MP samples revealed a more metallic surface, with 18.7% and 15.1%, respectively, than EP1 and EP2 samples with 13.8%. That said, EP3 and MP samples displayed a low oxygen amount, ~ 40 at%, while they had the highest metallic concentration. By opposite, the highest oxygen percentage was obtained for EP1 and EP2 conditions, ~ 44 at%, with a lower metallic percentage than MP and EP3. Finally, the amount of carbon was similar for all EP samples, ~ 40 at%, while MP had the highest C value of 44 at%. The high-resolution spectra of the different elements present on the surface were obtained (Table 6). The deconvolution of the O 1s high-resolution spectra evidenced differences. O1 (ref. 17) referred to metal oxide (Me-O_x), and O2 was related to the formation of hydroxides (Me-OH_n), while O3 and O4 were associated with physio-adsorbed water.

The oxygen species present on the surface were strongly correlated to the treatment used (Table 6). The ones found for EP1 conditions were significantly different from those present on the surface of the other conditions. Indeed, they displayed the highest O3 (33.0%) and O4 (10.4%) contributions, a relevant presence of O2 (40.0%) and the lowest O1 percentage (16.5%). In opposite, the EP3 sample exhibited the highest

contribution of metal oxides (O1) (53.4%), when compared to the other conditions, and the lowest area associated with O3 and O4, respectively, 13.8% and 2.6%. The EP2 and MP samples presented a similar composition of the O band, with an O1 contribution of $\sim 41\%$, and an O2 one of the same amount as O1. The oxidation state of the surface given by the ratio O_{metal}/(Fe + Mn) was similar for the EP2, EP3 and MP samples, ranging from 1 to 1.3, while the EP1 conditions displayed the lowest value (0.5). The latter presented a higher amount of O bound to hydroxides than to metal oxides. The EP2 conditions displayed the highest metallic oxide contribution, with an O/metal ratio of 1.3, and no O4 contribution, as well as the lowest WCA value, WCA_{EP2} = $53 \pm 2^\circ$, among all the conditions. These conditions also led to the highest $R_{\text{Fe/Mn}}$ compared to the other conditions, with visible grain boundary features (~ 9 nm step height). Furthermore, although the EP1 and EP3 conditions presented a mirror-like surface, EP1 showed numerous corrosion pits (Fig. 2a), while for the EP3 conditions, adsorption of organic residues from the viscous organic solution was observed. These two patterns can mislead electrochemical analysis, resulting in a falsely elevated corrosion rate due to the high presence of pits and slower dissolution, in the case of EP3. Therefore, EP2 exhibited the most promising chemistry and surface properties and will be used for further electrochemical and cytotoxicity assays compared to the reference sample, the MP conditions.

3.5. Electrochemical characterization

The results of PDP and EIS analyses of the EP2 and MP samples in both media, modified Hanks' solution and M199, are reported in Fig. 5. The pH of both solutions remained stable after electrochemical tests, consistently remaining within a range of 7.4 and 7.3; the detailed ionic composition of the solutions are presented in Table 3. The corrosion potential (E_{corr}), current density (i_{corr}), and corrosion rate (CR) were extracted from PDP curves (Fig. 5a). Regardless of the solution used, MP exhibited similar corrosion potentials ($E_{\text{corr, MP}} = -740$ mV). However, EP2 showed a significant difference in corrosion potentials when analyzed in M199 ($E_{\text{corr, EP2}} = -710$ mV) and in Hanks' ($E_{\text{corr, EP2}} = -750$ mV) solutions. Therefore, the EP and MP samples displayed significantly different corrosion current densities. Indeed, when analyzed in Hanks' solution (Fig. 5b, solid lines), the

Table 5 Chemical composition of EP1, EP2, EP3 and MP, highlighting Fe, Mn, C and O elements (at%). O_{metal}/(Fe + Mn) corresponds to the ratio between oxygen from metal oxide (peak at ~ 530 eV) and the total amount of detected metals

Sample	% Fe	% Mn	% C	% O	% (Fe + Mn)	O _{metal} ^a /(Fe + Mn)	R _{Fe/Mn} ($R_0^b = 6.7$)
EP1	11.0 ± 1.5	2.9 ± 0.1	40.9 ± 2.0	43.4 ± 1.9	13.8	0.5	3.8
EP2	12.7 ± 1.4	1.2 ± 0.4	40.7 ± 1.1	44.9 ± 1.4	13.8	1.3	10.6
EP3	16.1 ± 1.3	2.5 ± 0.1	39.4 ± 1.2	39.8 ± 0.9	18.7	1.1	6.4
MP	13.3 ± 0.2	1.8 ± 0.7	44.1 ± 1.2	37.2 ± 1.4	15.1	1.0	7.4

^a O_{metal} corresponds to the proportion of O1 area per absolute oxygen (Table 6). ^b R_0 is the ratio between Fe and Mn in the base material. Cl and N traces were found to be less than 0.1 at% under all conditions.



Table 6 Percentage of area for O 1s peaks (O1, O2, O3, O4) and binding energy (eV)

Sample	% area (O 1s peaks)			
	O1 529.7–530.1 eV	O2 531.2–531.5 eV	O3 532.3–532.5 eV	O4 533.5–533.9 eV
EP1	16.5 ± 0.8	40.0 ± 8.2	33.0 ± 9.7	10.4 ± 2.3
EP2	41.1 ± 6.7	43.8 ± 3.4	15.1 ± 8.9	–
EP3	53.4 ± 2.0	30.2 ± 0.7	13.8 ± 1.3	2.6 ± 1.4
MP	41.5 ± 4.7	37.6 ± 4.1	17.8 ± 4.6	3.1 ± 0.5

MP surfaces showed a lower corrosion current density (i_{corr} , $MP = 7.2 \pm 1.3 \mu\text{A cm}^{-2}$) than EP2 ($i_{corr, EP2} = 11.2 \pm 1.7 \mu\text{A cm}^{-2}$). While similar trends for cathodic branches were observed, the anodic dissolution one differed depending on the conditions. In fact, the EP2 ones displayed a faster anodic dissolution when compared to the MP ones, showing an overall corrosion resistance. Indeed, for Hanks' solution, the EP2 corrosion rate ($138 \pm 25 \mu\text{m per year}$) was twice as high as that of the MP samples ($45 \pm 24 \mu\text{m per year}$). Electrochemical measurements performed in M199 medium (dashed curves, Fig. 5b) showed an opposite trend to that observed for samples tested in Hanks' solution. It is noteworthy that the corrosion rate of MP showed no significant difference regardless of the solution.

Fig. 5c presents the Nyquist plots. Regardless of the medium, the EP2 surface displayed a more capacitive behavior than that of the MP sample in both solutions; in other words, the impedance values of EP2 reached higher values than those obtained for the MP samples. This behavior could be related to the adsorption of inorganic molecules, such as HPO_4^{2-} or other phosphate-based anions, forming a film at the surface that acts as a barrier against the corrosion process.⁷⁰ Analyzing the Bode plots (Fig. 5d), all the conditions presented a time constant for the medium frequency region ($\sim 100 \text{ Hz}$). The time constant in this region was attributed to localized corrosion of the substrate and the formation of corrosion products; the indication of another time constant at lower frequencies ($\sim 0.1 \text{ Hz}$) was associated

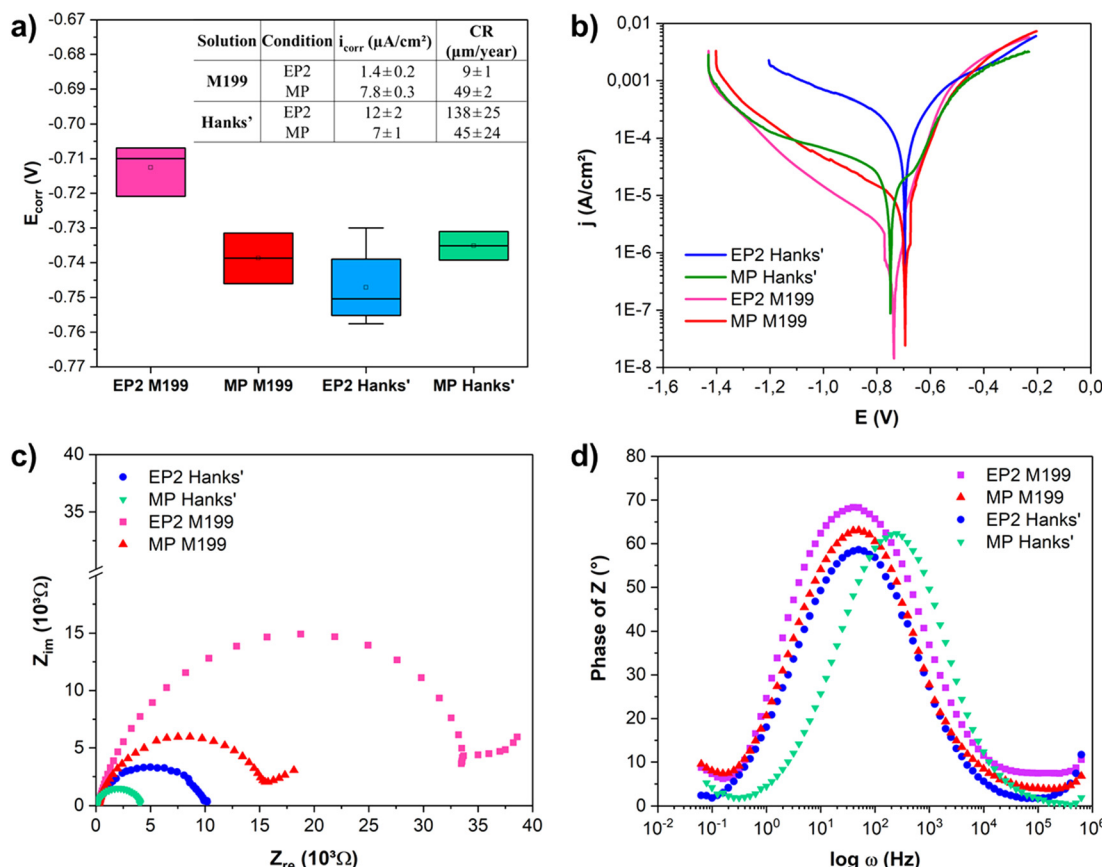


Fig. 5 Electrochemical measurements of samples after EP2 and MP treatments: a) OCP and kinetic parameters of PDP measurements for EP2 and MP conditions in Hanks' solution and M199, b) PDP curves, c) Nyquist plots, and d) Bode plots as a function of phase angle in modified Hanks' solution (green) and M199 medium (blue).

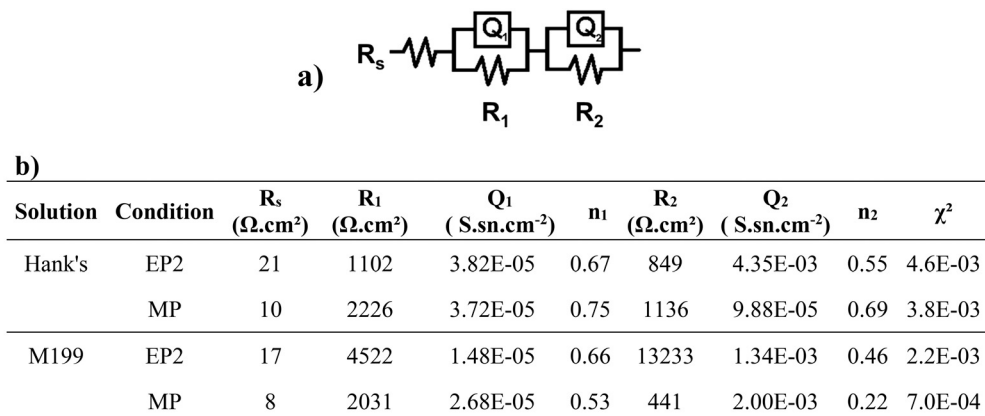


Fig. 6 Electrical equivalent circuit attributed to this study (a) and the fitting parameters of EP2 and MP samples in contact with Hanks' and M199 solutions (b) where R_s : the electrolyte resistance, R_1 : resistance of the oxide/corrosion product layer, Q_1 : capacitance of the oxide/corrosion product layer, n_1 : dimensionless coefficient associated with Q_1 , R_2 : resistance of the inner oxide layer, Q_2 : capacitance of the inner oxide/metal layer, and n_2 : dimensionless coefficient associated with Q_2 .

with species adsorption on the metallic substrate.⁷¹ Comparing the Bode plots, the time constant of the MP samples shifted to higher frequencies (around 500 Hz) when Hanks' solution was used. This shift indicates a faster corrosion process compared to the other conditions as seen for a Mg-based system.⁷²

Electrochemical results were fitted using an electric equivalent circuit (EEC) to quantify changes in impedance values (Fig. 6a). The EEC was composed of R_s , the electrolyte resistance, with R_1 and Q_1 (Q as the constant phase element, or CPE) representing the ohmic resistance of the oxide/corrosion product interface and the capacitance of this layer, respectively. The additional (R_2 and Q_2) components represent the resistance of the corrosion products adsorbed when in contact with the studied solutions, as well as the capacitance of the electric double layer. All fitted electric parameters are shown in Fig. 6. The lowest values of R_s were found for the MP conditions, with $R_{s, MP} = 10 \Omega$ for Hanks' solution and $R_{s, MP} = 8 \Omega$ for M199. Both conditions exhibited

a higher outer layer resistance R_1 than the corresponding R_2 (resistance of the inner passive oxide layer). For the EP2 conditions, the exponential n_1 , representative of the homogeneity of the system interface, was lower than 0.75 for both solutions. For the MP conditions, the same parameter presented a higher value when in contact with Hanks' solution (>0.7) than for M199 (0.2–0.5). Both conditions presented a deviation from the ideal capacitor which is related to the materials' surface chemical heterogeneities and roughness.⁷³

Although a different EP2 behavior was observed after Hanks' or M199 exposure, these conditions showed the highest charge transfer resistances, suggesting that their conductivity was low, whereas MP-treated samples exhibited lower corrosion resistance for both solutions.

XPS analyses were performed for EP2 and MP surfaces after exposure to both solutions as shown in Fig. 7. A lower C content, that is ~30 at%, was observed when compared to that of pristine conditions, which was around 40 at% (Table 5).

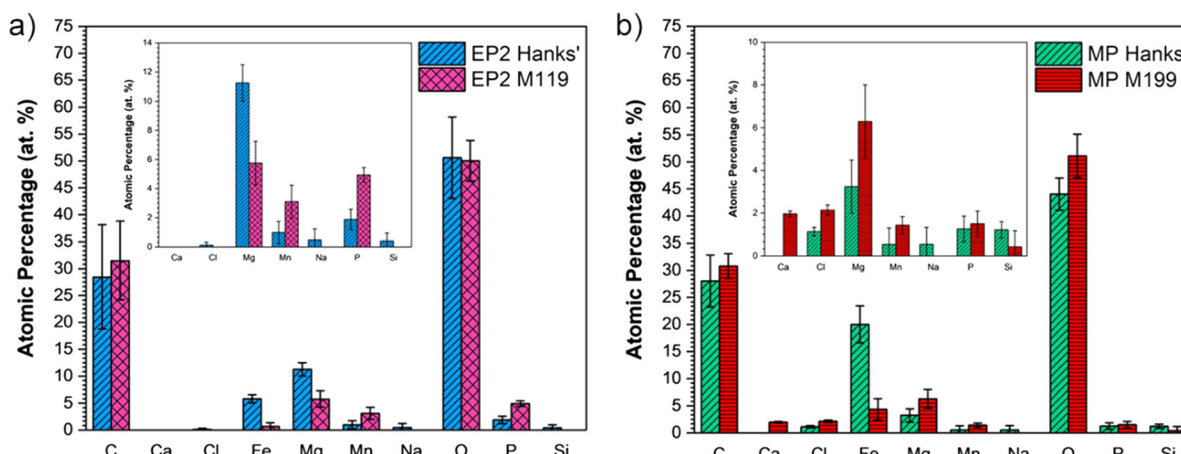


Fig. 7 Chemical composition (at%) of a) EP2 and b) MP conditions analyzed after the PDP test in modified Hanks' solution and M199 medium.

Compared to the chemical composition before corrosion assays, the oxygen content increased for all conditions (from 40 to 50 at%) except for EP2 analyzed in M199 solution. The MP conditions displayed the highest Fe amount on the surface after immersion in Hanks' solution, that is 20 ± 3 at% *versus* 5.8 ± 0.7 from EP2 analyzed in the same solution. Several elements present in both solutions, such as P and Mg, were found on EP2 and MP corroded surfaces.

As seen in Fig. 7, Cl, Na, and Si were found in lower amounts (<2 at%) under EP2 conditions when analyzed in modified Hanks' solution and in either solution for MP samples. On the other hand, Fe and Mn were detected under both conditions in either solution. No statistical difference was observed for Mn detected amounts and all samples presented a lower Fe presence, except for MP in modified Hanks' solution, where the values increased compared with the initial conditions. Interestingly, no Ca was detected under either conditions in modified Hanks' solution, despite both electrolytes containing Ca^{2+} (as shown in Table 3, Ca^{2+} Modified Hanks: 35 mg L^{-1} and

Ca^{2+} M199: 80 mg L^{-1}). Ca was only detected on the MP surfaces immersed in the M199 solution. Moreover, the presence of Mg and P was detected under all corroded conditions since these elements were also present in both chemical compositions of solutions (see Table 3).

Overall, the Fe/Mn ratio decreased after corrosion tests, except for the MP conditions in modified Hanks' solution, which presented a ratio $R_{\text{Fe/Mn}}$: 37 (at%/at%), which means five times higher than the initial conditions, *i.e.* MP. This is mainly attributed to the increase of Fe content since no significant statistical changes were observed for Mn amounts in MP samples before and after the PDP test in Hanks' solution. The low Mn content may be attributed to a preferential pickling of the Mn compounds when in contact with both solutions.¹⁷ Moreover, the increase of oxygen may be attributed to the oxidation and adsorption of degradation products originating from Hanks' solution, which often contains carbonate, phosphate, and hydroxides.^{74,75} Regarding the Cl content found in MP samples analyzed in both solutions, it might be related

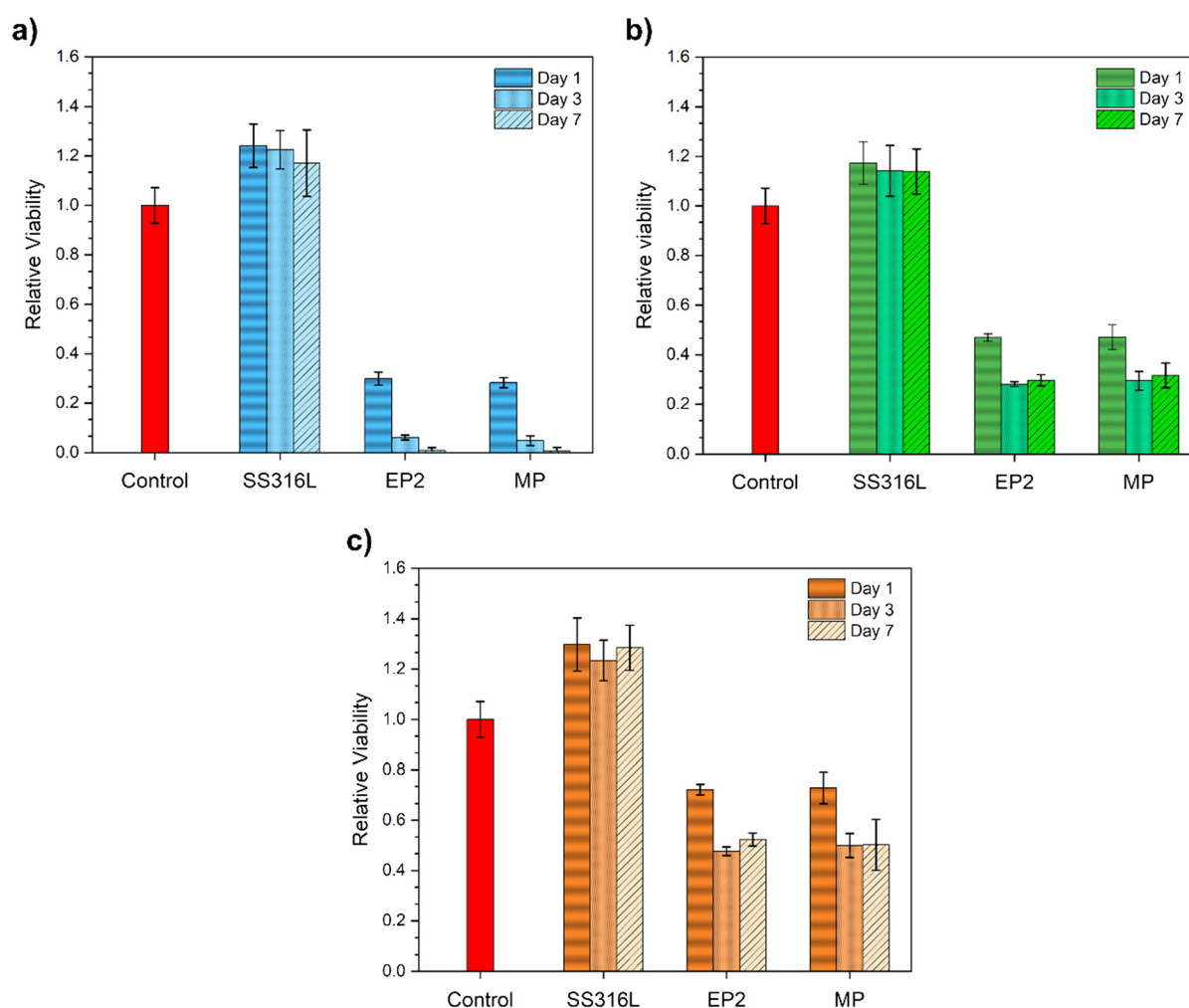


Fig. 8 Cell viability of Fe-13Mn-1.2C at different eluate concentrations: (a) 100%, (b) 10% and (c) 1%. The extracts from EP2 and MP samples, upon contact with HUVECs, were analyzed and compared to the standard culture medium (control) and extracts from SS 316 L. Cell viability was measured after 1 day of incubation by means of a resazurin salt solution assay. The graph shows the mean fluorescence \pm standard deviation, recorded from HUVECs treated with the different experimental conditions. Results have been normalized against the control conditions.



to the higher amount of metallic species under the pristine conditions (% Fe + Mn: 15.1 at%) leading to the formation of metallic chloride (FeCl_2) in the innermost layer at the surface, since Cl^- ions present a high ionic volume (approximately $23.3 \text{ cm}^3 \text{ mol}^{-1}$).⁷⁶ Moreover, the higher Fe content observed on MP surfaces exposed to Hanks' solution might explain the less capacitive behavior evident in Nyquist plots. This could indeed be attributed to the dissolution of metallic elements from the bulk to the oxide interface since the resistance R_2 (inner oxide layer) was lower than R_1 (resistance of the oxide/corrosion product layer). Since the metallic ions encounter more resistance, they remain in place while the cathodic reaction occurs at the outer interface, increasing the oxygen content.

3.6. Cell viability assays

The relative cell viability of EP2 and MP conditions was compared with that of the control and SS316 samples for 1, 3 and 7 days at different eluate concentrations (100%, 10% and 1%).

Regarding the viability of cells with 100% elution (pure extract), the SS316L conditions did not alter the HUVEC viability over time compared with the control. For the Fe-13Mn-1.2C alloy, both MP and EP2 conditions induced a decrease in cell viability, with no statistical difference between them. After 3 and 7 days, a decrease in cell viability was observed. Indeed, this behavior was expected, and it was related to the concentration of the degradation products of the medium in contact with the cells; longer cell contact (7 days *vs.* 1 day) with high-concentration degradation products (100% *vs.* 1%) was generally responsible for the lower viability^{21,77} for the HUVEC assays.

SS316L diluted extracts again exerted no effect on cell viability (Fig. 8b and c): in fact, cells showed viability comparable with the control conditions. As for the EP2 and MP samples, once again, their extracts induced cytotoxic effects on the treated HUVECs. However, an inverse correlation between extract concentrations and cytotoxic effects can be observed: increasing the dilution factor enhances the viability of the treated HUVECs. Moreover, for both conditions, no statistical difference was observed for EP2 and MP after 3 and 7 days in contact with the extract.

4. Discussion

Although degradable metallic materials are designed for gradual dissolution, their surface properties play a critical role in regulating corrosion and eliciting a host response at the initial interface between the medical implant and the biological environment.^{78,79} Natural and induced oxide layers have been demonstrated to enhance surface corrosion resistance and improve cell compatibility on biodegradable materials.⁸⁰ However, achieving control over the oxidation of biodegradable materials becomes a challenging endeavor when striving for simultaneous degradation and avoidance of cytotoxic effects in the human body.⁸¹ The passivation mechanism through electropolishing has undergone extensive investigation across various steel systems. The nature of electropolishing treatments

on biodegradable steel surfaces is elucidated based on the chemical composition of the electrolyte, surface morphology, and topography, as well as the cytotoxicity effects and electrochemical analyses for each EP and MP treatment studied in this research.

4.1. The influence of anodic dissolution on the electropolishing parameters and surface features

EP parameters were mainly related to the voltage–current curve where the limiting current region in LSV plots can be affected by, among other things, the electrolyte composition and the distribution and/or the concentration of electroactive species.^{82,83} The rate and reversibility of a reaction may be associated with variations in the peak observed in the current–potential graph. Indeed, samples tested in EP1 solution presented, at all scan rates, the highest current density peak (I_{crit}), even though no plateau was observed at 5 mV s^{-1} (Fig. 1b). The sharp increase in current density by increasing the scan rate from 20 to 100 mV s^{-1} could be associated with a fast reaction between the sample and the electrolyte due to the high concentration of acid in the solution, *i.e.*, perchloric acid.⁸⁴ At 100 mV s^{-1} , the development of a plateau confirms the formation of an oxide layer on the steel surface. Indeed, the main mechanism proposed for this electrolyte is based on the formation of a salt film by the ions extracted from the substrate (Fe^{2+} , $\text{Mn}^{2/4+}$) which triggers the formation of pits. Polar groups, such as ethanol, may cover the surface and not allow metallic ions such as Fe^{2+} and Mn^{4+} to dissolve in the bulk solution, leading to a heterogeneous electropolishing process and an undefined plateau (region C, Fig. 1a).

For EP2, perchlorate (ClO_4^-) and acetate (CH_3COO^-) anions can create a film through an adsorption process. Two hypothetical reactions of acetic acid and acetate ions with a divalent metallic element (Me^{2+}) can occur:



Acetic acid, only under specific conditions (*e.g.*, $T > 30 \text{ }^\circ\text{C}$) exhibited high activity in buffering the H^+ concentration at the metallic surface; furthermore, the I_{crit} current was shifted to the lower potential range of the mass transfer limiting conditions.⁸⁵ However, an increase in the current–potential curve without any plateau for 100 mV s^{-1} demonstrates that the reactions created under these conditions are too slow to follow this specific rate.⁸⁶

Samples treated in ChCl:EG solution (EP3) at room temperature and scan rates higher than 5 mV s^{-1} did not show a limiting current plateau, suggesting slow anodic dissolution, mainly controlled by an adsorption mechanism.

For all electropolishing conditions, the SEM images highlighted visible grain boundaries and a wavy surface morphology (Fig. 2). Indeed, a free-etched surface is expected



after EP treatment.³⁵ The etching mechanism occurs in the active dissolution region before reaching a diffusion-limited current (Fig. 1, region A). The crystal orientation influences the dissolution, resulting in distinct etch patterns,³¹ as observed in Fig. 3. The etching process stops at the formation of a passive oxide layer (Fig. 1, region B).⁶⁹ However, grain boundaries have already been observed in the case of electropolished pure Zn (ref. 46) and pure Fe.⁸⁷ While the observation of grain boundaries is generally considered a nonconformity in the electropolishing process, there is ongoing controversy in the literature regarding this aspect.^{30,49,88,89} Additionally, due to the limited number of published studies on the EP of bioabsorbable metallic alloys,^{46,87} specific mechanisms for this class of materials remain elusive.

4.2. Effect of the electropolishing process on the surface roughness and wettability

After the dissolution of the native oxide layer, a new surface with improved chemical homogeneity is expected to be formed.⁹⁰ The MP conditions exhibit low roughness values similar to the EP1 conditions, and both are smoother compared to the EP2 and EP3 conditions (Fig. 4a). In fact, the roughness is dependent on the mechanism by which the material is removed, dissolved, or treated.⁹¹ For abrasive mechanical polishing, the effect of surface roughness is determined by the particle size of the abrasive element and the normal load applied on the surface during the process.²⁶ For electropolishing, however, material removal is known to occur by two different mechanisms, which are anodic leveling (macro-smoothening) and micro-smoothening.^{31,92} The first is a function of the local charge distribution at the surface, where protuberances are preferentially dissolved. Micro-smoothening is induced by the suppression of the influence of surface defects (or crystallographic orientation) due to ionic salt formation, resulting in a diffusion-controlled process. However, the difference observed among the analyzed areas for EP2 and EP3 could be attributed to a microscale pattern and nanotopography. Liu *et al.*⁹³ observed that nanotopography can stimulate the initial osteoblast cell adhesion and spreading compared to the outer most surface chemistry. In fact, the authors concluded that the nanotopography and the chemical composition synergistically are the main mechanisms of cell differentiation. Therefore, distinct electrolytes produce different ionic salts, and thus, different surface roughness.³⁶ The presence of glycerol in EP1 and EP2 solutions promoted a significant decrease in surface roughness, R_a (Fig. 4a), and the formation of steps between adjacent grains with different orientations (Fig. 3g and h) compared to the EP3 sample (Fig. 4i). In the literature, similar trends have been observed on electropolished pure iron specimens where significant surface roughness variations were found by adding deionized water to the electrolyte composition.⁸⁷

The impact of micro-roughness on cell attachment is known,⁹⁴ but it must also be correlated with wettability and nano-topography,⁹⁵ *e.g.*, cell adhesion is enhanced on more hydrophilic surfaces. In this work, a range of water contact angles from 53° to 76° was found for all treated samples. Although samples treated in EP1 exhibited the smoothest surface among other conditions, the most hydrophilic character was obtained for EP2 ($WCA_{EP2} = 53^\circ \pm 2^\circ$). For stenting applications, a slightly hydrophilic surface (between 40° and 60°) has been suggested for improving the adhesion of endothelial cells (ECs).⁹⁶ In fact, highly hydrophilic surfaces bind proteins by weak forces and can lead to cell detachment, whereas hydrophobic surfaces promote EC adsorption but in a rigid and denatured state, resulting in conformational protein changes and reducing accessible sites to cell attachment.^{90,95,97}

4.3. Chemical composition after passivation: metallic oxides

EP2 and EP1 conditions displayed a higher contribution of the O2 band, associated with the presence of hydroxides, than the MP one, that is ~44%, ~40%, and ~38%, respectively. These hydroxides increased the surface wettability. Indeed, the EP2 surface displayed a more hydrophilic behavior than EP1 followed by MP, with $WCA_{EP2} = 53 \pm 2^\circ < WCA_{EP1} = 64 \pm 2^\circ < WCA_{MP} = 76 \pm 3^\circ$, respectively. Therefore, the hydrophilic behavior seems to be related to the OH moieties found on the surface, corresponding to the O2 peak, without neglecting the contributions of O3 and O4 peaks, associated with physiologically adsorbed water (Table 6). In contrast, the EP3 sample showed the highest contribution of the O1 peak (Me–O) and the lowest for the O2 peak, which may explain its low hydrophilicity character, $WCA_{EP3} = 72 \pm 5^\circ$, while exhibiting the lowest roughness values. Furthermore, EP3 displayed the highest concentration of metallic elements (Fe, Mn), with 18.7 at%, compared to the other conditions which exhibited a concentration range from 14–15% (Table 5). Finally, effective surface passivation was correlated with a metallic oxide enrichment, a high O_{metal}/metal ratio.

4.4. The effect of surface treatments on the corrosion behavior and biological performances

As illustrated in Fig. 8, cell viability was influenced by the extract concentrations, with the most dilute solution (Fig. 8c) leading to a twofold increase in cell viability by mitigating the concentration level, as observed in other studies.^{74,98,99} In addition, surprisingly, there was no statistical difference in HUVEC viability between the two conditions, even with varying eluate concentrations (Fig. 8). Indeed, if we consider that the samples present a different corrosion rate (section 3.5), the amount of Fe and Mn released should therefore be different and potentially lead to a cytotoxic effect. The corrosion rate of EP2 in M199 was $9 \pm 1 \mu\text{m per year}$, which is ~5 times less than that of MP (Fig. 5a, inset table). This difference in the corrosion rate, estimated by PDP tests, may be explained by the fact that EP2 degradation products



formed on the surface act as “protective” barriers against corrosion compared to those of MP. Indeed, XPS analyses (Fig. 7e and f) show that the phosphorus content is higher in EP2 than in MP, suggesting that EP2 is more prone to adsorbing phosphates, as corroborated by its high resistance to charge transfer observed in Nyquist diagrams (Fig. 5c).

The correlation between the lower corrosion resistance and the lower corrosion rate in MP-treated samples suggests a complex interaction between surface characteristics from the treatment and corrosion kinetics. While the treatment may not prevent corrosion initiation effectively, it could alter corrosion mechanisms or retard corrosion propagation once initiated. XPS analysis revealing a higher iron content implies greater permeability, mitigated by an anodically formed outer surface that restricts significant corrosion rate escalation. Thus, MP substrates show susceptibility to corrosion, albeit at a slower rate compared to EP2 conditions.

These findings underscore the importance of considering dynamic physiological conditions and larger sample volumes in toxicity assessments and corrosion studies for a more accurate representation of the material behavior in the human body. Indeed, assessing toxicity levels for soluble iron and manganese ions in static *in vitro* tests may be prone to overestimation, given the limited volume of the used medium (approximately 20 to 100 mL) compared to the total volume of human blood (6 L). In fact, toxic levels of metallic ions differ from *in vitro* to *in vivo* studies; therefore, concentrations of 3.5–5 $\mu\text{g mL}^{-1}$ for Fe and 0.03–0.06 $\mu\text{g mL}^{-1}$ for Mn in blood are known as critical concentrations.¹⁰⁰ The absence of dynamic flow, mimicking the conditions within the human body, further complicates the accuracy of such estimations.¹⁰¹

5. Conclusions

This study systematically investigated the influence of three distinct electrolytes and electropolishing conditions on the surface features of Fe-13Mn-1.2C biodegradable steel suitable for applications in a cardiovascular context. Mechanically polished samples were used as reference conditions, to compare the electropolishing effect of two acidic solutions and an ionic liquid-based one. Dissolution mechanisms occurring at the cathode/solution interface were discussed, as well as their effect on the properties of the treated surface.

1. Electropolishing was regulated mainly by two main mechanisms: a) a controlled mass transport observed for EP1 and EP2 electrolytes and b) an adsorption mechanism evident for the EP3 electrolyte. These findings suggest that EP1 and EP2 anodic dissolution was faster than that for EP3; EP1 showed the fastest anodic dissolution. Although surfaces subjected to EP1 and EP2 conditions had similar surface properties, those treated in EP2 displayed the highest ratio (1.3) of metal-bound O to metallic elements (Fe and Mn).

2. EP2 showed a passivated surface with reduced roughness ($R_a \sim 10 \text{ nm}$), a moderately hydrophilic behavior ($\text{WCA}_{\text{EP2}} = 53^\circ$), a chemical composition combining a

substantial proportion of metallic oxides (41.4% area of the O1 peak) with the highest contribution of metallic hydroxides (43.8% area of O2), and the absence of pitting and scattered degradation product heaps on the surface.

3. The significant corrosion resistance of EP2 samples in both modified Hanks' and M199 media could be attributed to the formation of a homogeneous oxide layer, hindering the diffusion of ions toward the material/medium interface.

4. No significant difference was observed in terms of cell viability studied for EP2 and MP conditions.

Further work should focus on the development of this process for tubular devices and on improving *in vitro* tests to more accurately simulate an endothelialization process.

Data Availability

OSF.io registration: <https://osf.io/dashboard>. Electropolishing Fe-based biodegradable metals for vascular applications: impact on surface properties, corrosion and cell viability https://osf.io/9dnve/?view_only=56104f08bb8b4d08bd64f54da56fdc33.

Conflicts of interest

There are no conflicts to declare.

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